CAMBRIDGE INTERNATIONAL EXAMINATIONS

Pre-U Certificate

MARK SCHEME for the May/June 2013 series

9790 BIOLOGY

9790/03

Paper 3 (Practical), maximum raw mark 80

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, which would have considered the acceptability of alternative answers.

Mark schemes should be read in conjunction with the question paper and the Principal Examiner Report for Teachers.

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Section A

Question	Sections	Indicative Material	Mark
1 (a)	ADC Interpretation of data	label lines to: (chloroplast) envelope / inner and/or outer membranes; accept chloroplast membrane(s) stroma; thylakoids / lamellae; accept grana (70S) ribosomes; starch (grain);	max 3
(b)	EPD Identifying limitations and sources of error	ice-cold to, prevent / reduce, (hydrolytic) enzyme action; that would, destroy / damage, the chloroplasts; buffer solution to maintain constant pH; keep enzyme activity constant / prevent denaturation; ref. to pH and chemiosmosis; sucrose solution has same water potential as, cell / cytoplasm / stroma / chloroplast; so chloroplasts remain intact / prevent chloroplasts bursting; ignore plasmolysis ref. to osmosis and water movement into the chloroplast (if suspended in water);	max 4
(c)	MMO Decisions EPD Improvements	DCPIP solution without any, leaf extract / chloroplasts; DCPIP may decolourise in the light / AW; to show, leaf extract containing chloroplasts is needed for colour change / DCPIP does not decolourise in light without chloroplasts / AW; boiled leaf extract; to show decolourising involves, enzymes / proteins; use, folded black card / aluminium foil, with DCPIP and leaf extract; accept 'put in the dark' to show light is necessary for colour change; AVP; e.g. further control(s) AVP; e.g. explanation for any extra control(s)	max 4

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(d)	PDO Recording	data recorded as a single table with percentage transmission in left hand column; accept light intensity ignore filter number informative column headings, correct units in column headings only; percentage transmission, time to decolourise (s), rate of photosynthesis / reduction of DCPIP (s ⁻¹) reject time unqualified results recorded to same degree of precision in each column; replicate(s) included and mean calculated and included; time recorded in seconds (not minutes and seconds);	
	MMO Collection	control(s) recorded in table; results show expected trend;	7
	ADC Display of calculation and reasoning	rate calculated correctly as 1 / t (× 1000); accept alternatives, e.g. a constant figure divided by the time taken	1
(e)	PDO Graph	axes correctly positioned (x-axis = percentage transmission, y-axis = rate of photosynthesis); accept time as ecf from 1 (d) axes scaled with ascending scales starting at 0,0; accept time / rate, only if filter number given axes with full titles and units; ecf if filter number and/or time points plotted accurately; result for control in dark (0%) plotted; points joined, clearly / neatly, by straight lines (unless conform to line/curve of best fit); reject if line goes beyond last plotted point	max 5

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(f)	ADC Description of patterns and trends	description of pattern from graph; e.g. rate increases as percentage transmission increases use of comparative data from, table / graph, to illustrate; ecf if time	2
	ADC Interpretation of data Making conclusions	light provides energy for, photosynthesis / LD stage; absorption of light by (named) chloroplast pigments; electrons / e ⁻ , are, energised / excited, and, leave, chlorophyll / photosystem II / reaction centre; ref. to electron carrier system in context; photolysis produces H ⁺ ions (and electrons); electrons / hydrogen ions (H ⁺) / protons, reduce DCPIP; independent variable is light intensity; light intensity is limiting factor; (if a plateau / levelling off) light intensity is no longer the limiting factor / some other factor is limiting; named factor(s); explanation of effect of named factor; e.g. temperature and damage to, proteins / carriers	max 6
(g)	ADC Making conclusions	any one of the following methods for disrupting membranes to max 1: suspend chloroplasts in, water / dilute solution / solution with a higher water potential; reject water ultrasound; named enzyme to digest, protein / phospholipid; heat shock; electric shock / electroporation; freeze-thawing; detergent; ethanol / organic solvent; acid / alkali; explanation to max 1 in terms of: envelopes / chloroplasts, break down (to release thylakoids); envelopes / chloroplast membranes, become more permeable (to allow electron acceptor into	
		organelles);	2

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(h)	ADC Making conclusions	penalise NAD once only	
	Making Conclusions	ref. to small quantity of NADP so is continually recycled;	
		if all, / most / many, electrons flow to DCPIP; none / few, available to reduce NADP;	
		no requirement for oxidised NADP; not dependent on Calvin cycle to recycle oxidised NADP; ref. to, non-cyclic photophosphorylation / ETC to	
		NADP; no production of ATP as no electron flow between PSII and PSI, so no energy for Calvin cycle;	
		reduced NADP / AW, is required for reduction stage of Calvin cycle; accept if no reduced NADP no Calvin cycle (if no reduced NADP) no conversion of PGA to,	
		PGAL / GP / TP / AW ;	max 3

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(i) Evaluation of procedures and data			
	Identifying limitations and sources of error	Suggesting improvements	
reliability repeatability	only one result per light intensity; accept ref. to number of replicates / 'do (more) repeats'	repeat at least three times and calculate mean (and SD); ref. to appropriate stats test, to test for correlation;	
	ref to anomalous results ; e.g. not able to check for them	repeat results that do not fit the trend; accept ref. to actual results	
end point / timing	end point is subjective / AW;	use apparatus that allows measurement of light transmitted through; reject 'use a colorimeter' unqualified	
	difficult to make sure the same end point is used each time;	better if can use a quantitative end point;	
	colour standard may have changed over time;	set up fresh colour standard each time;	
	low light intensities changing colour but slowly;	leave for longer than 10 minutes;	
	stated problem with start time;	standardise method for starting the timer;	
preparation of suspension	density of chloroplasts may not be the same in each tube;	method to standardise; e.g. stirring same number of times before taking each sample in melting point tubes / use magnetic stirrer	
	no stated volume of DCPIP;	standardise blue-green colour;	
	tubes set up at different times;	use a fresh mixture of chloroplast suspension (and DCPIP) each time;	
	denaturation / destruction, of proteins / AW (as temperature increased);		
	not a pure suspension of chloroplasts;	centrifuge;	
	very small volume in tubes;	use, test-tubes / specimen tubes / micropipettes ;	

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(i) Evaluation of procedures and data (continued)			
	Identifying limitations and sources of error	Suggesting improvements	
independent variable	external / stray / ambient, light; accept difficult to keep fixed distance for lamp	any suitable suggestion for keeping out stray light; e.g. dark chamber with single light source	
	dark filter is lifted / light filter is not;	standardise checking of colour / cover lamp with filter;	
	do not know actual light intensity;	use a light meter;	
uncontrolled variable	temperature was not constant / heat from bench lamp;	any suitable suggestion for maintaining constant temperature; e.g. heat screen / LED lamp	
	ref. to actual temperature(s);		
results	not enough / only six, filters / light intensities;	intermediate light intensities stated; reject ref. to range (as 0 to 100% included)	
	ref. to, number of points on the graph / pattern of results, in support;		
max 8			
		[Total: 45]	

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Section B

Question	Sections	Indicative Material	Mark
2 (a)	MMO Decisions Collection PDO Recording	drawing: large drawing that fills the space available with clear, unbroken lines; V-shape; primary lamellae shown clearly at least in part of drawing or as an inset; primary lamellae not all same length; and other detail, e.g. 'segmented pattern' of gill arch; annotations – suitable comments on shape / texture / colour / surface area of: gill arch; primary lamellae; secondary lamellae;	max 6
		scale indicated e.g. \times 2, \times 3;	1
(b)	PDO Recording	drawing large drawing to fill the space available; clear, unbroken lines;	
	MMO Collection	drawing to show secondary lamellae; label any two of the following;; secondary lamella(e) epithelium blood cells pillar cells cartilage endothelium / capillary nuclei	max 4
	ADC Interpretation of data Display of calculation and reasoning	magnification correctly calculated from measurements given; explanation: calibration of eyepiece graticule; maybe stated (as already known) and/or explained using stage micrometer measurement from slide using eyepiece	
		graticule ; measurement indicated on drawing ;	4

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(c)	MMO Decisions	take sections of gill to give LS of secondary lamellae / AW; dissect out secondary lamellae; measure (length and width) with, graticule / grid; measure maximum length of secondary lamella; measure depth; use radius to measure surface area; double for both sides;	max 3
(d) (i)	ADC Interpretation of data	idea that: standardise measurements to make valid comparisons between different fish (species);	1
(ii)	ADC Making conclusions	mackerel is most active of these species / AW; relatively more, muscle / active tissues; high rate of aerobic respiration; to provide, energy / ATP, required for swimming; needs high rate of uptake of oxygen; ref. to different concentrations of oxygen in water; any reason; e.g. different temperatures / depths / effects of currents / AW monkfish, is relatively inactive / maybe bottom dweller; uses anaerobic respiration for short bursts of activity; use of comparative figures from Table 2.1 in support of explanation; AVP;	max 3
(e)	MMO Decisions Collection	table with a column for features; comments on the following for each structure: qualitative comparison of width (of lumen); measurements of two or three taken from slide; qualitative comparison of thickness of walls; measurements of two or three taken from slide; relative numbers; shape; comparison of lining: presence / absence of ciliated epithelium; presence / absence of, endothelium / squamous epithelium; ref. to goblet cells / described; presence / absence, of other tissues in walls; e.g. smooth muscle, cartilage, elastic tissue, blood capillaries presence / absence of blood (cells) in lumen; AVP;	
		e.g. position in the slide	max 7

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(f)	MMO Decisions Collection	use of a table for comparisons; comments on the following: position of gas exchange surface; e.g. deep inside mammalian body vs in gill chamber / AW movement of respiratory medium; e.g. air travels through airways in mammalian lung vs water flows directly over gills in fish air flows tidally, water flow is unidirectional; both have a gas exchange surface with large surface area; gaseous exchange surfaces are alveoli and secondary lamellae; both have thin epithelium; both have short diffusion pathway;	
		secondary lamellae; both have thin epithelium; both have short diffusion pathway; both are well supplied with, capillaries / blood	
		(vessels); blood flows around alveoli; countercurrent flow in gills; both are moist; AVP;	
		e.g. ref. to surfactant AVP;	max 6

[Total: 35]