# AP Chem - Kolack 

## Week 3

Thermochemistry, atomic structure, $\mathrm{e}^{-}$config, periodic table, bonding

## Thermochemistry

## Energy

- The Universe is composed of matter and energy
- ENERGY is the capacity to do work
- POTENTIAL ENERGY is the energy of position (or composition)
- KINETIC ENERGY is the energy of an object in motion $=1 / 2 m v^{2}$
- Units are $\mathrm{kg}(\mathrm{m} / \mathrm{s})^{2}=$ Joules
- Note that since work $=$ force $\times$ distance, $\mathrm{N} \times \mathrm{m}=\left(\mathrm{kg} \mathrm{m} / \mathrm{s}^{2}\right) \times \mathrm{m}=\mathrm{J}$


## Kinetic energy



## Definitions

- THERMOCHEMISTRY is the study of energy changes that occur during chemical reactions
- Something loses energy; something else gains it
- The SYSTEM is the part of the Universe we are studying
- The SURROUNDINGS are the rest of the Universe


## Definitions (cont'd)

- An OPEN SYSTEM exchanges matter and energy with the surroundings
- A CLOSED SYSTEM exchanges energy only with the surroundings
- An ISOLATED system exchanges neither matter nor energy with the surroundings


## Definitions (cont'd)

- INTERNAL ENERGY $(\mathrm{U})$ is the total energy of the system (PE + KE)
- HEAT (q) is an energy transfer between the system and surroundings
thin wall

Surroundings

## More on internal energy

- Can't be measured (too many variables)
- Can still be useful as a STATE FUNCTION
- Value depends only on current state, not path taken to get there
- q and w aren't state functions


## Work

- Like heat, is an energy transfer between system and surroundings
- We only focus on PV work
- (and electrical work later)



## Laws

- Law of conservation of energy
- Energy can be exchanged, but neither created nor destroyed
- First law of thermodynamics
$-\Delta U=q+w$
- Heat absorbed by system, q>0
- Work done on system w > 0
- Heat given off by system $q<0$
- Work done by system w < 0


## Heat of reaction

- In an EXOTHERMIC reaction, heat is given off by the system to the surroundings ( $q$ is negative)
- In an ENDOTHERMIC reaction, heat is absorbed by the system from the surroundings ( $q$ is positive)


## Enthalpy (H)

- $\mathrm{H}=\mathrm{U}+\mathrm{PV}$
- We will not derive it, but $\Delta \mathrm{H}$ (enthalpy change) is the heat of reaction carried out at constant temperature and pressure with all work limited to PV work
- EXTENSIVE property - depends on how much is present
- Is a state function


## Enthalpy (cont'd)



## Problem-solving

- How much heat is given off or absorbed when $5.67 \mathrm{~mol} \mathrm{HCl}(\mathrm{g})$ is formed in the following reaction? $\mathrm{H}_{2}(\mathrm{~g})+\mathrm{Cl}_{2}(\mathrm{~g}) \rightarrow 2 \mathrm{HCl}(\mathrm{g}) \Delta \mathrm{H}=-184.6$ kJ
- $5.67 \mathrm{~mol} \mathrm{HCl} x(-184.6 \mathrm{~kJ} / 2 \mathrm{~mol} \mathrm{HCl})$
- = -523kJ
- (exothermic)


## Calorimetry

- The measurement of heat is called CALORIMETRY ("calorie measurement")
- $1 \mathrm{cal}=4.184 \mathrm{~J}$
- $1 \mathrm{Cal}=1000 \mathrm{cal}$
- Lab expts make assumptions



## Heat capacity

- The HEAT CAPACITY of a system is the quantity of heat required to change the temperature of the system by $1^{\circ} \mathrm{C}$ (or 1K)
- Units are either $\mathrm{J} /{ }^{\circ} \mathrm{C}$ or $\mathrm{J} / \mathrm{K}$
- MOLAR HEAT CAPACITY is the heat capacity of one mole of a substance
- SPECIFIC HEAT is the heat capacity of a 1 g sample


## Hess' Law

- HESS' LAW OF HEAT SUMMATION says that the heat of a reaction is constant, whether the reaction is one step or many steps
- Example:
given $\mathrm{CO}(\mathrm{g})+1 / 2 \mathrm{O}_{2}(\mathrm{~g}) \rightarrow \mathrm{CO}_{2}(\mathrm{~g}) \Delta \mathrm{H}=-283.0 \mathrm{~kJ}$ and C (graphite) $+\mathrm{O}_{2}(\mathrm{~g}) \rightarrow \mathrm{CO}_{2}(\mathrm{~g}) \Delta \mathrm{H}=-393.5 \mathrm{~kJ}$ what is $\Delta \mathrm{H}$ for C (graphite) $+1 / 2 \mathrm{O}_{2}(\mathrm{~g}) \rightarrow \mathrm{CO}(\mathrm{g})$ ?
- Reversing the first equation and adding it to the second gives us the equation we're looking for:
$\mathrm{CO}_{2}(\mathrm{~g}) \rightarrow \mathrm{CO}(\mathrm{g})+1 / 2 \mathrm{O}_{2}(\mathrm{~g})$
C (graphite) $+\mathrm{O}_{2}(\mathrm{~g}) \rightarrow \mathrm{CO}_{2}(\mathrm{~g})$
- Doing the same to the heats of reaction gives us the heat of reaction we're looking for: $-(-283.0 \mathrm{~kJ})+(-393.5 \mathrm{~kJ})=-110.5 \mathrm{~kJ}$


## Hess' Law (cont'd)



- Reverse the eq., reverse the $\Delta \mathrm{H}$
- Multiply the equation, multiply the $\Delta \mathrm{H}$


## Standard enthalpy

- STANDARD STATE of a liquid or solid or gas is the substance at 1 atm and the T of interest
- Just like altitude is given relative to sea level, STANDARD ENTHALPY OF FORMATION $\Delta H_{f}{ }^{\circ}$ is the enthalpy change that occurs in the formation of 1 mol of the substance from its elements in their standard states
- STANDARD ENTHALPY OF REACTION $\left(\Delta \mathrm{H}^{\circ}\right)$ is the enthalpy change when the reactants in their standard states form products in their standard states
$=\Delta H_{f}{ }^{\circ}$ (products) $-\Delta H_{f}^{\circ}{ }^{\circ}$ (reactants)


## end of section

## Review

- Reaction- the formation and breaking of chemical bonds
- Bond- a transfer or sharing of electrons


## Electrons

- Abbreviated $\mathrm{e}^{-}$
-What are they?
- How were they discovered?


## Early experiments

- William Crookes used an evacuated tube hooked up to a source of electricity (in 1879) and saw a
 fluorescent beam
- JJ Thomson (in 1897) determined the particles causing this fluorescence were negatively charged, based on their behavior in the presence of a magnetic field
- Thomson found the fluorescence was independent of the identity of the residual gas in the tube....the particles must therefore be a property of all matter


## Cathode rays



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## More on Thomson

- Thomson determined the mass to charge ratio $\left(\mathrm{m}_{\mathrm{e}} / \mathrm{e}=-5.686 \times 10^{-12}\right.$ $\mathrm{kg} / \mathrm{C}$ ) and decided (based on previous knowledge with $\mathrm{H}^{+}$) he had one of three situations:
- If the charge were similar to $\mathrm{H}^{+}$, then mathematically the mass of the particle must be much less than that of $\mathrm{H}^{+}$
- If the mass were similar to $\mathrm{H}^{+}$, then mathematically the charge must be much greater than that on $\mathrm{H}^{+}$
- Something in between the two extremes
- He suspected the first case to be true, but could not prove it


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## Explanations

- 1909 Millikan's oil drop experiment (next slide) showed that the rate of fall (velocity) of a charged oil drop could be varied in the presence of an electric field such that the charge could only in multiples of a fundamental unit ( $\mathrm{e}=-1.602 \times 10^{-19}$ )
- Thomson knew the e- had to be neutralized by + charges, but was unsure of the arrangement and devised the "raisin pudding" model (2 slides ahead)
- Ernest Rutherford (a student of Thomson's) studied radioactivity, a phenomenon of unstable heavy atoms giving off radiation during disintegration
- Rutherford bombarded metal foils with alpha particles $\left(\mathrm{He}^{2+}\right)$...most went through, but some were scattered at odd angles...not explained by the Thomson model of the atom...
- Can be explained with a nuclear model (three slides ahead)


## Millikan's Oil Drop Experiment

- George Stoney: names the cathode-ray particle the electron.
- Robert Millikan: determines a value for the electron's charge:

$$
e=-1.602 \times 10^{-19} \mathrm{C}
$$



## J. J. Thomson's Model of the Atom

- Thomson proposed an atom with a positively charged sphere containing equally spaced electrons inside.
- He applied this model to atoms with up to 100 electrons.


Helium atom, He


## Alpha Scattering "Gold Foil Expt": Rutherford's observations



Alpha
Scattering Experiment:

Rutherford's conclusions

If Thomson's model of the atom was correct, most of the alpha particles should have been deflected a little, like bullets passing through a cardboard target.


## Rearranging the periodic table

- Prior to 1914 , atomic numbers were given out simply by arranging the elements in order of mass...
- Now that the number of protons and electrons could be determined and the mass of them calculated, there was some additional mass to be explained.
- A neutral particle was postulated, the existence of which was proven in 1932 by Chadwick


## Proof

- A Crookes tube can also be constructed to detect positive ions drawn to the cathode (negative pole) (or deflected away from the anode (positive pole).
- These experiments ARE matter dependent.
- A mass analyzer (mass spectrometer) is based on this concept.



## A mass

 spectrum for
## mercury

Mass spectrum of an element shows the abundance of its isotopes. What are the three most abundant isotopes of mercury?


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## Electromagnetic (EM) radiation

- Composed of perpendicular electric and magnetic field waves (something that repeats as it progresses through space)
- The region of the full spectrum which WE call light is only a small portion (next slide)
- Wavelength $(\lambda)$ is the distance between equivalent wave points on adjacent peaks, expressed in meters



## Electromagnetic (EM) radiation



## A continuous spectrum



## Frequency

- Frequency $(v)$ is how many waves pass a fixed point per unit time, expressed in hertz $(\mathrm{Hz}, \mathrm{s}-1)$
- Frequency times wavelength $=$ speed, $\mathrm{c}=$ $2.99792458 \times 10^{8} \mathrm{~m} / \mathrm{s}$ (or just $3 \times 10^{8} \mathrm{~m} / \mathrm{s}$ )
- In 1900, Max Planck explained the behavior of both high and low frequency radiation by equating energy with frequency and a constant, $\mathrm{E}=\mathrm{hv}$, where h is Planck's constant $=6.626 \times 10^{-34}$ (think how small this number is)
- Planck's equation demonstrates that light comes only in discrete "packets" called quanta (singular = "quantum")


## Einstein

- In 1905, Einstein used Planck's theories to explain how electrons can
 result from
bombarding a sample with light...
- This plus the observation of line spectra when
 elements were heated lead to...


## The Photoelectric Effect



But not "any old" light will cause ejection of electrons ...

## A Line Spectrum



## The Bohr model of the atom

- Combined classical physics and quantum theory
- Different energy levels of electrons correspond to orbits of different distances from the nuclei of atoms
- The lowest energy level (that nearest the nucleus) is level 1 , the next is level 2 , and so on
- The electron energy levels $E_{n}=-B / n^{2}$, where $n$ is the energy level (an integer) and $B$ is a constant related to Planck's and the mass and charge of an electron (negative energy for an attractive force)


## The Bohr Model of hydrogen



## Experimental

- Bohr then explained line spectra as being due to the energy difference between 2 levels

$$
E_{f}=\frac{-B}{n_{j}^{2}} \quad \text { and } \quad E_{i}=\frac{-B}{n_{i}^{2}} \text { so } \quad \Delta E=B\left(\frac{1}{n_{j}^{2}}-\frac{1}{n_{i}^{2}}\right)
$$

- (That should be $n_{i}-n_{f}$ )


## Atoms

- When an atom has all its electrons in their lowest possible energy levels, the atom is in its GROUND STATE
- If energy has been supplied sufficient to promote an electron to a higher level, the atom is in an EXCITED STATE


## Energy levels and spectral lines for

## hydrogen



## Wave-particle duality

- If light can act like particles of matter, can particles act like waves?
- According to DeBroglie's theories, yes!

$$
A=\frac{h}{m v}
$$

## Particles as waves

- The Bohr model is mostly classical, but if the particles are treated as waves, quantum or wave mechanics are needed
- In 1926 Erwin Schrodinger developed a mathematical equation to describe the hydrogen atom (a wave equation, the solution to which is called a wave function
- According to Max Born, the square of the wave function $\left(\psi^{2}\right)$ gives the probability of finding an electron in a particular volume of space in an atom


## Particles as waves (cont'd)

- According to Werner Heisenberg, in fact, we cannot know both the exact position and motion of a tiny particle like an electron simultaneously... think of it this way- the act of measuring its position changes its motion, and vice versa

$$
\Delta x \Delta p \geq \frac{h}{4 \pi}
$$



## Quantum numbers

- Integral value parameters from the wave function of the hydrogen atom...
- A set of these three wave function quantum numbers (there is a fourth) is called an atomic orbital, a mathematical expression which allows us to visualize a 3D region in an atom where there is a significant probability of finding an electron


## Quantum numbers (cont'd)

- Principal quantum number (n)
- energy level
- 1,2,3,4,5,...
- Orbital angular momentum (azimuthal) quantum number (I)
- shape
$-0,1,2,3, \ldots(n-1)(s, p, d, f)$
- Magnetic quantum number ( $\mathrm{m}_{1}$ )
- direction
$-0,+/-1,+/-2, \ldots+/-$ I
- Spin quantum number $\left(\mathrm{m}_{\mathrm{s}}\right)$
-     + ½, - $1 / 2$


## Quantum Numbers: $n$

The principal quantum number ( $n$ ):

- Is independent of the other two quantum numbers.
- Can only be a positive integer ( $n=1,2,3,4, \ldots$ )
- The size of an orbital and its electron energy depend on the value of $n$.
- Orbitals with the same value of $n$ are said to be in the same principal shell.


## Quantum numbers: 1

The orbital angular momentum quantum number (I):

- Determines the shape of the orbital.
- Can have positive integral values from $0,1,2 \ldots(n-1)$
- Orbitals having the same values of $n$ and of $I$ are said to be in the same subshell.

| Value of $l$ | 0 | 1 | 2 | 3 |
| :--- | :--- | :--- | :--- | :--- |
| Subshell | $s$ | $p$ | $d$ | $f$ |

- Each orbital designation represents a different region of space and a different shape.


## Quantum numbers: $\mathrm{m}_{1}$

The magnetic quantum number ( $m_{1}$ ):

- Determines the orientation in space of the orbitals of any given type in a subshell.
- Can be any integer from -/ to +/
- The number of possible values for $m_{l}$ is $(2 /+1)$, and this determines the number of orbitals in a subshell.


## Electron spin: $\mathrm{m}_{s}$

- The electron spin quantum number ( $m_{s}$ ) explains some of the finer features of atomic emission spectra.
- The number can have two values: $+1 / 2$ and $-1 / 2$.
- The spin refers to a magnetic
 field induced by the moving electric charge of the electron as it spins.
- The magnetic fields of two electrons with opposite spins cancel one another; there is no net magnetic field for the pair.


## The 1s orbital

- The $1 s$ orbital $\left(n=1, I=0, m_{l}=0\right)$ has spherical symmetry.
- An electron in this orbital spends most of its time near the nucleus.



## Analogy to the 1s orbital



## The 2s orbital

- The $2 s$ orbital has two concentric, spherical regions of high electron probability.
- The region near the nucleus is separated from the outer region by a node-a region (a spherical shell in this case) in which the electron probability is zero.

(a)

(b)

(c)

(d)


## The three p orbitals



Three values of $\boldsymbol{m}_{\boldsymbol{l}}$ gives three $p$ orbitals

$p_{z}$ orbital

## The five d orbitals



## The seven f orbitals



## end of section

## Schrödinger

- Cannot be solved for multielectron atoms
- We must assume the orbitals are all hydrogen-like


## Differences

- In the H atom, all subshells ( $s, p, d, f$ ) of a principal shell are at the same energy
- This varies for other elements, though all orbitals within a subshell are at the same energy level. (2 slides ahead) (DEGENERATE orbitals)
- Orbital energies are lower in multielectron atoms than in H . (WHY????)
- In multielectron atoms, for higher numbered principal shells, some subshells have nearly identical energies. (WHY???)


## REMINDER

## ORBITALS

ARE
REGIONS OF PROBABILITY

## Orbital energy diagrams



## Electron configuration

- Describes the distribution of electrons among the various orbitals in the atom.
- Electron configuration is represented in two ways:
- $s p d f$ notation
- Numbers denote principal shell
- Letters denote subshell
- Superscripts denote number of electrons per subshell
- Electrons occupy orbitals of the lowest energy available
- Orbital diagram


Electrons occupy
the $n=1$ energy level.


## s p d f notation (cont'd)

- No two electrons can have the same set of 4 quantum numbers
- Thus, an orbital can accommodate at most 2 electrons (Wolfgang Pauli, 1926)- The Pauli Exclusion Principle
- Electrons enter empty orbitals whenever possible when faced with a set of identical-energy orbitals (Hund's rule)
- Electrons in half-filled orbitals have the same (parallel) spins
- Therefore, when drawing ground state electron configurations, the aufbau ("building up") principle is employed


## s p d f notation (cont'd)

- $1 s, 2 s, 2 p, 3 s, 3 p, 4 s$, $3 d, 4 p, 5 s, 4 d, 5 p, 6 s$, $4 f, 5 d, 6 p, 7 s, 5 f, 6 d$, $7 p$
- To remember the above, use the figure, or better yet, USE THE PERIODIC TABLE



## Using the periodic table to write e- configurations



## Exceptions to aufbau

- Cr and Cu are examples where the observed configuration (confirmed experimentally) is not as expected due to a hyperstability of a FILLED or HALFFILLED subshell


## Exceptions to the Aufbau principle (cont'd)

| Half-filled $d$ subshell plus half-filled $s$ subshell has slightly lower in energy than $s^{2} d^{4}$. | Sc |  | $3 d$ |  |  |  |  | $4 s$ | $[\operatorname{Ar}] 3 d^{1} 4 s^{2}$ |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  |  | $\uparrow$ |  |  |  |  | $4 \downarrow$ |  |
|  | Ti | [Ar] | $\uparrow$ |  |  |  |  | 41 | $[\operatorname{Arr}] 3 d^{2} 4 s^{2}$ |
|  | v | [Ar] | 4 |  | $\uparrow$ |  |  | 4 | $[\operatorname{Ar}] 3 d^{3} 4 s^{2}$ |
| Filled $d$ subshell plus half-filled $s$ subshell has slightly lower in energy than $s^{2} d^{9}$. |  | [Ar] |  |  | $\uparrow$ |  | $\uparrow$ | 4 | $[\mathrm{Ar}] 3 d^{5} 4 s^{1}$ |
|  | Mn | [Ar] | $\uparrow$ |  | $\uparrow$ | $\uparrow$ | $\uparrow$ | 41 | $[\mathrm{Ar}] 3 d^{5} 4 \mathrm{~s}^{2}$ |
|  | Fe | [Ar] | 4 |  | 4 | $\uparrow$ | $\uparrow$ | 41 | $[\mathrm{Ar}] 3 d^{6} 4 s^{2}$ |
|  | Co | [Ar] | 4 |  |  |  | $\uparrow$ | $4 \downarrow$ | $[\mathrm{Ar}] 3 d^{7} 7 s^{2}$ |
| More exceptions occur farther down the periodic table. They aren't always predictable, because energy levels get closer together. |  | [Ar] | 4. |  | $\uparrow \downarrow$ |  | $\uparrow$ | $4 \downarrow$ | $[\mathrm{Ar}] 3 d^{8} 4 s^{2}$ |
|  |  | [Ar] |  |  | $\uparrow 1$ |  | 4 | $\uparrow$ | $[\operatorname{Ar}] 3 d^{10} 4 s^{1}$ |
|  | Zn |  |  |  | १t |  | $41$ | $4 \downarrow$ | $[\mathrm{Ar}] 3 d^{10} 4 s^{2}$ |

## Exceptions to the Aufbau principle (cont'd)



## Example

- What is the electron configuration of Cs?
- Noble-gas-core (condensed) abbreviation: Another example of chemists' shorthand. The portion of the config. that corresponds to the electron config. of the nearest previous noble gas is replaced with a bracketed chemical symbol. It's easier to write ...

$$
\begin{aligned}
& (Z=3) \mathrm{Li}[\mathrm{He}] 2 \mathrm{~s}^{1} \\
& (Z=22) \mathrm{Ti}[\mathrm{Ar}] 4 \mathrm{~s}^{2} 3 \mathrm{~d}^{2}
\end{aligned}
$$

## Example

- Give the complete ground-state electron configuration of a strontium atom (a) in spdf notation and
(b) in the noble-gas-core abbreviated notation
(c) in the orbital box diagram notation.


## Valence electrons

- The outermost shell of electrons is known as the valence shell
- The inner electrons are the core electrons
- When writing the configuration of ions, add or remove electrons to/from the valence shell
- Often results in noble gas configuration
- EXCEPTION: Transition elements- outer $s$ electrons are lost first
- If two species have the same electron configuration, they are said to be isoelectronic


## Valence electrons and core electrons

- The valence shell is the outermost occupied principal shell. The valence shell contains the valence electrons.
- For main group elements, the number of valence shell electrons is the same as the periodic table group number ( 2 A elements: two valence electrons, etc.)
The period number is the same as the principal quantum number $n$ of the electrons in the valence shell.
- Electrons in inner shells are called core electrons.

Example: As
Five valence electrons, for which $\boldsymbol{n}=4$


## Electron configurations of ions

- To obtain the electron configuration of an anion by the aufbau process, we simply add the additional electrons to the valence shell of the neutral nonmetal atom.
- The number added often completes the shell.
- A nonmetal monatomic ion usually attains the electron configuration of a noble gas atom.
$\mathrm{O}^{2-}$ : [Ne]
$\mathrm{Br}^{-}$: [Kr]


## Electron configurations of ions (cont'd)

- A metal atom loses electrons to form a cation.
- Electrons are removed from the configuration of the atom.
- The first electrons lost are those of the highest principal quantum number.
- If there are two subshells with the same highest principal quantum number, electrons are lost from the subshell with the higher $l$.


## Electron configurations of ions (cont'd)

Atom
F $1 s^{2} 2 s^{2} 2 p^{5}$
$\mathrm{S}[\mathrm{Ne}] 3 s^{2} 3 p^{4}$
$\mathrm{Sr}[\mathrm{Kr}] 5 s^{2}$
$\mathrm{Sr}^{2+}[\mathrm{Kr}] 5 s^{2}$
$\mathrm{Ti}{ }^{4+}[\mathrm{Ar}] 4 s^{2} 3 \mathrm{c}^{R}$
$\mathrm{Fe}^{2+}[\mathrm{Ar}]-4 s^{2} 3 d^{6}$
(or)
[ Ne ]
[Ar]
[Kr]
[Ar]
[Ar] $3 d^{6}$


## $e^{-}$configuration of ions (cont'd)

## Table 8.3 Electron Configurations of Some Metal Ions

| Noble Gas |  |  | Pseudo-Noble Gas ${ }^{\text {a }}$ |  | $18+2^{\text {b }}$ | Various |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| $\mathrm{Li}^{+}$ | $\mathrm{Be}^{2+}$ | $\mathrm{Al}^{3+}$ | $\mathrm{Cu}^{+}$ | $\mathrm{Zn}^{2+}$ | $\mathrm{In}^{+}$ | $\mathrm{Cr}^{2+}$ : | $[\mathrm{Ar}] 3 d^{4}$ |
| $\mathrm{Na}^{+}$ | $\mathrm{Mg}^{2+}$ |  | $\mathrm{Ag}^{+}$ | $\mathrm{Cd}^{2+}$ | $\mathrm{Tl}^{+}$ | $\mathrm{Cr}^{3+}$ : | $[\mathrm{Ar}] 3 d^{3}$ |
| $\mathrm{K}^{+}$ | $\mathrm{Ca}^{2+}$ |  | $\mathrm{Au}^{+}$ | $\mathrm{Hg}^{2+}$ | $\mathrm{Sn}^{2+}$ | $\mathrm{Mn}^{2+}$. | $[\mathrm{Ar}] 3 d^{5}$ |
| $\mathrm{Rb}^{+}$ | $\mathrm{Sr}^{2+}$ |  |  |  | $\mathrm{Pb}^{2+}$ | $\mathrm{Mn}^{3+}$ : | [Ar] $3 d^{4}$ |
| Cs ${ }^{+}$ | $\mathrm{Ba}^{2+}$ |  |  |  | $\mathrm{Sb}^{3+}$ | $\mathrm{Fe}^{2+}$ : | $[\mathrm{Ar}] 3 d^{6}$ |
|  |  |  |  |  | $\mathrm{Bi}^{3+}$ | $\mathrm{Fe}^{3+}$ : | $[\mathrm{Ar}] 3 d^{5}$ |
|  |  |  |  |  |  | $\mathrm{Co}^{2+}$ : | $[\mathrm{Ar}] 3 d^{7}$ |
|  |  |  |  |  |  | $\mathrm{Co}^{3+}$ : | $[\mathrm{Ar}] 3 d^{6}$ |
|  |  |  |  |  |  | $\mathrm{Ni}^{2+}$ : | $[\mathrm{Ar}] 3 d^{8}$ |

[^0]
## Magnetism

- Diamagnetic atoms have all electrons paired
- Paramagnetic atoms have some unpaired electrons
- Ferromagnetism is the exceptionally strong attractions of a magnetic field for iron and a few other substances.
- Despite the fact that your instructor's thesis centered on this topic, we will not discuss it further


## Periodic properties

- Certain physical and chemical properties recur at regular intervals (Mendeleev, anyone?), and/or vary in regular fashion, when the elements are arranged according to increasing atomic number.
- Melting point, boiling point, hardness, density, physical state, and chemical reactivity are periodic properties.
- We will examine several periodic properties that are readily explained using electron configurations.
- Atomic radii
- Effective nuclear charge
- Covalent radii
- Ionic radii
- Ionization energy
- Electron affinity


## Periodic properties: atomic radius

- Half the distance between the nuclei of two atoms is the atomic radius.
Covalent radius: half the distance between the nuclei of two identical atoms joined in a molecule.
Metallic radius: half the distance between the nuclei of adjacent atoms in a solid metal.


## Atomic radius (cont'd)

- Atomic radius increases from top to bottom within a group.
- The value of $n$ increases, moving down the periodic table.
- The value of $n$ relates to the distance of an electron from the nucleus.
- Often in Angstroms ( $10^{-10} \mathrm{~m}$ ) or picometers ( $10^{-12} \mathrm{~m}$ )


## Atomic radius (cont'd)

- Atomic radius decreases from left to right within a period.
- Why? The effective nuclear charge $\left(\mathrm{Z}_{\text {eff }}\right)$ increases from left to right, increasing the attraction of the nucleus for the valence electrons, and making the atom smaller.


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## Atomic radii of the elements



## Example

- With reference only to a periodic table, arrange each set of elements in order of increasing atomic radius:
- (a) Mg, S, Si
- (b) As, N, P
- (c) As, Sb, Se


## Ionic radii

The ionic radius of each ion is the portion of the distance between the nuclei occupied by that ion.

Internuclear distance: 205 pm


## Ionic radii- cations

- Cations are smaller than the atoms from which they are formed; the value of $n$ usually decreases. Also, there is less electronelectron repulsion.



## Ionic radii- anions

- Anions are larger than the atoms from which they are formed.
- Effective nuclear charge is unchanged, but additional electron(s) increase electron-electron repulsion.
- Isoelectronic species have the same electron configuration; size
 decreases with effective nuclear charge.


## Some atomic and ionic radii



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## Example

- Refer to a periodic table, but not to the previous figure, and arrange the following species in the expected order of increasing radius:
$\mathrm{Ca}^{2+}, \mathrm{Fe}^{3+}, \mathrm{K}^{+}, \mathrm{S}^{2-}, \mathrm{Se}^{2-}$


## Ionization energy

- Ionization energy (IE) is the energy required to remove an electron from a ground-state gaseous atom.
- IE is usually expressed in kJ per mole of atoms.

$$
\begin{array}{ll}
\mathrm{M}(g) \rightarrow \mathrm{M}^{+}(g)+\mathrm{e}^{-} & \Delta H=I_{1} \\
\mathrm{M}^{+}(g) \rightarrow \mathrm{M}^{2+}(g)+\mathrm{e}^{-} \Delta H=I_{2} \\
\mathrm{M}^{2+}(g) \rightarrow \mathrm{M}^{3+}(g)+\mathrm{e}^{-} \Delta H=I_{3}
\end{array}
$$

## Trends in IE

- $I_{1}<I_{2}<l_{3}$
- Removing an electron from a positive ion is more difficult than removing it from a neutral atom.
- A large jump in / occurs after valence electrons are completely removed (why?).
- $I_{1}$ decreases from top to bottom on the periodic table.
- $n$ increases; valence electron is farther from nucleus.
- $I_{1}$ generally increases from left to right, with exceptions.
- Greater effective nuclear charge from left to right holds electrons more tightly.


## Selected ionization

## Table 8.4 Ionization Energies of Group 1A and Group 2A Elements, kJ/mol

|  | 1A | 2A |
| :---: | :---: | :---: |
|  | Li | Be |
| $I_{1}$ | 520 | 900 |
| $I_{2}$ | 7298 | 1757 |
|  | Na | Mg |
| $I_{1}$ | 496 | 738 |
| $I_{2}$ | 4562 | 1451 |
|  | K | Ca |
| $I_{1}$ | 419 | 590 |
| $I_{2}$ | 305 | 1145 |
| $I_{1}$ | Rb | Sr |
| $I_{2}$ | 403 | 550 |
|  | 2633 | 1064 |
| $I_{1}$ | Cs | Ba |
| $I_{2}$ | 376 | 503 |

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## Selected ionization energies



## First ionization energies



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## Example

- Without reference to the previous figure, arrange each set of elements in the expected order of increasing first ionization energy.
(a) $\mathrm{Mg}, \mathrm{S}, \mathrm{Si}$
(b) As, N, P
(c) As, Ge, P


## Electron affinity

Electron affinity $(E A)$ is the energy change that occurs when an electron is added to a gaseous atom:

$$
\mathrm{M}(\mathrm{~g})+\mathrm{e}^{-} \rightarrow \mathrm{M}^{-}(\mathrm{g}) \quad \Delta H=E A_{1}
$$

- A negative electron affinity means that the process is exothermic.
- Nonmetals generally have more affinity for electrons than metals do. (Nonmetals like to form anions!)
- Electron affinity generally is more negative or less positive on the right and toward the top of the periodic table.


## Selected electron affinities

|  | The halogens have a greater affinity for electrons than do the alkali metals, as expected. |  |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Tab | S | ecte | Ele |  |  |  |  |
| 1A | 2A | 3A | 4A | 5A | 6A | 7A | 8A |
| Li | Be | B | C | N | O | F | Ne |
| -60 | $>0$ | -27 | -154 | $\approx 0$ | -141 | -328 | $>0$ |
| Na |  |  |  |  | S | Cl |  |
| -53 |  |  |  |  | -200 | -349 |  |
| K |  |  |  |  | Se | Br |  |
| -48 |  |  |  |  | -195 | -325 |  |
| Rb |  |  |  |  | Te | I |  |
| -47 |  |  |  |  | -190 | -295 |  |
| Cs |  |  |  |  | Po | At |  |
| -46 |  |  |  |  | -183 | -270 |  |

## Other periodic properties (overview)

- Metallic character
- Increases top to bottom
-Decreases left to right
- Flame color
-Due to electronic transitions
- Oxidizing and reducing power
-Think of how easy it is to gain or lose electron(s)
- Acidic and basic character
-Ditto!


## Metals

- Metals have a small number of electrons in their valence shells and tend to form positive ions.
-For example, an aluminum atom loses its three valence electrons in forming $\mathrm{Al}^{3+}$.
- All s-block elements (except H and He ), all $d$ - and $f$-block elements, and some $p$ block elements are metals.


## Metallic Character

- Metallic character is related to atomic radius and ionization energy.
- Metallic character generally increases from right to left across a period, and increases from top to bottom in a group.

More

metallic
character


More
metallic
character

## Nonmetals

- Atoms of a nonmetal generally have larger numbers of electrons in their valence shell than do metals.
- Many nonmetals tend to form negative ions.
- All nonmetals (except H and He ) are p-block elements.
- Nonmetallic character generally increases right-to-left and increases bottom-totop on the periodic table (the opposite of metallic character).

More nonmetallic character


More nonmetallic character

## Metalloids

- A heavy stepped diagonal line separates metals from nonmetals; some elements along this line are called metalloids.
- Metalloids have properties of both metals and nonmetals.


| *Lanthanide series | $\mathbf{C e}$ | $\mathbf{P r}$ | $\mathbf{N d}$ | $\mathbf{P m}$ | $\mathbf{S m}$ | $\mathbf{E u}$ | $\mathbf{G d}$ | $\mathbf{T b}$ | $\mathbf{D y}$ | $\mathbf{H o}$ | $\mathbf{E r}$ | $\mathbf{T m}$ | $\mathbf{Y b}$ | $\mathbf{L u}$ |
| :--- | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| †Actinide series | $\mathbf{T h}$ | $\mathbf{P a}$ | $\mathbf{U}$ | $\mathbf{N p}$ | $\mathbf{P u}$ | $\mathbf{A m}$ | $\mathbf{C m}$ | $\mathbf{B k}$ | $\mathbf{C f}$ | $\mathbf{E s}$ | $\mathbf{F m}$ | $\mathbf{M d}$ | $\mathbf{N o}$ | $\mathbf{L r}$ |

## A Summary of trends

## More nonmetallic character

## More negative electron affinity

Increasing ionization energy


Increasing atomic radius

More metallic character
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## The noble gases

- The noble gases are on the far right of the periodic table between the highly active nonmetals of Group 7A and the very reactive alkali metals Group 1A.
- The noble gases rarely enter into chemical reactions because of their stable electron configurations.
- However, a few compounds of noble gases (except for He and Ne ) have been made.

Atoms emit energy when electrons drop from higher to lower energy states.

## Other trends-

 flame colorElements with low first ionization energies can be excited in a Bunsen burner flame, and often emit in the visible region of the spectrum.

Elements with high values of $\mathrm{IE}_{1}$ usually require higher temperatures for emission, and the emitted light is in the UV region of the spectrum.


## Oxidizing and reducing agents revisited

- The halogens (Group 7A) are good oxidizing agents.
- Halogens have a high affinity for electrons, and their oxidizing power generally varies with electron affinity.



## Oxidizing and reducing agents revisited (cont'd)

- The $s$-block elements are very strong reducing agents.
- All the IA metals and the heavier IIA metals will displace $\mathrm{H}_{2}$ from water, in part because of their low values of $I E_{1}$.
- A low $I E_{1}$ means that the metal easily gives up its electron(s) to hydrogen in water, forming hydrogen gas.



## end of section

## Reminder (again)

- As you remember, a chemical reaction is the formation and/or breaking of chemical bonds
- A chemical bond is a transfer or sharing of electrons


## Chemical bonds



- Chemical bonds are the forces that hold atoms together in compounds
(a) $\mathrm{H}_{2}{ }^{+}$
- Bonds are electrostatic forces; attractions between opposite charges and repulsions between like charges



## Chemical bonds (cont'd)



- Chemical bonds are the forces that hold atoms together in compounds
- When 2 atoms are brought together, there is an optimum distance where the unfavorable repulsions and the favorable attractions are maximized (see the potential energy diagram at left)
Chemical bonds (cont'd)
- Bonds are responsible for such physical properties as melting point and boiling point
- Bonds (and the type and number of atoms present) ultimately determine the shape of a molecule, and structure determines function


## Lewis theory

- Valence electrons participate in bonding
- Metals and nonmetals combine by transferring electrons, forming cations and anions involved in ionic bonds
- Nonmetals combine with each other by sharing electron pairs (overlapping orbitals), forming covalent bonds
- Complete transfer is a $100 \%$ ionic "bond"
- Equal sharing is a $100 \%$ covalent bond
- The above are two extremes


## Lewis theory (cont'd)

- When atoms gain, lose, or share electrons, atoms tend to acquire the electron configuration of a noble gas ("noble gas configuration")
- Remember, electrons are transferred, NOT protons, so the atom becomes an ion of the original element, not the noble gas itself... noble gas configurations are hyper-stable
- H, Li, and Be follow the duet rule (He configuration), while all other elements (except transition metals) follow the octet rule ( Ne , $\mathrm{Ar}, \mathrm{Kr}, \mathrm{Xe}, \mathrm{Rn}$ configuration with 8 (or 18) valence electrons)


## Lewis symbols

- A Lewis symbol is the element's symbol with "dots" on 4 sides of it to represent the valence electrons
- Electrons are shown "unpaired" whenever possible (although the idea of electron spin had not been developed when Lewis developed his theories)


## Lewis symbols (cont'd)

- In a Lewis symbol, the chemical symbol for the element represents the nucleus and core electrons of the atom.
- Dots around the symbol represent the valence electrons.
- In writing Lewis symbols, the first four dots are placed singly on each of the four sides of the chemical symbol. (Though spin was unknown at the time.)
- Dots are paired as the next four are added.
- Lewis symbols are used primarily for those elements that acquire noble-gas configurations when they form bonds.

| 1 A | 2 A | 3 A | 4 A | 5 A | 6 A | 7 A | 8 A |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |
| $\mathrm{Li} \cdot$ | $\mathrm{Be} \cdot$ | $\cdot \mathrm{B} \cdot$ | $\cdot \dot{\mathrm{C}} \cdot$ | $\cdot \dot{\mathrm{N}}:$ | $\cdot \dot{\mathrm{O}}:$ | $: \dot{\mathrm{E}}:$ | $: \stackrel{\ddot{\mathrm{N}} \mathrm{e}}{ }:$ |

## Reactions

- Reactions can be drawn showing the transfer of electrons
- Looking at tables of ionization energy and electron affinity along with the energies of some changes in state, lattice energies (2 slides ahead) and bond energies, the energy of a reaction can be calculated (Ex.- the enthalpy of formation of NaCl seven slides ahead) just like with Hess' law


## Ionic bonds and ionic crystals

- When atoms lose or gain electrons, they may acquire a noble gas configuration, but do not become noble gases.
- Because the two ions formed in a reaction between a metal and a nonmetal have opposite charges, they are strongly attracted to one another and form an ion pair.
- The net attractive electrostatic forces that hold the cations and anions together are ionic bonds.
- The highly ordered solid collection of ions is called an ionic crystal.


## Formation of a crystal of sodium chloride



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## Using Lewis symbols

to represent ionic bonding

- Lewis symbols can be used to represent ionic bonding between nonmetals and: the $s$-block metals, some $p$-block metals, and a few $d$-block metals.
- Instead of using complete electron configurations to represent the loss and gain of electrons, Lewis symbols can be used.



## Example

- Use Lewis symbols to show the formation of ionic bonds between magnesium and nitrogen.
- What are the name and formula of the compound that results?
- (worked out next slide)


## Example worked out

## Strategy

First, we determine the number of electrons that must be lost by magnesium atoms and gained by nitrogen atoms so that the resulting ions have noble-gas electron configurations.
Then we can combine Lewis symbols for the ions in the appropriate proportions to produce a neutral formal unit, from which we can deduce the compound's name and formula.

## Solution

Mg atoms (group 2A) lose their two valence electrons, and N atoms (group 5A) gain three additional valence electrons. To produce an electrically neutral formula unit, three Mg atoms must lose a total of six electrons and two N atoms must gain a total of six:


The compound is magnesium nitride, $\mathrm{Mg}_{3} \mathrm{~N}_{2}$.

## $\square$ Exercise

Use Lewis symbols to show the formation of ionic bonds between barium and iodine.
What are the name and formula of the compound that results?

## $\square$ Exercise

Use Lewis symbols to show the formation of ionic bonds in aluminum oxide.

## Energy changes in ionic compound formation

$$
\begin{array}{ll}
\mathrm{Na}(\mathrm{~g}) \rightarrow \mathrm{Na}^{+}(\mathrm{g})+\mathrm{e}^{-} & I E_{1}=+496 \mathrm{~kJ} / \mathrm{mol} \\
\mathrm{Cl}(\mathrm{~g})+\mathrm{e}^{-} \rightarrow \mathrm{Cl}^{-}(\mathrm{g}) & E A_{1}=-349 \mathrm{~kJ} / \mathrm{mol}
\end{array}
$$

- From the data above, it doesn't appear that the formation of NaCl from its elements is energetically favored. However ...
- ... the enthalpy of formation of the ionic compound is more important than either the first ionization energy or electron affinity.
- The overall enthalpy change can be calculated using a step-wise procedure called the Born-Haber cycle.


## Energy changes in <br> ionic compound formation (cont'd)

- The Born-Haber cycle is a hypothetical process, in which $\Delta H_{\mathrm{f}}$ is represented by several steps.
- What law can be used to find an enthalpy change that occurs in steps??


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## Example

- Use the following data to determine the lattice energy of $\mathrm{MgF}_{2}(\mathrm{~s})$ :
- enthalpy of sublimation of $\mathrm{Mg}:+146 \mathrm{~kJ} / \mathrm{mol}$;
$-I_{1}$ for Mg: +738kJ/mol;
$-I_{2}$ for Mg: +1451kJ/mol;
- bond-dissociation energy of $F_{2}(g)$ : $+159 \mathrm{~kJ} / \mathrm{mol} \mathrm{F}_{2}$;
- electron affinity of $F$ : $-328 \mathrm{~kJ} / \mathrm{mol} F$;
- enthalpy of formation of $\mathrm{MgF}_{2}(\mathrm{~s})$ :
-1124kJ/mol
- (worked out next slide)


## Example worked out

## Strategy

The approach we need here differs in three ways from the one used for $\mathrm{NaCl}(\mathrm{s})$. (1) The compound $\mathrm{MgF}_{2}(\mathrm{~s})$ has two anions for each cation in the crystal. Therefore, in the step where $F_{2}(g)$ dissociates, we need the bond-dissociation energy based on 1 mol of $\mathrm{F}_{2}(\mathrm{~g})$ in order to get 2 mol of $\mathrm{F}(\mathrm{g})$. Similarly, in the electron affinity step, we must produce 2 mol of $\mathrm{F}^{-}(\mathrm{g})$ rather than 1 mol . (2) Because the magnesium cation carries a $2+$ charge, we must include two ionization steps and both $I_{1}$ and $I_{2}$ in our calculation. (3) The enthalpy of formation of $\mathrm{MgF}_{2}(\mathrm{~s})$ is given, and our unknown is the lattice energy.

## Solution

The setup that follows incorporates all of the steps needed to determine the unknown lattice energy of $\mathrm{MgF}_{2}(\mathrm{~s})$ from the given data.

| Sublimation: | $\mathrm{Mg}(\mathrm{s}) \longrightarrow \mathrm{Mg}(\mathrm{g})$ | $\Delta H_{1}=+146 \mathrm{~kJ}$ |
| :---: | :---: | :---: |
| Bond dissociation: | $\mathrm{F}_{2}(\mathrm{~g}) \longrightarrow 2 \mathrm{~F}(\mathrm{~g})$ | $\Delta H_{2}=+159 \mathrm{~kJ}$ |
| First ionization: | $\mathbf{M g}(\mathrm{g}) \longrightarrow \mathrm{Mg}^{+}(\mathrm{g})+\mathrm{e}^{-}$ | $\Delta H_{3}=+738 \mathrm{~kJ}$ |
| Second ionization: | $\mathrm{Mg}^{+}(\mathrm{g}) \longrightarrow \mathrm{Mg}^{2+}(\mathrm{g})+\mathrm{e}^{-}$ | $\Delta H_{4}=+1451 \mathrm{~kJ}$ |
| Electron gain: | $2 \mathrm{~F}(\mathrm{~g})+2 \mathrm{e}^{-} \longrightarrow 2 \mathrm{~F}^{-}(\mathrm{g})$ | $\Delta H_{5}=2(-328 \mathrm{~kJ})$ |
| Unknown: | $\mathrm{Mg}^{2+}(\mathrm{g})+2 \mathrm{~F} \longrightarrow \mathrm{MgF}_{2}(\mathrm{~s})$ | $\Delta H_{6}=$ lattice energy |
| Overall: | $\mathrm{Mg}(\mathrm{s})+\mathrm{F}_{2}(\mathrm{~g}) \longrightarrow \mathrm{MgF}_{2}(\mathrm{~s})$ | $\Delta H_{f}^{\circ}=-1124 \mathrm{~kJ}$ |
| The enthalpy of formation is equal to the sum of the energies of the individual processes. | $\Delta H_{f}^{\circ}=-1124 \mathrm{~kJ}=(146+159+738$ | ) $\mathrm{kJ}+$ lattice energy |
| Rearranging the summed expression, we solve for the lattice energy. | Lattice energy $=(-1124-146-159$ | $+656) \mathrm{kJ}=-2962 \mathrm{~kJ}$ |
| The lattice energy is $-2962 \mathrm{~kJ} / \mathrm{mol} \mathrm{MgF} 2$ ( $\mathbf{s}$ ) . |  |  |

## Lewis structures of simple molecules

- A Lewis structure is a combination of Lewis symbols that represents the formation of covalent bonds between atoms.
- In most cases, a Lewis structure shows the bonded atoms with the electron configuration of a noble gas; that is, the atoms obey the octet rule. (H obeys the duet rule.)
- The shared electrons can be counted for each atom that shares them, so each atom may have a noble gas configuration.


## Lewis structures (cont'd)

- The shared pairs of electrons in a molecule are called bonding pairs.
- The bonding pair is represented by a dash (-).
- The other electron pairs, which are not shared, are called nonbonding pairs, or lone pairs.
- Two electrons from one element form a coordinate covalent bond (like in $\mathrm{H}_{3} \mathrm{O}^{+}$).



## Some illustrative compounds

Lewis
symbol:
Lewis
structure:
Name and formula:
Geometric shape:

4A



Methane, $\mathrm{CH}_{4}$

6A


- $\ddot{\mathrm{O}}:$


Water, $\mathrm{H}_{2} \mathrm{O} \quad$ Hydrogen fluoride, HF


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7A
7 A
$\cdot \stackrel{.}{\mathrm{F}}:$



- Note that the two-dimensional Lewis structures do not necessarily show the correct shapes of the three-dimensional molecules. Nor are they intended to do so.
- The Lewis structure for water may be drawn with all three atoms in a line: $\mathrm{H}-\mathrm{O}-\mathrm{H}$.
- We will learn how to predict shapes of molecules in Chapter 10.


## Multiple covalent bonds

- The covalent bond in which one pair of electrons is shared is called a single bond.
- Multiple bonds can also form:

In a double bond two pairs of electrons are shared.

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In a triple bond three pairs of electrons are shared.

Note that each atom obeys the octet rule, even with multiple bonds.
(Again, movement of a single electron should be represented by a single-headed arrow.)

## Polar covalent bonds and electronegativity

- Sharing is not always equal
- ELECTRONEGATIVITY (EN) is a measure of the ability of an atom to attract its bonding electrons to itself.
- EN is related to ionization energy and electron affinity.
- The greater the EN of an atom in a molecule, the more strongly the atom attracts the electrons in a covalent bond.

Increasing
Electronegativity generally increases from left to right within a period, and it generally increases from the bottom to the top within a group.


Increasing
electro-
negativity

## Pauling's electronegativities



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## Electronegativity (cont'd)

- The greater the difference in electronegativity the greater the ionic character...the more ionic character to the bond, the more polar it is.
- A diatomic molecule is completely nonpolar.
- Polarity is drawn using an arrow with a cross through the tail (3 slides ahead).
- "Partially positive" atoms are denoted by a "delta-plus" ( $\delta+$ ), partially negative atoms are denoted by a "delta-minus"( $\delta-)$.


## Electronegativity difference and bond type



## Depicting polar covalent bonds



## Example

- Use electronegativity values to arrange the following bonds in order of increasing polarity:
$\mathrm{Br}-\mathrm{Cl}, \mathrm{Cl}-\mathrm{Cl}, \mathrm{Cl}-\mathrm{F}, \mathrm{H}-\mathrm{Cl}, \mathrm{l}-\mathrm{Cl}$
- Worked out next slide


## Example worked out

## Strategy

We begin with the knowledge that the polar character of a bond is determined by electronegativity differences. For the individual electronegativities of the atoms, we turn to Figure 9.8.

## Solution

The electronegativities (EN) and electronegativity differences are

| EN: | 2.8 | 3.0 | 3.0 | 3.0 | 3.0 | 4.0 | 2.1 | 3.0 | 2.5 | 3.0 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | $\mathrm{Br}-\mathrm{Cl}$ | $\mathrm{Cl}-\mathrm{Cl}$ | $\mathrm{Cl}-\mathrm{F}$ | $\mathrm{H}-\mathrm{Cl}$ | $\mathrm{I}-\mathrm{Cl}$ |  |  |  |  |  |
| $\Delta \mathrm{EN}:$ | 0.2 | 0.0 | 1.0 | 0.9 | 0.5 |  |  |  |  |  |

The order of increasing polarity is therefore

$$
\mathrm{Cl}-\mathrm{Cl}<\mathrm{Br}-\mathrm{Cl}<\mathrm{I}-\mathrm{Cl}<\mathrm{H}-\mathrm{Cl}<\mathrm{Cl}-\mathrm{F} .
$$

## $\square$ Exercise 9.5A

Use electronegativity values to arrange the following bonds in order of increasing polarity:
$\mathrm{C}-\mathrm{Cl}, \mathrm{C}-\mathrm{H}, \mathrm{C}-\mathrm{Mg}, \mathrm{C}-\mathrm{O}, \mathrm{C}-\mathrm{S}$

## Exercise 9.5B

(a) Of the elements carbon, silicon, bromine, sulfur, nitrogen, chlorine, arsenic, fluorine, and selenium, which two form the covalent bond that has the most ionic character?
(b) Which two form the covalent bond that has the least ionic character?

## Writing Lewis structures: skeletal structures

- The SKELETAL STRUCTURE shows the arrangement of atoms.
- Lewis structures have terminal atoms drawn around a central atom....think of each atom as a four-sided box that must obey the octet (duet) rule.....draw dashes for single bonds, then add lone pairs of electrons to the terminal atoms to get an octet...then add lone pairs and/or form multiple bonds to central atoms as needed to account for the total number of valence electrons
- Hydrogen atoms are terminal atoms (bonded to only one other atom).
- The central atom of a structure usually has the lowest electronegativity.
- In oxoacids $\left(\mathrm{HClO}_{4}, \mathrm{HNO}_{3}\right.$, etc.) hydrogen atoms are usually bonded to oxygen atoms.
- Molecules and polyatomic ions usually have compact, symmetrical structures.
- $\mathrm{C}, \mathrm{N}, \mathrm{O}$, and S are often double bonded. C and N can be triple bonded.


## Example

## - Write the Lewis structure of nitrogen trifluoride, $\mathrm{NF}_{3}$.

## Solution

Step 1: Determine the number of valence electrons.

Step 2: Write a skeletal structure. The electronegativity of N is 3.0 ; that of F is 4.0. We expect a skeletal structure with N as a central atom and F as terminal atoms. The three nitrogen-to-fluorine bonds in this structure account for six electrons.

Step 3: Complete the octets of terminals atoms. We complete the octets of the F atoms by placing three lone pairs of electrons around each. This accounts for 18 additional electrons.

Step 4: Assign lone pairs to central atom(s). We have now assigned $6+18=24$ of the 26 valence electrons. We place the remaining two as a lone pair on the N atom.

In one N atom (group 5 A ) and three F atoms (group 7A), there are $5+(3 \times 7)=26$ valence electrons.


$: \stackrel{\ddot{\mathrm{F}}}{\bullet \rightarrow} \mathrm{M}-\ddot{\mathrm{F}}:$

- Write the Lewis structure of hydrazine, $\mathrm{N}_{2} \mathrm{H}_{4}$.


## Lewis structures are not always "right"The importance of experimental evidence

- The Lewis structure commonly drawn for oxygen is $: ~ \mathrm{O}=\mathrm{O}$ :
- But oxygen is paramagnetic, and therefore must have unpaired electrons.
- Lewis structures are a useful tool, but they do not always represent molecules correctly, even when the Lewis structure is plausible.



## Example

## Write a plausible Lewis structure for phosgene, $\mathrm{COCl}_{2}$.

Step 1: Determine the number of valence electrons.

Step 2: Write a skeletal structure. The electronegativities are 2.5 for C , 3.5 for O , and 3.0 for Cl . We expect C , the least electronegative atom, to be the central atom, and O and Cl to be terminal atoms attached to it. This skeletal structure accounts for six of the valence electrons.

Step 3: Complete the octets of terminal atoms. Place three lone pairs around the O atom and three lone pairs around each of the Cl atoms, for a total of 18 electrons. This completes the octets of the terminal atoms.

Step 4: Assign lone pairs to central atom(s). Up to this point, we have assigned $6+18=24$ of the 24 valence electrons, so there are none available to complete the octet of the central C atom. Therefore we must go to step 5.

Step 5: Form multiple bonds to complete octets of central atom(s). Complete the octet on the C atom by shifting a lone pair of electrons from the O atom to form a carbon-to-oxygen double bond. ( C and O are two atoms that are able to form a double bond.)

In one C atom (group 4A), one O atom (group 6A), and two Cl atoms $($ group 7A), there are $4+6+(2 \times 7)=24$ valence electrons.




## Formal charge

- Formal charge is the difference between the number of valence electrons in a free (uncombined) atom and the number of electrons assigned to that atom when bonded to other atoms in a Lewis structure.
- Formal charge is a hypothetical quantity; a useful tool in predicting reactivity.
- Usually, the most plausible Lewis structure is one with no formal charges.
- When formal charges are required, they should be as small as possible.
- Negative formal charges should appear on the most electronegative atoms. Makes sense, right?
- Adjacent atoms in a structure should not carry formal charges of the same sign.


## Formal charge illustrated



## Example

- Before, we wrote a Lewis structure for the molecule $\mathrm{COCl}_{2}$, shown here as structure (a). Show that structure (a) is more plausible than (b) or (c).

(a)

(b)

(c)


## Resonance: delocalized bonding

- When a molecule or ion can be represented by two or more plausible Lewis structures that differ only in the distribution of electrons, the true structure is a composite, a hybrid, of them.
- The different plausible structures are called resonance structures.
- The actual molecule or ion that is a hybrid of the resonance structures is called a resonance hybrid.
- Electrons that are part of the resonance hybrid are spread out over several atoms and are referred to as being delocalized.


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## Example

- Write three equivalent Lewis structures for the $\mathrm{SO}_{3}$ molecule that conform to the octet rule, and describe how the resonance hybrid is related to the three structures.

Step 1: Determine the number of valence electrons.
Step 2: Write a skeletal structure. The skeletal structure has sulfur, the atom of lower electronegativity, as the central atom.

Step 3: Complete the octets of terminal atoms. Place three lone pairs of electrons on each O atom to complete its octet.

Step 4: Assign lone pairs to central atom(s). We have already assigned all 24 valence electrons; there are none left to place as lone pairs on the central atom. Because the $S$ atom does not yet have an octet, we must go on to step 5 .

Step 5: Form multiple bonds to complete octets of central atom(s). Move a lone pair of electrons from any terminal $O$ atom to form a double bond to the central $S$ atom. Because the double bond can go to any one of the three terminal O atoms, we get three equivalent structures that differ only in the position of the double bond.

There are $6+(3 \times 6)=24$ valence electrons.



## Molecules that do not follow

## the octet rule

- Molecules with an odd number of valence electrons have at least one of them unpaired and are called free radicals.
- Some molecules have incomplete octets. These are usually compounds of $\mathrm{Be}, \mathrm{B}$, or Al ; they generally have some unusual bonding characteristics, and are often quite reactive.
- Some compounds have expanded valence shells, which means that the central atom has more than eight electrons around it.
- A central atom can have expanded valence if it is in the third period or lower (i.e., S, CI, P).


## Example

- Write the Lewis structure for bromine pentafluoride, $\mathrm{BrF}_{5}$.

Step 1: Determine the number of valence electrons.

Step 2: Write a skeletal structure. The skeletal structure has Br , the atom of lower electronegativity, as the central atom.

Both Br and F are in group 7A. All the atoms in the structure have seven valence electrons, and $7+(5 \times 7)=42$ valence electrons must appear in the Lewis structure.



Step 4: Assign lone pairs to central atom(s). Through step 3, we have assigned 40 of the 42 valence electrons; two remain to be placed. They can be shown as a lone pair on the bromine atom. In this representation, the Br atom has an expanded valence shell containing 12 electrons.

## Example

- Indicate the error in each of the following Lewis structures. Replace each by a more acceptable structure(s).
a. $: C \equiv \mathrm{~N}$ :



## Bond order and bond length

- Bond order is the number of shared electron pairs in a bond.
- $A$ single bond has $B O=1$, $a$ double bond has $\mathrm{BO}=2$, etc.
- Bond length is the distance between the nuclei of two atoms joined by a covalent bond.
- Bond length depends on the particular atoms in the bond and on the bond order.



## Bond energy

- Bond-dissociation energy ( $\boldsymbol{D}$ ) is the energy required to break one mole of a particular type of covalent bond in a gas-phase compound.
- Energies of some bonds can differ from compound to compound, so we use an average bond energy.



## Table 9.1 Some Representative Bond Lengths and Bond Energies

|  | Bond <br> Length, <br> pm | Bond <br> Energy, <br> $\mathbf{k J} / \mathbf{m o l}$ | Bond | Bond <br> Length, <br> $\mathbf{p m}$ | Bond <br> Energy, <br> $\mathbf{k J} / \mathbf{m o l}$ |
| :--- | :---: | :---: | :--- | :---: | :---: |
| $\mathrm{H}-\mathrm{H}$ | 74 | 436 | $\mathrm{C}-\mathrm{O}$ | 143 | 360 |
| $\mathrm{H}-\mathrm{C}$ | 110 | 414 | $\mathrm{C}=\mathrm{O}$ | 120 | $736^{\mathrm{b}}$ |
| $\mathrm{H}-\mathrm{N}$ | 100 | 389 | $\mathrm{C}-\mathrm{Cl}$ | 178 | 339 |
| $\mathrm{H}-\mathrm{O}$ | 97 | 464 | $\mathrm{~N}-\mathrm{N}$ | 145 | 163 |
| $\mathrm{H}-\mathrm{S}$ | 132 | 368 | $\mathrm{~N}=\mathrm{N}$ | 123 | 418 |
| $\mathrm{H}-\mathrm{F}$ | 92 | 565 | $\mathrm{~N}-\mathrm{O}$ | 110 | 946 |
| $\mathrm{H}-\mathrm{Cl}$ | 127 | 431 | $\mathrm{~N}=\mathrm{O}$ | 136 | 222 |
| $\mathrm{H}-\mathrm{Br}$ | 141 | 364 | $\mathrm{O}-\mathrm{O}$ | 120 | 590 |
| $\mathrm{H}-\mathrm{I}$ | 161 | 297 | $\mathrm{O}=\mathrm{O}$ | 145 | 142 |
| $\mathrm{C}-\mathrm{C}$ | 154 | 347 | $\mathrm{~F}-\mathrm{F}$ | 121 | 498 |
| $\mathrm{C}=\mathrm{C}$ | 134 | 611 | $\mathrm{Cl}-\mathrm{Cl}$ | 199 | 159 |
| $\mathrm{C} \equiv \mathrm{C}$ | 120 | 837 | $\mathrm{Br}-\mathrm{Br}$ | 228 | 193 |
| $\mathrm{C}-\mathrm{N}$ | 147 | 305 | $\mathrm{I}-\mathrm{I}$ | 266 | 151 |
| $\mathrm{C}=\mathrm{N}$ | 128 | 615 |  |  |  |
| $\mathrm{C} \equiv \mathrm{N}$ | 116 | 891 |  |  |  |

${ }^{\text {a }}$ Bond-dissociation energy for the bonds in diatomic molecules $\left(\mathrm{H}_{2}, \mathrm{HF}, \mathrm{HCl}, \mathrm{HBr}, \mathrm{HI}, \mathrm{N}_{2}, \mathrm{O}_{2}, \mathrm{~F}_{2}, \mathrm{Cl}_{2}, \mathrm{Br}_{2}\right.$ and $\mathrm{I}_{2}$ ) and average bond energies for the other bonds.
${ }^{\mathrm{b}}$ The value for the CO bond in $\mathrm{CO}_{2}$ is considerably different: $799 \mathrm{~kJ} / \mathrm{mol}$.
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## Trends in bond lengths and energies

- The higher the order (for a particular type of bond), the shorter and the stronger (higher energy) the bond.
- $\mathrm{A} \mathrm{N}=\mathrm{N}$ double bond is shorter and stronger than a $\mathrm{N}-\mathrm{N}$ single bond.
- There are four electrons between the two positive nuclei in $\mathrm{N}=\mathrm{N}$. This produces more electrostatic attraction than the two electrons between the nuclei in $\mathrm{N}-\mathrm{N}$.


## Example

- Estimate the length of (a) the nitrogen-to-nitrogen bond in $\mathrm{N}_{2} \mathrm{H}_{4}$ and (b) the bond in BrCl .
(a) The Lewis structure

is a plausible one: valence-shell octets for the N atoms, duets for the H atoms, and formal charges of 0 . The nitrogen-to-nitrogen bond is single, and from Table 9.1 we find its bond length to be 145 pm .
(b) To get the Lewis structure of BrCl , we imagine substituting one Br atom for one Cl atom in the Lewis structure of $\mathrm{Cl}_{2}$, arriving at the structure


The BrCl molecule contains a $\mathrm{Br}-\mathrm{Cl}$ single bond with a length approximately onehalf the $\mathrm{Cl}-\mathrm{Cl}$ bond length plus one-half the $\mathrm{Br}-\mathrm{Br}$ bond length. From Table 9.1, we get $\left[\left(\frac{1}{2} \times 199\right)+\left(\frac{1}{2} \times 228\right)\right]=214 \mathrm{pm}$.

## Calculations involving bond energies

For the reaction $\mathrm{N}_{2}(\mathrm{~g})+2 \mathrm{H}_{2}(\mathrm{~g}) \rightarrow \mathrm{N}_{2} \mathrm{H}_{4}(\mathrm{~g})$ to occur ...


## Flashback: alkenes and alkynes

- Hydrocarbons with double or triple bonds between carbon atoms are called unsaturated hydrocarbons.
- Alkenes are hydrocarbons with one or more $\mathrm{C}=\mathrm{C}$ double bonds.
- The simplest alkene is $\mathrm{C}_{2} \mathrm{H}_{4}$, ethene (ethylene).
- Alkynes are hydrocarbons that have one or more carbon-carbon triple bonds.
- The simplest alkyne is $\mathrm{C}_{2} \mathrm{H}_{2}$, ethyne (acetylene).


## Molecular models of ethene and ethyne

## space-filling models



Structural formula
$\mathrm{CH}_{2}=\mathrm{CH}_{2} \quad$ or $\quad \mathrm{H}_{2} \mathrm{C}=\mathrm{CH}_{2}$
Condensed structural formula
$\mathrm{H}: \mathrm{C}:: \mathrm{C}: \mathrm{H} \quad$ or $\quad \mathrm{H}-\mathrm{C} \equiv \mathrm{C}-\mathrm{H}$

## Fats and oils

- Triglycerides are composed of glycerol plus three long-chain carboxylic (fatty) acids
- What's the difference between a fat and an oil? Fats are solids at RT, Oils are liquids
- Unsaturated fats become saturated upon hydrogenation (Fig. 13.21)
- Your book also has a nice bromination picture in this chapter



## Polymers

- Polymers are compounds in which many identical molecules have been joined together.
- Monomers are the simple molecules which join together to form polymers.
- Often, the monomers have double or triple bonds.
- The process of these molecules joining together is called polymerization.
- Many everyday products and many biological compounds are polymers.


## Formation of polyethylene



Table 9.2 A Selection of Addition Polymers

| Monomer | Polymer | Polymer Name |
| :---: | :---: | :--- |
| $\mathrm{H}_{2} \mathrm{C}=\mathrm{CH}_{2}$ | Some Uses |  |

## Plastics

- Although plastics are marked recyclable 17 , most areas only recycle plastics 1 and 2



## end of section

## Flashback / flashforward

- We've learned how to write chemical formulas and line structures...
- We've learned how to show the locations of electrons in Lewis structures...
- How can we determine the actual shape of a molecule (and/or the geometry around atoms) from these things?


## Valence-Shell Electron-Pair Repulsion theory (VSEPR theory)

- Electron pairs (bonded and lone pairs) will orient themselves so that they are as far apart from one another as possible (the four electrons in a double bond and the six in a triple bond are each considered one "group")
- A lone pair takes up more space around an atom than a bonded pair



## Electron group geometry

- 2 electron pairs- linear
- 3 electron pairs- trigonal planar
- 4 electron pairs- tetrahedral
- 5 electron pairs- trigonal bipyramidal
- 6 electron pairs- octahedral


## $e^{-}$group geometry (cont'd)



## VSEPR notation

- central atoms are denoted "A"
- terminal atoms are denoted "X"
- Ione pairs are denoted "E"


Thus, water is $\mathrm{AX}_{2} \mathrm{E}_{2}$
X

## Molecular geometries

- $A X_{2}$ - linear
- $\mathrm{AX}_{3}$ - trigonal planar
- $\mathrm{AX}_{4}$ - tetrahedral
- $A X_{5}$ - trigonal bipyramidal
- $\mathrm{AX}_{6}$ - octahedral
- The $A X_{5}$ and $A X_{6}$ require an expanded valence shell and, therefore, the central atom must be a third-period or higher element.

> Molecular shape vs. electron group geometry... MOST BOOKS ARE CONFUSING!

- For structures with no lone pairs on the central atom (AXn), the molecular geometrySHAPE is the same as the electron-group geometry.
- When there are lone pairs, the molecular geometry SHAPE is derived from the electron-group geometry.
- In either case, the electron-group geometry is the tool we use to obtain the molecular geometry SHAPE.


## Molecular shape vs. electron group geometry

- The presence of lone pairs affects ONLY the SHAPE of the molecule (which is described in terms of locations of atoms), NOT the electron group geometry (sometimes referred to as the molecular geometry) (which takes into account lone pairs)
- $\mathrm{AX}_{2} \mathrm{E}$ vs $\mathrm{AX}_{3}$
- $\mathrm{AX}_{3} \mathrm{E}$ vs $\mathrm{AX}_{4}$
- $\mathrm{AX}_{2} \mathrm{E}_{2}$ vs $\mathrm{AX}_{4}$
- $\mathrm{AX}_{4} \mathrm{E}$ vs $\mathrm{AX}_{5}$
- $\mathrm{AX}_{3} \mathrm{E}_{2}$ vs $\mathrm{AX}_{5}$
- $\mathrm{AX}_{2} \mathrm{E}_{3}$ vs $\mathrm{AX}_{5}$
- $A X_{5} E$ vs $A X_{6}$
- $\mathrm{AX}_{4} \mathrm{E}_{2}$ vs $\mathrm{AX}_{6}$


## Shape and geometry



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## Shape and geometry (cont'd)

Table 10.1 (Part 2) VSEPR Notation, Electron-Group Geometry, and Molecular Geometry

| Number of Electron Groups | ElectronGroup Geometry | Number of Lone Pairs | VSEPR <br> Notation | Molecular Geometry |  | Example | Molecular Model |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| 4 | Tetrahedral | 0 | $\mathrm{AX}_{4}$ |  | $109.5^{\circ}$ | $\mathrm{CH}_{4}$ |  |

4
Tetrahedral $1 \quad \mathrm{AX}_{3} \mathrm{E}$

$109.5^{\circ}$
$\mathrm{NH}_{3}$

Trigonal
pyramidal

## Shape and geometry (cont'd)

Table 10.1 (Part 3) VSEPR Notation, Electron-Group Geometry, and Molecular Geometry

| Number <br> of | Electron- | Number <br> of |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Electron | Group | Lone | VSEPR | Molecular | Ideal |  |
| Groups | Geometry | Pairs | Notation | Geometry | Angles | Example |

4
Tetrahedral
2
$\mathrm{AX}_{2} \mathrm{E}_{2}$

$109.5^{\circ} \quad \mathrm{OH}_{2}$

5 Trigonal
bipyramida
0
$\mathrm{AX}_{5}$


Trigonal bipyramidal


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## Shape and geometry (cont'd)

Table 10.1 (Part 4) VSEPR Notation, Electron-Group Geometry, and Molecular Geometry

| Number of Electron Groups | ElectronGroup Geometry | Number of Lone Pairs | VSEPR <br> Notation | Molecular Geometry | Ideal Bond Angles | Example | Molecular Model |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |

5

| Trigonal <br> bipyramidal | 1 | $\mathrm{AX}_{4} \mathrm{E}$ |
| :---: | :---: | :---: |



5

$$
\begin{array}{cll}
\text { Trigonal } & 2 & \mathrm{AX}_{3} \mathrm{E}_{2} \\
\text { bipyramidal }
\end{array}
$$



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## Shape and geometry (cont'd)



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## Shape and geometry (cont'd)

Table 10.1 (Part 6) VSEPR Notation, Electron-Group Geometry, and Molecular Geometry

| Number <br> of | Electron- | Number <br> of |  |  |  |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Electron | Group | Lone | VSEPR | Molecular | Ideal <br> Bond |  |  |
| Groups | Geometry | Pairs | Notation | Geometry | Angles | Example | Molecular Model |
|  |  |  |  |  |  |  |  |

6 Octahedral

1
$\mathrm{AX}_{5} \mathrm{E}$


Square pyramidal

6
Octahedral
2


Square
planar

## Example

- Use the VSEPR method to predict the shape of the nitrate ion.


## Shape of methane


(a)


VSEPR notation: $\mathrm{AX}_{4}$
(b)

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## Molecular shape of water



## VSEPR notation: $\mathrm{AX}_{2} \mathrm{E}_{2}$

## Complex molecules

- For molecules with more than one central atom, the geometry and resulting shape around each atom must be evaluated


## Electronegativity revisited-polarity

 and dipole moment- Molecular dipoles
- Molecular shape and dipoles


## Polar molecules in an electric field



## Molecular shapes and dipole moments

- To predict molecular polarity:

1. Use electronegativity values to predict bond dipoles.
2. Use the VSEPR method to predict the molecular shape.
3. From the molecular shape, determine whether bond dipoles cancel to give a nonpolar molecule, or combine to produce a resultant dipole moment for the molecule. (Can you cut the molecule in half twice?)

- Note: Lone-pair electrons can also make a contribution to dipole moments.


## Example

- Explain whether you expect the following molecules to be polar or nonpolar:
(a) $\mathrm{CHCl}_{3}$
(b) $\mathrm{CCl}_{4}$

The distribution of bond dipoles in $\mathrm{CCI}_{4}$ is shown in Figure 10.9a. The resultant dipole in the downward direction produced by the three bond dipoles directed down and away from the center of the structure is just matched by the straight upward bond dipole at the top. Thus all the bond dipoles cancel, so there is no molecular dipole, and $\mathrm{CCl}_{4}$ is a nonpolar molecule.

(a)

(b)

A FIGURE 10.9 Molecular geometry and molecular dipoles
The bond dipoles are represented by black cross-based arrows $\longrightarrow \longrightarrow$
(a) The individual bond dipoles cancel, and there is no molecular dipole.
(b) All the bond dipoles point downward and combine to form a molecu-
lar dipole (red arrow).

## Example

- Of the two compounds NOF and $\mathrm{NO}_{2} \mathrm{~F}$, one has $\mu=1.81 \mathrm{D}$ and the other has $\mu=0.47 \mathrm{D}$. Which dipole moment do you predict for each compound? Explain.


In NOF, there are two bond dipoles and both point downward, leading to a net downward molecular dipole. In $\mathrm{NO}_{2} \mathrm{~F}$, the upward-pointing $\mathrm{N}-\mathrm{O}$ bond dipole opposes the other two, and consequently we expect a smaller molecular dipole. Our prediction is therefore NOF, $\mu=1.81 \mathrm{D}$, and $\mathrm{NO}_{2} \mathrm{~F}, \mu=0.47 \mathrm{D}$.

## BIG POINT

- Bonds are formed by the overlap of orbitals


## Bonding in $\mathrm{H}_{2} \mathrm{~S}$


(a) Isolated atoms


The hydrogen atoms' $s$ orbitals can overlap with the two halffilled $p$ orbitals on sulfur.

## Hybridization of atomic orbitals

- Often, the number of equivalent bonds around a central atom cannot be adequately explained using "conventional orbitals"
- $\mathrm{sp}^{3}$ orbitals
- $\mathrm{sp}^{2}$ orbitals
- sp orbitals
- d hybrids


## Important points of VB theory

- Most of the electrons in a molecule remain in the same orbital locations that they occupied in the separated atoms.
- Bonding electrons are localized in the region of AO overlap.
- For AOs with directional lobes (such as p orbitals), maximum overlap occurs when the AOs overlap end to end.
- VB theory is not without its problems ...


## Hybridization of atomic orbitals

VB theory: carbon should have just two bonds, and they should be about $90^{\circ}$ apart.
But $\mathrm{CH}_{4}$ has four $\mathrm{C}-\mathrm{H}$ bonds, $109^{\circ}$ apart.
Ground-state electron configuration of C :
[He]


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- We can hybridize the four orbitals holding valence electrons; mathematically combine the wave functions for the $2 s$ orbital and the three $2 p$ orbitals on carbon.
- The four AOs combine to form 4 new $s p^{3}$ hybrid AOs.
- The four hybrid AOs are degenerate (same energy) and each has a single electron (Hund's rule).


## The $\mathrm{sp}^{3}$ hybridization scheme



## Methane and ammonia



## $\mathrm{sp}^{2}$ hybridization

- Three $\boldsymbol{s} \boldsymbol{p}^{2}$ hybrid orbitals are formed from an $s$ orbital and two $p$ orbitals.
- The empty $p$ orbital remains unhybridized. It may be used in a multiple bond.
- The $s p^{2}$ hybrid orbitals are in a plane, $120^{\circ}$ apart.
- This distribution gives a trigonal planar molecular geometry, as predicted by VSEPR.



## The $\mathrm{sp}^{2}$ hybridization scheme in boron



## sp hybridization

- Two sp hybrid orbitals are formed from an s orbital and a $p$ orbital.
- Two empty $p$ orbitals remains unhybridized; the $p$ orbitals may be used in a multiple bond.
- The $s p$ hybrid orbitals are $180^{\circ}$ apart.
- The geometry around the hybridized atom is linear, as predicted by VSEPR.
sp hybridization in Be : [He]


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## sp hybridization in Be



# Hybrid orbitals involving d subshells 

- This hybridization allows for expanded valence shell compounds.
- By hybridizing one s, three $p$, and one $d$ orbital, we get five $\boldsymbol{s} \boldsymbol{p}^{3} \boldsymbol{d}$ hybrid orbitals.
- This hybridization scheme gives trigonal bipyramidal electron-group geometry.



## Hybrid orbitals involving d subshells (cont'd)

- By hybridizing one $s$, three $p$, and two d orbitals, we get five $\boldsymbol{s p}^{3} d^{2}$ hybrid orbitals.
- This hybridization scheme gives octahedral geometry.


## Predicting hybridization schemes

In the absence of experimental evidence, probable hybridization schemes can be predicted:

1. Write a plausible Lewis structure for the molecule or ion.
2. Use the VSEPR method to predict the electron-group geometry of the central atom.
3. Select the hybridization scheme that corresponds to the VSEPR prediction.
4. Describe the orbital overlap and molecular shape.

## Example

$s p^{3} d^{2}$ hybridization in I:



- lodine pentafluoride, $\mathrm{IF}_{5}$, is used commercially as a fluorinating agent- a substance that, via a chemical reaction, introduces fluorine into other compounds. Describe a hybridization scheme for the central atom, and sketch the molecular geometry of the $\mathrm{IF}_{5}$ molecule.

© FIGURE 10.19 Bonding scheme for iodine pentafluoride, $\mathrm{IF}_{5}$ The orbitals around the central I atom are six $s p^{3} d^{2}$ hybrid orbitals. One of them (red) is occupied by the LP electrons. The other five are the bonding orbitals. Each bond involves the overlap of an I $s p^{3} d^{2}$ hybrid orbital with a $2 p$ atomic orbital of a terminal F atom. Because of repulsions between the LP electrons and the bonding pairs, the (imaginary) plane at the base of the molecule is raised slightly above the I atom.

$$
\begin{gathered}
\text { Hybrid orbitals and multiple } \\
\text { bonds }
\end{gathered}
$$

- Orbitals cannot overlap the same region of space
- Double and triple bonds result from the overlap of multiple sets of orbitals in different regions of space

$$
\begin{aligned}
& \text { Hybrid orbitals and } \\
& \text { multiple covalent bonds }
\end{aligned}
$$

- Covalent bonds formed by the end-to-end overlap of orbitals are called sigma ( $\sigma$ ) bonds.
- All single bonds are sigma bonds.
- A bond formed by parallel, or side-by-side, orbital overlap is called a pi $(\pi)$ bond.
- A double bond is made up of one sigma bond and one pi bond.
- A triple bond is made up of one sigma bond and two pi bonds.


# VB theory for ethylene, $\mathrm{C}_{2} \mathrm{H}_{4}$ 


(a) The $\sigma$-bond framework
> $\pi$-bond has two lobes (above and below plane), but is one bond. Side overlap of $2 p-2 p$.

(b) The formation of a $\pi$-bond by the overlap of the half-filled $2 p$ orbitals

(c) Hybridization and bonding scheme

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## VB theory for acetylene, $\mathrm{C}_{2} \mathrm{H}_{2}$


(c) Hybridization and bonding scheme

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## Geometric isomers

- Same formula, different arrangement in space
- Geometric isomers are isomers that differ only in the geometric arrangement of certain substituent groups.
- Two types of geometric isomers include:
- cis: substituent groups are on the same side
- trans: substituent groups are on opposite sides
- cis- and trans- compounds are distinctly different in both physical and chemical properties.
- Usually formed across double bonds and in square planar compounds.


## Geometric isomerism in 2-butene


n-Butane does not have these isomers; why not??

## Example

- Is it possible to write a unique structural formula for 1,2-dichloroethene if we are told that the molecule is nonpolar?


## Molecular orbital theory

- An altexnative scheme to VB theory uses molécular orbitals.
- A molecular rerbital (MO) is a mathematical description of the region in a molecule where there is a high probability of finding electrons.
- In MO theory, molecular oxbitals are formed by the combination of atomic orbitals.

Characteristics of MOs

- Two atofic orbitals combine => two molecular orbitals result.
- Of each pair df molecular orbitals, one is a bonding molecuar orbital.
- The bonding orbitais at a lower energy than the separate atomic orbitas.
- Electrons in a bonding okital increase the stability of the molecule.
- The second orbital is an antrizonding orbital.
- The antibonding orbital is at a higker energy than the AOs.
- Electrons in an antibonding orbital decrease the stability of the molecule.
- There are nonbonding orbitals which we yill not discuss.




## MO diagrams of diatomic molecules of the



## Jargon

- The Highest Occupied Molecular Orbital is called the HOMO
- The Lowest Dnoccupied Molecular Orbital is called LUMO


## Example

- When in electron is removed from a $\mathrm{N}_{2}$ molecule, forming an $\mathrm{N}_{2}{ }^{+}$ion, the bond between the $N$ atoms is weakened. When an $\mathrm{O}_{2}$ molecule is ionized to $\mathrm{O}_{2}{ }^{+}$, the bond between O atoms is strengthened. Explainthis difference.


## Aromatic compounds

- Touched on briefly earlier...
- Many of the first benzene-like compounds discovered had pleasant odors, hence the name aromatic was applied to the compounds.
- Today an aromatic compound is one that has a ring structure and bonding characteristics related to those of benzene (more in Chapter 23).
- All organic compounds that are not aromatic are called aliphatic compounds.


## Aromatics - benzene

- In 1865, Kekulé proposed that benzene $\left(\mathrm{C}_{6} \mathrm{H}_{6}\right)$ has a cyclic structure, with a hydrogen atom attached to each carbon atom. Alternating single and double bonds join the carbon atoms.

- Modern view: there are two resonance hybrids of benzene.
- The pi-electrons are not localized between any particular carbon atoms, but are delocalized among all six carbon atoms.


## The $\sigma$-bonding framework in benzene



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## The $\pi$-bonding framework in benzene



## TABLE 10.3 Some Representative Aromatic Compounds

Name

Aniline
Structure

Benzoic acid

Bromobenzene


Nitrobenzene

Phenol

Toluene


## Typical Use(s)

Starting material for the synthesis of dyes, drugs, resins, varnishes, perfumes; solvent; vulcanizing rubber

Food preservative; starting material for the synthesis of dyes and other organic compounds; curing of tobacco

Starting material for the synthesis of many other aromatic compounds; solvent; motor oil additive

Starting material for the synthesis of aniline; solvent for cellulose nitrate; in soaps and shoe polish

Disinfectant; starting material for the synthesis of resins, drugs, and other organic compounds

Solvent; gasoline octane booster; starting material for the synthesis of benzoic acid, benzaldehyde, and many other organic compounds

## Band theory

- In the tree-electron model, a metal consists of more-or-less immobile metal ions in a crystal lattice, surrounded Dya "gas" of the valence electrons.



## Deformation of a metal <br> <empared to an ionic solid



## Explanation- band theory

- The free electron model is a classical theory, which is less satisfactory in many ways than a quantummechanical treatment of bonding in metals.
- Band theory is Aquantum-mechanical model.

The spacing between electron energy levels is so minute in metals that the levels essentially merge into a band.


## B Band theory (cont'd)

- When band is occupied by valence electrons, it is called a valence band. (Akin to the HOMO)
- In band theory the presence of a conduction band-a partially filled band of energy levels-is required for conductivity.
- Because the energy lexels in bands are so closely spaced, there are electronic transitions in a partially filled band that match in enexgy every component of visible light.
- Metals therefore absorb the light that falls on them and are opaque.
- At the same time electrons that have absorbed energy from incident light are very effective in radiating light of the same frequency-metals are highly reflective.


the end


[^0]:    ${ }^{\mathrm{a}}$ In the pseudo-noble gas configuration, all valence electrons are lost and the remaining $(n-1)$ shell has 18 electrons in the configuration $(n-1) s^{2}(n-1) p^{6}(n-1) d^{10}$.
    ${ }^{\mathrm{b}}$ In the $18+2$ configuration, $(n-1) s^{2}(n-1) p^{6}(n-1) d^{10} n s^{2}$, two valence electrons remain.
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