Survey of Elements

Hydrogen 1s¹
Naturally, loss of one e⁻ dominates 1s

Helium

 $1s^2$



- closed shell
- very small atom
 All noble gases have very high first I.E.'s which is in
 line with their chemical inertness

First short period of elements

Li, Be, B, C, N, O, F, Ne

Lithium

 $2s^1$



- like H, facile loss of one electron dominates the chemistry
- Li⁺ is found in solids and solution Li reacts with O₂, N₂, H₂O and many more...

Increase in nuclear charge as we add electrons to the same shell. I.E.'s reflect trends in shielding as we discussed at the beginning of the semester.

First Ionization Energies:

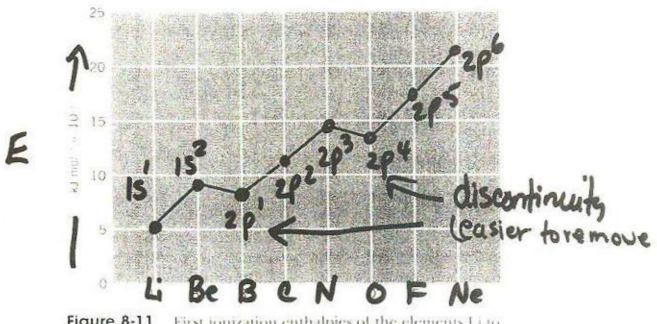


Figure 8-11 First ionization enthalpies of the elements Li to Ne. See also Fig. 2-1 L.

p electrons are better shielded than s electrons

Be has a high I.E. $2s^2$ The first and second I.E. are sufficiently high such that loss of both electrons to give Be²⁺ does not occur with even the most electronegative elements!

In reality these bonds are quite covalent

The elements after Be have even higher I.E.'s so there are no simple cations of B, C, N, O, F or Ne

 Carbon

Carbon $2s^22p^2$ 2s 2s 2p exists)

$$\{H-C\equiv C-H \xrightarrow{-2H^+} : C\equiv C:^{2-}\}$$

- True "non-metal" Found with single, double, triple bonds to itself and to N and O mainly
- Ability to bind to itself (catenate) is unparalleled among the elements responsible for polymer chain formation!

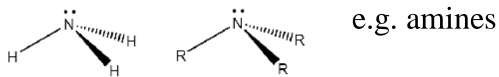
Nitrogen

 $2s^2 2p^3 \qquad \qquad 2s \qquad 2p$

- Forms anions like N³- that are stable with electropositive metals, Mⁿ⁺.
- N₂ is relatively unreactive Why?
 - (a) N≡N bond strength
 - (b) Electronic structure

(The lone pairs on the ends are not accessible – M.O. theory)

- compounds of N are covalent and usually involve three single bonds to other elements



but can involve triple bonds: C≡N, N≡N, etc.

Oxygen $2s^22p^4$ 2s 2p 2p

- two unpaired electrons in O atom according to Aufbau
- *two unpaired electrons on O_2 molecule M.O. theory predicted that the HOMO is a double degenerate π orbital: $\sigma_1^2 \sigma_2^2 \sigma_3^2 \pi_1^4 \pi_2^2$

(12 electrons in the O_2 valence shell; the last two are parallel due to Hund's Rule)

(p. 109 Chapter 3 on M.O. theory)

- O₂ is very reactive with these two unpaired electrons in the HOMO.
- Anions O^{2-} , O_2^- , O_2^{2-} exist in solid
- Covalent bonds exist in molecules CO, SO₃, etc.,
- OH exists in solids and solution

Fluorine 2s²2p⁶

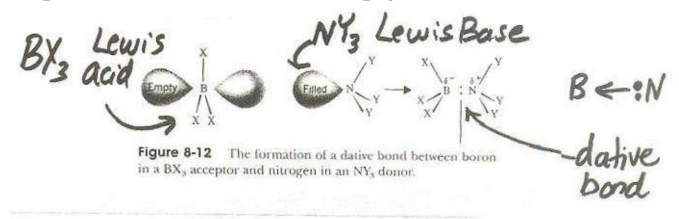
- one unpaired electron on the F atom
- extremely reactive due to low bond energy in F₂
 - → Why is the F-F bond not so stable?
- Ionic compounds containing F and covalent compounds with X F are well-known (the bonds, however, are quite polar)

Covalent Bonds

Three main points:

- 1. Be, B, C have fewer unpaired electrons than the number of bonds they typically form. e.g. C $2s^22p^2$ has only two unpaired electrons (recall promotion to hybrid states with 4 u.e.'s is proposed)
- 2. The elements of the first period Li \rightarrow F obey the octet rule $(2s^22p^6)$ is most stable electronic configuration after all) elements attempt to achieve an octet of electrons around them (octet rule doesn't apply strictly to elements after F)

3. In electron deficient systems such as Boron with only 3 valence electrons (BCl₃, *etc.*,) (too few electrons) there is a tendency for these molecules to accept electrons into their empty orbitals:



Note: B:N would mean complete sharing B^{δ} : $N^{\delta+}$ means a significant polarity exists from the original free states.

Second Short Period

Na, Mg, Al, Si, P, S, Cl, Ar $3s^1 \rightarrow 3s^2 4p^6$

Valence electron structures are similar to Li, Be, B, C, N, O, F, Ne, but their chemistry differs considerably (especially Si, P, S, Cl)

These elements are more like the rest of the heavier elements in their respective groups (columns)

Why? Five Main Reasons

1. First of all, it is not favorable to form $p\pi$ - $p\pi$ multiple bonds such as Si=Si, Si=O, P=P etc.,

This can be rationalized by considering that, in order to achieve good π - π overlap; atoms have to get close together. More repulsion is encountered due to filled 2s, 2p. The only way to stabilize such bonds as Si=Si, Si=P, Ge=Ge, Ge=C, P=P is to put bulky substituents on the atoms to make them have fewer bonds to other groups which makes them engage in a second bond to each other.

This doubly – bond phosphorus compound is kinetically stable.

- 2. When multiple bonding (σ/π combined) is exhibited for this period, the orbitals involved are the low-lying
- d orbitals. $p\pi d\pi$ bonding occurs as in O=PR₃ (We don't have to expect the octet rule to be strictly obeyed for elements like P)
- 3. The possibility of using 3d orbitals leads to more bonds than four in many cases PCl_5 , SF_6 , SiF_6^{2-} etc.,
- 4. The shapes of molecules and the nature of the bonds differ in compounds of Li→Ne versus Na→Ar. See VSEPR again in Chap. 3.
- 5. The compounds of cations and anions differ in these two periods

 $[Be(H_2O)_4]^{2+}$ vs. $[Mg(H_2O)_6]^{2+}$ Al³⁺ is very different from B³⁺, very electropositive – not very covalent. (Cl-Cl bond is actually stronger than F-F, so free Cl⁻ is not as prevalent. Cl₂ is much less reactive.

Rest of the Non-Transition Elements

As already stated, the first row of elements at the top of the periodic table does not serve as a reliable guide for the chemistry of the elements in those groups. The second short row Na—Ar is much more representative.

Two main points need to be clarified for chemistry of non-transition elements.

- 1. What are the differences in the row Li, Be, B, C, N, O, F, Ne and the remaining elements in the elements below each one (in the same group).
- 2. What are the regular variations after this? For example how do things change in the row Li→Ne in each group?

The regular variations that occur as you descend a group:

- (1) Metallic character of the elements
- (2) Properties of the oxide compounds
 - (a) Ionic vs. covalent Which ties in with:
 - (b) Acidic vs. basic

- (3) Properties of the halide compounds
 - (a) Molecular vs. ionic
 - (b) Ease of hydrolysis of the compound
- (4) Trends in covalency
- (5) Trends in structure
 Molecular vs. extended structures etc.,
- (6) Properties of the hydride compounds
- (7) Tendency for catenation (elements binding to another one itself)
- (8) Importance of pπ-pπ versus pπ-dπ bonding (or even $d\pi$ - $d\pi$)
- (9) The general strength of the covalent bonds to particular elements
- (10) The relative importance of low-valent versus highvalent oxidation states of the elements

The rest of the Chapter 8 is an introduction to what is covered in Chapters 9 - 22

Sec. 8-12 The Transition Elements of d and f blocks

Strict definition is:

"elements that have partly filled d or f sub-shells" Broader definition is:

Elements that form compounds with partly filled d or f sub-shells (includes Cu, Ag, Au which are d¹⁰)

Note:

Most of the community refer only to the "d-block" elements as "transition" elements. It is not really common to call the f block elements, transition elements.

f-block elements

- More commonly called "rare earth" elements
- two rows:

Lanthanides

Actinides

Lanthanides

$$La (Z=57) \to Lu (Z=71)$$

Comprise a row of 15 elements

→ lanthanides

Although La is: [Xe] $5d^16s^2$

And Lu is: $[Xe] 4f^{14}5d^{1}6s^{2}$,

They are not considered to be d-block elements because of their chemical and physical properties

Actinides

Ac
$$(Z=89) \rightarrow Lw (Z=103)$$
 (actinium) (Lawrencium)

When you get to Sc (Z=21) and Y (Z=39) the d orbitals become lower than s orbitals up on filling. As you continue to add electrons, the 4f level drops below both the d & s levels.

f orbitals < d < s in lanthanides & actinides

 Table 26-1
 Some Properties of Scandium, Yttrium, and the Lanthanides

Z	Name	Symbol	Electron Configuration	Valences	M³+ Radius (Å)	M ⁸⁺ Color
21	Scandium	Sc	[Ar]3d14s2	3	0.68	Colorless
39	Yttrium	Y	[Kr]4d\5s2	3	0.88	Colorless
57	Lanthanum	La	[Xe]5d16s2	3	1.06	Colorless
58	Cerium	Ce	$[Xe]4f^{1}5d^{1}6s^{2}$	3, 4	1.03	Colorless
59	Praseodymium	Pr	$[Xe]4\int^{3}6s^{2}$	3, 4	1.01	Green
60	Neodymium	Nd	[Xe]4f46s2	3	0.99	Lilac
61	Promethium	Pm	[Xe]4f56s2	3	0.98	Pink
62	Samarium	Sm	[Xe]4f66s2	2, 3	0.96	Yellow
63	Europium	Eu	[Xe]4f76s2	2, 3	0.95	Pale pink
64	Gadolinium	Gd	$[Xe]4f^{7}5d6s^{2}$	3	0.94	Colorless
65	Terbium	Tb	[Xe]4f96s2	3, 4	0.92	Pale pink
66	Dysprosium	Dy	$[Xe]4f^{10}6s^2$	3	0.91	Yellow
67	Holmium	Ho	[Xe]4f116s2	3	0.89	Yellow
68	Erbium	Er	[Xe]4f126s2	3	0.88	Lilac
69	Thulium	Tm	$[Xe]4f^{13}6s^2$	3	0.87	Green
70	Ytterbium	Yb	$[Xe]4f^{14}6s^2$	2, 3	0.86	Colorless
71	Lutetium	Lu	[Xe]4f145d6s2	3	0.85	Colorless

Actinides

Table 27-1 The Actinide Elements and Some of Their Properties

	Name		Electronic Structure ^a	Radii (Å)	
Z		Symbol	of Atom	M ^s	M ⁴⁺
89	Actinium	Ac	6d7s2	1.11	ent sa villavoran
90	Thorium	Th	$6d^27s^2$		0.90
91	Protactinium	Pa	$5f^26d7s^2$ or $5f^16d^27s^2$		0.96
92	Uranium	U	$5\int^{3}6d7s^{2}$	1.03	0.93
93	Neptunium	Np	$5\int^{5}7s^{2}$	1.01	0.92
94	Plutonium	Pu	$5f^67s^2$	1.00	0.90
95	Americium	Am	$5f^{7}7s^{2}$	0.99	0.89
96	Curium	Cm	$5f^{7}6d7s^{2}$	0.985	0.88
97	Berkelium	Bk	5f86d7s2 or 5f97s2	0.98	
98	Californium	Cf	$5f^{10}7s^2$	0.977	
99	Einsteinium	Es	$5f^{11}7s^2$		
100	Fermium	Fm	$5f^{12}7s^2$		
101	Mendelevium	Md	$5f^{13}7s^2$		
102	Nobelium	No	5f147s2		
103	Lawrencium	Lr	$5f^{14}6d7s^{2}$		
104	Rutherfordium	Rf	- W- E (1)		

[&]quot;Outside Rn structure.

The f orbitals have shapes that do not allow the electrons to shield each other very well from the effects of the increased nuclear charge. A 4f¹⁻¹⁴ filling involves a level n=4 that is fairly close to the nucleus.

As a result, there is a big shrinkage in the radii of the atoms and ions of the lanthanides La→Lu. The accumulation of successive shrinkage is called the lanthanide contraction.

Ramifications of Lanthanide Contraction

→Second and third row d-block transition elements have nearly the same size!

Zr Nb Mo Tc Ru Rh Pd Hf Ta W Re Os Ir Pt

The atoms in each pair are not as different in size as one would expect.

And size of atoms/ions helps determine chemical & physical properties!

Q. Why?

<u>A</u>. Valence electrons are involved in the processes that are what we call "chemistry". Their I.E.'s *etc.*, are somewhat determined by distance from the nucleus.

Properties of Transition-metals (elements)

- (1) All are metals
- (2) Practically all are hard, high melting, high boiling elements that are good conductors
- (3) They form alloys with each other
- (4) Many are highly electropositive such that they dissolve in mineral acids (oxidized)
- (5) They exhibit variable valence
- (6) They form paramagnetic compounds