

# Markscheme

# November 2015

# Chemistry

## **Standard level**

## Paper 2

14 pages



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## Subject Details: Chemistry SL Paper 2 Markscheme

### **Mark Allocation**

Candidates are required to answer **ALL** questions in Section A **[30 marks]** and **ONE** question in Section B **[20 marks]**. Maximum total = **[50 marks]**.

- **1.** A markscheme often has more marking points than the total allows. This is intentional.
- 2. Each marking point has a separate line and the end is shown by means of a semicolon (;).
- **3.** An alternative answer or wording is indicated in the markscheme by a slash (/). Either wording can be accepted.
- 4. Words in brackets ( ) in the markscheme are not necessary to gain the mark.
- **5.** Words that are <u>underlined</u> are essential for the mark.
- 6. The order of marking points does not have to be as in the markscheme, unless stated otherwise.

## Section A

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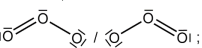
1.	(a)	(i)	$IO_3^-$ to $I_2$ : V/+5 to 0;	
			$I^{-}$ to $I_{2}$ : -I/-1 to 0;	
			Accept change in oxidation number $-5$ and $+1$ .	
			Penalize incorrect notation such as 5+ or 5 once only.	
		(ii)	Oxidizing agent: $IO_3^-$ /iodate and Reducing agent: $I^-$ /iodide;	[1]
	(b)	(i)	1.4 (%);	[1]
			Accept 1 (%).	
		(ii)	systematic;	
			dilute the orange juice; Accept other valid suggestions, eg compare with a standard (showing colour	[2]
			at equivalence) / look at mixture through a yellow filter / add more starch (for a	
			sharper colour change) / filter orange juice (through charcoal). Do not accept repeat the titration or alternative indicator.	
		(iii)	1.44×10 <sup>-5</sup> (mol);	[1]
	(C)	IO <sub>3</sub> -	$: 3C_6H_8O_6 / 1:3$ mole ratio;	
		$(1.44 \times 10^{-5} \times 3 =) 4.32 \times 10^{-5}$ (mol);		
			rd [2] for the correct final answer.	
		Awa	ard <b>[1 max]</b> for "4.80 x $10^{-6}$ (mol)" obtained from reversed ratio,3:1.	
	(d)	$(4.32 \times 10^{-5} \times 176.14 =) 7.61 \times 10^{-3} (g);$		
		Accept $M_r = 176$ and mass = $7.60 \times 10^{-3}$ (g).		

[1]

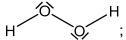
[2]

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2. (a) ।ō̄≡¯Ōı;



The coordinate bond may be represented as an arrow and the formal charges may be shown. Do not accept delocalized structure.



(J) ; [3]

Accept any combination of lines, dots or crosses to represent electron pairs.

- (b) O<sub>2</sub> < H<sub>2</sub>O<sub>2</sub> and O<sub>2</sub> has double bond/bond order of 2 (and H<sub>2</sub>O<sub>2</sub> has single bond/bond order of 1);
  Do not apply ECF from part (a).
- (c) Any value in the range 110° to <120°; Experimental value = 117°. Accept <120°. Do not accept > 109°.

3 negative charge centres/electron domains with 1 lone pair / lone pair-bond (pair) repulsion greater than bond (pair)-bond (pair) repulsion / lone pair occupies more space than bond (pair)/shared pair (so O–O–O angle reduced);

Do not apply ECF in this question.

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(a)  $C_3H_8(g) + 5O_2(g) \rightarrow 3CO_2(g) + 4H_2O(g);$ 3. [1] Ignore state symbols. bonds broken: 2(C-C)/694 + 8(C-H)/3304 + 5(O=O)/2490 / 6488 (kJ); (b) bonds made: 6(C=O)/4476 + 8(O-H)/3712 / 8188 (kJ); (6488 - 8188 =) - 1700 (kJ);[3] Ignore signs in M1 and M2. Award [3] for the correct final answer. Award [2] for +1700 (kJ). Accept values from 2016 data booklet to give 6494 (kJ) for M1, 8528 (kJ) for M2, and -2034 (kJ) for M3. (C)  $3C(s) + 3O_2(g) \rightarrow 3CO_2(g) / 3(-394) / -1182;$  $4H_2(g) + 2O_2(g) \rightarrow 4H_2O(g) / 4(-242) / -968$ ;  $C_{3}H_{8}(g) \rightarrow 3C(s) + 4H_{2}(g) / +104$ ;  $(-1182 + (-968) + 104 =) - 2046 (kJmol^{-1});$ [4] Award [4] for the correct final answer. Award **[3]** for +2046 / 2046 (kJ mol<sup>-1</sup>). part (b) values are based on average (bond enthalpy) values / part (c) values are (d) for specific compounds; [1] ability of an atom to attract (a pair of) electrons in a covalent bond/molecule / 4. (a) (i) ability of an atom to attract a shared pair of electrons; [1] Do not accept nucleus/element instead of atom. do not form bonds/compounds / do not share electrons / have (full/stable) (ii) octet / have full/stable outer shell; [1] Accept (chemically) inert / do not react / stable electron arrangements/ configurations.  $(Li \rightarrow Cs)$  atomic/ionic radius increases; (b) attraction between metal ions and delocalized electrons decreases; Accept metallic bonding gets weaker.  $(F \rightarrow I)$  London/dispersion/instantaneous induced dipole-induced dipole forces increase: Accept vdW/van der Waals' forces for London/dispersion forces. with increasing number of electrons/molar mass/surface area/size of electron cloud; [3 max] Do not accept "with increasing size" or "with increasing mass" only.

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## Section B

5.	(a)	(i)	Increasing the pressure, at constant temperature: decreases; more (gas) molecules/moles on the right / fewer (gas) molecules/moles on the left;	
			Increasing the temperature, at constant pressure: increases; (forward) reaction is endothermic;	[4]
		(ii)	(increasing) temperature <b>and</b> ( <i>K</i> <sub>c</sub> ) increases; Award <b>[0]</b> if both temperature and pressure stated.	[1]
		(iii)	equilibrium reached faster; no change in the concentration of reactants/products/yield (at equilibrium) / position of equilibrium is not affected; rates of forward and reverse reactions increase (equally); reduces activation energy;	
			no change in $K_c$ ;	[3 max]
	(b)	(i)	Reaction A: base <b>and</b> accepts a proton/H <sup>+</sup> ; Accept donates a pair of electrons.	
			<i>Reaction B</i> : acid <b>and</b> donates/loses a proton/H <sup>+</sup> ;	[2]
			Award [1] if base and acid identified correctly without reasons.	
		(ii)	Acid Base	
			Conjugate acid-base pair $1/2$ $H_2CO_3$ and $HCO_3^-$ ;	
			Conjugate acid-base pair $2/1$ HCO <sub>3</sub> <sup>-</sup> and CO <sub>3</sub> <sup>2-</sup> ;	
			Conjugate acid-base pair $1/2$ $H_3O^+$ and $H_2O$ ;	[2 max]
	(C)	(i)	<i>strong acid:</i> (assumed to be) completely/100 % dissociated/ionized <b>and</b> <i>weak acid:</i> partially dissociated/ionized;	[1]
		(ii)	Similarity: bubbling/effervescence/gas / heat/increase in temperature / solid dissolves;	
			<i>Difference:</i> strong acid more vigorous / faster reaction / greater temperature increase; <i>Accept converse statements for weak acid.</i>	[2]
		(iii)	10⁴(:1) / 10 <sup>-1</sup> :10 <sup>-5</sup> / 1:10 <sup>-4</sup> ; Do not accept inverse ratio, 1:10⁴.	[1]

[2]

(d) (i) Na **and** Mg: basic Al: amphoteric Do not accept amphiprotic.

> Si **to** Cl: acidic Ar: no oxide ;; Award **[2]** for three or four correct, award **[1]** for two correct. Award **[1]** for stating oxides become more acidic towards the right/chlorine or more basic towards the left/sodium. Do not penalize incorrect formulas of oxides.

(ii)  $Na_2O(s) + H_2O(l) \rightarrow 2NaOH(aq) / Na_2O(s) + 2HCl(aq) \rightarrow 2NaCl(aq) + H_2O(l);$ Accept a correct equation with any acid or acidic oxide.

 $SO_3(l) + H_2O(l) \rightarrow H_2SO_4(aq) / SO_3(l) + 2NaOH(aq) \rightarrow Na_2SO_4(aq) + H_2O(l);$  [2] Accept a correct equation with any metal hydroxide, metal oxide, metal carbonate or metal hydrogen carbonate. Do not accept equation with SO<sub>2</sub>.

Ignore state symbols. Accept ionic equations for M1 and/or M2. 6.

 (a) (i) use of colorimeter/colorimetry; measure change/decrease in intensity of (purple) colour; recording of colour intensity at regular time intervals / recording time needed for colour to disappear; calibration curve with known concentration; Accept any three points.

### OR

use of (analytical) balance/scale; change/decrease in mass of reaction mixture; recording of mass at regular time intervals / recording time needed for mass to become constant;

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#### OR

use of gas syringe / inverted gas tube; change/increase volume of carbon dioxide; recording of volume at regular time intervals / recording time needed for volume to become constant;

### OR

use of pH meter/probe; change/increase in pH of reaction mixture; recording of pH at regular time intervals / recording time needed for pH to become constant;

### OR

use of conductivity meter/probe; change/decrease in conductivity of reaction mixture; recording of conductivity at regular time intervals / recording time needed for conductivity to become constant;

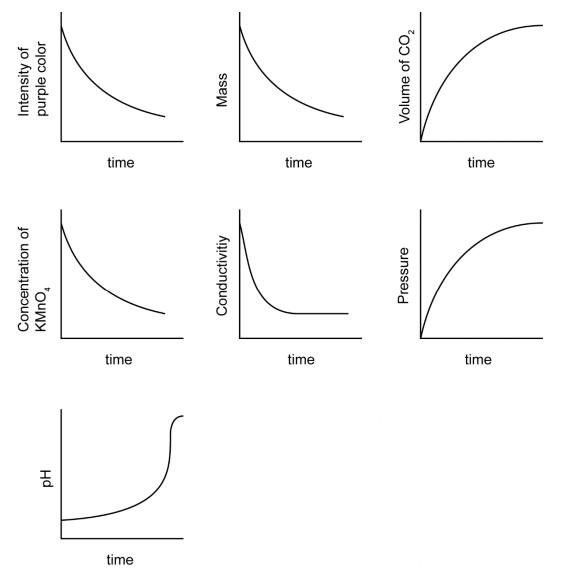
### OR

use of pressure sensor; change/increase in pressure of gas; recording of pressure at regular time intervals / recording time needed for pressure to become constant; (ii) axes labelled correctly; Units not required for axes.

> correct shape of curve; Curve must have a slope of a gradually decreasing magnitude (except the pH curve) but does not have to show the end of the reaction/plateau. Accept curve to start or end at zero or non-zero. Accept slight initial horizontal line for mass, volume and pressure curves due to slight solubility of  $CO_2$  released. Accept zero-order graphs. M2 can only be scored if M1 correct.

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Examples of graph:

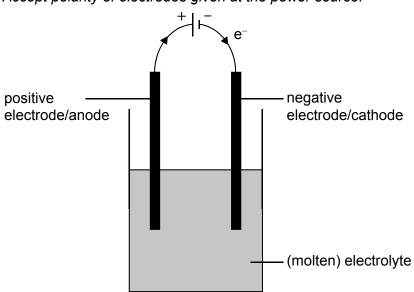


- (iii) rate = slope/gradient of tangent;
- (iv) (rate increases due to) increase in (average) <u>kinetic</u> energy/speed of the particles; increase in frequency of collisions/collisions per unit time; greater proportion/number of particles have energy  $\geq E_a$ ;

[2]

[1]

- (b) (i) Pb < Ni < Fe < Zn ;; Award [2] for the correct order. Award [1] for Zn > Fe > Ni > Pb as metals not listed in order of increasing reactivity. Award [1] if one error in the order.
  - (ii) Pb<sup>2+</sup> / lead(II) (ions); Do not accept Pb/lead.
- (c) power source and direction of e<sup>-</sup> movement;
  labelled +/positive electrode/anode and -/negative electrode/cathode and (molten) electrolyte/NiBr<sub>2</sub> (l);
  Accept polarity of electrodes given at the power source.



Negative electrode (cathode) :  $Ni^{2+} + 2e^- \rightarrow Ni(l)$ ; Positive electrode (anode):  $2Br^- \rightarrow Br_2(g) + 2e^- / Br^- \rightarrow \frac{1}{2}Br_2(g) + e^-$ ;

Award **[1 max]** for M3 and M4 if equations are given at wrong electrodes. Ignore state symbols and reversible sign. Allow e instead of e<sup>-</sup>.

- (d) (i) ionization and (bombardment) by high energy/fast moving electrons/electron gun (to form positive ions); acceleration and passing through electric field/potential difference/oppositely charged plates; deflection and passing through magnetic field/electromagnet; [3] Award [1] for naming 3 processes (ionization, acceleration, deflection) in the correct order with incorrect details.
  - to avoid collision with other particles (in the atmosphere) / allows ions to pass through unhindered (by air molecules);
    *Reference must be made to interaction with other particles.*

[1]

[2]

[4]

7. (a) (i) 
$$\operatorname{RBr}(l) + \operatorname{NaOH}(aq) \rightarrow \operatorname{ROH}(aq) + \operatorname{NaBr}(aq) / \operatorname{RBr}(l) + \operatorname{OH}^{-}(aq) \rightarrow \operatorname{ROH}(aq) + \operatorname{Br}^{-}(aq);$$
 [1]  
Ignore state symbols.

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(ii) 
$$(1.35 \times 10^{-2} - 7.36 \times 10^{-3} =) 6.14 \times 10^{-3} / 6.1 \times 10^{-3}$$
 (mol); [1]

(iii) (molar mass = 
$$\frac{0.842}{6.14 \times 10^{-3}}$$
 =) 137 (g mol<sup>-1</sup>); [1]

Accept 138.

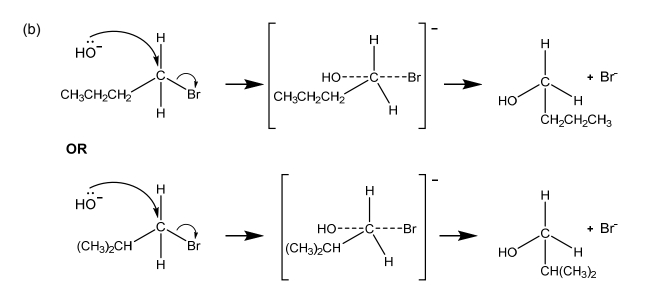
- (iv)  $(137 - 80 = 57 \text{ which corresponds to } C_4H_9, \text{ hence molecular formula}) C_4H_9Br;$ [1] Do not accept ECF from 7a(iii) for an impossible molecular formula, such as  $C_4H_{10}Br$ . Accept correct structural formula of one of the isomers as the molecular formula.
- (v) CH<sub>3</sub>CH<sub>2</sub>CH<sub>2</sub>CH<sub>2</sub>Br and primary; (CH<sub>3</sub>)<sub>2</sub>CHCH<sub>2</sub>Br and primary; CH<sub>3</sub>CHBrCH<sub>2</sub>CH<sub>3</sub> and secondary; (CH<sub>3</sub>)<sub>3</sub>CBr and tertiary; If primary, secondary or tertiary not stated, award [3] for four correct, [2] for three correct and [1] for two correct structural formulas. Penalize missing hydrogens once only. Accept either full or condensed structural formulas.

If C<sub>5</sub>H<sub>11</sub>Br was used, accept any correct structural formulas. CH<sub>3</sub>CH<sub>2</sub>CH<sub>2</sub>CH<sub>2</sub>CH<sub>2</sub>Br and primary; (CH<sub>3</sub>)<sub>2</sub>CHCH<sub>2</sub>CH<sub>2</sub>Br **and** primary; CH<sub>3</sub>CH<sub>2</sub>CH(CH<sub>3</sub>)CH<sub>2</sub>Br **and** primary; (CH<sub>3</sub>)<sub>3</sub>CCH<sub>2</sub>Br and primary: CH<sub>3</sub>CHBrCH<sub>2</sub>CH<sub>2</sub>CH<sub>3</sub> and secondary; CH<sub>3</sub>CH<sub>2</sub>CHBrCH<sub>2</sub>CH<sub>3</sub> and secondary; CH<sub>3</sub>CHBrCH(CH<sub>3</sub>)<sub>2</sub> and secondary; CH<sub>3</sub>CH<sub>2</sub>C(CH<sub>3</sub>)<sub>2</sub>Br and tertiary; [4] If primary, secondary or tertiary not stated, award [3] for four correct, [2] for three correct and [1] for two correct structural formulas. Penalize missing hydrogens once only. Accept either full or condensed structural formulas.

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curly arrow going from lone pair/negative charge on O in HO<sup>-</sup> to C; Do not allow curly arrow originating on H in HO<sup>-</sup> but do not penalize OH<sup>-</sup>.

curly arrow showing Br leaving; Accept curly arrow either going from bond between C and Br to Br in bromobutane or in the transition state.

representation of transition state showing negative charge, square brackets and partial bonds; Do not penalize if OH and Br are not at 180° to each other.

formation of products CH<sub>3</sub>CH<sub>2</sub>CH<sub>2</sub>CH<sub>2</sub>OH/(CH<sub>3</sub>)<sub>2</sub>CHCH<sub>2</sub>OH and Br<sup>-</sup>;

Penalize incorrect side-chain, missing hydrogens and/or incorrect bond linkages (eg OH–C instead of HO–C) only once in this question. Do not penalize missing hydrogens if already penalized in part 7 (a)(v). Award **[2 max]** if  $S_N$ 1 mechanism is given. [4]

[2]

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- (c) (i) CH<sub>3</sub>CH<sub>2</sub>CH<sub>3</sub> < CH<sub>3</sub>CHO < CH<sub>3</sub>CH<sub>2</sub>OH < CH<sub>3</sub>COOH;; Award [2] for correct order. Award [1] if one error in the order. Award [1] for CH<sub>3</sub>COOH > CH<sub>3</sub>CH<sub>2</sub>OH > CH<sub>3</sub>CHO > CH<sub>3</sub>CH<sub>2</sub>CH<sub>3</sub> as compounds are not listed in order of increasing boiling point.
  - (ii) CH<sub>3</sub>CH<sub>2</sub>CH<sub>3</sub> London/dispersion/instantaneous induced dipole-induced dipole forces
    CH<sub>3</sub>CHO dipole-dipole forces (and London/dispersion forces)
    CH<sub>3</sub>CH<sub>2</sub>OH H-bonding (and dipole-dipole and London/dispersion forces)
    CH<sub>3</sub>COOH H-bonding (and dipole-dipole and London/dispersion forces) ;;
    Award [2] for all four correct.
    Award [1] for two or three correct.

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H-bonding strongest / London/dispersion forces weakest / dipole-dipole stronger than London/dispersion / dipole-dipole weaker than H-bonding;

Accept vdW/van der Waals' forces for London/dispersion forces.

CH<sub>3</sub>COOH forms more/stronger H-bonds than CH<sub>3</sub>CH<sub>2</sub>OH / CH<sub>3</sub>COOH is more polar than CH<sub>3</sub>CH<sub>2</sub>OH; [4] Accept CH<sub>3</sub>COOH has more electrons/higher molar mass than CH<sub>3</sub>CH<sub>2</sub>OH.

(iii) CH<sub>3</sub>COOH; Accept either full or condensed structural formula.

orange to green;