

# QCE | **BIOLOGY** **UNITS 1&2**





# QCE | **BIOLOGY** **UNITS 1&2**



BIOZONE Learning Media respectfully acknowledges the traditional custodians of the lands where we work and the places in which we live.

We pay our respects to ancestors and to all First Nations elders:  
past, present and emerging.

---

## Copyright Notice

This Work is copyright. No part of this publication may be reproduced, stored in a retrieval system, or transmitted in any form or by any means, electrical, mechanical, photocopying or otherwise, without the permission of BIOZONE Learning Media Australia. Except as permitted under the Copyright Act 1968, for example, any fair dealing for the purposes of private study, research, criticism or review, subject to certain limitations. These limitations include: Restricting the copying to a maximum of 10% of this book; providing an appropriate notice and warning with all copies of the Work disseminated; taking all reasonable steps to limit access to these copies to people authorised to receive these copies; ensuring you hold the appropriate licences issued by the Copyright Agency Limited ("CAL"), supply a remuneration notice to CAL and pay any required fees. For details of CAL licences and remuneration notices please contact CAL at Level 12, 66 Goulburn Street, Sydney, 2000, Tel: 1800 066 844 or see: [www.copyright.com.au/licences-permission/educational-licences/](http://www.copyright.com.au/licences-permission/educational-licences/)

---

Second Edition 2025

ISBN 978-1-99-101435-1

Copyright ©2025 BIOZONE International Ltd

## Disclaimer

Although every care has been taken, Aboriginal and Torres Strait Islander people should be aware that this title may contain images of deceased persons.

The external weblinks (URLs) referenced in this book were correct at the time of publishing. However, due to the dynamic nature of the internet, some addresses may have changed, or cease to exist. While BIOZONE regrets any inconvenience that this may cause readers, no responsibility for any such changes or unforeseeable errors can be accepted by BIOZONE.

## Acknowledgements

BIOZONE wishes to thank and acknowledge the team for their efforts and contributions to the production of this title.

## Cover Photograph

Photo: Adobe Stock - Sven Taubert

Red tailed black cockatoo

### BIOZONE Learning Media Australia

P.O. Box 2841, Burleigh BC,  
QLD 4220, Australia

Phone: 07 5535 4896

Email: [sales@biozone.com.au](mailto:sales@biozone.com.au)

[www.BIOZONE.com/au](http://www.BIOZONE.com/au)

# QCE | BIOLOGY UNITS 1&2



## About the Authors



### Jillian Mellanby *Editor*

Jill began her science career with a degree in biochemistry and, after some time working in research institutes, became a science teacher, working in the UK and New Zealand. She spent many years as managing editor of a suite of science journals and has also written science articles for a public audience. She joined BIOZONE in late 2021.



### Kent Pryor *Author*

Kent has a BSc from Massey University majoring in zoology and ecology and taught secondary school biology and chemistry for 9 years before joining BIOZONE as an author in 2009.



### Sarah Gaze *Author*

Sarah has 16 years of experience as a Science and Chemistry teacher, recently completing M.Ed. (1st class hon) with a focus on curriculum, science, and climate change education. She has a background in educational resource development, academic writing, and art. Sarah joined the BIOZONE team at the start of 2022.



### Lissa Bainbridge-Smith *Author*

Lissa graduated with a Masters in Science (hons) from the University of Waikato. After graduation she worked in industry in a research and development capacity for eight years. Lissa joined BIOZONE in 2006 and is hands-on developing new curricula. Lissa has also taught science theory and practical skills to international and ESL students.

# Contents

Using This Worktext .....	vi
Using BIOZONE's Resource Hub.....	X

## Chapter 1: Basic Skills for QCE

<i>Key Skills and Knowledge</i> .....	1
<input type="checkbox"/> 1 How Do We Do Science?.....	2
<input type="checkbox"/> 2 Systems and Systems Models .....	4
<input type="checkbox"/> 3 Types of Data.....	5
<input type="checkbox"/> 4 Planning a Practical Investigation .....	6
<input type="checkbox"/> 5 Safety and Ethical Guidelines .....	8
<input type="checkbox"/> 6 Accuracy and Precision .....	10
<input type="checkbox"/> 7 Working with Numbers .....	12
<input type="checkbox"/> 8 Fractions, Percentages, and Ratios .....	13
<input type="checkbox"/> 9 Dealing With Large Numbers.....	14
<input type="checkbox"/> 10 Practising With Data.....	15
<input type="checkbox"/> 11 Apparatus and Measurement .....	16
<input type="checkbox"/> 12 Drawing Graphs .....	17
<input type="checkbox"/> 13 Interpreting Line Graphs .....	19
<input type="checkbox"/> 14 Correlation or Causation .....	20
<input type="checkbox"/> 15 Mean, Median, and Mode .....	21
<input type="checkbox"/> 16 What is Standard Deviation? .....	23
<input type="checkbox"/> 17 Reliability of the Mean.....	24
<input type="checkbox"/> 18 Detecting Bias in Samples .....	26
<input type="checkbox"/> 19 Statistical Tests: Which One to Use.....	27
<input type="checkbox"/> 20 Pearson Correlation Coefficient.....	28
<input type="checkbox"/> 21 Spearman Rank Correlation.....	30
<input type="checkbox"/> 22 Student's t Test.....	31
<input type="checkbox"/> 23 Chi-squared Test for Goodness of Fit .....	32
<input type="checkbox"/> 24 Did You Get it? .....	33

## Unit 1: Cells and Multicellular Organisms

### Topic 1: Cells as the basis of life

#### Chapter 2: Prokaryotic and Eukaryotic Cells

<i>Learning Outcomes</i> .....	34
<input type="checkbox"/> 25 The Cell is the Unit of Life .....	35
<input type="checkbox"/> 26 Types of Cells.....	36
<input type="checkbox"/> 27 What Are Cells Made Of?.....	37
<input type="checkbox"/> 28 What Cells Need for Survival.....	38
<input type="checkbox"/> 29 Prokaryotic Cells.....	39
<input type="checkbox"/> 30 Plant Cells.....	41
<input type="checkbox"/> 31 Identifying Structures in a Plant Cell.....	42
<input type="checkbox"/> 32 Animal Cells.....	43
<input type="checkbox"/> 33 Identifying Structures in an Animal Cell.....	44
<input type="checkbox"/> 34 Cell Structures and Organelles.....	45
<input type="checkbox"/> 35 Optical Microscopes.....	47
<input checked="" type="checkbox"/> 36 Preparing a Slide.....	49
<input type="checkbox"/> 37 Calculating Linear Magnification.....	51
<input type="checkbox"/> 38 Biological Drawings .....	52
<input type="checkbox"/> 39 Observing and Recording Using a Microscope.....	54
<input type="checkbox"/> 40 Electron Microscopes.....	55
<input type="checkbox"/> 41 Identifying Organelles .....	57
<input type="checkbox"/> 42 Did You Get it? .....	58

## Chapter 3: Cellular Differentiation and Specialisation

<i>Key Skills and Knowledge</i> .....	59
<input type="checkbox"/> 43 What are Stem Cells?.....	60
<input type="checkbox"/> 44 Cellular Differentiation.....	62
<input type="checkbox"/> 45 Mitosis and Cellular Differentiation.....	64
<input type="checkbox"/> 46 Applications of Stem Cells.....	65
<input type="checkbox"/> 47 Bioethical Issues Associated with Stem Cells .....	67
<input type="checkbox"/> 48 The Hierarchy of Life.....	68
<input type="checkbox"/> 49 Exploring Tissues and Organs.....	70
<input type="checkbox"/> 50 Respiratory and Circulatory Systems .....	72
<input type="checkbox"/> 51 Circulation and Digestive Interaction.....	74
<input type="checkbox"/> 52 Circulation and Excretory Interaction .....	76
<input type="checkbox"/> 53 Animals in Medical Research .....	78
<input type="checkbox"/> 54 Did You Get it? .....	79

## Chapter 4: Cell Membrane

<i>Key Skills and Knowledge</i> .....	80
<input type="checkbox"/> 55 The Plasma Membrane .....	81
<input type="checkbox"/> 56 Phospholipids and the Properties of Membranes .....	82
<input type="checkbox"/> 57 Proteins of the Plasma Membrane.....	83
<input type="checkbox"/> 58 How Do We Know? Membrane Structure .....	85
<input type="checkbox"/> 59 Cell Membrane Research.....	86
<input type="checkbox"/> 60 Modelling the Plasma Membrane.....	87
<input checked="" type="checkbox"/> 61 Diffusion.....	89
<input type="checkbox"/> 62 Diffusion and Cell Size .....	92
<input type="checkbox"/> 63 Comparing Cell Sizes .....	93
<input type="checkbox"/> 64 Investigating the Effect of Cell Size.....	94
<input type="checkbox"/> 65 Overcoming Limitations to Cell Size .....	96
<input type="checkbox"/> 66 Osmosis .....	97
<input checked="" type="checkbox"/> 67 Estimating Osmolarity of Cells .....	98
<input type="checkbox"/> 68 Water Relations in Plant Cells.....	99
<input type="checkbox"/> 69 Investigating Membrane Solubility and Diffusion .....	100
<input type="checkbox"/> 70 Active Transport .....	101
<input type="checkbox"/> 71 Ion Pumps and Cotransport.....	102
<input type="checkbox"/> 72 Cytosis.....	103
<input type="checkbox"/> 73 Active and Passive Transport Summary.....	105
<input type="checkbox"/> 74 Did You Get it? .....	106
<input type="checkbox"/> 75 Synoptic Question: Unit 1, Topic 1.....	107

### Topic 2: Exchange of nutrients and wastes

#### Chapter 5: Exchange of Nutrients and Wastes

<i>Key Skills and Knowledge</i> .....	109
<input type="checkbox"/> 76 Carbohydrates, Proteins, and Lipids.....	110
<input type="checkbox"/> 77 The Mammalian Circulatory System .....	111
<input type="checkbox"/> 78 Blood Vessels.....	112
<input type="checkbox"/> 79 Capillaries and Capillary Networks .....	113
<input type="checkbox"/> 80 Structure of the Mammalian Heart .....	115
<input type="checkbox"/> 81 The Digestive System .....	116
<input type="checkbox"/> 82 The Stomach and Small Intestine .....	117
<input type="checkbox"/> 83 Digestion, Absorption, and Transport.....	120
<input type="checkbox"/> 84 The Large Intestine.....	122
<input checked="" type="checkbox"/> 85 Investigating Amylase Activity .....	123
<input type="checkbox"/> 86 Nitrogenous Wastes in Animals.....	125

<input type="checkbox"/>	<b>87</b>	The Excretory System.....	126
<input type="checkbox"/>	<b>88</b>	Kidney Structure.....	127
<input type="checkbox"/>	<b>89</b>	Nephron Structure and Function.....	128
<input type="checkbox"/>	<b>90</b>	Organ and Tissue Transplantation.....	130
<input type="checkbox"/>	<b>91</b>	Did You Get it?.....	131

### Chapter 6: Internal Membranes and Enzymes

		<i>Key Skills and Knowledge</i> .....	132
<input type="checkbox"/>	<b>92</b>	Enzymes.....	133
<input type="checkbox"/>	<b>93</b>	Models of Enzyme Activity.....	134
<input type="checkbox"/>	<b>94</b>	How Enzymes Work.....	135
<input type="checkbox"/>	<b>95</b>	Factors Affecting Enzyme Activity.....	137
<input type="checkbox"/>	<b>96</b>	Enzyme Inhibition.....	139
<input checked="" type="checkbox"/>	<b>97</b>	Investigating Enzyme Activity.....	140
<input type="checkbox"/>	<b>98</b>	Achieving Metabolic Efficiency.....	142
<input type="checkbox"/>	<b>99</b>	Enzymes and Membranes.....	143
<input type="checkbox"/>	<b>100</b>	Enzymes and Disease.....	144
<input type="checkbox"/>	<b>101</b>	Did You Get it?.....	146
<input type="checkbox"/>	<b>102</b>	Synoptic Question: Unit 1, Topic 2.....	147

### Topic 3: Cellular energy, gas exchange and plant physiology

#### Chapter 7: Respiration and Mammalian Gas Exchange

		<i>Key Skills and Knowledge</i> .....	149
<input type="checkbox"/>	<b>103</b>	Metabolism and Life.....	150
<input type="checkbox"/>	<b>104</b>	ATP in Cells.....	151
<input checked="" type="checkbox"/>	<b>105</b>	Measuring Respiration.....	153
<input type="checkbox"/>	<b>106</b>	Cellular Respiration: Inputs and Outputs.....	155
<input type="checkbox"/>	<b>107</b>	Anaerobic Pathways.....	157
<input checked="" type="checkbox"/>	<b>108</b>	Investigating Fermentation in Yeast.....	158
<input type="checkbox"/>	<b>109</b>	Principles of Gas Exchange.....	160
<input type="checkbox"/>	<b>110</b>	The Human Gas Exchange System.....	161
<input type="checkbox"/>	<b>111</b>	The Lungs.....	162
<input type="checkbox"/>	<b>112</b>	Gas Transport in Humans.....	164
<input type="checkbox"/>	<b>113</b>	Did You Get it?.....	166

#### Chapter 8: Plant Gas Exchange and Transport Systems

		<i>Key Skills and Knowledge</i> .....	167
<input type="checkbox"/>	<b>114</b>	Energy Transformations in Cells.....	168
<input type="checkbox"/>	<b>115</b>	The Role of Photosynthesis.....	169
<input type="checkbox"/>	<b>116</b>	Chloroplasts.....	170
<input type="checkbox"/>	<b>117</b>	Photosynthesis: Inputs and Outputs.....	171
<input type="checkbox"/>	<b>118</b>	Investigating Photosynthetic Rate.....	173
<input type="checkbox"/>	<b>119</b>	Photosynthesis and Productivity.....	174
<input type="checkbox"/>	<b>120</b>	The Plant Body.....	175
<input type="checkbox"/>	<b>121</b>	Xylem.....	176
<input type="checkbox"/>	<b>122</b>	Phloem.....	177
<input type="checkbox"/>	<b>123</b>	Uptake at the Root.....	178
<input type="checkbox"/>	<b>124</b>	Transpiration.....	179
<input type="checkbox"/>	<b>125</b>	Gas Exchange and Stomata.....	181
<input type="checkbox"/>	<b>126</b>	Conditions for Photosynthesis.....	183
<input checked="" type="checkbox"/>	<b>127</b>	Investigating Plant Transpiration.....	184
<input type="checkbox"/>	<b>128</b>	Translocation.....	187
<input type="checkbox"/>	<b>129</b>	Plants and Technology.....	188
<input type="checkbox"/>	<b>130</b>	Did You Get it?.....	189
<input type="checkbox"/>	<b>131</b>	Synoptic Question: Unit 1, Topic 3.....	190

## Unit 2: Maintaining the Internal Environment

### Topic 1: Homeostasis- thermoregulation and osmoregulation

#### Chapter 9: Neural Homeostatic Controls

		<i>Key Skills and Knowledge</i> .....	192
<input type="checkbox"/>	<b>132</b>	Homeostasis.....	193
<input type="checkbox"/>	<b>133</b>	Negative feedback.....	194
<input type="checkbox"/>	<b>134</b>	Sensory Receptors.....	196
<input type="checkbox"/>	<b>135</b>	Nervous Regulation in Vertebrates.....	198
<input type="checkbox"/>	<b>136</b>	Neurons.....	199
<input type="checkbox"/>	<b>137</b>	Reflexes.....	201
<input type="checkbox"/>	<b>138</b>	Transmission of Nerve Impulses.....	202
<input type="checkbox"/>	<b>139</b>	Chemical Synapses.....	204
<input type="checkbox"/>	<b>140</b>	Integration at Synapses.....	206
<input type="checkbox"/>	<b>141</b>	Drugs at Synapses.....	207
<input type="checkbox"/>	<b>142</b>	Did You Get it?.....	208

#### Chapter 10: Hormonal Homeostatic Controls

		<i>Key Skills and Knowledge</i> .....	209
<input type="checkbox"/>	<b>143</b>	Types of Cell Signalling.....	210
<input type="checkbox"/>	<b>144</b>	Signalling Molecules.....	212
<input type="checkbox"/>	<b>145</b>	How Hormones Work.....	213
<input type="checkbox"/>	<b>146</b>	What is Signal Transduction?.....	214
<input type="checkbox"/>	<b>147</b>	Types of Signal Transduction.....	215
<input type="checkbox"/>	<b>148</b>	Action of Insulin.....	217
<input type="checkbox"/>	<b>149</b>	Hormone Regulation by Negative Feedback.....	218
<input type="checkbox"/>	<b>150</b>	Use of Hormones in the Dairy Industry.....	219
<input type="checkbox"/>	<b>151</b>	Did You Get It?.....	220

#### Chapter 11: Thermoregulation

		<i>Key Skills and Knowledge</i> .....	221
<input type="checkbox"/>	<b>152</b>	Mechanisms for Thermoregulation.....	222
<input type="checkbox"/>	<b>153</b>	Structural Features for Thermoregulation.....	223
<input type="checkbox"/>	<b>154</b>	Behavioural Responses for Thermoregulation.....	225
<input type="checkbox"/>	<b>155</b>	Physiological Mechanisms for Thermoregulation.....	227
<input type="checkbox"/>	<b>156</b>	Hormonal Mechanisms for Thermoregulation.....	229
<input type="checkbox"/>	<b>157</b>	Modelling Human Thermoregulation.....	231
<input type="checkbox"/>	<b>158</b>	Did You Get it?.....	233

#### Chapter 12: Osmoregulation

		<i>Key Skills and Knowledge</i> .....	234
<input type="checkbox"/>	<b>159</b>	What is Osmoregulation?.....	235
<input type="checkbox"/>	<b>160</b>	Osmoregulation in Intertidal Organisms.....	237
<input type="checkbox"/>	<b>161</b>	Osmoregulation in Bony Fish.....	238
<input type="checkbox"/>	<b>162</b>	Managing Fluid Balance on Land.....	239
<input type="checkbox"/>	<b>163</b>	ADH and Water Balance.....	241
<input type="checkbox"/>	<b>164</b>	Osmoregulation in Plants.....	242
<input checked="" type="checkbox"/>	<b>165</b>	Investigating Stomatal Density.....	245
<input type="checkbox"/>	<b>166</b>	Salt Tolerance in Plants.....	247
<input type="checkbox"/>	<b>167</b>	Did You Get it?.....	249
<input type="checkbox"/>	<b>168</b>	Synoptic Question: Unit 2, Topic 1.....	250

## Topic 2: Infectious disease and epidemiology

### Chapter 13: Infectious Disease

	<i>Key Skills and Knowledge</i> .....	252
<input type="checkbox"/>	<b>169</b> Infection and Disease .....	253
<input type="checkbox"/>	<b>170</b> Bacterial Diseases .....	256
<input type="checkbox"/>	<b>171</b> Fungal Diseases .....	258
<input type="checkbox"/>	<b>172</b> Protistan Diseases .....	259
<input type="checkbox"/>	<b>173</b> Viral Diseases .....	261
<input type="checkbox"/>	<b>174</b> HIV: An Example of a Viral Disease .....	262
<input type="checkbox"/>	<b>175</b> Prions .....	264
<input type="checkbox"/>	<b>176</b> Did You Get it? .....	265

### Chapter 14: Immune Response

	<i>Key Skills and Knowledge</i> .....	266
<input type="checkbox"/>	<b>177</b> The Nature of Antigens .....	267
<input type="checkbox"/>	<b>178</b> The Body's Defences: An Overview .....	269
<input type="checkbox"/>	<b>179</b> The Innate Immune Response .....	270
<input type="checkbox"/>	<b>180</b> Phagocytes and Phagocytosis .....	273
<input type="checkbox"/>	<b>181</b> The Lymphatic System .....	274
<input type="checkbox"/>	<b>182</b> Processing Antigens .....	275
<input type="checkbox"/>	<b>183</b> The Adaptive Immune Response .....	276
<input type="checkbox"/>	<b>184</b> Clonal Selection .....	278
<input type="checkbox"/>	<b>185</b> Antibodies .....	279
<input type="checkbox"/>	<b>186</b> Acquired Immunity .....	280
<input type="checkbox"/>	<b>187</b> Vaccines and Vaccination .....	282
<input type="checkbox"/>	<b>188</b> Vaccines Can Eliminate Infectious Disease ...	284
<input type="checkbox"/>	<b>189</b> Vaccine Development .....	285
<input type="checkbox"/>	<b>190</b> Long Term immune Response Data .....	286
<input type="checkbox"/>	<b>191</b> Physical Defences in Plants .....	288
<input type="checkbox"/>	<b>192</b> Chemical Defences in Plants .....	289
<input type="checkbox"/>	<b>193</b> Did You Get it? .....	291

### Chapter 15: Transmission and Spread of Disease

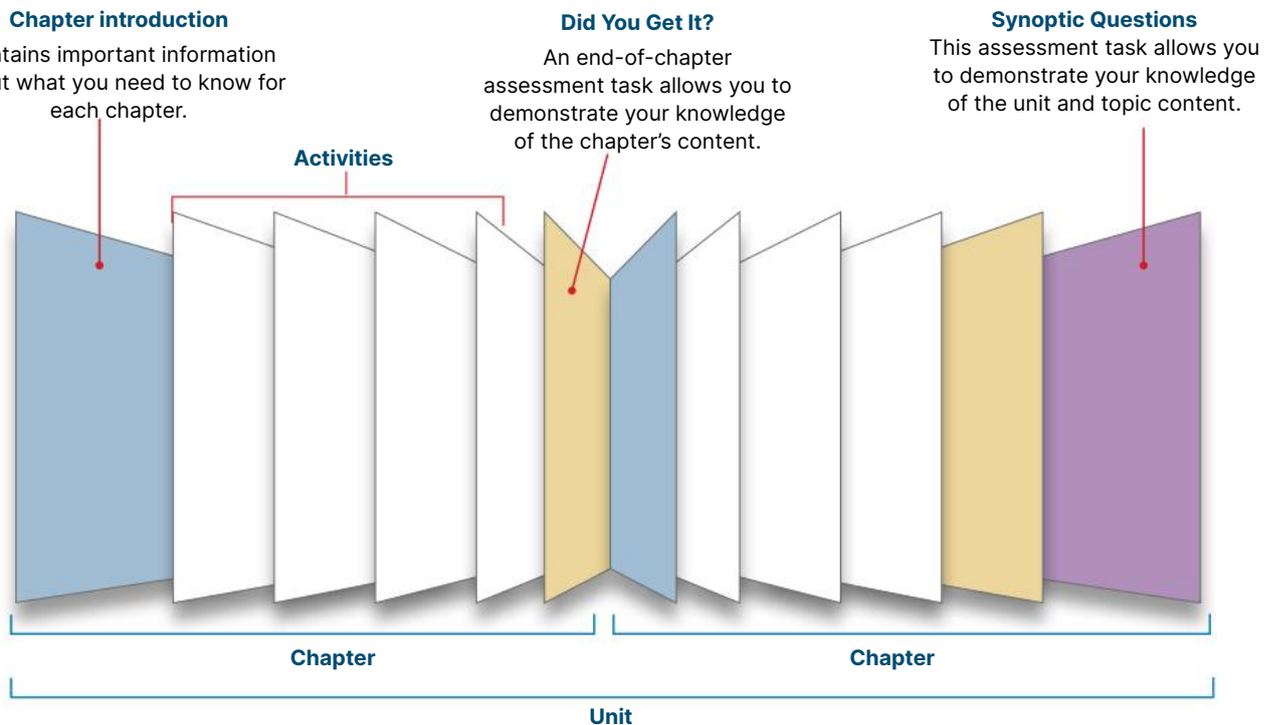
	<i>Key Skills and Knowledge</i> .....	292
<input type="checkbox"/>	<b>194</b> Transmission of Disease .....	293
<input type="checkbox"/>	<b>195</b> Testing Antibiotics .....	295
<input type="checkbox"/>	<b>196</b> Patterns of Disease .....	297
<input checked="" type="checkbox"/>	<b>197</b> The Effectiveness of Hand Washing .....	299
<input checked="" type="checkbox"/>	<b>198</b> Modelling Disease Outbreak and Spread .....	301
<input type="checkbox"/>	<b>199</b> Predicting Future Patterns of Disease .....	304
<input type="checkbox"/>	<b>200</b> Containing The Spread of Disease .....	306
<input type="checkbox"/>	<b>201</b> Biosecurity Measures to Protect Australia .....	308
<input type="checkbox"/>	<b>202</b> The Effectiveness of Health Programs .....	309
<input type="checkbox"/>	<b>203</b> Aboriginal Protocols in Medicine .....	311
<input type="checkbox"/>	<b>204</b> Did You Get it? .....	312
<input type="checkbox"/>	<b>205</b> Synoptic Question: Unit 2, Topic 2 .....	314

	Appendix 1: Equipment List .....	315
	Appendix 2: Glossary .....	316
	Image Credits .....	323
	Index .....	324

# Using This Worktext

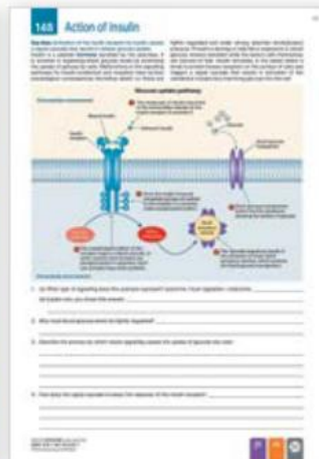
This edition of Biology for QCE Units 1 & 2 has been specifically written for the Queensland (QCE) Biology general senior syllabus (2025 version). The next few pages provide information about this resource and how to get the best use from it.

## Structure of a Unit and Chapter



### Introduction

- Provides a list of important key concepts for the chapter.
- Lists important key terms (vocab) for the chapter.
- Provides a check list of unit objectives for the chapter.
- Activities with SHE, SU, and SI components are identified.



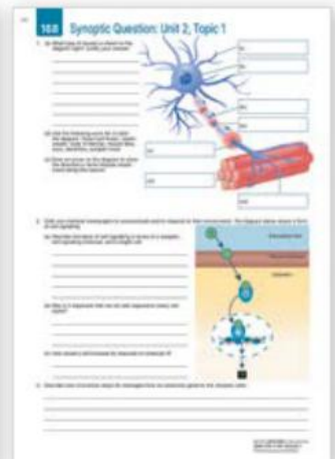
### Activities

- The KEY IDEA provides the focus for the activity.
- Annotated diagrams and photographs help you understand the content.
- Answering the questions helps you consolidate your understanding of the content.
- Use the material to revise for tests and exams.



### Chapter test

- End-of-chapter assessment tasks test your understanding of the biological terms and concepts covered within the chapter.
- Reviewing the answers can help you study for tests and exams.



### Synoptic question

- Synoptic questions conclude each unit and topic of study covered in the book.
- Use them to see how well you understand the content.
- Reviewing the answers can help you study for tests and exams.

## Chapter introductions

The chapter introductions contain useful information about what you need to know for QCE Biology. It identifies key concepts and learning outcomes (what you need to know), identifies important vocabulary, and provides a quick link to supporting resources on BIOZONE's **Resource Hub**. The key features of the chapter introduction are explained below.

The section of the worktext you are in is identified for easy navigation.

QR codes and bitly tags allow you to quickly navigate to helpful content (e.g. videos and models) on BIOZONE's **Resource Hub**.

Chapter number and chapter title are identified for quick navigation.

**Key concepts**  
These are the important key ideas for the chapter. Make sure you understand the concepts summarised here.

**Key terms**  
Important vocabulary you should understand and use in your course. Definitions are provided in the glossary at the back of the book.

**Learning outcomes**  
These provide a point by point summary of what you need to know or do by the end of the chapter.

**Check boxes**  
Use the check boxes to keep a record of which activities you need to complete and tick them off as you work through them.

**Activity number:**  
The activity number for each learning outcome is identified.

**Component coding**  
Colour panels and codes identify where Science as a Human Endeavour (SHE), Science Inquiry (SI), or Science Understanding (SU) occur in the chapter.

## Structure of the activity pages

Activities make up most of the worktext. Be sure to interact with all the elements on the page so you don't miss any valuable information. As you work through the material, answer the questions and complete the tasks provided. Inputting your answers will form a record of work which helps you remember what you have learned. It can also be used for revision at a later date.

**Key Idea:**  
This provides a focus for the activity and can be used as a summary take-home point of the activity.

**Activity number:**  
Identifies the activity number to help you navigate between activities.

**Introductory paragraph:**  
This provides background or introductory information to the topic.

More information about the topic is provided through explanatory text, images, diagrams, case studies, and data.

**QR codes:**  
These provide a quick link to interactive 3D models.

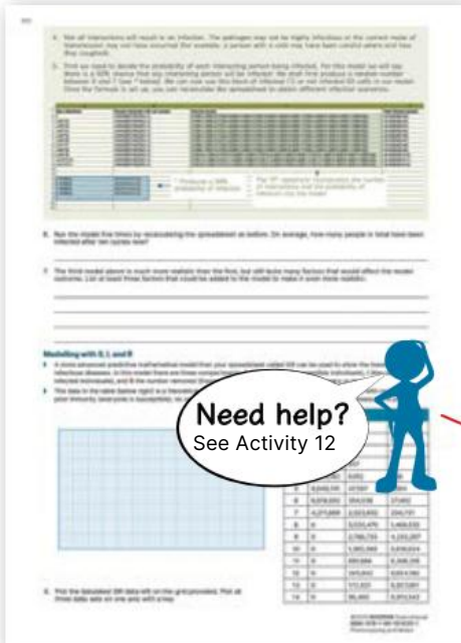
**Activity based questions:**  
Answering the questions helps reinforce your learning. Use your answers to review for tests and other assessments.

**Tab system:**  
Page tabs identify where components of the syllabus are embedded in an activity (see more on page viii). The grey tab indicates there is support material available on the **BIOZONE Resource Hub**.

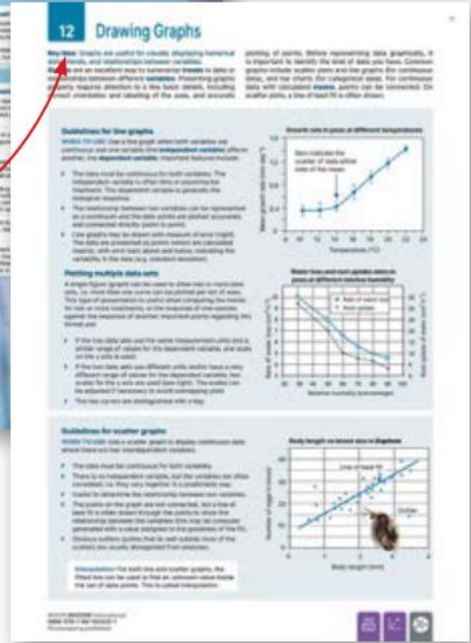
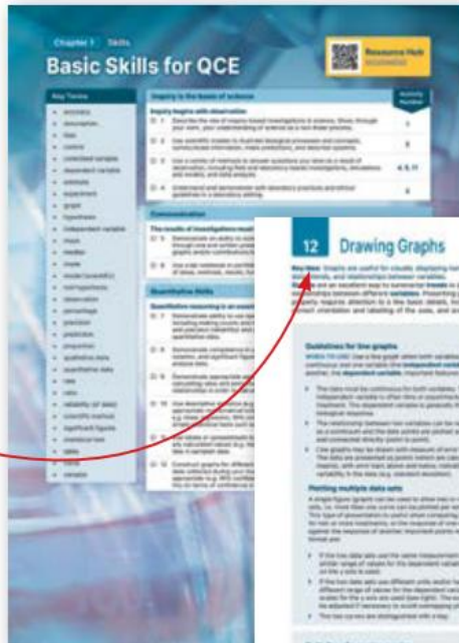


# Help with the basic maths skills

The first chapter in this worktext provides information and support to help you with some basic skills you will encounter as you work through the activities. It will help you with graphing, doing simple calculations (e.g. calculating mean, median, and mode, and rates), how to carry out common statistical tests and more. Your teacher may ask you to do certain science practice activities at specific times, but you can refer to the chapter at any time if you need help with maths skills.



**Need help?**  
See Activity 12



**Need help?**  
Some activities have a “Need help?” icon on the page. This icon lets you know that there is support for a maths skill in the Basic Skills for QCE chapter. For example, if you need help with deciding which graph to draw.

# Practical investigations

Practical investigations form an important component of the QCE Biology syllabus. Practical work provides opportunity for inquiry and investigation, and allows you to develop your manipulative skills. Practicals encourage the use of 21st century skills (collaboration and teamwork, communication, critical thinking) and provide opportunities to apply your skills in literacy and numeracy. An equipment list at the back of this worktext details the equipment needed to carry out the practical investigations.



# Using BIOZONE's Resource Hub

- ▶ Many activities have interesting resources, such as videos and 3D models, to help you understand the content. A grey tab (right) on the activity page indicates there is support on the BIOZONE **Resource Hub** for the activity.
- ▶ Navigate to the Resource Hub either by bookmarking the link below, or by utilising the bitly tag or QR code found on each chapter introduction (below, right).



**Step 1:** Navigate to the BIOZONE **Resource Hub**

**www.BIOZONEhub.com**

**Step 2:** Enter this code in the box displayed.

**QCE11-2-4351**

**Step 3:** Bookmark this page.



<https://bit.ly/3zocOuL>

Use this bitly tag or QR code to directly access the BIOZONE Resource Hub.

## Using the QR codes on activity pages

Some activities have QR codes on the pages (below). These link directly to informative and engaging 3D models. If your school does not let you use your phone in class, you can still access the models and data sets through the **Resource Hub**. Follow the steps above to access the resources through the **Resource Hub**.

**32 Animal Cells**

**Key Idea:** Animal cells are eukaryotic cells. They have many features in common with plant cells, but also have a number of unique features. Animal cells, unlike plant cells, do not have a number of unique features. In fact, some animal cells (such as phagocytes) are able to alter their shape for various purposes (e.g. engulfing foreign material). The diagram below shows the structure and organelles of a liver cell. It contains organelles common to most relatively unspecialised human cells. Note the differences between this cell and the generalised plant cell. The plant cells activity provides further information on the organelles listed here but not described.

**Vacuoles:** Smaller than those found in plant cells, in animal cells, vacuoles have minor roles in exocytosis and endocytosis.

**Smooth endoplasmic reticulum:** ER without ribosomes. It's a site for lipid and carbohydrate metabolism, including hormone synthesis.

**Nucleolus:** A dense, solid structure composed of crystalline protein and nucleic acid. They are involved in ribosome synthesis.

**Ribosomes:** These small structures may be free in the cytoplasm or associated with the endoplasmic reticulum (ER). Ribosomes in animal cells are 80S ribosomes.

**Rough endoplasmic reticulum:** A site of protein synthesis. The rough ER also synthesises new membranes, growing in place by adding proteins and phospholipids.

**Golgi apparatus (20–200 nm):** A series of flattened, disc-shaped sacs, stacked one on top of the other and connected with the ER. The Golgi stores, modifies, and packages proteins. It 'tags' proteins so that they go to their correct destination.

**Lysosomes:** A sac bounded by a single membrane. They are pinched off from the Golgi apparatus and contain and transport enzymes that break down food and foreign matter. Lysosomes show little internal structure but often contain fragments of material being broken down. Specialised lysosomes, are generally absent from plant cells.

**Nuclear pore:** A hole in the nuclear membrane allowing the nucleus to communicate with the rest of the cell.

**Tight junctions:** Join cells together in the formation of tissue.

**Nuclear membrane:** Double layered.

**Cytoplasm**

**Plasma (cell surface) membrane**

**Centrioles:** Structures with a centrosome associated with nuclear division. They are composed of microtubules, but appear as small, featureless particles, 0.25 µm diameter, under a light microscope. They are absent in higher plant cells and some protists.

**Chondrioid (or, chlorochlorid):** An organelle found in a double membrane system. The size of a cell depends on its metabolic activity.

1. What is the difference between vacuoles in plant and animal cells?

2. Name one structure or organelle present in generalised animal cells but absent from plant cells and describe its function:

3. The two photomicrographs, left, show several types of animal cells. Identify the features indicated by the letters A-C:

(a) \_\_\_\_\_

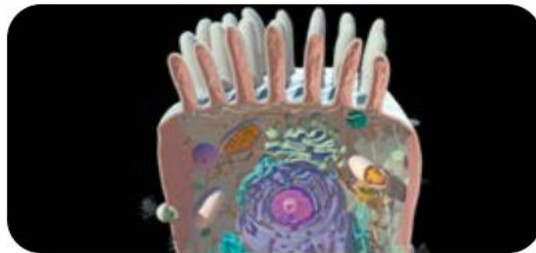
(b) \_\_\_\_\_

(c) \_\_\_\_\_

White blood cells and red blood cells

©2013 BIOZONE International  
ISBN: 978-1-99-101435-1  
Photocopying prohibited

Scan the **QR codes** on the activity pages. These link directly to informative and engaging 3D models. All models can be rotated and zoomed, and some contain informative annotations.



# Basic Skills for QCE



## Key Terms

- accuracy
- assumption
- bias
- control
- controlled variable
- dependent variable
- estimate
- experiment
- graph
- hypothesis
- independent variable
- mean
- median
- mode
- model (scientific)
- null hypothesis
- observation
- percentage
- precision
- prediction
- proportion
- qualitative data
- quantitative data
- rate
- ratio
- reliability (of data)
- scientific method
- significant figures
- statistical test
- table
- trend
- variable

## Inquiry is the basis of science

 Activity  
 Number

### Inquiry begins with observation

- |                            |  |          |
|----------------------------|--|----------|
| <input type="checkbox"/> 1 | Describe the role of inquiry-based investigations in science. Show, through your work, your understanding of science as a non-linear process.                                      | 1        |
| <input type="checkbox"/> 2 | Use scientific models to illustrate biological processes and concepts, communicate information, make predictions, and describe systems.  | 2        |
| <input type="checkbox"/> 3 | Use a variety of methods to answer questions you raise as a result of observation, including field and laboratory-based investigations, simulations and models, and data analysis. | 4, 5, 11 |
| <input type="checkbox"/> 4 | Understand and demonstrate safe practices and ethical guidelines in both laboratory and field settings.  | 5        |

## Communication

### The results of investigations must be communicated to peers to have value

- |                            |   |       |
|----------------------------|---|-------|
| <input type="checkbox"/> 5 | Demonstrate an ability to communicate the findings of your investigations through oral and written presentations, including lab reports, and through graphs and/or contributions to online resources. | 4, 12 |
| <input type="checkbox"/> 6 | Use a lab notebook or portfolio to organise your work and provide a record of ideas, methods, results, further questions, and references.   | 4, 12 |

## Quantitative Skills

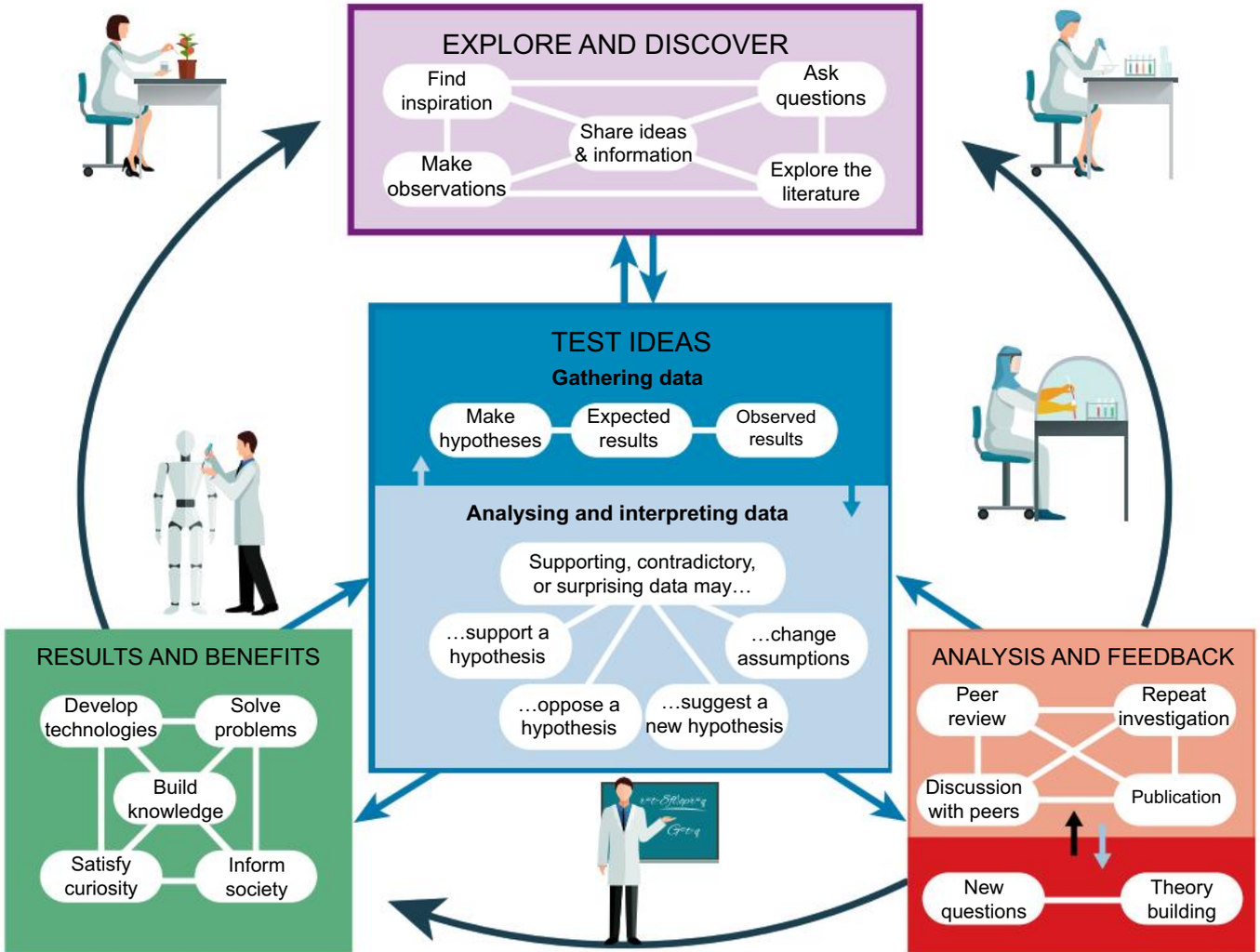
### Quantitative reasoning is an essential part of inquiry in biology

- |                             |   |              |
|-----------------------------|---|--------------|
| <input type="checkbox"/> 7  | Demonstrate ability to use basic mathematical skills to collect data, including making counts and measurements. Distinguish between accuracy and precision (reliability) and understand their importance when collecting quantitative data.   | 3, 4, 6      |
| <input type="checkbox"/> 8  | Demonstrate competence in use of ratios and proportions, scientific notation, and significant figures. Use estimation and calculation to analyse data.  | 7-10         |
| <input type="checkbox"/> 9  | Demonstrate appropriate application of mathematical routines to data, e.g. calculating rates and percentages. Interpret and manipulate mathematical relationships in order to calculate and predict values.   | 7-10         |
| <input type="checkbox"/> 10 | Use descriptive statistics (e.g. mean and standard deviation) and apply appropriate mathematical tools to analyse data and/or test hypotheses, e.g. linear regression, 95% confidence intervals, Spearman's rank correlation, and some simple statistical tests such as Student's t- and chi-squared. | 13-17, 19-23 |
| <input type="checkbox"/> 11 | Use tables or spreadsheets to organise different types of data, including any calculated values (e.g. means and standard deviation). Acknowledge bias in sampled data.  | 15-18        |
| <input type="checkbox"/> 12 | Construct graphs for different types of data, including logarithmic data and data collected during your investigations. Plot error in calculated values as appropriate (e.g. 95% confidence limits) and understand the value in doing this (in terms of confidence in the data).                      | 9, 12-17     |

# 1 How Do We Do Science?

**Key Idea:** The scientific method is a rigorous process of observation, measurement, and analysis that helps us to explain phenomena and predict changes in a system. Scientific knowledge is gained through a non-linear, dynamic process called the **scientific method**. The scientific method

is not a strict set of rules to be followed, but rather a way of approaching problems in a rigorous, but open-minded way. It involves inspiration and creativity, it is dynamic and context dependent, and usually involves collaboration. The **model** below is one interpretation of the scientific method.



## Citation and references

All scientific work acknowledges sources of information through citation and a list of references. Citations support the statements made in the text in context, and all citations are then listed alphabetically, or identified and referenced sequentially by number. Internet sites are dated and site author acknowledged. Thorough and accurate citation and referencing shows you have explored the topic, have evidence to support your work, and you are not taking credit for work that is not your own. Each publication sets its own particular referencing style and these can vary widely. In your own work, it is most important to be consistent.

Citation and reference by numbers

**Introduction**

Biological data are being produced at a phenomenal rate [1]. For example as of August 2000, the GenBank repository of nucleic acid sequences contain entries [2] and the database of proteins contained 88,166 [3]

**References**

1. Reichhardt T. It's sink or swim as a tidal wave of data approaches. *Nature* 1999;399(6736):517-20.

Citation and reference by authors

the long-term viability of a population. Individual fitness, resistance to disease and parasites, and the ability of populations to respond to environmental changes may decrease as a consequence of reduced genetic variation (Lacy 1997). Although "bottlenecks, or

Stearns. 1994. Selection against inbred Song Sparrows during a natural population bottleneck. *Nature* 372:356-357.

Lacy, R. C. 1997. Importance of genetic variation to the viability of mammalian populations. *Journal of Mammalogy* 78:320-335.

Author	Year	Title	Publication	Volume:	pages
Lacy, R. C.	1997	Importance of genetic variation to the viability of mammalian populations	<i>Journal of Mammalogy</i>	78	320-335

The style you choose is not as important as being consistent, thorough, and honest about drawing on other people's work. All the information needed to locate the reference should be included (above).



## Observations, questions, and hypotheses

- ▶ **Observation** is the beginning of any scientific investigation. Often the best investigations are based on a series of fortuitous or specific observations. For example, in 1765, Edward Jenner developed the first vaccination for smallpox after hearing that milkmaids who contracted cowpox (a harmless disease) never got smallpox. After observing a phenomenon, questions must be asked: What causes the phenomenon? Is it linked to other observations? Can it be manipulated?
- ▶ An observation may generate a number of plausible hypotheses. Scientific hypotheses are tentative testable explanations for observed phenomena. A hypothesis leads to one or more **predictions** about the way a system will behave so a research hypothesis is usually written to include a testable prediction, i.e. if X is true, then the effect of Y will be Z.
- ▶ For every **hypothesis**, there is a corresponding null hypothesis, i.e. a hypothesis of no difference or no effect. A null hypothesis allows a hypothesis to be tested statistically and can be rejected if the experimental results are statistically significant. Hypotheses are not static, but may be modified as more information becomes available.

**Example:** Two observations were made, as described below and used to produce a hypothesis:



**Observation 1:** Some caterpillar species are brightly coloured and appear to be conspicuous to predators (e.g. insectivorous birds). Predators appear to avoid these species. These caterpillars are often found in groups, rather than being solitary.



**Observation 2:** Some caterpillar species are cryptic in their appearance or behaviour. Their camouflage is so convincing that, when alerted to danger, they are difficult to see against their background. Such caterpillars are usually found alone.



**Hypothesis:** If bright colours indicate to predators that caterpillars are distasteful, birds will not eat them. The corresponding **null hypothesis** would be there is no difference in palatability between the bright and cryptically coloured caterpillars.

### Assumptions

Any biological investigation requires you to make **assumptions** about the biological system you are working with. Assumptions are features of the system (and your investigation) that you assume to be true but do not (or cannot) test. Possible assumptions about the biological system above are described in the box right:

- ▶ Insectivorous birds have colour vision.
- ▶ Caterpillars that look bright or cryptic to us, also appear that way to insectivorous birds.
- ▶ Insectivorous birds learn about the palatability of prey by tasting them.

1. What is the role of citation and correct referencing when reporting on scientific investigations?

---



---



---

2. Study the diagram opposite and write a paragraph on the scientific process and the role of surprising results in science. Attach it to this page. At the end of your course, reexamine what you wrote. Have your ideas changed?

---



---

3. Based on the hypothesis above, generate a prediction about the behaviour of insectivorous birds towards caterpillars:

---



---



---

4. During a routine preparation of bacterial colonies on agar plates, a laboratory assistant noticed that the colonies left overnight on the side of a bench near a heating unit grew faster than those left on the opposite side of the bench. The assistant decided to test this observation by experiment:

(a) State a hypothesis for the investigation: \_\_\_\_\_

(b) Generate a prediction based on the hypothesis: \_\_\_\_\_

# 2 Systems and System Models

**Key Idea:** Systems are assemblages of interrelated components working together. Models can be mathematical or visual representations of these systems.

A system is a set of interrelated components that work together. Energy flow in ecosystems (such as the Queensland rainforest on the right), gene regulation, interactions between organ systems, and feedback mechanisms are all examples of systems studied in biology. Modelling systems helps to understand how they work. A **model** is a representation of a system and is useful for breaking a complex system down into smaller parts that can be studied more easily. Often only part of a system is modelled. As scientists gather more information about a system, more data can be put into the model so that eventually it represents the real system more closely.

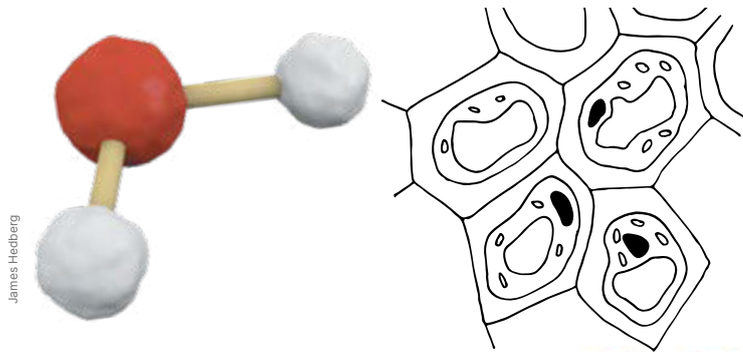


## Modelling systems

There are many different ways to model systems or their components. Often, seeing data presented in different ways can help us to understand it better. Some common examples of models are shown here.

### Visual models

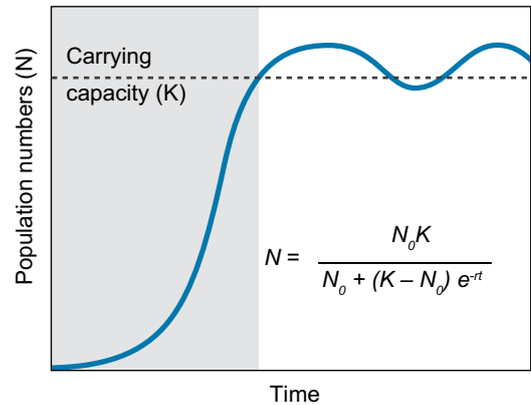
Visual models can include drawings, such as these plant cells (below right) or three dimensional physical or computer generated models. Three dimensional models can be made out of materials such as modelling clay and ice-cream sticks, like the model of a water molecule (below left).



James Hedberg

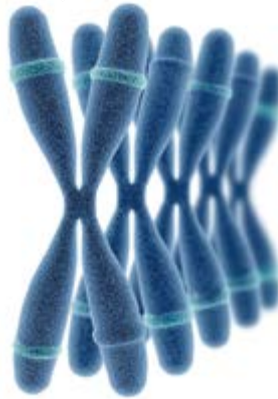
### Mathematical models

Displaying data in a **graph** or as a mathematical equation, as shown below for logistic growth, often helps us to see relationships between different parts of a system.



### Analogy

An analogy is a comparison between two things. Comparing a biological system to an everyday object can sometimes help us to understand it better. For example, the heart pumps blood in blood vessels in much the same way a fire truck pumps water from a fire hydrant through a hose. Similarly, the DNA in chromosomes is like a library. Extending that analogy further, the steps in baking a cake from a recipe book provide an analogy for how the instructions in DNA (the recipe) are translated into a specific protein (the cake).



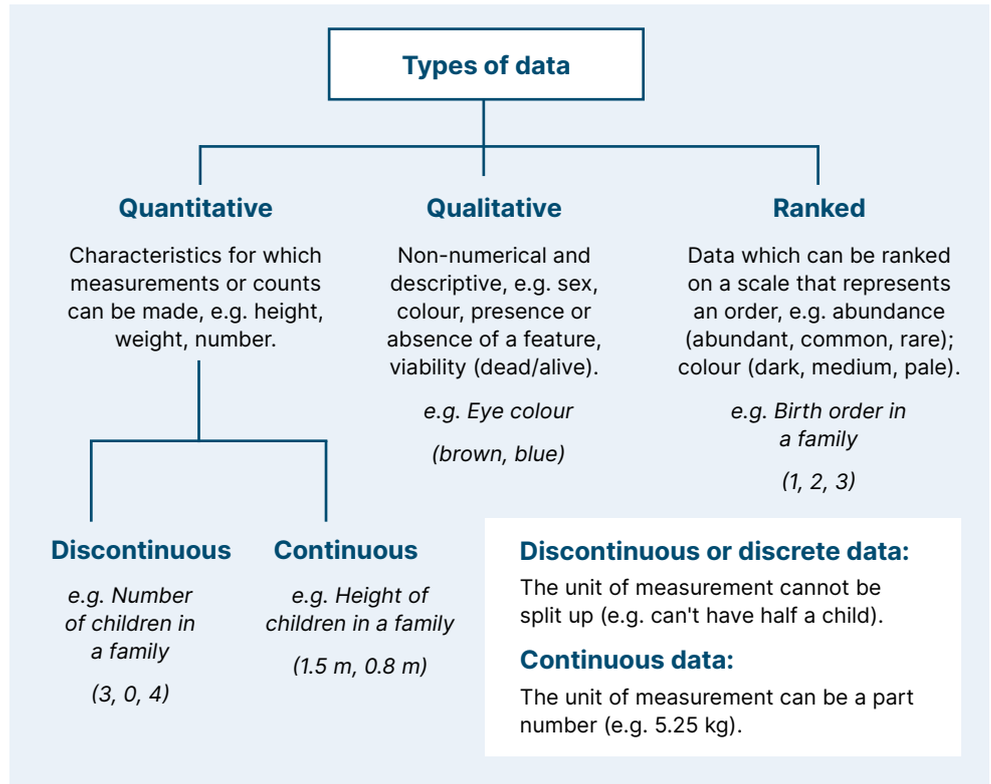
The DNA in chromosomes is like ..... a library of books

1. What is a system? \_\_\_\_\_  
\_\_\_\_\_
2. (a) What is a model? \_\_\_\_\_  
\_\_\_\_\_
- (b) Why do scientists often study one part of a system rather than the whole system? \_\_\_\_\_  
\_\_\_\_\_  
\_\_\_\_\_  
\_\_\_\_\_



**Key Idea:** Data is information collected during an investigation. Data may be quantitative, qualitative, or ranked.

When planning a biological investigation, it is important to consider the type of data that will be collected. It is best to collect **quantitative** or numerical data, because it is mathematically versatile and easier to analyse it objectively (without **bias**).



A: Skin colour



B: Eggs per nest



C: Tree trunk diameter



D: Rate of growth in seedlings

1. For each of the photographic examples A-D above, classify the data as quantitative, ranked, or qualitative:

(a) Skin colour: \_\_\_\_\_

(b) Number of eggs per nest: \_\_\_\_\_

(c) Tree trunk diameter: \_\_\_\_\_

(d) Rate of seedling growth: \_\_\_\_\_

2. Why is it best to collect quantitative data where possible in biological studies? \_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_

3. Give an example of data that could not be collected quantitatively and explain your answer: \_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_



# 4 Planning a Practical Investigation

**Key Idea:** Practical work carried out in a careful and methodical way makes analysis of the results much easier. A major part of any practical investigation is collecting the data. Practical work may be laboratory or field based. Typical laboratory based **experiments** involve investigating how a biological response is affected by manipulating a particular **variable**, e.g. temperature. The data collected for a

quantitative practical task should be recorded systematically, with due attention to safe practical techniques, a suitable quantitative method, and accurate measurements to an appropriate degree of **precision**. If your quantitative practical task is carried out well, and you have taken care throughout, your evaluation of the experimental results will be much more straightforward and less problematic.

## Carrying out your practical work



### Preparation

Familiarise yourself with the equipment and its set up. Calibrate equipment if necessary to give accurate measurements. Read through the methods and identify key stages and how long they will take.



### Execution and recording

Know how you will take your measurements and how often. Record your results systematically as you go in a log book. You could record results a hand-written **table** or in a spreadsheet. If using a data logger, data will be logged.



### Analysis and reporting

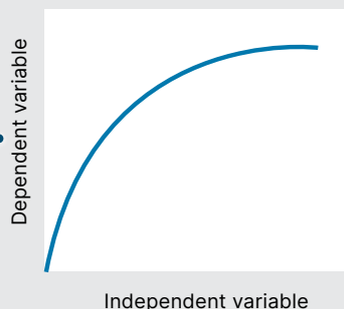
Analyse the data. Tables can summarise data. **Graphs** present the data to show patterns and **trends**. **Statistical tests** can determine the significance of results. Present your findings, e.g. as a poster, a digital presentation, or an oral report.

## Identifying variables

A variable is any characteristic or property able to take any one of a range of values. Investigations often look at the effect of changing one variable on another. It is important to identify all variables in an investigation: independent, dependent, and controlled, although there may be nuisance factors of which you are unaware. In all fair tests, only one variable is changed by the investigator.

### Dependent variable

- ▶ Measured during the investigation.
- ▶ Recorded on the y axis of the graph.



### Controlled variables

- ▶ Factors that are kept the same or controlled.
- ▶ List these in the method, as appropriate to your own investigation.

### Independent variable

- ▶ Set by the experimenter.
- ▶ Recorded on the graph's x axis.

## Experimental controls

A **control** refers to a standard or reference treatment or group in an experiment. It is the same as the experimental (test) group, except that it lacks the one variable being manipulated by the experimenter. Controls are used to demonstrate that the response in the test group is due a specific variable (e.g. temperature). The control undergoes the same preparation, experimental conditions, **observations**, measurements, and analysis as the test group. This helps to ensure that responses observed in the treatment groups can be reliably interpreted.



- ▶ The experiment above tests the effect of a certain nutrient on microbial growth. All the agar plates are prepared in the same way, but the control plate does not have the test nutrient applied.
- ▶ Each plate is inoculated from the same stock solution, incubated under the same conditions, and examined at the same set periods. The control plate sets the baseline; any growth above that seen on the control plate is attributed to the nutrient.



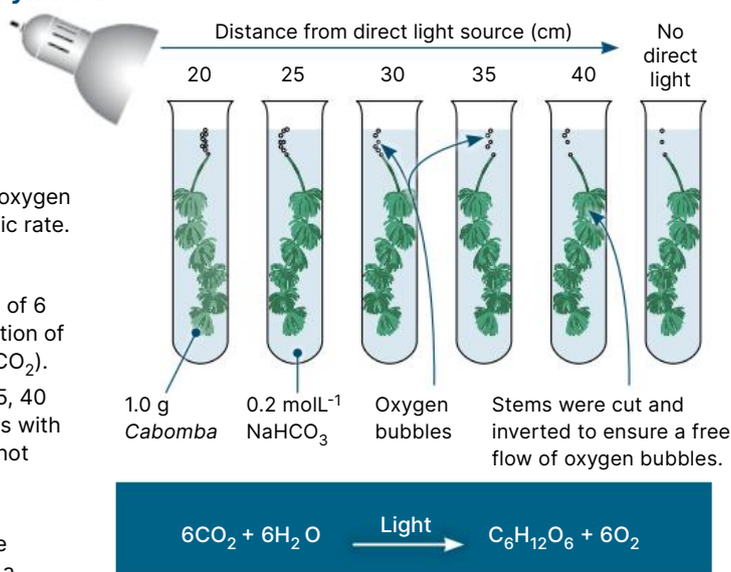
## Investigation: Effect of light on rate of photosynthesis

### Background

The aquarium plant, *Cabomba aquatica*, will produce a stream of oxygen bubbles when illuminated. The oxygen bubbles are a waste product of the process of photosynthesis (overall equation below right), which produces glucose ( $C_6H_{12}O_6$ ) for the plant. The **rate** of oxygen production provides an approximation of photosynthetic rate.

### The method

- ▶  $6 \times 1.0$  g of *Cabomba* stems were placed into each of 6 test-tubes filled with 10 mL room temperature solution of  $0.2 \text{ mol L}^{-1}$  sodium hydrogen carbonate (to supply  $CO_2$ ).
- ▶ Test tubes were placed at distances (20, 25, 30, 35, 40 cm) from a 60W light source (light intensity reduces with distance at a predictable rate). One test tube was not exposed to the light source.
- ▶ Before recording, the *Cabomba* stems were left to acclimatise to the new light level for 5 minutes. The bubbles emerging from the stem were counted for a period of three minutes at each distance.



1. Write a suitable aim for this experiment: \_\_\_\_\_  
\_\_\_\_\_
2. Write a possible hypothesis for this experiment: \_\_\_\_\_  
\_\_\_\_\_
3. (a) What is the independent variable in this experiment? \_\_\_\_\_  
(b) What is the range of values for the independent variable? \_\_\_\_\_  
(c) Name the unit for the independent variable: \_\_\_\_\_  
(d) How could you better quantify the independent variable? \_\_\_\_\_
4. (a) What is the dependent variable in this experiment? \_\_\_\_\_  
(b) Name the unit for the dependent variable: \_\_\_\_\_  
(c) What equipment might have made it easier to record the response of the dependent variable accurately? Predict when it would have been most needed: \_\_\_\_\_  
\_\_\_\_\_  
(d) What is the sample size for each treatment? \_\_\_\_\_  
(e) What could you change in the design of the experiment to guard against unexpected or erroneous results? \_\_\_\_\_  
\_\_\_\_\_
5. Which tube is the control for this experiment? \_\_\_\_\_
6. Identify two assumptions being made about this system:  
(a) \_\_\_\_\_  
(b) \_\_\_\_\_
7. Identify one variable that might have been controlled in this experiment, and how it could have been monitored: \_\_\_\_\_  
\_\_\_\_\_
8. How might you test the gas being produced is oxygen: \_\_\_\_\_  
\_\_\_\_\_

# 5 Safety and Ethical Guidelines

**Key Idea:** In practical work, research, and reporting you should act in accordance with safety and ethical guidelines. Scientific research, no matter what the level, should be carried out in accordance with ethical and safety guidelines. These guidelines apply to health and safety in the laboratory

and field, risk assessment, and correct use of equipment. Also, consider the ethical issues associated with animal welfare, privacy and personal information, and environmental impact. Ethical considerations also apply to reporting of data and honest use and acknowledgement of reference material.

## Safe practices in the laboratory

- ▶ Laboratories have three main hazards: chemicals, biological materials, and physical hazards.
- ▶ **Chemical hazards** are substances that can be harmful if ingested, absorbed, or inhaled, such as cleaning agents and reagents.
- ▶ **Biological hazards** include potentially harmful biological materials such as microbial samples and animal tissue.
- ▶ **Physical hazards** in can come from the environment or equipment, including incorrect equipment use, clutter, and slippery floors.



## Assessing and reducing risk in the lab



- ▶ Before you start any investigation or experiment, you should identify any potential hazards. Chemical safety data sheets identify the risks in using various chemicals.
- ▶ You should wear appropriate personal protection equipment (PPE) e.g. lab coat, gloves, safety glasses.
- ▶ Pay attention to warnings and hazard notices on chemicals and equipment.
- ▶ Be familiar with the safe use of the equipment you are going to use.
- ▶ Keep your work space clean and tidy.



1. (a) Identify potential health and safety issues associated with the dissection of the pig kidney in the photo (left):

---



---



---



---

(b) What has been done to reduce potential risks?

---



---



---



2. (a) Identify two potential safety or health hazards associated with the inspection of bacterial colonies in the photo (bottom left):

---



---



---

(b) What could be done to reduce these risks?

---



---



---



### Health and safety in fieldwork

Field studies present their own their own set of ethical and safety considerations. The Australian environment can be harsh, and bushland may contain wildlife, plants, and geographic features that may be hazardous.

- ▶ Assess the potential hazards of the area before beginning any field studies. Field studies may also require some follow-up laboratory work, especially if samples found in the field need to be identified or processed. In these cases, follow lab health and safety guidelines.
- ▶ Identify potential hazards before you start and become knowledgeable about their risks. In the field, this includes the weather as well as your surroundings. Be aware of hidden hazards such as wasp nests, stinging plants or animals, or territorial birds.



### Honesty and ethical issues

- ▶ If you are sampling or collecting live organisms, you must consider the environmental impact of any sampling procedures, return live organisms to the same place if possible, respect the natural environment, and handle animals in a way that minimises stress or damage to them. Plan your study to minimise your impact on the natural environment.
- ▶ Even if your results are not what you expected, you should report you results and data as they are. It is misleading and unethical to change your results to fit your hypothesis.
- ▶ Any intellectual property produced by others (e.g. photographs, data) should be acknowledged. Representing the work of others as your own is plagiarism.



Be meticulous in maintaining an accurate logbook, acknowledge all your sources, and reference cited works accurately. Act ethically and responsibly in all aspects of your research, including in the disposal of biological material.

3. Describe the potential ethical issues associated with each of the following investigative scenarios:

(a) A vegetation survey in a sensitive ecological area: \_\_\_\_\_

---



---

(b) A lab-based experimental investigation of salinity tolerance in shore crabs: \_\_\_\_\_

---



---

(c) Deriving values in a set of experimental measurements by interpolation because you missed a day of recordings: \_\_\_\_\_

---



---

4. Describe two reasons why acknowledgement of sources and correct reference of cited works is important:

(a) \_\_\_\_\_

---



---

(b) \_\_\_\_\_

---



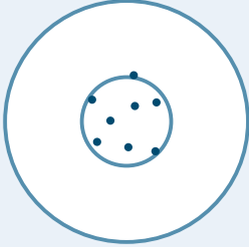
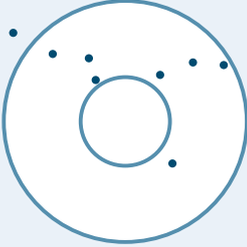
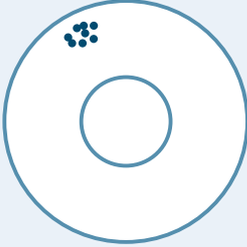
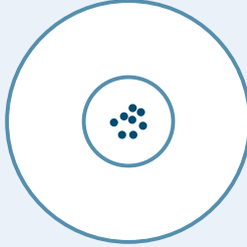
---

# 6

## Accuracy and Precision

**Key Idea:** Accuracy refers to the correctness of a measurement (how true it is to the real value). Precision refers to how close the measurements are to each other. **Accuracy** refers to how close a measured or derived value is to its true value. Simply put, it is the correctness of the measurement. **Precision** refers to how close repeated

measurements are to each other, i.e. the ability to be exact. A balance with a fault in it could give very precise (repeatable) but inaccurate (untrue) results. Data can only be reported as accurately as the measurement of the apparatus allows and is often expressed as **significant figures** (the digits in a number which express meaning to a degree of accuracy).

Accurate but imprecise	Inaccurate and imprecise	Precise but inaccurate	Accurate and precise
			
<p>The measurements are all close to the true value but quite spread apart.</p>	<p>The measurements are all far apart and not close to the true value.</p>	<p>The measurements are all clustered close together but not close to the true value.</p>	<p>The measurements are all close to the true value and also clustered close together.</p>
<p><b>Analogy:</b> The arrows are all close to the bullseye.</p>	<p><b>Analogy:</b> The arrows are spread around the target.</p>	<p><b>Analogy:</b> The arrows are all clustered close together but not near the bullseye.</p>	<p><b>Analogy:</b> The arrows are clustered close together near the bullseye.</p>
<p><b>Increasing precision</b> The accuracy of a measurement refers to how close the measured value is to the true value. The precision of a measurement relates to its repeatability. In most laboratory work, we usually assume a piece of equipment (e.g. a pipette) performs accurately, so precise measurement is the most important consideration. We can test precision by taking repeated measurements from individual samples. Precision and <b>reliability</b> are synonymous and describe how dependably an <b>observation</b> is the same when repeated.</p>			
<p><b>Increasing accuracy</b> Population studies present us with an additional problem. When a researcher makes measurements of some <b>variable</b> (e.g. fish length), they are usually trying to obtain an <b>estimate</b> of the true value for a parameter of interest (e.g. the <b>mean</b> size of fish). Populations are variable, so we can more accurately estimate a population parameter if we take a large number of random samples from the population.</p>		<p>A digital device such as the pH meter (above left) will deliver precise measurements, but its accuracy will depend on correct calibration. The precision of measurements taken with instruments such as callipers (above) will depend on the skill of the operator. Precise measurements provide reliable data.</p>	

1. Distinguish between accuracy and precision: \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_
2. Describe why it is important to take measurements that are both accurate and precise: \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_
3. A researcher is trying to determine the temperature at which an enzyme becomes denatured. Their temperature probe is incorrectly calibrated. Discuss how this might affect the accuracy and precision of the data collected:  
 \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_



## Accuracy and equipment

The accuracy of a measurement can be increased by increasing the number of measurements taken. For example, the accuracy of the mean mass of individuals in a population can be increased by increasing the number of individuals measured (i.e. increasing the sample size).

In many cases, the accuracy of the measuring equipment needs to be taken into account. For example, electronic balances may give readings to one or more decimal places based on their accuracy (laboratory balances may read to a hundred thousandths of a gram).

The **table** below illustrates the difference between a balance weighing to 0.1 of a gram and 0.01 of a gram.

Sample	1	2	3	4	5	Mean
Mass (g) (2 s.f)	1.1	1.2	1.4	1.2	1.3	1.2
Mass (g) (3 s.f)	1.12	1.23	1.44	1.19	1.28	1.25

The difference in mean mass is slight (just 0.05 g) but over larger samples or larger masses the differences can add up.

The table above also shows the importance of significant figures (s.f). The actual numerical mean for the second row is 1.252. However because we are measuring to three significant figures we cannot be sure of the final number, thus the answer must be given in the same significant figures as the measurements.

4. The period of a pendulum is based on the length of the pendulum and the mass at its end. Two students measure the time it takes for a pendulum to swing back and forth (its period). Student A measures three individual swings and calculates a mean value. Student B measures three sets of ten swings and calculates a mean. Each student measures the accuracy of the timer as 0.2 seconds. The results are shown below:

Student A	
	Time for swing (s)
1	2.7
2	2.1
3	2.5
Mean (1 swing)	

Student B	
Set	Time for ten swings (s)
1	20.3
2	20.1
3	19.8
Mean (10 swings)	
1 swing	

(a) Calculate the mean for each student's results and the time for one swing for student B.

(b) Explain why student B's results are more accurate than student A's: \_\_\_\_\_

\_\_\_\_\_

5. In a class of 20 students, the individual heights of the students in cm are: 135, 139, 141, 146, 147, 149, 156, 151, 158, 155, 156, 159, 161, 167, 162, 163, 161, 172, 171, 170.

(a) Calculate the mean height of the students: \_\_\_\_\_

(b) A person takes a sample of five of the students: 139, 151, 162, 172, 170. Calculate the mean of the sample and comment on its accuracy:

\_\_\_\_\_

(c) A second person takes a sample of ten of the students: 135, 146, 147, 156, 155, 156, 161, 167, 162, 170. Calculate the mean of the sample and comment on its accuracy:

\_\_\_\_\_

\_\_\_\_\_

## Reducing error

Sometimes, reducing error requires taking more measurements over a longer period of time. For example, hypothetical waves breaking on a shore do so with a relatively regular frequency. Recording the time between one wave breaking and the next depends when the wave is defined as breaking. This may be difficult to determine precisely for each individual wave and the waves may be breaking too quickly to allow enough time to elapse for recordings to be made accurately (especially if the timer is being started and stopped by a person).

To increase the accuracy of measuring the period between each wave, it is best to record the time for a number of waves to break (e.g. 10) and divide by that number to obtain the period between each wave. This has the effect of allowing for slight variations in the period and reduces the total error in the measurement.

**Example:** Actual wave period: 5.0 seconds.  
Accuracy of timer (i.e. reaction speed) 0.3 seconds

Measurements of individual periods (seconds): 5.4, 5.7, 5.7, 5.8, 4.5, 4.6, 5.7, 5.8, 5.1, 5.3, Mean: 5.4

In each measurement above, the error is about 0.3 seconds producing an error of up to 6.7% ( $0.3 \div 4.5 \times 100$ ) of the recorded value of a wave period.

If the time recorded for ten waves to break was 51.1 seconds, then the time for one wave to break is 5.1 seconds. The error is spread over the whole 51.1 seconds ( $0.3 \div 51.1$ ) and thus is much smaller at just 0.6% of the wave period.

# 7 Working with Numbers

**Key Idea:** Using correct mathematical notation and being able to carry out simple calculations and conversions are fundamental skills in biology.

Mathematics is used in biology to analyse, interpret, and

compare data. It is important that you are familiar with mathematical notation (the language of mathematics) and can confidently apply some basic mathematical principles and calculations to your data.

## Decimal and standard form

**Decimal form** (also called ordinary form) is the longhand way of writing a number (e.g. 15,000,000). Very large or very small numbers can take up too much space if written in decimal form and are often expressed in a condensed standard form. For example, 15,000,000 is written as  $1.5 \times 10^7$  in standard form.

In standard form a number is always written as  $A \times 10^n$ , where A is a number between 1 and 10, and n (the exponent) indicates how many places to move the decimal point. n can be positive or negative.

For the example above, A = 1.5 and n = 7 because the decimal point moved seven places (see below).

$$15\ 000\ 000 = 1.5 \times 10^7$$

Small numbers can also be written in standard form. The exponent (n) will be negative. For example, 0.00101 is written as  $1.01 \times 10^{-3}$ .

$$0.00101 = 1.01 \times 10^{-3}$$

Converting can make calculations easier. Work through the following example to solve  $4.5 \times 10^4 + 6.45 \times 10^5$ .

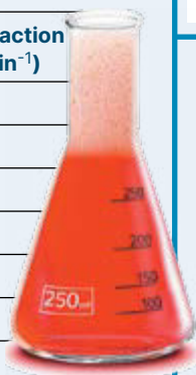
- Convert  $4.5 \times 10^4 + 6.45 \times 10^5$  to decimal form:  
\_\_\_\_\_
- Add the two numbers together: \_\_\_\_\_
- Convert to standard form: \_\_\_\_\_

## Rates

**Rates** are expressed as a measure per unit of time and show how a **variable** changes over time. Rates are used to provide meaningful comparisons of data that may have been recorded over different time periods.

Often rates are expressed as a **mean** rate over the duration of the measurement period, but it is also useful to calculate the rate at various times to understand how rate changes over time. The **table** below shows the reaction rates for a gas produced during a chemical reaction. A worked example for the rate at 4 minutes is provided below the table.

Time (Minute)	Cumulative gas produced (cm <sup>3</sup> )	Rate of reaction (cm <sup>3</sup> min <sup>-1</sup> )
0	0	0
2	34	17
4	42	4*
6	48	3
8	50	1
10	50	0



\* Gas produced between 2- 4 min:  $42 \text{ cm}^3 - 34 \text{ cm}^3 = 8 \text{ cm}^3$   
Rate of reaction between 2- 4 min:  $8 \div 2 \text{ min} = 4 \text{ cm}^3 \text{ min}^{-1}$

## Conversion factors and expressing units

Measurements can be converted from one set of units to another by the use of a conversion factor.

A conversion factor is a numerical factor that multiplies or divides one unit to convert it into another. Conversion factors are commonly used to convert non-SI units to SI units (e.g. converting pounds to kilograms). Note that mL and cm<sup>3</sup> are equivalent, as are L and dm<sup>3</sup>.

In the space below, convert 5.6 cm<sup>3</sup> to mm<sup>3</sup> (1 cm<sup>3</sup> = 1000 mm<sup>3</sup>):

4. \_\_\_\_\_

The value of a variable must be written with its units where possible. SI units or their derivations should be used in recording measurements: volume in cm<sup>3</sup>, dm<sup>3</sup>, or litre (L), mass in kilograms (kg) or grams (g), length in metre (m), time in seconds (s).

For example the rate of oxygen consumption would be expressed as: Oxygen consumption (cm<sup>3</sup>g<sup>-1</sup>s<sup>-1</sup>)

## Estimates

When carrying out mathematical calculations, typing the wrong number into your calculator can put your answer out by several orders of magnitude. An **estimate** is a way of roughly calculating what answer you should get, and helps you decide if your final calculation is correct.

Numbers are often rounded to help make estimation easier. The rounding rule is, if the next digit is 5 or more, round up. If the next digit is 4 or less, it stays as it is.

For example, to estimate  $6.8 \times 704$  you would round the numbers to  $7 \times 700 = 4900$ . The actual answer is 4787, so the estimate tells us the answer (4787) is probably right.

Use the following examples to practise estimating:

- $43.2 \times 1044$ : \_\_\_\_\_
- $3.4 \times 72 \div 15$ : \_\_\_\_\_  
\_\_\_\_\_
- $658 \div 22$ : \_\_\_\_\_

## Probability

Probability is how likely something is to happen. It is an important part of biology. Its uses include calculating the statistical significance of a difference between means or the probability of an event occurring.

The probability of an event ranges from 0 to 1. The sum of all probabilities equals 1.

**Product rule:** for independent events A and B the probability (P) of A and B occurring is  $P(A) \times P(B)$ . For example, the probability two children born one after the other both being male is  $0.5 \times 0.5 = 0.25$ .

**Sum rule:** For mutually exclusive events Y and Z the probability that one will occur (Y or Z) is  $P(Y) + P(Z)$ . E.g. in an Aa x Aa cross, the probability a person will have a dominant phenotype =  $0.25 + 0.5 = 0.75$ .



# 8

# Fractions, Percentages, and Ratios

**Key Idea:** Percentages and ratios are alternative ways to express fractions. All forms are commonly used in biology. The data collected in the field or laboratory are called raw data. Data are often expressed in ways that make them easy

to understand, visualise, and work with. Fractions, **ratios**, and **percentages** are widely used in biology and are often used to provide a meaningful comparison of sample data where the sample sizes are different.

### Fractions

- ▶ Fractions express how many parts of a whole are present.
- ▶ Fractions are expressed as two numbers separated by a solidus (/) (e.g. 1/2).
- ▶ The top number is the numerator. The bottom number is the denominator. The denominator can not be zero.
- ▶ Fractions are often written in their simplest form (the top and bottom numbers cannot be any smaller, while still being whole numbers). Simplifying makes working with fractions easier.



In a class of 20 students, five had blue eyes. This fraction is 5/20. To simplify this fraction, divide the numerator and denominator by a common factor (a number which both are divisible by). In this instance the lowest common factor is five (1/4). To add fractions with different denominators, obtain a common denominator, add numerators, then simplify.

### Ratios

- ▶ Ratios give the relative amount of two or more quantities, and provide an easy way to identify patterns.
- ▶ Ratios do not require units.
- ▶ Ratios are expressed as **a : b**.
- ▶ Ratios are calculated by dividing all the values by the smallest number.



Pea pod shape:  
Ratio = 2.95 : 1



Pea seed shape and colour:  
Ratio = 9 : 2.8 : 2.9 : 1

Example: Calculating phenotype ratios in Mendelian genetics.

### Percentages

- ▶ Percentages are expressed as a fraction of 100 (e.g. 20/100 = 20%).
- ▶ Percentages provide a clear expression of what **proportion** of data fall into any particular category, e.g. for pie graphs.
- ▶ Allows meaningful comparison between different samples.
- ▶ Useful to monitor change (e.g. % increase from one year to the next).

Volume of food colouring (cm <sup>3</sup> )	Volume of water (cm <sup>3</sup> )	Concentration of solution (%)
10	0	100
8	2	80
6	4	60
4	6	40
2	8	20
0	10	0



Example: Producing standards for a calibration curve.

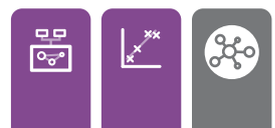
- (a) A student prepared a slide of the cells of an onion root tip and counted the cells at various stages in the cell cycle. The results are presented in the table (right). Calculate the ratio of cells in each stage (show your working):  
 \_\_\_\_\_  
 \_\_\_\_\_
- (b) Assuming the same ratio applies in all the slides examined in the class, calculate the number of cells in each phase for a cell total count of 4800.
- Simplify the following fractions:

Cell cycle stage	No. of cells counted	No. of cells calculated
Interphase	140	
Prophase	70	
Telophase	15	
Metaphase	10	
Anaphase	5	
Total	240	4800

(a) 3/9: \_\_\_\_\_ (b) 84/90: \_\_\_\_\_ (c) 11/121: \_\_\_\_\_

- In the fraction example pictured above 5/20 students had blue eyes. In another class, 5/12 students had blue eyes. What fraction of students had blue eyes in both classes combined?  
 \_\_\_\_\_
- The total body mass and lean body mass for women with different body types is presented in the table (right). Complete the table by calculating the % lean body mass column.

Women	Body mass (kg)	Lean body mass (kg)	% lean body mass
Athlete	50	38	
Lean	56	41	
Normal weight	65	46	
Overweight	80	48	
Obese	95	52	



# 9

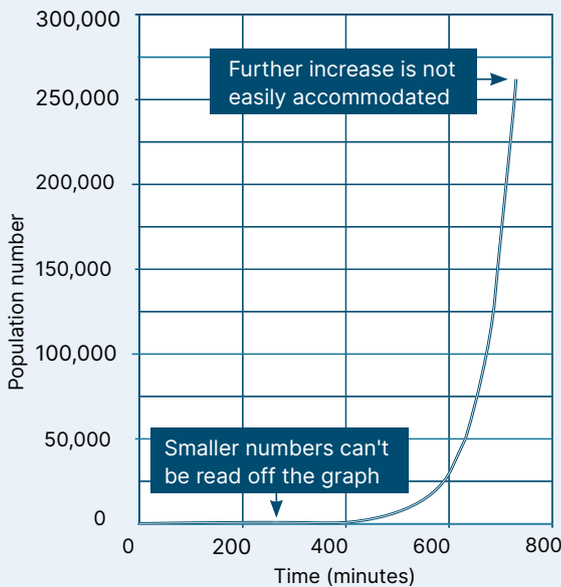
## Dealing with Large Numbers

**Key Idea:** Exponential functions are common in biology and may involve very large numbers. Exponential changes in numbers are defined by a function, which is simply a rule that allows us to calculate an output for any given input. In biology, numerical data indicating scale can often decrease or increase exponentially. Examples include the exponential

growth of populations, exponential decay of radioisotopes, and the pH scale. Exponential changes are defined by a function that allows us to calculate an output for any input. The numbers associated with exponential growth can be very large and are often log transformed. Log transformations of exponential numbers can make them easier to handle.

### Exponential function

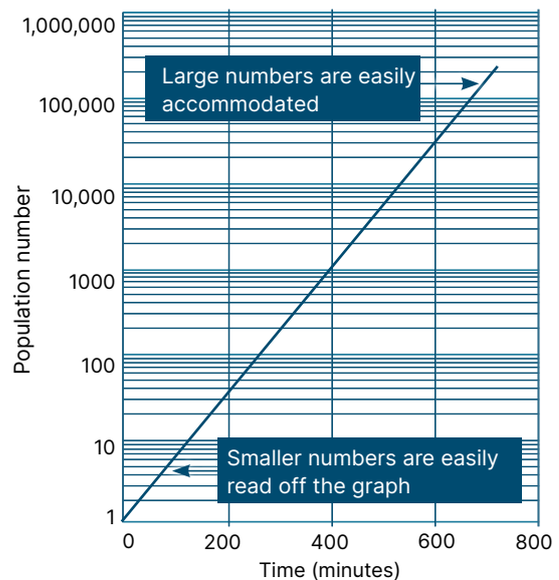
- ▶ Exponential growth occurs at an increasingly rapid rate in **proportion** to the growing total number or size.
- ▶ In an exponential function, the base number is fixed (constant) and the exponent is **variable**.
- ▶ The equation for an exponential function is  $y = c^x$ .
- ▶ Exponential growth and decay (reduction) are possible.
- ▶ Exponential changes in numbers are easy to identify because the curve has a J-shape appearance due to its increasing steepness over time.
- ▶ An example of exponential growth is the growth of a microbial population in an unlimited, optimal growth environment.



**Example:** Cell growth in a yeast culture where growth is not limited by lack of nutrients or build up of toxins.

### Log transformations

- ▶ A log transformation makes very large numbers easier to work with. The log of a number is the exponent to which a fixed value (the base) is raised to get that number. So  $\log_{10}(1000) = 3$  because  $10^3 = 1000$ .
- ▶ Both  $\log_{10}$  (common logs) and  $\log_e$  (natural logs or  $\ln$ ) are commonly used.
- ▶ Log transformations are useful for data where there is an exponential increase or decrease in numbers. In this case, the transformation will produce a straight line plot.
- ▶ To find the  $\log_{10}$  of a number, e.g. 32, using a calculator, key in  $\log(32) = .$  The answer should be 1.51.
- ▶ Alternatively, the untransformed data can be plotted directly on a log-linear scale (as below). This is not difficult. You just need to remember that the log axis runs in exponential cycles. The paper makes the log for you.



**Example:** The same yeast cell growth plotted on a log-linear scale. The y axis present 6 exponential cycles.

1. Why is it useful to plot exponential growth using semi-log paper? \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_
2. What would you do to show yeast exponential growth (left plot above) as a straight line plot on normal graph paper?  
 \_\_\_\_\_  
 \_\_\_\_\_
3. Log transformations are often used when a value of interest ranges over several orders of magnitude. Can you think of other examples of data from the natural world where the data collected might show this behaviour?  
 \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_



**Key Idea:** This activity allows you to practise working with data and applying the skills you have learned in previous activities.

1. Complete the transformations for each of the tables below. The first value is provided in each case.

(a) Photosynthetic rate at different light intensities

Light intensity (%)	Average time for leaf disc to float (min)	Reciprocal of time* ( $\text{min}^{-1}$ )
100	15	0.067
50	25	
25	50	
11	93	
6	187	

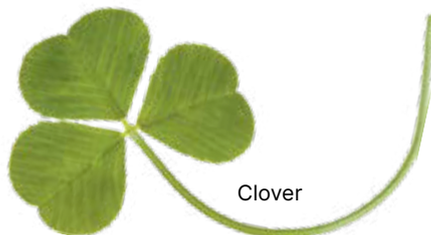
\* Reciprocal of time gives a crude measure of rate.

(b) Plant water loss using a bubble potometer

Time (min)	Pipette arm reading ( $\text{cm}^3$ )	Plant water loss ( $\text{cm}^3 \text{min}^{-1}$ )
0	9.0	–
5	8.0	0.2
10	7.2	
15	6.2	
20	4.9	

(c) Incidence of cyanogenic clover in different areas

Clover plant type	Frost free area		Frost prone area		Totals
	Number	%	Number	%	
Cyanogenic	124	78	26		
Acyanogenic	35		115		
Total	159				



(d) Frequency of size classes in a sample of eels

Size class (mm)	Frequency	Relative frequency (%)
0-50	7	2.6
50-99	23	
100-149	59	
150-199	98	
200-249	50	
250-299	30	
300-349	3	
Total	270	

2. Convert the following decimal form numbers to standard form:

(a) 8970 \_\_\_\_\_ (b) 0.046 \_\_\_\_\_ (c) 1,467,851 \_\_\_\_\_

3. Convert the following standard form numbers to decimal form:

(a)  $4.3 \times 10^{-1}$  \_\_\_\_\_ (b)  $0.0031 \times 10^{-2}$  \_\_\_\_\_ (c)  $6.2 \times 10^4$  \_\_\_\_\_

4. (a) The table on the right shows the nutritional label found on a can of chilli beans. Use the information provided to complete the table by calculating the percentage composition for each of the nutritional groups listed:

(b) How much of the total carbohydrate is made up of:

Dietary fibre? \_\_\_\_\_

Sugars? \_\_\_\_\_

(c) Manufacturers do not have to state the volume of water, which makes up the remainder of the serving size. What percentage of the can of beans is water?

\_\_\_\_\_  
\_\_\_\_\_

Chilli beans nutrition facts Serving size 1 cup (253 g)		
Amount per serving	% composition	
<b>Total fat</b>	8 g	
– Saturated fat	3 g	
<b>Total carbohydrate</b>	22 g	
– Dietary fibre	9 g	
– Sugars	4 g	
<b>Protein</b>	25 g	



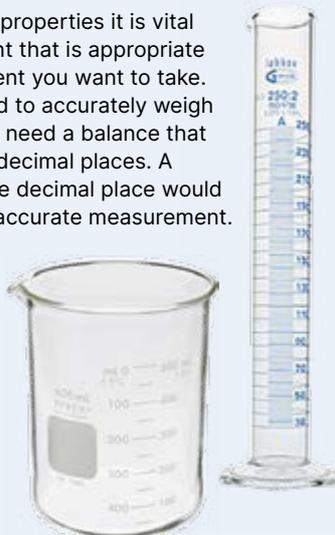
**Key Idea:** The apparatus used in experimental work must be appropriate for the experiment or analysis and it must be used correctly to eliminate experimental errors. Using scientific equipment can generate experimental errors.

These can be reduced by selecting the right equipment for what you want to measure and by using it correctly. Some error is inevitable, but evaluating experimental error helps to interpret and assess the validity of the results.

### Selecting the correct equipment

When measuring physical properties it is vital that you choose equipment that is appropriate for the type of measurement you want to take. For example, if you wanted to accurately weigh out 5.65 g of sucrose, you need a balance that accurately weighs to two decimal places. A balance that weighs to one decimal place would not allow you to make an accurate measurement.

Study the glassware (right). Which would you use if you wanted to measure 225 mL? The graduated cylinder has graduations every 10 mL whereas the beaker has graduations every 50 mL. It would be more accurate to measure 225 mL in a graduated cylinder.



### Percentage errors

Percentage error is a way of mathematically expressing how far out your result is from the ideal result. The equation for measuring percentage error is:

$$\frac{\text{experimental value} - \text{ideal value}}{\text{ideal value}} \times 100$$

For example, to determine the **accuracy** of a 5 mL pipette, dispense 5 mL of water from the pipette and weigh the dispensed volume on a balance.

The mass (g) = volume (mL). The volume is 4.98 mL.

$$\frac{\text{experimental value (4.98)} - \text{ideal value (5.0)}}{\text{ideal value (5.0)}} \times 100$$

The percentage error = -0.4% (the negative sign tells you the pipette is dispensing **less** than it should).

### Recognising potential sources of error



It is important to know how to use equipment correctly to reduce errors. A spectrophotometer measures the amount of light absorbed by a solution at a certain wavelength. This information can be used to determine the concentration of the absorbing molecule (e.g. density of bacteria in a culture). The more concentrated the solution, the more light is absorbed. Incorrect use of the spectrophotometer can alter the results. Common mistakes include incorrect calibration, errors in sample preparation, and errors in sample measurement.



A cuvette (left) is a small clear tube designed to hold spectrophotometer samples. Inaccurate readings occur when:

- ▶ The cuvette is dirty or scratched (light is absorbed giving a falsely high reading).
- ▶ Some cuvettes have a frosted side to aid alignment. If the cuvette is aligned incorrectly, the frosted side absorbs light, giving a false reading.
- ▶ Not enough sample is in the cuvette and the beam passes over, rather than through the sample, giving a lower absorbance reading.

1. Assume that you have the following measuring devices available: 50 mL beaker, 50 mL graduated cylinder, 25 mL graduated cylinder, 10 mL pipette, 10 mL beaker. What would you use to accurately measure:

(a) 21 mL: \_\_\_\_\_ (b) 48 mL: \_\_\_\_\_ (c) 9 mL: \_\_\_\_\_

2. Calculate the percentage error for the following situations (show your working):

(a) A 1 mL pipette delivers a measured volume of 0.98 mL: \_\_\_\_\_

\_\_\_\_\_

(b) A 10 mL pipette delivers a measured volume of 9.98 mL: \_\_\_\_\_

\_\_\_\_\_

(c) The pipettes used in (a) and (b) above both under-delivered 0.02 mL, yet the percentage errors are quite different. Use this data to describe the effect of volume on percentage error:

\_\_\_\_\_

\_\_\_\_\_



**Key Idea:** Graphs are useful for visually displaying numerical data, trends, and relationships between variables.

**Graphs** are an excellent way to summarise **trends** in data or relationships between different **variables**. Presenting graphs properly requires attention to a few basic details, including correct orientation and labelling of the axes, and accurate

plotting of points. Before representing data graphically, it is important to identify the kind of data you have. Common graphs include scatter plots and line graphs (for continuous data), and bar charts (for categorical data). For continuous data with calculated **means**, points can be connected. On scatter plots, a line of best fit is often drawn.

### Guidelines for line graphs

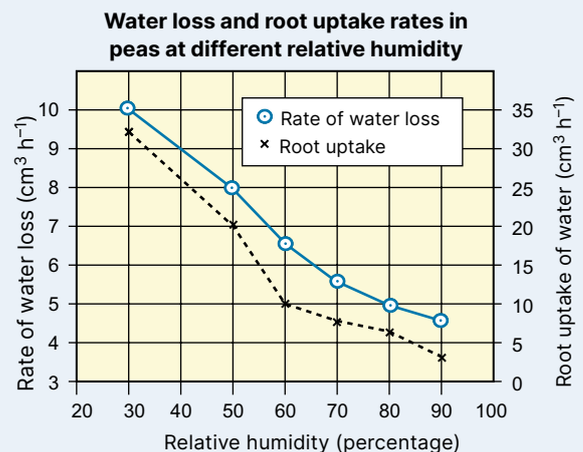
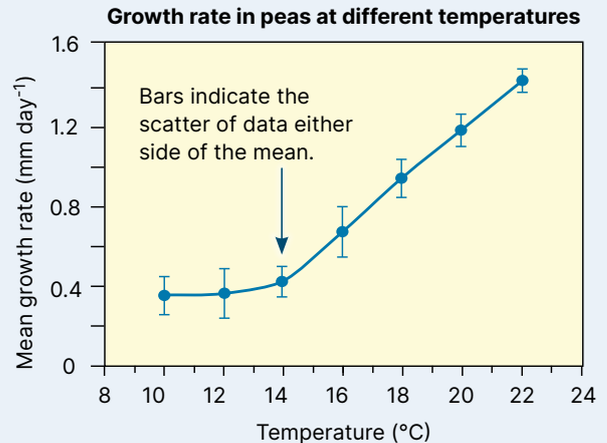
**WHEN TO USE:** Use a line graph when both variables are continuous and one variable (the **independent variable**) affects another, the **dependent variable**. Important features include:

- ▶ The data must be continuous for both variables. The independent variable is often time or experimental treatment. The dependent variable is generally the biological response.
- ▶ The relationship between two variables can be represented as a continuum and the data points are plotted accurately and connected directly (point to point).
- ▶ Line graphs may be drawn with measure of error (right). The data are presented as points (which are calculated means), with error bars above and below, indicating the variability in the data (e.g. standard deviation).

### Plotting multiple data sets

A single figure (graph) can be used to show two or more data sets, i.e. more than one curve can be plotted per set of axes. This type of presentation is useful when comparing the trends for two or more treatments, or the response of one species against the response of another. Important points regarding this format are:

- ▶ If the two data sets use the same measurement units and a similar range of values for the dependent variable, one scale on the y axis is used.
- ▶ If the two data sets use different units and/or have a very different range of values for the dependent variable, two scales for the y axis are used (see right). The scales can be adjusted if necessary to avoid overlapping plots.
- ▶ The two curves are distinguished with a key.

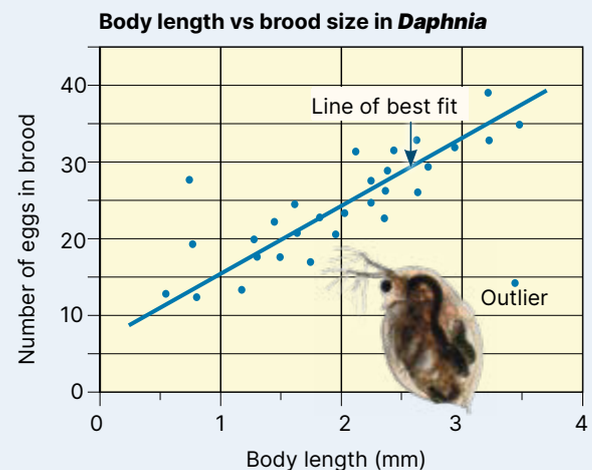


### Guidelines for scatter graphs

**WHEN TO USE:** Use a scatter graph to display continuous data where there are two interdependent variables.

- ▶ The data must be continuous for both variables.
- ▶ There is no independent variable, but the variables are often correlated, i.e. they vary together in a predictable way.
- ▶ Useful to determine the relationship between two variables.
- ▶ The points on the graph are not connected, but a line of best fit is often drawn through the points to show the relationship between the variables (this may be computer generated with a value assigned to the goodness of the fit).
- ▶ Obvious outliers (points that lie well outside most of the scatter) are usually disregarded from analyses.

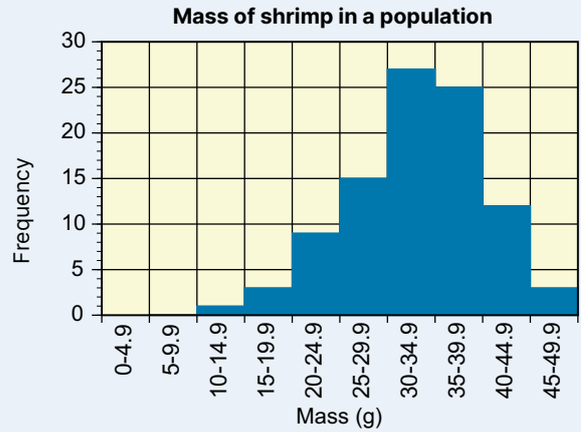
**Interpolation:** For both line and scatter graphs, the fitted line can be used to find an unknown value inside the set of data points. This is called interpolation.



**Guidelines for histograms**

**WHEN TO USE:** Use a histogram when one variable is continuous and the other is a frequency (counts). These plots produce a frequency distribution because the y-axis shows the number of times a measurement or value was obtained. Important features of histograms include:

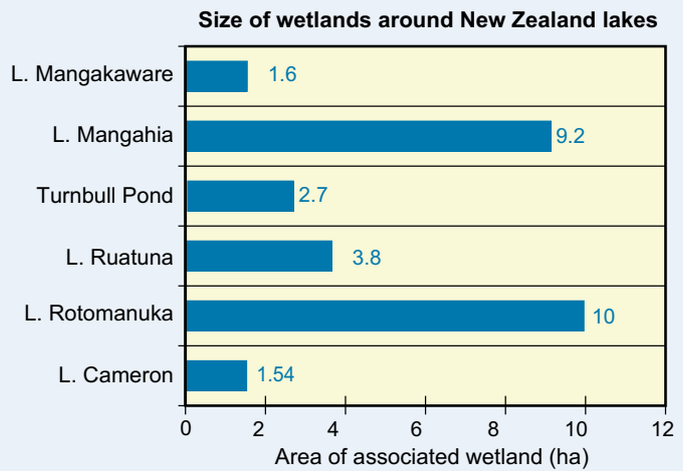
- ▶ The data are numerical and continuous (e.g. height or weight) so the bars touch.
- ▶ The x-axis usually records the class interval. The y-axis usually records the number of individuals in each class interval.



**Guidelines for bar and column graphs**

**WHEN TO USE:** Use a bar or column graph for data that are non-numerical and discrete (categorical) for one variable. There are no dependent or independent variables. Important features include:

- ▶ Data for one variable are discontinuous, non-numerical categories (e.g. place, colour, species), so the bars do not touch.
- ▶ Data values may be entered by the bars if you wish.
- ▶ Multiple sets of data can be displayed side by side to compare (e.g. males and females in the same age group).
- ▶ Axes may be reversed so that the categories are on the x axis, i.e. the bars can be vertical or horizontal. When they are vertical, these graphs are called column graphs.



1. Determine what type of graph is appropriate for each of the following examples:
  - (a) Arm span vs height in humans: \_\_\_\_\_
  - (b) Daily energy requirement for different species of deer: \_\_\_\_\_
  - (c) Number of fish of each size in a population: \_\_\_\_\_
  - (d) Mean volume of water used per person per day in different North American cities: \_\_\_\_\_
  - (e) Mean catalase reaction rate at different temperatures: \_\_\_\_\_
  - (f) Number of eggs per brood in different breeds of chickens: \_\_\_\_\_
  - (g) Mean monthly rainfall vs mean monthly temperature: \_\_\_\_\_
2. For the plots on the previous page:
  - (a) Use interpolation to determine the mean growth rate of pea seedlings at 17°C: \_\_\_\_\_
  - (b) Use interpolation to determine the number of eggs per brood in a 1.5 mm long *Daphnia*: \_\_\_\_\_
  - (c) Use interpolation to determine the rate of water loss in peas at 40% relative humidity: \_\_\_\_\_
3. Extrapolation, i.e. predicting a data value that lies outside the range of available data, is not recommended practice.
  - (a) Suggest why you should not extrapolate to find data values? \_\_\_\_\_  
\_\_\_\_\_
  - (b) Can you think of an example to illustrate your decision? \_\_\_\_\_  
\_\_\_\_\_

**Key Idea:** The equation for a straight line is  $y = mx + c$ . A line may have a positive, negative, or zero slope. The equation for a linear (straight) line on a **graph** is  $y = mx + c$  and can be used to calculate the gradient (slope) of a straight

line. It tells us about the relationship between  $x$  and  $y$  (how fast  $y$  is changing relative to  $x$ ). For a straight line, the rate of change of  $y$  relative to  $x$  is always constant. A line may have a positive, negative, or zero slope.

### Measuring gradients and intercepts

The equation for a straight line is written as:

**$y = mx + c$**

Where:

$y$  = the  $y$ -axis value

$m$  = the slope (or gradient)

$x$  = the  $x$ -axis value

$c$  = the  $y$  intercept (where the line cross the  $y$ -axis).

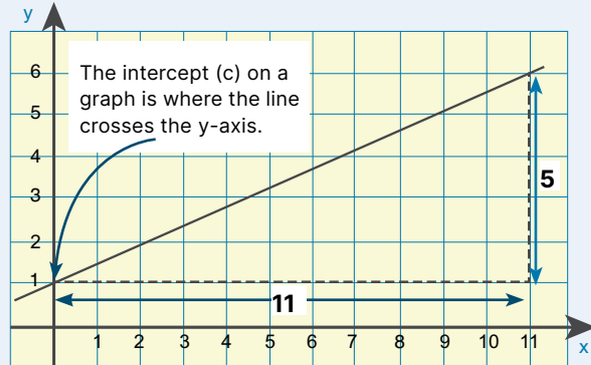
### Determining "m" and "c"

To find "c" just find where the line crosses the  $y$ -axis.

To find  $m$ :

1. Choose any two points on the line.
2. Draw a right-angled triangle between the two points on the line.
3. Use the scale on each axis to find the triangle's vertical length and horizontal length.
4. Calculate the gradient of the line using the equation:

$$\frac{\text{change in } y}{\text{change in } x}$$



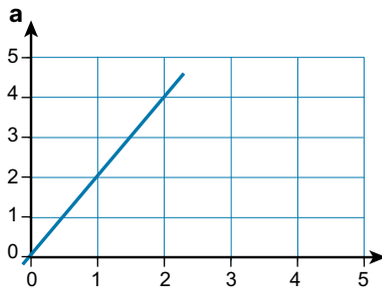
For the example above:

$c = 1$   
 $m = 0.45 (5 \div 11)$

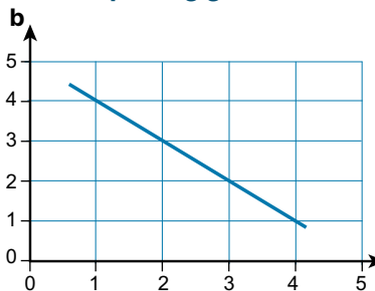
Once  $c$  and  $m$  have been determined you can choose any value for  $x$  and find the corresponding value for  $y$ .

For example, when  $x = 9$ , the equation would be:  
 $y = 9 \times 0.45 + 1$   
 $y = 5.05$

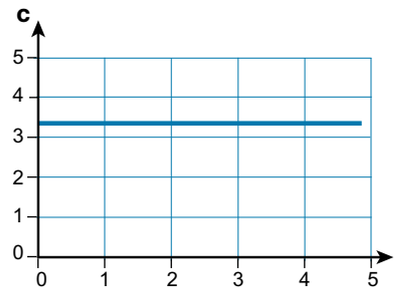
### Interpreting gradients



**Positive gradients:** the line slopes upward to the right ( $y$  is increasing as  $x$  increases).

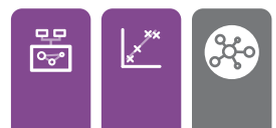
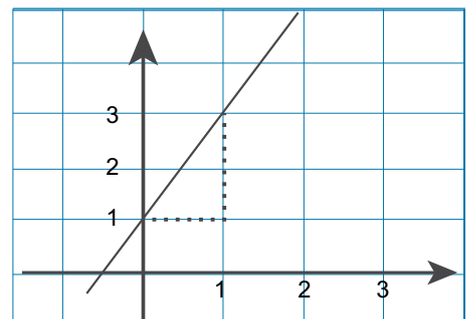


**Negative gradients:** the line slopes downward to the right ( $y$  is decreasing as  $x$  increases).



**Zero gradients:** the line is horizontal ( $y$  does not change as  $x$  increases).

1. State the gradient for graphs a, b, and c (above): (a) \_\_\_\_\_ (b) \_\_\_\_\_ (c) \_\_\_\_\_
2. For a straight line  $y = 3x + 2$ ,  
 (a) Identify the value of  $c$ : \_\_\_\_\_ (b) Determine  $y$  if  $x = 4$ : \_\_\_\_\_
3. For the graph (right):  
 (a) Identify the value of  $c$ : \_\_\_\_\_  
 (b) Calculate the value of  $m$ : \_\_\_\_\_  
 (c) Determine  $y$  if  $x = 2$ : \_\_\_\_\_  
 \_\_\_\_\_  
 (d) Describe the slope of the line: \_\_\_\_\_



# 14 Correlation or Causation

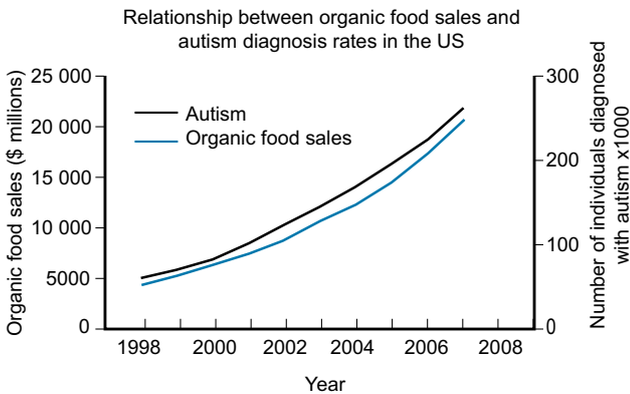
**Key Idea:** A correlation is a mutual relationship or association between two or more variables. A correlation between two variables does not imply that one causes change in the other. Researchers often want to know if two **variables** have any correlation (relationship) to each other. This can be achieved by plotting the data as a scatter graph and drawing a line of

best fit through the data, or by testing for correlation using a statistical test. The strength of a correlation is indicated by the correlation coefficient ( $r$ ), which varies between 1 and -1. A value of 1 indicates a perfect (1:1) relationship between the variables. A value of -1 indicates a 1:1 negative relationship and 0 indicates no relationship between the variables.

## Correlation does not imply causation

You may come across the phrase "correlation does not necessarily imply causation". This means that even when there is a strong correlation between variables (they vary together in a predictable way), you cannot assume that change in one variable caused change in the other.

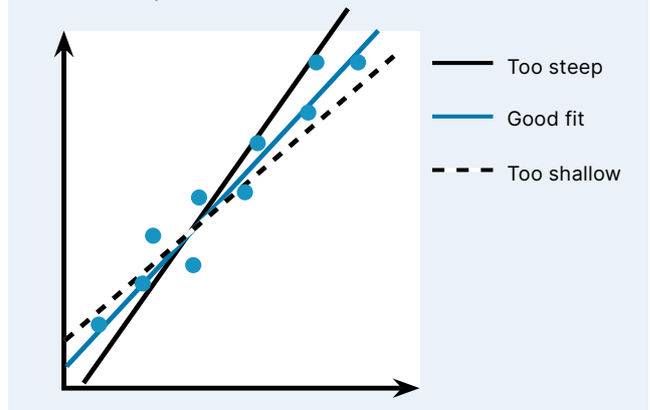
**Example:** When data from the organic food association and the office of special education programmes is plotted (below), there is a strong correlation between the increase in organic food and rates of diagnosed autism. However it is unlikely that eating organic food causes autism, so we can not assume a causative effect here.



## Drawing the line of best fit

Some simple guidelines need to be followed when drawing a line of best fit on your scatter plot.

- ▶ Your line should follow the **trend** of the data points.
- ▶ Roughly half of your data points should be above the line of best fit, and half below.
- ▶ The line of best fit does not necessarily pass through any particular point.
- ▶ The line of best fit should pivot around the point which represents the mean of the x and the mean of the y variables.



1. What does the phrase "correlation does not imply causation" mean? \_\_\_\_\_

---



---

2. A student measured the hand span and foot length measurements of 21 adults and plotted the data as a scatter graph (right).

(a) Draw a line of best fit through the data:

(b) Describe the results:

---



---



---

(c) Using your line of best fit as a guide, comment on the correlation between handspan and foot length:

---



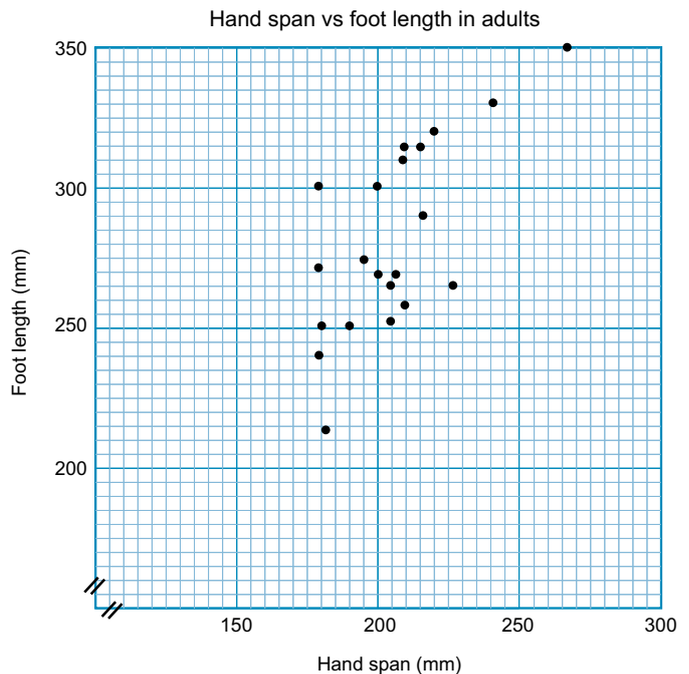
---



---



---



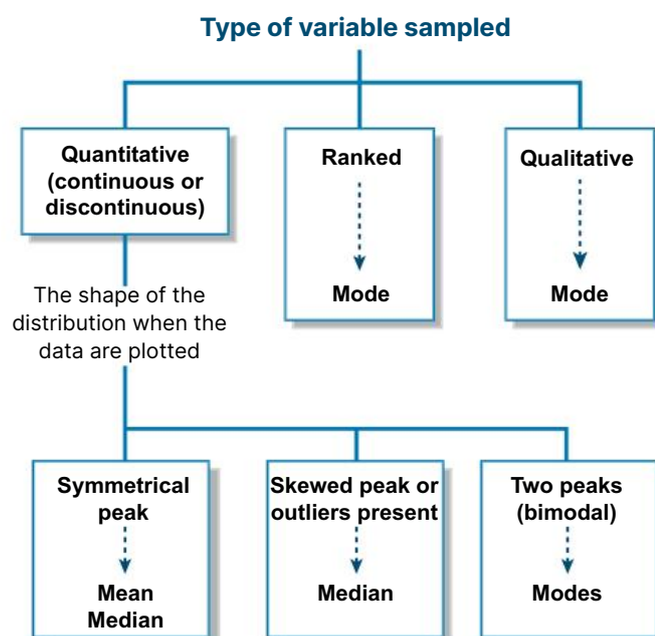
# 15 Mean, Median, and Mode

**Key Idea:** Mean, median, and mode are measures of the central tendency of data. The distribution of the data will determine which measurement of central tendency you use. Measures of a biological response are usually made from more than one sampling unit. In lab-based investigations, the sample size (the number of sampling units) may be as small as three or four (e.g. three test-tubes in each of

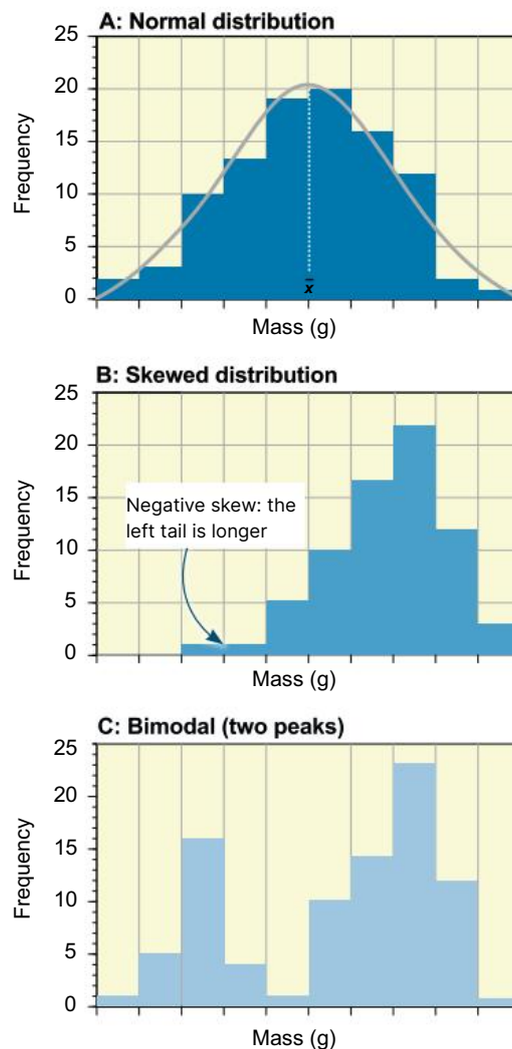
four treatments). In field studies, each individual may be a sampling unit, and the sample size can be very large (e.g. 100 individuals). It is useful to summarise data using descriptive statistics. Descriptive statistics, such as **mean**, **median**, and **mode**, can identify the central tendency of a data set. Each of these statistics is appropriate to certain types of data or distribution (as indicated by a frequency distribution).

## Variation in data

Whether they are obtained from **observation** or experiments, most biological data show variability. In a set of data values, it is useful to know the value about which most of the data are grouped, i.e. the centre value. This value can be the mean, median, or mode depending on the type of **variable** involved (see below). The main purpose of these statistics is to summarise important features of your data and to provide the basis for statistical analyses.



The shape of the distribution will determine which statistic (mean, median, or mode) best describes the central tendency of the sample data.



Statistic	Definition and use	Method of calculation
<b>Mean</b>	<ul style="list-style-type: none"> <li>The average of all data entries.</li> <li>Measure of central tendency for normally distributed data.</li> </ul>	<ul style="list-style-type: none"> <li>Add up all the data entries.</li> <li>Divide by the total number of data entries.</li> </ul>
<b>Median</b>	<ul style="list-style-type: none"> <li>The middle value when data entries are placed in rank order.</li> <li>A good measure of central tendency for skewed distributions.</li> </ul>	<ul style="list-style-type: none"> <li>Arrange the data in increasing rank order.</li> <li>Identify the middle value.</li> <li>For an even number of entries, find the mid point of the two middle values.</li> </ul>
<b>Mode</b>	<ul style="list-style-type: none"> <li>The most common data value.</li> <li>Suitable for bimodal distributions and <b>qualitative data</b>.</li> </ul>	<ul style="list-style-type: none"> <li>Identify the category with the highest number of data entries using a tally chart or a bar graph.</li> </ul>
<b>Range</b>	<ul style="list-style-type: none"> <li>The difference between the smallest and largest data values.</li> <li>Provides a crude indication of data spread.</li> </ul>	<ul style="list-style-type: none"> <li>Identify the smallest and largest values and find the difference between them.</li> </ul>

### When NOT to calculate a mean:

In some situations, calculation of a simple arithmetic mean is not appropriate.

#### Remember:

- DO NOT calculate a mean from values that are already means (averages) themselves.
- DO NOT calculate a mean of ratios (e.g. percentages) for several groups of different sizes. Go back to the raw values and recalculate.
- DO NOT calculate a mean when the measurement scale is not linear, e.g. pH units are not measured on a linear scale.





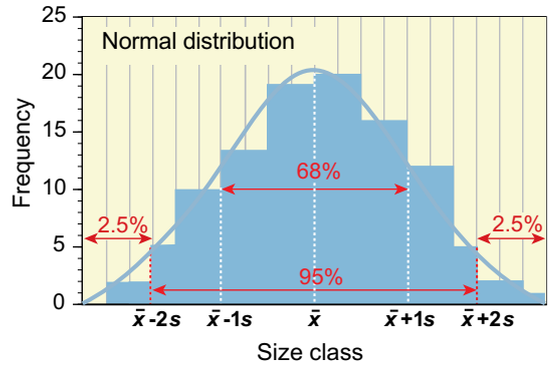
**Key Idea:** Standard deviation is used to quantify the variability around the central value and evaluate the reliability of estimates of the true mean.

While it is important to know the central tendency (e.g. mean) of a data set, it is also important to know how well this value

represents the data set as a whole. For a normal distribution, this can be evaluated using the standard deviation, which is a simple value that quantifies the spread in the data. If the standard deviation is small, more of the values will be clustered about the mean value.

### Standard deviation

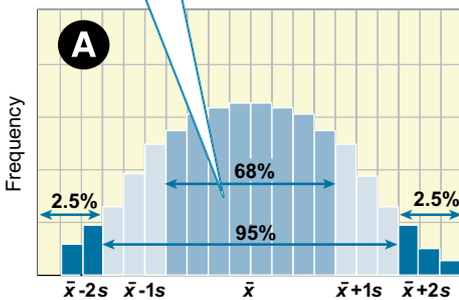
- ▶ Standard deviation is usually presented as  $\bar{x} \pm s$ . In normally distributed data, 68% of all data values will lie within one standard deviation (s) of the mean ( $\bar{x}$ ) and 95% of all values will lie within two standard deviations of the mean (right).
- ▶ Different sets of data can have the same mean and range, yet a different data distribution. In both the data sets below, 68% of the values lie within the range  $\bar{x} \pm 1s$  and 95% of the values lie within  $\bar{x} \pm 2s$ . However, in B, the data values are more tightly clustered around the mean.
- ▶ Standard deviation is easily calculated using a spreadsheet. Data should be entered as columns. In a free cell, type the formula for standard deviation and select the cells containing the data values, enclosing them in parentheses.



Histogram A has a larger standard deviation; the values are spread widely around the mean.

Both plots show a normal distribution with a symmetrical spread of values about the mean.

Histogram B has a smaller standard deviation; the values are clustered more tightly around the mean.

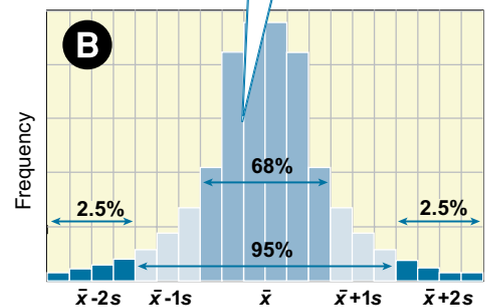


### Calculating s

$$s = \sqrt{\frac{\sum(x - \bar{x})^2}{n - 1}}$$

$\sum(x - \bar{x})^2$  = sum of squared deviations from the mean

n = sample size. n - 1 provides a unbiased s for small sample sizes (large samples can use n).



- Two sample data sets of rat body length have the same mean. The first data set has a much larger standard deviation than the second data set. What does this tell you about the spread of data around the mean in each case? Which data set is likely to provide the most reliable estimate of body length in the rat population being sampled and why?

---



---



---

- The data on the right shows the heights for 29 male swimmers.

(a) Calculate the mean for the data: \_\_\_\_\_

(b) Use manual calculation, a calculator, or a spreadsheet to calculate the standard deviation (s) for the data:

\_\_\_\_\_

(c) State the mean  $\pm 1s$ : \_\_\_\_\_

(d) What percentage of values are within 1s of the mean? \_\_\_\_\_

(e) What does this tell you about the spread of the data?

\_\_\_\_\_

\_\_\_\_\_



Raw data: Height (cm)

178	180	180	178	176
177	181	185	186	188
188	178	185	176	180
176	178	175	185	186
186	176	189	177	177
175	175	174	176	



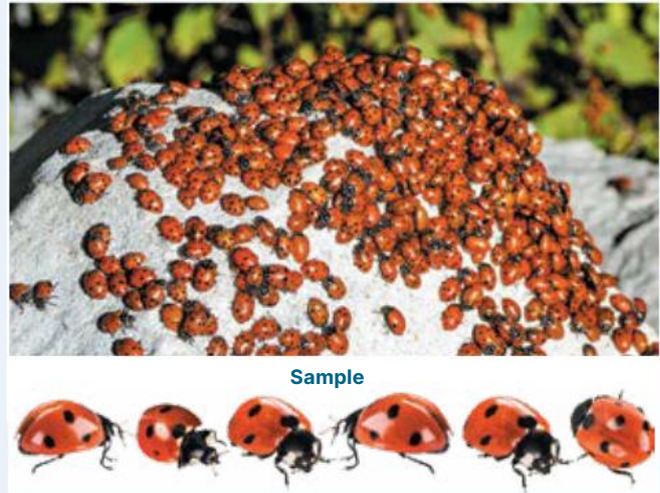
# 17 Reliability of the Mean

**Key Idea:** 95% confidence limits help to evaluate the reliability of the sample mean as an estimate of the population mean. You have already seen how to use the standard deviation ( $s$ ) to quantify the spread or dispersion in your data. Usually, you will also want to know how good your sample mean ( $\bar{x}$ )

is as an **estimate** of the true population mean ( $\mu$ ). You can do this by calculating the 95% confidence interval (95% CI). The mean  $\pm$  the 95% CI gives the 95% confidence limits. On average, 95 times out of 100, the true population mean will lie within the confidence limits.

## Reliability of the sample mean

- ▶ When we take measurements from samples of a larger population, we are using those samples as indicators of the **trends** in the whole population. Therefore, when we calculate a sample mean (the statistic), it is useful to know how close that value is to the true population mean ( $\mu$ ) for that attribute (the parameter).
- ▶ If you can determine the **reliability** of the sample mean, it will enable you to make inferences about the aspect of the population in which you are interested. Statistics based on samples and used to estimate population parameters are called inferential statistics.
- ▶ **Example:** If we calculated the mean number of spots from a sample of six ladybird beetles, how reliable is this statistic as an indicator of the mean number of spots in the whole population? We can find out by calculating the 95% confidence interval.



## Step 1: Calculate standard error (SE)

The standard error (SE) is simple to calculate and is usually a small value. Standard error is given by the standard deviation divided by the square root of  $n$ , where  $n$  is the sample size.

$$SE = \frac{s}{\sqrt{n}}$$

## Step 2: Use SE to calculate the 95% confidence interval

SE is required to calculate the 95% confidence interval (CI) of the mean. This is simple: just multiply SE by the value of  $t$  at  $P = 0.05$  (from a  $t$  table) for the appropriate degrees of freedom ( $df$ ) for your sample ( $n - 1$ ). Part of the  $t$  table is provided for you below.

$$95\% \text{ CI} = SE \times t_{P(n-1)}$$

## Critical values of Student's $t$ at $P = 0.05$

Use this table to calculate 95% confidence interval

df	$P$
	0.05
1	12.71
2	4.303
3	3.182
4	2.776
5	2.571
6	2.447
7	2.365
8	2.306
9	2.262
10	2.228
20	2.086
30	2.042
40	2.021
60	2.000
120	1.980
>120	1.960

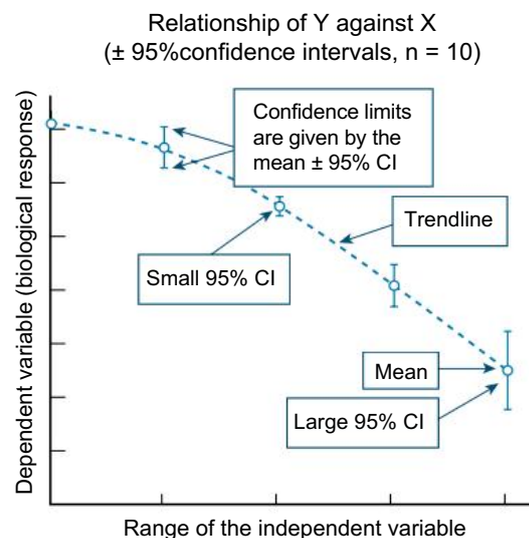
Value of  $t$  at  $n - 1 = 5$

As the sample becomes very large, the value of  $t$  becomes smaller. For very large samples,  $t$  is fixed at 1.96, so the 95% CI is slightly less than twice the SE.

Maximum value of  $t$  at this level of  $P$

## Step 3: Plotting your confidence intervals

Once you have calculated the 95% CI for the means in your data, you can plot them as error bars on your graph. Note that the 95% confidence limits are given by the value of the mean  $\pm$  95%CI. A 95% confidence limit (i.e.  $P = 0.05$ ) tells you that, on average, 95 times out of 100, the limits will contain the true population mean. Note that each of the plotted points represents a mean of 10 values.



All these statistics, including a plot of the data with Y error bars, can be calculated using a programme such as Microsoft Excel®.





**Clover root weevil**

- ▶ The clover root weevil (*Sitona lepidus*) is a pest of white clover pastures. The adults feed on clover leaves, while the larvae feed on clover nodules and roots, causing root loss and a reduction in nitrogen fixation.
- ▶ Research has indicated that different pastures have different susceptibility to infestation by clover root weevil (left). Armed with this knowledge, two students reasoned that the most susceptible grass type would have the greatest weevil population. The students chose five pasture types, and recorded the number of weevil larvae in each pasture type at six sample sites (sample area 1 m<sup>2</sup>). Their results are presented in the table below.

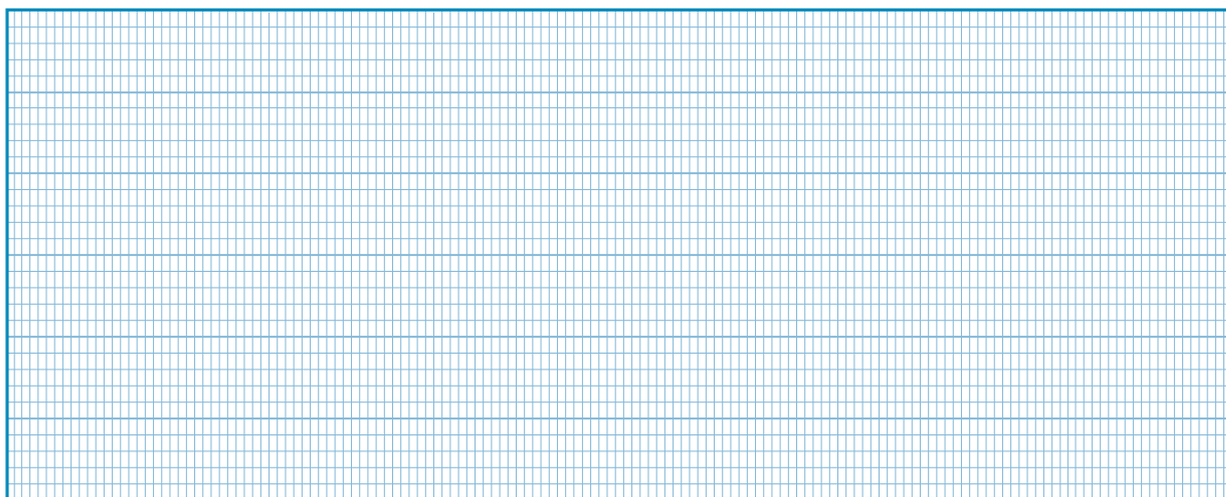
		Environment				
Sample		Perennial ryegrass	Fescue	White clover	Red clover	Chicory
<b>Number of weevils</b>	1	42	42	48	42	45
	2	45	46	54	46	44
	3	41	38	48	45	45
	4	42	41	52	42	38
	5	49	45	49	44	40
	6	43	44	52	44	47

1. Complete the table below by calculating the mean, standard deviation, standard error, and 95% confidence interval (95% CI) for each of the grass environments.

	Perennial ryegrass	Fescue	White clover	Red clover	Chicory
Mean					
Standard deviation					
Standard error					
95% CI					

2. Select the appropriate graph format and plot the means for each of the grass environments below. Include bars to show the 95% confidence intervals.

Need help?  
See Activity 12



3. Study your plot and decide if there are any significant differences between the abundance of clover root weevils in the five environments. Use your analysis to write a conclusion for the investigation below:

---



---



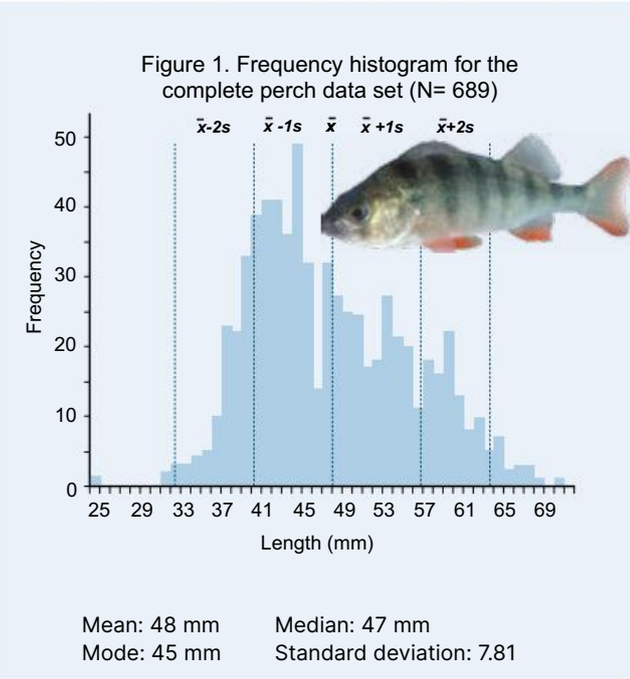
---

# 18 Detecting Bias in Samples

**Key Idea:** Sampling method can affect the results of a study, especially if it has an unknown bias.

**Bias** refers to the selection for or against one particular group in such a way that it can influence an investigation's results. Bias can occur when sampling is not random, and certain members of a population are under- or over-represented

relative to others in the population. Small sample sizes can also bias the results, which may not accurately reflect the population as a whole. Bias can be reduced by random sampling (sampling in which all members of the population have the same chance of being selected). Using appropriate collection methods and apparatus can also reduce bias.



- ▶ In this exercise, perch were collected and their body lengths (mm) were measured. Data are presented as a frequency histogram and with descriptive statistics (**mean, median, mode** and standard deviation).
- ▶ Figure 1 shows the results for the complete data set. The sample set was large (N= 689) and the perch were randomly sampled. The data are close to having a normal distribution.
- ▶ Figures 2 and 3 show results for two smaller sample sets drawn from the same population. The data collected in Figure 2 were obtained by random sampling but the sample was relatively small (N = 30). The person gathering the data displayed in Figure 3 used a net with a large mesh size to collect the perch.

1. (a) Compare the results for the two small data sets (Figures 2 and 3). How close are the mean and median to each other in each sample set?

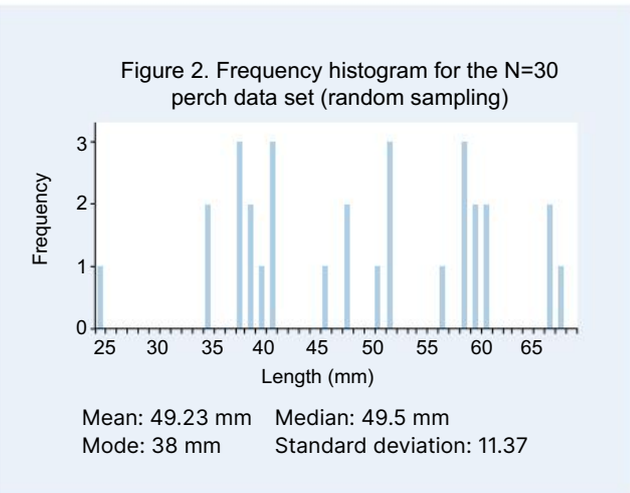
---

---

---

---

---



(b) Compare the standard deviation for each sample set:

---

---

---

---

(c) Describe how each of the smaller sample sets compares to the large sample set (Figure 1):

---

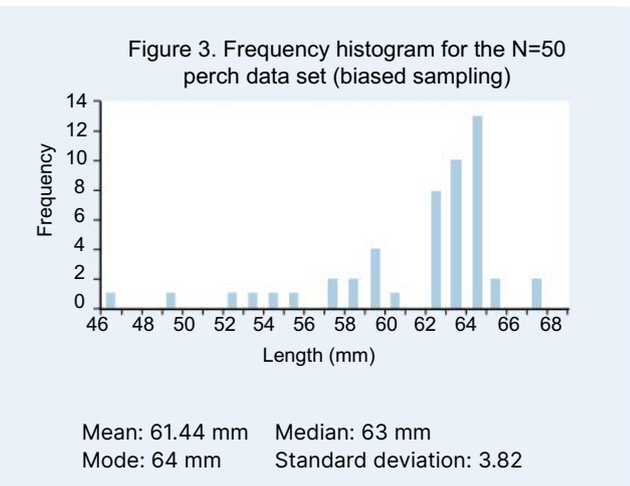
---

---

---

---

---



(d) Why do you think the two smaller sample sets look so different to each other?

---

---

---

---

---

---

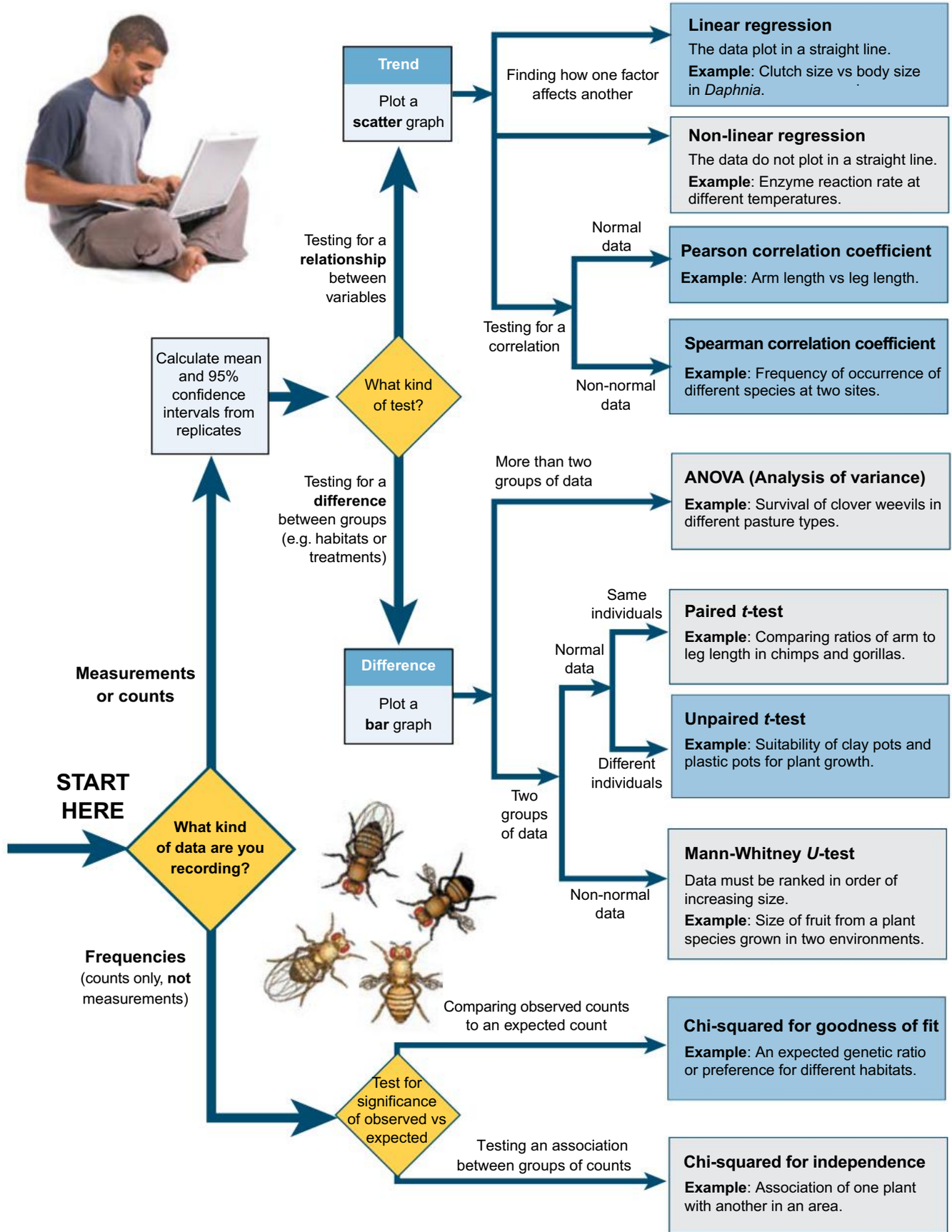
---

---



**Key Idea:** How your data is analysed depends on the type of data you have collected. Plotting your initial data can help you to decide what statistical analysis to carry out. Data analysis provides information on the biological

significance of your investigation. Never under-estimate the value of plotting your data, even at a very early stage. This will help you decide on the best type of data analysis. Sometimes, statistical analysis may not be required.



You may need to use these during your study

You will not be required to use these during your study

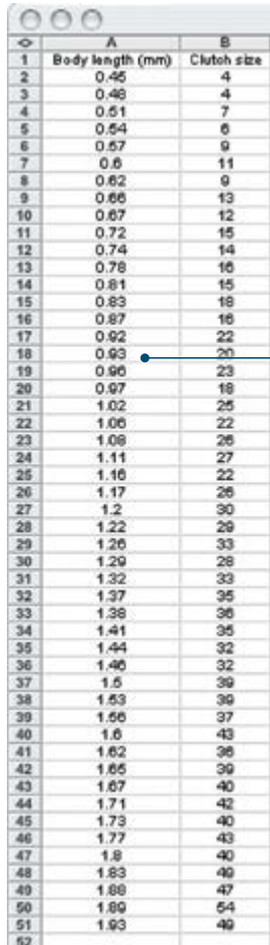


**Key Idea:** Correlation coefficients provide information on the strength of the relation between variables. The data can be used to produce a regression equation for the line of best fit. Pearson's correlation coefficient ( $r$ ) relates the value of one **variable** to another for a simple linear relationship. For a perfect correlation in which the change in one variable is perfectly matched or caused by a corresponding change in another variable,  $r = 1$  (positive correlation) or  $-1$  (negative correlation). If there is no correlation between variables,

then  $r = 0$ . The simplest way to calculate a correlation coefficient is to use a spreadsheet such as Microsoft Excel (below). Correlation can also be expressed as the  $R^2$  value, the coefficient of determination. This is the square of the correlation coefficient. It quantifies the **proportion** of variance in the **dependent variable** that is explained by the independent variable. Unlike  $r$ ,  $R^2$  has no direction (+/-) and determines how a **model** or equation fits the data.

## 1 Enter the data

Clutch size (number of eggs per female) was estimated for 50 females, and body length was measured to the nearest 0.01 mm for the same individuals to give 50 paired values. These data values were entered directly into Microsoft Excel®.



	A	B
1	Body length (mm)	Clutch size
2	0.45	4
3	0.48	4
4	0.51	7
5	0.54	6
6	0.57	9
7	0.6	11
8	0.62	9
9	0.65	13
10	0.67	12
11	0.72	15
12	0.74	14
13	0.78	16
14	0.81	15
15	0.83	18
16	0.87	16
17	0.92	22
18	0.93	20
19	0.96	23
20	0.97	19
21	1.02	25
22	1.05	22
23	1.08	26
24	1.11	27
25	1.16	22
26	1.17	26
27	1.2	30
28	1.22	29
29	1.26	33
30	1.29	28
31	1.32	33
32	1.37	35
33	1.38	36
34	1.41	35
35	1.44	32
36	1.46	32
37	1.5	39
38	1.53	39
39	1.55	37
40	1.6	43
41	1.62	38
42	1.65	39
43	1.67	40
44	1.71	42
45	1.73	40
46	1.77	43
47	1.8	40
48	1.83	49
49	1.88	47
50	1.89	54
51	1.93	49
52		



Brood pouch

Enter your data in columns with headings for the variables.

*Daphnia*



## Calculating a correlation coefficient 'r'

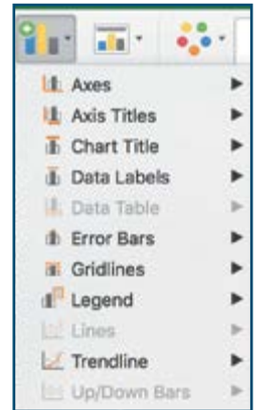
- Remember that correlation is not causation. Just because two variables (e.g. temperature and population number) are correlated it does not mean one causes the other.
- The simplest way to calculate a correlation coefficient is to use a spreadsheet such as Microsoft Excel®:
  - Enter the data into two columns.
  - In another cell type the formula `=CORREL(Range Column 1, Range Column 2)`.

## Clutch size vs body size in *Daphnia*

*Daphnia* is a small, freshwater crustacean common in water bodies throughout the world. In *Daphnia*, body size largely determines how many eggs and young can be carried (the clutch size). This is because the eggs are carried in a brood pouch, which physically limits the size of the clutch. Larger animals can also process more food. The strength of the relationship between body size and clutch size can be described using a correlation coefficient.

## 2 Graphing and fitting the regression

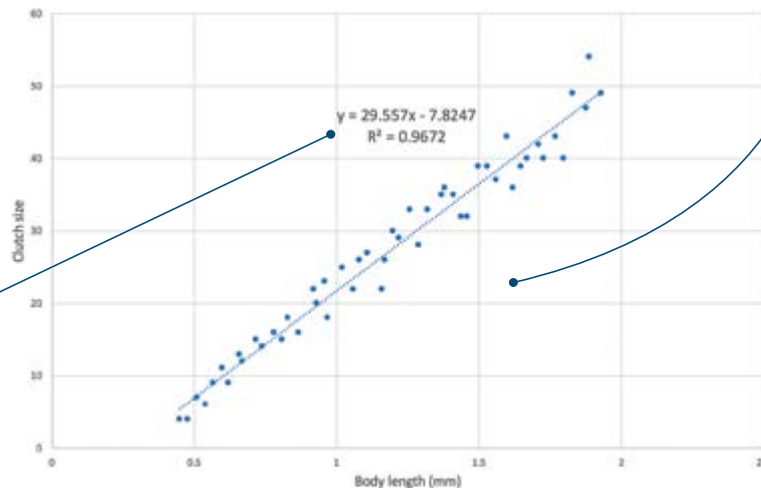
- Select the data columns: "Body length" and "Clutch size".
- Choose Insert > Chart > XY (Scatter). Choose the option with no line. A graph will appear on screen.
- Under the menu tab "Chart Design", choose from the drop down menu under "Add Chart Element" (right) to add titles for your axes and add a trendline (this is your regression line)
- Right click on the regression line itself to format the trendline. You can add the  $R^2$  value and regression equation if you wish.
- The  $R^2$  value describes how well the line fits the data. A value of 1 is a perfect fit. The equation describes the slope and intercept of the line. It allows you to predict values of the dependent variable.



In a cell beneath the columns type the formula `=CORREL(Range Column 1, Range Column 2)` to get the correlation coefficient.

Regression equation and  $R^2$ . The  $R^2 = 0.9672$ . This regression accounts for 96% of the scatter in the data.

The correlation coefficient ( $r = \sqrt{R^2}$ ) = 0.98, meaning a very strong relationship between the variables.



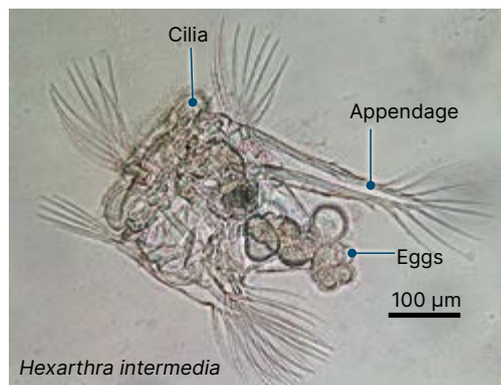
### A correlation example

Pearson's coefficient correlation can be calculated using the formula:

$$r = \frac{\sum xy - n\bar{x}\bar{y}}{ns_x s_y}$$

To find  $s_x$  and  $s_y$  use the population standard deviation:  $s = \sqrt{\frac{\sum(x - \bar{x})^2}{n}}$

*Hexarthra intermedia* (right) is a species of rotifer. Rotifers are small, ciliated animals found in freshwater ponds. Most feed on small algae. A study was carried out in order to understand how changes in their abundance might be related to seasonal changes in environmental factors. The data below records abundance per litre of pond water against pond temperature at the time that sample was taken. Note that the *Hexarthra* counts are not in whole numbers because number per litre was calculated from a larger, filtered sample volume.  $n = 25$ .



Data kindly supplied by Dr Ian Duggan, University of Waikato

Hexarthra no. (x) / per L	Temperature (y) / °C	$(x - \bar{x})^2$	$(y - \bar{y})^2$	xy
36.21	19.75			
33.76	17.53			
10.83	15.05			
1.88	14.40			
0.33	11.73			
2.40	11.05			
0.35	9.23			
0.08	8.75			
0.00	12.35			
0.04	13.13			
0.00	14.15			
0.21	14.63			
0.29	15.98			
5.72	19.63			
4.39	18.00			
7.42	19.80			
72.87	23.33			
443.38	23.30			
34.38	22.30			
147.58	25.88			
947.64	24.58			
573.47	22.90			
444.63	20.95			
338.25	21.10			
34.33	18.90			
$\bar{x} =$	$\bar{y} =$	$\sum(x - \bar{x})^2 =$	$\sum(y - \bar{y})^2 =$	$\sum xy =$
Standard deviation x =		Standard deviation y =		r =

- Complete the table above to calculate  $r$ :
- What does  $r$  tell you about the relationship between *Hexarthra* numbers and temperature? \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_

# 21 Spearman Rank Correlation

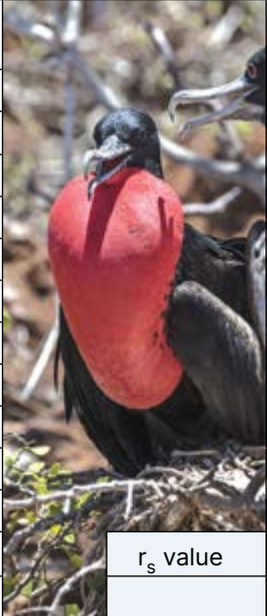
**Key Idea:** The Spearman rank correlation is a test used to determine if there is a statistical dependence (correlation) between two variables.

The Spearman rank correlation is appropriate for data that have a non-normal distribution (or where the distribution is not known) and assesses the degree of association between the X and Y **variables** (if they are correlated). For the test to

work, the values used must be monotonic i.e. the values must increase or decrease together or one increases while the other decreases. A value of 1 indicates a perfect correlation; a value of 0 indicates no correlation between the variables. The example below examines the relationship between the frequency of the drumming sound made by male frigatebirds (Y) and the volume of their throat pouch (X).

**Spearman's rank data for frigate bird pouch volume and drumming frequency**

Bird	Volume of pouch / cm <sup>3</sup>	Rank (R1)	Frequency of drumming sound / Hz	Rank (R2)	Difference (D) (R1-R2)	D <sup>2</sup>
1	2550		461			
2	<b>2440</b>	<b>1</b>	<b>473</b>	<b>6</b>	<b>-5</b>	<b>25</b>
3	2740		532			
4	2730		465			
5	3010		485			
6	3370		488			
7	3080		527			
8	4910		478			
9	3740		485			
10	5090		434			
11	5090		468			
12	5380		449			
Based on Madsen et al 2004					$\Sigma D^2$	



$r_s$  value

## Analysing the data

**Step one:** Rank the data for each variable. For each variable, the numbers are ranked in descending order, e.g. for the variable, volume, the highest value 5380 cm<sup>3</sup> is given the rank of 12 while its corresponding frequency value is given the rank of 2. Fill in the rank columns in the **table** above in the same way. If two numbers have the same rank value, then use the mean rank of the two values (e.g. 1+2 = 3. 3/2= 1.5).

**Step two:** Calculate the difference (D) between each pair of ranks (R1-R2) and enter the value in the table (as a check, the sum of all differences should be 0).

**Step three:** Square the differences and enter them into the table above (this removes any negative values).

**Step four:** Sum all the D<sup>2</sup> values and enter the total into the table.

**Step five:** Use the formula below to calculate the Spearman Rank Correlation Coefficient ( $r_s$ ). Enter the  $r_s$  value in the box above.

$$r_s = 1 - \left( \frac{6\Sigma D^2}{n(n^2-1)} \right)$$

### Spearman rank correlation coefficient

**Step six:** Compare the  $r_s$  value to the table of critical values (right) for the appropriate number of pairs. If the  $r_s$  value (ignoring sign) is greater than or equal to the critical value then there is a significant correlation. If

$r_s$  is positive then there is a positive correlation. If  $r_s$  is negative then there is a negative correlation.

Number of pairs of measurements	Critical value
5	1.00
6	0.89
7	0.79
8	0.74
9	0.68
10	0.65
12	0.59
14	0.54
16	0.51
18	0.48
20	0.45

- State the null hypothesis for the data set: \_\_\_\_\_
- Identify the critical value for the frigate bird data: \_\_\_\_\_
  - State if the correlation is positive or negative: \_\_\_\_\_
  - State whether the correlation is significant: \_\_\_\_\_
- In your class, gather data on heart rate (beats per minute measured by carotid or radial pulse) and breathing rate (breaths per minute). Use the Spearman rank coefficient to determine if there is a relationship between these variables. Complete your analysis and staple it to this page.



**Key Idea:** Differences between two populations (or sets of data) can be tested for significance using the Student's t-test. The Student's t-test is commonly used to compare two sample means, e.g. means for a treatment and a **control** in an experiment, or the means of some measured characteristic between two animal or two plant populations. It is a simple

test and useful for distinguishing real but marginal differences between samples. Usefully, the test remains robust even when sample sizes are small. A simple example outlining the steps in the Student's t-test is shown below. It compares data for a treatment and a control from a hypothetical experiment (the units are not relevant in this case, only the values).

### Steps in performing a Student's t-test

#### 1 Calculate summary statistics for the two data sets

Control (A)	Treatment (B)
6.6	6.3
5.5	7.2
6.8	6.5
5.8	7.1
6.1	7.5
5.9	7.3

$$n_A = 6, \bar{x}_A = 6.12, s_A = 0.496$$

$$n_B = 6, \bar{x}_B = 6.98, s_B = 0.475$$

$n_A$  and  $n_B$  are the number of values in the first and second data sets respectively (these do not need to be the same).

$\bar{x}$  is the mean.

$s$  is the standard deviation (a measure of scatter in the data).

#### 2 Set up and state your null hypothesis ( $H_0$ )

$H_0$ : there is no treatment effect. The differences in the data sets are the result of chance and they are not really different. The alternative hypothesis is that there is a treatment effect and the two sets of data are truly different.

#### 3 Decide if your test is one or two tailed

A one-tailed test looks for a difference only in one particular direction. A two-tailed test looks for any difference (+ or -). This tells you what section of the t table to consult. Most biological tests are two-tailed. Very few are one-tailed.

#### 4 Calculate the t statistic

For our sample data above the calculated value of t is -3.09. The degrees of freedom (df) are  $n_1 + n_2 - 2 = 10$ .

Calculation of the t value uses the variance which is simply the square of the standard deviation ( $s^2$ ). You may compute t using a spreadsheet but manual computation is not difficult (see activity 17). It does not matter if the calculated t value is a positive or negative (the sign is irrelevant).

The absolute value of the t statistic (3.09) well exceeds the critical value for  $P = 0.05$  at 10 degrees of freedom.

We can reject  $H_0$  and conclude that the means are different at the 5% level of significance.

If the calculated absolute value of t had been less than 2.23, we could not have rejected  $H_0$ .

1. (a) In an experiment, data values were obtained from four plants in experimental conditions and three plants in control conditions. The mean values for each data set (control and experimental conditions) were calculated. The t value was calculated to be 2.16. The null hypothesis was: "The plants in the control and experimental conditions are not different". State whether the calculated t value supports the null hypothesis or its alternative (consult t table below):

---



---

- (b) The experiment was repeated, but this time using 6 control and 6 "experimental" plants. The new t value was 2.54. State whether the calculated t value supports the null hypothesis or its alternative now:

---



---

2. Explain what you understand by statistical significance:

---



---



---



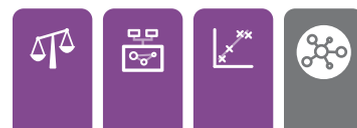
---



---

Table of critical values of t at different levels of P.

Degrees of freedom	Level of Probability		
	0.05	0.01	0.001
1	12.71	63.66	636.6
2	4.303	9.925	31.60
3	3.182	5.841	12.92
4	2.776	4.604	8.610
5	2.571	4.032	6.869
6	2.447	3.707	5.959
7	2.365	3.499	5.408
8	2.306	3.355	5.041
9	2.262	3.250	4.781
10	2.228	3.169	4.587
15	2.131	2.947	4.073
16	2.120	2.921	4.015
17	2.110	2.898	3.965
18	2.101	2.878	3.922
19	2.093	2.861	3.883
20	2.086	2.845	3.850
25	2.060	2.787	3.725
30	2.042	2.750	3.646
40	2.021	2.704	3.551
50	2.009	2.678	3.496
60	2.000	2.660	3.460
100	1.984	2.626	3.390



**Key Idea:** The chi-squared test ( $\chi^2$ ) for goodness of fit can be used for testing the significance of the differences between observed and expected outcomes.

The chi-squared test ( $\chi^2$ ), like the Student's *t* test, is a test for difference between data sets, but it is used when you are

### Using $\chi^2$ in Mendelian genetics

The predicted Mendelian ratios for the offspring of a cross between two *Drosophila* flies were 1:1:1:1 for each of the four following phenotypes: grey body-long wing, grey body-vestigial wing, ebony body-long wing, ebony body-vestigial wing. The observed results of the cross were not as predicted. The following numbers for each phenotype were observed in the offspring:

Observed results (O) of the example <i>Drosophila</i> cross			
			
Grey body, vestigial wing 88	Grey body, long wing 98	Ebony body, long wing 102	Ebony body, vestigial wing 112

Using  $\chi^2$ , the probability of this result being consistent with a 1:1:1:1 ratio could be tested. Worked example as follows:

#### STEP 1: Calculate the expected value (E)

In this case, this is the sum of the observed values divided by the number of categories ( $400 \div 4 = 100$ ).

#### STEP 2: Calculate the expected value (E)

The difference between the observed and expected values is calculated as a measure of the deviation from a predicted result. Since some deviations are negative, they are all squared to give positive values. This step is usually performed as part of a tabulation (next column, blue column).

#### STEP 3: Calculate the value of $\chi^2$

The calculated  $\chi^2$  value is given at the bottom right of the last column of the **table**. It is calculated using the formula below.

$$\chi^2 = \sum \frac{(O - E)^2}{E}$$

The completed table is shown at the top of the next column.

working with counts rather than measurements. The following worked example uses  $\chi^2$  to test the results of a genetic cross to see if departures from the predicted Mendelian **ratio** are significant. Raw counts should be used and large sample size is required for the test to be valid.

Category	O	E	O - E	(O - E) <sup>2</sup>	$\frac{(O - E)^2}{E}$
Grey, long wing	98	100	-2	4	0.04
Grey, vestigial wing	88	100	-12	144	1.44
Ebony, long wing	102	100	2	4	0.04
Ebony, vestigial wing	112	100	12	144	1.44

Total = 400

$$\chi^2 = \sum = 2.96$$

#### STEP 4: Calculate the degrees of freedom

The probability that any  $\chi^2$  value would be exceeded by chance depends on the degrees of freedom. This is one less than the total number of categories (this is the number that could vary independently without affecting the last value). Here:  $4 - 1 = 3$ .

#### STEP 5: Using the $\chi^2$ table

On the  $\chi^2$  table (produced in part below) with 3 degrees of freedom, the calculated value for  $\chi^2$  of 2.96 corresponds to a probability of between 0.2 and 0.5. This means that by chance alone, a  $\chi^2$  value of 2.96 could be expected between 20% and 50% of the time.

#### STEP 6: Interpreting the result

The probability of between 0.2 and 0.5 is higher than the 0.05 value that is generally regarded as significant. The null hypothesis cannot be rejected and we have no reason to believe that the observed ratios obtained differ significantly from the expected at  $P = 0.05$ .

**IMPORTANT NOTE:** When analysing Mendelian crosses, predicted ratios assume independent assortment of alleles, so significant departures from predicted ratios indicate the alleles are linked (on the same chromosome). Many Mendelian crosses involve ratios other than 1:1. For these, calculation of expected values is not simply a division of the total by the number of categories. Instead, the total must be apportioned according to the ratio. For example, for a total of 400 as above and a predicted 9:3:3:1 ratio, the total count must be divided by 16 and the expected values will be 225 : 75 : 75 : 25 in each category.

Table of critical values of  $\chi^2$  at different levels of probability

By convention, the critical probability for rejecting the null hypothesis ( $H_0$ ) is 5%. If the test statistic is less than the tabulated critical value for  $P = 0.05$ , we cannot reject  $H_0$  and the result is not significant. If the test statistic is greater than the tabulated critical value for  $P = 0.05$ , we reject  $H_0$  in favour of the alternative hypothesis.

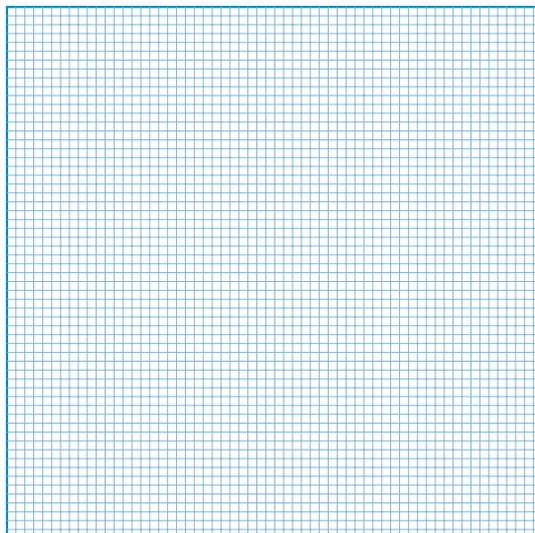
Degrees of freedom	Level of probability (P)									
	0.98	0.95	0.80	0.50	0.20	0.10	0.05	0.02	0.01	0.001
1	0.001	0.004	0.064	0.455	1.64	2.71	3.84	5.41	6.64	10.83
2	0.040	0.103	0.466	1.386	3.22	4.61	5.99	7.82	9.21	13.82
3	0.185	0.352	1.005	2.366	4.64	6.25	7.82	9.84	11.35	16.27
4	0.429	0.711	1.649	3.357	5.99	7.78	9.49	11.67	13.28	18.47
5	0.752	1.145	2.343	4.351	7.29	9.24	11.07	13.39	15.09	20.52

← Do not reject  $H_0$ 
Reject  $H_0$  →

# 24 Did You Get It?

- A balance has a calibration error of +0.04 g. A student weighs out 11.71 g of sodium hydroxide. Calculate the percentage error (show your working):
- 
- The table (below left) shows the rate of sweat production in an athlete on a stationary cycle.
    - Complete the table below to determine the rate of sweat loss in  $\text{cm}^3\text{min}^{-1}$ :
    - Choose an appropriate graph type and plot both cumulative sweat loss and rate of sweat loss on the grid below.

Time (minutes)	Cumulative sweat loss ( $\text{cm}^3$ )	Rate of sweat loss ( $\text{cm}^3\text{min}^{-1}$ )
0	0	
10	50	
20	130	
30	220	
60	560	



- Describe how the rate of sweat loss changes over time:

---



---



---

- Metabolic measurements were taken from seven Antarctic fish (*Pagothenia borchgrevinski*) affected by a gill disease, which increases the thickness of the gas exchange surfaces and affects oxygen uptake. The results of oxygen consumption of active fish with varying amounts of affected gill are tabulated right.

- Plot the data on the grid below right to show the relationship between oxygen consumption and the amount of gill affected by disease. Draw a line of best fit through the data.

Fish number	Percentage of gill affected	$\text{O}_2$ consumption ( $\text{cm}^3\text{g}^{-1}\text{h}^{-1}$ )
1	0	0.29
2	95	0.11
3	60	0.14
4	30	0.22
5	90	0.08
6	65	0.18
7	45	0.20

- Using a spreadsheet programme such as Microsoft Excel:

- Enter the data in columns and plot an XY scatter.
- Label your axes appropriately.
- Plot a regression line for the data.
- Display the  $R^2$  value and the regression equation.
- If you wish, print the graph and staple it here.

- How does the gill disease affect oxygen uptake? Use the results of your regression analysis to support your answer:

---



---



---



---



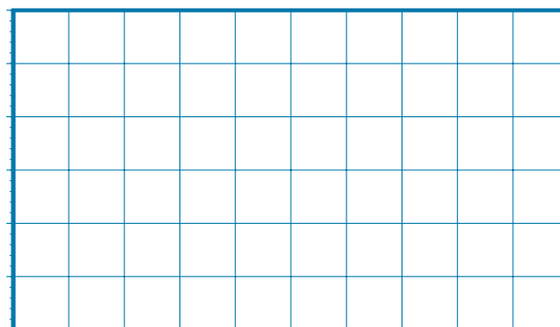
---



---



---





# Prokaryotic and Eukaryotic Cells

## Key Terms

- cell wall
- centrioles
- chloroplast
- cytoplasm
- electron micrograph
- electron microscope
- endoplasmic reticulum (ER)
- eukaryotic cell
- flagella
- Golgi
- light (=optical) microscope
- lysosome
- magnification
- mitochondrion
- nucleolus
- nucleus
- organelles
- plasma membrane
- prokaryotic cell
- resolution
- ribosome
- rough ER (rER)
- smooth ER (sER)
- stain
- vacuole

## Key Concepts

- ▶ The cell is the unit of life.
- ▶ Prokaryote and eukaryote cells are different.
- ▶ Organelles play important roles in cell functioning.
- ▶ Different types of microscopes can be used to study cells.

## Cells as the unit of life

### Activity Number

<input type="checkbox"/>	1	Explain how cells are the basic unit of life on Earth and outline the basic principles of the cell theory.	25
<input type="checkbox"/>	2	Name different types of cell and describe what cells are made of.	26-27
<input type="checkbox"/>	3	Describe the requirements for survival of all cells, including sources of energy, gases, and nutrients, and removal of wastes.	28
<input type="checkbox"/>	4	Describe the distinguishing features of prokaryotic cells, including small size, lack of a nucleus and membrane-bound organelles, and single circular chromosome. Explain that prokaryotes usually exist as single cells but may be colonial, with some specialisation of function.	29
<input type="checkbox"/>	5	Compare the features of prokaryotic and eukaryotic cells.	26, 29-33

## Eukaryotic cells have specialised organelles

<input type="checkbox"/>	6	Name the specialised organelles found in eukaryotic cells, including nucleus, mitochondria, rough and smooth endoplasmic reticulum, Golgi apparatus, lysosomes, vacuoles, and chloroplasts. Describe the role of these organelles in the functioning of the cell and the organism.	30-34
<input type="checkbox"/>	7	Compare and contrast the organelles in plant cells and animal cells.	30-34

## Studying cells

<input type="checkbox"/>	8	<b>SI:</b> Identify organelles visible in electron micrographs.	31, 33, 40-41
<input type="checkbox"/>	9	<b>SI:</b> Describe the structure and basic principles of light (optical) microscopes. Contrast light and electron microscopy in terms of magnification and resolution	35, 40
<input type="checkbox"/>	10	<b>SI:</b> Prepare wet mounts and use a light microscope to observe cells and identify structures and organelles in microorganism, plant, and animal cells. Calculate magnification and field of view.	36-37
<input type="checkbox"/>	11	<b>SI:</b> Produce annotated biological drawings of cells observed with a light microscope.	38-39

# 25 The Cell is the Unit of Life

**Key Idea:** All living organisms are composed of cells. Cells are broadly classified as prokaryotic or eukaryotic. The cell theory is a fundamental idea of biology. This idea,

that all living things are composed of cells, developed over many years and is strongly linked to the invention and refinement of the microscope in the 1600s.

## The cell theory

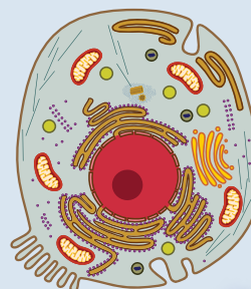
The idea that cells are fundamental units of life is part of the cell theory. The basic principles of the theory are:

- ▶ All living things are composed of cells and cell products.
- ▶ New cells are formed only by the division of pre-existing cells.
- ▶ The cell contains inherited information (genes) that are used as instructions for growth, functioning, and development.
- ▶ The cell is the functioning unit of life; all chemical reactions of life take place within cells.

## All cells show the functions of life

Cells use food (e.g. glucose) to maintain a stable internal environment, grow, reproduce, and produce wastes. The sum total of all the chemical reactions that sustain life is called metabolism.

- **Movement**
- **Respiration**
- **Sensitivity**
- **Growth**
- **Reproduction**
- **Excretion**
- **Nutrition**

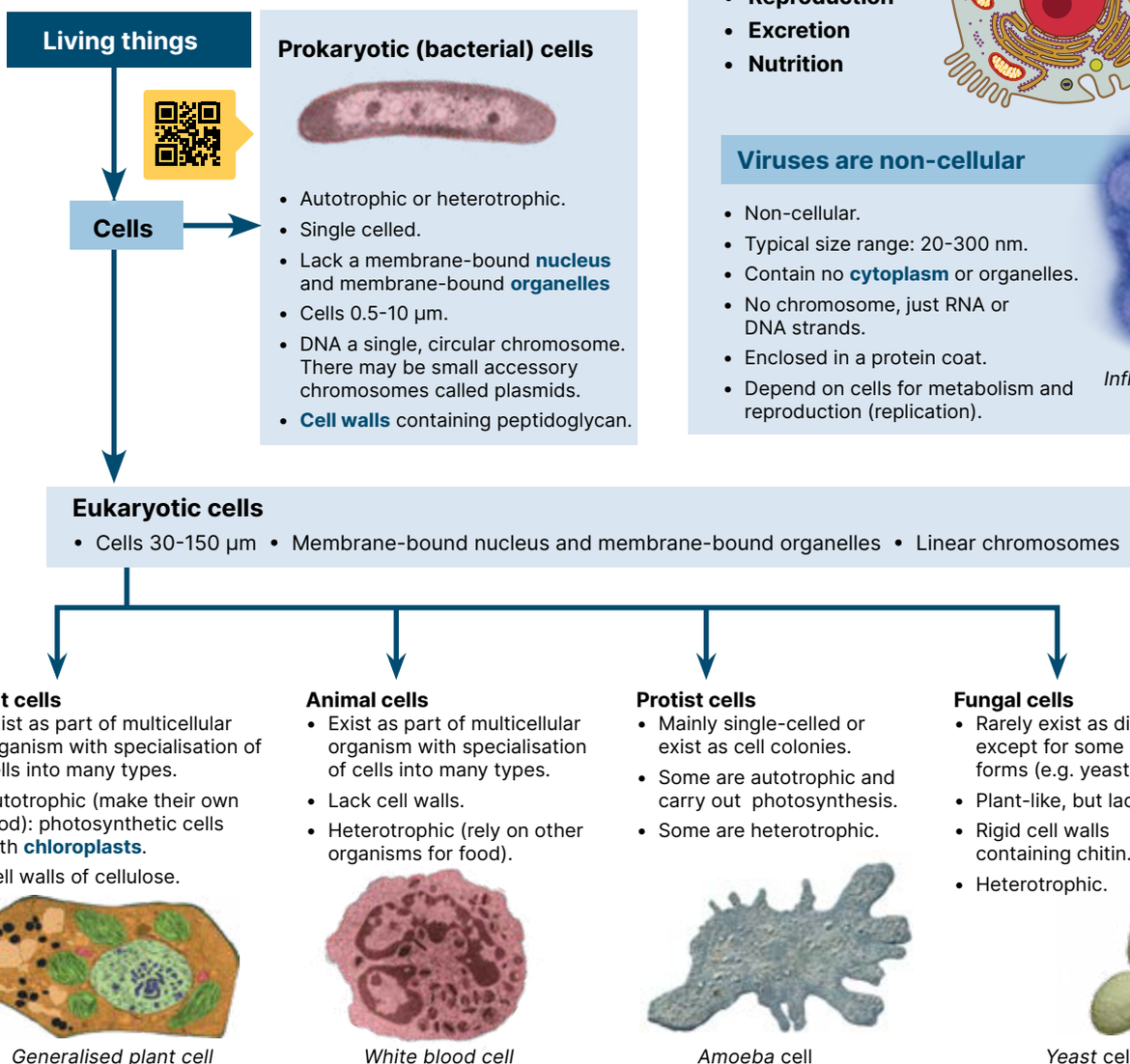


## Viruses are non-cellular

- Non-cellular.
- Typical size range: 20-300 nm.
- Contain no **cytoplasm** or organelles.
- No chromosome, just RNA or DNA strands.
- Enclosed in a protein coat.
- Depend on cells for metabolism and reproduction (replication).



Influenzavirus



1. What are the characteristic features of a prokaryotic cell? \_\_\_\_\_
2. What are the characteristic features of a eukaryotic cell? \_\_\_\_\_
3. Why are viruses considered to be non-cellular (non-living)? \_\_\_\_\_

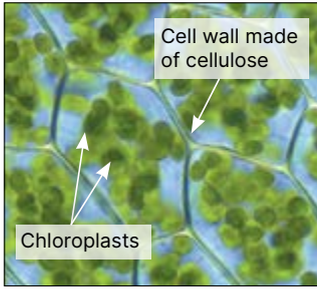
# 26

# Types of Cells

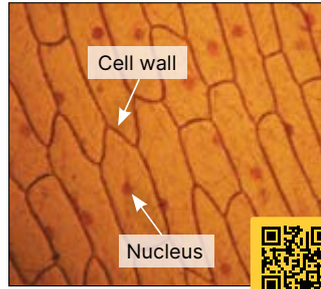
**Key Idea:** Cells come in a wide range of shapes and sizes. In multicellular organisms, cells are adapted for a specific role. Cells come in a wide range of types and forms. The images

below show a selection of cell types from the five kingdoms. Multicellular organisms typically have many specialised cell types, each of which performs a specific function.

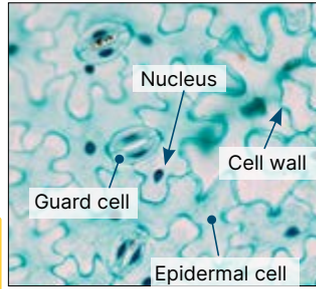
### Plant cells



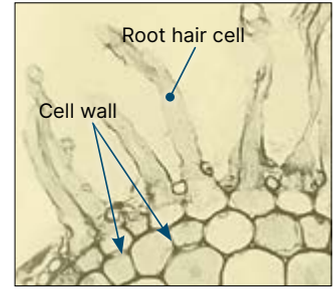
Palisade mesophyll cells



Epidermal cell

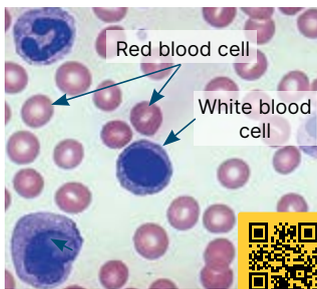


Guard cells and epidermal cells

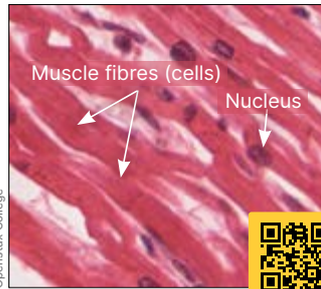


Root hair cell

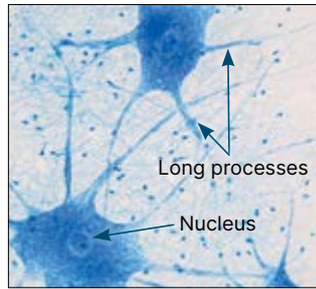
### Animal cells



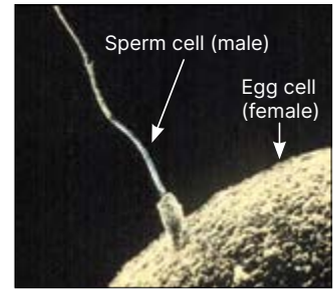
Blood cells



Muscle cells

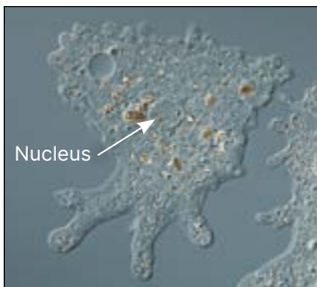


Nerve cells (neurons)

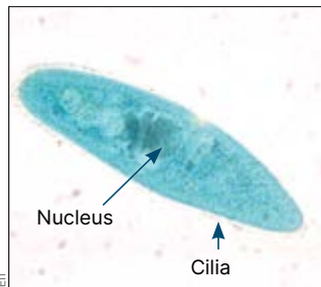


Reproductive cells

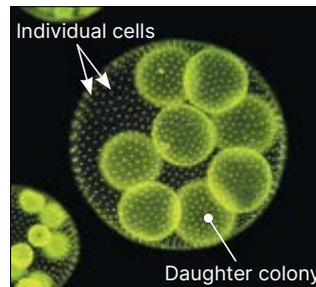
### Protists (single cells or colonies)



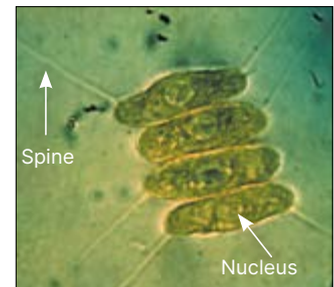
Amoeba



Paramecium

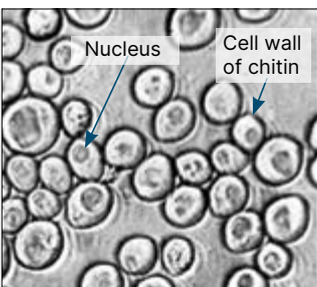


Volvox colony

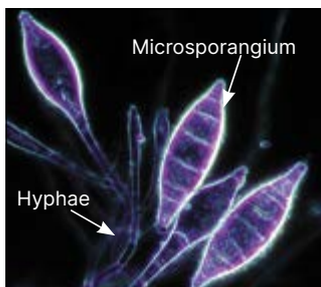


Scenedesmus colony

### Fungal cells

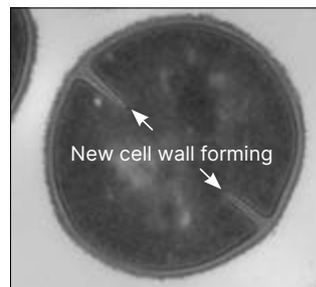


Yeast cells (*Saccharomyces*)

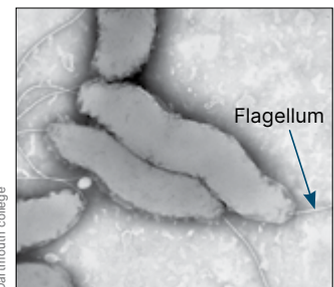


Microsporium cells

### Bacterial cells



Staphylococcus cell (dividing)



Campylobacter cell

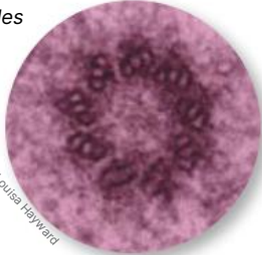
- Identify one distinguishing feature of each of the following cell types, based on what is (or is not) labelled above:
  - Plant cells: \_\_\_\_\_
  - Bacterial cells: \_\_\_\_\_
  - Fungal cells: \_\_\_\_\_
  - Animal cells: \_\_\_\_\_
- Both plants and animals have a large number of specialised cell types. Why do you think this is? \_\_\_\_\_

# What Are Cells Made Of?

**Key Idea:** The main components of a cell are water and compounds of carbon, hydrogen, nitrogen, and oxygen. Water is the main component of cells and organisms, providing an aqueous environment in which metabolic reactions can occur. Apart from water, most other substances in cells are compounds of carbon, hydrogen, oxygen, and nitrogen.

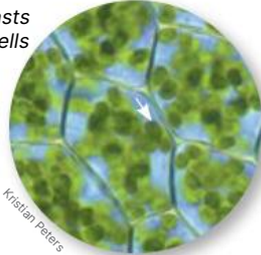
Carbon can combine with many other elements to form a large number of carbon-based (or organic) molecules. The organic molecules that make up living things can be grouped into four broad classes: carbohydrates, lipids, proteins, and nucleic acids. In addition, a small number of inorganic ions are also components of larger molecules.

Centrioles



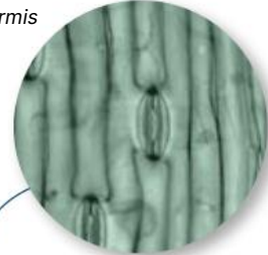
Louisa Hayward

Chloroplasts in plant cells



Kristian Peters

Plant epidermis

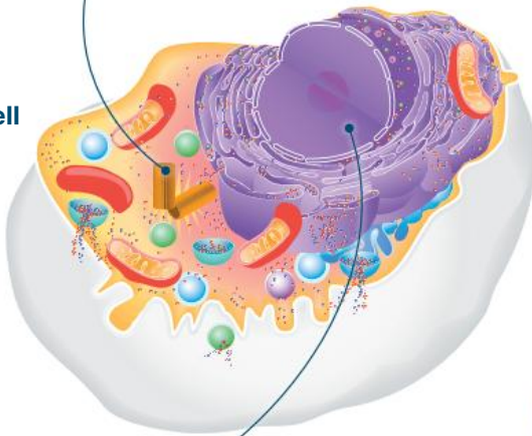


**Proteins** have an enormous number of structural and functional roles in plants and animals, e.g. as enzymes, structural materials (such as collagen), in transport, and movement (e.g. cytoskeleton and **centrioles**).

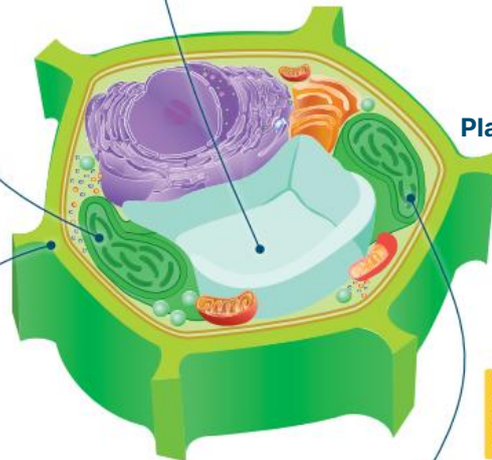
**Inorganic ions:** Dissolved ions participate in metabolic reactions and are components of larger organic molecules, e.g.  $Mg^{2+}$  is a component of the green chlorophyll pigment in the **chloroplasts** of green plants.

**Water** is a major component of cells: many substances dissolve in it and metabolic reactions occur in it. In plant cells, fluid pressure against the cell wall provides turgor, which supports the cell.

Animal cell



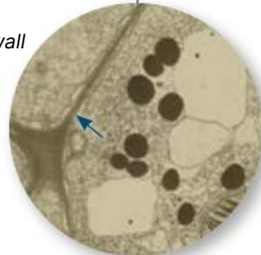
Plant cell



Chromosome



Plant cell wall



Chloroplast membranes



**Nucleotides and nucleic acids**  
Nucleic acids encode information for the construction and functioning of an organism (DNA and RNA). ATP, a nucleotide derivative, is the energy carrier of the cell.

**Carbohydrates** form the structural components of cells, e.g. cellulose **cell walls** (arrowed). They are important in energy storage and they are involved in cellular recognition.

**Lipids** provide a concentrated source of energy. Phospholipids are a major component of cellular membranes, including the membranes of organelles such as chloroplasts and mitochondria.

1. Given the components of cells above, predict some substances they need to remain functioning and the importance of those substance:

---



---



---



---



---

# What Cells Need for Survival

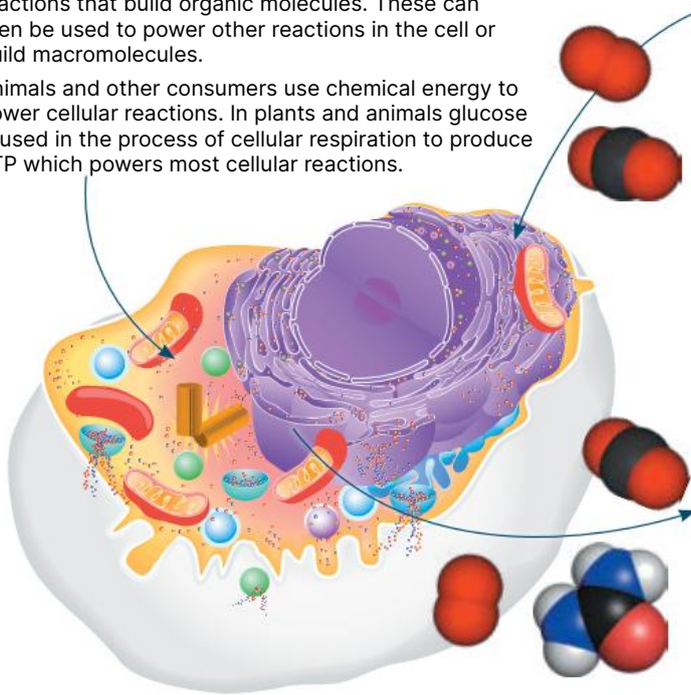
**Key Idea:** Cells have specific requirements for survival. These include obtaining nutrients and removing wastes. Cells require energy to power the reactions that build their

structures and maintain their functions. Cells also require a range of molecules and ions to build and maintain these structures and they need to be able to remove wastes.

## Cells need energy

Cells have evolved to use two basic forms of energy: light or chemical energy.

- Plant and algal cells containing **chloroplasts** and some bacteria use light from the Sun to power chemical reactions that build organic molecules. These can then be used to power other reactions in the cell or build macromolecules.
- Animals and other consumers use chemical energy to power cellular reactions. In plants and animals glucose is used in the process of cellular respiration to produce ATP which powers most cellular reactions.



## Cells require resources

Cells require molecules and ions to build macromolecules and help carry out cellular reactions.

- Carbon dioxide is needed by plants to build organic molecules during photosynthesis.
- Oxygen is needed by plants and animals as an electron acceptor at the end of cellular respiration.
- In plants, nitrates provide nitrogen, which is incorporated into amino acid molecules. Animals use these (by eating plants or plant eaters) to obtain building blocks for their proteins.
- Various metal ions are also needed. Some in relatively large amounts, e.g.  $\text{Na}^+$  is needed for nerve cell function in animals, while others are needed only in very small amounts.

## Cells need to remove wastes

Cells need to remove wastes generated during cellular reactions. What is regarded as a waste depends on the type of cell.

- Oxygen is a waste product of photosynthesis, but is required for cellular respiration.
- Other waste products include nitrogen wastes such as urea, ammonia, and uric acid (from metabolic processes).
- Most cellular reactions generate heat, which must be managed so that an organism does not overheat. In animals, metabolic heat is removed from cells by the blood and transferred to places where it can radiate into the environment (e.g. the skin).

## Cellular environments

The exact conditions a cell needs depends on many factors including whether the organism is unicellular or multicellular, and what environment it has evolved to survive in.

Jim Peaco, National Park Service PD



Some unicellular organisms (called thermophiles) can survive in temperatures as high as  $122^\circ\text{C}$ . Their enzymes can not function at the low temperatures experienced outside environments such as hot thermal pools.



Halophiles require environments with high salt concentrations (up to five times as concentrated as the sea). These cells are specially adapted to retain water. If placed in fresh water they quickly swell and burst.



Cells in multicellular organisms require the homeostatic environment provided by the organism. The organism provides an internal environment that provides the cells with nutrients, waste removal, and a relatively constant temperature.

- Why do cells need energy? \_\_\_\_\_  
\_\_\_\_\_
- Why must cells be able to remove wastes? \_\_\_\_\_  
\_\_\_\_\_
- Describe an example of where waste products of one cellular process can be used as a resource for another:  
\_\_\_\_\_  
\_\_\_\_\_



# 29

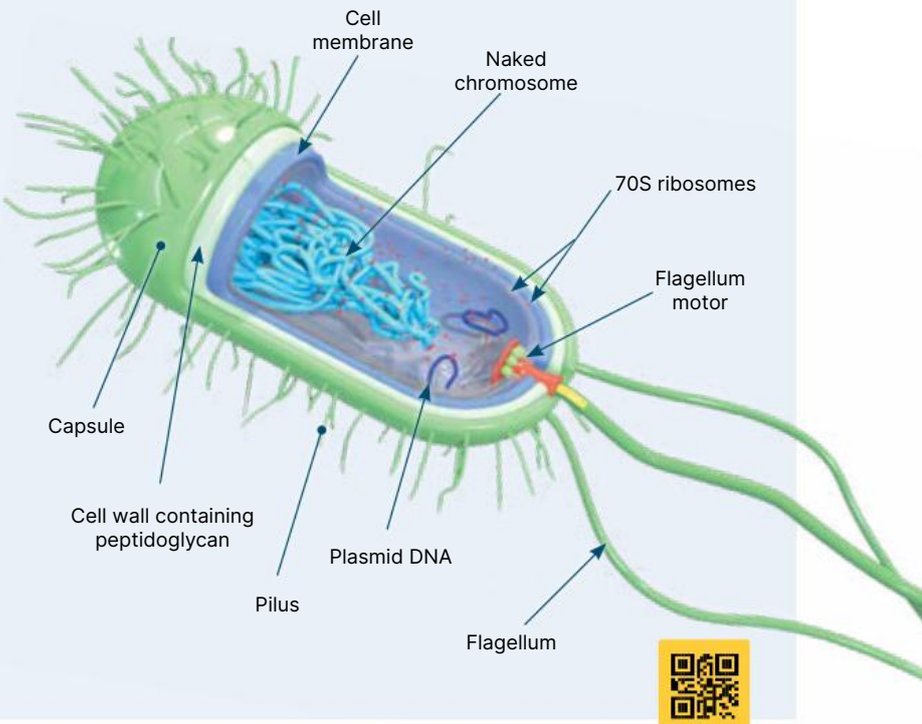
# Prokaryotic Cells

**Key Idea:** Prokaryotes are unicellular and have a relatively simple internal structure. Prokaryotes include the groups Bacteria and Archaea. They are much smaller and simpler than the cells of eukaryotes

and lack distinct **nucleus** and membrane-bound cellular organelles. Despite their simplicity compared to eukaryotes, prokaryotes are extremely diverse and can be found living in some of the most extreme environments on Earth.

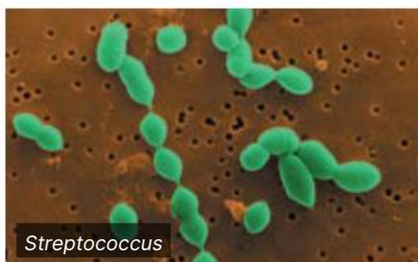
### Prokaryotic cells

- ▶ **Prokaryotic cells** are small (~0.5-10 μm) single cells. They lack any membrane-bound organelles.
- ▶ They are relatively unstructured with little cellular organisation. Their DNA, **ribosomes**, and enzymes are free floating within the cell's **cytoplasm**. The ribosomes (70S) are smaller than eukaryotic ribosomes.
- ▶ They have a single, circular chromosome of naked DNA (not associated with protein). They commonly have small, circular accessory chromosomes called plasmids.
- ▶ Some bacteria are photosynthetic. They have enzymes and light capturing membranes like those in eukaryotic **chloroplasts**.
- ▶ Prokaryotes have **cell walls**, but they are different in composition from the cell walls of eukaryotes.
- ▶ Examples of bacterial cells include the gut bacterium *Escherichia coli* and the cyanobacterium *Anabaena*.



### Prokaryote cell shapes

In terms of their appearance, there are only a few basic shapes found (shown below). The way in which members group together after division is often helpful in identifying certain species.



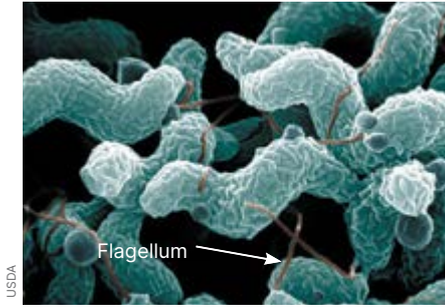
**Bacilli:** Rod-shaped bacteria that divide only across their short axis. Most occur as single rods, although pairs and chains are also found.

**Cocci:** usually round, but sometimes oval or elongated. When they divide, the cells stay attached to each other and may remain as pairs or clusters.

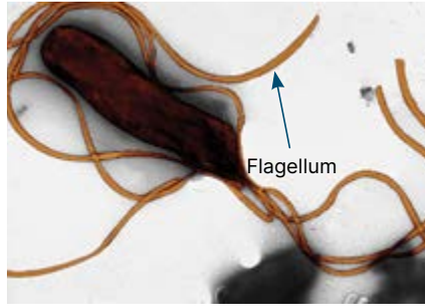
**Spirilla and vibrio:** Bacteria with one or more twists. Spirilla bacteria have a helical (corkscrew) shape. Bacteria that look like curved rods (comma shaped) are called vibrios.

1. Identify three distinguishing features of prokaryotes: \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_
2. Cyanobacterial cells (shown top) are photosynthetic. Describe features that enable them to capture and store energy:  
 \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_

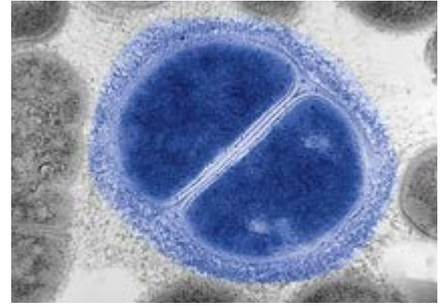
**Features of prokaryote cells**



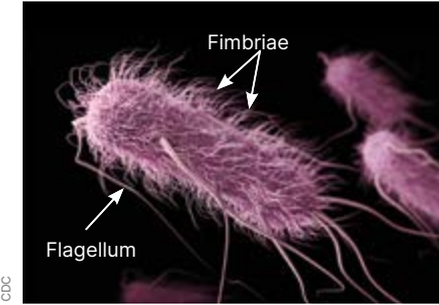
*Campylobacter jejuni* is a spiral bacterium responsible for foodborne intestinal disease. Note the single flagellum at each end for movement.



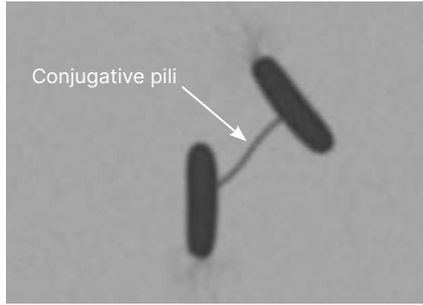
*Helicobacter pylori* is a comma-shaped vibrio bacterium that causes stomach ulcers in humans. This bacterium moves by means of multiple **flagella**.



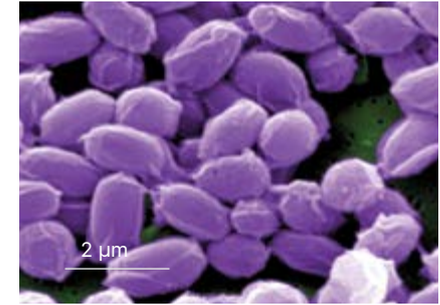
Bacteria usually divide by binary fission. During this process, DNA is copied and the cell splits into two cells, as seen in these gram positive cocci.



Some bacteria, such as *Escherichia coli*, a common gut bacterium (above), are covered with fimbriae which help it stick to surfaces (such as the gut wall). The flagellum helps it move about.



Bacteria are able to exchange genetic material via conjugative pili. The process, called conjugation, transfers plasmid DNA from one bacteria to another. This can help pass on useful genetic traits.



The scanning **electron micrograph** above shows endospores of *Bacillus anthracis* bacteria, which cause the disease anthrax. These heat-resistant spores can survive for many years and enable the bacteria to survive in a dormant state.

3. (a) Describe the function of flagella in bacteria: \_\_\_\_\_  
 \_\_\_\_\_
- (b) Explain how fimbriae differ structurally and functionally from flagella: \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_
4. (a) Describe the main method by which bacteria reproduce: \_\_\_\_\_  
 \_\_\_\_\_
- (b) Explain how conjugation differs from this usual method: \_\_\_\_\_  
 \_\_\_\_\_
5. What shape would you expect the following bacteria to be?
  - (a) *Staphylococcus aureus*: \_\_\_\_\_
  - (b) *Bacillus cereus*: \_\_\_\_\_
  - (c) *Spirillum volutans*: \_\_\_\_\_
6. Bacteria are extremely useful for human purposes and are used in a wide range of industries. Using the links in **BIOZONE's Resource Hub** or your own, find an industry where bacteria are used and give a brief description of their use:  
 \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_

# 30 Plant Cells

**Key Idea:** Plant cells are eukaryotic cells. They have features in common with animal cells, but also several unique features. **Eukaryotic cells** have a similar basic structure, although they may vary tremendously in size, shape, and function. Certain features are common to almost all eukaryotic cells, including their three main regions: a **nucleus**, surrounded by

a watery **cytoplasm**, which is itself enclosed by the **plasma membrane**. Plant cells are enclosed in a cellulose **cell wall**, which gives them a regular, uniform appearance. The cell wall protects the cell, maintains its shape, and prevents excessive water uptake. It provides rigidity to plant structures but permits the free passage of materials into and out of the cell.

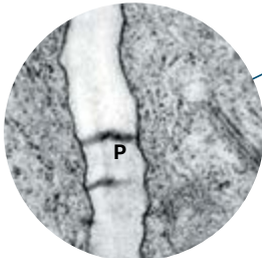
## Generalised plant cell

**Starch granule:** Carbohydrate stored in amyloplasts (specialised storage organelles).



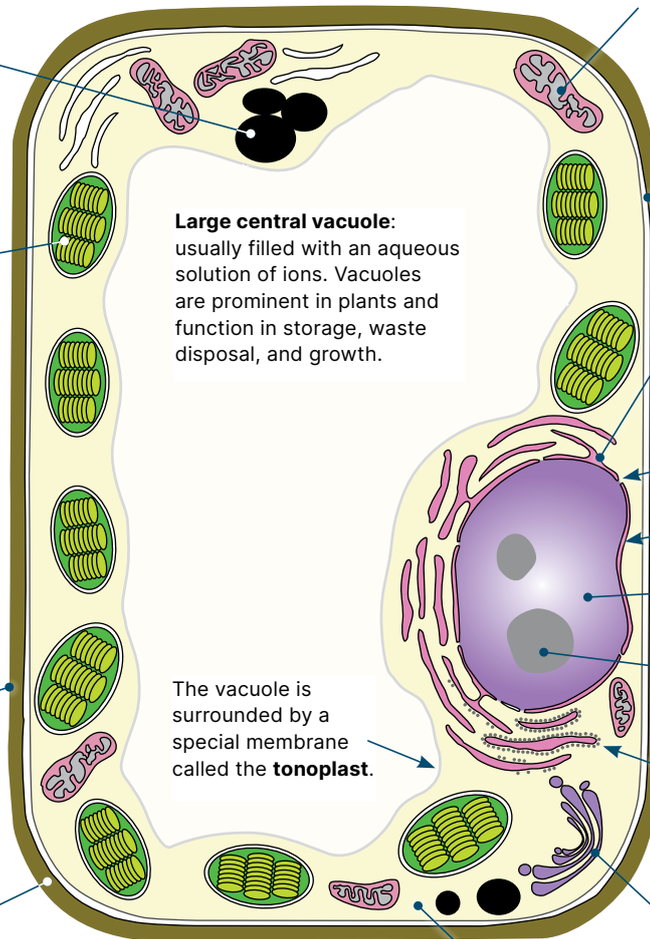
**Chloroplast:** Specialised organelles,  $2\ \mu\text{m} \times 5\ \mu\text{m}$ , containing the green pigment chlorophyll. Chloroplasts contain dense stacks of membranes within a fluid which is much like cytosol. They are the sites for photosynthesis and are found mainly in leaves. Chloroplasts are one of a group of double membraned organelles called **plastids**, which include amyloplasts (see above).

**Cell wall:** A semi-rigid structure outside the plasma membrane,  $0.1\ \mu\text{m}$  to several  $\mu\text{m}$  thick. It is composed mainly of cellulose. It supports the cell and limits its volume.



Alison Roberts

**Middle lamella** (seen here between adjacent cells left): The first layer of the cell wall formed during cell division. It contains pectin and protein, and provides stability. It allows the cells to form plasmodesmata (P), special channels that allow communication and transport to occur between cells.



**Large central vacuole:** usually filled with an aqueous solution of ions. Vacuoles are prominent in plants and function in storage, waste disposal, and growth.

The vacuole is surrounded by a special membrane called the **tonoplast**.

**Mitochondrion:**  $1.5\ \mu\text{m} \times 2\text{--}8\ \mu\text{m}$ . They are the cell's energy transformers, converting chemical energy into ATP.

**Plasma membrane:** Located inside the cell wall in plants, 3 to 10 nm thick.

**Endoplasmic reticulum (ER):** A network of tubes and flattened sacs. ER is continuous with the nuclear membrane and may be smooth or have attached ribosomes (rough ER).

**Nuclear pore:** 100 nm diameter

**Nuclear membrane:** a double layered structure.

**Nucleus:** A conspicuous organelle  $5\ \mu\text{m}$  diameter.

Nucleolus

**Ribosomes:** These small (20 nm) structures manufacture proteins. They may be free in the cytoplasm or associated with the surface of the endoplasmic reticulum.

Golgi apparatus

**Cytoplasm:** A watery solution containing dissolved substances, enzymes, and the cell organelles and structures.

- (a) What are the functions of the cell wall in plants? \_\_\_\_\_

\_\_\_\_\_

(b) Why is the middle lamella of the cell wall important? \_\_\_\_\_

\_\_\_\_\_
- What distinguishes the tonoplast and the plasma membrane? \_\_\_\_\_

\_\_\_\_\_
- (a) What structure takes up the majority of the space in the plant cell? \_\_\_\_\_

(b) What are its roles? \_\_\_\_\_
- Identify two structures in the diagram that are not found in animal cells: \_\_\_\_\_



# 31

## Identifying Structures in a Plant Cell

**Key Idea:** The position and appearance of the organelles in an electron micrograph can be used to identify them.

1. Study the diagrams on the other pages in this chapter to familiarise yourself with the structures found in eukaryotic cells. Identify the 11 structures in the cell below using the following word list: *cytoplasm, smooth endoplasmic reticulum, mitochondrion, starch granule, chromosome, nucleus, vacuole, plasma membrane, cell wall, chloroplast, nuclear membrane*

TEM

(a)

(b)

(c)

(d)

(e)

(f)

(g)

(h)

(i)

(j)

(k)

2. State how many cells, or parts of cells, are visible in the electron micrograph above: \_\_\_\_\_
3. Describe the features that identify this cell as a plant cell: \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_
4. (a) Explain where cytoplasm is found in the cell: \_\_\_\_\_  
 \_\_\_\_\_  
 (b) Describe what cytoplasm is made up of: \_\_\_\_\_  
 \_\_\_\_\_
5. Describe two structures, pictured in the cell above, that are associated with storage:
  - (a) \_\_\_\_\_  
 \_\_\_\_\_
  - (b) \_\_\_\_\_  
 \_\_\_\_\_

**Key Idea:** Animal cells are eukaryotic cells. They have many features in common with plant cells, but also have a number of unique features.

Animal cells, unlike plant cells, do not have a regular shape. In fact, some animal cells (such as phagocytes) are able to alter their shape for various purposes (e.g. engulfing

foreign material). The diagram below shows the structure and **organelles** of a liver cell. It contains organelles common to most relatively unspecialised human cells. Note the differences between this cell and the generalised plant cell. The plant cells activity provides further information on the organelles listed here but not described.

**Vacuoles:** Smaller than those found in plant cells. In animal cells, vacuoles have minor roles in exocytosis and endocytosis.

**Smooth endoplasmic reticulum:** ER without ribosomes. It is a site for lipid and carbohydrate metabolism, including hormone synthesis.

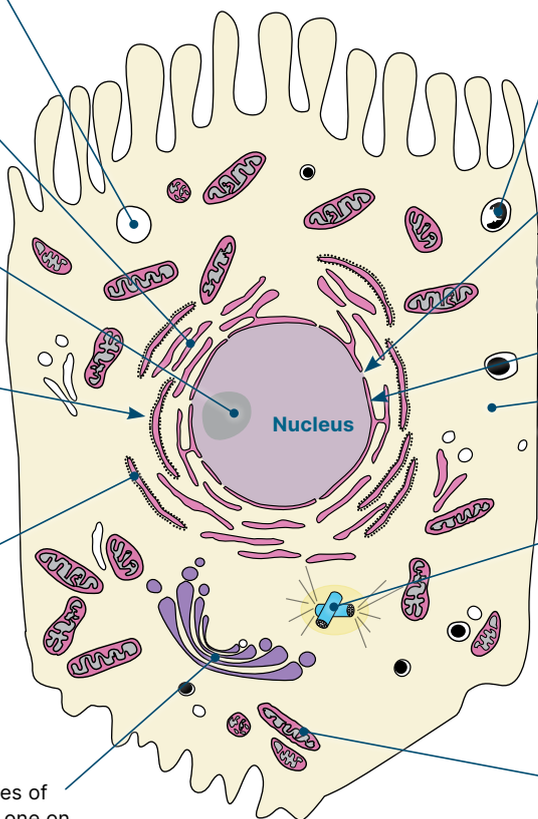
**Nucleolus:** A dense, solid structure composed of crystalline protein and nucleic acid. They are involved in ribosome synthesis.

**Ribosomes:** These small structures may be free in the cytoplasm or associated with the endoplasmic reticulum (ER). Ribosomes in animal cells are 80S ribosomes.

**Rough endoplasmic reticulum:** A site of protein synthesis. The rough ER also synthesises new membranes, growing in place by adding proteins and phospholipids.

**Golgi apparatus (20-200 nm):** A series of flattened, disc-shaped sacs, stacked one on top of the other and connected with the ER. The Golgi stores, modifies, and packages proteins. It 'tags' proteins so that they go to their correct destination.

## Generalised animal cell



**Lysosome:** A sac bounded by a single membrane. They are pinched off from the Golgi apparatus and contain and transport enzymes that break down food and foreign matter. Lysosomes show little internal structure but often contain fragments of material being broken down. Specialised lysosomes are generally absent from plant cells.

**Nuclear pore:** A hole in the nuclear membrane allowing the nucleus to communicate with the rest of the cell.

**Tight junctions:** Join cells together in the formation of tissues.

**Nuclear membrane:** Double layered.

**Cytoplasm**

**Plasma (cell surface) membrane**

**Centrioles:** Structures within a centrosome associated with nuclear division. They are composed of microtubules, but appear as small, featureless particles, 0.25 µm diameter, under a **light microscope**. They are absent in higher plant cells and some protists.

**Mitochondrion (pl. mitochondria):** An organelle bounded by a double membrane system. The number in a cell depends on its metabolic activity.



1. What is the difference between vacuoles in plant and animal cells? \_\_\_\_\_

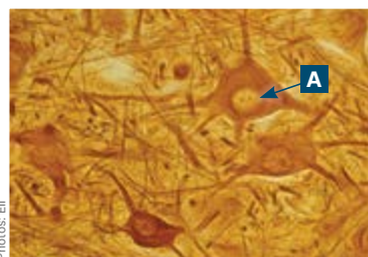
---



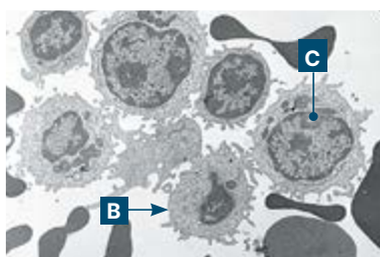
---

2. Name one structure or organelle present in generalised animal cells but absent from plant cells and describe its function: \_\_\_\_\_

---



Nerve cells in the spinal cord



White blood cells and red blood cells

3. The two photomicrographs, left, show several types of animal cells. Identify the features indicated by the letters A-C:

- (a) \_\_\_\_\_
- (b) \_\_\_\_\_
- (c) \_\_\_\_\_

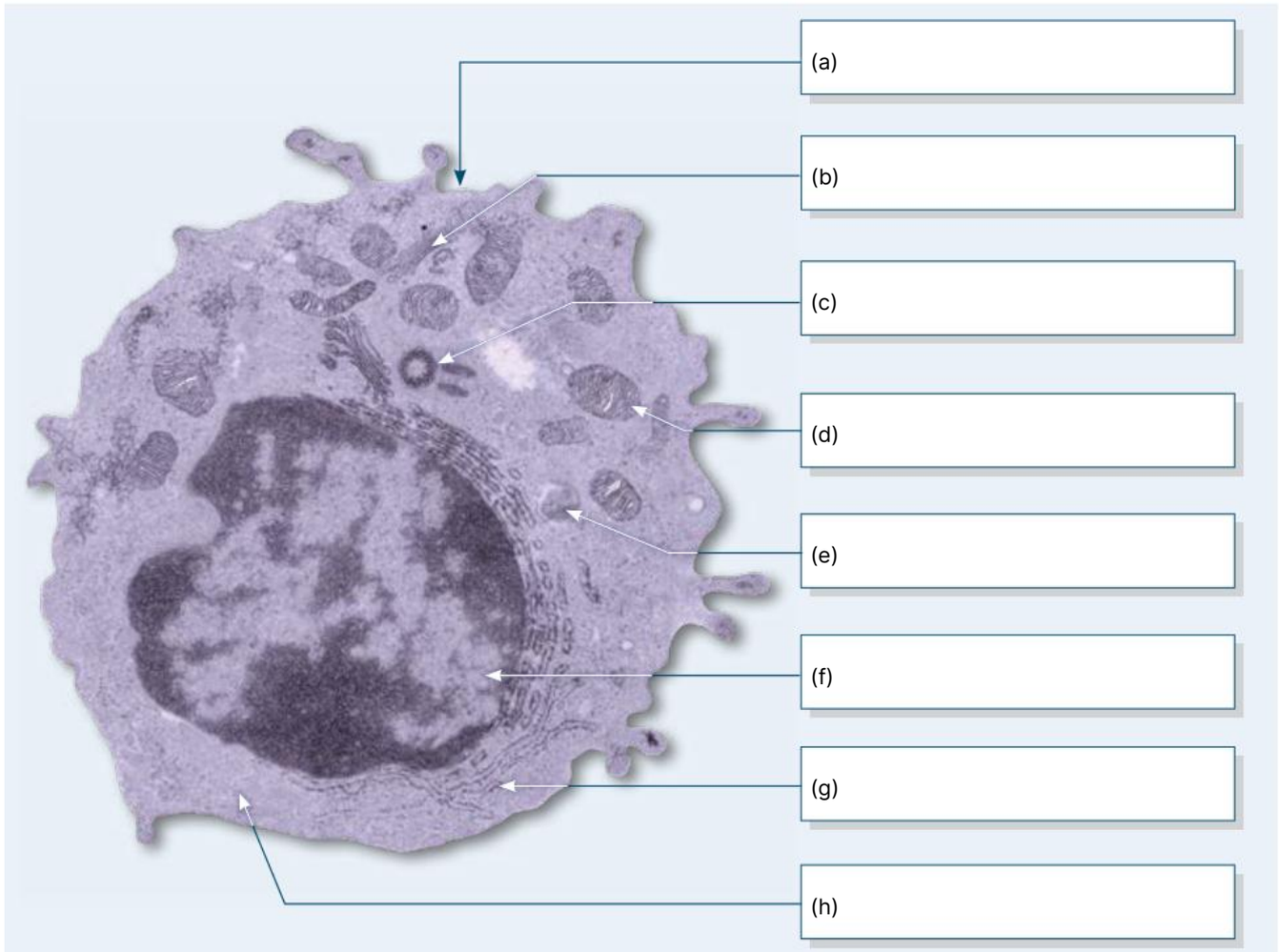
## 33

## Identifying Structures in an Animal Cell

**Key Idea:** The position of the organelles in an electron micrograph can result in variations in their appearance. Transmission electron microscopy (TEM) is the most frequently used technique for viewing cellular **organelles**.

When viewing TEMs, the cellular organelles may have quite different appearances depending on whether they are in transverse or longitudinal section.

1. Identify and label the structures in the animal cell below using the following list of terms: *cytoplasm, plasma membrane, rough endoplasmic reticulum, mitochondrion, nucleus, centriole, Golgi apparatus, lysosome*



2. Which of the organelles in the EM above are obviously shown in both transverse and longitudinal section?

\_\_\_\_\_

3. Why do plants lack any of the mobile phagocytic cells typical of animal cells? \_\_\_\_\_

\_\_\_\_\_

4. The animal cell pictured above is a lymphocyte. Describe the features that suggest to you that:

(a) It has a role in producing and secreting proteins: \_\_\_\_\_

\_\_\_\_\_

(b) It is metabolically very active: \_\_\_\_\_

\_\_\_\_\_

5. What features of the lymphocyte cell above identify it as eukaryotic? \_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_

**Key Idea:** Each type of organelle in a cell has a specific role. Not all cell types contain every type of organelle. The diagram below provides spaces for you to summarise

information about the **organelles** found in **eukaryotic cells**. The log scale of measurements (top of next page) illustrates the relative sizes of some cellular structures.

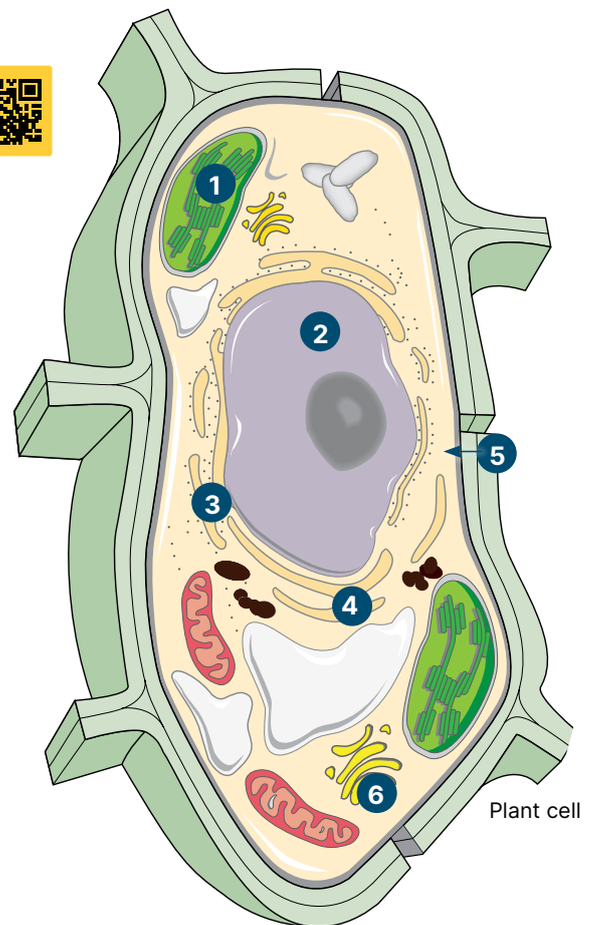
- Name this organelle:
  - Structure and location:
  - Function:
  - Is this organelle found in plant/animal/both cells?

- Name this organelle:
  - Structure and location:
  - Function:
  - Is this organelle found in plant/animal/both cells?

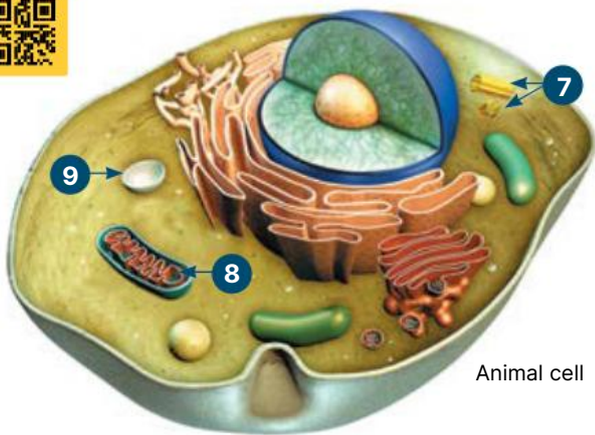
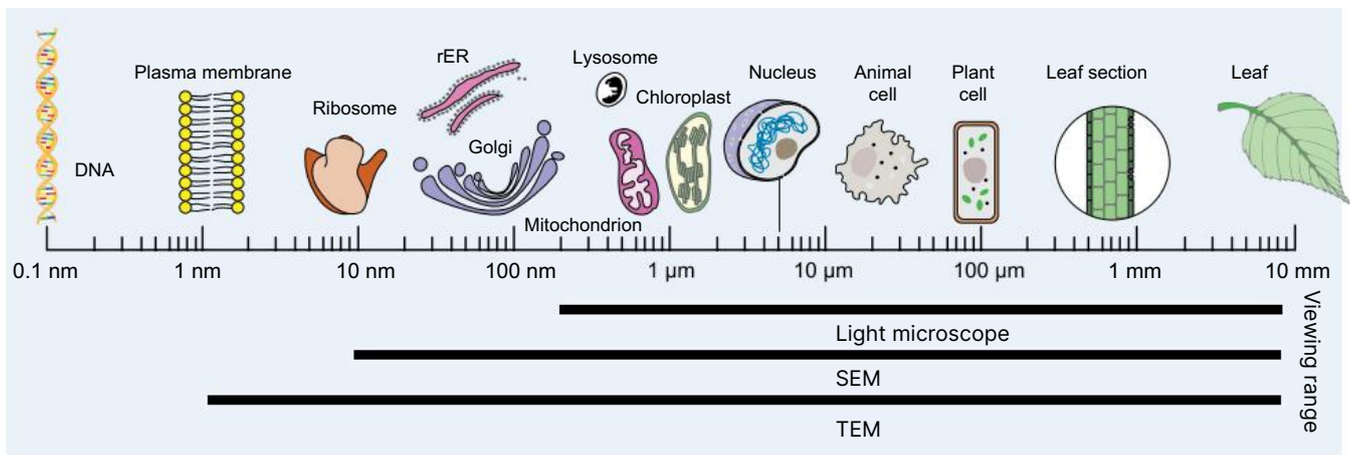
- Name this organelle:
  - Structure and location:
  - Function:
  - Is this organelle found in plant/animal/both cells?

- Name this organelle:
  - Structure and location:
  - Function:
  - Is this organelle found in plant/animal/both cells?

- Name this organelle:
  - Structure and location:
  - Function:
  - Is this organelle found in plant/animal/both cells?



- Name this organelle:
  - Structure and location:
  - Function:
  - Is this organelle found in plant/animal/both cells?



Animal cell

7. (a) Name this organelle:  
 (b) Structure and location:  
  
 (c) Function:  
  
 (d) Is this organelle found in plant/animal/both cells?

8. (a) Name this organelle:  
 (b) Structure and location:  
  
 (c) Function:  
  
 (d) Is this organelle found in plant/animal/both cells?

9. (a) Name this organelle:  
 (b) Structure and location:  
  
 (c) Function:  
  
 (d) Is this organelle found in plant/animal/both cells?

10. Use the scale at the top of the page and the information on previous activities to identify which of the organelles (1-9) can be seen through a light microscope:

---



---

11. Identify which of the organelles (1-9) require a TEM (transmission electron microscope) to be seen:

---



---

12. Identify one other structure in the plant cell not labelled on the previous page and describe its function:

---



---



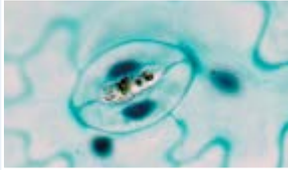
---

# 35 Optical Microscopes

**Key Idea:** Optical microscopes focus light through a series of lenses to magnify objects up to several hundred times. The light (or optical) microscope is an important tool in biology and using it correctly is an essential skill. High power compound **light microscopes** use visible light and

a combination of lenses to magnify objects up to several hundred times. The **resolution** of light microscopes is limited by the wavelength of light and specimens must be thin and mostly transparent so that light can pass through. No detail will be seen in specimens that are thick or opaque.

(a)



Stoma in leaf epidermis



(b)

(c)

(d)

Specimens viewed with a **compound light microscope** must be thin and mostly transparent so that light can pass through and structures can be seen. Modern microscopes are binocular, i.e. they have two adjustable eyepieces.

## Typical compound light microscope

**Word list:** In-built light source, arm, coarse focus knob, fine focus knob, condenser, mechanical stage, eyepiece lens, objective lens

(e)

(f)

(g)

(h)

### What is Magnification?

**Magnification** refers to the number of times larger an object appears compared to its actual size.

Magnification is calculated as follows:

$$\text{Objective lens power} \times \text{Eyepiece lens power}$$

(i)

(j)

(k)

(l)

Knob for the adjustment of the microscope on the arm



Apple seeds



(m)

## Dissecting microscope

**Word list:** Focus knob, stage, eyepiece lens, objective lens, eyepiece focus

### What is Resolution?

**Resolution** is the ability to distinguish between close together but separate objects. Examples of high and low resolution for separating two objects viewed under the same magnification are given below.

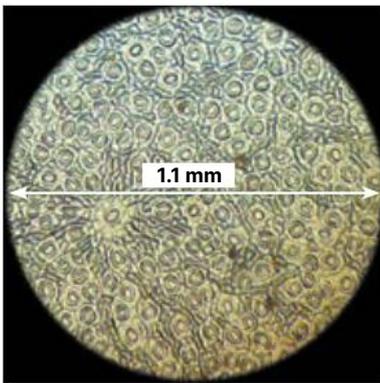
**High resolution** 

**Low resolution** 

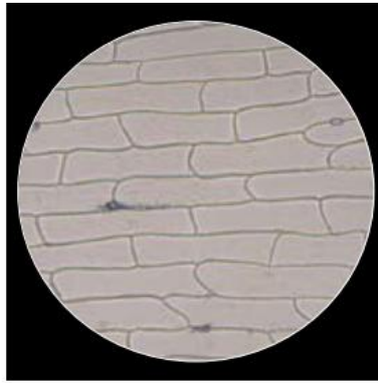
**Dissecting microscopes** are a special type of binocular microscope used for observations at low total magnification (X4 to X50), where a large working distance between the objectives and stage is required.

A dissecting microscope has two separate lens systems, one for each eye. Such microscopes produce a 3-D view of the specimen and are sometimes called stereo microscopes for this reason.

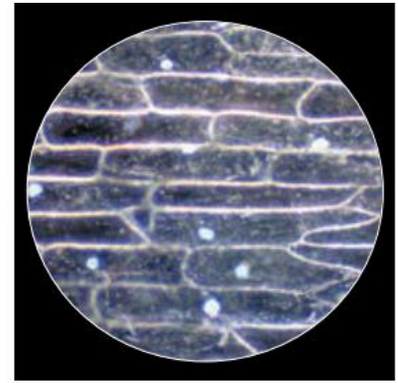




The field of view (FOV) of a microscope refers to the diameter of the area being viewed. The viewable area can be calculated from the diameter.



These onion epidermal cells are viewed with standard bright field lighting. Very little detail can be seen (only **cell walls**) and the cell nuclei are barely visible.



Dark field illumination is excellent for viewing specimens that are almost transparent. The nuclei of these onion epidermal cells are clearly visible.

Photos: EII

- Label the two photographs on the previous page, the compound light microscope (a) to (h) and the dissecting microscope (i) to (m). Use words from the lists supplied for each image.
- Determine the magnification of a microscope using:
  - 15 X eyepiece and 40 X objective lens: \_\_\_\_\_
  - 10 X eyepiece and 60 X objective lens: \_\_\_\_\_
- Describe the main difference between a compound light microscope and a dissecting microscope:
 

---



---
- What type of microscope would you use to:
  - Count stream invertebrates in a sample: \_\_\_\_\_
  - Observe cells in mitosis: \_\_\_\_\_
- Distinguish between magnification and resolution: \_\_\_\_\_  


---
  - Explain the benefits of a higher resolution: \_\_\_\_\_  


---
- Below is a list of ten key steps taken to set up a microscope and optimally view a sample. The steps have been mixed up. Put them in their correct order by numbering each step:
  - Focus and centre the specimen using the high objective lens. Adjust focus using the fine focus knob only.
  - Adjust the illumination to an appropriate level by adjusting the iris diaphragm and the condenser. The light should appear on the slide directly below the objective lens, and give an even amount of illumination.
  - Rotate the objective lenses until the shortest lens is in place (pointing down towards the stage). This is the lowest / highest power objective lens (delete one).
  - Place the slide on the microscope stage. Secure with the sample clips.
  - Fine tune the illumination so you can view maximum detail on your sample.
  - Focus and centre the specimen using the medium objective lens. Focus firstly with the coarse focus knob, then with the fine focus knob (if needed).
  - Turn on the light source.
  - Focus and centre the specimen using the low objective lens. Focus firstly with the coarse focus knob, then with the fine focus knob.
  - Focus the eyepieces to adjust your view.
  - Adjust the distance between the eyepieces so that they are comfortable for your eyes.

# 36 Preparing a Slide

**Key Idea:** Correctly preparing and mounting a specimen on a slide is important if structures are to be seen clearly under a microscope. A wet mount is suitable for most slides.

Specimens are usually prepared in some way before viewing in order to highlight features and reveal details. A wet mount is a temporary preparation in which a specimen and a drop of fluid are trapped under a thin coverslip. Wet mounts are used

to view thin tissue sections, live microscopic organisms, and suspensions such as blood. A wet mount improves a sample's appearance and enhances visible detail. Sections must be made very thin for two main reasons. A thick section stops light shining through making it appear dark when viewed. It also ends up with too many layers of cells, making it difficult to see any detail.

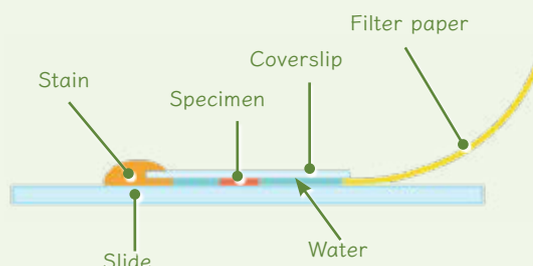
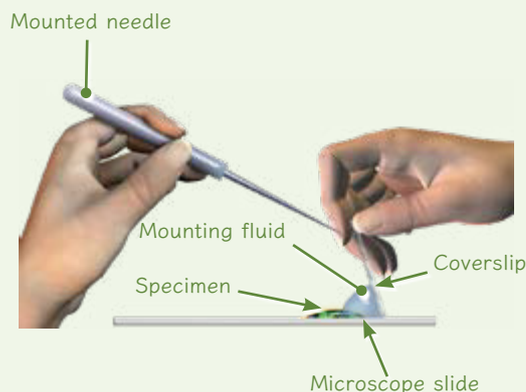
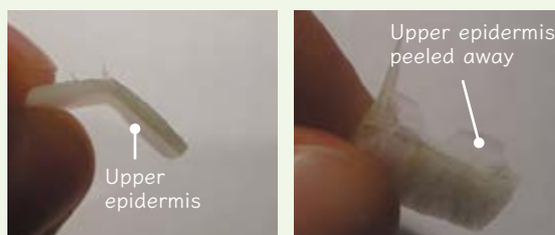
## Investigation 2.1 Preparing an onion slide

See appendix for equipment list.



**Caution is required when using scalpels or razors. Iodine stains skin and clothes, and irritates the eyes. You should wear protective eyewear.**

- Onions make good subjects for preparing a simple wet mount. Cut a square segment from a thick leaf of the bulb using a razor or scalpel.
- Bend the segment towards the upper epidermis until the lower epidermis and inner leaf tissue (the parenchyma) snaps so that just the upper epidermis is left attached.
- Carefully peel off the parenchyma from one side of the snapped leaf and then the other, leaving a peel of just the upper epidermis.
- Place peel in the centre of a clean glass microscope slide and cover it with a drop of water.
- Carefully lower a coverslip over the peel. A mounted needle can be used for better precision. This avoids including air in the mount.
- Use a small piece of tissue or filter paper to remove any excess water.
- Place the slide on the microscope tray. Locate the specimen or region of interest at the lowest magnification. Focus using the lowest magnification first (remembering to move the lens away from the slide) before switching to the higher magnifications.
- After viewing the slide under various magnifications, remove the slide and place it on the bench.
- At the edge of the coverslip place a small drop of iodine stain.
- On the opposite side of the coverslip use a piece of tissue or filter paper to draw the water out from under the coverslip. The iodine will be drawn under the coverslip.
- Replace the slide on the microscope and view the stained onion peel.
- Your teacher will have prepared slides of epithelial, connective, muscle, or nervous tissue available. View these under low and high power and note down any differences you see in the cells.



- Why must sections viewed under a microscope be very thin? \_\_\_\_\_
- Why do you think the specimen is covered with a coverslip? \_\_\_\_\_
- Why would no chloroplasts be visible in an onion epidermis cell slide? \_\_\_\_\_



**Stains and their uses**

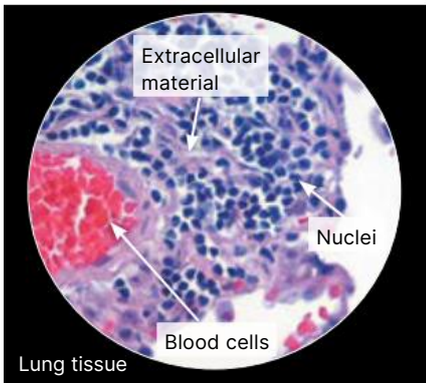
- ▶ Staining material for viewing under a microscope can make it easier to distinguish particular cell structures.
- ▶ **Stains** and dyes can be used to highlight specific components or structures. Stains contain chemicals that interact with molecules in the cell. Some stains bind to a particular molecule making it easier to see where those molecules are. Others cause a change in a target molecule, which changes their colour, making them more visible.
- ▶ Most stains are non-viable, and are used on dead specimens, but harmless viable stains can be applied to living material.

Some commonly used stains		
Stain	Final colour	Used for
Iodine solution	Blue-black	Starch
Crystal violet	Purple	Gram staining
Aniline sulfate	Yellow	Lignin
Methylene blue	Blue	Nuclei
Hematoxylin and eosin (H&E)	H=dark blue/violet E=red/pink	H=Nuclei E=Proteins



Photos: Eil

The light micrographs 1 and 2 (above) show how the use of a stain can enhance certain structures. The left image (1) is unstained and only the **cell wall** is easily visible. Adding iodine (2) makes the cell wall and nuclei stand out.



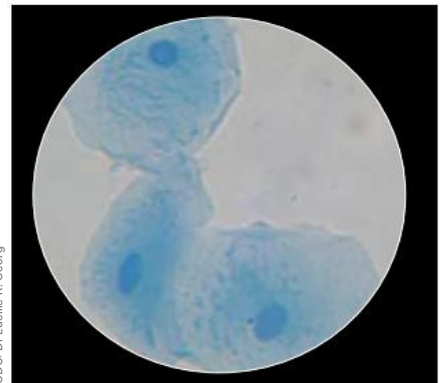
PloS

H&E stain is one of the most common stains for animal tissues. Nuclei stain dark blue, whereas proteins, extracellular material, and red blood cells stain pink or red.



Vinnoff

Viable stains do not immediately harm living cells. Trypan blue is a vital stain that stains dead cells blue but is excluded by live cells. It is also used to study fungal hyphae.



CDC: Dr. Lucille K. Georg

Methylene blue is a common temporary stain for animal cells, such as these cheek cells. It stains DNA and so makes the nuclei more visible.

4. Why is it necessary to focus on the lowest magnification first, before switching to higher magnifications?

---



---

5. Describe the difference the iodine stain made when viewing the onion cells under the microscope compared to when they were viewed without the stain:

---



---



---

6. What is the main purpose of using a stain? \_\_\_\_\_

---

7. What is the difference between a viable and non-viable stain? \_\_\_\_\_

---



---

8. Identify a stain that would be appropriate for distinguishing each of the following:

- (a) Live vs dead cells: \_\_\_\_\_ (c) Lignin in a plant root section: \_\_\_\_\_
- (b) Red blood cells in a tissue preparation: \_\_\_\_\_ (d) Nuclei in cheek cells: \_\_\_\_\_

## 37

## Calculating Linear Magnification

**Key Idea:** Magnification is how much larger an object appears compared to its actual size. It can be calculated from the ratio of image height to object height.

Microscopes produce an enlarged (magnified) image of an object allowing it to be observed in greater detail than is possible with the naked eye. **Magnification** refers to the number of times larger an object appears compared to its

actual size. Linear magnification is calculated by taking a ratio of the image height to the object's actual height. If this ratio is greater than one, the image is enlarged. If it is less than one, it is reduced. To calculate magnification, all measurements are converted to the same units. Often, you will be asked to calculate an object's actual size, in which case you will be told the size of the object and the magnification.

### Calculating linear magnification: A worked example

1 Measure the body length of the bed bug image (right). Your measurement should be 40 mm (not including the body hairs and antennae).

2 Measure the length of the scale line marked 1.0 mm. You will find it is 10 mm long. The magnification of the scale line can be calculated using equation 1 (below right).

The magnification of the scale line is **10** (10 mm ÷ 1 mm)

\*NB: The magnification of the bed bug image will also be 10x because the scale line and image are magnified to the same degree.

3 Calculate the actual (real) size of the bed bug using equation 2 (right):

The actual size of the bed bug is **4 mm**  
(40 mm ÷ 10 x magnification)



### Microscopy equations

$$1. \text{ Magnification} = \frac{\text{measured size of the object}}{\text{actual size of the object}}$$

$$2. \text{ Actual object size} = \frac{\text{size of the image}}{\text{magnification}}$$



1. The bright field microscopy image on the left is of onion epidermal cells. The measured length of the onion cell in the centre of the photograph is 52,000  $\mu\text{m}$  (52 mm). The image has been magnified 140 x. Calculate the actual size of the cell:

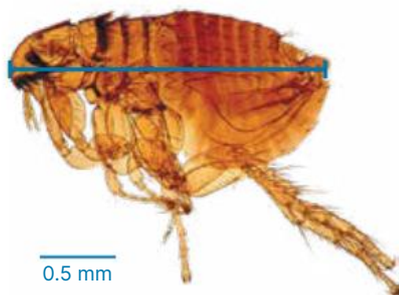
---



---



---



2. The image of the flea (left) has been captured using light microscopy.

(a) Calculate the magnification using the scale line on the image:

---



---



---

(b) The body length of the flea is indicated by a line. Measure along the line and calculate the actual length of the flea:

---



---



---



3. The image size of the *E. coli* cell (left) is 43 mm, and its actual size is 2  $\mu\text{m}$ . Using this information, calculate the magnification of the image:

---



---



---

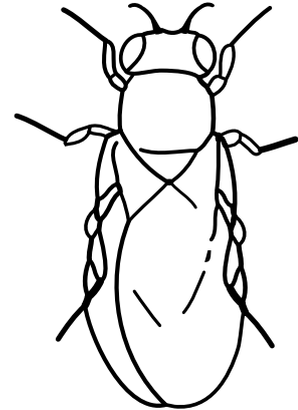


**Key Idea:** Good biological drawings provide an accurate record of the specimen you are studying and enable you to make a record of its important features.

Drawing is a very important skill to have in biology. Drawings record what a specimen looks like and give you an opportunity

to record its important features. Often drawing something will help you remember its features at a later date (e.g. in a test). Annotated drawings provide explanatory notes about the labelled structures, while plan diagrams label the main structures observed, but provide no additional detail.

- ▶ Biological drawings require you to pay attention to detail. It is very important that you draw what you actually see, and not what you think you should see.
- ▶ Biological drawings should include as much detail as you need to distinguish different structures and types of tissue, but avoid unnecessary detail which can make your drawing confusing.
- ▶ Attention should be given to the symmetry and proportions of your specimen. Accurate labelling, a statement of magnification or scale, the view (section type), and type of **stain** used (if applicable) should all be noted on your drawing.
- ▶ Some key points for making good biological drawing are described on the example below. The drawing of *Drosophila* (right) is well executed but lacks the information required to make it a good biological drawing.



This drawing of *Drosophila* is a fair representation of the animal, but has no labels, title, or scale.

All drawings must include a title. Underline the title if it is a scientific name.

→ **Copepod**

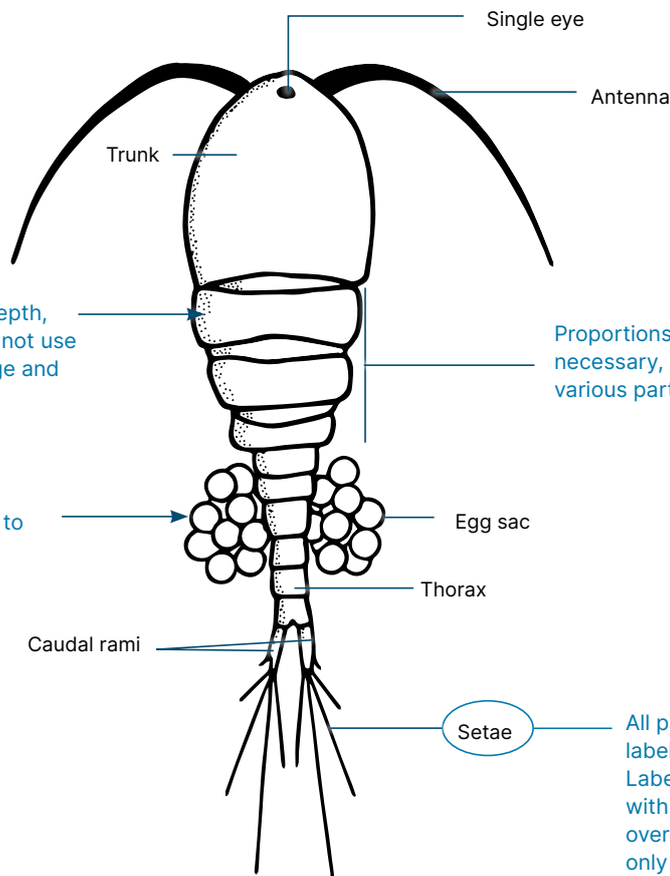
Centre your drawing on the page, not in a corner. This will leave room to place labels around the drawing.

If you need to represent depth, use stippling (dotting). Do not use shading as this can smudge and obscure detail.

Use simple, narrow lines to make your drawings.

Use a sharp pencil to draw with. Make your drawing on plain white paper.

Your drawing must include a scale or magnification to indicate the size of your subject.



Proportions should be accurate. If necessary, measure the lengths of various parts with a ruler.

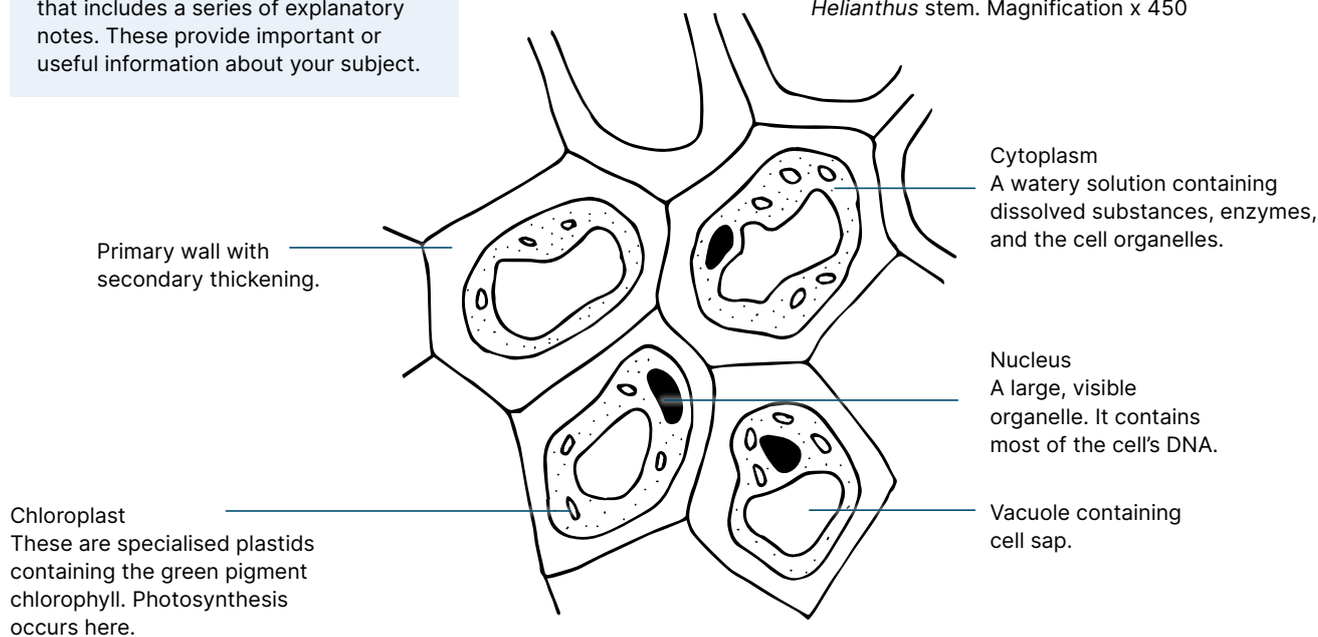
All parts of your drawing must be labelled accurately. Labelling lines should be drawn with a ruler and should not cross over other label lines. Try to use only vertical or horizontal lines.



### Annotated diagrams

An annotated diagram is a diagram that includes a series of explanatory notes. These provide important or useful information about your subject.

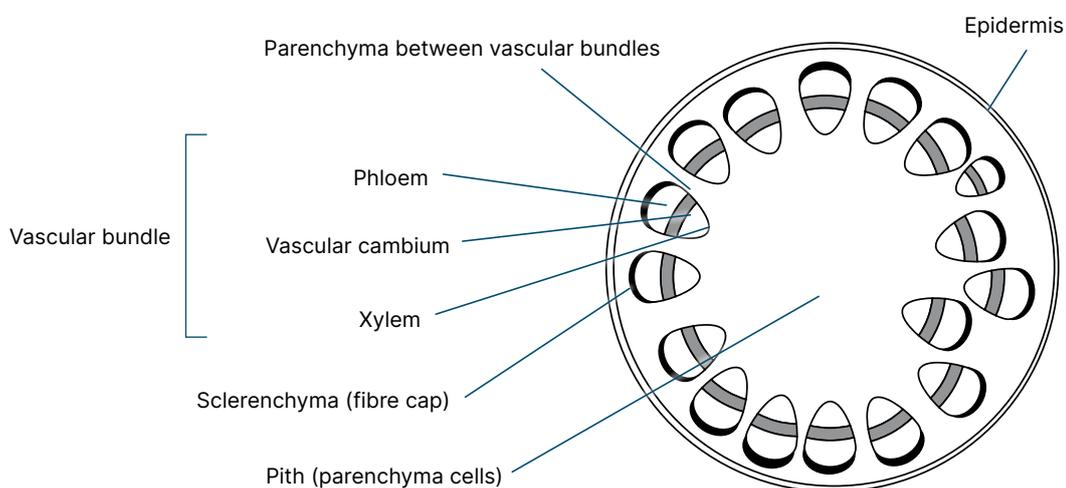
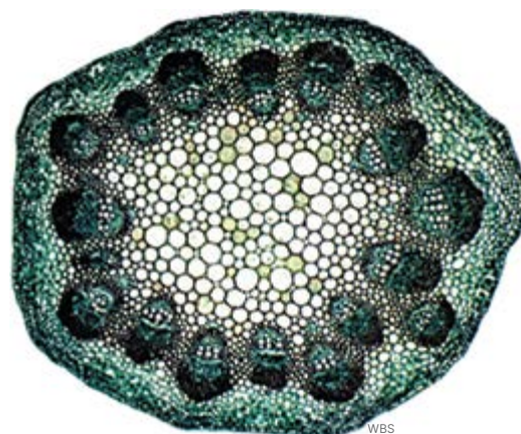
Transverse section through collenchyma of *Helianthus* stem. Magnification x 450



### Plan diagrams

Plan diagrams are drawings made of samples viewed with the naked eye, hand lens, or under a microscope at low or medium power. They are used to show the distribution of the different tissue types in a sample without any cellular detail. The tissues are identified, but no detail about the cells within them is included. The example here shows a plan diagram produced after viewing a light micrograph of a transverse section through a dicot stem.

Light micrograph of a transverse section through a dicot stem.

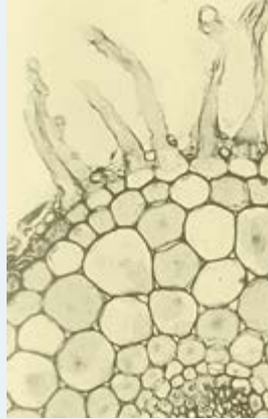


**Key Idea:** Attention to detail is vital when making accurate and useful biological drawings.

In this activity, you will practise the skills required to translate what is viewed into a good biological drawing.

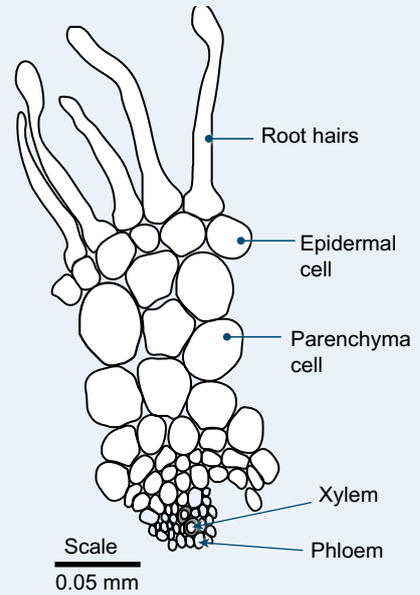


Above: Use relaxed viewing when drawing at the microscope. Use one eye (the left for right handers) to view and the right eye to look at your drawing.



Above: Light micrograph Transverse section (TS) through a *Ranunculus* root.

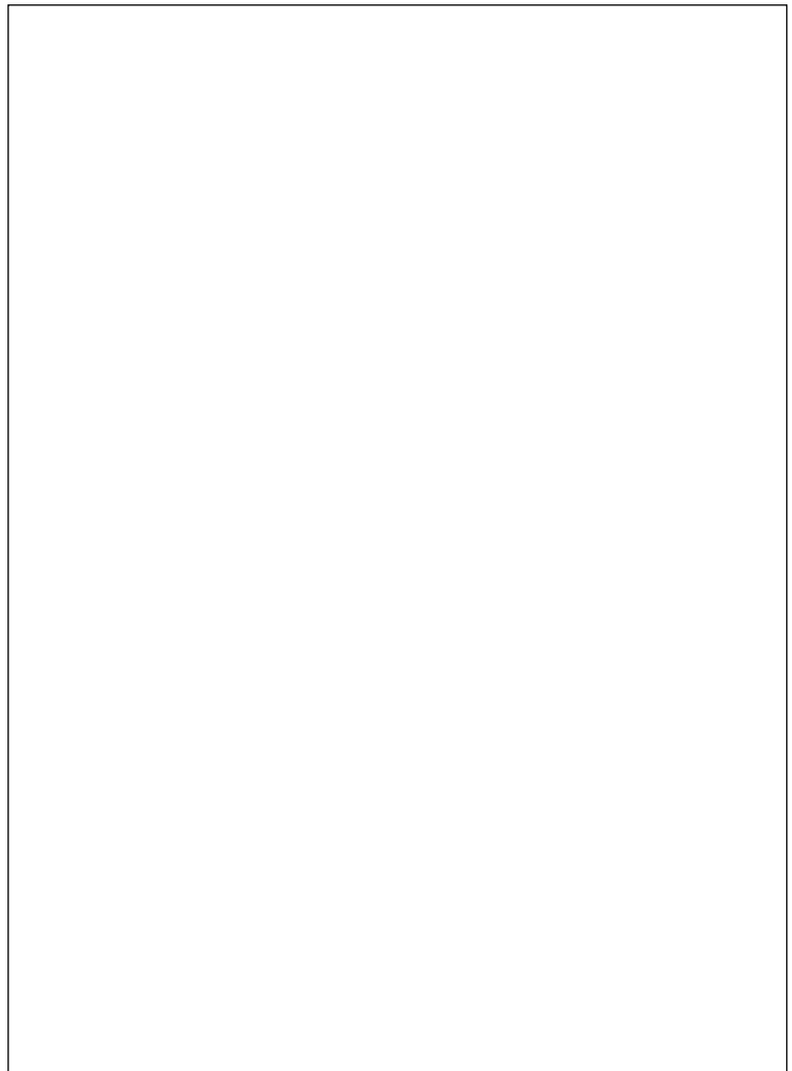
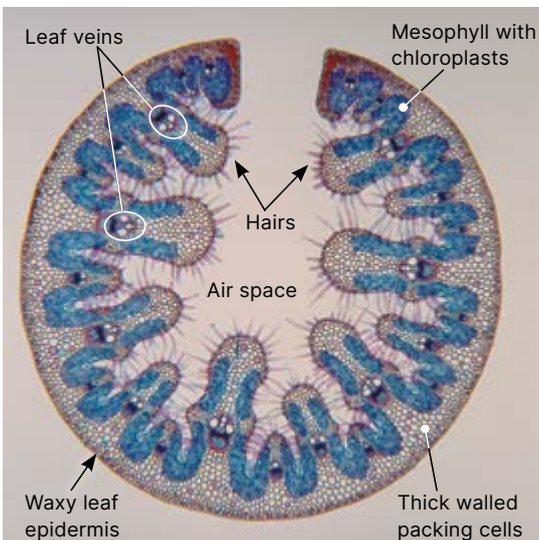
Right: A biological drawing of the same section.



1. During your course, you will study the features of cells and also make an investigation related to survival or an organism or species. You may need to identify and draw features of plant or animal tissues with a light microscope. Generally, only large organelles such as the nucleus and chloroplasts are easily seen at the magnifications typical of school microscopes (x 400).

In the space right, make a biological drawing of your own specimen or slide, or practise your drawing by making a plan diagram of the image below.

Below: A light micrograph of a leaf from the beach grass *Ammophila* below. The leaf is rolled inwards to reduce water loss.

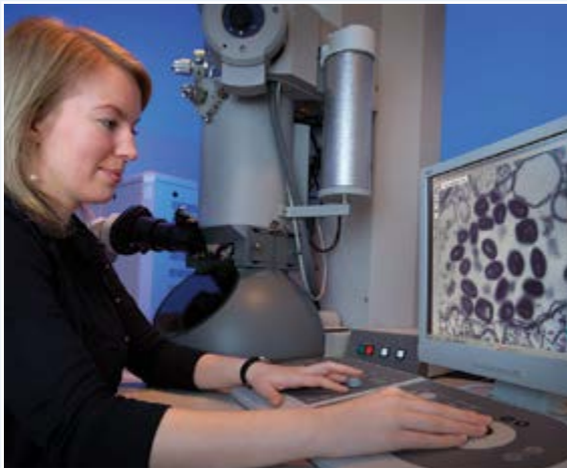


**Key Idea:** Electron microscopes use the short wavelengths of electrons to produce high resolution images of extremely small objects.

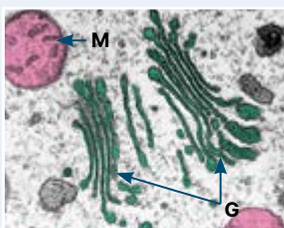
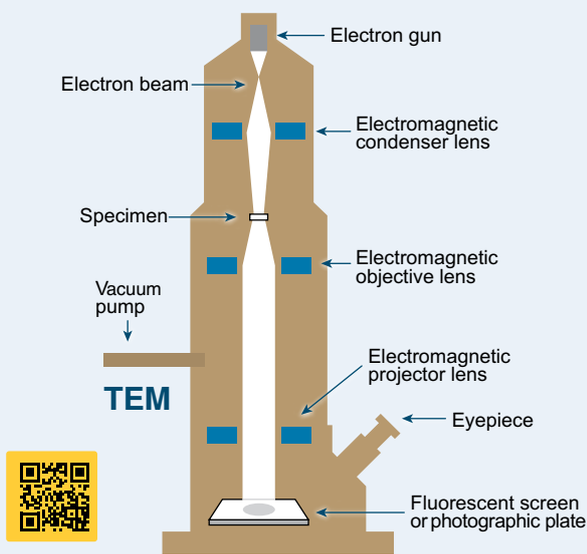
**Electron microscopes** (EMs) use a beam of electrons, instead of light, to produce an image. The higher **resolution** of electron microscopes is due to the shorter wavelengths of

electrons. There are two basic types of electron microscope: scanning electron microscopes (SEM) and transmission electron microscopes (TEM). In SEMs, the electrons are bounced off the surface of an object to produce detailed images of the external appearance. TEMs produce very clear images of specially prepared thin sections.

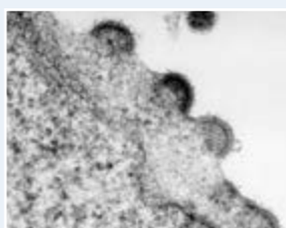
## Transmission electron microscope (TEM)



The transmission electron microscope is used to view extremely thin sections of material. Electrons pass through the specimen and are scattered. Magnetic lenses focus the image onto a fluorescent screen or photographic plate. The sections are so thin that they have to be prepared with a special machine, called an ultramicrotome, which can cut wafers to just 30 thousandths of a millimetre thick. It can magnify several hundred thousand times.



TEM photo showing the Golgi (G) and a mitochondrion (M).

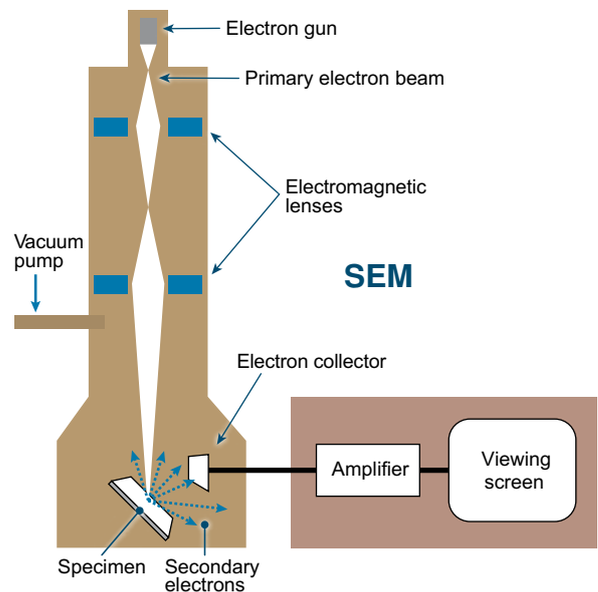


Three HIV viruses budding out of a human lymphocyte (TEM).

## Scanning electron microscope (SEM)



The scanning electron microscope scans a sample with a beam of primary electrons, which knocks electrons from the surface. These secondary electrons are picked up by a collector, amplified, and transmitted onto a viewing screen or photographic plate, producing a 3-D image. A microscope of this power easily obtains clear images of very small organisms such as bacteria, and small particles such as viruses. The image produced is of the outside surface only.



SEM photo of stoma and epidermal cells on the upper surface of a leaf.



Image of hair louse clinging to two hairs on a Hooker's sealion (SEM).

	Light microscope	Transmission electron microscope (TEM)	Scanning electron microscope (SEM)
<b>Radiation source:</b>	light	electrons	electrons
<b>Wavelength:</b>	400-700 nm	0.005 nm	0.005 nm
<b>Lenses:</b>	glass	electromagnetic	electromagnetic
<b>Specimen:</b>	living or non-living supported on glass slide	non-living supported on a small copper grid in a vacuum	non-living supported on a metal disc in a vacuum
<b>Maximum resolution:</b>	200 nm	1 nm	10 nm
<b>Maximum magnification:</b>	1500 x	250 000 x	100 000 x
<b>Stains:</b>	coloured dyes	impregnated with heavy metals	coated with carbon or gold
<b>Type of image:</b>	coloured, surface or section	monochrome, section	monochrome, surface only

1. Explain why electron microscopes are able to resolve much greater detail than a light microscope:

---



---



---

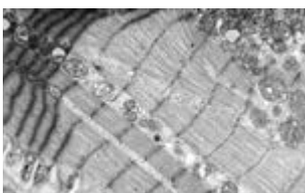
2. Which type of microscope [TEM, SEM, compound light microscope, or dissecting microscope] would you use for each of the following scenarios. Explain your choice in each case:

(a) Distinguishing extinct plant species on the basis of pollen surface features: \_\_\_\_\_

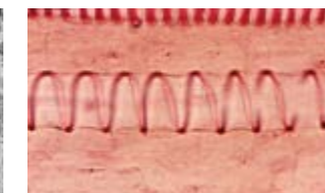
(b) Imaging the ultrastructure of a chloroplast: \_\_\_\_\_

(c) Performing a count of white blood cells from the blood of a person with an infection: \_\_\_\_\_

3. Identify which type of electron microscope (SEM or TEM) or optical microscope (compound light microscope or dissecting) was used to produce each of the images in the photos below (A-H):



Cardiac muscle



Plant vascular tissue



Mitochondrion



Plant epidermal cells

A. \_\_\_\_\_

B. \_\_\_\_\_

C. \_\_\_\_\_

D. \_\_\_\_\_



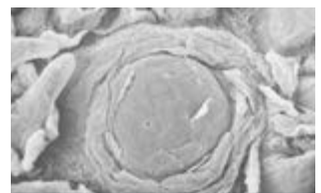
Head louse



Kidney cells



Body louse



Tongue papilla

E. \_\_\_\_\_

F. \_\_\_\_\_

G. \_\_\_\_\_

H. \_\_\_\_\_

4. Research and list some types of ultra high resolution microscopes: \_\_\_\_\_

---

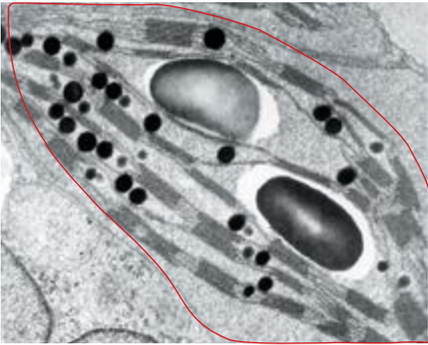
# 41

## Identifying Organelles

**Key Idea:** Cellular organelles can be identified in electron micrographs by their specific features.

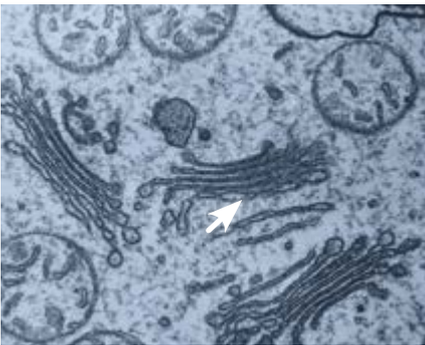
**Electron microscopes** produce a magnified image at high

**resolution** (distinguish between close together but separate objects). The transmission electron microscope (TEM) images below show the ultrastructure of some **organelles**.



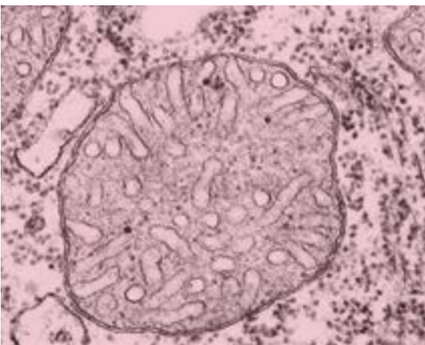
WWU

1. (a) Name the circled organelle: \_\_\_\_\_  
 (b) Which kind of cell(s) would this organelle be found in?  
 \_\_\_\_\_  
 \_\_\_\_\_  
 (c) Describe the function of this organelle: \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_



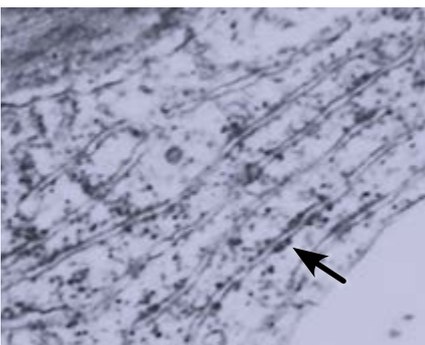
WWU

2. (a) Name this organelle (arrowed): \_\_\_\_\_  
 (b) State which kind of cell(s) this organelle would be found in:  
 \_\_\_\_\_  
 (c) Describe the function of this organelle: \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_



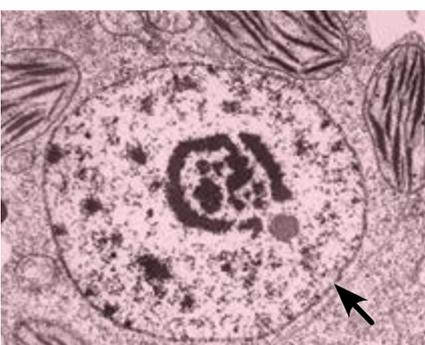
WWU

3. (a) Name the large, circular organelle: \_\_\_\_\_  
 (b) State which kind of cell(s) this organelle would be found in:  
 \_\_\_\_\_  
 (c) Describe the function of this organelle: \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_



WWU

4. (a) Name and label the ribbon-like organelle in this photograph (arrowed):  
 \_\_\_\_\_  
 (b) State which kind of cell(s) this organelle is found in:  
 \_\_\_\_\_  
 (c) Describe the function of this organelle: \_\_\_\_\_  
 \_\_\_\_\_  
 (d) Name the dark 'blobs' attached to the organelle you have labelled:  
 \_\_\_\_\_



BF

5. (a) Name this large circular organelle (arrowed): \_\_\_\_\_  
 (b) State which kind of cell(s) this organelle would be found in:  
 \_\_\_\_\_  
 (c) Describe the function of this organelle: \_\_\_\_\_  
 \_\_\_\_\_  
 (d) Label three features relating to this organelle in the photograph.

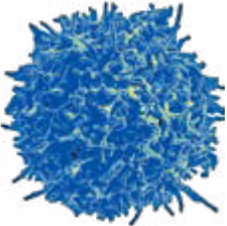
# 42

## Did You Get It?

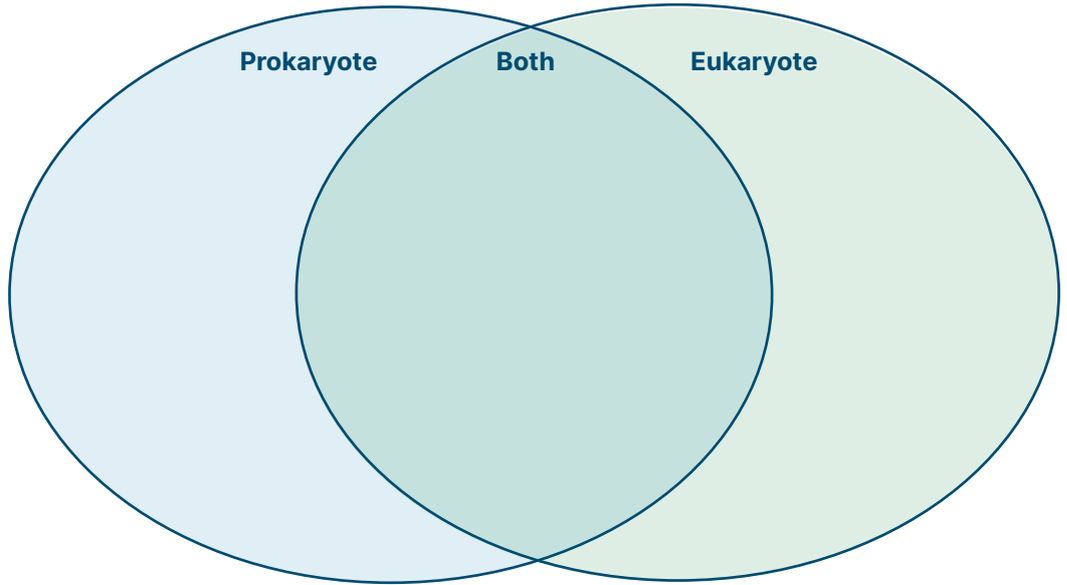
1. Use the Venn diagram below to compare prokaryote and eukaryote cells:



*E. coli* (prokaryote)



T lymphocyte (eukaryote)



2. (a) Identify organelle 1: \_\_\_\_\_

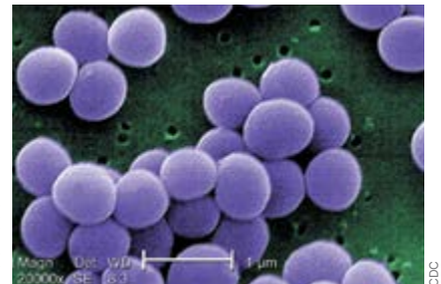
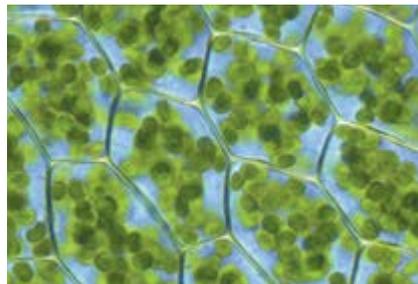
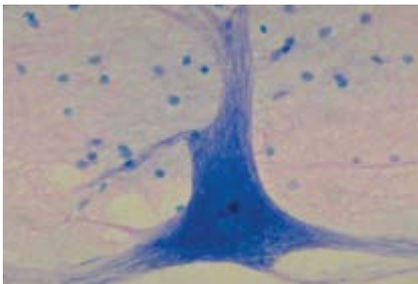
(b) The organelle in (a) is found in a plant cell / animal cell / both plant and animal cells:  
\_\_\_\_\_

(c) Identify organelle 2: \_\_\_\_\_

(d) The organelle in (a) is found in a plant cell / animal cell / both plant and animal cells:  
\_\_\_\_\_



3. For each of the following images of cells, identify the cell type (plant, animal, bacterial), give a brief reason for your decision and note any organelles or structures visible.



(a) \_\_\_\_\_  
\_\_\_\_\_  
\_\_\_\_\_

(b) \_\_\_\_\_  
\_\_\_\_\_  
\_\_\_\_\_

(c) \_\_\_\_\_  
\_\_\_\_\_  
\_\_\_\_\_

4. Write the number at the start of the sentence on the left in front of the correct sentence ending on the right:

- 1. Cells are the basic...
- 2. A cell is enclosed by a plasma membrane...
- 3. Plant cells have...
- 4. Animal cells do...
- 5. Eukaryotic cells contain many different types of organelle...
- 6. Each organelle carries out a specific function in the cell...
- 7. Prokaryotic cells...

- \_\_\_\_\_ ...such as photosynthesis or respiration.
- \_\_\_\_\_ ...a cell wall of cellulose.
- \_\_\_\_\_ ...do not contain membrane-bound organelles.
- \_\_\_\_\_ ...units of life.
- \_\_\_\_\_ ...not have a cell wall.
- \_\_\_\_\_ .. some of which are composed of membranes.
- \_\_\_\_\_ ...made of a phospholipid bilayer



# Cellular Differentiation and Specialisation

## Key Terms

- adult stem cell
- bioartificial tissue
- bioethics
- cell
- cell specialisation
- circulatory system
- connective tissue
- differentiation (cellular)
- digestive system
- embryonic stem cell
- epithelial tissue
- ethics
- excretory system
- hierarchical organisation
- gas exchange
- mitosis
- multipotent
- muscle tissue
- nervous tissue
- nutrition
- organ
- pluripotent
- potency
- respiratory system
- self renewal
- specialised cell
- stem cell
- system (body)
- tissue
- totipotent
- unipotent
- waste (removal)
- zygote

## Key Concepts

- ▶ Stem cells have the capacity to differentiate into different cell types.
- ▶ Multicellular organisms have a hierarchical structure from cells to organ systems.
- ▶ Organ systems interact with one another for maximum efficiency.

### Types and properties of stem cells

Activity Number

- |     |   |        |
|-----|---|--------|
| □ 1 | Describe the properties of stem cells, including self-renewal and potency. Describe the properties and features of different types of stem cell.  | 43     |
| □ 2 | Explain the process of cell differentiation and the role of stem cells in producing specialised cells that make up tissues and organs in multicellular organisms.   | 44     |
| □ 3 | Explain the process of mitosis in an animal cell with reference to the stages of interphase, prophase, metaphase, anaphase, telophase and cytokinesis.  | 45     |
| □ 4 | <b>SHE:</b> Explain the potential of pluripotent stem cells to develop into specialised cells that can be utilised for the restoration or substitution of failing organs and tissues. Discuss how advancements in technology can be used to produce pluripotent stem cells. | 43, 46 |
| □ 5 | <b>SI:</b> Explore the safety, ethics, and efficacy of stem cell technologies. Discuss the use of adult and embryonic stem cells (ASC and ESC) in medical technology. Analyse data and evaluate alternative perspectives on the use of stem cell research.                  | 46-47  |

### Multicellular organisms have a hierarchical structure

- |      |   |           |
|------|---|-----------|
| □ 6  | Using examples, describe the hierarchical structure of multicellular organisms, including reference to organelles, cells, tissues, organs, and organ systems.   | 48        |
| □ 7  | Explain how a hierarchical organisation builds structural complexity and contributes to the functional efficiency of the organism as a whole. Include reference to emergent properties. Discuss the presence of specialised cells, with varying organelle composition, adapted to allow each body system to function optimally.     | 48, 50-52 |
| □ 8  | Explore aspects of hierarchical organisation through a dissection of a mammalian organ such as a heart or kidney. Explain how the arrangement and interaction of tissues produce the organ's structure and contribute to its function.  | 49        |
| □ 9  | Recognise that organ systems cooperate and interact to deliver essential functions such as exchanging respiratory gases, circulating materials, obtaining nutrients, and removing waste products.   | 50-52     |
| □ 10 | <b>SHE:</b> Identify how scientists simplify complex biological systems into manageable components for study, noting that new properties emerge at each level of the biological hierarchy.  | 48        |
| □ 11 | <b>SHE:</b> Explain the importance of using animals in research as crucial for advancing scientific knowledge about multicellular organisms. Discuss the use of bioartificial tissue in the context of ethical treatment of animals as sentient beings, adhering to the three strategies of replacement, reduction, and refinement. | 53        |
| □ 12 | <b>SI:</b> Use photographs to compare epithelial, connective, muscle, and nervous tissues of the human body. Use photographs to view tissues from the respiratory, circulatory, excretory, digestive, and plant systems.  | 49        |

**Key Idea:** Stem cells are undifferentiated cells found in multicellular organisms. They are characterised by the properties of self renewal and potency.

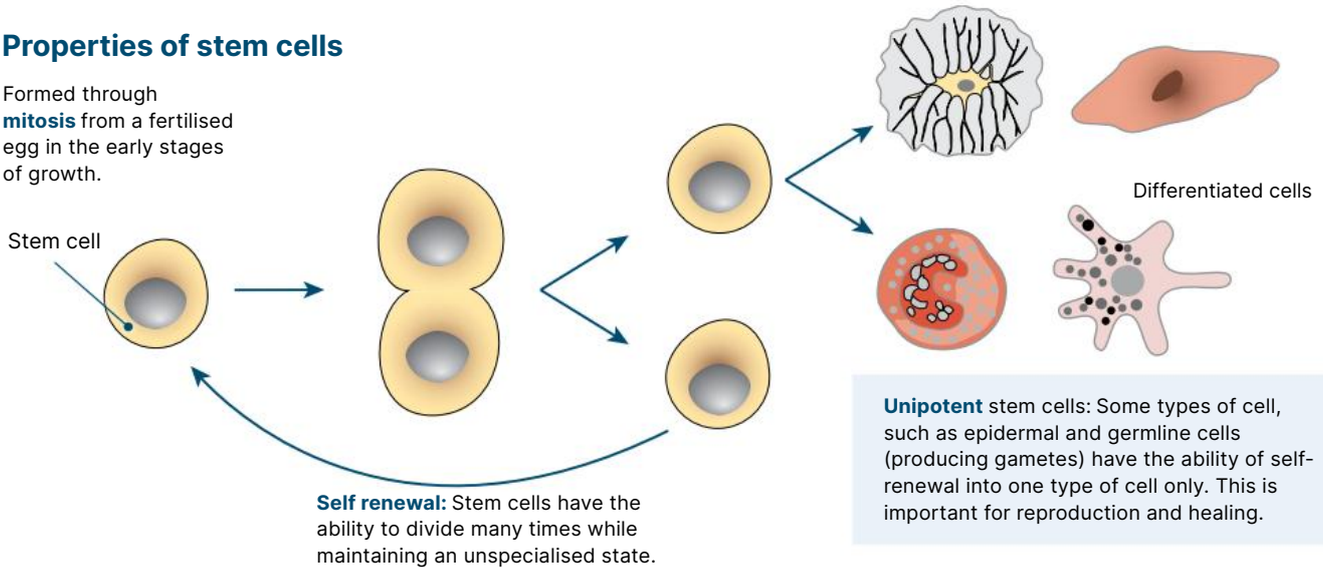
A **zygote** can differentiate into all the **cell** types of the body because its early divisions produce **stem cells**. Stem cells contain the entire genome and can divide repeatedly while

remaining unspecialised. They give rise to the many cell types that make up the **tissues** of a multicellular organism, i.e. the stem cells in bone marrow specialise to produce all the cell types that make up blood. These **multipotent** (or adult) stem cells are found in most organs, where they replace old or damaged cells and replenish cells throughout life.

## Properties of stem cells

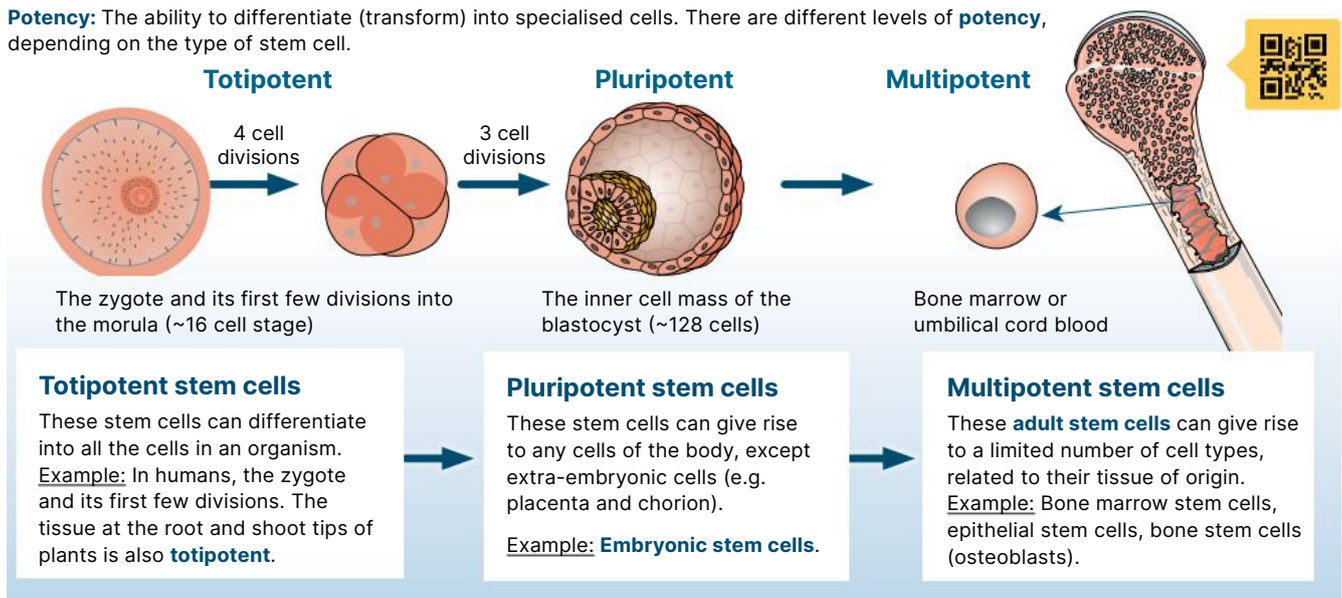
Formed through **mitosis** from a fertilised egg in the early stages of growth.

Stem cell



## Types of stem cells

**Potency:** The ability to differentiate (transform) into specialised cells. There are different levels of **potency**, depending on the type of stem cell.



1. Describe the two defining features of stem cells:

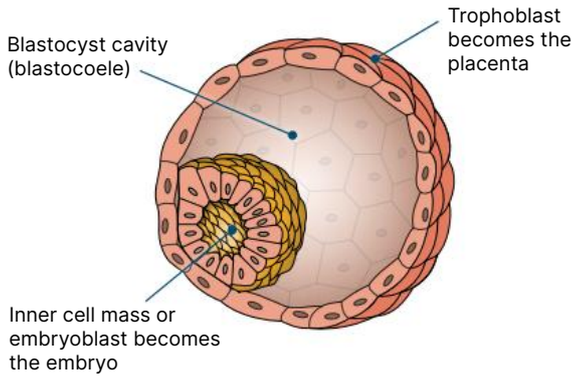
- (a) \_\_\_\_\_
- (b) \_\_\_\_\_

2. Describe the potency of stem cells and where they are found:

- (a) Totipotency: \_\_\_\_\_
- \_\_\_\_\_
- (b) Pluripotency: \_\_\_\_\_
- \_\_\_\_\_
- (c) Multipotency: \_\_\_\_\_
- \_\_\_\_\_



### Embryonic stem cells



- ▶ **Embryonic stem cells (ESC)** are derived from the inner cell mass of blastocysts (above). Blastocysts are 5 day old embryos consisting of a hollow ball of 50-150 cells.
- ▶ Cells derived from the inner cell mass are pluripotent. They can become any cells of the body, with the exception of placental cells.
- ▶ When cultured without any stimulation to differentiate, ESC retain their potency through multiple cell divisions. This means they have great potential for therapeutic use in regenerative medicine and tissue replacement.
- ▶ However, the use of ESC involves the deliberate creation and destruction of embryos and is therefore ethically unacceptable to many people.

### Adult stem cells



- ▶ **Adult stem cells (ASC)** are undifferentiated cells found in several types of tissues (e.g. brain, bone marrow, fat, and liver) in adults, children, and umbilical cord blood.
- ▶ Unlike ESCs, they are multipotent and can only differentiate into a limited number of cell types, usually related to the tissue of origin.
- ▶ There are fewer ethical issues associated with using ASC for therapeutic purposes, because no embryos are destroyed. For this reason, ASC are already widely used to treat a number of diseases including leukaemia and other blood disorders.
- ▶ In a recent development scientists have been able to genetically modify cells, through gene transfer, into 'reprogrammed' pluripotent cells (iPSCs). The disease-causing genes are removed and replaced with healthy genes. These iPSCs can then be differentiated into a wide variety of cells for implantation back into the patient.

3. Distinguish between embryonic stem cells and adult stem cells with respect to their potency:

---



---



---



---

4. Suggest how stem cells could be potentially useful for treating diseased or damaged organs:

---



---



---



---

5. Suggest some ways that technology has utilised the concepts of pluripotent cells in medicine:

---



---



---



---

6. Research has produced pluripotent stem cells from differentiated adult cells called induced pluripotent stem cells (iPSC). Suggest why using these cells in medicine generates fewer ethical issues than using other stem cell types:

---



---



---



---

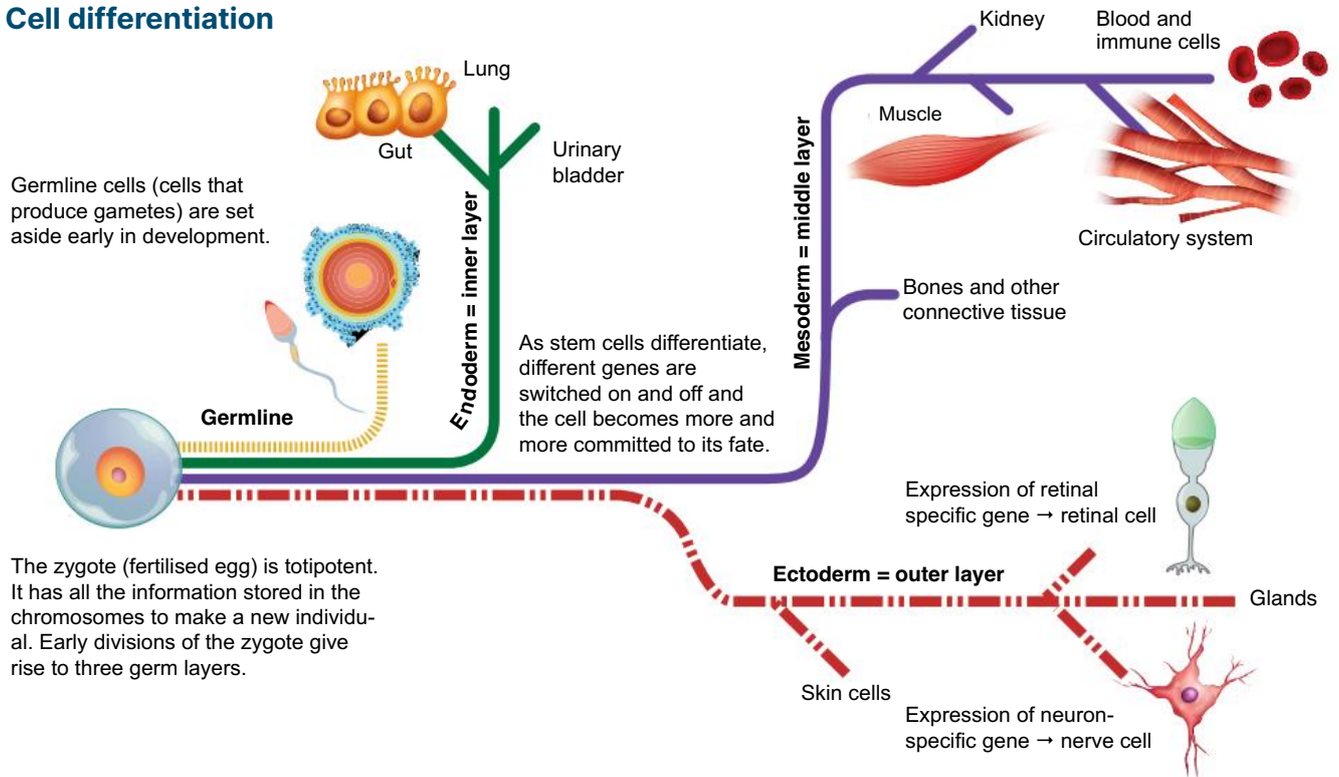
# Cellular Differentiation

**Key Idea:** A zygote divides by mitosis and produces all the cell types in the body by cellular differentiation. Specific patterns of gene switching determine what cell type develops.

Multicellular organisms consist of many different cell types, each specialised to carry out a particular role. A **zygote** and its first few divisions are **totipotent** and can differentiate to form any cell type in the body. During development, these cells divide and follow different developmental pathways, giving

rise to the three germ layers and the **specialised cells** that make up the **tissues** and **organs** of the body. This process by which more specialised cells develop from more generalised ones is called **cellular differentiation**. It is achieved through switching genes on and off in particular sequences. As a cell proceeds along its developmental pathway, its 'choices' become more limited. Once differentiated, it cannot (under normal circumstances) turn into another cell type.

## Cell differentiation



Germline cells (cells that produce gametes) are set aside early in development.

The zygote (fertilised egg) is totipotent. It has all the information stored in the chromosomes to make a new individual. Early divisions of the zygote give rise to three germ layers.

1. Multicellular organisms consist of many different cell types. Explain the role genes play to allow it to be possible for these cells all to arise from a single fertilised egg (zygote):

---



---



---



---

2. The zygote produces cells that differentiate in three cell lineages (germ layers). What types of cells or tissues do each of these lineages produce?

(a) Endoderm: \_\_\_\_\_

\_\_\_\_\_

(b) Mesoderm: \_\_\_\_\_

\_\_\_\_\_

(c) Ectoderm: \_\_\_\_\_

\_\_\_\_\_

3. Why can't a blood cell turn into a nerve cell? \_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_

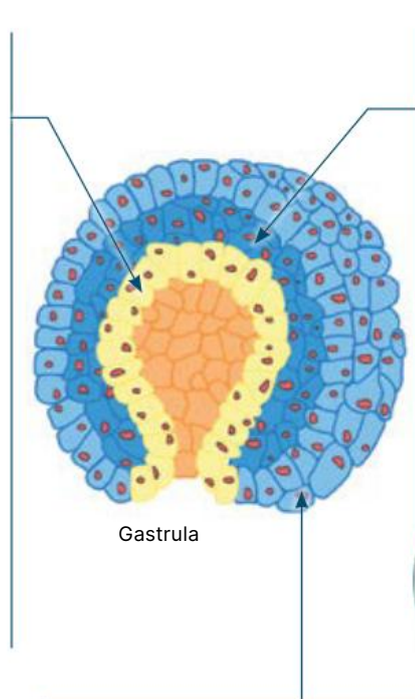
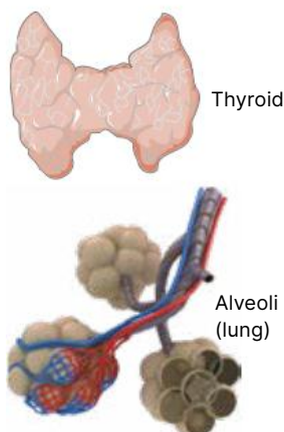


## Cell layers

Three distinct cell layers (germ layers) are produced in an early embryonic phase of development called the gastrula. These three cell layers (endoderm, mesoderm, and ectoderm) are the precursors of all adult cells and tissues. At this stage, the cells are now **multipotent** and considered **adult stem cells**. Some examples of cells and tissues formed from each layer are described below.

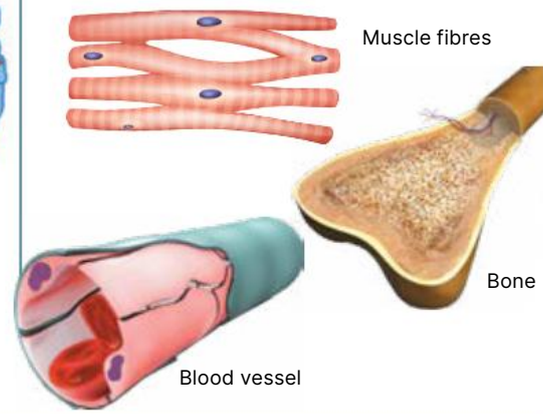
### The endoderm ●

- ▶ The endoderm is the innermost germ layer.
- ▶ In the early embryo, the endoderm forms the embryonic gut.
- ▶ It differentiates to form the **digestive system**, glands, and part of the **respiratory system**.



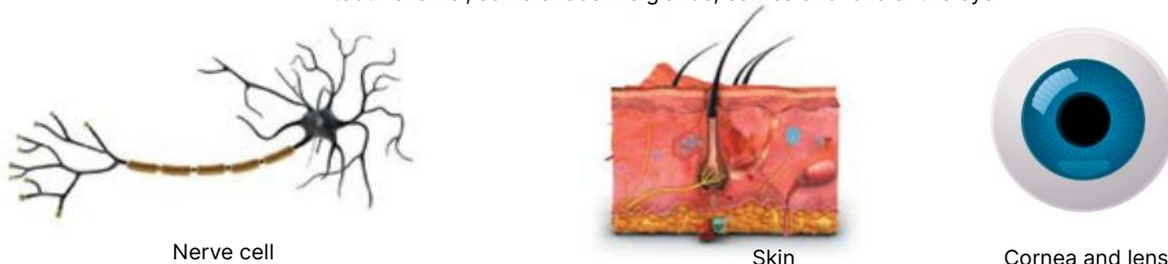
### The mesoderm ●

- ▶ The mesoderm is the middle germ layer, between the ectoderm and endoderm.
- ▶ The mesoderm differentiates to give rise to the muscles, **circulatory system** (heart and blood vessels), urinogenital system, dermis (inner skin layer), skeleton, and other supportive and **connective tissue**.



### The ectoderm ●

- ▶ The ectoderm is the outermost germ layer.
- ▶ In the fully developed embryo and adult human the ectoderm forms the brain and the **nervous system**, epidermis of skin (including hair, sweat glands and nails), tooth enamel, some endocrine glands, cornea and lens of the eye.



4. Evaluate how the process of cellular differentiation from each of the three germ layers (endoderm, mesoderm, and ectoderm) contributes to the complexity and diversity of cell types in multicellular organisms?

---



---



---



---



---

5. How does multipotency during the gastrula stage contribute to the variety of cell types in adult organisms?

---



---



---



---



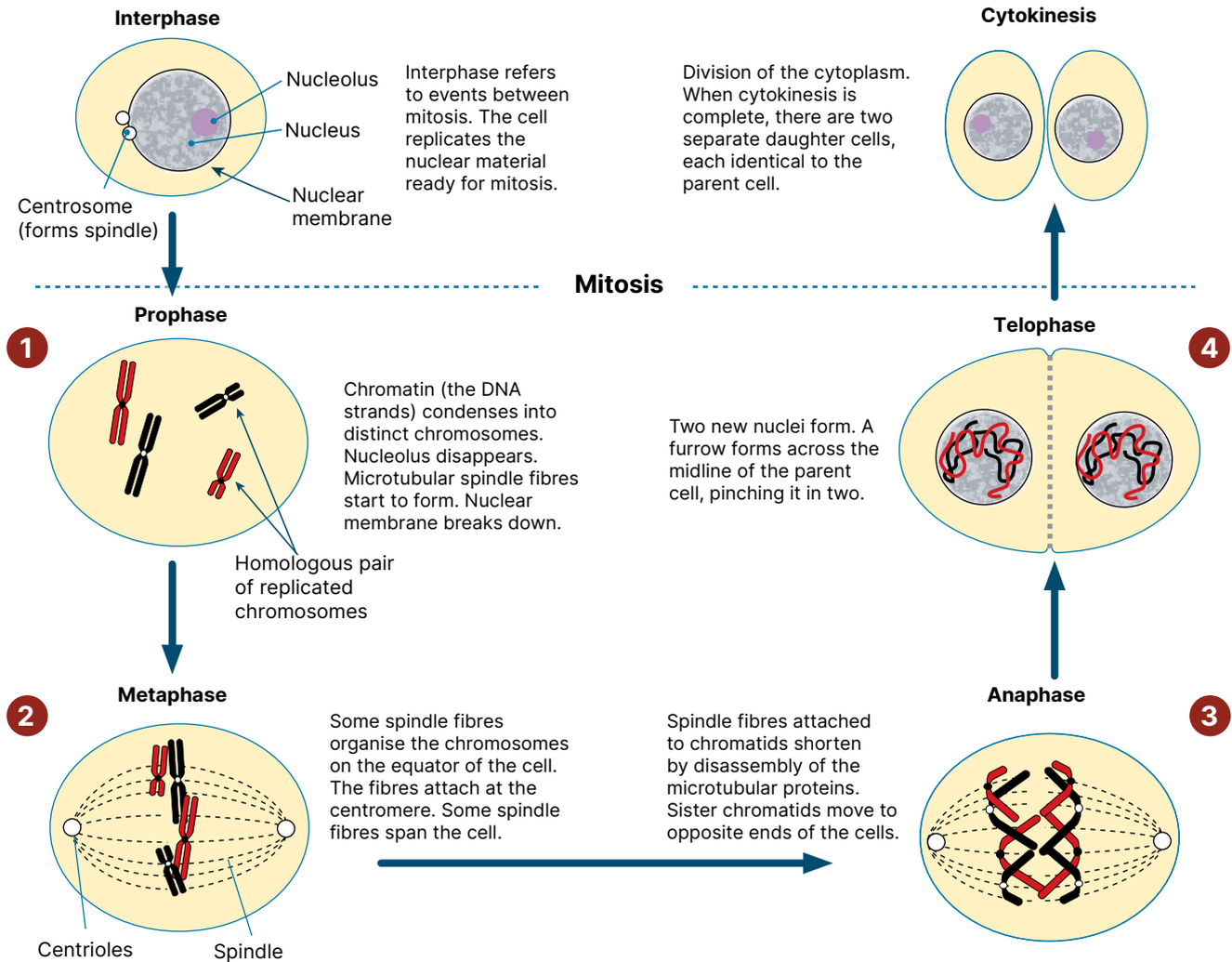
---

**Key Idea:** Somatic (body) cells divide by mitosis. Differentiation occurs as genes are turned on and off.

Mitosis refers to the separation (division) of the nuclear material and it precedes cell division. Its main purpose is the growth and repair of the body. There is no change of chromosome number and the daughter cells are genetically identical to the parent cell. Although mitosis is part of a

continuous cell cycle, it is divided into stages (prophase, metaphase, anaphase, and telophase) to help distinguish the processes involved. Cellular differentiation occurs when different parts of the DNA (genes) are expressed during the development of the cell. Thus, although the cells are genetically identical, they can be morphologically and functionally different.

## The cell cycle and stages of mitosis in an animal cell



An important difference between the cells in a growing embryo and the cells in an adult is that most of an adult's cells are differentiated. Only a few tissues, such as bone marrow, contain stem cells. Stem cells undergo an asymmetric cell division to generate particular cell types. One of the daughter cells has a finite capacity for cell division and begins to differentiate, while the other will remain as an undifferentiated stem cell, in order to generate more new cells.

- Define mitosis: \_\_\_\_\_
- What must occur before mitosis takes place? \_\_\_\_\_
- (a) What is the purpose of the spindle fibres? \_\_\_\_\_
- (b) Where do the spindle fibres originate? \_\_\_\_\_
- What is the difference in the purpose of mitosis in an embryo as opposed to an adult? \_\_\_\_\_

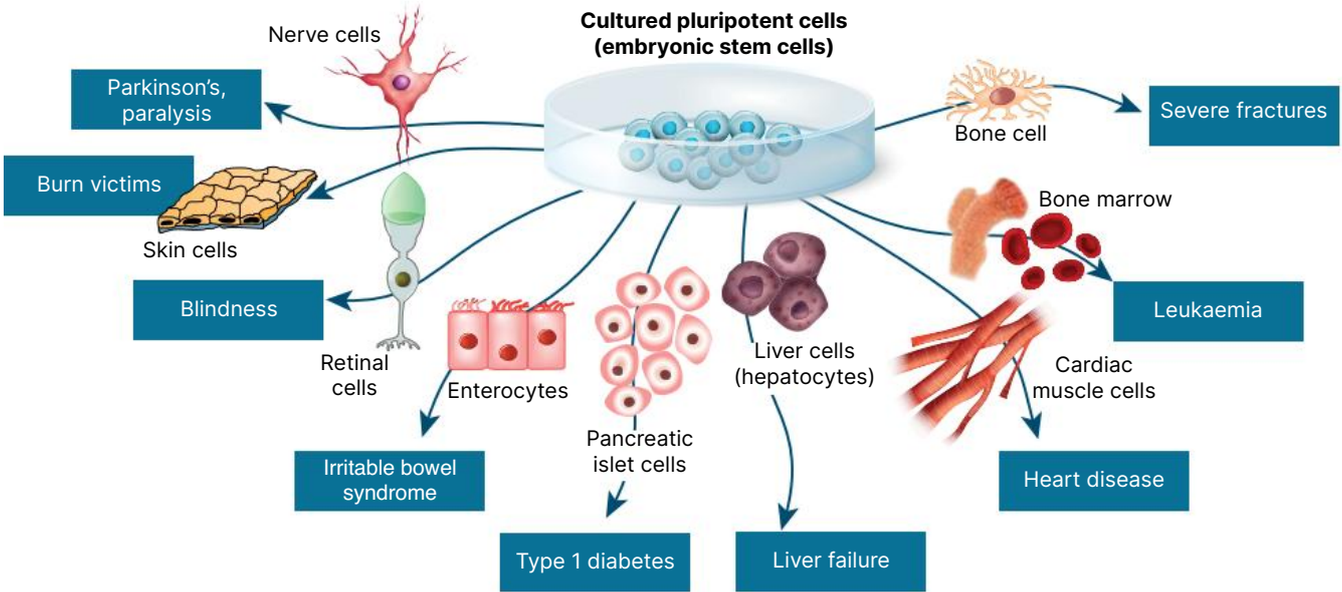
46

# Applications of Stem Cells

**Key Idea:** Stem cells have many potential medical applications, but technical difficulties must be overcome first.

**Stem cell** research is at an early stage and much is still to be learned about the environments required by cells to differentiate into specific cell types. The ability of **embryonic stem cells** (ESC) to differentiate into almost any cell type

means they have potential applications in replacing diseased or damaged cells (below). **Adult stem cells**, either from a donor or from the patient themselves, also have therapeutic uses. Donor stem cells must be matched for compatibility, while stem cells for autologous (self) transplants may require genetic correction before use (lower panels).

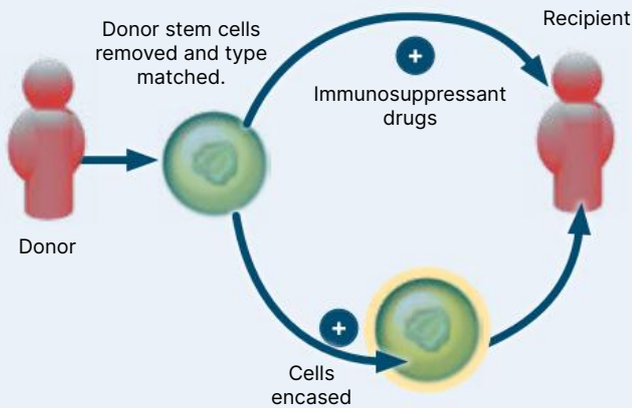


### Donor stem cells can be used to repair tissues

**Problem:** Immune system will attack the donor's cells.

**Solution:** Firstly, a donor with a **tissue** match is selected (the cell surface proteins on donor and recipient cells are the same or very similar). This reduces the risk of immune rejection. Secondly, the recipient will need to take immunosuppressant drugs to stop their immune system attacking the donated cells.

Another way to prevent immune rejection is to encase donor cells in a protective shell, isolating them from immune detection by the recipient. This is being investigated with respect to pancreatic cells and diabetes.

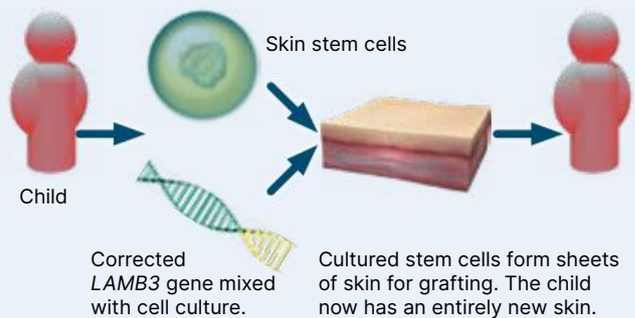


### A patient's cells can be corrected before use

**Problem:** Some diseases are the result of defective genes. Stem cells from the patient will carry these defective genes.

**Solution:** If the disease is due to a simple genetic fault, then the stem cells can be genetically corrected before use. Stem cells are isolated and cultured in the laboratory in the presence of the corrected gene. Corrected cells are identified and transplanted back into the patient, without immune rejection.

**Example:** In 2015, a young German child had a mutation in the **LAMB3** gene. His skin cells were not making the protein needed to hold cells together and his skin was falling off. His skin stem cells were genetically corrected and new skin was cultured and grafted back. More than a square metre of skin was grown and grafted on to the child.



1. Identify a problem with using stem cells from a donor to treat a recipient patient:

---



---



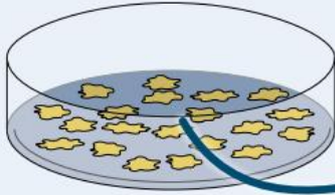
---



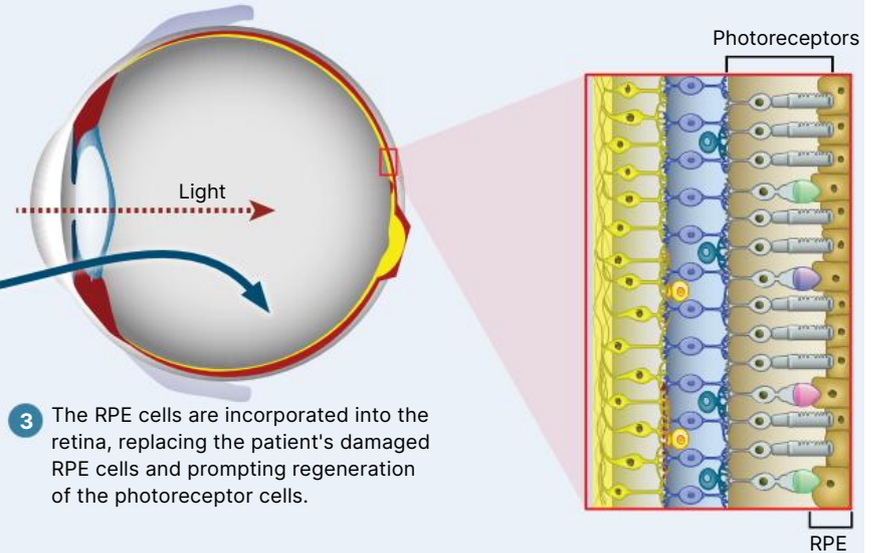
## Stem cells for Stargardt's disease

- ▶ Stargardt's disease is an inherited form of juvenile macular degeneration (a loss of the central visual field of the eye). The disease is associated with a number of different mutations and results in malfunction of the retinal pigment epithelium (RPE) cells, which normally nourish the retinal photoreceptor cells and protect the retina from excess light.
- ▶ Faulty RPE causes deterioration of the photoreceptor cells in the centre of the retina and progressive loss of central vision. This often begins between ages 6 and 12 and continues until a person is legally blind. Trials using stem cells have proved promising as a treatment, with impaired vision being corrected relatively quickly (within weeks).

- 1 Embryonic stem cells are cultured in the lab with proteins and vitamins so that they develop into retinal pigment epithelium (RPE) cells.



- 2 The RPE cells are injected just below the retina of the eye and above the choroid (the layer containing the blood vessels).



- 3 The RPE cells are incorporated into the retina, replacing the patient's damaged RPE cells and prompting regeneration of the photoreceptor cells.

2. Umbilical cord blood is promoted as a rich source of multipotent stem cells for autologous (self) transplants. Can you see a problem with using a baby's cord blood to treat a disease in that child at a later date?

---



---



---



---

3. (a) Explain the basis for correcting Stargardt's disease using stem cell technology:

---



---

- (b) There have also been stem cell therapy trials using the patient's own cells (e.g. bone marrow) to treat Stargardt's disease. What advantages might there be in using a patient's own cells and what difficulties might be involved?

---



---



---



---

4. Describe a technical difficulty associated with stem cell therapies when:

- (a) The stem cells come from a donor: \_\_\_\_\_

---



---

- (b) The stem cells used are ESC: \_\_\_\_\_

---



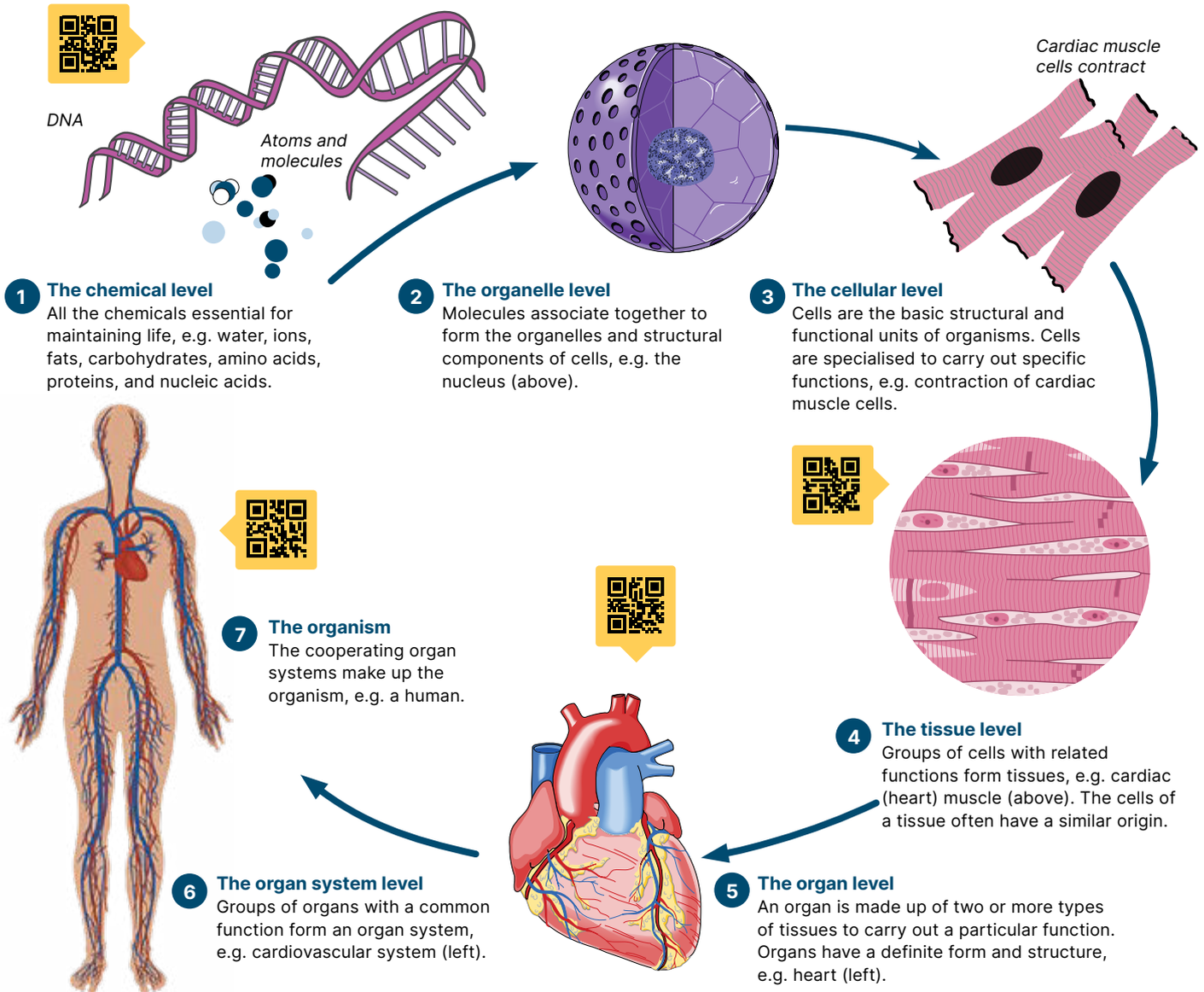
---



# 48 The Hierarchy of Life

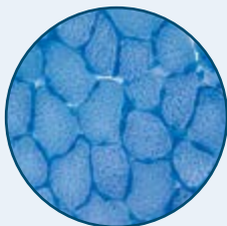
**Key Idea:** Structural organisation in multicellular organisms is hierarchical, with new properties arising at each level. Multicellular organisms are organised according to a hierarchy of structural levels, where each level builds on the one below it. At each level, new properties, absent at the simpler level, emerge. **Hierarchical organisation** allows **specialised cells**

to group together into **tissues** and **organs** to perform a specific function or set of related functions. This improves efficiency in the organism. Organisation and the emergence of new properties in complex systems are two of the defining features of living organisms. The diagrams following explain this hierarchical organisation for a human and a plant.



## Specialised cells make up tissues and organs

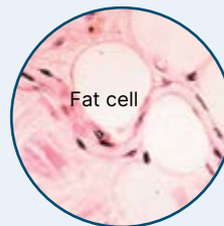
Specialised cells often have modifications or exaggerations to a normal cell feature to help them perform a particular task. They may have more (or fewer) of a particular organelle in order to perform their role most efficiently.



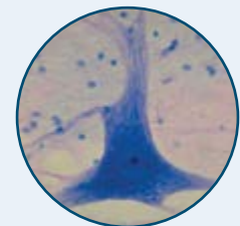
Muscle cells can contract (shorten) to bring about the movement of limbs and organs. Cardiac muscle cells have many mitochondria to supply their high energy needs.



Different types of blood cells have specific tasks. Red blood cells have no nucleus and contain molecules of haemoglobin for efficient oxygen transport.



Thin, flat epithelial cells line the walls of blood vessels (arrow). Large fat cells store lipid. Lipids act as energy storage, provide insulation and act as cushioning to protect organs.



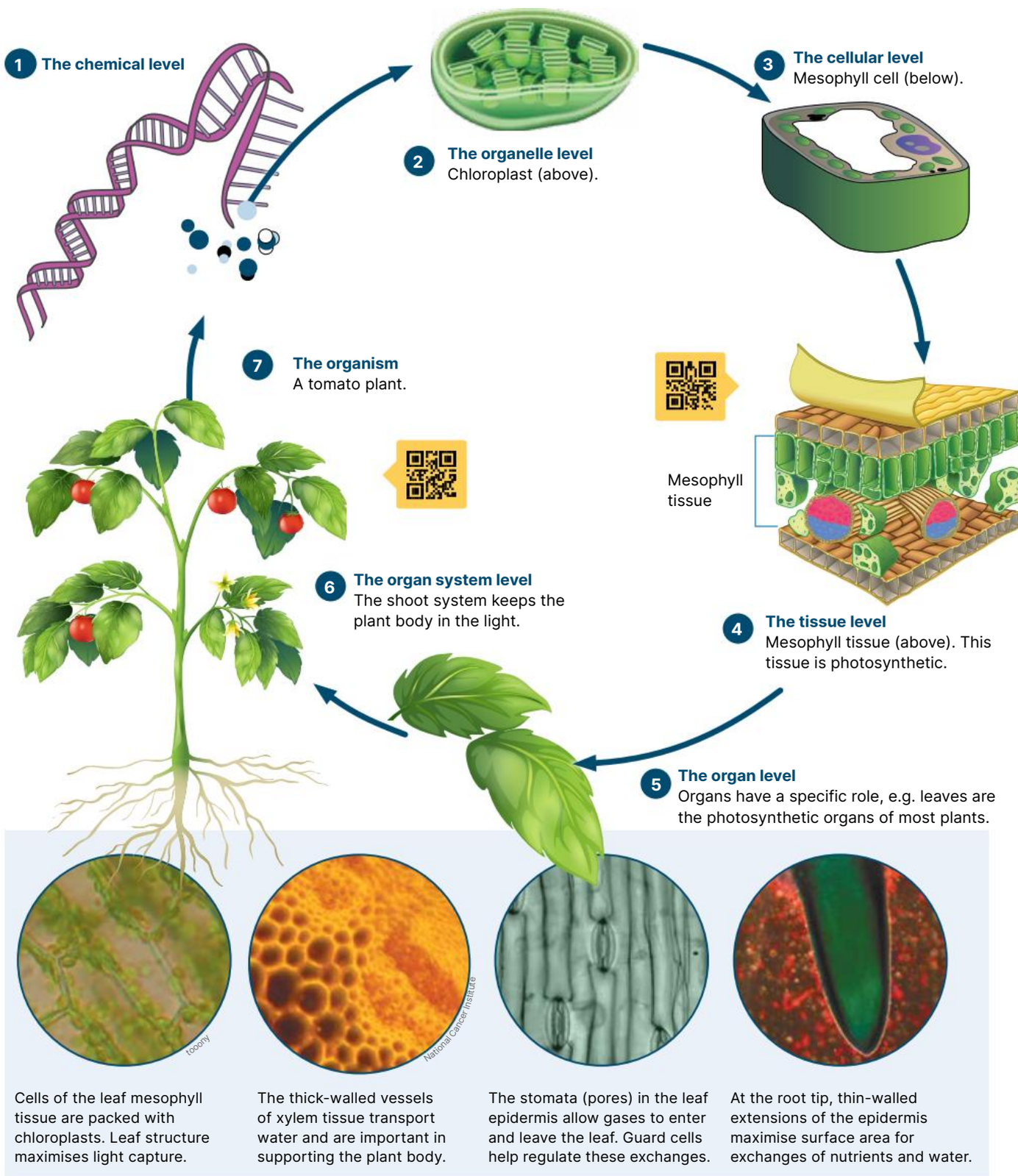
Nerve cells conduct impulses around the body enabling responses to the environment. Specialised structures known as axons and dendrites allow this to happen.



SU

SHE





1. Assign each of the following emergent properties to the level at which it first appears:

- (a) Metabolism: \_\_\_\_\_
- (b) Behaviour: \_\_\_\_\_
- (c) Replication: \_\_\_\_\_
- (d) Internal transport: \_\_\_\_\_
- (e) Surface protection: \_\_\_\_\_
- (f) Nutrient processing: \_\_\_\_\_

2. Explain how a hierarchical structure enables greater efficiency of function in the whole organism:

---



---



---

# Exploring Tissues and Organs

**Key Idea:** Tissues come together to make organs. You can see this in an isolated organ, such as a heart.

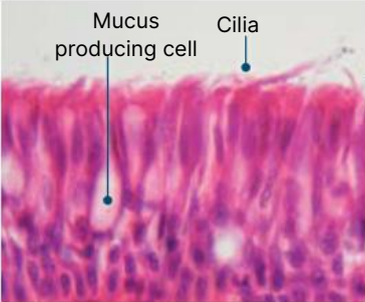
A **tissue** is a collection of related **cell** types that work together to carry out a specific function. Different tissues

come together to form **organs**. The cells, tissues, and organs

of the body interact to meet the needs of the entire organism. You can explore the different tissues that make up an organ by examining an isolated organ, such as a sheep's heart.

Muscle tissue	Epithelial tissue	Nervous tissue	Connective tissue
<ul style="list-style-type: none"> <li>▶ Contractile tissue.</li> <li>▶ Produces movement of the body or its parts.</li> <li>▶ Includes smooth, skeletal, and cardiac muscle.</li> </ul>	<ul style="list-style-type: none"> <li>▶ Lining tissue.</li> <li>▶ Covers the body and lines internal surfaces.</li> <li>▶ Can be modified to perform specific roles.</li> </ul>	<ul style="list-style-type: none"> <li>▶ Receives and responds to stimuli.</li> <li>▶ Makes up the structures of the nervous system.</li> <li>▶ Regulates function of other tissues.</li> </ul>	<ul style="list-style-type: none"> <li>▶ Supports, protects, and binds other tissues.</li> <li>▶ Contains cells in an extracellular matrix.</li> <li>▶ Can be hard or fluid.</li> </ul>

1. Research the type of tissue(s) that occur at the places in the body indicated below. State the types of tissues that occur in the spaces provided. Codes: E = epithelium, CT = connective tissue, SM = smooth muscle.

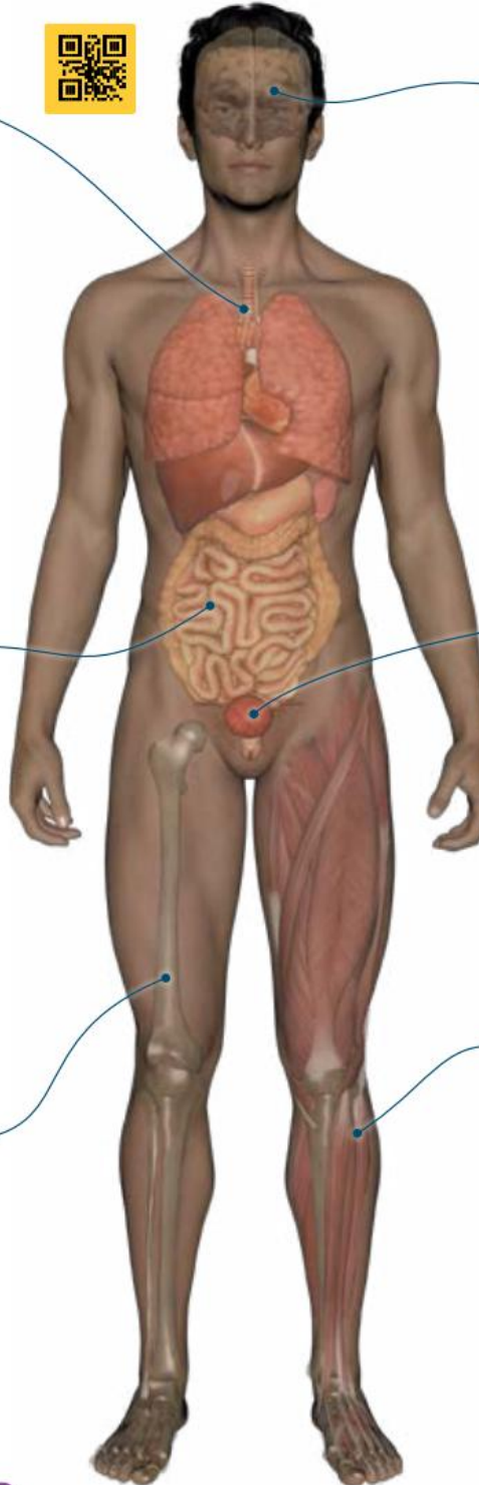


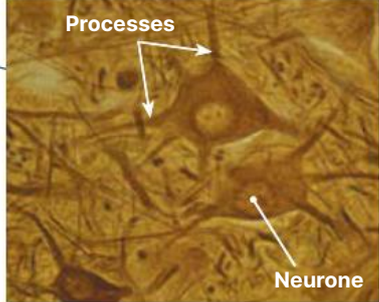
Upper respiratory tract:

---



---



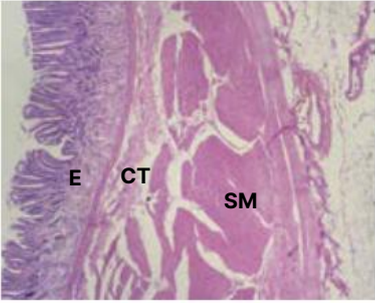


Brain:

---



---

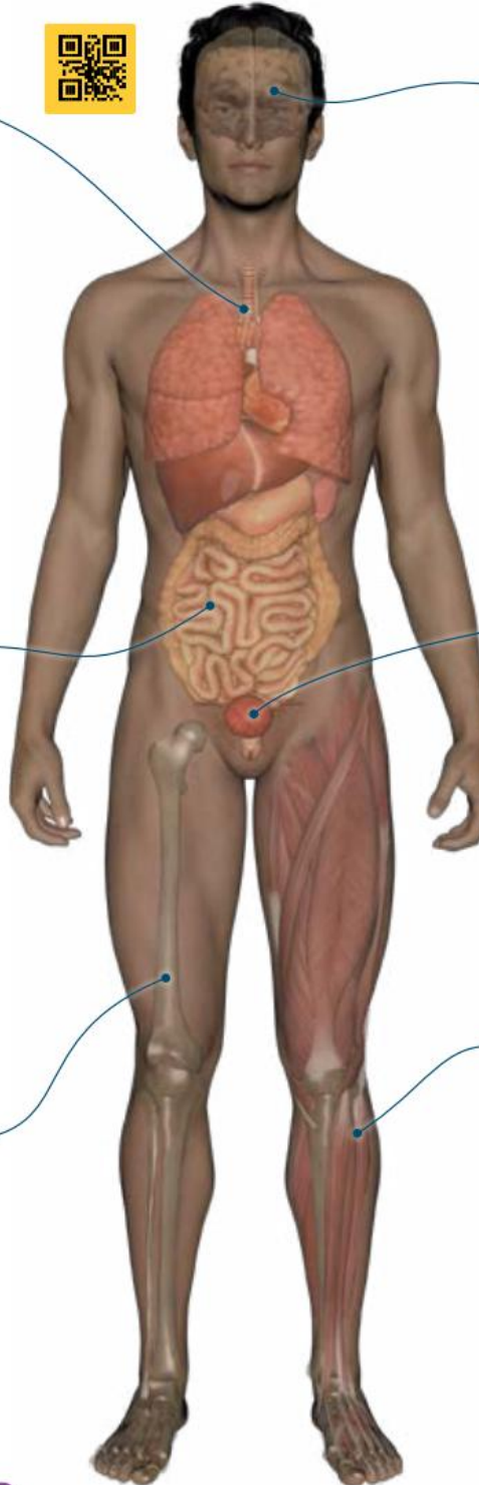


Digestive tract:

---



---



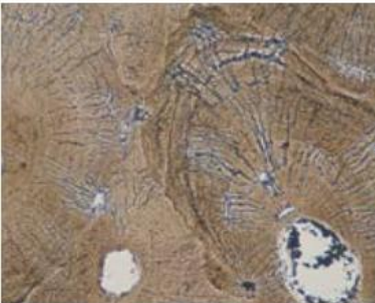


Bladder:

---



---

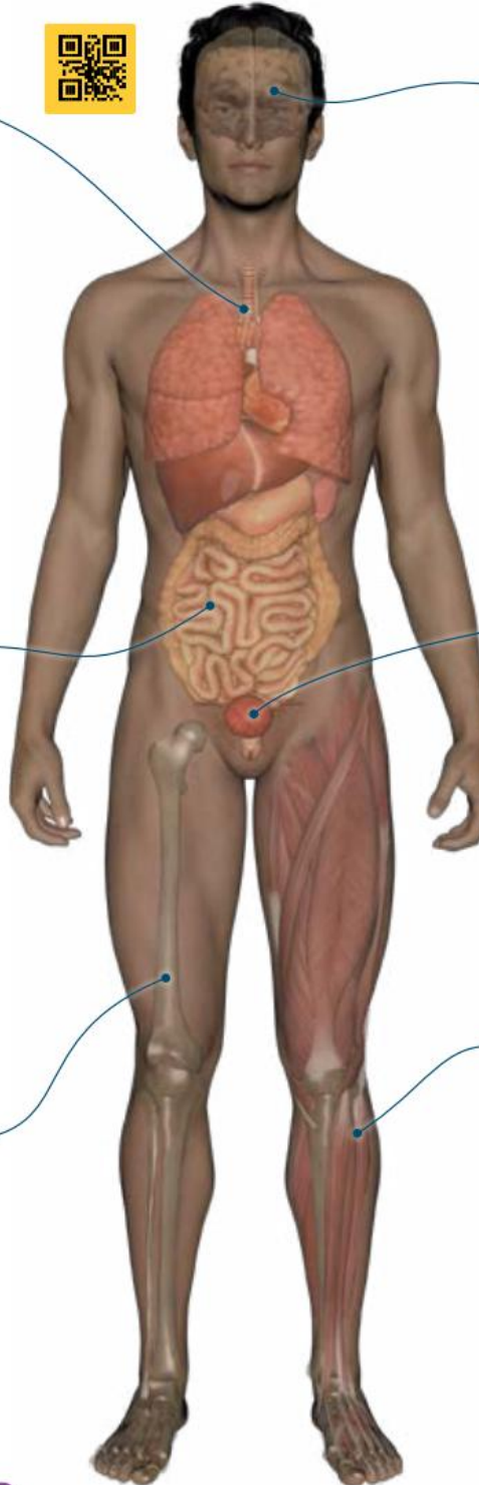


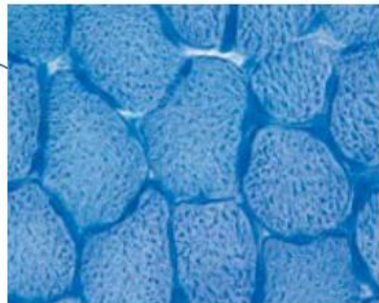
Bone:

---



---





Skeletal muscle:

---



---



**Key Idea:** The circulatory and gas exchange systems interact to provide the tissues with oxygen and remove carbon dioxide.

### Circulatory system

#### Function

Delivers oxygen ( $O_2$ ) and nutrients to all **cells** and **tissues**. Removes carbon dioxide ( $CO_2$ ) and other waste products of metabolism.  $CO_2$  is transported to the lungs.

#### Components

- ▶ Heart
- ▶ Blood vessels:
  - Arteries
  - Veins
  - Capillaries
- ▶ Blood

### Interaction between systems

In mammals, the **gas exchange** (or respiratory) system and cardiovascular system interact to supply oxygen and remove carbon dioxide from the body.

### Gas exchange system

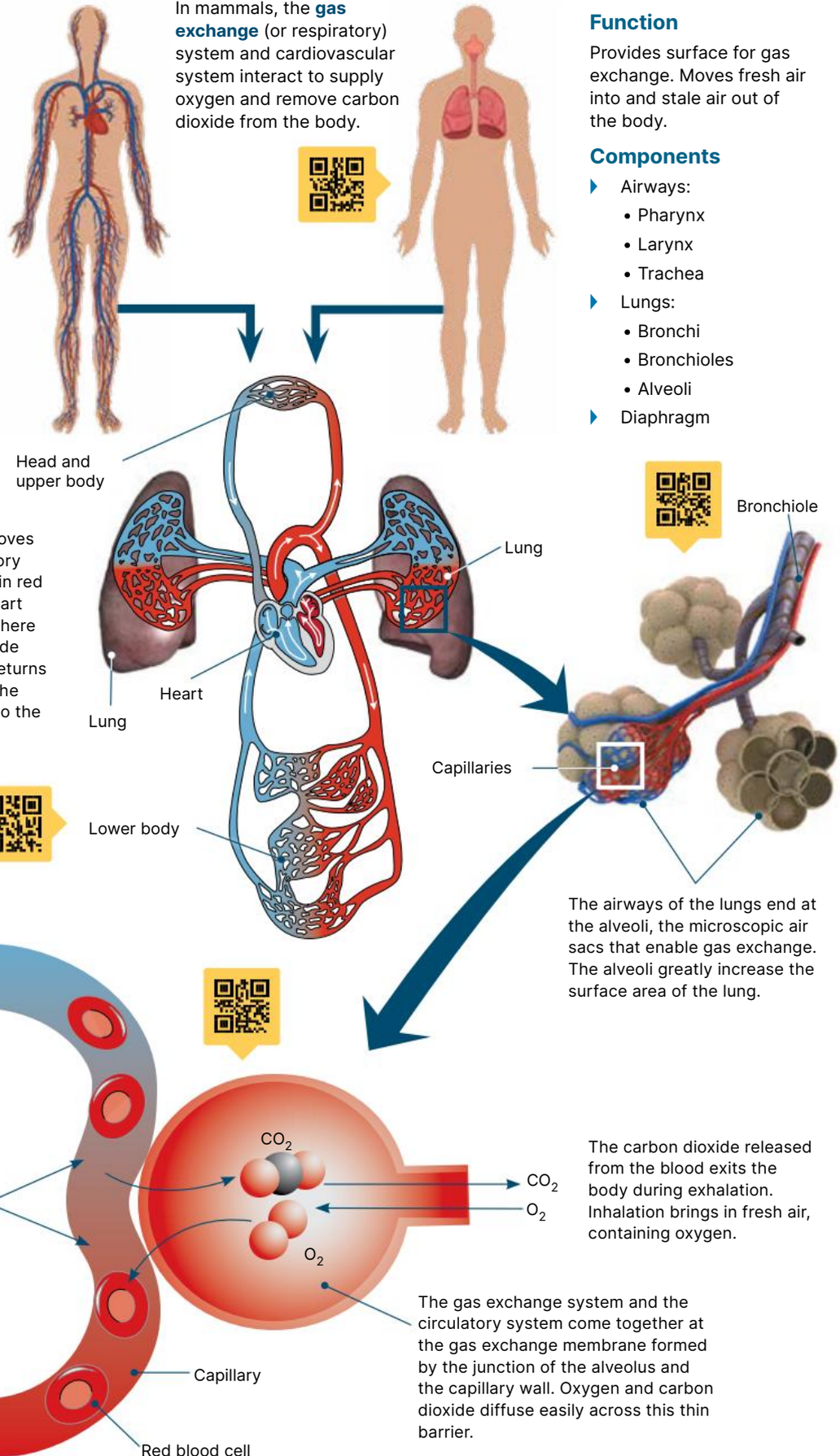
#### Function

Provides surface for gas exchange. Moves fresh air into and stale air out of the body.

#### Components

- ▶ Airways:
  - Pharynx
  - Larynx
  - Trachea
- ▶ Lungs:
  - Bronchi
  - Bronchioles
  - Alveoli
- ▶ Diaphragm

Oxygen ( $O_2$ ) from inhaled air moves from the lungs into the circulatory system and is transported within red blood cells to the heart. The heart pumps the blood to the body where  $O_2$  is released and carbon dioxide ( $CO_2$ ) is picked up. The blood returns to the heart and is pumped to the lungs where  $CO_2$  is released into the lungs to be breathed out.



The airways of the lungs end at the alveoli, the microscopic air sacs that enable gas exchange. The alveoli greatly increase the surface area of the lung.

The carbon dioxide released from the blood exits the body during exhalation. Inhalation brings in fresh air, containing oxygen.

The gas exchange system and the circulatory system come together at the gas exchange membrane formed by the junction of the alveolus and the capillary wall. Oxygen and carbon dioxide diffuse easily across this thin barrier.

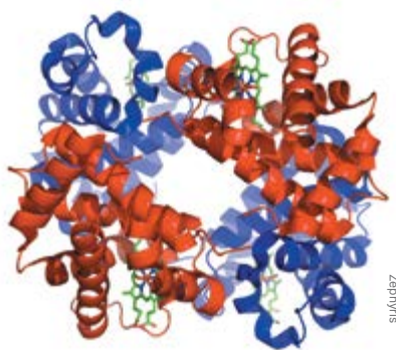


SU

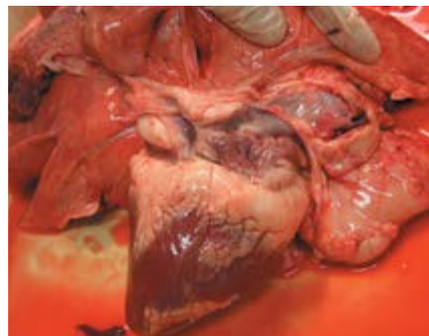




The response to exercise shows the close link between the circulatory and gas exchange systems. During exercise, breathing rate increases to provide more oxygen, which is carried by the blood to supply respiration (ATP generation) in working muscles. Heart rate increases to increase the rate at which oxygen is delivered to the tissues and carbon dioxide is returned to the lungs.



Oxygen is transported in red blood cells by the protein haemoglobin (above). In the capillaries of the lungs (high oxygen), haemoglobin binds oxygen tightly. In the tissues, higher carbon dioxide levels cause haemoglobin to release its oxygen.  $\text{CO}_2$  is carried in the blood as bicarbonate ( $\text{CO}_2 + \text{H}_2\text{O} \rightarrow \text{H}_2\text{CO}_3 \rightarrow \text{H}^+ + \text{HCO}_3^-$ ). In the lungs, this dissociates back into  $\text{CO}_2$  and water.



As with all **organ** systems, the circulatory and gas exchange systems are interdependent. Organs in the circulatory system (e.g. the heart) need oxygen to keep working and this is supplied by the lungs. If the heart were to stop beating, it and all other organs would quickly run out of oxygen. Similarly, if breathing were to stop, all organs would quickly run out of oxygen.

1. (a) What happens to the rate of blood flow during exercise? \_\_\_\_\_

(b) What happens to the breathing rate during exercise? \_\_\_\_\_

(c) How do the circulatory and gas exchange systems interact to accommodate the extra oxygen requirements of an exercising person?

---



---



---

2. Lung diseases affect rates of gas exchange in the lung. Suggest how this would affect the body:

---



---



---



---

3. (a) At which point in the body do the respiratory and circulatory systems directly interact?

---



---

(b) Explain what is happening at this point: \_\_\_\_\_

---



---



---

4. In your own words, describe how the circulatory system and respiratory system work together to provide the body with oxygen and remove carbon dioxide:

---



---



---



---



---



---

# 51 Circulation and Digestive Interaction

**Key Idea:** The circulatory and digestive systems interact to provide the body's tissues with nutrients.

## Circulatory system

### Function

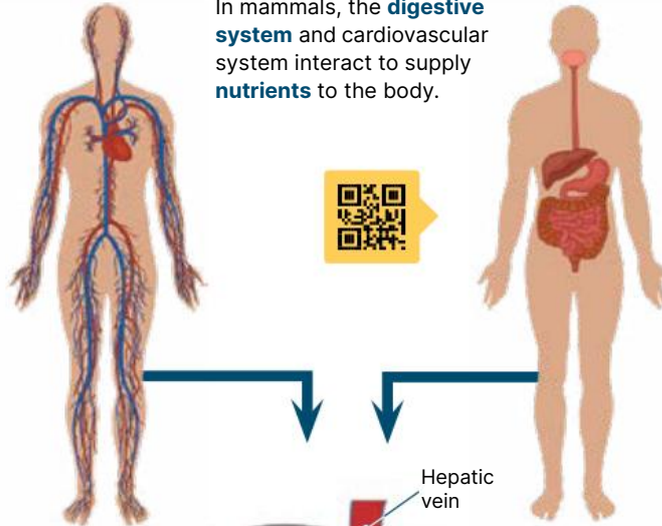
Delivers oxygen (O<sub>2</sub>) and nutrients to all **cells** and **tissues**. Removes carbon dioxide (CO<sub>2</sub>) and other waste products of metabolism.

### Components

- ▶ Heart
- ▶ Blood vessels:
  - Arteries
  - Veins
  - Capillaries
- ▶ Blood

## Interaction between systems

In mammals, the **digestive system** and cardiovascular system interact to supply **nutrients** to the body.



## Digestive system

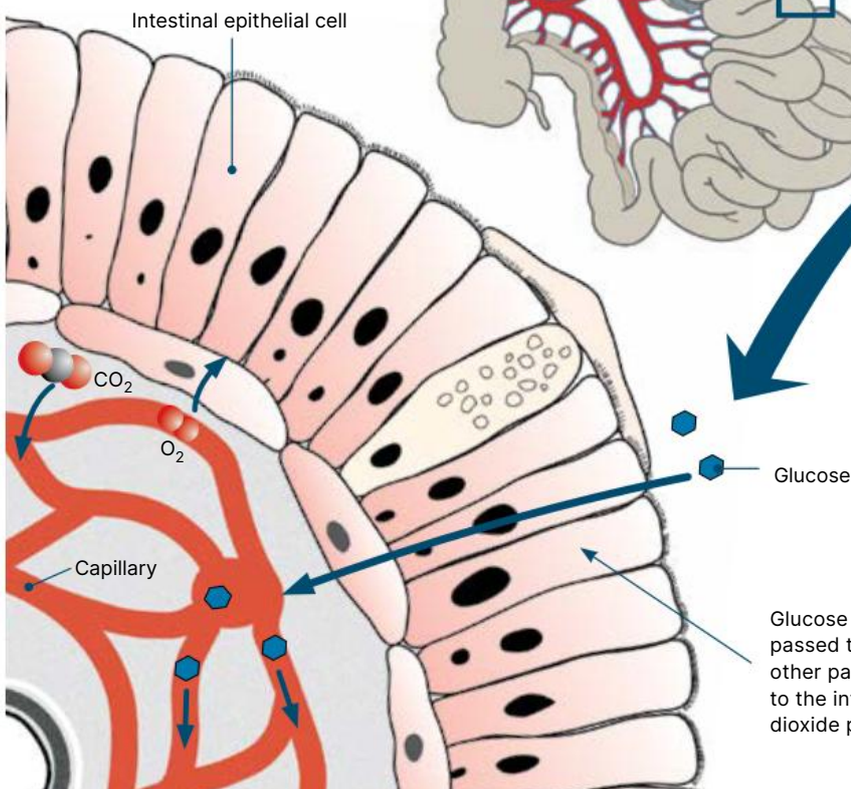
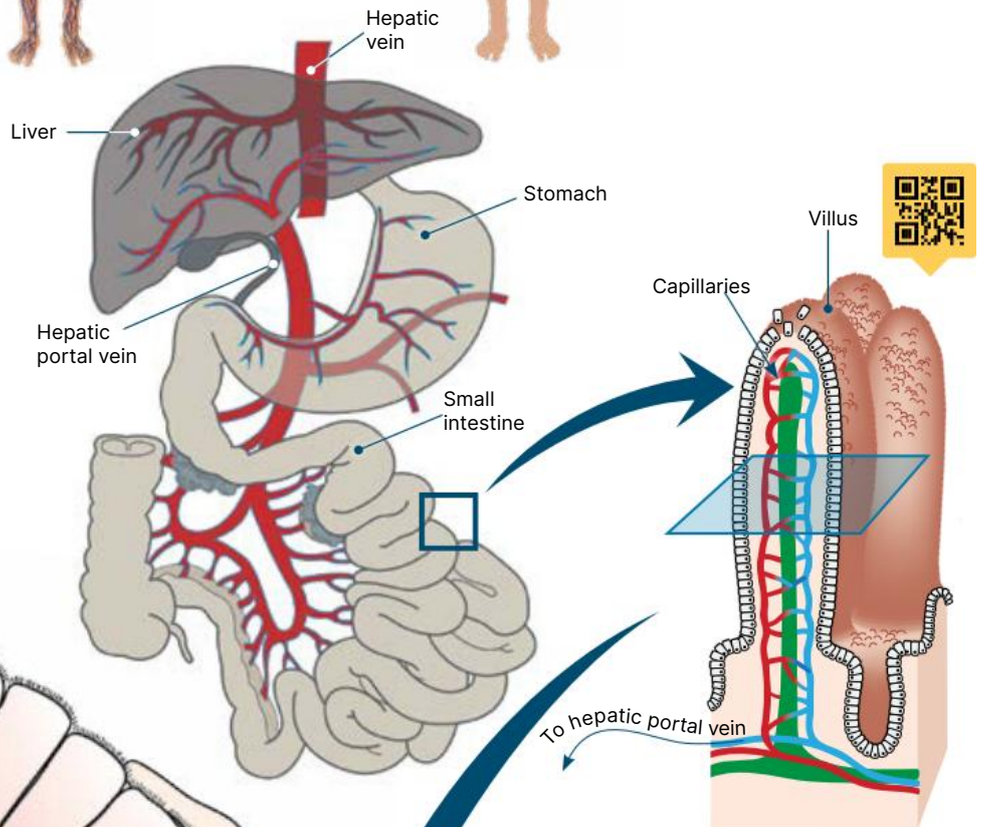
### Function

Digest food and absorb useful molecules from it, and eliminate undigested material.

### Components

- ▶ Mouth and pharynx
- ▶ Oesophagus
- ▶ Stomach
- ▶ Liver and gall bladder (accessory organs)
- ▶ Pancreas (accessory organ)
- ▶ Small intestine
- ▶ Large intestine

Food is digested in the stomach and small intestine and the nutrients are absorbed and passed to the circulatory system. The capillaries around the stomach and intestines collect nutrients and then drain to the hepatic portal vein, which carries the blood directly to the liver. The liver then processes this nutrient-rich blood, e.g. glucose is stored as glycogen. The hepatic vein then transports nutrients from the liver to supply the other tissues of the body.



Villi project into the lumen of the small intestine and absorb nutrients. Villi contain capillary networks which receive the nutrients and transport them to the hepatic portal system. The villi structure greatly increases the surface area of the intestine.

Glucose and other nutrient molecules are passed to the blood and transported to other parts of the body. Oxygen passes to the intestinal cells, while carbon dioxide passes into the blood.





Blood flow to the digestive tract increases steadily after a meal and remain elevated for about 2.5 hours, reaching a maximum after about 30 minutes. During exercise, blood flow in the digestive tract is reduced as it is redirected to the muscles.

Nutrients, e.g. minerals, sugars, and amino acids, are transported in the blood plasma to the liver. The liver receives nutrient-rich deoxygenated blood from the digestive system via the hepatic portal vein and oxygen rich blood from the hepatic artery.

Scarring of the liver tissue, or cirrhosis, can result in portal hypertension (high blood pressure). The scarred tissue obstructs blood flow in the liver. This causes pressure to build up in upstream blood vessels, resulting in swelling and possible haemorrhage.

1. How are nutrients transported in the blood? \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_
2. Explain how a liver cirrhosis affects the circulatory system: \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_
3. (a) At which two points in the body do the digestive and circulatory systems directly interact?  
 \_\_\_\_\_  
 \_\_\_\_\_
- (b) Explain what is happening at these points: \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_
4. (a) What happens to blood flow to the digestive tract after a meal? \_\_\_\_\_  
 \_\_\_\_\_
- (b) Explain why it is often recommended that a person should exercise within 2.5 hours of eating, or eat within half an hour of exercising to gain a most benefit from the exercise (in terms of muscle development):  
 \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_
5. In your own words, describe how the circulatory and digestive systems work together to provide the body with nutrients:  
 \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_

**Key Idea:** The circulatory and urinary systems interact to remove wastes from the body's tissues and helps maintain blood volume and pressure.

### Circulatory system

#### Function

Delivers oxygen ( $O_2$ ) and nutrients to all **cells** and **tissues**. Removes carbon dioxide ( $CO_2$ ) and other waste products of metabolism.

#### Components

- ▶ Heart
- ▶ Blood vessels:
  - Arteries
  - Veins
  - Capillaries
- ▶ Blood

### Interaction between systems

In mammals, the urinary system and cardiovascular system interact to remove metabolic **wastes** from the body.

### Urinary system

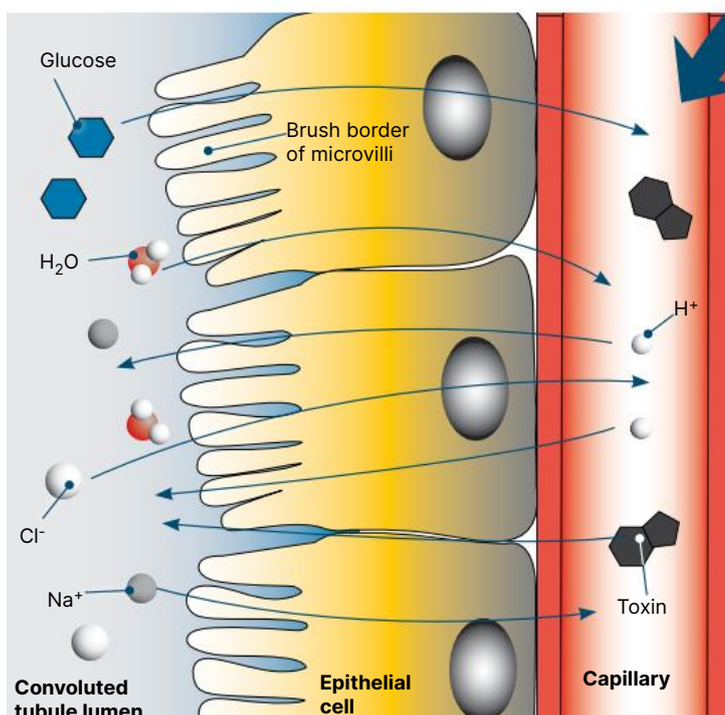
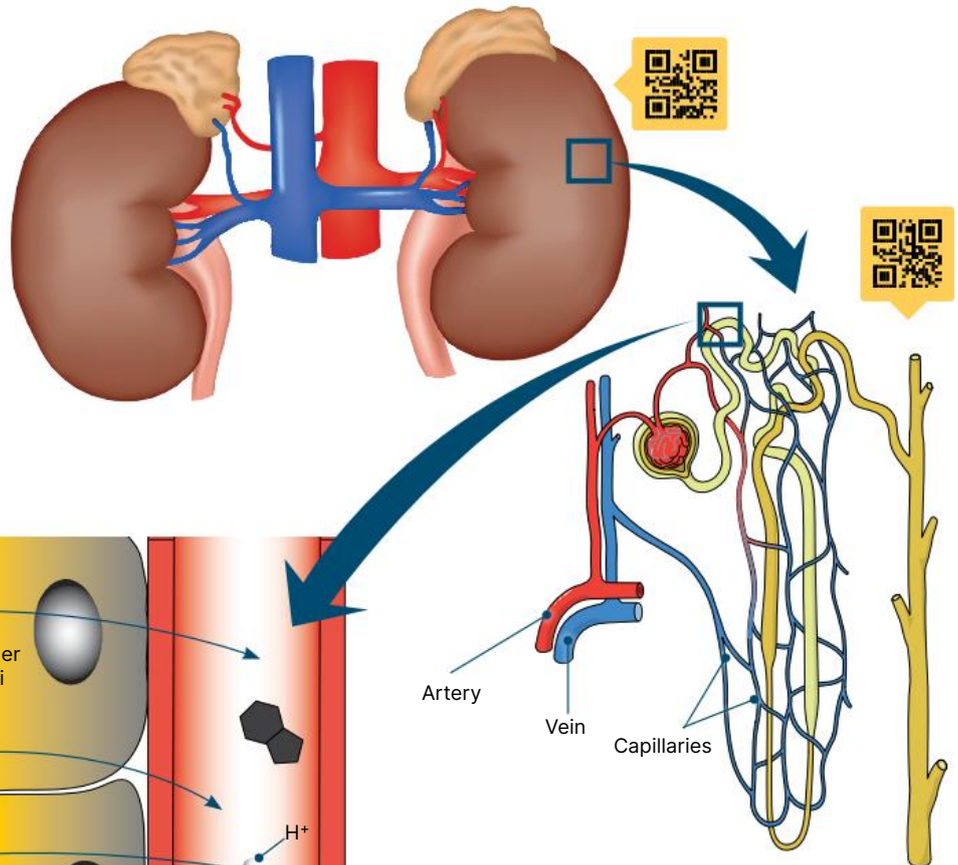
#### Function

Filters blood, retaining useful molecules and removing harmful ones. Regulates blood volume and ion content.

#### Components

- ▶ Kidneys
- ▶ Ureters
- ▶ Bladder
- ▶ Urethra

Metabolic wastes generated in the cells move into the blood plasma and are transported to the kidneys. Blood is forced at high pressure through the capillaries of the glomerulus, producing a filtrate, which is collected in the surrounding Bowman's capsule. Both useful and harmful molecules are contained in the filtrate although large proteins and blood cells are excluded.



The convoluted tubules through which the filtrate travels are surrounded by capillaries (above). As the filtrate moves along the tubules, epithelial cells reabsorb useful molecules or ions, e.g. glucose and sodium ions, back into the blood in the capillaries (left). Unwanted ions (e.g.  $H^+$ ) and toxins are also secreted into the filtrate by active transport and eliminated in the urine. The microvilli brush border of the epithelial cells allows for a greater surface area over which its activities can take place. These epithelial cells have many mitochondria to supply the active transport energy requirements.





Sodium chloride crystals

The kidneys regulate blood volume by regulating the ion content of the blood. Retention of ions help retain water (via osmosis). Various hormones carried in the blood stimulate the kidneys to increase water retention.



The regulation of blood volume is important for the body. Optimal blood volume enables the blood to flow at the correct rate through the capillaries, helps maintain the correct concentrations of electrolytes, and maintains the pressure needed for glomerular filtration.



Polycystic kidney disease

Chronic kidney disease makes it difficult for the body to regulate blood volume and this can promote hypertension. Hypertension can lead to atherosclerosis of the arteries and in turn heart disease, heart attack, or peripheral arterial disease.

1. (a) Explain why regulating salt content of the blood in turn regulates blood volume: \_\_\_\_\_

---



---

- (b) How would the active secretion of hydrogen ions into the filtrate (urine) help to regulate blood pH?

---



---

2. (a) Explain why kidney disease can make regulating blood volume and pressure difficult: \_\_\_\_\_

---



---

- (b) Explain why kidney disease can also lead to heart disease and heart attacks: \_\_\_\_\_

---



---

3. Use the graph right to answer the following questions:

- (a) What is the effect of reducing glomerular filtration rate on the risk of cardiovascular disease?

---



---

- (b) How many times greater is the risk of cardiovascular disease at a GFR of <15 compared to a GFR of >60?

---



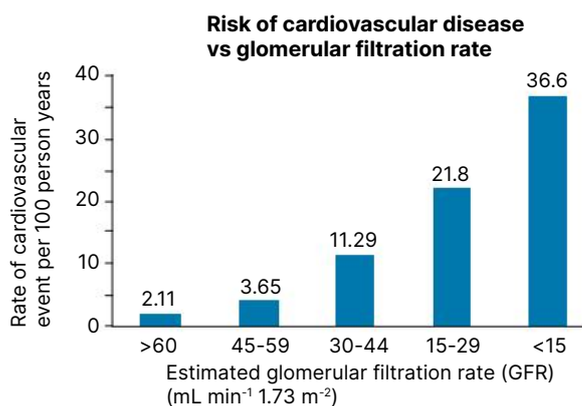
---

- (c) What might cause reduced GFR?

---



---



4. In your own words, describe how the circulatory system and urinary system work together to remove wastes and regulate the volume, pressure, and composition of the blood:

---



---



---



---



---

**Key Idea:** Using animals in scientific and medical research helps to understand and test new theories and medicines, but must be done in a way that minimises animal discomfort. The use of animals in scientific research has played a vital role in many advances in science and medicine. Animals (often specially bred rats or mice) are used as models of how a human system might work and are the basis for preliminary research that might later be applied to humans.

This includes developing theories and processes around stem cells and developing and using **bioartificial organs**. During this research, there is the chance that some of these animals may experience harm. How much depends greatly on the way they are treated and the research involved. Many countries have strict guidelines as to when and how animals can be used in research. These are often based around the 3Rs principle of replacement, reduction, and refinement.

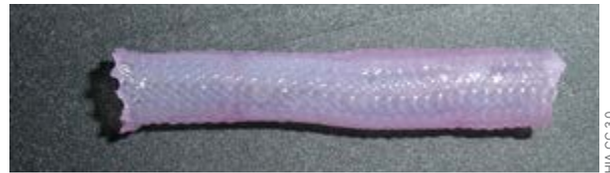
### The 3Rs of ethical research

The 3Rs are guiding principles for the use of animals in research, testing, and teaching. Animals should only be used when there are no alternatives, any harm must be measured against the benefits, and those harms must be minimised.

- ▶ **Replacement:** Can animals be replaced in the research? In many cases computer modelling can produce highly accurate results and can easily be used to identify unforeseen problems and refine tests. This reduces the need for animal testing, so that it occurs in the latter stages of the research (if at all).
- ▶ **Reduction:** Only the minimum number of animals needed to produce reliable or statistically significant data should be used.
- ▶ **Refinement:** Investigations should be reviewed to identify where the number of animals used could be reduced. This information can then be applied to refining subsequent investigations.

### Stem cells, bioartificial organs and ethics

- ▶ Bioartificial organs are grown using a scaffold implanted with **stem cells**. These are cultured to grow and differentiate around the scaffold to produce a functional three-dimensional **organ** (or part of an organ). Currently, only simple organs or sections of organs can be grown but research is constantly producing new developments (e.g. new materials and techniques).
- ▶ **Ethical** issues can arise from this kind of research. Often animals are used to practice and develop techniques and this can cause them considerable distress (see below).

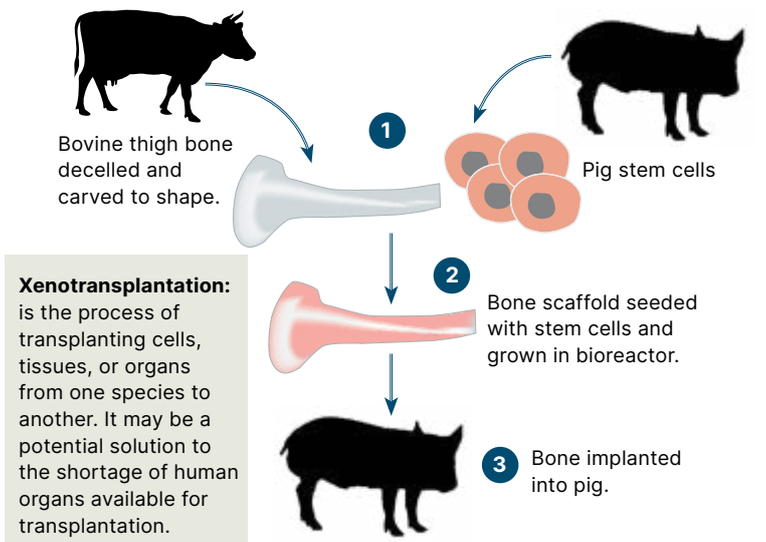


Engineered vascular tissue (blood vessel). The scaffold can be seen supporting the outer cells.

### Using bioartificial tissue to replace a jaw

Bones can be replaced using titanium but this means a loss of bone marrow, which plays an important role in the body. In 2016, researchers began trials on growing new bone for transplant.

- ▶ The experiment was carried out on Yucatan miniature pigs because the jaw structure is similar to humans. The researchers took part of a cattle thigh bone and removed all the **cells** and **tissue**, leaving just the bone structure. They then carved the bone into the shape of part of the pig mandible.
- ▶ This bone scaffold was seeded with pig stem cells and placed into a bioreactor. The cells developed into bone cells and grew into and around the scaffold (this took about 3 weeks). The matching part of the pig mandible was removed and the bioartificial jawbone implanted.
- ▶ Six months later the implants had fully grafted and the pigs were able to use their jaw normally.



1. As a group discuss why the experiment on pigs jaws above was carried out and its pros and cons. Discuss likely ethical issues in relation to the pigs and if you think the experiment was worth carrying out. Summarise your discussion below:




---

---

---

---

---

---

---

---

---

---

# Did You Get It?

1. Test your vocabulary by matching each term to its correct definition, as identified by writing the letter in the correct box.


- A** A type of cell that possesses the qualities of self renewal and potency.
- B** Ability to divide many times while maintaining an undifferentiated state.
- C** Able to give rise to any cells of the body, except extra-embryonic cells.
- D** The initial cell formed from the union of two gametes.
- E** Able to give rise a limited number of cell types, related to their tissue of origin.
- F** Ability to differentiate into specialised cell types.

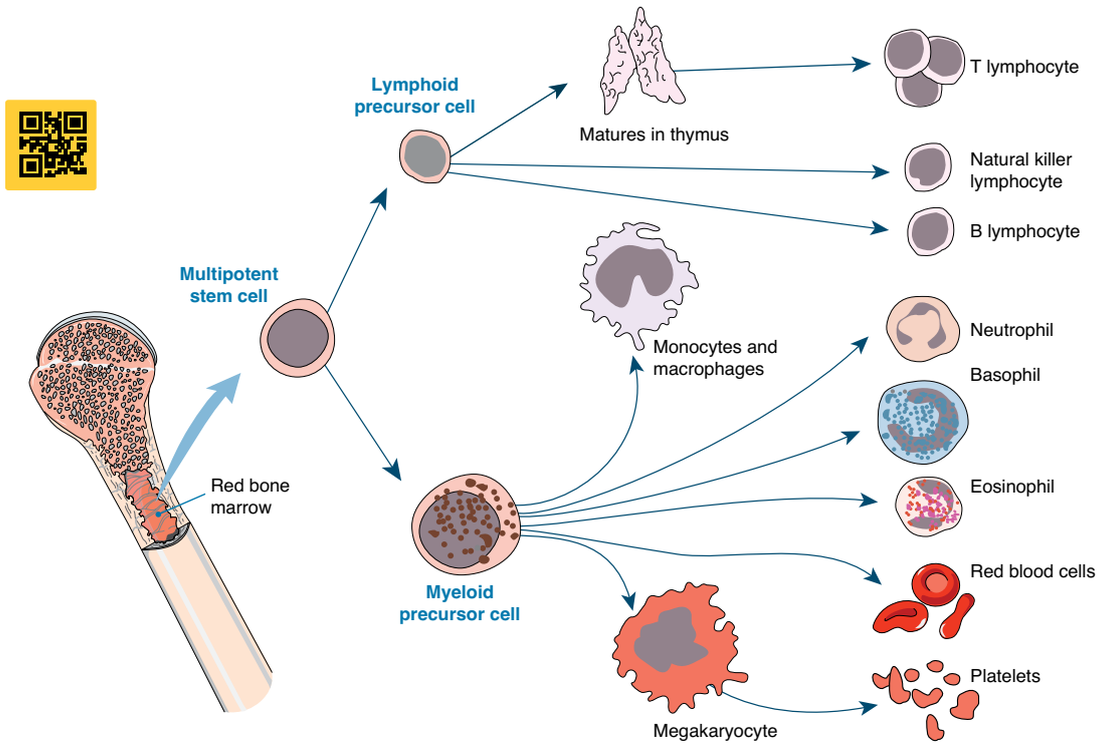
2. Give an example of each of the following stem cell types:

(a) Multipotent: \_\_\_\_\_

(b) Pluripotent: \_\_\_\_\_

(c) Totipotent: \_\_\_\_\_

3. Study the diagram of cellular differentiation below and answer the questions following as true or false:



(a) T lymphocytes can differentiate from a haematopoietic multipotent stem cell: \_\_\_\_\_

(b) Lymphoid precursor cells can produce red blood cells: \_\_\_\_\_

(c) Neutrophils mature from myeloid precursor cells: \_\_\_\_\_

4. (a) What is the link between cells, tissues, and organs? \_\_\_\_\_

\_\_\_\_\_

(b) What is the advantage of the hierarchical organisation of cells, tissues, organs, and systems? \_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_

# Cell Membrane



## Key Terms

- active transport
- carrier protein
- channel protein
- cholesterol
- concentration gradient
- diffusion
- endocytosis
- exocytosis
- facilitated diffusion
- fluid-mosaic model
- glycoprotein
- hydrophilic
- hydrophobic
- hypertonic
- hypotonic
- ion
- ion pump
- isotonic
- osmolarity
- osmosis
- partially permeable (= selectively-permeable)
- passive transport
- phospholipid
- plasma membrane
- polar
- plasmolysis
- protein channel:
- surface area: volume ratio
- turgor

## Key Concepts

- ▶ The cellular/plasma membrane has a fluid bilayer structure.
- ▶ Transport across cellular/plasma membranes can be passive or active.
- ▶ Proteins in or on the cellular membrane assist in the transport of specific molecules through it.

## The structure and role of the cell membrane

**Activity  
Number**

<input type="checkbox"/>	1	Describe the fluid mosaic model of membrane structure. Describe the significance of the phospholipid bilayer and cholesterol molecules.	55-56
<input type="checkbox"/>	2	Explain the general role of proteins in the plasma membrane, including protein channels, carrier proteins, and glycoproteins.	57
<input type="checkbox"/>	3	Construct or use a model to show the selectively permeable nature of the plasma membrane.	60
<input type="checkbox"/>	4	Describe the ongoing research to refine the fluid mosaic model, including the structure of membrane proteins, e.g. channel proteins.	59
<input type="checkbox"/>	5	<b>SI:</b> Describe the evidence for Singer and Nicolson's fluid mosaic model of membrane structure and explain how their understanding was assisted by advancements in technology.	58

## Passive transport across cell membranes

<input type="checkbox"/>	6	Describe the passive movement of molecules across membranes by diffusion and facilitated diffusion. Identify and explain factors regulating diffusion rates across membranes.	61
<input type="checkbox"/>	7	Investigate the direction of movement of materials across cell membranes.	61
<input type="checkbox"/>	8	Explain the importance of surface area to volume ratio in limiting cell size. Describe the effect of decreasing SA:V ratio on the efficient diffusion of materials to the interior of cells.	62
<input type="checkbox"/>	9	Explore a range of different sized mammalian cells to compare how size and shape relate to different functions.	63
<input type="checkbox"/>	10	Describe cellular adaptations for increasing cell surface area and diffusion surfaces.	65
<input type="checkbox"/>	11	Explain the movement of water across membranes by osmosis. Explain the terms hypotonic, isotonic, and hypertonic and the effects that solutions of different solute concentration can have on plant and animal cells.	66-68
<input type="checkbox"/>	12	Analyse secondary data investigating how the size and membrane solubility of molecules affects the rate of diffusion across the plasma membrane.	69
<input type="checkbox"/>	13	<b>SI:</b> Investigate the effect of surface area to volume ratio on diffusion in model cells.	64

## Active transport across cell membranes

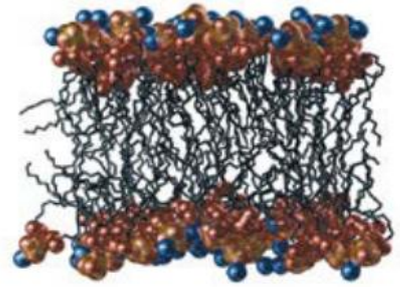
<input type="checkbox"/>	14	Distinguish between passive and active transport across cell membranes. Explain how the cell membrane maintains relatively stable internal conditions by active transport against a concentration gradient.	70, 73
<input type="checkbox"/>	15	Describe active transport across membrane using ion pumps, e.g. the sodium-potassium cotransporter (symport).	71
<input type="checkbox"/>	16	Distinguish between endocytosis and exocytosis, recognising both as active transport processes.	72

55

# The Plasma Membrane

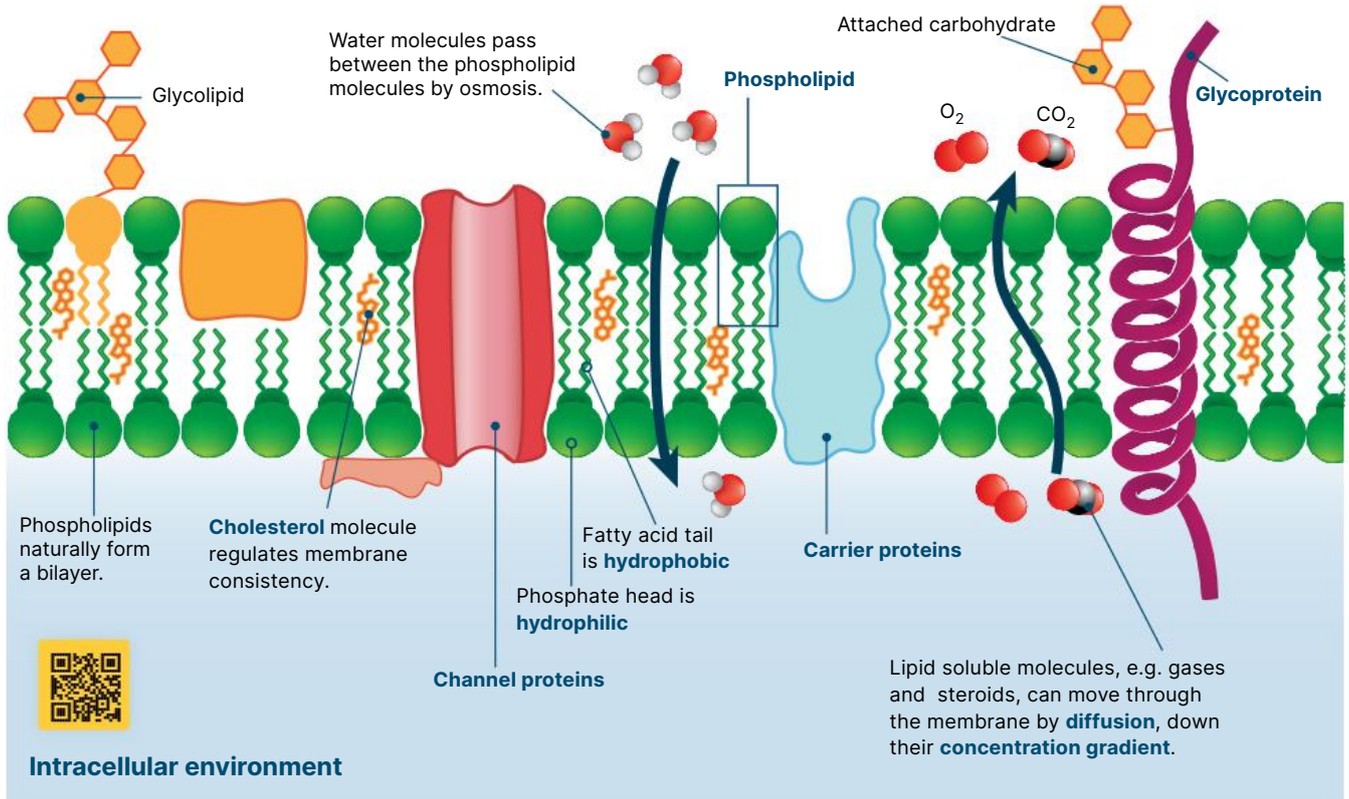
**Key Idea:** The plasma membrane is composed of a lipid bilayer with proteins moving freely within it. It is the partially permeable (also called semi-permeable or selectively permeable) boundary between the internal and external cell environments.

All cells have a **plasma membrane**, which forms the outer limit of the cell. A cell wall, if present, lies outside this and is quite distinct from it. Cellular membranes are also found inside eukaryotic cells as part of membranous organelles. The currently accepted model of the plasma membrane, called the **fluid-mosaic model** (below), describes a lipid bilayer with embedded proteins. This model was devised by Singer and Nicolson in 1972. The plasma membrane is a **partially permeable** barrier. It allows the passage of some molecules but not others. Many of the proteins embedded in the membrane are involved in the movement of molecules by transporting specific molecules (often large molecules or ions) across the membrane, often against their **concentration gradients**.



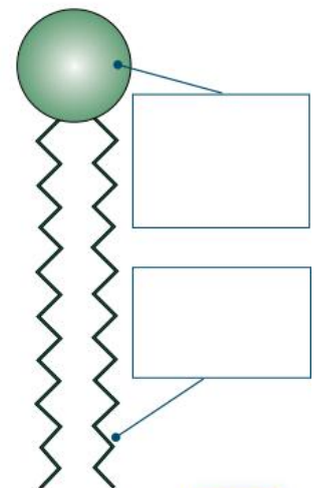
Molecular model showing how phospholipid molecules naturally orientate to form a bilayer.

## The fluid mosaic model of membrane structure



Based on a diagram in Biol. Sci. Review, Nov. 2009, pp. 20-21

- List the important components of the plasma membrane: \_\_\_\_\_  
\_\_\_\_\_
- Identify the kind of molecule on the diagram above that:
  - Can move through the plasma membrane by diffusion:  
\_\_\_\_\_
  - Forms a channel through the membrane: \_\_\_\_\_
- On the diagram (right) label the hydrophobic and hydrophilic ends of the phospholipid and indicate which end is attracted to water:  
\_\_\_\_\_
  - How does this structure make the phospholipid molecule behave?  
\_\_\_\_\_  
\_\_\_\_\_  
\_\_\_\_\_



# Phospholipids and the Properties of Membranes

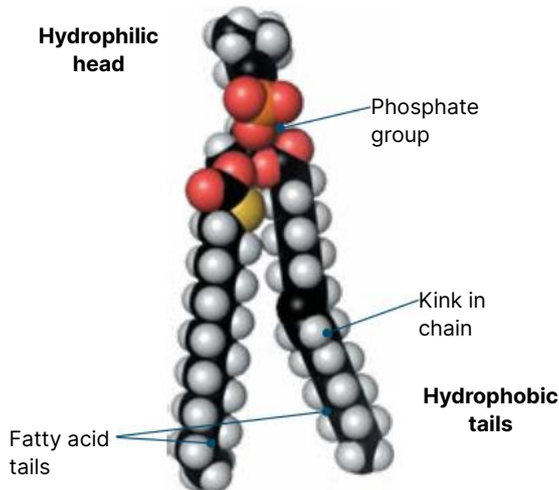
**Key Idea:** Phospholipids are important components of cellular membranes. They are made up of a hydrophilic head region and a hydrophobic tail region, making them amphipathic.

**Phospholipids** consist of a glycerol attached to two fatty acid chains and a phosphate ( $\text{PO}_4^{3-}$ ) group. Phospholipids

naturally form bilayers in aqueous solutions and are the main component of cellular membranes. The fatty acid tails can be saturated (straight chains) or unsaturated (kinked chains). The level of phospholipids with saturated or unsaturated tails affects the fluidity of the phospholipid bilayer.

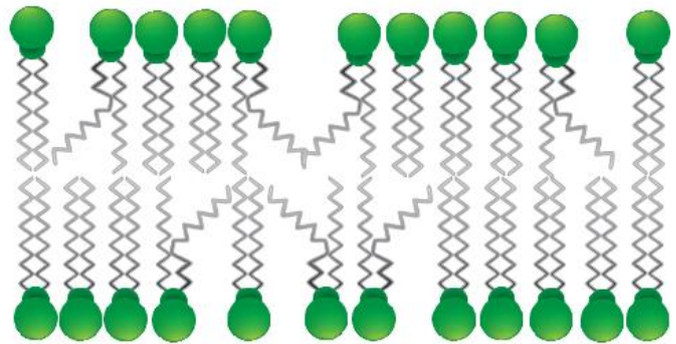
## Phospholipids

The phosphate end of the phospholipid is attracted to water (it is **hydrophilic**) while the fatty acid end is repelled (**hydrophobic**). In an aqueous environment, the hydrophobic ends turn inwards in the membrane to form a bilayer. Fatty acids containing double C=C bonds are unsaturated. This causes a "kink" in the chain.



## Phospholipids and membranes

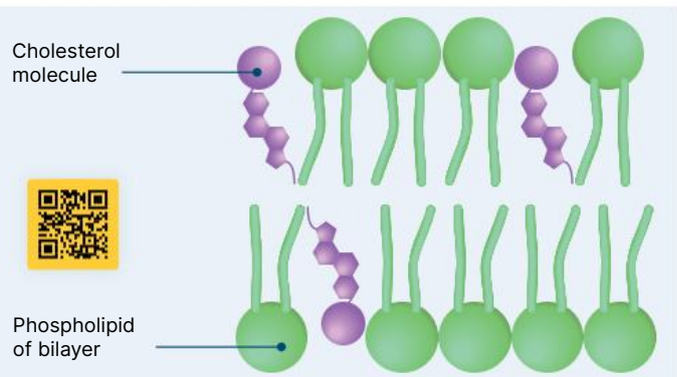
Phospholipids are amphipathic (have hydrophobic and hydrophilic regions). This means that they will spontaneously form bilayers when in aqueous environments and so form the outer boundary of cells and organelles. Modifications to the hydrophobic ends of the phospholipids regulate the fluidity of the bilayer. The greater the number of double bonds in the hydrophobic tails, the greater the fluidity of the membrane.



Membrane containing only phospholipids with saturated fatty acid tails (straight tails) and unsaturated fatty acid tails (bent). The fact that the phospholipids do not stack neatly together produces a more fluid membrane that may remain fluid even at low temperatures.

## Cholesterol in the cellular membrane

- ▶ **Cholesterol** molecules within the lipid bilayer help modulate the fluidity of the cellular membrane and maintain its stability.
- ▶ The polar head of the cholesterol molecule is attracted to the hydrophilic heads of the phospholipids: it reduces the fluidity in this region.
- ▶ Its hydrophobic tail aligns with the hydrophobic tails of the phospholipids and separates them.



1. (a) How do the properties of phospholipids contribute to their role in forming the structural framework of membranes?

---



---



---

- (b) Explain why phospholipid bilayers containing many phospholipids with unsaturated tails are particularly fluid:

---



---

2. Suggest how the cell membrane structure of an Arctic fish might differ from that of tropical fish species:

---



---



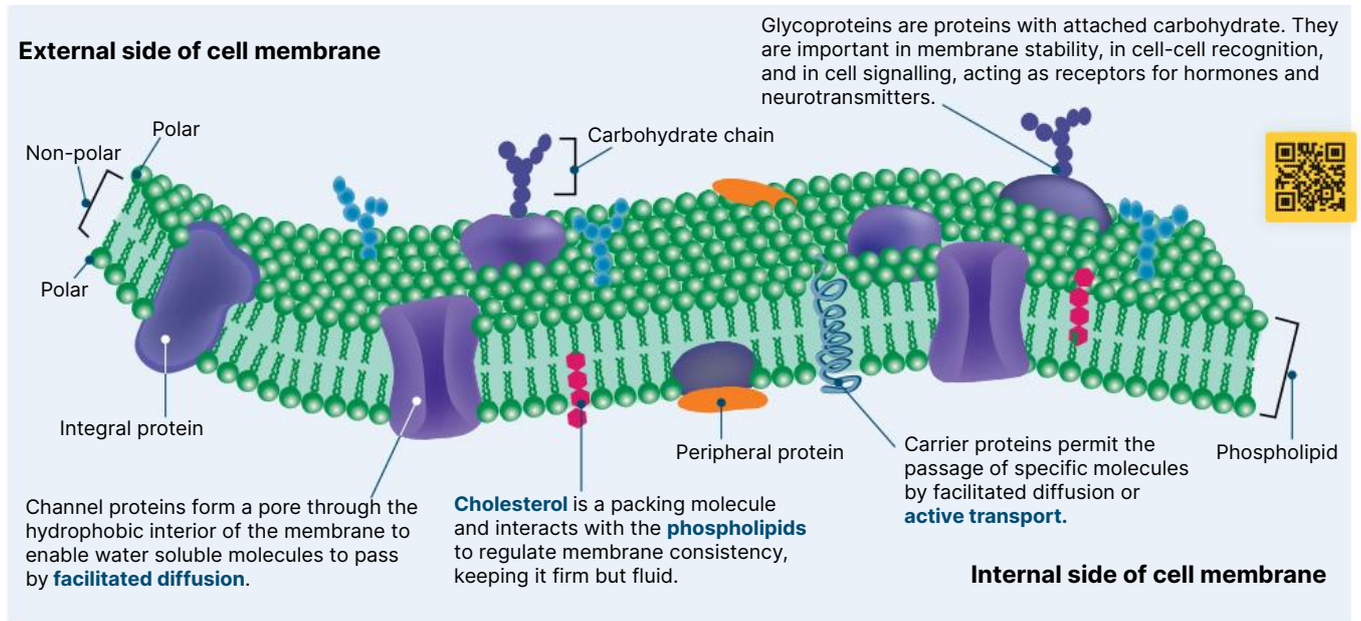
---

**Key Idea:** A cellular membrane is made of a phospholipid bilayer with different types of proteins embedded in it. The cell surface (or **plasma membrane**) encloses the cell's contents and regulates many of the cell's activities. Importantly, it controls what enters and leaves the cell by the use of **carrier proteins** and **channel proteins**. The structure and locations of membrane proteins enable them to perform their particular function in transport, cell signalling, or cell

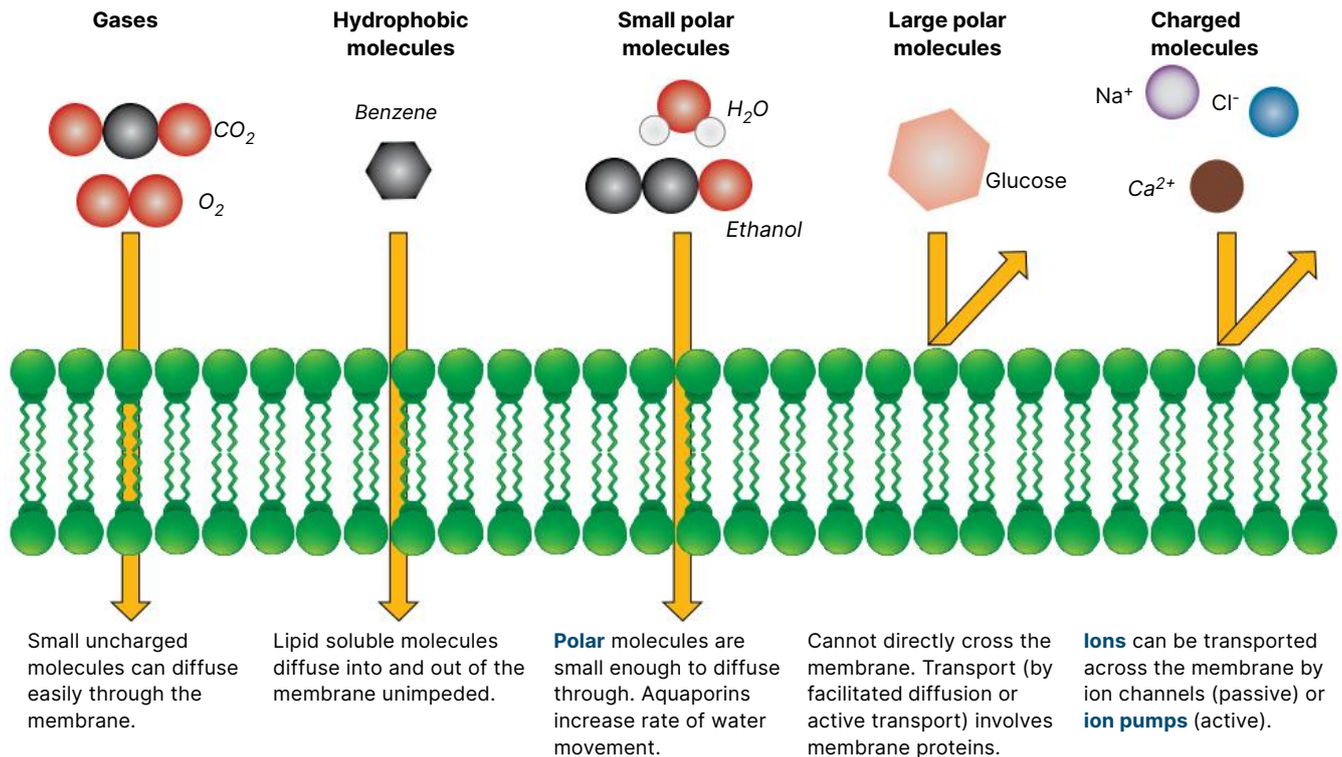
recognition. Proteins associated with the plasma membrane are either found embedded in the membrane, or found on the surface of the membrane. **Glycoproteins** are attached to the external side of the membrane and are involved in cell adhesion and cell recognition. Carrier proteins and channel proteins are transport proteins that allow substances to cross the membrane that are unable to utilise the processes of **diffusion** or **osmosis**.

## Glycoproteins

- ▶ Glycoproteins are membrane associated molecules composed of a carbohydrate component covalently bonded to an integral (embedded) protein.
- ▶ Glycoproteins facilitate attachment and adhesion to other cells. They also enable cell-to-cell communication and act as receptors for chemical signals. Other functions include memory consolidation in neuron cells, cell differentiation leading to changes in cell phenotype, and involvement in cancer growth.

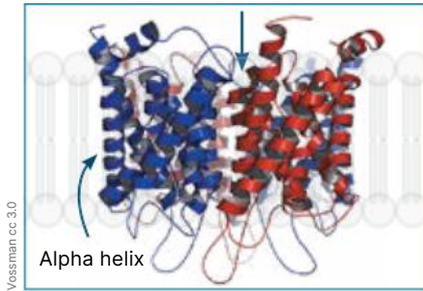


## What can cross a lipid bilayer?

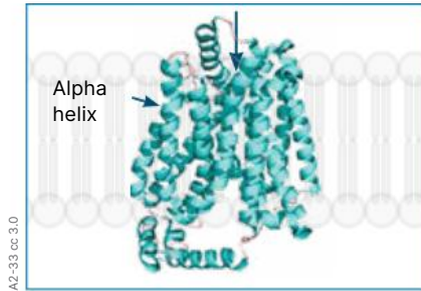


**What do proteins in the cell surface membrane really look like?**

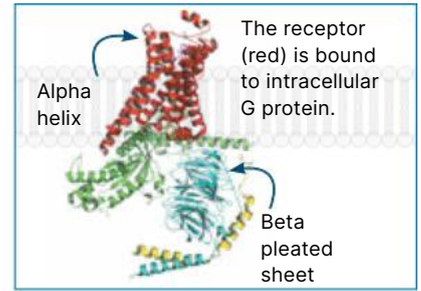
The structure of membrane proteins enables them to perform their particular function in transport, cell signalling, or cell recognition. The proteins are integral to the membrane, and often have parts of their structure projecting from both internal and external sides of the membrane. Note the two types of folding structure in membrane proteins: the alpha helix and the beta pleated sheet.



Aquaporins are a special type of channel protein that speed up the passage of water molecules across the membrane. Their tertiary structure creates a pore through the centre of the protein through which molecules can pass (arrow).



The GLUT1 glucose transporter is a carrier protein that facilitates the transport of glucose across the plasma membranes of mammalian cells. It increases the rate of glucose transport by 50,000X (high enough to supply the cell's energy needs).



G-protein coupled receptors are proteins involved in signalling pathways. A signal molecule binds to the receptor protein outside the cell to trigger a reaction involving intracellular G protein. In this example, the receptor binds to adrenaline.

1. What is the purpose of carrier proteins in the membrane? \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_
2. What is the purpose of channel proteins in the membrane? \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_
3. Identify the molecule(s) that:
  - (a) Can diffuse through the plasma membrane on their own:  
 \_\_\_\_\_
  - (b) Can diffuse through the membrane via channel proteins:  
 \_\_\_\_\_
  - (c) Must be transported across the membrane by carrier proteins:  
 \_\_\_\_\_
4. Describe the role of the following proteins in the plasma membrane:
  - (a) Aquaporins: \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_
  - (b) GLUT1 protein: \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_
  - (c) G protein: \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_

**Key Idea:** The freeze-fracture technique for preparing and viewing cellular membranes has provided evidence to support the fluid mosaic model of the plasma membrane.

Cellular membranes play many extremely important roles in cells and understanding their structure is central to understanding cellular function. Moreover, understanding the structure and function of membrane proteins is essential to understanding cellular transport processes, and cell recognition and signalling. Cellular membranes are far too small to be seen clearly using light microscopy, and certainly any detail is impossible to resolve. Since early last century, scientists have known that membranes were composed of a lipid bilayer with associated proteins. The original model of membrane structure, proposed by Davson and Danielli, was the unit membrane (a lipid bilayer coated with protein). This model was later modified by Singer and Nicolson after the discovery that the protein molecules were embedded within the bilayer rather than coating the outside. But how did they find out just how these molecules were organised?

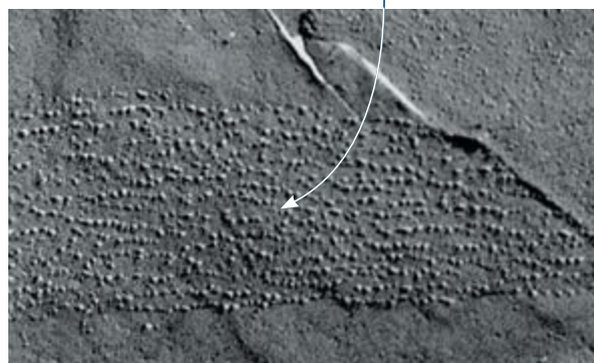
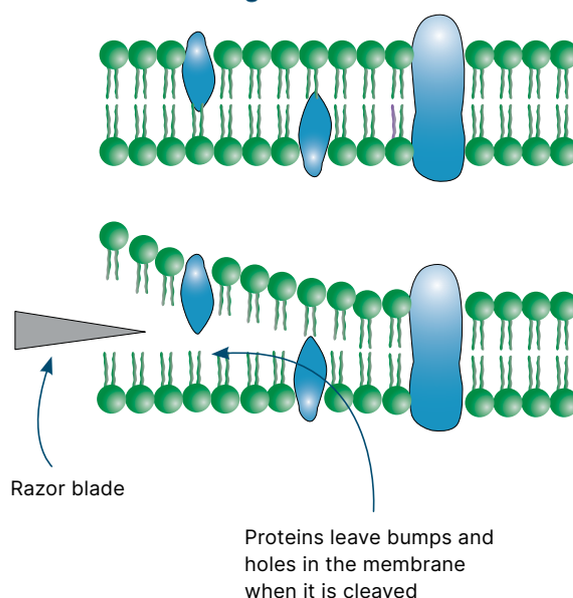
The answers were provided with electron microscopy, and one technique in particular – freeze fracture. As the name implies, freeze fracture, at its very simplest level, is the freezing of a cell and then fracturing it so the inner surface of the membrane can be seen using electron microscopy. Membranes are composed of two layers of **phospholipids** held together by weak intermolecular bonds. These split apart during fracture.

The procedure involves several steps:

- ▶ Cells are immersed in chemicals that alter the strength of the internal and external regions of the **plasma membrane** and immobilise any mobile macromolecules.
- ▶ The cells are passed through a series of glycerol solutions of increasing concentration. This protects the cells from bursting when they are frozen.
- ▶ The cells are mounted on gold supports and frozen using liquid propane.
- ▶ The cells are fractured in a helium-vented vacuum at  $-150$ . A razor blade cooled to  $-170^{\circ}\text{C}$  acts as both a cold trap for water and the fracturing instrument.
- ▶ The surface of the fractured cells may be evaporated a little to produce some relief on the surface (known as etching) so that a three-dimensional effect occurs.
- ▶ For viewing under an electron microscope (EM), a replica of the cells is made by coating them with gold or platinum to  $\sim 3$  nm thick. A layer of carbon around 30 nm thick is used to provide contrast and stability for the replica.
- ▶ The samples are then raised to room temperature and placed into distilled water or digestive enzymes, which separates the replica from the sample. The replica is then rinsed in distilled water before it is ready for viewing.

The freeze fracture technique provided the necessary supporting evidence for the current fluid mosaic model of membrane structure. When cleaved, proteins in the membrane left impressions that showed they were embedded into the membrane and not a continuous layer on the outside as earlier models proposed.

## Cleaving the membrane



50 nm

Photo: Louisa Howard and Chuck Daghljan, Dartmouth College

1. Explain how freeze-fracture studies provided evidence for our current model of membrane structure:

---



---



---



---

2. The Davson and Danielli model of membrane structure was the unit membrane; a phospholipid bilayer with a protein coat. Explain how the freeze-fracture studies showed this model to be flawed:

---



---



---



**Key Idea:** Research exploring the structure and function of channel proteins allows researchers to design membrane proteins with specific functions.

For the most part, the **fluid mosaic model** has generally proven to be correct. However, over time, as technology has improved, scientists have been able to study the components in more detail and a greater understanding of the model has

developed. An area of particular focus has been the protein components, including the structure and function of **channel proteins**. This information is valuable in understanding how the structure of a channel protein regulates the passage of molecules across the membrane. This knowledge can then be applied to the building of specific proteins that have particular characteristics.

### The contribution of Henderson and Unwin

Henderson and Unwin are two important contributors to membrane research and helped to refine the fluid mosaic model to incorporate a deeper understanding of membrane proteins. They are well known for their work on bacteriorhodopsin, a protein found in the purple membrane of Archaean cells. They were the first to provide the structure of a membrane protein *in situ* (in place) using electron microscopy. Their work (1975) revealed that the protein, which consists of three units, each made up of 7  $\alpha$ -helices (right), spanned the membrane. This was a major step in determining the structural make-up of membrane proteins.

### Studying channel proteins

Aquaporins are channel proteins that transport water across membranes. The first reported aquaporin, in 1986, was from the human red blood cell (RBC) membrane. Its function was discovered by selectively radiolabelling RBC membrane proteins with a water transport inhibitor. With the inhibitor present, water transport stopped, identifying the protein's role as a water channel. The structure of aquaporin was revealed some time later as four bundles of six  $\alpha$ -helices embedded in the cell membrane (below, right) surrounding a narrow channel. Supercomputer simulations identified the pathway of water through the channel. This research has been important in understanding the role of aquaporins in increasing the permeability of membranes to water.

### Future applications

Membrane proteins are relatively **hydrophobic** and their flexibility can make them unstable, so it can be difficult to extract them from the membrane in which they are embedded. These properties have made it difficult to study them. However, better imaging techniques and greater computing power have provided more detail about the structure and function of membrane proteins.

- ▶ Computer programs can now be used to predict how a membrane protein will fold once it has been synthesised. This has allowed molecular engineers to design and build artificial membrane proteins.
- ▶ One application of this is to improve the delivery of therapeutic drugs to target receptors (often membrane proteins). Building a protein allows researchers to study how a drug interacts with its target, and provides an opportunity to modify the drug for increased effectiveness.
- ▶ Another approach is to design new proteins with specific functions, e.g. to deliver specific substances across the membrane of a target cell.

1. Using the fluid mosaic model as an example, suggest why scientific models are often modified over time:

---



---



---

2. (a) Why has it been difficult to determine the structure of membrane proteins?

---



---

- (b) Explain the role in technological developments in furthering our understanding of membrane proteins:

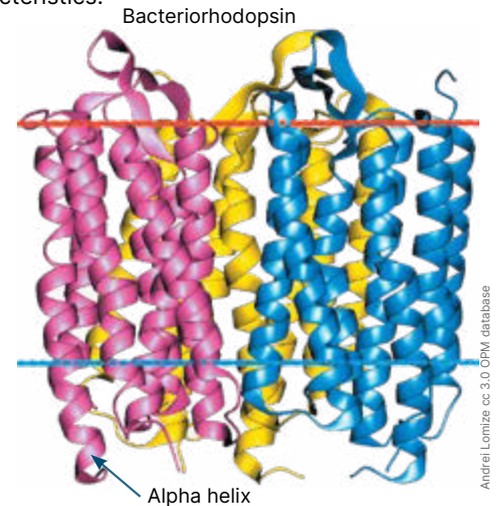
---



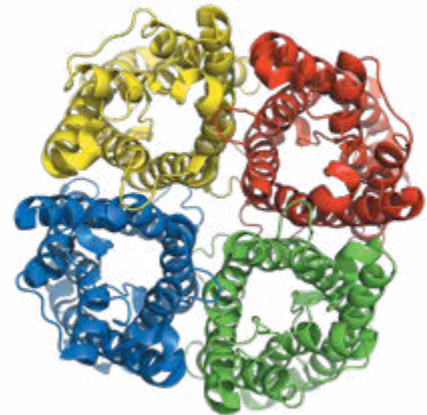
---



---



Bacteriorhodopsin acts as a light-driven proton pump in *Halobacterium salinarum*, converting light energy into a proton gradient, which is used to generate ATP.



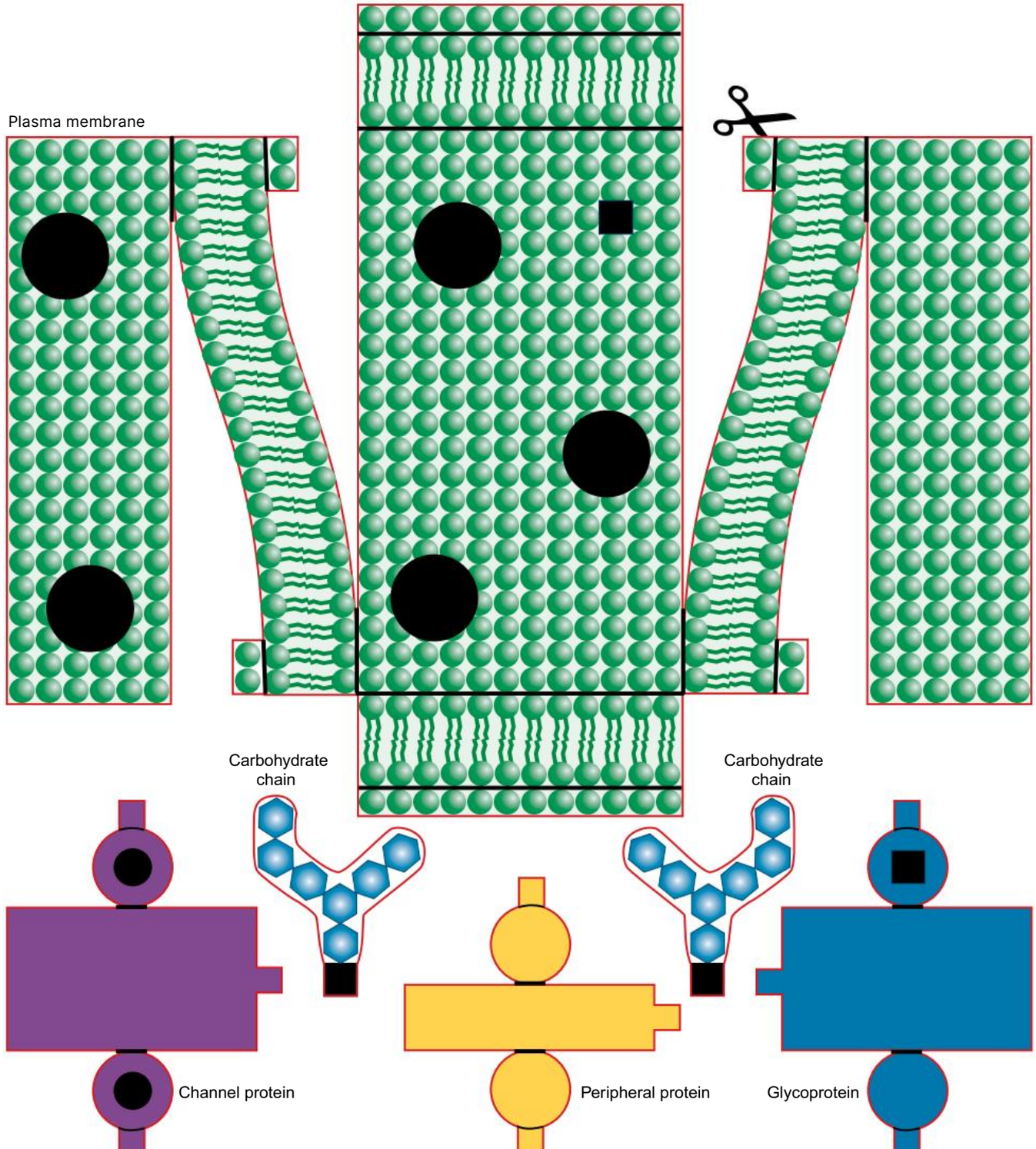
Aquaporin tetramer (4 units) viewed from the extracellular side. The pore is in the centre.

# Modelling the Plasma Membrane

**Key Idea:** Plasma membranes exist as dynamic 3D structures. **Plasma membranes** are often shown as two dimensional structures. Even when drawn to represent a three

dimensional structure, the nature of the plasma membrane may not be obvious. In this activity you will build a simple, three dimensional plasma membrane.

1. Cut out the plasma membrane along the red lines. Cut out the solid black circles. Fold along the black lines. Use clear tape to stick the sides together to produce a 3D, slightly curved box.
2. Cut out the three proteins along the red lines. Fold along the black lines and use clear tape to produce three cylinders.
3. Cut out both carbohydrate chains. Fold over the black squares. Stick one to the black square on the end of the **glycoprotein**. Stick the other to the black square on the plasma membrane surface to produce a glycolipid.
4. Slide the two transmembrane proteins into the channels created by cutting out the circles from the plasma membrane.
5. Slide the peripheral protein about halfway into the final hole. This completes your plasma membrane model.



**This page is deliberately left blank**

# 61 Diffusion

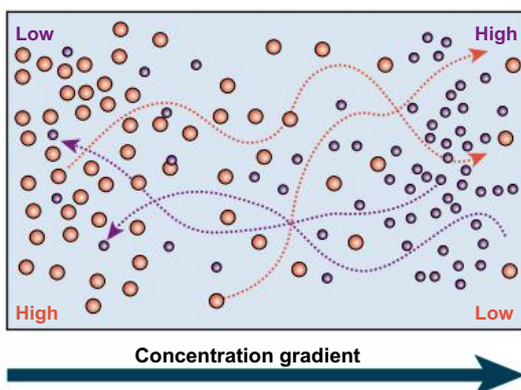
**Key Idea:** Diffusion is a type of passive transport where molecules move down a concentration gradient.

The molecules that make up substances are constantly moving about in a random way. This random motion causes them to disperse from areas of high to low concentration. This dispersal is called **diffusion** and it requires no energy. Each type of molecule moves down its own **concentration**

**gradient**. In biological systems, most diffusion occurs across membranes. Some molecules move freely (unassisted) across the membrane by simple diffusion. For other molecules, their diffusion is helped by proteins in the membrane. Diffusion is important in allowing cells to make exchanges with their extracellular environment (e.g. the blood and fluids that bathe them) and is crucial to the regulation of water content.

## What is diffusion?

Diffusion is the movement of particles down a concentration gradient. Diffusion is a passive process, meaning it needs no input of energy to occur. During diffusion, molecules move randomly about, eventually becoming evenly dispersed.

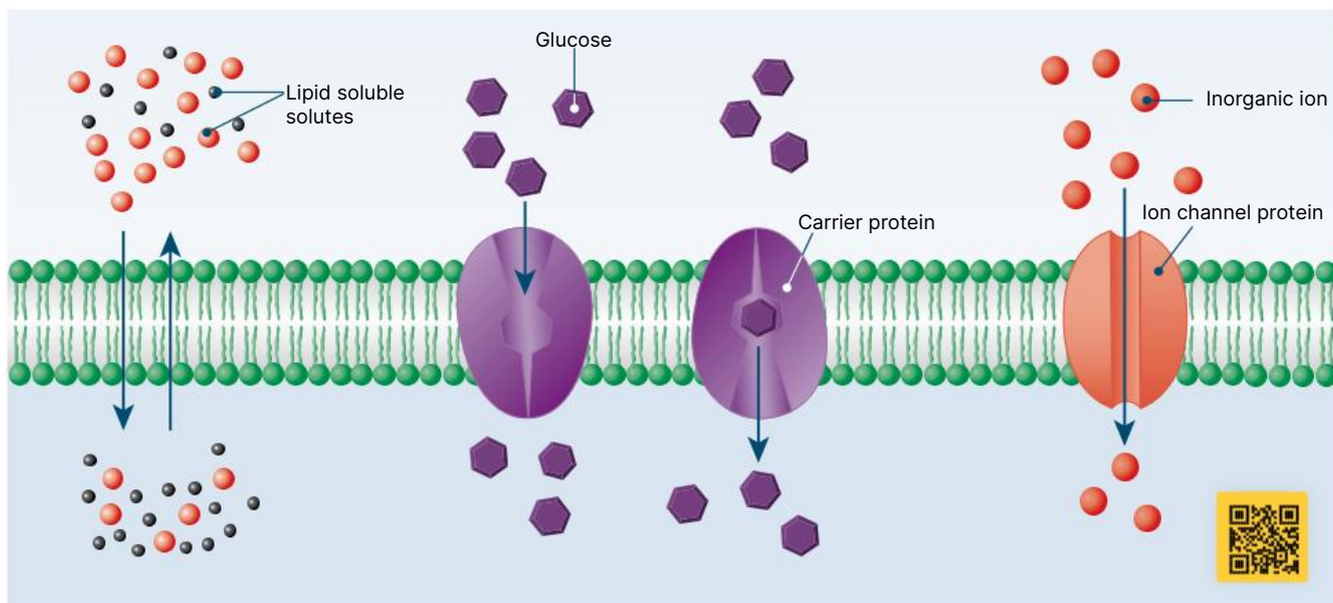


If molecules can move freely, they move from high to low concentration (down a concentration gradient) until evenly dispersed. Each molecule moves down its own concentration gradient, independent of the concentration of other types of molecule

## Factors affecting the rate of diffusion

<b>Concentration gradient</b>	The rate of diffusion is higher when there is a greater difference between the concentrations of two regions.
<b>The distance moved</b>	Diffusion over shorter distance occurs at a greater rate than over a larger distance.
<b>The surface area involved</b>	The larger the area across which diffusion occurs, the greater the rate of diffusion.
<b>Barriers to diffusion</b>	Thick barriers have a slower rate of diffusion than thin barriers.
<b>Temperature</b>	Particles at a high temperature diffuse at a greater rate than at a low temperature.
<b>Solubility</b>	Lipid-soluble or non-polar molecules pass across membranes more easily than polar materials, so their rates of diffusion are faster.
<b>Solvent density</b>	As the density of a solvent increases, the rate of diffusion decreases. Cellular dehydration adversely affects diffusion rates within cells.

## Diffusion across the membrane



### Simple diffusion

Molecules move directly through the plasma membrane without assistance or selectivity. **Example:**  $O_2$  diffuses into the blood and  $CO_2$  diffuses out. Diffusion gradients are maintained because substances are constantly being imported, made, or used by the cell.

### Facilitated diffusion involving carrier proteins

**Carrier proteins** in the membrane allow large, lipid-insoluble molecules that cannot cross the membrane by simple diffusion to be transported into the cell. **Example:** The transport of glucose into red blood cells.

### Facilitated diffusion involving channel proteins (hydrophilic pores)

**Channel proteins** (water-filled pores) allow selective permeability of substances such as inorganic **ions** to pass through. Aquaporins are special channel proteins for rapid diffusion of water. **Example:**  $K^+$  ions exiting nerve cells to restore resting potential.



1. What do the three types of diffusion described on the previous page all have in common?

---



---

2. How does facilitated diffusion differ from simple diffusion? \_\_\_\_\_

---



---

3. Why is carbon dioxide able to continually diffuse out of cells? \_\_\_\_\_

---



---

4. Why would a thin flat cell have a greater rate of diffusion to and from its centre than a thick spherical cell?

---



---



---



---

### Observing diffusion

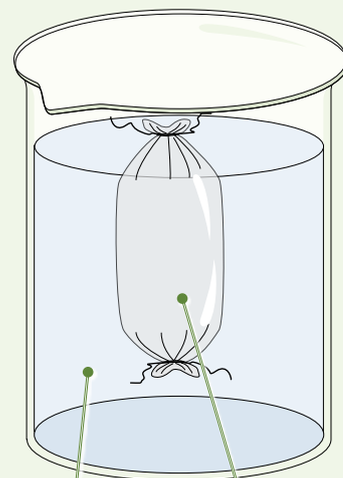
Diffusion through a **partially permeable** membrane can be modelled using dialysis tubing. The pores of the dialysis tubing determine the size of the molecules that can pass through. In the experiment described below, you will investigate how glucose will diffuse down its concentration gradient from a high glucose concentration to a low glucose concentration and demonstrate, via the model, the selective permeability of the plasma membrane.



#### Investigation 4.1 Simple diffusion across a membrane

See appendix for equipment list.

1. Add 200 mL of distilled water to a clean 200 mL beaker. Remove a 1 mL sample and place in a clean test tube. Use a glucose dipstick to test for the presence and concentration of glucose in the 1 mL sample. If glucose is present, the indicator window will change colour. The colour change can be compared against a reference to determine the concentration of glucose present.
2. Now add a few drops of Lugol's indicator to test for the presence of starch. Lugol's indicator contains iodine, and turns blue/black in the presence of starch.
3. Obtain a short section of dialysis tubing, approximately 10 cm long. Use thread or nylon line to tie off one end (or tie a knot in the tubing if long enough).
4. You may need to rinse the tubing under water to make it pliable enough to open.
5. Fill the dialysis tubing with 5 mL each of a 1% starch solution and a 10% glucose solution.
6. Remove a 1 mL sample and place in a clean test tube. Tie off the top of the dialysis tubing, rinse well with distilled water, then place in the beaker of distilled water.
7. Test for the presence and concentration of glucose and then starch in the sample from the dialysis tubing as in steps 1 and 2.
8. Leave the dialysis tubing in the distilled water for 30 minutes.
9. Remove 1 mL of water from the beaker and place in a clean test tube. Use a glucose dipstick to test for the presence and concentration of glucose. Test for the presence of the starch using Lugol's indicator.
10. Remove a 1 mL sample from the dialysis tubing and place in a clean test tube. Use a glucose dipstick to test for the presence and concentration of glucose. Test for the presence of the starch using Lugol's indicator.



Distilled water      Solution containing starch and glucose

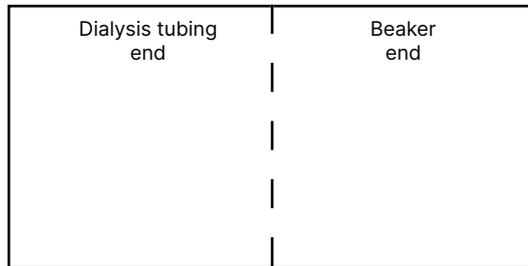
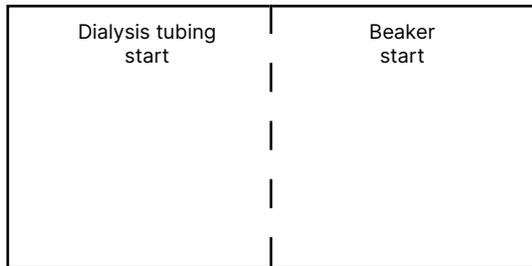
5. What is the aim of the experiment? \_\_\_\_\_  
 \_\_\_\_\_
6. What part of a cell does the dialysis tubing represent? \_\_\_\_\_
7. Why was it important to wash the dialysis tubing before placing it into the beaker of distilled water?  
 \_\_\_\_\_  
 \_\_\_\_\_

8. Complete the result table, right:

For relative concentration of glucose, use + for relatively low concentration and ++ for relatively high concentration.

	Beaker start	Dialysis tubing start	Beaker end	Dialysis tubing end
Starch (+/-)				
Glucose (relative concentration)				

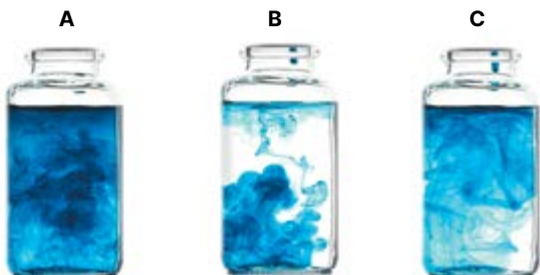
9. In the spaces provided (below) draw the distribution of starch and glucose at the start and at the end of the experiment. Use the coloured symbols shown under the table to represent starch and glucose:



10. Explain your results: \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_

11. Suggest how a cell could regulate the rate of facilitated diffusion of specific molecules:  
 \_\_\_\_\_  
 \_\_\_\_\_

12. Why is glucose able to continually diffuse into a cell? \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_



13. Study the images below. Place them in order of first event to last event. Explain your order of events in terms of diffusion:

\_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_

# 62 Diffusion and Cell Size

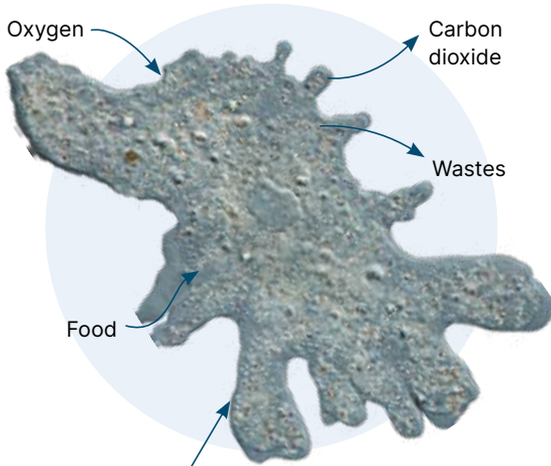
**Key Idea:** Diffusion is less efficient in cells with a small surface area relative to their volume than in cells with a large surface area relative to their volume.

Small objects, such as cells, have a large surface area relative to their volume and **diffusion** is an effective way to move materials in and out. As an object becomes larger, its **surface**

**area to volume ratio** is smaller and diffusion is no longer an effective way to transport materials to the inside. The effectiveness of diffusion is therefore the controlling factor determining how big an individual cell can become. In large, multicellular organisms, specialised systems deliver materials to the many cells that make up the tissues of the body.

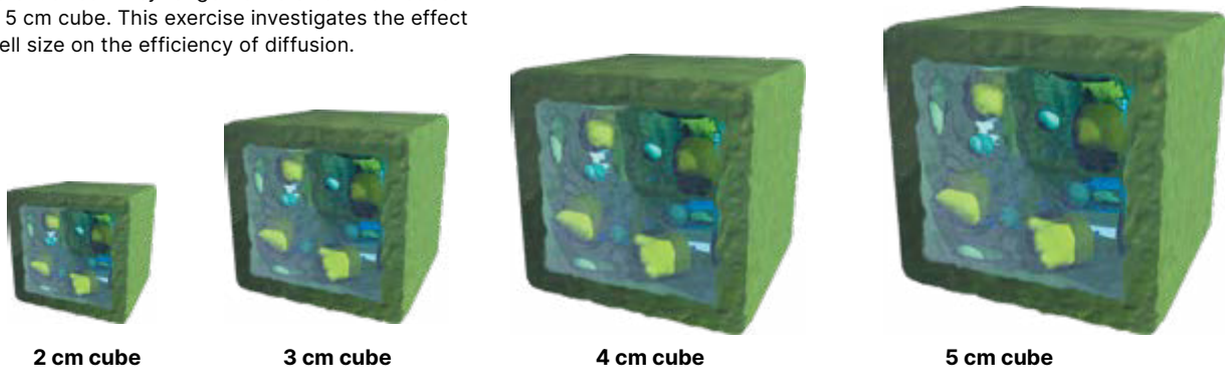
## Single-celled organisms

Single-celled organisms (e.g. *Amoeba*), are small and have a large surface area relative to the cell's volume. The cell's requirements can be met by the diffusion or **active transport** of materials into and out of the cell (below).



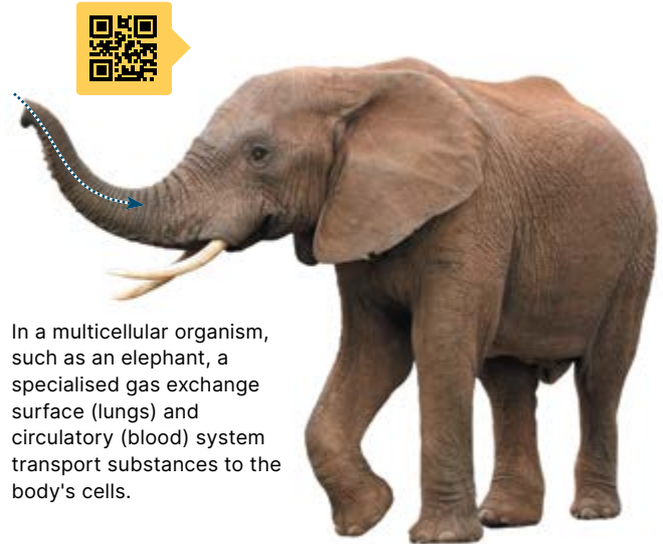
The **plasma membrane**, which surrounds every cell, regulates movements of substances into and out of the cell. For each square micrometre of membrane, only so much of a particular substance can cross per second.

The diagram below shows four hypothetical cells of different sizes. They range from a small 2 cm cube to a 5 cm cube. This exercise investigates the effect of cell size on the efficiency of diffusion.



## Multicellular organisms

Multicellular organisms (e.g. plants and animals) generally have a small surface area compared to their volume. They require specialised body systems to transport the materials they need to and from the cells and tissues in their body.



In a multicellular organism, such as an elephant, a specialised gas exchange surface (lungs) and circulatory (blood) system transport substances to the body's cells.

1. Calculate the volume, surface area and the ratio of surface area to volume for each of the four cubes above (the first has been done for you). When completing the table below, show your calculations.

Cube size	Surface area	Volume	Surface area to volume ratio
2 cm cube	$2 \times 2 \times 6 = 24 \text{ cm}^2$ (2 cm x 2 cm x 6 sides)	$2 \times 2 \times 2 = 8 \text{ cm}^3$ (height x width x depth)	$24 \text{ to } 8 = 3:1$
3 cm cube			
4 cm cube			
5 cm cube			



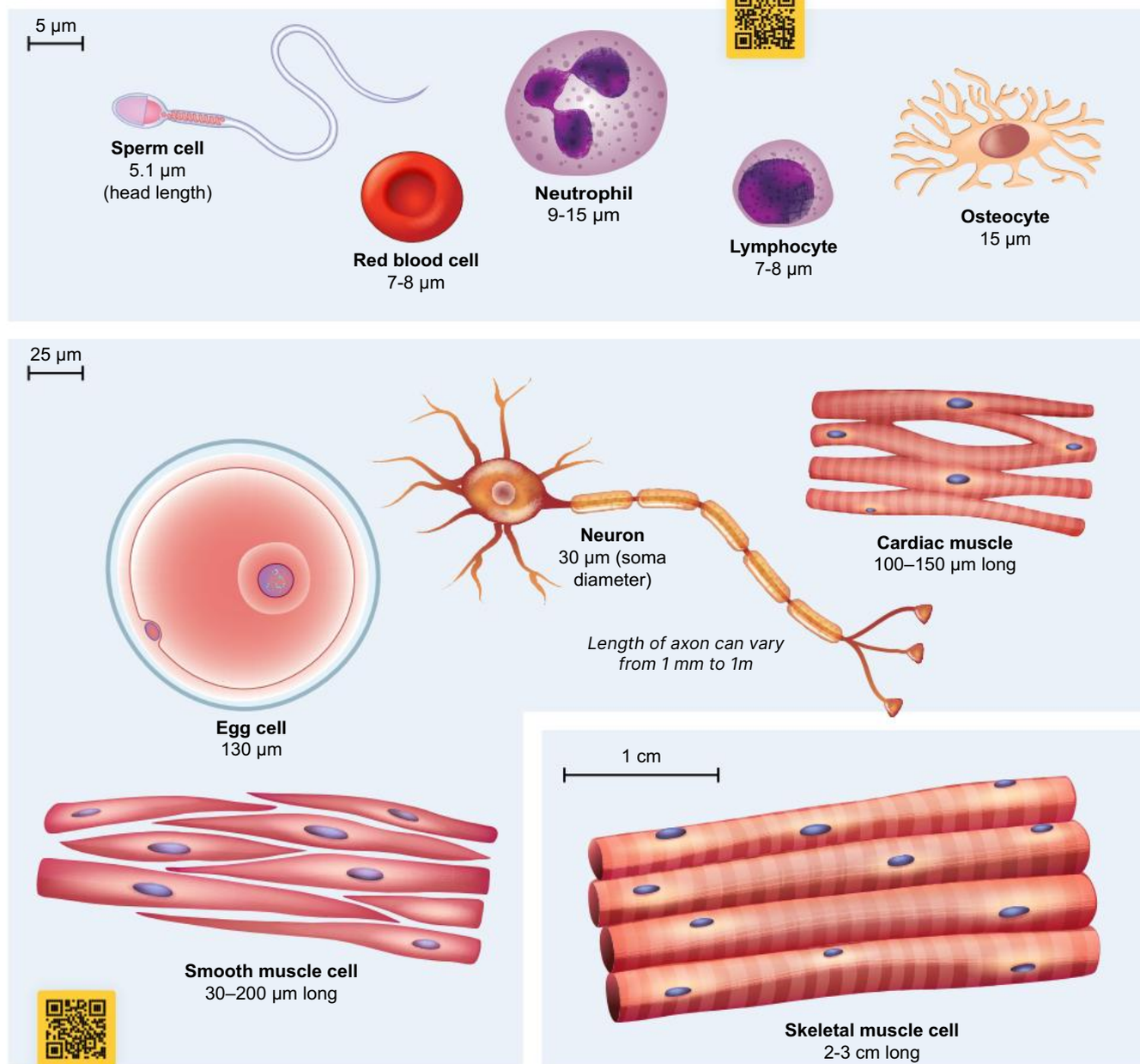
# 63 Comparing Cell Sizes

**Key Idea:** Different types of cells vary in size and this is a function of their specialisation.

Humans have around 200 different specialised cell types, and each type has adaptations to allow it to perform a particular role. One adaptation is cell size. Different types of

cells are different sizes. These vary from the tiny blood cells, which need to be small enough to fit through capillaries, to long neurons that need to connect to other cells throughout the body. Size is one factor that is integral to the cell's differentiation, and therefore its function.

## Typical sizes of various human cells



### Unit of length (international system)

Unit	Metres	Equivalent
1 meter (m)	1 m	= 1000 millimetres
1 millimetre (mm)	$10^{-3}$ m	= 1000 micrometres
1 micrometre ( $\mu\text{m}$ )	$10^{-6}$ m	= 1000 nanometres
1 nanometre (nm)	$10^{-9}$ m	= 1000 picometres

Micrometres are sometimes referred to as microns. Smaller structures are usually measured in nanometres (nm) e.g. molecules (1 nm) and plasma membrane thickness (10 nm).

1. (a) Compare length of the three muscle cells in  $\mu\text{m}$ :

---



---

- (b) Why do scientific diagrams require the inclusion of a scale as part of them?

---



---



---



---

# 64 Investigating the Effect of Cell Size

**Key Idea:** Diffusion is less efficient in cells with a small surface area relative to their volume than in cells with a large surface area relative to their volume.

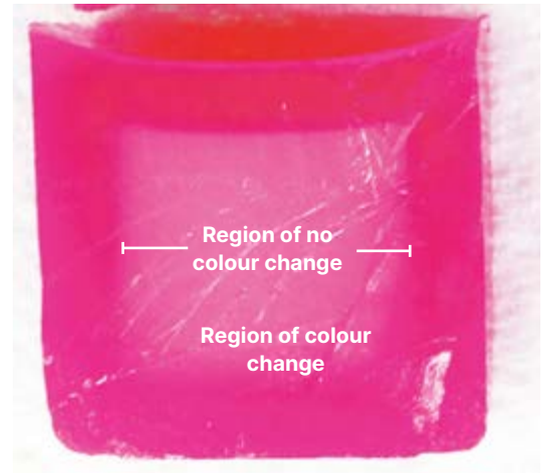
When an object (e.g. a cell) is small it has a large surface area in comparison to its volume. **Diffusion** is an effective way to transport materials (e.g. gases) into and out of the cell. As an object becomes larger, its surface area compared to its

volume is smaller and diffusion is no longer an effective way to transport materials to and from the inside. In this activity you will design an experiment to demonstrate the effect of **surface area: volume ratios** on diffusion in model cells. Think about how you will plan your investigation and analyse your data to obtain meaningful results. This will help you to make valid conclusions about your findings.

## Background information

Oxygen, water, cellular waste, and many nutrients are transported into and out of cells by diffusion. However, at a certain surface area to volume ratio, diffusion becomes inefficient. In this activity you will create model cells of varying sizes from agar and use them to test the relationship between cell size and rate or efficiency of diffusion.

- ▶ The diffusion of molecules into a cell can be modelled by using agar cubes infused with phenolphthalein indicator and soaked in sodium hydroxide (NaOH).
- ▶ Phenolphthalein is an acid/base indicator and turns pink in the presence of a base.
- ▶ As the NaOH diffuses into the agar, the phenolphthalein changes to a pink colour and thus indicates how far into the agar block the NaOH has diffused (right).
- ▶ By cutting an agar block into cubes of various sizes, it is possible to investigate the effect of cell size on diffusion.



A phenolphthalein-infused agar cube after exposure to NaOH.

## Equipment list



Glass beaker



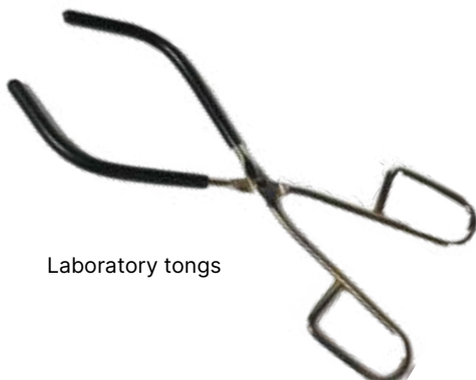
Paper towel



Timer



Agar blocks infused with phenolphthalein

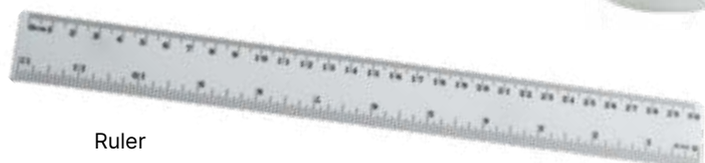


Laboratory tongs



Scalpel

Sodium hydroxide (NaOH) solution



Ruler





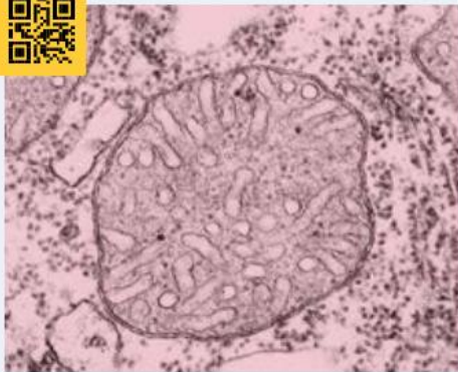
# Overcoming Limitations to Cell Size

**Key Idea:** Larger cells can maintain high surface area to volume ratios by having a non-spherical shape and membrane extensions. Organelles also increase functional efficiency. We have seen that cells must exchange materials with the extracellular environment in order to survive. The efficiency of these exchanges, which must occur across the **plasma**

**membrane**, is limited by the cell's **surface area to volume ratio**. Larger cells can maintain higher SA:V ratios by having a non-spherical shape and extensions of the membrane. Within the cell, the presence of organelles specialised to perform particular functions creates cellular compartments, which also improve functional efficiency in a larger cell.

### Cell size and functional efficiency

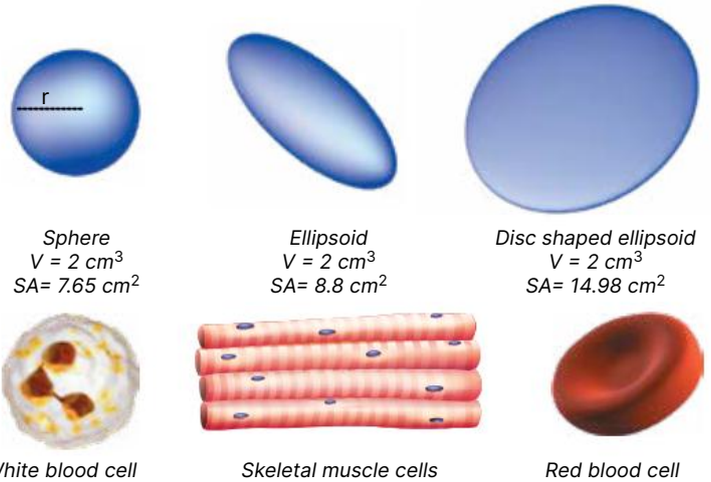
Cells have a wide range of sizes. Large eukaryotic cells may reach 100 µm in diameter, whereas bacteria typically only reach a tenth of that. Eukaryotic cells can remain efficient at larger sizes in part because they contain organelles, which concentrate associated materials (such as the reactants and enzymes in a metabolic pathway) into specific regions for specific purposes. These cellular compartments enable efficiency of function.



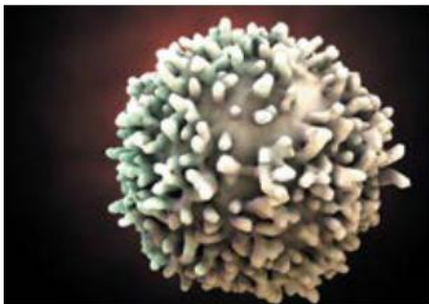
Cellular respiration occurs within the mitochondria, which has regions in which different reactions occur.

### Solving the size problem

One way of increasing a cell's surface area while retaining the same volume is to elongate the cell. An elongated sphere (an ellipsoid, e.g. a rod shaped cell) has a greater surface area than a sphere of the same volume. In this way, a cell can grow larger while still gaining the materials it needs. The cells of multicellular organisms are often highly specialised to maximise SA: V. The three images below are all to scale.



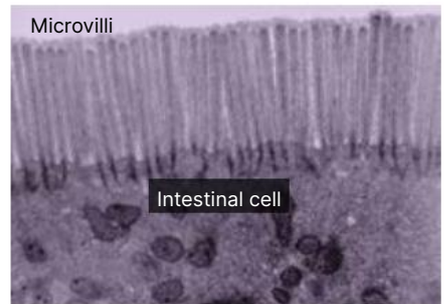
By flattening the ellipsoid along one axis and stretching it along the other two to form a disc, surface area increases while the volume remains the same.



Animal cells, such as this B cell (a type of white blood cell), often have extensions of the cell membrane providing a high surface area for transfer of materials.



Tissues are organised to increase surface area. Here, the intestinal wall is folded into projections called villi. Column-shaped intestinal cells line the surface of the villi.



The cell membrane of each intestinal cell is folded into numerous microvilli. These increase the surface area for absorbing nutrient and binding digestive enzymes.

- Use the formula  $4\pi r^2$  (where  $\pi = 3.14$ ) to calculate the surface area of a spherical cell with a radius (r) of:
  - 2 µm: \_\_\_\_\_
  - 5 µm: \_\_\_\_\_
  - 10 µm: \_\_\_\_\_
  - 30 µm: \_\_\_\_\_
- What happens to the SA:V ratio of a spherical cell as its volume increases? \_\_\_\_\_
  - How can eukaryotic cells overcome the restrictions of reduced SA:V as they become larger? \_\_\_\_\_

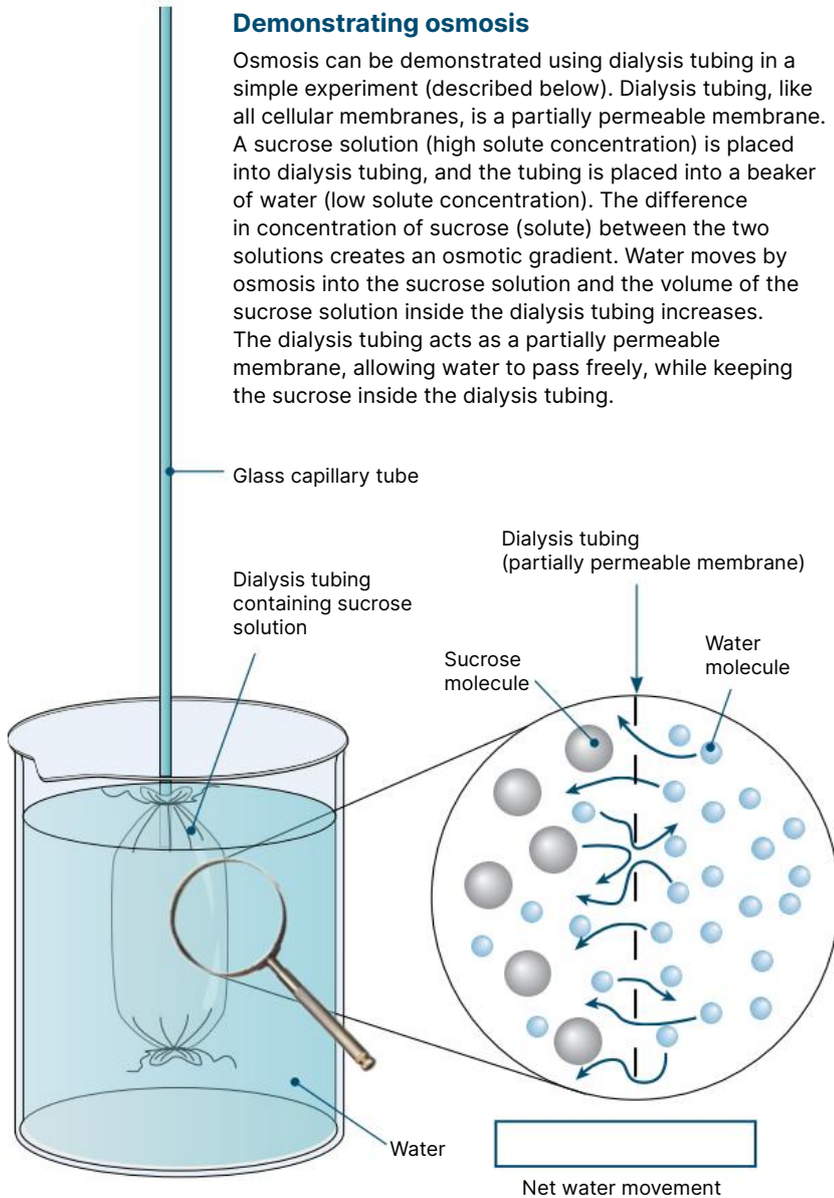
**Key Idea:** Osmosis is the diffusion of water molecules from a lower solute concentration to a higher solute concentration across a partially permeable membrane.

**Osmosis** is the **diffusion** of water molecules from regions of lower solute concentration (higher free water concentration) to regions of higher solute concentration (lower free water concentration) across a **partially permeable** membrane. A

partially permeable membrane allows some molecules, but not others, to pass through. Water molecules will diffuse across a partially permeable membrane until an equilibrium is reached and net movement is zero. The plasma membrane of a cell is an example of a partially permeable membrane. Osmosis is a passive process and does not require any energy input.

### Demonstrating osmosis

Osmosis can be demonstrated using dialysis tubing in a simple experiment (described below). Dialysis tubing, like all cellular membranes, is a partially permeable membrane. A sucrose solution (high solute concentration) is placed into dialysis tubing, and the tubing is placed into a beaker of water (low solute concentration). The difference in concentration of sucrose (solute) between the two solutions creates an osmotic gradient. Water moves by osmosis into the sucrose solution and the volume of the sucrose solution inside the dialysis tubing increases. The dialysis tubing acts as a partially permeable membrane, allowing water to pass freely, while keeping the sucrose inside the dialysis tubing.



### Osmotic potential

The presence of solutes (dissolved substances) in a solution increases the tendency of water to move into that solution. This tendency is called the osmotic potential or osmotic pressure. The more total dissolved solutes a solution contains, the greater its osmotic potential.

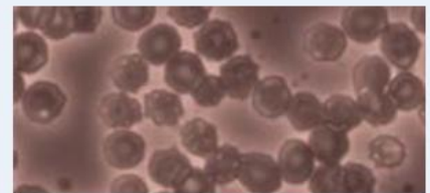
### Describing solutions

Osmosis is important when handling body tissues for medical transport or preparation. The tissue must be bathed in a solution with an **osmolarity** (a measure of solute concentration) equal to the tissue's to avoid a loss or gain of fluid in the tissue. Solutions separated by a partially permeable membrane are often described in terms of their solute concentrations relative to one another.

**Isotonic solution:** Having the same solute concentration relative to another solution (e.g. the cell's contents).

**Hypotonic solution:** Having a lower solute concentration relative to another solution.

**Hypertonic solution:** Having a higher solute concentration relative to another solution.



The red blood cells above were placed into a hypertonic solution. As a result, the cells have lost water and have begun to shrink, losing their usual discoid shape.

1. What is osmosis? \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_
2. (a) In the blue box on the diagram above, draw an arrow to show the direction of net water movement.  
 (b) Why did water move in this direction? \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_
3. What would happen to the height of the water in the capillary tube if the sucrose concentration was increased?  
 \_\_\_\_\_



**Key Idea:** Determining loss or gain of mass in tissues allows us to determine the osmolarity of the tissue's cells.

The **osmolarity** (a measure of solute concentration) of a cell or tissue can be estimated by placing part of the cell or

tissue into a series of solutions of known concentration and observing if the tissue loses (**hypertonic** solution) or gains (**hypotonic** solution) water. The solution in which the tissue remains unchanged indicates the osmolarity of the tissue.

## Investigation 4.2 Estimating osmolarity

See appendix for equipment list.

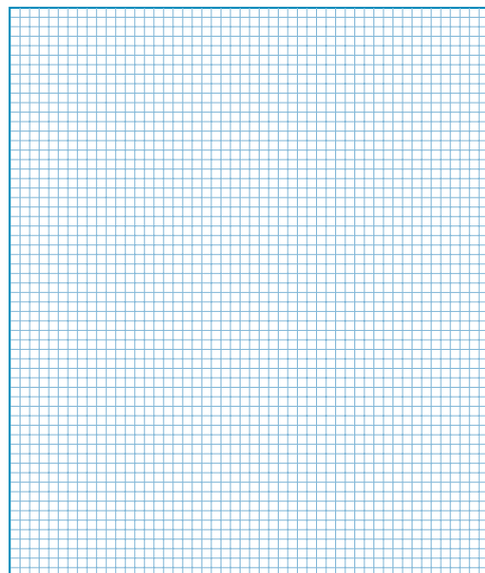
1. Prepare 6 beakers of sucrose ( $C_{12}H_{22}O_{11}$ , table sugar) solution with the concentrations of 0.0 (distilled water), 0.1, 0.2, 0.3, 0.4, and 0.5 mol L<sup>-1</sup> of sucrose (0, 34.2 g, 68.5 g, 102.6 g, 136.9 g, and 171.1 g per litre). Label the beakers so that they can be easily identified at the end of the experiment.
2. Peel a potato and cut it into 18 identical cubes 1 cm<sup>3</sup> (1 cm x 1 cm x 1 cm) or use a cork borer to produce 18 identical cylinders of potato. Pat the potato cubes dry with a paper towel.
3. Weigh three cubes together, record their mass in the table below under initial mass. Place the cubes in the beaker of distilled water.
4. Repeat step 3 with the other 15 potato cubes and concentrations. Make sure you identify each beaker so the cubes can be weighed at the end of the experiment.
5. Leave the potato cubes in the solutions for at least 40 minutes (or up to 24 hours).
6. Remove the potato cubes from the distilled water and pat dry with a paper towel. Weigh all three together and record their mass in the table below under final mass.
7. Repeat for all the other concentrations of sucrose.
8. Calculate the change in mass (if any) for all the concentrations. Then calculate the % change (+ or -) (this removes any error based on the masses of the potato cubes not being identical).
9. Plot the % change vs sucrose concentration on the grid provided.

**Need help?**  
See Activity 12



Sucrose concentration (mol/L)	Initial mass (I) (g)	Final mass (F) (g)
0.00		
Change (C) = (F-I) (g)		
% Change = (C/I x 100)		
0.1		
Change (C) = (F-I) (g)		
% Change = (C/I x 100)		
0.2		
Change (C) = (F-I) (g)		
% Change = (C/I x 100)		
0.3		
Change (C) = (F-I) (g)		
% Change = (C/I x 100)		
0.4		
Change (C) = (F-I) (g)		
% Change = (C/I x 100)		
0.5		
Change (C) = (F-I) (g)		
% Change = (C/I x 100)		

1. Use the grid below to draw a line graph of the sucrose concentration vs total % change in mass:



2. Use the graph to estimate the osmolarity of the potato (the point where there is no change in mass):  
\_\_\_\_\_
3. Which of the solutions are hypotonic? Which are hypertonic?  
\_\_\_\_\_  
\_\_\_\_\_



**Key Idea:** Plant cells in a hypertonic solution lose water and undergo plasmolysis. In a hypotonic solution, they gain water creating turgor pressure.

**Osmosis** across the **partially permeable** cell membrane is the main way by which water enters and leaves the cell.

### Osmosis and turgidity

When the watery contents of a plant cell push against the cell wall they create **turgor** (tightness) which helps to provide support for the plant body. When cells lose water, there is a loss of cell turgor and the plant will wilt. Complete loss of turgor from a cell is called **plasmolysis** and is irreversible. Two systems (cell and environment) with the same effective osmotic pressure are termed isotonic and there is no net movement of water molecules. However, when there is an osmotic gradient between the cell and environment there will be a net movement of water molecules down their concentration gradient. The diagram below shows two different situations: when a plant cell is in a **hypertonic** solution and when it is in a **hypotonic** solution.

When the external concentration of free water molecules is the same as that of the cytoplasm there is no net movement of water. Changing the tonicity of the external environment will cause a net movement of water into or out of the cell as water moves down its **concentration gradient**.

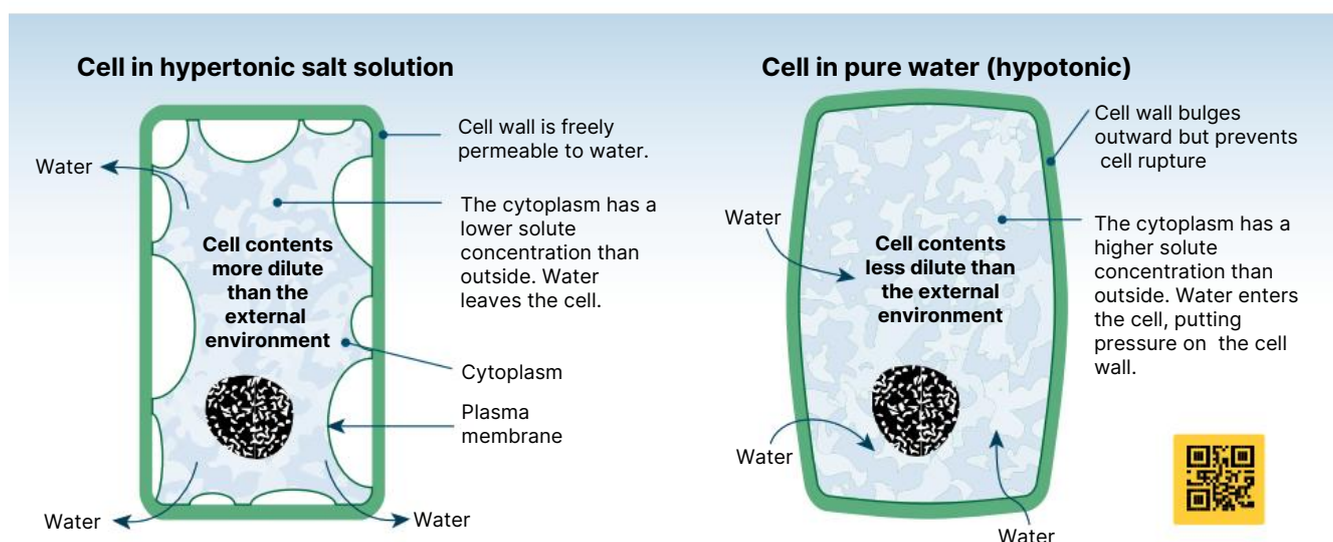


Wilted plant (cells have lost turgor)



Plant cells are turgid

MCC UW



### Plasmolysis in a plant cell

Tonicity is a measure of the osmotic pressure of a solution. In a hypertonic solution, the external free water concentration is lower than the free water concentration of the cell. Water leaves the cell and, because the cell wall is rigid, the cell membrane shrinks away from the cell wall. This is called **plasmolysis** and the cell becomes flaccid.

### Turgor in a plant cell

In a hypotonic solution, the external free water concentration is higher than the cell cytoplasm. Water enters the cell, causing it to swell tight. A wall (turgor) pressure is generated when the cell contents press against the cell wall. Turgor pressure increases until no more water enters the cell (the cell is **turgid**).

1. Identify the outcome of the following situations:

- (a) A plant cell is placed in a hypertonic solution: \_\_\_\_\_
- (b) A plant cell is placed in a hypotonic solution: \_\_\_\_\_
- (c) A plant cell in an isotonic solution: \_\_\_\_\_

2. (a) Explain the role of cell wall pressure in generating cell turgor in plants: \_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_

(b) Discuss the role of cell turgor in plants: \_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_

# Investigating Membrane Solubility and Diffusion

**Key Idea:** The rate of diffusion of molecules through the plasma membrane can be determined by measuring the change in light absorbance as a solution of red blood cells haemolyses.

How a cell behaves when suspended in a solution depends on whether or not the molecules or ions in the solution can cross the **plasma membrane**. If red blood cells (RBCs) are suspended in a concentrated solution of molecules that can

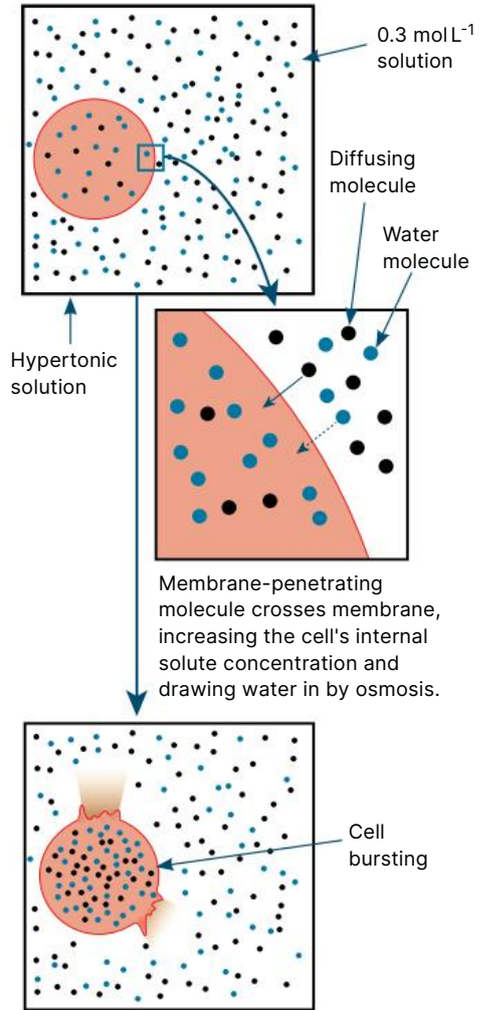
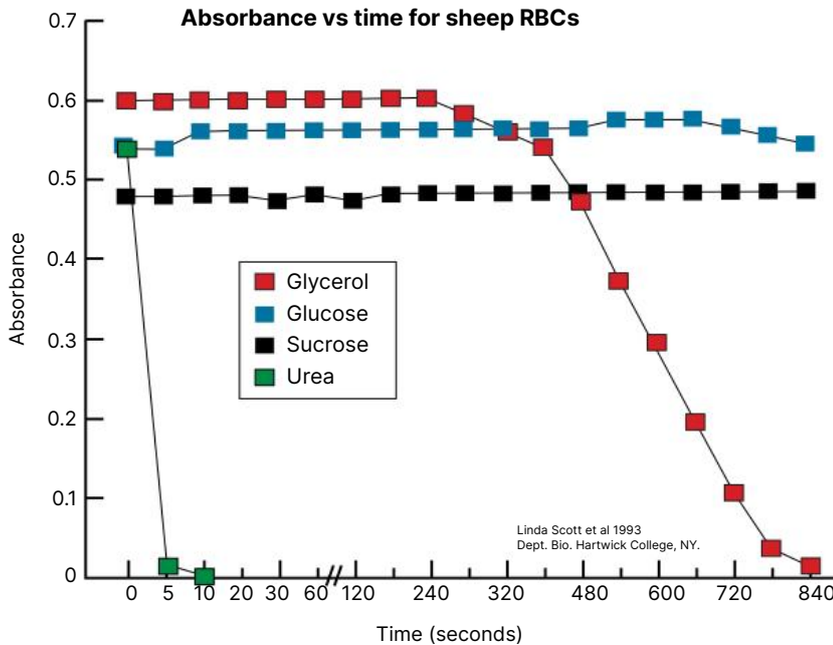
readily diffuse across the membrane, the molecules will enter the RBCs by diffusing down their **concentration gradient**. This will draw water into the RBCs (by **osmosis**) and burst (haemolyse). When the RBCs burst, the cellular material settles out of suspension and the solution becomes clear. By using a spectrophotometer to measure the rate at which the solution becomes clear, it is possible to determine the rate at which the molecules are crossing the plasma membrane.

## The aim

To investigate how the size and membrane solubility of molecules affects the rate of **diffusion** across the plasma membrane.

## The method

- ▶ 0.3 molL<sup>-1</sup> solutions of glucose, sucrose, urea, and glycerol were prepared (this concentration is greater than the cell internal concentration). A blank solution of distilled water was also prepared. Molecular weights (MW) are as follows: glucose (MW 180), sucrose (MW 342), urea (MW 60), and glycerol (MW 92).
- ▶ Both urea and glycerol readily diffuse across the plasma membrane. Glucose is transported across the membrane by a carrier protein.
- ▶ 3 mL of each solution was mixed with 0.1 mL of a sheep RBC suspension and added to cuvettes. The cuvettes were placed into a spectrophotometer and absorbance measured over 15 minutes. The results are plotted below:



Increased cellular volume causes the cell to burst (haemolysis). The cellular material settles out.

1. (a) Which molecule crosses the membrane the fastest? \_\_\_\_\_  
 (b) Which molecule appears to be unable to cross the plasma membrane? \_\_\_\_\_  
 (c) List the molecules in order of their ability to cross the plasma membrane (fastest to slowest):  
 \_\_\_\_\_
2. (a) What is the largest molecule used in the experiment? \_\_\_\_\_  
 (b) What is the smallest molecule used in the experiment? \_\_\_\_\_  
 (c) How does size affect the rate at which molecules can cross the plasma membrane? \_\_\_\_\_
3. Why don't the RBCs in the glucose solution haemolyse even though glucose is transported across the membrane?  
 \_\_\_\_\_



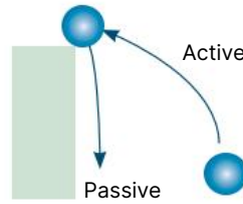
# Active Transport

**Key Idea:** Active transport uses energy to transport molecules against their concentration gradient across a partially permeable membrane.

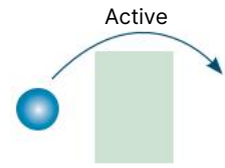
**Active transport** is the movement of molecules (or ions) from

regions of low concentration to regions of high concentration across a cellular membrane by a transport protein. Active transport needs energy to proceed because molecules are being moved against their **concentration gradient**.

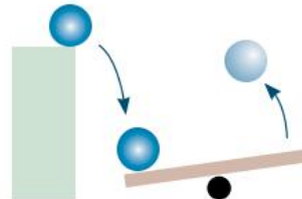
- ▶ The energy for active transport comes from ATP (adenosine triphosphate). Energy is released when ATP is hydrolysed (water is added) forming ADP (adenosine diphosphate) and inorganic phosphate (Pi).
- ▶ Transport (carrier) proteins in the membrane are used to actively transport molecules from one side of the membrane to the other (below).
- ▶ Active transport can be used to move molecules into and out of a cell.
- ▶ Active transport can be either primary or secondary. Primary active transport directly uses ATP for the energy to transport molecules. In secondary active transport, energy is stored in a concentration gradient. The transport of one molecule is coupled to the movement of another down its concentration gradient, ATP is not directly involved in the transport process.



A ball falling is a passive process (it requires no energy input). Replacing the ball requires active energy input.

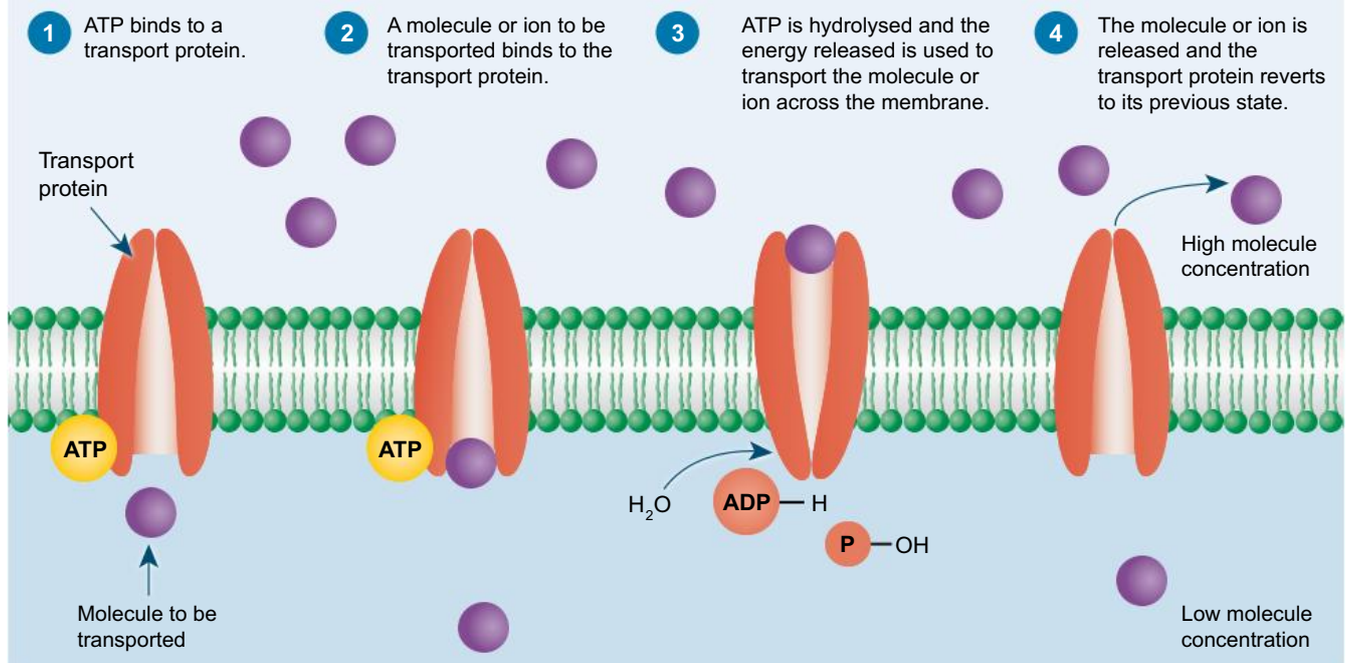


It requires energy to actively move an object across a physical barrier.



Sometimes the energy of a passively moving object can be used to actively move another. For example, a falling ball can be used to catapult another (left).

## Active transport



1. What is active transport? \_\_\_\_\_  
\_\_\_\_\_
2. Where does the energy for active transport come from? \_\_\_\_\_
3. What is the difference between primary active transport and secondary active transport?  
\_\_\_\_\_  
\_\_\_\_\_  
\_\_\_\_\_  
\_\_\_\_\_

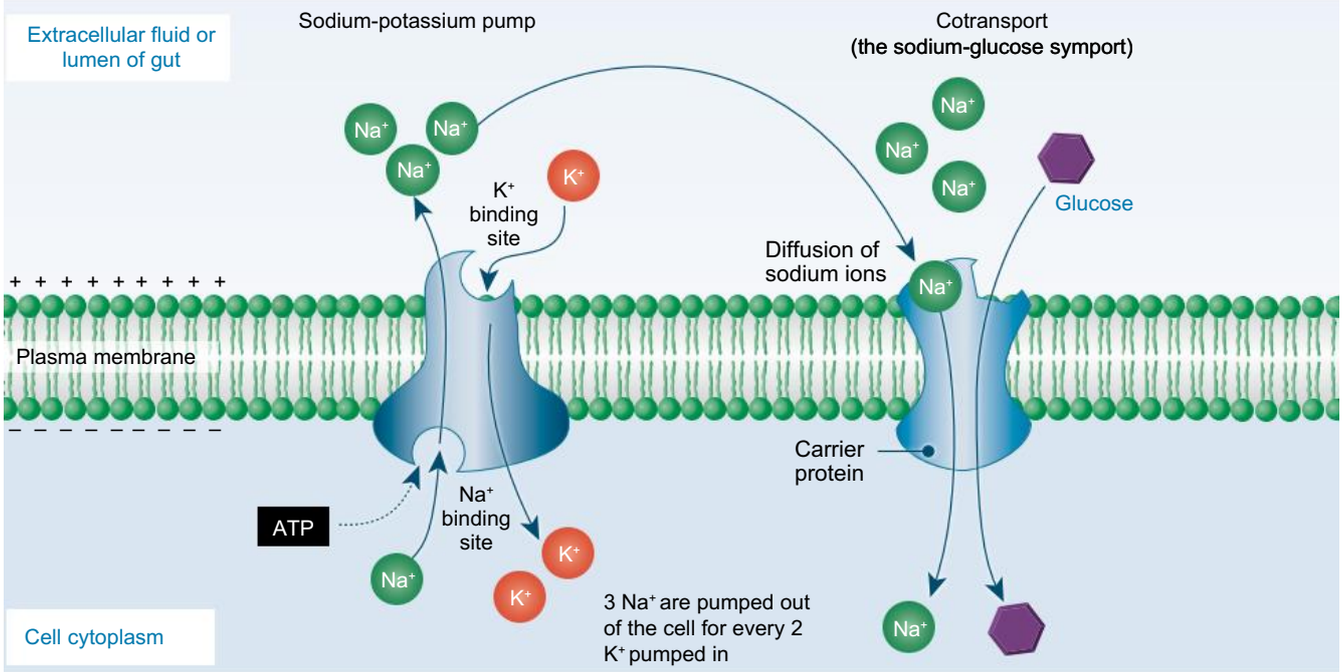


# 71 Ion Pumps and Cotransport

**Key Idea:** Ion pumps are transmembrane proteins that use energy to move ions and molecules across a membrane against their concentration gradient.

Sometimes molecules or **ions** are needed in concentrations that **diffusion** alone cannot supply to the cell, or they cannot diffuse across the **plasma membrane**. In this case **ion pumps** move ions (and some molecules) across the plasma

membrane. Proton pumps move  $H^+$  against a **concentration gradient** to create a potential difference across the membrane that can be used to do work. The sodium-potassium pump is found in almost all animal cells and is also common in plant cells. The concentration gradient created by ion pumps is often coupled to the transport of other molecules such as glucose across the membrane (below right).



### Sodium-potassium (Na<sup>+</sup>/K<sup>+</sup>) pump

The Na<sup>+</sup>/K<sup>+</sup> pump is a protein in the membrane that uses energy in the form of ATP to exchange sodium ions (Na<sup>+</sup>) for potassium ions (K<sup>+</sup>) across the membrane. The unequal balance of Na<sup>+</sup> and K<sup>+</sup> across the membrane creates large concentration gradients that can be used to drive transport of other substances (e.g. cotransport of glucose). The Na<sup>+</sup>/K<sup>+</sup> pump also helps to maintain the right balance of ions and so helps regulate the cell's water balance.

### Cotransport (coupled transport)

A gradient in sodium ions drives the active transport of glucose into intestinal epithelial cells. The specific transport protein couples the return of Na<sup>+</sup> down its concentration gradient to the transport of glucose into the intestinal epithelial cell across the cell membrane in contact with the gut lumen. Glucose diffuses from the epithelial cells across the opposite surface and is transported away in the blood. A low intracellular concentration of Na<sup>+</sup> (and therefore the concentration gradient) is maintained by a sodium-potassium pump.

- Why is ATP required for membrane pump systems to operate? \_\_\_\_\_
- (a) Explain what is meant by cotransport: \_\_\_\_\_
- (b) How is cotransport used to move glucose into the intestinal epithelial cells? \_\_\_\_\_
- (c) What happens to the glucose that is transported into the intestinal epithelial cells? \_\_\_\_\_
- (a) Does the sodium-potassium pump uses primary or secondary active transport? \_\_\_\_\_
- (b) Does the sodium-glucose symport uses primary or secondary active transport? \_\_\_\_\_
- (c) Describe one consequence of the extracellular accumulation of sodium ions: \_\_\_\_\_

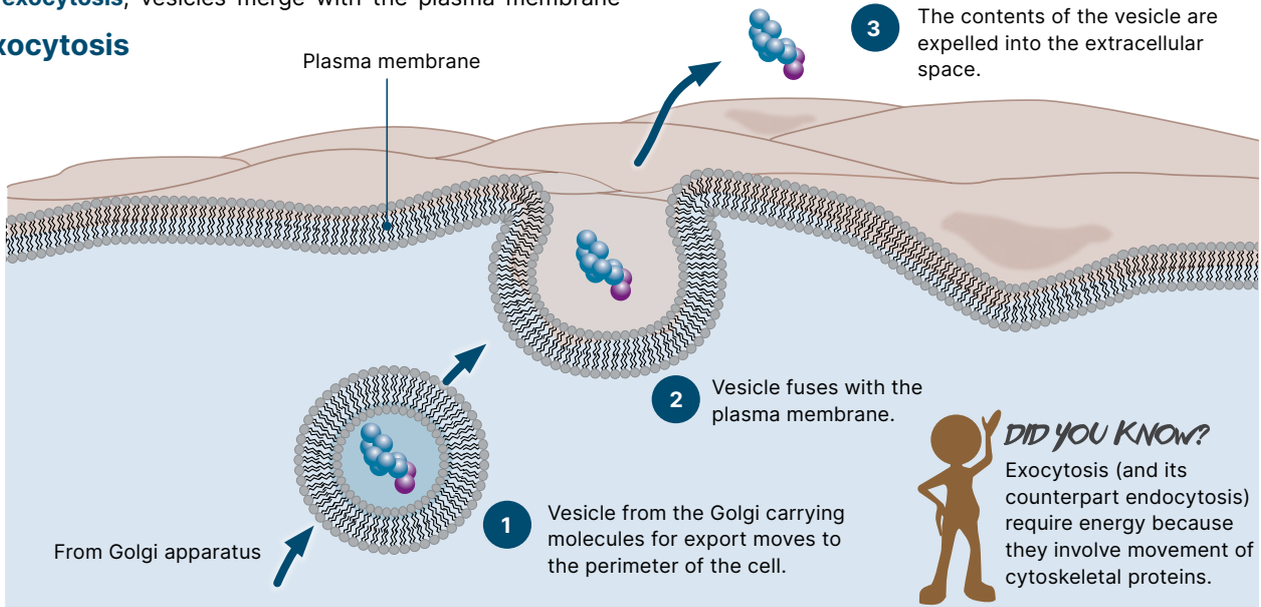
72

# Cytosis

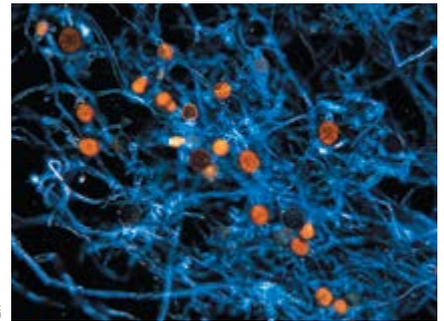
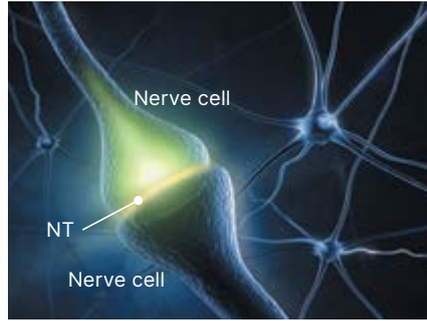
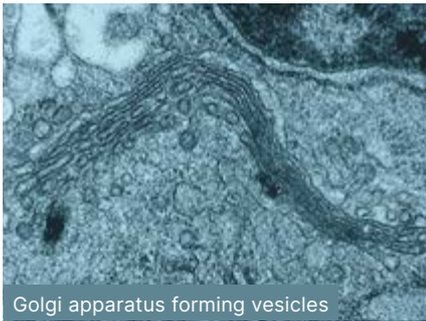
**Key Idea:** Cytosis is an energy demanding (active) transport mechanism involving the folding of the plasma membrane. Cytosis is an active process involving the **plasma membrane**. In **exocytosis**, vesicles merge with the plasma membrane

to export material from the cell. **Endocytosis** is a general term for engulfing of material by infolding of the plasma membrane.

## Exocytosis



**Exocytosis** (above) is an **active transport** process in which a secretory vesicle fuses with the plasma membrane and expels its contents into the extracellular space. In multicellular organisms, various types of cells (e.g. endocrine cells and nerve cells) are specialised to manufacture products, such as proteins, and then export them from the cell to elsewhere in the body or outside it.



The transport of Golgi vesicles to the edge of the cell and their expulsion from the cell occurs through the activity of the cytoskeleton. This requires energy (ATP).

Exocytosis is important in the transport of neurotransmitters (NT) into the junction (synapse) between nerve cells to transmit nervous signals.

Fungi and bacteria use exocytosis to secrete digestive enzymes, which break down substances extracellularly so that nutrients can be absorbed (by endocytosis).

- (a) What is the purpose of exocytosis? \_\_\_\_\_

\_\_\_\_\_

(b) How does it occur? \_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_
- Describe two examples of the role of exocytosis in cells:

(a) \_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_

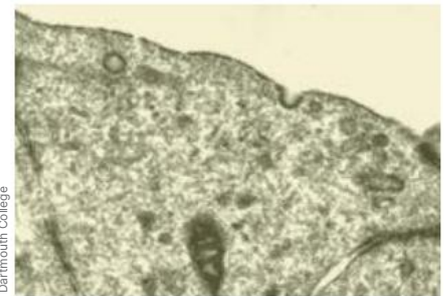
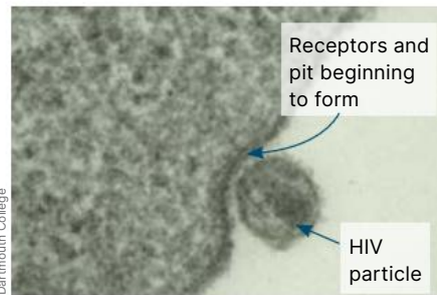
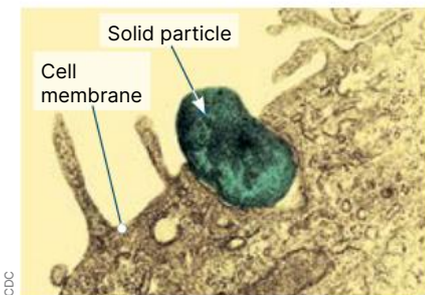
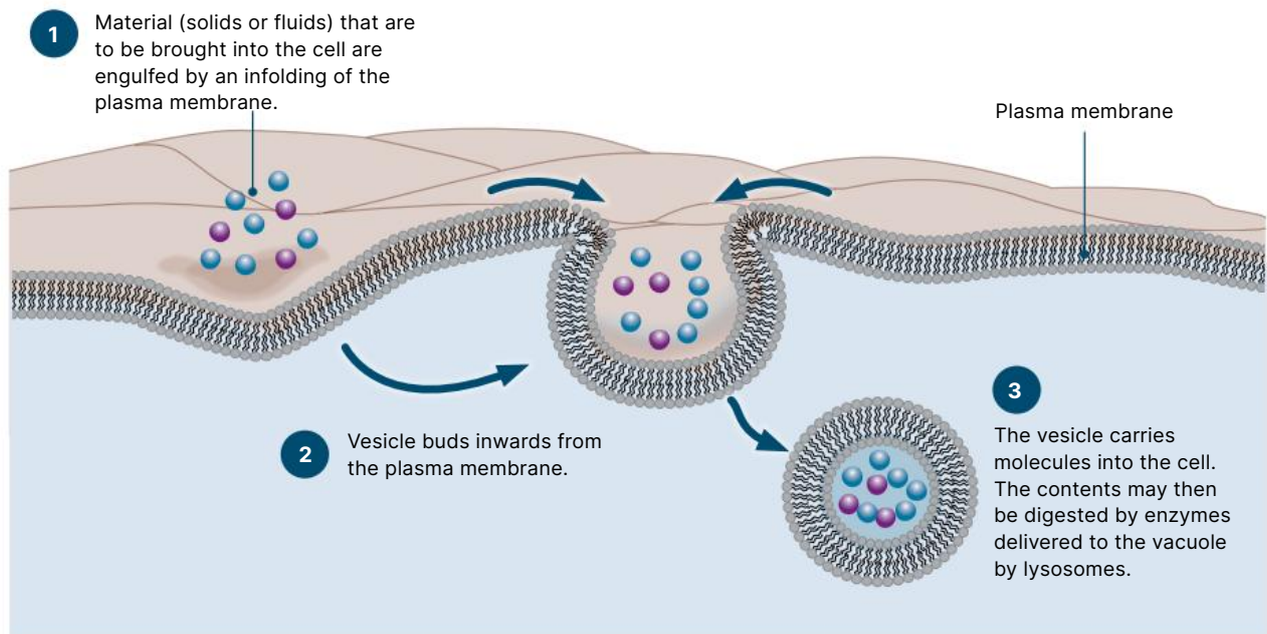
(b) \_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_

## Endocytosis

**Endocytosis** is a type of active transport in which the plasma membrane folds around a substance to transport it across the plasma membrane into the cell. The ability of cells to do this is a function of the fluid nature of the plasma membrane.



**Phagocytosis** (or 'cell-eating') involves the cell engulfing solid material to form large phagosomes or vacuoles (e.g. food vacuoles). It may be non-specific or receptor-mediated. Examples: Feeding in *Amoeba*, phagocytosis of foreign material and cell debris by neutrophils and macrophages.

**Receptor mediated endocytosis** is triggered when certain metabolites, hormones, or viral particles bind to specific receptor proteins on the membrane so that the material can be engulfed. Examples: The uptake of lipoproteins by mammalian cells and endocytosis of viruses (above).

**Pinocytosis** (or 'cell-drinking') involves the non-specific uptake of liquids or fine suspensions into the cell to form small pinocytic vesicles. Pinocytosis is used primarily for absorbing extracellular fluid. Examples: Uptake in many protozoa, some cells of the liver, and some plant cells.

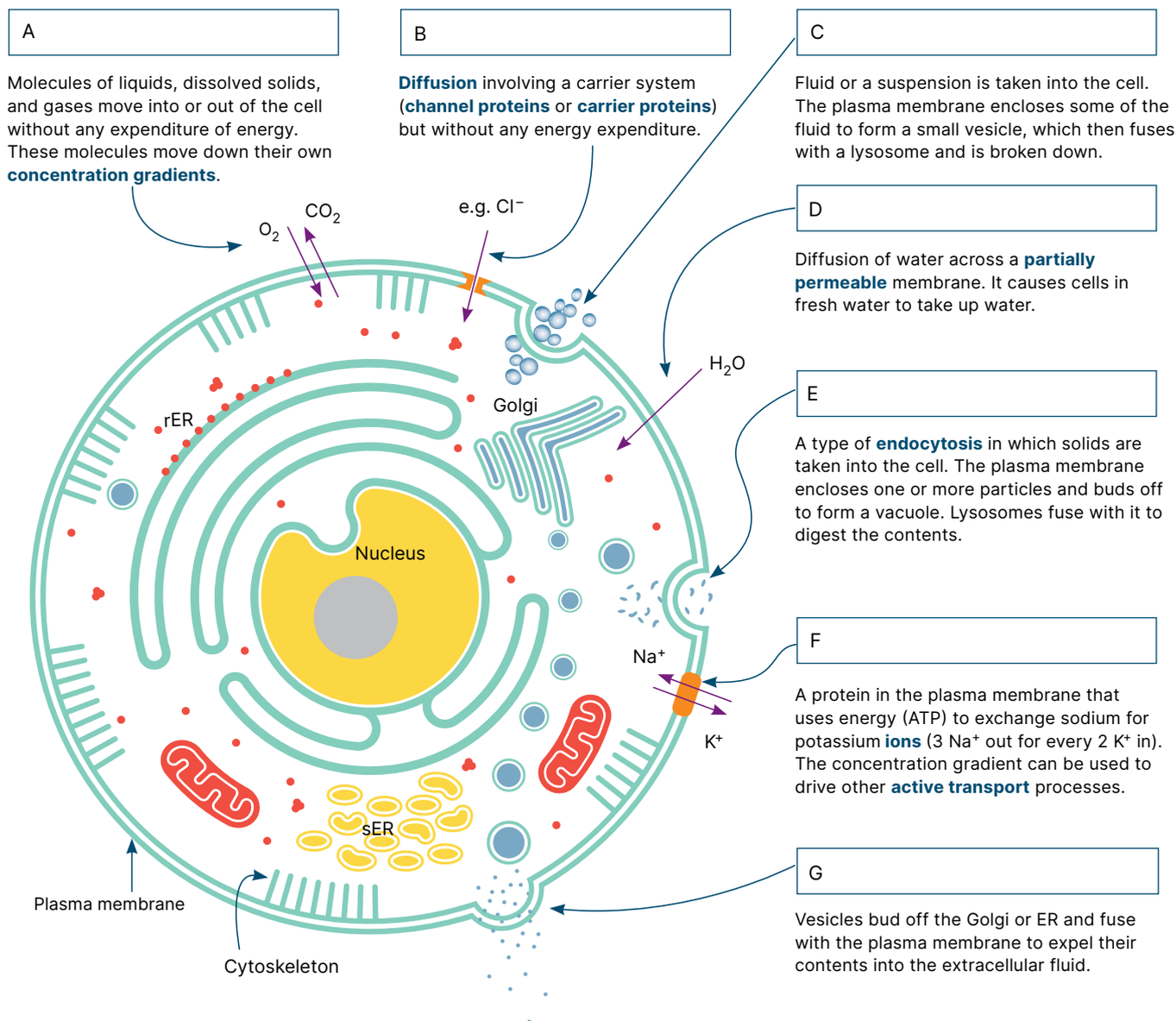
3. What is the purpose of endocytosis? \_\_\_\_\_
4. Is endocytosis active or passive transport? \_\_\_\_\_
5. Describe the following types of endocytosis:
  - (a) Phagocytosis: \_\_\_\_\_
  - (b) Receptor-mediated endocytosis: \_\_\_\_\_
  - (c) Pinocytosis: \_\_\_\_\_
6. Explain how the plasma membrane can form a vesicle: \_\_\_\_\_

## 73

## Active and Passive Transport Summary

**Key Idea:** Cells move materials into and out of the cell by either passive transport, which does not use energy, or by active transport which requires energy, usually as ATP.

The diagram below summarises the movement of material in and out of a cell. Use the information to complete the activity.



- Identify each of the processes (A-G) described in the diagram above in the spaces provided. Indicate whether the transport process is active or passive by using **A** for active and **P** for passive.
- Identify the transport mechanism involved in each of the following processes in cells:
  - Uptake of extracellular fluid by liver cells: \_\_\_\_\_
  - Capture and destruction of a bacterial cell by a white blood cell: \_\_\_\_\_
  - Movement of water into the cell: \_\_\_\_\_
  - Secretion of digestive enzymes from cells of the pancreas: \_\_\_\_\_
  - Synthesis of ATP via membrane-bound ATP synthase: \_\_\_\_\_
- In general terms describe the energy requirements of passive and active transport: \_\_\_\_\_

# 74

## Did You Get It?

1. Explain how the properties of the phospholipid molecule result in the bilayer structure of membranes:

---



---



---

2. Using the formulae: cuboid SA = 2(lh + lw + hw), cuboid volume = lwh, calculate the surface area to volume ratio of the following cell shapes:

(a) A cubic cell 6 μm x 6 μm x 6 μm: \_\_\_\_\_

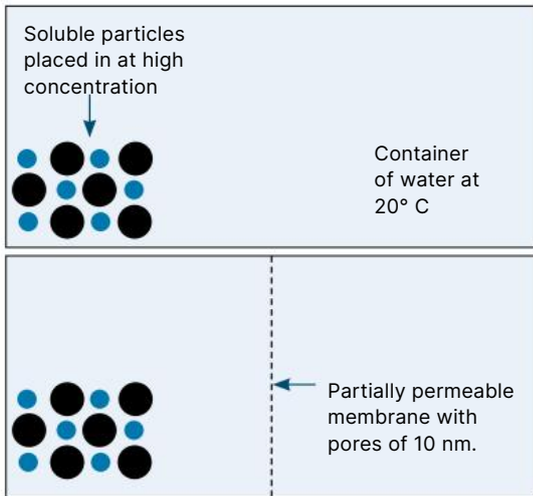
(b) A cuboid cell 1 μm x 12 μm x 5 μm: \_\_\_\_\_

(c) Which of these cells would exchange substances with its environment most efficiently and why: \_\_\_\_\_

---

3. Consider the two diagrams below. For each, draw in the appropriate box what you would expect to see after one hour.

● Particle with diameter of 5 nm    ● Particle with diameter of 20 nm



After one hour:

---

4. Describe the type of movement occurring at A, B, and C across the cell membrane:

---



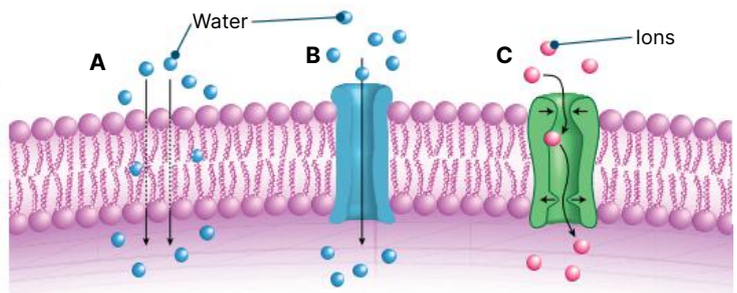
---



---



---



5. Compare the type of cell membrane transport in Q 4. with that occurring in a (sodium-potassium) ion pump:

---



---



---



---







# Exchange of Nutrients and Wastes

## Key Terms

- absorption
- ammonia
- amylase
- blood vessels
- Bowman's capsule
- capillaries
- circulatory system
- collecting duct
- digestion
- distal convoluted tubule
- enzyme
- excretion
- glomerulus
- kidney
- large intestine
- lipase
- Loop of Henle
- microvilli
- nephron
- organ trafficking
- protease
- proximal convoluted tubule
- small intestine
- stomach
- transplantation (organ)
- urea
- uric acid
- urine

## Key Concepts

- ▶ The mammalian circulatory system is responsible for the transport of oxygen and nutrients around the body.
- ▶ The human digestive system extracts nutrients from food and allows for the elimination of waste material.
- ▶ Nitrogenous waste is removed by the filtration system of the kidneys.

## Digestion, absorption, and circulation

### Activity Number

□ 1	Describe the basic structure and function of carbohydrates, proteins, and lipid biomolecules.	76
□ 2	Explain how structural features of exchange surfaces in the capillaries of the mammalian circulatory systems allow for efficient nutrient exchange and absorption.	77-79, 82-83
□ 3	Describe the basic structure and organisation of the circulatory system in a mammal, e.g. human. Explain how closed circulatory systems in mammals facilitate the efficient transport of materials (such as respiratory gases and nutrition) to and from all cells in the body.	77, 80
□ 4	Describe the basic structure and organisation of the digestive tract in a mammal, e.g. human, including the cells, tissues, and organs making up the different regions.	81, 84
□ 5	Identify the characteristics of the absorptive surfaces within the digestive system, e.g. the small intestine.	82
□ 6	Describe the role of enzymes in the extracellular chemical digestion of ingested food. Describe the source, substrate, products, and optimum pH for a range of digestive enzymes, including amylase, protease, and lipase.	82-83
□ 7	Describe how the intestinal villi and the structure of the intestinal epithelial cells themselves increase the surface area for the digestion and absorption of nutrients.	82-83
□ 8	<b>SI:</b> Investigate the effect of temperature and pH on the rate of reaction of amylase enzyme activity.	85

## Excretion of nitrogenous wastes

□ 9	Describe the different types of nitrogenous wastes produced by the breakdown of proteins.	86
□ 10	Describe the overall structure of the urinary system including kidneys, ureters, bladder, and urethra. Outline the structure and function of the mammalian kidney including the nephron and its associated capillary network.	87-88
□ 11	Describe in detail the function of each of the regions of the nephron in the production of urine. Include reference to the glomerulus, Bowman's capsule, proximal convoluted tubule, loop of Henle, distal convoluted tubule, and collecting duct.	89
□ 12	Explain urine formation and excretion of wastes by glomerular filtration (ultrafiltration) and selective reabsorption and secretion across the nephron membranes. Explain how the urine is concentrated.	89
□ 13	Discuss how the increased demand for transplant organs has led to illegal trafficking of organs and tissue, forced donation, and transplant tourism. Describe ethical concerns associated with these practices.	90
□ 14	<b>SHE:</b> Discuss how understanding of the human body system has advanced technology for tissue and organ transplants to treat disease.	90

# Carbohydrates, Proteins, and Lipids

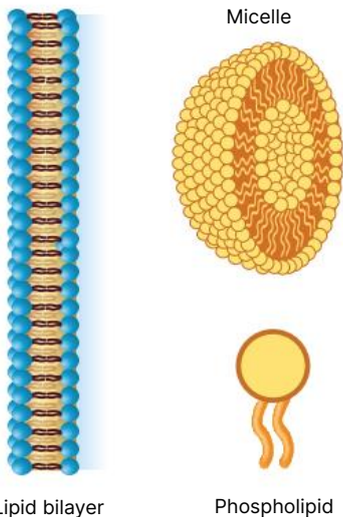
**Key Idea:** Carbohydrates are produced by plants through photosynthesis to store energy. The glucose biomolecule is subsequently transformed into more complex carbohydrates, as well as a wide range of proteins, and lipids.

Carbon has the unique ability to bond with a wide range of other elements, such as carbon, hydrogen, oxygen, and nitrogen, resulting in the formation of numerous carbon-based (organic) molecules. These organic molecules, known as biomolecules, are the building blocks of living organisms and can be broadly classified into three main categories:

carbohydrates, lipids, and proteins. The majority of organic molecules initially enter the food chain as uncomplicated glucose carbohydrates produced by photosynthetic organisms. These molecules undergo a sequence of cellular reactions, resulting in a variety of carbohydrates, proteins, and lipids distinguished by their complexity, size, and elemental composition. Animals consume these compounds as food in the process of nutrition, breaking them down into smaller components for utilization during **digestion**, with unneeded materials excreted as waste from the body.

## Lipids

Lipids, which are fatty compounds, have diverse functions within the body. Specifically, they are highlighted as integral components of cell membranes. In this role, lipids play a crucial part in regulating the passage of substances into and out of cells, contributing significantly to the overall functionality and integrity of cellular structures. Fats are a type of lipid that can be stored in the body and used as a ready source of energy.

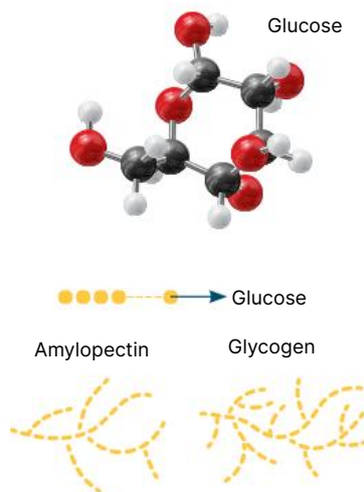


**Food examples:** Vegetable oils, oily fish, butter, nuts, avocados, olives.

Broken down by **lipase** enzymes

## Carbohydrates

Carbohydrates are categorised into four main groups based on their polymerisation level, which refers to how many sugar units are joined together. These groups are monosaccharides (single sugar units), disaccharides (two sugar units), oligosaccharides (a few sugar units), and polysaccharides (many sugar units). This classification system helps differentiate the different types of carbohydrates based on their structural complexity and functions in biological systems.

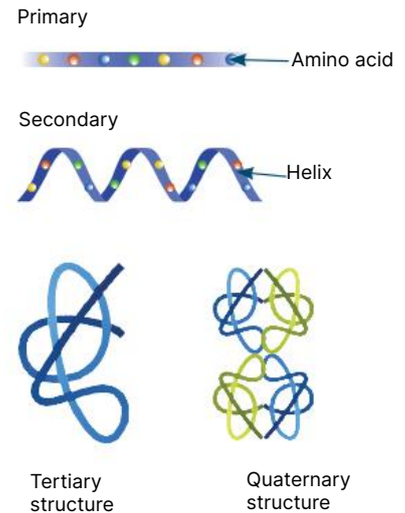


**Food examples:** Cereals, fruits, sugars, pasta, potatoes, bread.

Broken down by **amalyse** enzymes

## Proteins

Protein structures are constructed through a process where amino acids join together through condensation reactions to form peptide bonds. The specific order in which these amino acids are arranged in a protein is referred to as its primary structure. This primary structure is fundamental in determining the overall shape, function, and properties of the protein, showcasing the importance of the sequence of amino acids in defining the characteristics of a protein.



**Food examples:** meats, shellfish, dairy products, nuts, beans, eggs.

Broken down by **protease** enzymes

1. What are the four main groups into which carbohydrates are categorised based on their polymerisation level?

---



---

2. How are protein structures constructed?

---



---

3. What is the key role of lipids in the body?

---



---

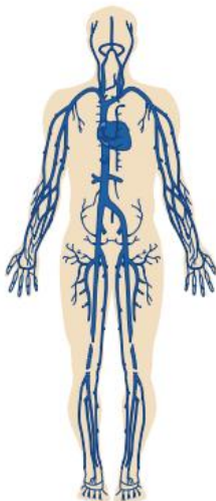
# The Mammalian Circulatory System

**Key Idea:** The mammalian circulatory system is a double circuit made up of a pulmonary circuit and a systemic circuit. The **blood vessels** of the **circulatory system** form a network of tubes that carry blood away from the heart, transport it to the tissues of the body, and then return it to the heart. The figure below shows a number of the basic circulatory routes. Mammals have a double circulatory system: a pulmonary

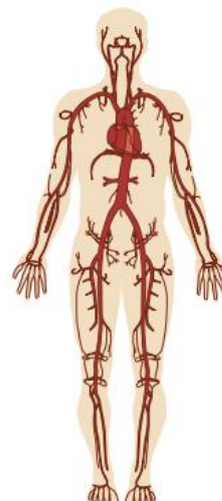
system, which carries blood between the heart and lungs, and a systemic system, which carries blood between the heart and the rest of the body. The systemic circulation has many subdivisions. Two important subdivisions are the coronary (cardiac) circulation, which supplies the heart muscle, and the hepatic portal circulation, which runs from the gut to the liver.

## Schematic overview of the human circulatory system

Deoxygenated blood (coloured blue below) travels to the right side of the heart via the vena cavae. The heart pumps the deoxygenated blood to the lungs where it releases carbon dioxide and receives oxygen. The oxygenated blood (coloured white below) travels via the pulmonary vein back to the heart from where it is pumped to all parts of the body. The venous system (figure, left) returns blood from the **capillaries** to the heart. The arterial system (figure right) carries blood from the heart to the capillaries. Portal systems carry blood between two capillary beds.



Venous system



Arterial system

**Pulmonary vein**  
Carries oxygenated blood back to the heart.

**Superior vena cava**  
Receives deoxygenated blood from the head and body.

**Right atrium**  
Receives deoxygenated blood via the superior and inferior vena cavae.

**Right ventricle**  
Pumps deoxygenated blood to the lungs.

**Inferior vena cava**  
Receives deoxygenated blood from the lower body and organs.

**Hepatic vein**  
Carries deoxygenated blood from the liver.

**Hepatic portal vein**  
Carries deoxygenated, nutrient rich blood from the gut for processing.

**Renal vein**  
Carries deoxygenated blood from the kidneys.

**Aorta**  
Carries oxygenated blood to the body. Anteriorly, it branches to form the carotid arteries supplying the head and neck.

**Pulmonary artery**  
Carries deoxygenated blood to the lungs.

**Left atrium**  
Receives oxygenated blood from the lungs.

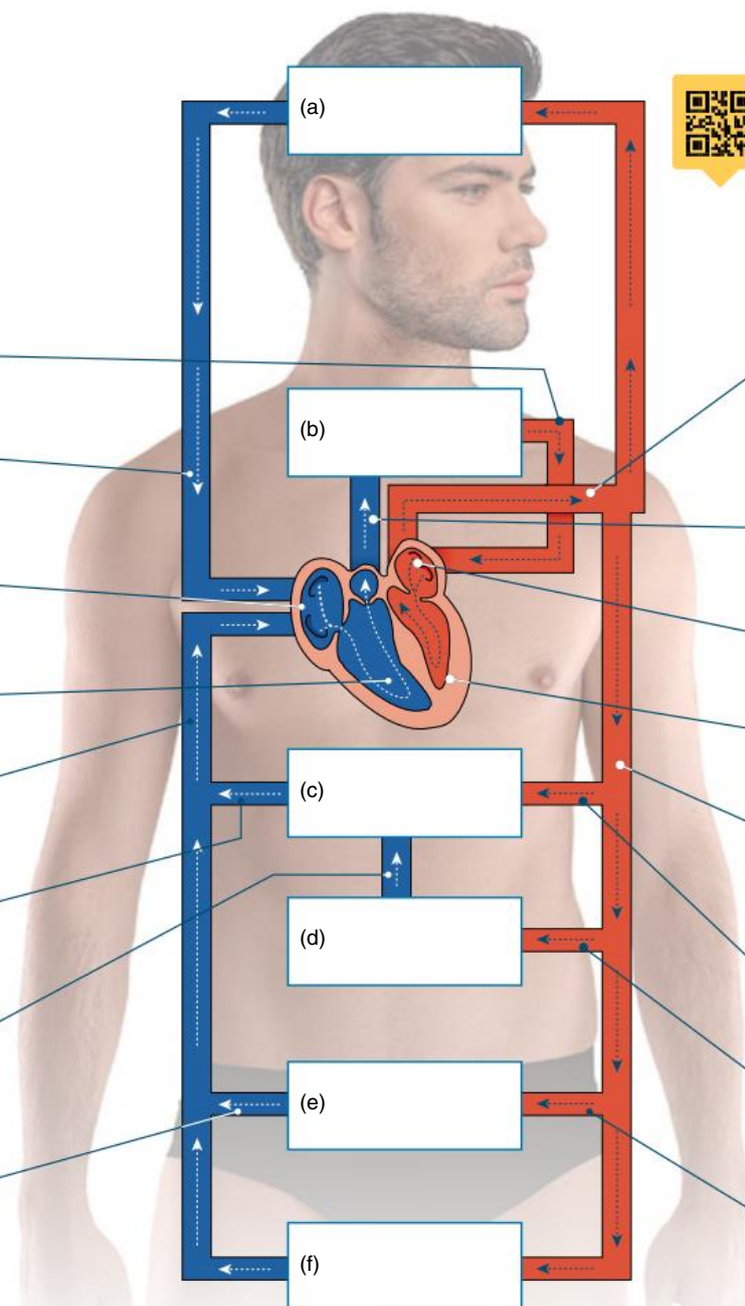
**Left ventricle**  
Pumps blood from the left atrium to the aorta.

**Abdominal aorta**  
Parallel to the inferior vena cava, branching to supply the organs of the abdominal cavity.

**Hepatic artery**  
Carries oxygenated blood to the liver.

**Mesenteric artery**  
Carries oxygenated blood to the gut.

**Renal artery**  
Carries oxygenated blood to the kidneys.



1. Complete the diagram above by labelling the boxes with the correct organs: *lungs, liver, head, intestines, genitals/lower body, kidneys.*

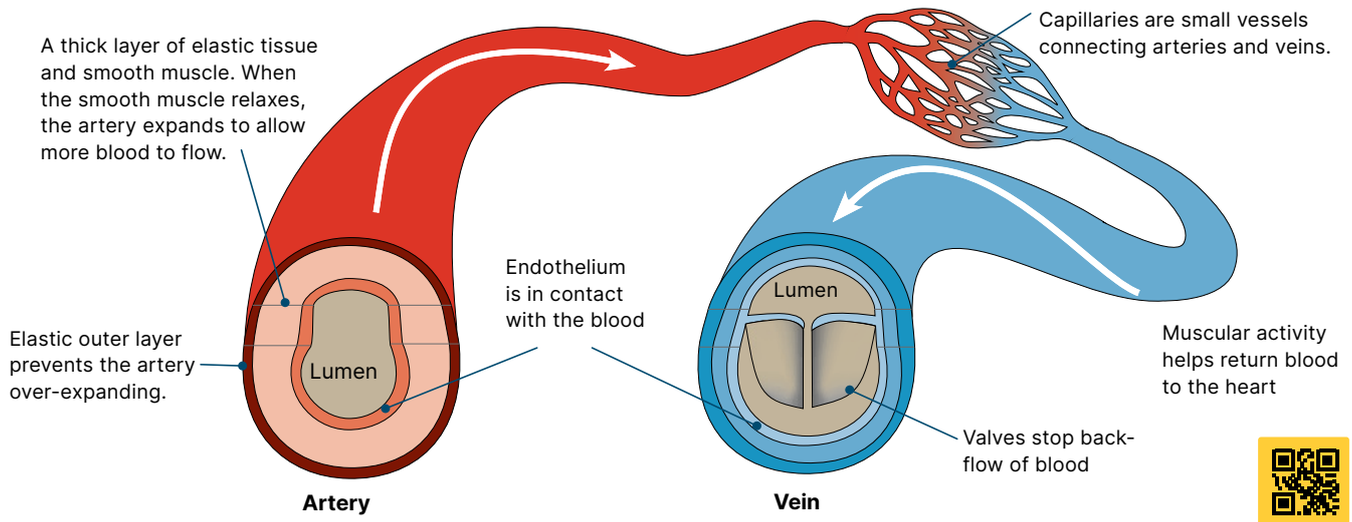
2. List the two blood vessels involved in the pulmonary circuit: \_\_\_\_\_



**Key Idea:** The blood vessels of the circulatory system connect the body's cells to the organs that exchange gases, absorb nutrients, and dispose of wastes.

In vertebrates, arteries are the **blood vessels** that carry blood away from the heart to the **capillaries** within the tissues. The large arteries that leave the heart divide into medium-sized (distributing) arteries. Within the tissues and organs, these distributing arteries branch to form arterioles, which deliver blood to capillaries. Blood flow to the tissues is altered by contraction (vasoconstriction) or relaxation (vasodilation)

of the blood vessel walls. Vasoconstriction increases blood pressure whereas vasodilation has the opposite effect. Veins are the blood vessels that return blood to the heart from the tissues. The smallest veins (venules) return blood from the capillaries to the veins. Veins and their branches contain about 59% of the blood in the body. The structural differences between veins and arteries are mainly associated with differences in the relative thickness of the vessel layers and the diameter of the lumen (space within the vessel). These, in turn, are related to the vessel's functional role.



## Arteries

- ▶ Arteries have an elastic, stretchy structure that enables them to withstand and maintain the high pressure of blood being pumped from the heart. At the same time, their ability to contract (a feature of the central muscle layer) helps regulate blood flow and pressure.
- ▶ Arteries nearer the heart have more elastic tissue to resist the higher pressures of the blood leaving the left ventricle. Arteries further from the heart have more muscle to help them maintain blood pressure.
- ▶ Between heartbeats, the elastic walls of the artery recoil, maintaining an even pressure despite the pulsing nature of blood flow.

## Veins

- ▶ Veins are made up of the same three layers as arteries but they have less elastic and muscle tissue, a relatively thicker external layer, and a larger, less defined lumen.
- ▶ Although veins are less elastic than arteries, they can still expand enough to adapt to changes in the pressure and volume of the blood passing through them. Blood flowing in the veins has lost a lot of pressure because it has passed through the narrow capillaries.
- ▶ The lower pressure flow means that many veins, especially those in the limbs, have valves to prevent backflow of the blood as it returns to the heart.

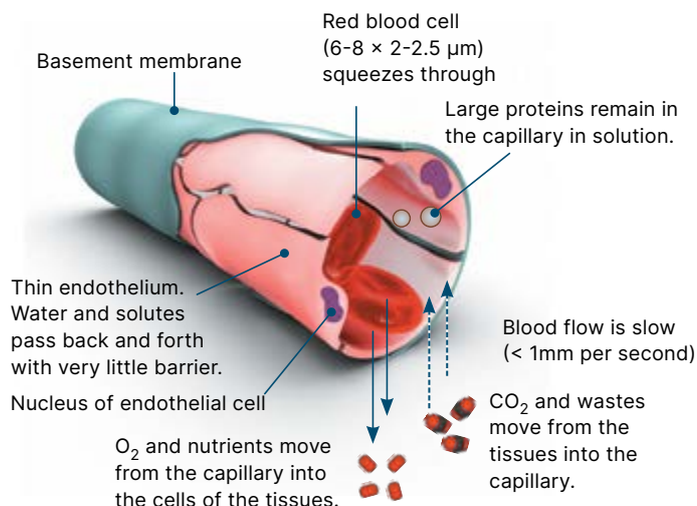
1. What is the function of blood vessels? \_\_\_\_\_  
\_\_\_\_\_
2. Why do the artery walls need to be thick with a lot of elastic tissue? \_\_\_\_\_  
\_\_\_\_\_
3. What is the role of valves in assisting the veins to return blood back to the heart? \_\_\_\_\_  
\_\_\_\_\_
4. How do the structural differences between arteries and veins relate to their functional roles? \_\_\_\_\_  
\_\_\_\_\_



# Capillaries and Capillary Networks

**Key Idea:** Capillaries are small, thin-walled vessels that allow the exchange of material between the blood and the tissues. In vertebrates, **capillaries** are very small **blood vessels** that connect arterial and venous circulation and allow efficient

exchange of nutrients and wastes between the blood and tissues. Capillaries form networks or beds and are abundant where metabolic rates are high. Fluid that leaks out of the capillaries has an essential role in bathing the tissues.



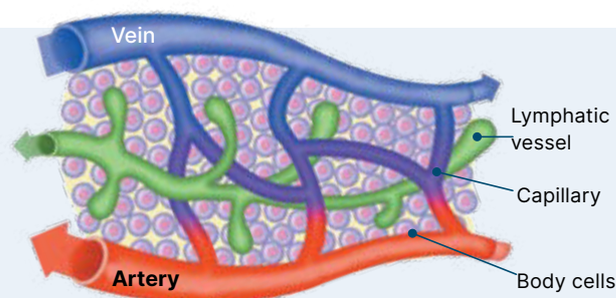
## Exchanges in capillaries

- ▶ Blood passes from the arterioles into the capillaries where the exchange of materials between the body cells and the blood takes place. Capillaries have a diameter of just 5-10 μm. The only tissue present is an endothelium of squamous epithelial cells. Capillaries are so numerous that no cell is more than 25 μm from any capillary.
- ▶ Blood pressure causes fluid to leak from capillaries through small gaps where the endothelial cells join. This fluid bathes the tissues, supplying nutrients and oxygen, and removing wastes (left).
- ▶ The density of capillaries in a tissue is an indication of that tissue's metabolic activity. For example, cardiac muscle has a high demand for blood flow and is well supplied with capillaries. Smooth muscle is far less active than cardiac muscle and does not need such an extensive blood supply.



## Blood, tissue fluid, and lymph

	Blood	Tissue fluid	Lymph
<b>Cells</b>	Red blood cells, white blood cells, platelets	Some white blood cells	White blood cells
<b>Proteins</b>	Hormones and plasma proteins	Some hormones and proteins	None
<b>Glucose</b>	High	None	Low
<b>Amino acids</b>	High	Used by body cells	Low
<b>Oxygen</b>	High	Used by body cells	Low
<b>Carbon dioxide</b>	Low	Produced by body cells	High



The fluid that leaks from the capillaries is called tissue fluid. Some of it returns to the blood at the venous end of the capillary bed, but some is drained by lymph vessels to form lymph.

- ▶ Blood transports nutrients, wastes, and respiratory gases to and from the tissues.
- ▶ Tissue fluid facilitates the transport of these between the blood and the tissues.
- ▶ Lymph drains excess tissue fluid and returns it to the general circulation. It has a role in the immune system.

1. What is the role of capillaries? \_\_\_\_\_

2. (a) Describe the structure of a capillary: \_\_\_\_\_

---



---



---

(b) Explain how the structure and position of capillaries (relative to the body's cells) is important in allowing the exchange of materials:

---



---



---



The flow of blood through a capillary bed is called microcirculation. In most parts of the body, there are two types of vessels in a capillary bed: the true capillaries, where exchanges take place, and a vessel called a vascular shunt, which connects the arteriole and venule at either end of the bed. The shunt diverts blood past the true capillaries when the metabolic demands of the tissue are low. When tissue activity increases, the entire network fills with blood.

3. Describe the structure of a capillary network:

---

---

---

---

---

---

---

---

---

---

4. Explain the role of the smooth muscle sphincters and the vascular shunt in a capillary network:

---

---

---

---

---

---

---

---

---

---

5. (a) Describe a situation where the capillary bed would be in the condition labelled A:

---

---

---

---

---

---

---

---

---

---

(b) Describe a situation where the capillary bed would be in the condition labelled B:

---

---

---

---

---

---

---

---

---

---

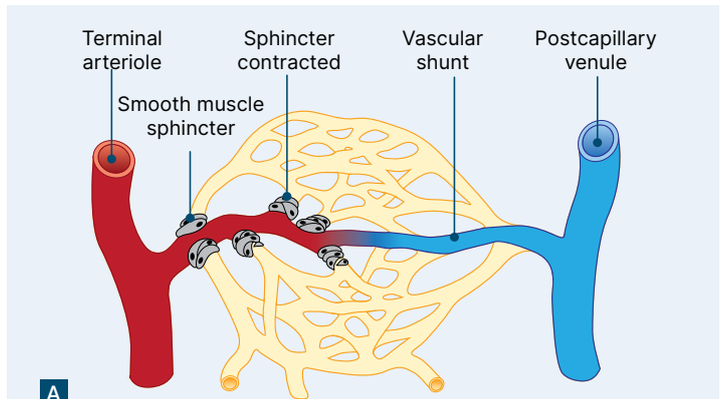
6. On the photograph, right, identify:

A: \_\_\_\_\_

B: \_\_\_\_\_

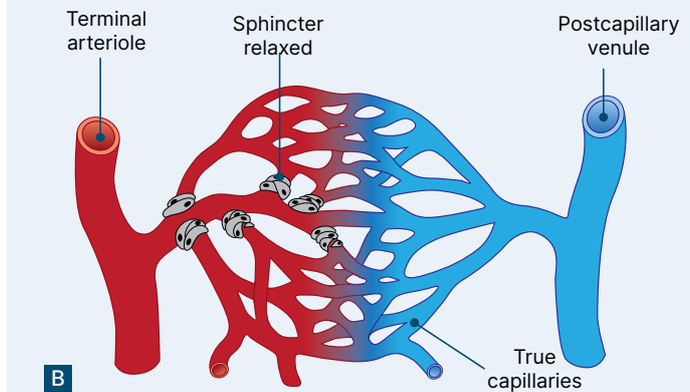
C: \_\_\_\_\_

D: \_\_\_\_\_



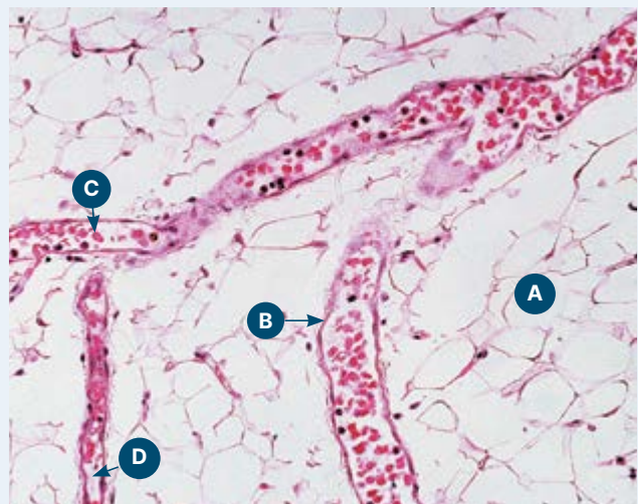
**A**

When the sphincters contract (close), blood is diverted via the vascular shunt to the postcapillary venule, bypassing the exchange capillaries.



**B**

When the sphincters are relaxed (open), blood flows through the entire capillary bed allowing exchanges with the cells of the surrounding tissue.



Capillaries supply all the tissues of the body, creating extensive networks to supply cells with the nutrients and oxygen they need and remove carbon dioxide and other metabolic wastes. This capillary is moving through fat tissue.

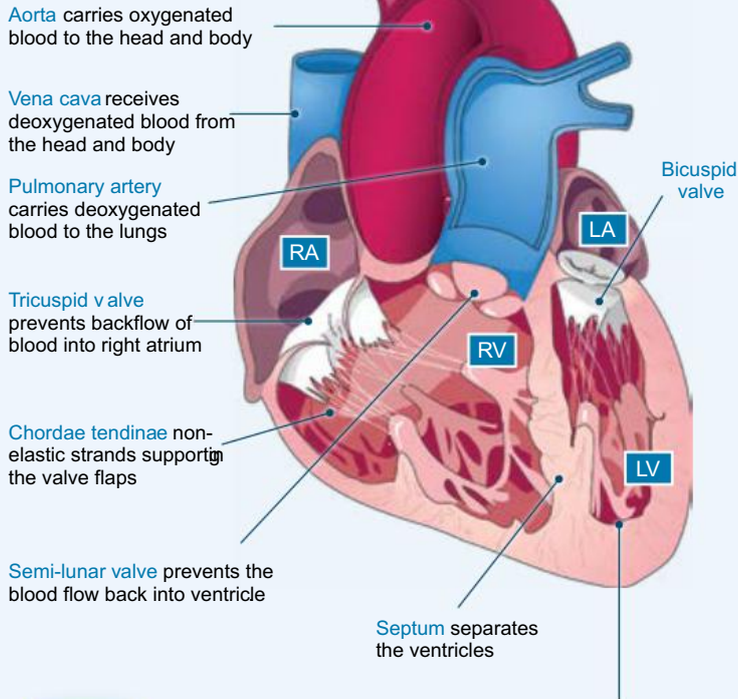
# Structure of the Mammalian Heart

**Key Idea:** Humans have a four chambered heart divided into left and right halves. It acts as a double pump. The heart is the centre of the human cardiovascular system. It is a hollow, muscular organ made up of four chambers (two atria and two ventricles) that alternately fill and empty of blood, acting as a double pump. The left side (systemic

circuit) pumps blood to the body tissues and the right side (pulmonary circuit) pumps blood to the lungs. The heart lies between the lungs, to the left of the midline, and is surrounded by a double layered pericardium of connective tissue, which prevents over distension of the heart and anchors it within the central compartment of the thoracic cavity.

## Human heart structure

(sectioned, anterior view)

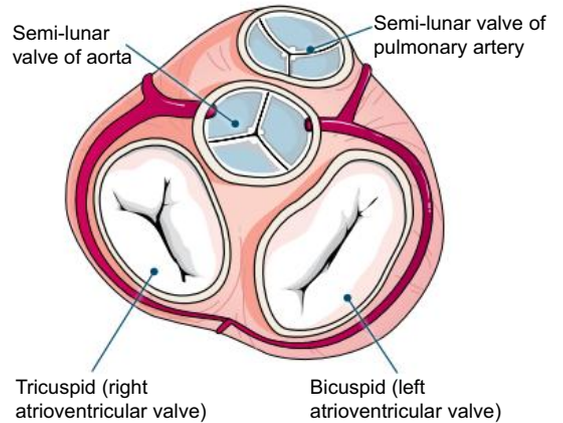


The heart is not a symmetrical organ. Although the quantity of blood pumped by each side is the same, the walls of the left ventricle are thicker and more muscular than those of the right ventricle. The difference affects the shape of the ventricular cavities, so the right ventricle is twisted over the left.

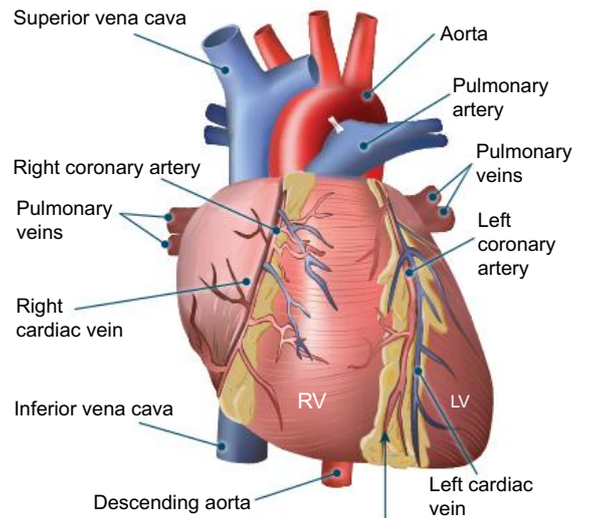
### Key to abbreviations

- RA** Right atrium: receives deoxygenated blood via the anterior and posterior vena cava
- RV** Right ventricle: pumps deoxygenated blood to the lungs via the pulmonary artery
- LA** Left atrium: receives blood returning to the heart from the lungs via the pulmonary veins
- LV** Left ventricle: pumps oxygenated blood to the head and body via the aorta

## Top view of a heart in section, showing valves

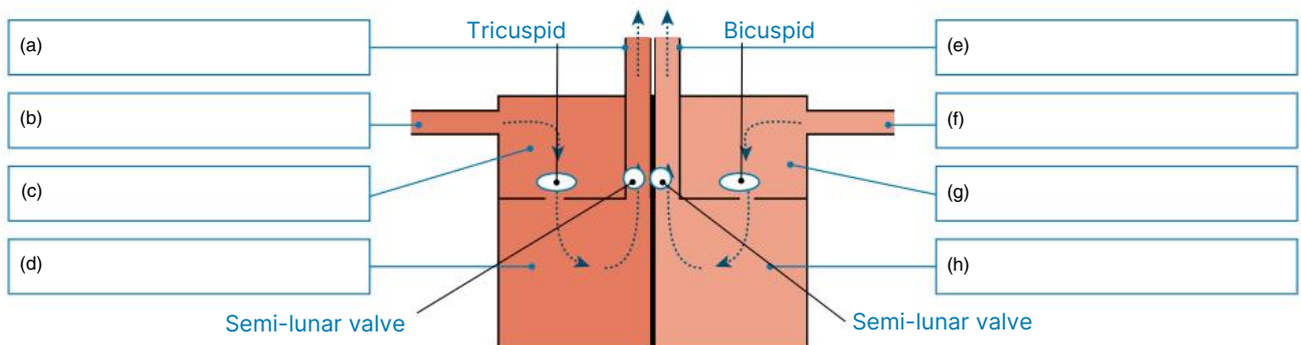


## Anterior view of heart to show coronary arteries



**Coronary arteries:** The high oxygen demands of the heart muscles are met by a dense capillary network. Coronary arteries arise from the aorta and spread over the surface of the heart supplying the cardiac muscle with oxygenated blood. Deoxygenated blood is collected by cardiac veins and returned to the right atrium via a large coronary sinus.

- In the schematic diagram of the heart, below, label the four chambers and the main vessels entering and leaving them. The arrows indicate the direction of blood flow. The four heart valves are labelled.



# 81 The Digestive System

**Key Idea:** The digestive tract is specialised to maximise the digestion of food, absorption of nutrients, and elimination of undigested material.

The human digestive system (gut) is a tubular tract, which is regionally specialised into a complex series of organs and glands. These work in sequence to maximise the efficiency with which food is processed. Collectively, the organs of

the digestive tract carry out the physical and chemical breakdown (**digestion**) of food, **absorption** of nutrients, and elimination of undigested material. The gut is a hollow, open-ended, muscular tube, and the food within it is essentially outside the body, having contact only with the cells lining the tract. Several accessory organs and glands lie external to the digestive tract. These secrete enzyme-rich fluids to the food to aid digestion.

**Salivary glands**

produce lubricating secretions with  $\alpha$ -**amylase**, which begins starch digestion

**Oesophagus**

**Gall bladder**

**Pancreas**

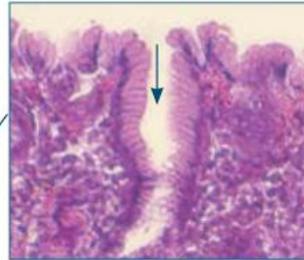
**Liver**

**Stomach**

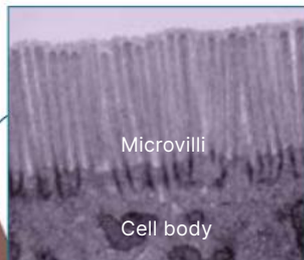
**Large intestine**

**Small intestine**

Gastric gland

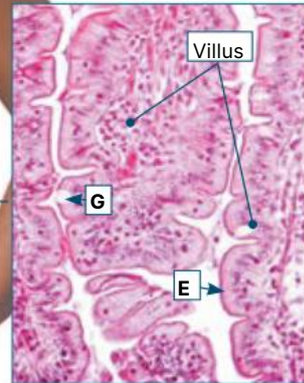


In the **stomach**, gastric glands contain parietal cells, which produce hydrochloric acid, and chief cells, which produce a protein-digesting enzyme. Scattered endocrine cells secrete a hormone to regulate gastric activity.



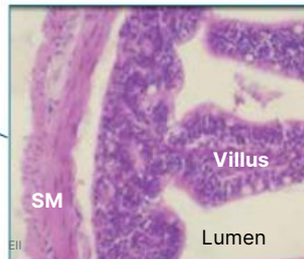
© Lisa Howard, Katherine Connolly Dartmouth College

Cells lining the walls on the **small intestine** (the intestinal epithelium) have microscopic extensions of the plasma membrane called **microvilli**. These form a brush border that increases the surface area for absorption of food molecules. Under lower power microscopy, it appears as a fuzzy edge.



Nephron

In the small intestine, the intestinal epithelial cells (E) and mucus-producing goblet cells (G) make up the epithelium lining the gut wall. The wall is folded into finger like projections called villi (*sing.* villus). These further increase the surface area of the intestine.



The entire gastrointestinal tract is supported by underlying connective tissue. Two layers of smooth muscle (SM), one running lengthwise and one running around the gut, encircle the tube, contracting in waves to move food through the gut. This process is called peristalsis.



- (a) How are villi formed? \_\_\_\_\_

\_\_\_\_\_

(b) What is the purpose of microvilli? \_\_\_\_\_

\_\_\_\_\_
- What is the purpose of the smooth muscle surrounding the intestine? \_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_



# The Stomach and Small Intestine

**Key Idea:** The stomach produces acid and a protein-digesting enzyme, which break food down into a slurry, called chyme. The **stomach** is a hollow, muscular organ between the oesophagus and **small intestine**. In the stomach, food is mixed in an acidic environment to produce a semi-fluid mixture called chyme. The low pH of the stomach destroys

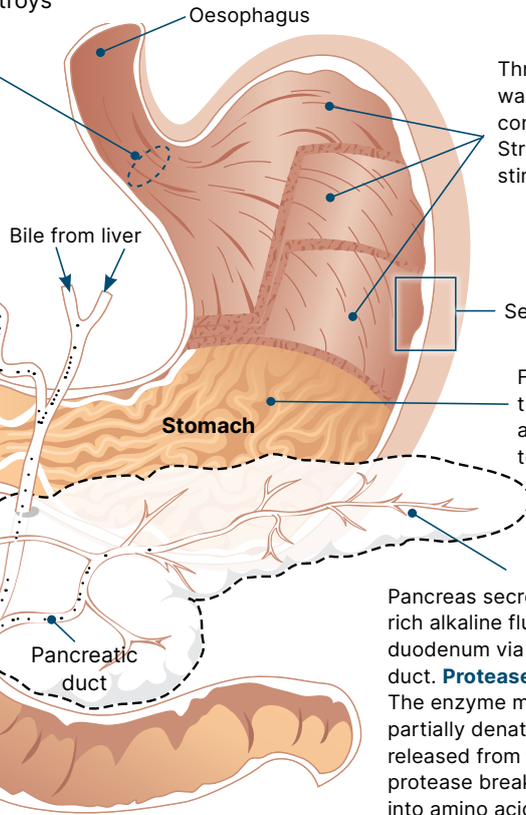
microbes, denatures proteins, and activates a protein-digesting **enzyme** precursor. There is very little absorption in the stomach, although small molecules (glucose, alcohol) are absorbed across the stomach wall into the surrounding **blood vessels**.



Cardiac sphincter (closes the junction between oesophagus and stomach). Prevents food moving back up oesophagus.

Three layered muscular wall mixes the stomach contents to produce chyme. Stretching the stomach wall stimulates gastric secretion.

The gall bladder stores bile, which is produced by the liver cells. Fat and acid in the duodenum stimulate release of bile from the gall bladder.



The stomach, pancreas, and mouth release the enzyme **lipase**. This enzyme works in combination with bile from the liver to break down fats and lipids.

Folds (rugae) in the stomach wall allow the stomach to expand to 1 L.

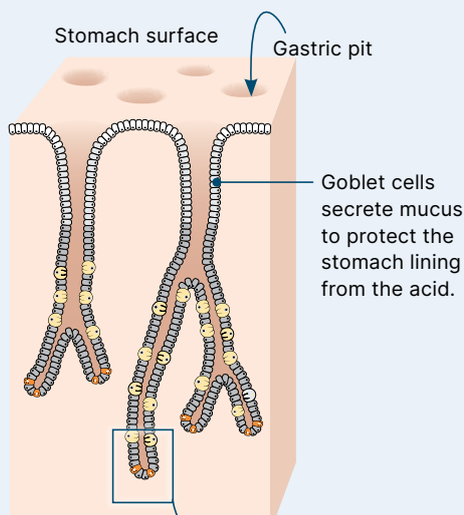
Pyloric sphincter (closes junction between stomach and duodenum).



Pancreas secretes an enzyme-rich alkaline fluid into the duodenum via the pancreatic duct. **Protease** is a key enzyme. The enzyme mixes in with partially denatured protein released from the stomach. The protease breaks the protein into amino acids, making them available for use in the body.

**Amylase** is also produced by the pancreas and secreted into the duodenum. This enzyme breaks down carbohydrates.

## Detail of a gastric gland (stomach wall)

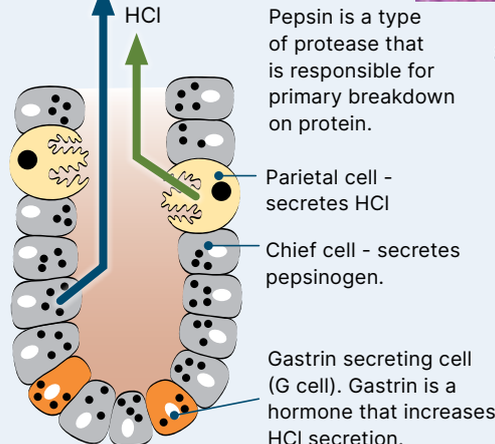


Right: High powered light micrograph of the stomach epithelium showing the gastric glands.



Pepsinogen (activated by HCl) → Pepsin

Pepsin is a type of protease that is responsible for primary breakdown on protein.



Parietal cell - secretes HCl

Chief cell - secretes pepsinogen.

Gastrin secreting cell (G cell). Gastrin is a hormone that increases HCl secretion.

### Stomach secretions

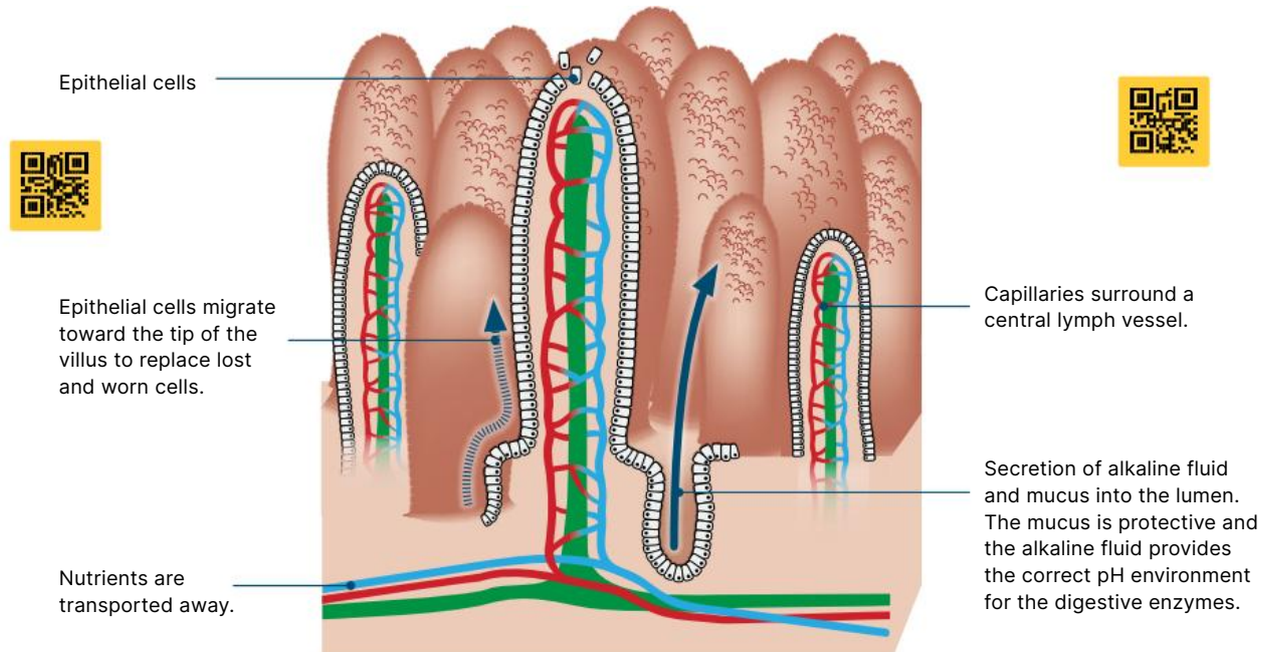
Gastric juice
Acid (HCl) secretion
Pepsin (optimal pH 1.5-2.0) Acts on proteins and breaks them down into peptides - short chains of amino acids (a protease)

In the stomach, gastric glands contain parietal cells, which produce hydrochloric acid, chief cells, which produce enzymes to break down protein, and endocrine cells.

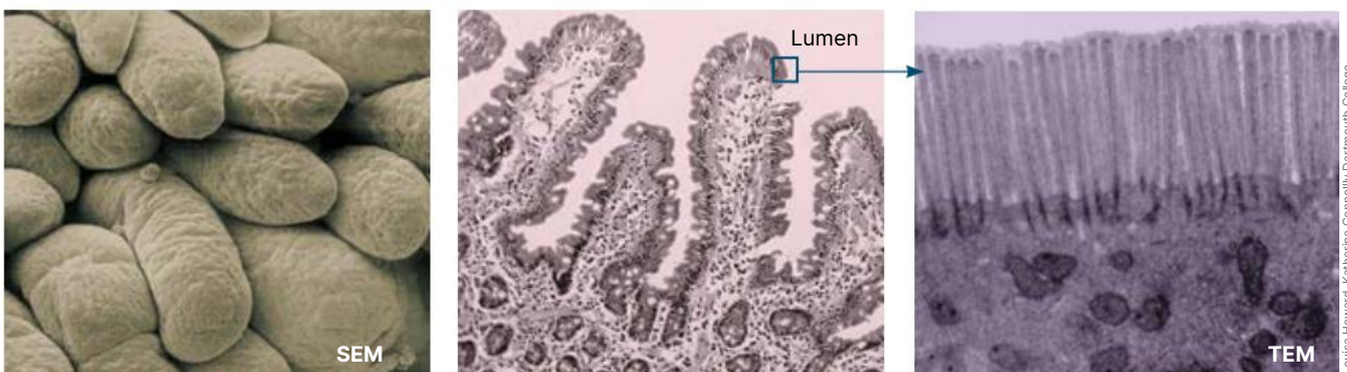


## The small intestine

- ▶ The small intestine receives the chyme directly from the stomach. It is divided into three regions, which are distinguished by the cell types present: the duodenum, where most chemical digestion occurs, and then the jejunum and the ileum. Most absorption occurs in the jejunum and ileum.
- ▶ The intestinal lining is folded into many intestinal villi, which project into the gut lumen (the space enclosed by the gut). The villi increase the surface area for nutrient absorption. The epithelial cells that make up the lining of each villus in turn have a brush-border of many **microvilli**, which are primarily responsible for nutrient absorption. The membrane of the microvilli is packed with enzymes that break down food molecules for absorption.
- ▶ Enzymes bound to the microvilli of the epithelial cells, and in the pancreatic and intestinal juices, break down fats, peptides, and carbohydrates (see tables below). The small molecules produced by this digestion are then absorbed into the underlying blood and lymph vessels.
- ▶ Tubular exocrine glands and goblet cells secrete alkaline fluid and mucus into the lumen, neutralizing the acidity of the chyme entering the small intestine from the stomach and protecting the lining of the intestine from damage.



**Photographs below:** The intestinal villi are shown projecting into the gut lumen in a scanning electron micrograph (left image) and in a light microscope image (centre image). The microvilli forming the brush border of a single intestinal epithelial cell are shown in the transmission electron micrograph (right image).



Enzymes in the small intestine break down food into small molecules that can be absorbed through the gut wall. Enzymes are present in the pancreatic juice added to the duodenum, in intestinal juice, and bound to the surfaces of the intestinal epithelial cells.

Enzymes in pancreatic juice		Enzymes in intestinal juice (IJ) and epithelium (E)	
Enzymes in duodenum (optimal pH)		Enzymes in small intestine (location, optimal pH)	
1. Pancreatic amylase (6.7-7.0)	1. Starch → maltose	1. Maltase (E, 6.0-6.5)	1. Maltose → glucose
2. Trypsin* (7.8-8.7)	2. Protein → peptides	2. Peptidases (proteases) (IJ, E, ~ 8.0)	2. Polypeptides → amino acids
3. Chymotrypsin* (7.8)	3. Protein → peptides	3. Sucrase (E, ~6.0)	3. Sucrose → fructose & glucose
4. Pancreatic lipase (8.0)	4. Fats → fatty acids & glycerol	4. Enteropeptidase (IJ 8.0)	4. Activates trypsin*
* secreted in an inactive form		*Once activated, trypsin activates chymotrypsin	

1. Summarise the structure and role of each of the following regions of the human digestive tract:

(a) Stomach: \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_

(b) Small intestine: \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_

2. (a) What is the purpose of the hydrochloric acid produced by the parietal cells of the stomach?

\_\_\_\_\_  
 \_\_\_\_\_

(b) Explain why protein-digesting enzymes (e.g. pepsin) are secreted in an inactive form and then activated after release:

\_\_\_\_\_  
 \_\_\_\_\_

3. Identify an endocrine cell in the stomach epithelium and state its purpose: \_\_\_\_\_

\_\_\_\_\_

4. How does the stomach achieve the mixing of acid and enzymes with food? \_\_\_\_\_

\_\_\_\_\_

5. (a) What is the purpose of the intestinal villi? \_\_\_\_\_

\_\_\_\_\_

(b) What is the purpose of the microvilli (brush border) on intestinal epithelial cells? \_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_

6. Identify two sites for secretion of enzymes active in the small intestine. Identify an enzyme produced there and its role:

(a) Site: \_\_\_\_\_ Enzyme: \_\_\_\_\_

Enzyme's role: \_\_\_\_\_

\_\_\_\_\_

(b) Site: \_\_\_\_\_ Enzyme: \_\_\_\_\_

Enzyme's role: \_\_\_\_\_

\_\_\_\_\_

(c) In general, do the enzymes act in acidic or alkaline conditions? \_\_\_\_\_

\_\_\_\_\_

(d) How is this pH environment generated? \_\_\_\_\_

\_\_\_\_\_

7. Suggest why the small intestine is so long: \_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_

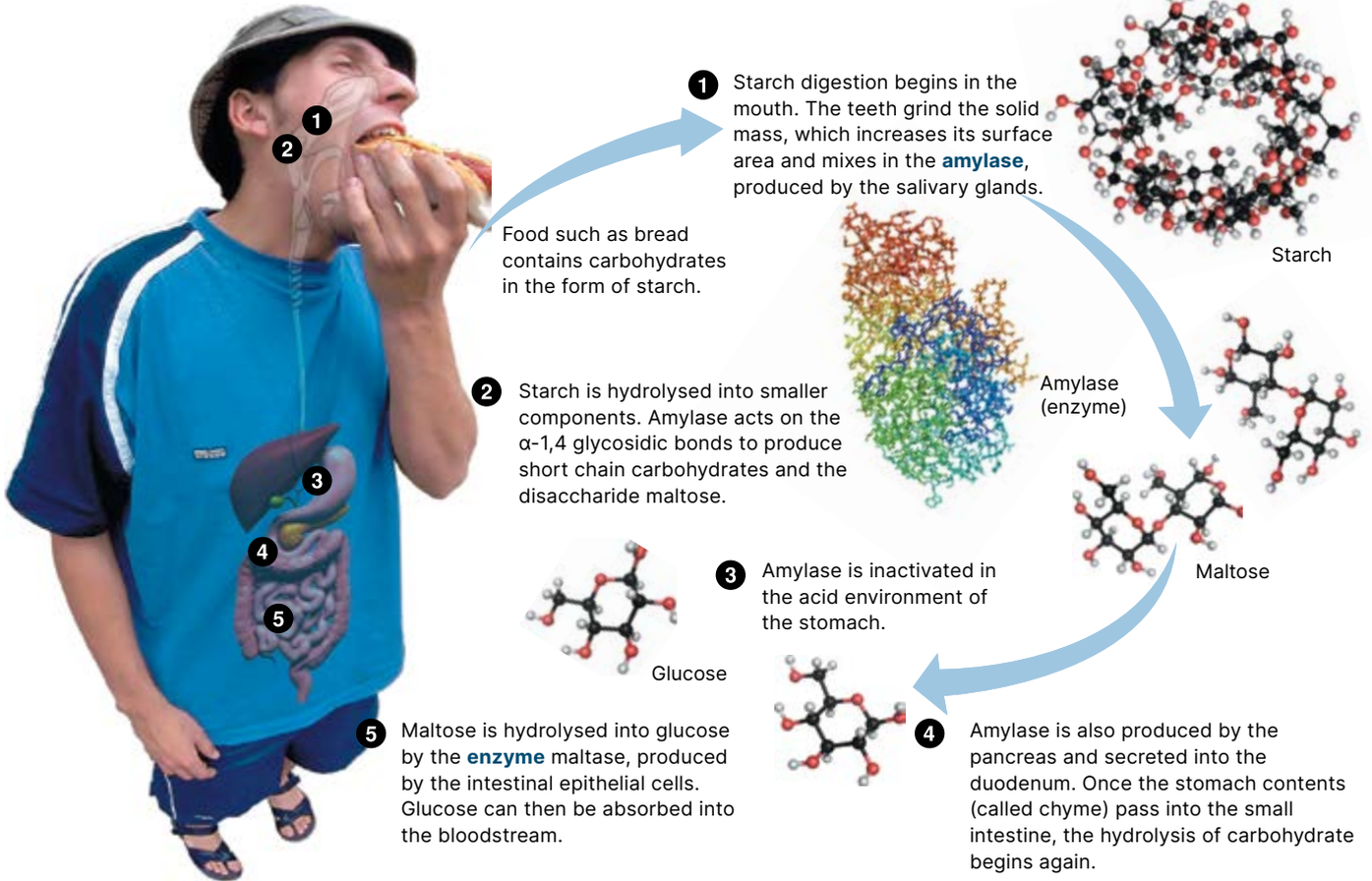
# Digestion, Absorption, and Transport

**Key Idea:** Food must be digested into components small enough to be absorbed by the body's cells and assimilated. Nutrient absorption involves both active and passive transport.

**Digestion** breaks down food molecules into small molecules that can pass through the intestinal lining into the underlying blood and lymph vessels. For example, starch is broken down first into maltose and short chain carbohydrates such as dextrose, before being hydrolysed to the simple sugar

glucose (below). Breakdown products of other foodstuffs include amino acids (from proteins), and fatty acids, glycerol, and acylglycerols (from fats). The passage of these molecules from the gut into the blood or lymph is called **absorption**. Nutrients are then transported directly or indirectly to the liver for storage or processing. After they have been absorbed nutrients can be assimilated, i.e incorporated into the substance of the body itself.

## Digestion of starch



1. Explain the roles of amylase and maltase in starch digestion: \_\_\_\_\_

---



---



---



---

2. Salivary and pancreatic secretions contain amylase. Why do two digestive organs produce the same enzyme? \_\_\_\_\_

---



---

3. Based on the diagram opposite, predict what would happen to nutrient absorption if the villi were damaged. Explain: \_\_\_\_\_

---



---

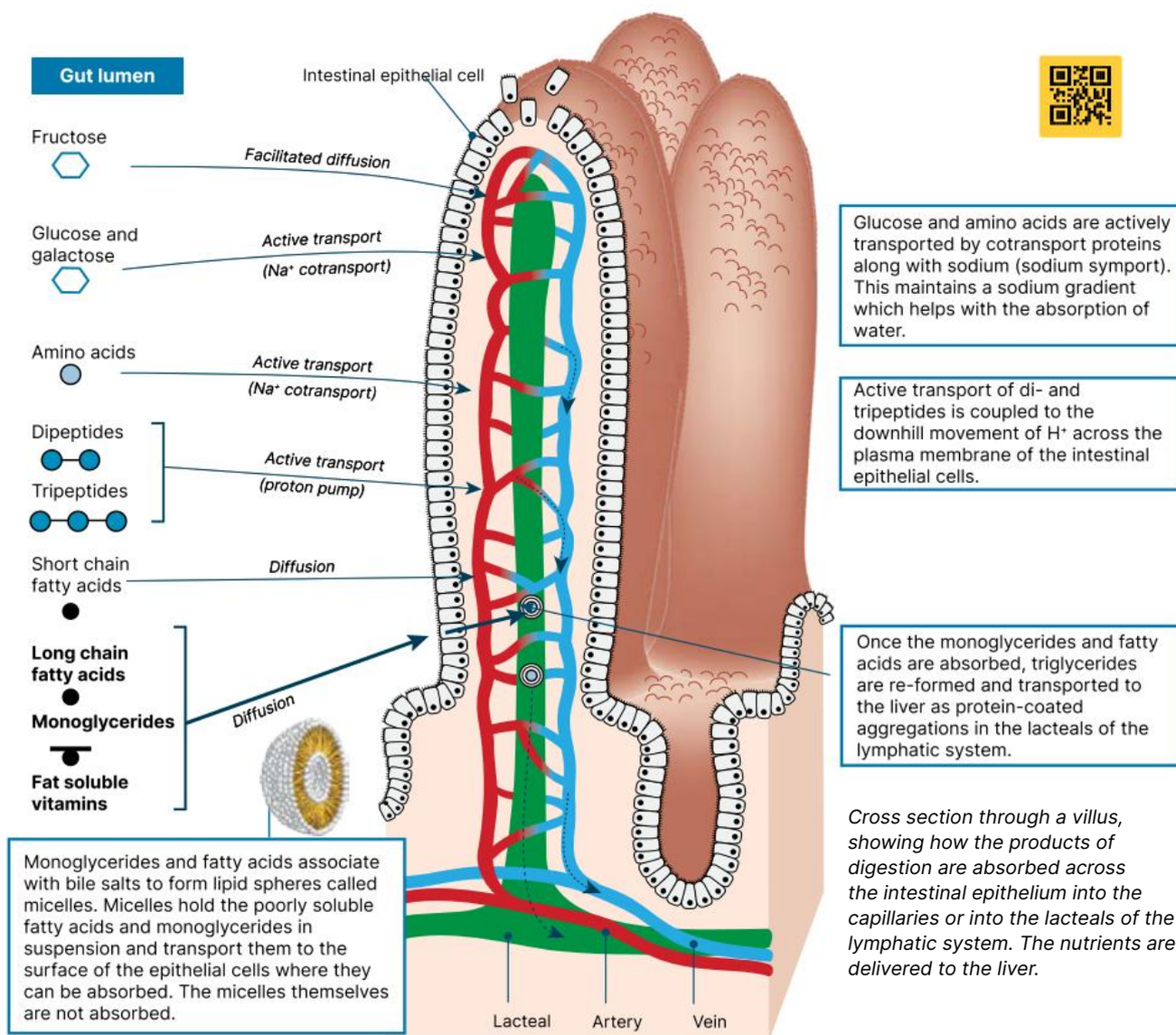


---



---

## Nutrient absorption by intestinal villi



- Describe how each of the following nutrients are absorbed by the intestinal villi:
  - Glucose: \_\_\_\_\_
  - Fructose: \_\_\_\_\_
  - Amino acids: \_\_\_\_\_
  - Di- and tripeptides: \_\_\_\_\_
- Describe the two purposes of the sodium symport in the intestinal epithelium: \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_
- What is the role of micelles in the absorption of lipids? \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_
- How are concentration gradients maintained for the absorption of nutrients by diffusion? \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_

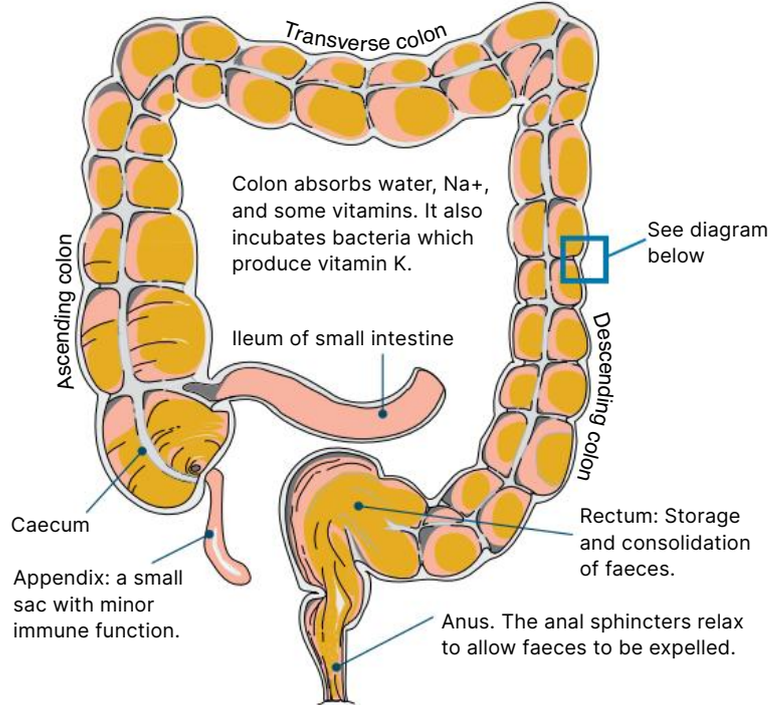
# 84

## The Large Intestine

**Key Idea:** The large intestine absorbs water and solidifies the indigestible material before passing it to the rectum. Undigested waste are egested as faeces from the anus. After most of the nutrients have been absorbed in the **small intestine**, the remaining semi-fluid contents pass into the

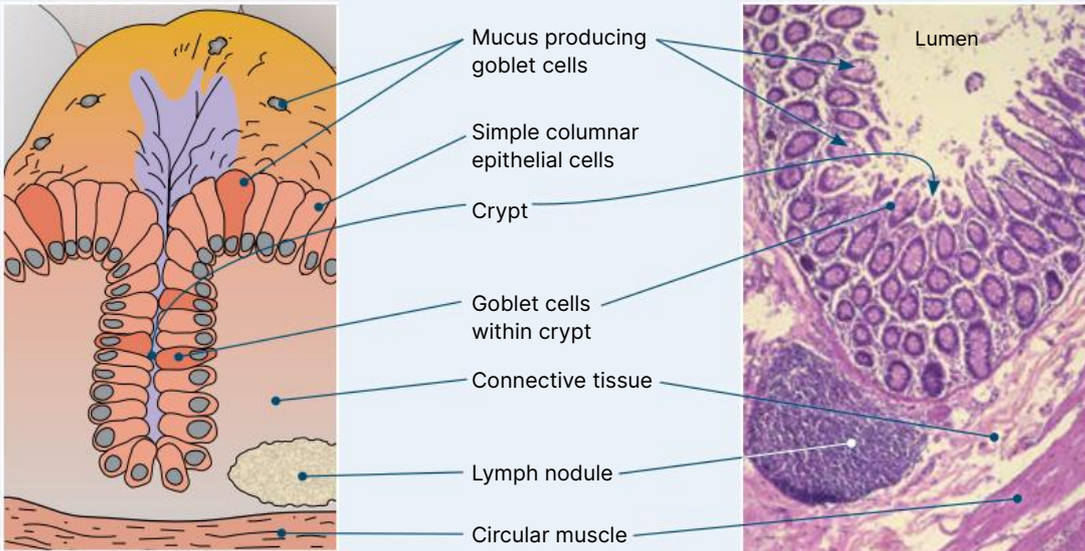
large intestine (consisting of the appendix, caecum, colon, and rectum). The large intestine's main role is to reabsorb water and electrolytes and to consolidate the waste material into faeces, which are eliminated from the anus in a process called egestion.

- ▶ After most of the nutrients have been absorbed in the small intestine, the remaining semi-fluid contents pass into the large intestine (appendix, cecum, and colon). This mixture includes undigested or indigestible food, (such as cellulose), bacteria, dead cells, mucus, bile, ions, and water. In humans and other omnivores, the large intestine's main role is to reabsorb water and electrolytes and consolidate the undigested material for egestion (elimination) from the anus.
- ▶ The rectum stores the waste faecal material before it is discharged out the anus. Fullness in the rectum produces the urge to defecate. If too little water is absorbed, the faeces will be watery as in diarrhoea. If too much water is absorbed the faeces will become compacted and difficult to pass.
- ▶ Defaecation is controlled by the anal sphincters, whose usual state is to be contracted (closing the orifice). Defaecation is under nervous control.



### Lining of the large intestine

The lining of the large intestine has a simple epithelium containing tubular glands (crypts) with many mucus-secreting cells. The mucus lubricates the colon wall and helps to form and move the faeces. In the photograph, some of the crypts are in XS and some are in LS.



Note the abundance of pale goblet cells.

1. What is the main purpose of the large intestine? \_\_\_\_\_
2. What are the effects of absorbing too little and too much water in the large intestine? \_\_\_\_\_

# Investigating Amylase Activity

**Key Idea:** Salivary amylase works optimally at the pH and temperature conditions of the human body. Enzyme activity outside these conditions decreases.

**Amylase** is a digestive **enzyme** that hydrolyses (breaks down) starch into maltose (a disaccharide) and glucose (a

monosaccharide). In mammals, amylase is secreted by the salivary glands into the saliva and by the pancreas into the **small intestine**. Like all enzymes, amylase works best under certain conditions. In this activity, you will investigate the effect of pH and temperature on amylase activity.



## Investigation 5.1 Investigating amylase activity

See appendix for equipment list.

- Obtain solutions of  $0.1 \text{ mol L}^{-1}$  iodine solution ( $\text{I}_2/\text{KI}$ ), 1% amylase, and 1% starch and buffer solutions to cover pH 4, 5, 6, 7, and 8. Iodine solution is a yellow/orange colour, but in the presence of starch, it turns a blue/black colour.
- Use a clean syringe to place a drop of iodine solution in each well of a two  $3 \times 4$  spotting plates.
- Add 1 mL of pH 4 buffer to a labelled test tube (TT4) and add 2 mL of amylase solution.
- Add 2 mL of the starch solution to TT4 and start a timer.
- Wait 10 seconds then use a clean syringe to add one drop of TT4 solution to the second well of the spotter plate (leave one well untouched as time 0). Return the solution in the syringe to TT4.
- Every 10 seconds add another one drop of solution from TT4 to another well on the spotting plate.
- Repeat until the iodine solution in the wells no longer changes colour. When this happens record the time as the time taken for the amylase solution to break down the starch.
- Repeat steps 2 to 7 with the rest of the buffer solutions (TT5, TT6, TT7, TT8).
- Record the results in the first two empty columns of the table below.



Spotting plate: each well contains a single drop of  $0.1 \text{ M}$  iodine solution (iodine dissolved in a solution of potassium iodide). Multiple spotting plates will accommodate the number of tests required.

- Why was it important to add the buffer and enzyme together before adding the starch?

---



---

- Complete your results table (right) by calculating the reaction rate for each pH ( $1 \div \text{seconds}$ ):

- (a) Graph the reaction rate vs pH on the grid.  
(b) Identify the pH where amylase activity was the highest:

---

- (c) Is this what you had expected? Explain:

---



---



---

- Some students repeated the experiment at pH 1. Each sample turned blue/black when added to the iodine even after five minutes of sampling. Explain what has happened here:

---



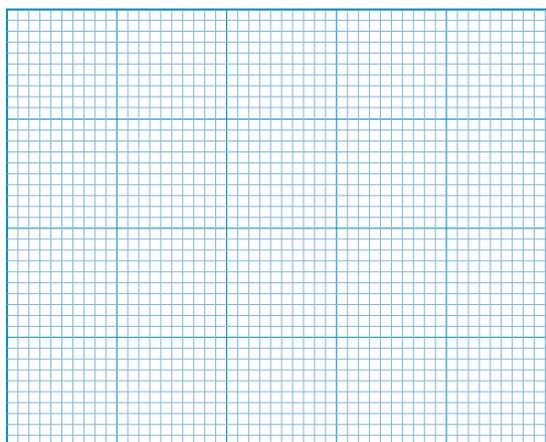
---



---

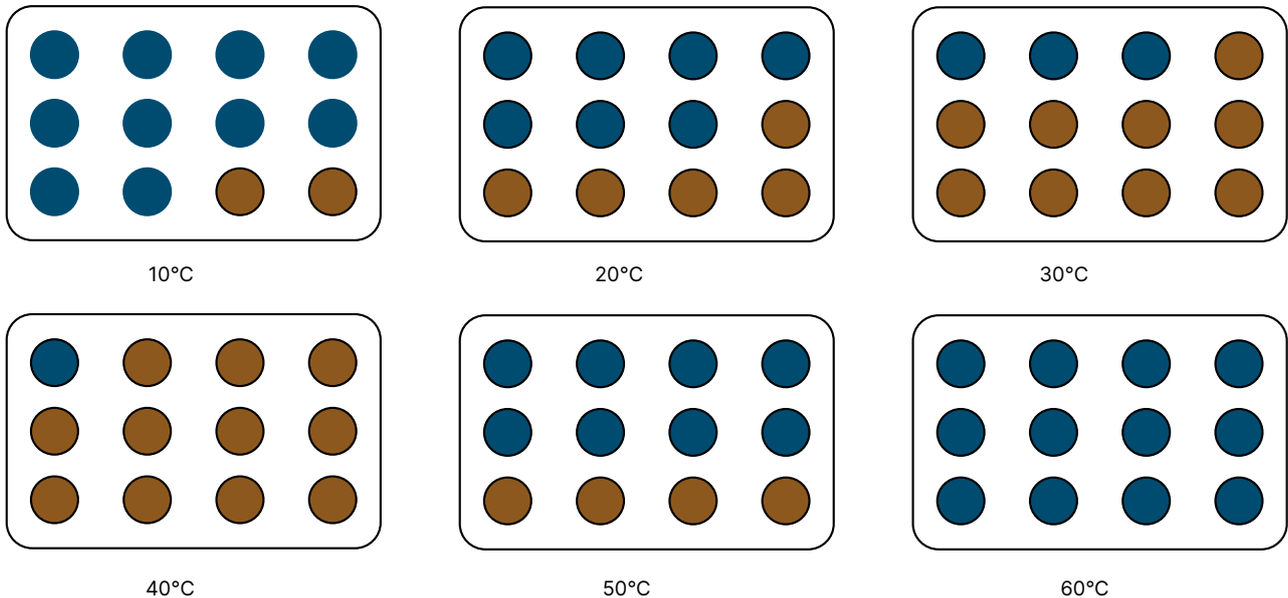
### Results

pH	Number of drops until no colour change occurred	Number of seconds until no colour change occurred	Rate of starch reduction (per second)
4			
5			
6			
7			
8			



## Effect of temperature on amylase activity

In an experiment, six students wanted to determine the temperature optimum for salivary amylase. Six spotting plates were set up by adding a single drop of 0.1 M iodine solution to each well. 2 cm<sup>3</sup> of 1% amylase solution and 1 cm<sup>3</sup> of a buffered pH 7 solution was added to each of 6 test-tubes. The test tubes were placed in water baths at the test temperatures (10, 20, 30, 40, 50, and 60°C ) and left for 5 minutes to equilibrate. Each student was responsible for investigating one temperature. Once the experimental temperature had been reached, 2 cm<sup>3</sup> of a 1% starch solution was added to the test-tube and a timer was started. After one minute, a plastic pipette was used to remove a small amount of solution. A single drop was added to the spotting plate and the colour change observed. Samples were repeated at one minute intervals until no colour change was seen. The results are shown below.



5. The students did not use any controls when they investigated the effect of pH on salivary amylase activity. What would a suitable control have been?

---

6. Why was the temperature investigation experiment carried out at pH 7? \_\_\_\_\_

---

7. Identify the temperature at which amylase shows no activity (the enzyme is denatured):

---

8. On the grid, plot the time taken for all the starch to be digested against temperature (do not plot 60°C):

9. Identify the optimum temperature for amylase:

---

10. Describe how temperature affects the activity of amylase:

---



---



---



---

11. Predict amylase activity below 10°C and give a reason for your prediction:

---



---



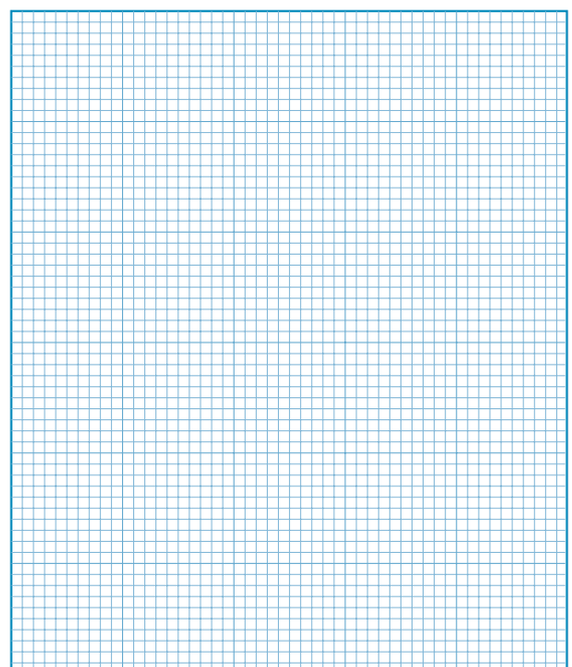
---



---



---

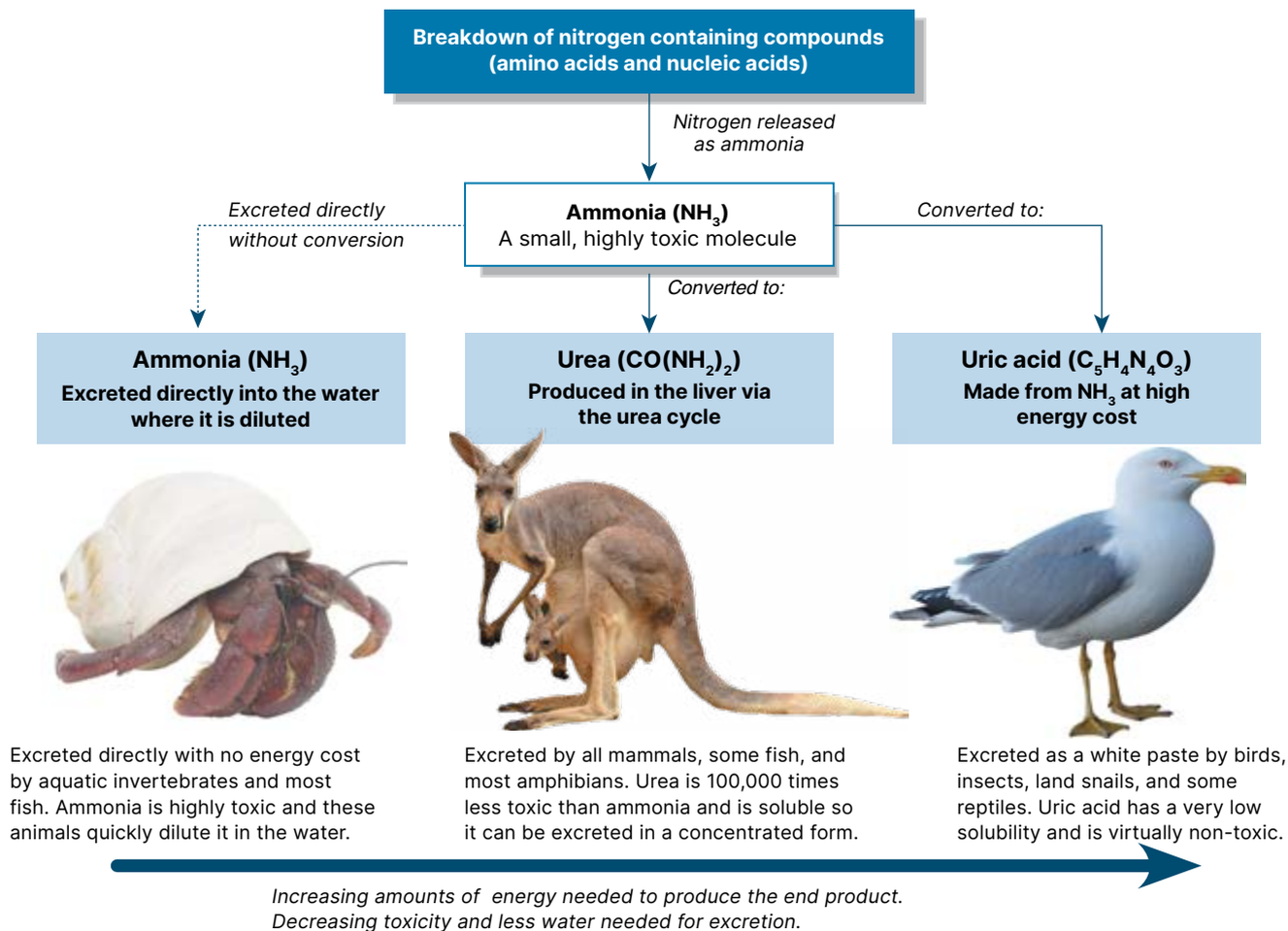


# Nitrogenous Wastes in Animals

**Key Idea:** Nitrogenous wastes are produced from the breakdown of nitrogen containing compounds. They must be excreted before they accumulate to toxic levels.

The process of removing the waste products of cellular metabolism is called **excretion**. These waste products include toxic nitrogenous wastes from the metabolism of amino acids and nucleic acids, as well as water,  $\text{CO}_2$ , and excess ions. The simplest breakdown product of nitrogen-containing compounds is **ammonia**, a highly toxic molecule that cannot be retained in the body for long. Most aquatic

animals excrete ammonia immediately into the water where it is washed away. Other animals convert the ammonia to a less toxic form (**urea** or **uric acid**) that can remain in the body for a short time before being excreted. The form of the excretory product in terrestrial animals depends on the organism type and life history. Terrestrial animals that lay eggs produce uric acid rather than urea, because it is non-toxic and very insoluble. It remains as an inert solid mass in the egg until hatching.



1. What is the main source of nitrogen-containing wastes in animals? \_\_\_\_\_

2. (a) Describe one advantage of uric acid as an excretory product (relative to urea and ammonia): \_\_\_\_\_

(b) Describe one disadvantage of ammonia as an excretory product: \_\_\_\_\_

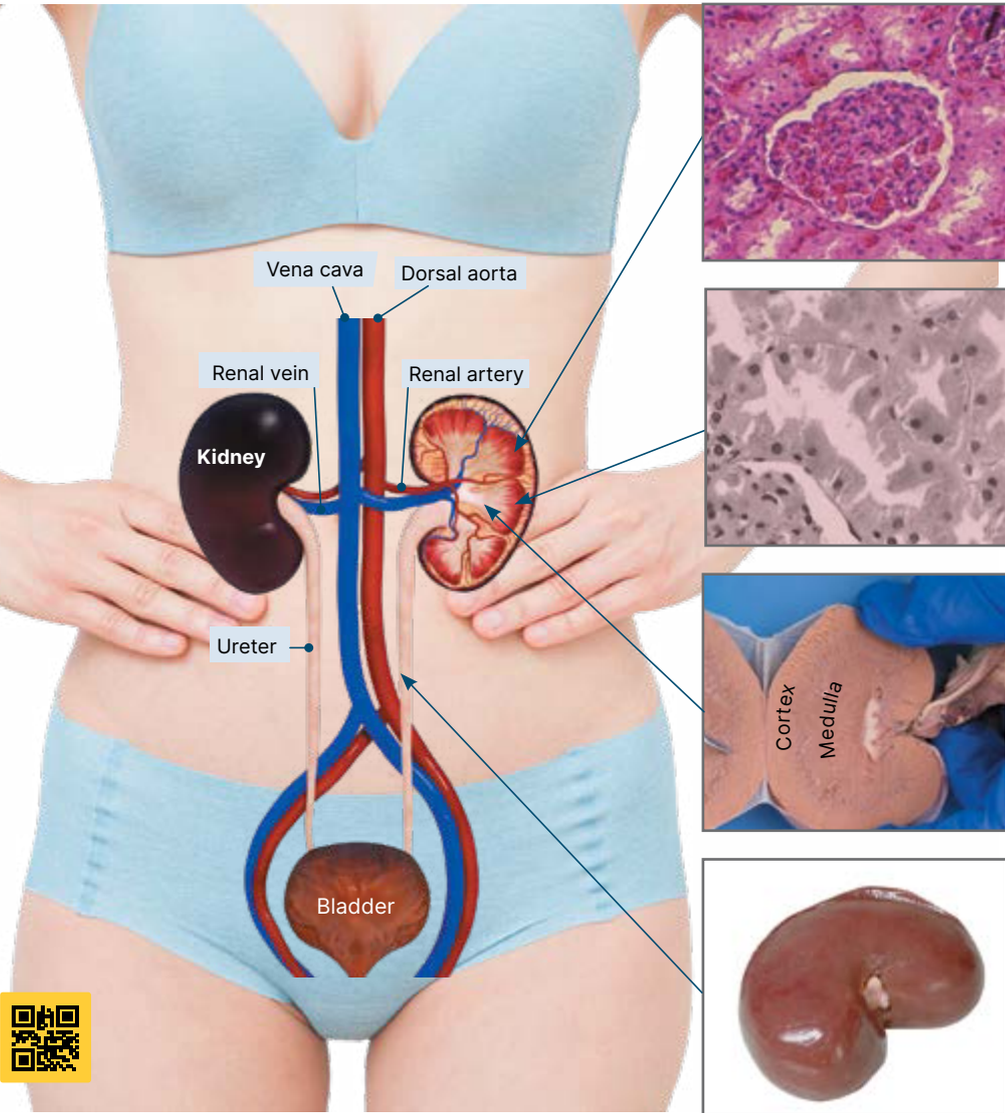
3. Discuss the relationship between the type of excretory product and an animal's environment: \_\_\_\_\_



# The Excretory System

**Key Idea:** The excretory system is responsible for removing metabolic wastes from the body by filtering the blood. The mammalian urinary system consists of the **kidneys** and bladder, and their associated **blood vessels** and ducts. The kidneys have a plentiful blood supply from the renal artery.

The blood plasma is filtered by the kidneys to form **urine**. Urine is produced continuously, passing along the ureters to the bladder. Mammalian kidneys are very efficient, producing a urine that is concentrated to varying degrees depending on fluid requirements at the time.



Blood is filtered in the kidneys by the **glomerulus**, a dense knot of capillaries. Blood is forced through them at high pressure, a process known as ultrafiltration. The filtrate is collected in the **Bowman's capsule** which surrounds the glomerulus.

The filtrate moves from Bowman's capsule to the convoluted tubules. These are lined with cuboidal epithelial cells, which have a brush border of **microvilli** to enhance absorption of substances from the filtrate. The glomerulus, capsule, and tubules form the functional unit structure of the kidney, the **nephron**.

The thousands of filtering elements of the kidney (the nephrons) are aligned and organised in a particular way in the kidney. The glomeruli and convoluted tubules are found in the outer region or cortex, while the "**loop of Henle**" is found in the inner region of medulla.

The filtrate passes to the renal ducts and then to the ureter and finally to the bladder. The kidney itself is bean shaped and is around 10 cm long in humans.



1. What is the purpose of the microvilli in the epithelial cells of the convoluted tubules? \_\_\_\_\_  
\_\_\_\_\_
2. (a) How is filtrate formed? \_\_\_\_\_  
\_\_\_\_\_  
\_\_\_\_\_
 

(b) How is the filtrate modified? \_\_\_\_\_  
\_\_\_\_\_
3. The circulation rate of blood through the renal artery is about  $1.2 \text{ L min}^{-1}$ , about one quarter of the heart's total output. Why does so much blood need to pass through the kidneys every minute?  
\_\_\_\_\_  
\_\_\_\_\_  
\_\_\_\_\_

# 88 Kidney Structure

**Key Idea:** In terrestrial vertebrates, the kidneys excrete nitrogenous waste and maintain water and solute balance. The central organs of the excretory system in humans and other mammals are the **kidneys**. They act as a selective filter of the blood, removing metabolic wastes while retaining useful substances, such as valuable ions and glucose. The kidneys receive blood under high pressure via the arterioles

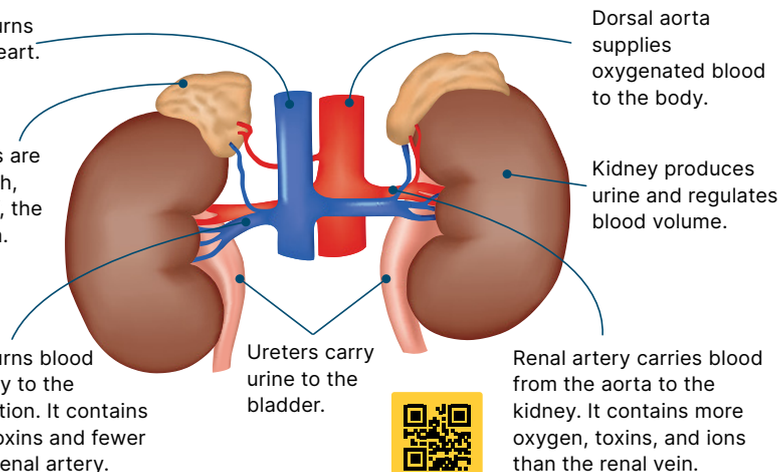
from the renal artery. This high pressure forces blood plasma out of the **capillaries**, forming a fluid called filtrate, which is then modified as it passes through the kidney to form the **urine**. Each day the kidneys filter about 180 L of plasma. Most of this is reabsorbed, leaving a daily urine output of about 1 L. By adjusting the composition of the fluid excreted, the kidneys help to maintain the body's internal chemical balance.

- ▶ The kidneys are bean shaped organs that lie at the back of the abdominal cavity to either side of the spine (below right).
- ▶ Human kidneys (right) are ~100–120 mm long and 25 mm thick. The precise alignment of the **nephrons** (the filtering elements of the kidney) and their associated **blood vessels** gives the kidney tissue a striped appearance (below). Each kidney contains more than 1 million nephrons. Nephrons are selective filter elements, which regulate blood composition and pH, and excrete wastes and toxins.

Vena cava returns blood to the heart.

Adrenal glands are associated with, but not part of, the urinary system.

Renal vein returns blood from the kidney to the venous circulation. It contains less oxygen, toxins and fewer ions than the renal artery.



## Kidney internal structure

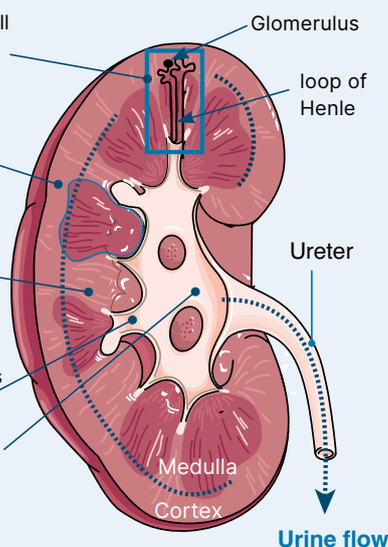
Nephrons are arranged with all the collecting ducts pointing towards the renal pelvis.

Outer cortex contains the renal corpuscles and convoluted tubules.

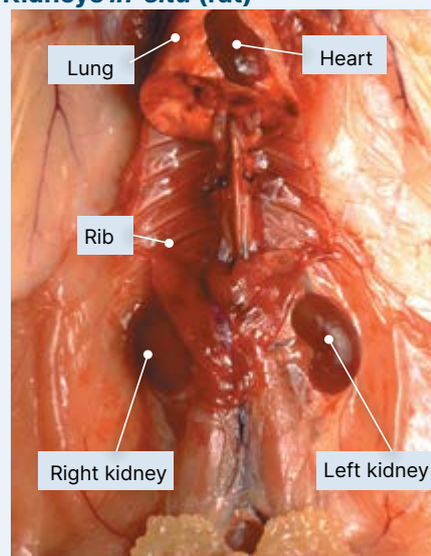
Inner medulla is organised into pyramids.

Urine flows from the pyramids towards the ureter.

Urine collects in a space near the ureter called the renal pelvis, before leaving the kidney via the ureter.



## Kidneys in-situ (rat)



1. What is the function of the kidney? \_\_\_\_\_
2. Calculate the percentage of the plasma reabsorbed by the kidneys: \_\_\_\_\_  
\_\_\_\_\_
3. The kidneys are located near the lower part of the ribcage. What do you think is the significance of this location?  
\_\_\_\_\_  
\_\_\_\_\_
4. Describe the location and orientation of the nephrons in a kidney: \_\_\_\_\_  
\_\_\_\_\_  
\_\_\_\_\_  
\_\_\_\_\_



# Nephron Structure and Function

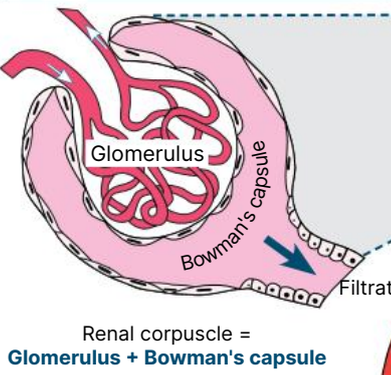
**Key Idea:** The functional unit of the kidney is the nephron. It is a selective filter element, comprising a renal corpuscle and its associated tubules and ducts.

Ultrafiltration, i.e. forcing fluid and dissolved substances through a membrane by pressure, occurs in the first part of the **nephron**, across the membranes of the capillaries and the glomerular capsule. The formation of the glomerular filtrate

depends on the pressure of the blood entering the nephron (below). If it increases, filtration rate increases; when it falls, glomerular filtration rate also falls. This process is precisely regulated so that glomerular filtration rate per day stays constant. The initial filtrate, now called **urine** is modified through secretion and tubular reabsorption according to body's needs at the time.

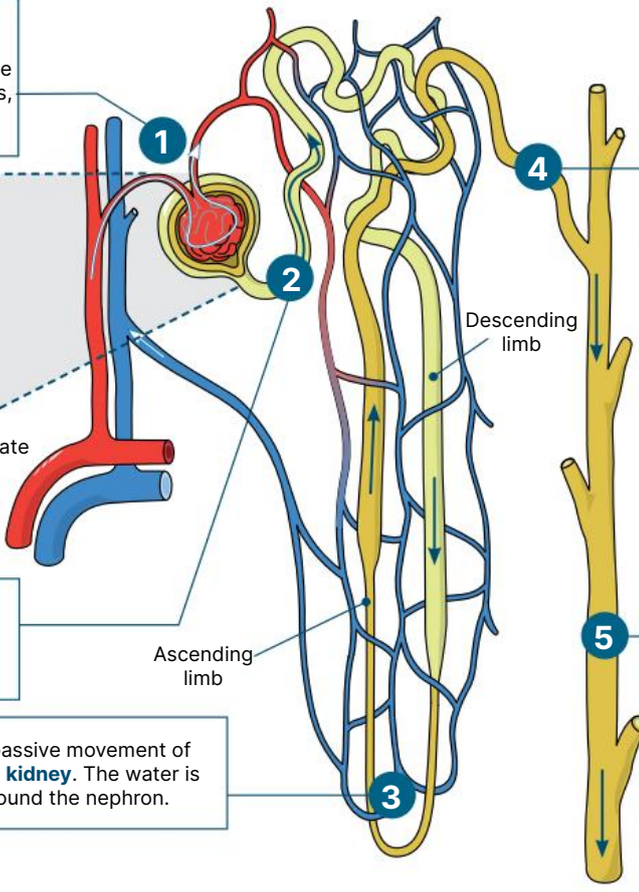


**Renal corpuscle:** Blood is filtered and the filtrate enters the convoluted tubule (enlargement below). The filtrate contains water, glucose, **urea**, and ions, but lacks cells and large proteins.



**Proximal convoluted tubule:** Reabsorption of ~ 90% of filtrate, including glucose and valuable ions.

**Loop of Henle:** Transport of salt and passive movement of water create salt gradient through the **kidney**. The water is transported away by blood vessels around the nephron.



**Distal convoluted tubule:** Further modification of the filtrate by active reabsorption and secretion of ions.

Blood  
 Filtrate (urine)  
 Blood vessels  
 around nephron

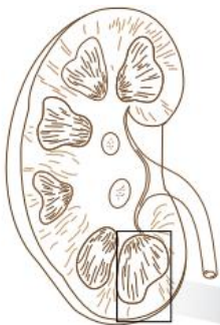
**Collecting duct:** Water leaves the filtrate (urine) by osmosis, making it more concentrated. The salt gradient established by the loop of Henle allows water to be removed along the entire length of the collecting duct.

1. What is the purpose of the nephron? \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_
2. Summarise the main activities in each of the five regions of the nephron:
  - (a) Renal corpuscle: \_\_\_\_\_
  - (b) Proximal (near) convoluted tubule: \_\_\_\_\_
  - (c) Loop of Henle: \_\_\_\_\_
  - (d) Distal (far) convoluted tubule: \_\_\_\_\_
  - (e) Collecting duct: \_\_\_\_\_
3. A kidney contains 1.5 million nephrons (filtering units). A person only needs 300,000 working nephrons to survive.
  - (a) What percentage of nephrons actually need to be working for a person to survive? \_\_\_\_\_
  - (b) Why is this important to someone with a damaged kidney? \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_



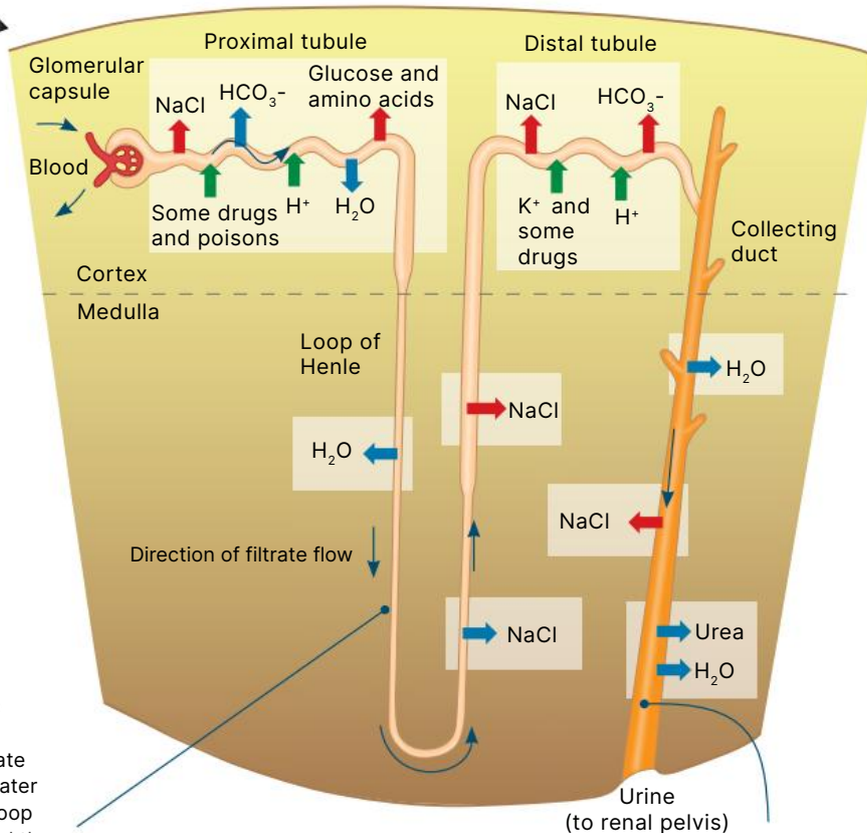
## Summary of activities in the kidney nephron

Urine formation begins by ultrafiltration of the blood, as fluid is forced through the capillaries of the glomerulus, forming a filtrate similar to blood but lacking cells and proteins. The filtrate is then modified by secretion and reabsorption to add or remove substances (e.g. ions). The processes involved in urine formation are summarised below for each region of the nephron (glomerulus, proximal convoluted tubule, **loop of Henle**, and distal convoluted tubule), and the collecting duct. The loop of Henle acts as a countercurrent multiplier, establishing and increasing the salt gradient through the medullary region. This is possible because the descending limb is freely permeable to water but the ascending limb is not.



Filtrate	
H <sub>2</sub> O	
Salts (NaCl, etc.)	
HCO <sub>3</sub> <sup>-</sup> (bicarbonate)	
H <sup>+</sup>	
Urea	
Glucose; amino acids	
Some drugs	

Reabsorption	
Active transport	→
Passive transport	⇨
Secretion (active transport)	⇦



The thick ascending limb of the loop of Henle pumps out sodium and chloride ions from the filtrate. This produces a high solute concentrate in the interstitial space. This in turns draws water by osmosis from the descending limb of the loop Henle into the interstitial space. The water and the ions are transported away by the capillaries. The countercurrent flow within the descending and ascending limbs multiplies the osmotic gradient between the tubular fluid and the interstitial space.



Reabsorption of a small amount of urea from the urine helps to maintain the osmotic gradient for the removal of water.

- (a) What is the purpose of the salt gradient in the kidney? \_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_

(b) How is this salt gradient produced? \_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_
- (a) The kidneys of desert mammals are adapted to conserve water. One of these adaptations is in the length of the loop of Henle. Would a desert mammal have a longer or shorter loop of Henle than a non-desert adapted mammal?

\_\_\_\_\_

(b) Explain why: \_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_
- Where in the nephron is water reabsorbed? \_\_\_\_\_

# Organ and Tissue Transplantation

**Key Idea:** There are many more people needing organ and tissue transplants than there are donors. And not all organs are obtained ethically.

The demand for new organs often fails to match the number available from donors. Illegal or unethical sale of human

organs can be a problem and organ trafficking is worth big money throughout the world. Because humans can survive without two kidneys, poor people can be exploited for organ donation and given inappropriate aftercare. Transplant tourism is a real phenomenon and can involve large sums of money.

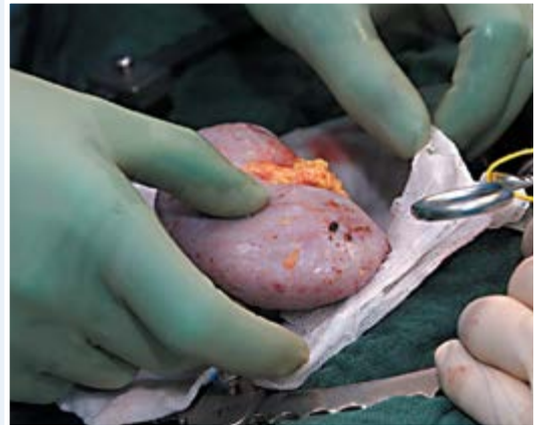
**Organ trafficking** is the (illegal) sale and purchase of human organs for **transplantation**. The most commonly trafficked organ is the kidney. Donors can live perfectly well with just one kidney, so they can be bought or sold more easily, especially from the poor or desperate.

There are tens of thousands of people throughout the world needing organ transplants. In many countries, these organs are obtained from donors, usually after their death. Waiting lists are based on need, tissue matches, and other medical information. However, there are many more people waiting for transplants than there are organ donors. Some people are prepared to pay large sums of money for an organ transplant, either to move them up the waiting list or avoid the usual processes and obtain preferential treatment through a private clinic.



- ▶ Guidelines on organs donation set up by the WHO include that organ donation should be altruistic (for no commercial gain). In 2013, Australia legalised financial compensation for organ donors. This is limited and aimed at compensating the donor for taking time off work while recovering.
- ▶ Some people argue that organs should be able to be bought and sold legally. This would allow low-income people to benefit financially and lets people needing a transplant to quickly get on with their lives. However most medical professions reject this on the basis of ethical issues, which include the exploitation of the poor or desperate and the treatment of human organs as commodities to be bought and sold.
- ▶ Transplant tourism is the overseas travel of people to places where they can acquire a needed organ. It refers mainly to the buying and selling of organs as a purely commercial venture.
- ▶ Obtaining organs from paid (often exploited) donors raises the risk of spread of disease, including hepatitis and HIV. There can also be a lack of follow up care, including anti-rejection drugs, so the transplant may fail. Studies have shown there are a higher number of failed operations from illegal organ transplants than from legal surgeries.

- ▶ Transplantation of a healthy kidney from a donor is the preferred treatment for end-stage kidney failure. The organ is usually taken from a person who has just died, although kidneys can also be taken from living donors. The damaged kidneys are left in place and the new kidney transplanted into the lower abdomen (right). If recipients comply with medical requirements (e.g. correct diet and medication) over 85% of kidney transplants are successful.
- ▶ The two major problems associated with kidney transplants are lack of donors and tissue rejection. Cells from donor tissue have different antigens to those of the recipient, and the recipient's immune system will attack the new kidney, recognising it as foreign.
- ▶ Tissue-typing and the use of immunosuppressant drugs helps to decrease organ rejection rates. In the future, the transplant of genetically modified organs from other species may help to solve the problems of supply and immune rejection.



1. In Australia, there are more than 1000 people waiting for a kidney transplant but fewer than 250 live donors. As a group, discuss the issues surrounding organ transplants, including the lack of willing donors and how this might be solved. Is financial compensation ethical? How far should compensation go? Should people be able to sell their own organs? Summarise your discussion below:

---



---



---



---



---



---



---

1. Test your vocabulary by matching each term to its correct definition, as identified by writing the letter in the correct box.

- (i) amylase
- (ii) excretion
- (iii) glomerulus
- (iv) kidney
- (v) loop of Henle
- (vi) small intestine
- (vii) urea

- A** Part of the gut that receives chyme directly from the stomach. Here, digestive enzymes are added, which break down food and food molecules reabsorbed into the blood.
- B** Part of the kidney nephron between the proximal and distal convoluted tubules. Its function is to create a salt gradient through the medullary region of the kidney.
- C** The primary nitrogenous excretory product of mammals.
- D** Enzyme produced by the salivary glands and pancreas that breaks down starch into smaller maltose molecules.
- E** Elimination of the waste products of metabolism.
- F** Bean shaped excretory organ in vertebrates, which removes and concentrates metabolic wastes from the blood.
- G** The collection of capillaries within Bowman's capsule in the kidney where ultrafiltration of the blood plasma occurs.

2. (a) What structures from the small intestine of a mammal are shown in the photograph (right)?

\_\_\_\_\_

(b) What is their function? \_\_\_\_\_

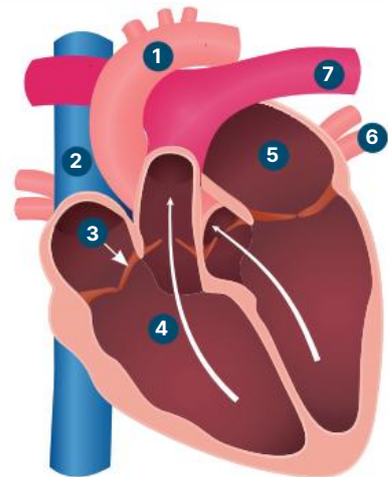
\_\_\_\_\_

\_\_\_\_\_



3. Identify and name the numbered features in the heart diagram, (right):

- (a) 1: \_\_\_\_\_
- (b) 2: \_\_\_\_\_
- (c) 3: \_\_\_\_\_
- (d) 4: \_\_\_\_\_
- (e) 5: \_\_\_\_\_
- (f) 6: \_\_\_\_\_
- (g) 7: \_\_\_\_\_



4. The drawing, right, depicts nephrons from the kidney.

- (a) How many nephrons are shown? \_\_\_\_\_
- (b) Label the diagram to show glomerulus, Bowman's capsule, proximal convoluted tubule, distal convoluted tubule, loop of Henle, collecting duct.
- (c) Draw arrows to indicate the direction of urine flow.
- (d) In a different colour, label where the reabsorption of glucose would occur.
- (e) What significant feature is missing from this diagram?

\_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_





# Internal Membranes and Enzymes

## Key Terms

- activation energy
- active site
- anabolic reaction
- catabolic reaction
- catalyst
- coenzyme
- cofactor
- competitive inhibition
- denaturation
- enzyme
- extracellular
- induced fit model
- intracellular
- lock-and-key model
- metabolic pathway
- metabolism
- noncompetitive inhibition
- optimum (for enzyme)

## Key Concepts

- ▶ Enzymes are biological catalysts that are used to speed up biochemical reactions.
- ▶ Enzymes have optimum conditions under which they best control reactions and are affected by temperature, pH and concentrations of reactants and products.
- ▶ Failure of enzymes to work correctly can cause human metabolic conditions.

## Enzymes are protein catalysts

### Activity Number

- |     |  |       |
|-----|--|-------|
| □ 1 | Explain the meaning of enzymes as biological catalysts and define the terms substrate and active site.   | 92    |
| □ 2 | Describe models for enzyme function, including reference to the enzyme-substrate complex, enzyme-product complex, and product formation.   | 92-93 |
| □ 3 | Compare and contrast the induced-fit and the lock-and-key models of enzyme function and explain why the lock and key model was changed.  | 93    |
| □ 4 | Explain how enzymes catalyse reactions by lowering the activation energy and identify this on a plot of the progress of the reaction against the free energy. Describe the structure and role of the active site, explaining its importance to the specificity of the enzyme to its substrate(s). Explain the role of cofactors. | 94    |

## Factors affecting enzyme activity

- |     |   |        |
|-----|---|--------|
| □ 5 | Analyse how enzyme activity can be affected by pH, temperature, and the concentrations of reactants (substrates) and products.                      | 95     |
| □ 6 | <b>SI:</b> Investigate how enzyme reaction rates can be affected by pH, temperature, and the concentrations of reactants (substrates) and products. | 95, 97 |

## Enzymes and metabolism

- |      |  |     |
|------|--|-----|
| □ 7  | Using examples, describe the effects of inhibitors on the rate of enzyme-controlled reactions. Include reference to competitive and non-competitive inhibition and identify these on graphs of reaction rate vs substrate concentration.                         | 96  |
| □ 8  | Explain the role of end-product inhibition (negative feedback) in regulating biochemical pathways.   | 98  |
| □ 9  | Explain, using an example, how the arrangement of internal membranes controls biochemical processes through enzyme attachment and contributes to the functional efficiency of the cell.  | 99  |
| □ 10 | <b>SI:</b> Explore the effect of enzyme failure on metabolic reactions. Describe the metabolism of phenylalanine and the effect on this metabolic pathway if enzymes fail to work correctly. Link examples of metabolic disease to absence of essential enzymes. | 100 |

**Key Idea:** Enzymes are biological catalysts. The active site is critical to this functional role.

Most **enzymes** are globular proteins. Enzymes are biological **catalysts** because they speed up biochemical reactions, but the enzyme itself remains unchanged. The substrate in a

reaction binds to a region of the enzyme called the **active site**, which is formed by the precise folding of the enzyme's amino acid chain. Enzymes control **metabolic pathways**. One enzyme will act on a substance to produce the next reactant in a pathway, which will be acted on by a different enzyme.

### The active site

An enzyme acts on a specific chemical called a substrate. The substrate binds to a specific part of the enzyme called the active site.

The shape and chemistry of the active site is specific to an enzyme and is a function of the polypeptide's tertiary structure (the way the protein folds up). The amylase shown here breaks starch (a large molecule made of repeating glucose units) into smaller pieces with 2-3 glucose units.

Extremes of temperature or pH can alter the enzyme's active site and lead to loss of function. This process is called **denaturation**.

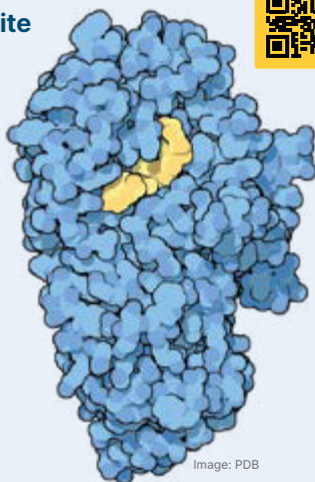


Image: PDB  
Amylase (blue) with bound glucose (yellow) at the active site

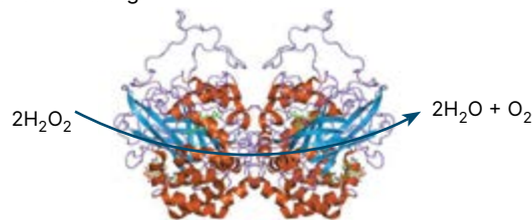
### Enzymes can be intracellular or extracellular

Enzymes can be defined based on where they are produced relative to where they are active.

An **intracellular enzyme** is an enzyme that performs its function within the cell that produces it. Most enzymes are intracellular enzymes, e.g. respiratory enzymes.

**Example:** Catalase.

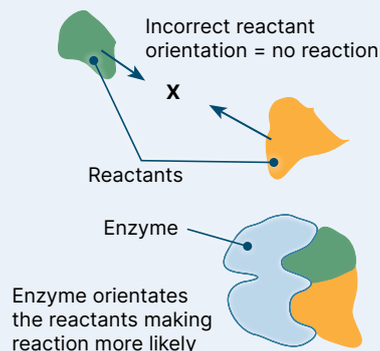
Many metabolic processes produce hydrogen peroxide, which is harmful to cells. Catalase converts hydrogen peroxide into water and oxygen (below) to prevent damage to cells and tissues.



Catalase

### Substrates collide with an enzyme's active site

For a reaction to occur, reactants must collide with sufficient speed and with the correct orientation. Enzymes enhance reaction rates by providing a site for reactants to come together in such a way that a reaction will occur. They do this by orientating the reactants so that the reactive regions are brought together. They may also destabilise the bonds within the reactants making it easier for a reaction to occur.



An **extracellular enzyme** is an enzyme that functions outside the cell from which it originates (i.e. it is produced in one location but active in another).

**Examples:** Amylase and trypsin.

Amylase is a digestive enzyme produced in the salivary glands and pancreas in humans. However, it acts in the mouth and small intestine respectively to hydrolyse starch into sugars.

Trypsin is a protein-digesting enzyme and hydrolyses the peptide bond immediately after a basic amino acid (e.g. arginine). It is produced in an inactive form (called trypsinogen) and secreted into the small intestine by the pancreas. It is activated in the intestine by the enzyme enteropeptidase to form trypsin. Active trypsin can convert more trypsinogen to trypsin.

1. (a) What is meant by the active site of an enzyme and relate it to the enzyme's tertiary structure:

---



---



---

- (b) Why are enzymes specific to one substrate (or group of closely related substrates)?

---



---

2. How do substrate molecules come into contact with an enzyme's active site?

---

3. (a) Suggest why digestion (the breakdown of large macromolecules) is largely performed by extracellular enzymes:

---

- (b) Why would an extracellular enzyme be produced and secreted in an inactive form?

---

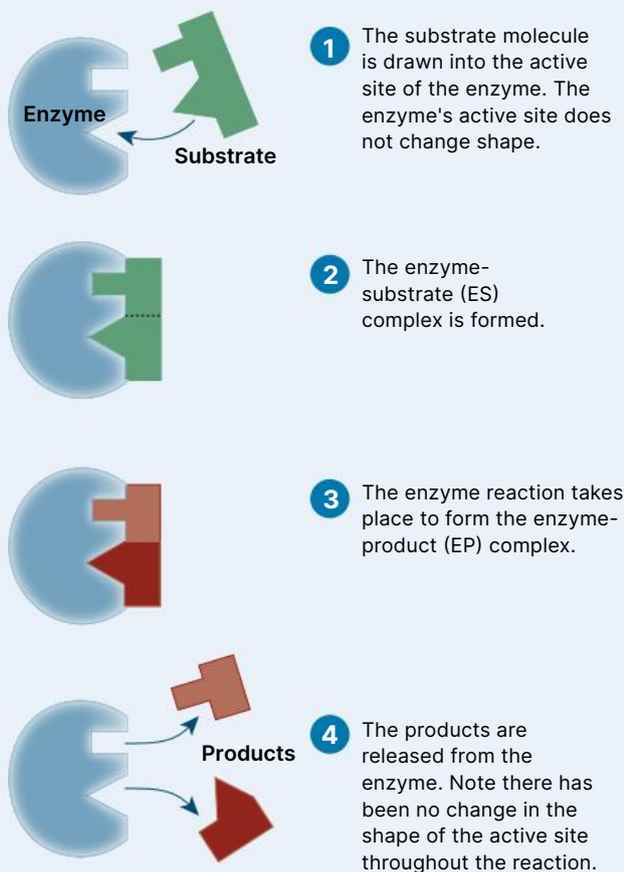
# 93 Models of Enzyme Activity

**Key Idea:** Enzymes catalyse reactions by providing a reaction site for a substrate. The model that describes the behaviour of enzymes the best is the induced fit model.

The initial model of **enzyme** activity was the lock and key model proposed by Emil Fischer in the 1890s. Fischer proposed enzymes were rigid structures, similar to a lock, and the substrate was the key. While some aspects of

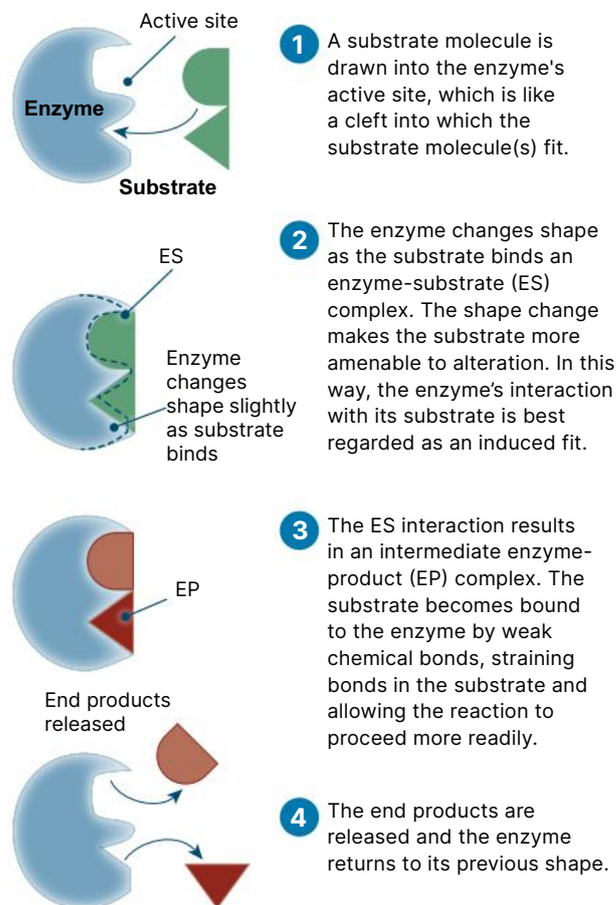
Fischer's model were correct, for example, substrates align with enzymes in a way that is likely to make a reaction more likely, the model has been adapted as techniques to study molecular structures have developed. The current '**induced-fit**' model of enzyme function is supported by studies of enzyme inhibitors, which show that enzymes are flexible and change shape when interacting with the substrate.

## The lock and key model of enzyme function



The **lock and key model** proposed in 1894 suggested that the (perfectly fitting) substrate was simply drawn into a matching site on the enzyme molecule. If the substrate did not perfectly fit the **active site**, the reaction did not proceed. This model was supported by early X-ray crystallography studies but has since been modified to recognise the flexibility of enzymes (the induced fit model).

## The current induced fit model



Once the substrate enters the active site, the shape of the active site changes to form an active complex. The formation of an ES complex strains substrate bonds and lowers the energy required to reach the transition state. The induced-fit model is supported by X-ray crystallography, chemical analysis, and studies of enzyme inhibitors, which show that enzymes are flexible and change shape when interacting with the substrate.

1. Describe the key features of the 'lock and key' model of enzyme action and explain its deficiencies as a working model:

---



---



---



---

2. How does the current 'induced fit' model of enzyme action differ from the lock and key model?

---



---



---



# 94 How Enzymes Work

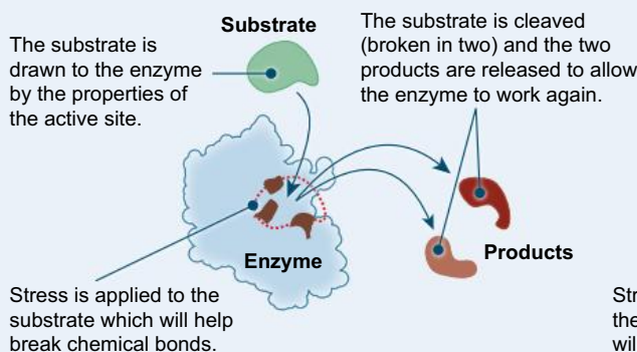
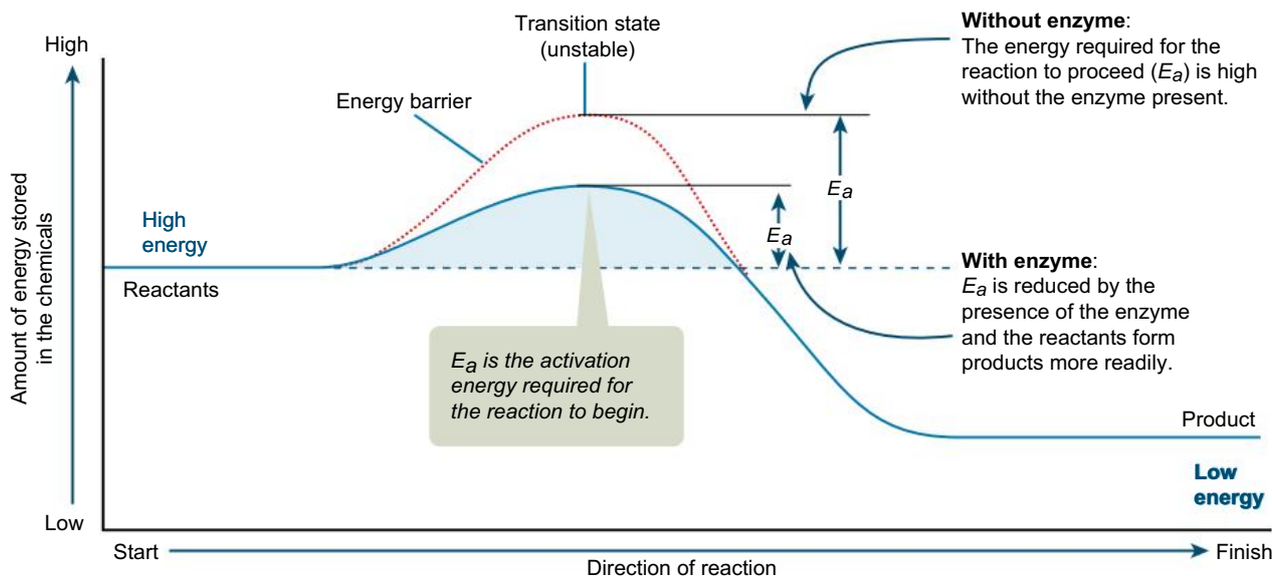
**Key Idea:** Enzymes increase the rate of biological reactions by lowering the reaction's activation energy.

Chemical reactions in cells are accompanied by energy changes. The amount of energy released or taken up is directly related to the tendency of a reaction to run to completion (for all the reactants to form products). Any reaction needs to raise the energy of the substrate to an unstable transition

state before the reaction will proceed (below). The amount of energy needed to do this is the **activation energy** ( $E_a$ ). **Enzymes** lower the  $E_a$  by destabilising bonds in the substrate so that it is more reactive. Enzyme reactions can break down a single substrate molecule into simpler substances (**catabolic reactions**), or join two or more substrate molecules together (**anabolic reactions**).

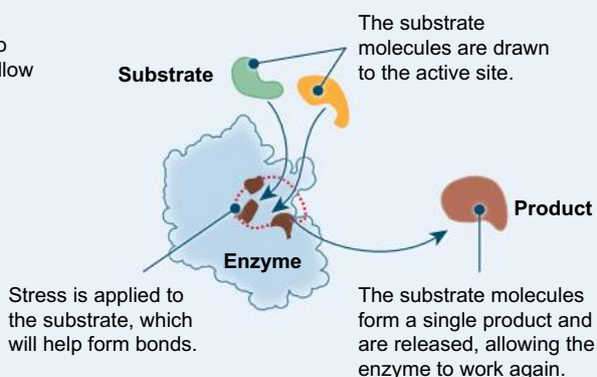
## Lowering the activation energy

The presence of an enzyme simply makes it easier for a reaction to take place. All **catalysts** speed up reactions by influencing the stability of bonds in the reactants. They may also provide an alternative reaction pathway, thus lowering the activation energy ( $E_a$ ) needed for a reaction to take place (see the graph below).



### Enzymes can catalyse the breakdown of molecules

Some enzymes can cause a single substrate molecule to be drawn into the **active site**. Chemical bonds are broken, causing the substrate molecule to break apart to become two separate molecules. Reactions that break down complex molecules into simpler ones are called catabolic reactions and involve a net release of energy (they are exergonic). Example: *digestion*.



### Enzymes can catalyse the building of molecules

Some enzymes can cause two substrate molecules to be drawn into the active site. Chemical bonds are formed, causing the two substrate molecules to form bonds and become a single molecule. Reactions that build more complex molecules and structures from simpler ones are called anabolic reactions and involve a net use of energy (they are endergonic). Example: *protein synthesis*.

1. Why do reactants need energy added to them in order for them to react? \_\_\_\_\_

---



---



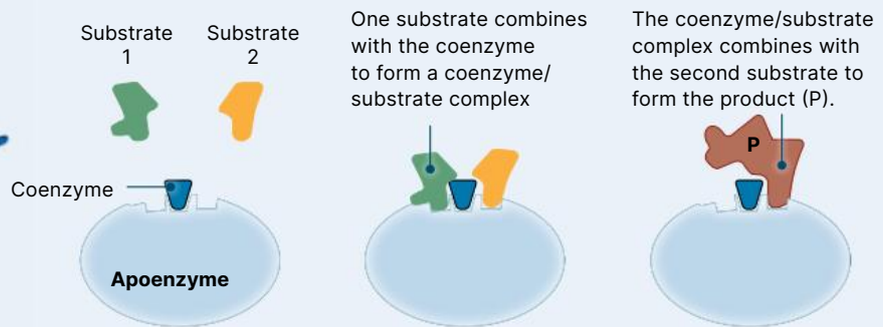
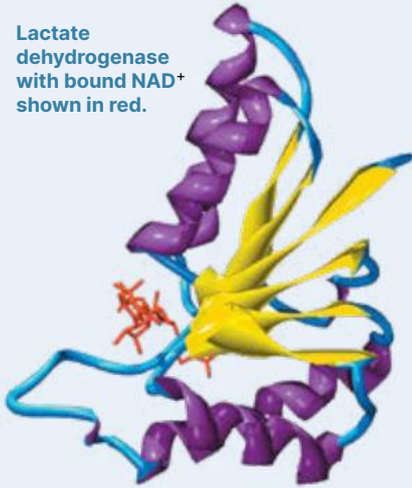
---



## Enzymes and cofactors

Nearly all enzymes are made of protein. Some enzymes are functional protein-only molecules, but many require additional non-protein components, called **cofactors**, to function. Cofactors can be subclassified as either inorganic ions (e.g.  $Zn^{2+}$ ) or complex organic molecules called **coenzymes** (many of which are vitamins). Many enzymes need several cofactors in order to function. Where a cofactor is needed for enzyme function, the enzyme (protein) component is called the apoenzyme. The cofactor often completes the active site or makes the active site more reactive by assisting enzyme-substrate interactions.

**Lactate dehydrogenase with bound  $NAD^+$  shown in red.**



If cofactors are not permanently bound to the enzyme, they can detach after the reaction to participate in other reactions. Neither the apoenzyme nor the cofactor has catalytic activity on its own. **Example:** dehydrogenases + NAD. NAD is the coenzyme form of the vitamin niacin ( $B_3$ ). Many coenzymes are vitamin derivatives.

2. How do enzymes lower the activation energy for a reaction? \_\_\_\_\_

---



---



---

3. Why are enzymes referred to as "biological catalysts"? \_\_\_\_\_

---



---

4. Describe the difference between digestion and protein synthesis in terms of the energy released or required :

---



---



---

5. What is a cofactor? \_\_\_\_\_

---



---

6. Describe the difference between the two different broad categories of cofactors: \_\_\_\_\_

---



---



---

7. How do cofactors enable an enzyme's catalytic activity? \_\_\_\_\_

---



---



---

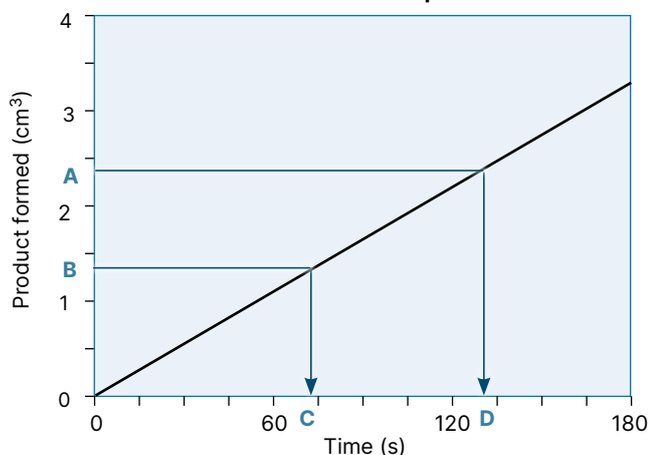
# Factors Affecting Enzyme Activity

**Key Idea:** Enzymes operate most effectively within a narrow range of conditions. The rate of enzyme-catalysed reactions is influenced by both enzyme and substrate concentration.

**Enzymes** usually have an **optimum** set of conditions (e.g. of pH and temperature) under which their activity is greatest. Many plant and animal enzymes show little activity at low

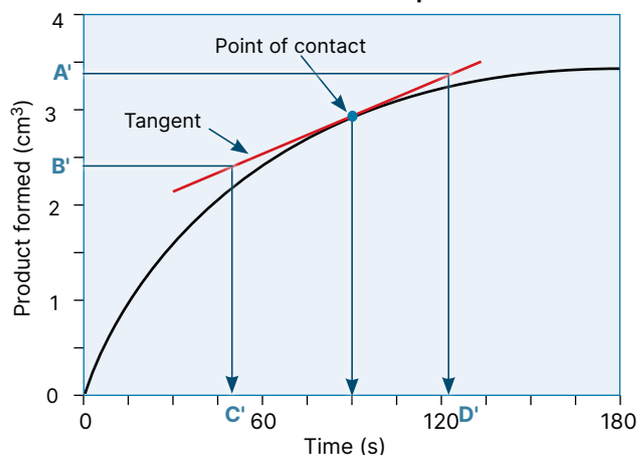
temperatures. Enzyme activity increases with increasing temperature, but falls off after the optimum temperature is exceeded and the enzyme is denatured. Extremes in pH can also cause **denaturation**. Within their normal operating conditions, enzyme reaction rates are influenced by enzyme and substrate concentration in a predictable way.

Reaction rate: Graph 1

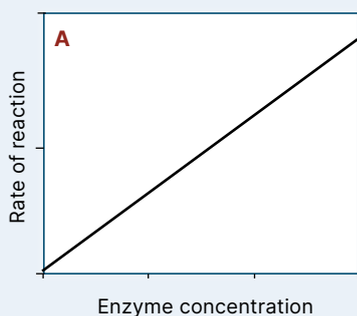


The rate of a reaction can be calculated from the amount of product produced during a given time period. For a reaction in which the rate does not vary (graph 1) the reaction rate calculated at any one point in time will be the same. For example:  $B \div C = A \div D = A - B \div D - C = (\Delta p / \Delta t)$  (the change in product divided by the change in time).

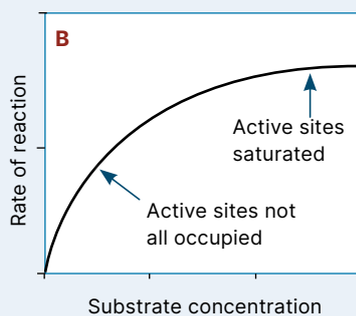
Reaction rate: Graph 2



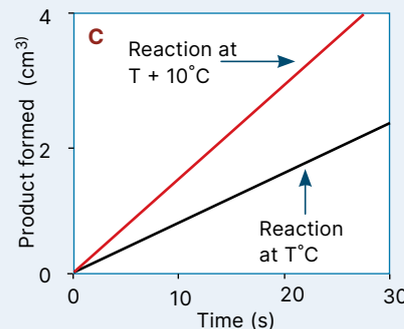
In a reaction in which the rate varies (graph 2) a reaction rate can be calculated for any instantaneous moment in time by using a tangent. The tangent must touch the curve at only one point. The gradient of the tangent can then be used to calculate the rate of reaction at that point in time ( $A' - B' \div D' - C'$ ).



Given an unlimited amount of substrate, the rate of reaction will continue to increase as enzyme concentration increases. More enzyme means more reactions between substrates can be catalysed in any given time (graph A).



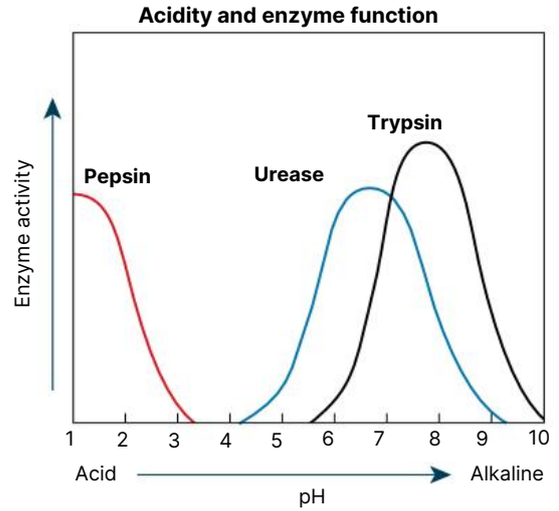
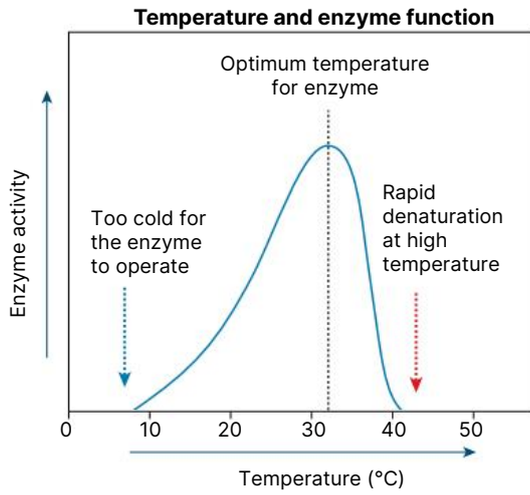
If there is unlimited substrate but the enzyme is limited, the reaction rate will increase until the enzyme is saturated, at which point the rate will remain static (graph B).



The effect of temperature on a reaction rate is expressed as the temperature coefficient, usually given as the  $Q_{10}$ .  $Q_{10}$  expresses the increase in the rate of reaction for every rise of 10°C.  $Q_{10} = \text{rate of reaction at } (T + 10^\circ\text{C}) / \text{rate of reaction at } T$ , where T is the temperature in °C (graph C).

- Calculate the reaction rate in graph 1: \_\_\_\_\_
- For graph 2:
  - The reaction rate at 90 seconds: \_\_\_\_\_
  - The reaction rate at 30 seconds: \_\_\_\_\_
- What must be happening to the reaction mix in graph 1 to produce the straight line (constant reaction rate)?  
\_\_\_\_\_  
\_\_\_\_\_
  - Explain why the reaction rate in graph 2 changes over time: \_\_\_\_\_  
\_\_\_\_\_





Higher temperatures speed up all reactions, but few enzymes can tolerate temperatures higher than 50–60°C. The rate at which enzymes are denatured (change their shape and become inactive) increases with higher temperatures. The temperature at which an enzyme works at its maximum rate is called the optimum temperature.

Like all proteins, enzymes are denatured by extremes of pH. Within these extremes, most enzymes have a specific pH range for optimum activity. For example, digestive enzymes are specific to the region of the gut where they act: pepsin in the acid of the stomach and trypsin in the alkaline small intestine. Urease catalyses the hydrolysis of urea at a pH near neutral.

4. (a) Describe the change in reaction rate when the enzyme concentration is increased and the substrate is not limiting:

---



---

(b) Suggest how a cell may vary the amount of enzyme present: \_\_\_\_\_

---



---

5. Describe the change in reaction rate when the substrate concentration is increased (with a fixed amount of enzyme):

---



---

6. (a) Describe what is meant by an optimum temperature for enzyme activity: \_\_\_\_\_

---



---

(b) Explain why most enzymes perform poorly at low temperatures: \_\_\_\_\_

---



---

(c) For graph C on the previous page, calculate the  $Q_{10}$  for the reaction: \_\_\_\_\_

---



---

7. (a) State the optimum pH for each of the enzymes:

Pepsin: \_\_\_\_\_ Trypsin: \_\_\_\_\_ Urease: \_\_\_\_\_

(b) Explain how the pH optima of each of these enzymes is suited to its working environment: \_\_\_\_\_

---



---



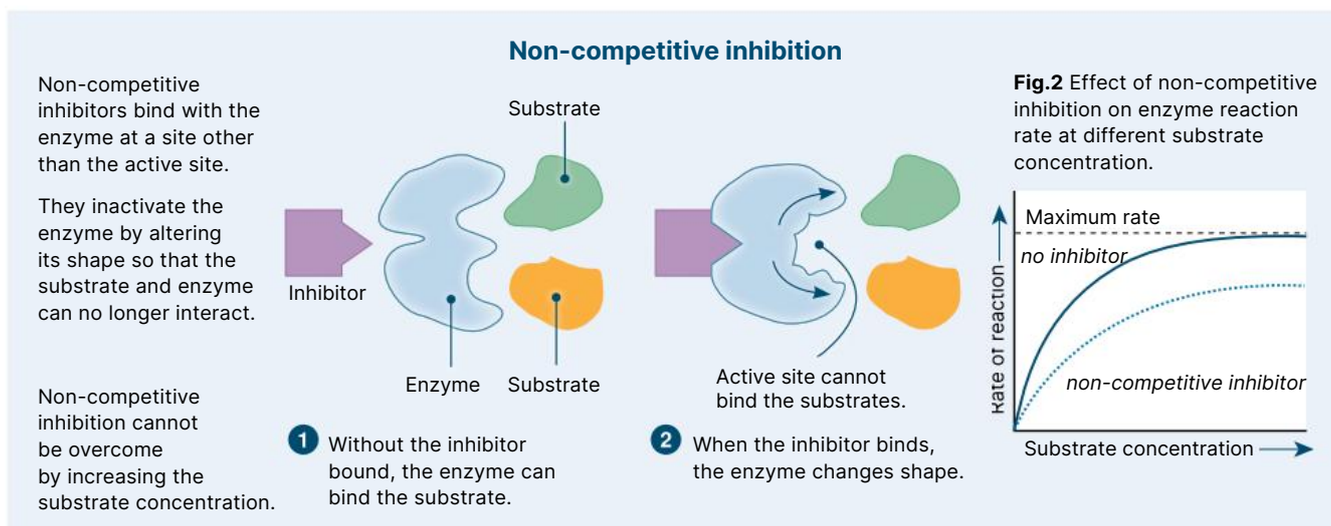
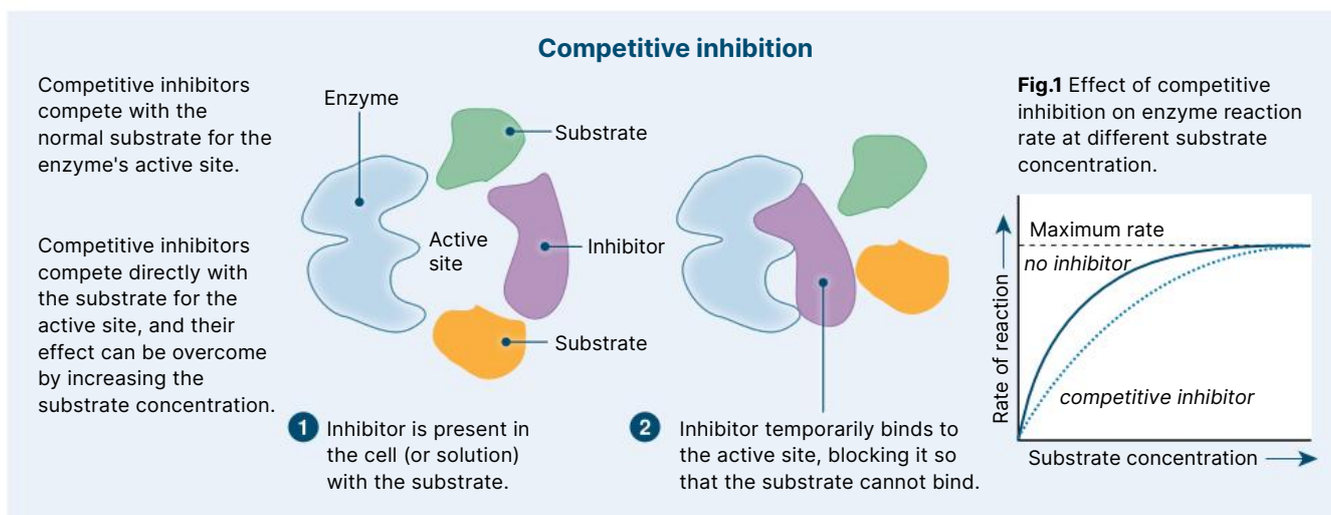
---

# 96 Enzyme Inhibition

**Key Idea:** Enzyme activity can be reduced or stopped by inhibitors. These may be competitive or non-competitive.

**Enzyme** activity can be stopped, temporarily or permanently, by chemicals called enzyme inhibitors. **Competitive inhibitors** compete directly with the substrate for the active

**site** and their effect can be overcome by increasing the concentration of available substrate. A **non-competitive inhibitor** does not occupy the active site, but distorts it so that the substrate and enzyme can no longer interact.



1. Distinguish between competitive and non-competitive inhibition: \_\_\_\_\_

---



---



---

2. (a) Compare and contrast the effect of competitive and non-competitive inhibition on the relationship between the substrate concentration and the rate of an enzyme controlled reaction (figures 1 and 2 above):

---



---



---



---

(b) Suggest how you could distinguish between competitive and non-competitive inhibition in an isolated system:

---



---

**Key Idea:** The factors affecting peroxidase activity can be measured using the indicator guaiacol.

**Enzymes** control all the metabolic activities required to sustain life. Changes to environmental conditions (e.g. pH or temperature) may alter an enzyme's shape and functionality.

This may result in decreased activity or complete loss of activity if the enzyme is denatured. In this activity you will use the information provided and your own understanding of enzymes to design an experiment to investigate factors affecting enzyme activity.

### Background

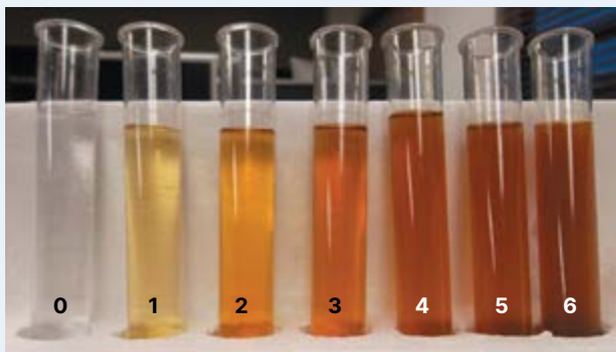
Hydrogen peroxide ( $H_2O_2$ ) is a toxic by-product of respiration and must be broken down in order to avoid cellular damage.

**Peroxidase** acts in the presence of naturally occurring organic reducing agents (electron donors) to catalyse the breakdown of  $H_2O_2$  into water and oxidised organic substrates.



Like all enzymes, the activity of peroxidase is highest within specific ranges of pH and temperature, and activity drops off or is halted altogether when the conditions fall outside of the optimal range. The conversion of  $H_2O_2$  is also influenced by other factors such as the levels of substrate and enzyme.

The effect of peroxidase on  $H_2O_2$  breakdown can be studied using a common reducing agent called guaiacol. Oxidation of guaiacol (as in the equation above) forms tetraguaiacol, which is a dark orange colour. The rate of the reaction can be followed by measuring the intensity of the orange colour as a function of time.



Increasing levels of oxygen production over time (minutes)

A time-colour palette is shown above. You can use it as a reference against which to compare your own results from the investigation below. The palette was produced by adding a set amount of peroxidase to a solution containing hydrogen peroxide and water. The colour change was recorded at set time points (0-6 minutes).

### Investigation 3.7 Investigating peroxidase activity

See appendix for equipment list.

1. Prepare six substrate tubes by 7 mL of distilled water, 0.3 mL of 0.1%  $H_2O_2$  solution, and 0.2 mL of prepared guaiacol solution to a boiling tube. Cover the tubes with parafilm and mix.
2. Prepare six enzyme tubes by adding 6.0 mL of prepared buffered pH solution (one of pH 3, 5, 6, 7, 8, and 10) and 1.5 mL of prepared turnip peroxidase solution. Cover the tubes with parafilm and mix.
3. Combine the contents of substrate and enzyme tubes and cover with parafilm. Mix and place back on the rack.
4. Begin timing immediately. Record the colour change every minute (1-6 based on the colour palette above).
5. You can take photos with your phones or keep a written record of the colour changes.

Colour reference number							
	0 min	1 min	2 min	3 min	4 min	5 min	6 min
pH 3							
pH 5							
pH 6							
pH 7							
pH 8							
pH 10							

1. The colour palette (above) shows the relative amounts of tetraguaiacol formed when guaiacol is oxidised. How can this be used to determine enzyme activity?



2. Graph your results on the grid (right).
3. (a) Describe the effect of pH on peroxidase activity:

---



---



---



---

- (b) Was there a colour change at pH 10? Explain the result at this pH and relate it to the enzyme's structure and the way it interacts with its substrate:

---



---



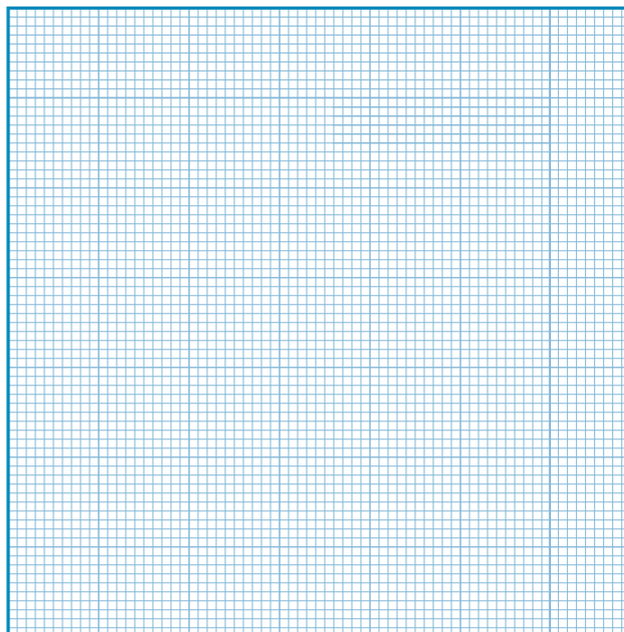
---



---



---



4. In your experiment, the rate of enzyme activity is measured by comparing against a ranked colour palette. How could you have measured the results more quantitatively?
- 
5. How might the results be affected if you did not begin timing immediately after mixing the contents of the enzyme and substrate tubes together?
- 
- 
6. Why is peroxidase written above the arrow in the equation for enzymatic breakdown of  $H_2O_2$ ?
- 
- 
7. Based on the information provided, and your answer to question 4, design an experiment to investigate the effect of lead nitrate (an enzyme inhibitor) on the activity of turnip peroxidase. You may also want to do some research on the internet to help. Summarise your method as step by step instructions below. Note how you will record and display the data and calculate the reaction rate. Include reference to any limitations or sources of potential error in your design:

---



---



---



---



---



---



---



---



---



---



---



---



---



---



---



---



---



---



---



---



---



---

# Achieving Metabolic Efficiency

**Key Idea:** Metabolic pathways are linked biochemical reactions that occur within organisms to maintain life. Metabolic reactions often occur as a linked series in which each step in the pathway relies on the completion of a previous step and each step is controlled by specific **enzymes**. The end product of one enzyme-controlled step provides the

substrate for the next, so failure of one step causes failure of all later steps. **Metabolic pathways** are tightly controlled to prevent energy being wasted. This energy conservation is termed metabolic efficiency. Metabolic reactions are often localised within specific organelles so that all the components of the pathway are kept together.

## Cellular compartments assist efficiency

To increase metabolic efficiency, regions within a cell or an organelle are compartmentalised (separated) by membranes. Particular metabolic reactions are restricted to certain regions where all the necessary metabolic components are located. Having compartments within the cell and within organelles prevents interference between different reaction pathways and enables radically different reaction environments to be accommodated within different organelles.

### Example: cellular respiration in the mitochondrion

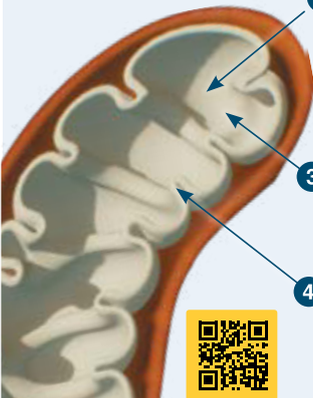
The membrane system of the mitochondrion divides it into several regions. Glycolysis takes place outside of the mitochondrion, in the cell's cytoplasm, but the remaining steps take place in different specialised regions of the mitochondrion. This helps to regulate movement of substrates and end-products and therefore reaction rates, increasing efficiency of the process (below).

#### 1 Cytoplasm (outside the mitochondrion): Glycolysis

2 **Matrix:** Link reaction. Link reaction enzymes (e.g. pyruvate dehydrogenase complex) are in the matrix.

3 **Matrix:** Krebs cycle. Krebs cycle enzymes (e.g. fumarase) are in the matrix.

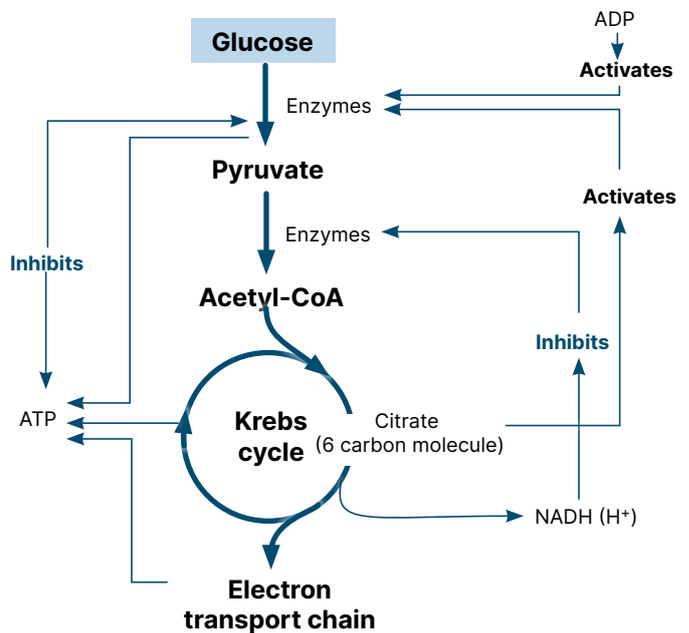
4 **Cristae:** Electron transport chain. Membrane-bound enzymes include ATP synthase



## Achieving efficiency by inhibition

Many metabolic pathways are controlled by feedback inhibition (negative feedback loop). The pathway is stopped when there is a build-up of end product (or certain intermediate products). The build-up stops the enzymes in the pathway from working and allows the cell to shut down a pathway when it is not needed. This conserves the cell's energy, so it is not manufacturing products it does not need.

Both linear pathways (e.g. glycolysis), and cyclic pathways (e.g. the Krebs cycle) can be regulated this way (below).



1. What does metabolic efficiency mean? \_\_\_\_\_

\_\_\_\_\_

2. Describe how cells achieve metabolic efficiency through:

(a) Compartmentalising: \_\_\_\_\_

\_\_\_\_\_

(b) Feedback inhibition: \_\_\_\_\_

\_\_\_\_\_

3. What would happen if cells could not regulate their metabolic pathways? \_\_\_\_\_

\_\_\_\_\_

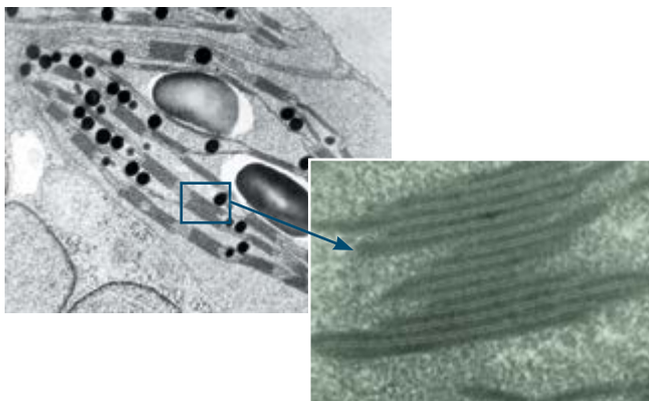


# 99 Enzymes and Membranes

**Key Idea:** Increasing the surface area for enzyme attachment increases the number of enzymes that can be present and so also increases the rate of biochemical reactions.

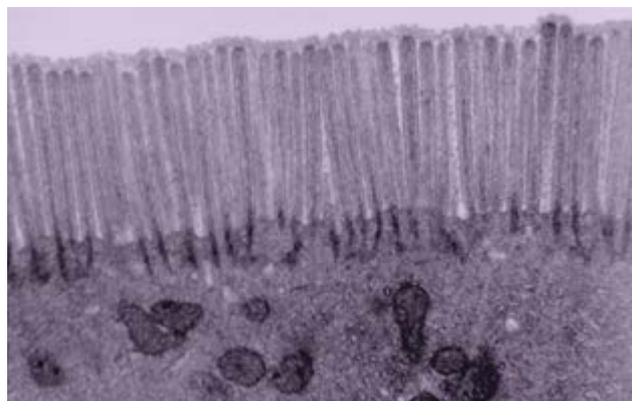
Chemical reactions in organisms must occur at a relatively fast rate. If too slow, the products produced will never be available in the quantities needed to maintain life. One way

to increase reaction rate is to concentrate certain **enzymes** in areas where their substrates are also concentrated. Another way is to increase the surface area for attachment of enzymes so that the number of reactions occurring over any amount of time can be increased. This occurs in many membranous organelles and also in the gut microvilli.



Dartmouth College

Chloroplasts have internal membranes (thylakoid membranes) organised into stacks called grana. Embedded into the membranes are the protein complexes that capture light and catalyse the light dependent reactions of photosynthesis. The large surface area provided by the thylakoid membranes increases the membrane area for protein attachment and thus the amount of light captured. It also increases the area for associated enzymes that use the captured light to move protons across the membrane and generate ATP.



Intestinal epithelial cells are found lining the villi of the intestinal wall. The cell surface projecting out to the intestinal lumen (space) is covered with microvilli (projections of the plasma membrane). These increase the surface area for absorbing molecules from food in the lumen and increase the surface area for attachment of enzymes that carry out the final stages of carbohydrate digestion. There are also transporter proteins embedded in the membrane. These transport the products of this digestion into the cell.



The enzymes of the electron transport chain in cellular respiration are embedded in the internal membranes of the mitochondria. The internal membrane is folded into structures called cristae. These increase the surface area for enzyme attachment and allow a much greater reaction rate.



The endoplasmic reticulum is a large region of folded membrane that is attached to the nuclear membrane. Numerous types of enzymes are embedded in the membranes of this organelle, including ribosomes, which catalyse protein synthesis (on the rough endoplasmic reticulum).

- How do cells increase cellular membrane surface area? \_\_\_\_\_  
\_\_\_\_\_
- (a) How does increasing surface area of cellular membranes help to increase reaction rates in cells?  
\_\_\_\_\_  
\_\_\_\_\_
  - Why is increasing the reaction rate important in cells? \_\_\_\_\_  
\_\_\_\_\_  
\_\_\_\_\_

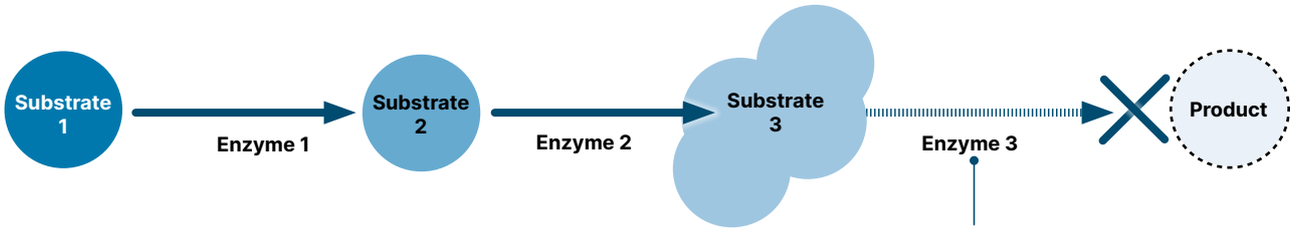
# 100 Enzymes and Disease

**Key Idea:** When enzymes fail to work correctly and do not catalyse the biochemical reactions of metabolism, serious metabolic disorders can occur.

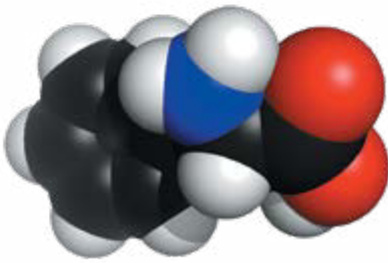
**Metabolism** is all the chemical activities of life. **Enzymes** play an important part in metabolism. They catalyse the reactions that convert one substrate into another. If any of the enzymes in a **metabolic pathway** fail to work correctly then

intermediate substrates or metabolic waste products can build up in the body and cause numerous different metabolic disorders. Many of these disorders can be identified by simple blood tests. The heel prick test or Guthrie test carried out on newborns tests for several metabolic and genetic disorders. An example of a well studied metabolic pathway, the metabolism of phenylalanine, is described below.

## A simple metabolic pathway



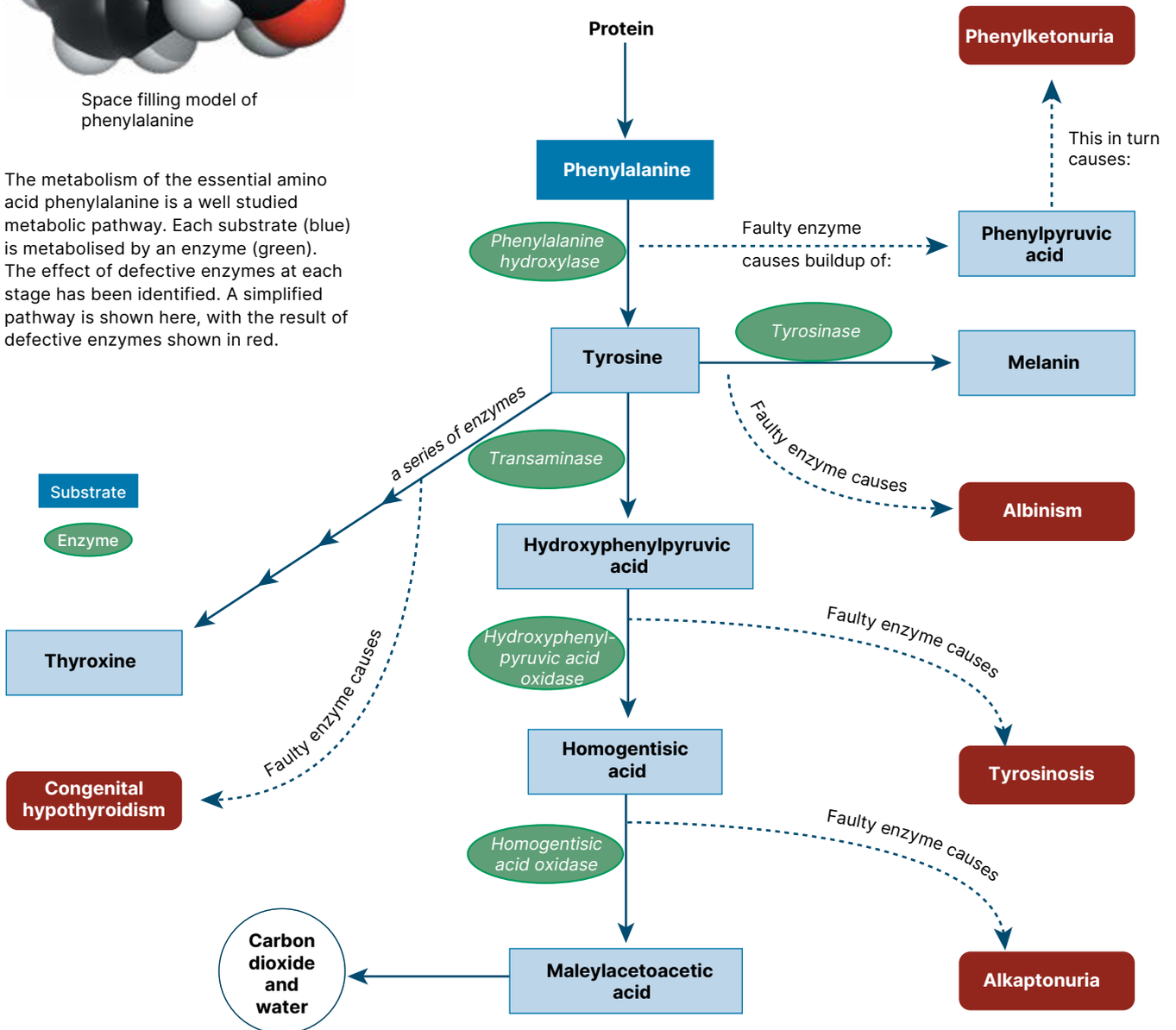
Enzyme 3 is not expressed or is nonfunctional. The reaction series cannot progress to the end product, so levels of substrate 3 build up. If substrate 3 is a toxic substance, this can have serious health effects.



Space filling model of phenylalanine

The metabolism of the essential amino acid phenylalanine is a well studied metabolic pathway. Each substrate (blue) is metabolised by an enzyme (green). The effect of defective enzymes at each stage has been identified. A simplified pathway is shown here, with the result of defective enzymes shown in red.

## The metabolism of phenylalanine





Melanin is the pigment that gives skin, hair, and eyes their colour. It is formed from the metabolism of the amino acid tyrosine. Lack of melanin results in albinism, a condition where there is little or no pigmentation (above). The most common cause of albinism is a faulty enzyme in the pathway that converts tyrosine into melanin.



Phenylketonuria (PKU) is an example of a metabolic disorder that occurs when there is an error in a metabolic pathway. Babies born with PKU are missing the enzyme needed to catalyse the first step in the pathway that metabolises the essential amino acid phenylalanine. Without the enzyme, phenylalanine cannot be converted to the next substrate, tyrosine, and it is metabolised to toxic derivatives, which cause central nervous system damage. Children with PKU tend to have lighter skin and hair than people without the disorder.

Newborn babies are tested for a number of genetic disorders, including PKU, soon after birth. Blood is collected from a heel prick on to a Guthrie card (above) and tested. The prognosis is good if the disease is detected early, and a low phenylalanine diet is followed throughout life. People with PKU must also take supplements to provide the amino acids that would otherwise be lacking in a low-phenylalanine diet (e.g. tyrosine, which is normally derived from phenylalanine and which is needed for brain function).

- Identify four end products of the normal metabolism of phenylalanine: \_\_\_\_\_  
\_\_\_\_\_
- In the metabolic pathway of the metabolism of phenylalanine, identify the faulty enzyme that results in:
  - Albinism: \_\_\_\_\_
  - Phenylketonuria: \_\_\_\_\_
  - Tyrosinosis: \_\_\_\_\_
  - Alkaptonuria: \_\_\_\_\_
- Why do people with phenylketonuria have light skin? \_\_\_\_\_  
\_\_\_\_\_  
\_\_\_\_\_
- Using the metabolism of phenylalanine as an example, discuss the role of enzymes in metabolic pathways:  
\_\_\_\_\_  
\_\_\_\_\_  
\_\_\_\_\_  
\_\_\_\_\_
- The conditions illustrated in the diagram are due to too much or too little of a chemical in the body. For each condition listed below, state which chemical causes the problem and whether it is absent or present in excess:
  - Albinism: \_\_\_\_\_
  - Phenylketonuria: \_\_\_\_\_
  - Congenital hypothyroidism: \_\_\_\_\_

1. Write a definition for the terms below:

(a) Denaturation: \_\_\_\_\_  
 \_\_\_\_\_

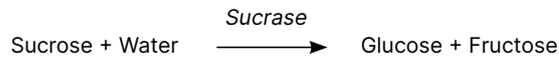
(b) Induced fit model: \_\_\_\_\_  
 \_\_\_\_\_

(c) Catalyst: \_\_\_\_\_  
 \_\_\_\_\_

2. A specific enzyme has an optimum temperature of 30°C. Explain what this means and what would happen if the enzyme was in temperatures above and below this:

\_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_

3. Study the enzymatic word equation below and answer the following questions:



(a) Identify the substrate: \_\_\_\_\_

(b) Identify the products: \_\_\_\_\_

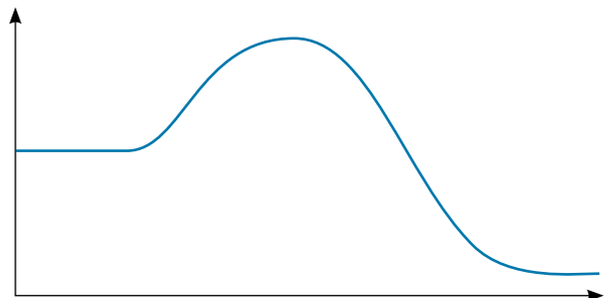
(c) Identify the enzyme: \_\_\_\_\_

4. Some heavy metals act as irreversible competitive inhibitors. Why does this make them dangerous poisons?

\_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_

5. (a) Label the graph right with axes and the following labels: *Reactants*, *products*, *activation energy*, *transition state*.

(b) Assume the reaction has had no enzyme added. Draw the shape of the graph when an enzyme is added to the reaction mix.

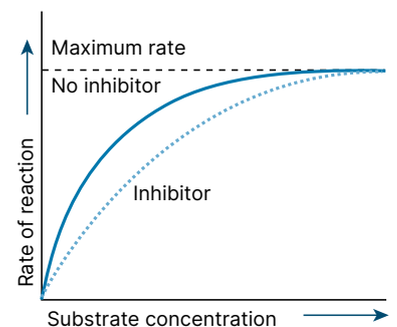
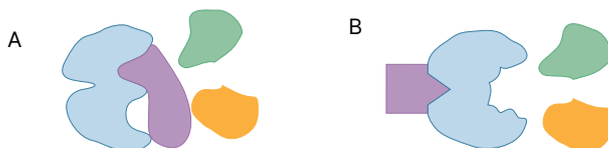


6. The graph (right) shows the effect of an enzyme inhibitor in enzyme reaction rate.

(a) Does the graph show competitive inhibition or non-competitive inhibition?

\_\_\_\_\_

(b) Identify the diagram below that illustrates your choice in (a): \_\_\_\_\_



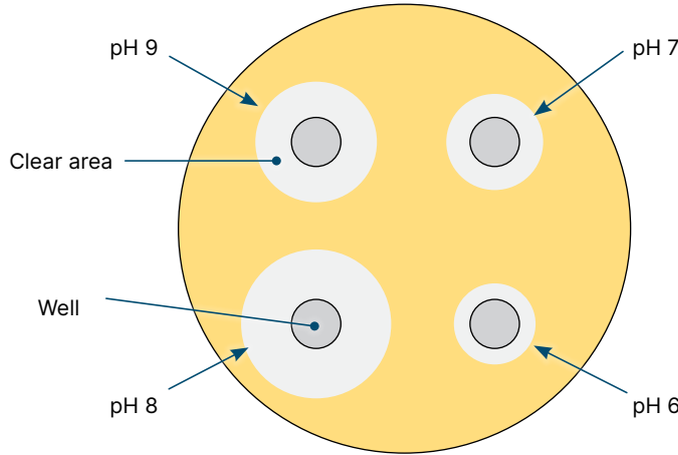
7. Identify two ways organelles can increase the rate and efficiency of metabolic reactions:

\_\_\_\_\_  
 \_\_\_\_\_



5. A student wanted to investigate the effect of pH on a peptidase (a protease) produced in the small intestine of a human. She used an agar plate containing protein. This made the agar plate cloudy. Digestion of the protein by the peptidase makes the agar clear.

- The agar plate was made with four equal sized wells into which the peptidase could be added. Four different mixtures of peptidase at different pH were produced and added to the wells.
- The set up was incubated at a constant temperature for 4 hours.
- The diagram below shows the results:



(a) Suggest a temperature for incubation of the agar plate for best results: \_\_\_\_\_

(b) Why did you choose this temperature? \_\_\_\_\_  
 \_\_\_\_\_

(c) Which pH produced the best result? \_\_\_\_\_

(d) Explain why: \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_

6. Enzymes are essential for catalysing biological reactions. Describe the induced fit model of enzyme action:

\_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_

7. Identify the following statements as true (T) or false (F):

(a) Enzymes are biological catalysts. They lower the activation energy of a reaction: \_\_\_\_\_

(b) Competitive inhibition is when an inhibitor binds to a site other than the active site: \_\_\_\_\_

(c) The induced fit model states that the enzyme changes shape when a substrate fits into the active site: \_\_\_\_\_

(d) End product inhibition causes a feedback loop that escalates the outcome of the loop: \_\_\_\_\_

8. In the space below draw an annotated diagram to show how a metabolic pathway can be regulated by enzymes and what happens if an enzyme does not work correctly.



# Respiration and Mammalian Gas Exchange

## Key Terms

- aerobic respiration
- alveoli
- anaerobic respiration
- anabolic reaction
- ATP
- capillaries
- catabolic reaction
- cellular respiration
- electron transport chain
- enzyme
- fermentation
- gas exchange
- glycolysis
- haemoglobin
- Krebs cycle
- lung
- metabolism
- mitochondria
- respiratory gases
- respiratory system
- respirometer

## Key Concepts

- ▶ Cellular respiration provides the energy required for the survival of an organism.
- ▶ Cellular respiration can occur with and without oxygen through the use of different biochemical pathways.
- ▶ The gas exchange system in humans allows for efficient exchange of respiratory gases.

## Obtaining energy for life

### Activity Number

- |     |   |     |
|-----|---|-----|
| □ 1 | Define metabolism as all the chemical processes occurring within a living organism to maintain life. Distinguish between catabolism and anabolism.  | 103 |
| □ 2 | Explain the role of ATP (adenosine triphosphate) as an energy carrier in cells. Explain why organisms need to respire, recalling the universal role of ATP in metabolism.   | 104 |
| □ 3 | Explain how cellular respiration functions as energy transformation processes. Include reference to the relationship between the raw materials and products of the process. Explain diagrams and schematics of energy transfer. | 104 |

## Cellular respiration

- |      |  |     |
|------|--|-----|
| □ 4  | Recognise that organisms can respire aerobically and anaerobically.  | 104 |
| □ 5  | Investigate respiration in germinating seeds.  | 105 |
| □ 6  | Recognise that cellular respiration is an enzyme-controlled series of chemical reactions and that aerobic respiration requires oxygen.   | 106 |
| □ 7  | Identify the main steps in the complete oxidation of glucose by aerobic cellular respiration: glycolysis, Krebs cycle, and electron transport chain.   | 106 |
| □ 8  | Summarise the reactions of aerobic respiration in a word equation and in a chemical equation, including the ATP yield.   | 106 |
| □ 9  | Describe anaerobic (without oxygen) pathways for ATP generation in eukaryotes: lactic acid fermentation in mammalian muscle and alcoholic fermentation in yeast and plant roots. Compare the energy yield from aerobic and anaerobic pathways. | 107 |
| □ 10 | Measure the outputs of fermentation in yeast grown on different substrates.  | 108 |

## Gas exchange

- |      |   |         |
|------|---|---------|
| □ 11 | Distinguish between cellular respiration and gas exchange and explain why organisms must exchange respiratory gases with their environment and why they need specialised gas exchange surfaces. Name the raw materials and waste products of cellular respiration.                                  | 109     |
| □ 12 | Describe the gross structure of the human gas exchange system, explaining how the cells, tissues, and organs function together to exchange respiratory gases between the blood and the environment.   | 110     |
| □ 13 | Describe the structural features of gas exchange surfaces and explain the significance of surface area : volume ratio to the exchange of materials, such as respiratory gases, with the environment.  | 111     |
| □ 14 | Explain how respiratory gases move across the gas exchange membrane. Use data presented in diagrams and schematics to predict the direction in which materials will be exchanged between the alveoli and capillaries and between the capillaries and the tissues of the body (e.g. muscle, organs). | 112     |
| □ 15 | <b>SI:</b> Investigate a range of adaptations that allow gas exchange to occur efficiently in animals.  | 109-112 |

# 103 Metabolism and Life

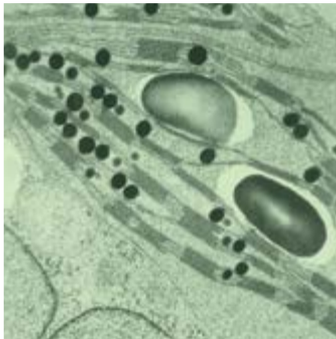
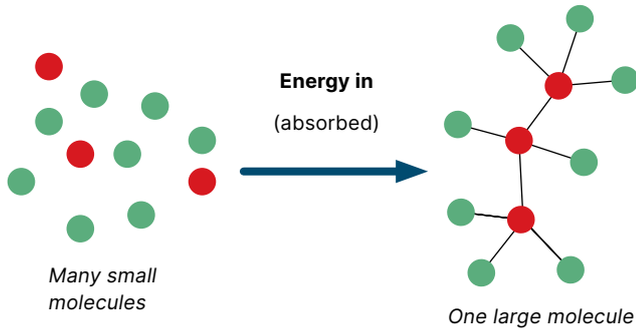
**Key Idea:** Enzymes catalyse all of the body's metabolic reactions. Homeostasis helps to maintain the physiological conditions required for optimum enzyme activity.

**Metabolism** is defined as all the chemical processes occurring within a living organism to maintain life. **Enzymes** catalyse each of these metabolic reactions. Recall that enzymes have a narrow range of physical conditions for optimum activity. Regional specialisation within the body or its cells

helps to provide these conditions. Outside the optimum range, enzyme activity drops off or stops. When this occurs, metabolic pathways are impaired and there are detrimental or even fatal effects on the organism. For example, acidosis is a condition in which blood pH falls. Enzyme activity is impaired and this depresses organ function. Homeostasis maintains narrow physiological conditions, such as pH, so that metabolic reactions can proceed normally.

## Anabolic reactions

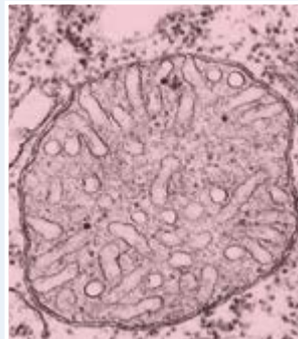
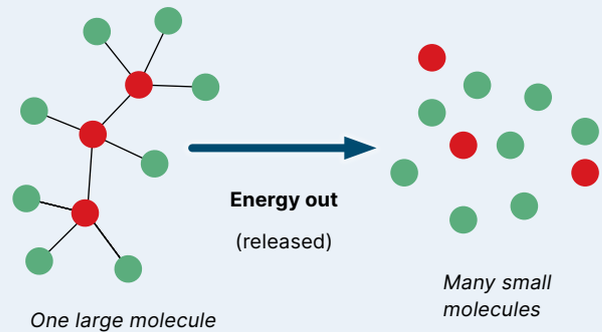
- ▶ **Anabolic reactions** are reactions that result in the production (synthesis) of a more complex molecule from smaller components or smaller molecules. During anabolic reactions, simple molecules are joined to form a larger, more complex molecule.
- ▶ Anabolic reactions need a net input of energy to proceed. They are called endergonic reactions.



Photosynthesis is an example of an anabolic reaction pathway (many reactions are involved). Plants carry out photosynthesis in organelles called chloroplasts (left). Photosynthesis is an anabolic process because it converts carbon dioxide and water into glucose. Energy from the sun is required to drive photosynthesis.

## Catabolic reactions

- ▶ **Catabolic reactions** are reactions that break down large molecules into smaller components.
- ▶ Catabolic reactions involve a net release of energy. They are called exergonic reactions.
- ▶ The energy released from catabolic reactions can be used to drive other metabolic processes.



**Cellular respiration** is an example of a catabolic reaction pathway. Glucose is broken down in a series of reactions to release carbon dioxide, water, and ATP (energy). The energy in ATP is used to fuel other activities in the cell. Most of the reactions of cellular respiration take place in the **mitochondria** (left).

1. What is an anabolic reaction? \_\_\_\_\_  
\_\_\_\_\_
2. (a) What is a catabolic reaction? \_\_\_\_\_  
\_\_\_\_\_
 

(b) Why are catabolic reactions considered to be the opposite to anabolic reactions? \_\_\_\_\_  
\_\_\_\_\_
3. Acidosis is a drop of blood pH to below 7.35 (normal blood pH is 7.35-7.45), which can be caused by kidney malfunction, alcohol intoxication, or starvation. Explain how acidosis might affect enzyme functioning and metabolism: \_\_\_\_\_  
\_\_\_\_\_  
\_\_\_\_\_

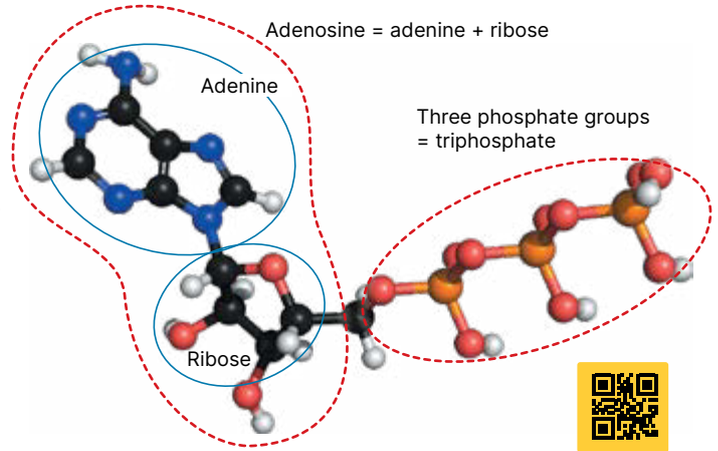
**Key Idea:** ATP transports chemical energy within the cell for use in metabolic processes.

All organisms require energy to allow them to perform the metabolic processes required to function and reproduce. This energy is obtained by **cellular respiration**, a set of metabolic reactions which ultimately convert biochemical energy

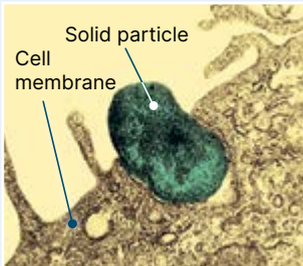
from 'food' into the nucleotide **adenosine triphosphate (ATP)**. ATP is considered to be a universal energy carrier, transferring chemical energy within the cell for use in metabolic processes such as biosynthesis, cell division, cell signalling, thermoregulation, cell mobility, and active transport of substances across membranes.

## Adenosine triphosphate (ATP)

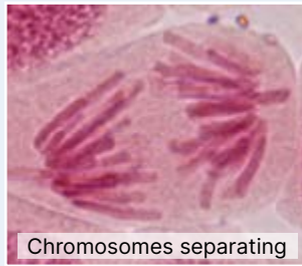
- ▶ The ATP molecule consists of three components: a purine base (adenine), a pentose sugar (ribose), and three phosphate groups that attach to the 5' carbon of the pentose sugar. Adenine + ribose form adenosine (the 'A' in ATP). The structure of ATP is shown right.
- ▶ The bonds between the phosphate groups contain electrons in a high energy state which store a large amount of energy. This is released during ATP hydrolysis. Typically, hydrolysis is coupled to another cellular reaction to which the energy is transferred. The end products of the reaction are adenosine diphosphate (ADP) and an inorganic phosphate (Pi).
- ▶ Note that energy is released during the formation of bonds during the hydrolysis reaction, not the breaking of bonds between the phosphates (which requires energy input).



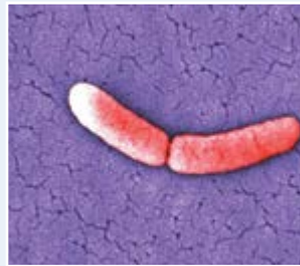
### ATP powers life processes in the cell



The energy released from the removal of a phosphate group of ATP is used for active transport of molecules and substances across the plasma membrane e.g. phagocytosis (above) and other active transport processes.



Mitosis, as seen in the stained onion cell above, requires ATP to proceed. Formation of the mitotic spindle and chromosome separation both require the energy provided by ATP hydrolysis to occur.

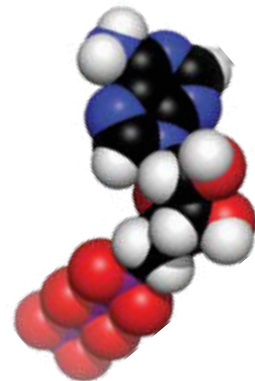


ATP is required when bacteria divide by binary fission (above). For example, ATP is required in DNA replication and to synthesise components of the peptidoglycan cell wall.



Not all of the energy released in the oxidation of glucose is captured in ATP. The rest is lost as heat. This heat energy can be used to maintain body temperature. Thermoregulatory mechanisms such as shivering and sweating also use ATP.

1. What process produces ATP in a cell?  
\_\_\_\_\_
2. On the space filling model of ATP shown right, label adenine, ribose, and the phosphate groups:  
\_\_\_\_\_
3. Explain why thermoregulation requires the expenditure of energy:  
\_\_\_\_\_  
\_\_\_\_\_  
\_\_\_\_\_  
\_\_\_\_\_
4. Describe one other process in a cell that requires ATP: \_\_\_\_\_  
\_\_\_\_\_  
\_\_\_\_\_  
\_\_\_\_\_



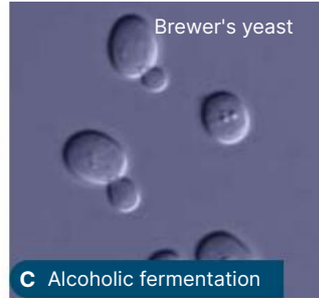
## Aerobic and anaerobic pathways for ATP production



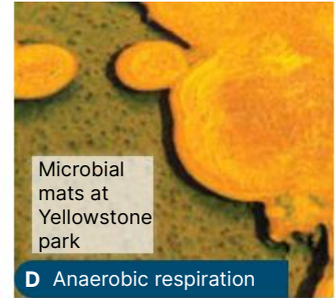
**A** Aerobic respiration



**B** Lactic acid fermentation



**C** Alcoholic fermentation



**D** Anaerobic respiration

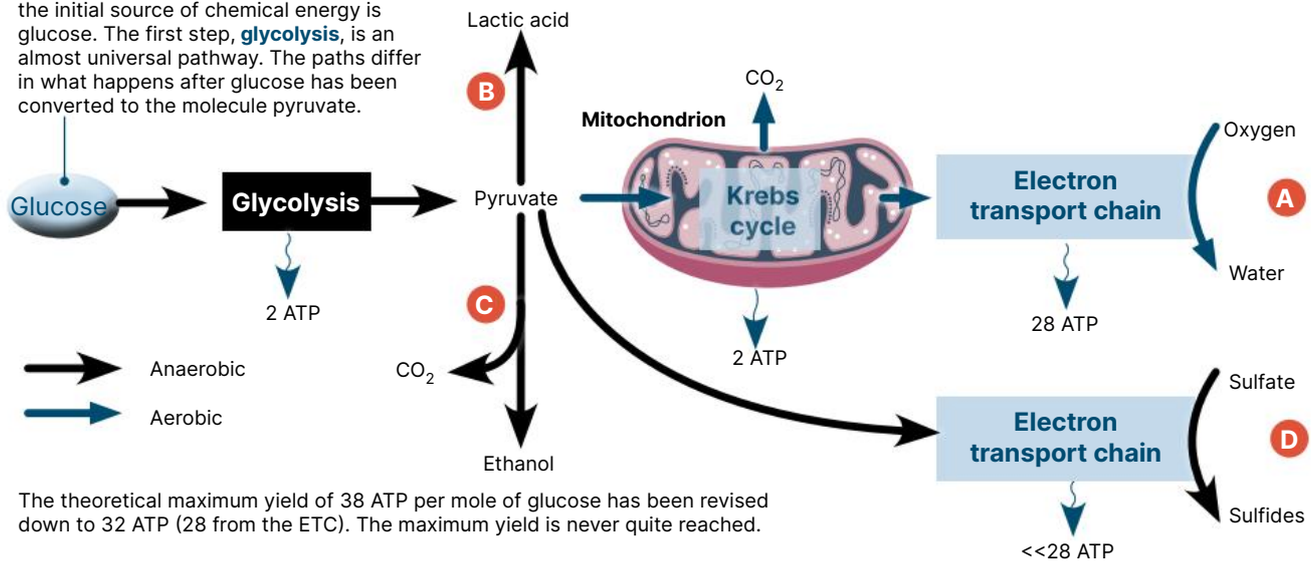
**Aerobic respiration** produces the energy (as ATP) needed for metabolism. The rate of aerobic respiration is limited by the amount of oxygen available. In animals and plants, most of the time the oxygen supply is sufficient to maintain aerobic **metabolism**. Aerobic respiration produces a high yield of ATP per molecule of glucose (**path A**).

During maximum physical activity, when oxygen is limited, **anaerobic respiration** provides ATP for working muscle. In mammalian muscle, metabolism of a respiratory intermediate produces lactate, which provides fuel for working muscle and produces a low yield of ATP. This process is called **lactic acid fermentation** (**path B**).

The process of brewing utilises the anaerobic metabolism of yeasts. Brewer's yeasts preferentially use anaerobic metabolism in the presence of excess sugars. This process, called **alcoholic fermentation**, produces ethanol and CO<sub>2</sub> from the respiratory intermediate pyruvate. It is carried out in vats that prevent entry of O<sub>2</sub> (**path C**).

Many bacteria and archaea are anaerobic, using molecules other than oxygen, e.g. nitrate or sulfate, as a terminal electron acceptor of their **electron transport chain**. These electron acceptors are not as efficient as oxygen (less energy is released per oxidised molecule) so the energy (ATP) yield from anaerobic respiration is generally quite low (**path D**).

In most energy-yielding pathways, the initial source of chemical energy is glucose. The first step, **glycolysis**, is an almost universal pathway. The paths differ in what happens after glucose has been converted to the molecule pyruvate.



The theoretical maximum yield of 38 ATP per mole of glucose has been revised down to 32 ATP (28 from the ETC). The maximum yield is never quite reached.

5. What do all the ATP yielding pathways above have in common? \_\_\_\_\_
  
6. Distinguish between anaerobic pathways in eukaryotes (e.g. yeasts) and anaerobic respiration in anaerobic microbes: \_\_\_\_\_
  
7. When brewing alcohol, why is it important to prevent entry of oxygen to the fermentation vats? \_\_\_\_\_
  
8. Rank the following processes from lowest to highest in terms of ATP produced (use = for processes you consider equal in yield): *lactic acid fermentation, anaerobic respiration, alcoholic fermentation, aerobic respiration, glycolysis*. \_\_\_\_\_

# 105 Measuring Respiration

**Key Idea:** Respiration is the process by which cells convert energy in glucose to usable energy, which is stored in the molecule ATP. The process uses oxygen, which can be quantified using a simple respirometer.

A **respirometer** can be used to measure the amount of oxygen consumed by an organism during **cellular respiration** and so can be used to measure **respiration** rate. A simple

respirometer is shown in the diagram below. The carbon dioxide produced during respiration is absorbed by the potassium hydroxide. As the oxygen is used up, the coloured bubble in the glass tube moves. Measuring the movement of the bubble (e.g. with a ruler or taped graph paper) allows an estimation of the change in volume of gas and therefore the rate of cellular respiration.



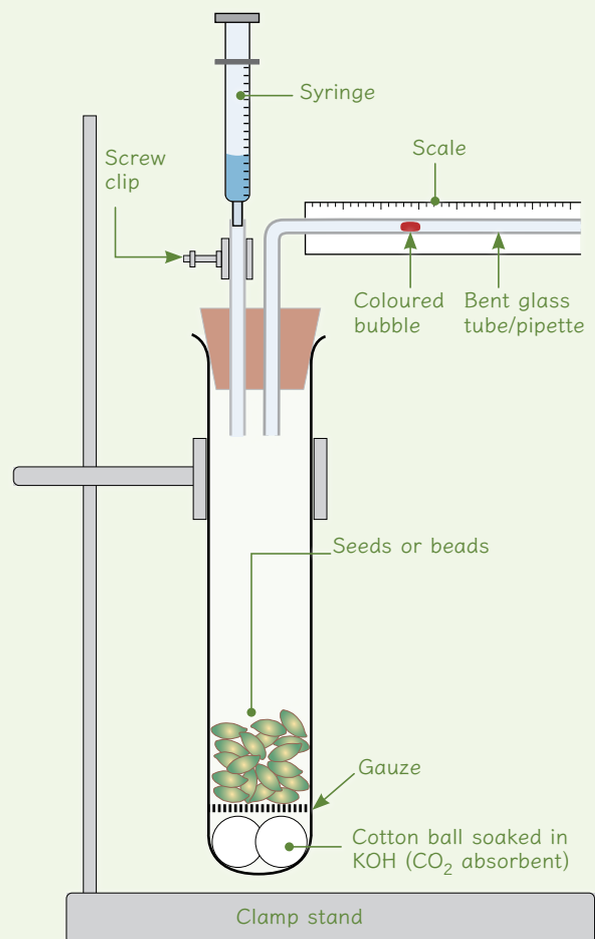
## Investigation 7.1 Measuring respiration in germinating seeds

See appendix for equipment list.



**Caution is required when handling potassium hydroxide as it is caustic and can cause chemical burns. You should wear protective eyewear and gloves.**

1. Work in groups of four to set up three respirometers using the setup shown right as a guide.
2. Collect three boiling tubes and place two cotton balls in the bottom of each. Label the tubes A, B, and C.
3. Use a dropper to add 15% potassium hydroxide (KOH) solution on to the cotton balls until they are saturated (there should be no liquid in the boiling tube). Add the same amount of KOH to the cotton balls in each boiling tube.
4. Place gauze on top of the cotton balls in each tube. This prevents the KOH coming into contact with the seeds and killing them.
5. Quarter fill tube A with germinated bean seeds. These seeds will be damp because they have been germinated under damp paper towels for four days.
6. Quarter fill tube B with ungerminated (dry) seeds.
7. Quarter fill tube C with glass beads.
8. Place a two-hole stopper firmly in each boiling tube. In one hole insert a bent glass tube or bent pipette. In the second hole insert a tube that can be clamped shut using a screw clip.
9. Use a dropper or fine pipette to place a drop of coloured liquid into the bent tube/pipette of each set up. Attach a syringe to the clamped tube. Open the screw clip and use the syringe to draw the coloured bubble into the middle of the bent tube/pipette.
10. Place all three tubes in a water bath at 25°C. Secure them with a clamp stand or in racks.
11. Leave the apparatus to equilibrate for 10 minutes.
12. At the end of the equilibration period, close the screw clip on the boiling tubes. Mark the position of the bubble with a marker pen. This is your time zero position.
13. Start the timer.
14. Use a ruler or the pipette's scale (if there is one) to measure the distance the coloured bubble moved at 5, 10, 15, 20, and 25 minutes.
15. Record your results on the table at the top of the next page.



Respirometers of this sort are very sensitive to poor procedure because the volumes involved are so small.

Be very careful with your set-up and when taking readings. Have one person responsible taking the measurements of the bubble movement.



Time (minutes)	Distance bubble moved (mm)		
	Germinated seeds (A)	Ungerminated seeds (B)	Glass beads (C)
0			
5			
10			
15			
20			
25			

1. What is the purpose of the test tube with the beads?

---



---

2. (a) Calculate the corrected distance the bubble moved in tubes A and B by subtracting the distance moved in tube C from each value. Record these values in the table below.

(b) Use the corrected distance the bubble moved to calculate the rate of respiration. Record this in the table below:

Time (minutes)	Corrected distance bubble moved (mm)		Rate (mm/min)	
	Germinated seeds (A)	Ungerminated seeds (B)	Germinated seeds (A)	Ungerminated seeds (B)
0				
5				
10				
15				
20				
25				

(c) Plot the rate of respiration on the grid (right). Include appropriate titles and axis labels:

(d) What does your plot show?

---



---



---



---



---



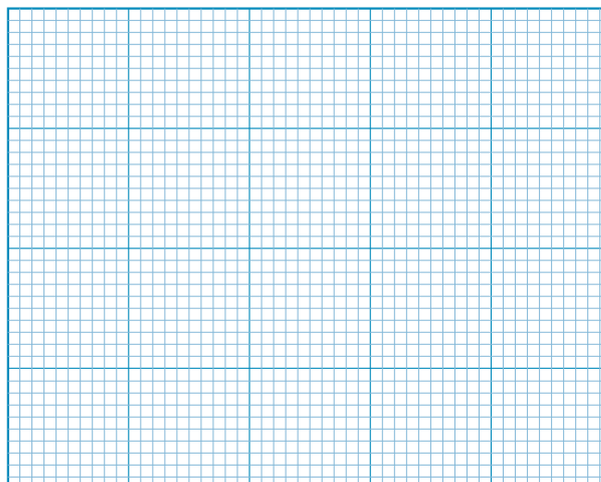
---



---



---



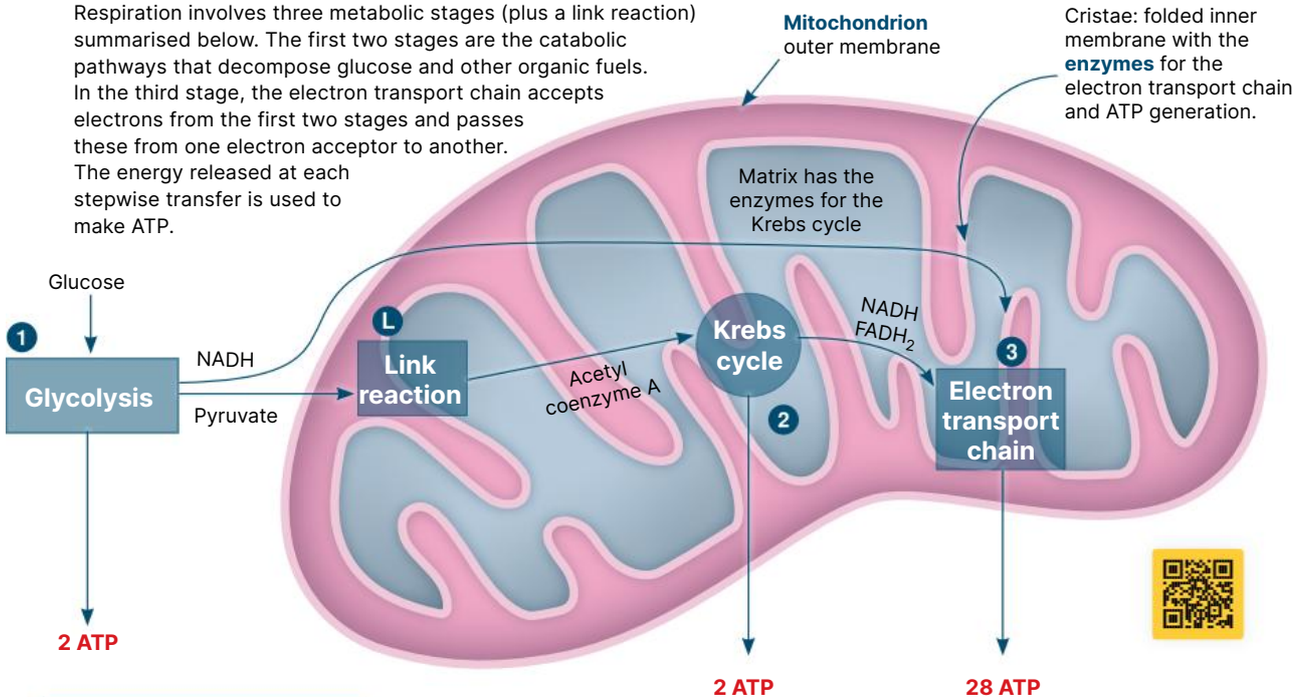
# Cellular Respiration: Inputs and Outputs

**Key Idea:** During cellular respiration, the energy in glucose is transferred to ATP in a series of enzyme controlled steps. The oxidation of glucose is a catabolic, energy yielding pathway. The breakdown of glucose and other organic fuels to simpler molecules is coupled to **ATP** synthesis. **Glycolysis** and the **Krebs cycle** supply electrons to the **electron transport chain** (ETC) which drives oxidative phosphorylation. The conversion of pyruvate (the end

product of glycolysis) to acetyl CoA links glycolysis to the Krebs cycle. Most of the ATP generated in cellular respiration is produced by oxidative phosphorylation when NADH + H<sup>+</sup> and FADH<sub>2</sub> donate electrons to the electron carriers in the ETC. At the end of the chain, electrons are passed to molecular oxygen, reducing it to water. Electron transport produces an H<sup>+</sup> gradient. As H<sup>+</sup> flows down its gradient, ATP synthase produces ATP which can be used by the cell.

## Overview of cellular respiration

Respiration involves three metabolic stages (plus a link reaction) summarised below. The first two stages are the catabolic pathways that decompose glucose and other organic fuels. In the third stage, the electron transport chain accepts electrons from the first two stages and passes these from one electron acceptor to another. The energy released at each stepwise transfer is used to make ATP.



The older stated theoretical maximum of 38 ATP per mole of glucose has now been revised down to 32 ATP (28 from the ETC). Inefficiencies in the process reduce the yield.



- Describe precisely in which part of the cell the following take place:
  - Glycolysis: \_\_\_\_\_
  - Krebs cycle reactions: \_\_\_\_\_
  - Electron transport chain: \_\_\_\_\_
- Write a word equation for the general equation for cellular respiration: \_\_\_\_\_
- How many ATP molecules are produced from one glucose molecule during aerobic respiration? \_\_\_\_\_
  - If one mole of glucose contains 2870 kJ of energy, and one mole of ATP releases 30.7 kJ of energy during a reaction, what is the percentage of energy in glucose that is available for the body to use?  
 \_\_\_\_\_  
 \_\_\_\_\_
- What is the purpose of NADH and FADH<sub>2</sub> in cellular respiration? \_\_\_\_\_
- Name three functions of glycolysis in cellular respiration: \_\_\_\_\_  
 \_\_\_\_\_



### Steps in cellular respiration

#### Glycolysis

Glycolysis is the beginning of cellular respiration. It takes glucose and produces two pyruvate molecules, each of which can then enter the Krebs cycle. Glycolysis initially uses two ATP but produces four ATP. NADH is produced for use in the electron transport chain. **The numbers shown are for one glucose molecule.**

#### Link reaction

The link reaction removes CO<sub>2</sub> from pyruvate and adds coenzyme A, producing the 2C molecule acetyl coenzyme A, which enters the Krebs cycle. NADH is also produced and flows to the electron transport chain.

#### Krebs cycle

In the Krebs cycle, acetyl coenzyme A is attached to the 4C molecule oxaloacetate and coenzyme A is released. Oxaloacetate is eventually remade in a cyclic series of reactions that produce more NADH and FADH<sub>2</sub> for the electron transport chain. Two ATP are also made by substrate level phosphorylation.

#### Electron transport chain (ETC)

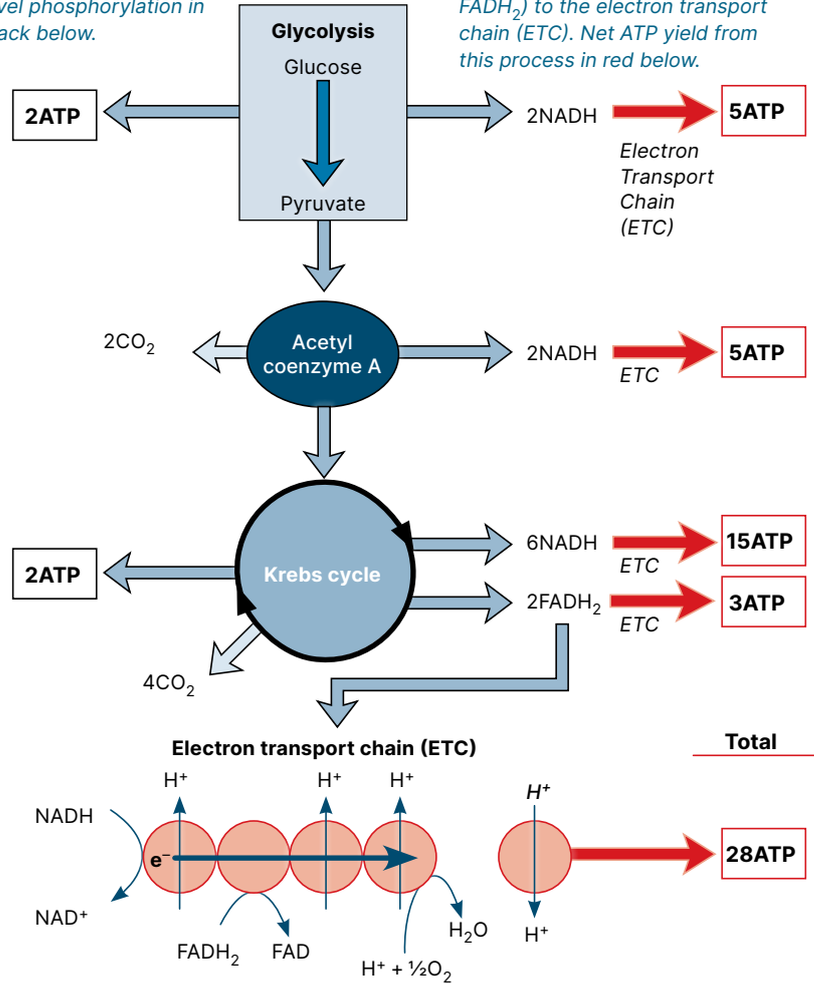
Electrons carried by NADH and FADH<sub>2</sub> are passed to a series of electron carrier enzymes embedded in the inner membrane of the mitochondria. The energy from the electrons is used to pump H<sup>+</sup> ions across the inner membrane from the matrix into the intermembrane space. These are allowed to flow back to the matrix via the enzyme ATP synthase which uses their energy to produce ATP. The electrons are coupled to H<sup>+</sup> and oxygen at the end of the electron transport chain to form water.

#### Substrate level phosphorylation

An enzyme transfers a phosphate group directly from a substrate (such as glucose) to ADP to form ATP. Net ATP yield from substrate level phosphorylation in black below.

#### Oxidative phosphorylation

Glucose is oxidised in a series of reduction and oxidation reactions that provide the energy to form ATP. This is achieved by the flow of reducing power (as NADH and FADH<sub>2</sub>) to the electron transport chain (ETC). Net ATP yield from this process in red below.



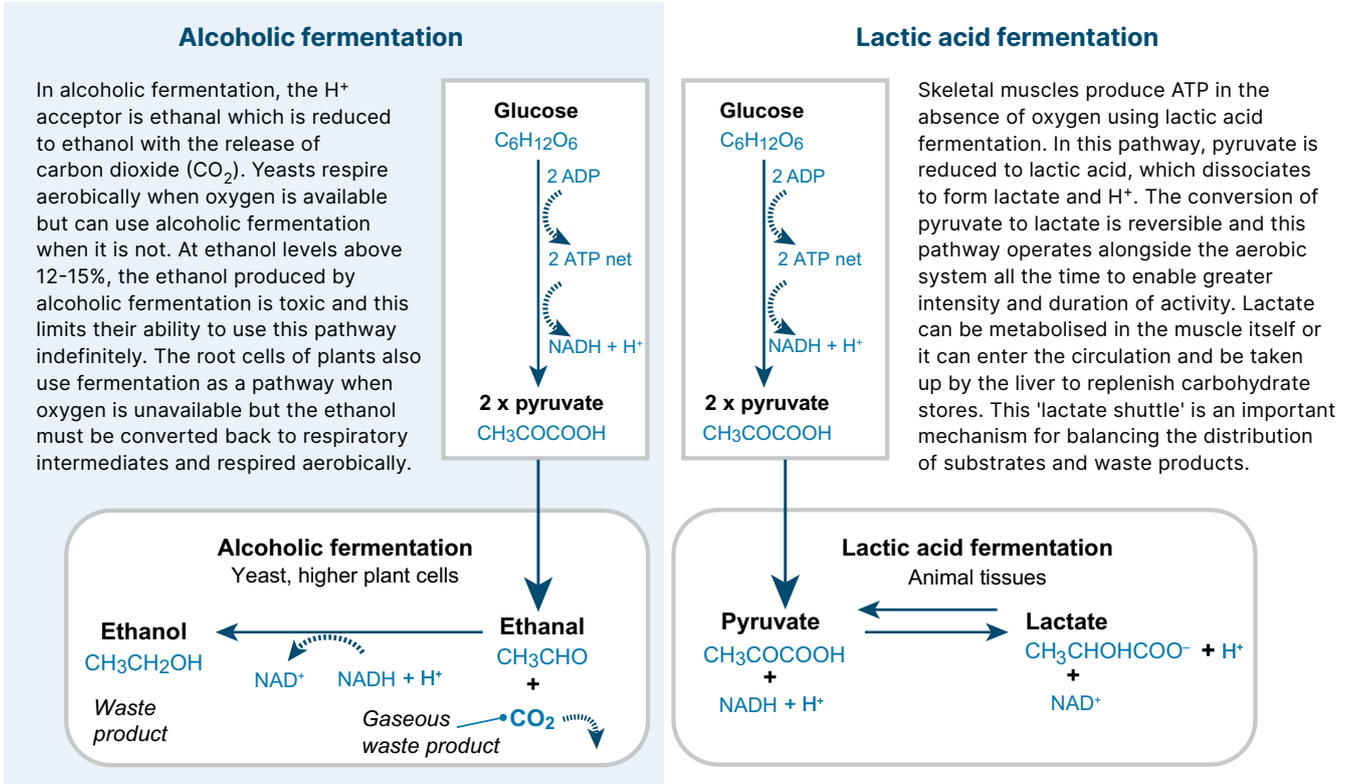
- Name two functions of the Krebs cycle in cellular respiration: \_\_\_\_\_
- (a) What is substrate level phosphorylation? \_\_\_\_\_  
 (b) How many ATP are produced this way during cellular respiration (per molecule of glucose)? \_\_\_\_\_
- (a) What is oxidative phosphorylation? \_\_\_\_\_  
 (b) How many ATP are produced this way during cellular respiration (per molecule of glucose)? \_\_\_\_\_
- Which parts of cellular respiration produce CO<sub>2</sub>? \_\_\_\_\_
- Describe how ATP is produced in the electron transport chain: \_\_\_\_\_

# Anaerobic Pathways

**Key Idea:** Glucose can be metabolised aerobically and anaerobically to produce ATP. The ATP yield from aerobic processes is higher than from anaerobic processes.

**Aerobic respiration** occurs in the presence of oxygen. Organisms can also generate **ATP** when oxygen is absent by

using a molecule other than oxygen as the terminal electron acceptor for the pathway. In alcoholic **fermentation** in yeasts, the electron acceptor is ethanal. In lactic acid fermentation, which occurs in mammalian muscle even when oxygen is present, the electron acceptor is pyruvate itself.



The alcohol and CO<sub>2</sub> produced from alcoholic fermentation form the basis of the brewing and baking industries. In baking, the dough is left to ferment and the yeast metabolises sugars to produce ethanol and CO<sub>2</sub>. The CO<sub>2</sub> causes the dough to rise.



Yeasts are used to produce almost all alcoholic beverages, e.g. wine and beers. The yeast used in the process breaks down the sugars into ethanol (alcohol) and CO<sub>2</sub>. The alcohol produced is a metabolic by-product of fermentation by the yeast.



The lactate shuttle in vertebrate skeletal muscle works alongside the aerobic system to enable maximal muscle activity. Lactate moves from its site of production to regions within and outside the muscle, e.g. liver, where it can be respired aerobically.

Andreas Braakhuis, Wintec

1. Describe the key difference between aerobic respiration and fermentation: \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_
2. Why is the efficiency of these anaerobic pathways so low? \_\_\_\_\_  
 \_\_\_\_\_
3. Why can't alcoholic fermentation go on indefinitely? \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_



# 108 Investigating Fermentation in Yeast

**Key Idea:** Brewer's yeast preferentially uses alcoholic fermentation when there is excess sugar. The  $\text{CO}_2$  released can be collected as a measure of fermentation rate.

Brewer's yeast is a facultative anaerobe (meaning it can respire aerobically or use **fermentation**). One would expect glucose to be the preferred substrate, as it is the starting molecule in **cellular respiration**, but brewer's yeast can

use a variety of sugars, including disaccharides (two unit sugars), which can be broken down into single units. The rate at which yeast (*Saccharomyces cerevisiae*) metabolises carbohydrate substrates is influenced by temperature, solution pH, and type of carbohydrate available. High levels of sugars suppress **aerobic respiration** in yeast, instead utilising fermentation in the presence of excess substrate.

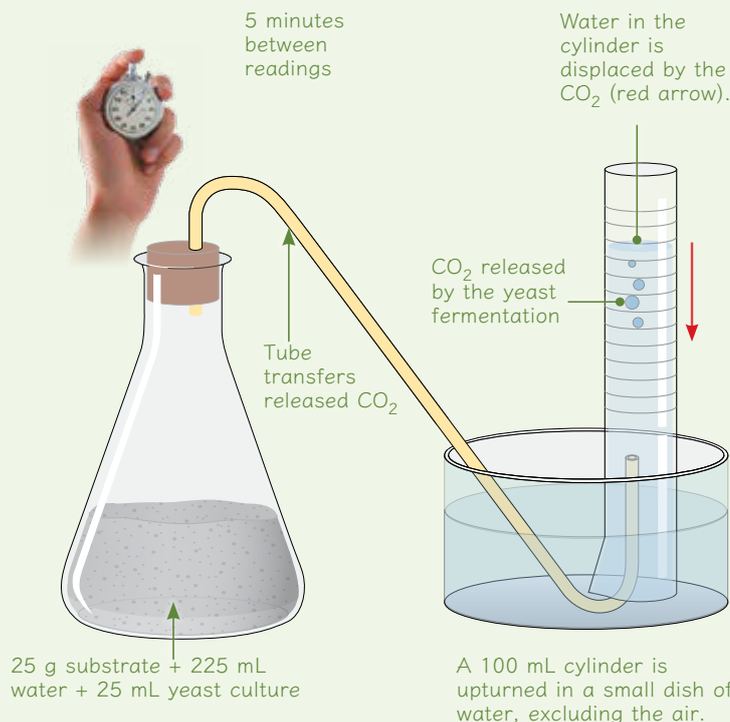


## Investigation 7.2 Investigating fermentation in yeast

See appendix for equipment list.

Work in pairs for this activity. Your teacher will assign you a substrate to investigate.

1. Make a yeast culture by dissolving 10 g of active yeast into 50 mL of water at  $24^\circ\text{C}$ .
2. In a conical flask boil 225 mL of tap water then cool to room temperature ( $24^\circ\text{C}$ ). This removes any dissolved oxygen from the water.
3. Add 25 g of substrate (glucose, maltose, sucrose, lactose, or none). Stir carefully to dissolve (stirring too vigorously will cause oxygen to dissolve back into the water).
4. Then add 25 mL of the source yeast culture to the conical flask solution.
5. Add a thin layer of paraffin oil over the solution in the conical flask to create an anaerobic environment.
6. Stopper the conical flask and set up a measuring cylinder to capture any gas as in the diagram right.
7. Start timing and record the change in gas volume every five minutes for 1 hour. Record the results for your substrate in the table. Pool data as a class and use it to complete the table below.



Substrate \ Time (min)	Cumulative volume of carbon dioxide collected (mL)				
	None	Glucose	Maltose	Sucrose	Lactose
0					
5					
10					
15					
20					
25					
30					
35					
40					
45					
50					
55					
60					

1. Write the equation for the fermentation of glucose by yeast:

\_\_\_\_\_

\_\_\_\_\_

2. Using the final values (60 minutes) collected from the class, calculate the rate of  $\text{CO}_2$  production per minute for each substrate:

(a) None: \_\_\_\_\_

(b) Glucose: \_\_\_\_\_

(c) Maltose: \_\_\_\_\_

(d) Sucrose: \_\_\_\_\_

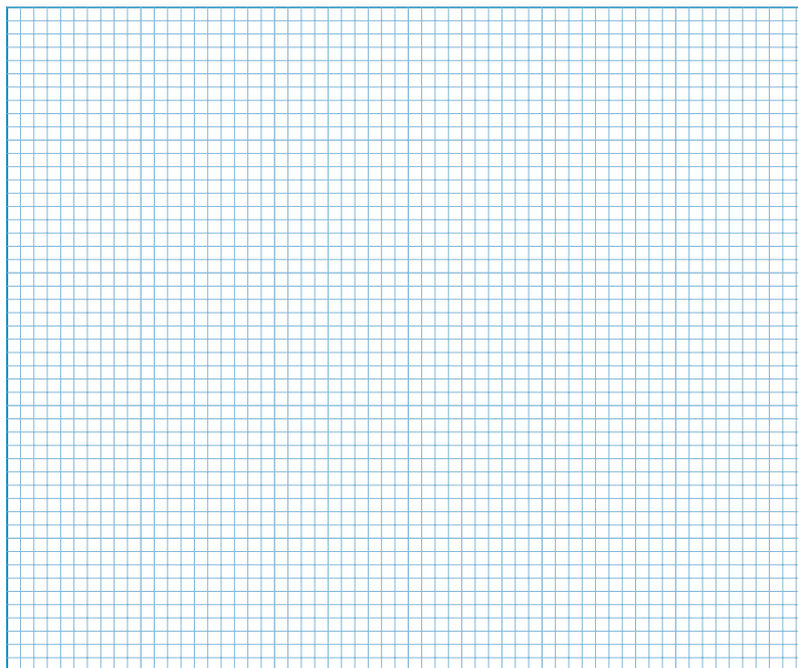
(e) Lactose: \_\_\_\_\_



SU



3. Use the tabulated data to plot an appropriate graph of the results on the grid provided:



Need help?  
See Activity 12



4. Identify the independent variable: \_\_\_\_\_

5. (a) Identify the dependent variable: \_\_\_\_\_

(b) Name the unit for the dependent variable: \_\_\_\_\_

6. (a) Summarise the results of the fermentation experiment: \_\_\_\_\_

---



---

(b) Which substrate produced the most CO<sub>2</sub>, and explain why: \_\_\_\_\_

---



---

(c) Were fermentation rates lower on maltose and sucrose than on glucose? Was this what you expected? Suggest an explanation (you may have to do some research on these molecules to find out the answer):

---



---



---

(d) Did any substrate produce no CO<sub>2</sub>? Can you suggest why? \_\_\_\_\_

---



---



---

7. Predict what would happen to CO<sub>2</sub> production rates if the yeast cells were respiring aerobically:

---



---



---

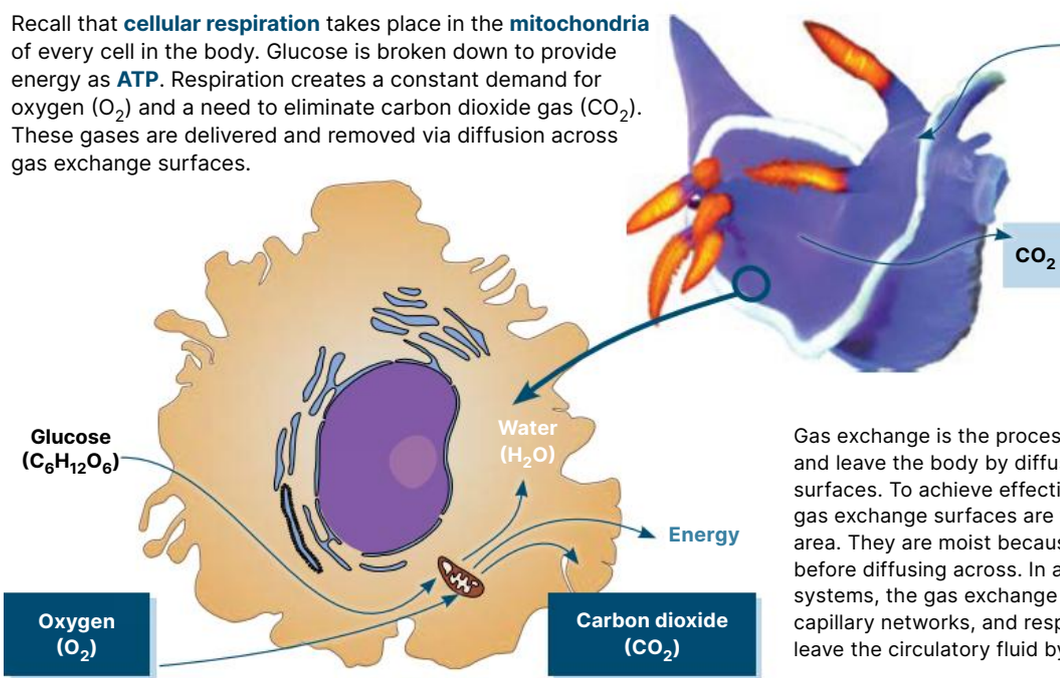
# 109 Principles of Gas Exchange

**Key Idea:** Animal gas exchange systems are suited to the animal's environment, body form, and metabolic needs. To meet the demands of aerobic metabolism, organisms must exchange gases with the environment. Some organisms can exchange gases directly across their body surface, but most

organisms have specialised **gas exchange** systems adapted to function in their specific environment. The type and complexity of the exchange system reflects the demands of **metabolism** for gas exchange (oxygen delivery and carbon dioxide removal) and the environment (aquatic or terrestrial).

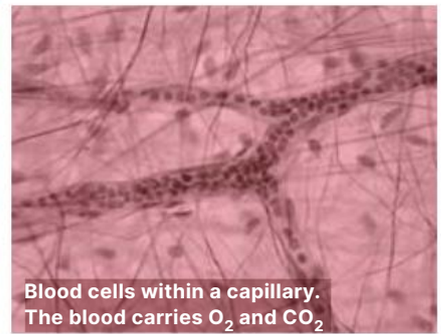
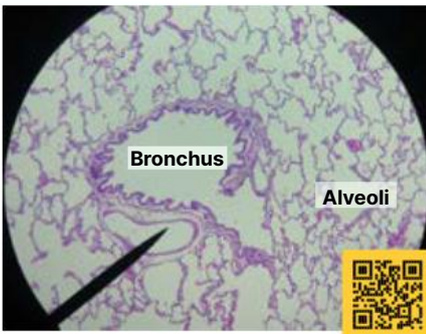
## Cellular respiration and gas exchange are linked

Recall that **cellular respiration** takes place in the **mitochondria** of every cell in the body. Glucose is broken down to provide energy as **ATP**. Respiration creates a constant demand for oxygen ( $O_2$ ) and a need to eliminate carbon dioxide gas ( $CO_2$ ). These gases are delivered and removed via diffusion across gas exchange surfaces.



Flat organisms, such as this sea slug, use the body surface as the gas exchange surface. Most multicellular organisms have specialised gas exchange systems.

Gas exchange is the process by which gases enter and leave the body by diffusion across gas exchange surfaces. To achieve effective gas exchange rates, gas exchange surfaces are thin with a high surface area. They are moist because gases must dissolve before diffusing across. In animals with gas exchange systems, the gas exchange surfaces lie close to capillary networks, and respiratory gases enter and leave the circulatory fluid by diffusion.



- ▶ In mammalian lungs, the **alveoli** (microscopic air sacs) provide a large surface area for gas exchange. The walls of the alveoli are only one cell thick (tissue section above) and are covered by capillaries (model, centre).
- ▶ **Respiratory gases** move across the gas exchange surface by diffusion. Effective gas exchange relies on maintaining a concentration gradient for gas diffusion.
- ▶ Oxygen is transported away from the gas exchange surface by the blood (above right), reducing its concentration relative to the environmental side of the gas exchange surface.  $CO_2$  is transported to the gas exchange surface, increasing its concentration relative to the environmental side of the membrane. It then diffuses out of the blood, across the membrane, and into the external environment.

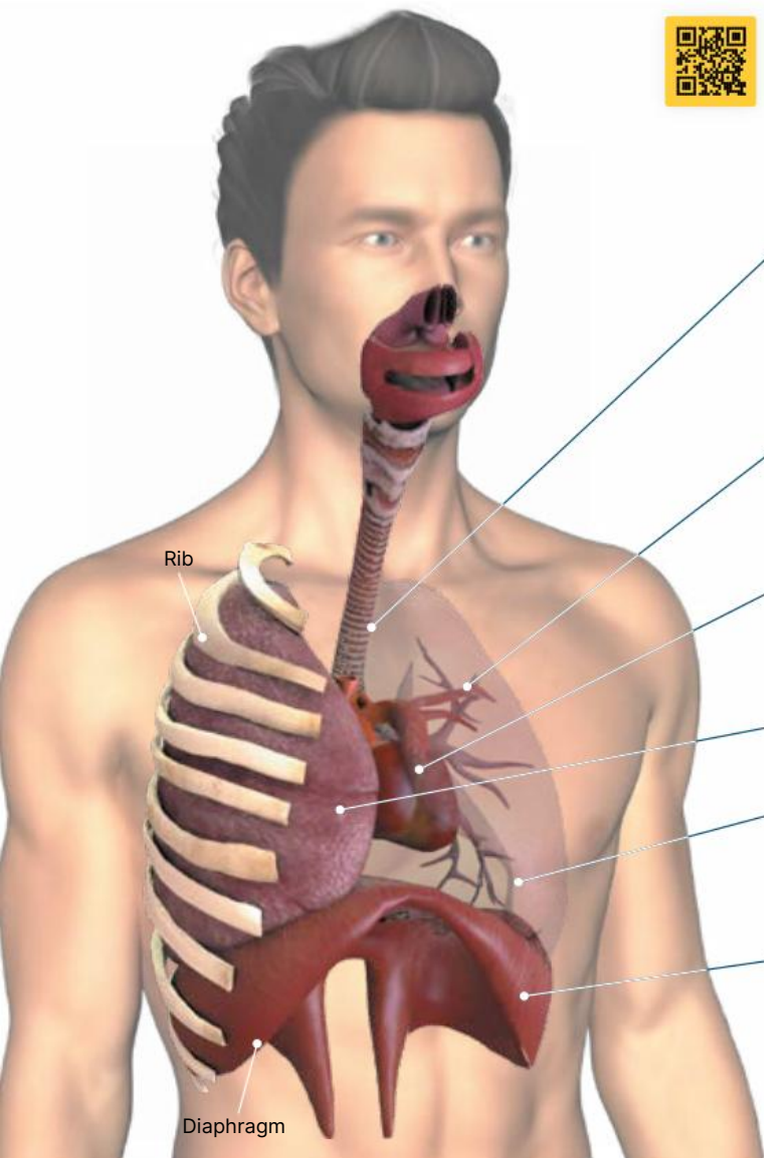
1. What is the purpose of gas exchange? \_\_\_\_\_
2. How are gases exchanged with the environment? \_\_\_\_\_
3. How are gradients for diffusion maintained in a simple organism (one without a gas exchange system)? \_\_\_\_\_
4. How are gradients for diffusion maintained in an organism with a gas exchange system? \_\_\_\_\_



**Key Idea:** The tissues and organs of the human gas exchange system work together to enable the exchange of gases between the body's cells and the environment.

The **gas exchange** system consists of the passages of the mouth and nose, the trachea, and the tubes and air sacs of

the lungs. Cooperation with the muscles of the diaphragm and ribcage contribute to its function. Each region is specialised to perform a particular role in the organ system's overall function, which is to exchange **respiratory gases** (O<sub>2</sub> and CO<sub>2</sub>) between the body's cells and the environment.



The trachea (windpipe) transfers air to the **lungs**. It is strengthened with C-shaped bands of stiff cartilage. The trachea divides into two bronchi, also supported by cartilage bands.

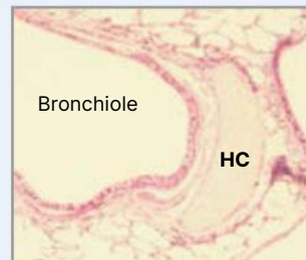
Bronchioles branch from the bronchi and divide into progressively smaller branches. Cartilage is gradually lost as the bronchioles decrease in diameter.

The cardiac notch in the left lung makes space for the heart.

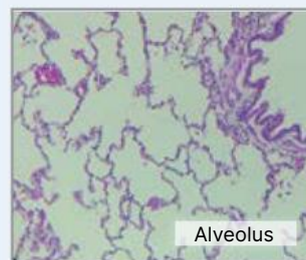
The right lung is slightly larger than the left. It takes up 55-60% of the total lung volume.

The smallest respiratory bronchioles subdivide into the alveolar ducts. The alveoli are found at the end of these.

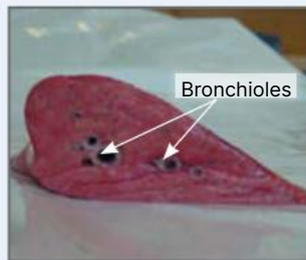
The diaphragm is a dome shaped muscle that works with the intercostal muscles of the ribcage to bring about lung ventilation (breathing). When it contracts, it moves down, reducing pressure in the lung so that air flows in.



Rings of hyaline cartilage (HC) provide support for the trachea, bronchi, and the larger bronchioles.



The lungs contain air spaces surrounded by alveolar epithelial cells (pneumocytes), forming alveoli (air sacs), where gas exchange takes place. The alveoli receive air from tubes, called bronchioles.



1. Name three types of cells in the respiratory system and their function:

- (a) \_\_\_\_\_
- \_\_\_\_\_
- (b) \_\_\_\_\_
- \_\_\_\_\_
- (c) \_\_\_\_\_
- \_\_\_\_\_

2. What is the primary organ of gas exchange? \_\_\_\_\_

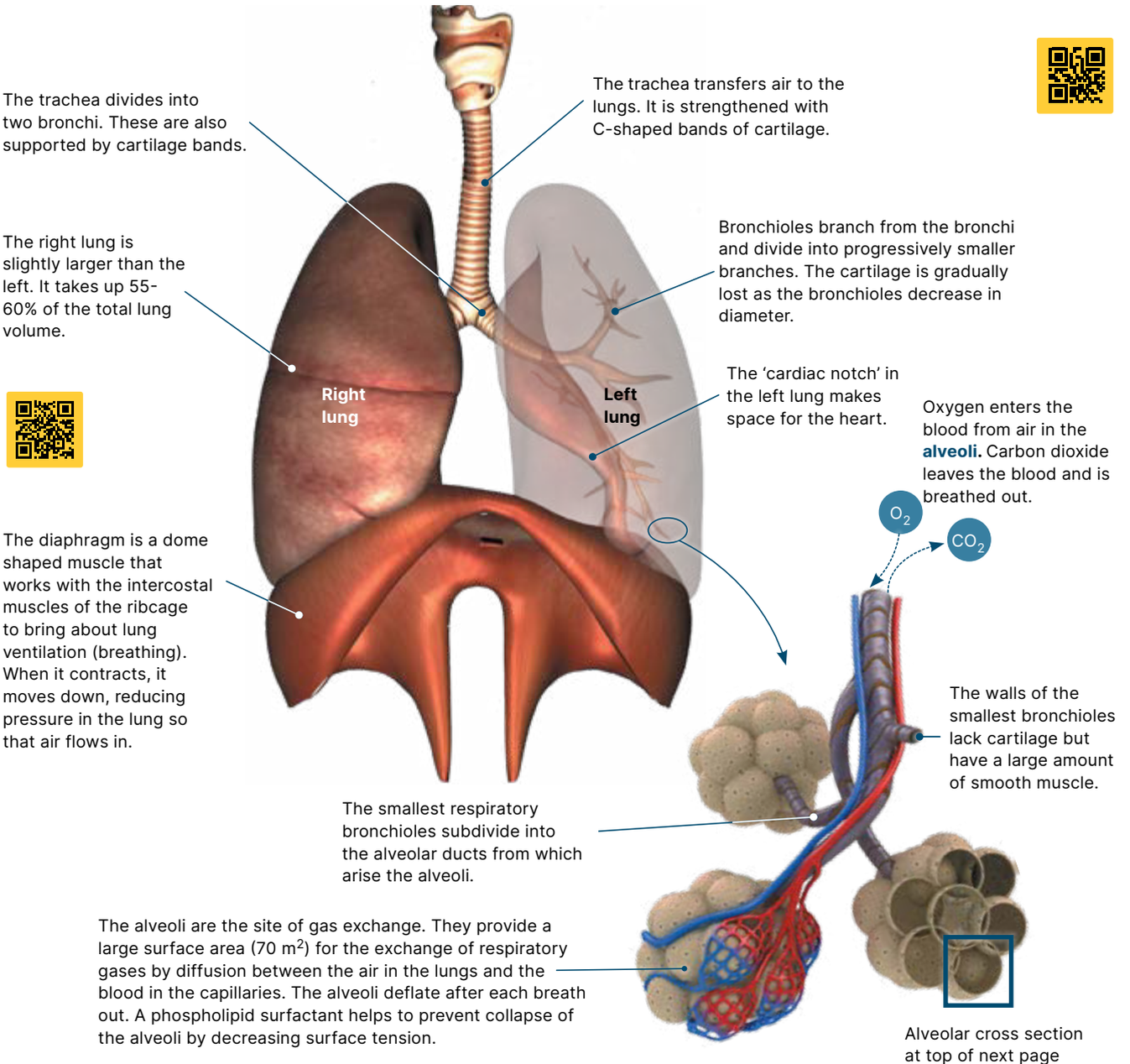
3. Which cells form the alveoli? \_\_\_\_\_

4. What is the purpose of the hyaline cartilage in the gas exchange system? \_\_\_\_\_



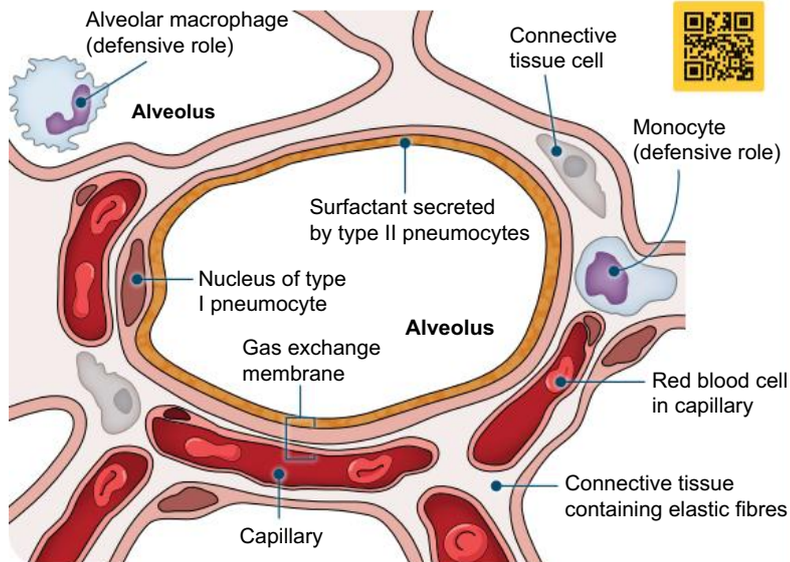
**Key Idea:** Lungs are internal sac-like organs connected to the outside by a system of airways. The smallest airways end in thin-walled alveoli, where gas exchange occurs. The **respiratory system** includes all the structures associated

with exchanging **respiratory gases** with the environment. In mammals, the **gas exchange** organs are paired lungs connected to the outside air by way of a system of tubular passageways: the trachea, bronchi, and bronchioles.



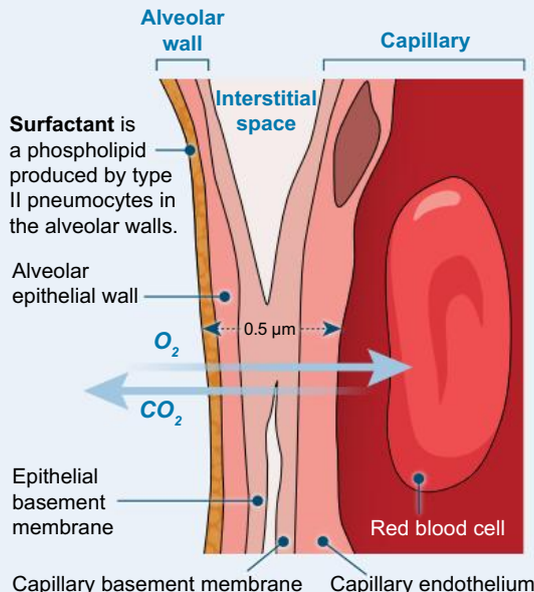
1. What is the purpose of the trachea, bronchi, and bronchioles? \_\_\_\_\_
2. What is the purpose of the diaphragm? \_\_\_\_\_
3. (a) Explain how the basic structure of the human gas exchange system provides such a large area for gas exchange: \_\_\_\_\_
- (b) In what region of the lung does the actual exchange of gases take place? \_\_\_\_\_

**Cross section through an alveolus**



The physical arrangement of the alveoli to the capillaries through which the blood moves. The alveolus is lined with a thin, single-celled layer of Type I pneumocytes (alveolar epithelial cells), across which gases are exchanged. Type II pneumocytes secrete surfactant to reduce surface tension of the alveoli to prevent them collapsing during exhalation. Phagocytes (monocytes and macrophages) are present to protect the lung tissue. Elastic connective tissue gives the alveoli their ability to expand and recoil.

**The gas exchange membrane**



The gas exchange membrane is the layered junction between the alveolar epithelial cells, the endothelial cells of the capillary, and their associated basement membranes (thin connective tissue layers under the epithelia). Gases move freely across this membrane.

4. Describe the structure and purpose of the alveolar-capillary (gas exchange) membrane:

---

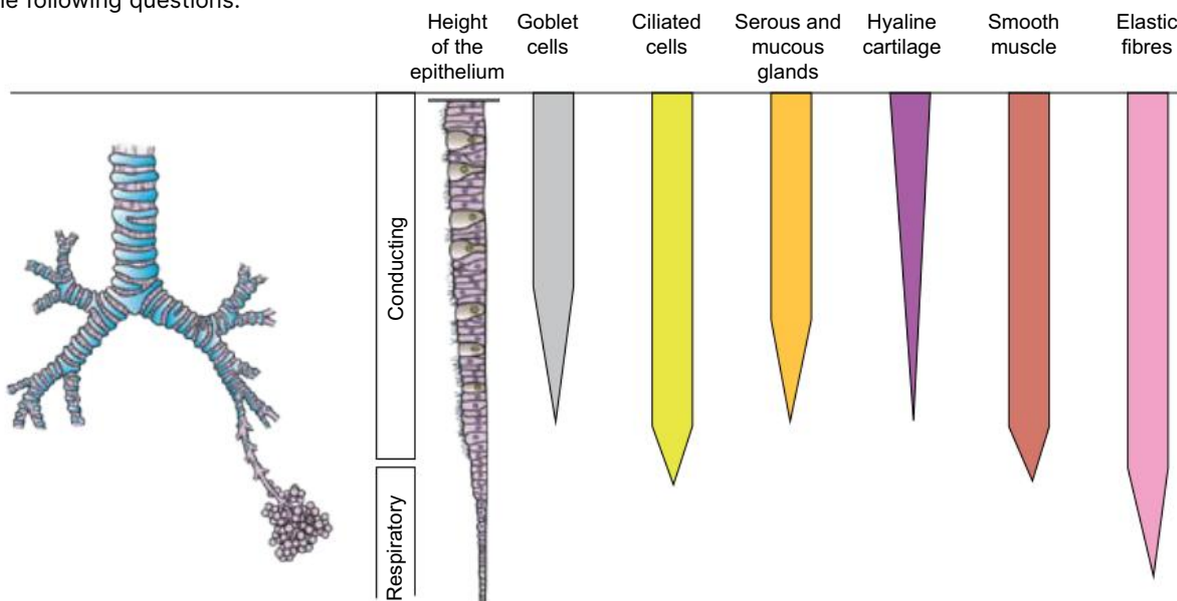


---



---

5. The diagram below shows the different types of cells and their positions and occurrence in the lungs. Use it to answer the following questions:



(a) Why does the epithelium become very thin in the respiratory zone? \_\_\_\_\_

---



---

(b) Why would elastic fibres be present in the respiratory zone, whereas hyaline cartilage is not? \_\_\_\_\_

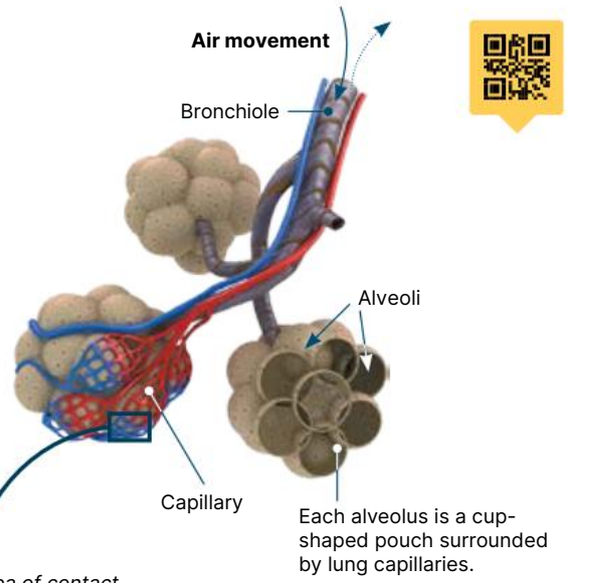
---

# 112 Gas Transport in Humans

**Key Idea:** Haemoglobin is a respiratory pigment in red blood cells which binds oxygen and increases the efficiency of its transport and delivery to tissues throughout the body.

The transport of **respiratory gases** around the body is the role of the blood and its respiratory pigment. Most of the carbon dioxide in the blood is carried as bicarbonate in the plasma. Oxygen does not dissolve in blood easily, so in vertebrates, e.g. humans, it is transported throughout the body chemically bound to the respiratory pigment **haemoglobin** (Hb) inside the red blood cells.

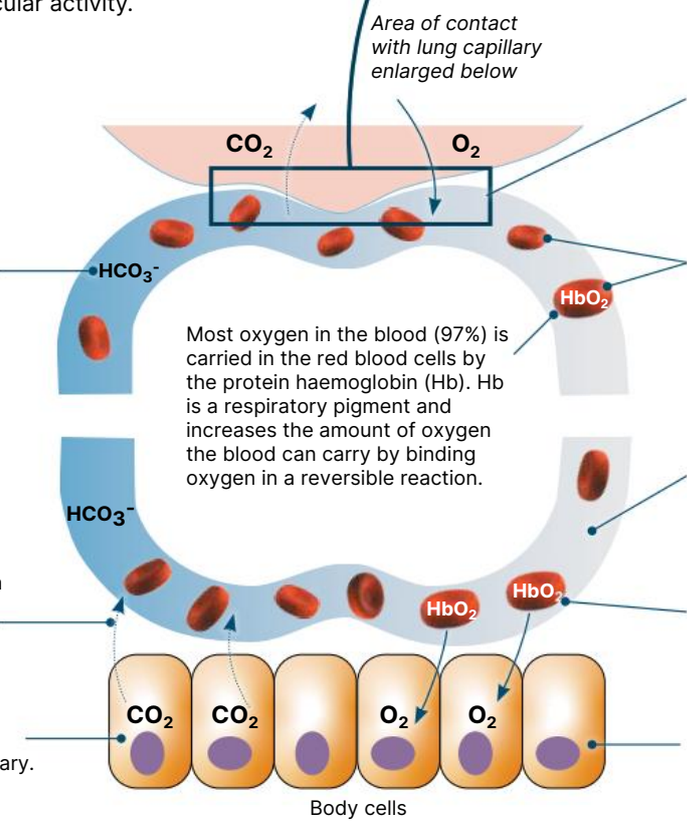
In the muscles, oxygen from haemoglobin is transferred to and retained by **myoglobin**, a molecule that is chemically similar to haemoglobin except that it consists of only one heme-globin unit. Myoglobin has a greater affinity for oxygen than haemoglobin and acts as an oxygen store within muscles, releasing the oxygen during periods of prolonged or extreme muscular activity.



Most CO<sub>2</sub> in the blood (85%) is carried as bicarbonate (HCO<sub>3</sub><sup>-</sup>) formed in the red blood cells from CO<sub>2</sub> in a reversible, enzyme-catalysed reaction. HCO<sub>3</sub><sup>-</sup> diffuses out of the red blood cells and into the plasma where it contributes to the buffer capacity of the blood.

When CO<sub>2</sub> levels rise too quickly, H<sup>+</sup> can accumulate in the blood, reducing pH. This provides a strong stimulus to increase breathing rate.

Carbon dioxide diffuses from the body's cells into the capillary.



Gas exchange membrane: Formed by the epithelial cells of the alveolus and capillary together. It is only 0.5 μm thick so gases diffuse rapidly across.

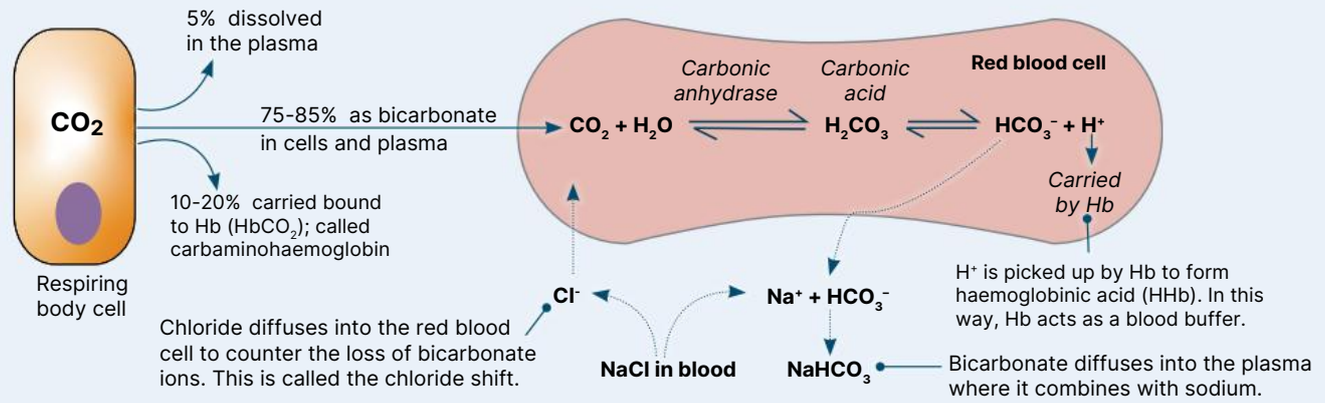
When oxygen levels are high (lungs and surrounding blood vessels) haemoglobin binds with a lot of oxygen (the Hb is saturated).

Body tissue capillary: The capillaries in the tissues are very close to the body's cells, allowing for rapid diffusion back and forth.

When carbon dioxide levels are high (body tissues) haemoglobin releases its oxygen.

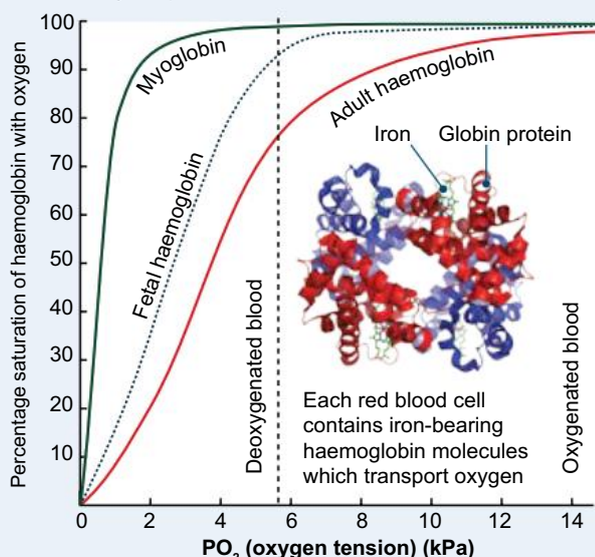
Oxygen diffuses into the body's cells from the capillary.

## Transport of carbon dioxide in the blood



## Respiratory pigments and the transport of oxygen

Fig. 1: Dissociation curves for haemoglobin and myoglobin at normal body temperature for fetal and adult human blood.



- ▶ The most important factor determining how much oxygen is carried by haemoglobin (Hb) is the level of oxygen in the blood. The greater the oxygen tension, the more oxygen will combine with Hb.
- ▶ This relationship can be illustrated in an **oxygen-haemoglobin dissociation curve** (left). In the lung capillaries (high  $O_2$ ), a lot of oxygen is picked up and bound by Hb. In the tissues (low  $O_2$ ), oxygen is released.
- ▶ Myoglobin in skeletal muscle has a very high affinity for oxygen and will take up oxygen from Hb in the blood. It can therefore act as an oxygen store.
- ▶ Fetal Hb has a high affinity for oxygen and carries 20-30% more than maternal Hb.
- ▶ The release of oxygen to the tissues is enhanced by the effect of pH. As pH increases (lower  $CO_2$ ), more oxygen combines with Hb. As the blood pH decreases (higher  $CO_2$ ), Hb binds less oxygen and releases more to the tissues. This is called the Bohr effect.



- (a) Identify two regions in the body where oxygen levels are relatively high: \_\_\_\_\_

\_\_\_\_\_

(b) Identify two regions where carbon dioxide levels are relatively high: \_\_\_\_\_

\_\_\_\_\_
- (a) What is the function of haemoglobin? \_\_\_\_\_

\_\_\_\_\_

(b) Explain the significance of the reversible binding of oxygen by haemoglobin: \_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_
- (a) How is haemoglobin saturation affected by the oxygen level in the blood? \_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_

(b) What is the significance of this relationship to oxygen delivery to the tissues? \_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_
- At low blood pH, less oxygen is bound by haemoglobin and more is released to the tissues:

(a) Name this effect: \_\_\_\_\_

(b) What is its significance? \_\_\_\_\_

\_\_\_\_\_
- (a) Compare the affinity of myoglobin and haemoglobin for oxygen: \_\_\_\_\_

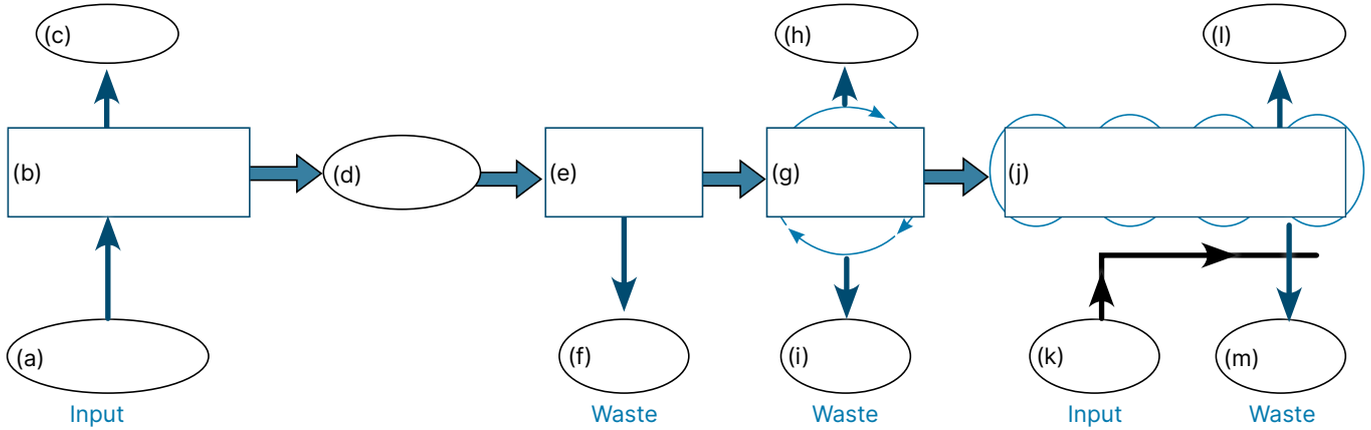
\_\_\_\_\_

(b) Why is the very high affinity of myoglobin for oxygen important? \_\_\_\_\_

\_\_\_\_\_

# 113 Did You Get It?

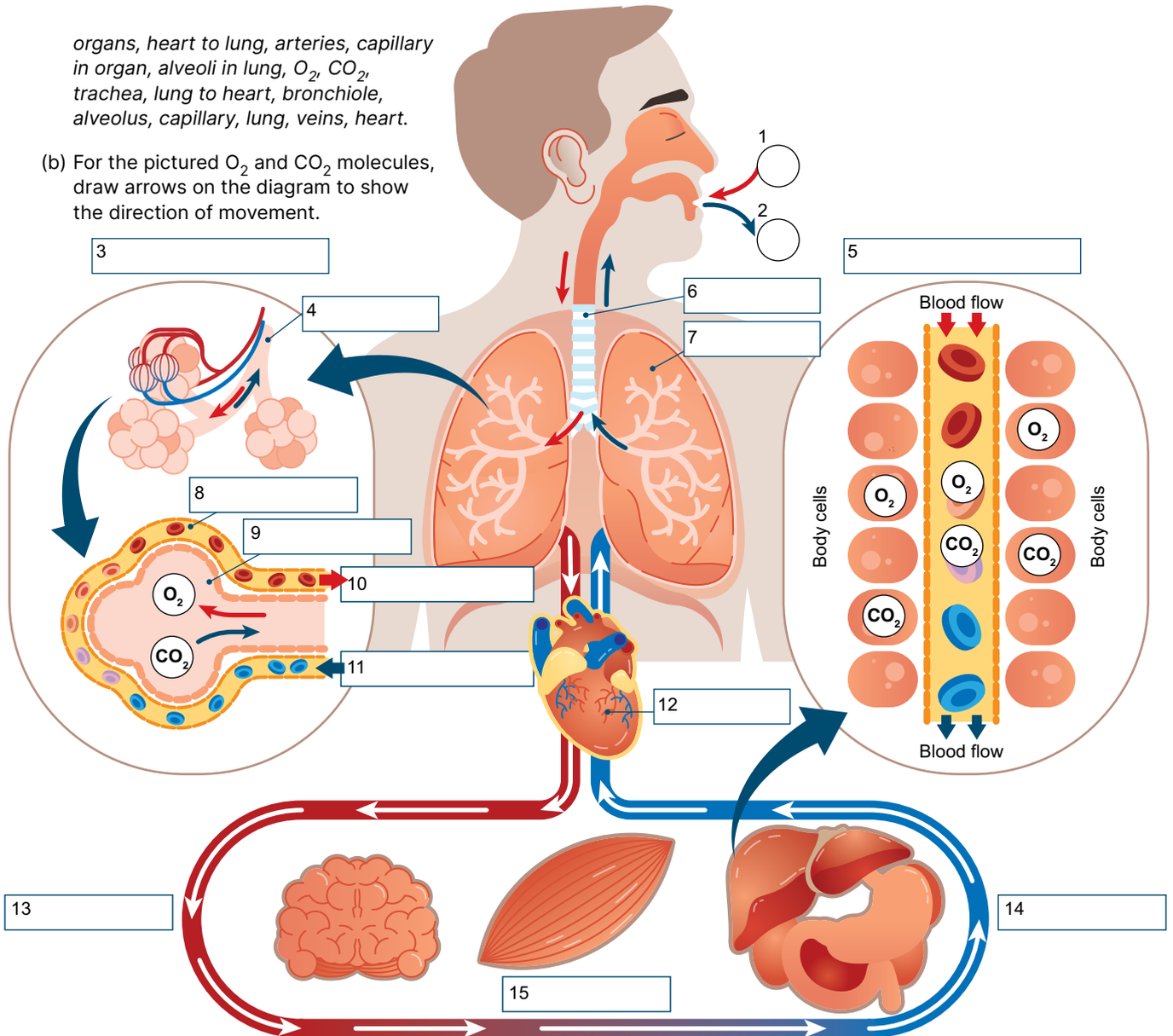
- Cellular respiration is a continuous, integrated process. A simple diagram of the process in a eukaryote is shown below.
  - In the diagram, fill in the rectangles with the process and the ovals with the substance used or produced. Use the following word list (some words can be used more than once): pyruvate, glycolysis, glucose, oxygen ( $O_2$ ), link reaction, electron transport chain (ETC), Krebs cycle, ATP, carbon dioxide ( $CO_2$ ), water ( $H_2O$ )
  - Add in a pathway to show fermentation. Write the two possible products of this pathway in eukaryotes.



- Label the diagram below (1-15) using the following word list:

*organs, heart to lung, arteries, capillary in organ, alveoli in lung,  $O_2$ ,  $CO_2$ , trachea, lung to heart, bronchiole, alveolus, capillary, lung, veins, heart.*

- For the pictured  $O_2$  and  $CO_2$  molecules, draw arrows on the diagram to show the direction of movement.





# Plant Gas Exchange and Transport Systems

## Key Terms

- artificial leaf
- autotroph
- bionic leaf
- Calvin cycle
- chlorophyll
- chloroplast
- cohesion-tension
- companion cell
- cuticle
- epidermis
- grana
- guard cells
- leaf
- light dependent phase
- light independent phase
- phloem
- photosynthesis
- sieve plate
- sieve tube
- stem
- stomata
- stroma
- thylakoids
- translocation
- transpiration
- vascular tissue
- xylem

## Key Concepts

- ▶ Plants are autotrophs and use sunlight to make their own food through the process of photosynthesis.
- ▶ Some plant cells contain specific structures for efficient gas exchange and nutrient transport.

## Photosynthesis

Activity Number

□ 1	Describe photosynthesis as an enzyme-controlled series of reactions which use light energy to synthesise glucose. Describe the ecological role and importance of plants as producers. Analyse the conditions required for photosynthesis.	114-115, 126
□ 2	Describe features of chloroplasts related to their role in photosynthesis.	116
□ 3	Summarise photosynthesis in both a word equation and a balanced chemical equation, identifying raw materials and end products. Identify and explain the key stages of photosynthesis, including the light dependent reactions and the light independent reactions (the Calvin cycle).	117
□ 4	Analyse secondary data from an investigation on the effect of light on the photosynthesis rate. Describe the role of Rubisco in the photosynthetic reaction.	118-119

## Gas exchange in plants

□ 5	Describe the role of stomata and guard cells in controlling the movement of gases (oxygen, carbon dioxide, and water vapour) in plants.	125
□ 6	Describe the structure and arrangement of leaf cells in a generalised plant and explain how it facilitates the exchange of gases between the environment and the plant tissues.	125
□ 7	Explain the relationship between photosynthesis and the main tissues of the leaves (spongy and palisade mesophyll, epidermis, cuticle, and vascular bundles).	125
□ 8	<b>SI:</b> Explore adaptations in plants to allow efficient gas exchange	120, 125

## Vascular tissue and plant transport

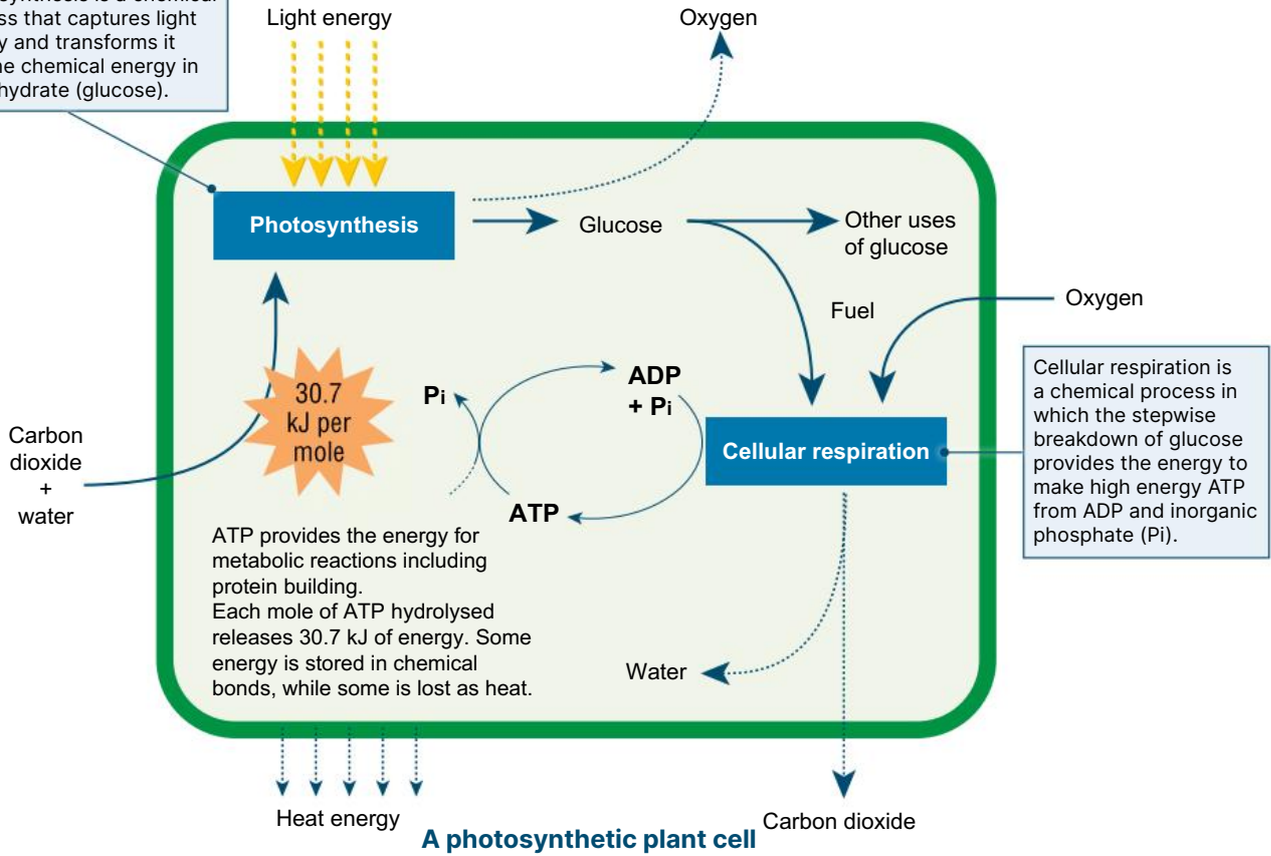
□ 9	Describe and contrast the structure of xylem and phloem tissue. Include reference to vessels and tracheids in xylem and sieve tubes, sieve plates, and companion cells in phloem; relate the structure of these elements to their function.	121-122
□ 10	Explain the movement of water and dissolved minerals through the xylem (the transpiration stream). Include reference to transpiration pull, the cohesion-tension hypothesis, root uptake, and root pressure.	123-124
□ 11	Collect and analyse data to show how light, temperature, wind, and humidity affect transpiration rate.	127
□ 12	Explain the source to sink transport of the sugars and some minerals via translocation in the phloem.	128
□ 13	<b>SHE:</b> Evaluate how scientists use their understanding of plant photosynthesis to develop new technologies.	129
□ 14	<b>SI:</b> Explore the structure and function of plant phloem as an adaptation to allow efficient transport of dissolved sugars.	122, 128
□ 15	<b>SI:</b> Investigate transpiration rates in different plants or under different conditions, collecting data with a potometer.	127
□ 16	<b>SI:</b> Explore how understanding photosynthesis can be used to design new artificial and bionic leaves that convert solar energy into liquid fuel.	129

# 114 Energy Transformations in Cells

**Key Idea:** The energy from sunlight is captured and stored as glucose, which powers the production of ATP in the process of cellular respiration. Hydrolysis of ATP provides the energy for the chemical reactions in living systems.

Energy flow in the cell of an **autotroph** (a plant) is shown below. Note that ATP has a central role in acting as an energy carrier to power metabolic reactions. Some of the energy is lost as heat during these reactions.

Photosynthesis is a chemical process that captures light energy and transforms it into the chemical energy in carbohydrate (glucose).



### DID YOU KNOW?

It takes energy to break bonds, so how does the hydrolysis of ATP provide energy for metabolic reactions?

The hydrolysis of ATP is linked to the formation of a reactive intermediate, which can be used to do work. The reactions that make the energy in ATP available occur virtually simultaneously, so the reaction is simplified to omit the intermediates:



- How does ATP act as a supplier of energy to power metabolic reactions? \_\_\_\_\_
- (a) Identify the ultimate source of energy for most autotrophs: \_\_\_\_\_  
 (b) Identify a group of autotrophic organisms that do not use this source of energy: \_\_\_\_\_
- Identify the ultimate source of energy for most heterotrophs: \_\_\_\_\_
- In what way are the processes pictured above (photosynthesis and cellular respiration) connected?  
 \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_



# The Role of Photosynthesis

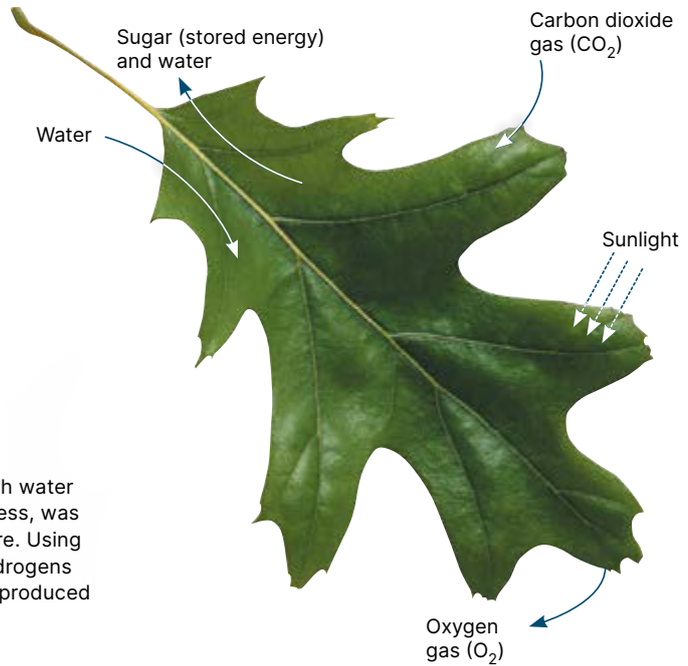
**Key Idea:** Photosynthesis is the chemical process in which autotrophs use sunlight to produce carbohydrates.

**Photosynthesis** is the process by which plants capture light energy and use it to fix (convert) the carbon in CO<sub>2</sub> into carbohydrates (e.g. glucose). The carbohydrate is used

by the plant to power ATP production and build its body. Plants (and other photosynthetic organisms) carry out this process without input from other organisms, so they are called producers (as opposed to consumers, which depend on energy and carbon from other organisms).

## Photosynthesis and producers

- ▶ A producer (or **autotroph**) is an organism that can make its own food.
- ▶ Plants, algae, and some bacteria are producers.
- ▶ Most producers use the energy in sunlight to make their food. The process by which they do this is called photosynthesis. Photosynthesis transforms sunlight energy into chemical energy.
- ▶ The chemical energy is stored as glucose, and the energy is released when the glucose undergoes further metabolic processes.
- ▶ The inputs and outputs of photosynthesis are shown on the leaf diagram (right).



### DID YOU KNOW?

The evolution of oxygenic photosynthesis, in which water is split to provide the hydrogens to drive the process, was responsible for our current oxygen rich atmosphere. Using water (rather than hydrogen sulfide) to supply hydrogens provided far more energy for ATP production and produced oxygen gas as a waste product.



Photosynthesis by marine algae provides oxygen and absorbs carbon dioxide. Most algae are microscopic but some, like this kelp, are large.



Depending on the plant, 0.1% to 8% of the light intercepted is used in photosynthesis. Typically crop plants use about 1%-2%.



Producers, such as grasses, make their own food, and are also the ultimate source of food and energy for consumers, such as this cow.

1. (a) What is a producer? \_\_\_\_\_  
 \_\_\_\_\_  
 (b) Name some organisms that are producers: \_\_\_\_\_  
 \_\_\_\_\_
2. Where do producers get their energy from? \_\_\_\_\_  
 \_\_\_\_\_
3. Why are producers so important in an ecosystem? \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_



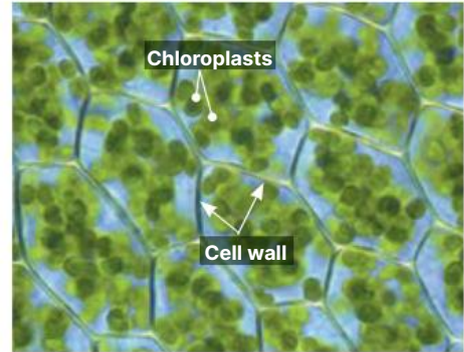
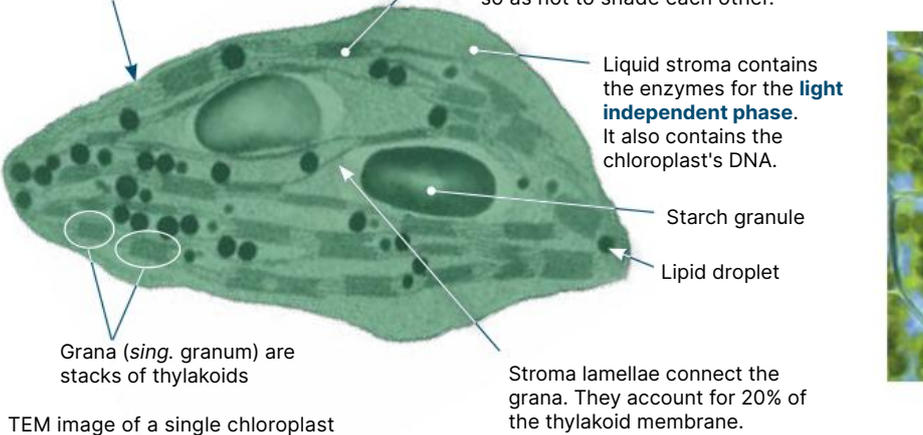
# 116 Chloroplasts

**Key Idea:** Chloroplasts have a complicated internal membrane structure. They are the site of photosynthesis in plant cells. **Chloroplasts** are the specialised plastids in which **photosynthesis** occurs. A mesophyll (photosynthetic) leaf cell contains between 50-100 chloroplasts. The chloroplasts are generally aligned so that their broad surface runs parallel to the cell wall to maximise the surface area available for

light absorption. Chloroplasts have an internal structure characterised by a system of membranous structures called **thylakoids** arranged into stacks called **grana**. Special pigments, called **chlorophylls** and carotenoids, are bound to the membranes as part of light-capturing photosystems. They absorb light of specific wavelengths and thereby capture the light energy.

## The structure of a chloroplast

Chloroplast is enclosed by a double membrane envelope (inner and outer membrane)



Chloroplasts visible in plant cells

TEM image of a single chloroplast

1. Label the transmission electron microscope image of a chloroplast below:

2. (a) Where is chlorophyll found in a chloroplast? \_\_\_\_\_

\_\_\_\_\_

(b) Why is chlorophyll found there? \_\_\_\_\_

\_\_\_\_\_

3. Explain how the internal structure of chloroplasts helps absorb the maximum amount of light:

\_\_\_\_\_

\_\_\_\_\_

4. Explain why plant leaves appear green: \_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_



**Key Idea:** Photosynthesis is the process by which light energy is used to convert CO<sub>2</sub> and water into glucose and oxygen.

**Photosynthesis** is of fundamental importance to living things because it transforms sunlight energy into chemical energy stored in molecules, releases free oxygen gas, and absorbs carbon dioxide (a waste product of cellular metabolism). Photosynthesis has two phases, the **light dependent phase** and the **light independent phase**. In the reactions

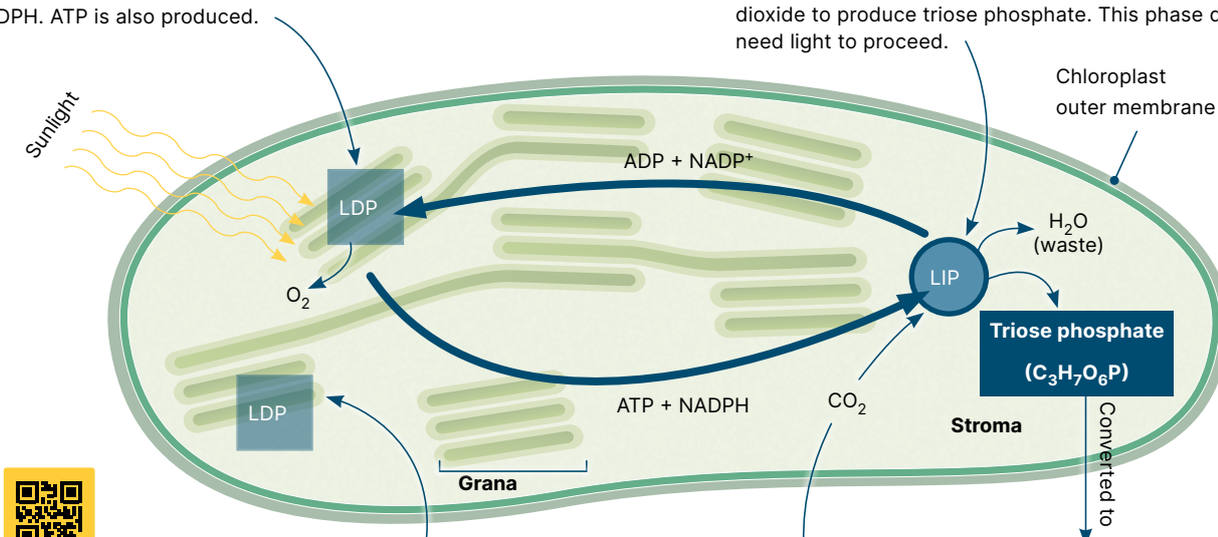
of the light dependent phase, light energy is converted to chemical energy (ATP and NADPH). This phase occurs in the **thylakoid** membranes of the **chloroplasts**. In the reactions of the light independent phase, the chemical energy is used to synthesise carbohydrate. This phase occurs in the **stroma** of chloroplasts. In photosynthesis, water is split and electrons are transferred together with hydrogen ions from water to CO<sub>2</sub>, reducing it to triose phosphates (converted to sugars).

## Light dependent phase (LDP): Thylakoid

In the first phase of photosynthesis, chlorophyll captures light energy which is used to split water, producing O<sub>2</sub> gas (waste) and H<sup>+</sup> ions that are transferred to the molecule NADPH. ATP is also produced.

## Light independent phase (LIP): Stroma

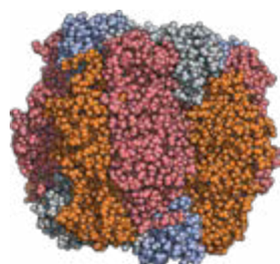
The second phase of photosynthesis occurs in the stroma and uses NADPH and ATP to drive a series of enzyme controlled reactions (the **Calvin cycle**) that fix carbon dioxide to produce triose phosphate. This phase does not need light to proceed.



The light dependent phase occurs in the thylakoid membranes of the grana.

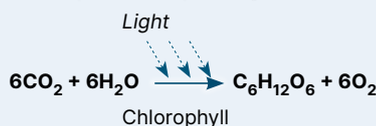
CO<sub>2</sub> from the air provides raw materials for glucose production.

Monosaccharides, e.g. glucose, and other carbohydrates, lipids, and amino acids.



Rubisco is the central enzyme in the LIP of photosynthesis (carbon fixation) catalysing the first step in the Calvin cycle. However, it is remarkably inefficient, processing just three reactions a second. To compensate, rubisco makes up almost half the protein content of chloroplasts.

### The general equation for photosynthesis



1. Explain how the light-dependent reaction and the light independent reaction are linked:

---



---



---

2. Write a word equation for photosynthesis: \_\_\_\_\_

3. What is Rubisco and what is its role? \_\_\_\_\_

---

4. State the fate of the following molecules involved in photosynthesis:

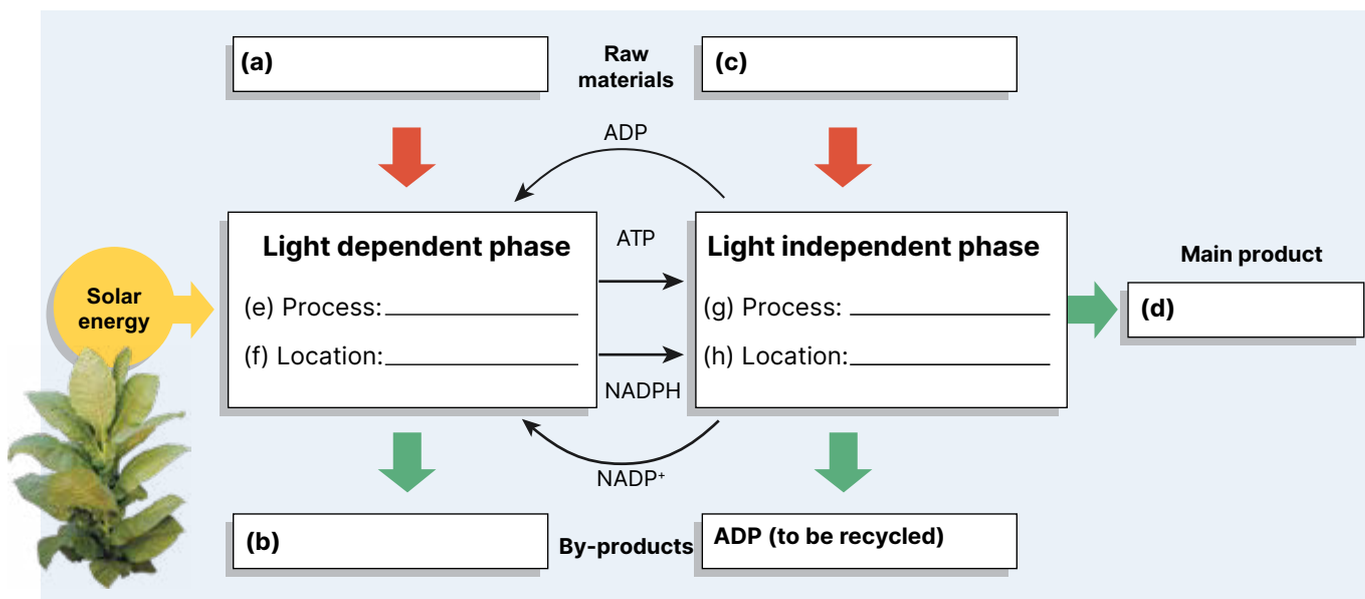
(a) Carbon dioxide: \_\_\_\_\_

(b) Oxygen: \_\_\_\_\_

(c) Hydrogen: \_\_\_\_\_



5. Use the information on the previous page to fill in the diagram below. Fill in the raw material (inputs), products (outputs), and state what is happening at each phase and where the phase takes place (occurs).



6. In two experiments, radioactively-labelled oxygen (shown in blue) was used to follow oxygen through the photosynthetic process. The results of the experiment are shown below:



From these results, what would you conclude about the source of the oxygen in:

(a) The carbohydrate produced? \_\_\_\_\_

(b) The oxygen released? \_\_\_\_\_

7. Name the products that triose phosphate is converted into: \_\_\_\_\_

8. Describe what happens during:

(a) The light dependent phase of photosynthesis: \_\_\_\_\_

\_\_\_\_\_

(b) The light independent phase of photosynthesis: \_\_\_\_\_

\_\_\_\_\_

9. What is the function of each of the following in photosynthesis:

(a) ATP: \_\_\_\_\_

(b) NADPH: \_\_\_\_\_

(c) Light: \_\_\_\_\_

(d) Chlorophyll: \_\_\_\_\_

(e) Carbon dioxide: \_\_\_\_\_

(f) Water: \_\_\_\_\_

# Investigating Photosynthetic Rate

**Key Idea:** Measuring the production of oxygen provides a simple means of measuring the rate of photosynthesis.

The rate of **photosynthesis** can be investigated by measuring the substances involved in photosynthesis. These include

measuring the uptake of carbon dioxide, the production of oxygen, or the change in biomass over time. Measuring the rate of oxygen production provides a good approximation of the photosynthetic rate and is relatively easy to carry out.

**The aim**

To investigate the effect of light intensity on the rate of photosynthesis in an aquatic plant, *Cabomba aquatica*.

**The method**

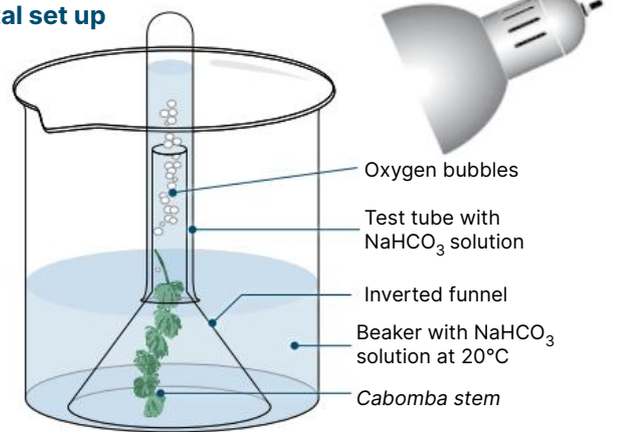
0.8-1.0 grams of *Cabomba* stem were weighed on a balance. The stem was cut and inverted to ensure a free flow of oxygen bubbles.

The stem was placed into a beaker filled with a solution containing 0.2 mol L<sup>-1</sup> sodium hydrogen carbonate (to supply carbon dioxide). The solution was at approximately 20°C. A funnel was inverted over the *Cabomba* and a test tube filled with the sodium hydrogen carbonate solution was inverted on top to collect any gas produced.

The beaker was placed at distances (20, 25, 30, 35, 40, 45, 50 cm) from a 60W light source and the light intensity measured with a lux meter at each interval.

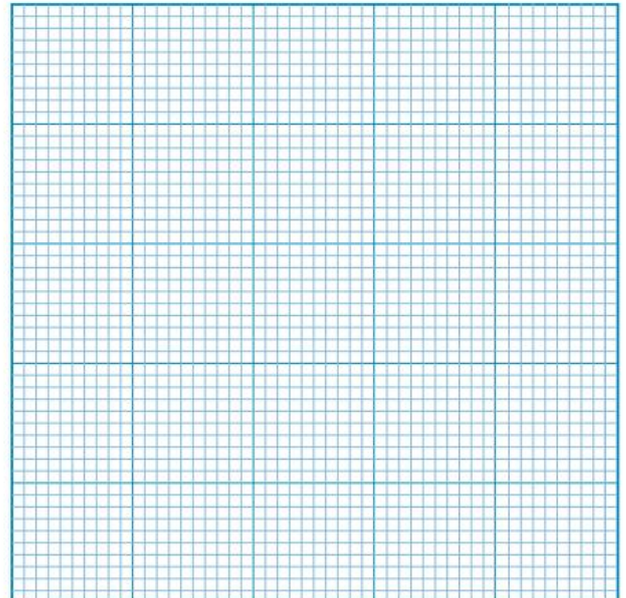
Before recording data, the *Cabomba* stem was left to acclimatise to the new light level for 5 minutes. Because the volumes of oxygen gas produced are very low, bubbles were counted for a period of three minutes at each distance.

**Experimental set up**



**The results**

Light intensity (lx) (distance)	Bubbles counted in three minutes	Bubbles per minute
5 (50 cm)	0	
13 (45 cm)	6	
30 (40 cm)	9	
60 (35 cm)	12	
95 (30 cm)	18	
150 (25 cm)	33	
190 (20 cm)	35	



1. Complete the table above by calculating the rate of oxygen production (bubbles of oxygen gas per minute):
2. Use the data to draw a graph of the bubble produced per minute vs light intensity:
3. Although the light source was placed set distances from the *Cabomba* stem, light intensity in lux was recorded at each distance rather than distance per se. Explain why this would be more accurate:

---



---



---

4. The sample of gas collected during the experiment was tested with a glowing splint. The splint reignited when placed in the gas. What does this confirm about the gas produced?

---

5. What could be a more accurate way of measuring the gas produced in the experiment?

---



---



---



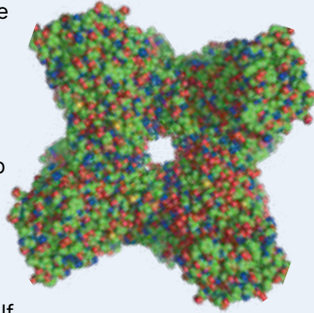
# 119 Photosynthesis and Productivity

**Key Idea:** Increasing productivity by improving photosynthetic efficiency could help to increase food and fuel production. The glucose produced in **photosynthesis** can be converted into biomass, which can be used for food or fuel (e.g. biodiesel). However, photosynthesis is an inefficient process. Productivity (i.e the rate of biomass production) can be

improved by manipulating biotic factors such as light or CO<sub>2</sub> levels and growing plants in a greenhouse. More recently scientists have begun to look at other ways to increase photosynthesis to meet the increasing demand for food and fuel required by our growing population. Many solutions focus around improving Rubisco performance.

## Rubisco

- ▶ Rubisco enzyme (right) catalyses the first major step in carbon fixation.
- ▶ Rubisco activity is very inefficient, processing just three reactions per second.
- ▶ This inefficiency makes the first step the rate limiting step of the entire photosynthetic pathway.
- ▶ To compensate for its inefficiency, Rubisco is present in high levels within a plant, it makes up almost half the protein content of chloroplasts.



## Suggestions for improving productivity

- ▶ Improving the catalytic activity of Rubisco.
- ▶ Increasing the amount of Rubisco in the leaf.
- ▶ Enhancing the CO<sub>2</sub> concentration around Rubisco.
- ▶ Improving Rubisco's affinity for CO<sub>2</sub>.
- ▶ Enhancing chloroplast electron transport rate.
- ▶ Increasing the thermostability of Rubisco Activase, an enzyme involved in activating Rubisco.
- ▶ Gene insertion from more efficient plants.

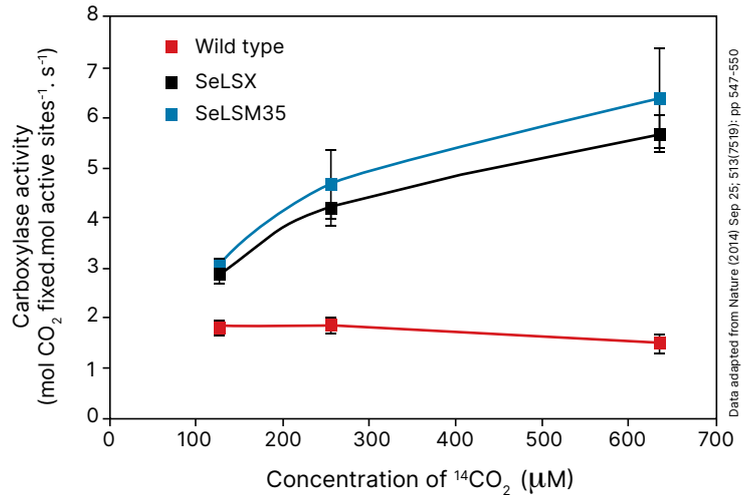
## Improving Rubisco activity

Rubisco catalyses two reactions; one is carbon fixation and the other is a reaction with oxygen. This dual processing capability reduces the amount of time it can spend fixing carbon, and so reduces photosynthesis productivity. Researchers have investigated a number of ways to improve productivity, including the genetic modification of Rubisco.

Cyanobacteria have a CO<sub>2</sub> concentrating mechanism (CCM), which allows them to produce a CO<sub>2</sub> rich environment around Rubisco. This improves photosynthetic productivity and there is less oxygen reacting with Rubisco.

Two genes (SeLSX and SeLSM35) from the cyanobacterium *Synechococcus elongatus* PCC7942 were transplanted into the DNA of the chloroplasts of the tobacco plant. The graph on the right shows the carbon fixation results of the two modified tobacco plants compared against the wild type.

Mean carboxylase activity (± standard deviation) as a measure of carbon dioxide fixation in tobacco plants.



1. Why has it become important for researchers to look at ways of boosting photosynthesis productivity?

---



---

2. (a) How do cyanobacteria improve their photosynthetic activity?

(b) Describe the effect of the transplanted cyanobacteria genes on Rubisco activity in tobacco plant:

---



---



---

3. In small groups choose a different mechanism for improving photosynthetic efficiency (and therefore productivity) from the list at the top of the page. Research the mechanism and success to date and report your findings to the class.



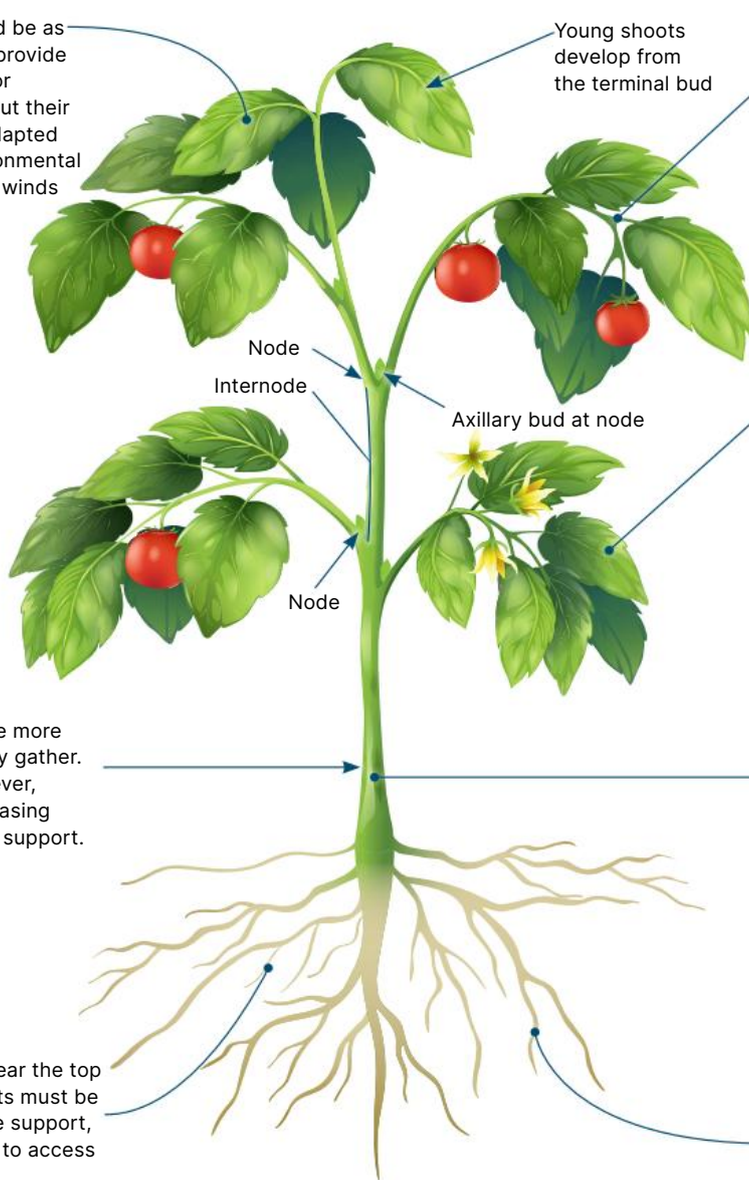
# 120 The Plant Body

**Key Idea:** The plant body consists of connected shoot and root systems. The shoot system collects carbon dioxide, oxygen, and light and produces sugars. The root system collects water and nutrients from the soil.

As terrestrial organisms, plants have two interdependent systems to take advantage of and to solve the problems of living on land. The shoot system, consisting of stems, leaves and reproductive structures, has evolved to collect carbon dioxide, oxygen and light, and to disperse pollen and seeds. The root system has evolved to collect water and nutrients

from the soil and to provide anchorage to the ground or substrate. These systems are integrated to form the closely linked support and transport systems. If a plant is to grow to any size, it must have ways to hold itself up against gravity and to move materials around its body. **Vascular tissues (xylem and phloem)** link all plant parts. Water and minerals are transported in the xylem, while manufactured food is transported in the phloem. All plants rely on fluid pressure within their cells (turgor) to give some support to their less rigid structures e.g. leaves and flowers.

Ideally, leaves should be as large as possible to provide the maximum area for gathering sunlight. But their structure must be adapted to the specific environmental conditions, e.g. high winds or low rainfall.



Functions of the stems:

Functions of the leaves:

Materials transported around the plant:

Specific functions of xylem:

Specific functions of phloem:

Functions of the roots:

The taller a plant, the more light it can potentially gather. Tall structures, however, require an ever increasing amount of structural support.

Most nutrients are near the top of the soil layer. Roots must be structured to provide support, while still being able to access this nutrient layer.

- In the boxes provided in the diagram above:
  - Describe the main functions of the leaves, roots and stems (remember that the leaves themselves have leaf veins).
  - List the materials that are transported around the plant body.
  - Describe the functions of the transport tissues: xylem and phloem.
- Name the solvent for all the materials that are transported around the plant: \_\_\_\_\_
- What factors are involved in determining how tall a plant could potentially grow? \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_



# 121 Xylem

**Key Idea:** The xylem is involved in water and mineral transport in vascular plants.

**Xylem** is the principal water conducting tissue in vascular plants. It is also involved in conducting dissolved minerals and in supporting the plant body. As in animals, tissues in plants are groupings of different cell types that work together for a common function. In flowering plants, xylem

tissue is composed of five cell types: tracheids, vessels, xylem parenchyma, sclereids (short sclerenchyma cells), and fibres. The tracheids and vessel elements form the bulk of the tissue. They are heavily strengthened and are the conducting cells of the xylem. Parenchyma cells are involved in storage, while fibres and sclereids provide support. When mature, xylem is dead.

- (a) What cells conduct the water in xylem?

\_\_\_\_\_

\_\_\_\_\_
- (b) What other cells are present in xylem tissue and what are their roles?

\_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_
- (a) How does water pass between vessels?

\_\_\_\_\_

\_\_\_\_\_
- (b) How does water pass between tracheids:

\_\_\_\_\_

\_\_\_\_\_
- (c) Which cell type do you think provides the most rapid transport of water and why?

\_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_
- (d) Why do you think the tracheids and vessel elements have/need secondary thickening?

\_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_
- How can xylem vessels and tracheids be dead when mature and functional?

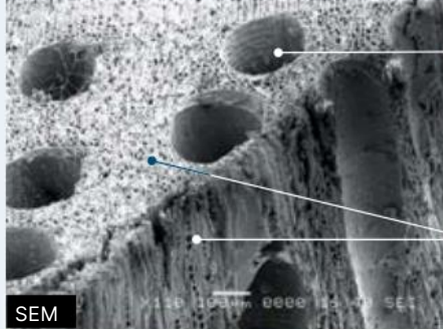
\_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_

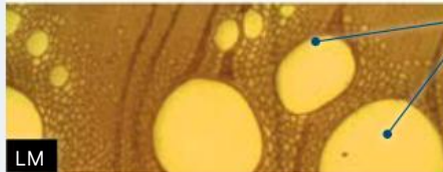
\_\_\_\_\_

\_\_\_\_\_



Water moves through the continuous tubes made by the vessel elements of the xylem.

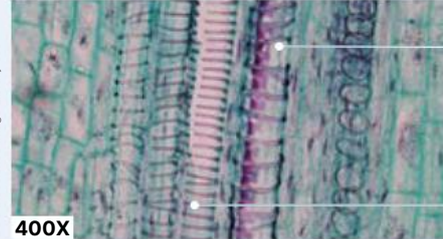
Smaller tracheids are connected by pits in the walls but do not have end wall perforations.



Vessels

Xylem is dead when mature. Note how the cells have lost their cytoplasm.

As shown in these SEM and light micrographs of xylem, the tracheids and vessel elements form the bulk of the xylem tissue. They are heavily strengthened and are involved in moving water through the plant. The transporting elements are supported by parenchyma (packing and storage cells) and sclerenchyma cells (fibres and sclereids), which provide mechanical support to the xylem.



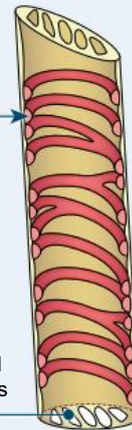
The xylem cells form continuous tubes through which water is conducted.

Spiral thickening of lignin around the walls of the vessel elements give extra strength and rigidity.

**Vessel element**

**Diameter ~ 500 µm**  
Secondary walls of cellulose are laid down after the cell has elongated or enlarged and lignin is deposited to add strength. This thickening is a feature of tracheids and vessels.

Vessels connect end to end. The end walls of the vessels are perforated to allow rapid water transport.



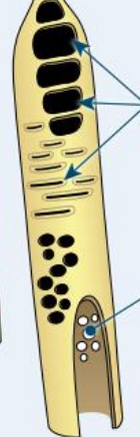
**Tip of tracheid**

**Diameter ~80 µm**

Pits and bordered pits allow transfer of water between cells but there are no end wall perforations.

No cytoplasm or nucleus in mature cell.

Tracheids are longer and thinner than vessels.



Vessel elements and tracheids are the two water conducting cell types in the xylem of flowering plants. Tracheids are long, tapering hollow cells. Water passes from one tracheid to another through thin regions in the wall called pits. Vessel elements are much larger cells with secondary thickening in different patterns (e.g. spirals). Vessel end walls are perforated to allow efficient conduction of water.



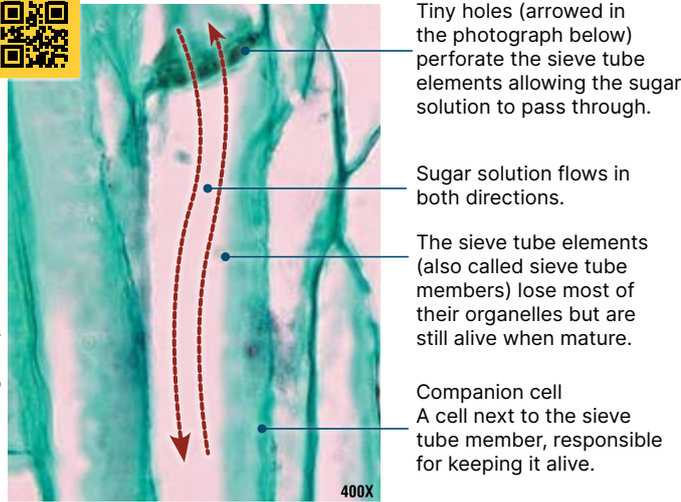
# 122 Phloem

**Key Idea:** Phloem is the principal food (sugar) conducting tissue in vascular plants, transporting dissolved sugars around the plant.

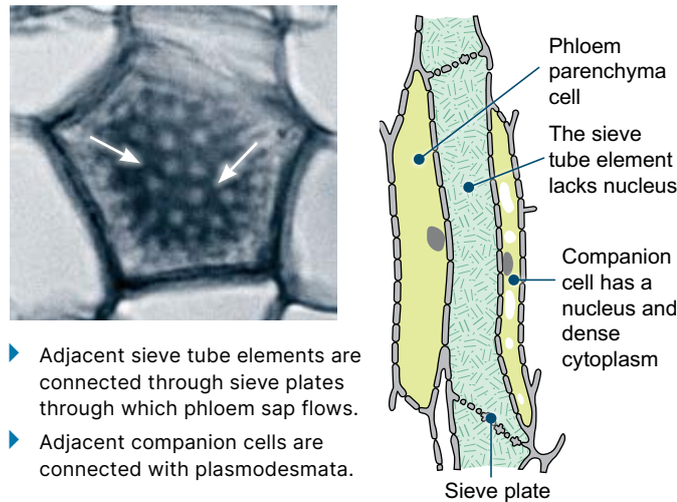
Like **xylem**, **phloem** is also a complex tissue, made up of a variable number of cell types. The bulk of phloem tissue is made up of the **sieve tubes** (sieve tube elements and sieve

cells) and their companion cells. The **sieve tubes** are the main conducting cells in phloem and are closely associated with the companion cells which support them. Parenchyma cells, concerned with storage, occur in phloem, and strengthening fibres and sclereids (short sclerenchyma cells) may also be present. Unlike xylem, phloem is alive when mature.

## Longitudinal section through a sieve tube end plate



## Transverse section through a sieve tube end plate

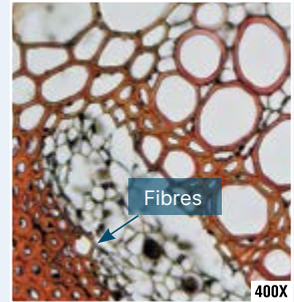


- ▶ Adjacent sieve tube elements are connected through sieve plates through which phloem sap flows.
- ▶ Adjacent companion cells are connected with plasmodesmata.

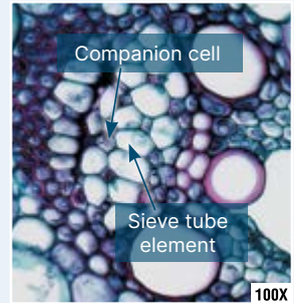
## The structure of phloem tissue

Phloem is alive at maturity and functions in the transport of sugars and minerals around the plant. Like xylem, it forms part of the structural vascular tissue of plants.

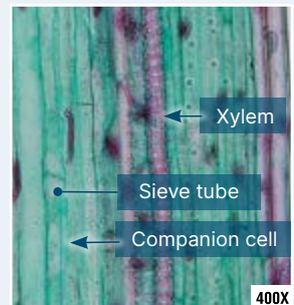
Fibres are associated with phloem as they are in xylem. Here, they are seen in cross section where you can see the extremely thick cell walls and the way the fibres are clustered in groups.



In this cross section through the vascular bundle of a corn stem, the smaller companion cells can be seen lying alongside the sieve tube members. It is the sieve tube elements that, end on end, produce the sieve tubes. They are the conducting tissue of phloem. They have reduced cytoplasm and organelles.



In this longitudinal section of a corn stem, each sieve tube element has a thin companion cell associated with it. Companion cells retain their nucleus and have many mitochondria. The cells control the metabolism of the sieve tube member next to them, and also have a role in the loading and unloading of sugar into the phloem.



- (a) What is the conducting cell type in phloem? \_\_\_\_\_

(b) What other cell type is associated with these conducting cells? \_\_\_\_\_

(c) Describe two roles of these associated cells: \_\_\_\_\_
- Mature phloem is a live tissue, whereas xylem (the water transporting tissue) is dead when mature. Why is it necessary for phloem to be alive to be functional, whereas xylem can function as a dead tissue?

\_\_\_\_\_

\_\_\_\_\_
- What is the role of fibres and sclereids in phloem? \_\_\_\_\_

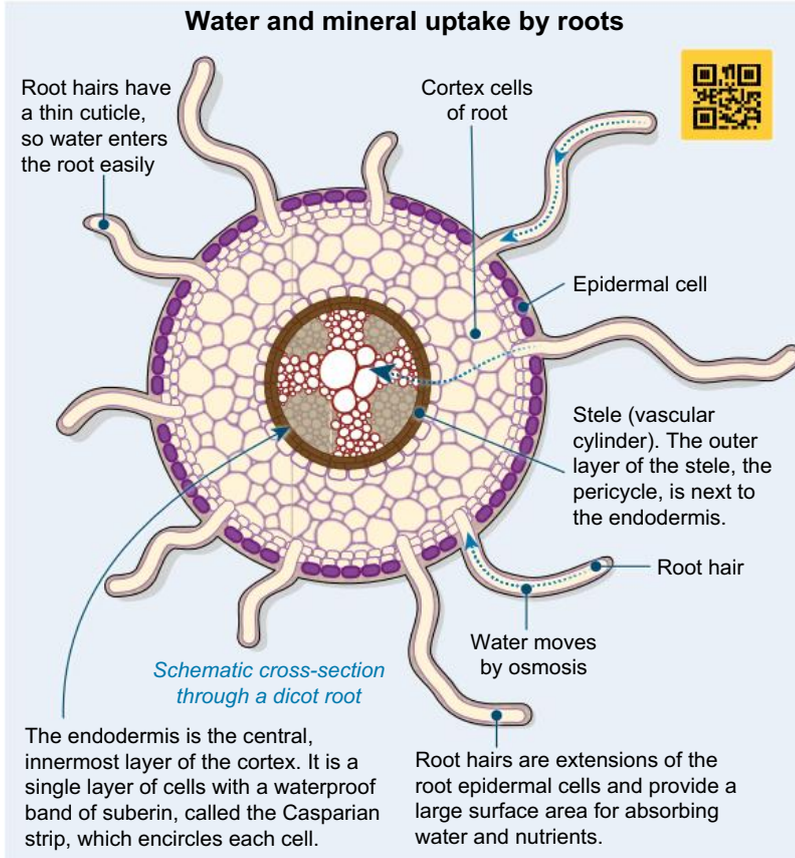
\_\_\_\_\_
- What are the large open cells next to the phloem in the centre photo above right? \_\_\_\_\_



# 123 Uptake at the Root

**Key Idea:** Water uptake by the root is a passive process. Mineral uptake can be passive or active. Plants need to take up water and minerals constantly. They must compensate for the continuous loss of water from the leaves and provide the materials the plant needs to make food. The uptake of water and minerals is mostly restricted

to the younger, most recently formed cells of the roots and the root hairs. Water uptake occurs by osmosis, whereas mineral ions enter the root by diffusion and active transport. Pathways for water movements through the plant are outlined below.



1. (a) What two mechanisms do plants use to absorb nutrients?

---



---



---

(b) Describe the two main pathways by which water moves through a plant:

---



---



---

2. Plants take up water constantly to compensate for losses due to transpiration. Describe a benefit of a large water uptake:

---



---



---



---

3. (a) How does the Casparian strip affect the route water takes into the stele?

---



---



---

(b) Why might this feature be an advantage in terms of selective mineral uptake?

---



---



---



---



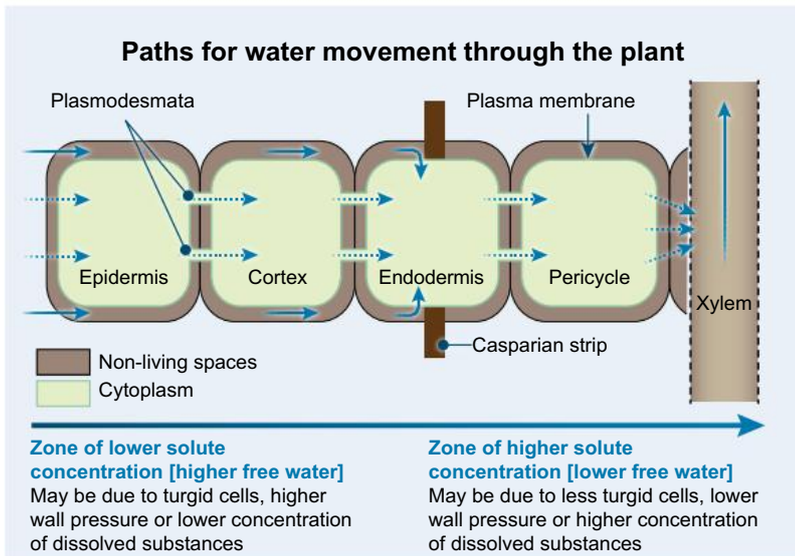
---



---



---



- ▶ The uptake of water and minerals is mostly restricted to the younger, most recently formed cells of the roots and the root hairs.
- ▶ Water moves into the roots because the solute concentration is higher in the root tissue than in the soil. When transpiration rates are too low to 'pull in' water, due to lost leaves in winter or high environmental humidity, active transport through specific transport proteins in the root hair cell membranes can increase the root pressure.
- ▶ Some water moves through the plant tissues via cytoplasmic connections between cells (the plasmodesmata), but most passes through the free spaces outside the plasma membranes of the cells.



# 124 Transpiration

**Key Idea:** Water moves through the xylem primarily as a result of evaporation from the leaves and the cohesive and adhesive properties of water molecules.

Plants lose water all the time. Approximately 99% of the water a plant absorbs from the soil is lost by evaporation from the leaves and stem. This loss, mostly through stomata, is called **transpiration** and the flow of water through the plant is called the transpiration stream. Plants rely on a gradient in solute

concentration that increases from the roots to the air to move water through their cells. Water flows passively from soil to air along this gradient of increasing solute concentration. The gradient is the driving force for the movement of water up a plant. Transpiration benefits the plant because evaporative water loss cools the plant and the transpiration stream helps the plant to take up minerals. Factors contributing to water movement are described below.

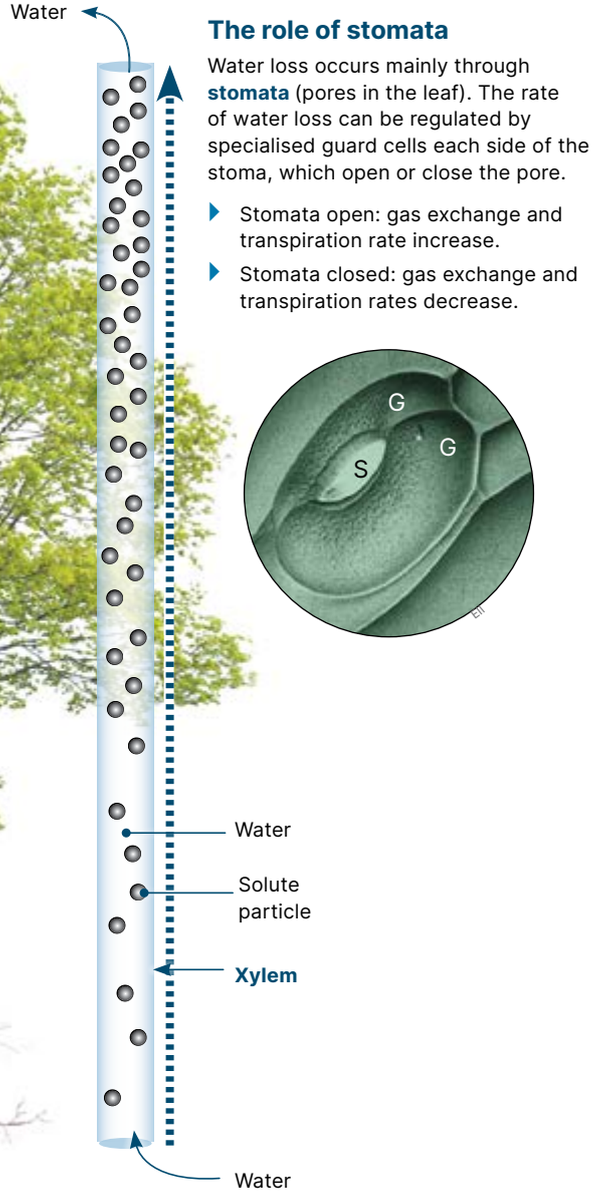
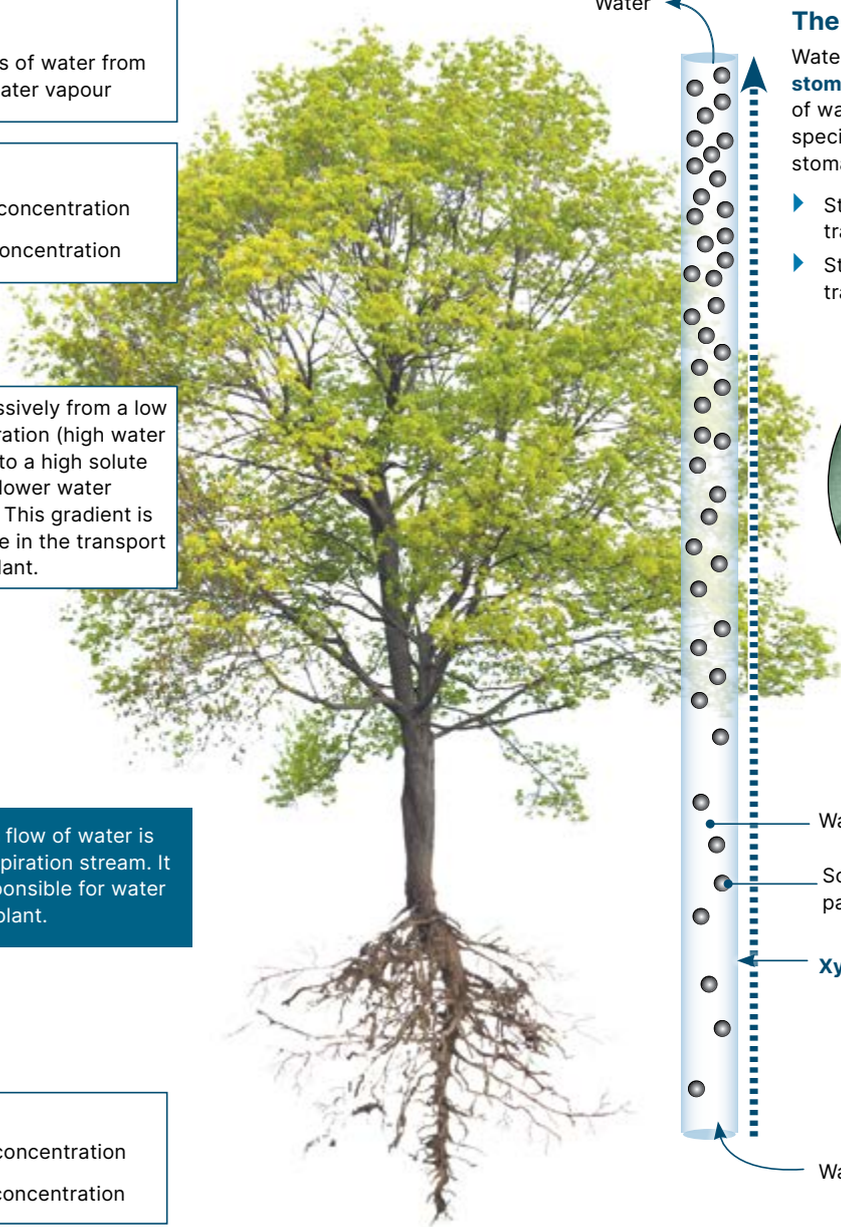
**Air**  
Evaporative loss of water from the leaves as water vapour

**Leaves**  
Highest solute concentration  
Lowest water concentration

Water flows passively from a low solute concentration (high water concentration) to a high solute concentration (lower water concentration). This gradient is the driving force in the transport of water up a plant.

The continuous flow of water is called the transpiration stream. It is primarily responsible for water moving up the plant.

**Soil**  
Highest water concentration  
Lowest solute concentration



1. (a) What is transpiration? \_\_\_\_\_

\_\_\_\_\_

(b) Describe one benefit of the transpiration stream for a plant: \_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_

2. How does the plant regulate the amount of water lost from the leaves? \_\_\_\_\_

\_\_\_\_\_



## Processes involved in moving water through the xylem

### 1 Transpiration pull

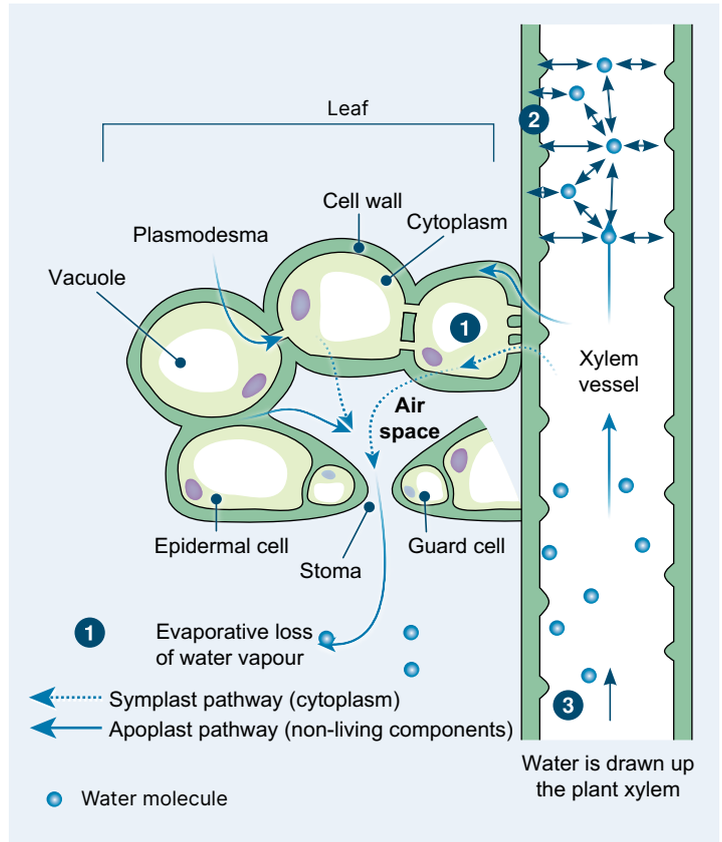
Water is lost from the air spaces by evaporation through stomata and is replaced by water from the mesophyll cells. The constant loss of water to the air (and production of sugars) creates a solute concentration in the leaves that is higher than elsewhere in the plant. Water is pulled through the plant along a gradient of increasing solute concentration.

### 2 Cohesion-tension

The transpiration pull is assisted by the special cohesive properties of water. Water molecules cling together as they are pulled through the plant. They also adhere to the walls of the xylem (adhesion). This creates one unbroken column of water through the plant. The upward pull on the cohesive sap creates a tension (a negative pressure). This helps water uptake and movement up the plant.

### 3 Root pressure

Water entering the stele from the soil creates a root pressure; a weak 'push' effect for the water's upward movement through the plant. Root pressure can force water droplets from some small plants under certain conditions (guttation), but generally it plays a minor part in the ascent of water.



3. (a) What would happen if too much water was lost from the leaves? \_\_\_\_\_

(b) When might this happen? \_\_\_\_\_

4. Describe the three processes that assist the transport of water from the roots of the plant upward:

(a) \_\_\_\_\_

(b) \_\_\_\_\_

(c) \_\_\_\_\_

5. The maximum height water can move up the xylem by cohesion-tension alone is about 10 m. How then does water move up the height of a 40 m tall tree?

\_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_

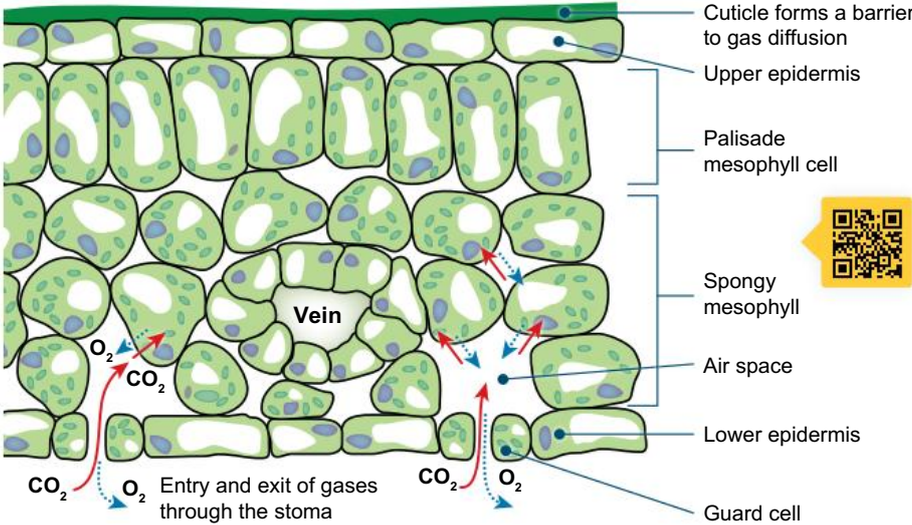
\_\_\_\_\_

# 125 Gas Exchange and Stomata

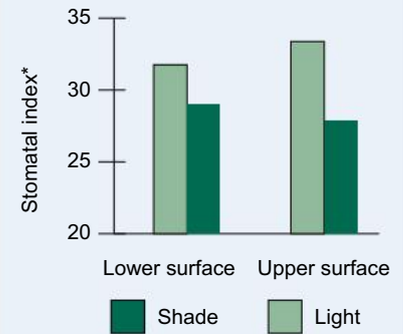
**Key Idea:** Gas exchange through stomata is associated with water losses. Guard cells help regulate these water losses. The leaf epidermis of angiosperms is covered with tiny pores, called **stomata**. Angiosperms have many air spaces between the cells of the stems, leaves, and roots. These air spaces are continuous and gases are able to move freely through

them and into the plant's cells via the stomata. Each stoma is bounded by two **guard cells**, which together regulate the entry and exit of gases (including water vapour). Although stomata permit gas exchange between the air and the photosynthetic cells inside the leaf, they are also the major routes for water loss through **transpiration**.

## Gas exchanges and the function of stomata



The number of stomata is influenced by the environment



\*Stomatal index is the percentage number of stomata compared to all the epidermal cells in a unit area of leaf.

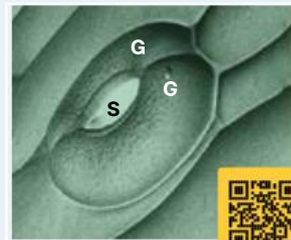
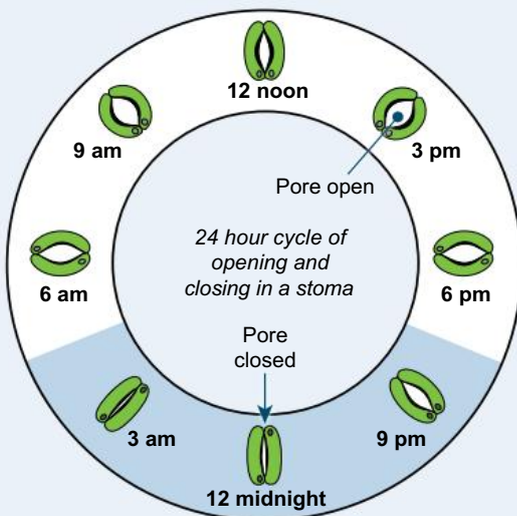
An increase in light intensity on mature leaves increases the number of stomata developing on young leaves.

### Net gas exchange in a photosynthesising dicot leaf

- ▶ Gases enter and leave the leaf through stomata. Inside the leaf (as illustrated for a dicot, above), the large air spaces and loose arrangement of the spongy mesophyll facilitate the diffusion of gases and provide a large surface area for gas exchanges.
- ▶ Respiring plant cells use oxygen (O<sub>2</sub>) and produce carbon dioxide (CO<sub>2</sub>). These gases move in and out of the plant and through the air spaces by diffusion.
- ▶ When the plant is photosynthesising, the situation is more complex. Overall there is net consumption of CO<sub>2</sub> and net production of oxygen. Fixation of CO<sub>2</sub> maintains a gradient in CO<sub>2</sub> concentration between the atmosphere (high) and the leaf tissue (low).
- ▶ Oxygen is produced in excess of respiratory needs and diffuses out of the leaf. These net exchanges are indicated by the arrows on the diagram.

## The cycle of opening and closing of stomata

The opening and closing of stomata shows a daily cycle that is largely determined by the hours of light and dark.



The image left shows a scanning electron micrograph (SEM) of a single stoma from the leaf epidermis of a dicot.

Note the guard cells (G), which are swollen tight and open the pore (S) to allow gas exchange between the leaf tissue and the external environment.

### Factors influencing stomatal opening

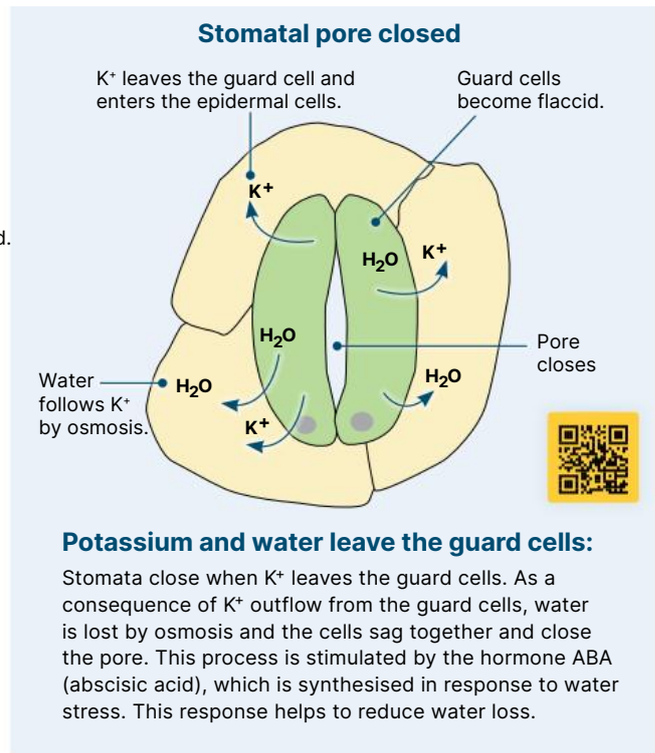
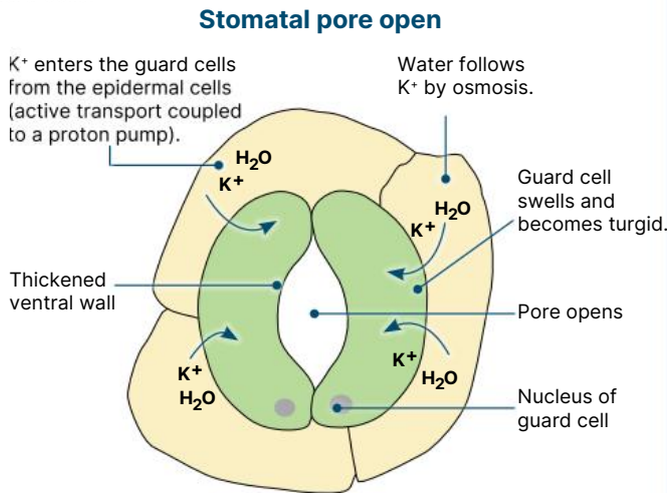
Stomata	Guard cells	Daylight	CO <sub>2</sub>	Soil water
Open	Turgid	Light	Low	High
Closed	Flaccid	Dark	High	Low

The opening and closing of stomata depends on environmental factors, the most important being light, CO<sub>2</sub> concentration in the leaf tissue, and water supply. Stomata tend to open during daylight in response to light, and close at night (left and above). Low CO<sub>2</sub> levels also promote stomatal opening. Conditions that induce water stress cause the stomata to close, regardless of light or CO<sub>2</sub> level.



## Guard cells

The guard cells on each side of a stoma control the diameter of the pore by changing shape. When the guard cells take up water by osmosis they swell and become turgid, opening the pore. When the guard cells lose water, they become flaccid and the pore closes. By this mechanism a plant can control the amount of gas entering, or water leaving, the plant. The changes in turgor pressure that open and close the pore result mainly from the reversible uptake and loss of potassium ions (and thus water) by the guard cells.



### Potassium and water enter the guard cells

Stomata open when the guard cells actively take up  $K^+$  from the neighbouring epidermal cells. As a consequence, water enters the cell by osmosis and they swell and become turgid. The walls of the guard cells are thickened more on the inside surface (the ventral wall) than the outside wall, so that when the cells swell they buckle outward, opening the pore.

### Potassium and water leave the guard cells:

Stomata close when  $K^+$  leaves the guard cells. As a consequence of  $K^+$  outflow from the guard cells, water is lost by osmosis and the cells sag together and close the pore. This process is stimulated by the hormone ABA (abscisic acid), which is synthesised in response to water stress. This response helps to reduce water loss.

1. Describe two adaptive features of leaves:

- (a) \_\_\_\_\_
- (b) \_\_\_\_\_

2. For a terrestrial flowering plant, with no special adaptations for water conservation (a mesophyte):

(a) Describe the net gas exchanges between the air and the cells of the mesophyll in the dark (no photosynthesis):

\_\_\_\_\_

(b) Explain how this situation changes when a plant is photosynthesising: \_\_\_\_\_

\_\_\_\_\_

3. Describe two ways in which the continuous air spaces through the plant facilitate gas exchange:

- (a) \_\_\_\_\_
- (b) \_\_\_\_\_

4. Outline the role of stomata in gas exchange in a flowering plant: \_\_\_\_\_

\_\_\_\_\_

5. (a) Explain how the guard cells open the stomata: \_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_

(b) Explain how the guard cells close the stomata: \_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_

# Conditions for Photosynthesis

**Key Idea:** Photosynthesis requires light to proceed. It produces glucose molecules, which are stored as starch.

**Photosynthesis** produces glucose which is stored in plant leaves as granules of starch (a glucose polymer) within the photosynthesising cells. In most plants, the leaves are green

due to the presence of **chloroplasts**. In some plants (often those labelled as ornamentals) the leaves can be variegated and carry white or often red patches. Sometimes the entire leaf can be red (e.g. Japanese maples). How does this affect the production of starch in the leaf?



The vast majority of plants have green leaves or stems. The green colour comes from the **chlorophyll** pigment, which absorbs red and blue light and reflects green (A). Some plants have white leaf patches (B) while others may have other colours, most commonly red (C).

### Testing for starch

Students investigated photosynthesis in the three types of leaves above (A, B, C) in an experiment outlined below:

- ▶ Three plants A, B, C, (as above) were placed in darkness for 48 hours. One leaf (L) was then removed from plant A.
- ▶ Several leaves from each plant were then covered with aluminium foil to block the light and the plants were placed back into the light for 24 hours.
- ▶ During this time, leaf L was placed into a test tube containing ethanol at its boiling point for ten minutes to remove any pigments from the leaf. This produced a white leaf. The leaf was then placed back into hot water for 20 seconds to soften it.
- ▶ Leaf L was tested for starch by applying iodine solution. The applied solution remained brown indicating the absence of starch.
- ▶ After 24 hours, a covered and uncovered leaf from each plant was randomly selected. Each leaf was tested for the presence of starch as described above.
- ▶ The results are shown below



Plant with foil covered leaves

Iodine test result		
Plant	Uncovered leaf	Covered leaf
A	Intense blue/black over all leaf	Applied solution remains brown
B	Less intense blue/black over most of the leaf. More intense near centre.	Applied solution remains brown
C	Intense blue/black over middle regions of leaf. Brown on outer edges	Applied solution remains brown

1. Explain the results found in each plant:

- (a) A: \_\_\_\_\_  
 \_\_\_\_\_
- (b) B: \_\_\_\_\_  
 \_\_\_\_\_
- (c) C: \_\_\_\_\_  
 \_\_\_\_\_

2. Why was leaf L tested for starch immediately after removing the plant from the dark? \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_

3. What two things does the experiment show the plants need for photosynthesis (as shown by the production of starch):  
 \_\_\_\_\_  
 \_\_\_\_\_



# 127 Investigating Plant Transpiration

**Key Idea:** The relationship between the rate of transpiration and the environment can be investigated using a potometer. In this activity, you will investigate the effect of different

environmental conditions on **transpiration** rate using a **potometer**. You will use the results to predict the kinds of conditions that cause the greatest water losses.

## The potometer

A potometer is a simple instrument for investigating transpiration rate (water loss per unit time). The equipment is simple to use and easy to obtain. A basic potometer, such as the one shown right, can easily be moved around so that transpiration rate can be measured under different environmental conditions.

Some physical conditions investigated are:

- Humidity or vapour pressure (high or low)
- Temperature (high or low)
- Air movement (still or windy)
- Light level (high or low)
- Water supply

It is also possible to compare the transpiration rates of plants with different adaptations e.g. comparing transpiration rates in plants with rolled leaves vs rates in plants with broad leaves. If possible, experiments like these should be conducted simultaneously using replicate equipment. If conducted sequentially, care should be taken to keep the environmental conditions the same for all plants used.



A potometer attached to a data logger

Pasco

## Investigation 8.1 Investigating plant transpiration

See appendix for equipment list.

1. Four different conditions that influence transpiration will be tested: room conditions (ambient), wind, bright light, and high humidity.
2. Before starting, your teacher will decide if your group is to test one of these conditions (and which one) and pool class data for all four.
3. Set up the potometer and plant as in the diagram. It is best if the plant leaves used are large and few (4-6 leaves) rather than small and many. Alternatively the plant can be placed in a 250 mL conical flask with 200 mL of water and a thin layer of cooking oil floated on top. This is weighed before the experiment and then every 3 minutes (or as the experiment requires). The difference in mass in grams is equal to the volume of water transpired in mL.
4. After setting up the potometer, let the apparatus equilibrate for 10 minutes, and then record the position of the air bubble in the pipette (or the mass of the equipment for the alternative method). This is time 0 and position 0.
5. The plant can now be exposed to one of the four conditions. Record results in Table 1.
6. For the ambient environment the equipment can be placed on the bench away from bright light or wind. Record the net movement of the bubble every 3 minutes for 30 minutes.
7. For the high wind environment the equipment can be placed on the bench in front of a fan set on a moderate speed (away from bright light). Record the net movement of the bubble every 3 minutes for 30 minutes.
8. For the bright light environment, the equipment can be placed on the bench in front of a bright light (about



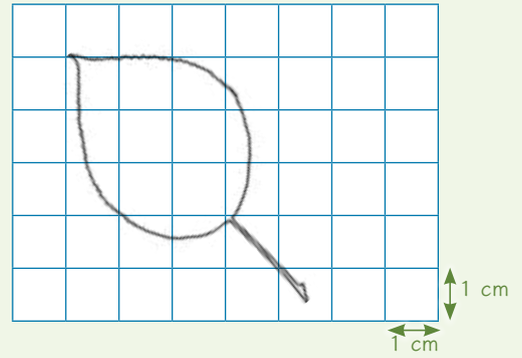
SU

SI



40 cm away). Record the net movement of the bubble every 3 minutes for 30 minutes.

9. For the high humidity environment the equipment can be placed on a bench away from bright light, in a sealed plastic bag with 2-3 sprays of water from a spray bottle. Record the net movement of the bubble every 3 minutes for 30 minutes.
10. It is important that for fair comparison of transpiration the area of leaf used in each environment (or by different groups) should be calculated and the volume of water lost per square centimetre compared ( $\text{mL cm}^{-2}$ ).
11. Leaf area can be measured by tracing the leaves onto graph paper and counting the squares, or by tracing or photocopying the leaves onto a paper of a known mass per area, then cutting out the shapes and weighing them. For both methods, multiply by 2 for both leaf surfaces.
12. Once the area of the leaf is calculated the transpiration (water lost) in  $\text{mL cm}^{-2}$  can be calculated for each time recording and record in Table 2.



**Table 1. Potometer readings (in mL water loss)**

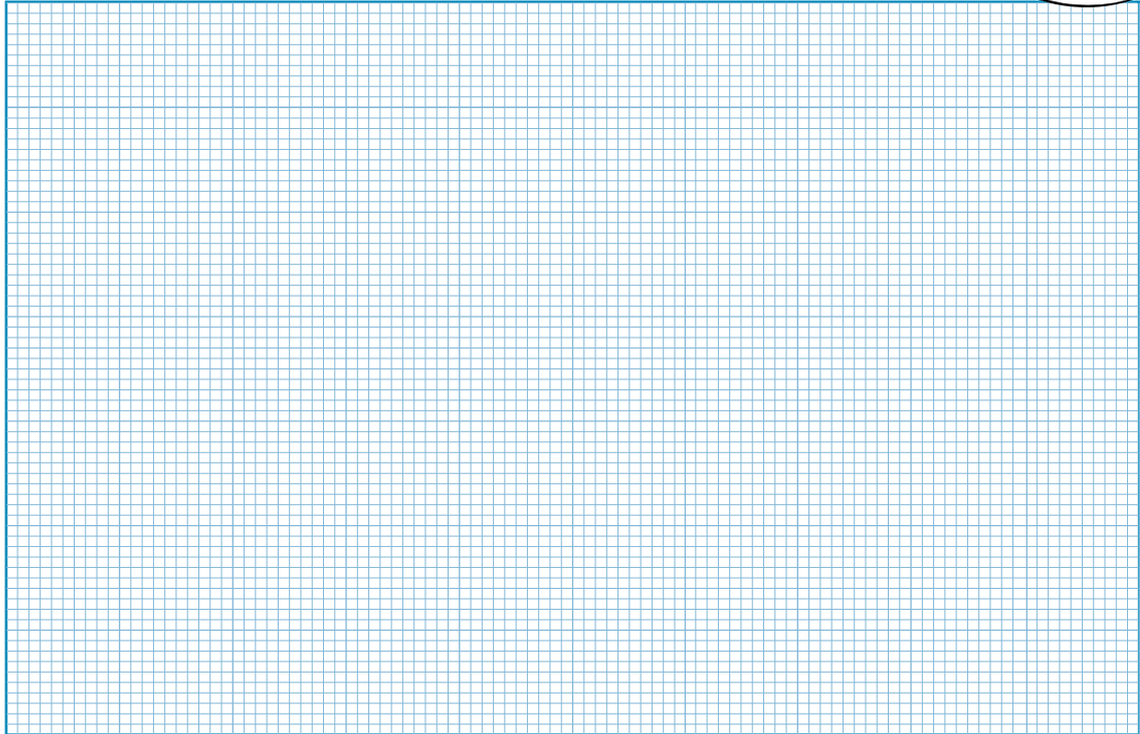
Time (min)	0	3	6	9	12	15	18	21	24	27	30
Treatment											
Ambient											
Wind											
High humidity											
Bright light											

**Table 2. Potometer readings in mL per  $\text{cm}^2$**

Time (min)	0	3	6	9	12	15	18	21	24	27	30
Treatment											
Ambient											
Wind											
High humidity											
Bright light											

1. Measure the area of the leaves you used: \_\_\_\_\_
2. Why is comparing water loss per square cm over time more important than just comparing the water loss over time?  
 \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_

3. Plot the data in Table 2 on the grid provided:



4. Identify the independent variable: \_\_\_\_\_

5. (a) Identify the control: \_\_\_\_\_

(b) Explain the purpose of including an experimental control in an experiment:

\_\_\_\_\_  
\_\_\_\_\_  
\_\_\_\_\_

6. (a) Which factors increased water loss? \_\_\_\_\_

(b) How does each environmental factor influence water loss? \_\_\_\_\_

\_\_\_\_\_  
\_\_\_\_\_  
\_\_\_\_\_  
\_\_\_\_\_

7. From your results predict how each of the following conditions might influence transpiration:

(a) Low humidity (e.g. dry desert): \_\_\_\_\_

(b) Low light levels (e.g. overcast day): \_\_\_\_\_

(c) Hot dry winds: \_\_\_\_\_

8. How might different types of plants affect the results? \_\_\_\_\_

\_\_\_\_\_  
\_\_\_\_\_

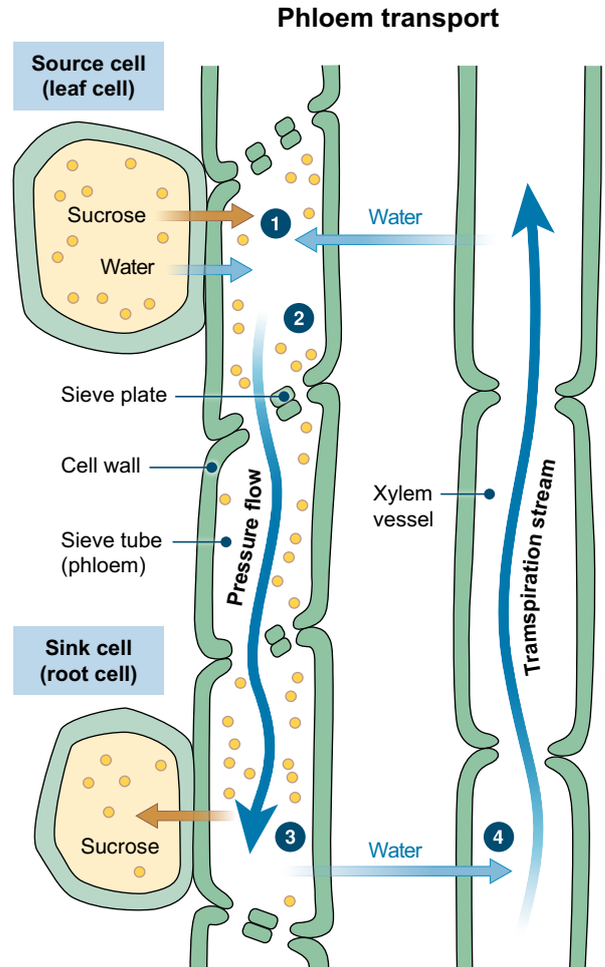
**Key Idea:** Phloem transports the organic products of photosynthesis (sugars) through the plant by translocation. In angiosperms, the sugar moves through the phloem **sieve-tube** members, which are arranged end-to-end and perforated with **sieve plates**. Apart from water, **phloem** sap contains mainly sucrose (up to 30%). It may also contain minerals, hormones, and amino acids in transit around the plant. Movement of sap in the phloem is from a source (a plant

organ where sugar is made or mobilised) to a sink (a plant organ where sugar is stored or used). Loading sucrose into the phloem at a source involves energy expenditure. We know this because it is slowed or stopped by high temperatures or respiratory inhibitors. In some plants, unloading the sucrose at the sinks also requires energy, although in others, diffusion alone is sufficient to move sucrose from the phloem into the cells of the sink organ.

## Phloem transport

Phloem sap moves from source to sink at rates as great as  $100 \text{ m h}^{-1}$ , which is too fast to be accounted for by cytoplasmic streaming. The most acceptable model for phloem movement is the mass flow hypothesis (also known as the pressure flow hypothesis). Phloem sap moves by bulk flow, which creates a pressure (hence the term "pressure-flow"). The key elements in this model are outlined below and right. For simplicity, the cells that lie between the source (and sink) cells and the phloem sieve-tube have been omitted.

- 1 Loading sugar into the phloem increases the solute concentration inside the sieve-tube cells. This causes the sieve-tubes to take up water by osmosis.
- 2 The water uptake creates a hydrostatic pressure that forces the sap to move along the tube, just as pressure pushes water through a hose.
- 3 The pressure gradient in the sieve tube is reinforced by the active unloading of sugar and consequent loss of water by osmosis at the sink (e.g. root cell).
- 4 Xylem recycles the water from sink to source.



Source: Modified after Campbell Biology 1993



**Measuring phloem flow**

Aphids can act as natural phloem probes to measure phloem flow. The sucking mouthparts (stylet) of the insect penetrates the phloem sieve-tube cell. While the aphid feeds, it can be severed from its stylet, which remains in place and continues to exude sap. Using different aphids, the rate of flow of this sap can be measured at different locations on the plant.

1. (a) From what you know about osmosis, explain why water follows the sugar as it moves through the phloem:

---



---



---



---

(b) What is meant by 'source to sink' flow in phloem transport? \_\_\_\_\_

---



---

2. Why does a plant need to move food around, particularly from the leaves to other regions?

---



---



---

# 129 Plants and Technology

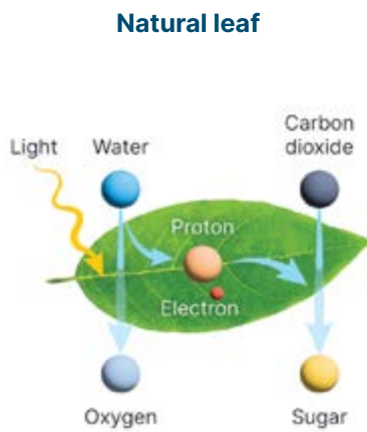
**Key Idea:** Plant processes can be replicated with technology to produce valuable products for human use.

Plants convert light energy into a form of chemical energy which they can use for cellular functioning through the process of **photosynthesis**. **Artificial leaves** and **bionic leaves** are new technologies developed to mimic photosynthesis by

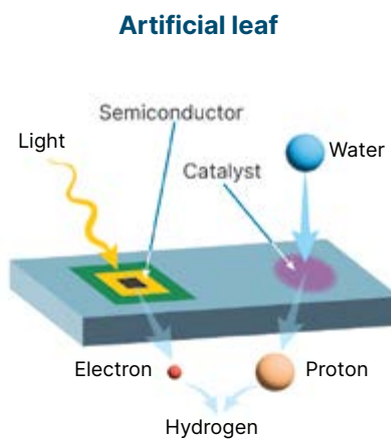
harvesting light energy and producing sustainable hydrogen biofuel. This can be further processed into carbon-based fuels, plastics, and medicines by co-opting bacteria. Although the technology is only in prototype phase and not yet scaled up for commercial use, artificial photosynthesis has the potential to convert a limitless energy source into useful products.

## Copying photosynthesis

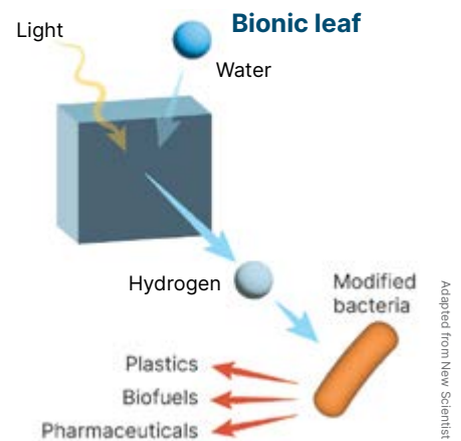
- ▶ Photosynthesis, used by plants to produce glucose, has been successfully imitated by technology on a small scale. Various mechanical and chemical techniques are used to produce an artificial leaf. This is able to transform light energy into an alternative storable energy form.
- ▶ Another technology, the bionic leaf, involves genetically modified bacteria using the outputs of the artificial leaf, specifically hydrogen, and converting it into biofuel in the form of alcohol.
- ▶ These processes must be scalable to be commercially feasible. Broadening the range of products produced, such as medicines, is an important goal.



At a cellular level, plant leaves use chloroplasts to convert carbon dioxide and water into glucose and oxygen, transforming light energy into stored chemical energy. This conversion process occurs in every photosynthetic cell. It has an efficiency of approximately 1% in converting light energy into chemical energy.



Artificial 'leaves' harness electrons from light energy through a silicon solar panel and a semiconductor. They extract protons from water using a chemical catalyst made of nickel and cobalt. The electrons and protons are then merged to generate hydrogen.



Bionic 'leaves' continue the conversion of hydrogen generated from artificial leaves by utilising genetically modified bacteria (*Ralstonia eutropha*). The bacteria use a number of processes that result in the production of beneficial carbon-based products such as biofuels. By altering the bacterial DNA further, it is possible to create specialised substances such as medicines, pesticides, and fertilisers.

1. Summarise steps in the process of copying photosynthesis through artificial leaves and bionic leaves:

---



---



---



---



---

2. Suggest some potential benefits and challenges associated with scaling up the artificial photosynthesis process for commercial use?

---



---



---



---

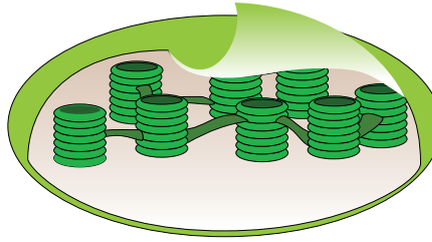


---



# 130 Did you Get It?

- Label the following features of a chloroplast on the diagram below:  
*granum, stroma, thylakoid disc, stroma lamellae.*
  - Indicate on the diagram where the light dependent and light independent reactions occur.



2. Test your vocabulary by matching each term to its correct definition by writing the letter in the correct box.

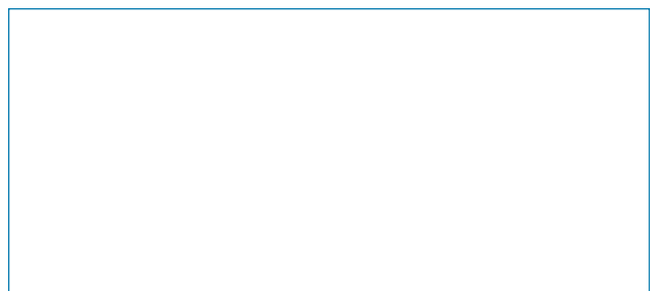
(i) cohesion-tension	<input type="checkbox"/>	<b>A</b> Device used for investigating the rate of transpiration.
(ii) guard cells	<input type="checkbox"/>	<b>B</b> The loss of water vapour by plants, mainly from leaves via the stomata.
(iii) phloem	<input type="checkbox"/>	<b>C</b> Specialised cells either side of the stoma, which open or close the pore.
(iv) potometer	<input type="checkbox"/>	<b>D</b> Vascular tissue that conducts water and minerals from the roots to the rest of the plant.
(v) stomata	<input type="checkbox"/>	<b>E</b> Pores in the leaf surface through which gases and water vapour can pass
(vi) transpiration	<input type="checkbox"/>	<b>F</b> Partial explanation for the movement of water up the plant in the transpiration stream.
(vii)xylem	<input type="checkbox"/>	<b>G</b> Tissue that conducts dissolved sugars in vascular plants. Largely made up of sieve tubes and companion cells.

3. Transpiration in a hydrangea shoot was investigated using a potometer. The experiment was set up and the plant left to stabilise (environmental conditions: still air, light shade, 20°C). The plant was then placed in different environmental conditions and the water loss was measured each hour. Finally, the plant was returned to original conditions, allowed to stabilise and transpiration rate measured again. The results are presented below:

Experimental conditions	Temperature (°C)	Humidity (%)	Transpiration rate (g h <sup>-1</sup> )
(a) Still air, light shade, room temperature	20	70	1.20
(b) Moving air, light shade	20	70	1.60
(c) Still air, bright sunlight	23	70	3.75
(d) Still air and dark, moist chamber	19.5	100	0.05

- What conditions acted as the control in this experiment? \_\_\_\_\_
- Which factors increased transpiration rate and why? \_\_\_\_\_  
\_\_\_\_\_  
\_\_\_\_\_
- Why did the plant have such a low transpiration rate in humid, dark conditions? \_\_\_\_\_  
\_\_\_\_\_  
\_\_\_\_\_

4. The leaf below was left in light for 24 hours then tested for starch. In the space provided draw a diagram of the leaf below to show where you would expect to find starch:





4. (a) The photograph shows a structure on a plant leaf. Identify the structure arrowed in the photograph:



(b) When is this structure likely to be open (day or night)?

(c) Explain the steps and structures involved in the opening of the stoma to allow carbon dioxide gas to enter the plant leaf:

---



---



---

5. Compare and contrast the structure and function of the phloem and xylem tissue in vascular plants:

---



---



---



---



---



---



---



---



---



---

6. Discuss how ATP is formed as a product of catabolic reaction and then facilitates anabolic reactions, using cellular respiration and photosynthesis as examples. Use the box to draw a simple diagram showing the processes:

---



---



---



---



---



---



---



---



---



---

# Neural Homeostatic Controls



## Key Terms

- acetylcholine
- action potential
- chemoreceptor
- depolarisation
- effector
- homeostasis
- hyperpolarisation
- mechanoreceptor
- motor neuron
- myelin
- negative feedback
- nerve impulse
- neuron
- nociceptor
- node of Ranvier
- photoreceptor
- proprioceptor
- refractory period
- response
- resting potential
- sensory neuron
- sensory receptor
- stimulus (pl. stimuli)
- summation
- synapse
- synaptic integration
- thermoreceptor
- threshold potential
- transducer

## Key Concepts

- ▶ Homeostasis allows the body to maintain a constant physiological state independent of the external environment.
- ▶ The nervous system helps maintain homeostasis by receiving and processing environmental stimuli and responding to the information.

## Principles of homeostasis

### Activity Number

<input type="checkbox"/> 1	Interpret information from models of negative feedback stabilising systems.	133
<input type="checkbox"/> 2	Describe the role of sensory receptors in detecting stimuli. Classify receptors based on the stimuli to which they respond, including chemoreceptors, photoreceptors, thermoreceptors, mechanoreceptors, and nociceptors.	134
<input type="checkbox"/> 3	Describe the role of effectors (muscles and glands) in bringing about the response to stimuli.	132, 135
<input type="checkbox"/> 4	<b>SHE:</b> Explain that homeostasis (steady state) involves a stimulus-response model in which change in the internal or external environment (the stimulus) is detected and appropriate responses occur via negative feedback.	132-133
<input type="checkbox"/> 5	<b>SI:</b> Explain, giving examples, how negative feedback systems operate.	133-134

## Structure and function of neurons

<input type="checkbox"/> 6	Explain the role of the nervous system in receiving stimuli through sensory receptors and responding to those stimuli through effectors. Identify pathways for neural coordination in a mammal, including reference to the role of the central nervous system.	135
<input type="checkbox"/> 7	Identify the cells that transmit nerve impulses from sensory receptors to the central nervous system and from the central nervous system to effectors.	136
<input type="checkbox"/> 8	Distinguish between sensory and motor neurons in terms of their structure and function. Include reference to dendrites, soma (cell body), axon, myelin sheath, nodes of Ranvier, axon terminal, and synapse.	136
<input type="checkbox"/> 9	Describe the role of reflexes in providing rapid responses to stimuli. Giving examples, distinguish between monosynaptic and polysynaptic reflex arcs.	137

## Transmission of nerve impulses

<input type="checkbox"/> 10	Describe the generation and transmission of nerve impulses in mammals.	138
<input type="checkbox"/> 11	Describe the structure of a chemical synapse (e.g. a cholinergic synapse). Identify pre- and post-synaptic neurons, vesicles containing neurotransmitters (e.g. acetylcholine), and the synaptic cleft.	139
<input type="checkbox"/> 12	Describe impulse transmission across a synapse (e.g. a cholinergic synapse) to include the role of calcium, diffusion of the neurotransmitter (e.g. acetylcholine), and generation of an action potential in the post-synaptic cell. Explain how the effect of the neurotransmitter depends on the neurotransmitter involved, its position in the nervous system, and the properties of the post-synaptic cell.	139
<input type="checkbox"/> 13	Explain synaptic integration and the role of synapses in summation and control of nervous system responses.	140
<input type="checkbox"/> 14	Explain, using examples, how synaptic transmission can be affected by drugs and poisons (e.g. snake venom).	141

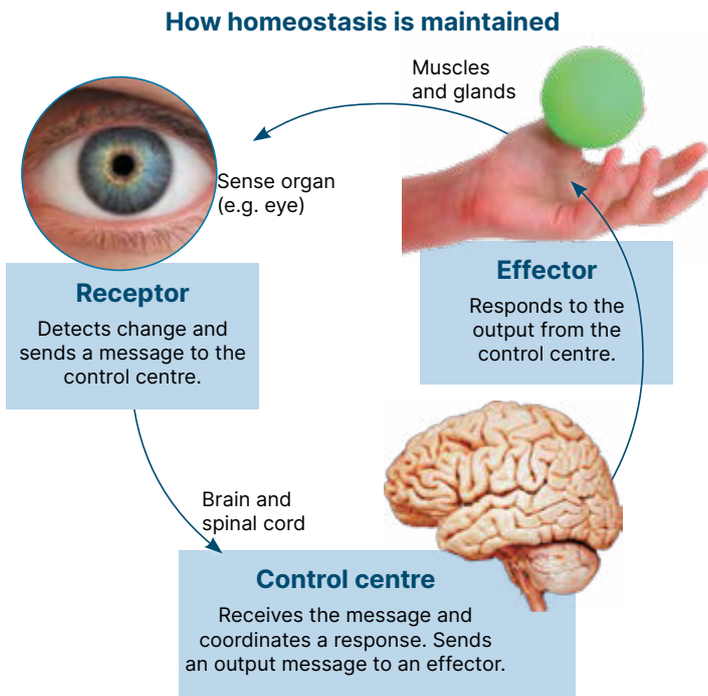
**Key Idea:** Homeostasis is the process of sustaining a constant physiological state within the body, regardless of fluctuations in the external environment.

Organisms maintain a relatively constant physiological state, called **homeostasis**, despite changes in their environment. Any change in the environment to which an organism responds is called a **stimulus**. Environmental stimuli are constantly changing, so organisms must adjust their behaviour and physiology constantly to maintain homeostasis. This requires the coordinated activity of the body's organ systems. Homeostatic mechanisms prevent potentially harmful deviations from the steady state, and keep the body's internal conditions within strict limits.

Homeostasis is required to maintain constant body temperature, at about 37°C. Similarly, you must regulate blood sugar (glucose) levels and blood pH, water and electrolyte balance, and blood pressure. Your body's organ systems coordinate to carry out these tasks.

### Maintaining homeostasis: the stimulus-response model

To maintain homeostasis, the body must detect stimuli through **receptors**, process this sensory information in a control centre, and respond to it appropriately via an **effector**. The responses provide new feedback to the receptor. These three components are illustrated below.



### Homeostasis analogies



The analogy of a temperature setting on a heat pump is a good way to explain how homeostasis is maintained. A heat pump has sensors (a receptor) to monitor room temperature. It also has a control centre to receive and process the data from the sensors. Depending on the data it receives, the control centre activates the effector (heating/cooling unit), switching either on or off.

When the room is too cold, the heating unit switches on, and the cooling unit is off. When it is too hot, the heating unit switches off and the cooling unit is switched on. This system maintains a constant temperature, similar to homeostasis in the body.



The analogy of staying upright on a mountain bike, using body weight, arms, pedals, brakes, and steering, demonstrates that many homeostasis systems have multiple mechanisms to maintain a steady state.

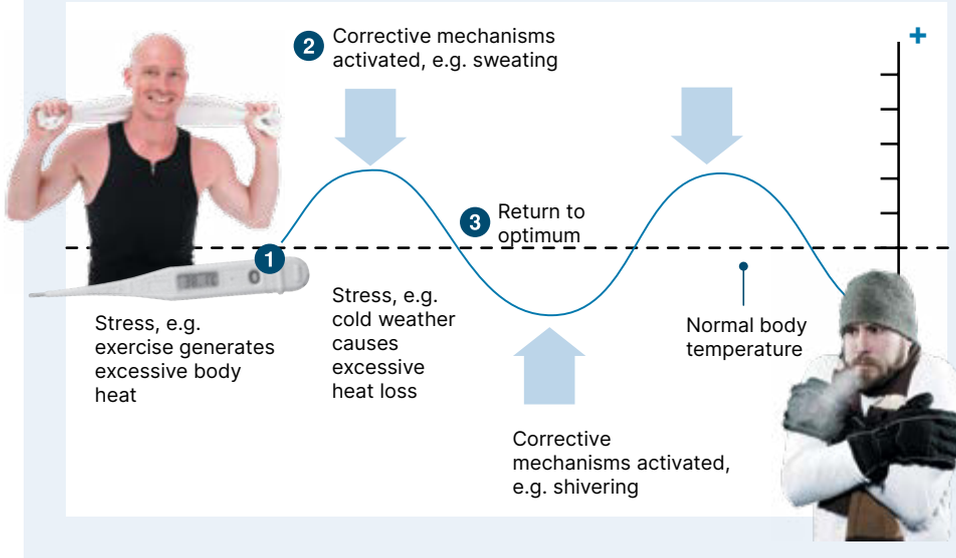
1. What is homeostasis? \_\_\_\_\_  
\_\_\_\_\_
2. What is the role of the following components in maintaining homeostasis:
  - (a) Receptor: \_\_\_\_\_  
\_\_\_\_\_
  - (b) Control centre: \_\_\_\_\_  
\_\_\_\_\_
  - (c) Effector: \_\_\_\_\_  
\_\_\_\_\_

# 133 Negative Feedback

**Key Idea:** Negative feedback mechanisms detect departures from a desired set point and act to restore the steady state. **Negative feedback** is a regulatory mechanism that maintains the body's **homeostasis** by detecting deviations from a set point and acting to restore those set point conditions.

Negative feedback mechanisms act to dampen variations and so have a stabilising effect on biological systems. Most body systems achieve homeostasis through negative feedback. Body temperature, blood pressure, and blood glucose levels are all controlled by negative feedback mechanisms.

## Negative feedback in temperature regulation

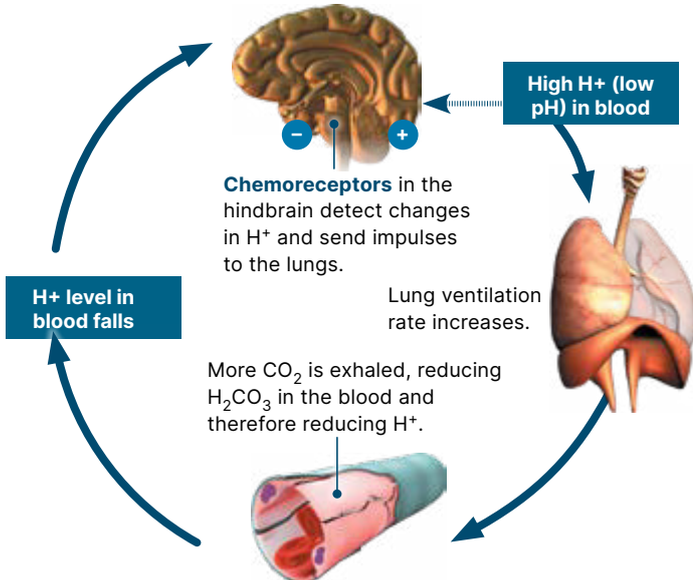


The diagram (left) shows how temperature is regulated by negative feedback mechanisms.

- 1 A stressor, e.g. exercise, takes the internal environment away from optimum.
- 2 Stress is detected by receptors and corrective mechanisms (e.g. sweating or shivering) are activated.
- 3 Corrective mechanisms act to restore optimum conditions.

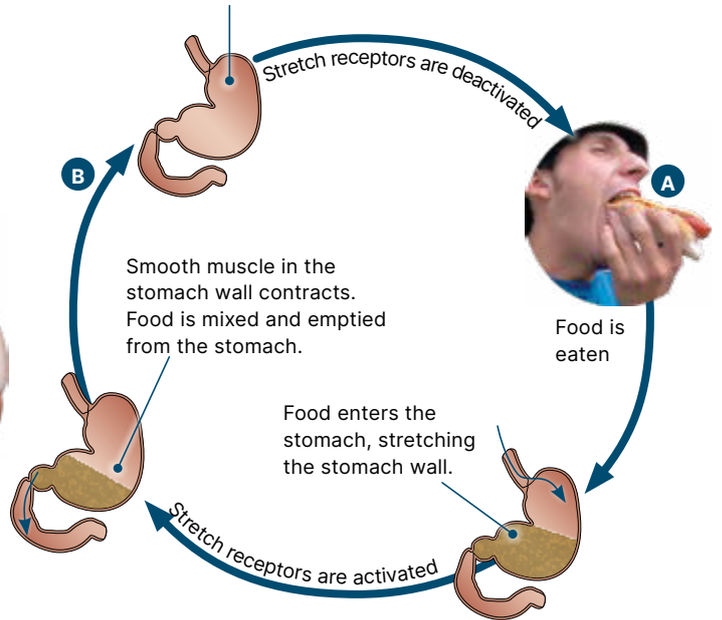
## Negative feedback in blood pH

Regulation of ventilation rate helps to maintain blood pH between 7.35 and 7.45. Low blood pH stimulates increased breathing rate, which reduces  $H^+$  via exhalation. This reduces sensory input to the medulla and breathing returns to normal.



## Negative feedback in stomach emptying

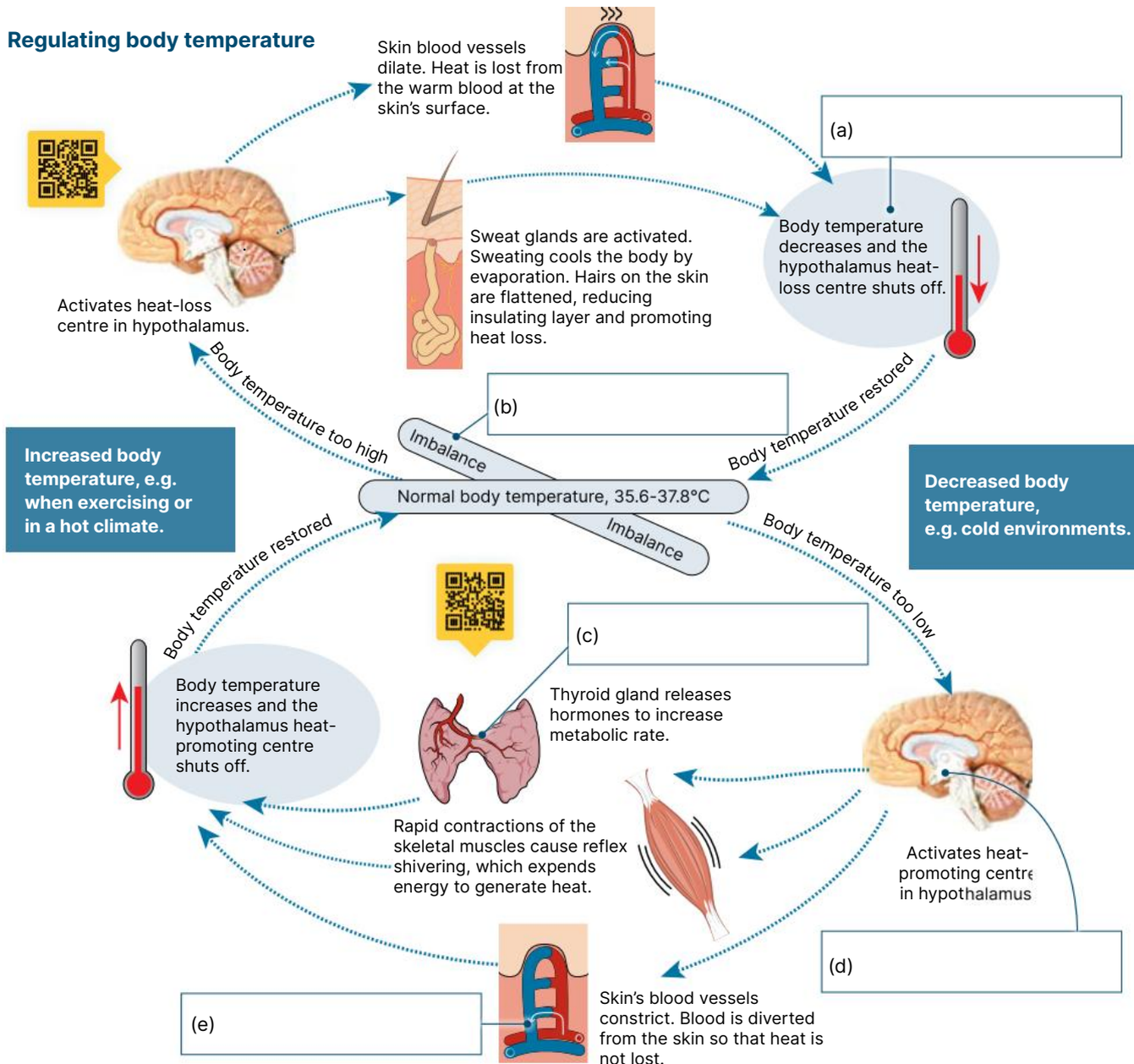
Empty stomach. Stomach wall is relaxed.



1. How do negative feedback mechanisms maintain homeostasis in a variable environment? \_\_\_\_\_
2. On the diagram of stomach emptying:
  - (a) State the stimulus at A: \_\_\_\_\_ State the response at B: \_\_\_\_\_
  - (b) Name the effector in this system: \_\_\_\_\_
  - (c) What is the steady state for this example? \_\_\_\_\_

### How is body temperature regulated?

- ▶ In humans, the temperature regulation centre is a region of the brain called the hypothalamus. It has **thermoreceptors** that monitor core body temperature and has a 'set-point' temperature of 36.7°C.
- ▶ The hypothalamus acts like a thermostat. It registers changes in the core body temperature and also receives information about temperature changes from thermoreceptors in the skin. It then coordinates nervous and hormonal **responses** to counteract the changes and restore normal body temperature, as shown in the feedback diagram below. When normal temperature is restored, the corrective mechanisms are switched off.

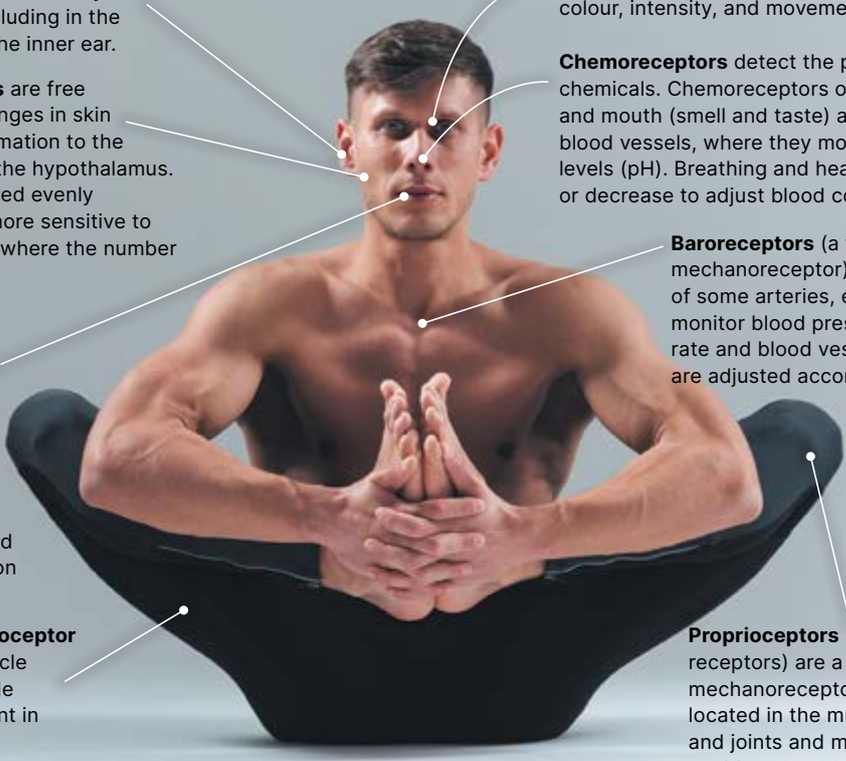


- In the diagram above showing the regulation of body temperature:
  - Identify the stimulus: \_\_\_\_\_
  - Identify the effectors: \_\_\_\_\_
  - Identify the control centre: \_\_\_\_\_
  - What structure(s) would you add to represent the receptors? \_\_\_\_\_
- Label the diagram above by appropriately adding the labels: stimulus, receptors, control centre, and effectors.
- How do the effectors restore body temperature when it increases above the set point? \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_

# 134 Sensory Receptors

**Key Idea:** Sensory receptors allow the body to respond to a range of stimuli in the internal and external environments. A **stimulus** is any physical or chemical change in the environment capable of provoking a **response** in an organism. Organisms respond to stimuli in order to survive. Stimuli may

be either external (outside the organism) or internal (within its body). Some of the **sensory receptors** that animals use to detect stimuli are shown below. Sensory receptors respond only to specific stimuli, so the sense organs an animal has determines how it perceives the world.



**Mechanoreceptors** respond to physical (mechanical) pressure or distortion. They are found throughout the body including in the skin, muscles and joints, and the inner ear.

Hot and cold **thermoreceptors** are free nerve endings that detect changes in skin temperature and provide information to the temperature control centre in the hypothalamus. Thermoreceptors are not located evenly around the body. The skin is more sensitive to temperature changes in areas where the number of thermoreceptors is denser.

**Nociceptors** are sensory **neurons** activated by noxious, potentially damaging stimuli, including extremes in temperature and pressure and toxic chemicals. External nociceptors are located in the skin, cornea, mouth and nose, Internal nociceptors are located in several organs. Nociception is experienced as pain.

The muscle spindle is a **proprioceptor** that monitors the state of muscle contraction and enables muscle to maintain its length (important in posture and muscle tone).

**Photoreceptor** cells in the eyes detect colour, intensity, and movement of light.

**Chemoreceptors** detect the presence of chemicals. Chemoreceptors occur in the nose and mouth (smell and taste) and in certain blood vessels, where they monitor blood CO<sub>2</sub> levels (pH). Breathing and heart rate increase or decrease to adjust blood composition.

**Baroreceptors** (a type of mechanoreceptor) in the walls of some arteries, e.g. aorta, monitor blood pressure. Heart rate and blood vessel diameter are adjusted accordingly.

**Proprioceptors** (position receptors) are a type of mechanoreceptor. They are located in the muscles, tendons, and joints and monitor limb position, stretch, and tension.



Temperature and pain are detected by nerve endings in the skin. Deep tissue injury is sometimes felt on the skin as referred pain.



Humans rely heavily on hearing when learning to communicate; without it, speech and language development are more difficult.



The vibration receptors in the limbs of arthropods are sensitive to movement: either sound or vibration (from struggling prey).



The chemosensory Jacobson's organ in the roof of the mouth of reptiles (e.g. snakes) enables them to detect chemical stimuli.



Breathing and heart rates are regulated in response to sensory input from internal chemoreceptors.



Baroreceptors and osmoreceptors act together to maintain blood pressure and volume.



Many insects, such as these ants, rely on chemical sense for location of food and communication.



Stimulation of nociceptors in the eye often cause rapid blinking or watering of the eye to remove the harmful stimulus.



SU

SI



## Stimuli activate ion channels

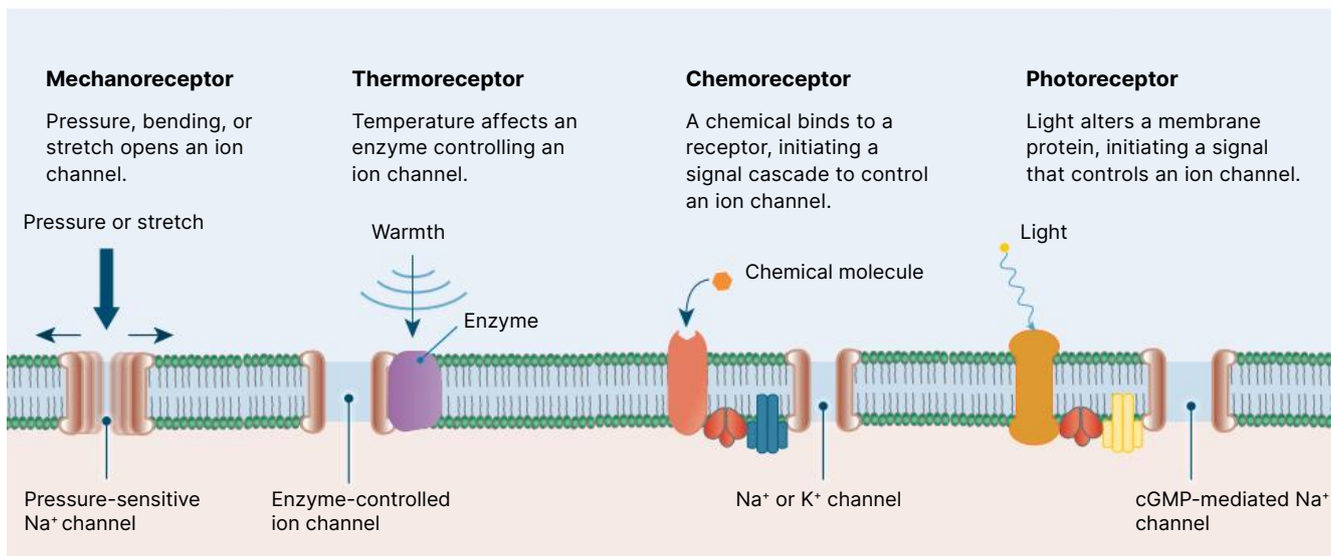
Sensory receptors respond to stimuli by producing an electrical (or chemical) discharge. In this way they act as biological **transducers**, converting the energy from a stimulus into an electrochemical signal. They can do this because the stimulus opens (or closes) ion channels and leads to localised changes in membrane potential called receptor potentials. These receptor potentials lead to **nerve impulses**, which can then be interpreted by the central nervous system (e.g. as pain, light, smell etc). The membrane potential is the difference in the concentrations of ions on opposite sides of a cellular membrane. The diagram below shows the four types of ion channels activated in stimulated receptors.



D. Fankhauser, University of Cincinnati, Clermont College

## The Pacinian corpuscle

Pacinian corpuscles (left) are pressure receptors in the deep tissues of the body. They are relatively large but structurally simple, consisting of a sensory nerve ending (dendrite) surrounded by a capsule of connective tissue layers. Pressure deforms the capsule, stretching the nerve ending and leading to a receptor potential and then a nerve impulse. This is interpreted by the central nervous system as pressure.



1. What is a stimulus? \_\_\_\_\_  
\_\_\_\_\_
2. Why is it important for an organism to be able to respond to a stimuli? \_\_\_\_\_  
\_\_\_\_\_  
\_\_\_\_\_
3. Decide if the following stimuli are internal and/or external and name the type of receptor involved in its detection:
  - (a) Light: \_\_\_\_\_
  - (b) Blood pH: \_\_\_\_\_
  - (c) Degree of muscle stretch: \_\_\_\_\_
  - (d) Temperature: \_\_\_\_\_
  - (e) Pain: \_\_\_\_\_
  - (f) Body position: \_\_\_\_\_
4. Explain how sensory receptors act as biological transducers and why this is important: \_\_\_\_\_  
\_\_\_\_\_  
\_\_\_\_\_  
\_\_\_\_\_

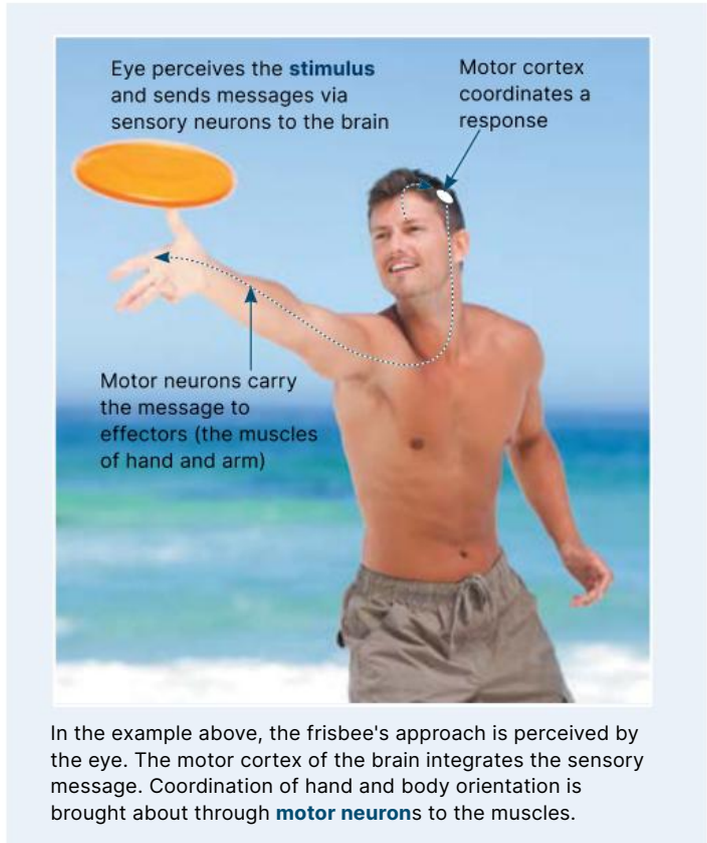
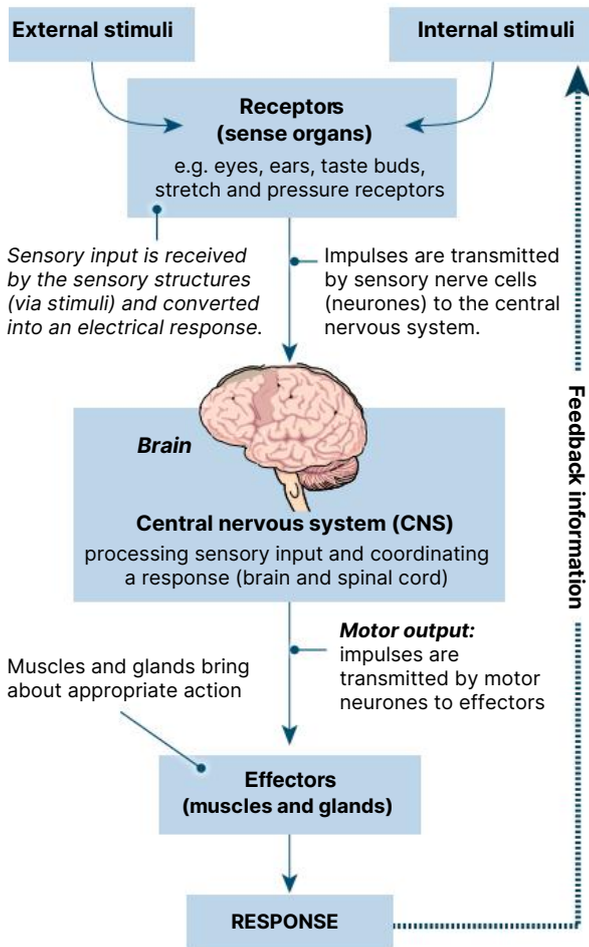
# 135 Nervous Regulation in Vertebrates

**Key Idea:** The nervous and endocrine systems work together to maintain homeostasis. Neurons of the nervous system transmit information as nerve impulses to the central nervous system, which coordinates appropriate responses to stimuli. In humans, the nervous and endocrine (hormonal) systems work together to regulate the internal environment and maintain **homeostasis** in a fluctuating environment. The

nervous system includes cells called **neurons** (nerve cells) which are specialised to transmit information in the form of electrochemical impulses (**action potentials**). The nervous system is a signalling network with branches carrying information directly to and from specific target tissues. Impulses can be transmitted over considerable distances and the **response** is very precise and rapid.

## Coordination by the nervous system

The vertebrate nervous system consists of the central nervous system (brain and spinal cord), and the nerves and receptors outside it (peripheral nervous system). Sensory input to receptors comes via stimuli. Information about the effect of a response is provided by feedback mechanisms so that the system can be readjusted. The basic organisation of the nervous system can be simplified into a few key components: the **sensory receptors**, a central nervous system processing point, and the **effectors**, which bring about the response.



Comparison of nervous and hormonal control		
	Nervous control	Hormonal control
Communication	Impulses across synapses	Hormones in the blood
Speed	Very rapid (within a few milliseconds)	Relatively slow (over minutes, hours, or longer)
Duration	Short term and reversible	Longer lasting effects
Target pathway	Specific (through nerves) to specific cells	Hormones broadcast to target cells everywhere
Action	Causes glands to secrete or muscles to contract	Causes changes in metabolic activity

1. Identify the three basic components of a nervous system and describe their role:

- (a) \_\_\_\_\_
- (b) \_\_\_\_\_
- (c) \_\_\_\_\_

2. Comment on the significance of the differences between the speed and duration of nervous and hormonal controls:

\_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_

# 136 Neurons

**Key Idea:** Neurons conduct electrical impulses from sensory receptors along axons to other neurons or to effector cells.

**Neurons** (nerve cells) transmit **nerve impulses**. Neurons have a recognisable structure with a cell body (soma) and long processes (dendrites and axons). Most neurons in the peripheral nervous system (nerves outside the brain

and spinal cord) are also supported by a fatty insulating sheath of **myelin**. Information, in the form of electrochemical impulses, is transmitted along neurons from receptors to a coordination centre and then to **effectors**. The speed of impulse conduction depends primarily on the axon diameter and whether or not the axon is myelinated.

**Motor neuron**  
Transmits impulses from the CNS to effectors (muscles or glands).

**Axon:** A long extension of the cell transmits the nerve impulse. Axons may be very long and, in the peripheral nervous system, many are myelinated.

Labels: Dendrite, Cell body, Axon hillock (generation of action potential), Myelin sheath, Axon terminal, Impulse direction.

**Axon terminals** are synaptic knobs. These release neurotransmitters, which carry the impulse between neurons or between a neuron and an effector.

Sense organ (in this case a pressure receptor) in the skin.  
Two axonal branches, one central (to the CNS) and one peripheral (to the sensory receptor). The axons of sensory neurones tend to be short.

**Sensory neuron**  
Transmits impulses from sensory receptors to the central nervous system (CNS), i.e. brain or spinal cord.

**Cell body** contains the organelles to keep the neuron alive and functioning.

Labels: Axon terminal, Node of Ranvier, Axon surrounded by myelin sheath.

**Interneuron**  
Located in the CNS and carry impulses from sensory to motor neurons (as in reflexes).

Labels: Axon terminal, Axon.

**Dendrites:** Bushy extensions of the cell body, specialised to receive stimuli.

1. Describe the basic structure of a neuron:

---

---

---

---

---

---

2. (a) Describe the structural differences between a motor and a sensory neuron:

---

---

---

---

---

---

---

---

---

---

---

---

---

---

---

---

---

---

---

---

---

---

---

---

---

---

(b) Describe a functional difference between a motor and a sensory neuron:

---

---

---

---

---

---

---

---

---

---

---

---

---

---

---

---

---

---

---

---

---

---

---

---

---

---

3. Explain why the axons of relay neurons are short, whereas those of motor neurons may be very long:

---

---

---

---

---

---

---

---

---

---

---

---

---

---

---

---

---

---

---

---

---

---

---

---

---

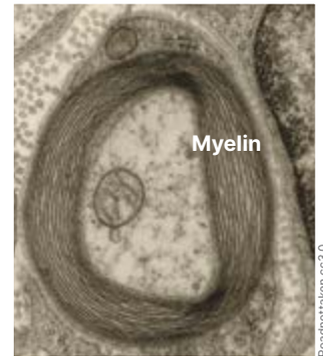
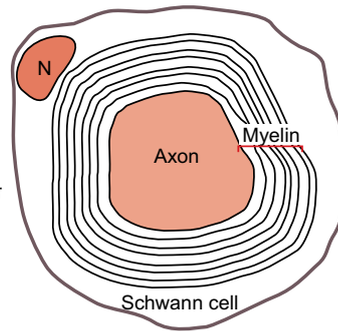
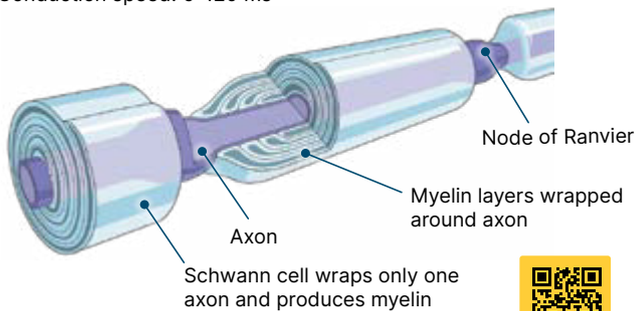
---



## Myelinated neurons

Where conduction speed is important, the axons of neurons are sheathed within a lipid-rich substance called myelin. Outside the CNS, in the peripheral nervous system, myelin is produced by specialised cells called Schwann cells. At intervals along myelinated axons, there are gaps between neighbouring Schwann cells and their sheaths called **nodes of Ranvier**. Myelin acts as an insulator, increasing the speed at which nerve impulses travel because it forces the impulse to "jump" from one uninsulated region to the next.

Diameter: 1-25  $\mu\text{m}$   
Conduction speed: 6-120  $\text{ms}^{-1}$



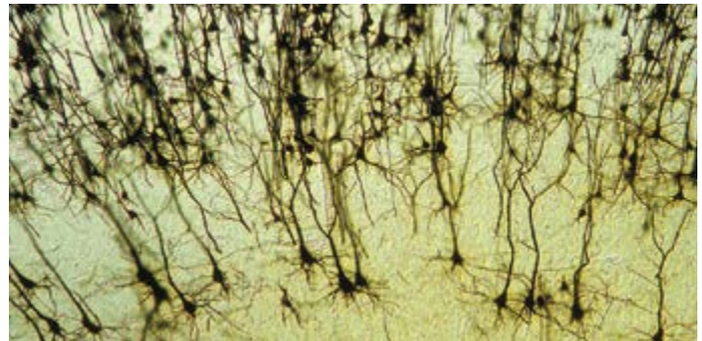
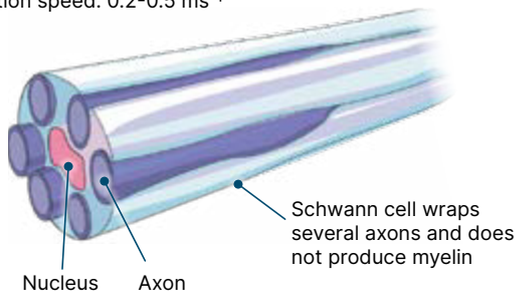
ReidHottelken cc3.0

Drawing (above left) and TEM cross section (above right) through a myelinated axon. N = nucleus of Schwann cell.

## Non-myelinated neurons

Non-myelinated axons are more common in the CNS where the distances travelled are less than in the peripheral nervous system. Here, the axons are protected by Schwann cells, but there is no myelin produced. Impulses travel more slowly because the nerve impulse is propagated along the entire axon membrane, rather than jumping from node to node as in myelinated neurons.

Diameter: <1  $\mu\text{m}$   
Conduction speed: 0.2-0.5  $\text{ms}^{-1}$



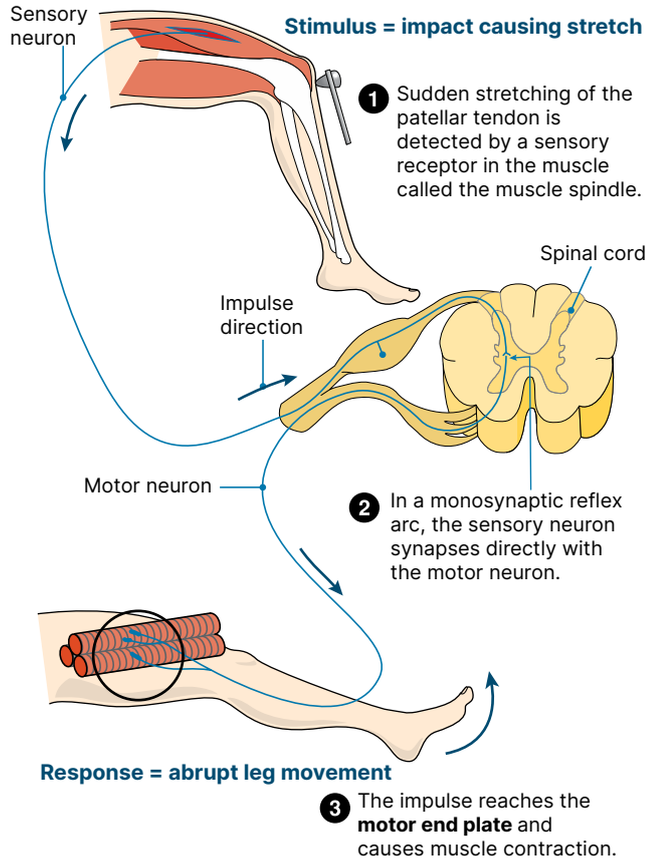
Unmyelinated pyramidal neurons in the cerebral cortex of the brain.

3. (a) What do neurons do? \_\_\_\_\_  
\_\_\_\_\_
- (b) How does this differ from supporting cells (e.g. Schwann cells)? \_\_\_\_\_  
\_\_\_\_\_
4. What is the purpose of the synaptic knobs at axon terminals? \_\_\_\_\_  
\_\_\_\_\_  
\_\_\_\_\_
5. (a) What is the function of myelination in neurons? \_\_\_\_\_  
\_\_\_\_\_
- (b) What cell type produces the myelin sheath in the peripheral nervous system? \_\_\_\_\_
- (c) Explain how an action potential travels in a myelinated neuron: \_\_\_\_\_  
\_\_\_\_\_
- (d) How does this differ from its travel in a non-myelinated neuron? \_\_\_\_\_  
\_\_\_\_\_
- (e) Why do motor neurons outside the CNS tend to be myelinated? \_\_\_\_\_  
\_\_\_\_\_

**Key Idea:** A reflex is an involuntary response to a stimulus. A reflex is an automatic **response** to a **stimulus** involving a small number of **neurons** and a central nervous system (CNS) processing point (usually the spinal cord, but sometimes the brain stem). This type of circuit is called a reflex arc. Reflexes permit rapid responses to stimuli. They are classified

according to the number of CNS **synapses** involved. Monosynaptic reflexes involve only one CNS synapse (e.g. knee jerk reflex), whereas polysynaptic reflexes involve two or more (e.g. pain withdrawal reflex). Both are spinal reflexes. The pupil reflex (opening and closure of the pupil) is an example of a cranial reflex.

### Knee-jerk reflex: A monosynaptic reflex arc



The patella (knee jerk) (left) reflex is a simple deep tendon reflex used to test the function of the femoral nerve and spinal cord segments L2-L4. It helps to maintain posture and balance when walking.



The pupillary light reflex refers to the rapid expansion or contraction of the pupils in response to the intensity of light falling on the retina. It is a polysynaptic cranial reflex and can be used to test for brain death.



Normal newborns exhibit a number of primitive reflexes in response to particular stimuli. These reflexes disappear within a few months of birth as the child develops. Primitive reflexes include the grasp reflex (above left) and the startle or Moro reflex (above right) in which a sudden noise will cause the infant to throw out its arms, extend the legs and head, and cry. The rooting and sucking reflexes are further examples of primitive reflexes.



1. Reflexes do not require conscious thought to occur. How does this provide a survival advantage?

---



---

2. (a) Describe the difference between a monosynaptic and a polysynaptic reflex arc:

---



---



---

(b) Which would produce the most rapid response, given similar length sensory and motor pathways? Explain:

---

3. What might be the survival advantage of primitive reflexes in newborns?

---



---



---



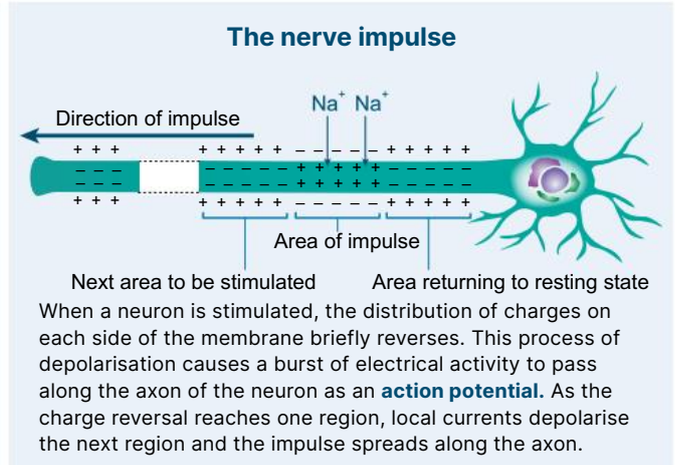
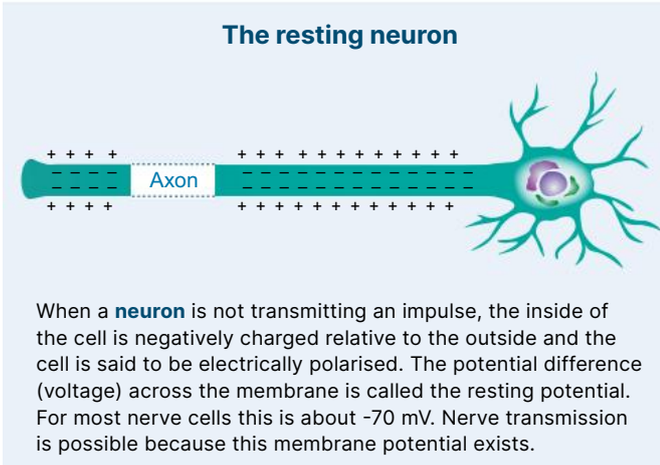
---

# 138 Transmission of Nerve Impulses

**Key Idea:** A nerve impulse occurs in response to a stimulus and involves the transmission of a membrane depolarisation along the axon of a neuron.

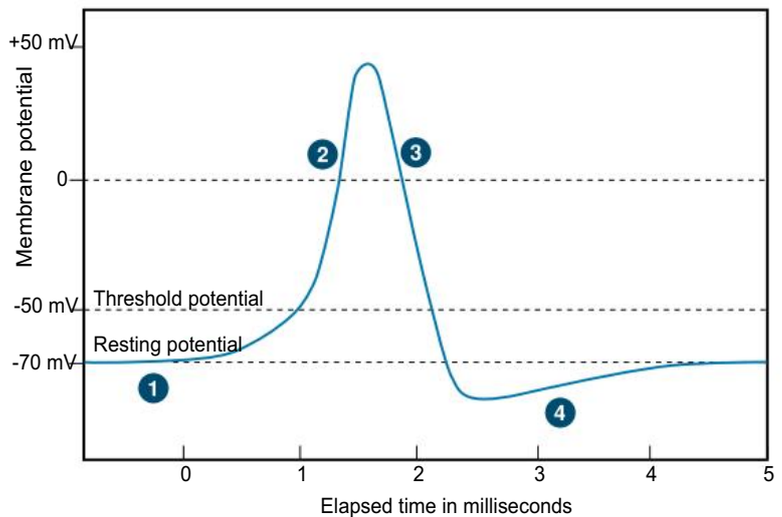
The plasma membrane of cells, including **neurons**, contain sodium-potassium ion pumps which actively pump sodium ions ( $\text{Na}^+$ ) out of the cell and potassium ions ( $\text{K}^+$ ) into the cell. The action of these ion pumps in neurons creates a separation of charge (a potential difference or voltage) either side of the membrane and makes the cells electrically excitable. It

is this property that enables neurons to transmit electrical impulses. The resting state of a neuron, with a net negative charge inside, is maintained by the sodium-potassium pumps, which actively move two  $\text{K}^+$  into the neuron for every three  $\text{Na}^+$  moved out (below left). When a nerve is stimulated, a brief increase in membrane permeability to  $\text{Na}^+$  temporarily reverses the membrane polarity (a **depolarisation**). After the **nerve impulse** passes, the sodium-potassium pump restores the **resting potential**.

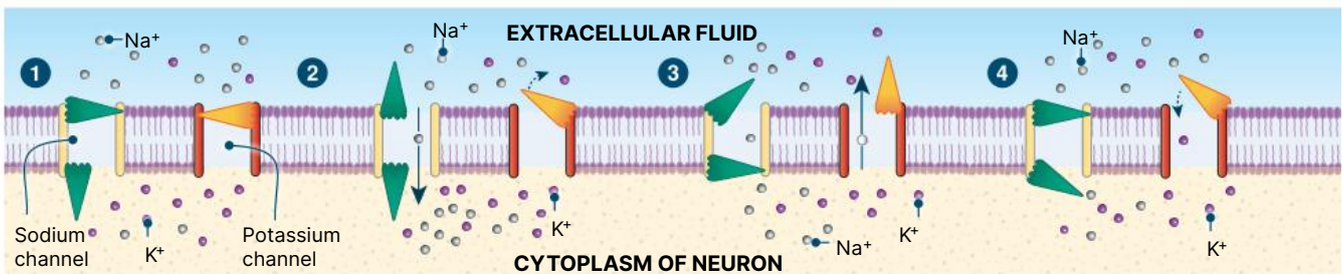


The depolarization in an axon can be shown as a change in membrane potential (in millivolts). A stimulus must be strong enough to reach the **threshold potential** before an action potential is generated. This is the voltage at which the depolarisation of the membrane becomes unstoppable.

- ▶ When at rest, voltage-gated  $\text{Na}^+$  channels are closed (1).
- ▶ Voltage-gated  $\text{Na}^+$  channels open and the membrane depolarises as  $\text{Na}^+$  floods into the cell (2).
- ▶ Voltage gated  $\text{Na}^+$  channels close and voltage-gated  $\text{K}^+$  channels open, allowing  $\text{K}^+$  to move out of the cell (3).
- ▶ A delay in closing voltage gated  $\text{K}^+$  causes **hyperpolarisation**.  $\text{Na}^+/\text{K}^+$  pumps restore the membrane potential (4). During this **refractory period**, the nerve cannot respond, so nerve impulses are discrete.



## Voltage-gated ion channels and the course of an action potential



**Resting state:**

Voltage activated  $\text{Na}^+$  and  $\text{K}^+$  channels are closed. Negative interior is maintained by the  $\text{Na}^+/\text{K}^+$  pump.

**Depolarisation:**

Voltage activated  $\text{Na}^+$  channels open and there is a rapid influx of  $\text{Na}^+$  ions. The interior of the neuron becomes positive relative to the outside.

**Repolarisation:**

Voltage activated  $\text{Na}^+$  channels close and the  $\text{K}^+$  channels open;  $\text{K}^+$  moves out of the cell, restoring the negative charge to the cell interior.

**Returning to resting state:**

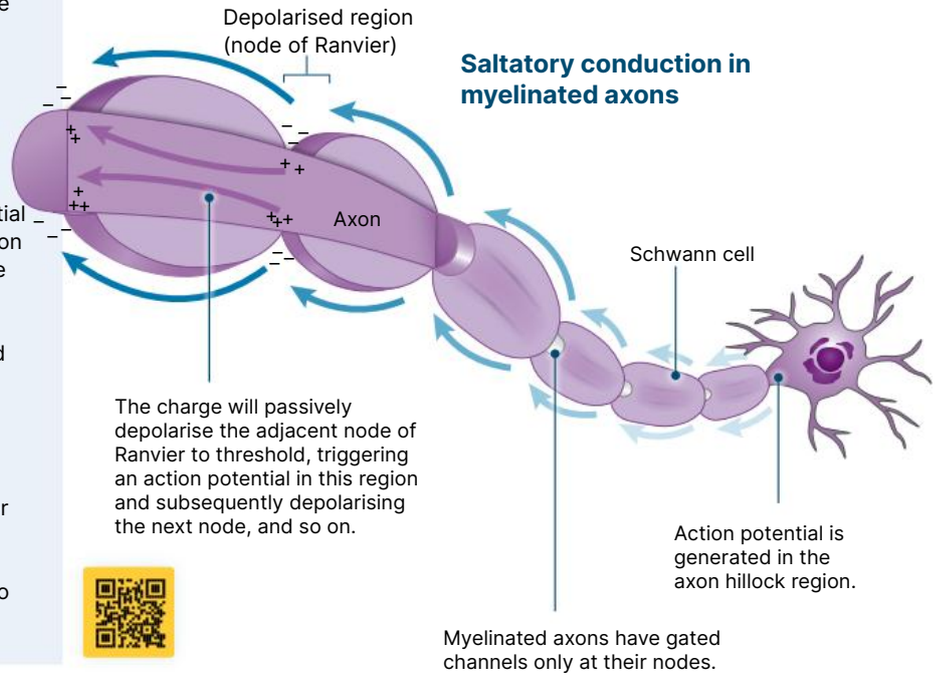
Voltage activated  $\text{Na}^+$  and  $\text{K}^+$  channels close and the  $\text{Na}^+/\text{K}^+$  pump restores the original balance of ions, returning the neuron to its resting state ( $3\text{Na}^+$  out for  $2\text{K}^+$  in).



Axon myelination is a feature of vertebrate nervous systems and it enables them to achieve very rapid speeds of nerve conduction.

In a myelinated neuron, action potentials are generated only at the nodes, which is where the voltage gated channels occur. The axon is insulated so the action potential at one node is sufficient to trigger an action potential in the next node and the impulse 'jumps' along the axon (called saltatory conduction). This contrasts with a non-myelinated neuron in which voltage-gated channels occur along the entire length of the axon.

As well as increasing the speed of conduction, the **myelin** sheath reduces energy expenditure because the area over which depolarisation occurs is less (and therefore also the number of sodium and potassium ions that need to be pumped to restore the resting potential).



1. What is an action potential? \_\_\_\_\_  
\_\_\_\_\_
2. Describe the movement of voltage-gated channels and ions associated with:
  - (a) Depolarisation of the neuron: \_\_\_\_\_
  - (b) Repolarisation of the neuron: \_\_\_\_\_
3. Summarise the sequence of events in a neuron when it receives a stimulus sufficient to reach threshold:
  - (i): \_\_\_\_\_  
\_\_\_\_\_
  - (ii): \_\_\_\_\_  
\_\_\_\_\_
  - (iii): \_\_\_\_\_  
\_\_\_\_\_
  - (iv): \_\_\_\_\_  
\_\_\_\_\_
4. (a) Explain why the nerve impulse in a myelinated neuron jumps along the axon from node to node:  
\_\_\_\_\_  
\_\_\_\_\_
  - (b) How does myelination reduce the energetic costs of impulse conduction?  
\_\_\_\_\_  
\_\_\_\_\_
5. How is the resting potential restored in a neuron after an action potential has passed? \_\_\_\_\_  
\_\_\_\_\_
6. Explain how the refractory period influences the direction in which an impulse will travel: \_\_\_\_\_  
\_\_\_\_\_

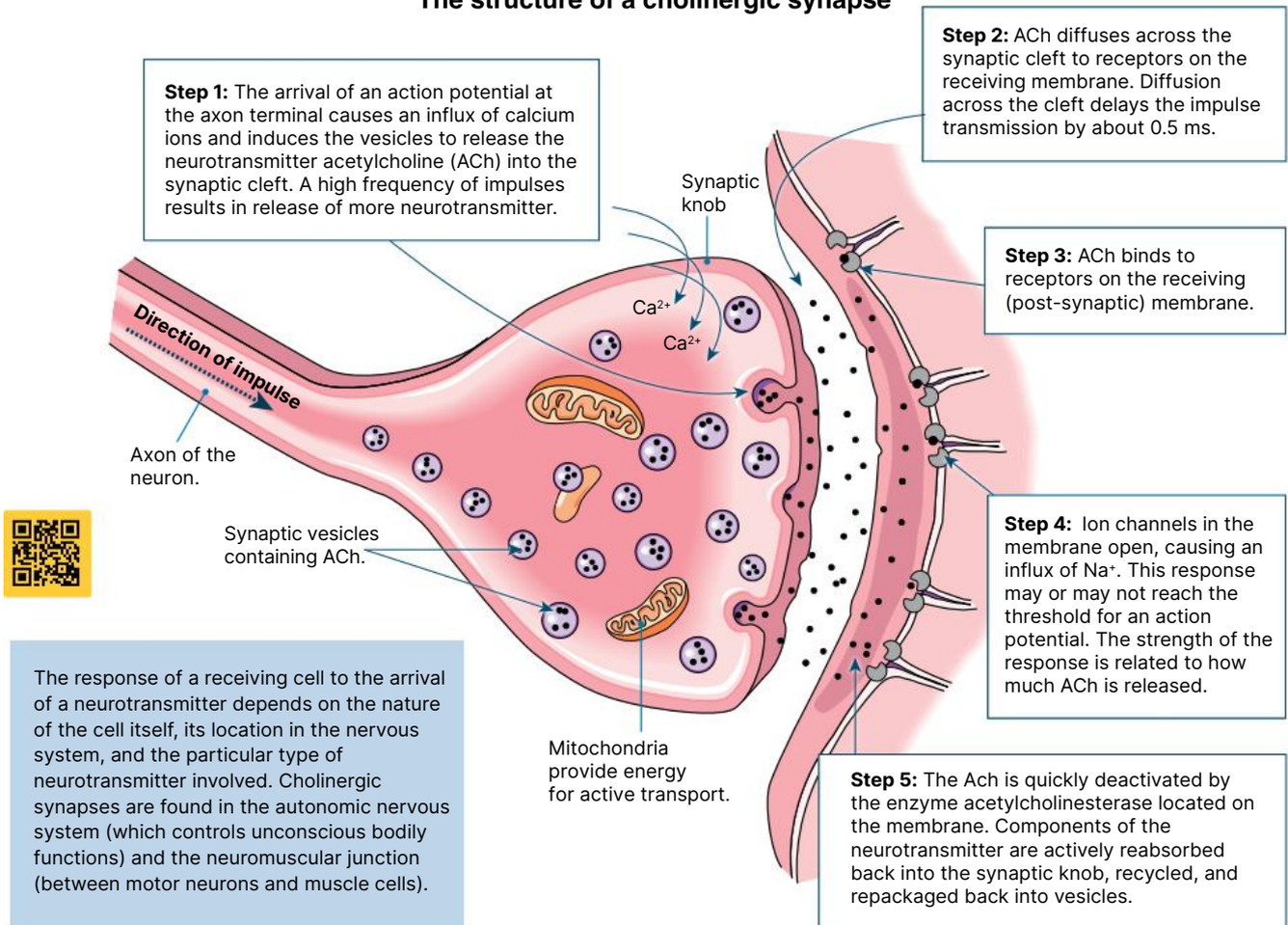
# 139 Chemical Synapses

**Key Idea:** Synapses are junctions between neurons, or between neurons and receptor or effector cells. Nerve impulses are transmitted across synapses.

**Action potentials** are transmitted across junctions called **synapses**. Almost all synapses in vertebrates are chemical synapses, which involve the diffusion of a signal molecule or neurotransmitter from one cell to another. Chemical synapses can occur between two **neurons**, between a receptor cell and a neuron, or between a neuron and an **effector** (e.g. muscle fibre or gland cell). The synapse consists of the axon terminal (synaptic knob), a gap called the synaptic cleft, and

the membrane of the post-synaptic (receiving) cell. Arrival of an action potential at the axon terminal causes release of the neurotransmitter, which diffuses across the cleft and produces an electrical **response** in the post-synaptic cell (an example of signal transduction). Cholinergic synapses are named for the neurotransmitter they release, **acetylcholine (ACh)**. In the example pictured below, ACh results in **depolarisation** (excitation) of the post-synaptic neuron. Unlike electrical synapses, in which transmission can occur in either direction, transmission at chemical synapses is always in one direction (unidirectional).

## The structure of a cholinergic synapse



- (a) What is a synapse? \_\_\_\_\_

\_\_\_\_\_

(b) What defines a cholinergic synapse? \_\_\_\_\_
- What causes the release of neurotransmitter into the synaptic cleft? \_\_\_\_\_

\_\_\_\_\_
- Why is there a brief delay in impulse transmission across the synapse? \_\_\_\_\_

\_\_\_\_\_
- What determines the strength of the response in the receiving cell? \_\_\_\_\_

\_\_\_\_\_

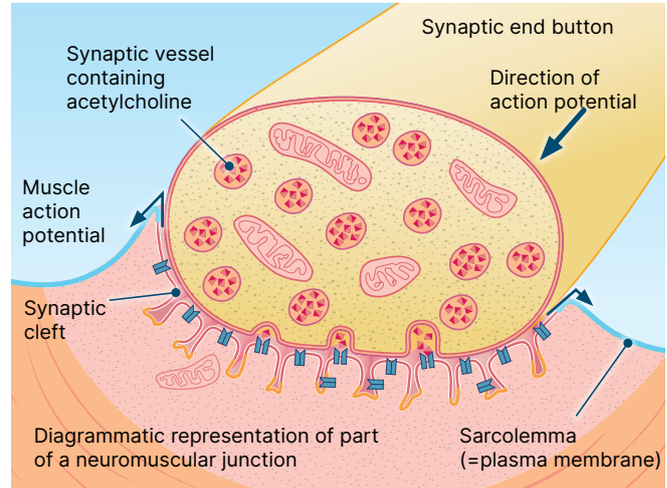
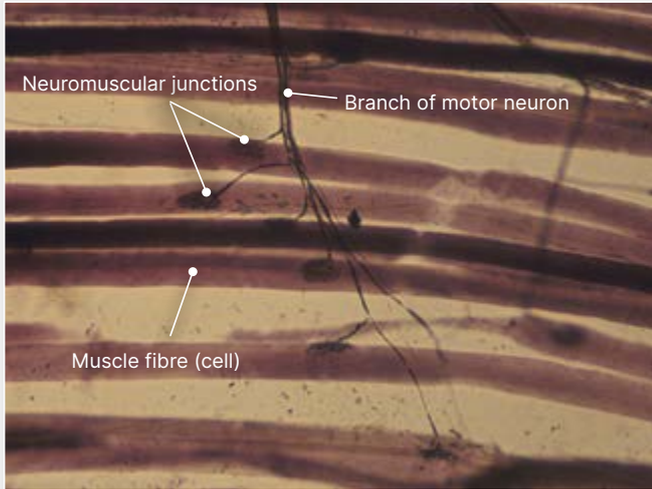
\_\_\_\_\_



### The neuromuscular junction

The neuromuscular junction is a specialised cholinergic synapse between a **motor neuron** and a muscle fibre. Functionally, they operate in the same way as the excitatory cholinergic synapse pictured opposite.

- ▶ Arrival of an action potential at the neuromuscular junction results in depolarisation of the muscle fibre membrane (the sarcolemma) and this results in contraction of the muscle fibre.
- ▶ For a muscle fibre to contract, it must receive a **threshold** stimulus in the form of an action potential. Action potentials are carried by motor neurons from the central nervous system to the muscle fibres they supply. The arrival of an action potential at the neuromuscular junction results in release of the neurotransmitter acetylcholine and contraction of the fibre.
- ▶ The response of a single muscle fibre is all-or-none, meaning it contracts maximally or not at all. This differs from the graded response that can occur with transmission between neurons.



Axon terminals of a motor neuron supplying a muscle. Axon branches end on the sarcolemma (plasma membrane) of a muscle fibre at regions called neuromuscular junctions. Each fibre receives a branch of an axon, but one axon may supply many muscle fibres. A motor neuron and all the fibres it innervates is called a motor unit.

When an action potential arrives at the neuromuscular junction on a muscle cell (fibre), it causes release of acetylcholine, which diffuses across the synaptic cleft to stimulate an action potential in the sarcolemma. The action potential travels throughout the muscle fibre causing muscle contraction.

5. What factors determine the response of the post-synaptic cell? \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_
6. (a) How is the neurotransmitter deactivated? \_\_\_\_\_  
 \_\_\_\_\_  
 (b) Why do you think it is important for the neurotransmitter to be deactivated soon after its release?  
 \_\_\_\_\_  
 (c) Why is transmission at chemical synapses unidirectional and what is the significance of this?  
 \_\_\_\_\_  
 \_\_\_\_\_
7. (a) In what way is the neuromuscular junction (above) similar to the cholinergic synapse described opposite?  
 \_\_\_\_\_  
 \_\_\_\_\_  
 (b) In what ways are these two synaptic junctions different? \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_

# 140 Integration at Synapses

**Key Idea:** Synapses play a pivotal role in the ability of the nervous system to respond appropriately to stimulation and to adapt to change by integrating all inputs.

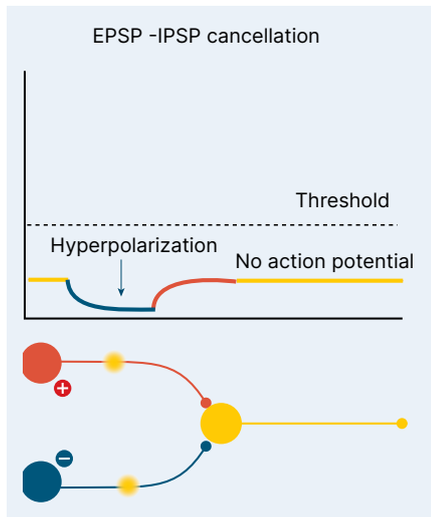
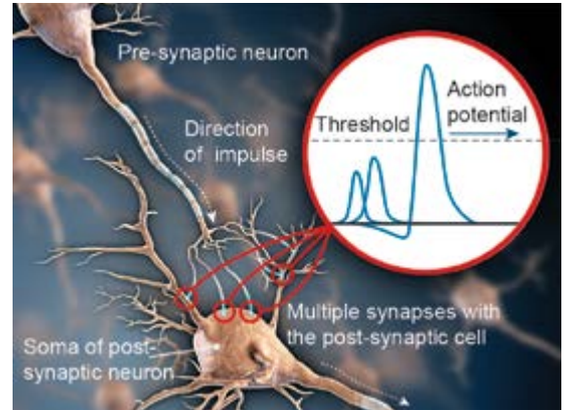
The nature of synaptic transmission in the nervous system allows the integration (interpretation and coordination) of inputs from many sources. These inputs can be excitatory

(causing **depolarisation**) or inhibitory (making an **action potential** less likely). It is the sum of all excitatory and inhibitory inputs that leads to the final **response** in a post-synaptic cell. **Synaptic integration** is behind all the various responses we have to stimuli. It is also the most probable mechanism by which learning and memory are achieved.

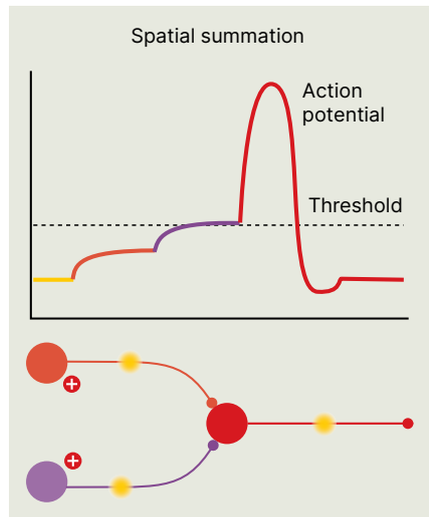
## Summation at synapses

Graded postsynaptic responses may sum to produce an action potential. Impulse transmission across chemical **synapses** has several advantages, despite the delay caused by neurotransmitter diffusion. Chemical synapses transmit impulses in one direction to a precise location and, because they rely on a limited supply of neurotransmitter, they are subject to fatigue (inability to respond to repeated stimulation). This protects the system against overstimulation.

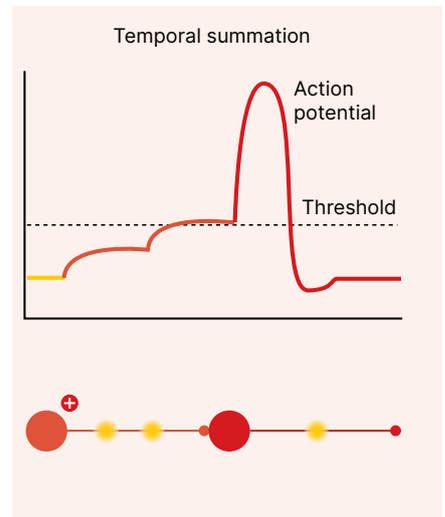
Synapses allow inputs from many sources to be integrated. The response of a post-synaptic cell is often not strong enough on its own to generate an action potential. However, because the strength of the response is related to the amount of neurotransmitter released, subthreshold responses can sum together to produce a response in the post-synaptic cell. This additive effect is called **summation**. Summation can be temporal or spatial (below).



If an inhibitory signal reaches a synapse at the same time as an excitatory signal, the changes to membrane potential cancel out, producing no response.



Impulses from spatially separated axon terminals may arrive simultaneously at different regions of the same post-synaptic **neuron**. The responses from the different places sum to produce an action potential.



Several impulses may arrive at the synapse in quick succession from a single axon. The individual responses sum to reach **threshold** and produce an action potential in the post-synaptic neuron.

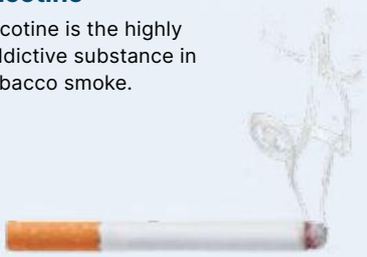
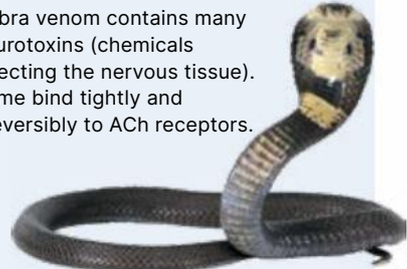
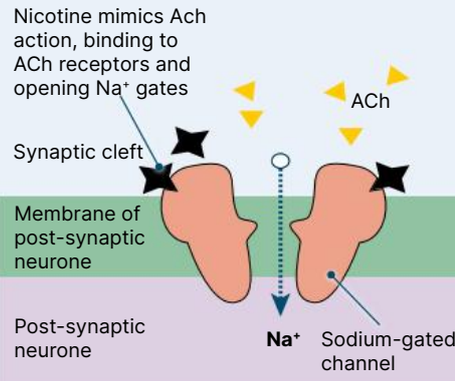
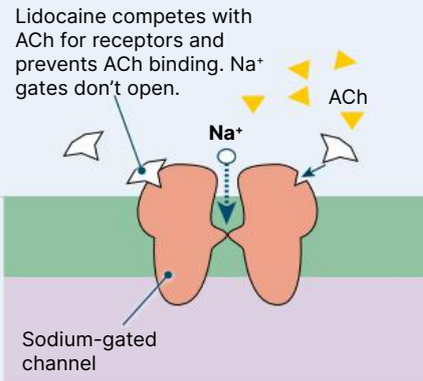
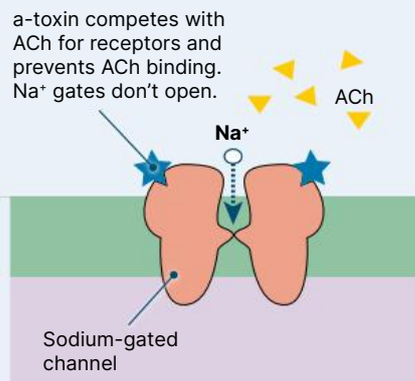
1. Explain the purpose of nervous system integration: \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_
1. (a) Explain what is meant by summation: \_\_\_\_\_  
 \_\_\_\_\_  
 (b) In simple terms, distinguish between temporal and spatial summation: \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_
2. How does hyperpolarisation reduce the chance of an action potential occurring? \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_

# 141 Drugs at Synapses

**Key Idea:** Drugs may increase or decrease the effect of neurotransmitters at synapses.

Drugs may act at **synapses** either mimicking or blocking the usual effect of a neurotransmitter (whether it be excitatory or inhibitory). Drugs that increase the usual

effect of a neurotransmitter are called agonists while those that decrease their effect are called antagonists. Many recreational and therapeutic drugs work through their action at synapses, controlling the **response** of the receiving cell to incoming **action potentials**.

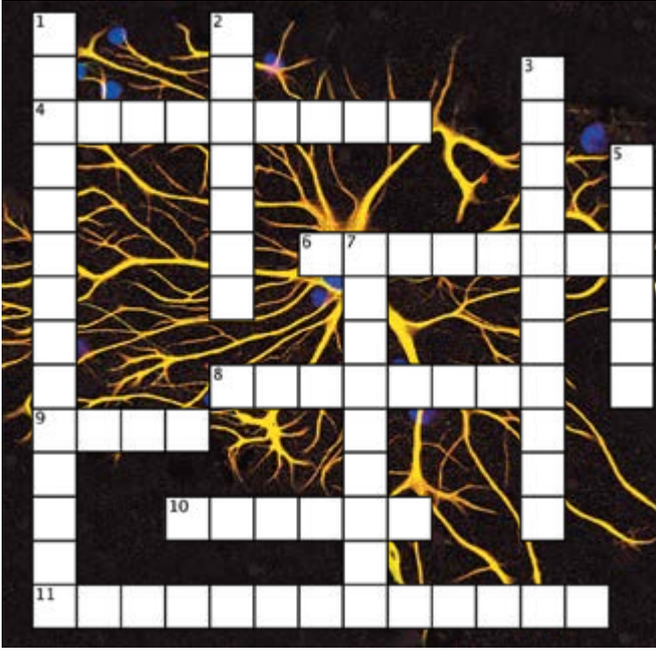
<h3>Nicotine</h3> <p>Nicotine is the highly addictive substance in tobacco smoke.</p> 	<h3>Lidocaine</h3> <p>Lidocaine is a quick-acting local anaesthetic used to block pain during minor surgery or dental work.</p> 	<h3>Cobra venom</h3> <p>Cobra venom contains many neurotoxins (chemicals affecting the nervous tissue). Some bind tightly and irreversibly to ACh receptors.</p> 
<p>Nicotine mimics ACh action, binding to ACh receptors and opening Na<sup>+</sup> gates</p>  <p>Synaptic cleft</p> <p>Membrane of post-synaptic neurone</p> <p>Post-synaptic neurone</p> <p>Na<sup>+</sup> Sodium-gated channel</p>	<p>Lidocaine competes with ACh for receptors and prevents ACh binding. Na<sup>+</sup> gates don't open.</p>  <p>Na<sup>+</sup></p> <p>Sodium-gated channel</p>	<p>a-toxin competes with ACh for receptors and prevents ACh binding. Na<sup>+</sup> gates don't open.</p>  <p>Na<sup>+</sup></p> <p>Sodium-gated channel</p>
<p><b>Effect:</b> Agonistic</p> <p><b>Result:</b> Action potential generation</p> <p>Nicotine acts as an agonist at nicotinic synapses (autonomic ganglia and the motor end plate). It binds to and activates ACh receptors on the postsynaptic membrane (e.g. of a muscle cell). This opens sodium gates, leading to a sodium influx and membrane depolarisation.</p>	<p><b>Effect:</b> Antagonistic</p> <p><b>Result:</b> Sensory inhibition</p> <p>Lidocaine binds to the ACh receptors on <b>sensory neurons</b> and prevents ACh binding. No depolarisation occurs, so no action potential is generated on the post-synaptic neuron. Pain signals are not generated.</p>	<p><b>Effect:</b> Antagonistic</p> <p><b>Result:</b> Muscular paralysis</p> <p>Toxins in cobra venom bind to ACh receptors and prevent ACh binding to receptors on the plasma membrane of muscle cells. As a result, sodium channels remain closed and no action potentials are produced. They can cause muscular paralysis, respiratory failure, and death.</p>

1. Explain the difference between an agonistic and antagonistic drug: \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_
2. Nicotine and cobra venom both bind to acetylcholine receptors. Explain why their effects are different:  
 \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_
3. Explain why lidocaine is used as a local anaesthetic: \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_



# 142 Did You Get It?

1. Complete the crossword below:



GerryShaw CC 3.0

**Across**

- 4. A self propagating nerve impulse is called an action potential.
- 6. Extension of the nerve cell body specialised to receive stimuli.
- 8. A specialised cell that detects stimuli and responds by producing a nerve impulse.
- 9. Long extension of the nerve cell which transmits the nerve impulse to another cell.
- 10. A cell specialised to transmit electrical impulses.
- 11. An organ system made up of a network of specialised cells or neurons, which coordinates responses and transmits signals between parts of the body (2 words).

**Down**

- 1. A temporary change in membrane potential caused by influx of sodium ions.
- 2. The gap between neighbouring neurons or between a neuron and an effector.
- 3. These synapses release acetylcholine.
- 5. This lipid-rich substance surrounds and insulates the axons of nerves in the peripheral nervous system.
- 7. Motor nerves carry impulses from the central nervous system to these.

2. (a) Label the components of this neuron (below right) using the following word list: *cell body, axon, dendrites, node of Ranvier*.

(b) Is this neuron myelinated or unmyelinated? \_\_\_\_\_

(c) Explain your answer:

\_\_\_\_\_

\_\_\_\_\_

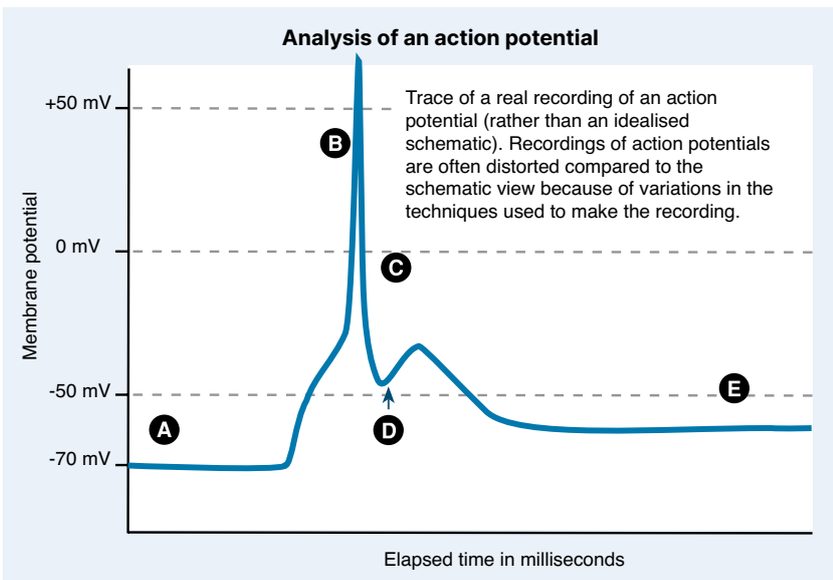
\_\_\_\_\_

(d) In what form do electrical signals travel in this cell?

\_\_\_\_\_



3. (a) The graph below shows a recording of the changes in membrane potential in an axon during transmission of an action potential. Match each stage (A-E) to the correct summary provided below.



- Membrane depolarisation (due to rapid Na<sup>+</sup> entry across the axon membrane).
- Hyperpolarisation (an overshoot caused by the delay in closing of the K<sup>+</sup> channels).
- Return to resting potential after the stimulus has passed.
- Repolarisation as the Na<sup>+</sup> channels close and slower K<sup>+</sup> channels begin to open.
- The membrane's resting potential.

(b) What is the resting potential of the axon? \_\_\_\_\_

(c) What is the maximum voltage reached by the action potential? \_\_\_\_\_



# Hormonal Homeostatic Controls

## Key Terms

- downregulation
- extracellular receptor
- first messenger
- hormone
- hydrophilic signal
- hydrophobic signal
- intracellular receptor
- pheromone
- second messenger
- signal molecule
- signal transduction
- target cell
- transcription factor
- upregulation

## Key Concepts

- ▶ Cells use signals (chemical messengers) to communicate and to respond to changes in their environment.
- ▶ Hormones are chemical signalling molecules produced in one part of the body that may bring about a change in another part of the body.

## Hormones are signalling molecules

### Activity Number

- |     |   |          |
|-----|---|----------|
| □ 1 | Explain what is meant by a signal molecule (or ligand) and, in a general way, explain the effect of signal molecules on target cells. Describe how the transmission of nerve impulses at synapses, immune responses, and hormonal regulation all involve cell signalling. | 143      |
| □ 2 | Explain how a cell's sensitivity to a specific hormone is directly related to the number of receptors it displays for that hormone. Interpret diagrams of upregulation and downregulation and predict the effects of each.  | 143      |
| □ 3 | Identify types of signalling molecules and their roles in regulating the development, behaviour, and physiology of organisms.   | 144      |
| □ 4 | Explain the function of hormones and describe how hormones are produced and distributed to target cells. Explain why only target cells are affected by a specific hormone whereas other cells are unaffected  | 143, 145 |
| □ 5 | Describe, using examples, how the stimulus for release of a hormone may be a substance in the blood (humoral), a nerve impulse (neural), or another hormone (hormonal).   | 145      |
| □ 6 | <b>SHE:</b> Using an example, describe the role of negative feedback in regulating the release of hormones. Using the example of growth hormone release and regulation, explain how some hormones work antagonistically to control a homeostatic process.                 | 149      |
| □ 7 | <b>SI:</b> Explain how hormones are used in the dairy industry to increase milk production and reduce costs. Discuss the risks and ethical concerns with this practice.   | 150      |

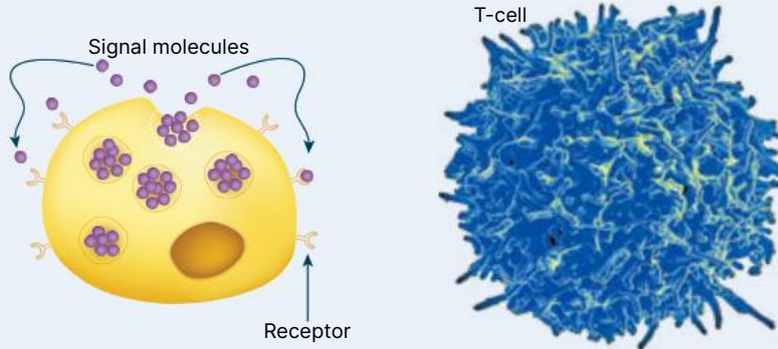
## Signal transduction

- |      |   |         |
|------|---|---------|
| □ 8  | Describe the stimulus-response model with respect to cell signalling to include reception, transduction, and cellular response.   | 146     |
| □ 9  | Distinguish between signal transduction involving hydrophilic signals (e.g. adrenaline) and hydrophobic signals (e.g. steroids such as cortisol). Include reference to differences in how the signal molecule is received by receptors and how transduction is initiated. | 147     |
| □ 10 | For hydrophilic signal molecules, recognise the role of protein-coupled receptors, second messengers, and phosphorylation cascades in producing the cellular response (names of molecules not required).  | 147-148 |
| □ 11 | For hydrophobic signal molecules, recognise the role of nuclear receptors (transcription factors) which are activated when the signal molecule binds.   | 147     |

# 143 Types of Cell Signalling

**Key Idea:** Cells use signals (chemical messengers) to communicate and to respond to changes in their environment. In order to communicate and respond to changes in their environment, cells must be able to send, receive, and process signals. Chemical signals are called **signal molecules** or ligands. In order for a signal to have an effect on a cell it must be able to bind to the cell and bring about a response.

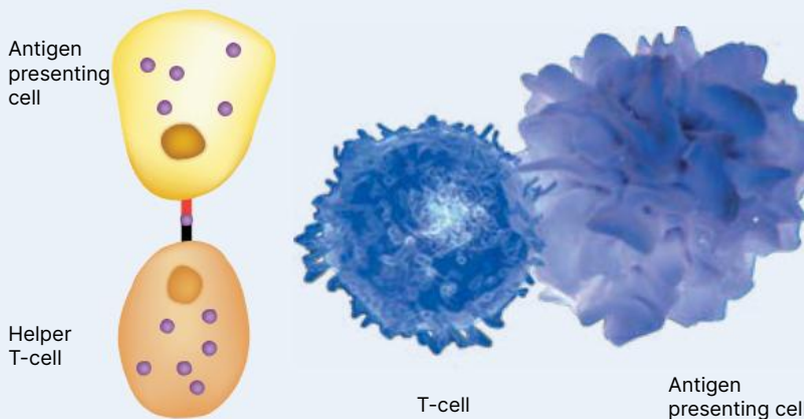
Cells with the receptors to bind a particular signal molecule are called **target cells** for that signal. If a cell does not have the specific receptor, then it is unaffected by the chemical signal. Cells can alter their sensitivity to a chemical signal by altering the number of receptors on the cell surface. Chemical signals can be classified based on how far they travel to cause an effect. Some act locally, others act over long distances.



## Autocrine signalling

Cells can produce and react to their own signals. This type of signalling is important during growth and development and in the functioning of the immune system.

**Example:** In vertebrates, the presence of a foreign antibody causes T-cells to produce a growth factor to stimulate their own production. The increased number of T-cells helps to fight the infection.

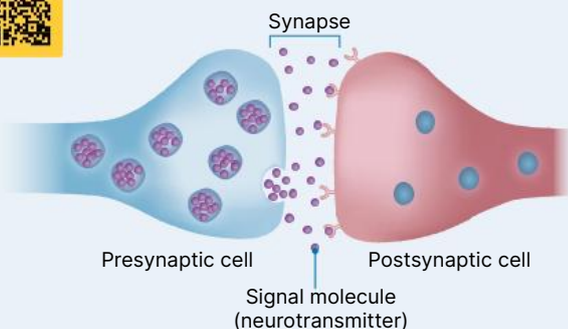


## Cell-to-cell communication

Cell-to-cell communication involves cells interacting directly with one another. There are two forms: 1) communication via special channels between adjacent cells and 2) two cells bind to and communicate with each other because they have complementary proteins on their surfaces.

**Example:** Plasmodesmata are microscopic channels that run through the cell wall of adjacent plant cells. Signal molecules can pass through to the next cell.

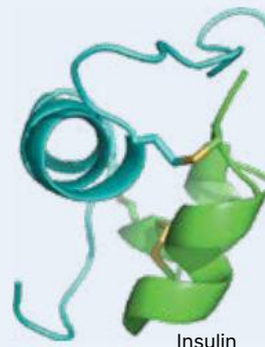
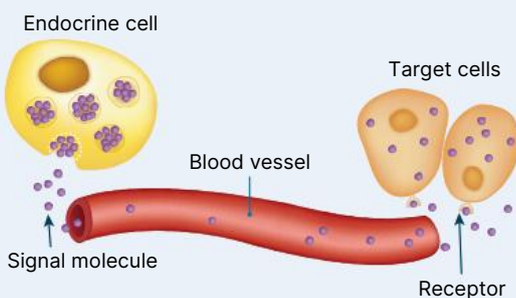
**Example:** In the immune system, antigen presenting cells present antigens to helper T-cells for destruction.



## Signalling by local regulators

Some cell signalling occurs between cells that are close together. The signal molecule binds to receptors on a nearby cell causing a response. Both neurotransmitters and cytokines (small molecules produced by a range of different cells) are involved in this type of local regulation.

**Example:** Neurotransmitters released from a nerve cell travel across the synapse (gap) to another cell to cause a response.



## Endocrine signalling

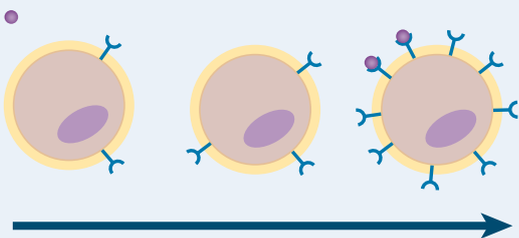
A signal is carried in the bloodstream to target cells, often some distance away. Endocrine signalling may involve **hormones** (released from the cells of endocrine glands) or cytokines as the signalling molecule although cytokines are also important in local regulation and circulate in more variable concentrations than hormones.

**Example:** Insulin from the pancreas stimulates the cellular uptake of glucose.



## Upregulation and downregulation

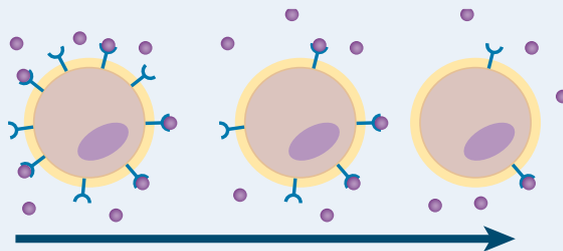
A cell's response to a chemical signal, such as a hormone, not only depends on the presence of the correct receptor, but also on the number of receptors present. The more receptors a cell has for a specific signalling molecule, the greater the cellular response. The increase in a cell's response to a stimulus is called **upregulation**, whereas a reduced response to a stimulus is called **downregulation**.



Upregulation over time

**Upregulation** increases a cell's sensitivity to a specific hormone. It can occur by production of more receptors, or by decreasing the rate at which existing receptors are broken down. Upregulation generally occurs when the concentration of a signal molecule is very low. An increased number of receptors increases the chances of interacting with the signal to bring about a response.

**Example:** During the last trimester of pregnancy there is an increase in uterine oxytocin receptors. Binding of oxytocin induces uterine contraction during labour helping to expel the fetus.



Downregulation over time

**Downregulation** decreases a cell's sensitivity to a signal molecule. Receptor numbers are decreased by reducing the production of receptor proteins or increasing the rate at which existing receptors are broken down. This reduces the receptors available to interact with the signal molecule. Downregulation protects against receptor over-stimulation. It generally occurs when the levels of a signal molecule are very high or when there has been long term exposure to a particular signal molecule.

**Example:** Prolonged high blood glucose decreases insulin production in people with type 2 diabetes (insulin resistance).

1. Explain the purpose of cell signalling: \_\_\_\_\_

---



---



---

2. Identify the components shared by all types of communication involving chemical signalling: \_\_\_\_\_

---



---



---

3. (a) How can a cell alter its response to a signal molecule? \_\_\_\_\_

---



---



---

(b) Use examples to contrast upregulation and downregulation and explain why these mechanisms are important: \_\_\_\_\_

---



---



---



---



---



---



---



---

# 144 Signalling Molecules

**Key Idea:** Signalling molecules are widespread in nature. The effect they have is highly varied and they may act over short or long distances to cause a response.

Signalling molecules bind to specific receptors to cause a response in a **target cell**. There are a huge number of different **signal molecules**, and each has a specific effect.

Some (such as **hormones**) tend to be slow acting and long lasting, while others (such as neurotransmitters) take effect rapidly, but the effect is short lived as the chemical is quickly broken down. Most signals affect cells within the organism, but **pheromones** are secreted into the environment and may travel over long distances to influence members of the same species.



Gibberellins break dormancy in seeds

Plant hormones (phytohormones) have important roles in plant growth and development (e.g. stem elongation, breaking dormancy, and fruit fall). Plant hormones are transported around the plant by the plant's vascular tissue. They often work together and the response varies depending on the relative concentrations of each.



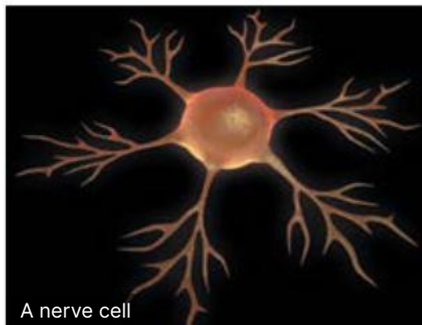
Hormones help animals prepare for and adjust to seasonal changes

In mammals, hormones are secreted by endocrine glands (e.g. the pituitary) and carried in the blood to target cells. Hormones are very potent, and effective at low concentrations. Hormonal responses tend to be slow (because it takes time for the signal to reach its target) and generally long lasting because they induce metabolic changes.



Lymphocytes produce cytokines

Cytokines are a large group of peptides and small proteins involved in coordinating the response of cells in the immune system both within their immediate vicinity or over large distances. Cytokines are produced by a wide range of cells, including immune cells and endothelial cells. They include interferons, interleukins, and tumour necrosis factors.



A nerve cell

Neurotransmitters are chemicals that carry signals between nerve cells or between a nerve cell and another type of cell such as a muscle or gland. Neurotransmitters act on the cell immediately next to it. They are released into a synapse (gap between the cells) and bind to receptors on the receiving cell. The response is rapid and short lived.



A bee swarm

Pheromones are chemical signals released into the external environment. They are widely used by many animals, especially social insects and mammals. Pheromones have different purposes (e.g. aggression, aggregation, reproduction, territoriality) but all act to generate a specific response in members of the same species (conspecifics). A gland at the base on the honeybee abdomen secretes Queen Mandibular Pheromone, which attracts worker bees to swarm. This behaviour is important when establishing a new hive. In mammals, pheromones are used to signal sexual receptivity and attract mates. Special receptors in the nasal cavity detect the pheromones. Mammals often curl the upper lip (flehmen) to expose the receptors.



Stallion exposing receptors

1. Compare how hormones are transported in plants and animals: \_\_\_\_\_  
\_\_\_\_\_
2. (a) Identify a type of long lasting chemical signal: \_\_\_\_\_  
(b) Identify a type of short lasting chemical signal: \_\_\_\_\_
3. What are the benefits of using both short and long acting signals when coordinating physiology and behaviour?  
\_\_\_\_\_  
\_\_\_\_\_  
\_\_\_\_\_
4. How do pheromones differ from the other types of chemical signals? \_\_\_\_\_  
\_\_\_\_\_

# How Hormones Work

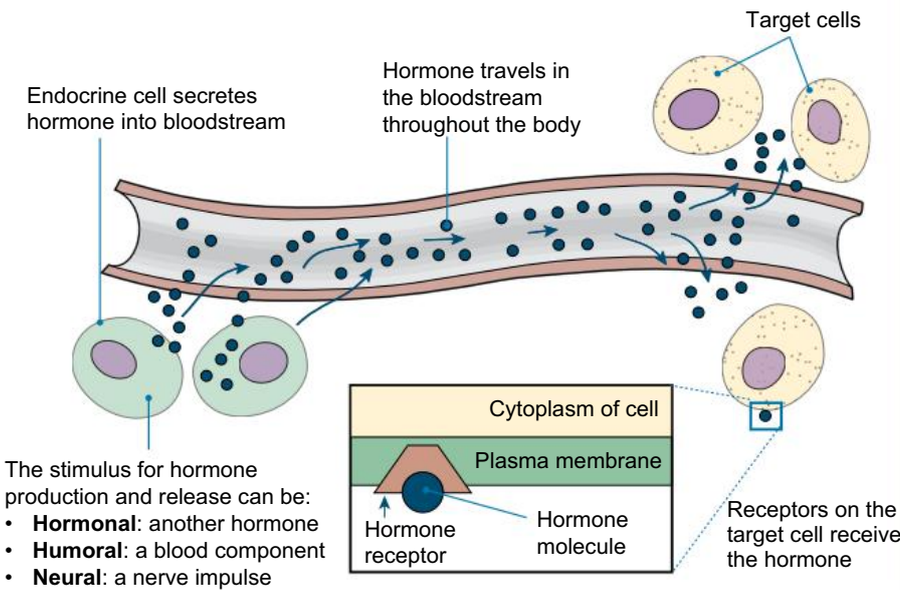
**Key Idea:** The endocrine system regulates physiological processes by releasing blood-borne chemical messengers called hormones, which interact with target cells.

Endocrine signalling has an important role in maintaining homeostasis. The endocrine system is made up of endocrine cells (organised into endocrine glands) and the **hormones**

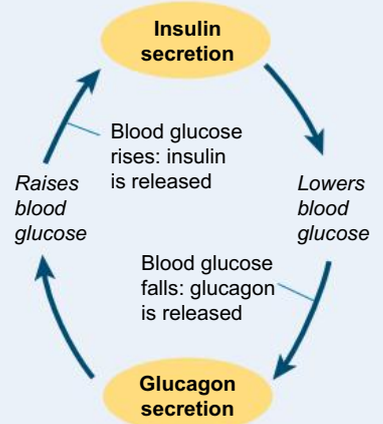
they produce. Hormones are potent chemical regulators. They are produced in very small quantities but can exert a very large effect on metabolism. Endocrine glands secrete hormones directly into the bloodstream rather than through a duct or tube. The basis of hormonal regulation through negative feedback is described below.

## How hormones work

Endocrine cells produce hormones and secrete them into the bloodstream where they are distributed throughout the body. Although hormones are sent throughout the body, they affect only specific **target cells**. These target cells have receptors on the plasma membrane which recognise and bind the hormone (see inset, below). The binding of hormone and receptor triggers the response in the target cell. Cells are unresponsive to a hormone if they do not have the appropriate receptors.



## Antagonistic hormones



The effects of one hormone are often counteracted by an opposing hormone. Feedback mechanisms adjust the balance of the two hormones to maintain a physiological function.

**Example:** insulin acts to decrease blood glucose and glucagon acts to raise it.

- (a) What is a hormone? \_\_\_\_\_

\_\_\_\_\_

(b) Why can a hormone only influence specific target cells even though all cells may be exposed to the hormone?

\_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_
- (a) Describe how antagonistic hormones act to maintain homeostasis: \_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_

(b) Is the stimulus for the release of insulin humoral, hormonal, or neural? \_\_\_\_\_

(c) Use the example of blood glucose to explain the role of feedback mechanisms in adjusting hormone levels:

\_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_



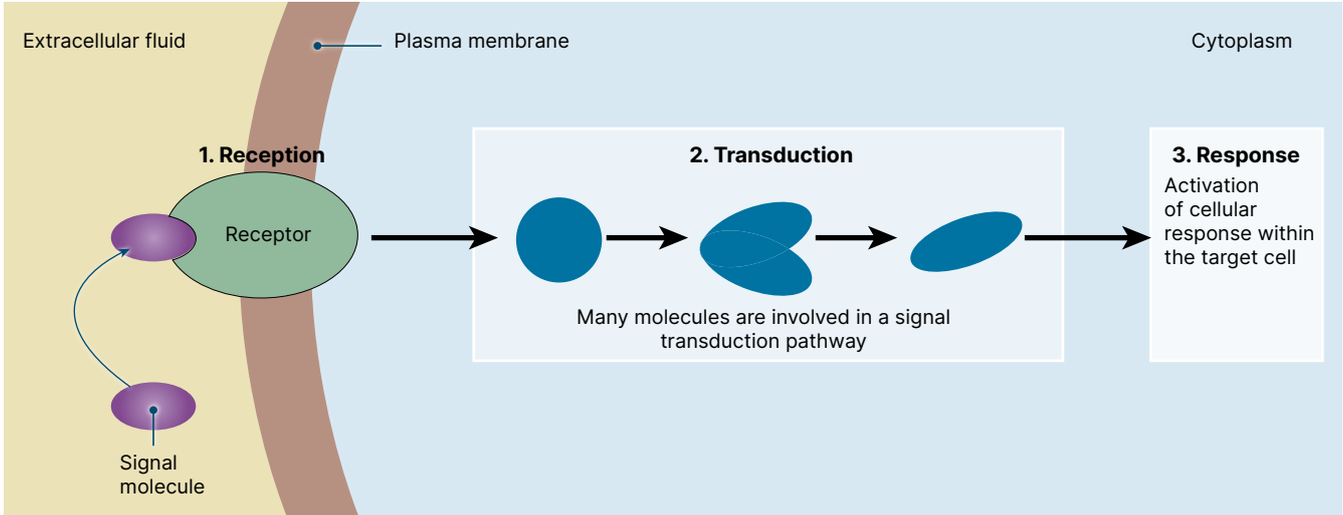
# What is Signal Transduction?

**Key Idea:** Signal transduction is the conversion of an external signal to a functional change within the cell through a series of biochemical reactions.

**Signal transduction** is the process by which molecular signals are transmitted from outside the cell to inside, bringing about a cellular response. The transduction involves an external **signal molecule** binding to a receptor and triggering a series

of biochemical reactions, which lead to a specific cellular response. The series of biochemical reactions is often called a cascade and usually involves phosphorylation (charging) of a number of molecules in a sequence. The type of response varies and may include a change in metabolism (activating a pathway), gene expression (to produce a specific protein), or membrane permeability (to allow entry of specific molecules).

## An overview of signal transduction

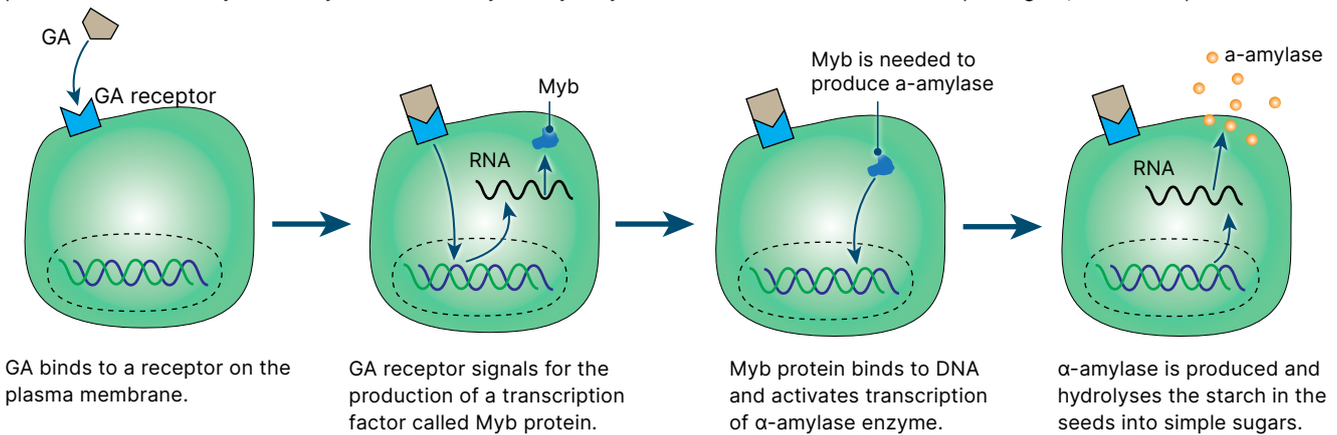


Signal transduction can be broken into three main steps:

- ▶ **Reception:** An extracellular signal molecule binds to its receptor on a **target cell**.
- ▶ **Transduction:** The activated receptor triggers a chain of biochemical events within the cell. Many different enzymes are involved, and the entire reaction is often called a signalling cascade.
- ▶ **Response:** The signal cascade results in a specific cellular response.

## Gibberellic acid activation of $\alpha$ -amylase: An example of a cellular response

In plants, the **hormone** gibberellic acid (GA) is involved in seed germination. GA acts as a signal molecule to stimulate the production of the enzyme  $\alpha$ -amylase. The  $\alpha$ -amylase hydrolyses (breaks down) starch into simple sugars, which the plant can use.



1. Name the three stages of signal transduction and describe what occurs at each stage:

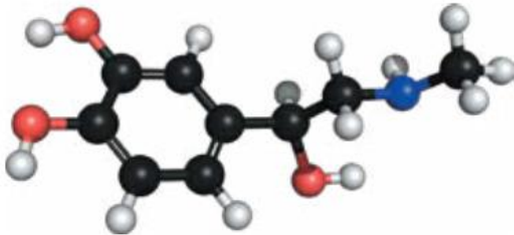
- (a) \_\_\_\_\_
- \_\_\_\_\_
- (b) \_\_\_\_\_
- \_\_\_\_\_
- (c) \_\_\_\_\_
- \_\_\_\_\_

# 147 Types of Signal Transduction

**Key Idea:** The majority of cell signals bind to extracellular receptors to exert their effect. However some cell signals are able to pass through the plasma membrane and bind directly to intracellular receptors within the cell to exert their effect. Cell receptors fall into two broad classes. **Extracellular receptors** bind **signal molecules** outside of the cell. The

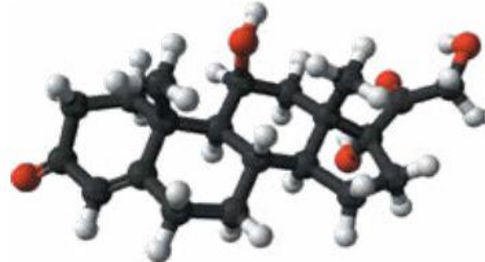
signal molecule does not have to pass across the plasma membrane to cause a cellular response. Most cell receptors are extracellular receptors. **Intracellular receptors** bind signal molecules that have passed into the cell directly across the plasma membrane. Intracellular receptors may be located in the cytoplasm or on the nucleus.

## Hydrophilic signal molecules are received by extracellular receptors

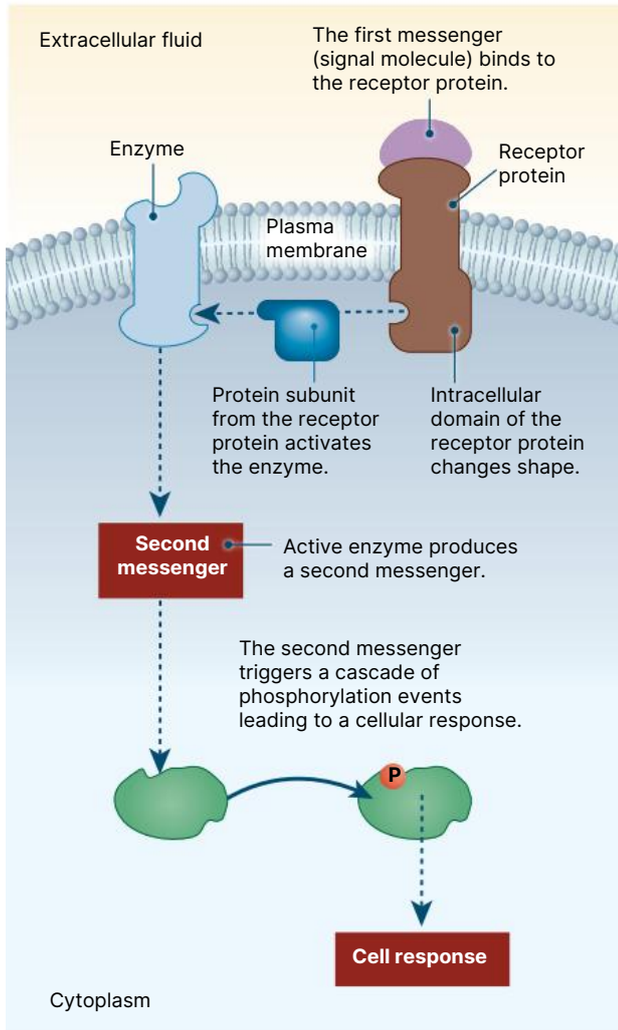


Adrenaline accelerates heart rate and is involved in the fight or flight response

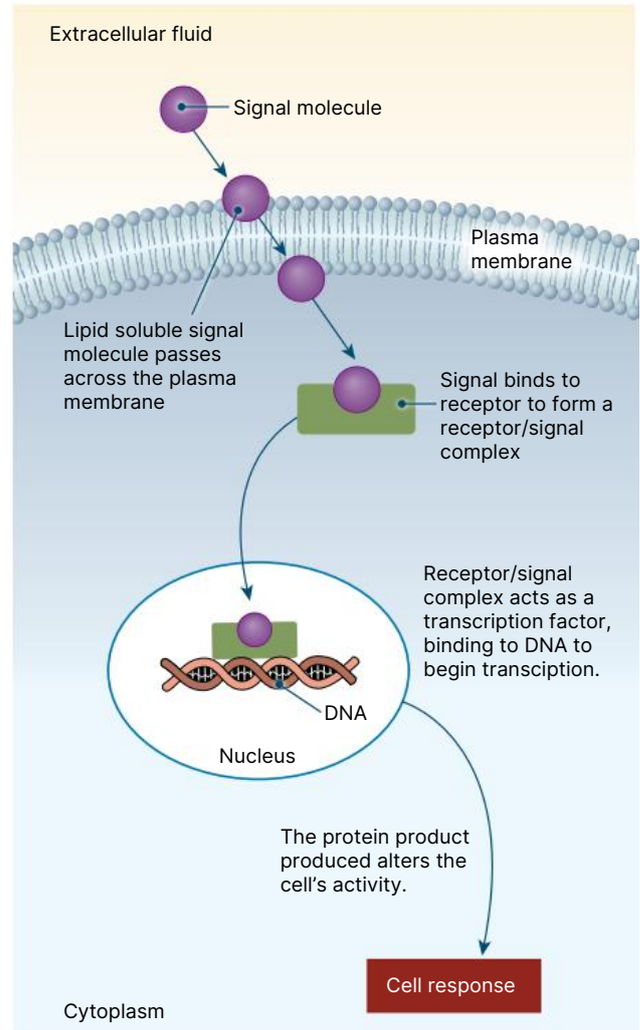
## Hydrophobic signal molecules are received by intracellular receptors



Cortisol is involved in glucose metabolism and response to stress



**Hydrophilic signal molecules** are water soluble and so cannot cross the plasma membrane. They exert their effect via an extracellular receptor. **Hydrophilic signals** include water soluble **hormones** such as adrenaline. The signal molecule is the **first messenger**. Binding activates the extracellular receptor, triggering a sequence of biochemical reactions, including activation of a second messenger. As a result, the original signal is amplified, and there is a cellular response.



**Hydrophobic signal molecules** diffuse freely across the plasma membrane and into the cytoplasm of **target cells**. In the example, above the signal molecule binds to a receptor in the cytoplasm to form a receptor/signal complex. The complex moves to the cell nucleus where it acts as a **transcription factor** to control the expression of specific genes. Steroid hormones, such as cortisol and sex hormones, are examples of **hydrophobic signal** molecules.



1. Describe the differences between an intracellular receptor and an extracellular receptor:

---



---



---

2. What must a signal molecule do in order to activate a receptor? \_\_\_\_\_

---

3. In terms of their ability to cross the plasma membrane, describe the difference between a hydrophobic signal molecule and a hydrophilic signal molecule:

---



---



---

4. (a) Outline the process when signal transduction occurs via an extracellular receptor: \_\_\_\_\_

---



---



---



---

(b) Describe the differences between a first messenger and a second messenger: \_\_\_\_\_

---



---



---

5. Outline the process when signal transduction occurs via an intracellular receptor: \_\_\_\_\_

---



---



---



---

6. The diagram on the right represents a cell signalling process.

(a) Does this diagram represent an extracellular or intracellular signalling process? Explain your answer:

---



---



---

(b) What type of receptor is B? \_\_\_\_\_

(c) What does A represent? \_\_\_\_\_

(d) Would A be hydrophobic or hydrophilic? Explain your answer:

---



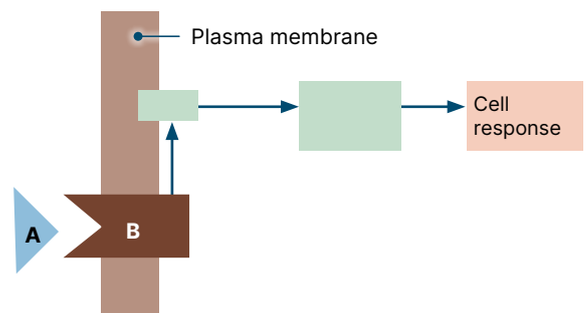
---



---

(e) Another receptor molecule C sits beside B and has a cup shaped receptor. Would molecule A cause a response from this receptor?

---

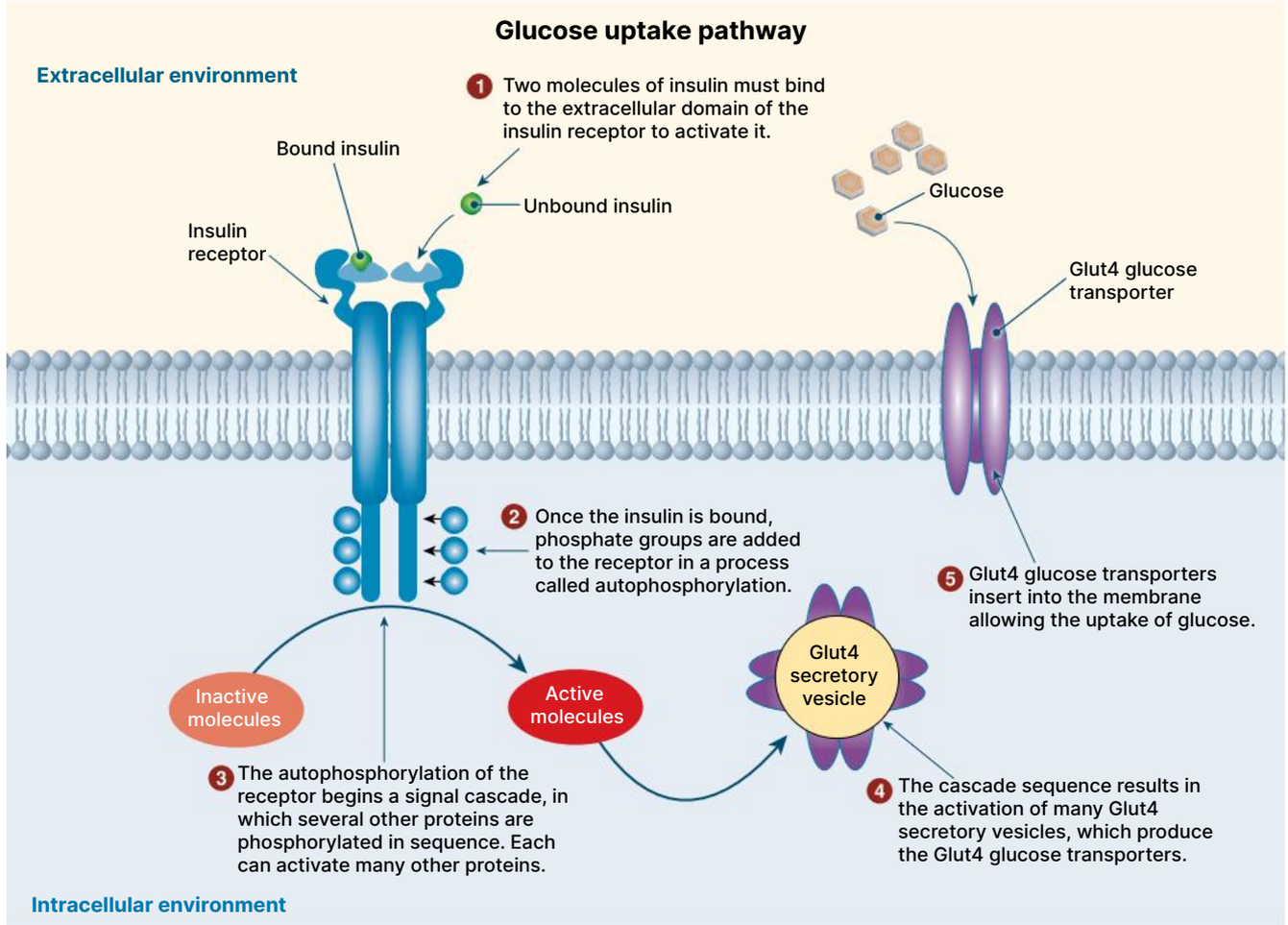


# Action of Insulin

**Key Idea:** Activation of the insulin receptor by insulin causes a signal cascade that results in cellular glucose uptake.

Insulin is a peptide **hormone** secreted by the pancreas. It is involved in regulating blood glucose levels by promoting the uptake of glucose by cells. Malfunctions in the signalling pathways for insulin production and reception have serious physiological consequences (including death) so these are

tightly regulated and under strong selection (evolutionary) pressure. If insulin is lacking or cells fail to respond to it, blood glucose remains elevated while the body's cells themselves are starved of fuel. Insulin circulates in the blood where it binds to protein kinase receptors on the surface of cells and triggers a signal cascade that results in activation of the membrane transporters that bring glucose into the cell.



- (a) What type of signalling does this example represent? autocrine / local regulation / endocrine: \_\_\_\_\_

(b) Explain why you chose this answer: \_\_\_\_\_

\_\_\_\_\_
- Why must blood glucose levels be tightly regulated? \_\_\_\_\_

\_\_\_\_\_
- Describe the process by which insulin signalling causes the uptake of glucose into cells:

\_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_
- How does the signal cascade increase the response of the insulin receptor? \_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_



# 149 Hormone Regulation by Negative Feedback

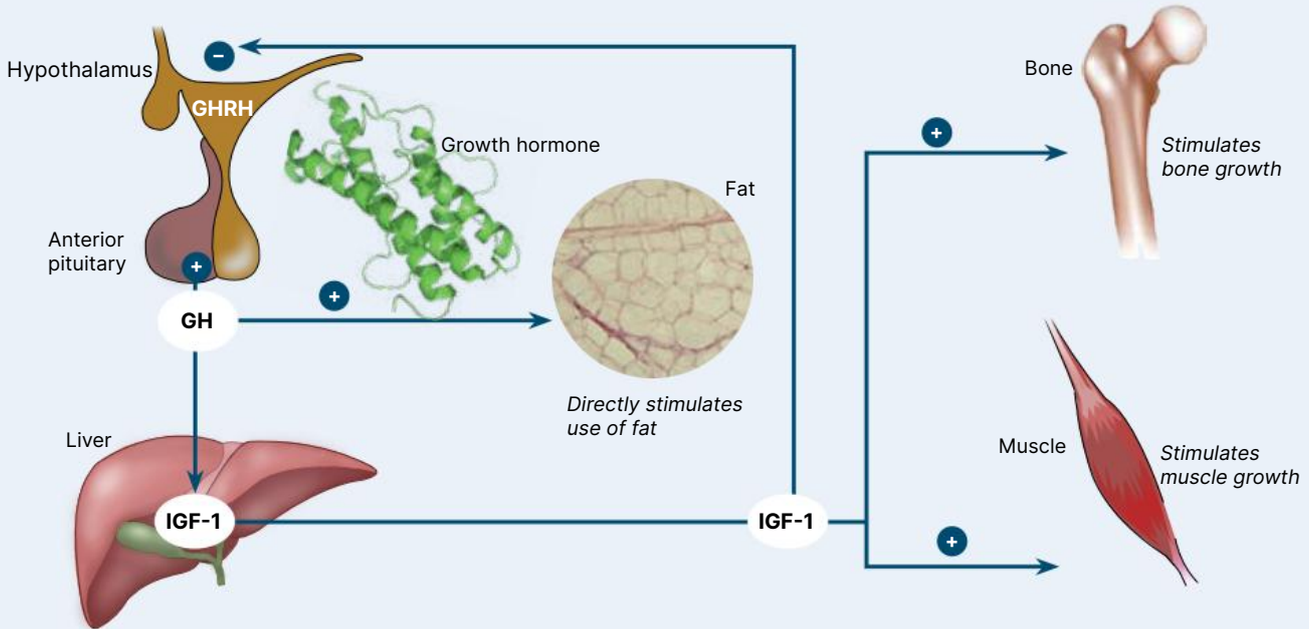
**Key Idea:** Growth hormone is required for normal growth and development. Its levels are tightly regulated through negative feedback mechanisms to prevent growth disorders.

Growth **hormone** (GH) is a peptide hormone that stimulates growth, cell reproduction, and cell regeneration through endocrine signalling. GH is released from the pituitary gland at the base of the brain and circulates in the blood where it binds to two adjacent receptors on liver cells and stimulates

the production of a protein called insulin-like growth factor (IGF-1). GH levels are regulated by negative feedback (a regulatory mechanism in which a stimulus input causes an opposite output in order to maintain a steady state). GH has a critical role in human development and malfunctions in its regulation can result in a number of serious conditions including growth deficiencies (low GH) and abnormal tissue growth resulting in tumours or gigantism (excessive GH).

## The effects and regulation of growth hormone

- ▶ Growth hormone (GH) is released in response to GHRH (growth hormone releasing hormone) from the hypothalamus of the brain. GH acts both directly and indirectly to affect metabolic activities associated with growth.
- ▶ GH directly stimulates metabolism of fat, but its major role is to stimulate the liver and other tissues to secrete IGF-1 (Insulin-like Growth Factor 1) and through this stimulate bone and muscle growth.
- ▶ GH secretion is regulated via negative feedback. High levels of IGF1 suppress GHRH secretion (and therefore GH secretion from the anterior pituitary).



1. Describe the metabolic effects of growth hormone: \_\_\_\_\_

---



---

2. What is the role of negative feedback in regulating the secretion of growth hormone?

---



---



---



---



---

3. In a separate but related control pathway, high levels of IGF-1 also stimulate the release of GHIH, an inhibiting hormone from the hypothalamus. GHIH suppresses secretion of GH. In the space on the right, construct a negative feedback pathway to show how GHIH regulates GH secretion:



# Use of Hormones in the Dairy Industry

**Key Idea:** Recombinant bovine somatotropin (rBST) can be injected into dairy cows to increase milk production. However, there are numerous concerns about its effects on livestock and its use is not permitted in Australia. Cows produce **hormones** naturally, some of which promote growth and influence milk production. In some countries, synthetic (man-made) versions of these hormones are used

by farmers to promote livestock growth or increase milk production in dairy cows. An example of this is recombinant bovine somatotropin (rBST), also called recombinant bovine growth hormone (rBGH). However, there are concerns over the welfare of animals injected with rBST and its use is banned in some countries. Several countries, including Australia, do not allow rBST to be used in dairy cows.

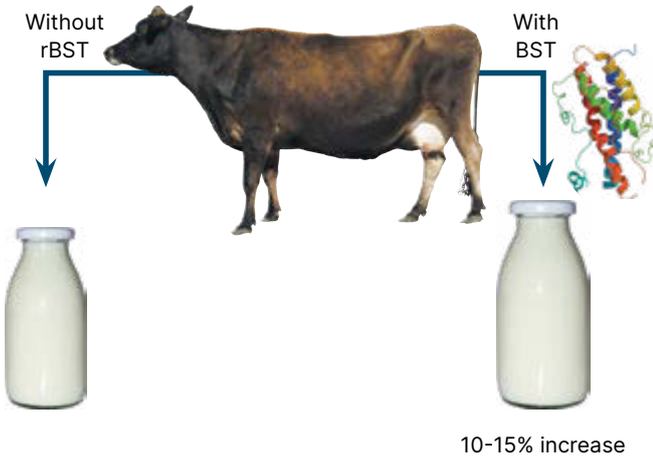
**rBST**

- ▶ Bovine somatotropin (BST) is a hormone produced naturally in cows from the anterior pituitary (a gland at the base of the brain). BST is important in normal growth and development.
- ▶ In the 1930s and 1940s, BST researchers found that BST extracted from the pituitary gland of dead cows and injected into cows increased milk production.
- ▶ In the 1980s, recombinant BST (rBST) was produced using biotechnology methods, allowing large quantities of rBST to be produced in *E.coli* bacteria.
- ▶ The first commercial sales of rBST in the USA occurred in 1993.

**Pros and cons of using rBST**

BST regulates milk production in lactating cows by directing more nutrients to be utilised to make milk. This increases milk production. Usually milk production peaks 60-90 days into lactation and then begins to fall. However, peak production can be extended by injecting cows with rBST. The use of rBST:

- ▶ Increases milk production by 10-15%.
- ▶ Increases the efficiency of milk production.
- ▶ Decreases the environmental impact of dairy farming (through improvements in productivity and efficiency).
- ▶ Reduces the economic cost of dairying.
- ▶ Although some potential health effects in humans have been raised (e.g. increased risk of cancer) as yet there is no evidence linking rBST to negative health effects in humans. The fact that the hormone is destroyed in the human gut is considered to reduce any potential health risks.



Some countries (e.g. the USA) permit the use of rBST in dairy herds. However, its use is banned in several countries (including Australia) because of animal health issues. These include:

- ▶ 24% increase in mastitis (painful inflammation of the mammary gland). Mastitis (right) is treated by antibiotics and could contribute to increasing antibiotic resistance.
- ▶ 40% reduction in fertility.
- ▶ 55% increase in lameness.



Mohammad Golkar cc4.0

1. (a) What effect does BST hormone have on milk production? \_\_\_\_\_

\_\_\_\_\_

(b) Describe how it exerts its effect: \_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_

2. Discuss the advantages and disadvantages of using rBST in the dairy industry and decide if you think its ban in Australia is justified.

\_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_



# Did You Get It?

1. (a) The molecules labelled A-C are signalling molecules. Identify the signal molecule that will bind to the receptor shown:

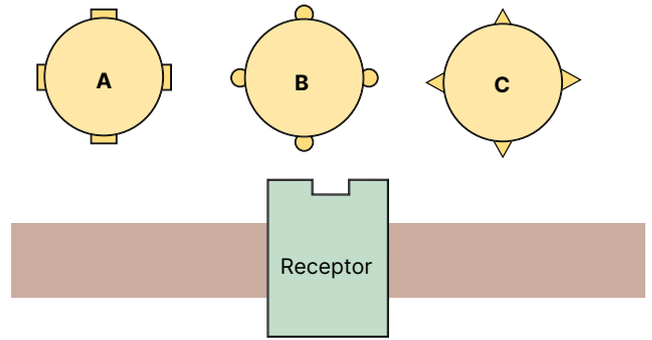
\_\_\_\_\_

(b) What prevents the other two signal molecules from binding to this receptor?

\_\_\_\_\_  
\_\_\_\_\_

(c) Why is it important that not all cells react to every signal molecule?

\_\_\_\_\_  
\_\_\_\_\_



2. The decrease in the expression of an extracellular receptor after its increased activation by a signal molecule is called:

\_\_\_\_\_

3. How does growth hormone (GH) affect metabolic activities associated with growth?

\_\_\_\_\_  
\_\_\_\_\_  
\_\_\_\_\_

4. Choose the correct answer from the choices below:

Hormones are relatively long lived signals, which travel through the blood. This type of signalling is called:

- (a) Autocrine signalling
- (b) Cell-to-cell communication
- (c) Local regulation
- (d) Endocrine signalling

5. Choose the correct answer from the choices below:

Where do intracellular receptors bind signal molecules that have passed into the cell?

- (a) Cytoplasm
- (b) Extracellular fluid
- (c) Nucleus
- (d) Plasma membrane

6. Provide a definition for each of the following terms:

(a) Hormone: \_\_\_\_\_

\_\_\_\_\_

(b) Signal molecule: \_\_\_\_\_

\_\_\_\_\_

(c) Target cell: \_\_\_\_\_

\_\_\_\_\_

(d) Signal transduction: \_\_\_\_\_

\_\_\_\_\_

7. (a) Does hormone signalling produce short-term or long-term effects compared to nervous signalling?

\_\_\_\_\_

(b) Does hormone signalling produce fast or slow responses compared to nervous signalling?

\_\_\_\_\_

# Thermoregulation



## Key Terms

- aestivation
- brown fat
- endotherm
- hibernation
- insulation
- kleptothermy
- panting
- sweating
- thermogenesis
- thermoregulation
- thyroxine
- torpor
- vasoconstriction
- vasodilation

## Key Concepts

- ▶ Animals have different physiological methods for maintaining their internal temperature.
- ▶ Hormones play a role in helping moderate body temperature.

## Thermoregulation

### Activity Number

- |     |  |          |
|-----|--|----------|
| □ 1 | Explain what is meant by thermoregulation. Explain the importance of maintaining body temperature in a homeotherm.   | 152-53   |
| □ 2 | Explain how feedback mechanisms control thermoregulation of the body. Explain how sweating, shivering, vasoconstriction, and vasodilation all help maintain body temperature within acceptable limits. | 153, 158 |
| □ 3 | <b>SI:</b> Describe the structural, behavioural, and physiological methods used by endotherms to both lose or retain body heat as needed.  | 152-156  |

## Thermoregulatory responses

### Science understanding

- |     |   |         |
|-----|---|---------|
| □ 4 | Explain how structural mechanisms directly or indirectly contribute to thermoregulation in endotherms.  | 152-153 |
| □ 5 | Describe how thermoregulatory behaviours in endotherms are related to reducing the energetic costs of maintaining a constant temperature.   | 154     |
| □ 6 | Identify thermoregulatory behaviours in endotherms, including kleptothermy (e.g. huddling), and behaviours that enable animals to offset the high energetic costs of thermoregulation at certain times of the year (hibernation, aestivation, and torpor).  | 154     |
| □ 7 | Using examples, describe and explain physiological mechanisms for thermoregulation. Include autonomic (vasomotor) control of blood flow (vasoconstriction and vasodilation), shivering, evaporative heat loss from body surfaces, countercurrent heat exchangers, and thermogenesis (heat production from metabolism in brown fat). | 155     |
| □ 8 | Explain the role of hormones (thyroid hormones and insulin) in regulating the metabolic generation of heat.   | 156     |
| □ 9 | <b>SHE:</b> Describe how models of heat loss in humans can be used to design clothing to allow for better thermoregulation in different situations.   | 157     |

# 152 Mechanisms for Thermoregulation

**Key Idea:** Endotherms regulate their body temperature to within narrow limits by controlling heat exchanges with the environment and generating heat from metabolism. This is termed thermoregulation.

Heat exchanges with the environment occur via conduction, radiation, and evaporation. To maintain a relatively constant body temperature, **endotherms** must balance heat losses

and gains. **Thermoregulation** is achieved through a variety of mechanisms: structural (physical attributes of the body), behavioural, and physiological (mechanisms at the metabolic level). These mechanisms allow an organism to maintain a body temperature that is optimum for functioning. For endotherms generating their body heat through metabolism, thermoregulation represents a high energy cost.

## Thermoregulation mechanisms to prevent heat loss

The fairy penguin can be found in cold water and surrounding coastal areas around southern regions of Australia.

### Physiological:

Involuntary shivering when cooler core body temperature activates muscles.

**Structural:** No external ear flaps.

**Behavioural:** Feather fluffing to increase insulation layer when on land.

**Structural:** Overlapping feathers form waterproof layer to aid **insulation**.

### Behavioural:

Wings and legs held close to body.

**Structural:** Compact body shape with short legs.



The orca is found in oceans world-wide, including around many coastal areas of Australia.

**Structural:** Heavily insulated surfaces of vascularised fat called blubber (up to 60% of body thickness).



**Physiological:** Countercurrent heat exchangers between venous and arterial blood in the flippers and tail fluke reduce heat loss.

## Thermoregulation mechanisms to increase heat loss

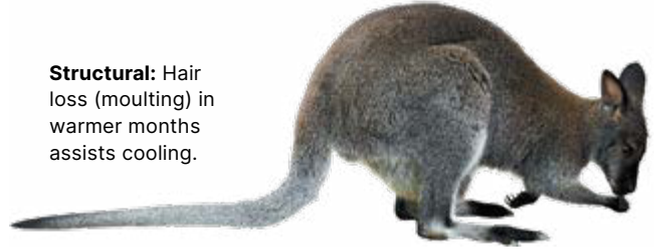
Kangaroos and wallabies live in the hot arid grassland areas of Australia.

### Physiological:

Evaporation from vascular rich mouth and tongue, enabled by **panting**.

**Structural:** Ears with large surface area aid cooling.

**Structural:** Hair loss (moulting) in warmer months assists cooling.



**Behavioural:** Poorly insulated parts of the body are licked to aid evaporative heat loss.

Kangaroos cannot sweat. **Sweating** cools by evaporation. Animals tend to rely on panting or sweating for cooling, not both.

The emu lives in both open dry plains and tropical forest areas of Australia.

### Structural:

Sparse feathers on the neck aid heat loss.

**Physiological:** Lungs function as evaporative coolers, removing heat energy into exhaled water vapour when the birds pant.

### Behavioural:

Shade seeking and swimming during hot days.

**Structural:** Black tipped feathers capture heat while lighter feathers underneath insulate the body. This allows the bird to be active in the hot parts of the day without overheating.

**Behavioural:** Stretches wings to allow air to circulate around body.



1. Endotherms use a number of different mechanisms to thermoregulate, although not every mechanism is present in every species. Determine if the following are structural, behavioural, or physiological mechanisms:

- Seeking shade: \_\_\_\_\_
- Countercurrent heat exchange systems: \_\_\_\_\_
- Presence of fur, feathers or hair: \_\_\_\_\_
- Presence of a blubber layer: \_\_\_\_\_
- Generation of heat through metabolism: \_\_\_\_\_
- Panting: \_\_\_\_\_
- Alteration of circulation pattern: \_\_\_\_\_



SU

SI

# Structural Features for Thermoregulation

**Key Idea:** Structural features for thermoregulation are those involving how the animal is built, such as long or short ears, blubber, or fur.

Structural mechanisms for **thermoregulation** are those that arise from the physical structure of the animal, its size and build, and features of the skin and body coverings. A common

structural feature for thermoregulation in mammals is fur (except in some marine mammals). In birds, this role is taken by feathers. Physiological adaptations usually accompany structural features, such as the ability to moult from a winter to summer coat and the ability to raise and lower the hair or feathers to increase or decrease **insulation**.

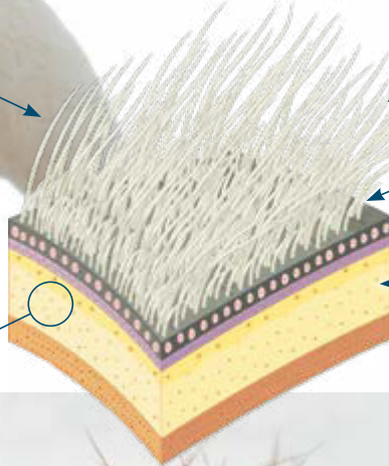
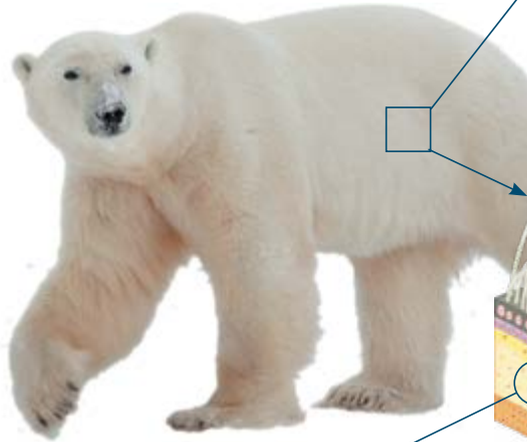
## Keeping warm

Preventing heat loss is important for small animals and those in very cold environments. This includes animals that live in the polar regions and marine animals.

### Fur

Fur (hair) is a mammalian characteristic. It is a very good insulator in terrestrial environments. There is often a double layer of fur, an outer long layer and an inner (and very thick) insulating layer. Oils in the fur help to waterproof it. Both fur and feathers work by trapping air close to the body, warming it and preventing it escaping.

Sea otters are one of the furiest mammals with more than 124,000 hairs per cm<sup>2</sup> in their inner coat.



### Polar bear fur

**Guard hairs:** Long and transparent. Scatter light, enhancing camouflage but also allow some light through to heat the skin.

**Dense underfur:** Traps air and prevents body heat from escaping.

**Dark pigment:** Helps the absorption of heat from sunlight.

**Fat layer:** Thick fat layer aids insulation especially when swimming, when the fur loses much of its insulation value.

### Blubber and fat

Blubber is found in marine mammals including whales and seals. It is highly vascularised (many blood vessels) fat tissue just below the skin and surrounds most of the body, insulating it. Fat (adipose tissue) is less vascularised and has less structure but acts in a similar way by trapping heat in the body. Fat is divided into white fat (insulation and energy storage) and **brown fat** (heat generation)



### Feathers

Feathers evolved primarily as insulation rather than as a flight surface. They still carry out the function of insulation exceedingly well. Down feathers (left) are found below the outer feathers. They are such effective insulators that birds such as emperor penguins are at risk of over heating when out of water. Down is used by humans as an insulator in bedcovers, jackets, and sleeping bags.

## Keeping cool

As much as animals need to keep warm in some environments, they need to keep cool in others. Structural features to help prevent overheating include large exposed areas of highly vascularised skin, localisation of fat stores, and fur and feather colouration.



### Ears

The external ears of desert mammals provide a structural mechanism to assist heat loss. Elephants, jack rabbits, and fennec foxes have huge ears relative to their body size, and the ears are covered in highly vascularised skin. As blood passes through the blood vessels it loses heat to the environment. Having large ears helps to catch a breeze which increases evaporative cooling. Elephants are able to flap their ears, further increasing their cooling ability.



### Feathers

Not only do feathers keep heat in but they can be just as effective at keeping heat out. Desert birds often have dark coloured feathers, especially on their dorsal surface (back). These absorb heat near their surface, preventing it reaching the skin. A breeze, especially when flying, can then easily remove the heat. Feathers on the breast and belly are often thinner to increase heat loss. The bald head of the vulture above may also have some thermoregulatory function.



### Localisation

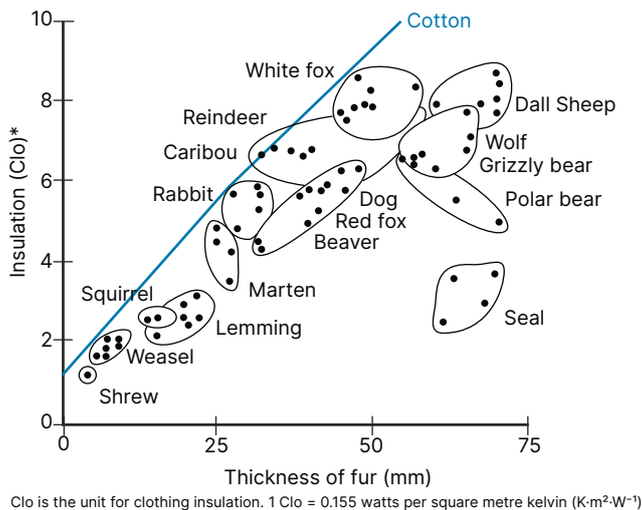
One strategy to help lose heat is to localise fat stores and hair cover. Instead of having fat reserves evenly distributed over the body they are concentrated in certain areas, such as the back hump in camels. This reduces the insulating effect of the fat to a small region on the back (which may also help reduce heat absorption from the sun). Hair is found mainly on the dorsal surface to protect the skin from the sun but the belly may be relatively naked.



1. Why are feathers good insulators? \_\_\_\_\_  
\_\_\_\_\_
2. How do feathers help protect a desert bird from overheating? \_\_\_\_\_  
\_\_\_\_\_  
\_\_\_\_\_
3. (a) In what way is blubber different from fat? \_\_\_\_\_  
\_\_\_\_\_  
\_\_\_\_\_
 

(b) How do blubber and fat help an animal retain heat? \_\_\_\_\_  
\_\_\_\_\_  
\_\_\_\_\_
4. How does fur or hair help to insulate the body? \_\_\_\_\_  
\_\_\_\_\_  
\_\_\_\_\_
5. In what way can ears be used as thermoregulatory structures? \_\_\_\_\_  
\_\_\_\_\_  
\_\_\_\_\_
6. How does localising structures such as fur and fat deposits help in the regulation of body temperature? \_\_\_\_\_  
\_\_\_\_\_  
\_\_\_\_\_

7. Study the graph below:



- (a) What is the relationship between fur (hair) length and insulation effect: \_\_\_\_\_  
\_\_\_\_\_
- (b) Seal fur appears to provide little insulation. Why would this not affect the seal in a cold environment?  
\_\_\_\_\_  
\_\_\_\_\_

# 154 Behavioural Responses for Thermoregulation

**Key Idea:** Thermoregulation is related to energy balance and animals have behavioural responses to reduce energy consumption when energy sources are scarce.

Animals have many different behavioural responses to help them regulate their body temperature. These may be very simple, such as moving out of the sun into the shade, or they may be more complex, such as **hibernation** over winter or periods of **torpor**. **Endotherms** expend large amounts of

energy to maintain a high constant body temperature and high metabolic rate. Periods of reduced activity and low body temperature, such as occur during hibernation, conserve large amounts of energy and enable survival through periods when food is scarce. Such energy savings are particularly important for many small endotherms because they lose heat very quickly and their per gram metabolic costs are much higher than for larger animals.

Bernard DUPONT CC 2.0



Regulating body temperature can be as simple as moving in or out of the shade. In some cases the shade can be carried around with as shown by these cape ground squirrels.



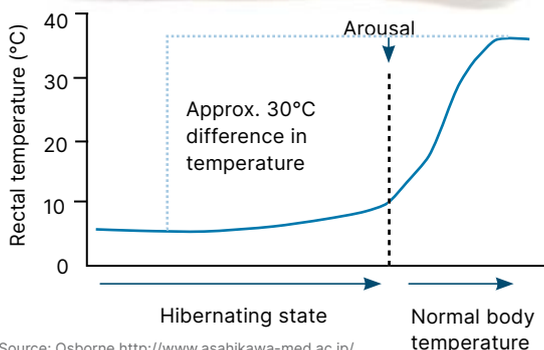
Many animals huddle together in cold conditions to conserve body heat. This behaviour is called **kleptothermy** and it occurs in both endotherms and ectotherms.



**Panting** evaporates water from the lining of the oral cavity. In dogs, it is an important way to lose accumulated heat because they have sweat glands only on the pads of their feet.

## Hibernation

- ▶ Hibernation is a prolonged (usually seasonal) state of reduced activity and metabolic depression, during which body temperature drops. It markedly reduces energy expenditure, allowing the animal to survive winter.
- ▶ Short daylength, low temperatures, and low food availability are strong cues for entering hibernation.
- ▶ The graph (below) shows body temperature in golden hamsters during hibernation. Note the difference between the animal's normal and hibernating body temperature (~30°C).
- ▶ During hibernation, metabolic activity (blood flow to the brain and respiration rate) significantly decreases. It increases to a maximum during arousal and tapers off once normal body temperature is achieved. The elevated metabolic rate during the arousal period speeds up arousal and rapidly clears waste products from the body.



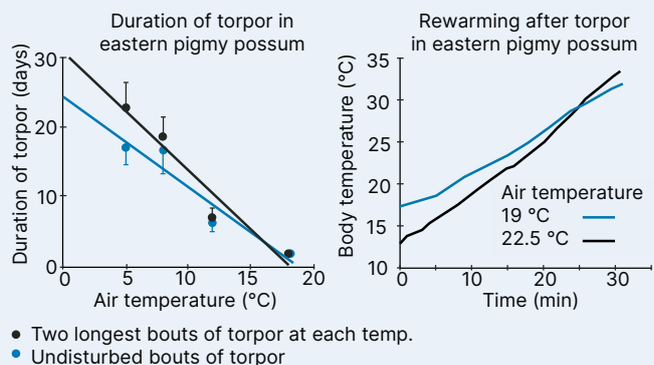
Source: Osborne <http://www.asahikawa-med.ac.jp/dept/mc/phys1/profiles/osborne.html>

## Torpor

- ▶ Some animals reduce their metabolic activity on a daily (or rather nightly) basis during their sleep. This is called torpor. The eastern pigmy possum is found throughout the eastern coast and south of Australia. It weighs up to 43 grams. During winter, it carries out daily torpor. The period of the torpor depends on the air temperature. The possum may remain in a torpid state for up to 35 days at a time during winter hibernation and its body temperature may fall as low as 1 °C.



Phil Spark CC 2.0



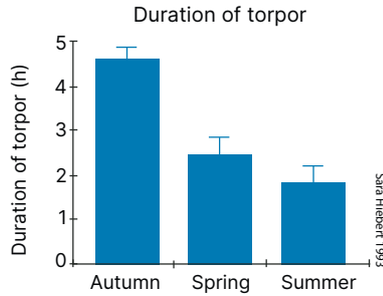
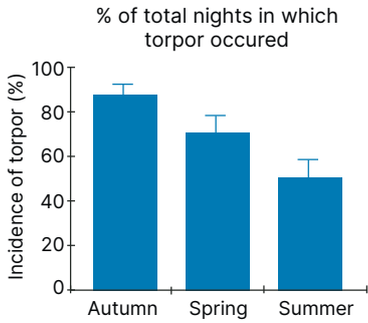
- Two longest bouts of torpor at each temp.
- Undisturbed bouts of torpor

Source: Australian Journal of Zoology, 1993



## Aestivation

- ▶ **Aestivation** is a form of hibernation or torpor that occurs during the warmer months of the year rather than the cooler months. In endotherms, it is physiologically difficult to distinguish from torpor and is essentially an arbitrary label used to distinguish torpor during warmer months.
- ▶ It occurs in many mammals including echidnas, dunnarts (a small mouse-sized marsupial), possums, and bats. Many mammals that aestivate enter daily torpor for most months of the year and are constantly active only during the most favourable months (the hottest or coolest months depending on the environment).
- ▶ Aestivation is often a response to a lack of food in arid environments. It reduces the need to expend energy keeping cool and can reduce the amount of water lost due to evaporation (by between 20-40% in dunnarts). Aestivation also occurs in some birds, although it is much rarer. Rufous hummingbirds (below) enter a nocturnal torpor during the summer months to reduce thermoregulatory energy expenditure.



1. What is the difference between hibernation, torpor, and aestivation? \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_
  
2. (a) What are the survival advantages of hibernation? \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_
- (b) What are the common environmental cues triggering hibernation and why? \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_
  
3. (a) What happens to the body temperature of the golden hamster during hibernation? \_\_\_\_\_  
 \_\_\_\_\_
- (b) Why does this change in temperature occur? \_\_\_\_\_  
 \_\_\_\_\_
- (c) Explain why blood flow to the brain and respiration rate may peak during arousal from hibernation:  
 \_\_\_\_\_  
 \_\_\_\_\_
  
4. (a) How does air temperature affect the length of bouts of torpor in the eastern pigmy possum?  
 \_\_\_\_\_  
 \_\_\_\_\_
- (b) Why does torpor enhance survival of small endotherms in cold conditions? \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_
  
5. What is often the trigger for aestivation? \_\_\_\_\_  
 \_\_\_\_\_

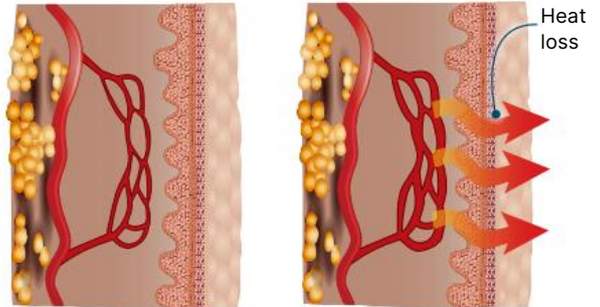
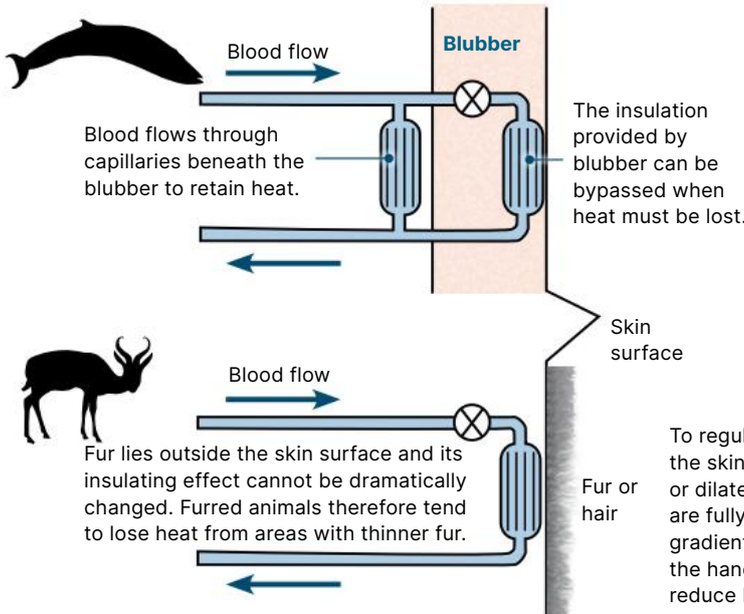
# 155 Physiological Mechanisms for Thermoregulation

**Key Idea:** Temperature can be regulated and maintained by internal mechanisms that control energy use and blood flow. Physiological mechanisms are internal mechanisms that affect how the body operates. Mechanisms of physiological

**thermoregulation** include the use of energy resources (e.g. metabolising fat), changing aspects of metabolism (redirecting chemical reactions), and changing aspects of blood flow (**vasoconstriction** and countercurrent flows).

## Regulating blood flow to the skin

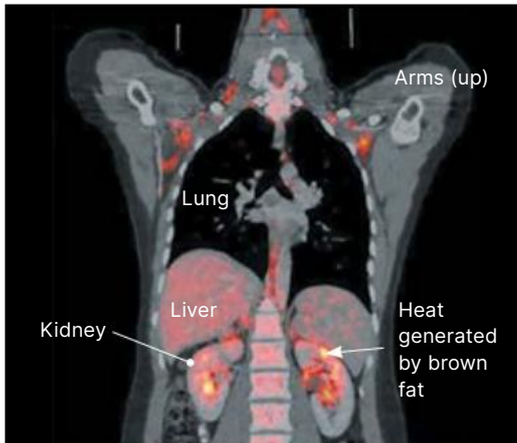
The blubber in marine mammals provides good **insulation** against heat loss but presents a problem in warmer waters or during exertion when a lot of metabolic heat is generated. In these situations, blood flows through the blubber to the skin surface where excess heat is dissipated. Cold adapted land mammals have insulation outside the skin and have thinly covered areas on the face and feet, where heat can be lost during exertion.



**Vasoconstriction**  
**Vasodilation**  
*Constriction of a the capillaries (vasoconstriction) restricts blood flow. Vasodilation allows blood to flow through the capillaries and transport heat from the body to the skin.*

To regulate heat loss or gain from the skin, the blood vessels beneath the skin's surface constrict (vasoconstriction) to reduce blood flow or dilate (**vasodilation**) to increase blood flow. When blood vessels are fully constricted, there may be as much as a 10°C temperature gradient from the outer to inner layers of the skin. Extremities such the hands and feet have additional vascular controls, which can reduce blood flow to them in times of severe cooling. These controls are mediated through the vasomotor centre in the hindbrain and are autonomic (occur without conscious thought).

## Uncoupling H<sup>+</sup> flow and ATP generation



In some cells, such as the **brown fat** cells of mammals, 'uncoupling' proteins in the inner mitochondrial membrane act as channels, allowing protons to pass directly to the matrix of the mitochondria without being used to generate ATP. This allows the energy of the proton gradient to be dissipated, generating body heat (bright spots above).

## Evaporative heat loss



Evaporative heat loss is a effective way of losing heat. In humans this happens via the production of sweat from sweat glands in the skin. The sweat absorbs heat from the skin which causes it to evaporate. People in warm climates tend to sweat in a more uniform way than those who live in cooler climates. People not acclimatised to warm climates can sweat up to 2 L h<sup>-1</sup> less than those in warmer climate and the sweat usually beads up and drips off the body.

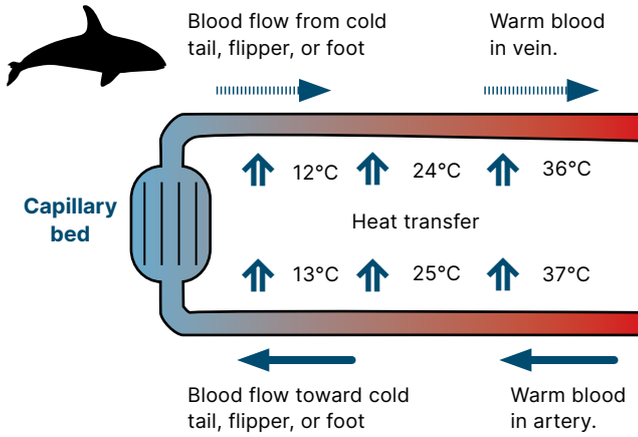
Not all animals are able to sweat but still carry out evaporative heat loss. Dogs pant, using saliva on the tongue for evaporative cooling. Kangaroos lick their forearms, which have blood vessels close to the surface of the skin. Heat is transferred from the skin to the saliva which absorbs the heat and evaporates.

1. Explain why cold adapted terrestrial mammals have regions of the body with thinner fur: \_\_\_\_\_



**Countercurrent heat exchangers**

In a counter current system, arteries coming from the body and carrying warm blood lie alongside veins returning from the skin or limbs, which carry cooler blood. Heat is transferred from the arteries to the veins before it is lost from the skin or limbs. Mammals in cold environments use countercurrent exchange to reduce heat losses to the environment. Those in hot environments use countercurrent exchange in the opposite way: to cool arterial blood supply to the brain during intense activity.



**Shivering**

Shivering is an involuntary response that occurs when the body is exposed to cold temperatures. When the body's temperature drops below the set temperature of the hypothalamus, signals are sent to the skeletal muscles to begin rapid rhythmic contractions. This increases respiration in the muscles which results in heat energy being lost during the exothermic reactions that breakdown glucose. This lost heat warms the body.

Shivering also occurs during a fever. The hypothalamus increases the body's set temperature. In order to help the body reach this new higher temperature, the muscles begin shivering.



Vasoconstriction and goosebumps (left) and shivering (right) are involuntary responses low to temperature.

2. (a) Explain how countercurrent heat exchangers help retain body heat in marine mammals:

---



---



---



---

(b) Explain the thermoregulatory changes a marine mammal would make when moving from colder to warmer waters:

---



---



---

3. (a) What is the purpose of sweating and how does it achieve its effect? \_\_\_\_\_

---



---

(b) Why does a dab of methanol or ethanol on the skin feels cold, even if the liquid is at room temperature?

---



---

4. How do the blood vessels help to regulate the amount of heat lost from the skin and body?

---



---



---



---

5. (a) How does shivering warm the body? \_\_\_\_\_

---



---

(b) When might shivering occur? \_\_\_\_\_

# 156 Hormonal Mechanisms for Thermoregulation

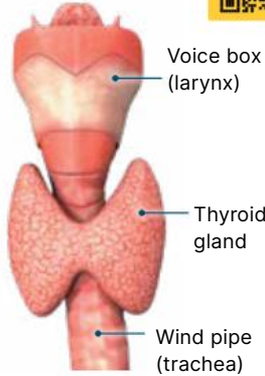
**Key Idea:** Hormones regulate many aspects of metabolism. Over or under production of hormones can affect metabolic aspects of thermoregulation, as can a lack of cellular response to those hormones. The hypothalamus has a central role in **thermoregulation**.

It registers changes in core body temperature and coordinates nervous and hormonal responses to restore normal body temperature. **Thyroxine**, produced by the thyroid gland, is an important hormone in thermoregulation. Insulin also has a thermoregulatory role but this is less well understood.

## The thyroid gland and thermoregulation



- ▶ The thyroid gland is a butterfly shaped endocrine gland located just below the Adam's apple at the front of the trachea. The thyroid secretes several hormones, but the main hormone produced is thyroxine ( $T_4$ ).
- ▶ Thyroid hormones have many functions including regulating metabolism, growth and development, and body temperature (below).



## Hyperthyroidism and temperature regulation

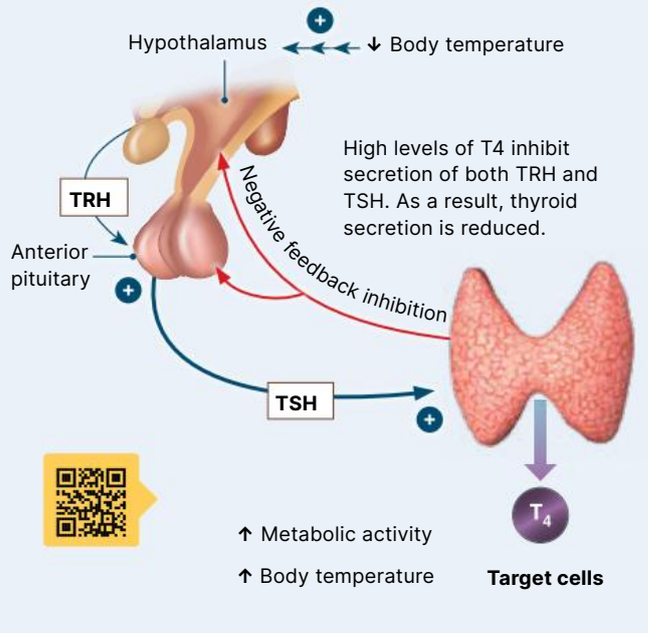
- ▶ One of the effects of  $T_4$  is to speed up metabolic activity in cells. The increase in metabolic activity also results in the production of heat and, under normal conditions, this is one of the mechanisms by which the body raises body temperature.
- ▶ The negative feedback regulation of  $T_4$  production can be disrupted by hyperthyroidism, a condition where the thyroid makes too much  $T_4$ . This can disrupt temperature regulation.



The most common cause of hyperthyroidism is Graves' disease, characterised by an enlarged thyroid (goitre) and bulging eyes (above). In Graves' disease, the negative feedback loop is bypassed because a protein called thyroid stimulating immunoglobulin (TSI) binds directly to the thyroid and stimulates  $T_4$  production. Because  $T_4$  production is independent of TSH production, the usual regulatory mechanisms are ineffective.

## Negative feedback regulates thyroxine production

- ▶ Thyroxine ( $T_4$ ) production is controlled by negative feedback. This mechanism involves two parts of the brain, the hypothalamus and the pituitary gland.
- ▶ Low body temperature stimulates the hypothalamus to secrete thyrotropin releasing hormone (TRH), which in turn stimulates cells in the anterior pituitary to secrete thyroid stimulating hormone (TSH).
- ▶ TSH acts on the thyroid gland, causing it to produce thyroid hormones, including  $T_4$  (thyroxine).  $T_4$  binds to target cells, increasing their metabolic activity and producing heat.
- ▶ High levels of circulating thyroid hormones inhibit production of TRH and TSH. As a result, thyroid secretion is reduced. When the level of thyroid hormones drops below a certain threshold, TRH and TSH production begins again.

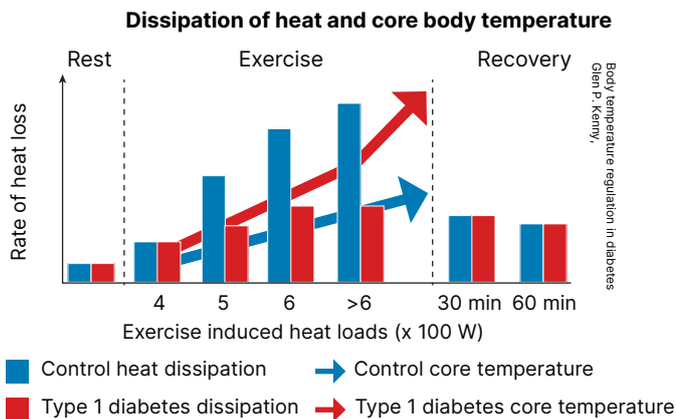
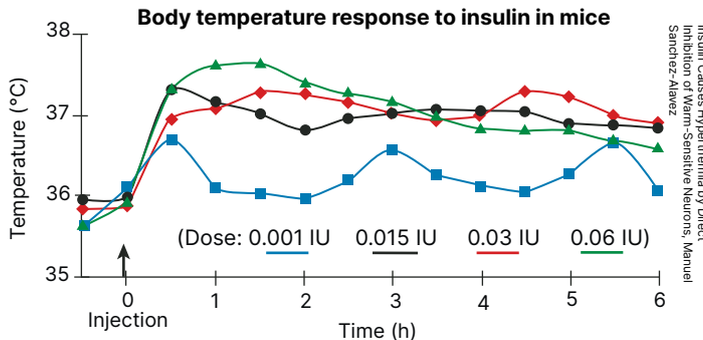


1. How is  $T_4$  involved in temperature regulation? \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_
2. Explain how  $T_4$  production is regulated by negative feedback: \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_



### Insulin and thermoregulation

- ▶ The hormone insulin is normally linked to the regulation of blood glucose. Insulin signals to cells to take up glucose from the blood. The cells then use the glucose to produce ATP for use in metabolic processes. However, research has also shown that insulin can directly affect core body temperature.
- ▶ Experiments have shown that when insulin is injected into the preoptic area of the hypothalamus there is a rapid hyperthermic (increased temperature) response (above right). This was caused by the stimulation of **thermogenesis** in **brown fat** (see page 227). This may help to explain why diabetics (whose cells do not produce insulin or do not respond to insulin) often have difficulty with thermoregulation.
- ▶ Insulin also indirectly affects thermoregulation. Studies have shown that people with type 1 diabetes (a lack of insulin production) and type 2 diabetes (insulin resistance) often suffer from problems with thermoregulation, including poor control over **vasodilation** or **vasoconstriction**. In particular, type 2 diabetes is associated with impaired vasodilation and this can severely affect the ability to dissipate heat. Type 1 and 2 diabetes are also associated with impaired sweat production, which again also affects the ability to dissipate heat.



3. Why do high levels of T4 not inhibit its production from the thyroid gland in a person with Graves' disease?

---



---



---



---

4. (a) Would you expect someone with an overactive thyroid gland to feel hot or cold? \_\_\_\_\_

(b) Would you expect someone with an underactive thyroid gland to feel hot or cold? \_\_\_\_\_

5. What was the effect of injecting insulin into the preoptic area of the hypothalamus? \_\_\_\_\_

---



---

6. (a) Describe the relationship between exercise induced heat loads and core body temperature:

---



---

(b) How does this differ between people with and without type 1 diabetes? \_\_\_\_\_

---



---



---



---

7. Why does impaired vasodilation affect thermoregulation in people with diabetes? \_\_\_\_\_

---



---



---

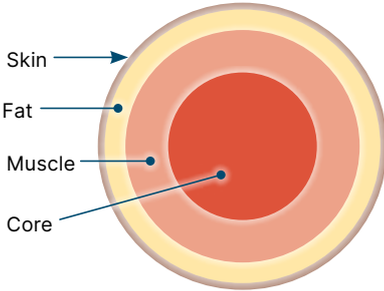


---

# 157 Modelling Human Thermoregulation

**Key Idea:** Modelling heat loss can help us design better clothes to reduce chances of hypo- or hyperthermia. Humans evolved in the warm climate of Africa. As a result, humans are better adapted for dissipating heat than conserving it. For example, we have little body hair, and are generally tall and thin, both adaptations for losing heat.

To keep warm, we wear clothes and live in houses, but this can result in us being too hot in some situations (try running a marathon in a polar jacket). By modelling how we thermoregulate, we can design clothes and environments that never feel too warm or too cold.



## A model of thermal inputs and outputs in humans

Modelling **thermoregulation** begins with a simple model of a human in cross section. The model has four layers: the core, muscle, fat, and skin. The model must then account for the rate at which each of these layers produces and dissipates heat. The models become more complicated as different thermoregulatory mechanisms are added (e.g. heat loss by increased blood flow to the skin) and different areas of the body are taken into account (e.g. torso vs legs).

Once a mathematical model for this is produced, we can investigate the effect of changing the environment or adding clothes on the rate of heat loss (right).

<b>Input</b> <b>Environment</b> • Temperature • Relative humidity • Wind speed <b>Exertion</b> • Physical activity <b>Clothing</b>	<b>Energy balance</b>	<b>Convection</b> • Blood flow	<b>Conduction</b>	<b>Heat generation</b> • Basal metabolism • Exertional metabolism (muscle)	<b>Output</b> <b>Distribution of temperatures around the body</b>	
	<b>Thermoregulation</b>	<b>Shivering</b> • Heat generation of the muscle	<b>Vasoconstriction</b> • Decrease blood flow to the skin	<b>Vasodilation</b> • Increase blood flow to the skin		<b>Sweating</b> • Heat loss due to evaporation
	<b>Heat transfer to the environment</b>	<b>Convection</b>	<b>Radiation</b>	<b>Respiration</b>		<b>Perspiration</b>
	<b>Circadian rhythm</b>	<b>Basal metabolism</b>	<b>Skin blood flow</b>			

Adapted from Gnu Umnikrishnan et al 2020



Moisture wicking clothing has become an important part of sports wear. Older fabric trapped sweat, producing a hot humid environment close to the skin. Moisture wicking clothing rapidly moves moisture away from the skin to the outer surface where it is evaporated, leaving a cooler inner surface. Cycling vests are designed to prevent extreme cooling caused by air movement on the front of the torso, but allow for air flow and cooling over the rear of the torso, which is sheltered from the wind.

Understanding how humans thermoregulate allows us to develop safer working conditions. Working in extremely hot conditions can cause heat stress. Developing clothing that can help regulate body temperature can help this. Astronauts working in space wear body suits through which cool water cycles to regulate their body temperature. Similarly working in extremely cold environments can leave a person at risk of hypothermia.

Thermoregulation provides a good example of the misinterpretation of data. It is often stated that humans lose up to 40% of their body heat through their head. This is a misinterpretation of data released from a study in the 1950s. It found that without a covering, heat lost from the head amounts to 40% of the heat lost from the body of a *fully clothed person*. If the person was unclothed then the heat lost from the head was proportional to the surface area of the head compared to the body, which is about 10%.

1. Why does modelling the different layers of the body help understand thermoregulation?

---



---



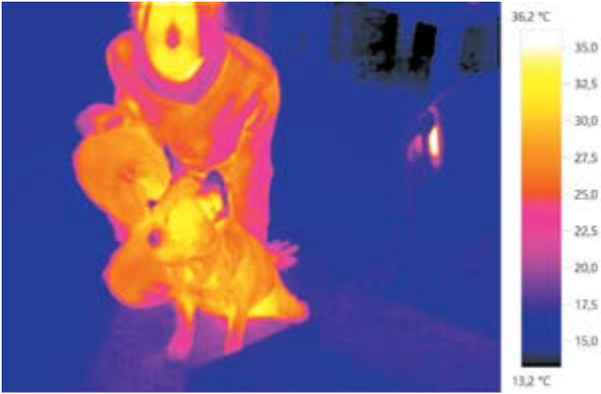
---



---



### Thermal imaging for better thermoregulation



Thermal imaging cameras have become a useful tool in studying human thermoregulation and designing clothing to help humans regulate their body temperature.

Infrared light can be detected by a special infrared camera. Warm objects give off infrared lightwaves. The hotter an object is, the shorter the wavelengths of infrared light it gives off. Once detected by the camera, the differences in these wavelengths can be interpreted as temperature. From the image on the left, it can be seen that uncovered parts of the body, like the face, show up as the warmest (brightest). Where heat is being lost from through the clothing can also be seen.

Clothing with high **insulation** or heat reflective layers will show as dark blue or black in thermal cameras.

#### Temperature and dexterity

Many jobs rely on manual dexterity. Cold fingers make it hard to carry out even simple tasks (imagine zipping up a polar jacket wearing polar grade gloves in a blizzard)! Models of finger skin temperature and dexterity are important in designing gloves and equipment that will provide the best dexterity under a wide range of temperatures and conditions.

The data below shows how dexterity (measured by the number of pins and washers assembled on a Purdue pegboard) changes with skin temperature.

Skin temperature (ST)    Finger dexterity (FD)

ST	FD	ST	FD
8.5	19.5	16	25
10	20.5	16.5	25.5
11	20.5	15.5	26
8.5	21	15	26
10	21.5	17	24.5
10	22	17.5	26
10.5	23	26	27
12	23	18.5	27
12	23.5	20	27
12.5	23.5	22.5	26
14	24	24.5	28
14	24.5	28.5	28
13.5	25	31	29
14.5	25.5	31.5	28.5

Adapted from Arctic Medical Research 1995

- Enter the data below left into a spreadsheet (in two columns only). The instructions below are for Microsoft Excel, but most spreadsheets work in a similar way. Select the data and choose to plot a scatter graph. Select one of the data points on the graph and right click to bring up the options menu. Select **Add trendline**. Under **Trendline options** click **Logarithmic**. Under **Forecast** type 10 periods in both **forward** and **backward**. Then click **Display equation on chart**. The equation can be used to calculate dexterity for any point along the graph

(a) Is there an optimal skin temperature for finger dexterity?

\_\_\_\_\_

(b) Why is forecasting the data beyond 40°C not useful?

\_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_

(c) How might this kind of data and model be useful for clothing designers and health and safety legislators?

\_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_

- How does thermal imaging help in the development of both insulating and cooling clothing?

\_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_

- What role does moisture wicking clothing play in sports wear?

\_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_

1. Provide a definition for each of the following terms:

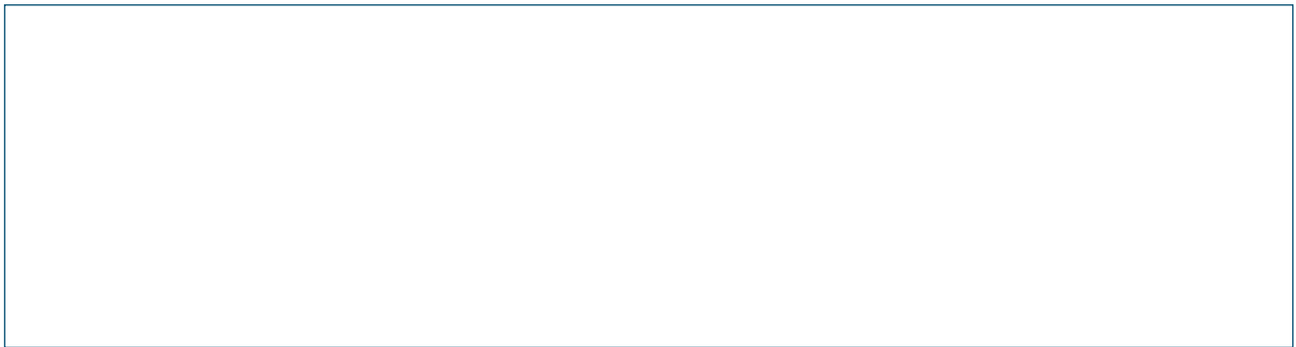
(a) endotherm: \_\_\_\_\_  
 \_\_\_\_\_

(b) hibernation: \_\_\_\_\_  
 \_\_\_\_\_

(c) thermoregulation: \_\_\_\_\_  
 \_\_\_\_\_

(d) torpor: \_\_\_\_\_  
 \_\_\_\_\_

2. Use a diagram to show how the human body maintains its constant temperature. Use the terms *sweating*, *shivering*, *vasodilation* and *vasoconstriction*.

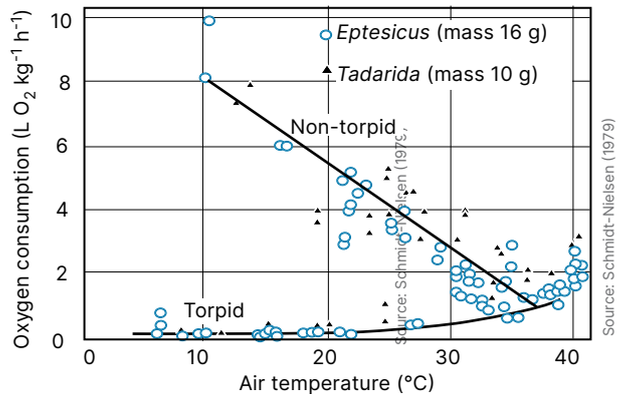


3. The plot to the right shows the per mass rate of oxygen consumption in two species of small North American bats. The rate of oxygen consumption is a measure of the energy being used.

The bats are of similar size, but *Eptesicus* hibernates in the northern part of its range, whereas *Tadarida* migrates south but does not hibernate. With reference to the plot:

(a) What happens to the per mass rate of oxygen consumption in bats that enter torpor (relative to that when active)?

\_\_\_\_\_  
 \_\_\_\_\_



(b) What happens to the per mass rate of oxygen consumption in bats that enter torpor (relative to that when active)?

\_\_\_\_\_  
 \_\_\_\_\_

4. Polar bears and penguins live in similar polar environments. They have both converged on certain structural features that help them remain warm in polar conditions.

List three structural features that are similar in polar bears and penguins and describe how these features help them maintain a stable body temperature:

(a) \_\_\_\_\_  
 \_\_\_\_\_

(b) \_\_\_\_\_  
 \_\_\_\_\_

(c) \_\_\_\_\_  
 \_\_\_\_\_

# Osmoregulation



## Key Terms

- ABA (abscisic acid)
- ADH (anti-diuretic hormone)
- halophyte
- hydrophyte
- kidneys
- mesophyte
- metabolism
- osmoconformer
- osmoregulation
- osmoregulator
- osmosis
- stomata
- urine
- xerophyte

## Key Concepts

- ▶ Many organisms must regulate the concentration of water in their bodies.
- ▶ Bony fish have adapted to live in either saltwater or freshwater environments and have mechanisms that allow them to maintain their internal water balance.
- ▶ Plants have adaptations that allow them to live in a variety of climatic conditions.

## Principles of osmoregulation

### Activity Number

- |                          |   |  |         |
|--------------------------|---|--|---------|
| <input type="checkbox"/> | 1 | Explain what is meant by osmoregulation and explain why organisms need a specific balance of water and ions in their bodies.   | 159     |
| <input type="checkbox"/> | 2 | Explain the osmoregulatory problems of animals in freshwater, salt water, and on land.   | 159     |
| <input type="checkbox"/> | 3 | Distinguish between osmoconformers and osmoregulators and explain why all freshwater organisms are osmoregulators. Explain how water and ions enter and leave cells. | 159-160 |

## Osmoregulatory responses in animals

- |                          |    |   |              |
|--------------------------|----|---|--------------|
| <input type="checkbox"/> | 4  | Analyse and interpret data related to osmoregulation, e.g. in an intertidal organism.   | 160          |
| <input type="checkbox"/> | 5  | Describe the close relationship between osmoregulation and excretion and relate this to the organs involved (with reference to fish and mammals in particular).   | 161-162      |
| <input type="checkbox"/> | 6  | Describe and explain structural mechanisms for osmoregulation, including the role of excretory system (kidneys in mammals and kidneys and gills in fish).   | 159, 161-162 |
| <input type="checkbox"/> | 7  | Describe and explain behavioural responses for osmoregulation including drinking, shade-seeking (to lower evaporative losses), and habitat (salinity) selection.  | 162          |
| <input type="checkbox"/> | 8  | Using examples, describe and explain physiological mechanisms for osmoregulation including metabolism of glucose and fat, active secretion and reabsorption of ions, concentration of urine, and reabsorption of water from the gut and nasal passages. | 161-162      |
| <input type="checkbox"/> | 9  | Explain how the hormone ADH and feedback systems regulate urine volume in the mammalian kidney and explain its role in water balance.   | 163          |
| <input type="checkbox"/> | 10 | <b>SI:</b> Investigate structural, behavioural, and physiological mechanisms that maintain water balance in different species   | 159-164      |

## Osmoregulatory responses in plants

### Science understanding

- |                          |    |   |     |
|--------------------------|----|---|-----|
| <input type="checkbox"/> | 11 | Describe and explain the main osmoregulatory problem facing most plants.  | 164 |
| <input type="checkbox"/> | 12 | Identify and explain the various mechanisms that maintain water balance in plants of terms of structural features (stomata, vacuoles, cuticle) and hormonal mechanisms (synthesis and action of ABA in response to water stress). Include reference to hydrophytes, mesophytes, xerophytes, and halophytes. | 164 |
| <input type="checkbox"/> | 13 | <b>SI:</b> Investigate the distribution of stomata in plants adapted to different environments. Relate differences to different osmoregulatory challenges.  | 165 |
| <input type="checkbox"/> | 14 | <b>SI:</b> Investigate tolerance limits for salt balance by testing the effect of salt level on plant growth.   | 166 |

# 159 What is Osmoregulation?

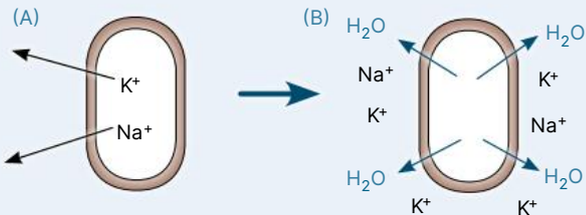
**Key Idea:** Osmoregulation is the process by which organisms regulate the concentrations of salts and water in their bodies. The chemical reactions sustaining life require a specific osmotic environment. Organisms in which the osmolarity of the body fluids is different from the osmolarity of the environment must regulate their salt (solute) and water levels through a process called **osmoregulation**. As a

result of disease or a change in environment, organisms may sometimes experience osmotic stress (an abnormal concentration of dissolved solutes). Osmotic stress disrupts the steady state and can be fatal if the imbalance is prolonged. Osmoregulators are able to maintain the osmolarity of their body fluids independently of the environment. How they do this depends on the type of environment they inhabit.

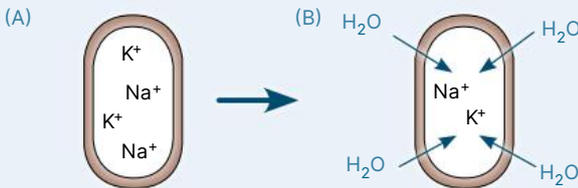
## Ion and water fluxes in bacteria

Recall that water will diffuse from regions of lower solute concentration (higher free water concentration) to regions of higher solute concentration (lower free water concentration). Water enters a bacterial cell by osmosis. The bacteria can regulate water fluxes by retaining salts or pumping them out of the cell. They respond to osmotic stress by rapidly accumulating ions or organic solutes via membrane transporters which are stimulated by increases in osmolarity in the environment.

When the solute level is lower outside the cell than inside it, a bacterium will pump salts out of the cell (A) so that water will tend to leave by **osmosis** (rather than enter (B)).



If the solute level is higher outside the cell, a bacterium will retain salts (A), so that water will tend to enter (rather than leave) the cell (B).



## Water fluxes in different environments



The type of external environment an organism lives in will dictate the natural direction of water movement. For marine bony fish, the concentration of solutes is higher in the environment than in the body fluids of the fish. The fish loses water across the skin and gills, and it must be replaced or the fish will experience osmotic stress.



The body fluids of freshwater fish contains a higher solute concentration than the fresh water environment. The fish gains water as it flows across the gills during gas exchange. A freshwater fish must remove the excess water from its body or it will experience osmotic stress.

1. What is osmoregulation? \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_
2. How does the bacterium reduce the amount of water entering the cell? \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_
3. The two diagrams (right) depict a bony fish in a marine environment and a bony fish in a freshwater environment.
  - (a) On each diagram, draw arrows to show the direction of water movement:
  - (b) In the marine environment, is the fish body hypertonic or hypotonic to the seawater? \_\_\_\_\_  
 In the freshwater environment, is the fish body hypertonic or hypotonic to the water? \_\_\_\_\_

Fish body	Fish body
Gills	Gills
Marine environment	Freshwater environment



## Osmoconformers and osmoregulators

Organisms have differing abilities to respond to environmental change. **Osmoconformers** match the osmolarity of their environments and do not regulate their water and ion fluxes. In contrast, **osmoregulators** maintain relatively constant water and ion concentrations independently of the environment.

### Osmoconformers

Not all organisms osmoregulate. Those that do not are called osmoconformers. In osmoconformers, the osmolarity of the body fluids (number of solute particles per litre) fluctuates with the osmolarity of the environment. However, the solute composition of body fluids may be different from those in the water.

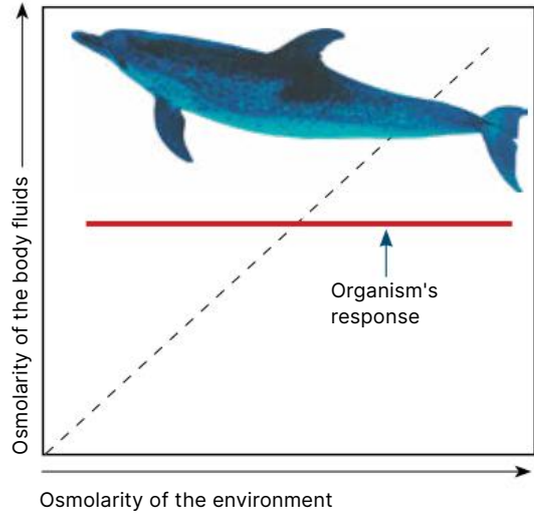
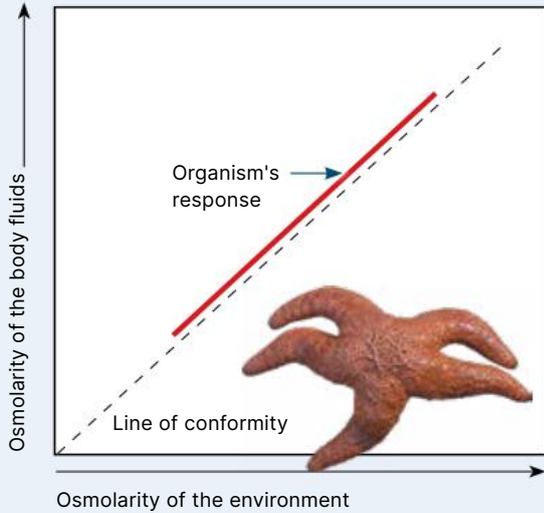
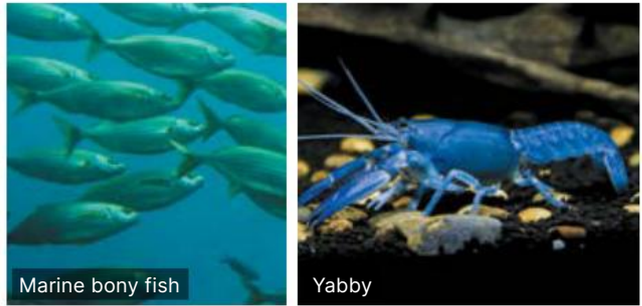
All osmoconformers are marine (although not all marine organisms are osmoconformers)! Most marine invertebrates are osmoconformers and many rely on a relatively stable external osmotic environment for survival. Deviations outside of their tolerance range are fatal. However, some intertidal species can tolerate greater fluctuations in the osmotic environment, such as occur with the frequent dilutions of seawater as the tide changes.



### Osmoregulators

Animals that regulate their salt and water fluxes independently of the environment, such as bony fish and marine mammals, are osmoregulators. Osmoregulation requires large amounts of energy. Marine bony fish lose water osmotically and counter the loss by drinking salt water and excreting the excess salt across the gill surfaces. Marine mammals produce a **urine** that is high in both salt and urea.

Freshwater animals, such as the freshwater crayfish (yabby), below right, have body fluids that are osmotically more concentrated than the water they live in and all are osmoregulators. Water tends to enter their tissues by osmosis and must be expelled to avoid flooding the body. Freshwater crayfish osmoregulate by excreting the excess water.



4. (a) Use the graphs above to explain how osmoconformers differ from osmoregulators: \_\_\_\_\_

---



---



---



---



---

(b) Describe one advantage and one disadvantage of being an osmoconformer: \_\_\_\_\_

---



---

# Osmoregulation in Intertidal Organisms

**Key Idea:** A line of best fit or a linear regression can be used to determine if there is a relationship between an organism's ability to regulate salt and water levels in different concentrations of saltwater.

Intertidal species are subjected to varying levels of salinity as the tides come in and out and sea water mixes with

freshwater. Species of intertidal crabs vary widely in their ability to regulate their salt and water levels in the face of environmental fluctuations. A student investigated the effect of increasing seawater dilution on the cumulative weight gain of a common rock crab. The methodology and results are provided below.

### The aim

To determine the effect of seawater concentration on crab weight gain.



Table 1. Cumulative weight gain in crabs at two seawater concentrations

Time (min)	Cumulative weight gain (mg) in 75% seawater			Cumulative weight gain (mg) in 50% seawater		
	Crab 1	Crab 2	Crab 3	Crab 1	Crab 2	Crab 3
3	3.8	4.0	4.0	5.6	6.2	5.8
6	8.0	8.3	7.7	11.2	11.6	11.9
9	11.5	11.0	9.5	17.0	17.6	17.2
12	14.8	15.1	15.2	23.5	23.6	24.0
15	18.9	19.5	19.7	29.0	28.2	28.6
18	23.5	22.9	23.8	33.0	32.5	32.7
21	26.5	26.9	26.7	37.5	37.6	39.0
24	31.5	32.0	31.2	43.1	43.5	43.6
27	35.0	35.5	35.5	48.0	48.1	47.5
30	40.0	40.1	41.2	53.0	52.6	52.8

### The method and results

- ▶ Six common rock crabs were used in the experiment.
- ▶ Three were placed in seawater dilution of 75:25 (75% seawater) and three were placed in a seawater dilution of 50:50 (50% seawater).
- ▶ Cumulative weight gain (mg) in each of the six crabs was measured at 3 minute intervals over a period of 30 minutes.

### The results

The results are presented in the table (right).

1. Plot the raw data for each individual crab for each seawater concentration as two scatter plots on the grid. Use different colours to distinguish the data sets.
2. Evaluate the strength of the relationship between the concentration of seawater and crab weight gain. You can do this in two ways:
  - (a) Draw a line of best fit for the data at each seawater concentration.
  - (b) Enter the data on a spreadsheet (time vs weight gain for each seawater concentration) and calculate a correlation coefficient ( $r$ ) for each data set. You can also plot the data and fit a regression line for each data set. Show the regression equation and the  $R^2$ . Print the graph and attach it here.
  - (c) How well does your chosen method fit the data?

---



---



---



---

(d) Why does a regression provide more information? \_\_\_\_\_

---

3. (a) Explain what is happening at each seawater concentration: \_\_\_\_\_

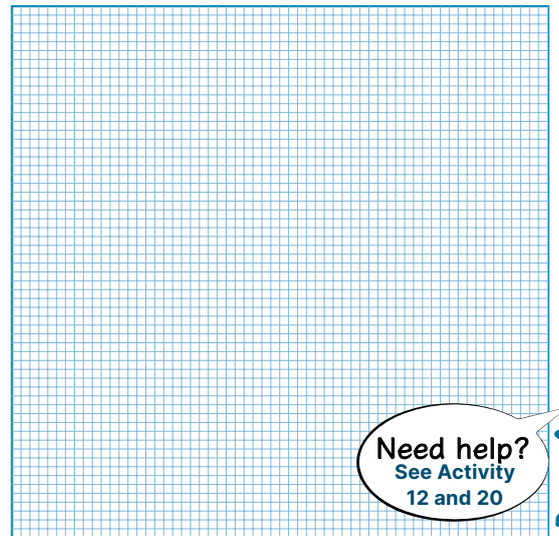
---



---

(b) What does this experiment suggest about the osmoregulatory ability of this crab species? \_\_\_\_\_

---



# 161 Osmoregulation in Bony Fish

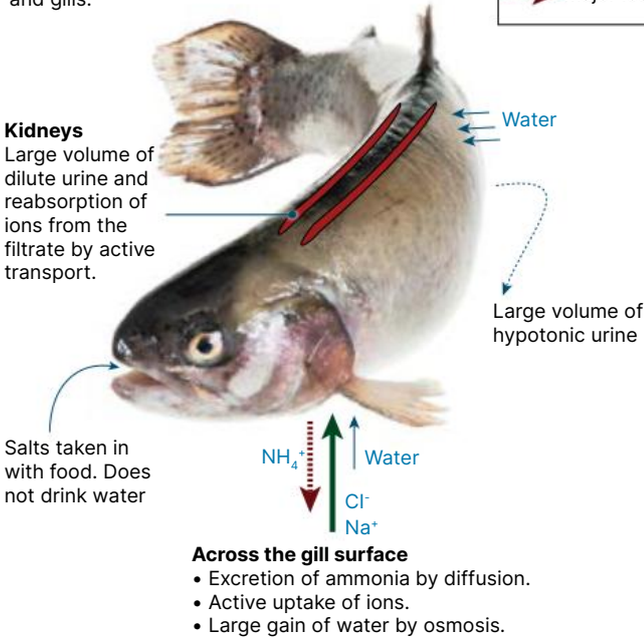
**Key Idea:** Gills are the primary organs for osmoregulation in fish, which must balance water and ion fluxes.

Bony fish face contrasting osmoregulatory problems depending on their environment. In fish, **osmoregulation** and excretion of ions and nitrogenous wastes are closely linked. Fish **kidneys** are unable produce a **urine** that is

more concentrated than the body fluids and nearly all their nitrogenous waste is excreted via diffusion across the gills, which also have an important role in salt balance. In freshwater fish, excess water is lost in copious amounts of dilute urine. In marine fish, the kidneys produce a scanty urine with the same osmolality as the blood.

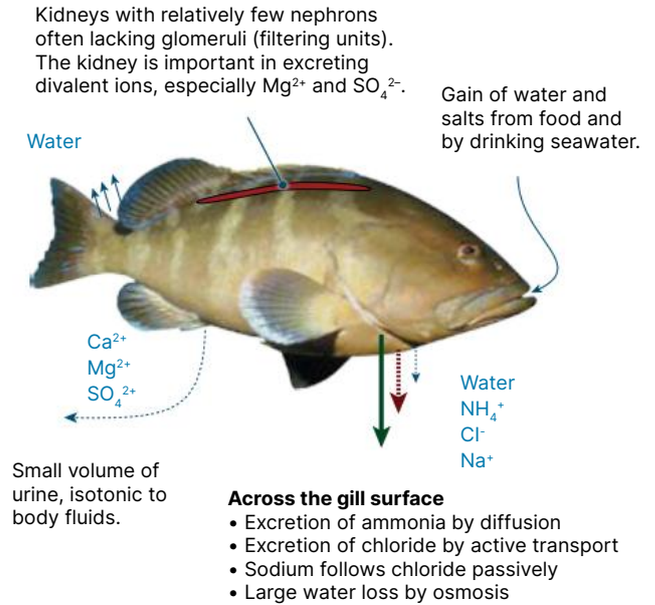
## Freshwater bony fish

Body fluids more concentrated than the freshwater. Water gained across the skin and gills.



## Marine bony fish

Body fluids 25-33% the ion concentration of seawater. Water lost across the skin and gills.



Freshwater fish gain water by **osmosis** so must excrete this excess water as well as nitrogenous waste. Their kidneys produce large volumes of dilute urine, which results in loss of valuable ions. To compensate, the kidneys reabsorb salts from the filtrate and the gills take up ions from the water.

Marine fish lose water to the environment by osmosis and must drink to replace this lost fluid. The extra salts they take in by drinking are excreted by active transport across the gill surfaces. Fish cannot concentrate the urine so the urine is isotonic and scanty. Ammonia is lost by diffusion across the gills.

1. Describe the contrasting problems of excretion and osmoregulation for bony fish in fresh and salt water environments:

---



---



---

2. (a) Explain why marine bony fish must drink vast quantities of salt water:

---



---

(b) Explain why freshwater fish do not drink water at all:

---



---

3. (a) How are the gills involved in osmoregulation in a freshwater fish?

---



---

(b) How are the gills involved in osmoregulation in a marine bony fish?

---



# Managing Fluid Balance on Land

**Key Idea:** Terrestrial animals have adaptations to obtain enough water to maintain their fluid and ion balance, either through drinking or the metabolism of foodstuffs.

All organisms, whether terrestrial or aquatic, must maintain their water and solute concentrations at levels that support their life processes. For animals on land, the main challenges to fluid and ion regulation arise from a dependence on water,

which is often in short supply. The water an animal loses must be replaced by an equal volume. These fluxes make up the animal's water budget. Water losses through evaporation from the skin and lungs, and in **urine** and faeces are balanced by water gains through eating and drinking. Animals show specific adaptations for obtaining and conserving water in an environment where water loss is a constant problem.

## Obtaining water



Most animals obtain the majority of their water by drinking. Some, such as the emu, drink infrequently, but when they do drink they consume a large volume of water. For many large predators (e.g. lions) living in dry environments, obtaining water from the food they eat is an important source of water.



Many of Australia's arid-adapted marsupials, e.g. the bettong and many of the macropods, have low water requirements and rarely drink, obtaining much of their water from oxidation of dry foods (producing ATP, CO<sub>2</sub>, and water). The rest comes from the small amount of water present in the food.



Amphibians can take up water directly through the skin, which is water permeable. When they need water, they can acquire it by **osmosis** while submerged or resting on a damp surface. Desert adapted frogs burrow underground and spread wax over their skin from epidermal wax glands.



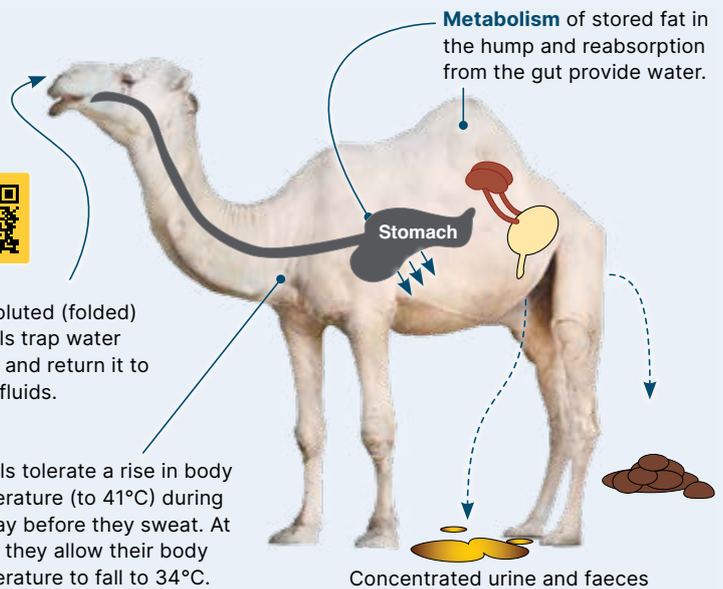
## Conserving and losing water in camels

- ▶ All mammals excrete nitrogenous wastes as urea and lose water through urine, faeces, and evaporative losses from the skin and lungs. Arid-adapted mammals all show similar adaptations to their dry environment. All have long loops of Henle in the kidney and produce a very concentrated urine. Many have reduced sweat glands to conserve water in dry environments.
- ▶ Camels have further adaptations for life in desert conditions, most directly related to water balance. For example, camels (being ruminants) can retain relatively large volumes of water in the gut, but most will need to regularly visit a water supply. When they have access to water, a dehydrated camel can drink up to 200 L in 3 minutes. In most animals, this would cause osmotic shock, but the camel's red blood cells can withstand huge fluctuations in body water content.



Convolved (folded) nostrils trap water vapor and return it to body fluids.

Camels tolerate a rise in body temperature (to 41°C) during the day before they sweat. At night, they allow their body temperature to fall to 34°C.



1. Name four ways in which water can be obtained: \_\_\_\_\_

\_\_\_\_\_

2. How does metabolism provide water for the body's activities? \_\_\_\_\_

\_\_\_\_\_



General adaptations associated with major routes of water loss in animals (vertebrates and arthropods)	
<b>An insulated body covering</b>	Body coverings reduce but do not totally eliminate water loss. They form a moderate barrier against water loss and may be thickened or covered with insulation or wax to limit water loss.
<b>Using metabolic water</b>	All animals produce water through metabolism. Metabolising fat yields more water than metabolising carbohydrate and this can be used to maintain water balance when no liquid water is available.
<b>Changing behaviour</b>	Behavioural strategies to reduce water losses, such as seeking shade, are common and may be associated with physiological adaptations to take up water from humid environments.
<b>Minimising losses from the excretory system</b>	Water balance and excretion are tightly linked because excretion of nitrogenous wastes represents a major route for water loss. Being able to produce a concentrated urine is a feature of terrestrial life.



Salleem Hameed cc 2.5

DannyS cc 3.0

The permeability of the waxy cuticle of desert adapted arthropods is very low, greatly reducing water loss across the body surface. Insects excrete nitrogenous wastes as uric acid with low water loss, and many are able to take up water from the air when humidity increases. Desert darkling beetles (above) can collect water on their hardened forewings.

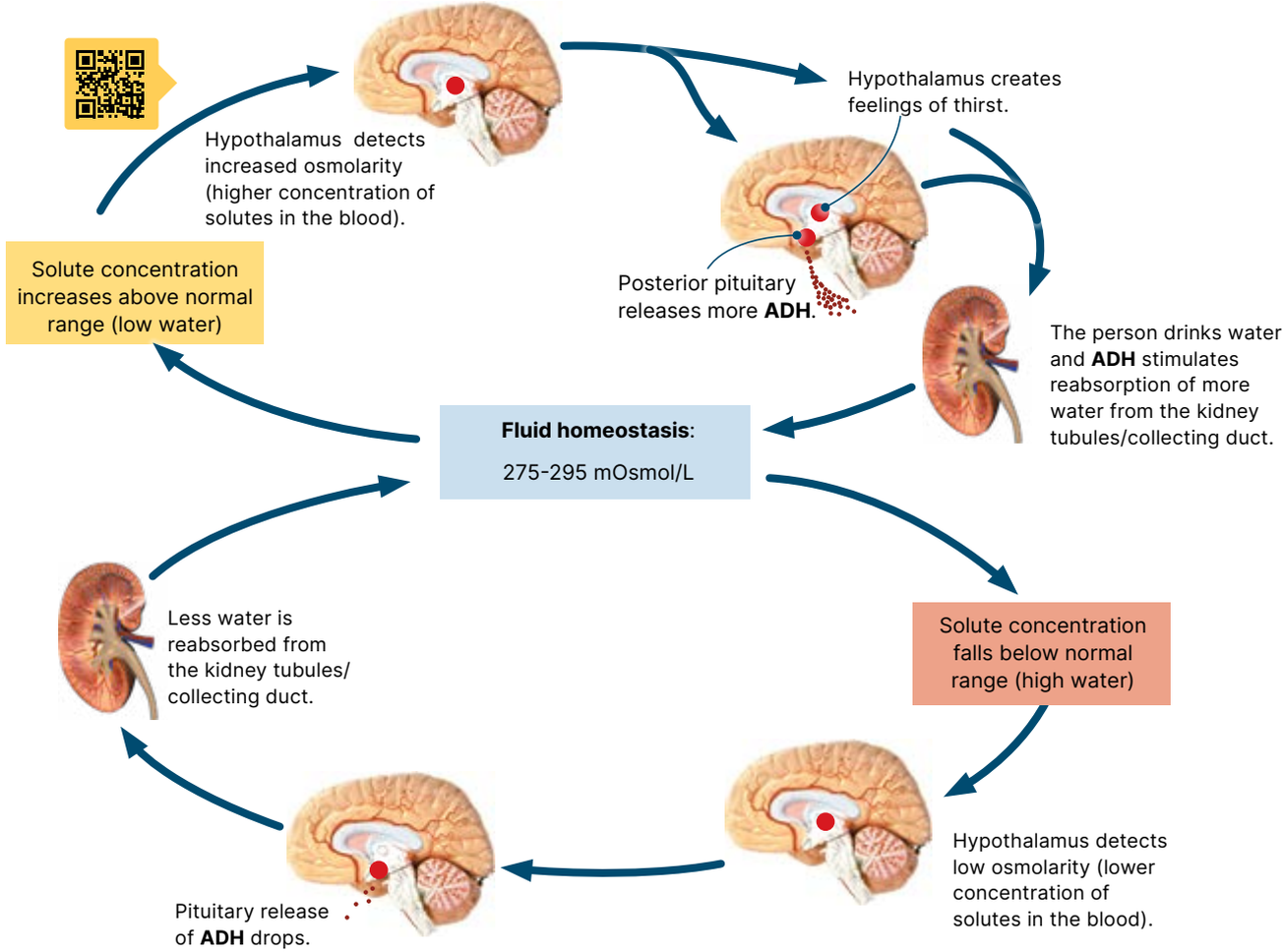
Birds and reptiles share many strategies for water conservation (a reflection of their shared ancestry). Scales in reptiles and feathers in birds (which are modified scales) reduce water losses from the body surface. Neither sweats and although water is lost in breathing, excretion of nitrogen as uric acid results in minimal loss of water. Marine birds and reptiles also have salt glands to excrete excess salts gained from eating and drinking. Shade-seeking behaviour reduces water losses during the day. In omnivorous birds (e.g. emus) reabsorption of water from food is a major source of water. Desert reptiles, such as goannas, obtain most of their water from their food and some can also absorb water across the nasal epithelium.

- Identify three ways in which animals lose water to the environment: \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_
- What features of fluid and ion homeostasis do reptiles and birds share? \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_
- Identify two ways in which each of the following animals conserves water:
  - Arthropod: \_\_\_\_\_
  - Amphibian: \_\_\_\_\_
  - Reptile: \_\_\_\_\_
  - Bird: \_\_\_\_\_
  - Mammal: \_\_\_\_\_
- In humans, water intake generally equals water losses over the course of a day. During a 24 hour period a student obtained 2500 mL of water from eating, drinking, and metabolism. They produced 1500 mL of urine in the same period.
  - Calculate the percentage of total water gains lost as urine: \_\_\_\_\_
  - Suggest how the remaining water is lost: \_\_\_\_\_  
 \_\_\_\_\_

# 163 ADH and Water Balance

**Key Idea:** Antidiuretic hormone (ADH) helps maintain water balance by regulating water absorption by the kidneys. The body regulates fluid balance in response to how much water is gained or lost. One mechanism by which fluid balance is maintained is by varying the volume of water reabsorbed by the **kidneys** and so also the volume and concentration of

**urine.** This involves a hormone called **antidiuretic hormone (ADH)**. Osmoreceptors in the hypothalamus monitor blood osmolarity (solute concentration) and send messages to the pituitary gland, which regulates the amount of ADH released. ADH promotes the reabsorption of water from the kidney tubules and collecting ducts, regulating urine volume.



1. What effect does ADH have on the kidneys? \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_
2. How do negative feedback mechanisms operate to regulate blood volume and urine output: \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_
3. Predict whether a high fluid intake would increase or decrease ADH production: \_\_\_\_\_
4. (a) Diabetes insipidus is a type of diabetes caused by the a lack of ADH. Based on what you know about the role of ADH in kidney function, describe the symptoms of this disease:  
 \_\_\_\_\_  
 \_\_\_\_\_  
 (b) How would diabetes insipidus be treated? \_\_\_\_\_  
 \_\_\_\_\_

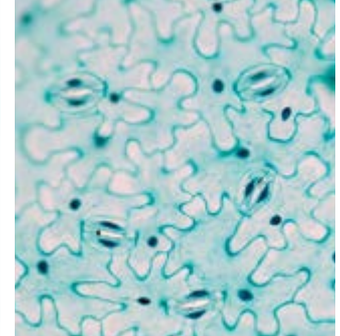
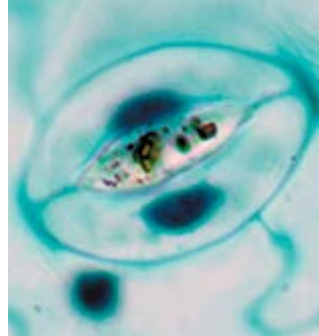
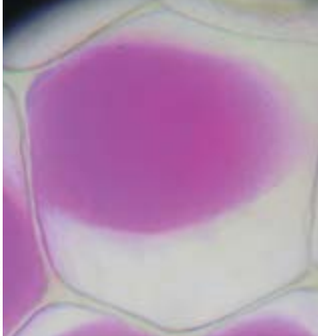


# 164 Osmoregulation in Plants

**Key Idea:** Plants in a range of habitats show a variety of adaptations for maintaining water balance. The adaptations and mechanisms plants use to maintain water balance are in part related to the environment in which they grow. Plants growing in what could be called average conditions (moderate to warm, humid environments) show

generalised adaptations for water conservation and are called **mesophytes** (meso = middle). In contrast, plants growing in extreme environments (aquatic, semi-aquatic, dry, or salty environments) have adaptations to survive these conditions. Many of their adaptations help to minimise water loss.

## Structures that help osmoregulation



Recall that a large part of the plant cell is taken up by the central vacuole (pink above). The vacuole plays an important role in maintaining turgor pressure and maintaining the shape of the cell. If too much water is lost from the cell the vacuole shrinks and the plant wilts as its cells lose their structure.

Leaves have a waxy cuticle that covers the epidermis and limits evaporation from the leaf surface. Plants adapted to dry environments (**xerophytes**) have thicker cuticles than those that are not (e.g. mesophytes). The cuticle is often thicker on the top of the leaf in mesophytes but more even in xerophytes.

Recall that **stomata** allow CO<sub>2</sub> to enter the leaf but this also allows water to escape. Mesophytes tend to have more stomata on the leaf undersides which reduces water loss via transpiration without restricting the entry of CO<sub>2</sub>. In times of water stress (drought) stomata are closed to reduce water loss.

Recall the role of turgor changes in guard cells and the opening and closing of stomata. The hormone **abscisic acid** (ABA) is synthesised in response to water stress and regulates stomatal closure. When water is low, ABA levels increase and K<sup>+</sup> and Cl<sup>-</sup> leave the guard cells. Water follows by **osmosis** and the guard cells flop together, closing the pore.

## Leaf and root structure of hydrophytes

- ▶ **Hydrophytes** are plants adapted to aquatic or semi-aquatic environments. Because of their watery environment they do not need to prevent water loss.
- ▶ The stomata are present in high numbers on the upper surface of the leaf only. The waxy cuticle on the leaf is designed to repel water off the leaf rather than to prevent drying out.
- ▶ The emergent leaves of hydrophytes are broad to increase the area for photosynthesis and also to aid flotation. Submerged leaves tend to be thin and divided. Any water lost from their broad surface by transpiration or evaporation is easily replaced.
- ▶ The root system of hydrophytes is much smaller than those of mesophytes. This is because its main function is to anchor the plant rather than a dual role of also obtaining water from the soil.



1. Explain why mesophytes do not show many adaptations for preventing water loss: \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_
2. Contrast the differences in the position of stomata in mesophytes and hydrophytes giving reasons for the differences.  
 \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_
3. How do plants reduce water loss in times of drought? \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_



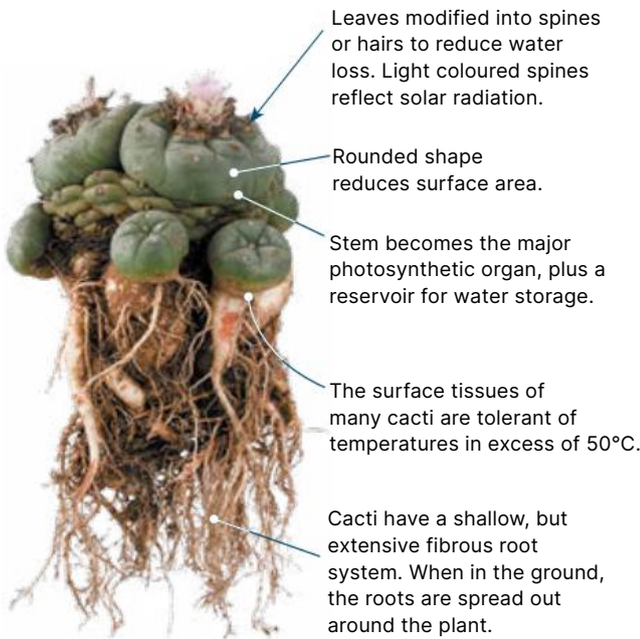
## Adaptations of xerophytes

Plants adapted to dry conditions are called xerophytes. Most xerophytes are found in deserts and semi-arid areas, but they may be found in humid environments, provided that their roots are in dry micro-environments (e.g. the roots of epiphytic plants that grow on tree trunks or branches).

- ▶ Xerophytes, such as cacti and Australia's sclerophyll plants, have a number of adaptations (called xeromorphic adaptations) that allow them to conserve water and survive in dry environments. These adaptations include small, hard leaves (sclerophylls), an epidermis with a thick cuticle, sunken stomata, succulence, and permanent or temporary absence of leaves.
- ▶ Many xerophytes have a succulent morphology. Their stems are often thickened and retain a large amount of water in the tissues, e.g. aloe.
- ▶ Many xerophytes have a low surface area to volume ratio, reducing the amount of water lost through transpiration.

### Adaptations in cacti

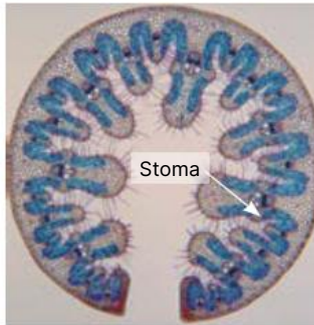
Desert plants, such as cacti (below), must cope with low or sporadic rainfall and high transpiration rates.



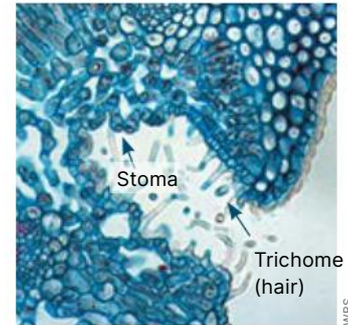
Acacia trees have deep root systems, which allows them to draw water from sources deep underground.



An outer surface coated in fine hairs traps air close to the surface and reduces transpiration.



Coastal grasses, e.g. marram grass (above), have curled leaves. Stomata are sunken in pits, creating a moist microclimate around the pore, which reduces transpiration rate.



Oleander has a thick multi-layered epidermis and the stomata are sunken in trichome-filled pits on the leaf underside which restrict water loss.

4. What is a xeromorphic adaptation? \_\_\_\_\_
5. Describe three xeromorphic adaptations of plants that reduce water loss:
  - (a) \_\_\_\_\_
  - (b) \_\_\_\_\_
  - (c) \_\_\_\_\_
6. How do sunken stomata reduce water losses via transpiration? \_\_\_\_\_
7. How does a low surface area to volume ratio in a plant such as a cactus reduce water loss? \_\_\_\_\_

## Adaptations of halophytes

- ▶ **Halophytes** are plants adapted to growing in saline (salty) environments. Mangroves are halophytes and show many of their typical adaptations. They are shrubs or small trees and grow in water-logged substrates in estuarine environments. Not only is their growth environment high in salt, but the salinity level fluctuates with tidal flows. Evaporation of water at low tide can greatly increase the salt levels compared to when the tide is in.
- ▶ The high salt environment would kill most other kinds of plants because high extracellular salt levels cause water to leave the cells. Mangroves overcome this by storing salt in their cell vacuoles and maintaining a high concentration of solutes in the cell cytoplasm. This reverses the osmotic gradient and maintains the transpiration stream.
- ▶ Australia has 45 species of mangroves. They are found in a range of habitats including tropical, subtropical, and sheltered temperate coastal rivers, as well as estuaries, bays, and marine shorelines. They grow in the upper part of the intertidal zone, but may extend further inland to form salt marshes and coastal wetland communities. Almost half of Australia's mangrove forests are located in Queensland.



BoundaryRider cc: 3.0

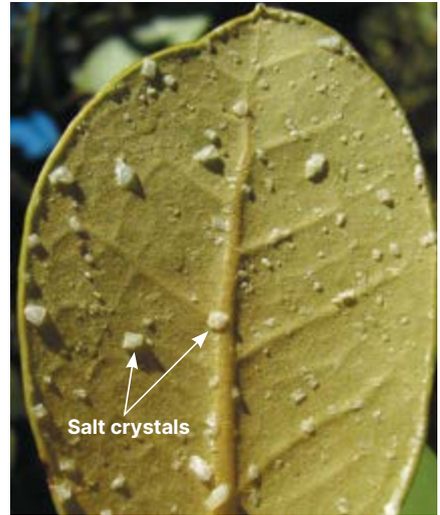
Mangroves in Queensland



RA



RA



Mangroves have specialised "breathing roots" called pneumatophores. They protrude above the surface of the mud and allow the mangrove to obtain oxygen even when the tide is in. A waxy coating of suberin on the root cells excludes 97% of salt from the water taken up by the roots.

Mangrove leaves are adapted for conserving water. They are covered in a thick waxy cuticle or dense hairs to reduce water loss. Sunken stomata limit water loss via transpiration. Leaves may also store water or orientated to minimise water loss in the hottest part of the day.

Some species of mangroves can secrete salt through salt glands in the surface layer of the leaves. This active transport process requires energy expenditure. Other mangrove species store salt in older leaves before they fall from the tree, taking the salt with them.

8. Describe a physiological problem associated with living in a high salt environment:

---



---



---

9. Describe three methods by which various mangrove species solve the problem of a high salt environment:

(a) \_\_\_\_\_

\_\_\_\_\_

(b) \_\_\_\_\_

\_\_\_\_\_

(c) \_\_\_\_\_

\_\_\_\_\_

# 165 Investigating Stomatal Density

**Key Idea:** The density and distribution of leaf stomata in different plant species are related to the rate of water loss. Different plant species have different leaf shapes and structures and these can be correlated with the environment

in which they are found. Comparing the leaf area and stomatal density of different plant species helps to explain observed differences in transpiration rate but factors in the environment, such as shading and wind, are also important.

## Plant species show different leaf shapes and structures associated with their environments

### Aloe (agave)

A succulent



Tropical species with thick, fleshy leaves. Physiology allows it to fix CO<sub>2</sub> during the night and keep stomata closed during the day.

### Pine

A conifer



Temperate species with thin, needle like leaves and a thick waxy leaf cuticle. Stomata are sunken into pits.

### Eucalyptus

An Australian gum tree



Sub-tropical drought tolerant species with a deep root systems and waxy leaves that hang downwards.

### Sunflower

A perennial dicot with large leaves



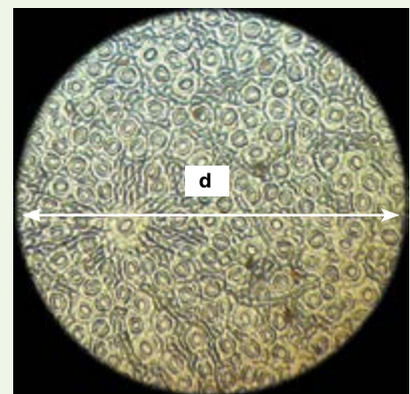
Widespread cultivated North American dicot with a showy flower head and very large soft leaves.



## Investigation 12.1 Comparing stomatal density

See appendix for equipment list.

1. Your teacher will have up to four leaf types from four dicot plants adapted to different environments, or you may need to obtain samples of your own.
2. The number of stomata per mm<sup>2</sup> on the surface of a leaf can be determined by counting the stomata visible under a microscope. Use clear nail varnish to paint over the lower surface of a leaf. Leave it to dry. This creates a layer with impressions of the leaf surface.
3. Carefully peel off the nail varnish layer and place on a clean microscope slide.
4. Calculate the diameter of the area viewable under a microscope using the field number divided by the magnification of the objective lens (for example if the objective lens magnification is 40, and the field number 18, then  $18 \div 40 = 0.45$  mm diameter. The area viewable is then  $\pi r^2$ ).
5. You could also use a micrometer to measure the diameter of the field of view or use a thin clear ruler.
6. Place the slide with the nail varnish layer on it under the microscope and count the number of stomata you see. If there are too many stomata then count one quarter of the field of view and multiply by four. Do this in several places. Enter your results in the table and calculate a mean.
7. You should also take note of where the stomata are on the leaf (are they scattered randomly or in specific places?)
8. Repeat on the upper surface of the leaf.
9. Repeat for the other leaf types.



$$\begin{aligned} \text{Area} &= \pi \times (d \div 2)^2 \\ &= \pi r^2 \end{aligned}$$



	Number of stomata per mm <sup>2</sup> lower surface					Number of stomata per mm <sup>2</sup> upper surface				
	Count number						Count number			
Plant name/type	1	2	3	4	Mean	1	2	3	4	Mean

1. (a) Write an aim for the investigation: \_\_\_\_\_

\_\_\_\_\_

(b) Write a hypothesis for the investigation: \_\_\_\_\_

\_\_\_\_\_

2. Complete the table above:

3. (a) Which plant has the highest stomatal density? \_\_\_\_\_

(b) Which plant has the lowest stomatal density? \_\_\_\_\_

4. (a) Is there a relationship between the number of stomata per mm<sup>2</sup> and the type of leaf or plant?

\_\_\_\_\_

(b) Explain your answer: \_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_

5. (a) Where are the majority of stomata located in a typical dicot leaf? \_\_\_\_\_

(b) Suggest why this might be the case: \_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_

6. Explain your results in terms of the environment the plants are adapted for and the need to regulate water loss:

\_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_

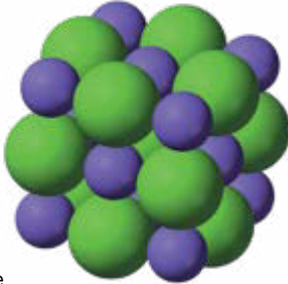
\_\_\_\_\_

# 166 Salt Tolerance in Plants

**Key Idea:** Different species have differing tolerances to abiotic factors. Levels outside an organism's tolerance limits can reduce its chances of survival. Salt tolerance in plant species can be measured experimentally.

Tolerance limit is the ability to live within a certain range of abiotic factors (such as temperature or rainfall). For plants,

salt tolerance is important because soil salinity affects plant growth and productivity. Above certain soil salinities, plants will not germinate or grow. In many regions of Australia, soil salinity is increasing (salinisation) lowering productivity. Human activities, such as removal of natural vegetation and poor irrigation management, contribute to salinisation.



Sodium chloride

Sodium chloride (NaCl) is an important contributor to soil salinisation. Increased soil salt concentrations have several negative effects on plants:

- ▶ High salt increases osmotic stress and decreases the ability of a plant to take up water.
- ▶ Large uptakes of  $\text{Na}^+$  and  $\text{Cl}^-$  have a negative effect on growth by impairing metabolic processes and decreasing photosynthetic efficiency.

Plants can cope with NaCl to varying degrees. Some excrete excess salt onto their leaves (e.g. mangroves and salt grass) and others compartmentalise the salts (often in vacuoles).

## The aim

Determine how temperature and salt levels influence the growth of salt grass (*Distichlis spicata*).

Salt grass (*Distichlis spicata*)



Matt Lavin CC 2.0

Salt grass (*Distichlis spicata*) is a native American plant found in a number of habitats including coastlines, desert scrub and marshes. It is very salt tolerant and capable of growing in very salty soils. Excess salt is excreted from its tissues onto the leaf surfaces.

## The method

- ▶ Salt grass seeds were germinated in quartz sand and transplanted into their solutions of salt water when the fourth leaf appeared.
- ▶ The salt water concentrations were prepared by diluting sea water to obtain the following concentrations: 306, 612, 1834, and 2448 ppm.
- ▶ A nutrient enrichment solution was added to each sea water solution after dilution. The pH was adjusted to 5.7-6.0.
- ▶ Samples were prepared in duplicate.
- ▶ A control was grown in nutrient solution only.
- ▶ The plants were raised in greenhouses at either 12.7°C or 21.1°C.
- ▶ Material was harvested (as two cuttings) dried and weighed. The data are presented in the table below.

## The results

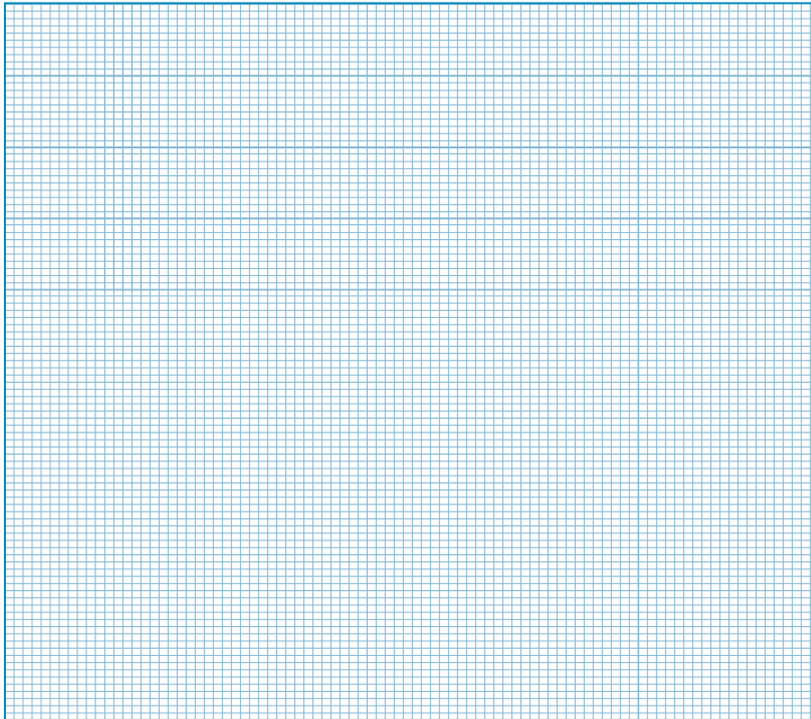
**Table 1: Effect of salt water concentration and temperature on salt grass growth**

Concentration of sea water (ppm)	Weight of dry matter Average of 2 cultures (first cutting) (g)		Weight of dry matter Average of 2 cultures (second cutting) (g)		Total weight of dry matter (first cutting + second cutting) (g)	
	12.7°C	21.1°C	12.7°C	21.1°C	12.7°C	21.1°C
0 ppm + complete nutrient solution	47.0	21.2	49.0	25.3	96.0	46.5
306 ppm + dilute nutrient solution	14.4	8.3	21.2	12.5		
612 ppm + dilute nutrient solution	14.3	7.7	20.0	11.1		
1834 ppm + dilute nutrient solution	12.7	7.7	13.1	9.5		
2448 ppm + dilute nutrient solution	11.3	3.4	10.5	3.6		

Data: Ahi and Powers (1938) Salt tolerance of plants at various temperatures. Plant Physiol. 13 : 767-789.

1. Complete the table above by calculating the total weight of dry matter for each temperature and salt concentration. The first one has been done for you:
2. (a) Plot the total weight of dry matter at each temperature as a line graph on the grid (following page):





(b) Describe how salt water concentration and temperature affect salt grass growth: \_\_\_\_\_

---



---



---



---

3. How does salt grass remove excess salts and suggest how might this help it to survive in saline conditions?

---



---



---

4. Wheat and rice are important food crops. The graph on the right shows their salt tolerance compared to two salt adapted species. Based on the data for salt grass, and assuming that salinisation will continue to be a problem, how might increasing soil salinity affect productivity of these important food crops in Australia?

---



---



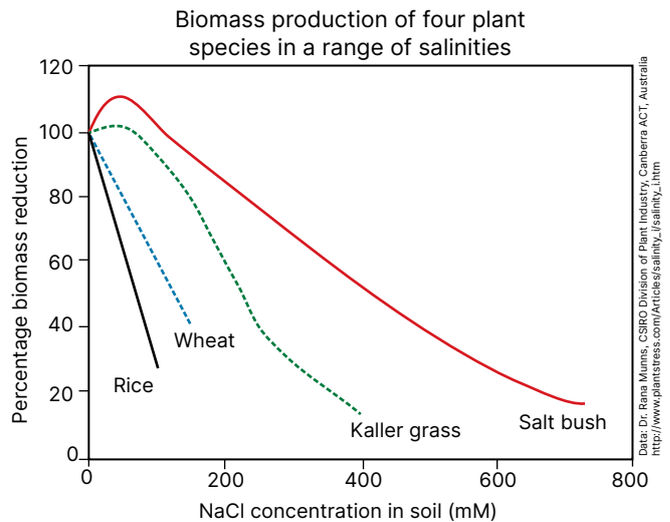
---



---



---



5. How could the development of salt tolerant plants (e.g. through genetic modification) benefit Australian farmers?

---

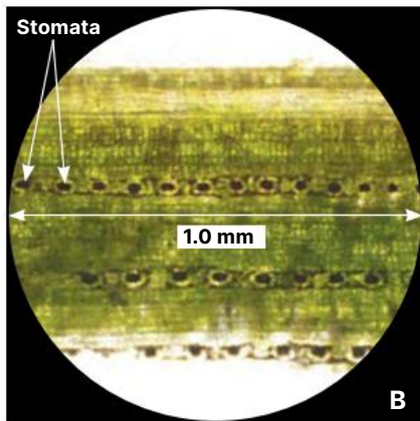
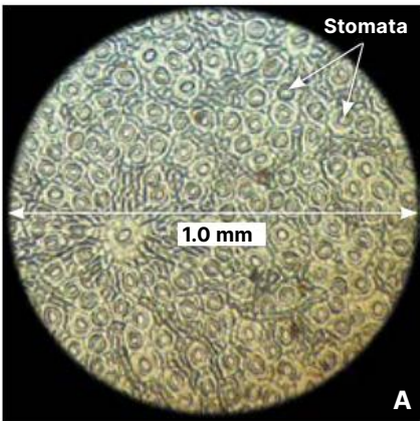


---

# 167 Did You Get It?

- The following examples give ways in which organisms osmoregulate. Decide whether they are examples of structural (morphological), behavioural or physiological adaptations:
  - The secretion of ADH in response to low blood volume: \_\_\_\_\_
  - The stomata on xerophyte leaves are sunken: \_\_\_\_\_
  - A freshwater fish takes up ions across the gill surface: \_\_\_\_\_
  - The roots of mangrove trees are covered in a waxy substance to prevent salt uptake: \_\_\_\_\_
  - Camels metabolise stored fat in their hump to produce water: \_\_\_\_\_
  - A goanna seeks shade during the hottest part of the day: \_\_\_\_\_
  - Secretion of salt from salt glands in a mangrove: \_\_\_\_\_

2. The light micrographs below show the surface of leaves of plants adapted to different environments. Calculate the number of stomata per  $\text{mm}^2$  for each leaf:



- Leaf A: \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_
- Leaf B: \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_

(c) Which of these plants (A or B) is more likely to be adapted to dry environments?  
 \_\_\_\_\_

3. The graph below shows the volume of urine collected from a subject after drinking  $1000 \text{ cm}^3$  of distilled water. The subject's urine was collected at 25 minute intervals over a number of hours.

(a) Explain the difference in the volume of urine collected at 25 minutes and 50 minutes:

\_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_

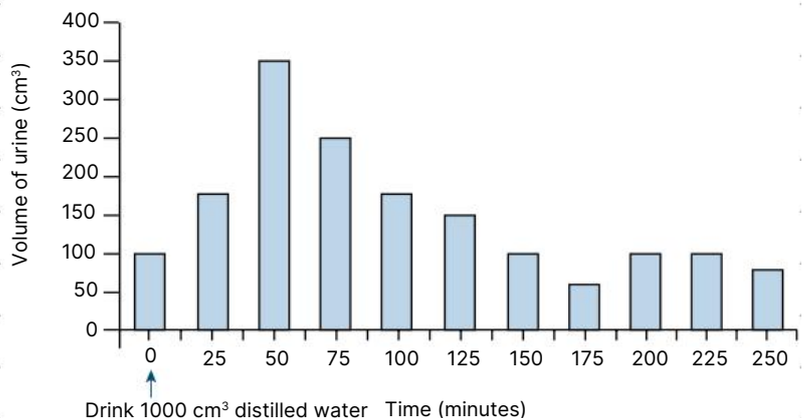
\_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_



(b) Explain how ADH production, urine concentration and production, and kidney permeability would be affected by drinking  $1000 \text{ cm}^3$  of water:

\_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_

# 168 Synoptic Question: Unit 2, Topic 1

1. (a) What type of neuron is shown in the diagram right? Justify your answer:

---

---

---

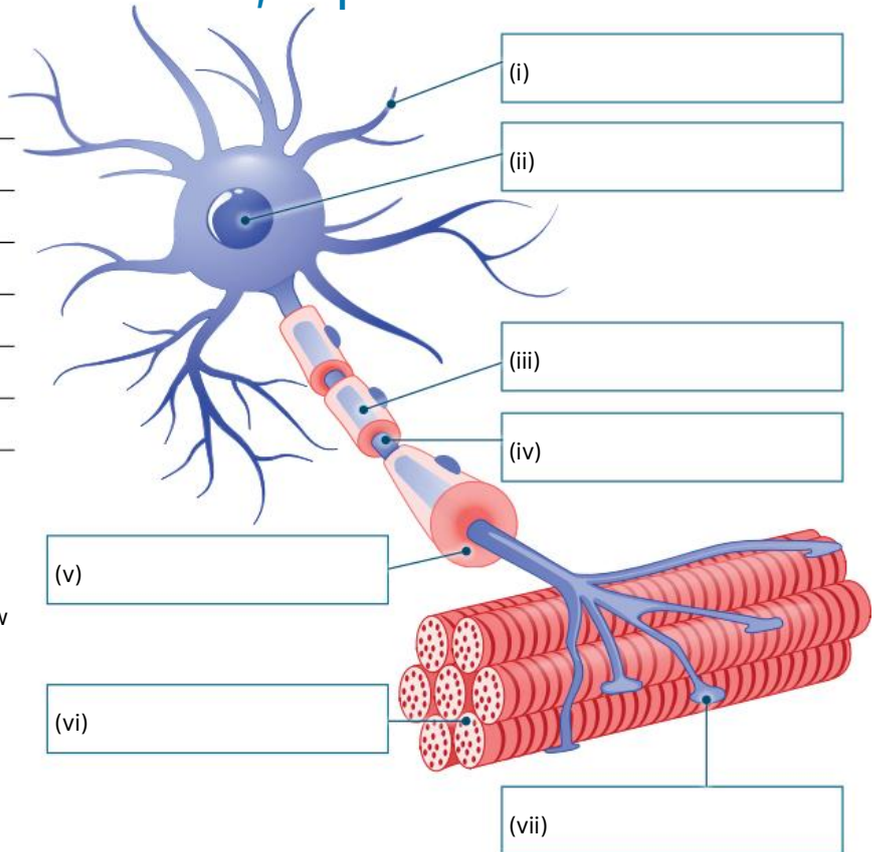
---

---

---

---

---



(b) Use the following word list to label the diagram: *Soma (cell body), myelin sheath, node of Ranvier, muscle fibre, axon, dendrites, synaptic knob.*

(c) Draw an arrow on the diagram to show the direction a nerve impulse would travel along this neuron:

(v)

(vi)

(vii)

2. Cells use chemical messengers to communicate and to respond to their environment. The diagram below shows a form of cell signalling.

(a) Describe the basis of cell signalling in terms of a receptor, cell signalling molecule, and a target cell:

---

---

---

---

---

---

---

---

(b) Why is it important that not all cells respond to every cell signal?

---

---

---

---

---

---

---

---

(c) How would a cell increase its response to molecule A?

---

---

---

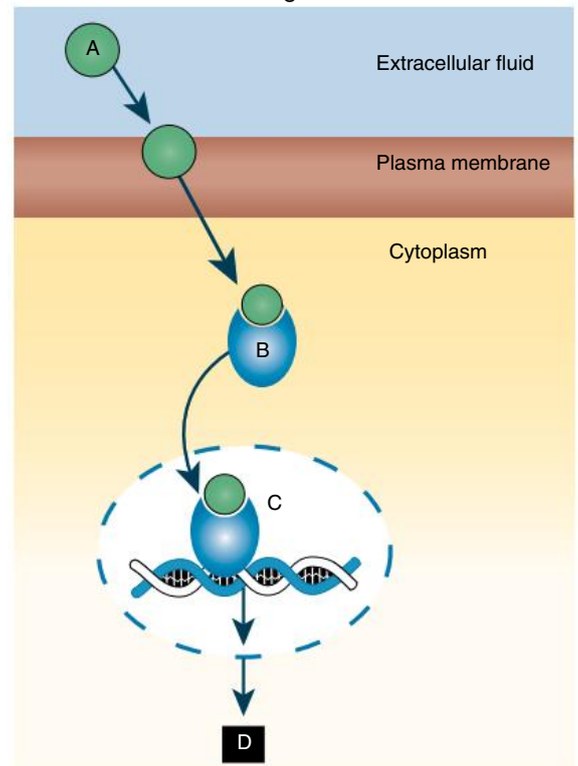
---

---

---

---

---



3. Describe how a hormone relays its messages from an endocrine gland to the receptor cells:

---

---

---

---

---

---

---

---

4. (a) Using thermoregulation as an example, explain why maintaining homeostasis is crucial for ensuring metabolic reactions can proceed:

---



---



---



---

- (b) Most mammals use fur to help retain body heat. Humans have very little body hair. Give a structural and a behavioural way in which humans can retain body heat:

---



---

- (c) An experiment was performed with three volunteers with the same body shape and dimensions. Volunteer A was given good insulated clothing, including an insulating knitted hat, volunteer B was also given well insulated clothing but no insulated hat, volunteer C wore only a light shirt and pants. The three stood in a freezer room ( $-18^{\circ}\text{C}$ ) for five minutes while their core body temperature was monitored. Explain what will happen to the body temperature of these three volunteers and explain why:

---



---



---



---



---



---



---



---



---



---



5. Plants living in arid or saline environments have many adaptations to conserve water. Explain how adaptations in xerophytes and halophytes help regulate water balance:

---



---



---



---



---



---



---



---



---



---



---



---



---



---



---



---



---



---



---



---



---



---



# Infectious Disease

## Key Terms

- adherence factors
- bacteria
- bacterial disease
- capsid
- cellular (pathogen)
- disease
- fungal disease
- fungus (fungi)
- germ theory
- host
- infectious disease
- invasion factors
- Koch (Robert)
- non-cellular (pathogen)
- non-infectious disease
- Pasteur (Louis)
- pathogen
- pathogenesis
- prion
- Semmelweis (Ingaz)
- toxins
- vector
- viral disease
- virus

## Key Concepts

- ▶ Infectious diseases are caused by pathogens, while non-infectious diseases result from genetic, environmental, or lifestyle factors
- ▶ Pathogens can be classified as non-cellular such as prions, misfolded proteins, and viruses; or cellular such as bacteria, fungi, and protists.
- ▶ Pathogens have adaptations to enable them to gain entry to hosts.

## Pathogens and Disease

### Activity Number

<input type="checkbox"/> 1	Understand what is meant by disease. Distinguish between infectious diseases and non-infectious diseases and give examples.	169
<input type="checkbox"/> 2	Describe the nature of pathogens and explain their role in infectious disease. Identify and characterise types of pathogens, including viruses, prions, bacteria, fungi, protists and parasites.	170
<input type="checkbox"/> 3	Explain what is meant by pathogenesis. Identify and describe virulence factors (factors produced by pathogen that add to their effectiveness) including adherence factors, invasion factors, capsules, toxins, and life cycle changes.	170
<input type="checkbox"/> 4	<b>SHE:</b> Explore the historical understanding of germ theory, disease and its transmission, including the work of both Koch and Semmelweis.	169

## Types of Pathogens

### Science understanding

<input type="checkbox"/> 5	Describe features of bacterial pathogens, including relevant virulence factors and adaptations specific to the pathogen involved. Diseases include tuberculosis, cholera, tetanus, <i>Salmonella</i> poisoning, and crown gall in plants.	171
<input type="checkbox"/> 6	Describe features of fungal pathogens, including relevant virulence factors and adaptations specific to the pathogen involved. Diseases include amphibian chytrid fungus disease.	172
<input type="checkbox"/> 7	Describe features of protistan pathogens, including relevant virulence factors and adaptations specific to the pathogen involved. Diseases include malaria (protozoan) and <i>Phytophthora</i> dieback (oomycetes or water moulds, not to be confused with fungal moulds).	173
<input type="checkbox"/> 8	Describe features of viral pathogens, including relevant virulence factors and adaptations specific to the pathogen involved. Diseases include HIV/AIDS, influenza, Ross River virus, measles, and Ebola.	174-175
<input type="checkbox"/> 9	Describe features of prion pathogens, including relevant virulence factors and adaptations specific to the pathogen involved. Diseases include variant Creutzfeldt–Jakob disease (vCJD), kuru, BSE, and scrapie.	176

# 169 Infection and Disease

**Key Idea:** The term disease encompasses any disorder in the structure or function of an animal or plant. It may be caused by an infectious agent or a spontaneous change in the body. **Disease** can be divided into two main types: **infectious disease** and **non-infectious disease**. Infectious disease

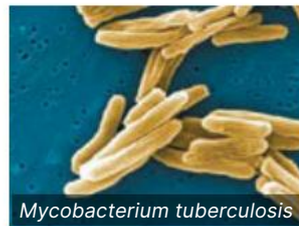
## Infectious disease

The method by which disease develops is called **pathogenesis**. The pathogens causing infectious diseases may be **cellular** (e.g. bacteria and **protists**) or **non cellular** (e.g. viruses and **prions**). The adaptations of the pathogens can influence the attachment (**adherence factors**) and entry (**invasion factors**) to the host. Pathogens produce molecules (called virulence factors) that make them more effective at infecting their **host**. These virulence factors are commonly antigenic (capable of causing an immune reaction in the host).

## Cellular pathogens

### Bacterial pathogens

Pathogenic bacteria can be transmitted through food, water, air, or by direct contact. Although bacteria have historically caused widespread and devastating diseases, the discovery and use of antibiotics and aseptic techniques have significantly reduced these.



*Mycobacterium tuberculosis*

CC0: Janice Haney Cair

### Fungal pathogens

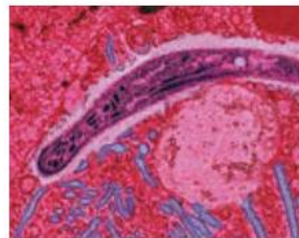
Pathogenic **fungi** are more common in plants than in animals. They spread by spores and the infections they cause are generally chronic (long-lasting) infections because fungi grow relatively slowly.



Fungal infection of toenails

### Protistan pathogens

**Protists** are a large and diverse group of eukaryotes. A number of species are significant pathogens of animals or plants. Pathogenic protists have very complex life cycles, often involving a number of different hosts and several different life stages.



## Non-cellular pathogens

### Viral pathogens

A virus is a highly infectious pathogen that infects living cells (including bacterial cells) and uses the cell's metabolic machinery to replicate. Viruses consist of a protein envelope surrounding the nuclear material that can be either DNA or RNA.

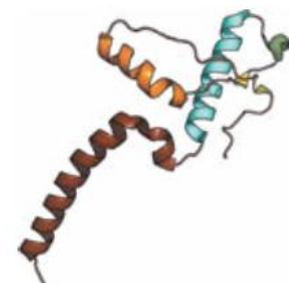


Ebola virus

NIAID CC-2.0

### Prions

Prions are misfolded infectious proteins that have pathogenic properties. They contain no genetic material so do not replicate in the usual way. Instead, an infectious prion binds to a normal protein and causes it to change shape and become infectious. Prions cause degenerative nervous diseases in mammals including scrapie in sheep, BSE in cattle, and kuru in humans.



is caused by infectious agents called **pathogens**, which include **bacteria** and **viruses**. Non-infectious disease is caused by damage or changes to the body which may occur because of genetic defects which may be congenital (e.g. type 1 diabetes) or caused by environment effects.

## Non-infectious disease

Non infectious diseases are not transmitted from person to person and are often caused by genetic or lifestyle factors.

### Environmental diseases

These are caused by external factors such as a lack of sunlight or vitamins. These diseases also include diseases caused by lifestyle choices, such as drinking too much alcohol or smoking. Environmental diseases include cancer, diseases of affluence and various social diseases (below):



Lung cancer

Emmanuelm cc3.0

### Cancer

Cancer is caused by mutations in the genes that keep cell growth and division in check. Cancer causes the cells to grow and divide continuously, causing tumors and affecting organ function.

### Western Diseases

This is a relatively new term that encompasses a wide range of environmental diseases more commonly seen in relatively wealthy (and often Western) societies. They include obesity, mental health issues, and cardiovascular diseases. They are often said to contrast so-called "poverty diseases", such as severe malnutrition.



### Inherited diseases

These are genetic diseases caused by mutations (DNA errors) carried by the parents and passed to the offspring. They may also occur spontaneously during the development of the sperm or egg. Inherited diseases include Huntington's disease and cystic fibrosis (right).



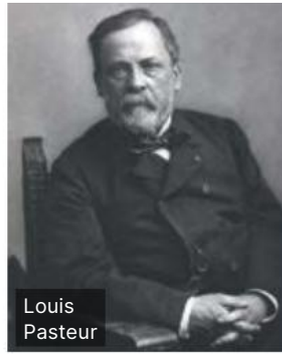
### Autoimmune diseases

Occasionally the immune system of the body begins to attack parts of its own body. These diseases include type 1 diabetes and multiple sclerosis. Type 1 diabetics must inject insulin because their insulin-producing cells have been destroyed (right).

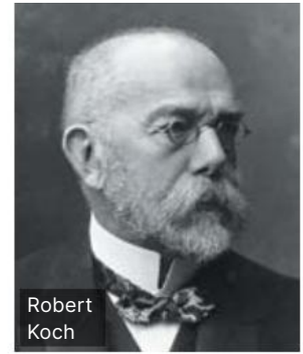


## Life arises from life

- ▶ The experiments of Louis **Pasteur** and Robert **Koch** contributed to our understanding of microbiology and infectious disease.
- ▶ For a long time people thought that new life could spontaneously evolve from non-living matter (spontaneous generation) and that clouds of poisonous gas caused disease (Miasma Theory). The work of two scientists, Louis Pasteur (French) and Robert Koch (German), methodically disproved both of these theories. Their work contributed significantly to what we know about microbiology and the spread of infectious disease today.



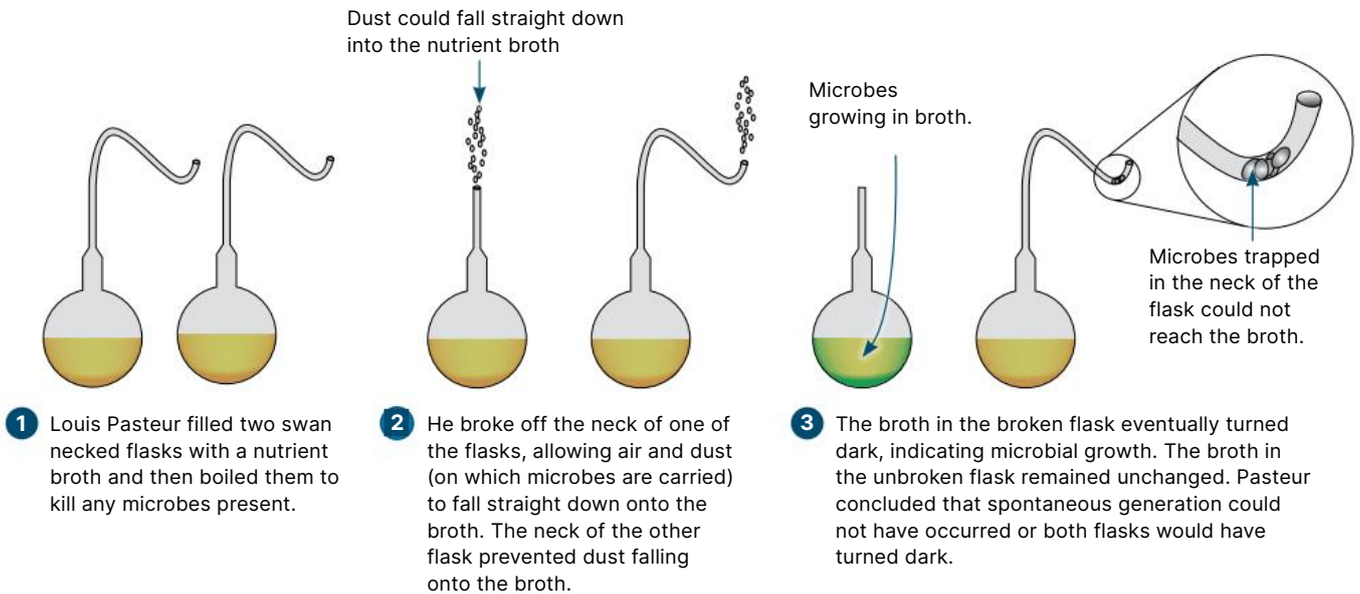
Louis Pasteur



Robert Koch

### Pasteur's swan neck flask experiment

Until the 1800s it was generally believed that new life could arise spontaneously from non-living matter. For example, when a piece of meat was left on a bench, maggots appeared a day or two later. Under spontaneous generation theory the maggots had spontaneously generated from the components of the meat. Louis Pasteur disproved this theory when he carried out his very simple swan neck flask experiments (below).



1. A student set up an experiment to replicate Pasteur's experiment. Nutrient broth was added to two test tubes and the tubes were sealed. Both tubes were heated for several minutes over a Bunsen burner. After cooling, tube 1 was uncovered, and tube 2 was left covered. The students observations are described in the table.

(a) Did the student collect qualitative or quantitative data?

\_\_\_\_\_

(b) Explain your reason for your answer in 2(a):

\_\_\_\_\_

\_\_\_\_\_

Test tube	Observation of the broth at day 1	Observation of the broth at day 10
1 uncovered	Clear	Cloudy
2 covered	Clear	Cloudy

2. Suggest a reason why the student's results were different from Pasteur's results: \_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_

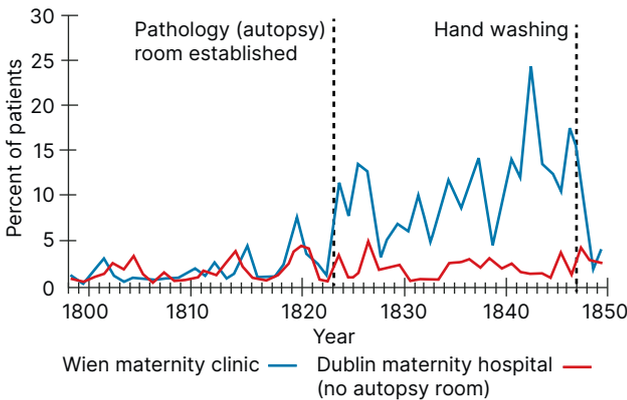
\_\_\_\_\_

## Germ theory and disease transmission

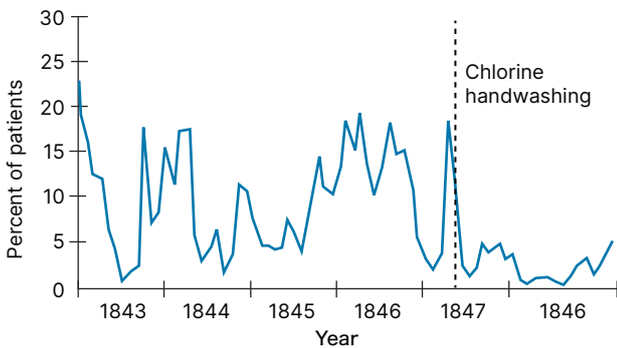
**Germ theory**, the idea that infections are caused by infectious microscopic agents, was developed around the middle of the nineteenth century. Since the development of the modern scientific method, based on controlled experiment and observation, we are now able to identify and treat a vast range of diseases.

Ingaz **Semmelweis** was an assistant professor at the Vienna General Hospital in which there were two maternity clinics. The first clinic taught medical students and the second taught midwives. Semmelweis noticed that the first clinic always had higher mortality rates in the mothers from fever after childbirth (postpartum) than the second clinic. He realised that the medical students were carrying some sort of infectious agent from the bodies they were dissecting in the autopsy room to the maternity ward, but that the midwives were not. He instigated a policy of hand washing with a solution of chlorinated lime. The result was an almost immediate substantial drop in mortality.

**Mortality by postpartum fever in Wien maternity clinic and Dublin maternity hospital**



**Mortality by postpartum fever**



**Robert Koch** showed that a specific disease was caused by a specific pathogenic (disease-causing) agent. He developed what are now known as Koch's postulates.

### Koch's postulates

**1**

Pathogenic microorganisms are isolated from a dead animal.

**2**

The microorganisms are injected into a healthy animal.

**3**

The disease is reproduced in the second animal. Microorganisms are isolated.

**4**

Isolated pathogenic microorganisms are identical to original pathogens.

Koch isolated bacteria from a diseased animal, then injected them into a healthy animal, causing it to exhibit identical symptoms to the first. This demonstrated that a specific infectious disease (e.g. anthrax) was caused by a specific microorganism (*Bacillus anthracis*). Koch used the procedure to identify the bacteria that caused anthrax and tuberculosis.

Koch's findings are summarised as Koch's postulates:

1. The same pathogen must be present in every case of the disease.
2. The pathogen must be isolated from the diseased host and grown in pure culture.
3. The pathogen from the pure culture must then cause the disease when it is inoculated into a healthy, susceptible animal.
4. The pathogen must be isolated from the inoculated animal and shown to be the original organism.

3. (a) What is the difference between infectious and non-infectious disease? \_\_\_\_\_  
 \_\_\_\_\_
- (b) What are the two main types of pathogens and give examples? \_\_\_\_\_  
 \_\_\_\_\_
4. What was the significance of Koch's contribution to germ theory? \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_
5. Describe the evidence supporting Semmelweis' theory that an infectious agent was responsible for postpartum fever:  
 \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_

# 170 Bacterial Diseases

**Key Idea:** Pathogenic bacteria are responsible for some of the world's most devastating diseases of plants and animals. Relatively few of the world's bacterial species cause **disease**. Those that do (the pathogenic **bacteria**), have a range of adaptations that enable them to penetrate the defences of a **host** and cause infection (below). **Bacterial diseases** are

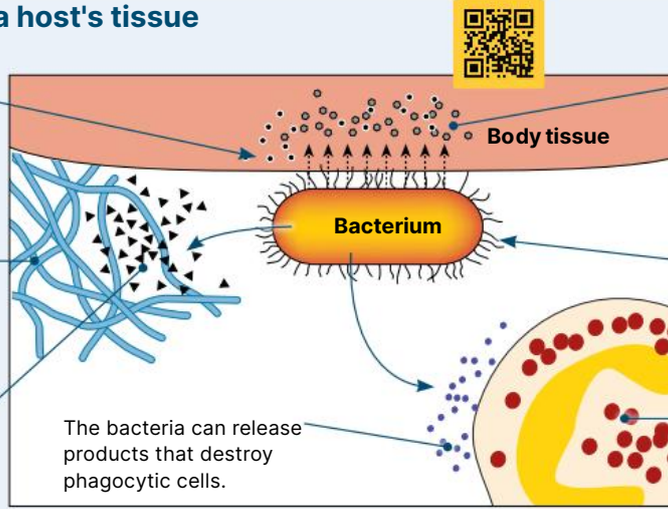
commonly transmitted through food, water, air, or by direct contact. The natural source of infection of a disease varies from species to species, ranging from humans and other organisms, to sewage or contaminated water. Much of our control of bacterial disease is achieved through identifying reservoirs of infection and limiting the routes of transmission.

## How bacteria invade a host's tissue

**Toxins:** Bacterial **toxins** can act locally to promote bacterial invasion (e.g. the enzymes that degrade collagen), or they may have cytotoxic activity and destroy cells directly.

**Fibrin:** Fibrous threads of protein are deposited when blood clots. This action by the host effectively limits the movement of **pathogens** in infected areas.

Enzymes are released that break down fibrin, allowing the bacteria greater freedom of movement.

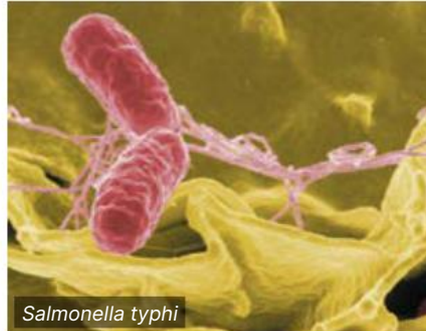


The bacterium releases enzymes that degrade the connective tissue of the host, allowing the spread of infection.

**Fimbriae** are fine, threadlike extensions from the bacterial cell. The bacteria use them to attach to the mucous membranes (**adherence factors**) and directly attack the host tissues.

**Phagocyte:** These white blood cells are very effective in identifying and destroying foreign cells such as pathogens.

## Methods of bacterial transmission



**Foodborne bacterial diseases:** Bacterial foodborne illnesses are caused by consuming food or beverages contaminated with bacteria or their toxins. Examples include Salmonella food poisoning and Campylobacter infection. Symptoms of bacterial food poisoning include fever, abdominal cramps, and diarrhoea. Some are associated with consuming raw or undercooked poultry.

**Waterborne bacterial diseases:** Waterborne bacterial pathogens are responsible for a number of serious diarrhoeal illnesses, including typhoid and cholera. Transmission of these diseases is usually through faecal contamination of drinking water. The fever and diarrhoea associated with such diseases is responsible for hundreds of thousands of deaths annually in countries with poor sanitation.

**Airborne bacterial diseases:** Airborne pathogens are transmitted on dust particles or droplets when people cough, sneeze, or exhale. Vaccination against certain airborne bacteria has been highly successful. Whooping cough (above) is a potentially fatal respiratory disease caused by the bacterial pertussis toxin. The prevalence of this disease has declined dramatically following the introduction of immunisation programmes.

1. Describe the specific adaptations of bacteria that contribute to their ability to cause disease:

---



---



---



---



---

2. What are the most common ways in which bacteria spread? \_\_\_\_\_

---



---



### Examples of bacterial disease

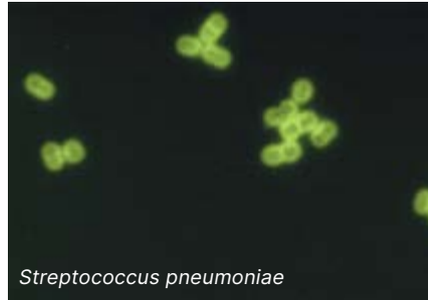
#### Tetanus

Tetanus is a condition characterised by prolonged, strong contractions of the skeletal muscles. It is caused by the toxin tetanospasmin, produced by the bacteria *Clostridium tetani* an anaerobic bacterium commonly found in the soil. Infection is usually through a puncture wound that becomes contaminated with soil or dirt.



#### Bacterial meningitis

Bacterial meningitis is most commonly caused by the bacterial pathogens *Neisseria meningitidis* and *Streptococcus pneumoniae* (below). The bacteria infect the membranes around the brain (the meninges) causing headaches, fever, rashes and sometimes death. The fatality rate is between 10%-20%.



#### Plant Crown Gall

Crown gall is a tumour-like growth in plants caused by the soil bacterium *Agrobacterium tumefaciens*. The gall is produced when the bacterium transfers a circular piece of DNA (the tumour-inducing or *Ti* plasmid) to the plant cell. *A. tumefaciens* is now commonly used in biotechnology to insert new genes into plants for genetic modification.



3. Why is immunisation often the best option for controlling airborne bacterial diseases? \_\_\_\_\_

---

---

---

4. Why is it not good practice to chop vegetables on the same chopping board as is used to prepare raw chicken?

---

---

---

5. Why are there often outbreaks (or risks of outbreak) of bacterial diseases such as typhoid and cholera after large scale natural disasters, such as large earthquakes or tsunamis?

---

---

---

---

6. How can simple measures, such as washing hands before eating, reduce the incidence of bacterial diseases?

---

---

7. Explain why *Agrobacterium tumefaciens* is of particular interest to scientists: \_\_\_\_\_

---

---

8. How is tetanus caused and what are the common ways of contracting the disease?

---

---

---

---

---

# 171 Fungal Diseases

**Key Idea:** Pathogenic fungi are rare in animals, but they can cause infections that are long lasting and difficult to treat. All **fungi** are heterotrophic, requiring organic compounds for energy and carbon. They may be parasitic or saprotrophic, obtaining nutrition by the extracellular digestion of living or dead organic matter. Very few fungi are **pathogenic** to animals, although thousands of fungal species are plant pathogens. They spread by spores and the infections they cause are generally chronic (long-lasting) infections because

fungi grow relatively slowly. **Fungal diseases** are categorised into three broad groups (below), the most common being superficial infections of the skin. Of great concern recently is the spread of a fungal pathogen in amphibian populations. Amphibian chytrid fungus **disease** has been linked to the dramatic decline in amphibian populations globally. Amphibians rely on their skin for osmoregulation and oxygen uptake, so they are particularly vulnerable to pathogens that compromise its integrity.

## Types of fungal infection



CDC

**Systemic Infections** are usually ones that occur deep inside the body, affecting internal organs, such as the lungs, bones, heart, and urinary tract. They often start in the lungs by inhalation of the spores and spread throughout the body. e.g. candidiasis in the kidney above.



**Cutaneous (superficial) infection:** Infection that affects the skin, hair, nails, genital organs, and inside of mouth. Contracted through contact with spores, e.g. trichosporosis infection of the toenails above. They are slow growing and difficult to treat.



CDC

**Subcutaneous Infection:** Rare infection of the fatty connective tissue beneath the skin. Contracted through direct implantation of the spores into the skin via a scratch or puncture wound, e.g. sporotrichosis (above) caused by the fungus *Sporothrix schenckii*.

## Chytridiomycosis

- ▶ Chytridiomycosis is a waterborne disease of amphibians caused by the fungi *Batrachochytrium dendrobatidis*. It disperses via motile spores called zoospores, which enter the host via the skin, although much of how new hosts are infected is still unknown. It can be fatal to infected frogs within 10-18 days.
- ▶ *Batrachochytrium dendrobatidis* is found to various parts of Australia, notably on the east coast, Adelaide, south-west Western Australia and the Kimberley region of WA.
- ▶ Chytridiomycosis has been implicated in the dramatic population declines of frog species including *Litoria nannotis* (waterfall frog), and *Litoria rheocola* (common mistfrog). It is also implicated in the extinction of at least four species of Australian frog.



Forrest, Brem CC 2.5. Riders of a Modern-Day Ark. Gewin V. PLoS Biology 6(1), e24. doi:10.1371/journal.pbio.0060024.

Frog killed by chytridiomycosis. Note the reddening of the skin which is characteristic of the disease.

1. Describe two features of fungal diseases: \_\_\_\_\_  
\_\_\_\_\_
2. Why is it often difficult to treat a fungal infection? \_\_\_\_\_  
\_\_\_\_\_  
\_\_\_\_\_
3. (a) Suggest one way in which chytridiomycosis causes death in infected frogs: \_\_\_\_\_  
(b) Suggest why amphibians are so vulnerable to chytridiomycosis: \_\_\_\_\_  
\_\_\_\_\_  
(c) Frogs are popular aquarium pets in many countries. How might this be contributing to the spread of the disease?  
\_\_\_\_\_  
\_\_\_\_\_

# Protistan Diseases

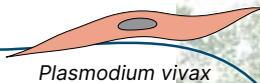
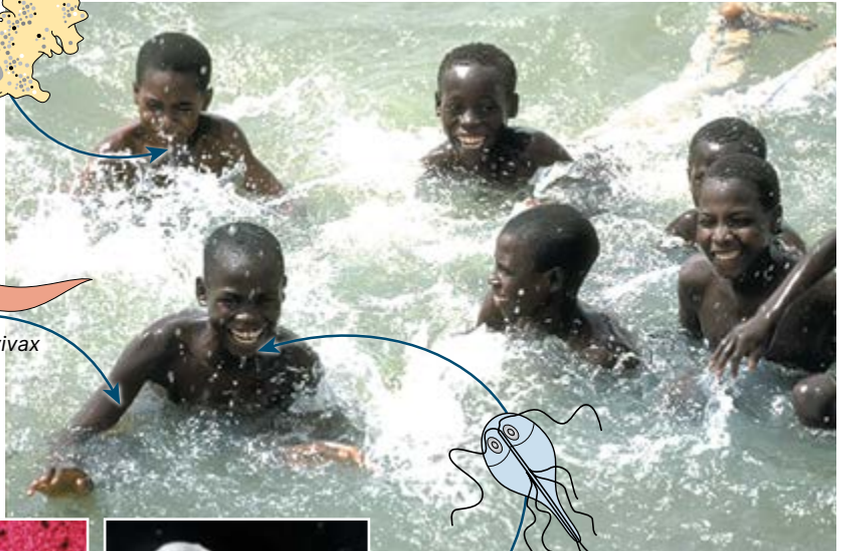
**Key Idea:** Protistsans are a large and extremely diverse group of eukaryotic organisms. A number of species are significant pathogens of animals or plants.

The **protists** are a group of unicellular or colonial eukaryotes. Most inhabit water or soil habitats and relatively few cause **disease**. However, a number of species are highly specialised **pathogens**. These include species of the parasitic genus

*Plasmodium*, which cause malaria in humans, and species of the oomycete genus *Phytophthora*, which cause devastating dieback and blight in a number of plant species. Pathogenic protists have very complex life cycles, often involving a number of different **hosts** and several different life stages. Both oomycetes and plasmodia, for example, have infective motile stages as well as resistant resting stages.

### Amoebae

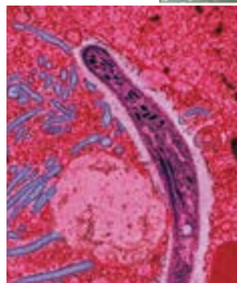
Amoebae move by extending projections of their cytoplasm. Several pathogenic amoebae infect humans and feed mainly on red blood cells. People become infected with the pathogen for amoebic microencephalitis while swimming in warm bodies of fresh water or hot springs, when the waterborne cysts pass across mucous membranes and infect blood, brain, and spinal cord. It's almost always fatal.



### Apicomplexa

These protozoans are not mobile and tend to be intracellular parasites. They use special enzymes to penetrate the host's tissues. They have complex life cycles involving transmission between several host species. Apicomplexans include *Plasmodium*, which is spread by mosquito **vectors** and causes malaria.

*Plasmodium* sporozoite stage moving through the cytoplasm of the intestinal epithelia.

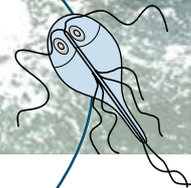


Ute Frevert, PloS



Giardia trophozoite, SEM

CDC



### Flagellates

Flagellates are usually spindle-shaped, with flagella projecting from the front end. Their whiplike motion pulls the cells through their environment. *Giardia* (left) is found in the small intestine of mammals. It is passed in the faeces and its life cycle alternates between an actively swimming trophozoite (left) and an infective, resistant cyst.

### Phytophthora dieback

Phytophthora dieback is caused by the soil-borne water mould *Phytophthora cinnamomi*. Although originally classified as a fungus, *Phytophthora* is now included in the Protista. Flagellated zoospores enter the plant near the growing tip of the roots where they germinate, produce fungal-like hyphae, and absorb carbohydrates. This eventually destroys the internal structure of the roots and causes the death of the plant.

*Phytophthora cinnamomi* is one of the world's most invasive plant pathogens. In Australia, it is responsible for the dieback of Eucalyptus trees especially in the Jarrah Forest bioregion of Western Australia. *Phytophthora* dieback can be treated with various fungicides including phosphite salts (e.g. calcium phosphite).

Edward L. Barnard, Florida Department of Agriculture and Consumer Services, Bugwood.org



*Phytophthora* infection in a pine tree. Note the stunted growth. High water tables and excess irrigation provide suitable conditions for root infections.



*Phytophthora* infection in a pine tree showing rotted area near roots. Once infected, water flow through the xylem is reduced via wilt-inducing toxins.

John H. Giant, USDA Forest Service, Bugwood.org

1. Several parasitic protozoans causing diseases in humans use other animal species as hosts for part of their life cycle. Identify the host (including class and genus) that is involved in part of the life cycle for malaria:

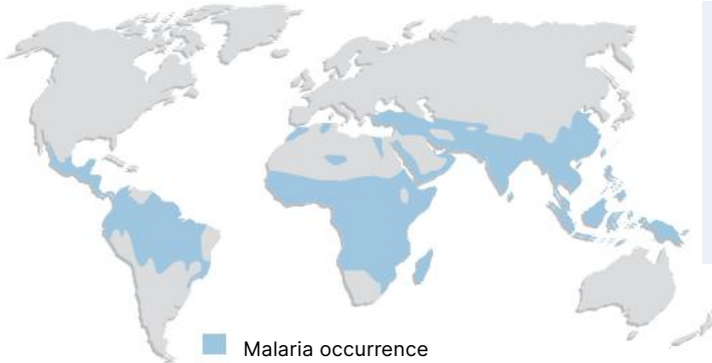
\_\_\_\_\_

2. Why does infection by *Phytophthora cinnamomi* cause stunted growth and death in plants? \_\_\_\_\_

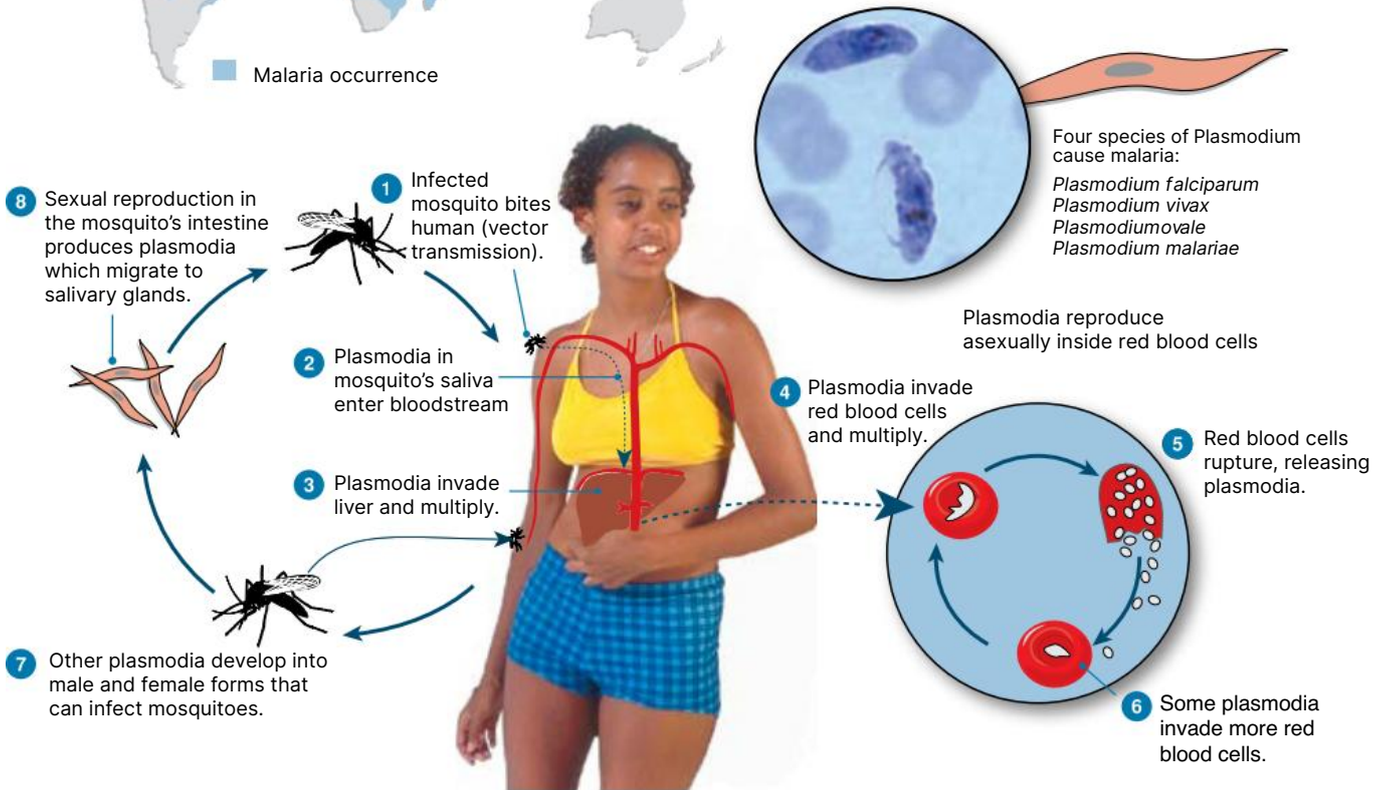
\_\_\_\_\_

## Malaria is caused by a protistan parasite

- ▶ Malaria is a disease caused by protistan parasites of the genus *Plasmodium*. The plasmodia have a life cycle involving two hosts, *Anopheles* mosquitoes, which act as a vector for transmission of the parasite, and humans. Humans become infected when bitten by mosquitoes infected with the protozoans. In their human host, the plasmodia infect red blood cells (RBCs) and multiply inside the cells by asexual reproduction.
- ▶ Four *Plasmodium* species cause malaria, ranging in severity from relatively mild to fatal. *Falciparum* malaria is the most severe because it affects red blood cells of all ages. Destruction of the RBCs results in a condition called haemolytic anaemia (loss of RBCs through lysis). Infected blood cells also become sticky and then block blood vessels to vital organs such as the kidneys and brain.



Malaria is a major health problem in tropical regions where the climate is warm and wet enough to support breeding populations of the mosquito vector. Malaria affects more than 300 million people a year in equatorial regions (left). Cases in Australia only result when infected travellers return from these regions, although Northern Australia could harbour malarial mosquitoes if the global climate warms significantly.



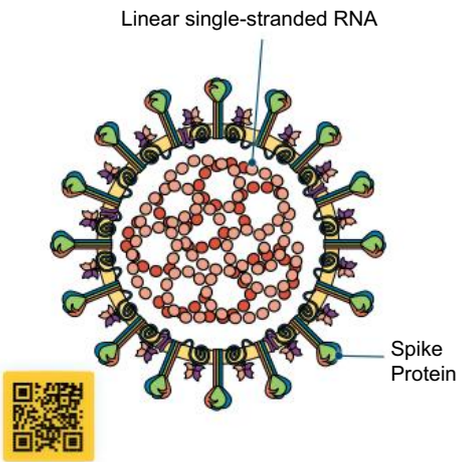
- How does a *Plasmodium* parasite enter the body? \_\_\_\_\_  
 \_\_\_\_\_
- What aspects of the biology of this pathogen could make it difficult to control? \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_
- (a) What biological factors are important in the global occurrence of malaria? \_\_\_\_\_  
 \_\_\_\_\_  
 (b) What measures could be cost effective in controlling the number of new malaria infections? \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_
- Why is global warming expected to increase the geographical range of malaria? \_\_\_\_\_  
 \_\_\_\_\_

# 173 Viral Diseases

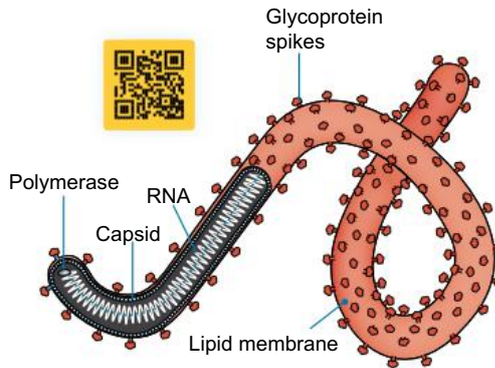
**Key Idea:** A virus is an infectious, highly specialised intracellular parasite. They are acellular and non-living.

**Viruses** are exclusively **disease-causing agents (pathogens)**, which replicate (reproduce themselves) only inside the living cells of other organisms (**host**). Viruses are acellular, meaning they are not made up of cells, so they do not conform to the existing criteria for live organisms. A typical virus contains genetic material (DNA or RNA) encased in a protein coat (**capsid**). Some viruses have an additional membrane, called

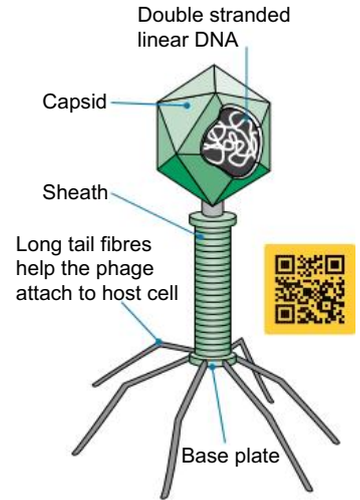
an envelope, surrounding the capsid. Many viruses have glycoprotein receptor spikes on their envelopes. These are **adherence factors** and are adaptations that allow viruses to attach to targeted host cells. Viruses also have **invasion factors**, adaptations that allow them to invade a host cell, and break through the protective cell membrane and wall (in bacteria and plants). Viruses vary greatly in their appearance and the type of host they infect (below). Although often called microorganisms, viruses are not classed as lifeforms.



Structure of SARS-CoV-2, a coronavirus causing Covid-19.



Structure of Ebola virus, an RNA filovirus that causes Ebola haemorrhagic fever.



Structure of Lambda phage, a bacteriophage that infects E.coli.

<p>HIV budding from a lymphocyte</p> <p>HIV</p>	<p>Tobacco mosaic virus (TMV)</p>	<p>Ebola virus</p>	<p>Bacteriophages (arrowed) infect bacteria. They use tail fibres to attach to the host cell and a contractile region below the capsid to inject their DNA into the cell.</p>
<p>After replication, new viral particles (virions) leave the host cell to infect more cells. In animals, enveloped viruses bud from the host cell, e.g. HIV (above left). Plant viruses cannot bud from the host cell due to the rigid cell wall. Instead, plant viruses, e.g. TMV (above right), move through the plasmodesmata connecting plant cells.</p>		<p>Viruses cause a wide variety of human diseases, e.g. colds, influenza, chickenpox, measles, Mpox, Covid-19 and life-threatening diseases such as Ebola (above).</p>	

1. What is the significance of viruses being non-living? \_\_\_\_\_
2. Describe the basic structure of a generalised virus, identifying the features they all have in common: \_\_\_\_\_
3. Describe the purpose of the following:
  - (a) Glycoprotein spikes: \_\_\_\_\_
  - (b) A bacteriophage's tail fibres: \_\_\_\_\_
  - (c) Protein capsid: \_\_\_\_\_



# 174 HIV: An Example of a Viral Disease

**Key Idea:** The human immunodeficiency virus (HIV) infects lymphocyte cells, eventually causing AIDS, a fatal disease, which acts by impairing immune system function.

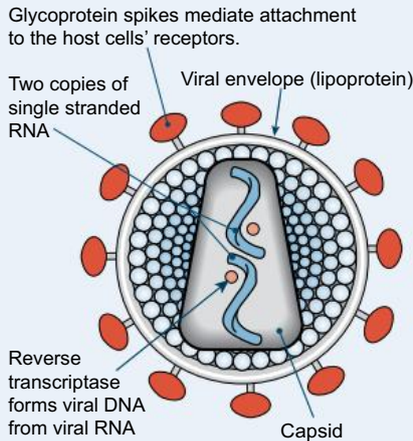
HIV (human immunodeficiency **virus**) is a retrovirus (a type of viral **pathogen**) which binds to the CD4 receptor on the surface of T helper cells (a type of white blood cell); these are central to cellular immunity and coordinate the immune response. HIV causes immune deficiency by replicating

inside T helper cells and destroying them. Over time, a **disease** called AIDS (acquired immunodeficiency syndrome) develops and the immune system progressively loses its ability to fight infection. HIV is transmitted from person to person in body fluids such as blood, vaginal secretions, semen, breast milk, and across the placenta. Unless there are skin cuts, the risk of HIV transmission and infection through close contact between people remains very low.

## HIV infects T helper cells

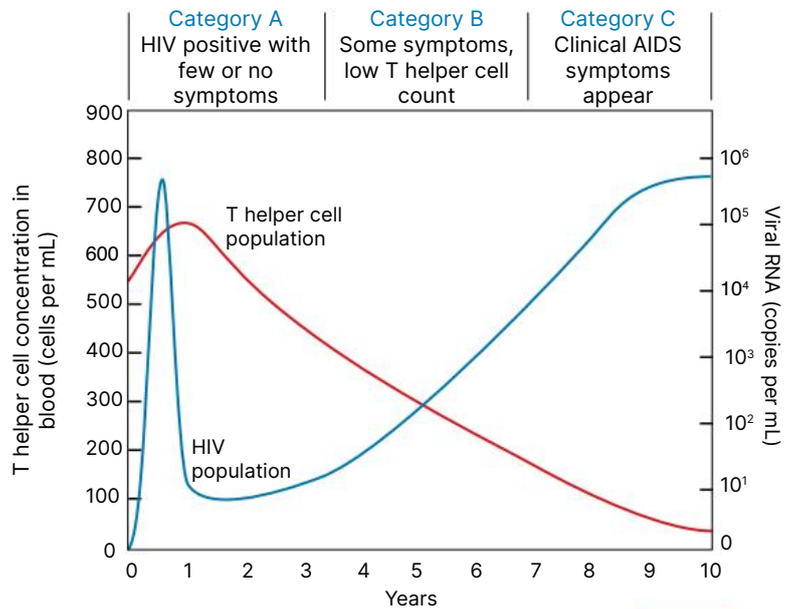
HIV infects T helper cells by fusing its lipid bilayer with that of the host cell. It uses the cells to replicate itself in great numbers, then the newly formed viral particles exit the cell to infect more T helper cells. Many T helper cells are destroyed by viral replication. Because of their role in cellular immunity, T helper cell destruction recruits more T-lymphocytes, accelerating the infection of new cells.

Once the T helper cell population becomes depleted, the immune system's ability to fight infection is severely compromised.



Structure of HIV

The graph below shows the relationship between the level of HIV infection and the number of T helper cells. AIDS is only the end stage of an HIV infection. Shortly after the initial infection, HIV antibodies appear in the blood. There are three clinical categories during progression of the disease. The progressive reduction of T helper cells results in almost no immunity against any other pathogens. Mild illness, such as a respiratory cold, can be fatal for HIV/AIDS patients in the final stages.



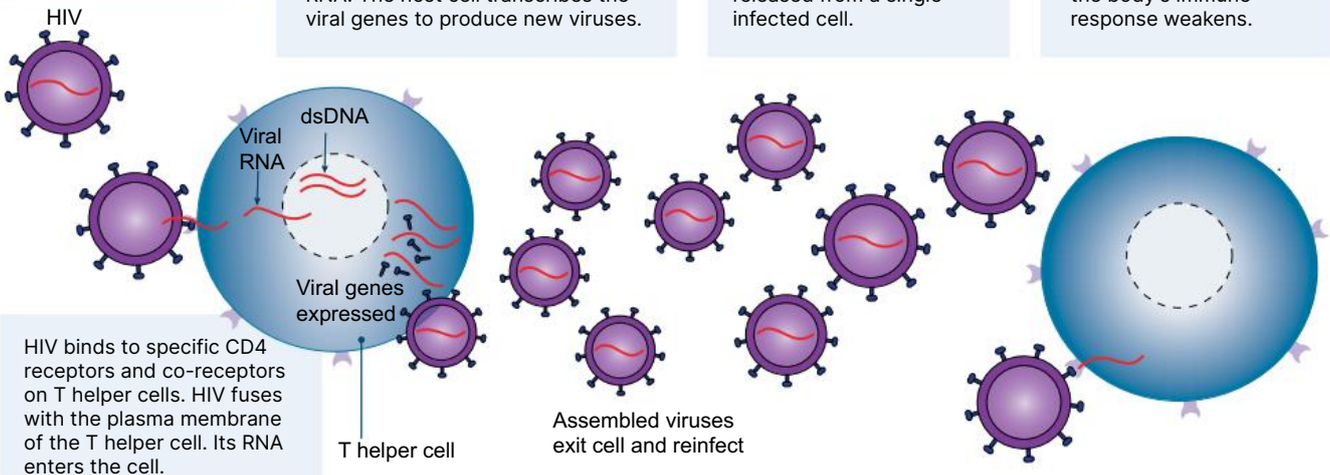
## HIV uses the cellular machinery of T helper cells to replicate

The genetic material of HIV is a single strand of RNA.

HIV hijacks the T helper cells' machinery to replicate itself. Reverse transcriptase produces double stranded DNA (dsDNA) from the viral RNA. The host cell transcribes the viral genes to produce new viruses.

The new HIV particles bud from the T helper cell. Between 1000 and 3000 new HIV particles can be released from a single infected cell.

The HIV particles mature and infect more T helper cells. As more T helper cells become infected, the body's immune response weakens.



HIV binds to specific CD4 receptors and co-receptors on T helper cells. HIV fuses with the plasma membrane of the T helper cell. Its RNA enters the cell.

Assembled viruses exit cell and reinfect



**AIDS: The end stage of an HIV infection**

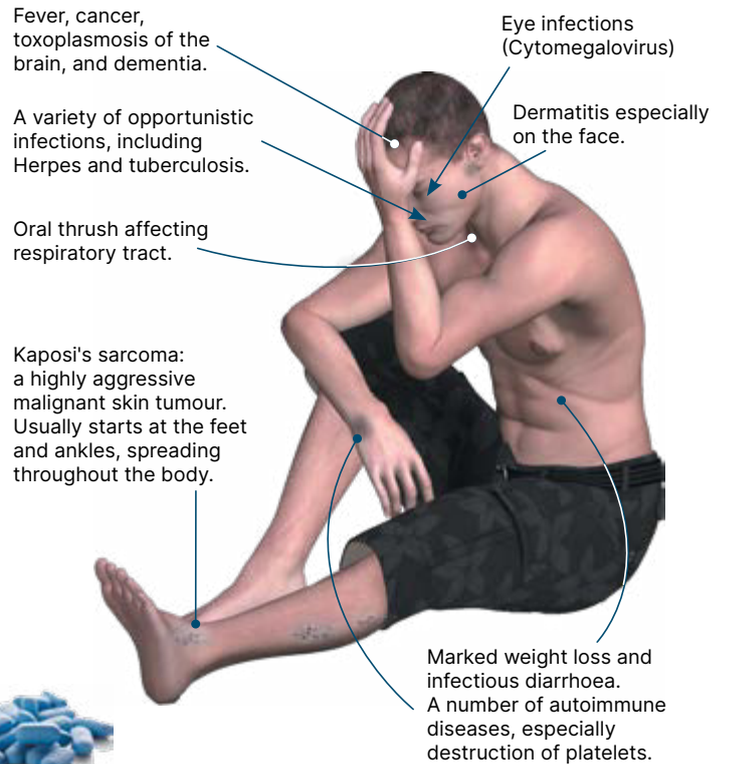
HIV/AIDS is a spectrum of disorders (right) arising as a consequence of impaired immune function, which prevents the body detecting and destroying pathogens or damaged cells. People with healthy immune systems can fight off the challenges of pathogens and are able to detect and destroy damaged (pre-cancerous) cells. However, people with HIV are susceptible to all pathogens because their resistance to disease is so low. What's more, loss of the T cell population compromises the ability of HIV-infected people to detect and destroy pre-cancerous cells. Rare cancers are a common symptom of HIV/AIDS.

**Medications**

Antibiotics can be used to treat some of the infections contracted due to the reduced immune system, e.g. tuberculosis, but they cannot be used to treat the HIV infection itself because antibiotics are ineffective against viruses.

Although there is currently no cure for HIV/AIDS, some antiretroviral drugs can slow the progress of the disease by interfering with the replication of HIV and slowing the advance of the disease.

Pre-exposure prophylaxis (PrEP) medications can be prescribed to people who do not have HIV, but are at risk of contracting it. Correct use of PrEP, along with other precautions, can reduce the risk of contracting HIV by up to 99%.



1. (a) What type of cells does HIV infect? \_\_\_\_\_  
 (b) How does HIV recognise this type of cell? \_\_\_\_\_  
 \_\_\_\_\_  
 (c) What is the role of reverse transcriptase in HIV replication?  
 \_\_\_\_\_  
 \_\_\_\_\_
2. Study the graph on the previous page showing how HIV affects the number of T helper cells. Describe how the viral population changes with the progression of the disease:  
 \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_
3. (a) What effect does HIV have the cells of the immune system? \_\_\_\_\_  
 \_\_\_\_\_  
 (b) Describe the effect of this change on the long-term health of a person with HIV: \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_
4. (a) Why is the purpose of antibiotics in treatment of HIV/AIDS? \_\_\_\_\_  
 \_\_\_\_\_  
 (b) Why are antibiotics ineffective against the HIV infection itself? \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_

# 175 Prions

**Key Idea:** Prions are misfolded infectious proteins that can propagate by causing misfolding in the original protein type. Until recently, all **pathogens** were thought to contain some form of nucleic acid. We now know that particular proteins, called **prions**, are capable of causing infection. Prions have been spread by eating contaminated meat and, because they resist normal sterilisation methods, they can be spread on

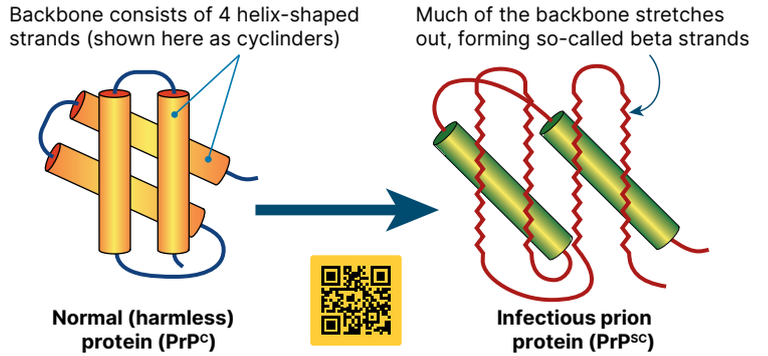
surgical instruments. Prions are produced by mutations in the gene coding for a normal cell protein (PrP). They cause a group of degenerative nervous **diseases** in mammals called transmissible spongiform encephalopathies (TSE). They include scrapie in sheep, BSE in cattle, and kuru in humans. Different mutations of the PrP gene are responsible in each case of disease.



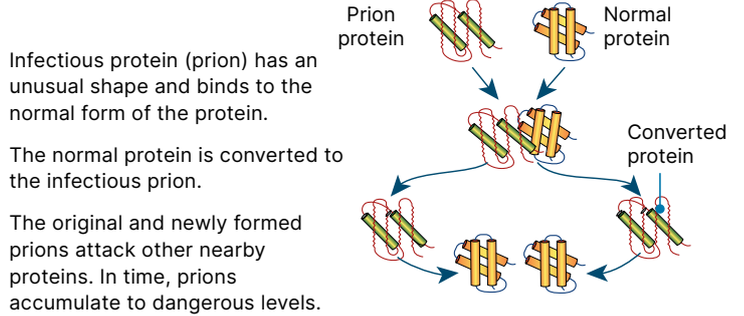
The disease kuru first brought prion diseases to prominence in the 1950s. It occurred in the geographically isolated tribes in the Fore highlands of Papua New Guinea. These people were eating the brain tissue of dead relatives for religious reasons. Normal and infectious prions have the same primary structure, so go unchallenged by the immune system. If the infectious prion is from another species (as in vCJD) there is an initial immune response but this is shut down as the infectious protein converts more and more of the body's own PrP<sup>C</sup> to PrP<sup>Sc</sup>.

## Infectious prion proteins

A shape change transforms the harmless protein into an infectious prion. The change may be caused by a point mutation in the encoding gene. The normal (common) form of the protein is denoted PrP<sup>C</sup>, whereas the abnormal form is denoted PrP<sup>Sc</sup> (after scrapie, the prototype prion).



## Propagation of the prion protein



Prion diseases of humans (and cause)	
All these diseases are characterised by dementia and loss of coordination. There may be other symptoms as well.	
▶	Kuru (infection through cannibalism)
▶	Variant Creutzfeldt-Jacob Disease (vCJD) (infection)
▶	Classical Creutzfeldt-Jacob Disease (infection or mutation)
▶	Fatal Familial Insomnia (inherited mutation)

1. What is the main feature of prions distinguishing them from other infectious agents? \_\_\_\_\_
2. How does a prion's mode of transmission make it a successful agent of disease? \_\_\_\_\_
3. What is the source of infection for people with variant CJD? \_\_\_\_\_
4. How did the cultural practices of highland tribes in PNG enable the spread of kuru? \_\_\_\_\_
5. An epidemic of BSE in the UK in the 1990s had its origin in the practice of processing waste parts of cattle (particularly nervous tissue) and recycling them into cattle feed. Infected cattle subsequently entered the human food chain and were linked to cases of vCJD. Explain why it is poor practice to process an animal and feed it back to the same species: \_\_\_\_\_

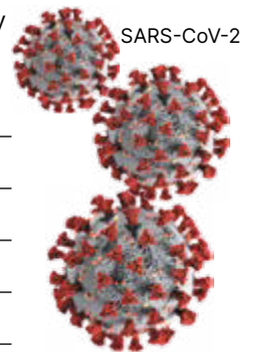
1. Test your vocabulary by matching each term to its correct definition, as identified by writing the letter in the correct box.

- |                            |                          |  |
|----------------------------|--------------------------|--|
| (i) prion                  | <input type="checkbox"/> | <b>A</b> Any disease caused by the invasion of a host by a pathogen which grows and multiplies in the body and is transmissible to others.                             |
| (ii) bacteria              | <input type="checkbox"/> | <b>B</b> A type of disease that cannot be transmitted between individuals.   |
| (iii) disease              | <input type="checkbox"/> | <b>C</b> Infectious proteins that can cause abnormal folding of normal cellular proteins, leading to neurodegenerative diseases.                                       |
| (iv) infectious disease    | <input type="checkbox"/> | <b>D</b> A disease-causing organism.   |
| (v) non-infectious disease | <input type="checkbox"/> | <b>E</b> Single celled microorganisms surrounded by a cell wall containing the substance peptidoglycan. Some are pathogens responsible for serious diseases in humans. |
| (vi) pathogen              | <input type="checkbox"/> | <b>F</b> A non-cellular obligate intracellular parasite, requiring a living host to reproduce. Does not respond to antibiotics.  |
| (vii) virus                | <input type="checkbox"/> | <b>G</b> An abnormal condition of the body when bodily functions are impaired.   |

2. The table below lists some infectious diseases. Complete the table by naming the type of pathogen that causes the disease (bacteria, virus, protist), and the symptoms of the disease. You may need to do some extra research.

Disease	Type of pathogen	Symptoms of disease
Cholera		
Malaria		
TB		
HIV/AIDS		
Smallpox		
Measles		

3. SARS-CoV-2 is the virus responsible for the Covid-19 disease that started as a pandemic in early 2020. Explain how the glycoprotein spikes act as an adherence factor and the spike protein acts as an invasion factor:




---



---



---



---



---

4. Bacteriophages are viruses that infect bacteria. They can be used to diagnose certain bacterial diseases in much the same way as testing the effect of antibiotics on bacteria. Study the photo below, it shows the effect of a bacteriophage (a type of virus that infects bacteria) and an antibiotic (polymyxin B) on cholera bacteria.



(a) What evidence is there that the bacteriophage was effective at killing the cholera bacteria and how can you tell?

---



---

(b) Explain the process that the bacteriophage used to kill the bacteria:

---



---



---



---



# Immune Response

## Key Terms

- adaptive immune response
- allergens
- antibodies
- antigens
- B lymphocytes (cells)
- clonal selection
- complement system
- defensins
- immune response
- immunity
- inflammation
- inflammatory response
- innate response
- killer (cytotoxic) T cells
- macrophages
- major histocompatibility complex (MHC)
- memory cells
- natural killer cells
- neutrophils
- non-self antigen
- pathogens
- phagocytes
- prostaglandins
- self-antigens
- T lymphocytes (cells)
- toxins (plant)
- vaccine
- vaccination
- vasodilation

## Key Concepts

- ▶ Vertebrates defend against pathogens through three lines of defence.
- ▶ The body has an innate and an adaptive immune response.
- ▶ Plants use physical barriers and chemical defences to protect against pathogens.

## Innate Immunity

	Activity Number
□ 1 Explain how bacterial and viral pathogens can stimulate the immune system of a host by acting as antigens. Explain how the body distinguishes self from non-self, including the role of the major histocompatibility complex (MHC) and the processing of antigens.	177, 182
□ 2 Describe non-specific (innate) defences in humans and describe the nature and role of skin, mucous membranes, and body secretions.	178
□ 3 Discuss the inflammatory response, including the roles of prostaglandins and vasodilation, phagocytosis, and natural killer cells.	179-180
□ 4 Describe how prostaglandin production is controlled by COX enzymes in the inflammatory response.	179
□ 5 Explain the role of the complement (non-specific) system, including the function of neutrophils and macrophages.	179
□ 6 Describe passive and active defences in plants, including reference to physical and chemical barriers, defensins, toxins, and nastic responses.	191-192

## Adaptive and acquired immunity

□ 7 Explain the adaptive immune response in vertebrates (e.g. humans). Describe cell-mediated immunity (T lymphocytes) and humoral (B lymphocyte and antibody-mediated) immunity, identifying the specific white blood cells involved in each case.	183
□ 8 Describe clonal selection and the basis of immunological memory. Explain how the immune system is able to respond to the large range of potential antigens.	184
□ 9 Explain antibody production, including how B lymphocytes bring about humoral (antibody-mediated) immunity to specific antigens.	185
□ 10 Analyse the similarities and differences between passive and active immunity for both naturally and artificially acquired immunity.	186
□ 11 Explain the principles of vaccination, including reference to the primary and secondary response to infection and the role of these.	187
□ 12 Explain the role of herd immunity and its relationship to vaccination rate.	187
□ 13 Compare individual and population immunities of different geographical and demographical populations, including response to vaccination.	188
□ 14 Interpret long term immune response data. Examine short and long term patterns of immunity within the context of vaccination practices.	190
□ 15 <b>SHE:</b> Investigate a range of case studies contributing to the immune response and vaccine development.	189-190
□ 16 <b>SI:</b> Explore how the work of scientists, including Rosalyn Yalow and Peter C Doherty, has contributed to our understanding of vaccines and immunity.	189
□ 17 <b>SI:</b> Explore stages of vaccine development in the context of the Covid-19 vaccine.	189

# 177 The Nature of Antigens

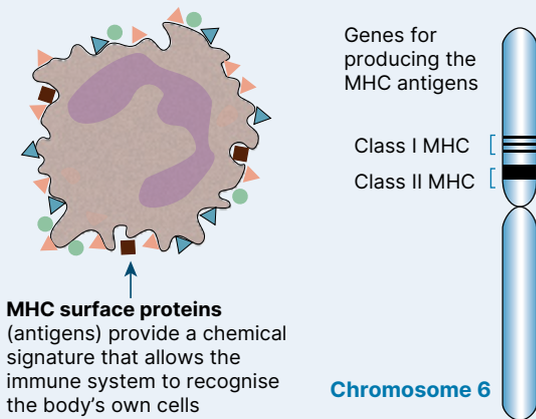
**Key Idea:** Antigens are substances capable of producing an immune response. It is important that the body can distinguish its own tissues from foreign material so that it does not attack itself.

An **antigen** is any substance that produces an **immune response**. Most antigens are **non-self** antigens, i.e. they are foreign and originate from outside the organism (e.g. bacteria or viruses). Sometimes an organism will react to its own cells

and tissues. Antigens that originate from within the body are called **self-antigens**. Normally, because of the development of self-tolerance, the body recognises and does not attack its own tissues. In some instances, the immune system may mistakenly destroy its own tissues. Such a response is called an autoimmune disorder. Allergens are a specific type of antigen that produce a vigorous hypersensitive allergic response from the immune system.

## Distinguishing self from non-self

- ▶ Every type of cell has unique protein markers (antigens) on its surface. The type of antigen varies greatly between cells and between species. The immune system uses these markers to identify its own cells (self) from foreign cells (non-self). If the immune system recognises the antigen markers, it will not attack the cell. If the antigen markers are unknown, the cell is attacked and destroyed.
- ▶ In humans, the system responsible for this property is the **major histocompatibility complex (MHC)**. The MHC is a cluster of tightly linked genes on chromosome 6. These genes code for protein molecules (MHC antigens) that are attached to the surface of body cells. The main role of MHC antigens is to bind to antigenic fragments and display them on the cell surface so that they can be recognised by the cells of the immune system.
- ▶ Class I MHC antigens are found on the surfaces of almost all human cells. Class II MHC antigens occur only on **macrophages** and **B lymphocytes** (B cells) of the immune system.



## Tolerance towards foreign bodies

- ▶ The human body has a very large population of resident microbes. Under normal conditions, *E.coli* in the gut form a protective layer preventing the colonisation of pathogenic bacteria. The microbial cells have foreign antigens but they are not attacked by the immune system because tolerance (the prevention of an immune response) has developed.
- ▶ During pregnancy, specific features of the self recognition system are suppressed to allow the mother to tolerate a nine month relationship with a foreign body (the fetus).



*E.coli*

## Intolerance to tissue transplants

The MHC is responsible for the rejection of tissue grafts and organ transplants. Foreign MHC molecules on the transplanted tissue are viewed as antigenic, causing the immune system to respond and the tissue to be rejected. To minimise rejection, attempts are made to match the MHC of the organ donor to that of the recipient as closely as possible. Immunosuppressant drugs are also used to minimise the immune response.



Kidney transplant

- (a) What is an antigen? \_\_\_\_\_

\_\_\_\_\_
- (b) Distinguish between non-self antigens and self antigens: \_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_
- (c) Why is it important that the body detects foreign antigens? \_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_

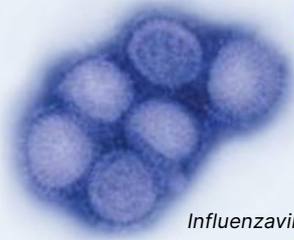
## Types of antigens

### Non-self antigens

Any foreign material provoking an immune response is termed a non-self antigen. Disease-causing organisms (**pathogens**) such as bacteria, viruses, and fungi are non-self antigens. The body recognises them as foreign and will attack and destroy them before they cause harm.



cdc



Influenzavirus

Pathogens have ways of avoiding detection. Mutations result in new surface antigens, delaying the immune response and allowing the pathogen to reproduce in its host undetected for a time (e.g. the flu virus, above). Some pathogens, e.g. the malaria-causing *Plasmodium*, switches off its surface antigens in order to enter cells undetected.

### Self antigens

The body is usually tolerant of its own antigens. However, sometimes the self-tolerance system fails and the body attacks its own cells and tissues as though they were foreign. This can result in an autoimmune disorder in which tissue is destroyed, grows abnormally, or changes in function.

Autoimmune disorders, such as multiple sclerosis and rheumatoid arthritis, may be triggered by infection. The similarity of the pathogen and self antigens is thought to be behind this failure of self recognition.



Type 1 diabetes is the result of autoimmune destruction of the insulin-producing pancreatic cells. Patients must inject insulin to maintain normal blood glucose levels.

### Allergens

Antigens that cause allergic reactions are called allergens. An allergic reaction is a very specific type of immune response in which the immune system overreacts to a normally harmless substance. An allergic response can produce minor symptoms (itching, sneezing, rashes, swelling) or life-threatening anaphylaxis (respiratory and cardiovascular distress).

Common allergens include dust, chemicals, mould, pet hair, food proteins, or pollen grains.



The swelling on the foot in the left of the photograph is a result of an allergic reaction to a bee sting.

Kent Pryor

2. How can pathogens avoid detection by the immune system? \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_
3. (a) What is the nature and purpose of the major histocompatibility complex (MHC)? \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_  
 (b) Why is a self-recognition system important? \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_
4. (a) What is immune tolerance? \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_  
 (b) When might tolerance to foreign antigens be beneficial or necessary? \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_
5. Using examples, describe what happens when the body develops an inappropriate response to:
  - (a) Self-antigens: \_\_\_\_\_  
 \_\_\_\_\_
  - (b) Normally non-antigenic substances: \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_

# The Body's Defences: An Overview

**Key Idea:** The human body has a tiered system of defences that provides resistance against disease. The human body has a suite of physical, chemical, and biological defences against **pathogens**, collectively called resistance. The first line of defence consists of external barriers to prevent pathogen entry. If this fails, a second line of defence targets any foreign bodies that enter. Lastly, the specific **immune response** provides targeted defence against the pathogen. The defence responses of the body fall into two broad categories: the **innate** and the **adaptive**

**immune** responses. The innate (or non-specific) response (the first and second lines of defence) protects against a broad range of non-specific pathogens. This response is present in all animals. It involves blood proteins (e.g. complement), **inflammation**, and phagocytic white blood cells. The adaptive (or specific) immune response (the third line of defence) is specific to identified pathogens and is present only in vertebrates. It involves defence by specific **T lymphocytes** (cellular immunity) as well as **antibodies**, which neutralise foreign **antigens** (humoral immunity).

Most microorganisms find it difficult to get inside the body. If they succeed, they face a range of other defences that protect the body.

The natural populations of harmless microbes living on the skin and mucous membranes inhibit the growth of most pathogenic microbes.

Microorganisms are trapped in sticky mucus and expelled by cilia (tiny hairs that move in a wavelike fashion).

## 1st line of defence (Innate)

Also called the primary defence, and comprises the skin and mucous membranes. The skin is the body's largest organ. Unlike other epithelial membranes, it is a dry membrane, whereas mucous membranes (mucosa) are moist due to secretions.

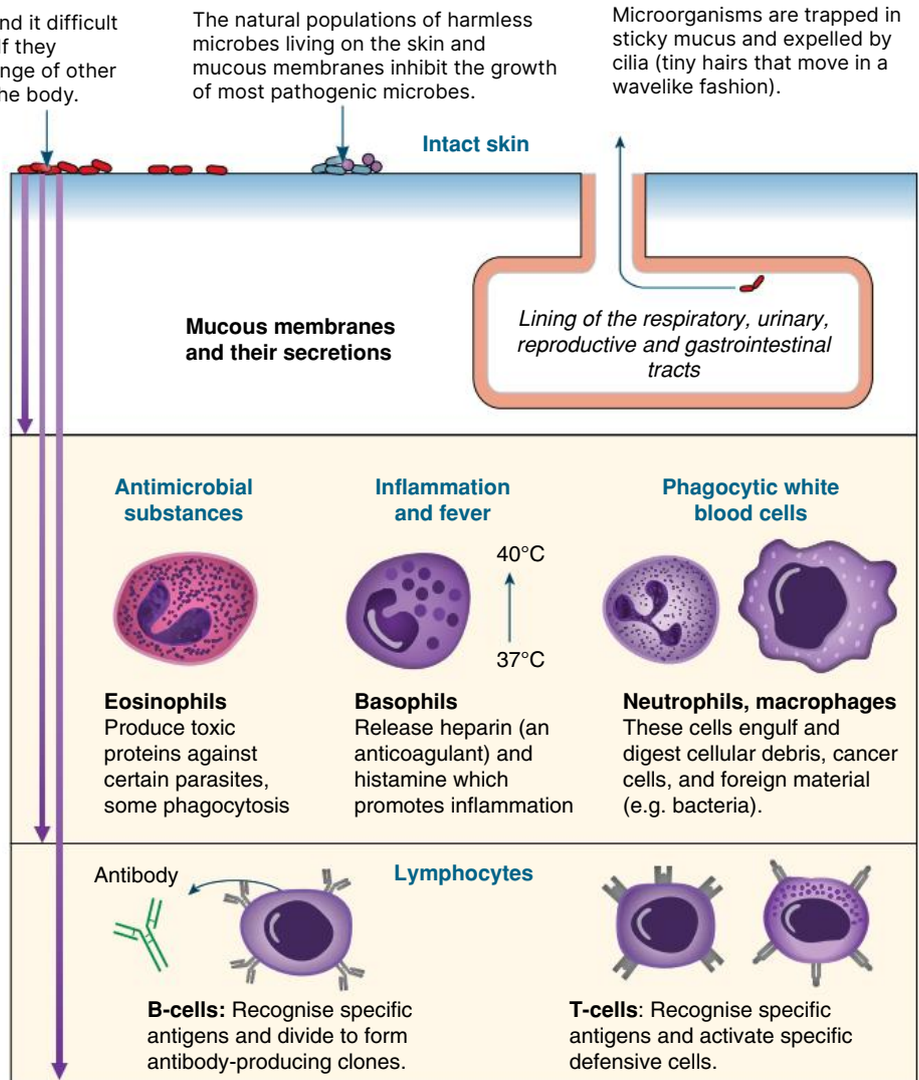
## 2nd line of defence (Innate)

A range of defence mechanisms operate inside the body to inhibit or destroy pathogens. These responses react to the presence of any pathogen, regardless of which species it is. White blood cells are involved in most of these responses.

The 2nd line of defence includes the complement system, whereby blood plasma proteins work together to bind pathogens and induce inflammation to help fight infection.

## 3rd line of defence (Adaptive)

Once the pathogen has been identified by the immune system, lymphocytes (specialised white blood cells) launch a range of specific responses to the pathogen, including the production of defensive proteins called antibodies. Each type of antibody is produced by a B cell lymphocyte clone and is specific against a particular antigen.



1. What are the differences between the innate and adaptive immune responses?

---



---



---



---

2. How does having a tiered defence help protect an organism from a pathogen?

---



---



---



---



**Key Idea:** The innate immune response provides a rapid response to contain and destroy pathogens. Inflammation is an important part of the response.

The innate immune system provides protection against a pathogen, even if it has never encountered it before. The **innate response** is very fast and provides general protection (it is not **antigen** specific), but does not provide long lasting

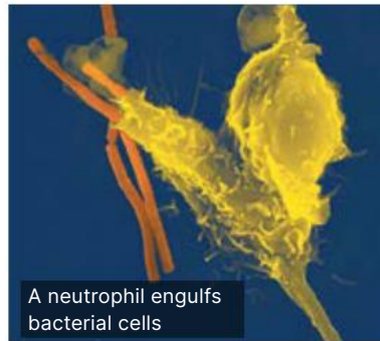
**immunity**. Many different cells and processes are involved. The primary outcome is to destroy and remove the cause of infection. This is achieved through containing the infection through **inflammation** and then recruitment of immune cells to destroy the **pathogen**. During this process a series of biochemical reactions (the complement system) are activated to destroy the pathogen and recruit immune cells to the site.

### Phagocytic cells of the innate immune system



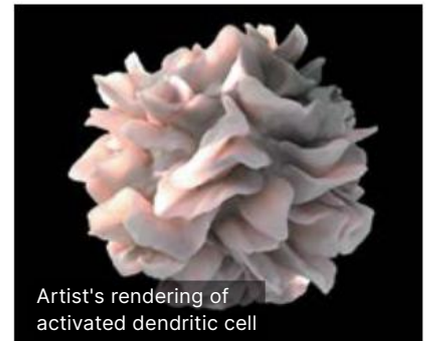
#### Macrophage

**Macrophages** are very large and are highly efficient **phagocytes**. They are found throughout the body and move using an amoeboid movement (above) to hunt down and destroy pathogens. Macrophages also have a role in recruiting other immune cells to an infection site.



#### Neutrophil

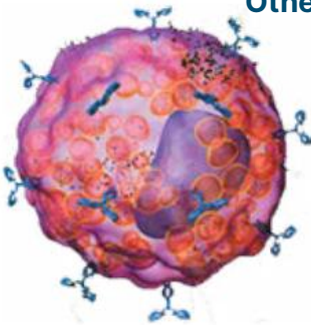
**Neutrophils** are the most abundant type of phagocyte and are usually the first cells to arrive at the site of an infection. They contain toxic substances that kill or inhibit the growth of bacteria and fungal pathogens. Neutrophils release cytokines which amplify the immune response and recruit other cells to the infection site.



#### Dendritic cell

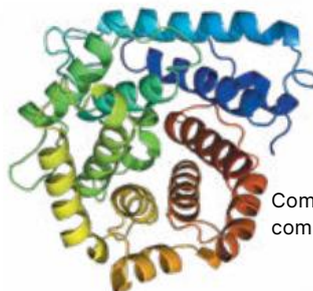
Dendritic cells are present in tissues that are in contact with the external environment (e.g. skin, and linings of the nose, lungs, and digestive tract). They act as messengers between the innate and **adaptive immune** system by presenting **antigen** materials to the **T lymphocytes** of the immune system.

### Other cells and processes of the innate immune response



#### Mast cells

Mast cells contain a lot of histamine, a chemical involved in both inflammation and allergic responses. When activated, histamine is released from the mast cell causing the blood vessels to dilate and become leaky. The increased permeability allows phagocytes to reach the site of infection.



#### Complement proteins

The **complement system** comprises a number of different proteins. The proteins circulate as inactive precursors until they are activated. Complement proteins have three main roles: phagocytosis, attracting macrophages and neutrophils to the infection site, and rupturing the membranes of foreign cells.



#### The process of inflammation

The inflammatory process is a protective response to pathogen invasion. It has several functions: (1) to destroy the cause of the infection and remove it and its products from the body; (2) if this fails, to limit the effects on the body by confining the infection to a small area; (3) replacing or repairing tissue damaged by the infection.

1. Outline the role of the following phagocytes in the innate immune response:

(a) Macrophages: \_\_\_\_\_

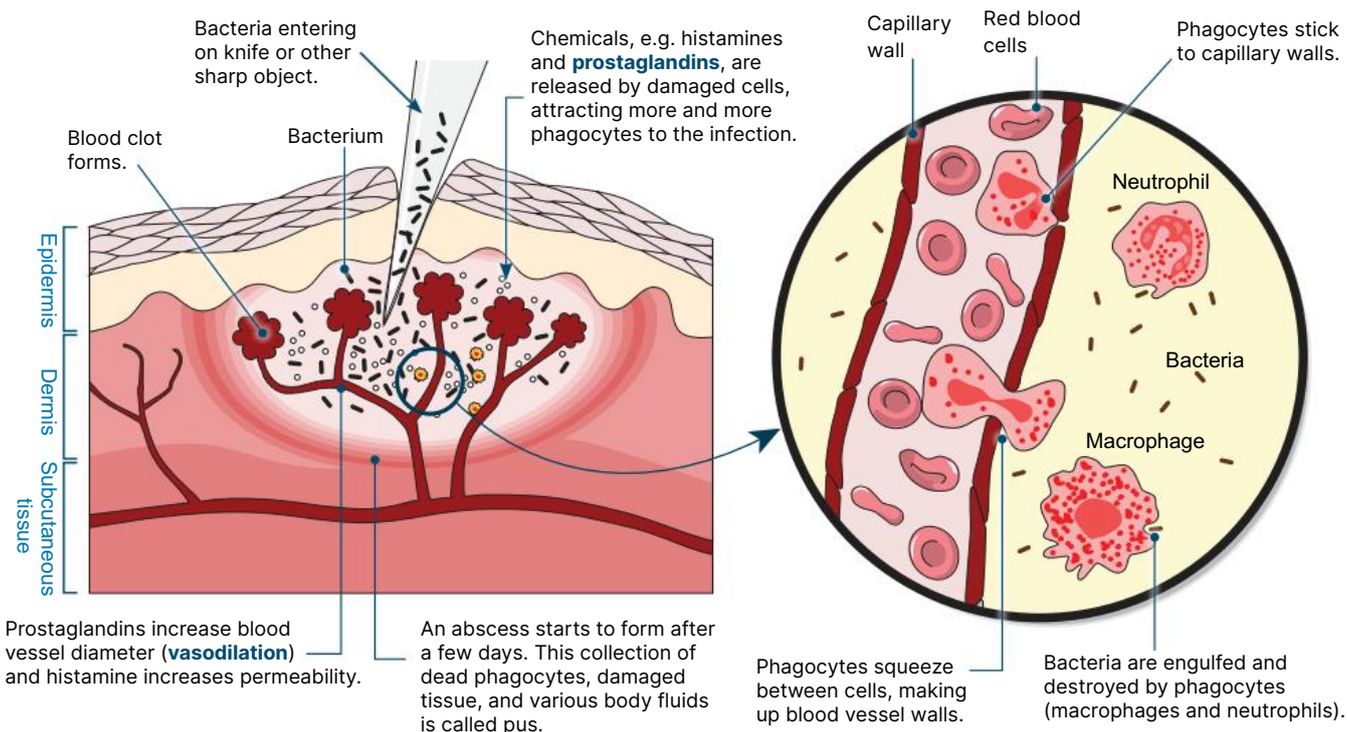
(b) Neutrophils: \_\_\_\_\_

(c) Dendritic cells: \_\_\_\_\_

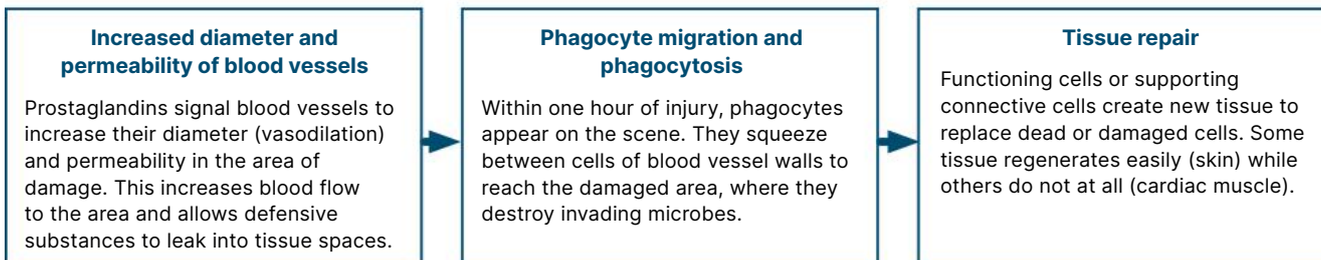


## The inflammatory response

Inflammation is a defensive response to damage. The inflammation process involves pain, redness, heat, and swelling. Damage to the body's tissues can be caused by physical agents, e.g. sharp objects, heat, radiant energy, or electricity; microbial infection, or chemical agents, e.g. gases, acids, and bases. The damage triggers a defensive response called inflammation. The **inflammatory response** is generally beneficial and the process of inflammation can be divided into three distinct stages (described below).

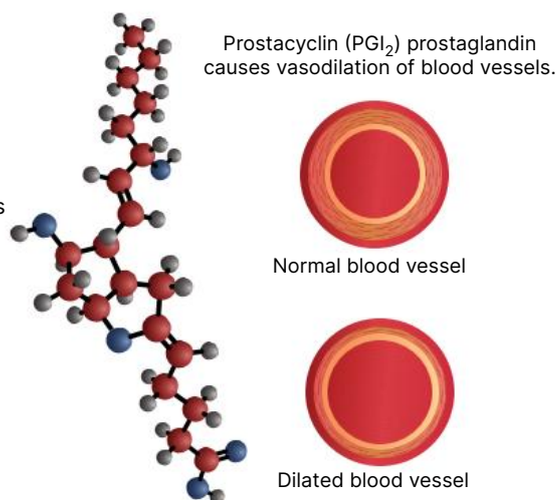


### Stages in the inflammatory response



### Prostaglandins, vasodilation, and inflammation

- ▶ Prostaglandins are lipid structures that act as hormones. There are many types of prostaglandins, each with a distinct function, but many prostaglandins serve as signals to drive the immune response.
- ▶ Prostaglandins are produced by cells in close proximity to where they are needed. When tissue is damaged or infected, specific prostaglandins are involved in producing the inflammatory response (above).
- ▶ Many prostaglandins act as powerful vasodilators, causing the smooth muscle of blood vessels to relax so that the vessels dilate (open up) which allows defensive substances and white blood cells to flow more freely to the damaged or infected tissue and initiate repairs.

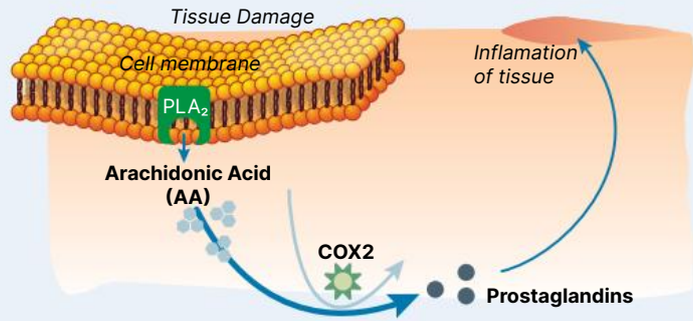


2. Why is vasodilation and permeability an important response to inflammation? \_\_\_\_\_
- \_\_\_\_\_
- \_\_\_\_\_
- \_\_\_\_\_

### COX, prostaglandins, and painkillers

- ▶ PLA<sub>2</sub> is an enzyme that hydrolyses phospholipids in the cell membrane, releasing arachidonic acid (AA).
- ▶ Arachidonic acid is a precursor for the synthesis of prostaglandins.
- ▶ Initial prostaglandin production is regulated by a two-step enzyme reaction. Cyclooxygenase-1 (COX-1) enzyme maintains a baseline level of prostaglandins necessary for normal physiological functions.
- ▶ Upon injury or inflammation, the cyclooxygenase-2 (COX-2) enzyme is induced, leading to increased prostaglandin production to mediate the inflammatory response.
- ▶ Non-steroidal anti-inflammatory drugs (NSAIDs) are commonly taken as painkillers and interfere with prostaglandin production (diagram right).

### Prostaglandin production in response to tissue damage



NSAIDs such as aspirin and ibuprofen reduce pain and swelling by blocking the effects of the COX enzymes, interfering with the prostaglandin production pathway. This reduces prostaglandin levels and their pain and inflammatory effect in the body.

### Natural killer cells and inflammation

- ▶ **Natural killer cells (NK)** are a type of white blood cell that rapidly respond to target and destroy tumour cells and virus-infected cells. They also have an important role in homeostasis of the immune system and controlling the inflammatory response.
- ▶ Infection or injury triggers NK cells to produce pro-inflammatory molecules called cytokines. Inflammation helps to isolate the threat (pathogen) or begin the healing process. However, if the inflammatory response is too intense or carries on for too long, healthy tissues can become damaged, so the inflammatory response must be regulated.
- ▶ NK cells are also involved in moderating or reducing inflammation. They can signal for anti-inflammatory cytokines to be produced. These reduce the inflammatory response once the threat is under control and make sure the inflammatory response doesn't last longer than necessary, reducing the risk of long term damage.
- ▶ NK cells also control inflammation by destroying over-active immune cells that may be causing excessive inflammation.



Low grade inflammation can last for months or even years; this is called chronic inflammation. This prolonged response can affect nearly every body system. Day-to-day symptoms may include sleep disruption, weight fluctuations, and gastro-intestinal problems. Chronic inflammation is also linked to the onset of diseases such as type 2 diabetes, arthritis, cardiovascular disease, dementia, and some cancers.

3. Outline the three stages of inflammatory response and identify the beneficial role of each stage:
  - (a) \_\_\_\_\_
  - \_\_\_\_\_
  - (b) \_\_\_\_\_
  - \_\_\_\_\_
  - (c) \_\_\_\_\_
  - \_\_\_\_\_
4. What triggers the production of cyclooxygenase-2 (COX-2) and what is its purpose? \_\_\_\_\_
- \_\_\_\_\_
- \_\_\_\_\_
- \_\_\_\_\_
5. Explain why it is important that NK cells have an anti-inflammatory effect: \_\_\_\_\_
- \_\_\_\_\_
- \_\_\_\_\_
- \_\_\_\_\_

# Phagocytes and Phagocytosis

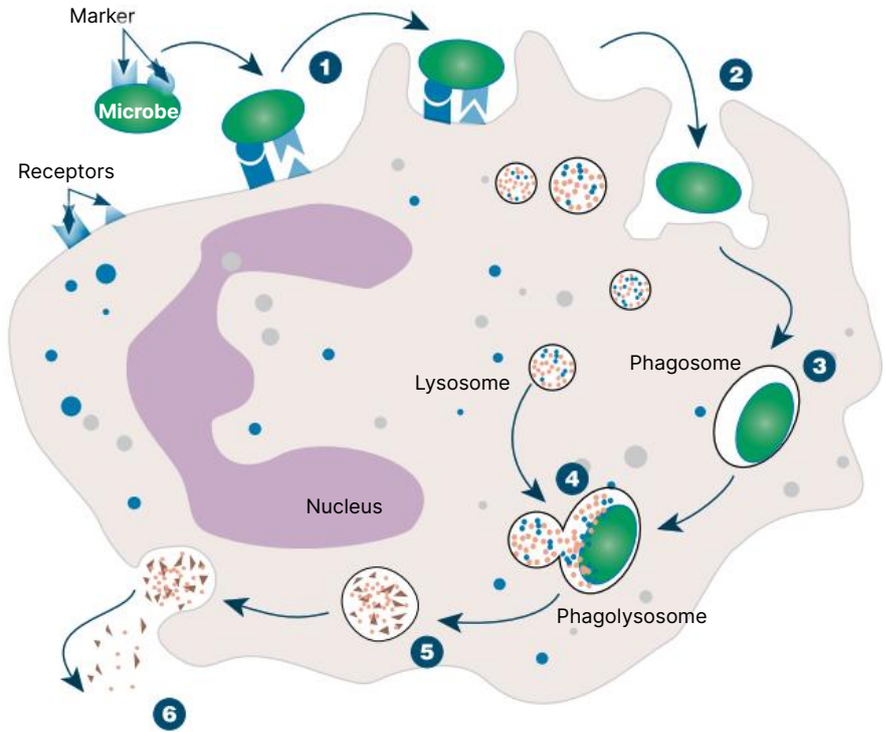
**Key Idea:** Phagocytes are mobile white blood cells that ingest microbes and digest them by phagocytosis.

Phagocytosis is the process by which a cell engulfs another cell or particle. Cells which do this are called phagocytes. All types of **phagocytes** (e.g. **neutrophils**, dendritic cells, and **macrophages**) are white blood cells. These specialised cells have receptors on their surfaces that can detect antigenic

material, such as microbes. They then ingest the microbes and digest them by phagocytosis. As well as destroying microbes, phagocytes also release substances called cytokines that help to coordinate the overall response to an infection. Macrophages and dendritic cells also play a role in antigen presentation in processing and presenting antigens from ingested microbes to other cells of the immune system.

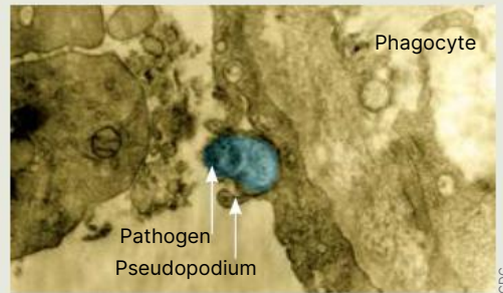
## Stages in phagocytosis and destruction of a pathogen

- 1 Detection and interaction**  
Microbe coated in chemical markers is detected by the phagocyte, which attaches to it. Chemical markers coating the foreign material, e.g. a bacterial cell, mark it as a target for phagocytosis.
- 2 Engulfment**  
The markers trigger engulfment of the microbe by the phagocyte. The microbe is taken in by endocytosis.
- 3 Phagosome forms**  
A phagosome forms, enclosing the microbe in a membrane.
- 4 Fusion with lysosome**  
Phagosome fuses with a lysosome containing digestive enzymes. The fusion forms a phagolysosome.
- 5 Digestion**  
The microbe is broken down into its chemical constituents.
- 6 Discharge**  
Indigestible material is discharged from the phagocyte.



### Amoeboid movement in phagocytes

In order to move independently, the phagocytes force their cytoplasm out to form a pseudopodium (false foot), which moves them forward in space and wraps around the pathogen to engulf them (right). Phagocytes can alter their shape to fit between the thin walls of the capillaries and out into interstitial fluid to trap pathogens. Neutrophils mostly remain inside blood vessels, while macrophages are only found at very low levels in the blood as most remain in body tissues, where they mature.



1. Identify the white blood cells capable of phagocytosis: \_\_\_\_\_  
\_\_\_\_\_
2. Explain the role of chemical markers and phagocyte receptors in enhancing phagocytosis:  
\_\_\_\_\_  
\_\_\_\_\_  
\_\_\_\_\_
3. What is the purpose of phagocytosis and how is involved in internal defence?  
\_\_\_\_\_  
\_\_\_\_\_  
\_\_\_\_\_

# 181 The Lymphatic System

**Key Idea:** Defensive white blood cells are transported in lymph through the lymphatic system and are concentrated in the lymph nodes.

The lymphatic system is a network of tissues and organs that collects the tissue fluid leaked from the blood vessels and returns it to the heart. The lymphatic system has an important

role in immunity because the fluid transported around the body by the lymphatic system (lymph) is rich in infection-fighting white blood cells. The thymus is a primary lymphoid organ and the site of T lymphocyte (also called T-cell) maturation. Secondary lymphoid tissues (spleen and lymph nodes) are important as the site of lymphocyte (T and B **lymphocyte**) activation.

## Components of the lymphatic system

### Tonsils

A collection of secondary lymphoid tissues in the throat. They provide defence against ingested or inhaled pathogens and produce activated B and T lymphocytes/cells.

### Thymus

A primary lymphoid organ located above the heart. It is large in infants and shrinks after puberty to a fraction of its original size. Important for maturation of T-lymphocytes.

### Spleen

The largest mass of lymphatic tissue in the body. It stores and releases blood in case of demand (e.g. in severe bleeding), produces mature B-cells and antibodies, and removes antibody-coated antigenic material.

### Lymph nodes

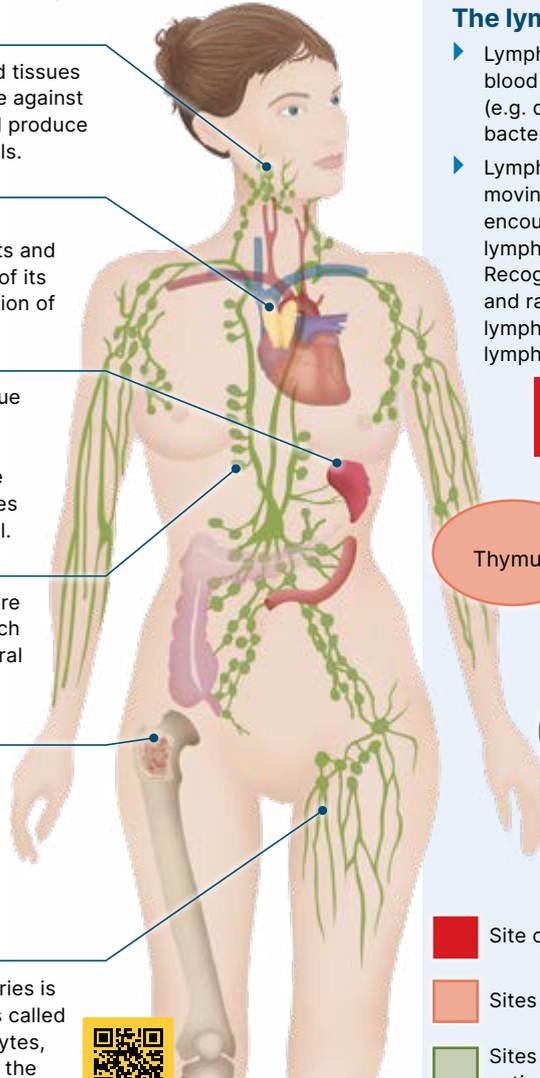
Ovoid masses of lymph tissue where lymphocytes are concentrated. Each node receives lymph through several incoming and outgoing vessels.

### Red bone marrow

A primary lymphoid tissue where all the different kinds of blood cells (including white blood cells) are produced by cellular differentiation from stem cells. B lymphocytes also mature here.

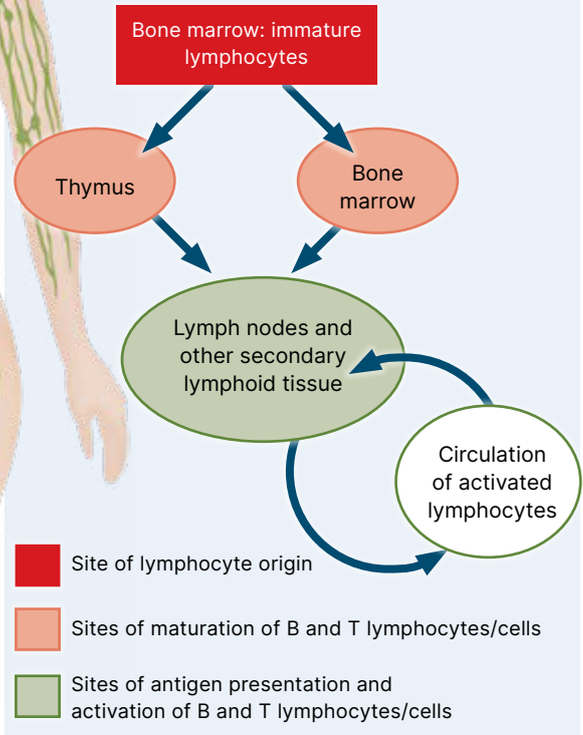
### Lymphatic vessels

When the fluid leaking from capillaries is picked up by lymph capillaries, it is called lymph. The lymph, carrying leucocytes, flows in lymphatic vessels through the secondary lymphoid tissues.



## The lymphatic system and immunity

- ▶ Lymphocytes (T and B cells) are types of white blood cells. They are important in fighting infection (e.g. destroying the microbes causing viral or bacterial infections).
- ▶ Lymphocytes in circulation are constantly moving between sites where antigens may be encountered. The antigens are presented to T lymphocytes in the secondary lymphoid tissues. Recognition of the antigen leads to activation and rapid increase in number of both T and B lymphocytes. After several days, antigen-activated lymphocytes begin leaving the lymphoid tissue.



1. What is the general role of the lymphatic system in immunity? \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_

2. (a) What is the role of lymph nodes in the immune response? \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_

(b) Suggest why lymph nodes become swollen when someone has an infection?  
 \_\_\_\_\_  
 \_\_\_\_\_



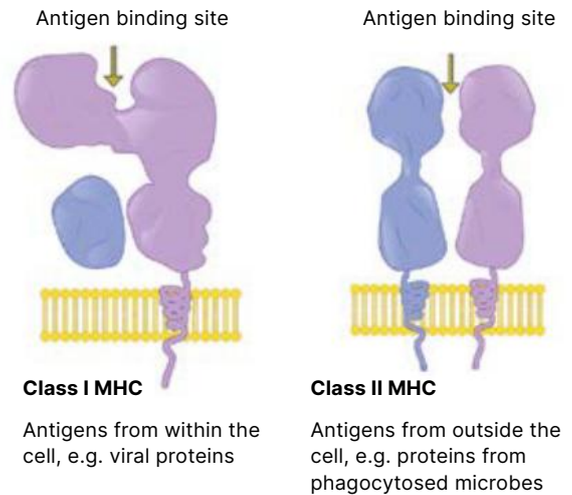
# Processing Antigens

**Key Idea:** Antigen processing prepares and displays antigens for presentation to the T lymphocytes of the immune system. **Antigen** presenting cells (APCs) process and present antigens for recognition by **T lymphocytes**. During antigen processing, the APC digests the foreign antigen into smaller peptide fragments which are then displayed on the surface of

the APC by **MHC** receptors. The **immune response** evoked by the T lymphocytes depends on which MHC receptor (MHC I or MHCII) is activated. Antigen presentation is necessary for T lymphocytes to recognise infection or abnormal growth and activate other cells of the immune system. Dendritic cells, **macrophages**, and **B lymphocytes** are APCs.

## The role of MHC receptors

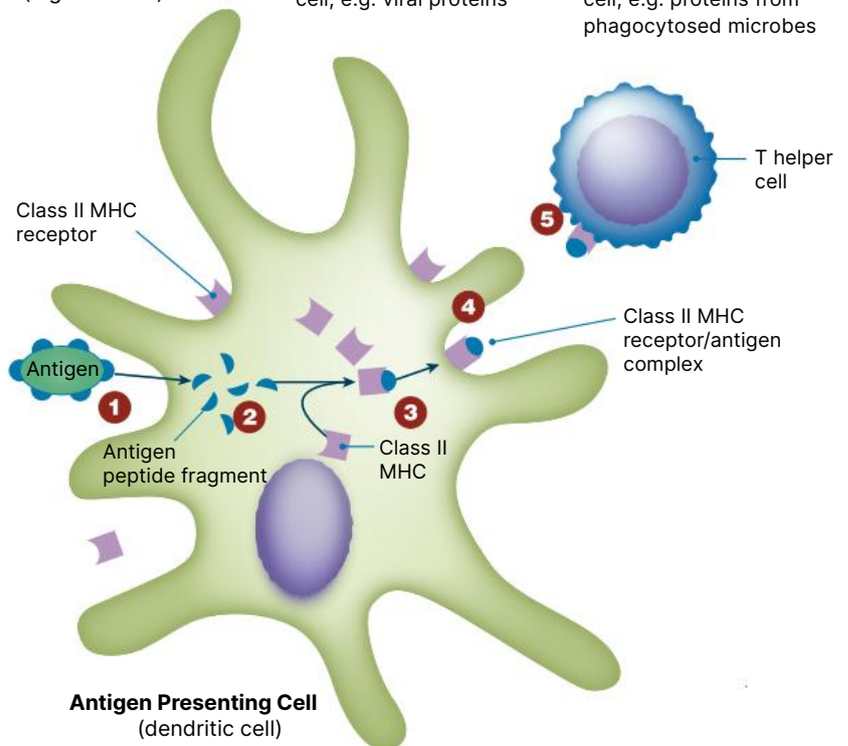
- ▶ Recall there are two types of MHC receptors, class I and class II (right). Both have similar functions in that they display antigens on cell surfaces so that they can be recognised and processed by the T lymphocytes of the immune system. T lymphocytes can only recognise antigens if they are displayed by the MHC receptors. MHC receptors presenting no foreign antigens are ignored by T lymphocytes, and are recognised as "**self**". Only MHC receptors with foreign antigens bound to them will attract T lymphocytes and evoke an immune response.
- ▶ The two classes of MHC receptors display different types of antigens. Class I MHC receptors display antigens from intracellular pathogens (e.g. viruses). Class II MHC receptors display antigens from **pathogens** that have been phagocytosed (e.g. bacteria).



## An overview of antigen processing

The diagram on the right represents antigen processing of an extracellular peptide antigen via a class II MHC receptor.

- 1 An APC encounters an antigen.
- 2 The antigen is engulfed via phagocytosis and digested into short peptide fragments.
- 3 Class II MHC receptors bind the fragments and form a MHC-antigen complex.
- 4 The MHC-antigen complex is displayed on the surface of the APC.
- 5 A receptor on the T helper cell recognises the peptide as foreign. It binds and a series of events stimulate the adaptive immune response.



1. What is the purpose of antigen processing? \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_
2. Why do MHC receptors with no antigenic peptide bound not cause an immune response?  
 \_\_\_\_\_  
 \_\_\_\_\_
3. Describe the differences between class I and class II MHC receptors:  
 \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_

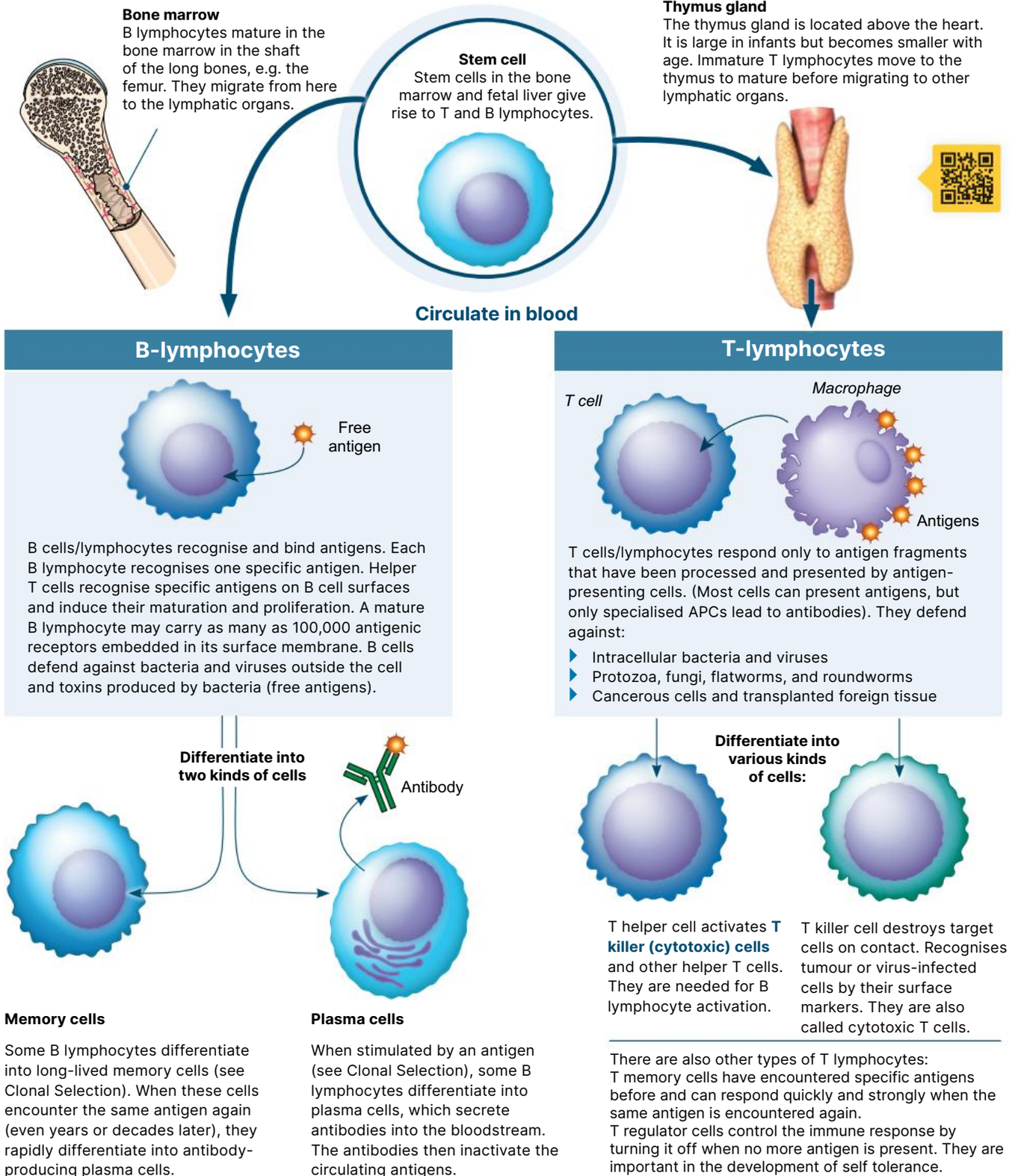


# 183 The Adaptive Immune Response

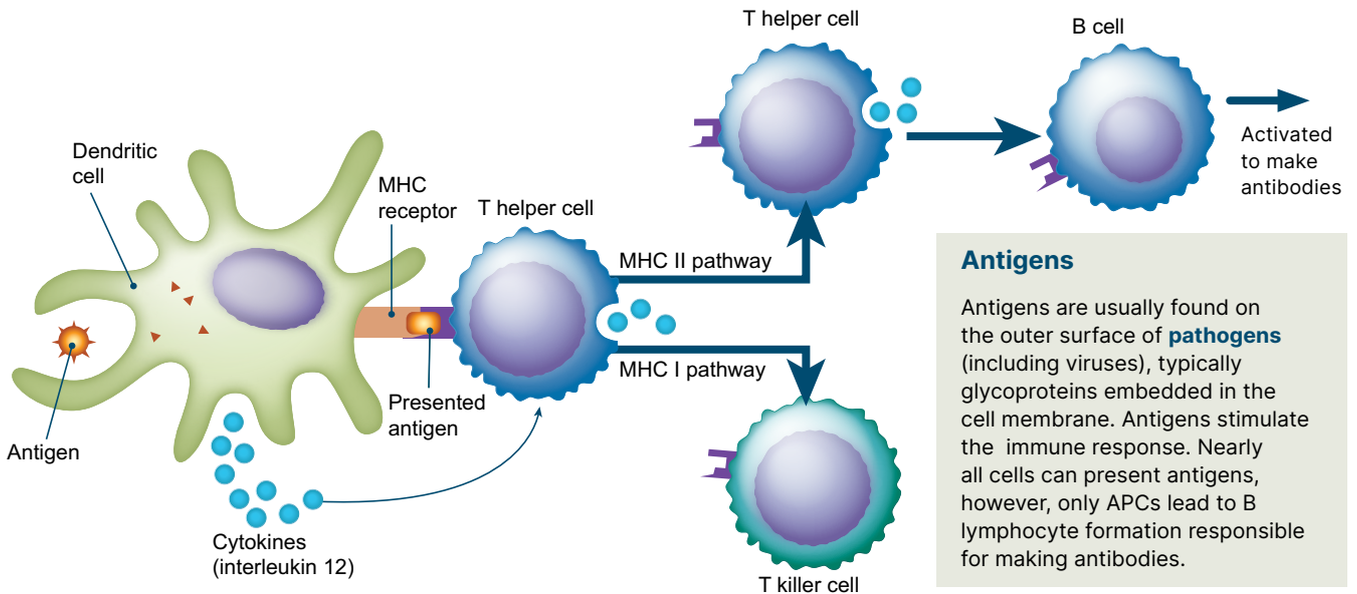
**Key Idea:** Antigens, such as the cell walls of microbial cells, when processed by antigen-presenting cells, activate the B and T cells of the immune system against specific pathogens. There are two main components of the **adaptive immune** system: the humoral and the cell-mediated responses. They work separately and together to protect against disease. The humoral immune response is associated with the serum (the non-cellular part of the blood) and involves the action of **antibodies** secreted by **B lymphocytes**. Antibodies are

found in extracellular fluids including lymph, plasma, and mucus secretions and protect against viruses, and bacteria and their **toxins**. The cell-mediated immune response is associated with the production of specialised lymphocytes called **T cells/lymphocytes**. **Antigens** are recognised by T cells only after antigen processing. The antigen is first engulfed by an antigen-presenting cell, which processes the antigen and presents it on its surface. T helper cells can then recognise the antigen and activate other cells of the immune system.

## Lymphocytes and their functions



## Dendritic cells stimulate the activation and proliferation of lymphocytes



- ▶ Dendritic cells (DC) are antigen-presenting cells. Immature DC originate in bone marrow and migrate through the body to lymph nodes. When a DC encounters an antigen, it presents it to a T helper cell, stimulating it to secrete chemicals called cytokines. Cytokines stimulate the activation and proliferation (rapid increase in number) of T lymphocytes, activating the immune system against that specific antigen. T helper cells go on to stimulate the production of antibody-producing B lymphocytes.
- ▶ Dendritic cells with **MHC I** receptors stimulate the production of T killer cells (also called cytotoxic T cells) (with CD8 receptors).
- ▶ Dendritic cells with MHC II receptors stimulate the production of T helper cells (with CD4 receptors).

1. Where do B lymphocytes and T lymphocytes originate (before maturing)? \_\_\_\_\_
2. (a) Where do B lymphocytes mature? \_\_\_\_\_  
 (b) Where do T lymphocytes mature? \_\_\_\_\_
3. Describe the nature and general action of the two major divisions in the immune system:
  - (a) Humoral immune system: \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_
  - (b) Cell-mediated immune system: \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_
4. Explain how an antigen causes the activation and proliferation of T and B lymphocytes/cells, including the role of dendritic cells:
 

\_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_
5. In what way do dendritic cells act as messengers between the innate and the adaptive immune systems?
 

\_\_\_\_\_

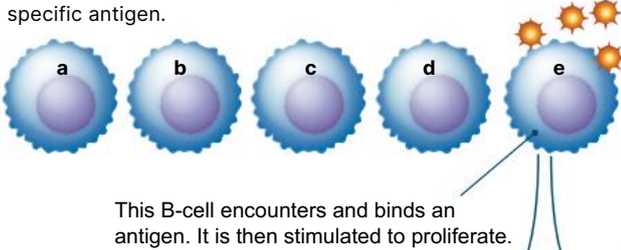
\_\_\_\_\_
6. Describe the function of each of the following cells in the immune system response:
  - (a) T helper cells: \_\_\_\_\_
  - (b) T killer cells: \_\_\_\_\_

# 184 Clonal Selection

**Key Idea:** Clonal selection theory explains how lymphocytes can respond to a large and unpredictable range of antigens. The **clonal selection** theory explains how the immune system can respond to the large and unpredictable range of potential **antigens** in the environment. The diagram below

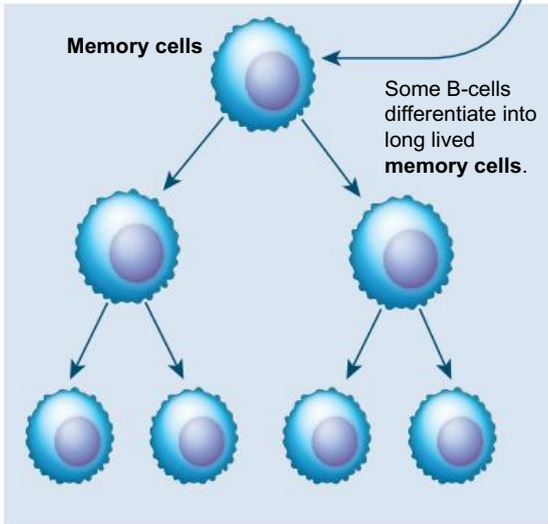
describes clonal selection after antigen exposure for **B lymphocytes**. In the same way, a **T lymphocyte** stimulated by a specific antigen will multiply and develop into different types of T lymphocytes. Clonal selection and differentiation of lymphocytes provide the basis for immunological memory.

Five (a-e) of the many B lymphocytes generated during development. Each one can recognise only one specific antigen.

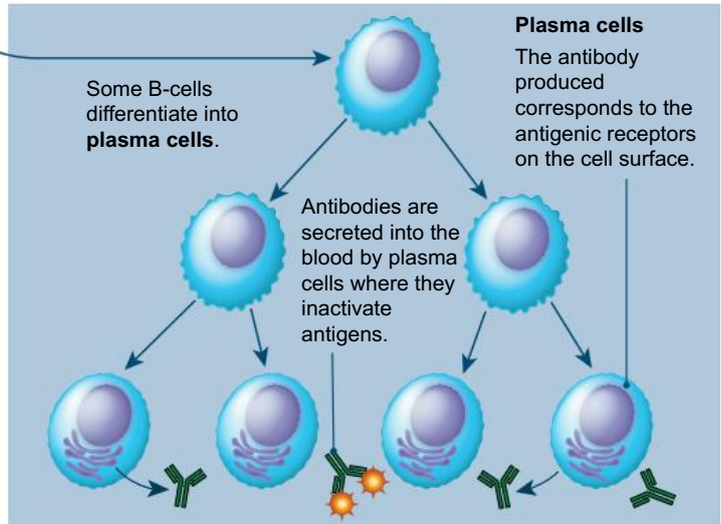


### Clonal selection theory

Millions of B lymphocytes form during development. Antigen recognition is randomly generated, so collectively they can recognize many antigens, including those that have never been encountered. Each B lymphocyte has receptors on its surface for specific antigens and produces antibodies that correspond to these receptors. When a B lymphocyte encounters its antigen, it responds by proliferating and producing many clones that produce the same kind of antibody. This is called clonal selection because the antigen selects the B lymphocytes that will proliferate.



Some B lymphocytes differentiate into long lived memory cells. These are retained in the lymph nodes to provide future immunity (immunological memory). If the antigen returns a second time, memory B cells react more quickly and vigorously than the first time the antigen appeared.



Plasma cells secrete antibodies specific to the antigen that stimulated their development. Each plasma cell lives for only a few days, but can produce about 2000 antibody molecules per second. During development, any B lymphocytes that react to the body's own antigens are destroyed in a process that leads to self tolerance (acceptance of the body's own tissues).

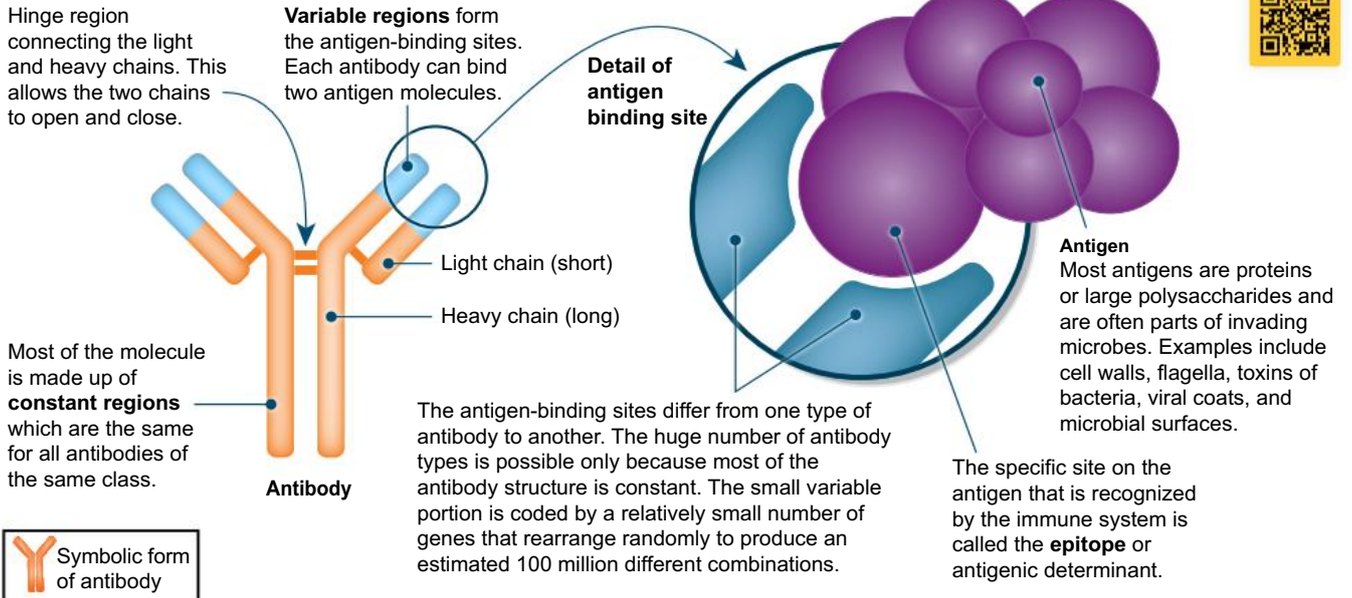
- Describe how clonal selection results in the proliferation of one particular B cell clone: \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_
- What is the function of the plasma cells in the immune system response?  
 \_\_\_\_\_
  - What is the significance of B lymphocytes producing antibodies that correspond to their antigenic receptors?  
 \_\_\_\_\_  
 \_\_\_\_\_
- Explain the basis of immunological memory: \_\_\_\_\_  
 \_\_\_\_\_
  - Why are B memory cells able to respond so rapidly to an encounter with an antigen long after an initial infection?  
 \_\_\_\_\_  
 \_\_\_\_\_

# 185 Antibodies

**Key Idea:** Antibodies are large, Y-shaped proteins, made by plasma cells, which destroy specific antigens.

**Antibodies** and **antigens** play key roles in the response of the immune system. Antigens are foreign molecules which promote a specific **immune response**. Antigens include pathogenic microbes and their **toxins**, as well as substances such as pollen grains, blood cell surface molecules, and the

surface proteins on transplanted tissues. Antibodies (or immunoglobulins) are proteins made in response to antigens. They are secreted from B lymphocytes/cells into the plasma where they can recognise, bind to, and help destroy antigens. There are five classes of antibodies, each plays a different role in the immune response. Each type of antibody is specific to only one particular antigen.



## How antibodies inactivate antigens

**Acting as agglutinins**

Soluble antigens

Antibodies can act as agglutinins and cause antigens to bind together, forming inactivated clumps.

**Acting as antitoxins**

Toxins

Antibodies can act as antitoxins by binding to toxins and neutralizing them.

**Enhancing phagocytosis**

Chemical marker

Phagocyte

Antibody

Antigen/bacteria

Tags foreign cells for destruction by phagocytes.

1. Describe the structure of an antibody, identifying the specific features of its structure that contribute to its function:

---

---

---

2. Explain how the following actions by antibodies enhance the immune system's ability to stop infections:

(a) Acting as agglutinins: \_\_\_\_\_

---

(b) Acting as antitoxins: \_\_\_\_\_

---

(c) Tagging foreign cells with chemical markers: \_\_\_\_\_

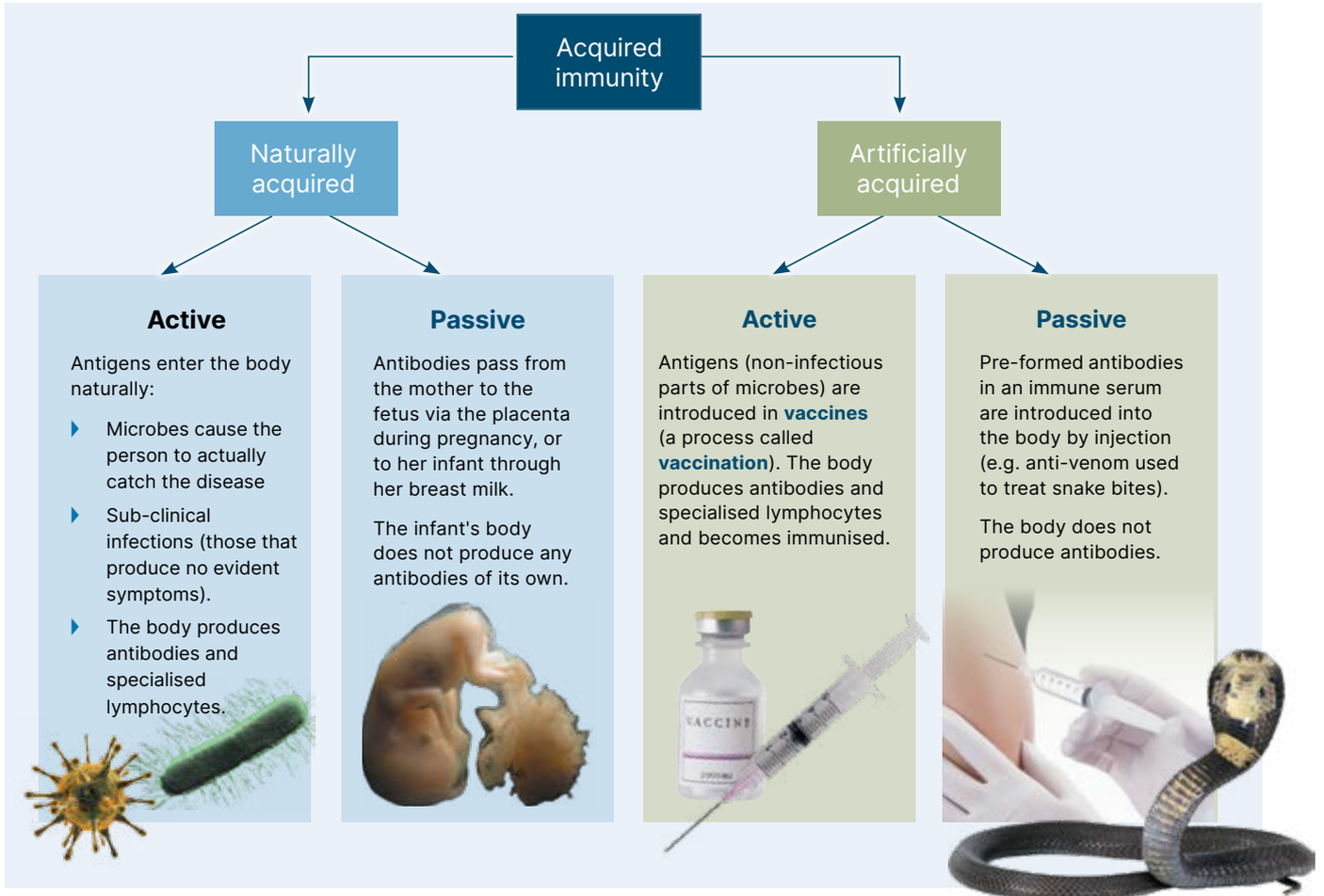
---



# 186 Acquired Immunity

**Key Idea:** Acquired immunity is a resistance to specific pathogens acquired over the life-time of an organism. We are born with natural or innate resistance which provides non-specific immunity to certain illnesses. In contrast, acquired immunity is protection developed over time to specific **antigens**. Active immunity develops after the

immune system responds to being exposed to microbes or foreign substances. Passive **immunity** is acquired from gaining preformed **antibodies** without exposure to the antigen. Immunity can be naturally acquired, through natural exposure to microbes, or artificially acquired as a result of medical treatment (below).



- (a) What is meant by passive immunity? \_\_\_\_\_

\_\_\_\_\_

(b) Distinguish between naturally and artificially acquired passive immunity and give an example of each:

\_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_
- (a) Why does a newborn baby need to have received a supply of maternal antibodies prior to birth?

\_\_\_\_\_

\_\_\_\_\_

\_\_\_\_\_

(b) Why is this supply supplemented by antibodies in breast milk? \_\_\_\_\_

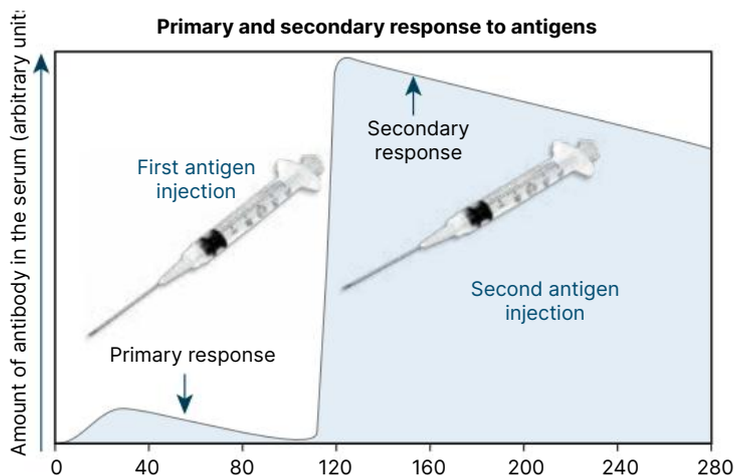
\_\_\_\_\_
- (a) What is active immunity? \_\_\_\_\_

\_\_\_\_\_



## Primary and secondary response to antigens

- ▶ When the **B lymphocytes** encounter antigens and produce antibodies, the body develops active immunity against that antigen.
- ▶ The initial response to antigenic stimulation, caused by the sudden increase in B lymphocyte clones, is called the primary response. Antibody levels as a result of the primary response peak a few weeks after the response begins and then decline. However, because the immune system develops an immunological memory of that antigen, it responds much more quickly and strongly when presented with the same antigen subsequently (the secondary response).
- ▶ This forms the basis of immunisation programmes where one or more booster shots are provided following the initial vaccination.



(b) Distinguish between naturally and artificially acquired active immunity and give an example of each:

---



---



---

4. (a) Describe two differences between the primary and secondary responses to presentation of an antigen:

---



---



---

(b) Why is the secondary response so different from the primary response?

---

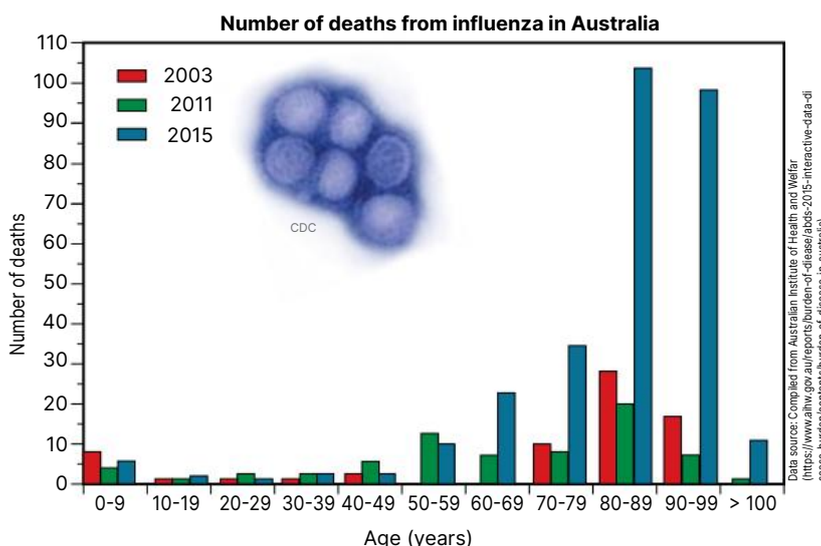


---



---

5. Some diseases do not affect all members of a population equally. Socioeconomic factors, age, sex, ethnicity and where someone lives can influence how a disease affects a particular individual or population. The data (right) shows deaths from influenza (flu) in Australia by age over three years.



(a) Do you think the data shows an age related effect for influenza deaths? Explain your reasoning:

---



---



---



---



---

(b) Suggest what could be done to help reduce the number of influenza deaths:

---



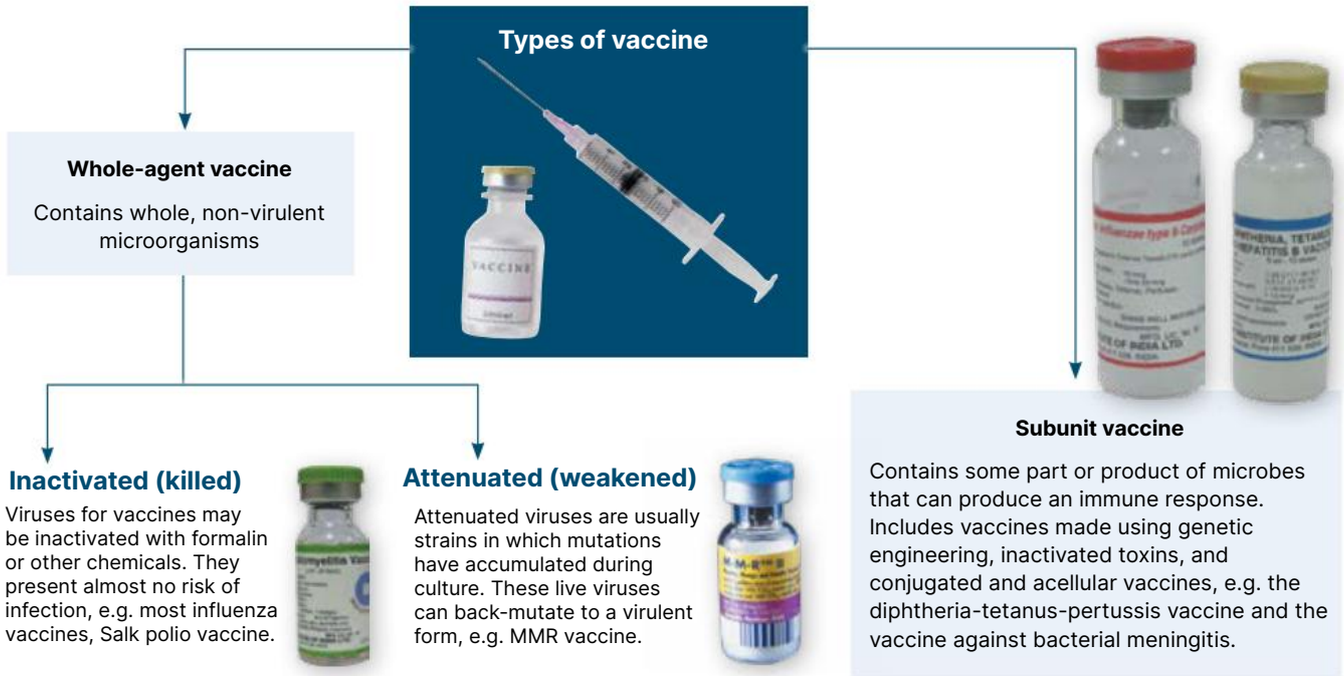
---

# 187 Vaccines and Vaccination

**Key Idea:** A vaccine is a suspension of antigens that is deliberately introduced into the body to protect against disease. If enough of the population are vaccinated, herd immunity provides protection to unvaccinated individuals.

A **vaccine** is a preparation of a harmless foreign **antigen** that is deliberately introduced into the body to protect against a specific disease. The antigen in the vaccine is usually some part of the **pathogen** and it triggers the immune system to

produce **antibodies** against the antigen, but it does not cause the disease. The immune system remembers its response and will produce the same antibodies if it encounters the antigen again. If enough of the population are vaccinated, herd **immunity** (indirect protection) provides unvaccinated individuals in the population with a measure of protection against the disease. There are two basic types of vaccine, subunit vaccines and whole-agent vaccines (below).



## Why are vaccinations given?



Vaccines against common diseases are given at various stages during childhood according to an immunisation schedule. **Vaccination** has been behind the decline of some once-common childhood diseases, such as mumps and measles.



Most vaccinations are given in childhood, but adults may be vaccinated against a disease, e.g. TB, tetanus, if they are in a high risk group, e.g. the elderly or farmers, or to provide protection against seasonal diseases such as influenza.



Tourists may need specific vaccines if the country they are visiting has a high incidence of a certain disease. For example, travellers to South America should be immunised against yellow fever, a disease that does not occur in many other countries.

1. (a) What is a vaccine? \_\_\_\_\_

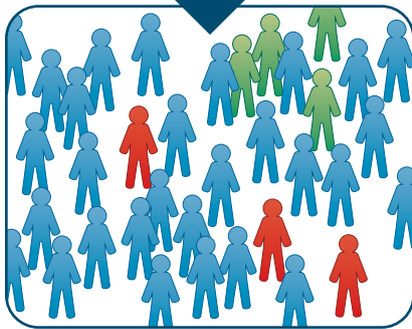
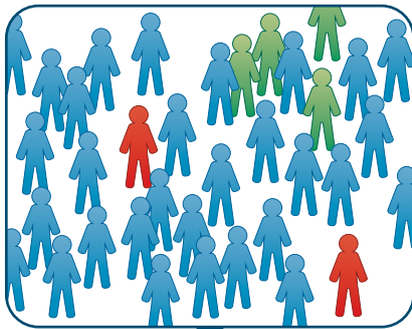
(b) Provide some examples of when vaccinations are needed: \_\_\_\_\_



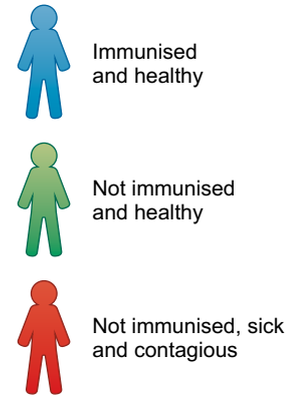
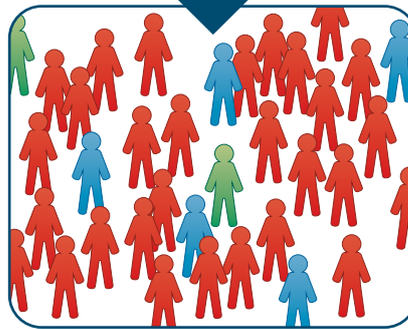
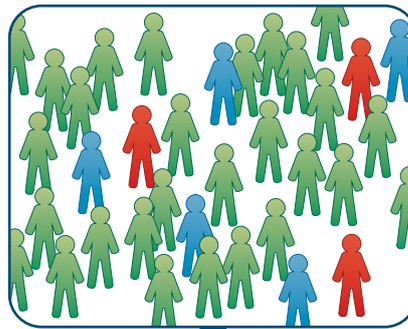
## Vaccination can provide herd immunity

- ▶ Herd immunity occurs when the vaccination of a significant portion of a population provides some protection for individuals who have not developed immunity, e.g. have not been vaccinated and are not immunised. In order to be effective for any particular disease, a high percentage of the population needs to be vaccinated against that disease. High vaccination rates make it difficult for the disease to spread because there are very few susceptible people in the population.
- ▶ Herd immunity is important for people who cannot be vaccinated, e.g. the very young, people with immune system disorders, or people who are very sick, such as cancer patients.

High herd immunity: Most of the population is immunised. The spread of the disease is limited. Only a few people are susceptible and become infected.



Low herd immunity: Only a small proportion of the population is immunised. The disease spreads more readily through the population, infecting many more people.



### DID YOU KNOW?

The level of vaccination coverage to obtain herd immunity differs for each disease. Highly contagious diseases (e.g. measles) need a much higher vaccine uptake (95%) than a less contagious disease such as polio (80-85%).



2. Attenuated viruses provide long term immunity to their recipients and generally do not require booster shots. Why do you think attenuated viruses provide such effective long-term immunity when inactivated viruses do not?

---



---



---



---

3. (a) What is herd immunity? \_\_\_\_\_

---



---

- (b) Why are health authorities concerned when the vaccination rates for an infectious disease fall?

---



---

4. Some members of the population are unable to be vaccinated. Give an example and explain why herd immunity is very important to them:

---



---



---

# 188 Vaccines Can Eliminate Infectious Disease

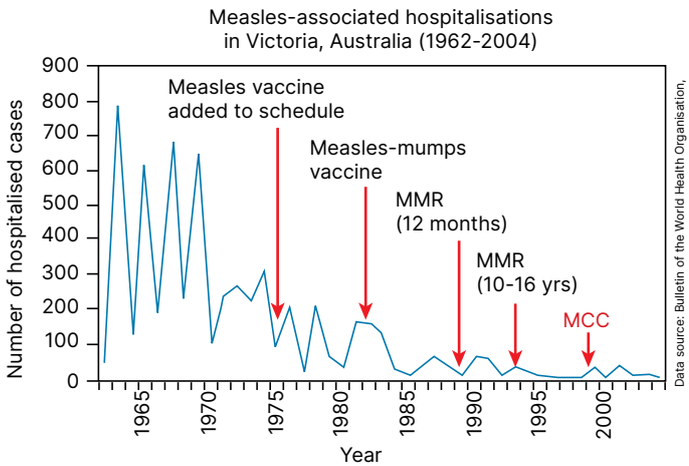
**Key Idea:** Vaccination programmes have been successful in the global eradication of smallpox, but other diseases (such as influenza) are more difficult to eradicate.

To date, the only infectious disease globally eradicated has been smallpox. Several factors led to this success. Smallpox is easily identifiable by its characteristic rash making surveillance and containment of infected patients easier.

It has no other natural carriers so once immunisation rates reached a critical level, its spread through the population was limited. Other diseases can be more difficult to eradicate. This is especially true for diseases that have a long period between infection and the symptoms showing (e.g. TB) or diseases caused by **pathogens** with high rates of mutation (e.g. *influenzavirus* or HIV).

## Measles elimination in Australia

Measles is a highly contagious disease, one infected person can infect 12-18 people during their infectious period. In 2014, the World Health Organisation (WHO) announced measles had been eliminated from Australia. High **vaccination** rates contributed to its elimination. However, measles still occurs in other countries so it could be reintroduced if an infected traveller entered Australia. Maintaining high levels of vaccination is important in preventing its reintroduction.



The graph above shows the role of vaccination in reducing measles hospitalisations in the state of Victoria. MMR is the introduction of the measles/mumps/rubella vaccine. MCC (measles control campaign) was an extensive mass vaccination and monitoring campaign.

## The challenges of eradicating disease



CDC

Whooping cough (above) is a respiratory disease caused by the bacterium *Bordetella pertussis*. Despite high vaccination rates, whooping cough is increasing in Australia. Several factors may be contributing to this.

- ▶ Until 1997, a whole **vaccine** was used. It contained hundreds of different antigens and provided protection against many strains of the pertussis pathogen. In 1999, an acellular vaccine, which does not contain the whole pathogen, was introduced (inset above). It only contains five antigens and so provides less protection.
- ▶ New strains of *B. pertussis* are evolving, and the new vaccine is not effective against them.
- ▶ More adults who were vaccinated against whooping cough in childhood are contracting the disease. This suggests the effectiveness of the vaccine declines over time.

1. The graph above provides long term immunity data for measles in Victoria. Use this data to provide evidence for the role of vaccination programmes in eliminating measles from Australia:

---



---



---



---

2. What could happen if vaccination rates for measles fell too low? \_\_\_\_\_

---

3. (a) Why could the change to a new vaccine have affected the rates of whooping cough in Australia? \_\_\_\_\_

---



---

(b) Why do you think a new vaccine was introduced? \_\_\_\_\_

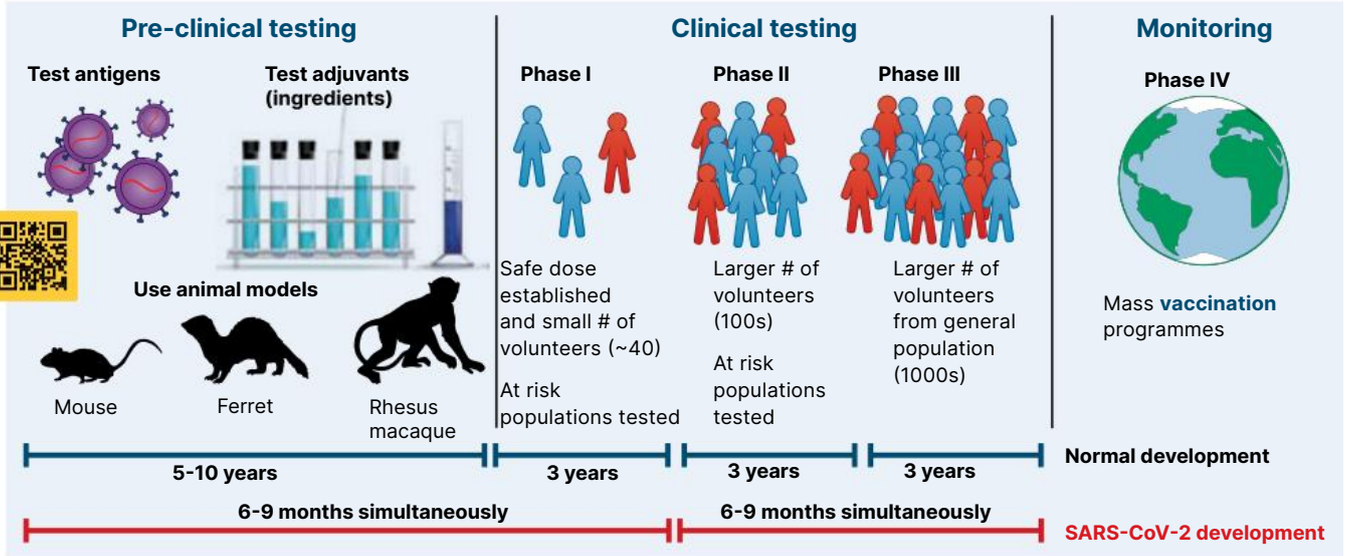


# Vaccine Development

**Key Idea:** Vaccine development typically proceeds through the same stages but this can be accelerated if the need for the vaccine is urgent.

The development of a Covid-19 **vaccine** became a global priority when the seriousness of the virus was recognised. In

order to produce the vaccine fast enough for it to be useful, vaccine development was fast tracked. A number of Covid-19 vaccines have undergone the same rigorous development and testing process as other vaccines, but over a shorter time period. A number have been approved as safe to use.



## Contributions to vaccine development

Advancing science understanding often involves collaboration across various fields. Vaccine development is one such scientific undertaking that involves scientists building their understanding on the findings of others. Some notable Nobel Prize recipients were recognised for their work in the immunity and vaccine areas. In 1977, American physicist Rosalyn Sussman Yalow received the Nobel Prize in Physiology or Medicine for her work on a radio-immunoassay (RIA) technique that enhanced the detection and monitoring of infectious and hormone-related diseases. Importantly, she was able to show how tiny protein particles stimulated the formation of antibodies. In 1996, Queensland immunologist Peter C Doherty and Swiss scientist Rolf M Zinkernagel were both honoured with the Nobel Prize for their contributions to enhancing our knowledge of the cell-mediated immune response, specifically the recognition of virus infected cells by the immune system. Their work was important for further vaccine development.



Rosalyn Yalow

US Information Agency Public Domain

1. Explain how the Covid-19 vaccines were developed in a much shorter time than most other vaccines:

---

---

---

---

2. Transparency and peer review is an important part of research. Early research results were released from various Covid-19 vaccination development teams so that other scientists could replicate the experiments.

(a) Why is peer review an important part of scientific methodology (process)? \_\_\_\_\_  
\_\_\_\_\_  
\_\_\_\_\_

(b) What are some potential issues of media publishing early results on Covid-19 vaccination development before the research has been properly evaluated?  
\_\_\_\_\_  
\_\_\_\_\_  
\_\_\_\_\_



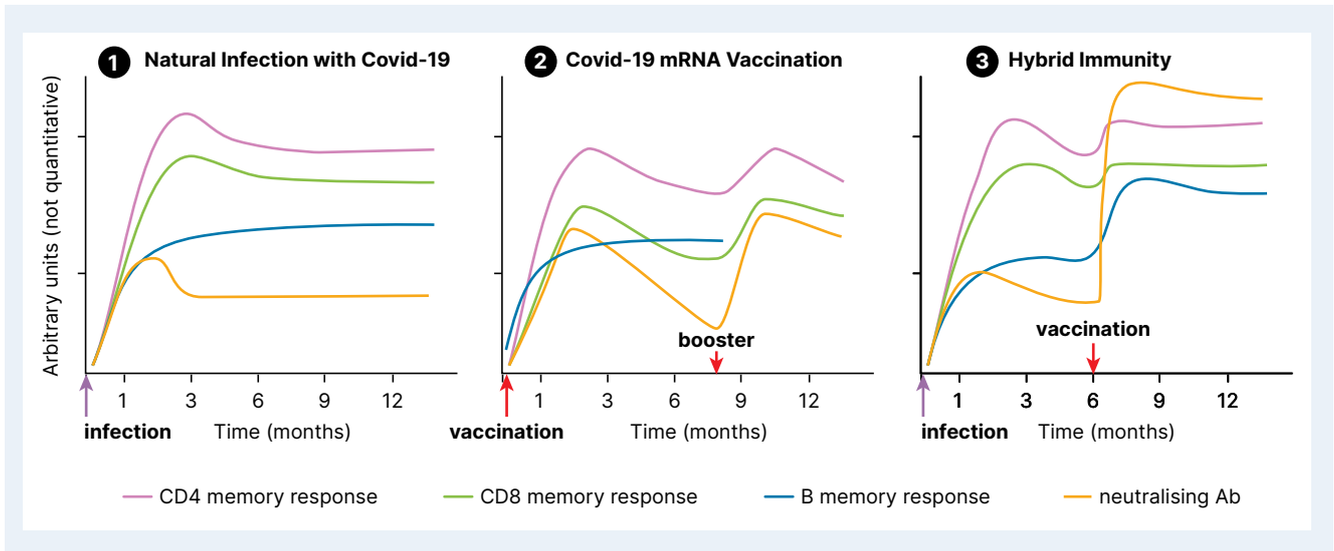
**Key Idea:** Many vaccines require regular boosting to maintain consistent and effective immunity.

Viruses that mutate rapidly each season into different variants, such as flu (influenza) and Covid-19 (SARS-CoV-2) require regular vaccination to provide effective protection from infection, serious symptoms, and consequent spread. As time since vaccination increases, the vaccine-induced **immunity** declines, especially if new viral mutants are circulating in the community. Each season, **vaccines** are

developed to match the predicted variants of virus that are likely to present 6-12 months in the future, especially in the winter months. The southern hemisphere tends to be 6 months behind the northern hemisphere in variant spread. Vaccines for many childhood illness also wear off over time, which is why immunisation schedules include boosters to provide long term protection. Immunologists use immune response data, like the examples in this activity, to help set effective vaccination schedules.

## Comparing natural infection and mRNA vaccination for Covid-19

- ▶ Different immunity blood cells activate a different, and sometimes partially overlapping, response to infection and vaccination as part of the immune system.
- ▶ Natural infection and vaccination cause different long-term responses in acquired immunity, but both provide protection. Both show a rapid increase in immunity followed by a dip after a peak. In the case of Covid-19 / SARS-CoV-2 the immunity dip continues to fall with mRNA vaccination, whereas the immunity from infection plateaus and remains stable for at least a year.
- ▶ Immune response can be determined using a range of different markers. The graphs below show the responses of four different markers as measurements of the immune response. You do not need to know the specifics of these markers, just the general trends produced.
- ▶ The graphs below show the immune response from natural Covid-19 infection, after Covid-19 vaccination, and following Covid-19 infection and a vaccination (hybrid immunity).



Data source: Petrone et al. (2023) 'The Importance of Measuring SARS-CoV-2-Specific T-Cell Responses in an Ongoing Pandemic' CC BY 4.0

1. How does the immune response from natural immunity differ from the immunity provided by the COVID-19 vaccine?

---

---

---

---

---

---

---

---

2. Study graphs 2 and 3 above. Is getting a COVID-19 booster after the initial vaccination more effective than getting vaccinated after recovering from a natural COVID-19 infection? Explain your reasoning.

---

---

---

---

---

---

---

---

## Long term immune response to the whooping cough vaccine

Whooping cough (pertussis) is a highly contagious bacterial infection affecting the respiratory system. It often begins with cold-like symptoms followed by a cough that gets worse and may last for months. Young children are most often affected. Whooping cough can cause serious health problems, and sometimes death, especially in unvaccinated young babies and children.

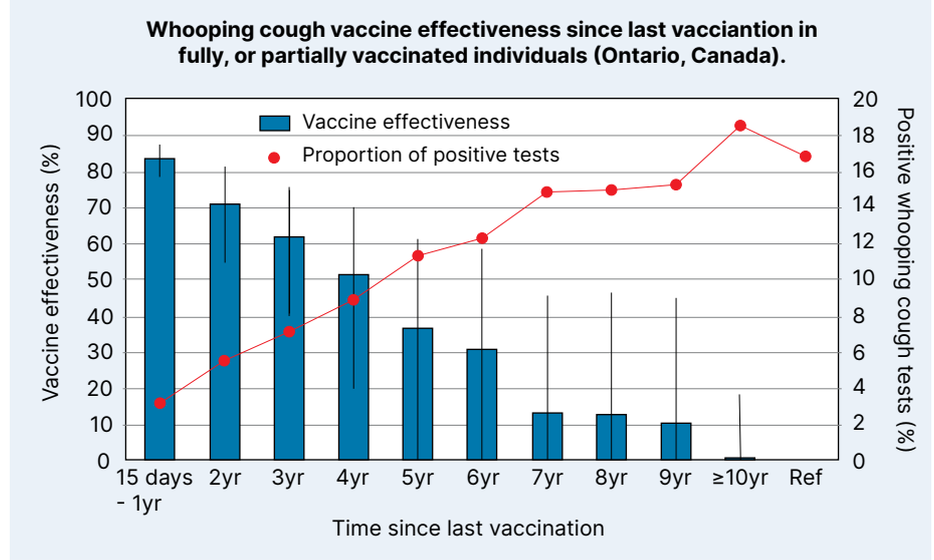
A vaccine for whooping cough was introduced into the Australian immunisation schedule in 1942, significantly reducing whooping cough infections and deaths. The current Australian vaccination schedule recommends vaccination against the disease for children at 4, 6 and 18 months, 4 years of age, and in adolescents at 11–13 years of age. Boosters for adults are recommended at ages 50 years and 65 years as well as pregnant women (ideally between 20–32 weeks).

The graph on the right shows the effectiveness of the whooping cough vaccine over time in participants with up-to-date or partial vaccination, in Ontario, Canada.

Results are measured from the last vaccine, and are compared to unvaccinated individuals (Ref on the graph). Error bars represent 95% CI.

The proportion of positive tests measures whooping cough cases within the study's duration.

Ontario's immunisation schedule recommends the vaccine is given at 3, 5, 7 and 19 months, and again at ages 7 and 17 years of age.



Data source: Schwartz K.L. et al. CMAJ November 01 2016; 188 (16): E399-E406; DOI: https://doi.org/10.1503/cmaj.160193

3. Study the graph (above) of whooping cough vaccine effectiveness over time.

(a) Describe how vaccine effectiveness (blue bars) changes over time: \_\_\_\_\_

---



---



---



---

(b) Describe how the proportion of positive tests for whooping cough (red line) changes over time:

---



---



---

(c) Describe the relationship between vaccine effectiveness and positive whooping cough tests: \_\_\_\_\_

---



---



---

4. (a) Why do you think the immunisation schedule includes whooping cough boosters for children at age 11-13 and then again for adults?

---



---



---

(b) What do you think would happen to whooping cough infections if the boosters were not given?

---



---

# 191 Physical Defences in Plants

**Key Idea:** Physical defences against pathogens and predators include physical barriers such as thick outer bark and leaf cuticle, and mechanical defences such as nastic responses in plants.

Most plants have a tiered system of defences to provide resistance against disease. Physical defences are the first line of defence against **pathogens**. Physical defences are

a non-specific form of defence; they are always present and are designed to stop as many pathogens as possible from entering the plant. Physical barriers to protect plants from pathogens include spikes, thorns, hairs, and thick waxy cuticles on leaves. Some plants have behavioural mechanisms that can protect once stimulated.

## Physical defences in plants

- ▶ Plants have physical defences that protect them from infection by pathogens.
- ▶ Many trees and shrubs are covered in bark. This tough external covering forms a physical barrier, which pathogens find difficult to cross.
- ▶ The addition of pectin (a type of carbohydrate) and a layer of cellulose (the cell wall) adds strength and has the added benefit of providing extra layers to protect against infection.
- ▶ Many plants have leaves covered in a waxy cuticle. The cuticle is another physical barrier, reducing pathogen entry into the plant tissue via the leaf.



Juniper wattle

### Woody stems and bark

Trees and woody shrubs, like the juniper wattle, have thick coverings of waxy suberin, which limits pathogen entry to the inner tissues. The spiny, reduced leaves protect it from being eaten, but also limit the pathways for pathogen entry.



Hoya, an Australian native

### Impervious waxy cuticle

All plants have a waxy cuticle, even if it is thin. The cuticle is made up of fatty acids so all leaves are negatively charged and hydrophobic. These properties repel many spores and microbes and make the leaf environment less suitable for fungal invasion.



Unstimulated leaf



Disturbed leaf

## Nastic antipredator mechanism of the Mimosa plant

- ▶ The sensitive plant (*Mimosa pudica*) has long leaves composed of small leaflets. When a leaf is touched, it collapses and its leaflets fold together. Strong disturbances cause the entire leaf to droop from its base. This response takes only a few seconds and is caused by a rapid loss of turgor pressure from the cells at the bases of the leaves and leaflets.
- ▶ The message that the plant has been disturbed is passed quickly around the plant by electrical signals (changes in membrane potential), not by plant hormones (as occurs in tropisms). The response can be likened to the nerve impulses of animals, but it is much slower. After the disturbance is removed, turgor is restored to the cells, and the leaflets slowly return to their normal state.
- ▶ The adaptive value of these responses can dislodge any attacking insects or deterring browsers. However, there is an energetic cost to the plant, and regular leaf folding can inhibit photosynthesis.

1. What features of physical barriers make them an important first line of defence against pathogens?

---



---

2. What are the similarities between the way human skin and plant bark reduce pathogen entry?

---



---

3. Some plants leaves secrete wax in response to the presence of pathogens. How is this different from the other physical barriers in plants?

---



---

4. What is the purpose of the mimosa plant leaf collapse?

---



---

**Key Idea:** Plants have chemical defences to defend against pathogens. Some mechanisms are always present while others are stimulated by an attack.

Living organisms are under constant attack from **pathogens**. As a result, plants have evolved a wide range of chemical defences to protect themselves from pathogens and limit the damage they can do. Plants have **innate** or non-specific defences only. The plants contain peptides called **defensins**,

present in the common ancestor of both plants and animals, that exhibit both antibacterial and antimicrobial functions. The chemical defences of plants not only protect them from attack by pathogens, but include the production of **toxins** that may also stop animals eating them or inhibit the growth of other plants. Many herbs produce toxins with antimicrobial properties. These compounds are sometimes extracted for human use.

## Chemical defence strategies in plants: plant toxins

- ▶ Many plants produce a range of antimicrobial and antifungal toxins and enzymes to kill or inhibit the growth of pathogens. Some of these compounds cover the surface of the plant, killing pathogens before they enter the plant. Other compounds act internally.
- ▶ The type of toxin depends on the plant. Cyanide-based toxins are found in apricot seeds, flax seeds (*Linum usitatissimum*), bamboo shoots, lima beans, and sorghum, evolving independently in different plant families. The cyanide toxin is stored safely in the plant by being covered with a glucose 'coat'. When the plant is eaten, the digestive process of the animal removes the coat, the toxin is activated, and the animal is exposed to the harmful toxin.

### Passive defences

**Passive defences** are always present and are not the result of contact with a pathogen or grazer. Plants have both physical and chemical defences to deter pathogens. For example, the thick waxy surface of many leaves (right) acts as a physical barrier to limit pathogen entry. However, if the physical defence is breached, the chemical defences protect the plant against further damage.



Apricots and stones



Sorghum

Flaxseed  
*Linum usitatissimum*



### Hemlock

- ▶ The hemlock plant (*Conium maculatum*) (image, right) is a member of the carrot family but, unlike the carrot, it contains a powerful toxin called coniine, deadly to humans and animals in small doses. Around 8 leaves would contain a fatal dose for humans. The seeds and roots are even more toxic than the leaves.
- ▶ The toxin is structurally similar to nicotine and blocks the same nicotinic acetylcholine receptors on neuromuscular junctions of neurons (which control skeletal muscle contractions).
- ▶ Poisoning by the hemlock toxin causes failure of the respiratory muscles and eventually results in death. Victims fall into a coma if breathing is not artificially carried out.

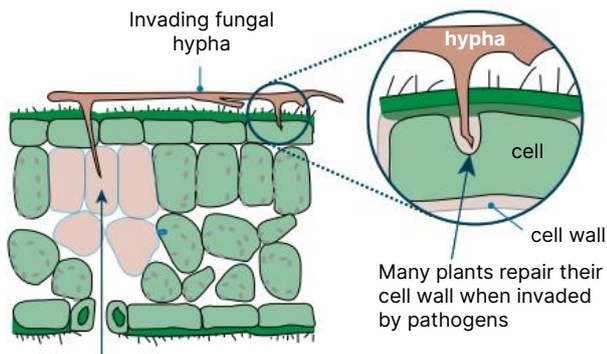


Hemlock

1. Why do plants produce toxins as a defence mechanism? \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_
2. How do plant toxins such as cyanide-based compounds protect plants from threats?  
 \_\_\_\_\_  
 \_\_\_\_\_
3. Why is hemlock toxin so harmful to animals? \_\_\_\_\_  
 \_\_\_\_\_

## Chemical defence strategies in plants: defensins

- ▶ Once infected, a plant responds actively to prevent any further damage. Active defences are invoked only after a pathogen has been recognised, or after wounding or attack by a herbivore. This makes biological sense because active defences are costly to produce and maintain.
- ▶ Active plant defences work via a variety of mechanisms, but the key innate immune system is based on substances called **defensins**. These are composed of peptides (chains of amino acids) and vary in structure in different plant species. Defensins are also present in animals and fungi, which indicates they evolved in a common ancestor to all three groups.
- ▶ Defensins have a range of antibacterial and antifungal functions including slowing pathogen growth, damaging fungal membranes, puncturing the bacterial cell wall, disrupting metabolism and the cell cycle of pathogens, or killing cells by release of reactive oxygen species such as hydrogen peroxide ( $H_2O_2$ ). Some defensins can also disrupt DNA replication and transcription during reproduction of pathogens.
- ▶ Recent research has uncovered some types of defensin chemicals that can also deter insect herbivory by acting as enzyme inhibitors that interfere with carbohydrate metabolism, causing insect growth retardation.



Many plants produce an enzyme-activated hypersensitive response led by defensin substances when invaded by pathogens. This leads to the production of reactive nitric oxide and cell death. Cell death (apoptosis) in the infected region limits the spread of the pathogen.

### Plant galls



Another chemical plant defence strategy involves sealing off infected areas giving rise to abnormal swellings called galls (oak gall, left and bulls-eye galls on a maple leaf, right). These galls limit the spread of the parasite or the infection in the plant. The irritation that initiates gall formation can arise from viruses, fungi, bacteria, insects or even other plants. Some plants can even develop a mutualistic relationship by providing the gall as a microhabitat for an insect. For example, the fig wasp is a pollinator of the fig which develops a gall where the insect safely broods.

4. (a) Distinguish between passive and active defence mechanisms in plants: \_\_\_\_\_

---



---



---

(b) Why are most plant defensive chemicals produced only after a pathogen is detected? \_\_\_\_\_

---



---



---

5. How are galls effective in reducing the spread of infection in some plants? \_\_\_\_\_

---



---



---

6. What are plant defensins and what other groups have them? \_\_\_\_\_

---



---



---

7. How do defensins act as a defence strategy in plants? \_\_\_\_\_

---



---



---



---



---

1. Contrast the innate and the adaptive immune responses with reference to the basic action and the cells involved:

---

---

---

---

2. The photograph on the right shows the effect of a pathogen infecting a human.



(a) Name the defensive response occurring: \_\_\_\_\_

(b) What is happening to the blood vessels at this location? \_\_\_\_\_

(c) Name the substance responsible for the change in the blood vessels: \_\_\_\_\_

(d) What type of cell is the substance released from? \_\_\_\_\_

3. Contrast physical and chemical defence strategies in plants: \_\_\_\_\_

---

---

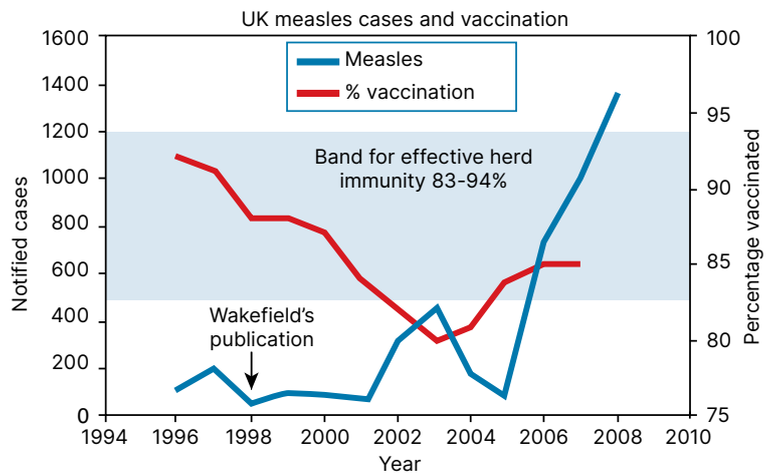
---

---

---

---

4. In 1998, Dr Andrew Wakefield and his colleagues published a paper linking the measles, mumps, and rubella vaccine (MMR) to an increase in autism rates. As a result, the uptake of the MMR vaccine in the UK dropped, and several measles outbreaks occurred. Dr Wakefield's paper was later retracted by the journal in which it was published as it was found to be flawed in several aspects, e.g. sample size of only 12, with no control group. Since the publication of Wakefield's paper, 20 large scale epidemiologic studies into MMR and autism have been carried out in several countries. All have shown that the MMR vaccine does not cause autism. However, the damage was done, and health authorities must continue to convince the public that the vaccine is safe.



The graph above shows the number of measles cases in the UK, together with percentage vaccination, 1994-2008.

(a) What happened to MMR vaccination rates after the publication of Wakefield's study? \_\_\_\_\_

---

---

(b) What is the trend in measles cases in the UK since 2006? \_\_\_\_\_

---

---

(c) Give a likely explanation for this trend: \_\_\_\_\_

---

---

---



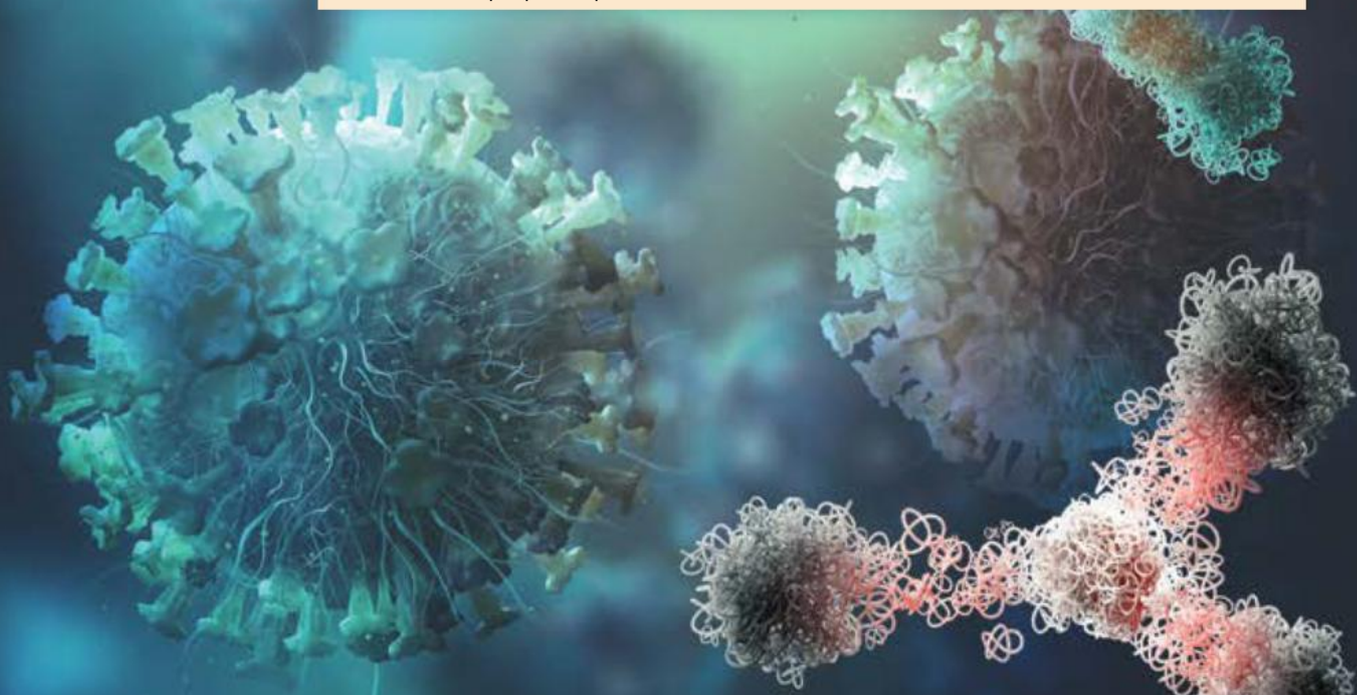
# Transmission and Spread of Disease

- Key Terms**
- antibiotic
  - antiseptic
  - biosecurity
  - bush medicine
  - contact tracing
  - disease
  - epidemic
  - host
  - infectious disease
  - immunisation
  - outbreak
  - pandemic
  - pathogen
  - quarantine
  - spread (disease)
  - transmission (disease)
  - vaccination
  - vaccine
  - vector

- Key Concepts**
- ▶ Many factors contribute to the transmission and spread of disease through populations.
  - ▶ Data can be used to predict outbreaks, identify their source, infer transmission modes, and evaluate control strategy effectiveness.
  - ▶ Biosecurity precautions reduce the likelihood of disease transmission between countries.

Transmission of disease		Activity Number
<input type="checkbox"/> 1	Describe modes of disease transmission, including direct contact, contact with body fluids, contaminated food, contaminated water and disease-specific vectors.	194
<input type="checkbox"/> 2	<b>SI:</b> Investigate the effect of antibiotics on the growth of a microorganism in either a laboratory or virtual context.	195
<input type="checkbox"/> 3	<b>SI:</b> Compare different handwashing techniques for removal of microorganisms from the hands.	197
<input type="checkbox"/> 4	<b>SI:</b> Investigate, giving examples, how the transmission of disease is aided by the regional and global movement of organisms.	194, 196

Spread of disease and epidemiology		Activity Number
<input type="checkbox"/> 5	Model disease outbreak and spread in a digital simulation.	198
<input type="checkbox"/> 6	Identify and explain the interrelated factors that affect spread of disease, including the transmission mechanism, the immunity status of the population, and the mobility and density of individuals in the population.	199
<input type="checkbox"/> 7	Evaluate strategies to control the spread of disease, including public health measures.	200
<input type="checkbox"/> 8	<b>SHE:</b> Research the long-standing use of natural antiseptics and bush medicines by First Nations peoples to prevent and treat infections.	203
<input type="checkbox"/> 9	<b>SHE:</b> Explain how Australia can safeguard its agriculture industry and environment by implementing quarantine measures.	201
<input type="checkbox"/> 10	<b>SI:</b> Investigate the effectiveness of health programs in preventing and eradicating infectious diseases.	202
<input type="checkbox"/> 11	<b>SI:</b> Explore the use of natural antiseptics and bush medicines by First Nations peoples to prevent and treat infections.	203



# 194 Transmission of Disease

**Key Idea:** Infectious disease can spread rapidly within and between regions given the right conditions.

The human body, like that of other large animals, is constantly exposed to a wide range of potential parasites and **pathogens**. **Transmission** and **spread** of a pathogen

depends on its rate of growth, the density of the **host** population, the mobility of the host population, and the mode of transmission. The transmission of **infectious diseases** can be virtually eliminated by observing hygienic practices, and by providing adequate sanitation.

## Transmission and spread



Most pathogens, once inside the body, multiply rapidly, producing symptoms and making the host infectious within a few days. Others take longer to present symptoms. The infectious period can last from a few days to weeks, but in some cases the host may be infectious for long periods of time.



Human cities can contain millions of people, often living very closely together. In these congested conditions, infectious diseases can spread rapidly, especially if sanitation or personal hygiene is poor, or if seasonal weather produces conditions favourable for spreading the pathogen. High speed transport can help spread a pathogen around a region very quickly.



The mode of transmission affects how quickly a pathogen spreads. Direct person to person contact (i.e. touching) is a slower method of spreading, while spreading via mucus droplets coughed into the air or by animal **vectors** (such as mosquitoes) can help a pathogen spread quickly.

## Portals of entry



### Respiratory tract

The mouth and nose are major entry points for pathogens, particularly airborne viruses, which are inhaled from the expelled mucus of infected people.

Examples: tuberculosis (TB), whooping cough, meningococcal meningitis, influenza, measles, rubella, chickenpox.

*Salmonella typhi* causes typhoid fever



Influenzavirus



### Gastrointestinal tract

Food and water are often contaminated with microorganisms, but most of these are destroyed in the stomach.

Examples: cholera, typhoid fever, mumps, hepatitis A, poliomyelitis, salmonellosis, giardia.

*Clostridium tetani* causes tetanus; found in soil.



### Breaking the skin surface

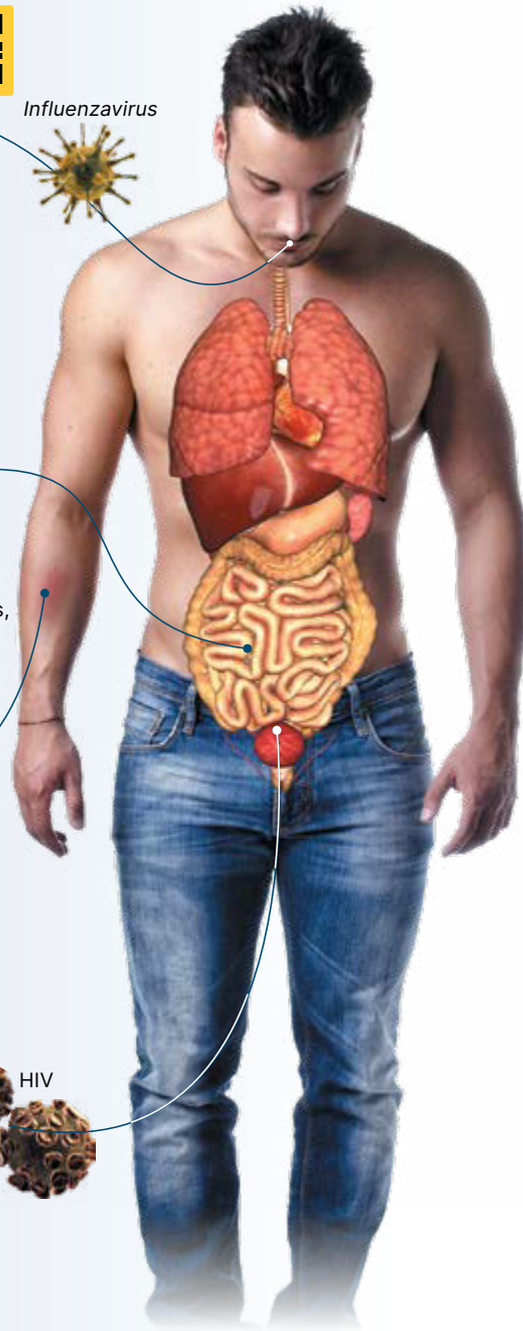
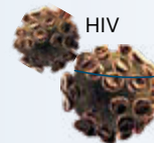
The skin provides an effective barrier to most pathogens, but cuts and abrasions allow pathogens to penetrate.

Examples: tetanus, gas gangrene, hepatitis B, rabies, malaria, and HIV.

### Urinogenital openings

Urinogenital openings provide entry points for the pathogens responsible for sexually transmitted infections (STIs) and other opportunistic infections (e.g. thrush).

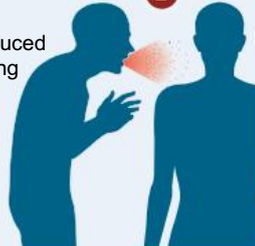
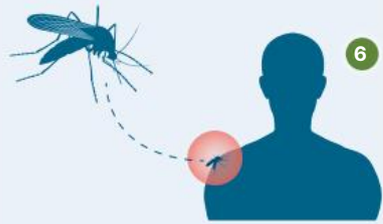
Examples: gonorrhoea, HIV.



1. Why can disease spread quickly in congested human cities? \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_
2. Why would transmission by direct touch be slower than transmission by coughing or sneezing?  
 \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_



## Modes of transmission

Direct contact	Indirect contact	Vector transmission
<p>Direct person to person contact can occur when an infectious person touches or exchanges fluids with another.</p>  <p>1 Person to person transmission (e.g. touching, kissing).</p> <p>2 Contact with droplets produced when coughing or sneezing.</p> 	<p>Indirect transmission occurs when an infected person infects another without having direct contact with that person.</p>  <p>3 Faecal-oral transmission, e.g. someone not washing hands properly after using the toilet.</p> <p>4 In some cases the pathogen can become airborne. Transmission can occur by breathing or coughing.</p>  <p>5 Transmission by contact with a contaminated object, including food and water, touched by an infected person.</p>	<p>Vectors carry a pathogen from one person to another, or from an animal to a human. Different vectors carry different pathogens.</p>  <p>6 Vector-borne pathogens are commonly carried by insects but sometimes other animals (e.g. ticks). They not only infect a person but can transfer the pathogen from person to person.</p> <p>A well know example is the Zika virus transmitted by the <i>Aedes</i> mosquito.</p>

### Examples of transmission

 <p>1 2 5 Ebola</p>	 <p>1 2 5 Chickenpox</p>	 <p>2 Tuberculosis</p>	 <p>6 Lyme disease</p>	 <p>1 3 5 Norovirus</p>
 <p>1 6 Zika</p>	 <p>1 2 4 5 Influenza</p>	 <p>1 3 E. Coli</p>	 <p>1 HIV</p>	 <p>1 5 Streptococcus</p>

### Transmission of SARS-CoV-2

- ▶ The SARS-CoV-2 virus, which causes COVID-19, can be spread in the tiny liquid particles given off when an infected person coughs, sneezes, speaks, or breathes.
- ▶ Transmission can be by direct or indirect contact. Droplets may land on surfaces that are then touched by others. The virus needs to be breathed in or make contact with the mouth or nose. A person may touch a contaminated surface with their hands then touch their nose or use their hands to eat and so become infected by the virus.
- ▶ The modes of transmission of SARS-CoV-2 make wearing masks and washing hands before eating important factors in slowing the spread of the virus.



Coughing or sneezing into your hands is an ideal way of spreading viruses.

3. Why is coughing into your elbow more effective at stopping the spread of a virus than coughing into your hands?

---



---



---

4. How are vector-borne pathogens transmitted?

---

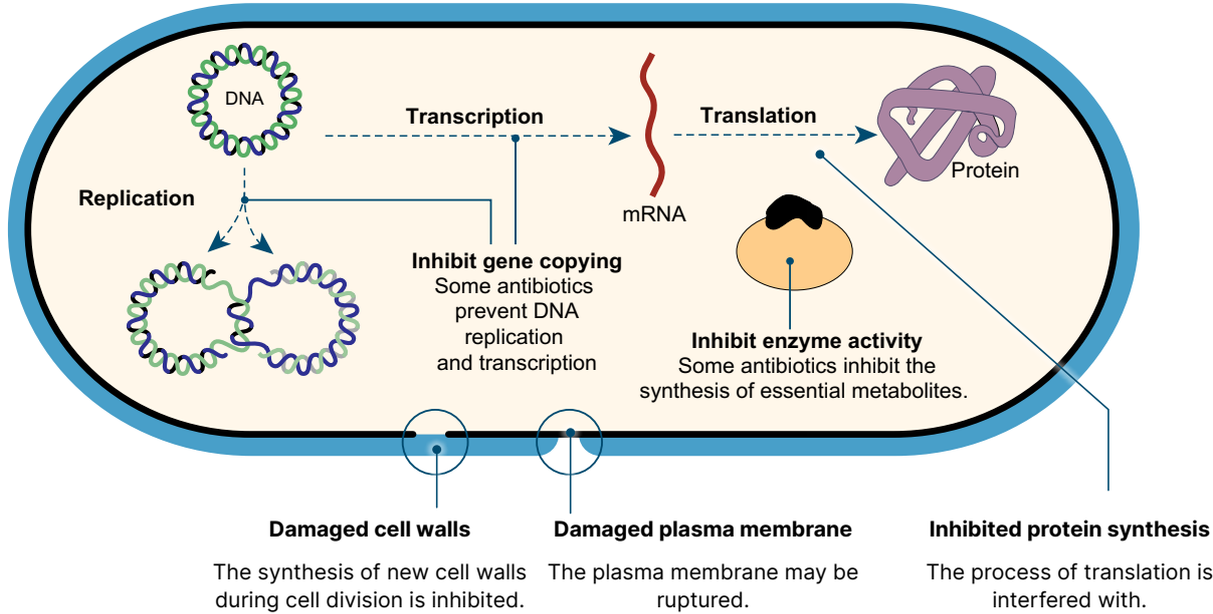


---

**Key Idea:** Antibiotics are antimicrobial chemicals that kill bacteria (bactericidal) or inhibit their growth (bacteriostatic). **Antibiotics** are chemicals that act against bacterial infections by either killing the bacteria (bactericidal action) or preventing them from growing (bacteriostatic action). Antibiotics interfere with bacterial growth by disrupting key

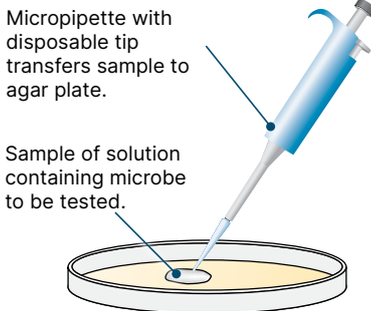
aspects of bacterial metabolism (below). Antibiotics are ineffective against viruses because viruses lack the structure and metabolic machinery that antibiotics target. Antibiotics are produced naturally by bacteria and fungi to kill or inhibit competitors or **pathogens**, but most modern antibiotics are semi-synthetic modifications of these natural compounds.

## How antibiotics work

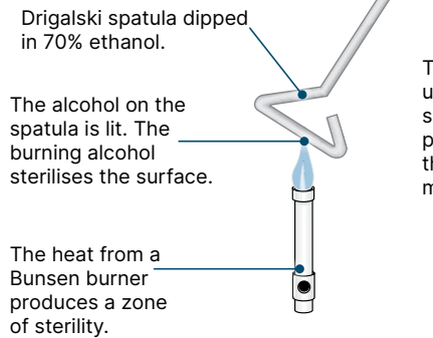


## Testing effectiveness of antibiotics

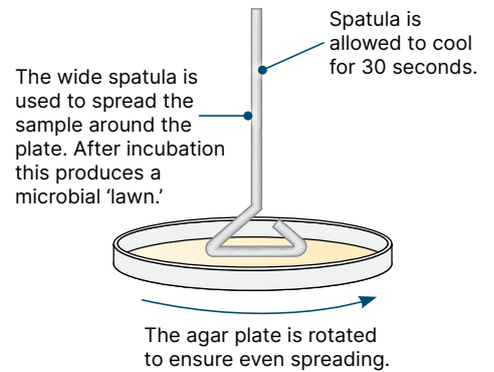
### 1 Pipette a sample



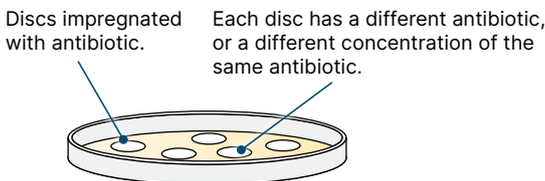
### 2 Sterilise the spatula



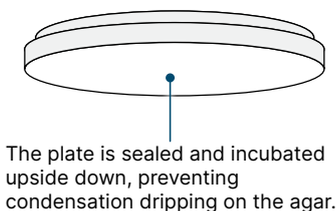
### 3 Spread the sample



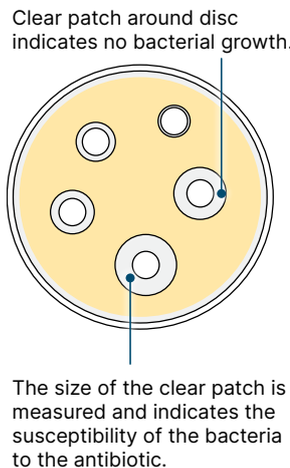
### 4 Antibiotic discs



### 5 Incubate plate



### 6 Analyse results

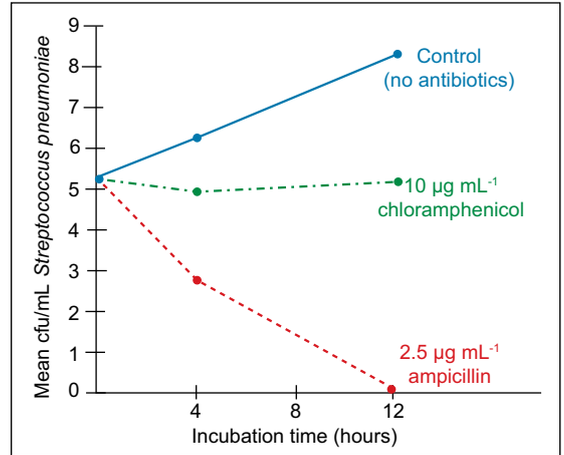


1. Why are viruses not affected by antibiotics? \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_
2. Distinguish between bacteriostatic and bactericidal: \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_

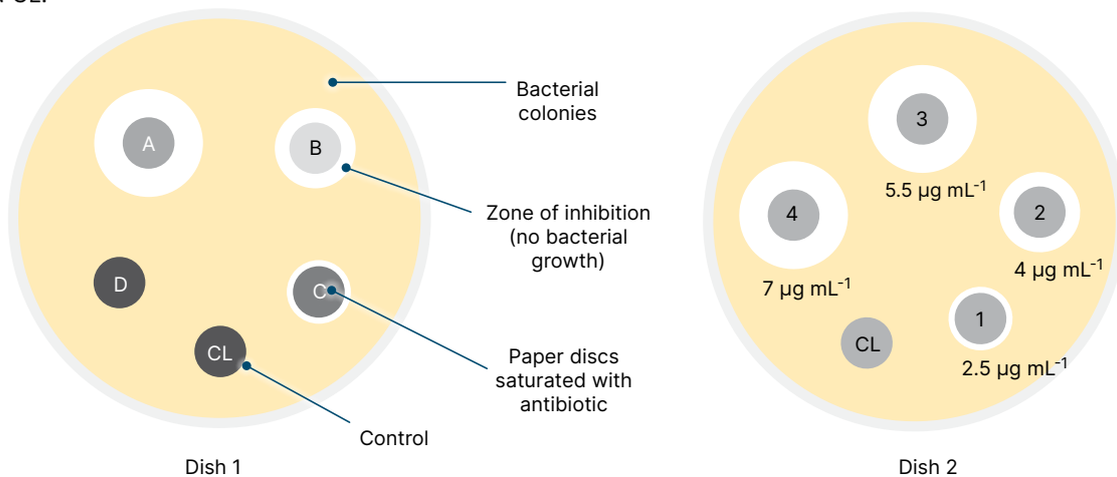
3. The graph (right) shows the effects of two antibiotics. Identify the antibiotic with a bacteriostatic action and the antibiotic with a bactericidal action. Explain your choice:

Bacteriostatic: \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_

Bactericidal: \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_



4. Two students carried out an experiment to determine the effect of antibiotics on bacteria. They placed discs saturated with antibiotic on petri dishes evenly coated with bacterial colonies. Dish 1 contained four different antibiotics labelled A to D and a control labelled CL. Dish 2 contained four different concentrations of a single antibiotic and a control labelled CL.



- (a) Which was the most effective antibiotic on Dish 1? \_\_\_\_\_
- (b) Which was the most effective concentration on Dish 2? \_\_\_\_\_
- (c) Explain your choice for question 4(b): \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_

5. Referring to the procedure for testing antibiotics on the opposite page:

- (a) Why is the agar plate incubated upside down? \_\_\_\_\_  
 \_\_\_\_\_
- (b) Why is the spatula dipped in alcohol and heated? \_\_\_\_\_
- (c) How would you measure the clear zone around the antibiotic discs? \_\_\_\_\_  
 \_\_\_\_\_

**Key Idea:** Studying the prevalence and spread of a disease gives insights into its origins and how to combat it.

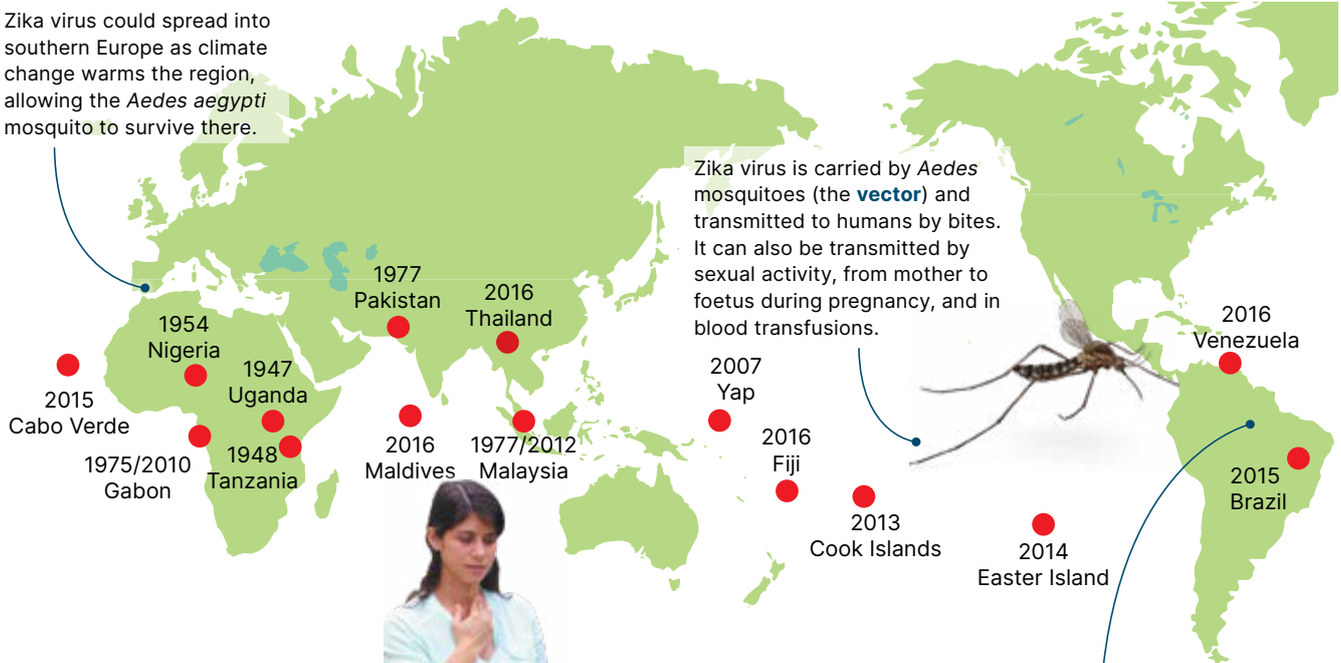
**Diseases** present at constant low levels in a population or region are known as endemic diseases. Occasionally there may be a sudden increase in the prevalence of a particular disease. On a local level this is known as an **outbreak**. When

an **infectious disease** spreads rapidly through a nation and affects large numbers of people it is called an **epidemic**. On rare occasions a new kind of disease will appear and spread to other countries. The rapid spread of a disease throughout the world is a **pandemic**. Examples of pandemic diseases include Covid-19, HIV/AIDS, influenza, and Zika virus.

### Zika virus: An example of global disease spread and its containment

Zika virus was first isolated from the area of the Zika Forest in Uganda in 1947. Since then it has spread slowly across the globe with outbreaks in the Americas in 2015 and 2016. Zika causes a mild fever and rash that is not usually serious in adults. However, in the last few years, infection of pregnant women by Zika has been linked to microcephaly (small head and brain) in their newborns.

Zika virus could spread into southern Europe as climate change warms the region, allowing the *Aedes aegypti* mosquito to survive there.



Zika virus is carried by *Aedes* mosquitoes (the **vector**) and transmitted to humans by bites. It can also be transmitted by sexual activity, from mother to foetus during pregnancy, and in blood transfusions.

The severe effects of Zika on foetal development prompted world health authorities to begin an awareness campaign to limit Zika's spread and reduce the risk of people contracting it. The campaign focussed on prevention, and included travel advisories in unaffected countries, as well as awareness campaigns in affected countries.



Insect repellent should be used, particularly if wearing clothing that exposes the skin.

Zika virus became an important international concern in 2015 and 2016 in the lead up to, and during, the 2016 Rio de Janeiro Olympics. Concerns focussed on the movement of spectators, tourists, and athletes, and the spread of the disease around the globe as people returned home after the events.



Reducing areas where water can stagnate reduces mosquito breeding sites.

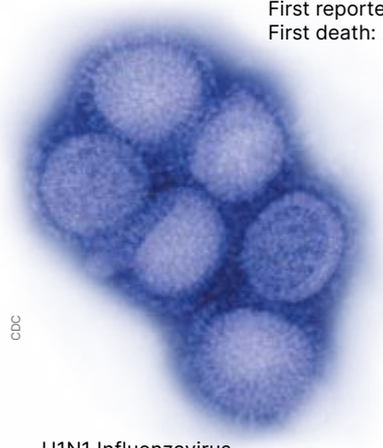
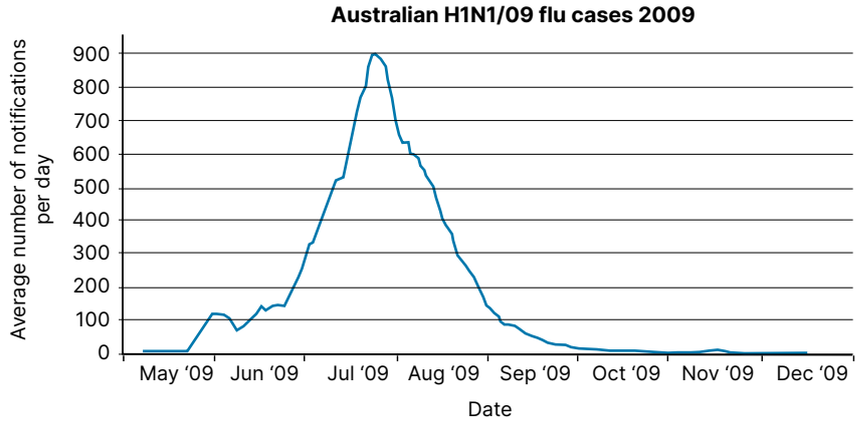
People are advised to wear long sleeves and pants to prevent mosquito bites.

- (a) Which general direction has Zika virus spread across the globe? \_\_\_\_\_  
 (b) Describe the area that Zika virus appears to be generally confined to and explain this:  
 \_\_\_\_\_  
 \_\_\_\_\_
- How is Zika virus transmitted? \_\_\_\_\_  
 \_\_\_\_\_
- Describe how the spread of Zika virus can be reduced: \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_

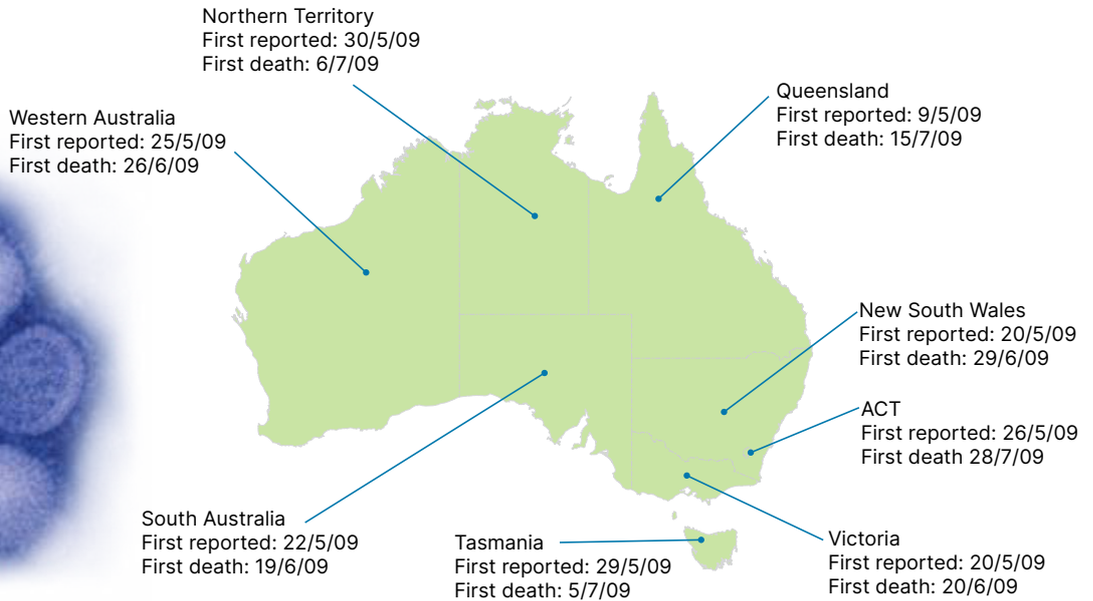
## H1N1/09: the 2009 pandemic

In 2009, the H1N1/09 subtype of the influenza virus (formerly known as swine flu), crossed the species barrier to humans (a zoonoses) in Mexico and from there spread to the rest of the world. Cases in Australia were first reported in May 2009 and quickly rose to a peak in late July. Total cases for 2009 were over 37,000. The 2009 H1N1/09 pandemic was the first influenza pandemic of the 21st century. The strain is now included in the annual flu **vaccination**.

Epidemiologists gather data on the number of infected people (morbidity) and the number of people that have died (mortality) within a population. These data help to establish the incidence (number of new cases per unit time) and prevalence (number of infected people expressed as a proportion of the population) of the disease in the population at any given time. Aetiology is the study of the cause of a disease. It can assist in pinpointing the origin of new diseases, such as the H1N1/09 influenza virus (swine flu).



Spread of H1N1/09 flu across Australia



4. In which Australian state was H1N1/09 first reported? \_\_\_\_\_
5. (a) Why was H1N1/09 classed as a pandemic? \_\_\_\_\_  
\_\_\_\_\_
- (b) What was the most likely origin of H1N1/09 spread in Australia? Explain your answer:  
\_\_\_\_\_  
\_\_\_\_\_
- (c) Suggest likely reasons for the rapid fall-off in the incidence of H1N1/09 in Australia after August 2009:  
\_\_\_\_\_  
\_\_\_\_\_
6. Why is it important to establish the incidence of a disease when it begins to spread through a community?  
\_\_\_\_\_  
\_\_\_\_\_  
\_\_\_\_\_

**Key Idea:** Hand washing effectiveness can affect microbial transmission to the body.

As humans, we spend much of our time manipulating objects with our hands, so it follows that our hands are covered with the microorganisms found in our environment. These microbes can then be easily transferred by touch to our

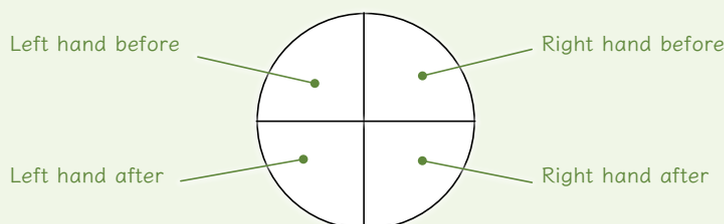
mouths, such as when eating, or to other people, such as when we hand them an object. Hand washing after contact with potentially contaminated material reduces the chance of transmitting microbes to our internal environment or to others. In the practical below you will obtain data on the effectiveness of handwashing.



### Investigation 15.1 Investigating the effectiveness of handwashing

See appendix for equipment list.

- The class will be divided into thirds. One third will wash their hands with warm water. One third will wash their hands with soap and warm water and one third will use hand sanitiser. Your teacher will place you into one of these groups. Do not wash your hands until step 5!
- Each person in the group should take a nutrient agar plate and use a marker pen to label the edge of the lid of the plate with name, the incubation temperature (e.g. 30°C), and which group you are in.
- Then use the marker pen to divide the plate lid into quarters and label them as shown below:



- Open the lid and press the tips of your middle and fore fingers from your left hand in the 'Left hand before' quarter. Hold them there for 5 seconds. Then press the tips of your middle and fore fingers from your right hand in the 'Right hand before' quarter. Hold them there for 5 seconds. Close the lid.
- Now wash your hands using the regime assigned to your group (water, soap and water, hand sanitiser). Dry your hands if needed with a clean paper towel.
- Open the lid of the agar plate again and press the tips of your middle and fore fingers from your left hand in the 'Left hand after' quarter. Hold them there for 5 seconds. Then press the tips of your middle and fore fingers from your right hand in the 'Right hand after' quarter. Hold them there for 5 seconds. Close the lid and seal it with clear tape.
- Incubate the plate at your chosen incubation temperature, lid down, for 24 hours.
- Retrieve the agar plates and observe the four different quarters. Count and record the number of bacterial colonies on the plate in each half (before and after). Do this for all the plates in your assigned group. If you only have a small number in your group, just enter the data you have. Calculate the mean number the colonies before and after (below).
- Compare your means with means from the other groups in the class.

1. (a)

Your technique: _____	Plate number										Mean
Number of colonies before washing hands											
Number of colonies after washing hands											

(b) Handwashing technique: \_\_\_\_\_ Mean colonies before: \_\_\_\_\_ Mean colonies after: \_\_\_\_\_

(c) Handwashing technique: \_\_\_\_\_ Mean colonies before: \_\_\_\_\_ Mean colonies after: \_\_\_\_\_

2. Which technique appears to have the greater ability to remove bacteria from your hands? Explain why:

---



---





# 198 Modelling Disease Outbreak and Spread

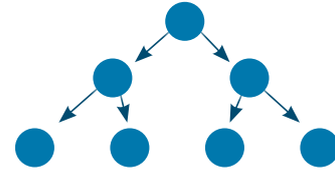
**Key Idea:** Being able to model the spread of a disease can help predict where, when, and how it will spread.

Modelling how a **disease** spreads can help preparation for an eventual **outbreak**. Elements of the model must account for

how infectious a **pathogen** is and for how long, the density and mobility of the population, and even the level of mortality of infected people. These models can be used to test the effectiveness of public health measures.

## Modelling a disease

- ▶ A spreadsheet can be used to model the spread of disease. There are also numerous online models that can be used.
- ▶ In the most simple model (right) whenever an infected person meets another, a new infection occurs. The number of interactions at each infection cycle affects the spread of the disease.
- ▶ Using a spreadsheet, you will first model an infected person meeting (and infecting) two other people. In this model, once the infected person has infected two people they are no longer infectious.



A simple infection model. One person infects two, who infect two more...



## Investigation 15.2 Modelling disease outbreak and spread

See appendix for equipment list.

1. Working in pairs, enter the following into a spreadsheet:

	A	B
1	New infections	Total infections
2	1	=SUM(\$A\$2:A2)
3	=A2*2	=SUM(\$A\$2:A3)
4	=A3*2	=SUM(\$A\$2:A4)
5	=A4*2	=SUM(\$A\$2:A5)
6	=A5*2	=SUM(\$A\$2:A6)
7	=A6*2	=SUM(\$A\$2:A7)
8	=A7*2	=SUM(\$A\$2:A8)
9	=A8*2	=SUM(\$A\$2:A9)

One infection cycle. Copy this down to row 12 (10 cycles of interactions).

1. How many new infections are there per infection cycle after 10 infection cycles? \_\_\_\_\_
2. How many infected people are there in total after 10 infection cycles? \_\_\_\_\_

2. Now set the interactions per infected person to 3 ( $A2*3$ ) and reset the model.

3. How many new infections are there per cycle of infection after 10 infection cycles? \_\_\_\_\_
4. How many infected people are there after 10 cycles of infection? \_\_\_\_\_

3. We can now extend the model by adding in a little randomness. The number of people interacting with each infected person may not always be the same. In our extended model, we shall randomise the number of people interacting to between 1 and 4.

	A	B	C
1	New infections	People interacted with per person	Total infected people
2	1	=RANDBETWEEN(1,4)	=SUM(\$A\$2:A2)
3	=A2*B2	=RANDBETWEEN(1,4)	=SUM(\$A\$2:A3)
4	=A3*B3	=RANDBETWEEN(1,4)	=SUM(\$A\$2:A4)
5	=A4*B4	=RANDBETWEEN(1,4)	=SUM(\$A\$2:A5)
6	=A5*B5	=RANDBETWEEN(1,4)	=SUM(\$A\$2:A6)
7	=A6*B6	=RANDBETWEEN(1,4)	=SUM(\$A\$2:A7)
8	=A7*B7	=RANDBETWEEN(1,4)	=SUM(\$A\$2:A8)
9	=A8*B8	=RANDBETWEEN(1,4)	=SUM(\$A\$2:A9)
10	=A9*B9	=RANDBETWEEN(1,4)	=SUM(\$A\$2:A10)

← Add new infections to total from previous row (cycle)

← Generates a random number between 1 and 4

← Calculates the total number of people infected

5. Run the model five times by recalculating the spreadsheet using the **recalculate** or **calculate now** option (depending on your spreadsheet). On average, how many people in total have been infected after ten cycles? \_\_\_\_\_



- Not all interactions will result in an infection. The pathogen may not be highly infectious or the correct mode of transmission may not have occurred (for example, a person with a cold may have been careful where and how they coughed).
- First we need to decide the probability of each interacting person being infected. For this model we will say there is a 50% chance that any interacting person will be infected. We shall first produce a random number between 0 and 1 (see \* below). We can now use this block of infected (1) or not infected (0) cells in our model. Once the formula is set up, you can recalculate the spreadsheet to obtain different infection scenarios.

A	B	C	D
1 New infections	People interacted with per person	Infected people	Total infected people
2 1	=RANDBETWEEN(1,4)	=IF(B2=1,\$D\$15,IF(B2=2,\$D\$15+\$D\$16,IF(B2=3,\$D\$15+\$D\$16+\$D\$17,IF(B2=4,\$D\$15+\$D\$16+\$D\$17+\$D\$18)))	=SUM(\$A\$2:A2)
3 =A2*C2	=RANDBETWEEN(1,4)	=IF(B3=1,\$D\$15,IF(B3=2,\$D\$15+\$D\$16,IF(B3=3,\$D\$15+\$D\$16+\$D\$17,IF(B3=4,\$D\$15+\$D\$16+\$D\$17+\$D\$18)))	=SUM(\$A\$2:A3)
4 =A3*C3	=RANDBETWEEN(1,4)	=IF(B4=1,\$D\$15,IF(B4=2,\$D\$15+\$D\$16,IF(B4=3,\$D\$15+\$D\$16+\$D\$17,IF(B4=4,\$D\$15+\$D\$16+\$D\$17+\$D\$18)))	=SUM(\$A\$2:A4)
5 =A4*C4	=RANDBETWEEN(1,4)	=IF(B5=1,\$D\$15,IF(B5=2,\$D\$15+\$D\$16,IF(B5=3,\$D\$15+\$D\$16+\$D\$17,IF(B5=4,\$D\$15+\$D\$16+\$D\$17+\$D\$18)))	=SUM(\$A\$2:A5)
6 =A5*C5	=RANDBETWEEN(1,4)	=IF(B6=1,\$D\$15,IF(B6=2,\$D\$15+\$D\$16,IF(B6=3,\$D\$15+\$D\$16+\$D\$17,IF(B6=4,\$D\$15+\$D\$16+\$D\$17+\$D\$18)))	=SUM(\$A\$2:A6)
7 =A6*C6	=RANDBETWEEN(1,4)	=IF(B7=1,\$D\$15,IF(B7=2,\$D\$15+\$D\$16,IF(B7=3,\$D\$15+\$D\$16+\$D\$17,IF(B7=4,\$D\$15+\$D\$16+\$D\$17+\$D\$18)))	=SUM(\$A\$2:A7)
8 =A7*C7	=RANDBETWEEN(1,4)	=IF(B8=1,\$D\$15,IF(B8=2,\$D\$15+\$D\$16,IF(B8=3,\$D\$15+\$D\$16+\$D\$17,IF(B8=4,\$D\$15+\$D\$16+\$D\$17+\$D\$18)))	=SUM(\$A\$2:A8)
9 =A8*C8	=RANDBETWEEN(1,4)	=IF(B9=1,\$D\$15,IF(B9=2,\$D\$15+\$D\$16,IF(B9=3,\$D\$15+\$D\$16+\$D\$17,IF(B9=4,\$D\$15+\$D\$16+\$D\$17+\$D\$18)))	=SUM(\$A\$2:A9)
10 =A9*C9	=RANDBETWEEN(1,4)	=IF(B10=1,\$D\$15,IF(B10=2,\$D\$15+\$D\$16,IF(B10=3,\$D\$15+\$D\$16+\$D\$17,IF(B10=4,\$D\$15+\$D\$16+\$D\$17+\$D\$18)))	=SUM(\$A\$2:A10)
11 =A10*C10	=RANDBETWEEN(1,4)	=IF(B11=1,\$D\$15,IF(B11=2,\$D\$15+\$D\$16,IF(B11=3,\$D\$15+\$D\$16+\$D\$17,IF(B11=4,\$D\$15+\$D\$16+\$D\$17+\$D\$18)))	=SUM(\$A\$2:A11)
12 =A11*C11	=RANDBETWEEN(1,4)	=IF(B12=1,\$D\$15,IF(B12=2,\$D\$15+\$D\$16,IF(B12=3,\$D\$15+\$D\$16+\$D\$17,IF(B12=4,\$D\$15+\$D\$16+\$D\$17+\$D\$18)))	=SUM(\$A\$2:A12)
13			
14			
15	=RAND()		
16	=IF(A15>0.5,1,0)		
17	=RAND()		
18	=IF(A17>0.5,1,0)		
19			
20			
21			

\* Produces a 50% probability of infection

The "IF" statement incorporates the number of interactions and the probability of infection into the model

- Run the model five times by recalculating the spreadsheet as before. On average, how many people in total have been infected after ten cycles now?
- The third model above is much more realistic than the first, but still lacks many factors that would affect the model outcome. List at least three factors that could be added to the model to make it even more realistic:

**Modelling with S, I, and R**

- A more advanced predictive mathematical model than your spreadsheet called SIR can be used to show the transmission of infectious diseases. In this model there are three compartments: **S** (the number of susceptible individuals), **I** (the number of infected individuals), and **R** the number removed (those who have been removed through recovery or death).
- The data in the table (below right) is a theoretical example. It assumes a closed system (e.g. a single state with no travel), no prior immunity (everyone is susceptible), no vaccine, and no physical distancing or other precautionary measures in place.

Need help?  
See Activity 12

Week	S	I	R
0	7,000,000	2	0
1	6,999,986	15	1
2	6,999,881	113	8
3	6,999,090	847	65
4	6,993,162	6352	488
5	6,948,741	47,597	3664
6	6,618,002	354,538	27,462
7	4,271,669	2,523,602	204,731
8	0	5,533,470	1,466,532
9	0	2,766,735	4,233,267
10	0	1,383,368	5,616,634
11	0	691,684	6,308,318
12	0	345,842	6,654,160
13	0	172,921	6,827,081
14	0	86,460	6,913,542

- Plot the tabulated SIR data left on the grid provided. Plot all three data sets on one axis with a key

9. Describe the relationship between the three compartments (S,I,and R) over time:

---



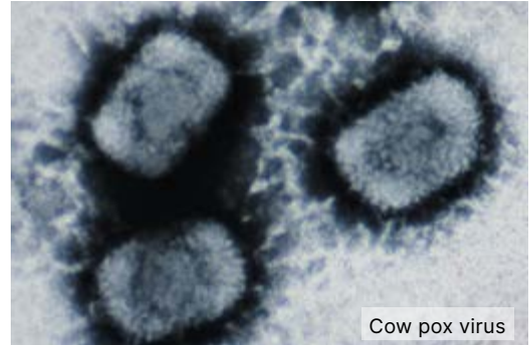
---



---

**Mathematical models and vaccination**

Mathematical models of the effect of **vaccination** on populations have been used since the mid-1700s. In 1760, Swiss mathematician Daniel Bernoulli published a study on the effect of **immunisation** with cowpox (against smallpox) on the life expectancy of the immunised population. Around the time of the First World War, Ronald Ross produced mathematical models to show that malaria could be controlled without removing every last mosquito. These mathematical models are the basis for many vaccination programmes and show why herd immunity is an important aim of public health programmes.



- ▶ All that is needed for the incidence of disease to decline is that every case or primary infection should generate less than one other case or secondary infection (on average).
- ▶ The number of secondary infections caused by an infectious individual is denoted as R. **R<sub>0</sub>** (R nought) is the basic reproductive number of the pathogen. It is the number of secondary infections caused by a primary infection introduced into a wholly susceptible population.
- ▶ **R<sub>0p</sub>** is the basic reproductive number under vaccination. It is the number of secondary infections caused by a primary infection introduced into a population where a proportion (*p*) of the population is vaccinated. For a perfect vaccination that confers life long immunity:

$$R_{0p} = (1 - p) R_0$$

- ▶ *p<sub>c</sub>* is the critical vaccination proportion that will achieve eradication. To achieve this proportion, the basic reproductive number under vaccination (**R<sub>0p</sub>**) must be just less than 1, so:

$$p_c = 1 - \frac{1}{R_0}$$

Calculating *p<sub>c</sub>* requires estimates of R<sub>0</sub> (right)

**Estimates of R<sub>0</sub> for populations and dates**

Infection	Location	Date	R0
Measles	Senegal	1964	18
Smallpox	West Africa	1960	2.3
Mumps	UK	1987	8
Rubella	USA	1967	6
Influenza	UK	2010	1.5
Covid-19	Global	2019	~2.5

10. How are mathematical and computer models useful in controlling disease?

---



---

11. Calculate the critical vaccination proportion for each of the following diseases:

- (a) Measles: \_\_\_\_\_
- (b) Influenza: \_\_\_\_\_
- (c) Covid-19: \_\_\_\_\_

12. Why is the *p<sub>c</sub>* of measles so high? \_\_\_\_\_

---



---

13. Calculating the R<sub>0</sub> of Covid-19 was a critical step in controlling the spread of the virus. What does the R<sub>0</sub> of Covid-19 tell us? Could the disease have been controlled without a vaccine, or is vaccination the only way to stop its spread?

---



---



---

# 199 Predicting Future Patterns of Disease

**Key Idea:** Predicting future disease outbreaks relies on monitoring current disease episodes and using population statistics to identify potentially vulnerable groups of people.

**Disease outbreaks** have occurred throughout history. The plague that spread through Europe in the 1400s (the Black Death) and again in the mid 1600s (the Great Plague) were

some of the most devastating outbreaks ever. The Spanish flu (1918) was the last great worldwide **pandemic** prior to the 2020 Covid-19 pandemic. Improved global cooperation and more effective action by health authorities have improved **infectious disease** containment and significantly reduced the potential death rate resulting from Covid-19.



Predicting future disease is often a case of identifying diseases in animals that could cross over to human **hosts**. Many infectious diseases have an animal origin including various strains of influenza (e.g. avian flu H5N1 and swine flu H1N1). Identifying these **pathogens** in animals, especially livestock and poultry living in close proximity to humans, can help prepare for possible disease **outbreaks**.



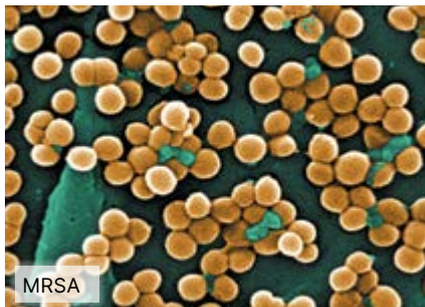
Population density is important to how quickly an infectious disease can spread. Cities generally have very high population densities, with some reaching densities of over 20,000 per km<sup>2</sup>. Disease spreads most quickly in areas with poor living conditions, poor sanitation, and low levels of immunity. For example, the Spanish flu initially spread quickly due to the cramped confines of military camps and hospital wards.



How quickly an infectious disease spreads also depends of the population's mobility. The 1918 Spanish flu spread around the world due to infected troops returning home and taking the disease with them. Part of predicting where and when diseases will occur is being able to predict the movements of groups of people. For example people moving from rural areas to cities may transport potential pathogens from livestock.



The spread of disease also depends on the mode of **transmission**. Is it spread by air borne particles or by touch? The most feared scenario is an airborne pathogen that is highly contagious, has a long infectious period, and is ultimately deadly.



Resistance to **antibiotics** is becoming a greater problem with many bacterial strains becoming extremely difficult to treat. Plans need to be in place for if (or when?) a highly resistant pathogenic bacteria begins to spread through the general population.



Climate change is already being taken into account to predict where possible outbreaks may occur. For example, malaria made spread further north and south from the tropics as the climate becomes more favourable, due to climate change, for its mosquito **vectors**.

1. Explain how each of the following are important in predicting where the next epidemic may originate:

(a) Pathogen in livestock related to human diseases: \_\_\_\_\_

\_\_\_\_\_

(b) Population density: \_\_\_\_\_

\_\_\_\_\_

(c) Global travel networks: \_\_\_\_\_

\_\_\_\_\_

(d) Resistance to antibiotics: \_\_\_\_\_

\_\_\_\_\_

(e) Climate change: \_\_\_\_\_

\_\_\_\_\_



## Seasonal patterns

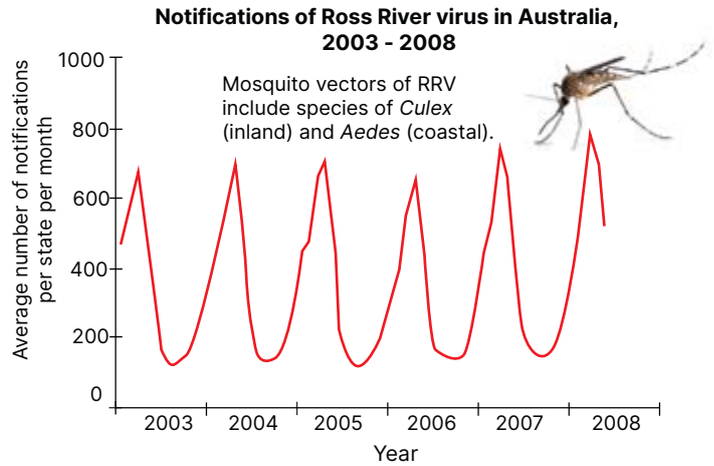
Some diseases are seasonal, showing patterns of increased or decreased prevalence at specific times of the year. Influenza (the flu) commonly becomes more prevalent in the winter of both the Southern and Northern Hemispheres. Ross River fever becomes more common during the summer/autumn rainy season (January to March). The seasonal patterns of these diseases allow for simple predictions of when most cases will occur. For example, health authorities prepare for increased influenza cases in winter by offering the latest **vaccines**.

### Ross River virus

Ross River virus (RRV) is endemic to Australia, Papua New Guinea, and several other islands of the South Pacific and causes Ross River fever. RRV is transmitted by mosquitoes and may have natural reservoirs in native Australian mammals. In Australia, most RRV infections occur in the wetter tropical areas of QLD, WA, and NT. There have been several large outbreaks since the first noted outbreak in 1928. The largest outbreak occurred in 1979-80 across the Western Pacific, affecting 60,000 people.

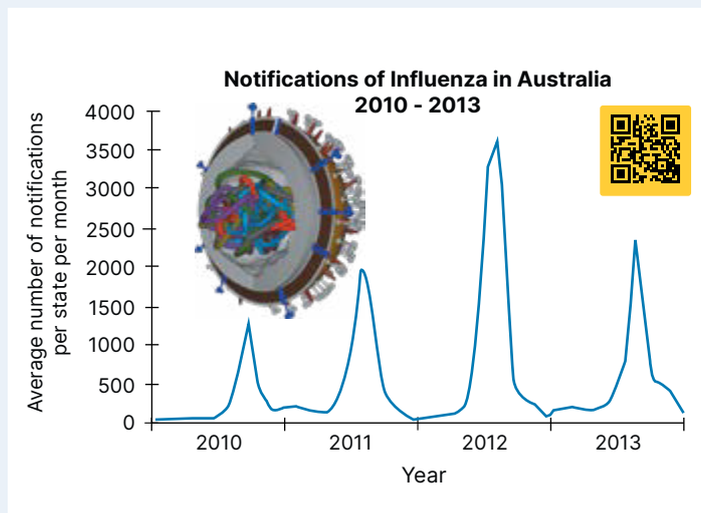
### Ross River fever

Symptoms of Ross River fever include fever, rash, and arthritic-like symptoms causing extreme pain in various joints. In most cases, the disease lasts about a month but can reoccur over a period of years. Currently a blood test is the only way to confirm Ross River fever and there is no vaccine.



### Influenza

Although it is often looked upon as simply a bad cold, influenza (commonly called 'the flu') has arguably caused more human deaths than any other disease throughout history. Influenza occurs in seasonal **epidemics**, usually infecting around five million people globally, and causing the death of between 250,000 and 500,000 people annually. The influenza virus continually changes in two ways. Antigenic drift involves small cumulative genetic changes over time. Occasionally the virus will also undergo antigenic shift, producing an entirely new strain that causes a pandemic. The latest of these is the H1N1/09 strain, commonly referred to as swine flu. This strain caused around 14,000 deaths globally. Compare this the Spanish flu pandemic of 1918-1920, during which possibly up to 100 million people died globally (more than all deaths during World War I and possibly as many as World War II).



2. (a) How is the Ross River virus transmitted to people? \_\_\_\_\_

(b) Suggest why cases of Ross River fever are more prevalent in January to March:

\_\_\_\_\_

\_\_\_\_\_

(c) There was a large mosquito outbreak in 2013 in Newcastle, NSW. Predict the effect of this on the prevalence of Ross River fever:

\_\_\_\_\_

3. (a) What time of the year is the influenza virus most prevalent? \_\_\_\_\_

(b) Why must people receive the flu vaccine every year if they want to remain protected against influenza?

\_\_\_\_\_

\_\_\_\_\_

4. Why is it useful to track the incidence of diseases such as Ross River fever and influenza?

\_\_\_\_\_

\_\_\_\_\_

# 200 Containing the Spread of Disease

**Key Idea:** Preventing the entry and spread of pathogens is important in protecting a country's population and industries from infectious diseases.

Many factors can influence the **spread of disease**, including the social climate, diet, general health, and access to medical care. Human intervention and modification of behaviour, including **vaccination**, can reduce the **transmission** rate of

some diseases and inhibit their spread. Global air travel and international trade in commodities has increased the risk that diseases of humans, livestock, and crops will be spread between countries. Persistence of pathogens: the ability of them to remain within a host organism for an extended period, often evading the immune system, can also affect how disease spreads.



**Quarantine**

Transmission of disease can be reduced by adopting 'safe' behaviours, such as isolation of people already infected, or establishing **quarantine** procedures for people who have been exposed to infection. The quarantine lasts until hosts are no longer infectious.



**Contact tracing**

If virus infected patients are able to recall the people and places they were in contact with in the period during which they were contagious, contact tracers can test and/or quarantine those people. This can potentially stop further spread.



**Disinfecting**

Disinfectants and sterilisation techniques, e.g. autoclaving (above), destroy pathogenic microbes before they have the opportunity to infect. The use of these techniques in medicine has significantly reduced post operative infections and associated deaths.



**Lockdown and social distancing**

During the Covid-19 **pandemic** in early 2020, many countries implemented lockdown procedures where businesses and schools were closed, and small groups remained isolated in 'bubbles' at home. Public gatherings were very limited and people outside their bubbles needed to adhere to social distancing to avoid spread.



**Hygiene**

Appropriate personal hygiene practices reduce the risk of infection and transmission. Soap may not destroy the pathogens but washing will dilute and remove them from the skin. Masks can also prevent the spread of airborne viruses and were mandatory in many businesses and work places during the Covid-19 pandemic.



**Vaccination**

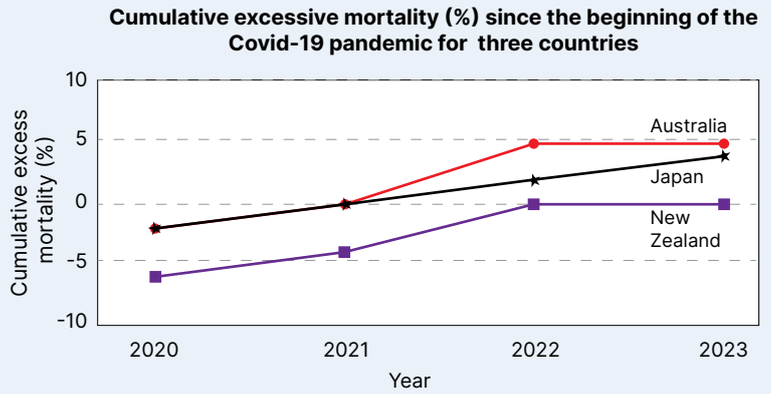
Vaccination schedules form part of public health programmes. Vaccination and **vaccine** development is one of the most effective ways of preventing transmission of contagious diseases. If most of the population is **immunised**, herd immunity limits outbreaks to sporadic cases and prevents **epidemics**.

1. (a) What is the link between persistence of pathogen and length of quarantine required?  
 \_\_\_\_\_  
 \_\_\_\_\_
- (b) What role does contact tracing play in controlling the spread of infectious diseases?  
 \_\_\_\_\_
- (c) What is herd immunity and how does vaccination contribute to it?  
 \_\_\_\_\_  
 \_\_\_\_\_
2. Discuss the importance of reducing the prevalence of disease compared to trying to contain an outbreak. Provide examples from the Covid-19 pandemic to support your argument.  
 \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_



### Covid 19 and excessive mortality rates

- ▶ The first confirmed case of COVID-19 in Australia was reported on January 25, 2020. By March 12, there were 140 confirmed cases in the country. A series of public health and social measures were put in place to slow the spread of the virus.
- ▶ A number of health measures including vaccinations and anti-viral medicines have been used to reduce serious illness or death in Covid-19 infected patients. However, data shows mortality (death) rates are above what would typically be expected within the Australian population.
- ▶ The graph (right) shows the percentage difference between the cumulative number of deaths since 1 January 2020 and the cumulative expected number of deaths based on previous years (2015-2019).

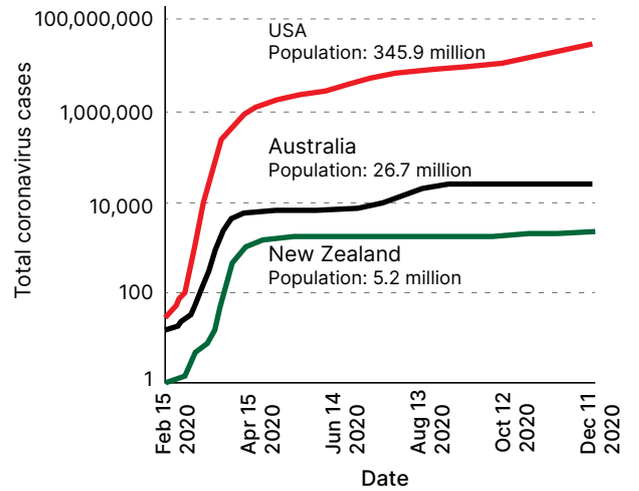


Data source: Australian Institute of Health and Welfare  
<https://www.aihw.gov.au/reports/australias-health/covid-19>

### Covid 19: Different countries, different outcomes

- ▶ During the early stages of the pandemic, some countries were very successful in slowing or containing the spread of the virus. Graphically, this was shown by a flattening of the infection curve.
- ▶ In some countries the virus spread widely and quickly. This is illustrated by a steep, exponential increase on a graph.
- ▶ The way governments, health departments, and populations responded to the disease was important in the pattern of Covid-19 spread.
- ▶ The graph (right) shows the number of confirmed cases for three countries: Australia, New Zealand, and the US in the first year of the pandemic.
- ▶ Although these countries have very different populations and recorded their first Covid-19 infection at different times, the data can be used to see how well each country contained the spread of disease.

Comparison of Covid-19 cases by country (Feb 15 2020 - Dec 11 2020)



Data source: Worldometer <https://www.worldometers.info/coronavirus/>

3. Study the graph showing cumulative excess mortality rates during the Covid-19 pandemic.

(a) What trend does the data show? \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_

(b) Compare Australia's excessive mortality rates against those of Japan and New Zealand:  
 \_\_\_\_\_  
 \_\_\_\_\_

(c) How could the data be used to study the effect of Covid-19 on death rates? \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_

4. Study the graph showing the number of Covid-19 cases in the USA, Australia, and New Zealand.

(a) All three countries showed exponential increase in Covid-19 cases during one period. When was this?  
 \_\_\_\_\_

(b) Which country was most successful at controlling Covid-19 spread? Explain your answer: \_\_\_\_\_  
 \_\_\_\_\_

**Key Idea:** Biosecurity measures are used to protect the valuable agricultural industry and biodiversity of Australia. **Biosecurity** is important for Australia which, because of its relative isolation, has managed to exclude many of the **pathogens** that infect plants and animals in other countries. Precautions such as **quarantine**, which isolates exposed

individuals that may be infected, as well as screening of imported produce and international travellers, help to limit the entry of **diseases** into Australia. Quarantine is distinct from isolation, which aims to contain disease by isolating an already infected person.

### Biosecurity in Australia

Australia has strict biosecurity rules and measures in place to prevent the entry of pests and diseases into the country. For most people, this is most visible in airports when entering Australia from overseas. Passengers disembarking from aircraft are repeatedly reminded of what can and can't be brought into the country. Large fines can be instantly given to people who ignore the warnings. Inspection officers commonly use X-ray machines and detector dogs to check passengers' luggage for prohibited goods, especially fresh food or animal and plant materials.

Biosecurity inspections are also made on goods entering on cargo ships. Shipping containers are inspected for unwanted plants or animals that may have got into the container when loading. This is common in fresh food containers (e.g. fruits). If pests are found, the containers may be turned away. Inspection of many goods occurs at the home port before they are loaded on to cargo ships.

### Processes used to protect Australian agriculture and biodiversity

**Surveillance and Monitoring:** Essential activities conducted at high-risk locations, complementing border control efforts that concentrate on mitigating potential biosecurity risks at airports, seaports, and mail centers, with detector dogs playing a valuable role in these operations.

**Examination:** Opening and testing for specific materials, such as drugs, can ascertain if they are banned items.

**Clearance activities:** Inspection of cargo, including careful observation of shipping containers can identify harmful items.

**Risk analysis:** Make decisions on importing depending on likely risk of items.



Björn Christian Tarrissen CC3.0

### The role of quarantine

When organisms are brought into Australia they must undergo a quarantine period to monitor health and ensure no pests or diseases are in or on the organism. Quarantine may also apply to travellers who have been in contact with infected persons or have returned from places known to have disease **outbreaks**. These quarantine procedures were used during the SARS epidemic in 2003, the swine flu **pandemic** in 2009, and the Covid-19 pandemic in 2020.

The equine industry is an important part of the Australian economy. Live horses brought into the country are quarantined to check for diseases that may affect the industry. In August 2007, equine influenza was discovered in horses at the Eastern Creek Quarantine Station, the first time it had entered Australia. Somehow, it escaped quarantine, possibly through human error, and spread throughout NSW, reaching as far as Gatton and Warwick in Queensland. Due to strict non-movement orders, the outbreak was contained by February 2008.



Equine influenza is highly infectious and so easily spread.

1. Why is biosecurity important for Australia? \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_
2. Why is pre-inspecting goods at the home port a useful biosecurity measure?  
 \_\_\_\_\_  
 \_\_\_\_\_
3. How does quarantine prevent the spread of disease? \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_
4. How does Australia's geographic position help prevent the entry and spread of disease in Australia?  
 \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_

# 202 The Effectiveness of Health Programs

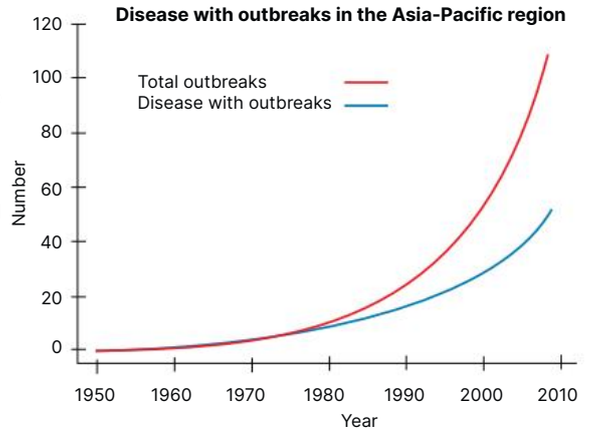
**Key Idea:** Public health programmes have successfully been used in disease pandemics.

Public health **vaccination** programs are successful due to global cooperation, investment, and the development of effective **vaccines** that provide immunity and inhibit **disease**

**transmission.** Mandatory vaccination requirements for travel also ensure widespread coverage and protection against diseases, contributing to the success of these programs. Successful global vaccination programs have eradicated or almost eliminated some once common **infectious diseases.**

## Vulnerability of disease health programmes in Asia-Pacific

- ▶ The Asian region encompasses about 30% of the Earth's land area and 60% of the human population. With a vast range of countries, environments, beliefs, and political systems, producing a single actionable plan in the case of a **pandemic** is a difficult task in Asia.
- ▶ Most of the world's largest and most densely populated cities are found in Asia making it particularly vulnerable to **epidemics.** The total number of diseases with **outbreaks** and the total number of outbreaks has increased rapidly in the Asia-Pacific region since the 1950s (right).
- ▶ The outbreak of SARS (Severe Acute Respiratory Syndrome) in 2003 showed that the public health systems and cooperation of nations throughout Asia needed strengthening. SARS appeared in China in November 2002, but Chinese officials did not inform the World Health Organisation (WHO) until February 2003. This lack of coordination resulted in SARS spreading much further and infecting many more people than it might have otherwise.
- ▶ Since then, several other epidemics have swept through Asia, including the avian flu H5N1, of which the first human-human transmission was recorded in 2006. Since then, millions of domestic poultry have been culled in order to control the disease. A new subtype H7N9 was reported in 2013 in China.



Flu vaccine stockpiles



Panic buying is common in uncertain times

The cost of vaccines and health care, especially in poorer countries, is a major barrier to preventing outbreaks. A key part of containing outbreaks is to ensure developing countries have access to vaccines and treatments. However, it is not reasonable to expect a nation with a low GDP to spend a large percentage of its income on medical equipment it might not need.

A lack of health infrastructure is another barrier to the prevention of outbreaks. For example, India spends just 3% of its GDP on health care whereas Australia spends more than 10% of its GDP of health care. This can mean that if a disease outbreak does occur in a vulnerable country, health care services may not be able to meet the demand for resources and treatments.

Managing information is an important part of any pandemic management plan. People can now get up-to-date information on a disease from the internet. It is important for health authorities to maintain communication with the public. This can help to avoid speculation and panic, which may result in people taking inappropriate and ineffective actions.

1. Why is the Asia region particularly vulnerable to epidemics and pandemics?

---



---



---

2. Why is maintaining communication with the public so important during a pandemic?

---



---



3. In small groups, discuss how countries in the Asia region can collectively prepare for a pandemic. Summarise your discussion as bullet points below.

---



---



---



## Smallpox: the virus that was eradicated

- ▶ Smallpox was a viral disease that affected human populations for around 3000 years. Evidence of its early existence came from examination of Egyptian mummies and ancient Chinese texts.
- ▶ The disease covered the host in disfiguring pustules. Fever, vomiting, and mouth sores accompanied the rash. More than 3 out of 10 infected people died and those who survived were often left with permanent scars, went blind, or became infertile.
- ▶ Smallpox spread across Europe, Asia, and Africa as a result of trade expansion in the Middle Ages. The age of exploration caused it to spread with devastating consequences to New World countries of the Americas and Oceania.
- ▶ Estimates suggest that smallpox killed between 300-500 million people worldwide, 90% of which were under the age of 10. Combined efforts on a worldwide scale allowed complete eradication of smallpox by 1977.

### Combating smallpox

- ▶ The first attempts at public health control of smallpox used a practice called variolation. Researchers have found evidence that this technique started as early as 200 BCE. It involved rubbing a small amount of smallpox pus from an infected person onto a healthy person. This person then became infected, but with a milder case. Once recovered, they had acquired immunity to the disease. It is important to note that although this practice used principles of immunisation, the practitioners had no scientific concept of viruses or immunity at the time.
- ▶ Observers noticed that women working closely with cows had a lower incidence of infection with smallpox. In 1796, the first vaccine, a dose of cowpox virus, was given to a sample of the population who then became immune to smallpox. By the 19th century, a related virus, *Vaccinia*, was substituted for cowpox virus in smallpox vaccines.
- ▶ Worldwide vaccination programmes began soon after, and smallpox vaccination certificates were required for travel to some countries.
- ▶ Smallpox was eliminated in Europe by 1900 but small pockets existed elsewhere in the world and had the potential to cause outbreaks worldwide.
- ▶ The World Health Organization began a smallpox eradication programme in 1959. Due to massive global cooperation and investment, smallpox was eventually eradicated. The last case was in 1977.



CDC/James Hicks

Child in Bangladesh covered with smallpox pustules in 1973.

### Polio: another vaccination program success

- ▶ Polio is a highly infectious disease caused by the polio virus. It results in death or paralysis and mainly affects children.
- ▶ In 1955, Jonas Salk, an American virologist, developed an inactivated (dead) polio vaccine (IPV). A global vaccination programme led by WHO reduced annual cases from 58,000 to under 200 in 6 years.
- ▶ Around the same time, Albert Sabin developed a live, attenuated vaccine (OPV). This could be taken orally, which made distribution and dispensing to remote areas of the world very easy. This vaccine inhibited transmission as well as offering immunity.
- ▶ Due to worldwide efforts, polio is now eradicated in most countries but low vaccination rates in central Africa and Indonesia means that people there are still at risk from the disease.



Oral polio (OPV) vaccine being administered as part of program.

Julien Hamels CC 2.0

4. How did the practice of variolation help control of smallpox? \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_
5. How was the first smallpox vaccine developed in 1796 and how did it help combat the disease?  
 \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_
6. What advantage does the live attenuated oral polio vaccine (OPV) have over the inactivated polio vaccine (IPV)?  
 \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_  
 \_\_\_\_\_

# 203 Aboriginal Protocols in Medicine

**Key Idea:** Traditional knowledge and property rights need to be acknowledged when developing traditional medicines for commercial use.

Indigenous Australian people have long used native plants for traditional medicines. Most modern medicinal drugs are manufactured industrially on a vast scale and the majority are based on chemicals made naturally in living organisms.

As more **pathogens** become resistant to current medicines, it is important that researchers explore the natural world for new chemicals that could potentially be used against these pathogens and possibly act as new treatments for cancer. However, the production and use of medicines derived from plants used in traditional medicine raises many ethical and legal questions over who actually owns the knowledge.

- ▶ Australian flora is rich in plants that produce chemically active compounds such as aromatics, alkaloids, tannins, and oils. In the 50-60 thousand years that Aboriginal Australians have inhabited Australia, they have identified and used a wide range of plants that have medicinal value. The traditional use of these plants is commonly called **bush medicine**.



Mark Marathou CC 3.0

Gumbi gumbi (*Pittosporum angustifolium*) is traditionally used to treat coughs, colds, and even eczema. Research shows it may have use as an antiviral, and blood pressure regulator.



John Jennings CC 2.0

Tea tree (*Melaleuca*) oil is well known for its **antiseptic** properties. Vapour from crushed leaves can be used to treat headaches. Tea brewed from it helps relieve coughs and colds.



Geoff Dierrin CC 4.0

The leaves of emu bush (*Eremophila longifolia*) have been used to treat wounds, headaches, and chest pains. Smoke from the leaves can create a sterile environment.

## Smoke bush

- ▶ The smoke bush plant (*Conospermum*) is traditionally used in Aboriginal bush medicine. In 1960, specimens were collected by the US Cancer Institute and tested. No useful properties for use against cancer were found and the samples were put in storage. In the 1980s, following the spread of HIV, the samples were tested for use against the virus. The chemical, conocurovone, was found and shown to destroy HIV, in low concentrations.
- ▶ A patent was filed in the US in 1993 and in Australia in 1994. The patents gave the US Government the exclusive rights to developed compounds from the smokebush plant and to license them to other companies.
- ▶ Via the West Australian Government, commercial rights were licensed to AMRAD, an Australian pharmaceuticals company. No acknowledgement of traditional Aboriginal knowledge was included. There was also a possibility that the traditional knowledge holders could be excluded from using any sort of plant product.
- ▶ This is an example of what has been termed 'biopiracy', in which a company applies for a patent to a traditional resource that can stop any traditional use of the resource while making profit of their own.



Casillier CC 3.0

1. In groups of 3-4 investigate traditional Aboriginal medicines and identify ways to protect traditional knowledge and property rights. Useful links can be found in **BIOZONE's Resource Hub**. Report back to your class on a traditional medicine (not mentioned above) you have identified, whether it has any possible commercial uses or potential for development, and how traditional knowledge and rights will be acknowledged or protected. Summarise the results of your investigation below:

---



---



---



---



---



---



---



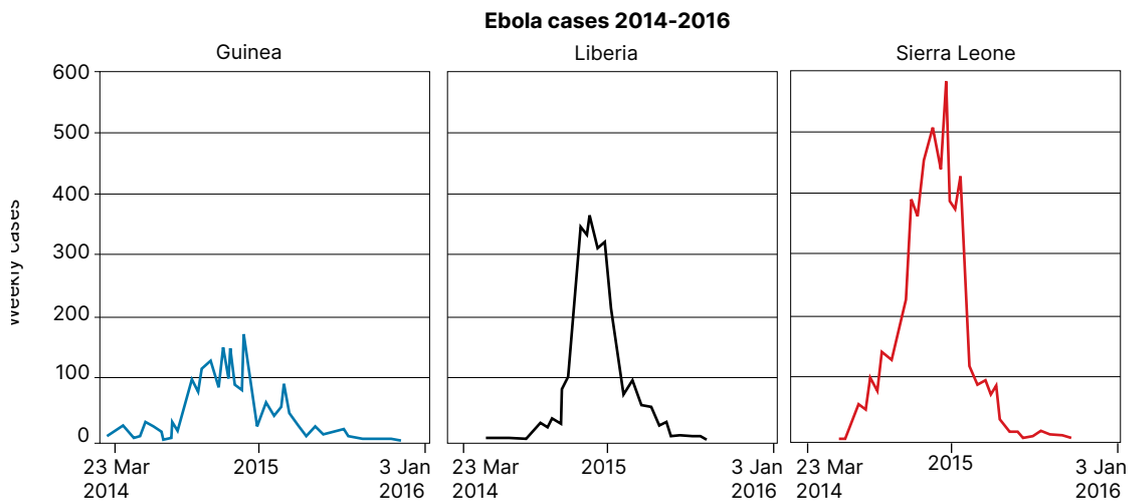
# 204 Did You Get It?

1. Match each term to its definition by writing the correct letter in the box:

- (i) antibiotic
- (ii) contact tracing
- (iii) vaccine
- (iv) quarantine
- (v) outbreak
- (vi) disease transmission

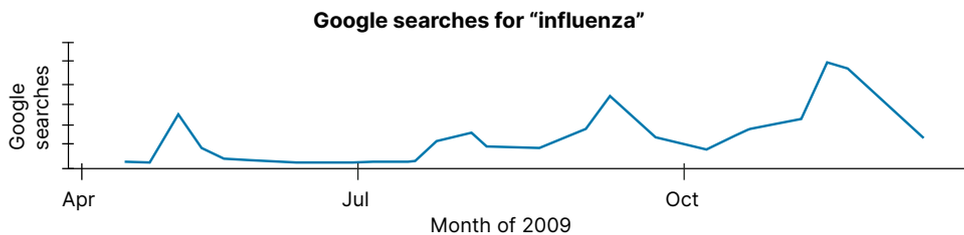
- A** A period or place of isolation in which an individual who may have been exposed to a disease is placed in order to prevent them from possibly spreading a disease.
- B** A biological preparation that provides active acquired immunity to a particular disease.
- C** A public health strategy used to identify and monitor individuals who may have come into contact with an infected person.
- D** The process by which an infectious agent, such as a virus or bacteria, is passed from one individual to another, leading to the spread of the disease.
- E** A type of medication that is used to treat bacterial infections by either killing the bacteria or inhibiting their growth.
- F** The sudden occurrence of a large number of disease cases in a particular place.

2. Study the graphs of the 2014–2015 West Africa Ebola outbreak below:



- (a) In which country did Ebola first appear? \_\_\_\_\_
- (b) Which country had the greatest number of cases of Ebola? \_\_\_\_\_
- (c) What was the highest number of new cases reported per week? \_\_\_\_\_
- (d) When and where did this occur? \_\_\_\_\_

3. Various health intelligence networks e.g. the Global Public Health Intelligence Network, monitor internet searches in order to determine if a disease outbreak is imminent. For example, the graph below shows the number of Google searches including the word 'influenza' for 2009:



How would monitoring the number of internet searches about diseases or symptoms help identify and locate potential outbreaks?

---



---



---



---



(b) What effect could the high number of whooping cough cases have on young children who have not yet completed their vaccination schedule?

---



---

(c) How could the rates of whooping cough be reduced in Australia? \_\_\_\_\_

---



---

4. It is important to maintain high standards of hygiene in medical environments (e.g. hospitals or aged care facilities) to prevent the spread of pathogens. Handwashing with soap and water reduces the number of bacteria present, but it is not always convenient or possible to do this. The use of alcohol-based sanitisers has become a common alternative. The data, right, shows the effect of handwashing or alcohol sanitiser on reducing bacterial load on the fingers of a group of intensive care nurses. 204 samples were taken from the nurses' fingers to determine the base level of contamination (shown by growth of bacterial colonies on agar). The nurses were then split into two groups (soap or alcohol rub). After they had cleaned and dried their hands, the fingers were pressed onto agar to determine the remaining bacterial load.

The effect of hand wash versus alcoholic hand rub on the disinfection of hands.

Bacterial growth on agar plates	Untreated hands (n = 204)	After soap & water wash (n =102)	After alcohol rub (n =102)
No growth or scanty growth (< 20 colonies)	16	51	91
Moderate growth (20-100 colonies)	136	44	5
Heavy growth (> 100 colonies)	52	7	0

Data source: Mäkelä, M et al. (2005) *Infection - J. Clin. Microbiol.* Vol 9(3).

(a) Did the two treatments reduce bacterial contamination?

---

(b) Which treatment was most effective: \_\_\_\_\_

---

(c) What evidence supports your choice? \_\_\_\_\_

---



---

(d) Repeat this experiment for yourselves. What did you find? Attach your results to this page.

5. Hong Kong is a densely packed region, with 7 million people within 1,104 km<sup>2</sup>. In 1997, there was an outbreak of avian influenza virus (bird flu) in Hong Kong. All 18 humans infected had been in recent contact with live domestic fowl (e.g. chickens) in markets. Six of the 18 infected people died. Authorities ordered the slaughter of all live chickens within Hong Kong (1.6 million birds) and stopped the import of more birds. No further cases of bird flu in humans were reported.

(a) Health authorities suspected that the chickens were the source of the virus. Based on the information above decide if you think they were correct in their hypothesis and explain your reasoning:

---



---



---



---



---



---

(b) Based on the evidence, do you think that the large scale slaughter of chickens and the ban on bird imports were justified? Explain your reasoning:

---



---



---



---

# Appendix 1: Equipment List

## 2: Prokaryote and Eukaryote Cells

### INVESTIGATION 2.1 Preparing an onion slide

Per student/group:

- Light microscope
- Onion/onion leaf
- Glass microscope slides
- Coverslips
- Scalpel or razor
- Iodine stain
- Filter paper/tissue paper

## 4: Cell Membrane

### INVESTIGATION 4.1 Simple diffusion across a membrane

Per student/group:

- 200 mL beaker
- 1 mL pipette
- Glucose dipsticks
- Lugol's indicator
- 4 x test tubes
- Dialysis tubing
- Thread or nylon line
- Distilled water
- 1% starch solution
- 10% glucose solution
- Timer or watch

### INVESTIGATION 4.2 Estimating osmolarity

Per student/group:

- 6 x 500 mL beakers
- Balance and equipment to weigh sugar
- Table sugar or lab sucrose
- Potato
- Cork borer or scalpel
- Paper towels
- Marker pen

## 5: Exchange of Nutrients and Wastes

### INVESTIGATION 5.1 Investigating amylase activity

Per class:

- Buffer solutions at pH 4, 5, 6, 7, 8
- 0.1 mol/L iodine solution (I<sub>2</sub>/KI)
- 1% amylase solution
- 1% starch solution

Per student/group/pH

- Timer
- 1 mL pipette
- 2 mL pipette
- Clean syringe
- Test tube
- Spotting plate

## 6: Internal Membranes and Enzymes

### INVESTIGATION 6.1 Investigating peroxidase activity

Per student/group:

- 13 x boiling tubes
- 42 mL distilled water
- 1.8 mL 0.1% H<sub>2</sub>O<sub>2</sub> solution
- 1.2 mL prepared guaiacol solution
- Parafilm
- 6 mL of each pH buffered solution (pH 3, 5, 6, 7, 8, 10)
- 9 mL turnip peroxidase solution
- Test tube rack
- Timer

## 7: Respiration and Mammalian Gas Exchange

### INVESTIGATION 7.1 Measuring respiration in germinating seeds

Per student/group:

- 3 x boiling tubes
- Marker pen
- 6 x cotton balls
- 15% KOH solution
- 2 x eye dropper or plastic pipette
- 3 x gauze pieces
- Germinated bean seeds (enough to fill one quarter of the boiling tube)
- Ungerminated bean seeds (enough to fill one quarter of the boiling tube)
- Glass beads (enough to fill one quarter of the boiling tube)
- 3 x 2-hole tube stoppers
- 3 x bent glass tubes or pipettes
- 3 x tubes (must be able to be clamped shut)
- 3 x screw clips
- A few drops of coloured liquid
- 3 x syringes (must fit tube with screw clamp attached)
- 3 x clamp stands or rack
- Water bath (25°C)
- Ruler
- Timer

### INVESTIGATION 7.2 Investigating yeast fermentation

Per student/group:

- 1 x 100 mL beaker
- 10 g of active yeast
- 50 mL tap water at 24°C
- 25 g of substrate (glucose, maltose, sucrose, or lactose)
- 1 x glass stirring rod
- 1 x conical flask (to hold 275 mL)
- Paraffin oil
- Single hole stopper
- Tubing
- 1 x 100 mL measuring cylinder
- 1 x small basin to hold inverted cylinder
- Stopwatch

## 8: Plant Gas Exchange and Transport Systems

### INVESTIGATION 8.1 Investigating plant transpiration

Per pair/group:

- 250 mL conical flask with rubber bung
- Petroleum jelly
- 1 cm<sup>3</sup> pipette
- Clamp stand
- Leafy plant shoot
- Water
- Cooking oil (for optional set up)
- Timer or watch
- Lamp, or plastic bag and water spray bottle, or fan

## 12: Osmoregulation

### INVESTIGATION 12.1 Comparing stomatal density

Per pair/group:

- Variety of leaf types
- Clear nail varnish
- Microscope slide
- Light microscope (with eyepiece micrometer if available)

## 15: Transmission and Spread of Disease

### INVESTIGATION 15.1 Investigating the effectiveness of handwashing

Per group:

- Warm water
- Soap
- Hand sanitiser

Per individual

- 1 x nutrient agar plates
- Marker pen
- Paper towels
- Incubator (if using)

### INVESTIGATION 15.2 Modelling disease outbreak and spread

Per student/group:

- Computer
- Spreadsheet application (e.g. Excel)

# Appendix 2: Glossary

## A

### ABA (abscisic acid)

Plant hormone with functions in seed and bud dormancy, growth, and stomatal closure.

### absorption

The process by which nutrients are taken up by the body (from intestine to blood).

### accuracy

The correctness of a measurement; how close a measured value is to the true value.

### acetylcholine

Neurotransmitter in both the central and peripheral nervous systems and the only neurotransmitter used in the motor division of the somatic nervous system.

### action potential

A potential difference produced across the plasma membrane of nerve or muscle cells when they are stimulated.

### activation energy

The minimum amount of energy needed to initiate a chemical reaction.

### active site

Region of an enzyme where the substrate binds and undergoes a chemical reaction.

### active transport

The movement of molecules or ions across a cell membrane against a concentration gradient, requiring an expenditure of energy.

### adaptive immune response

The antigen-specific immune response, responsible for immunological memory.

### ADH (anti-diuretic hormone)

The hormone released in response to low blood volumes, high sodium levels and low fluid intake.

### adherence factors

Molecules produced by pathogens that aid in their attachment to host cells, facilitating infection.

### adult stem cell

An unspecialised cell found in the body after development that can give rise to one or more different types of specialised cells.

### aerobic respiration

A form of cellular respiration that requires oxygen to accept electrons during the breakdown of carbohydrate.

### aestivation

A prolonged dormancy in hot and dry conditions. Occurs in a variety of animals including land snails and lung fish.

### allergen

A substance causing an immune response in which the immune system overreacts to a normally harmless substance

### alveoli

Air sacs in the lungs through which carbon dioxide and oxygen are exchanged between the blood and the air.

## ammonia

Highly toxic nitrogenous molecule produced from the breakdown of amino acids.

### amylase

An enzyme responsible for breaking down complex carbohydrates (starches) into simpler sugars like maltose, aiding in the digestion of carbohydrates.

### anabolic reaction

A chemical reaction that constructs large, complex molecules from simpler molecules.

### anabolism

The metabolic process where smaller molecules are combined to form larger, more complex molecules, requiring energy input.

### anaerobic respiration

The process of producing energy from glucose without oxygen, leading to the production of lactic acid or alcohol.

### antibiotic

Chemical substance capable of destroying or reducing the growth of microorganisms, especially bacteria and fungi.

### antibody

A Y-shaped protein produced by B cells in response to an antigen.

### antigen

A molecule that is recognised by the immune system as foreign.

### antiseptic

A substance that prevents the growth of disease-causing microorganisms.

### artificial leaf

A human-made device that mimics the process of natural photosynthesis to convert sunlight into energy, typically using materials like semiconductors to generate fuel.

### assumption

A statement that is assumed to be true but is not (or cannot be) tested.

## ATP

An organic compound that serves as an energy source for metabolic processes.

## B

### B lymphocyte (B cell)

An antibody-producing lymphocyte responsible for humoral immunity.

### bacteria

Member of a large group of microorganisms collectively known as prokaryotes. Cells have a cell wall but lack organelles.

### bacterial disease

Illnesses caused by pathogenic bacteria, leading to a range of symptoms and health issues.

### bias

The influence on the outcome of an experiment due to unintentional or intentional methodology that favours a specific result.

## bioartificial tissue

Engineered tissue created by combining cells with biomaterials to mimic the structure and function of natural tissues for medical purposes.

### bioethics

The ethical, legal, and social, issues involved in biological studies.

### bionic leaf

An artificial system that combines biological components like bacteria with synthetic materials to enhance photosynthetic efficiency and potentially produce renewable fuels.

### biosecurity

Procedures and measures put in place to reduce or stop the entrance of unwanted or pest organisms into a country.

### blood vessels

Tubular structures that form a network throughout the body, carrying blood to and from the heart. Includes arteries, veins, and capillaries.

### Bowman's capsule

A cup-shaped structure in the kidney's nephron that surrounds the glomerulus and is involved in the initial filtration of blood to form urine.

### brown fat

Type of body fat that regulates body temperature, converting energy in glucose directly to heat.

### bush medicine

Traditional remedies and treatments derived from plants and natural resources.

## C

### Calvin cycle

The light-independent phase of photosynthesis during which chemical reactions convert carbon dioxide into sugars.

### capillaries

Thin blood vessels with walls only one cell thick. Capillaries make networks through organs and tissues to allow blood to flow close to all cells for exchange of materials.

### capsid

The protein shell that surrounds and protects the genetic material of a virus, crucial for its structure and function.

### carrier protein

A protein with a function to transport small molecules (or other proteins) through biological membranes.

### catabolism

The metabolic process whereby large molecules are broken down into smaller components, releasing energy.

### catabolic reaction

The breakdown of large, complex molecules into smaller, simpler molecules.

### catalyst

A substance that modifies and increases the rate of a chemical reaction without being consumed in the process.

**cell**

The smallest biological unit that can survive on its own. It is the base unit of all living organisms.

**cell specialisation**

The process where cells develop specific functions and structure in an organism due to the switching on and off of specific genes.

**cell wall**

The rigid outermost cell layer found in plants and certain algae, bacteria, and fungi but absent from animal cells.

**cellular pathogen**

Pathogen (disease causing agent) that is a living cell. Includes some bacteria, fungi, and protists.

**cellular respiration**

The series of metabolic reactions that oxidise organic molecules to produce ATP.

**centrioles**

A cylindrical cell structure made of microtubules which exists as part of the centrosome, found in most eukaryotic cells.

**channel protein**

A protein that allows the transport of specific substances across a cell membrane.

**chemoreceptor**

A sensory receptor that produces an action potential in response to a chemical signal.

**chlorophyll**

A green photosynthetic pigment found primarily in the chloroplasts of algae and plants, essential to photosynthesis.

**chloroplast**

An organelle within the cells of plants and green algae that contains chlorophyll and is the site of photosynthesis.

**cholesterol**

A lipid molecule found in the cell membrane that helps regulate membrane fluidity and stability.

**circulatory system**

System in which blood and plasma is transported around the body inside blood vessels.

**clonal selection**

A theory for how B and T cells are selected to target specific antigens invading the body.

**coenzyme**

A non-protein compound that binds to an enzyme to initiate or aid in its function.

**cofactor**

Substance essential for the correct operating of an enzyme. They bind to the enzyme and may complete the active site or make it more reactive.

**cohesion-tension hypothesis**

The hypothesis that explains how water is transported in plants to extreme heights against the force of gravity.

**collecting duct**

Tubule in the kidney nephron where urine is concentrated as water leaves the tubule by osmosis.

**companion cell**

Specialised plant cells located alongside sieve tube elements in phloem, providing metabolic support and maintaining the function of the sieve tubes for the transport of sugars and other nutrients.

**competitive inhibition**

A form of enzyme inhibition in which an inactive molecule reversibly binds to the active site, preventing the actual substrate from binding (*cf. non-competitive inhibition*).

**complement system**

A group of proteins that enhance the ability of antibodies and phagocytic cells to clear pathogens from the body.

**concentration gradient**

The difference in the amounts of a dissolved substance on either side of a membrane or in two areas of a biological system.

**connective tissue**

Tissue the supports, protects, or gives structure to other tissues and organs.

**contact tracing**

The process of identifying and monitoring individuals who may have come into contact with an infected person.

**control**

A sample in an experiment that is not exposed to the experimental factor/treatment.

**controlled variable**

Any factor that is constant or unchanged throughout the course of an experiment.

**cuticle**

The waxy coating over the epidermis of a leaf.

**cytoplasm**

The watery solution within a cell, including dissolved substances, enzymes, and cell organelles (except for the nucleus).

**D****defensins**

Antimicrobial peptides that play a role in the innate immune response by destroying pathogens.

**denaturation**

The alteration of a protein shape resulting in a loss of function.

**dependent variable**

The variable being tested in a scientific experiment. The response variable.

**depolarisation**

The change in a cell's membrane potential, such that it becomes less negative.

**differentiation**

The normal process by which a less specialised cell undergoes maturation to become a more specialised cell.

**diffusion**

The net movement of molecules from a region of high concentration to one of lower concentration.

**digestion**

The process of breaking down large, insoluble molecules of food into smaller, water-soluble molecules, which can then be absorbed by the body.

**digestive system**

An organ system consisting of the central gastrointestinal tract and associated organs that are responsible for digestion.

**disease**

An abnormal condition of the body when bodily functions are impaired.

**distal convoluted tubule**

A segment of the kidney nephron where further reabsorption and secretion of substances occur before urine is formed.

**downregulation**

The process of reducing or suppressing a stimulus on a cell by reducing the receptors for the stimulus.

**E****effector**

A structure or organ that responds to a stimulus received from a receptor.

**electron micrograph**

A photograph or image of a specimen taken using an electron microscope, which produces images using a beam of electrons.

**electron microscope**

A microscope that uses electron beams to produce high resolution images.

**electron transport chain**

A series of protein complexes that transfer electrons from donors to acceptors across a membrane via redox reactions.

**embryonic stem cell**

A stem cell derived from the early stages of an embryo, which is capable of differentiating into any type of body cell.

**endocytosis**

A process of cellular ingestion by which the plasma membrane folds inward to bring substances into the cell (*cf. exocytosis*).

**endoplasmic reticulum**

A membranous network found in eukaryotic cells, composed of ribosome-studded (rough) and ribosome-free (smooth) regions.

**endotherm**

Animal that is able to produce and maintain its own internal heat.

**enzyme**

Globular proteins that act as biological catalysts for specific reactions.

**epidemic**

The rapid spread of disease to a large number of people in a given population within a short period of time.

**epidermis**

In botany, the single layer of cells that cover the leaves, flowers, roots and stems.

**epidermal tissue**

The tissues (in plants) that make up the outer layer of a plant organ e.g. the leaf.

**epithelial tissue**

The tissues (in animals) that form the covering of all body surfaces..

**estimate**

A rough calculation or determination of a value.

**ethics**

A set of moral obligations that define right and wrong in scientific practices and decisions.

**eukaryotic cell**

A cell that contains a membrane-bound nucleus and organelles.

**excretion**

The process by which organisms expel metabolised waste products and other toxic substances from their body.

**excretory system**

The organ system that removes metabolic wastes and toxins from the body.

**exocytosis**

Secretion of intracellular molecules, contained within membrane-bound vesicles, to the outside of the cell by fusion of vesicles with the plasma membrane (*cf. endocytosis*).

**experiment**

A contrived situation used as a method of investigating causal relationships among variables, or to test a hypothesis.

**extracellular**

Located or occurring outside a cell or cells.

**extracellular receptor**

Receptor located on the outside of the plasma membrane that will respond to an extracellular signal molecule.

**F****facilitated diffusion**

The passive movement of molecules along the concentration gradient.

**fermentation**

An anaerobic metabolic process by which a carbohydrate, such as starch or a sugar, is converted into an alcohol or an acid..

**first messenger**

An extracellular substance that binds to a cell surface receptor and initiates intracellular activity.

**flagellum (pl. flagella)**

A microscopic hair-like structure involved in the locomotion of a cell..

**fluid mosaic model**

The accepted model of a membrane structure, which proposes a double phospholipid bilayer in which proteins and cholesterol are embedded.

**fungal disease**

Fungal diseases are caused by pathogenic fungi, leading to infections that can affect humans, animals, and plants.

**fungus**

Any member of a spore-producing group of eukaryotes that are consumers and have a cell wall made of chitin.

**G****gas exchange**

The process by which oxygen is taken in and carbon dioxide is released from the body, typically occurring in the lungs or through the skin in some organisms.

**germ theory**

States that microorganisms, such as bacteria and viruses, can cause infectious diseases in humans and other organisms.

**glomerulus**

A ball of capillaries surrounded by Bowman's capsule in the nephron, serving as the site of filtration in vertebrate kidneys.

**glycolysis**

The metabolic pathway that converts glucose into pyruvate.

**glycoprotein**

Proteins in the cell membrane that have attached carbohydrate chains, playing roles in cell recognition and cell-cell interactions.

**Golgi**

An organelle found in eukaryotic cells that packages and transports molecules from the endoplasmic reticulum to their destination.

**granum (pl., grana)**

Stack of thylakoids found in the stroma of chloroplasts, where the light-dependent reactions of photosynthesis take place.

**graph**

A diagram displaying numerical information in a way that can be used to identify trends in the data.

**guard cells**

The two cells that flank the stomatal pore and regulate the pore's opening and closing.

**H****haemoglobin**

A large iron-containing protein, which transports oxygen in the blood of vertebrates.

**halophyte**

A plant that can grow and survive in environments with a high salt content (low availability of free water).

**hibernation**

A state of minimal metabolic activity entered into to conserve energy, usually in colder months.

**hierarchical organisation**

The arrangement of cells, tissues, organs, and organ systems in a structured manner that contributes to the overall function and efficiency of a multicellular organism.

**homeostasis**

The steady-state physiological condition of the body.

**hormone**

Chemical messengers secreted directly into the blood, where they circulate to exert specific effects on target tissues and organs.

**host**

An organism that harbours another organism or a pathogen, providing an environment for the pathogen to live and reproduce.

**hydrophilic**

Interacting readily with, or attracted to, water (*cf. hydrophobic*).

**hydrophilic signal**

Signal molecule blocked by the plasma membrane. Attached to receptors on the outside of cells (extracellular).

**hydrophobic**

Not readily interacting with water;

a molecule or substance that does not dissolve in or mix with water (*cf. hydrophilic*).

**hydrophobic signal**

Signal molecule that can pass through the plasma membrane and attach to intracellular receptors.

**hydrophyte**

A plant that grows either partly or totally submerged in water.

**hyperpolarisation**

The change in a cell's membrane potential, such that it becomes less negative.

**hypertonic**

A solution with higher osmotic pressure than another solution; a solution that, when surrounding a cell, will cause the cell to lose water (*cf. hypotonic*).

**hypothesis**

A tentative explanation, proposition, or set of propositions capable of being tested by scientific experimentation.

**hypotonic**

A solution with lower osmotic pressure than another solution; a solution that, when surrounding a cell, will cause the cell to take up water (*cf. hypertonic*).

**IJ****immune response**

How the human body recognises and defends itself against bacteria, viruses, and substances that appear foreign and harmful to the body.

**immunisation** *see vaccination*

**immunity**

Inherited, acquired, or induced resistance to infection by a specific pathogen.

**independent variable**

The variable that is changed or controlled in a scientific experiment, and is assumed to have an effect on the dependent variable.

**induced fit model**

A model for the interactions between an enzyme and substrate involving conformational change in the enzyme.

**inflammation**

A defensive response to damage caused by physical agents, microbial infections or chemical agents. The inflammation process involves pain, redness, heat and swelling.

**inflammatory response**

The response that triggers white blood cells to an injury site after damage to body tissues.

**infectious disease**

A disease caused by a communicable or transmissible pathogen.

**innate response**

The immediate, non-specific defence mechanism of the immune system against pathogens.

**insulation**

Material (e.g. fur, feathers) that trap air and stop heat escaping from the body.

**intracellular**

Occurring or situated within a cell or cells.

**intracellular receptor**

Globular proteins, often transcription factors, activated by intracellular signal molecules.

**invasion factors**

Illnesses caused by pathogenic agents like bacteria, viruses, and parasites that can be transmitted from one individual to another.

**ion**

An electrically charged atom or molecule resulting from the loss or gain of electrons.

**ion pump**

Membrane proteins that is capable of transporting ions against a concentration gradient using the energy from ATP.

**isotonic**

Referring to a solution that, when surrounding a cell, causes no net movement of water into or out of the cell.

**K****kidney**

In vertebrates, one of a pair of excretory organs where blood filtrate is formed and processed into urine.

**killer (cytotoxic) T cell**

A type of T lymphocyte that kills infected cells, cancer cells, and damaged cells.

**kleptothermy**

A thermoregulatory behaviour in which individuals group together to share or benefit from the body heat of others.

**Koch (Robert)**

Robert Koch was a prominent German scientist who made significant contributions to microbiology and is known for his work on infectious diseases.

**Krebs cycle**

A cycle of aerobic catalysed reactions in respiration occurring within mitochondria. Generates ATP and reducing power.

**L****large intestine**

In vertebrates, the last part of the gastrointestinal tract and digestive system, comprising the caecum, colon, and rectum.

**leaf**

The lateral appendage of the plant vascular system. The primary site of photosynthesis and gas exchange.

**light (optical) microscope**

An optical instrument with lenses that refract visible light to magnify images of specimens.

**light dependent phase**

The phase of photosynthesis during which light energy is converted into chemical energy through chemical reactions.

**light independent phase**

See **Calvin cycle**.

**lipase**

An enzyme that breaks down fats (lipids) into fatty acids and glycerol, aiding in the digestion and absorption of fats.

**lock-and-key model**

Model of enzyme action in which the

active site is specifically structured to fit the substrate. The substrate binds to the active site in order for the reaction to take place.

**loop of Henle**

Part of the kidney nephron between the proximal convoluted tubule and the distal convoluted tubule. Its function is to create a gradient in salt concentration through the medullary region of the kidney.

**lungs**

A pair of gas exchange organs located on either side of the chest cavity in mammals.

**lysosome**

A membrane-enclosed sac of hydrolytic enzymes found in the cytoplasm of animal cells and some protists.

**M****macrophage**

A large phagocytic leucocyte, which engulfs and devours invading cells.

**magnification**

The amount or degree of visual enlargement of an observed object.

**major histocompatibility complex**

A tightly linked cluster of genes that code for cell surface proteins essential for the adaptive immune system.

**mass flow hypothesis**

A hypothesis that describes the movement of sap in plant phloem through diffusion gradients and hydrostatic pressure.

**matrix (of mitochondria)**

The space within the inner membrane of a mitochondrion.

**mean**

The sum of the data divided by the number of data entries; a measure of central tendency in a normal distribution.

**mechanoreceptor**

A sensory receptor sensitive to mechanical energy, i.e. stretching or bending.

**median**

The middle number in an ordered sequence of numbers. For an odd number of values, it is the average of the two middle numbers.

**memory cells**

Long lived cells derived from B cells that are retained in the lymph nodes and provide future immunity by rapidly differentiating in plasma cells when an infection is re-encountered.

**mesophyte**

Terrestrial plant adapted to neither a particularly dry nor particularly wet environment, i.e. a mesophytic environment of adequate water and moderate temperatures.

**metabolic pathway**

A linked series of chemical reactions occurring within a cell.

**metabolism**

The chemical processes occurring within a living cell/organism that sustain life.

**microvilli**

Tiny projections on the surface of cells in

the small intestine that increase surface area for absorption of nutrients during digestion.

**mitochondrion (pl., mitochondria)**

An organelle in eukaryotic cells that serves as the site of cellular respiration.

**mitosis**

The phase of the cell cycle resulting in nuclear division.

**mode**

The value that occurs most often in a data set.

**model**

A conceptual, mathematical or physical representation of a real-world phenomenon.

**motor neuron**

A nerve cell that carries impulses away from the central nervous system to an effector muscle. May be non-myelinated or myelinated.

**multipotent**

The ability (of stem cells) to develop into more than one cell type related to the tissue of origin; adult stem cells.

**muscle tissue**

Tissue composed of muscle cells (fibres) specialised for contracting.

**myelin**

A lipid-rich substance surrounding and insulating the axon of nerve fibres.

**N****natural killer cells:**

A type of lymphocyte that can identify and destroy virus-infected or cancerous cells without prior sensitisation.

**negative feedback**

In physiology, a primary mechanism of homeostasis where a change in a variable triggers a response that counteracts the initial change.

**nephron**

The tubular excretory unit of the vertebrate kidney.

**nerve impulse**

Waves of depolarization that create action potentials moving along the axon length.

**nervous tissue**

Tissue composed of nerves cell. Specialised for conducting signals.

**neuron**

Cell of the nervous system that transmits information to other nerve cells.

**neutrophils**

A type of white blood cell that plays a key role in the innate immune response by engulfing and destroying pathogens.

**nociceptor**

Sensory receptor, normally a free nerve ending, that responds to damage or potentially damaging stimuli as pain.

**node of Ranvier**

The gaps occurring at intervals along the myelin sheath.

**non-cellular pathogen**

A pathogen that is not a living cell.

**non-competitive inhibition**

Enzyme inhibition where the inhibitor

binds to the enzyme at a region other than the active site (*cf. competitive inhibition*). Includes viruses and prions.

#### **non-infectious disease**

Disease that is not transmissible, with the exception of inheritance, and is often caused by genetic or lifestyle factors. Also referred to as non-communicable disease (NCD).

#### **nonself antigen**

Antigens that do not originate in an organism's own body

#### **nucleolus**

A specialised spherical structure in the nucleus, consisting of chromosomal regions containing ribosomal RNA (rRNA); the site of rRNA synthesis and ribosomal assembly

#### **nucleus**

The organelle of a eukaryotic cell that contains the genetic material in the form of chromosomes, made up of chromatin.

#### **null hypothesis**

The hypothesis of no difference or no effect that will be statistically accepted or rejected by testing against another alternative hypothesis, subject to a given level of error.

#### **nutrition**

The process by which organisms obtain and utilise food for growth, energy, and maintaining health.

### **O**

#### **observation**

The act of seeing and noting an occurrence in the object or substance being studied.

#### **optimum (enzyme)**

The conditions under which a particular enzyme is most active.

#### **organ**

A group of different tissues working together to perform a particular function.

#### **organelles**

subcellular structures with one or more specific jobs to perform in the cell.

#### **osmoconformer**

An organism that maintains its internal osmolarity isotonic to the environment it is in.

#### **osmolarity**

The concentration of a solution expressed as the total number of solute particles per litre; number of moles of solute per litre.

#### **osmoregulation**

The maintenance of the internal balance of water and dissolved materials when in a changing external environment.

#### **osmoregulator**

An organism that maintains a constant internal osmolarity that is not influenced by the external environment.

#### **osmosis**

The diffusion of free water across a selectively permeable membrane.

#### **outbreak (disease)**

A sudden increase in the occurrence of a disease in a particular time and place.

### **P**

#### **pandemic**

A disease that is continuing to spread and which has become prevalent throughout most of the world.

#### **panting**

Quick, short breaths that draw air in and out over the moist linings of the tongue and mouth in order to increase evaporation and help reduce body heat.

#### **partially permeable**

A membrane that is permeable to the small molecules of water and certain solutes but does not allow the passage of large solute molecules.

#### **passive transport**

The diffusion of a substance across a biological membrane with no expenditure of energy.

#### **Pasteur (Louis)**

Louis Pasteur. French chemist and microbiologist who founded many principles of modern microbiology, disproving spontaneous generation, developing the germ theory of diseases, and the process of pasteurisation.

#### **pathogen**

Microorganism that causes disease.

#### **pathogenesis**

The process by which a disease develops and progresses within an organism, involving the interaction between the host and the pathogen.

#### **percentage**

The number of items or part of a whole expressed as a number out of 100.

#### **phagocyte**

A cell that protects the body by engulfing and ingesting harmful foreign particles, bacteria, and dead or dying cells.

#### **pheromone**

A chemical produced by an individual and released into the environment that has an effect on the behaviour of another individual of the same species.

#### **phloem**

Living plant vascular tissue that transports sugar and other nutrients.

#### **phospholipid**

A lipid composed of glycerol joined to two fatty acids and a phosphate group. Phospholipids form bilayers that function as biological membranes.

#### **photoreceptor**

A receptor that responds to light.

#### **proprioceptor**

Receptors in the muscles, tendons and joints that monitor limb position, stretch and tension.

#### **photosynthesis**

A process used by green plants, algae, and some bacteria to convert light energy into chemical energy (carbohydrate).

#### **plasma membrane**

The membrane at the boundary of every cell that acts as a selective barrier, regulating the cell's chemical composition.

#### **plasmolysis**

A phenomenon in walled cells in which the cytoplasm shrivels and the plasma membrane pulls away from the cell wall;

occurs when the cell loses water to a hypertonic environment.

#### **pluripotent**

Ability (of a stem cell) to give rise to all the cells of the adult body, but not extra-embryonic tissues such as the placenta.

#### **polar**

A molecule that carries a partial positive charge on one side and a partial negative charge on the other.

#### **potency**

The ability of a stem cell to differentiate into different cell types.

#### **precision**

The consistency of results when measurements or tests are repeated. Precision is independent of accuracy.

#### **prediction**

What is expected to happen if the hypothesis of an experiment or scenario is true.

#### **prion**

One of a group of infectious proteins that cause several neuro-degenerative diseases in humans and other animals including CJD in humans and BSE in cattle.

#### **prokaryotic cell**

A type of cell lacking a membrane-enclosed nucleus or other cell organelles.

#### **proportion**

The part in comparison to the whole, often expressed as a fraction.

#### **prostaglandins**

Lipid compounds that play a role in inflammation, immune response, and the regulation of various physiological processes.

#### **protease**

Enzyme able to breakdown proteins by hydrolysis of peptide bonds.

#### **protein channel**

Specialised proteins in the cell membrane that create passageways for specific molecules or ions to move across the membrane.

#### **proximal convoluted tubule**

The initial segment of the kidney nephron where reabsorption of water, ions, and nutrients occurs after filtration in the glomerulus.

### **Q**

#### **qualitative data**

Non-numerical data that describes qualities or characteristics.

#### **quantitative data**

Numerical data expressing a certain quantity, amount, or range.

#### **quarantine**

Disease control mechanism to isolate exposed individuals that *may* be infected.

### **R**

#### **rate**

How quickly one variable is changing with respect to another variable. E.g. the volume of gas produced in a reaction per unit of time.

**ratio**

The relationship between two numbers expressed as how many times bigger one value is compared to the other.

**refractory period**

The period during a nerve impulse when the sodium channels are inactivated and the neurone cannot respond.

**reliability (of data)**

The degree of consistency of a measurement (see *precision*).

**resolution**

A microscope's ability to distinguish detail.

**respiratory gases**

Gas involved in respiration and gas exchange, oxygen and carbon dioxide.

**respiratory system**

The organ system responsible for breathing and gas exchange in the body.

**respirometer**

A device used to measure the rate of oxygen consumption or carbon dioxide production in organisms.

**response**

Any behaviour of a living organism that results from an external or internal stimulus

**resting potential**

The potential difference across the cell membrane of a neurone when there is no impulse passing.

**ribosome**

A complex of rRNA and protein molecules that function as a site of protein synthesis in the cytoplasm.

**rough endoplasmic reticulum (rER)**

The portion of the endoplasmic reticulum with ribosomes attached.

**S****scientific method**

The use of an ordered, repeatable method to investigate, manipulate, gather, and record data.

**second messenger**

A molecule that relays signals from receptors on the cell surface to a target molecule inside the cell.

**self renewal**

The process of giving rise to indefinitely more cells of the same cell type.

**self antigen**

Antigen produced in the organism's own body and is recognised as such by the immune system.

**Semmelweis (Ingaz)**

Hungarian physician, recognised for his pioneering work in hand hygiene and infection control practices in healthcare settings.

**sensory neuron**

A nerve cell concerned with transmitting impulses from sensory receptors to other neurones.

**sensory receptor**

A specialised cell that detects stimuli and responds by producing an electrical discharge.

**sieve plate:**

A perforated structure found in phloem that separates individual sieve tube elements, allowing for the movement of sugars and other nutrients between cells.

**sieve tube**

A specialised plant cell involved in transporting organic compounds like sugars through the phloem, with perforated sieve plates facilitating the flow of nutrients.

**signal molecule**

Molecules that are responsible for transmitting information between cells in the body.

**significant figures**

Any of the digits of a number, beginning with the first non zero digit farthest to the left and ending with the last digit farthest to the right. Usually stated to a specific number of significant figures (e.g. 2).

**signal transduction**

A mechanism that converts a mechanical or chemical stimulus into a specific cellular response.

**small intestine**

Part of the gastrointestinal tract where most of the absorption of nutrients occurs.

**smooth endoplasmic reticulum (sER)**

The portion of the endoplasmic reticulum that lacks ribosomes.

**specialised cell**

A cell that has developed the characteristics needed to perform particular functions.

**spread (disease)**

The dissemination of a disease-causing agent within a population or environment.

**stain**

A chemical that binds to parts of the cell and allows those parts to be seen more easily under a microscope.

**statistical test**

A calculation used to determine if the null hypothesis is to be accepted or rejected. The greater the significance level of the test the greater the likelihood the outcome of the calculation will represent the true situation.

**stem**

An organ in vascular plants that supports the above-ground parts of the plant, transports water and dissolved substances, and produces new living tissue.

**stem cell**

An undifferentiated cell, characterised by self renewal and potency.

**stimulus**

A physical or chemical change in the environment which causes a response in an organism.

**stomach**

An organ of the digestive system that stores food and begins protein digestion.

**stoma (plural, stomata)**

A microscopic pore in the epidermis of leaves and stems that allows gas exchange between the plant and the environment.

**stroma**

The fluid surrounding the membranous stacks within the chloroplast.

**summation**

The cumulative effect of a number of nerve impulses.

**surface area : volume ratio**

The amount of surface area per unit volume of an object.

**sweating**

Fluid is excreted via the sweat glands of mammals, and the evaporation cools the body.

**synapse**

The point of communication between one nerve cell and another or between a nerve cell and a non-neuronal target cell such as muscle.

**synaptic integration**

The receiving and processing of signals at a synapse and the generation of a nerve impulse (either all or none).

**system (body)**

A group of organs working together to perform complex functions and maintain homeostasis.

**T****T lymphocytes (T cell)**

A type of lymphocyte that plays a central role in the adaptive immune response.

**table**

A way of presenting data in a structured format that allows relationships and trends to be easily recognized.

**target cell**

Cells that responds to a specific messenger.

**thermogenesis**

The production of heat in the body by the conversion of chemical energy into heat.

**thermoregulation**

The maintenance of internal body temperature within a tolerable range.

**thermoreceptor**

A receptor in the dermis that detects changes in the skin temperature outside the normal range.

**threshold potential**

The critical potential that must be reached before an action potential is initiated in an excitable cell.

**thylakoids**

Membrane-bound sacs containing chlorophyll; the site of the light-dependent reactions of photosynthesis.

**thyroxine**

Hormone secreted into the blood by the thyroid gland. Plays a role in the rate of metabolism.

**tissue**

An integrated group of cells with a common structure, function, or both.

**torpor**

A state of reduced physiological activity in which the body temperature and metabolic activity lowers.

**totipotent**

Ability (of a stem cell) to give rise to all the cells of the adult body and extra-embryonic tissues such as the placenta.

**toxins**

Harmful substances produced by pathogens, such as bacteria and fungi, that can damage cells and tissues in the host organism.

**toxins (plant)**

Plant toxins are poisonous substances produced by plants that can be harmful to animals or humans if ingested or exposed to them.

**transcription factor**

Proteins that bind to specific DNA sequences to regulate transcription.

**transducer**

Sensory receptor that responds to stimuli by producing an electrical (or chemical) discharge converting the energy from a stimulus into an electrochemical signal.

**translocation**

The transport of organic nutrients in the phloem of vascular plants.

**transmission (disease)**

The process by which a disease is passed from one person or organism to another.

**transpiration**

The evaporative loss of water from a plant.

**transplantation (organ)**

The surgical procedure of transferring an organ from one person (donor) to another (recipient) to replace a damaged or failing organ, often crucial for saving lives.

**trend**

A pattern observed in data showing that there may be a predictable relationship between data values.

**turgor**

Distension or rigidity of plant cells, resulting from fluid pressure against the rigid cell wall.

**UVW****unipotent**

Unipotent stem cells can only differentiate into one type of cell, crucial for specific functions like reproduction and tissue repair.

**upregulation**

The process of increasing a cell's response to a stimulus by increasing the receptors for the stimulus.

**urea**

Relatively nontoxic and soluble nitrogenous waste excreted by mammals and amphibians.

**uric acid**

White, paste-like nitrogenous waste product, with a low solubility, excreted by birds, reptiles, and insects.

**urine**

Fluid containing metabolic wastes which, in vertebrates, collects in the urinary bladder.

**vaccination**

Inoculation with a vaccine to help the immune system develop immunity to a particular disease.

**vaccine**

A substance used to stimulate the production of antibodies and provide immunity against specific disease.

**vacuole**

A membrane-bounded vesicle whose function varies in different kinds of cells.

**variable**

A measurable property that changes over time or can take on different values.

**vascular tissue**

Plant tissue consisting of cells joined into tubes that transport water and nutrients.

**vasoconstriction**

The process by which blood flow to the skin is decreased by constriction of the capillaries.

**vasodilation**

The process by which blood flow to the skin is increased by expansion of the capillaries.

**vector**

An organism or artificial vehicle that is capable of transferring disease to another organism.

**viral disease**

Illnesses caused by viruses, which can infect various organisms, including humans, animals, and plants.

**virus**

A non-cellular obligate intracellular parasite, requiring a living host to reproduce.

**waste (removal)**

The process of eliminating metabolic by-products and other unnecessary substances from the body to maintain internal balance and health.

**XYZ****xerophyte**

A plant adapted to an arid climate.

**xylem**

Vascular plant tissue consisting of tubular dead cells that conduct water and minerals up the plant from the roots.

**zygote**

A fertilised egg cell.

## Questioning Terms

The following terms are often used when asking questions in examinations and assessments.

- Analyse: Interpret data to reach stated conclusions.
- Annotate: Add brief notes to a diagram, drawing or graph.
- Apply: Use an idea, equation, principle, theory, or law in a new situation.
- Calculate: Find an answer using mathematical methods. Show the working unless instructed not to.
- Compare: Show similarities between two or more items, referring to both (or all) of them throughout.
- Construct: Represent or develop in graphical form.
- Contrast: Show differences. Set in opposition.
- Define: Give the precise meaning of a word or phrase as concisely as possible.
- Derive: Manipulate a mathematical equation to give a new equation or result.
- Describe: Define, name, draw annotated diagrams, give characteristics of, or an account of.
- Design: Produce a plan, object, simulation or model.
- Determine: Find the only possible answer.
- Discuss: Show understanding by linking ideas. Where necessary, justify, relate, evaluate, compare and contrast, or analyse.
- Distinguish: Give the difference(s) between two or more items.
- Draw: Represent by means of pencil lines. Add labels unless told not to do so.
- Estimate: Find an approximate value for an unknown quantity, based on the information provided and application of scientific knowledge.
- Evaluate: Assess the implications and limitations.
- Explain: Provide a reason as to how or why something occurs.
- Identify: Find an answer from a number of possibilities.
- Illustrate: Give concrete examples. Explain clearly by using comparisons or examples.
- Interpret: Comment upon, give examples, describe relationships. Describe, then evaluate.
- List: Give a sequence of answers with no elaboration.
- Outline: Give a brief account or summary. Include essential information only.
- Predict: Give an expected result.
- Solve: Obtain an answer using numerical methods.
- State: Give a specific name, value, or other answer. No supporting argument or calculation is necessary.
- Suggest: Propose a hypothesis or other possible explanation.
- Summarise: Give a brief, condensed account. Include conclusions and avoid unnecessary details.

## Image Credits

**We acknowledge the generosity of those who have provided photographs for this edition:**

• Louisa Hayward (Dartmouth College) for the centriole image • Kristian Peters • Waikato Institute of Technology for the photo of the athletes being monitored on cycles • Louisa Howard, Katherine Connolly Dartmouth College • D. Fankhauser, University of Cincinnati, Clermont College • Silas Pryor • Andrea Braakhuis, Wintec • Berkshire Community College Bioscience Image Library •

**We also acknowledge the photographers that have made their images available through Wikimedia Commons under Creative Commons Licences 1.0, 2.0, 2.5, 3.0, or 4.0:**

• James Scott • Olaboy • Suseno • GerryShaw • James Hedberg • Mnoif • Vossman • A2-33 • Andrei Lomize OPM database • Zephyris • MCC UW • Dartmouth College • tooony • Alison Roberts • CDC Dr Lucille K. Georg • PloS • Vossman • PDB • Matthias Zepper • National Cancer Institute • HIA • Nephron • McKDandy • bauer • Ragesoss • Lavin • Roadnottaken • Sebastian80 • Sichy007 • Mohammad Golkar • Bernard DUPONT • Phil Spark • Ryan Bushby • Ildar Sagdejev • HG6996 • Bahudhara • Nhobgood • AlbertKok • SaleemHameed • DannyS • BoundaryRider • NIAID • Matt Lavin • Sparks • CDC: Janice Haney Carr • Emmanuelm • Ute Frevert, Plos • Edward L. Barnard, Florida Department of Agriculture and Consumer Services, Bugwood.org • John H. Ghent, USDA Forest Service, Bugwood.org • looksclose • Volker Brinkmann PLOS • Bjørn Christian Tørrisse • Danny Cho • Julien Harne • Mark Marathon • John Jennings • Geoff Derrin • Casliber • Obli • Forrest Brem

**Contributors identified by coded credits are:**

BF: Brian Finerran (Uni. of Canterbury) • CDC: Centers for Disease Control and Prevention, Atlanta, USA • EII: Education Interactive Imaging • JDG: John Green (University of Waikato), • KP: Kent Pryor • NASA: National Aeronautics and Space Administration • NIH: National Institute of Health • RA: Richard Allan • RCN: Ralph Cocklin • WBS: Warwick Silvester (Uni. of Waikato) • USDA: United States Department of Agriculture WMU: Waikato Microscope Unit. • NOAA: National Oceanic and Atmospheric Administration • NIAID National Institute of Allergy and Infectious Diseases • USAF: United States Air Force

**Royalty free images, purchased by BIOZONE International Ltd, are used throughout this workbook and have been obtained from the following sources:**

• Adobe Stock, stock.adobe.com • iStock images • Dollar Photo Club • Corel Corporation from various titles in their Professional Photos CD-ROM collection • Image stills from Sketchfab • IMSI (International Microcomputer Software Inc.) images from IMSI's MasterClips® and MasterPhotosTM Collection, 1895 Francisco Blvd. East, San Rafael, CA 94901-5506, USA • ©1996 Digital Stock, Medicine and Health Care collection, ©Hemera Technologies Inc, 1997-2001 • © 2005 JupiterImages Corporation www.clipart.com • ©1994 ©Digital Vision • Gazelle Technologies Inc. • ©1994-1996 Education Interactive Imaging (UK) • PhotoDisc®, Inc. USA, www.photodisc.com. We also acknowledge the following clipart providers: TechPool Studios, for their clipart collection of human anatomy: Copyright ©1994, TechPool Studios Corp. USA (some of these images have been modified); Totem Graphics, for clipart; Corel Corporation, for vector art from the Corel MEGAGALLERY collection.

# Index

95% confidence interval 24

## A

Aboriginal medicine 311  
 Accuracy, measurements 10-11  
 Acetylcholine, as a neurotransmitter 204  
 Acquired immunity 280-281  
 Action potential 204  
 Activation energy, enzyme 135  
 Active immunity 280  
 Active site, of enzyme 133  
 Active transport 101-105  
 Adaptations, for water balance 240  
 Adaptive immune response 269, 276-279  
 Adenosine triphosphate (ATP) structure 151  
 Adult stem cell 61  
 Aerobic pathways 152  
 Aerobic respiration 152, 155-156  
 Aestivation 226  
 AIDS 263  
 Albinism 144-145  
 Alcoholic fermentation 152, 157  
 Allergens 268  
 Alveolus, structure 163  
 Ammonia, as waste 125  
 Amylase activity 123-124  
 Anabolic reaction 135, 150  
 Anaerobic pathways 152, 157  
 Anaerobic respiration 152  
 Analogy (as a model) 4  
 Anaphase 64  
 Animal cell 35-36  
 - features of 43-44  
 - organelles 43-44, 46  
 Annotated diagrams, rules 53  
 Antagonistic hormones 213  
 Antibiotics, testing effectiveness 295-296  
 Antibody 269  
 - structure 279  
 Antidiuretic hormone 241  
 Antigen 267, 269  
 - processing 275  
 - response to 281  
 - types 268  
 Apparatus, selection of 16  
 Aquaporin protein 84  
 Arterial circulatory system 111  
 Arteries 112  
 Assumptions 3  
 ATP production 152  
 ATP  
 - and metabolism 151  
 - structure 151  
 Autocrine signalling 210  
 Autotrophs 169

## B

B-lymphocytes 276  
 Bacterial cell 35-36, 39-40  
 Bacterial diseases 256-257  
 Bar graph 17  
 Baroreceptors 196  
 Bias, in sampling 26  
 Bimodal distribution 21  
 Biological catalyst 135  
 Biological drawing, rules 52-53  
 Biological stains 50

Biosecurity 308  
 Blood 113  
 Blood glucose, regulation 213, 217  
 Blood pH, maintaining 194  
 Blood vessels, types 112-114  
 Body systems, interaction of 72-77  
 Body temperature, maintaining 195  
 Bohr effect 165  
 Brown fat, and thermoregulation 227, 230

## C

Calvin cycle 171  
 Capillaries 112-114  
 Capillary networks 113-114  
 Carbohydrates 37  
 Carbon dioxide, transport of 164  
 Catabolic reaction 135, 150  
 Catalyst, biological 135  
 Causation 20  
 Cell shapes  
 - bacterial 39  
 - prokaryotes 39  
 Cell signalling, types 210-211  
 Cell size  
 - and diffusion 92, 94-95  
 - compared 93  
 Cell theory 35  
 Cell to cell signalling 210  
 Cell types 35-36  
 Cell, composition of 37  
 Cellular differentiation 62-63  
 Cellular environments 38  
 Cellular functions 38  
 Cellular respiration 168  
 - and gas exchange 160  
 - measuring 153-154  
 - overview 155-156  
 Central nervous system 198  
 Central tendency measures 21-22  
 Chemical defences, plants 289-290  
 Chemoreceptors 196-197  
 Chi squared test 32  
 Chloroplast  
 - and photosynthesis 171  
 - structure 170  
 Cholinergic synapse 204  
 Circulatory system  
 - interactions 72-77  
 - mammalian 111-115  
 Citations, how to 2  
 Clonal selection 278  
 Cofactors, and enzymes 136  
 Cohesion-tension 180  
 Column graph 17  
 Compartmentalisation, and enzyme activity 142  
 Complement protein 270  
 Compound microscope 47  
 Concentration gradient 89  
 Connective tissue 70  
 Continuous data 5  
 Control centre 193  
 Controlled variable 6  
 Conversion factors 12  
 Correlation 20  
 Correlation vs causation 20

Cotransport 102  
 Countercurrent heat loss 228  
 Covid-19  
 - transmission 294, 306-307  
 - vaccine development 285  
 - vaccine effects 286-287  
 Cytokinesis 64  
 Cytosis 103-104

## D

Data transformation 12, 15  
 Data, types 5  
 Decimal form 12  
 Defence, against disease 269  
 Defensins 290  
 Denaturation, enzyme 137  
 Dendrites 199  
 Dendritic cell 270, 277  
 Dependent variable 6  
 Developmental cell layers 63  
 Dialysis tubing 97  
 Diffusion  
 - influence of cell size 92, 94-95  
 - types 89-91  
 Digestive enzymes 118  
 Digestive system  
 - interactions 74-75  
 - mammalian 116-125  
 Discontinuous data 5  
 Discrete data 5  
 Disease outbreak, modelling 301-303  
 Disease patterns 301-305  
 Disease spread 306-307  
 Disease transmission 293-294  
 - theories 255-256  
 Disease  
 - AIDS 263  
 - bacterial 256-257  
 - containment 306-307, 309-310  
 - fungal 258  
 - HIV 262-263  
 - infectious 253, 284  
 - influenza 305  
 - malaria 260  
 - metabolic 144-145  
 - non-infectious 253  
 - polio 310  
 - prion 264  
 - protistan 259-260  
 - smallpox 310  
 - spread of 297-298  
 - viral 261-263  
 Dissecting microscope 47  
 Distribution, of data 21  
 Downregulation 211  
 Drawing, for biology 52-54  
 Drugs, effect on nerve transmission 207

## E

Ectoderm 63  
 Effector 193, 198  
 Electron microscopes 55-56  
 Electron transport chain 152, 155-156  
 Embryonic stem cell 61  
 Endocrine signalling 210  
 Endocytosis 104  
 Endoderm 63  
 Energy transformation, in cells

168  
 Enzyme 133-141  
 - activity 135-141  
 - cofactor 136  
 - denaturation of 137  
 - inhibition of 138  
 - models of action 134-135  
 - structure 133  
 Enzymes  
 - metabolic efficiency 142-143  
 - and metabolism 150  
 - and photosynthesis 171, 174  
 - digestive 118  
 - investigating activity 123-124  
 - role in disease 144-145  
 Epithelial tissue 70  
 Error  
 - reduction of 11  
 - sources of 16  
 Estimates 12  
 Ethical guidelines 8-9  
 Ethics  
 - of medical research 78  
 - of stem cell use 67  
 - of transplants 130  
 Eukaryotic cell 35-37, 39-46  
 Evaporative heat loss 227  
 Excretory system, mammalian 126-129  
 Exocytosis 103  
 Experimental control 6  
 Exponential functions 14  
 Extracellular enzyme 133  
 Extracellular receptor 215

## F

Facilitated diffusion 89  
 Feedback mechanisms, and homeostasis 194-195  
 Fermentation pathways 152, 158  
 Fish, osmoregulation 238  
 Fluid balance, in land animals 239  
 Fluid mosaic model, of membrane structure 81  
 Fractions 13  
 Fungal cell 35-36  
 Fungal disease 258

## G

G-protein 84  
 Gas exchange membrane 163  
 Gas exchange system  
 - human 161-165  
 - interactions 72-73  
 Gas exchange, principles 160  
 Gas transport, humans 164-165  
 Gastric gland 117  
 Germ layers 63  
 Gibberellic acid 214  
 Glucagon 213  
 Glucose transporter protein 84  
 Glycolysis 152, 155-156  
 Gradient, of a line 19  
 Graphs, types 17-18  
 Guard cell 179, 181-182

## H

Haemoglobin 164-165  
 Haemolysis 100  
 Halophytes 244, 247-248  
 Hand washing, effectiveness

- 299-300  
Health programs, effectiveness of 309-310  
Heart structure 114  
Herd immunity 283  
Hibernation 225  
Hierarchical organisation, of life 68-69  
Histogram 17  
HIV 262-263  
Homeostasis, defined 193  
Hormonal control 198  
Hormone regulation 218  
Hormones  
- actions of 213  
- and dairy industry 219  
- and thermoregulation 229-230  
- and water balance 241  
- as signal molecules 212  
Hydrophytes 242  
Hypertonic solution 97, 99  
Hypothesis 3  
Hypotonic solution 97, 99
- I**  
Immunisation response 281, 286-287  
Immunity  
- acquired 280-281, 286-287  
- and lymphatic system 274  
Independent variable 6  
Indigenous practices, medicine 311  
Induced fit model, of enzyme function 134  
Infectious disease 253, 284  
- containment 309-310  
Inflammation response 270-271  
Influenza spread 298, 305
- Inhibition, of enzyme activity 138, 142  
Innate immune response 269  
Inorganic ions 37  
Insulation, and thermoregulation 223  
Insulin 213  
- action of 217  
- and thermoregulation 230  
Intercept, of a line 19  
Interneuron 199  
Interphase 64  
Intestinal villi, role in absorption 118  
Intracellular enzyme 133  
Intracellular receptor 215  
Iodine test, for starch 183  
Ion channels, and sensory reception 197  
Ion pumps 102  
Isotonic solution 97
- K**  
Kidney,  
- function 128-129  
- structure 127-128  
Knee jerk reflex 201  
Koch, Robert 255-256  
Krebs cycle 152, 155-156
- L**  
Lactic acid fermentation 152, 157  
Large intestine 122  
Levels of organisation, biological 68-69  
Light and photosynthesis 173
- Light dependent phase, of photosynthesis 171-172  
Light independent phase, of photosynthesis 171-172  
Line graph 17, 19  
Line of best fit 20  
Linear magnification, calculating 51  
Link reaction 155-156  
Lipid bilayer 81-83  
Lipids 37  
Local signalling 210  
Lock and key model, of enzyme function 134  
Log transformations 14  
Loop of Henle 129  
Lungs, structure 162-163  
Lymph 113  
Lymphatic system 274  
Lymphocyte 274, 275, 278  
- functions 276-277
- M**  
Macrophage 270  
Magnification, in microscopy 47, 51  
Major histocompatibility complex (MHC) 267, 275  
Malaria 260  
Mast cell 270  
Mean, calculating 21  
Measles, vaccination 284  
Mechanoreceptors 196-197  
Median, calculating 21  
Medical research, and animals 78  
Membrane potential 202  
Membranes, and enzymes 143  
Memory cells 276, 278  
Mesoderm 63  
Metabolic disorders 144-145  
Metabolic efficiency 142  
Metabolic pathways, role of enzyme 142  
Metabolism, defined 150  
Metaphase 64  
Microscopes  
- comparison 56  
- electron 55-56  
- optical 47-48  
Microvilli, role in absorption 118, 121  
Mitochondrion, and cellular respiration 155  
Mitosis, stages 64  
Mode, calculating 21  
Models, types of 4  
Monosynaptic reflex 201  
Motor neuron 199  
Multipotent stem cell 60  
Muscle tissue 70  
Myelination, of neurons 200  
Myoglobin 164
- N**  
Nastic response 288  
Natural killer cells 272  
Negative feedback 194-195, 218  
- and thermoregulation 229-230  
Nephron, structure and function 128-129  
Nerve impulse, transmission 202-203  
Nervous control 198  
Nervous regulation 198
- Nervous tissue 70  
Neuromuscular junction 205  
Neurons 199-200  
Neutrophil 270  
Nitrogenous waste 125  
Nociceptors 196  
Node of Ranvier 200  
Non-infectious disease 253  
Non-self antigens 268  
Normal distribution 21  
Nucleic acids 37  
Nucleotides 37
- O**  
Observations 3  
Optical microscopes 47-48  
Optimal condition, for enzyme activity 137  
Organ transplants 130  
Organelles  
- bacterial cell 39  
- eukaryotic cell 41-46  
- identifying 57  
- prokaryotic cell 39  
Organs 68, 70-71  
Osmoconformers 236  
Osmoregulation 242  
- defined 235  
- strategies 235-244  
- plants 242  
Osmoregulators 236-237  
Osmosis 97-98, 99  
- and red blood cells 100  
Osmotic potential 97  
Oxidative phosphorylation 156  
Oxygen transport 165-166  
Oxygen-haemoglobin dissociation curve 166
- P**  
Pacinian corpuscle 197  
Passive immunity 280  
Passive transport 89-91, 105  
Paster, Louis 254  
Patella reflex 201  
Pathogen spread 293-294  
Pathogen, types 253  
Pearson correlation coefficient 28-29  
Percentage error 16  
Percentages 13  
Peroxidase, activity of 140-141  
pH, and enzyme activity 123, 138, 140  
Phagocytes 273  
Phagocytosis 273  
Phenylalanine metabolism 144  
Phenylketonuria 144-145  
Pheromones 212  
Phloem 177  
- role in translocation 187  
Phospholipids  
- role in membranes 82  
- structure 81-82  
Photoreceptors 196-197  
Photosynthesis 168-174  
- and light 183  
- effect of light 7  
- factors affecting 173-174  
- stages 171-172  
Physical defences, plants 288  
Plan diagrams, rules 53  
Plant cell 35-36  
- and tonicity 99  
- features of 41-42
- organelles 41-42, 45  
Plant systems 175  
Plants  
- adaptations for water loss 242-244  
- defence mechanisms 288-290  
- salt tolerance 247-248  
Plasma cells 276, 278  
Plasma membrane  
- evidence 85-86  
- model 87  
- permeability 81, 83  
- structure 81-87  
Plasmolysis 99  
Pluripotent stem cell 60  
Polio 310  
Potometer 184  
Practical investigation, planning 6-7  
Precision, measurements 10-11  
Predictions 3  
Prion disease 264  
Probability 12  
Probability table 31  
Producers 169  
Productivity, photosynthesis 174  
Prokaryotic cell 35-36  
Prophase 64  
Proprioceptors 196  
Prostaglandins, and inflammation 271-272  
Proteins 37  
- role in membranes 83-84  
Protist cell 35-36  
Protistan disease 259-260  
Pupil reflex 201
- Q**  
Qualitative data 5  
Quantitative data 5  
Quarantine 308
- R**  
Range, calculating 21  
Ranked data 5  
Rates, calculating 12  
Ratios 13  
Receptor 193, 198  
- types 215  
Referencing, how to 2  
Reflex, types 201  
Refractory period 202  
Reliability of the mean 24  
Resolution, in microscopy 47  
Respirometer 153  
Resting potential 202  
Root pressure 180  
Root structure 178  
Roots, uptake at 178  
Rubisco enzyme 171, 174
- S**  
Safety guidelines 8-9  
Salt tolerance, plants 247-248  
Saltatory conduction, in nerves 203  
Sampling bias 26  
Scanning electron microscope (SEM) 55-56  
Scatter graph 17  
Scientific method 2  
Self-antigens 268  
Sensory neuron 199  
Sensory receptors, types 196-197

- Shivering, and thermoregulation 228
- Sieve tubes 177, 187
- Signal molecules 212
- Signal transduction 214-215
- Simple diffusion 89-90
- Skewed distribution 21
- Slide, preparation of 49-50
- Small intestine 117-118
- Small intestine, increasing absorption 118
- Smallpox 310
- Sodium-potassium pump 102
- Spearman rank correlation 30
- Specialisation, of cells 62-63
- Stains, for microscopy 50
- Standard deviation, calculating 23
- Standard error, calculating 24
- Standard form 12
- Starch
- digestion of 120-121
  - testing for 183
- Stargardt's disease 66
- Startle reflex 201
- Statistical test 27-32
- Statistical test
- selection of 27
- Stem cells 62
- ethics of 67, 78
  - types 60-61
  - uses of 65-66
- Stimulus 196, 198
- and homeostasis 193
- Stomach 117
- Stomach emptying, control of 194
- Stomata
- and gas exchange 181-182
  - and transpiration 179
  - role in water loss 242
- Stomatal density, and water loss 245-246
- Student's t Test 31
- Substrate concentration, enzyme activity 137
- Substrate level phosphorylation 156
- Summation, and nerve transmission 206
- Surface area to volume ratio, and diffusion 94-96
- Synapse, chemical 204-205
- Synapse, integration at 206
- Synaptic cleft 204
- Systems 4
- T**
- T-helper cells, and HIV 262
- T-lymphocytes 276
- Telophase 64
- Temperature, and enzyme activity 124, 138
- Thermoreceptors 196-197
- Thermoregulation 195
- and technology 231-232
  - behavioural adaptations 225-226
  - hormonal control 229-230
  - mechanisms 222
  - physiological adaptations 227-228
  - structural adaptations 222-223
- Thyroid gland, and thermoregulation 229
- Tissue fluid 113
- Tissue transplants 130
- Tissue types, human 70-71
- Tissues 68, 70-71
- Tolerance, immunological 267
- Tonicity, and plant cells 99
- Torpor 225
- Totipotent stem cell 60
- Toxins, plant 289
- Translocation 187
- Transmission electron microscope (TEM) 55-56
- Transmission, of disease 293-294
- Transpiration 179-180
- measuring 184-186
- Transpiration pull 180
- Transplants 130
- rejection to 267
- Turgor, plant cell 99
- U**
- Unipotent stem cell 60
- Upregulation 211
- Urea, as waste 125
- Uric acid, as waste 125
- Urinary system, interactions 76-77
- V**
- Vaccination 284
- Covid-19 285-287
  - effects 309-310
  - response 281
- Vaccine development 285
- Vaccine, types 282
- Variables, types 6, 21
- Vascular tissue, plants 175-178
- Vasoconstriction 227
- Vasodilation 227
- Veins 112
- Venous circulatory system 111
- Viral disease 261-263
- Virus 35
- types 261
- Voltage-gated ion channel 202
- W X Y Z**
- Water 37
- Water balance, hormonal control 241
- Whooping cough, vaccination against 284, 287
- Xenotransplantation 78
- Xerophytes 243
- Xylem 175, 180
- Yeast, fermentation 158
- Zika virus 297