

**Assessment Schedule – 2019****Chemistry: Demonstrate understanding of bonding, structure, properties and energy changes (91164)****Evidence Statement**

Q	Evidence				Achievement	Merit	Excellence																
ONE (a)	<table border="1"> <thead> <tr> <th>Solid</th> <th>Type of solid</th> <th>Type of particle</th> <th>Attractive forces between particles</th> </tr> </thead> <tbody> <tr> <td>Na(s) (sodium)</td> <td>metal / metallic</td> <td>atoms /cations (or metal nuclei) in sea of delocalised electrons</td> <td>metallic bond</td> </tr> <tr> <td>NaI(s) (sodium iodide)</td> <td>ionic</td> <td>ions</td> <td>ionic bond / electrostatic attraction between (oppositely charged) ions</td> </tr> <tr> <td>I<sub>2</sub>(s) (iodine)</td> <td>(covalent) molecular</td> <td>molecules</td> <td>(weak) intermolecular (forces)</td> </tr> </tbody> </table>	Solid	Type of solid	Type of particle	Attractive forces between particles	Na(s) (sodium)	metal / metallic	atoms /cations (or metal nuclei) in sea of delocalised electrons	metallic bond	NaI(s) (sodium iodide)	ionic	ions	ionic bond / electrostatic attraction between (oppositely charged) ions	I <sub>2</sub> (s) (iodine)	(covalent) molecular	molecules	(weak) intermolecular (forces)				<ul style="list-style-type: none"> <li>Two rows or two columns correct.</li> </ul>		
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(b)	<p>Sodium is a metallic solid made up of atoms in 3D lattice held together by non-directional metallic bonds (or cations non-directionally electrostatically attracted to a surrounding sea of electrons). When a force (or pressure) is applied, the atoms / layers can move without breaking / disrupting these non-directional bonds; thus the structure can change shape.</p> <p>NaI is made up of alternating positive ions / Na<sup>+</sup> ions, and negative ions / I<sup>-</sup> ions, ionically bonded in a 3D lattice. NaI is not malleable because if pressure is applied to an ionic lattice, it forces ions with the same charge next to each other; they repel each other and break the structure.</p>				<ul style="list-style-type: none"> <li>Describes structure of sodium.</li> </ul> OR Recognises metallic bonding as non-directional.	<ul style="list-style-type: none"> <li>Describes malleability and Links this to non-directional metallic bonding of sodium.</li> </ul>	<ul style="list-style-type: none"> <li>Describes brittleness of sodium iodide and links it to directional ionic bonding / repulsion of like charged ions.</li> </ul>	<ul style="list-style-type: none"> <li>Comprehensively explains malleability of sodium and brittleness of sodium iodide.</li> </ul>															

(c)	<p>Iodine is a non-polar (covalent) molecular substance made up of I<sub>2</sub> molecules held together by weak intermolecular forces.</p> <p>Iodine is soluble in cyclohexane, but does not easily dissolve in water.</p> <p>For iodine in water, the iodine-water attractions are not strong enough to overcome both the iodine-iodine / solute-solute and the strong water-water / solvent-solvent attractions.</p> <p>For iodine in cyclohexane, the iodine-cyclohexane attractions are strong enough to overcome iodine-iodine / solute-solute and cyclohexane-cyclohexane / solvent-solvent attractions because all attractive forces are similar (nonpolar).</p>	<ul style="list-style-type: none"> <li>• Recognises I<sub>2</sub> as a non-polar molecule.</li> <li>• Identifies iodine as (more) soluble in cyclohexane and insoluble/less soluble in water.</li> </ul>	<ul style="list-style-type: none"> <li>• Links attractions (or lack of) of water OR cyclohexane for non-polar iodine to solubility.</li> </ul>	<ul style="list-style-type: none"> <li>• Comprehensively explains iodine's solubility in cyclohexane and insolubility (low solubility) in water linking polarity, strength of attraction and overcoming / not overcoming existing bonding within the solvent or solid.</li> </ul>
(d)	<p>The melting of ice is endothermic, as (intermolecular) bonds are being broken as water changes from solid to liquid. This requires energy to be absorbed from the surroundings/the drink. This causes the temperature of the drink to decrease.</p>	<ul style="list-style-type: none"> <li>• Identifies the melting of ice is endothermic / absorbs or requires energy / breaks bonds.</li> </ul>	<ul style="list-style-type: none"> <li>• Links bond breaking to energy being absorbed from the surroundings / drink / environment.</li> </ul>	<ul style="list-style-type: none"> <li>• Full explanation that links to the cooling of the drink.</li> </ul>

<b>NØ</b>	<b>N1</b>	<b>N2</b>	<b>A3</b>	<b>A4</b>	<b>M5</b>	<b>M6</b>	<b>E7</b>	<b>E8</b>
No response; no relevant evidence.	1a	2a	3a	4a	2m	3m	2e with up to one minor error/omission	3e with up to one minor error/omission

Q	Evidence				Achievement	Merit	Excellence												
TWO (a)(i)	<table border="1"> <thead> <tr> <th>Molecule</th> <th>CH<sub>4</sub></th> <th>NCl<sub>3</sub></th> <th>OF<sub>2</sub></th> </tr> </thead> <tbody> <tr> <td>Lewis Structure</td> <td> <pre>       H         H - C - H               H           </pre> </td> <td> <pre>       .. .. ..       :Cl- N -Cl:                       :Cl:           ..           </pre> </td> <td> <pre>       .. ..       :F: O :F:       .. ..           </pre> </td> </tr> <tr> <td>Name of Shape</td> <td>tetrahedral</td> <td>trigonal pyramidal</td> <td>bent / v-shaped / angular</td> </tr> </tbody> </table>				Molecule	CH <sub>4</sub>	NCl <sub>3</sub>	OF <sub>2</sub>	Lewis Structure	<pre>       H         H - C - H               H           </pre>	<pre>       .. .. ..       :Cl- N -Cl:                       :Cl:           ..           </pre>	<pre>       .. ..       :F: O :F:       .. ..           </pre>	Name of Shape	tetrahedral	trigonal pyramidal	bent / v-shaped / angular	<ul style="list-style-type: none"> <li>Two Lewis structures correct.</li> </ul> OR <ul style="list-style-type: none"> <li>Two shapes correct.</li> </ul>		
Molecule	CH <sub>4</sub>	NCl <sub>3</sub>	OF <sub>2</sub>																
Lewis Structure	<pre>       H         H - C - H               H           </pre>	<pre>       .. .. ..       :Cl- N -Cl:                       :Cl:           ..           </pre>	<pre>       .. ..       :F: O :F:       .. ..           </pre>																
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(ii)	<p>Bond angle is determined by the number of electron density regions around the central atom, which are arranged into a position to minimise repulsion by having maximum separation.</p> <p>All molecules have 4 electron density regions / areas of negative charge around the central atom which arrange with maximum separation into a tetrahedral shape / geometry with a bond angle of (approx.) 109.5° / 109°.</p> <p>In CH<sub>4</sub> all of the electron pairs are bonded, and so the shape of the molecule is also tetrahedral.</p> <p>In NCl<sub>3</sub> three of the electron pairs are bonded and one is non-bonding. The observed shape of the molecule is trigonal pyramidal.</p> <p>In OF<sub>2</sub>, due to the presence of two non-bonding pairs of electrons / regions (or two bonding regions) on the central atom, OF<sub>2</sub> has an observed shape that is bent / v-shaped / angular.</p>				<ul style="list-style-type: none"> <li>Identifies the 4 regions of negative charge for each molecule.</li> </ul> OR <ul style="list-style-type: none"> <li>Identifies the bonding and non-bonding pairs for one molecule.</li> </ul> <ul style="list-style-type: none"> <li>Recognises that electrons density regions are arranged in a position of maximum separation / minimal repulsion.</li> </ul>	<ul style="list-style-type: none"> <li>Links number of areas of negative charge (including bonding / nonbonding) around the central atom to the shape of TWO molecules using minimise repulsion / maximum separation.</li> </ul> OR <ul style="list-style-type: none"> <li>Links four regions on the central atom for two or more molecules to a bond angle of 109.5° using minimise repulsion / maximum separation.</li> </ul>	<ul style="list-style-type: none"> <li>Justifies the shape of all molecules by referring to all factors that influence shape and bond angle.</li> </ul>												
(b)(i)  (ii)	<p>CHCl<sub>3</sub> is polar. NH<sub>3</sub> is polar.</p> <p>In CHCl<sub>3</sub>, there are two types of bond, C–H and C–Cl, each polar, due to the difference in electronegativity between C and H and C and Cl atoms. These dipoles have different polarities / sizes as H and Cl have different electronegativities. (Despite the tetrahedral arrangement appearing symmetrical) the different (sized) bond dipoles do not cancel each other out, so CHCl<sub>3</sub> is polar.</p> <p>In NH<sub>3</sub>, the three N–H bonds are polar, i.e. have a dipole, due to the difference in electronegativity between N and H atoms. These (equally sized) dipoles are arranged in a non-symmetrical trigonal pyramidal shape, resulting in the bond dipoles not cancelling each other out, so NH<sub>3</sub> is polar.</p>				<ul style="list-style-type: none"> <li>Identifies polarity of both molecules.</li> </ul> <ul style="list-style-type: none"> <li>Identifies bonded atoms have different electronegativity (values).</li> </ul>	<ul style="list-style-type: none"> <li>Links bond polarity to electronegativity differences between bonded atoms for all bonds in one molecule.</li> </ul> OR <ul style="list-style-type: none"> <li>Uses lack of symmetry for NH<sub>3</sub> OR differing bond dipoles for CHCl<sub>3</sub> to link molecule polarity to dipoles not cancelling in one molecule.</li> </ul>	<ul style="list-style-type: none"> <li>Justifies polarity of both molecules by referring to differences in electronegativity, dipoles, and non-symmetrical arrangement of NH<sub>3</sub> dipoles.</li> </ul>												

(c)	<p>Bond breaking</p> $2 \times \text{C} - \text{C} = 348 \times 2 = 696$ $8 \times \text{C} - \text{H} = 413 \times 8 = 3304$ $5 \times \text{O} = \text{O} = 495 \times 5 = 2475$ <p style="text-align: right;">Total = <b>6475</b></p> <p>Bond making</p> $8 \times \text{O} - \text{H} = 463 \times 8 = 3704$ $6 \times \text{C} = \text{O} = 6x$ $\Delta_r H = \Sigma \text{Bond energies (bonds broken)} - \Sigma \text{Bond energies (bonds formed)}$ $6475 - 3704 - 6x = -2056 \text{ kJ mol}^{-1}$ $6x = +2056 + 6475 - 3704 = 4827$ $x = 805 \text{ kJ mol}^{-1}$	<ul style="list-style-type: none"> <li>• Correctly calculates total bonds broken.</li> </ul> <p>OR</p> <ul style="list-style-type: none"> <li>• Correctly identifies bonds made in a formula.</li> </ul>	<ul style="list-style-type: none"> <li>• Correct process with minor error / omission.</li> </ul>	<ul style="list-style-type: none"> <li>• Correct answer with unit.</li> </ul>				
NØ	N1	N2	A3	A4	M5	M6	E7	E8
No response; no relevant evidence.	1a	2a	3a	4a	2m	3m	2e	3e

Q	Evidence	Achievement	Merit	Excellence
THREE (a)(i) (ii)	<p>Exothermic as <math>\Delta_r H</math> is negative.</p> <p>To be able to conduct electricity, there needs to be mobile/free moving charged particles. Graphite, C(s), is an extended covalent network solid. Each carbon atom is covalently bonded to 3 other carbon atoms in hexagonal layers. This leaves one delocalised electron per carbon atom that is mobile and able to carry a charge, so graphite conducts electricity.</p> <p>Carbon dioxide is a covalent molecule. The molecules are held together by weak intermolecular forces, so it is a gas at room temperature. There are no free moving ions or electrons in their structure. Therefore, it can't conduct electricity.</p>	<ul style="list-style-type: none"> <li>• Correct term with reason in (i). accept energy/heat is lost OR <math>\Delta H</math>/enthalpy change is negative.</li> <li>• Recognises mobile charged particles needed for conductivity.</li> </ul> <p>OR</p> <p>Identifies graphite is a covalent network made of atoms and CO<sub>2</sub>(g) consists of molecules.</p>	<ul style="list-style-type: none"> <li>• Links 2D network/layered covalent <b>network</b> structure of graphite to free moving electrons which conduct electricity OR molecules of CO<sub>2</sub> to being uncharged / not having a free moving charged particle and therefore not conducting electricity.</li> </ul>	<ul style="list-style-type: none"> <li>• Justifies the conductivity of graphite and carbon dioxide in terms of structure and bonding providing / not providing free moving charged particles.</li> </ul>
(b)(i)  (ii)	$n(\text{O}_2(\text{g})) = \frac{1804.5}{1203} = 1.5 \text{ moles}$ $m = n \times M = 1.5 \times 32 = 48.0 \text{ g}$ $n = \frac{m}{M} = \frac{100}{40.3} = 2.48$ $\text{Energy} = \Delta_r H \times n = \frac{-1203}{2} \times 2.48 = -1492.5 \text{ kJ} / -1493 \text{ kJ}$ <p>-1490 kJ (3 sf)</p> <p>(either positive or negative values accepted)</p>	<ul style="list-style-type: none"> <li>• ONE step of calculation correct.</li> <li>• ONE step of calculation correct.</li> </ul>	<ul style="list-style-type: none"> <li>• Correct answer.</li> <li>• Correct answer.</li> </ul>	<p>BOTH correct answers with units.</p>

(c)	$65\,000 \times 60 = 3\,900\,000 \text{ g}$ $n(\text{Al}) = \frac{3\,900\,000}{27} = 144\,444 \text{ moles}$ $\Delta_r H = \frac{144\,444}{4} \times 3350$ $= 121\,000\,000 \text{ kJ} = 1.21 \times 10^8 \text{ kJ (rounded to 3sf)}$ <p>OR</p> $\frac{65000}{27} = 2407 \text{ moles}$ $\Delta_r H = \frac{2407}{4} \times 3350 = 2\,015\,862 \text{ kJ} \times 60$ $= 121\,000\,000 \text{ kJ} = 1.21 \times 10^8 \text{ kJ (rounded to 3sf)}$	<ul style="list-style-type: none"> <li>• ONE step of process correct.</li> </ul>	<ul style="list-style-type: none"> <li>• Process correct with minor error.</li> </ul>	<ul style="list-style-type: none"> <li>• Calculates energy with units (kJ or kJ h<sup>-1</sup>) and 3 sf.</li> </ul>
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NØ	N1	N2	A3	A4	M5	M6	E7	E8
No response; no relevant evidence.	1a	2a	3a	4a	2m	3m	2e	3e

### Cut Scores

Not Achieved	Achievement	Achievement with Merit	Achievement with Excellence
0 – 7	8 – 13	14 – 18	19 – 24