



Chemistry B (Salters)

Advanced GCE F335

Chemistry by Design

Mark Scheme for June 2010

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Mark schemes should be read in conjunction with the published question papers and the Report on the Examination.

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Q	Jestic	n	Expected Answers	Marks	Additional Guidance
1	а	i	dinitrogen oxide / nitrogen(I) oxide / dinitrogen(I) oxide \checkmark	1	ALLOW dinitrogen monoxide IGNORE gaps
	а	ii	N I O dative bond ✓ completely correct ✓ shape – linear ✓ (depends on diagram – see advice)	3	To score first mark there must be (only) two electrons of the same symbol between the nitrogen and oxygen ALLOW (for this mark) if they are oxygen's electrons. To score the second mark there must be alternating dots and crosses for the elements' electrons as one moves from N to N to O ALLOW shared electron pairs horizontally (eg •+) ALLOW alternatives to linear, eg "straight" or 180 IGNORE 'planar' IF diagram is wrong, use it to determine shape mark: • No diagram no mark • No lone pairs on central N: linear, etc • One or two lone pairs/single electrons: bent (NOT triangular), allow 120+2 or 109+2 as appropriate
	b	i	+5 🗸 +1 🗸	2	5, 1 does not score. 5+, 1+ scores 1
	b	ii	10 $H^+ + 2NO_3^- + 8 e^- \rightarrow N_2O + 5 H_2O$ 10 AND 5 \checkmark 8 \checkmark <i>Mark separately</i>	2	Each piece of additional material in the equation CON s a mark
	b	iii	 (nitrogen / nitrate) gain of electrons ✓ oxidation number / state <u>of nitrogen</u> goes down / goes from (+)5 to (+)1 (or ecf from b(i), provided this is a fall) ✓ 	2	'gain of electrons' need not be qualified but any other reagents quoted apart from nitrogen/nitrate are CON IGNORE answers in terms of oxygen lost IGNORE what has gained electrons Answers can both be on same line NB - 1(b)(i) answer line is shown on the screen to allow for ecf
	b	iv	nitrogen (in compounds) <u>/nitrate(s)</u> : is less available to plants / crops is needed by plants / crops is a fertliliser makes plants / crops grow (<i>AW</i>) ✓	1	For the mark, nitrate or 'nitrogen' is needed (ALLOW N but NOT N ₂), AND an indication of its availability / need by plants OR fertiliser function IGNORE '(nitrogen) is reduced', implying less of it

Question	Expected Answers	Marks	Additional Guidance
C	33% N OR 1:2 by moles (stated or implied) OR two-thirds oxygen \checkmark NO ₂ \checkmark no ecf from the wrong working	2	Answer alone scores 1 (NOT 2) if first marking point is not scored IGNORE Multiples of NO ₂ (eg N ₂ O ₄) but working can score
d i	$\begin{array}{c c} \hline Mark separately \\ \hline H & $	2	ALLOW CH ₂ etc for first mark must be full structural to score second mark ALLOW dashes or numbers on R, but no other representation of acid side-chain $H = \begin{array}{c} 0 \\ 0 \\ 0 \\ 0 \\ 0 \\ 0 \\ 0 \\ 0 \\ 0 \\ 0 $
d ii	instantaneous (dipole) – induced dipole/ permanent (dipole) – permanent dipole ✓ intermolecular bonds are similar <i>stated or implied</i> OR imb formed are stronger than / similar to those broken ✓	2	ALLOW either or both types as imb for either compound or between compounds (ALLOW permanent dipole – induced dipole between). Hydrogen bonds are CON. NOT abbreviations of bond descriptions for <i>this</i> mark IGNORE references to molecules being non-polar eg 'both have' give this mark, even if imb stated is wrong but reference to
	Mark separately		covalent bonds is CON to this mark

Question		n	Expected Answers	Marks	Additional Guidance
	e	i	Image: Weight of the second	3	ALLOW carboxylic acid groups in any orientation Ignore 'drafting' lines, give BOD if possible no ecf for non-skeletal structures for more than one compound CON O–H on first appearance, then allow as ecf – including 2(a) –OH connected through the H is a CON only the first time it occurs OH OH acceptable unacceptable up to bar of H beyond bar of H
	e	ii	190 and 226 ✓ 190 x 100/226 = 84/84.1% ✓	2	Full marks for correct answer with no / inaccurate working shown. 64.6/65 with no working scores 1 Second mark is for writing a ratio of two numbers (including eg '100 + 2x63') and correctly evaluating the answer (but NOT if answer >100%) ALLOW 100% ALLOW 2 or more sig figs (allow any value between 84 and 84.1)
			Total	22	

Question	Expected Answers	Marks	Additional Guidance
2 a		2	Ignore 'drafting' lines, give BOD if possible ALLOW other skeletal representations, including O of OH must be attached to carbon (see rule in 1ei) ALLOW ecf on O–H if it occurred in 1ei and mark was not awarded because of this NOT three-dimensional representations with wedges and dashes
	2-methylpropan-2-ol / methylpropan-2-ol \checkmark Mark separately – no ecf		IGNORE dashes, commas, gaps in name ALLOW 'methly' but no other mis-spellings
b	carbon with OH is attached to three other carbons / methyl (groups) / alkyl (groups) / R (groups) OR carbon with OH ✓	1	It must be clear that the carbon is being referred to ALLOW 'central carbon' instead of 'carbon with OH'
bi	from: orange/yellow ✓ to: green/blue ✓ butanone ✓ <i>Mark separately</i>	3	 DO NOT ALLOW other colours apart from mixtures or shades of those given ALLOW butan-2-one (ignore dashes, brackets commas and gaps) IGNORE formula

Question	Expected Answers		Additional Guidance	
C	$\int_{0}^{\delta_{H}} \int_{0}^{\delta_{H}} $	3	 Hydrogen bond can be represented by a dashed line but NOT a solid line (unless labelled as 'hydrogen bond') ALLOW 'OH' for O–H Representation of alcohols can be in any way that indicates their structures (ALLOW missing Hs), provided OH groups are clear. ALLOW ecf from wrong alcohol structure in 2a ALLOW ambiguous attachment of alkyl groups (eg via Hs) but not OH (see rule in 1di) Ignore 'drafting' lines, give BOD if possible If only an incorrectly positioned hydrogen bond is drawn (eg to alkyl H) it scores zero out of three. ButIf there is more than one hydrogen bond: Incorrectly positioned hydrogen bonds Mark the best hydrogen bond A 'square' of two hydrogen bonds can be considered for all except the first mark 	

Question		on	Expected Answers	Marks	Additional Guidance
	d		instantaneous (dipole) – induced dipole bonds/forces ✓	3	NOT abbreviations for this mark ALLOW 'Van der Waals' (ignore capitals) other bonds accounting for the difference are CON ALLOW 'it' for 't-butanol'
			(intermolecular bonds) are weaker in t-butanol / less energy (or heat) required to: break (intermolecular bonds) in t-butanol / separate molecules in t-butanol <i>ORA</i> ✓		IGNORE less/fewer imb or 'less likely to form' ALLOW second mark for any one of the following described as weaker: just 'intermolecular bonds / forces' OR any named intermolecular bond (including hydrogen bonds or different ones for the two alcohols) OR abbreviated (eg 'id-id' / imb)
			t-butanol molecules/chains OR t-butanol: can't get as close together / don't line up / don't pack/stack together so well/so easily / less areas of contact <i>ORA</i> ✓		Mark third mark separately IGNORE more / less branched or linear IGNORE atoms

Question	Expected Answers		Additional Guidance
Question	Expected Answers $H_{3}C$ CH_{3} $H_{3}C$ CH_{3} C	4 4	Additional GuidanceFirst arrow must start at lone pair and point between O and Hor at Hsecond arrow must start on (or above or below) bond (NOTon H) and point towards Cl (see box)third arrow must start on bond and point towards Ofourth arrow must start on lone pair and point between C ⁺ and Cl or at C ⁺ Any wrong arrows in excess of four are each CON to onecorrect arrowFirst and third arrows can be straightSingle-headed arrows are CON only the first time they appearin an otherwise correct situation; accept after that
	\downarrow CH ₃		
	Mark separately – no ecf		

Question		Expected Answers	Marks	Additional Guidance
1		CH ₃ CH ₂ OCH ₂ CH ₃ ✓	6	Please annotate by ticking each point scored. Always look for information on page 8 and mark appropriately. Page 8 is available as a thumbnail on the left
		IR (two marks) no peak above $3000 / 3200 / 3640 \text{ (cm}^{-1})$ OR no peak at $3600 - 3640 / 3200 - 3600 \checkmark$ no OH / not alcohol \checkmark NMR (one mark plus QWC) QWC is scored for relating NMR evidence to structure, as described below <i>Look first for:</i> (protons identified) CH ₃ (–C) / methyl AND O(–)CH ₍₂₎ (or in		Accept any clear structural formula of ethoxyethane, including C ₂ H ₅ OC ₂ H ₅ IGNORE name Ignore 'drafting' lines, give BOD if possible IGNORE references to other peaks <i>Mark two IR marks separately. Can score if the structure is</i> <i>wrong.</i> <i>Note brackets carefully</i> : H ₃ in first and O in second are essential - NOT CHO for second one but CH–O and CH ₂ O are acceptable. IGNORE any shifts quoted
		words or indicated on structure) \checkmark The QWC mark is then scored from this response if 'CH ₃ CH ₂ OCH ₂ CH ₃ ' (or more displayed) structure given. Place tick under pencil icon \checkmark		ALLOW 'protons' stated or implied (eg 'methyl group')
		 OR if the above are absent, incomplete or wrong, look for: two proton / hydrogen environments ✓ QWC is awarded here if there is an indication of two environments on the <i>correct</i> structure (using the formula or describing it in words) ✓ Place tick under pencil icon 		For QWC here ALLOW references to symmetry or 'two C_2H_5 '
		splitting (one mark): general: indication that no. of peaks is one more than the no. of protons on the adjacent <u>carbon</u> OR specific: identification of one of $CH_3 CH_2$ (ie triplet for hydrogens on C next to CH_2 or quartet for hydrogens on C next to CH_3) (ignore anything incorrect) \checkmark		Either way, the idea of number of protons / hydrogens on <i>adjacent</i> (<i>AW</i>) carbon (IGNORE adjacent environment) must be there to score the mark. IGNORE specific if general correct Can score splitting mark if the structure is wrong NOT just 'n+1 rule' without explanation
		Total	22	

Que	Question		Expected Answers	Marks	Additional Guidance
3	a	i	Endothermic (forward reaction), (high / increasing temp moves) equilibrium (position) to right / towards products ✓	4	IGNORE references to rate ALLOW 'reverse reaction is exothermic' IGNORE 'moves in / favours endothermic direction'
			(high) pressure pushes <u>equilibrium</u> (position)* to the left \checkmark more moles / molecules / particles on the right <i>ORA</i> \checkmark		NOT 'more atoms' or 'more products'
			(ignore wrong references) ✓		(can assume <i>high</i> pressure or temperature since given in the question)
			*'position' must be mentioned <i>once</i> . Award one of these marks without 'position' but for both marks it must be mentioned once.		
	a	ii	(the energy source must) not produce carbon dioxide / CO_2 OR fossil fuels produce carbon dioxide / CO_2 (<i>AW</i>) \checkmark	1	ALLOW "It" for the Sun, so allow, eg, "It does not form CO ₂ "
	b	i	$Kc = [CO]^2 [O_2] / [CO_2]^2 \checkmark$	1	Must have square brackets; NO mark if p symbols. In top line: may have multiplication sign, must not have plus sign. IGNORE state symbols
	b	ii	4 x 10 ⁻²⁰ ✓ 1sf ✓ mol dm ⁻³ ✓ <i>Mark separately</i>	3	ALLOW ecf for first and third marks from $b(i)$ UNLESS plus sign used The $(b)(i)$ answer is shown on the screen to facilitate ecf Award sf mark if the number is to 1 sf and is the correct or incorrect result of any calculation shown. units: ALLOW mol/dm ³ 4 x 10 ⁻²⁰ on answer line scores 2
	С	i	+172 (number with sign) ✓✓	2	+188, 172 and –172 score one mark; nothing else does

Question		on	Expected Answers	Marks	Additional Guidance
	С	ii	T= 566000/172 ✓	2	ALLOW ecf from c(i) negative temperatures are CON second mark is for manipulation and correct statement of unit;
			= 3290 K ✓		no ect from errors in first mark(i.e. 3.29 K scores zero) ALLOW 'Kelvin' and lower-case 'k'
					negative answers score zero allow 2 or more sf: 3300, 3291, 3290.7 etc correct answer with no working scores 2
	d	i	$Ca(OH)_2 + CO_2 \rightarrow CaCO_3 + H_2O \checkmark$	1	ALLOW Ca(OH) ₂ + $2CO_2 \rightarrow Ca(HCO_3)_2$ IGNORE state symbols. Brackets in formulae must be correct Anything extra on either side is CON
		ii	acid-base ✓	1	ALLOW any unambiguous indication of the answer eg circling any others indicated are CON
		iii	uses a lot of Ca(OH) ₂ / large amounts of solid (or CaCO ₃) formed OR CO ₂ emitted in manufacture of Ca(OH) ₂ (AW) \checkmark	1	Must have idea of 'large amount' to score in this way IGNORE cost, expense, damage to environment etc
			Total	16	

Question		on	Expected Answers	Marks	Additional Guidance
4	а	i	1/13 x 44 = 3.38 g Correct M_r values 44 and 13 (or 78/6) ✓ Correct manipulation of recognisable M_r values and evaluation (with ecf) ✓	2	Full marks for correct answer ALLOW two or more sf 0.56(4) scores 1
	а	ii	(it will be the value of the) highest mass / $\frac{m}{z}$ peak / molecular ion (peak) / M ⁺ (peak) peak furthest to the right \checkmark	1	NOT just "highest peak" IGNORE base peak
	b	i	H = C = C = C = C = C = C = C = C = H = 120	3	structure need not be the correct shape (eg it can be linear) NOT 3-dimensional structure (showing dashes and wedges) any one 180 angle and any one 120±2 angle should be illustrated NB NOT $\stackrel{180}{\bigcirc}$ ALLOW $\stackrel{180}{\bigcirc}$ (extra wrong angles are CON to one correct angle mark each) If there are errors in parts of this structure, bond angle marks can still be given for the correct parts. However, wrong structures (eg Kekulé benzene) score zero marks out of three.
	b	ii	Alkene / C=C groups / double bonds / unsaturated groups / alkyne AND react with HBr / undergo addition reactions ✓	1	IGNORE references back to 'structure in (i)' IGNORE references to decolorising / reaction with bromine 'substitution' is CON

Q	Question		Expected Answers	Marks	Additional Guidance
	С		accounts for: bond angle same / 120 ✓	4	ignore that it is evidence for the hexagon alone (rather than a symmetrical hexagon)
			three bonds OR three groups / sets of electrons around each carbon \checkmark		NOT pairs of electrons
			does not account for: bond lengths equal \checkmark		
			single bonds longer than double bonds \checkmark		
			Mark separately		
	d	i	delocalised (electrons) ✓ one electron from each carbon ✓	4	QWC 'delocalised' must be spelt correctly to score first mark ALLOW 'delocalized' or derivations such as 'delocalisation'
			two rings ✓		'Above and below' in last point will cover the 'two'
			above and below <u>carbon atoms / carbon ring</u> \checkmark		IGNORE 'either side of C atoms' ALLOW second two marking points from a
			Mark separately		diagram

Qu	esti	on	Expected Answers	Marks	Additional Guidance
	d	ii		6	Please annotate each point scored with a tick
			electron is excited / jumps up energy levels AND (as a result of) light / (UV) radiation / energy / photons \checkmark frequency (absorbed) depends on energy change OR (Δ)E = hv \checkmark		IGNORE references to d-shells must be energy change, not just energy, for example: separation of energy levels / size of gap between levels / difference in energy (between levels) / the excitation energy
			dyes / coloured compounds / 'compounds containing more benzene rings' (<i>AW</i>) <u>absorb</u> in the visible / <u>absorb</u> light ✓ QWC ✓ – see guidance		IGNORE 'absorb energy' QWC scored if second marking point is made in words and first marking point made Place QWC tick by 'pencil' symbol or cross if not awarded
			<i>plus two from:</i> energy levels are closer / energy gap is smaller / excitation energy is smaller when there is: more delocalisation more conjugation more than one benzene ring larger chromophore ✓		max 2 (out of six) if 'emission by dropping down energy levels (<i>AW</i>)' mentioned. Highlight in yellow the words that imply this Must imply 'emission', otherwise IGNORE 'electrons dropping back'
			frequency of uv (radiation / light) is greater than visible / light ORA \checkmark (dyes) transmit / reflect (NOT emit) the complementary colour \checkmark		ALLOW wavelength smaller ALLOW 'complimentary'
	е		one ✓ 6.4 – 8.2 (ppm) ✓	2	
			Mark separately		

Que	estic	n	Expected Answers	Marks	Additional Guidance
	f	i	bromobenzene ✓	1	ALLOW 1 - bromobenzene with or without dash ALLOW 'bromo-benzene' or 'bromo benzene' no other spelling errors
		ii	C_6H_6 + $Br_2 \rightarrow C_6H_5Br$ + $HBr \checkmark$	1	ALLOW skeletal formulae for aromatic compounds IGNORE state symbols IGNORE 'Fe' / 'FeBr ₃ ' / conditions over arrow Any other additions are CON
		III	electrophile is a (partially) positively charged / electron deficient (species) \checkmark (electrophile) accepts a pair of electrons / forms (covalent) bond \checkmark bromine (molecule) is polarised (or diagram) / forms Br ^{δ^+} AND positive end (<i>AW</i>) attacks / forms bond / reacts / substitutes / is electrophile \checkmark	3	ALLOW Br+ (formed) and attacks / is electrophile
			Total	28	

Question		ion	Expected Answers	Marks	Additional Guidance
5	а	i	one COOH group ringed ✓	1	ALLOW ring to cross C-C bond anywhere but not to include next carbon away from COOH
		ii	Any two from: amine ✓ (secondary) amide ✓ imine ✓	2	IGNORE ketone*, carbonyl*, amino* ALLOW alkene ALLOW primary and secondary amines as two groups (but both 'primary' and 'secondary' must be stated to score more than one mark) *extra incorrect groups (apart from these) are each CON to a mark gained
		111		1	Any other arrows are CON ALLOW other unambiguous indications of chiral carbon NB lack of an arrow is 'NR'
	b		H ₂ N H H H H H H H H H H H H H H H H H H H	2	Arrows in excess of two each CON a correct one
	С		it fits into the <u>active/receptor</u> site \checkmark blocks site / 'prevents substrate (<i>AW</i>) from binding' \checkmark	3	Idea of fit needed ('similar shape' and 'binds / bonds to active site' covers this)
			(trimetrexate): has arene ring / is aromatic OR does not have: C=N / double bond / alkene (<i>AW</i>) ✓		IGNORE references to methyl group

Qu	Question		Expected Answers	Marks	Additional Guidance
	d		$CH_3CI / chloromethane (ALLOW methyl chloride) \checkmark$	3	Max 1 for reagents if extra incorrect reagents are included (wrong name is CON to correct formula and vice versa)
			AICl ₃ / aluminium chloride ✓		only accept 'reflux' if one other mark scored
			reflux OR anhydrous conditions OR ionic liquid (solvent) ✓		
	е	i	$HA \implies H^+ + A^- OR$	1	must be equilibrium sign
			$HA + H_2O = H_3O^+ + A^- \checkmark$		NOT [H ⁺] or [A ⁻]
					State symbols, apart from '(aq)' ((<i>l</i>) for water) are CON
	е	11	HA = H + A	1	ALLOW HA + H ₂ O \implies H ₃ O + A
			conjugate acid conjugate base 🗸		
					UR $\Pi A + \Pi_2 U = \Pi_3 U + A$
					ALLOW if arrow, rather than equilibrium sign
			[H ⁺] [A ⁻]/[HA] √	1	Must have square brackets ALLOW multiplication sign
			Lil Kiklind .		(or dot) on top but NOT plus
					NOT signs outside brackets
					ALLOW $[H_3O^+]$ for $[H^+]$ but NOT $[H_2O]$ on bottom
					State symbols, apart from '(aq)' are CON
		iv	(–log 5 x 10 ⁻³) = 2.3 ✓	1	ALLOW more sf than 2.3 if it rounds to 2.3
		v	$[H^+] = \sqrt{(5.0 \times 10^{-3} \times 0.1)} \checkmark (= 2.24 \times 10^{-2})$	2	Must say ' $[H^+]$ =' or ' H^+ =' to score this mark on its own or
					where ecf is considered
			pH = $-\log [H^+] = 1.65/1.7/1.66$ (early rounding to 0.022) \checkmark		Allow ecf from first mark if working or evaluation of $[H^+]$ is
					present and $[H^{\dagger}]$ is smaller than 5 x 10 ⁻²
					ALLOW more sf than 1.65/1.66 if it rounds to 1.65/1.66
					Correct answer with no working scores 2
		vi	concentration of acid at equilibrium = concentration of acid initially	2	
		VI	$(AW) \checkmark$	2	
					Second mark depends on first
			$[H^+]$ or 2.24 x 10 ⁻² compared with 0.1 or compared with [HA] is		
			not negligible (AW) 🗸		

Qu	Question		Expected Answers	Marks	Additional Guidance
	f		$[H^{+}] = 3.98 / 4 / 4.0 \times 10^{-8} \checkmark$ $\frac{[A^{\circ}]}{[HA]} = K_{a} / [H^{+}] = 1.25 / 1.26 / 1.3 \times 10^{5} \checkmark$	2	must say ' $[H^+]$ =' or ' H^+ =' to score this mark on its own or where ecf is considered Allow ecf from first mark if value of $[H^+]$ is present and $[H^+]$ is smaller than 5 x 10 ⁻² ALLOW more sf on ratio (eg 125594/125628) Correct answer with no working scores 2
	g	i	hydrogencarbonate ✓	1	ALLOW 'hydrogen carbonate' IGNORE 'bicarbonate' ALLOW (IV) after name, but no other numbers
		ii	([H ⁺] increases) so equilibrium (position) moves to left / equilibrium moves to form more CO ₂ \checkmark	3	Must be in terms of equilibrium
			excess / reservoir / large concentrations / large amounts of (CO2 and) $HCO_3^-\checkmark$		ALLOW 'salt' or 'A ⁻ ' for HCO ₃ ⁻
			pH (virtually) unchanged (<i>AW</i>) ✓ Mark separately		constancy of pH scores this mark
	h	I	circle (or Na (with or without '+')) surrounded by three or more bent or triangular shapes \checkmark circle shown as '+' (or Na ⁺ shown) and H and O atoms labelled on at least one shape, with at least one H labelled δ + and one O labelled δ - and O pointing to central ion \checkmark	2	$\begin{array}{c} \overset{\delta +}{H} \\ & \overset{\delta -}{O - H} \\ & & & \\ & &$
	h	ii	ion-dipole ✓	1	IGNORE anything else (eg ionic dipole)

Question		ion	Expected Answers		Additional Guidance
	h		Iattice enthalpy / energy (change)√ Na ⁺ (g) + HCO ₃ ⁻ (g) √ Na ⁺ (aq) + HCO ₃ ⁻ (aq) Na ⁺ (aq) + HCO ₃ ⁻ (aq) Image: NaHCO ₃ (s)	3	ALLOW ΔH _{LE} , ΔH _{sol(n)' solution} ALLOW 'enthalpy change of lattice formation' but NOT 'enthalpy change of lattice' ALLOW 'solution enthalpy' IGNORE: • anything after correct answers in the bottom two boxes (eg 'of hydrogencarbonate') • 'sum of' before 'lattice enth' and 'enth of soln' • negative signs in lattice enthalpy box • 'gaseous ions' in top box Please remember to check each time that there is no writing on pages 23 and 24. These will not show as 'Additional objects' but they show up for this question on the 'thumbnails' to the left as 'item 2' and 'item 3'. Annotations made in other questions will not show here. Please ensure that some annotation (default: cross) appears on the top left corner of
			Total	32	pages 23 and 24.

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