

Cambridge International AS & A Level

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CEI NUI	TRE CANDIDATE NUMBER	
ВІС	.OGY	9700/33
Pap	r 3 Advanced Practical Skills 1 O	ctober/November 2023
		2 hours
You	nust answer on the question paper.	

You will need: The materials and apparatus listed in the confidential instructions

INSTRUCTIONS

- Answer all questions. •
- Use a black or dark blue pen. You may use an HB pencil for any diagrams or graphs. •
- Write your name, centre number and candidate number in the boxes at the top of the page. •
- Write your answer to each question in the space provided.
- Do not use an erasable pen or correction fluid. •
- Do not write on any bar codes. •
- You may use a calculator.
- You should show all your working and use appropriate units.

INFORMATION

- The total mark for this paper is 40.
- The number of marks for each question or part question is shown in brackets []. •

For Examiner's Use	
1	
2	
Total	

This document has **16** pages. Any blank pages are indicated.

1 Yeast cells contain enzymes that hydrolyse sucrose into reducing sugars, as shown in Fig. 1.1.

enzymes in yeast cells sucrose — reducing sugars

Fig. 1.1

You will investigate the activity of the enzymes in yeast cells. The yeast cells will be immobilised in sodium alginate beads.

You are provided with the materials shown in Table 1.1.

labelled	contents	hazard	volume/cm ³
Y	yeast cell suspension	none	30
Α	sodium alginate solution	harmful irritant	20
С	calcium chloride solution	harmful irritant	30
S	sucrose solution	none	100
В	Benedict's solution	harmful irritant	30

If any solution comes into contact with your skin, wash off immediately under cold water.

It is recommended that you wear suitable eye protection.

You will investigate the activity of yeast enzymes by using different numbers of beads of immobilised yeast cells in sucrose solution, S.

Carry out step 1 to step 9 to immobilise the yeast cells in sodium alginate beads.

- step 1 Put 10 cm^3 of **A** into a beaker.
- step 2 Stir the yeast cell suspension **Y** with a glass rod.
- step 3 Put 10 cm^3 of **Y** into the beaker containing **A** and mix well.
- step 4 Put 20 cm^3 of **C** into another beaker.
- step 5 Use a 10 cm^3 syringe to collect 10 cm^3 of the mixture of **A** and **Y**.
- step 6 Hold this syringe over the beaker containing 20 cm^3 of **C** (step 4), as shown in Fig. 1.2.





- step 7 Hold the barrel of the syringe with one hand while slowly pressing down on the plunger with the other hand so that a drop of the mixture is released into solution **C**. The drop will form a bead.
- step 8 Repeat step 7 to make at least 31 beads.

The immobilised yeast beads must be left in the beaker for 5 minutes.

step 9 After 5 minutes tip the beads and the solution into a Petri dish.

You will test the activity of the yeast enzymes by using different numbers of beads (1, 2, 4, 8 and 16) in sucrose solution.

Carry out step 10 to step 20.

- step 10 Label five beakers 1, 2, 4, 8 and 16 and label five test-tubes 1, 2, 4, 8 and 16.
- step 11 Put 1, 2, 4, 8 or 16 beads into each of the appropriately labelled beakers, as shown in Fig. 1.3.





- step 12 Put 10 cm^3 of sucrose solution, **S**, into each of the beakers containing the beads.
- step 13 Start timing and leave for 5 minutes. While you are waiting set up a water-bath ready for step 14 and step 19.

In step 19 you will use the water-bath to carry out the test for reducing sugars using Benedict's solution, **B**.

(a) (i) State the temperature you will use for the water-bath.

temperature[1]

- step 14 Heat the water-bath to the temperature stated in (a)(i).
- step 15 At the end of 5 minutes (step 13) stir the contents of each beaker.
- step 16 Use a syringe to transfer 2 cm³ of the solution from beaker 1 into the test-tube labelled 1.
- step 17 Repeat step 16 for each of the beakers and test-tubes labelled 2, 4, 8 and 16.
- step 18 Put 2 cm³ of Benedict's solution, **B**, into each of the test-tubes labelled 1, 2, 4, 8 and 16.
- step 19 Put test-tube 1 into the water-bath at the temperature stated in (a)(i) and time how long before the appearance of the first colour change. If there is no colour change after 2 minutes, stop timing and record as 'more than 120'.

Record your result in (a)(ii).

step 20 Repeat step 19 for the other test-tubes, 2, 4, 8 and 16.

(ii) Record your results in an appropriate table.

(iii)	State the independent variable in this investigation.
	[1]
(iv)	State one main source of error in this investigation.
	[1]
(v)	A student set up a beaker as a control experiment. The result of the control experiment showed that the sucrose was hydrolysed by an enzyme.
	Suggest what substances the student put in the beaker for the control experiment.
	[1]

[5]

(vi) The procedure described in step 1 to step 20 investigated the effect of changing the number of yeast beads on the rate of hydrolysis of sucrose.

Describe how you would modify the procedure to investigate the effect of changing the concentration of the sucrose solution on the rate of hydrolysis of sucrose.

[2]

(b) Pectinases are enzymes that are used to increase the volume of fruit juice extracted from apples. Pectinases can be immobilised and placed inside a container with apple pulp as shown in Fig. 1.4. The fruit juice is then collected.



Fig. 1.4

A scientist carried out an investigation to determine the effect of temperature on the activity of immobilised pectinase by measuring the volume of fruit juice extracted.

All other variables were kept constant.

The results are shown in Table 1.2.

immobilised pectinase		
temperature/°C	volume of fruit juice/cm ³	
30	38	
40	54	
50	79	
60	98	
70	62	
80	4	

Table 1.2

(i) Plot a graph of the data in Table 1.2 on the grid in Fig. 1.5.

Use a sharp pencil.



[4]

(ii) Use your graph in Fig. 1.5 to estimate the volume of fruit juice extracted at 55 °C.

Show on your graph how you obtained your answer.

volume of fruit juice = cm³ [2]

(iii) Describe the change in volume of fruit juice shown by your graph in Fig. 1.5.

The scientist repeated the investigation replacing the immobilised pectinase with the same amount of pectinase but not immobilised in beads (free pectinase).

The results are shown in Table 1.3.

free pectinase (not immobilised)		
temperature/°C volume of fruit juice / cm ³		
30	52	
40	85	
50	98	
60	41	
70	12	
80	4	

Table 1.3

(iv) Suggest an explanation for why more fruit juice was obtained when using the free pectinase between the temperatures of 30 °C and 50 °C than when using immobilised pectinase.

(v) Suggest an explanation for why more fruit juice was obtained when using immobilised pectinase between the temperatures of 60°C and 70°C than when using the free pectinase.

[2]
[Total: 22]

- 2 K1 is a slide of a stained transverse section through a plant leaf.
 - (a) (i) Draw a large plan diagram of the whole leaf on K1. Use a sharp pencil.

Use **one** ruled label line and label to identify the epidermis.

[5]

(ii) Observe the cells in the epidermis of the leaf on K1.

Select a line of **four** adjacent epidermal cells.

Each cell must touch at least **one** of the other epidermal cells.

- Make a large drawing of this line of **four** cells.
- Use **one** ruled label line and label to identify the cell wall.

Fig. 2.1 is a photomicrograph of a stained transverse section of a leaf from a different type of plant.



Fig. 2.1

(b) Identify three observable differences between the leaf on **K1** and the leaf shown in Fig. 2.1.

Record these **three** observable differences in an appropriate table.

Fig. 2.2 is an enlarged version of the transverse section of the leaf shown in Fig. 2.1.



Fig. 2.2

(c) (i) Measure line **P** and line **Q** to find the length of vascular bundle **P** and the length of vascular bundle **Q** as shown in Fig. 2.2.

length of vascular bundle **P** =

length of vascular bundle **Q** =

(ii) Calculate the percentage difference in length between vascular bundle **P** and vascular bundle **Q**.

Show your working and give your answer to **two** significant figures.

answer =%[2]

[2]

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