## UNIVERSITY OF CAMBRIDGE INTERNATIONAL EXAMINATIONS

GCE Advanced Subsidiary Level and GCE Advanced Level

## MARK SCHEME for the October/November 2008 question paper

## 9700 BIOLOGY

9700/32

Paper 32 (Advanced Practical 2), maximum raw mark 40

This mark scheme is published as an aid to teachers and candidates, to indicate the requirements of the examination. It shows the basis on which Examiners were instructed to award marks. It does not indicate the details of the discussions that took place at an Examiners' meeting before marking began.

All Examiners are instructed that alternative correct answers and unexpected approaches in candidates' scripts must be given marks that fairly reflect the relevant knowledge and skills demonstrated.

Mark schemes must be read in conjunction with the question papers and the report on the examination.

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Question	Expected Answers		Additional Guidance	Marks
Draw and I	abel ONE cell in distilled water		2 MMO collection	+
1 (a) (i)	one cell drawn (at high power), two lines for	or cell wall;	Ignore low power. Reject two or more cells together.	
	correct cell structure and cell wall and nuc	<u>cleus</u> labelled correctly;	Rej. if have additional organelles mitochondria, chloroplasts, Golgi.	[2]
Present yo	Present your observations from the slides made from distilled water, T1 and T2		2 MMO decisions, 2 PDO recording	
1 (a) (ii)	single table, all cells drawn, column headings: solution/slide/(distilled) water/ <b>W</b> and <b>T1</b> and <b>T2</b> ; to left/across top, observations/feature/e.g. to right/ underneath/clear what is recorded in the	Or when only drawings given three drawings, labelled (distilled) water/W, T1 and T2; clear that cell walls and cell membranes are all different (for water, T1 and T2);  T1 cell membranes/cytoplasm pulled	No outer boundary needed for table.  Reject cells shrink or become smaller. Accept vacuole shrinking or drawn.	
	boxes; T1 cell membranes/cytoplasm pulled away from cell wall/plasmolysed; T2 granules/particles in cell/more plasmolysed/destroyed/stained/coloured e.g. brown/black/AW;	away from cell wall/plasmolysed; <b>T2</b> granules/particles in cell/ <u>more</u> plasmolysed/destroyed/stained/coloured e.g. brown/black/AW;	Allow any description that cells have been destroyed/cell membranes ruptured/disorganised/leakage of cell.  Reject cell walls broken down.	[4]

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Explain ob	oservations from water, T1 and T2.				
1 (a) (iii)	Idea of  1. high/less negative water potential to lower/more negative water potential/down water potential gradient  Any two of:  2. (in water)idea of water has moved in/no net movement;  3. (in T1/T2) idea of water has moved out;  4. (in T2/lead nitrate) killed/destroyed cells/toxic/effect described/AW;	AND by osmosis at any point;	the same as water pote so reject pt1 if wrong wa Ignore hypotonic and hy correct context if used.	ypertonic but must be in e candidate's own results.	[1]
Identify tw	o sources of error in this experiment			2 ACE interpretation	1
1 (a) (iv)	Two from evaporation from solutions/concentration of cells left different lengths of time/too short AVP; volume/no. of drops used; or different or different onions/parts of onion/	a time/not long enough;	Reject not immersed.  Reject should be same time –not an error.  Reject amount.	Mark for any correct.  Reject improvements.	[2 max]

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Suggest	how you could modify the experiment to investigate the effect of lead nitrate.		3 ACE improvements	
1 (b)	more/serial dilution concentrations of lead nitrate; Then any <b>TWO</b> from at least 3 specified lead nitrate concentrations; repeat each concentration/more than one strip (per concentration); keep the time the same/give an example of a time/longer time; keep the volume <b>AND</b> method /use graduated pipette/no.of drops the same/AW; same onion/same part/fresh; detailed measurement method/use of eyepiece graticule to measure plasmoylsed cells/count number of plasmolysed cells in a sample of 20 or more;	Reject shorter time.	·	[1]

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Complete	the Table 1.2	by calculating the missing values	PDO display	
1 (c) (i)	<u>6,81;</u>		A whole numbers only and both correct	[1]
	oh of concent	ration of sodium chloride against the percentage plasmolysis of the cells	PDO layout	
1 (c) (ii)	parcentage	80		[3]

0	x-axis conc, mol dm <sup>-3</sup> /M or molar/mole(s)/l or per litre	AND y-axis percentage/% plasmolysis;	Rej. mol/dm <sup>-3</sup> and mol dm <sup>3</sup> .	[1]
S/P	scale as shown/y axis 25 to 2cms, allow no 0 marked	AND plotting crosses or dot in circle ONLY AND 0.0, 0.2 and 0.6 and 1.0 plotted correctly; no larger then <b>X</b> or <b>O</b> plots 0.2 must be on horizontal line, 0.2 and 0.6 and 1.0 between the horizontal lines. Ignore incorrect calculated mean plots i.e. 0.4 and 0.8.	Rej. blobs in or out of circle.	[1]
L	either ruled lines joining each point or smooth curve thro go to 0	ugh 0; no thicker than no feathery line, line must	Rej. any extrapolation beyond either axis.	[1]

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Question	Expected Answers	Additional Guidance	Marks
State cond	entration at which 50% plasmolysis occurred	1 ACE interpretation	
1 (c) (iii)	take reading from candidates own graph, AND must have units;	Allow two decimal places. Ecf units from graph.	[1]
'The more	concentrated the solution the more plasmolysed the cells become' draw	2 ACE conclusion	
conclusion	n include whether the data support the hypothesis and produce a revised		
hypothesis	s if necessary		
1 (d)	General statement :		
	Either support or no support or partial support for the hypothesis or writes a	Needs clear statement.	
	conclusion which states the hypothesis;	Reject supports conclusion.	
	quotes 2 sets of figs. with both axes; <b>OR</b>		
	idea that up to 0.4 /low concentration only small % plasmolysed/or % plasmolysis	Idea of correct relationship may quote figures to	
	does not increase evenly with increasing concentration/or levels off at high	get same idea.	
	concentration;		
		Reject all/100% plasmolysed.	[2]

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Question	Expected Answers	Additional Guidance					
Draw a LA	RGE, LOW-POWER plan diagram of photomicrograph fig 2.1. (artery)	1 MN	MO collection, 3 PDO layout				
2 (a) (i)	sharp, clear unbroken lines, height no more than two thirds the length; no cells, no shading, larger than 6 cm in any direction; at least three lines (plus very thin inner layer if shown); uneven all the way round and one solid inner line;	Outer two lines <b>only</b>					

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Use this info	rmation to calculate the actua	l width of the	e lumen.			2 M	MO co	ollectio	n, 1 P	DO re	ecordir	ng, 1 F	DO di	splay	
6	Each division on stage scale is ( First mark Reject any measurements giver		nts 1 and 2. <i>F</i>	Accept units o	r divisions	S.								-	
First Mark	No. of eyepiece grat. W	- 7	7	1.	4		2	:8			2	29			
Second Marl	<del>-  </del>	4.5	9.0	9	18	7	18	25	36	7	18	25	36	1	
	No on stage micrometer Z	5	10	5	10	2	5	7	10	2	5	7	10		
Third Mark	Show logical reasoning	and then W, strictly the c	d and allow n or W and the orrect reason Ignore answe . if additional	er and units. figs. even if x	ough not	OR Z x V AND divided by Y. followed by x W									
Fourth Mark	Need NOT be the correct answer			) and 999 wit correct.  Rejec	•	OR answer between 0.1 and 0.99 with mm; Allow standard form if correct. Reject metres.							[4]		
	s are for – <u>collecting</u> the corrects for <u>recording</u> – use of the corre		ird mark is fo	or <u>display</u> – sh	owing cle	ar reasc	ning ii	n the c	alcula	tion.				•	
Suggest how	an error in measuring the lur	nen could od	cur								1 A	CE in	terpret	ation	
n	not knowing where edge is/lumen irregular shape/preparation squashed/only 1 measurement/thicknesses of lines(stage micrometer)/between divisions on eyepiece graticule/one scale line is not at edge of lumen/focussing of both scales/lining up the scales;				on	gnore pa	arallax	error							[1]

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Compare and contras	t specimens Fig. 2.4 and 2	2.5.	2 MMO collection 1 PDO recording 2 ACE interpretation	n
2 (b) (i)	The state of the part of the p	1020		
	nn diagram/ruled boxes cor	nnected, correctly headed;	Must have at least one similarity.	
comparative statements	Fig. 2.4		Accept tubes/vessels as alternative to lumen.	
comparative statements	Fig. 2.4  lumen/central space;	Fig. 2.5		
comparative statements	Fig. 2.4	nnected, correctly headed;	Accept tubes/vessels as alternative to lumen.	
both have	Fig. 2.4  lumen/central space; larger,	Fig. 2.5 smaller;	Accept tubes/vessels as alternative to lumen. Reject ref. to Fig. 2.4 having cells – not visible. Reject uses	
both have lumens number (lumen/tubes) cells /cell walls/end	Fig. 2.4  lumen/central space; larger,  single/one,	Fig. 2.5  smaller; more/lots;	Accept tubes/vessels as alternative to lumen. Reject ref. to Fig. 2.4 having cells – not visible. Reject uses	

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Suggest or	ne feature which indicates the Fig 2.5 is a			ACE conclusion	
2 (b) (ii)	have cell walls/xylem/phloem/sieve tube (	element)/companion cell/pits/rings;	Ignore cellulose, lignin, vessel on own. Reject sieve plates.		
Make a lab	elled drawing of 5 representative cells.		1 MMO collection, 3 MMO decisions		
2 (b) (iii)	<ul> <li><u>5</u> shown on Fig.;</li> <li>drawn 3 diverse cells;</li> <li><u>3</u> different sizes;</li> <li>at least 1 cell drawn with bands/parts of bands/pits;</li> </ul>	AND longer than wide;	Reject point 1 if more than 5 marked or drawn. Entire cells or open tubes. Ignore labels.	Reject points 2, 3 and 4 if more than 2 TS or textbook.  Max 1 point, 1 only	[1]
					[3]
	(b) The whole specimen in Fig. 2.2 is repeated below without the Fig. 2.5 shows a longitudinal section of a specimen from a Fig. 2.4 and Fig. 2.5 are not reproduced at the same scale.				