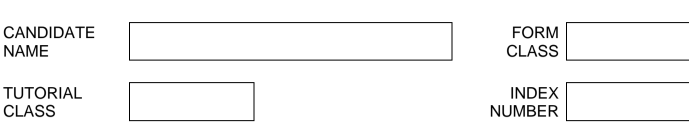
# **Anglo-Chinese Junior College**

JC2 Biology Preliminary Examination Higher 2



## BIOLOGY

Paper 4 Practical

Candidates answer on the Question Paper. Additional Materials: As listed in the Confidential Instructions.

### READ THESE INSTRUCTIONS FIRST

Write your Name, Class and Index number in the spaces on all the work you hand in. Give details of the practical shift and laboratory, where appropriate in the boxes provided. Write in dark blue or black pen. You may use an HB pencil for any diagrams or graphs.

You may use an HB pencil for any diagrams or graphs.

Do not use staples, paper clips, glue or correction fluid.

Answer **all** questions in the spaces provided on the Question Paper.

The use of an approved scientific calculator is expected, where appropriate. You may lose marks if you do not show your working or if you do not use appropriate units.

At the end of the examination, fasten all your work securely together. The number of marks is given in brackets [] at the end of each question or part question.

For Examiners' use		
1	/ 12	
2	/ 23	
3	/ 20	
Total	/ 55	



Shift
Laboratory

**9744/04** 06 August 2024 2 hours 30 minutes

#### Answer **all** questions.

1 In some countries, certain plants have seasonal growth. During the summer plants transport sucrose from the leaves to store it as starch in the roots.

Table 1.1 summarises the changes in the leaves and roots during the year.

#### Table 1.1

season	leaves	roots
summer	leaves synthesise sucrose	sucrose stored as starch
winter no leaves		starch stored
spring	leaves growing	starch converted to glucose

You are required to identify the source of four plant extracts. These have been taken from

- a root in winter
- a root in spring
- phloem sap from the stem in summer
- phloem sap from the stem in winter
- (a) Use the information in Table 1.1 to predict which substances you would expect to be present in each of the four plant extracts, then complete Table 1.2.

Key:  $\checkmark$  (tick) substance present in plant extract

X (cross) substance absent from plant extract

source of plant extract	substances present in each of the plant extracts		
Source of plant extract	starch	sucrose	glucose
root in winter			
root in spring			
phloem sap from the stem in summer			
phloem sap from the stem in winter			[0]

[2]

One sample was taken from each of the four plant extracts shown in Table 1.2.

You are required to identify from which plant extract each of the four samples **S1**, **S2**, **S3** and **S4** was taken.

You are provided with:

- Benedict's solution, in a container labelled **B**
- iodine solution, in a container labelled I
- hydrochloric acid, in a container labelled H
- sodium hydrogencarbonate powder, in a container labelled S
- a source of heated water.

Hydrochloric acid H and sodium hydrogencarbonate S are irritants, Benedict's solution B is harmful and iodine solution I is a stain. Suitable eye protection should be worn. If any of these reagents come into contact with your skin, wash off immediately under cold water.

#### Do not carry out any tests until you have read the instructions on pages 3 to 5.

(b) Describe the two tests that show that starch and glucose are present in a plant extract.

test for starch: [1] test for glucose: [2] For the test for sucrose, refer to steps 1 to 9.

- 1 Before starting the test for sucrose, perform the test for glucose you have described in (b).
- (c) Explain why the test for glucose must be conducted before the test for sucrose can be carried out.

[2]

- **2** To start the test for sucrose, put  $2.0 \text{ cm}^3$  of a fresh sample into a test-tube.
- **3** Put 2.0 cm<sup>3</sup> of **H** into the same test-tube.
- 4 Shake the test-tube gently to mix the contents.
- 5 Put the test-tube in a boiling water-bath. Leave the test-tube for 2 minutes.
- 6 After 2 minutes, remove the test-tube from the water-bath and put it in a test-tube rack.
- 7 Leave the test-tube to cool for a further 3 minutes.
- 8 After 3 minutes, put a small amount of **S** into the test-tube. The mixture will fizz and rise up the test-tube. Continue to add small amounts of **S** until there is no more fizzing.

Note: there may be a little of S left in the bottom of the test-tube. This will not affect the results.

9 Perform the test for glucose you have described in (b) on the solution from step 8.

#### Proceed as follows.

Use the beaker labelled **hot water** to collect approximately 400 cm<sup>3</sup> of hot water from where it is provided in the laboratory. Heat the water to boiling, if needed.

Select the appropriate reagents from those provided and carry out suitable tests to identify the samples **S1**, **S2**, **S3** and **S4**.

(d) Record your observations in an appropriate table. You do **not** need to make conclusions on the presence or absence of each substance in the samples tested.

[4]

(e) Complete Table 1.3 to match the samples, S1, S2, S3 and S4, with each plant extract.

Table 1	1.3
---------	-----

source of plant extract	sample
a root in winter	
a root in spring	
phloem sap from the stem in summer	
phloem sap from the stem in winter	

[1]

[Total: 12]

[Turn over

2 Yeast contains an enzyme that will break down hydrogen peroxide into oxygen and water. The loss of mass resulting from the release of oxygen can be measured.

You will investigate the effects of the concentration of hydrogen peroxide on the rate of enzymatic activity.

You are provided with:

- Yeast cell suspension, in a container labelled Y
- 1.0% hydrogen peroxide solution, in a container labelled P
- Distilled water, in a container labelled W

Hydrogen peroxide solution P is an irritant and is harmful. Suitable eye protection should be worn. If P comes into contact with your skin, wash them off immediately under cold water.

Each time that you take a sample of yeast cell suspension Y, you should make sure that it is mixed thoroughly by stirring it with a glass rod. You should also collect Y from below the surface, so as to minimise the volume of froth collected.

(a) (i) You will carry out proportional dilutions of the 1.0% hydrogen peroxide solution P to obtain a range of concentrations in which the concentration of hydrogen peroxide is reduced by 0.2% between each successive dilution.

You will prepare 10.0 cm<sup>3</sup> of each concentration, using **P** and **W**.

Using a table in the space below, show how you will prepare the different concentrations of hydrogen peroxide solution. One of the concentrations should include 1.0% hydrogen peroxide.

[3]

#### 7

#### Read steps 1–6 before starting the investigation. Proceed as follows.

- 1 Prepare the concentrations of hydrogen peroxide solution according to the table in **(a)(i)**, in the vials provided. Label the vials where appropriate.
- 2 Use the electronic mass balance to weigh the mass of the reaction mixture in subsequent steps.
- **3** Put 5.0 cm<sup>3</sup> of yeast suspension **Y** into the vial containing 1.0% hydrogen peroxide solution. Start timing immediately and proceed to step **4** without delay.
- **4** Weigh the mass of the reaction mixture, including the weight of the vial, using the set-up shown in Fig. 2.1.

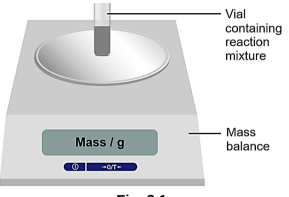


Fig. 2.1

- 5 After five minutes, weigh the mass of the reaction mixture in the vial again.
- 6 Repeat steps **3 5** with each of the other concentrations of hydrogen peroxide that you prepared in step **1**. You do **not** need to wait to complete the measurement for one concentration of hydrogen peroxide before starting on the next concentration.
  - (ii) Record your results in Table 2.1. Calculate the change in mass and the percentage change in mass. No workings are required.

#### Table 2.1

concentration of hydrogen peroxide / %	initial mass of reaction mixture / g	final mass of reaction mixture / g	change in mass / g	percentage change in mass
1.0				

[4]

[Turn over] H2 9744 Paper 4 Preliminary Examination

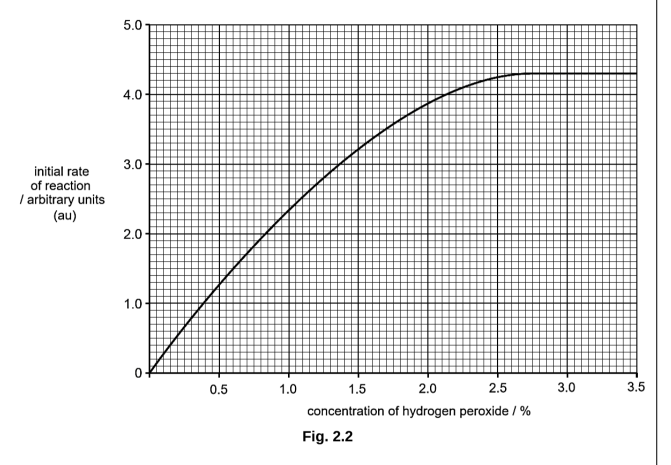
(iii)	Suggest a suitable control for this experiment to show that it is an enzyme that catalyses the break down of hydrogen peroxide.
	[1]
(iv)	Other than the lack of a suitable control, describe <b>two</b> modifications to this method that would increase the confidence in your results.
	[2]

(b) A student wanted to determine the Michaelis-Menten constant (K<sub>m</sub>) for the enzyme-catalysed break down of hydrogen peroxide by the enzyme found in yeast, enzyme Y.

 $K_m$  is the substrate concentration at which the reaction rate is 50% of the maximum rate of reaction ( $V_{max}$ ).

 $K_m$  gives an indication of the affinity an enzyme has for its substrate.

The student measured the initial rate of reaction at different concentrations of hydrogen peroxide. The results are shown in Fig. 2.2.



(i) Use the graph in Fig. 2.2 to estimate the Michaelis-Menten constant (K<sub>m</sub>).

Show your working on the graph and in the space below.

(ii) The  $K_m$  value for another enzyme, **Z**, is 2.5%.

State which enzyme, **Z** or **Y**, has a **lower affinity** for its substrate.

Give a reason for your answer.



......[2]

(c) Living, respiring yeast cells also contain enzymes which reduce methylene blue, a blue dye commonly used for the staining of biological samples, turning it colourless. In non-respiring cells, the reduction of methylene blue does not occur.

It is thought that respiration in yeast cells is inhibited by a high concentration of sodium chloride solution in the immediate environment.

The half maximal inhibitory concentration ( $IC_{50}$ ) is a measure of the concentration of a particular inhibitory substance that is needed to inhibit a given biological process by 50%.

Plan an investigation, based on observing the colour of yeast cells mixed with methylene blue, to measure the  $IC_{50}$  value of sodium chloride solution on the respiration of yeast.

You have been provided with the following which you must use:

- prepared sodium chloride solutions with concentrations ranging from 0.1% to 2.0%
- yeast cell suspension, containing glucose as the respiratory substrate
- methylene blue solution
- microscope with an eyepiece graticule.

You may select from the following apparatus and plan to use appropriate additional apparatus:

- normal laboratory glassware, e.g. test-tubes, boiling tubes, beakers, measuring cylinders, graduated pipettes, glass rods, etc.
- syringes and Pasteur pipettes
- timer, e.g. stopwatch
- microscope slides and cover slips.

Your plan should:

- have a clear and helpful structure such that the method you use is able to be repeated by anyone using it
- identify the dependent variable and the independent variable
- identify the variables you will need to control
- use the correct technical and scientific terms
- indicate how the results will be recorded and analysed.

You can consider all steps in the procedure to be low risk and there is therefore no need to include reference to any safety measures in your plan.

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[8]

L1 is a slide of a stained transverse section through a leaf of a land plant that is affected by a fungal infection. The fungal infection affects the upper leaf surface of this plant.

(a) (i) Use your microscope to observe the different tissues in the region of slide L1 shown by the darkly shaded area in Fig. 3.1. The observed area should include at least one vascular bundle and be affected by fungal infection.

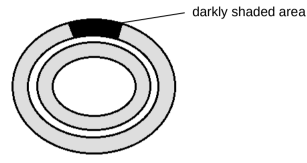


Fig. 3.1

Draw a large plan diagram of the part of the leaf shown by the shaded area in Fig. 3.1.

A plan diagram shows the arrangement of different tissues, including their correct shapes and proportions. No cells should be drawn.

Your drawing should also include any fungal tissue observed. Use **one** ruled label line and label to identify the fungal tissue.

(ii) Observe one vascular bundle of the section on L1.

Select one large xylem vessel element and three adjacent smaller cells.

Each smaller cell must touch the large xylem vessel element and at least one of the other smaller cells.

Make a large, labelled drawing of these four cells.

[5]

(iii) Fig. 3.2 is a photomicrograph of a stained transverse section of part of a leaf from a different species of plant.

Observe the photomicrograph in Fig. 3.2 and the section on **L1** to identify differences between them.

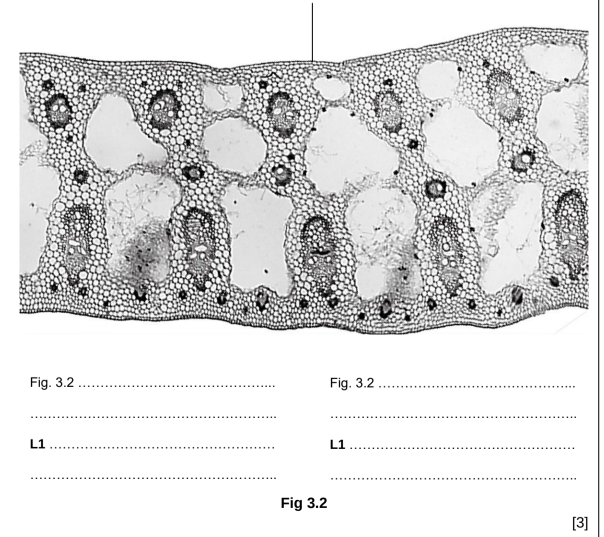
Fig. 3.2 has been annotated to describe **one** of these differences. A label line has been used to indicate the feature that is different.

Complete Fig. 3.2 by:

- identifying and annotating two more differences between the micrograph in Fig. 3.2 and the section on L1
- using a label line to identify the feature that is different.

Fig. 3.2 Upper epidermis is continuous and not disrupted

L1 Upper epidermis is disrupted by fungal tissue



(b) A scientist investigated changes in the mean width of stomata in the leaves of a plant growing in hot, dry conditions. The scientist measured the widths of stomata at different times of day, from 02:00 hours to 22:00 hours. Fig. 3.3 shows where the scientist measured the width of each stoma.

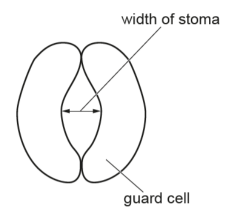
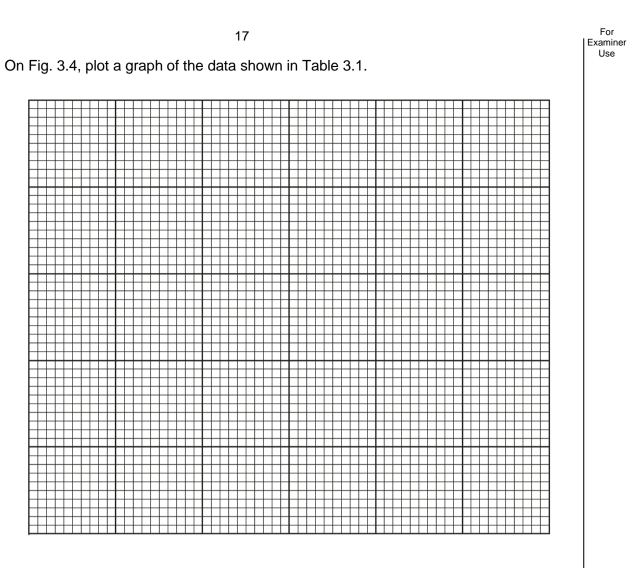


Fig. 3.3

The scientist calculated the mean width of stomata for each time of day. The results are shown in Table 3.1.

time of day	mean width of stomata
/ hours	/ arbitrary units (au)
02:00	86
04:00	36
07:00	4
15:00	2
22:00	95



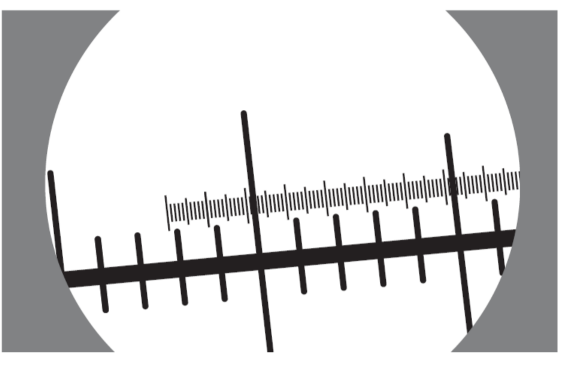


[4]

(i)

[Turn over] H2 9744 Paper 4 Preliminary Examination (ii) Fig. 3.5 shows a stage micrometer scale that is being used to calibrate an eyepiece graticule.

The length of one division on this stage micrometer is **0.02 mm**.





Using the eyepiece graticule shown in Fig 3.5, the width of a guard cell measures **3 eyepiece graticule divisions**.

Calculate the actual width, in micrometres ( $\mu$ m), of this guard cell.

Show all the steps in your calculation, including the appropriate units.

actual width of guard cell = ......  $\mu m$  [3]