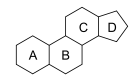
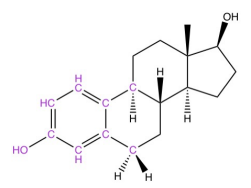
#### **AS A REMINDER:**

#### More examples for representation of molecules



Steroid (C<sub>17</sub>)

#### 1. Estradiol



Female hormone

All purple atoms are in the same plane

#### Types of C:

 $CH_3 - Methyl$ 

 $CH_2$  – Methylene

CH – Methine



- Quaternary carbon

#### **Formal Charge**

- Convention to keep track of charges
- $\sum$  (sum of) of formal charges on all atoms in a molecule = overall charge on molecule

#### Rules for calculating formal charge

- Add number of protons in nucleus
- Subtract number of inner shell electrons
- Subtract number of unshared electrons
- Subtract ½ of the number of shared outer shell electrons

#### **Examples:**

#### 1. Nitrite anion

Overall charge on the nitrite anion is = -1

# Single bonded oxygen:

- +8 (number of protons)
- -2 (1s electrons)
- -6 (unshared electrons)

 $\frac{1}{2}$  x 2 = -1 (1/2 of shared electrons)

#### Central N:

- +7 (number of protons)
- $-2 (1s e^{-})$
- -2 (unshared e<sup>-</sup>)
- -3 (1/2 shared e<sup>-</sup>)
- = 0

#### 2. Methyl anion

Overall charge on the methyl anion is = -1

# Formal Charge on Carbon

- +6 (number of protons)
- -2 (1s electrons)
- -2 (unshared electrons)
- $\frac{1}{2} \times 6 = -3$  (1/2 of shared electrons)

#### 3. Methyl radical

Overall charge on the methyl anion is =  $\mathbf{0}$ Very unstable since it doesn't have an inert gas configuration

## Formal Charge on Carbon

- +6 (number of protons)
- -2 (1s electrons)
- 1 (unshared electrons)
- $\frac{1}{2} \times 6 = \frac{-3}{0}$  (1/2 of shared electrons)

## 4. Methyl cation

- (sp<sup>2</sup> hybridized carbon, planer shape)
- can be reactive intermediate in principle

Overall charge on the methyl anion is = +1

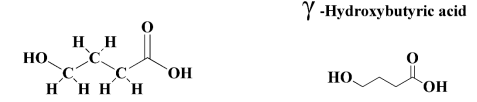
#### Formal Charge on Carbon

- +6 (number of protons)
- -2 (1s electrons)
- 0 (unshared electrons)
- $\frac{1}{2}$  x 6 = -3 (1/2 of shared electrons)

# Empty p orbital



#### **DRAWING CHEMICAL STRUCTURES**



Open chain form

Bond line form

$$MF = C_4H_8O_3$$

NB:The above compound can also be represented in the following forms, resulting from the free rotation of single bonds (sigma).

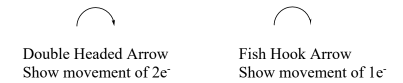
Example:

Note: Single bonds, in general, have free rotation

**Resonance Structures:** Different drawings (or pictures) of the same molecule made by moving electrons but not atoms

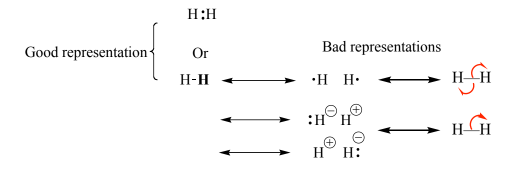
- Move the electrons, keeping the position of the atoms same
- Good resonance structures:
  - o Maintain inert gas configuration around each atom
  - Avoid separation of charges
- Avoid like-charges on adjacent atoms

- Double headed arrow (←→ ) is used indicate resonance forms
- Fish Hook and double headed arrows are used to show electron movement



#### **Examples**

#### 1. Hydrogen gas, H<sub>2</sub>



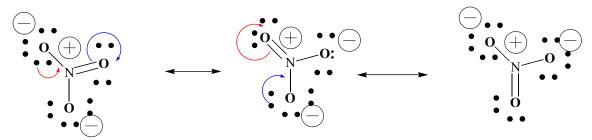
**NB:** In the bad representations, non-inert gas configuration and extra charges have been created

#### 2. Sodium Nitrite, NaNO<sub>2</sub>



Nitrite anion is reactive in both O atoms. Electrons are delocalized in more than one atom – both O atoms has -1/2 charge and contains partial double and single bond character.

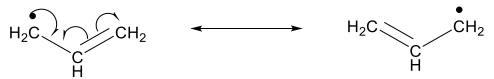
#### 3. Sodium Nitrate, NaNO<sub>3</sub>



# NB: No inert gas configuration disrupted No extra charge created

- The O atoms contain partial single and double bond characteristics (each O has -2/3 charge)

#### 4. Allyl Radical



The radical is relatively stable due to resonance.

#### 5. Propyne cation

**6.** CH<sub>4</sub> Methane – below are POOR resonance structures – additional charges or unshared electrons (not inert gas configuration)

# • CH<sub>3</sub>

but methyl radical – can be reactive intermediate in principle

but methyl anion – can be a reactive intermediate in principle

an be a reactive iple

#### 5. 1,2-Dichlorobenzene

# **BENZENE**

### **Intermolecular Forces: (forces present between molecules)**

- Attractive intermolecular forces:
  - i) **Hydrogen bonding** strongest on per atom basis (e.g. base recognition in forming DNA helix) (also in RNA)
  - ii) **Dipole-dipole interaction** (Intermediate strength)
  - iii) **London forces** (temporary dipole; hydrophobic bonding) weakest on per atom basis distortion of inner shells.

#### **Electronegativity:**

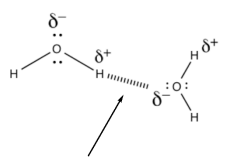
- An atom's desire for electrons (negative charge).
- On the periodic table, electronegativity increases as you go from left to right (up to inert gases, which are not electronegative) and as you go from down to up

- Halogens (F, Cl, Br, I) are highly electonegative
  - o i.e. Fluorine is the most electronegative atom (wants to gain the inert gas configuration of Ne) and is small (has few electrons)
- It influences acidity of H's attached, as well as the intermolecular forces between molecules.

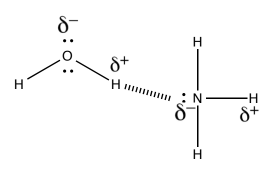
#### **Hydrogen Bonding:**

- Strongest intermolecular attractive force
- Need H directly attached to a very electronegative atom (N, O, F, Cl, Br, I)
  - Known as donors
- Very electronegative atom needs a lone pair of electrons (N, O, F, Cl, Br, I)
  - o Known as acceptors

e.g. H-O-H (water)

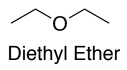


Hydrogen bond



- Can hydrogen bond to itself
  - -Has H directly attached to oxygen
- Has a high boiling points relative to its size due to hydrogen bonding
- Can dissolve in water very well

- Oxygen is electronegative and it is sp<sup>3</sup> hybridized
- The partial positive charge on H and the partial negative charge on O lead to their attraction
- Results in high boiling point (100 C) and high melting point by self-association
- HF, H<sub>2</sub>O and NH<sub>3</sub> form hydrogen bonds
- 1. Water is a liquid at RT while ammonia is a gas
- 2. Oxygen is more e-neg than nitrogen, so the protons on water have a higher positive partial charge than the protons on ammonia
- 3. In an ammonia solution, water would be the hydrogen bond donor and ammonia would be the acceptor
- 4. Water dissolves ammonia very well up to 18M



- Cannot hydrogen bond to itself
  - Has no H directly attached to oxygen (No donor)
  - Can H-bond to water because it has an acceptor
- Has a low boiling point
- Will not dissolve in water very well (although a little bit will be dissolved)

Can't form H-bonds with itself (not a H-bond donor) Lone pairs on O can form H-bonds with water (H-bond acceptor) Poorly soluble in water

#### Dimethylformamide

Can't form H-bonds with itself (not a H-bond donor) Lone pairs on O and N can form H-bonds with water (H-bond acceptor) Infinitely soluble (miscible) in water

Note: The more H-bonds it can form, the more soluble it is in water

#### **Dipole-Dipole Interactions:**

Partial positive charge is the "plus" end, partial negative charge is the arrow head

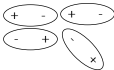
1. Methane; CH<sub>4</sub>



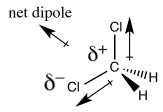
- C and H have ~same electronegativity
- Non-polar (net-zero ~dipole); gas at room temperature
- Low BP -164 °C (this is relatively low compared to water at 100°C)
- Low MP -182 °C
- 2. Chloromethane, methyl chloride; CH<sub>3</sub>Cl

- H and C have similar electronegativity values (non-polar bond)
- Cl is very electronegative due to the fact that it only needs one electron to get inert gas configuration.
- Electron density is pulled toward the chlorine atom, creating a net dipole toward chlorine atom. A net dipole is the vector sum of individual bond dipoles.
- Has a higher MP and BP than methane

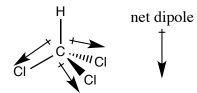
Dipoles in different molecules tend to line-up temporarily with each other (partial positive / negative charge on the molecule) – causes molecules to "stick" to each other



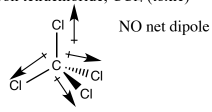
3. Dichloromethane, methylene chloride; CH<sub>2</sub>Cl<sub>2</sub>



- Liquid at room temperature BP 40 °C MP 95 °C
- More polar than chloromethane
- Not miscible with water
- 4. Trichloromethane, chloroform; CHCl<sub>3</sub>



- More polar than methylene chloride BP 61 °C MP 64 °C
- Higher than dichloromethane due to dipole dipole interaction
- 5. Tetrachloromethane, carbon tetrachloride; CCl<sub>4</sub> (toxic)



- Non-polar molecule (net-zero dipole)
- Has temporary dipoles since chlorine is polarizable (see below), BP ~77
- Historically used as a dry-cleaning fluid