



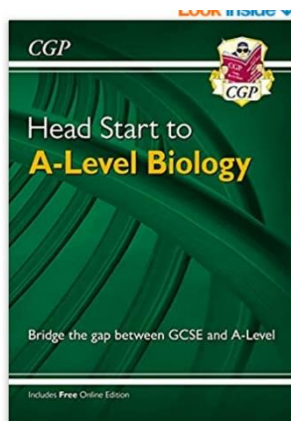
Denbigh School

A Level Biology

GCSE to A Level SUMMER WORK

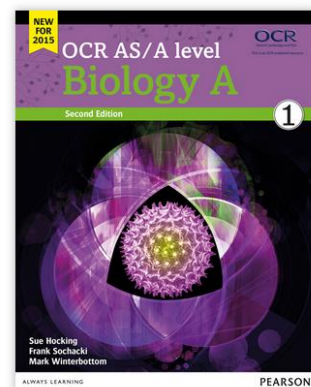
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**Summer work to prepare you for A Level
Biology in Sept'24**

GCSE to A Level Transition – Summer work

The purpose of this work is to prepare you for your A Level Biology course starting in September. You will be reviewing what you know from GCSE and building on this to ensure you are ready to begin a course in A Level Biology.

You will have a Baseline Assessment test at the beginning of the course in September – completing this work is important!

Tasks:

1. Purchase and work through Head Start to A Level Biology

This book will enable you to review key topics covered at GCSE that are built upon at A Level. We expect you to spend time everything week reviewing the contents of this book. All of the other activities will support this.

You will have a baseline assessment during your 2nd lesson in September, this book will enable you to effectively prepare for it.

To do: Work through the book preparing revision flash cards – YOU WILL HAVE A TEST ON ALL THE CONTENT OF THE BOOK!

2. Read through Appendix 1 – Textbook pages (pages 10-21) focussing on investigation skills –

You will **NEED** to purchase a copy of this textbook ready for lessons in September – see details on front cover

Practical and investigational skills form a large part of the course and will be examined at the end of the course, so it is important you are able to demonstrate and apply your knowledge and understanding of these skills. This activity builds on the skills you have already developed.

To do: Complete the following tasks on A4 paper to be shown to your teachers during your very first lesson in September – keep it safe.

- a. Make a table showing the different units and correct abbreviations for length, area, volume and mass.
- b. Find a leaf and produce a biological drawing of it labelling some of its features.
- c. Complete questions 1-3 on page 15
- d. Draw a suitable graph of the data in table 7 on page 15 including range bars
- e. Complete question 1 on page 17
- f. Complete questions 1-6 on page 19

3. Keyword Glossary

Start creating a keyword glossary – you could aim to learn a word a day

Start with these

Activation energy	Glycoprotein
Allele	Hydrolysis (reaction)
Antigen-presenting cell	Induced fit (hypothesis)
ATP	Messenger RNA (mRNA)
Atrioventricular valves	Optimum (temperature / pH)
Base-pairing rules	Oxyhaemoglobin
Benedict's test	Partially permeable membrane
Binomial system	Peptide bond
Buffer	Phospholipid
Carbohydrate	Plasmolysis
Cell signalling	Reducing sugar
Centriole	Semi-conservative replication
Channel protein	Sinoatrial node (SAN)
Chromatin	Stoma (pl: stomata)
Competitive inhibitor	Thylakoid (membrane)
Condensation (reaction)	Transpiration
Crenation	Triglyceride
Denaturation	Turgid
Deoxyribose (sugar)	Ultrastructure
Dipeptide	Vaccine
Endocytosis	Vesicle
Enzyme–substrate complex	Water potential (ψ)
Fluid mosaic (model)	Xerophyte

4. Additional activities – Books, Movies, Research activities, Websites – Appendix 2

This task is designed to further enhance your skills in preparation for A level success. The more you prepare the better you will cope.

If you have any questions, then please email us and we will try to get back to you before the start of the new school year.

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By the end of this topic, you should be able to demonstrate and apply your knowledge and understanding of:

- experimental design, including how to solve problems set in a practical context
- identification of variables that must be controlled, where appropriate
- evaluation that an experimental method is appropriate to meet the expected outcomes

Solving problems in a practical context

In your written examination you may be asked, as part of a particular question, how you could test a prediction or investigate a hypothesis or question. This is an example of solving problems in a practical context.

Although this would be a written paper and you would not have to actually carry out the investigation, you should suggest a procedure that is possible.

- You should:
- be able to state which apparatus, equipment and techniques would be needed for the proposed experiment.
 - apply your scientific knowledge relating to that topic.
 - identify and state the independent and dependent variables and the variables that need to be controlled.
 - evaluate the proposed method to see if it would do the job and provide an answer to the question. It is quite likely that your proposed method would not provide a full answer and that is fine as long as you can recognise this and say so in your evaluation.

An example of a problem

Is the growth of the single-celled green alga *Pleurococcus* affected by its geographical position, e.g. north-facing or south-facing aspect? What factors might influence its distribution?

Applying some biological knowledge to the problem

Many living things are unevenly distributed both between and within ecosystems. Many factors affect their distribution. These may be temperature, habitat, availability of water, minerals, food, space and mates; light intensity, pollution and competition with other organisms for those limited resources.

Pleurococcus is a single-celled, photosynthetic green alga. It looks like green dust and you see it on vertical surfaces such as walls and tree trunks. You may notice that there is often more on the north-facing side of these surfaces or it may be more abundant in shaded and damp areas.

As it is photosynthetic you might expect it to grow more where light intensity is greater. However, it may be damaged by high light intensities or high temperatures, or be susceptible to desiccation, in which case it would grow more in shaded areas. It is living and so will need some water.

Observations have indicated that *Pleurococcus* may have greater abundance and distribution in cooler areas with lower light intensity, i.e. in areas with a north-facing aspect. However, you cannot draw any conclusions unless you carry out some systematic investigations.

Experimental design

Think about the type of data you will be collecting and whether you have a suitable statistical test for analysing that type of data. You would need to sample many trees in different locations. If you tied a piece of string to form a transect, around the tree trunk and then used a compass to find North, you could sample around the trunk, by placing mini **quadrats** (of sides 10 cm) at intervals around the circumference where the string is, and give a score of 0–10 for density of *Pleurococcus* (see topic 4.2.2 for more on using transects and quadrats for sampling plants).

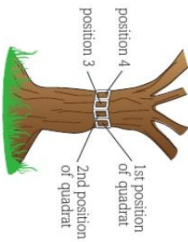


Figure 1 Using a transect and quadrat to sample the densities of *Pleurococcus* around a tree trunk.

Variables

- The independent variable (IV) is the aspect – whether north-, south-, east- or west-facing tree surface.
- The dependent variable (DV) is the density of *Pleurococcus* resulting from the different aspects.

Variables to be controlled

- For example:
- species of tree
 - ecosystem, whether a field or a wood
 - sampling height above ground
 - time of day/same day, so the weather and ambient temperature are the same
 - the same person to assess density, as it is subjective.

What will you do with the data?

You could visually represent the data by constructing a bar chart for each tree, as shown in Figure 2.

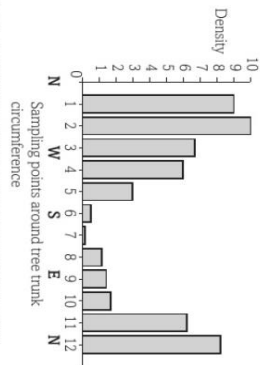


Figure 2 The distribution of *Pleurococcus* around an oak tree in a field, measured at noon during June.

Evaluation of the experimental method

- There are limitations in this design:
- We have only sampled one tree, of one species, in one location.
 - We have not sampled any other vertical surfaces such as walls.
 - We have not used data loggers that can be left for a period of time to monitor the varying conditions.
 - The data have not been analysed statistically to see if the difference between density on the north- and south-facing sides of the trees is significant.
 - Even if we see a correlation between variables, for example light intensity and *Pleurococcus* distribution, correlation between two variables does not necessarily mean that one is causing the other.

Further investigations

Many experimental investigations lead to other questions that need investigating.

The data here show that the distribution of *Pleurococcus* is uneven but this does not solve the problem of what factors may cause this uneven distribution. We can make educated guesses, or hypotheses, as to the causes but we would need to investigate further. These further investigations would also have to be evaluated.

Could it be light intensity? We could use a light meter to measure light intensity at the sampling areas and also look at the data on the bar chart to see if there is any pattern or correlation between light intensity and *Pleurococcus* distribution. Evaluation points. This would have to be done on the same day and at the same time of day, on a cloudy day and on a sunny day, and at the same sampling height.

Could it be temperature? Light heats surfaces so we might expect the temperature to be higher on the south-facing side of the tree trunk. Evaluation points. We could measure the

temperature at each sampling area around the trunk, at the same sampling height, at the same time of day, this could be done for a cloudy day and a sunny day.

Could it be water availability? We could tape test tubes around the tree trunk and leave them to collect rain water that runs off the tree trunk. Evaluation points: Each tube would have to be left in place for the same length of time, at the same sampling height, and on the same days of the year. The tubes would have to be collected at the same time and covered to prevent evaporation, and then the water content measured by mass or volume.

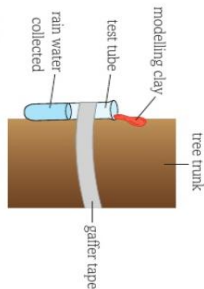


Figure 3 Collecting the water running off a tree trunk.

Could it be predation or infection? Does anything eat *Pleurococcus*? Do any microorganisms infect *Pleurococcus*? We might need to research to find this out and then examine the tree or its location to see if organisms might be infecting or eating *Pleurococcus*.

LEARNING TIPS

If you are stating that light is a possible factor that affects an organism's distribution, refer to the *intensity* of light. Note that italics are used for proper names of living organisms. If you were writing *Pleurococcus* for example in your field note book, by hand, you would underline it. As this is the generic name, it begins with an upper case letter.

DID YOU KNOW?

Pleurococcus is a genus of algae and has been said to be the most abundant organism on the planet. If you use an artist's fine paint brush you can put a little of the green powdery *Pleurococcus* onto a microscope slide and examine it under low and high power. This is a eukaryotic organism, what features of its cell structure can you identify?

Questions

- 1 Suggest a more objective way of assessing the density of *Pleurococcus* on the bark of tree trunks.
- 2 Write a list of equipment you would need to carry out the investigation outlined above on *Pleurococcus* distribution.
- 3 Suggest improvements to this investigation, to reduce its limitations.
- 4 What are the possible sources of errors in this investigation?



Implementing an investigation

By the end of this topic, you should be able to demonstrate and apply your knowledge and understanding of:

- how to use a wide range of practical apparatus and techniques correctly
- appropriate units for measurement
- presenting observations and data in an appropriate format

Using practical apparatus and techniques correctly

Throughout your course, you will carry out several practical investigations, some of which are outlined in this book. Bear in mind that our knowledge and ideas about biology stem from practical investigations that gather data to support hypotheses that then become theories or models.

You will already have carried out practical investigations for GCSE Science and will be familiar with a range of equipment and apparatus and be aware of how to use it safely.

In your written examination you may be asked about the use of apparatus in a practical investigation. Table 1 lists some of the apparatus and techniques that you should use during your course, as well as giving examples of suitable practical activities and areas of the specification that they cover.

Type of practical activity	Skills and techniques	Examples of suitable practical activities	Specification section
Light microscopy	<ul style="list-style-type: none"> • Prepare and stain material for slides • Use microscopes at a range of magnifications • Use a graticule and measure specimens • Produce annotated scientific drawings 	<ul style="list-style-type: none"> • Study structure of plant, animal and prokaryotic cells • Study stages of mitosis • Observe plasmolysis and crenation • Observe a range of tissues 	<ul style="list-style-type: none"> • Cells • Exchange and transport • Homeostasis (A Level only) • Respiration (A Level only) • Photosynthesis (A Level only)
Dissection	<ul style="list-style-type: none"> • Safely use dissecting instruments • Make annotated drawings 	<ul style="list-style-type: none"> • Dissect mammalian heart • Dissect mammalian kidney • Dissect plant stems 	<ul style="list-style-type: none"> • Homeostasis (A Level only) • Exchange and transport
Sampling techniques	<ul style="list-style-type: none"> • Sampling techniques used in fieldwork • Make annotated scientific drawings 	<ul style="list-style-type: none"> • Calculate species diversity 	<ul style="list-style-type: none"> • Biodiversity • Ecosystems (A Level only)
Rates of enzyme-controlled reactions	<ul style="list-style-type: none"> • Use a range of apparatus to record quantitative measurements • Use a range of glassware to make serial dilutions • Use data loggers to collect data or use computer software to process data 	<ul style="list-style-type: none"> • Effects of temperature, pH, substrate and enzyme concentration on rate of enzyme-catalysed reactions 	<ul style="list-style-type: none"> • Enzymes • Homeostasis (A Level only)
Colorimeter or potentiometer	<ul style="list-style-type: none"> • Use colorimeter to record quantitative data • Use potentiometer 	<ul style="list-style-type: none"> • Effect of temperature on membrane permeability • Rate of enzyme-catalysed reaction • Investigate the factors affecting rate of transpiration 	<ul style="list-style-type: none"> • Enzymes • Membranes • Exchange and transport

Table 1 Apparatus and techniques used in A Level Biology.

continued

Type of practical activity	Skills and techniques	Examples of suitable practical activities	Specification section
Chromatography or electrophoresis	<ul style="list-style-type: none"> • Thin layer or paper chromatography to separate biological compounds • Gel electrophoresis 	<ul style="list-style-type: none"> • Analyse chlorophyll • Separate and identify a mixture of amino acids • Separate DNA fragments produced by treatment with restriction enzymes 	<ul style="list-style-type: none"> • Biological molecules • Photosynthesis (A Level only) • Nucleic acids, genetic manipulation
Microbiological techniques	<ul style="list-style-type: none"> • Aseptic techniques • Use of solid and liquid culture media • Colony morphology • Serial dilutions 	<ul style="list-style-type: none"> • The effect of antibiotics on microbial growth 	<ul style="list-style-type: none"> • Cloning and biotechnology (A Level only) • Genetic manipulation (A Level only)
Transport into and out of cells	<ul style="list-style-type: none"> • Serial dilutions • Data logging 	<ul style="list-style-type: none"> • Investigate water potential of plant tissue, such as potato tuber 	<ul style="list-style-type: none"> • Cells • Membranes
Qualitative testing	<ul style="list-style-type: none"> • Use qualitative reagents to identify biological molecules 	<ul style="list-style-type: none"> • Test for biological molecules, such as proteins, lipids, sugars and starch 	<ul style="list-style-type: none"> • Biological molecules
Investigation using a data logger or computer modelling	<ul style="list-style-type: none"> • Use ICT 	<ul style="list-style-type: none"> • Investigate DNA structure using RasMol 	<ul style="list-style-type: none"> • Nucleic acids
Investigate plant and animal responses	<ul style="list-style-type: none"> • Safe and ethical use of organisms to measure plant and animal responses and physiological functions • Use spirometer 	<ul style="list-style-type: none"> • Investigate tropism in plants • Investigate growth requirements of bacteria • Measure human pulse rate at rest and after exercise • Investigate breathing rate and oxygen uptake by human at rest and during exercise • Use <i>Drosophila</i> for genetic investigations 	<ul style="list-style-type: none"> • Plant and animal responses (A Level only) • Exchange and transport
Research skills	<ul style="list-style-type: none"> • Use online sources and books to research topics • Correctly cite sources of information 	<ul style="list-style-type: none"> • Investigate respiration in yeast <i>Saccharomyces cerevisiae</i> 	<ul style="list-style-type: none"> • All topic areas

Table 1 Apparatus and techniques used in A Level Biology (continued). Note that the types of practical activity listed are organised according to the practical activity groups (PAGs) referred to in the specification.

Appropriate units for measurement

In many practical investigations you are likely to be measuring something. It is important that you use the correct units and the correct symbols or abbreviations.

Below are some of the units you may use, with their correct symbols, e.g. kilograms (kg), metres (m), seconds (s), joules (J) or kilopascals (kPa) for energy, kilopascals (kPa) for pressure or water potential. However, the actual unit used depends on what you are measuring. If you are measuring the diameter of a cell, micrometres (μm) would be appropriate, but if measuring the height of a tree, metres would be a more appropriate unit. For certain studies involving energy flow through ecosystems, the units might be gigajoules per hectare per year ($\text{GJ ha}^{-1} \text{yr}^{-1}$).

Prefix	Order of magnitude
nano-	10^{-9}
micro-	10^{-6}
milli-	10^{-3}
centi-	10^{-2}
kilo-	10^3
mega-	10^6
giga-	10^9
tera-	10^{12}
peta-	10^{15}

Table 2 Prefixes denoting orders of magnitude.

DID YOU KNOW?

A googol is the name given to a number of order magnitude 10^{100} , which is 10 with 100 zeros after it. And 10^{999} is called a googolplex. Both these names were invented by a nine-year-old child, the son of a mathematician.

Unit	Abbreviation	Number of metres
kilometre	km	1000
metre	m	1
centimetre	cm	0.01
millimetre	mm	0.001
micrometre	μm	0.000 001
nanometre	nm	0.000 000 001

Table 3 Units for length. SI base unit = metre.

Unit	Abbreviation	Number of square metres
kilometres squared	km^2	1 000 000
hectare	ha	10 000
centimetres squared	cm^2	0.0001
millimetres squared	mm^2	0.000 001

Table 4 Units for area.

Unit	Abbreviation	Number of centimetres cubed
cubic decimetres	dm^3	1000
cubic centimetres – also called millilitres	cm^3 or ml	1
cubic millimetres – also called microlitres	mm^3 or μl	0.001

Table 5 Units for volume.

Unit	Abbreviation	Number of grams
metric tonne	t	1 000 000
kilogram	kg	1000
gram	g	1
milligram	mg	0.001
microgram	μg	0.000 001

Table 6 Units for mass.

LEARNING TIP
Always state the units you are using when describing a quantity.

Presenting your observations and data

If you have been observing a structure, such as an organ or organ system via dissection, a labelled drawing is the way to present this. When you study transport in animals (Chapter 3.2) you will have the opportunity to dissect a mammalian heart and make annotated drawings of your observations. A labelled drawing is also the way to present observations of cells or tissues on a microscope slide. In Chapter 2.1 you will have several opportunities to make such annotated drawings from microscope slides, in the correct way.

Besides drawings, figures, graphs and diagrams are also visual representations of observations and results of investigations. Topics 1.1.3 and 1.1.4 deal with different types of graphs and diagrams.

Tables

Often the best way to present initial data from an investigation is in a table – see Table 7 for an example:

- The table must have a clear title to inform the reader.
- The table should be ruled off.
- The independent variable should be in the first column (to the left side of the table).
- Each column should have an informative heading and the units for the quantities shown should be in the column heading, not in the column itself.

- You can tabulate data that are not quantitative, such as colour of reagents used in tests and the inference (what it tells you).
- If the data are quantitative, the same number of decimal places should be used for all the values in one particular column.
- If replicates have been carried out there should be a column for each and a column for the calculated mean values.
- The mean values should be calculated to the same number of decimal places or to one more decimal place than those of the raw data values, but all the mean values in a column must be to the same number of decimal places.

Temperature ($^{\circ}\text{C}$)	Rate of hydrolysis of starch (mg s^{-1})			Mean rate of hydrolysis of starch (mg s^{-1})
	1	2	3	
10	11.54	11.36	11.43	11.44
20	21.90	21.59	22.01	21.83
30	35.30	36.00	35.85	35.72
40	36.54	37.01	36.97	36.84

Table 7 Rates of digestion of starch by the enzyme amylase, obtained from goat saliva, at different temperatures.**Questions**

A student investigated the digestion of triglyceride (fat) by the enzyme lipase. He wanted to investigate the effect of increasing temperature on the rate of reaction. The enzyme-catalysed reaction produces fatty acids and these lower the pH. This change in pH can be detected by an indicator, such as bromothymol blue, which is blue at pH 7.6, green at pH 7.0 and yellow at pH 6.0. The time taken for the indicator to change to yellow can be measured and so the rate of digestion can be determined. The student presented his data in a table as shown below.

Time taken for indicator to become yellow (secs)	Temperature		
	1	2	3
454	476	468	10 $^{\circ}\text{C}$
287	295	305	15 $^{\circ}\text{C}$
210	208	212	20 $^{\circ}\text{C}$
121	123	126	25 $^{\circ}\text{C}$
105	110	109	30 $^{\circ}\text{C}$
68	63.5	65.5	35 $^{\circ}\text{C}$

- State six ways in which this table can be improved.
- Calculate the mean rates of reaction for these data. Calculate rate as 1000 divided by time taken for indicator to become yellow. (We use 1000/7 rather than 1/7 to calculate the rate, so that the numbers in the calculation are more user friendly. As long as all values are treated in this way, the relative rate of reaction is the same. In effect $1/7 \times 10^3$.)
- Present these data in a properly constructed table.
- Comment on the range of temperatures used in this investigation.
- What are the limitations of this investigation in terms of determining the end point of the indicator?
- Suggest how this investigation could be improved and include suggestions for other ways of measuring the fall in pH.

Analysis of data 1: Qualitative and quantitative data

By the end of this topic, you should be able to demonstrate and apply your knowledge and understanding of:

- processing, analysing and interpreting qualitative and quantitative experimental results
- use of appropriate mathematical skills for analysis of quantitative data
- appropriate use of significant figures

KEY DEFINITIONS
qualitative data: data that does not involve quantity (numbers),
quantitative data: data that does involve quantity (numbers),
significant figures: the digits of a number that have a meaning and contribute to the number's precision.

Processing, analysing and interpreting results

When you carry out tests to indicate the presence of glucose, starch, lipids or proteins (see topic 2.2.12 for more about these food tests), you will obtain **qualitative data**. You can represent such findings in a table and indicate the colour observed and the inference – this tells us whether a substance is present or not.

Benedict's reagent is used to test for reducing sugar (if positive, reagent changes from blue to red when heated); iodine/KI solution tests for starch (if positive, a blue-black colour is seen); ethanol emulsion test indicates the presence of lipids if a white emulsion is seen; biuret reagent indicates the presence of protein by a purple/mauve colour.

To make your data **quantitative**, for example to see how much glucose is in a particular drink, you would need to make up a range of glucose solutions of known concentrations, using serial dilution. You would then carry out a Benedict's test, keeping certain variables constant, such as:

- volume of reagent
- volume of solution being tested
- temperature at which heated
- length of time for heating.

Food tested	Colour/observation at end of test		Inference
	Benedict's reagent at 80 °C for 10 min	Iodine/KI solution	
bread	blue	black	contains starch and protein
potato	blue	black	contains starch and protein
apple	red	brown	contains reducing sugar and protein
cheese	blue	brown	contains lipid and protein
chicken	blue	white	contains lipid and protein

Table 1 Results of tests carried out on a variety of foods.

Using mathematical skills to analyse quantitative data

Think about the measurement of water uptake by a potometer, as described in Chapter 3.3. If measurements are taken at different ambient temperatures we can see the effect of temperature on the rate of water uptake and therefore on the rate of transpiration.

Ambient temperature (°C)	Distance travelled by air bubble in 10 min (mm)			Rate of uptake of water ($\mu\text{l s}^{-1}$)
	1	2	3	
10	12.5	13.0	13.5	13.0
20	28.0	27.5	27.3	27.6
30	45.0	47.0	46.0	46.0
40	55.5	56.5	55.7	55.9

Table 2 Mean rate of water uptake ($\mu\text{l s}^{-1}$) in a leafy stemware maize, *Acer pseudoplatanus*, shoot in a potometer, at different ambient temperatures.

Calculating the volume of water taken up

If you are told, for example, that the diameter of the bore of the capillary tube is 2.5 mm and the air bubble travelled 24 mm in 10 minutes, you can calculate the rate of uptake of water in $\mu\text{l per minute}$ or per second.

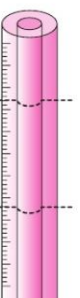


Figure 1 Calculating the volume of water in a section of capillary tube.

If the diameter is 2.5 mm then the radius is 1.25 mm.

L indicates the length moved by the air bubble, so the space in this cylinder is the same as the volume of water taken up by the shoot.

The formula for calculating the volume of a cylinder, V , is $V = \pi r^2 L$.

So the volume of water taken up by the shoot in 10 minutes is $[3.142 \times (1.25)^2 \times 24] \text{ mm}^3$
 $= 117.825$
 $= 118 \mu\text{l}$

LEARNING TIP

Notice that the number in this calculated example has been rounded to a whole number. This is because you can only read this scale to one decimal place. You could express the answer as 117.8 μl , but to no more than one decimal place. mm^3 is not incorrect as a unit but μl is more often used.

Now to calculate rate of uptake, which is volume taken up per unit time.

If 118 μl is taken up in 10 minutes, then the rate of uptake is $118/10 = 11.8 \mu\text{l min}^{-1}$.

You could also express this in terms of volume taken up per second, which would be $118/600 = 0.20 \mu\text{l s}^{-1}$.

Calculating a median value

Suppose you measure the lengths of the leaves on a branch of a shrub. Their measurements in mm are:

62, 65, 75, 83, 55, 78, 77, 68, 57, 58, 54, 66, 72, 80, 48, 71, 72, 62, 49, 81.

The **arithmetic mean** is 66.7 mm.

The range is from 48 to 83 mm.

There are 10 numbers from 48 to 66 and 10 numbers from 68 to 83. The **median** is therefore 67 (between 66 and 68). This is correct even though there are no leaves of 67 mm in the sample.

Appropriate use of significant figures

In some cases we do not need a detailed answer or very precise number. When you work out an answer on your calculator you do not need to express it to 10 decimal places so you round it off to a certain number of decimal places.

Another method is to round it off using **significant (meaningful) figures**.

From the column in Table 2 showing the rate of transpiration, in the second row where the rate is 0.23, 2 is the most significant digit because it tells you that the rate is about $0.2 \mu\text{l s}^{-1}$. The second number, 3, is the next significant figure. It tells us that the rate is faster than $0.2 \mu\text{l s}^{-1}$. This therefore gives a more accurate and precise indication of the value of the rate calculated. Because this is a calculated value, it can be expressed to one more decimal place than the values in the other columns that were obtained by reading the apparatus and were therefore limited by the precision of the apparatus. The calculated values in this column in Table 2 are all to two significant figures.

As a general rule, the calculated values, in order to be significant, can be to one more decimal place than the values in the columns from which the calculation was made.

The following are not significant figures: leading zeros, trailing zeros and digits derived by calculation and giving several decimal places, which therefore give *far* greater precision than the original data or the instrument used for measurement.

Questions

- Express the following to two significant figures:
 - 5.374641
 - 1.6457836
 - 0.9853421
 - 150
 - 0.6780000
- In an investigation using a potometer, the bubble of air moved 65 mm along the capillary tube in 15 minutes. The diameter of the bore of the capillary tube was 2 mm. Calculate the rate of water uptake by the plant in $\text{mm}^3 \text{s}^{-1}$ ($\mu\text{l s}^{-1}$).
- Suggest how you could adapt the use of the biuret test for protein to make it quantitative.

By the end of this topic, you should be able to demonstrate and apply your knowledge and understanding of:

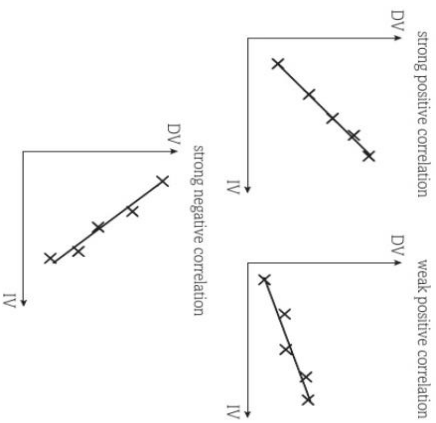
- plotting and interpreting suitable graphs from experimental results

There is a variety of graphs and each type has specific uses, but each communicates information visually. In a written examination you may be given a table of data and be asked to graph those data. You may also be asked to:

- make deductions from graphical data
- draw conclusions from graphical data
- evaluate the data or its presentation (see next topic).

Line graphs

Line graphs are used to see if there is any correlation between two variables where the data are continuous.



- Figure 1** Examples of correlation between two variables.
- They involve a vertical y-axis and a horizontal x-axis forming a grid.
 - Each axis should have a suitable linear scale and be labelled with quantities and units.
 - The independent variable (IV) is usually plotted along the x-axis and the dependent variable (DV) along the y-axis.

More than one curve can be drawn on the same set of axes, so comparisons can be made and a picture of what is happening during an investigation or observed phenomenon can be seen.

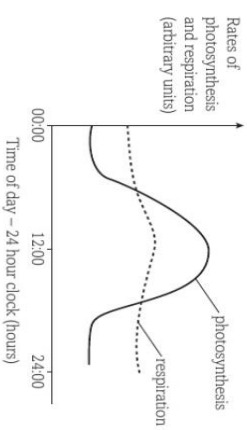


Figure 4 Graph showing the changes in rates of photosynthesis and respiration in a small pond over a 24-hour period during May.

- The rate of reaction can be calculated from the slope of a curve showing the progress of the reaction over time.

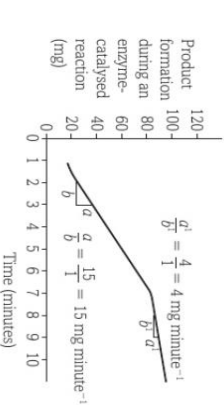


Figure 5 Calculating the rate of an enzyme-catalysed reaction from the slope of a graph

Scattergrams

Also called scatter diagrams or scatter plots, scattergrams are used when investigating the relationship between two naturally changing variables. For example, several plots can be made showing mean blood cholesterol level and death rates from heart disease and stroke in various countries. No line needs to be drawn, but the pattern of the plots can show if there is any correlation.

Bar graphs

Bar graphs are used to investigate relationships when the independent variable is categorical and the dependent variable is continuous, e.g. the concentration of Vitamin C (DV) in different fruit drinks (IV).

- The bars should be of the same width and equally spaced.
- If mean values are shown on the bars, the range bars can also be shown.
- If the data sets being compared have been analysed statistically, the error bars can be shown. If there is overlap it indicates that any apparent difference is not significant.

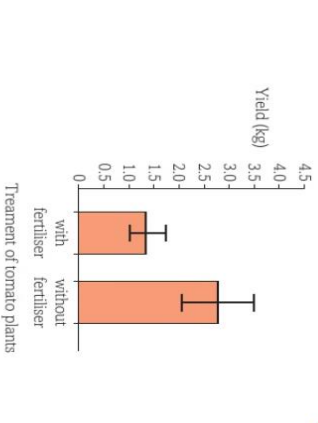


Figure 6 Comparison of yield of tomatoes grown with and without fertiliser. Error bars do not overlap, showing that the difference between these two data sets is significant.

Histograms

Histograms can be used for showing quantitative data organised into classes. For example, if we measured the height of a large number of human adults we may categorise the data, for example those between 140 and 149 cm and those between 150 and 159 cm. The number of people within each class shows the frequency. The class or category that contains the greatest frequency is the **mode**.

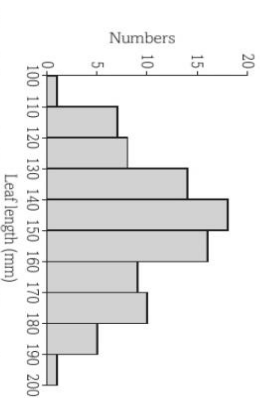


Figure 7 Histogram showing frequency of leaf length in sweet chestnut.

Questions

Which type of graph would you draw to display each of the following types of data?

- Lengths of leaves on a tree branch.
- Effect of changing pH on enzyme activity.
- Sugar content of different types of biscuits.
- Effect of light intensity on rate of photosynthesis.
- Collagen content of skin and age in humans.
- Amino acid content of beef and cheese.

By the end of this topic, you should be able to demonstrate and apply your knowledge and understanding of:

- how to evaluate results and draw conclusions
- the identification of anomalies in experimental measurements
- Limitations in experimental procedures
- the refining of experimental design by suggestion of improvements to the procedures and apparatus
- precision and accuracy of measurements and data, including margins of error, percentage errors and uncertainties in apparatus

What can we conclude from these data?

- It would appear that statins are more effective at lowering blood cholesterol level than a patient making lifestyle changes.
- Lifestyle changes appear to lower blood cholesterol, although the improvement was not as marked as in the group taking statins. However, the SD of this group was greater after treatment so some may not have shown any improvement.

KEY DEFINITIONS

accuracy: how close a measured or calculated value is to the true value.

anomaly: result that does not fit the expected trend or pattern.

precision: the closeness of agreement between measured values obtained by repeated measurements.

Evaluating results and drawing conclusions

You may be shown data and asked to evaluate them or to comment on a conclusion drawn from the data.

For example, in Table 1 are data about changes in blood cholesterol levels. One group of patients was given cholesterol-lowering drugs, called statins. Another group of patients within the same GP practice decided to try to lower their blood cholesterol levels by taking more exercise and altering their diet.

Patient group	Mean blood cholesterol level (mmol dm ⁻³) (±SD)	
	Before treatment	6 months after treatment began
Group A – treated with statins (n = 12)	6.36 (±1.58)	4.21 (±0.19)
Group B – treated with lifestyle change (n = 12)	5.95 (±1.34)	4.87 (±1.60)

Table 1 Blood cholesterol levels of groups of patients in a GP practice.

Statins inhibit an enzyme in the liver from making cholesterol. The guidelines set by NICE (National Institute for Health and Care Excellence) in 2008 stated that people should have a blood cholesterol level of 5.2 mmol dm⁻³ or less. In 2014 the guidelines were changed to 4.0 mmol dm⁻³.

Identifying anomalies in data

You have been trained to identify **anomalies** in data. These are results that do not fit the expected pattern. Seeing an anomaly can be an exciting moment, providing evidence that your expectation is wrong and a scientific breakthrough could be staring you in the face. On the other hand it could be due to a piece of grit in your detector or a leaky flask in your incubator. If you are certain that an anomalous piece of data was produced due to a failure in the experimental procedure, you might be justified in removing it before analysing the data. However, you must never discard data simply because they do not correspond with your expectation. By repeating the experiment and amassing more data one of two things could happen. If the anomaly was the result of an experimental error or was simply a very unusual result from naturally-occurring variation it will disappear as the repeat measurements produce a mean in line with expectation. On the other hand, if the anomaly was in fact telling you something surprising about the system you are investigating it will be confirmed by repeat observations and your Nobel Prize is just around the corner.

Limitations in experimental procedures

- It is not always possible to control all extraneous variables.
- Some investigations would be unethical, such as deliberately damaging an area of children's brains to study the effects on their development.
- Results obtained from studying a small population cannot be generalised to the whole population.
- The resolution of the instruments and equipment used may impose limitations.
- The degree of **accuracy** of measurements may lead to limitations.

- Using a small sample size or having too few replicates is also a limitation, as it is difficult to see if the data are reliable; therefore a large enough sample or enough replicates should be used where possible.
- Not leaving a reaction for long enough to fully complete will give misleading data; therefore we should make sure that reactions are given long enough to complete.
- Not allowing reactants to reach the required temperature before adding them together will reduce validity; reactants should be placed, in their tubes, into a water bath to reach the required temperature before they are mixed.
- Some investigations that rely on questioning people or observing them in particular situations may be limited, because only certain types of people will volunteer to take part or people will behave differently when they think they are being observed.

- Lack of equipment to objectively measure something, such as a colour change, is a limitation as the observation is subjective and may change depending on the investigator.
- Limitations in equipment such as using a beaker of hot water for a waterbath; the investigation can be improved by using a thermostatically controlled water bath, with a thermometer to check the temperature, so as to maintain the desired temperature throughout the reaction.

Errors

Errors or experimental uncertainties arise because there are:

- inadequacies and imperfections in experimental procedures
- lapses of judgement by the experimenter
- limits to resolution, **precision** or accuracy of measuring apparatus.

Random errors due to judgement errors made by the experimenter are reduced when the procedure is repeated several times.

Systematic errors may be inherent in the equipment and are repeated at every replicate. However, if the percentage error is known, a calculation can be done to determine the margin of error.

LEARNING TIP

Be clear about the difference between **accuracy** and **precision**. A thermometer is inaccurate if it gives readings that are 5 °C above the true temperature but it could still be precise if it gives very consistent readings. By recalibrating an inaccurate instrument you can correct this and make accurate measurements. Still confused? An analogy might help: A precise archer will have her arrows tightly clustered somewhere on the target. By recalibrating her sight she can become accurate and precise and will have her arrows clustered on the bulls-eye!

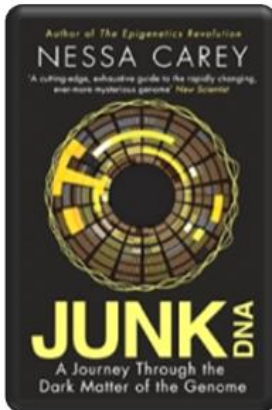
Questions

- 1 A digital stopwatch can measure to the nearest 0.1 s. Explain why using this stopwatch to measure a reaction for 5 minutes is more accurate than using it to measure the reaction times of humans, which are around 0.3 s duration.
- 2 In school laboratories, thermometers filled with alcohol rather than mercury are used for safety reasons. They are precise and have an impressive resolution of 0.2 °C. However, the overall calibration could be up to 1 °C out. If you used one of these thermometers to measure the temperature of a water bath at 38 °C, within what range would the real temperature be?
- 3 Explain why using a gas syringe to collect oxygen given off from a well-illuminated aquatic plant for 5 minutes, is better than counting the bubbles of oxygen produced during 5 minutes.

Appendix 2 – Additional activities

Book Recommendations

Kick back this summer with a good read. The books below are all popular science books and great for extending your understanding of Biology

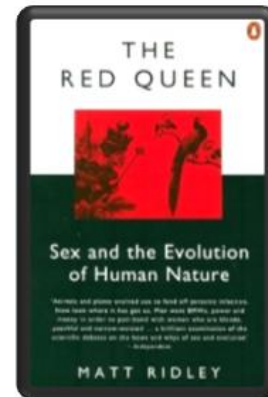


Junk DNA

Our DNA is so much more complex than you probably realize, this book will really deepen your understanding of all the work you will do on Genetics. Available at [amazon.co.uk](https://www.amazon.co.uk)

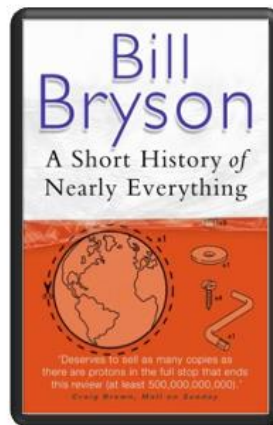
The Red Queen

Its all about sex. Or sexual selection at least. This book will really help your understanding of evolution and particularly the fascinating role of sex in evolution. Available at [amazon.co.uk](https://www.amazon.co.uk)

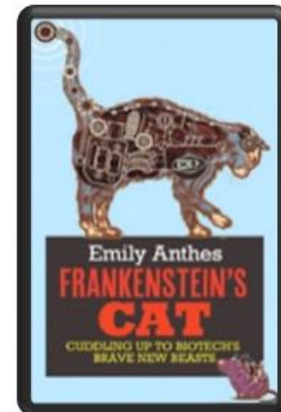
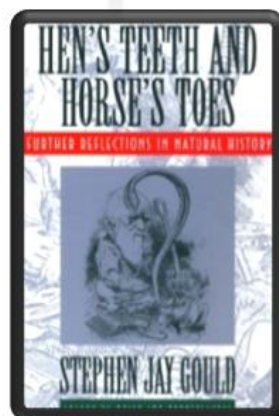


A Short History of Nearly Everything

A whistle-stop tour through many aspects of history from the Big Bang to now. This is a really accessible read that will re-familiarise you with common concepts and introduce you to some of the more colourful characters from the history of science! Available at [amazon.co.uk](https://www.amazon.co.uk)



Studying Geography as well? **Hen's teeth and horses toes** Stephen Jay Gould is a great Evolution writer and this book discusses lots of fascinating stories about Geology and evolution. Available at [amazon.co.uk](https://www.amazon.co.uk)



An easy read..

Frankenstein's cat

Discover how glow in the dark fish are made and more great Biotechnology breakthroughs. Available at [amazon.co.uk](https://www.amazon.co.uk)

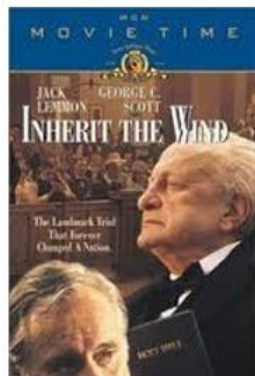
Movie Recommendations

Everyone loves a good story and everyone loves some great science. Here are some of the picks of the best films based on real life scientists and discoveries. You won't find Jurassic Park on this list, we've looked back over the last 50 years to give you our top 5 films you might not have seen before. Great watching for a rainy day.



Inherit The Wind (1960)

Great if you can find it. Based on a real life trial of a teacher accused of the crime of teaching Darwinian evolution in school in America. Does the debate rumble on today?



Gorillas in the Mist (1988)

An absolute classic that retells the true story of the life and work of Dian Fossey and her work studying and protecting mountain gorillas from poachers and habitat loss. A tear jerker.

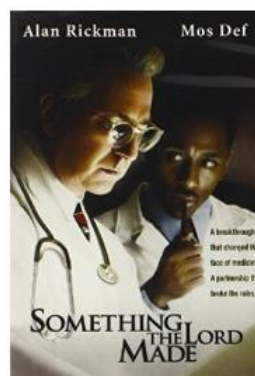
Andromeda Strain (1971)

Science fiction by the great thriller writer Michael Crichton (he of Jurassic Park fame). Humans begin dying when an alien microbe arrives on Earth.



Lorenzo's Oil (1992)

Based on a true story. A young child suffers from an autoimmune disease. The parents research and challenge doctors to develop a new cure for his disease.



Something the Lord Made (2004)

Professor Snape (the late great Alan Rickman) in a very different role. The film tells the story of the scientists at the cutting edge of early heart surgery as well as issues surrounding racism at the time.

There are some great TV series and box sets available too, you might want to check out: Blue Planet, Planet Earth, The Ascent of Man, Catastrophe, Frozen Planet, Life Story, The Hunt and Monsoon.

Movie Recommendations

If you have 30 minutes to spare, here are some great presentations (and free!) from world leading scientists and researchers on a variety of topics. They provide some interesting answers and ask some thought-provoking questions. Use the link or scan the QR code to view:

A New Superweapon in the Fight Against Cancer

Available at :

http://www.ted.com/talks/paula_hammond_a_new_superweapon_in_the_fight_against_cancer?language=en

Cancer is a very clever, adaptable disease. To defeat it, says medical researcher and educator Paula Hammond, we need a new and powerful mode of attack.



Why Bees are Disappearing

Available at :

http://www.ted.com/talks/marla_spivak_why_bees_are_disappearing?language=en

Honeybees have thrived for 50 million years, each colony 40 to 50,000 individuals coordinated in amazing harmony. So why, seven years ago, did colonies start dying en-masse?

Why Doctors Don't Know About the Drugs They Prescribe

Available at :

http://www.ted.com/talks/ben_goldacre_what_doctors_don_t_know_about_the_drugs_they_prescribe?language=en

When a new drug gets tested, the results of the trials should be published for the rest of the medical world — except much of the time, negative or inconclusive findings go unreported, leaving doctors and researchers in the dark.



Growing New Organs

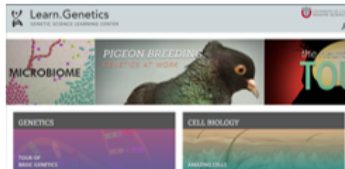
Available at :

http://www.ted.com/talks/anthony_atalla_growing_organs_engineering_tissue?language=en

Anthony Atalla's state-of-the-art lab grows human organs — from muscles to blood vessels to bladders, and more.

Science websites

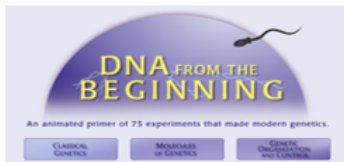
These websites all offer an amazing collection of resources that you should use again and again through out your course.



Probably the best website on Biology....

Learn Genetics from Utah University has so much that is pitched at an appropriate level for you and has lots of interactive resources to explore, everything from why some people can taste bitter berries to how we clone mice or make glow in the dark jelly fish.

<http://learn.genetics.utah.edu/>



DNA from the beginning is full of interactive animations that tell the story of DNA from its discovery through to advanced year 13 concepts.

One to book mark!
<http://www.dnaftb.org/>



In the summer you will most likely start to learn about Biodiversity and Evolution. Many Zoos have great websites, especially London Zoo. Read about some of the case studies on conservation, such as the Giant Pangolin, the only mammal with scales.
<https://www.zsl.org/conservation>



At GCSE you learnt how genetic diseases are inherited. In this virtual fly lab you get to breed fruit flies to investigate how different features are passed on.

<http://sciencecourseware.org/vcise/drosophila/>



Ok, so not a website, but a video you definitely want to watch. One of the first topics you will learn about is the amazing structure of the cell. This BBC film shows the fascinating workings of a cell... a touch more detailed than the "fried egg" model you might have seen.

http://www.dailymotion.com/video/xh0kb_the-hidden-life-of-the-cell_shortfilms

If this link expires – google "BBC hidden life of the cell"