STRUCTURE & BONDING

Introduction: The physical properties (boiling point, conductivity, strength) of a substance depend on its structure and type of bonding. Bonding determines the structure.

TYPES OF BOND

CHEMICAL BONDS (strong bonds)
- IONIC (or electrovalent)
- COVALENT
- DATIVE COVALENT (or COORDINATE)
- METALLIC

PHYSICAL BONDS (weak bonds)
- induced dipole-dipole interactions (London forces)
- permanent dipole-dipole interactions (both the above are examples of van der Waals’ forces)
- hydrogen bonds

FORMATION OF IONS FROM ATOMS

Positive ions
- known as cations
- formed when electrons are removed from atoms
- are smaller than the original atom
- the energy associated with the process is known as the ionisation energy (IE).

1st I.E.
The energy required to remove one mole of electrons (to infinity) from one mole of gaseous atoms to form one mole of gaseous positive ions.

\[ \text{Na}(g) \longrightarrow \text{Na}^+(g) + e^- \quad \text{or} \quad \text{Mg}(g) \longrightarrow \text{Mg}^+(g) + e^- \]

There are as many ionisation energy steps as there are electrons in the atom.

2nd I.E.
\[ \text{Mg}^+(g) \longrightarrow \text{Mg}^{2+}(g) + e^- \quad \text{and so on} \]

Notes
- successive ionisation energies get larger as the proton : electron ratio increases.
- big jumps in value occur when electrons are removed from shells nearer the nucleus - less shielding so more energy is needed to overcome the attraction.

\[ 1st \text{ I.E.} \quad 500 \text{kJmol}^{-1} \quad 2nd \text{ I.E.} \quad 900 \text{kJmol}^{-1} \quad 3rd \text{ I.E.} \quad 6000 \text{kJmol}^{-1} \]

The 3rd electron must have been in a shell nearer the nucleus - In Group 2

- if the IE values are very high, covalent bonding is favoured (e.g. beryllium).
**Negative ions**
- known as **anions**
- larger than the original atom due to electron repulsion in outer shell
- formed when electrons are added to atoms
- energy is released as the nucleus pulls in an electron
- this energy is the **electron affinity**.

**Electron Affinity**

The energy change when one mole of gaseous atoms acquires one mole of electrons (from infinity) to form one mole of gaseous negative ions.

\[ \text{e.g. } \text{Cl}(g) + e^- \rightarrow \text{Cl}^-(g) \text{ and } \text{O}(g) + e^- \rightarrow \text{O}^-(g) \]

The greater the effective nuclear charge (ENC) the easier an electron is pulled in.

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**Q.1** Write out equations representing the ....

- 1st I.E. of Li
- 1st I.E. of Al
- 1st I.E. of F
- 2nd I.E. of Na
- 2nd I.E. of F
- 3rd I.E. of Li
- 4th I.E. of Al
- 21st I.E. of Rb

---

**Q.2** Write out equations representing the ....

- 1st E.A. of Br
- 2nd E.A. of Br
- 1st E.A. of N

---

**Q.3** In which group would you find elements with the following successive I.E.’s ?

- 577 1820 2740 11600 14800
- 418 3070 4600 5860 7990
- 736 1450 7740 10500 13600
THE IONIC (OR ELECTROVALENT) BOND
‘An electrostatic attraction between positive and negative ions’

Formation
Ionic bonds tend to be formed between elements whose atoms need to “lose” electrons to gain the nearest noble gas electronic configuration (ngeom) and those needing to gain electrons. Electrons are transferred from one atom to the other.

**Sodium** (Na) → Na⁺ + e⁻ and **Chloride** (Cl) → Cl⁻ + e⁻

```
| Sodium | 1s² 2s² 2p⁶ 3s¹ | 1s² 2s² 2p⁶ |
| Chloride | 1s² 2s² 2p⁶ 3s² 3p⁵ | 1s² 2s² 2p⁶ 3s² 3p⁶ |
```

or
```
| Sodium | 2,8,1 | 2,8 |
| Chloride | 2,8,7 | 2,8,8 |
```

- 1 electron is transferred from the 3s orbital of sodium to the 3p orbital of chlorine
- both species end up with an ‘octet’ of electrons in their outer shell
- the resulting ions are held together in a crystal lattice by electrostatic attraction

**MgCl₂** Because magnesium atoms have two outer shell electrons they can combine with two chlorine atoms by the transfer of one electron to each atom to form one Mg²⁺ and two Cl⁻ ions

```
| Mg (2,8,2) | 2 x Cl (2,8,7) | Mg²⁺ | 2 x Cl⁻ |
```

**Q.4** Show how the following combine to form ionic compounds.

a) Na and O  b) Mg and O  c) Mg and F  d) Al and O
Predicting the charge on an ion

**Simple ions**

Can be predicted from their position in the Periodic Table

<table>
<thead>
<tr>
<th>Group</th>
<th>electrons in outer shell</th>
<th>charge on ion</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>1</td>
<td>+</td>
</tr>
<tr>
<td>2</td>
<td>2</td>
<td>2+</td>
</tr>
<tr>
<td>3</td>
<td>3</td>
<td>3+</td>
</tr>
<tr>
<td>6</td>
<td>6</td>
<td>2-</td>
</tr>
<tr>
<td>7</td>
<td>7</td>
<td>-</td>
</tr>
</tbody>
</table>

**d block elements**

Transition elements (eg iron) can have more than one ion. The use of a Roman numeral identifies which ion is present.

iron(II) $\text{Fe}^{2+}$
iron(III) $\text{Fe}^{3+}$

**Complex ions**

Some groups possess a charge.

- nitrate $\text{NO}_3^-$
- sulphate $\text{SO}_4^{2-}$
- carbonate $\text{CO}_3^{2-}$
- ammonium $\text{NH}_4^+$

**SOME COMMON IONS**

| 1 | hydrogen  | H$^+$  | chloride  | Cl$^-$  |
|   | sodium    | Na$^+$ | bromide   | Br$^-$  |
|   | potassium | K$^+$  | iodide    | I$^-$   |
|   | lithium   | Li$^+$ | hydroxide | OH$^-$  |
|   | rubidium  | Rb$^+$ | nitrate   | NO$_3^-$|
|   | caesium   | Cs$^+$ | nitrite   | NO$_2^-$|
|   | copper(I) | Cu$^+$ | hydrogencarbonate | HCO$_3^-$ |
|   | silver(I) | Ag$^+$ | hydrogen sulphate | HSO$_4^-$ |
|   | ammonium  | NH$_4^+$ |                                      |
| 2 | calcium   | Ca$^{2+}$ | sulphate | SO$_4^{2-}$ |
|   | barium    | Ba$^{2+}$ | sulphite | SO$_3^{2-}$ |
|   | magnesium | Mg$^{2+}$ | sulphide | S$^{2-}$ |
|   | zinc      | Zn$^{2+}$ | oxide | O$^{2-}$ |
|   | iron(II)  | Fe$^{2+}$ | carbonate | CO$_3^{2-}$ |
|   | cobalt    | Co$^{2+}$ | copper(II) | Cu$^{2+}$ |
|   | manganese(II) | Mn$^{2+}$ |                                      |
| 3 | aluminium | Al$^{3+}$ | phosphate | PO$_4^{3-}$ |
|   | iron(III) | Fe$^{3+}$ |                                      |
GIANT IONIC LATTICES

*bonding*  • oppositely charged ions held in a regular 3-d lattice by electrostatic attraction
• ions pack together in the most efficient way so there is little wasted space
• the arrangement of ions in a lattice depends on the relative sizes of the ions

\[
\begin{align*}
\text{NaCl} & \quad \begin{array}{c}
\text{Na}^+ \\
\text{Cl}^-
\end{array} \\
\text{CsCl} & \quad \begin{array}{c}
\text{Cs}^+ \\
\text{Cl}^-
\end{array}
\end{align*}
\]

The Na\(^+\) ion is small enough relative to the Cl\(^-\) ion to fit in the spaces so that both ions occur in every plane.

Each Na\(^+\) is surrounded by 6 Cl\(^-\) (co-ordination number = 6) and each Cl\(^-\) is surrounded by 6 Na\(^+\) (co-ordination number = 6).

**Physical properties of ionic compounds**

*melting pt*  Very high  A large amount of energy must be put in to overcome the strong electrostatic attractions and separate the ions.

*strength*  Very brittle  Any dislocation leads to layers moving and similarly charged ions being next to each other. The repulsion splits the crystal.

*electrical*  • do not conduct electricity when solid - ions are held strongly in the lattice
• conduct electricity when molten or in aqueous solution - the ions become mobile and conduction takes place.

*solubility*  • insoluble in non-polar solvents
• soluble in water as it is a polar solvent and stabilises the separated ions
• energy is needed to overcome the electrostatic attraction and separate the ions
• stability is achieved by polar water molecules surrounding the ions

*Diagram*
COVALENT BONDING

**Definition**
- consists of a **shared pair of electrons**; each atom supplies one electron
- atoms are held because their nuclei are attracted to the shared electrons

- **Average bond enthalpy** is a measurement of the strength of a covalent bond; the stronger the bond, the higher its value (see later work)

**Formation**
- between atoms of the same element; (N₂, O₂, diamond and graphite)
- between atoms of different elements on RHS of the periodic table; (CO₂, SO₂).
- when one of the elements is in the middle of the table; (e.g. C, Si)
- head-of-the-group elements with high ionisation energies, (e.g. Be in BeCl₂)

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**METHANE**
- each needs one electron
- needs four electrons
- four covalent single bonds are formed

**WATER**
- atoms share electrons in order to complete their ‘octet’ of electrons
- some don’t achieve an ‘octet’ as they haven’t enough electrons - Al in AlCl₃
- others share only some - if they share all their ‘octet’ is exceed - NH₃ and H₂O
- atoms of elements in the 3rd period onwards can exceed their ‘octet’ because they are not restricted to eight electrons in their ‘outer shell’ - S in SF₆

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**Q.5** Show how the covalent bonding is arranged in the following molecules

<p>| | |</p>
<table>
<thead>
<tr>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>a)</td>
<td>H₂</td>
</tr>
<tr>
<td>b)</td>
<td>Cl₂</td>
</tr>
<tr>
<td>c)</td>
<td>O₂</td>
</tr>
<tr>
<td>d)</td>
<td>N₂</td>
</tr>
<tr>
<td>e)</td>
<td>NH₃</td>
</tr>
<tr>
<td>f)</td>
<td>SiCl₄</td>
</tr>
<tr>
<td>g)</td>
<td>BF₃</td>
</tr>
<tr>
<td>h)</td>
<td>SF₆</td>
</tr>
<tr>
<td>i)</td>
<td>PCl₅</td>
</tr>
<tr>
<td>j)</td>
<td>CO₂</td>
</tr>
</tbody>
</table>
## Structures containing covalent bonds

### 1. Simple molecules

- **bonding**: Atoms are joined together **within** the molecule by covalent bonds.
- **electrical**: Don’t **conduct** electricity as they have no mobile ions or electrons.
- **solubility**: Tend to be more soluble in organic solvents than in water; some are **hydrolysed**
- **boiling pt**: Low - the forces between molecules (intermolecular forces) are weak (known as **van der Waals forces** - see below)

Attractions between molecules increases as the molecules get more electrons.

<table>
<thead>
<tr>
<th></th>
<th>CH₄</th>
<th>C₂H₆</th>
<th>C₃H₈</th>
</tr>
</thead>
<tbody>
<tr>
<td>e.g.</td>
<td>-161°C</td>
<td>- 88°C</td>
<td>-42°C</td>
</tr>
</tbody>
</table>

as forces are weak, little energy is required to separate molecules from each other so... **boiling points are LOW**

### Dipole-dipole interactions (van der Waals’ forces)

#### Types

- **Induced** dipole-dipole interactions - London (or Dispersion) Forces
- **Permanent** dipole-dipole interactions

#### 1. Induced dipole-dipole interactions

- **Origin**
  - electrons in atoms or molecules are moving at high speeds in orbitals
  - it is possible for more electrons to be on one side of an atom/molecule
  - this forms a dipole where one side is slightly negative; the other slightly positive
  - a dipole in one atom/molecule can then induce a dipole in a neighbouring one

  ![Dipole Diagram]

  For an instant there are more electrons on the right side of the atom - a dipole is formed

  The dipole on one atom induces (causes) dipoles to form on other atoms. The atoms are then attracted to each other by their oppositely charged ends

- **Result**
  - atoms/molecules become attracted to each other
  - this makes them harder to separate and gives them higher boiling points

- **Trends**
  - the more electrons there are in an atom/molecule the bigger the effect

- **Examples**
  - layers in graphite are held together by weak van der Waals’s forces so it is soft
  - the boiling point of noble gases increases down the group

<table>
<thead>
<tr>
<th>Element</th>
<th>He</th>
<th>Ne</th>
<th>Ar</th>
<th>Kr</th>
<th>Xe</th>
</tr>
</thead>
<tbody>
<tr>
<td>No. of Electrons</td>
<td>2</td>
<td>10</td>
<td>18</td>
<td>36</td>
<td>54</td>
</tr>
<tr>
<td>Boiling point / °C</td>
<td>-269</td>
<td>-246</td>
<td>-186</td>
<td>-152</td>
<td>-108</td>
</tr>
</tbody>
</table>
Some molecules have boiling points much higher than one would expect!

**Electronegativity**

‘The ability of an atom to attract the pair of electrons in a covalent bond to itself.’

**Non-polar bond**

- similar atoms have the same electronegativity
- they will both pull on the electrons to the same extent
- the electrons will be equally shared

**Polar bond**

- different atoms have different electronegativities
- one will pull the electron pair closer to its end
- it will be slightly more negative than average, $\delta^-$
- other will be slightly less negative, or more positive, $\delta^+$
- a dipole is induced and the bond is said to be polar
- the greater the electronegativity difference, the greater the bond polarity.

**Pauling Scale**

- a scale for measuring electronegativity
- values increase across periods
- values decrease down groups
- fluorine has the highest value

<table>
<thead>
<tr>
<th></th>
<th>H</th>
<th>Li</th>
<th>Be</th>
<th>B</th>
<th>C</th>
<th>N</th>
<th>O</th>
<th>F</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>2.1</td>
<td>1.0</td>
<td>1.5</td>
<td>2.0</td>
<td>2.5</td>
<td>3.0</td>
<td>3.5</td>
<td>4.0</td>
</tr>
<tr>
<td>Na</td>
<td>0.9</td>
<td>1.2</td>
<td>1.5</td>
<td>1.8</td>
<td>2.1</td>
<td>2.5</td>
<td>3.0</td>
<td>Br</td>
</tr>
<tr>
<td>K</td>
<td>0.8</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>2.8</td>
</tr>
</tbody>
</table>

**Q.6** Predict the polarity in the following bonds; where applicable, draw in the $\delta^+$ and $\delta^-$

- a) $S—Cl$
- b) $S—O$
- c) $N—O$
- d) $C—O$
- e) $F—Cl$
- f) $C—Cl$
- g) $C—C$
Polar molecules

- some molecules are polar if they contain polar bonds
- the molecules will be polar if they have a NET DIPOLE MOMENT
- it is a bit like balanced forces
- non-polar molecule dipoles in bonds within the molecule ‘cancel each other’
- polar molecule dipoles do not ‘cancel each other out’

\[
\begin{align*}
\text{H}^+ \text{Cl}^- & \quad \text{NO NET DIPOLE} \\
\text{H}^+ \text{Cl}^- & \quad \text{NO NET DIPOLE}
\end{align*}
\]

Experiment

- place a liquid in a burette
- allow a narrow stream to run out
- place a charged rod next to the flow
- polar molecules will be attracted

<table>
<thead>
<tr>
<th>POLAR</th>
<th>NON POLAR</th>
</tr>
</thead>
</table>

Q.7 Which of the following molecules are polar?

| a) F₂ | b) CH₃Cl | c) H₂S | d) C₂H₅OH | e) NH₃ |

2. Permanent dipole-dipole interaction

Occurrence

- between molecules containing polar bonds in addition to the basic induced forces
- the extra attraction between dipoles means that more energy must be put in to separate molecules
- get higher boiling points than expected for a given mass

Q.8 Find the boiling points of the hydrides of elements in Groups 4, 5, 6, and 7.

Plot four lines (one for each group) on a graph of boiling pt. v. molecular mass.

On the graph, state what is unusual about the values for NH₃, H₂O and HF?
HYDROGEN BONDING

Formation
- an extension of dipole-dipole interaction giving even higher boiling points
- bonds between H and the three most electronegative elements, F, O and N are extremely polar
- the small sizes of H, F, N and O concentrates the partial charges in a small volume thus leading to a high charge density
- intermolecular attractions are greater, leading to even higher boiling points

Hydrogen fluoride (HF)

WATER

Ice
- each water molecule is hydrogen bonded to 4 others in a tetrahedral formation
- ice has a “diamond-like” structure
- it is a simple molecular lattice
- its volume is larger than the liquid water making it
- when ice melts, the structure collapses slightly and molecules close together
- they then move a little further apart as they get more energy (warmer)
- this is why water has a maximum density at 4°C and ice floats.

Liquid water
- intermolecular hydrogen bonding gives higher than expected boiling point
- extra attraction between molecules just below the surface gives a high surface tension
  and causes the meniscus to be the shape it is

Viscosity
The greater the hydrogen bonding in alcohols, the greater the viscosity
propan-1-ol propan-1,2-diol propan-1,2,3-triol
CH₃CH₂CH₂OH CH₃CH(OH)CH₂OH CH₂(OH)CH(OH)CH₂OH

INCREASING VISCOSITY
2. GIANT COVALENT LATTICES (covalent networks) - DIAMOND, GRAPHITE and SILICA

bonding
Many atoms joined together in a regular array by large numbers of covalent bonds

Diamond each carbon atom is joined to four others - Co-ordination No. = 4
Graphite each carbon atom is joined to three others - Co-ordination No. = 3

melting point
Very high - structures are made up of a large number of covalent bonds, all of which need to be broken if the atoms are to be separated.

strength
Diamond and silica (SiO₂)
hard exists in a rigid tetrahedral structure

Graphite
soft consists of layers which are attracted by weak induced dipole-dipole interactions
layers can slide over each other
it used as a lubricant and in pencils

electrical
Do not conduct electricity as they have no mobile ions or electrons. BUT...

Graphite conducts electricity
• each atom only uses three of its outer shell electrons for bonding to other atoms
• remaining electron can move through layers allowing the conduction of electricity
• carbon atoms in diamond use all four electrons for bonding so have no free ones

3. MOLECULAR SOLIDS

Iodine
At room temperature, iodine is a grey solid. However, on gentle warming it produces a purple vapour. This is because iodine is composed of diatomic molecules (I₂) existing in an ordered molecular crystal. Each molecule is independent and attracted by weak induced dipole-dipole interactions.

Therefore, little energy is required to separate the iodine molecules.
DATIVE COVALENT (CO-ORDINATE) BONDING

**Theory**

- differs from a covalent bond only in its formation
- both electrons of the shared pair are provided by one species (donor) and it shares the electrons with the acceptor
- donor species will have lone pairs in their outer shells
- acceptor species will be short of their “octet” or maximum.

**Lewis Base :- a lone pair donor  Lewis Acid :- a lone pair acceptor**

**Formation**  

ammonium ion, \( \text{NH}_4^+ \)

The lone pair on N is used to share with the hydrogen ion which needs two electrons to fill its outer shell. The N now has a +ive charge as it is now sharing rather than owning two electrons.

**Boron trifluoride-ammonia**  \( \text{NH}_3\text{BF}_3 \)

Boron has an incomplete shell in BF\(_3\) and can accept a share of a pair of electrons donated by ammonia. The B becomes -ive as it is now shares a pair of electrons (i.e. it is up one electron) it didn’t have before.

**Q.9** Why does BF\(_3\) react with NH\(_3\) but not with CH\(_4\) or AlH\(_3\)?

**Q.10** Show the formation of dative covalent bonds between  

a) \( \text{H}_2\text{O} \) and \( \text{H}^+ \)  
b) \( \text{AlCl}_3 \) and \( \text{Cl}^- \)  
c) \( \text{PCl}_5 \) and \( \text{Cl}^- \)
Fajan's Rules

• not all ionic compounds have high melting points
• some covalently bonded compounds have higher than expected boiling points this is due to dipoles in their structure
• reason: in many substances the bonding is not 100% ionic or covalent

The ideal ionic compound has completely separate, spherical ions and the electron densities are apart from each other.

If the positive ion has a high charge density it can distort the negative ion by attracting the outer shell electrons to give an area of electron density between the two species ... a bit like a covalent bond

The feasibility of formation of covalent bonds is predicted using Fajan’s Rules.

The rules

A compound is more likely to be covalent if the ...

<table>
<thead>
<tr>
<th>CATION</th>
<th>small size</th>
<th>“highly polarising”</th>
<th>attracts electrons in the anion</th>
</tr>
</thead>
<tbody>
<tr>
<td>ANION</td>
<td>large size</td>
<td>“highly polarisable”</td>
<td>will be easily distorted</td>
</tr>
</tbody>
</table>

N.B. Just because a substance is less likely to be covalent according to Fajan’s Rules doesn’t mean it will be ionic; it will remain covalent but have some ionic character.

Examples

Changes in bond type of chlorides as the positive charge density increases due to higher charge (across Period 3) or larger size (down Group 1)

<table>
<thead>
<tr>
<th></th>
<th>charge</th>
<th>ionic rad.</th>
<th>m.pt./°C</th>
<th>solubility</th>
<th>bonding</th>
</tr>
</thead>
<tbody>
<tr>
<td>Period 3</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>NaCl 1+</td>
<td>0.095nm</td>
<td>808</td>
<td>soluble</td>
<td>ionic</td>
<td></td>
</tr>
<tr>
<td>MgCl₂ 2+</td>
<td>0.065nm</td>
<td>714</td>
<td>soluble</td>
<td>ionic</td>
<td></td>
</tr>
<tr>
<td>AlCl₃ 3+</td>
<td>0.050nm</td>
<td>180</td>
<td>hydrolysed</td>
<td>covalent</td>
<td></td>
</tr>
<tr>
<td>SiCl₄ 4+</td>
<td>0.041nm</td>
<td>-70</td>
<td>hydrolysed</td>
<td>covalent</td>
<td></td>
</tr>
<tr>
<td>Group 1</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>LiCl 1+</td>
<td>0.060nm</td>
<td>soluble</td>
<td>some cov. character</td>
<td></td>
<td></td>
</tr>
<tr>
<td>NaCl 1+</td>
<td>0.095nm</td>
<td>soluble</td>
<td>ionic</td>
<td></td>
<td></td>
</tr>
<tr>
<td>KCl 1+</td>
<td>0.133nm</td>
<td>soluble</td>
<td>ionic</td>
<td></td>
<td></td>
</tr>
<tr>
<td>RbCl 1+</td>
<td>0.148nm</td>
<td>soluble</td>
<td>ionic</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Q.11 Which ion / species in each pair favours covalency?

a) Li⁺ and Na⁺  
b) Li⁺ and Be²⁺  
c) B³⁺ and Al³⁺  
d) F⁻ and I⁻  
e) C and Pb  
f) Sn²⁺ and Sn⁴⁺
METALLIC BONDING

Involves a lattice of positive ions surrounded by delocalised electrons

**Formation**

Metal atoms achieve stability by “off-loading” electrons to attain the electronic structure of the nearest noble gas. These electrons join up to form a **mobile cloud** which prevents the newly-formed positive ions from flying apart due to repulsion between similar charges.

Atoms arrange themselves in regular close packed 3-dimensional crystal lattices.

The outer shell electrons of each atom leave to join a mobile “cloud” or “sea” of electrons which can roam throughout the metal. The electron cloud binds the newly-formed positive ions together.

Metallic **bond strength depends** on:

- number of outer electrons donated
- the size of the metal atom/ion.

The melting point is a measure of the attractive forces within the metal.

**electrical**

Conduct electricity as there are mobile electrons.

**strength**

The delocalised electron cloud binds the “ions” together making metals...

- **malleable** can be hammered into sheets
- **ductile** can be drawn into rods

**melting pt.** High. Ease of separation depends on the density of the electron cloud and ionic size/charge.

<table>
<thead>
<tr>
<th>PERIODS</th>
<th>Na (2,8,1)</th>
<th>&lt;</th>
<th>Mg (2,8,2)</th>
<th>&lt;</th>
<th>Al (2,8,3)</th>
</tr>
</thead>
<tbody>
<tr>
<td>m.pt</td>
<td>98°C</td>
<td></td>
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**reason**

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<th>Na (2,8,1)</th>
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<th>K (2,8,8,1)</th>
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<th>Rb</th>
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