

Dr. Diab Qadah

Department of Chemistry Birzeit University

CHEM 141

Chapter 8





Chapter 8

Electron Configuration and Chemical Periodicity





Electron Configuration and Chemical Periodicity

8.1 Characteristics of Many-Electron Atoms

8.2 The Quantum-Mechanical Model and the Periodic Table

8.3 Trends in Three Atomic Properties

8.4 Atomic Properties and Chemical Reactivity





Figure 8.1 The effect of electron spin.

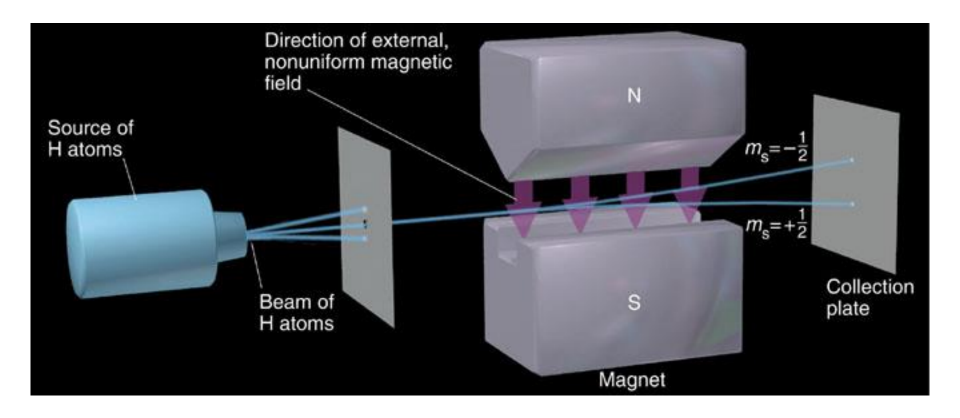






Table 8.1 Summary of Quantum Numbers of Electrons in Atoms

Name	Symbol	Permitted Values	Property
principal	n	positive integers (1, 2, 3,)	orbital energy (size)
angular momentum	l	integers from 0 to <i>n</i> -1	orbital shape (The <i>l</i> values 0, 1, 2, and 3 correspond to s, p, d, and f orbitals, respectively.)
magnetic	m_l	integers from $-l$ to 0 to $+l$	orbital orientation
spin	$m_{_S}$	+½ or -½	direction of e ⁻ spin



Quantum Numbers and The Exclusion Principle

Each electron in any atom is described completely by a set of *four* quantum numbers.

The first three quantum numbers describe the orbital, while the fourth quantum number describes electron spin.

Pauli's **exclusion principle** states that *no two electrons in the same atom can have the same four quantum numbers*.

An atomic orbital can hold a *maximum of two electrons* and they must have *opposing spins*.



Factors Affecting Atomic Orbital Energies

- The energies of atomic orbitals are affected by
 - nuclear charge (Z) and
 - shielding by other electrons.
- A higher nuclear charge increases nucleus-electron interactions and lowers sublevel energy.
- Shielding by other electrons reduces the full nuclear charge to an effective nuclear charge (Z_{eff}).
 - $Z_{\rm eff}$ is the nuclear charge an electron actually experiences.
- Orbital shape also affects sublevel energy.





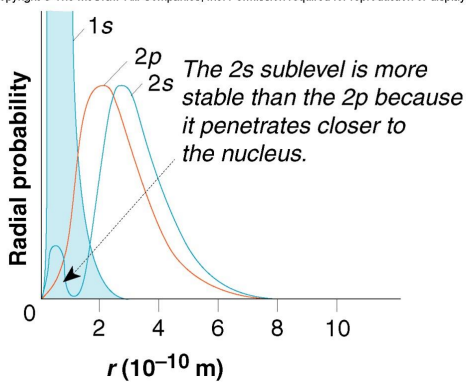
Shielding and Orbital Energy

- Electrons in the same energy level shield each other to some extent.
- Electrons in *inner* energy levels shield the outer electrons very effectively.
 - The farther from the nucleus an electron is, the lower the $Z_{\rm eff}$ for that particular electron.



Figure 8.2 Penetration and sublevel energy.

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Orbital shape causes electrons in some orbitals to "penetrate" close to the nucleus.

Penetration increases nuclear attraction and decreases shielding.



Splitting of Levels into Sublevels

Each energy level is split into *sublevels* of differing energy. Splitting is caused by penetration and its effect on shielding.

For a given *n* value, a lower *l* value indicates a lower energy sublevel.

Order of sublevel energies: s



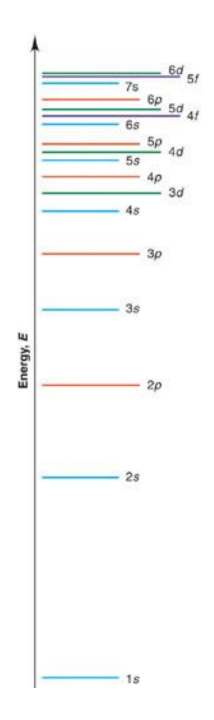


Figure 8.3

Order for filling energy sublevels with electrons.

In general, energies of sublevels increase as n increases (1 < 2 < 3, etc.) and as l increases (s).

As *n* increases, some sublevels overlap.

Electron Configurations and Orbital Diagrams

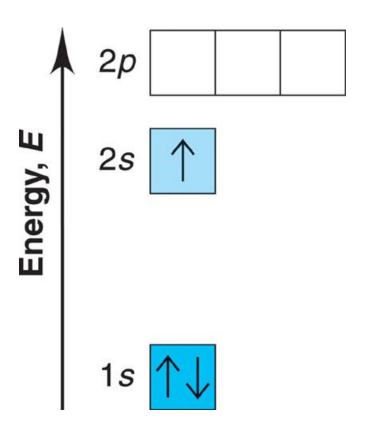
Electron configuration is indicated by a shorthand notation:

$$nl \stackrel{\#}{\longleftarrow}$$
 # of electrons in the sublevel as s, p, d, f

Orbital diagrams make use of a box, circle, or line for each orbital in the energy level. An arrow is used to represent an electron *and* its spin.



A vertical orbital diagram for the Li ground state.



Building Orbital Diagrams

The **aufbau principle** is applied – electrons are always placed in the lowest energy sublevel available.

$$H(Z=1) 1s^1$$
 1s

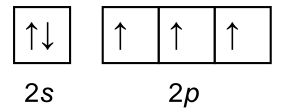
The **exclusion principle** states that each orbital may contain a maximum of 2 electrons, which must have opposite spins.

He
$$(Z=2)$$
 1s² 1s

Building Orbital Diagrams

Hund's rule specifies that when orbitals of equal energy are available, the lowest energy electron configuration has the maximum number of unpaired electrons with parallel spins.

$$N(Z=7) 1s^2 2s^2 2p^3$$





Determining Quantum Numbers from Orbital Diagrams

PROBLEM: Write a set of quantum numbers for the third electron and a set for the eighth electron of the F atom.

PLAN: Identify the electron of interest and note its level (n), sublevel, (l), orbital (m_l) and spin (m_s) . Count the electrons in the order in which they are placed in the diagram.

SOLUTION:

F (
$$Z = 9$$
) $1s^2 2s^2 2p^5$

1s $2s$ $2p$

For the 3rd electron: n = 2, l = 0, $m_l = 0$, $m_s = +\frac{1}{2}$

For the 8th electron: n = 2, l = 1, $m_l = -1$, $m_s = -\frac{1}{2}$





Partial Orbital Diagrams and Condensed Configurations

A *partial orbital diagram* shows only the highest energy sublevels being filled.

A *condensed electron configuration* has the element symbol of the *previous* noble gas in square brackets.

Al has the condensed configuration [Ne] $3s^23p^1$



Table 8.2 Partial Orbital Diagrams and Electron Configurations* for the Elements in Period 3.

Atomic Number	Element		al Orbital Diagram nd 3 <i>p</i> Sublevels Only)	Full Electron Configuration [†]	Condensed Electron Configuration
11	Na	3s	3 <i>p</i>	$[1s^22s^22p^6]$ $3s^1$	[Ne] 3s ¹
12	Mg	$\uparrow\downarrow$		$[1s^22s^22p^6]$ 3s ²	[Ne] $3s^2$
13	Al	$\uparrow\downarrow$	\uparrow	$[1s^22s^22p^6] \ 3s^23p^1$	[Ne] $3s^2 3p^1$
14	Si	$\uparrow\downarrow$	\uparrow \uparrow	$[1s^22s^22p^6] 3s^23p^2$	[Ne] $3s^2 3p^2$
15	P	$\uparrow\downarrow$		$[1s^22s^22p^6] 3s^23p^3$	[Ne] $3s^2 3p^3$
16	S	$\uparrow\downarrow$	$\uparrow\downarrow$ \uparrow \uparrow	$[1s^22s^22p^6] 3s^23p^4$	[Ne] $3s^2 3p^4$
17	Cl	$\uparrow\downarrow$	$\uparrow\downarrow\uparrow\uparrow\downarrow$	$[1s^22s^22p^6] 3s^23p^5$	[Ne] $3s^2 3p^5$
18	Ar	$\uparrow\downarrow$		$[1s^22s^22p^6] 3s^23p^6$	[Ne] $3s^2 3p^6$

^{*}Colored type indicates the sublevel to which the last electron is added.

[†]The full configuration is not usually written with square brackets; they are included here to show how the [Ne] designation arises.

Electron Configuration and Group

Elements in the same group of the periodic table have the same outer electron configuration.

Elements in the same group of the periodic table exhibit similar chemical behavior.

Similar outer electron configurations correlate with similar chemical behavior.



Figure 8.4 Condensed electron configurations in the first three periods.

8A 1A (18)(1) 2 H He 2A **3A** 4A 5A 6A **7A** 1*s*¹ $1s^{2}$ (2)(13)(14)(15)(16)(17)10 3 8 Period Li N 0 Ne Be [He] $2s^22p^1$ [He] $2s^22p^2$ [He] $2s^22p^3$ [He] $2s^22p^4$ [He] $2s^22p^5$ [He] $2s^22p^6$ [He] 2s1 [He] 2s² 11 12 13 14 15 16 17 18

Si

AI

S

[Ne] $3s^23p^1$ [Ne] $3s^23p^2$ [Ne] $3s^23p^3$ [Ne] $3s^23p^4$ [Ne] $3s^23p^5$ [Ne] $3s^23p^6$

CI

Ar

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Na

[Ne] 3s1

Mg

[Ne] $3s^2$

Table 8.3 Partial Orbital Diagrams and Electron Configurations* for the Elements in Period 4.

Atomic Number	Element	Partial Orbital Diagram (4 <i>s</i> , 3 <i>d</i> , and 4 <i>p</i> Sublevels Only)	Full Electron Configuration	Condensed Electron Configuration			
19	K	4s 3d 4p ↑	$1s^2 2s^2 2p^6 3s^2 3p^6 4s^1$	$[Ar] 4s^1$			
20 21	Ca Sc	$ \begin{array}{c cccc} $	$1s^{2}2s^{2}2p^{6}3s^{2}3p^{6}4s^{2}$ $1s^{2}2s^{2}2p^{6}3s^{2}3p^{6}4s^{2}3d^{1}$	[Ar] $4s^2$			
22	Ti	$\uparrow \downarrow \qquad \uparrow \qquad \uparrow \qquad $	$1s^2 2s^2 2p^6 3s^2 3p^6 4s^2 3d^2$	$[Ar] 4s^2 3d^2$			
23	V	$\uparrow \downarrow \qquad \qquad \uparrow \qquad \uparrow \qquad \uparrow \qquad \qquad \boxed{}$	$1s^2 2s^2 2p^6 3s^2 3p^6 4s^2 3d^3$	$[Ar] 4s^2 3d^3$			
24	Cr	$\begin{array}{ c c c c c c c c c c c c c c c c c c c$	$1s^22s^22p^63s^23p^64s^13d^5$	$[Ar] 4s^1 3d^5$			
25	Mn	$\begin{array}{ c c c c c c c c c c c c c c c c c c c$	$1s^22s^22p^63s^23p^64s^23d^5$	$[Ar] 4s^2 3d^5$			
26	Fe	$\uparrow \downarrow \qquad \uparrow \downarrow \uparrow $	$1s^22s^22p^63s^23p^64s^23d^6$	$[Ar] 4s^2 3d^6$			
27	Co	$\uparrow \downarrow \qquad \uparrow \downarrow \uparrow \downarrow \uparrow \qquad \uparrow \qquad \uparrow$	$1s^2 2s^2 2p^6 3s^2 3p^6 4s^2 3d^7$	$[Ar] 4s^2 3d^7$			



^{*}Colored type indicates the sublevel to which the last electron is added.

Table 8.3 Partial Orbital Diagrams and Electron Configurations* for the Elements in Period 4.

Atomic Number	Element		Partial Orbital Diagran 4 <i>s,</i> 3 <i>d,</i> and 4 <i>p</i> Sublev		Full Electron Configuration	Condensed Electron Configuration			
28 29 30 31	Ni Cu Zn Ga	↑↓ ↑↓ ↑↓	$\begin{array}{c ccccccccccccccccccccccccccccccccccc$		$1s^{2}2s^{2}2p^{6}3s^{2}3p^{6}4s^{2}3d^{8}$ $1s^{2}2s^{2}2p^{6}3s^{2}3p^{6}4s^{1}3d^{10}$ $1s^{2}2s^{2}2p^{6}3s^{2}3p^{6}4s^{2}3d^{10}$ $1s^{2}2s^{2}2p^{6}3s^{2}3p^{6}4s^{2}3d^{10}4p^{1}$	[Ar] $4s^23d^8$ [Ar] $4s^13d^{10}$ [Ar] $4s^23d^{10}$ [Ar] $4s^23d^{10}4p^1$			
32	Ge	$\uparrow\downarrow$	$\boxed{\uparrow\downarrow \uparrow\downarrow \uparrow\downarrow \uparrow\downarrow \uparrow\downarrow \uparrow\downarrow}$	\uparrow \uparrow	$1s^2 2s^2 2p^6 3s^2 3p^6 4s^2 3d^{10} 4p^2$	[Ar] $4s^2 3d^{10} 4p^2$			
33	As	$\uparrow\downarrow$	$\uparrow\downarrow\uparrow\uparrow\downarrow\uparrow\uparrow\downarrow\uparrow\uparrow\downarrow\uparrow\uparrow\downarrow$	\uparrow \uparrow \uparrow	$1s^22s^22p^63s^23p^64s^23d^{10}4p^3$	[Ar] $4s^2 3d^{10} 4p^3$			
34	Se	$\uparrow\downarrow$	$\uparrow\downarrow \uparrow\downarrow \uparrow\downarrow \uparrow\downarrow \uparrow\downarrow \uparrow\downarrow$	$\uparrow\downarrow$ \uparrow \uparrow	$1s^22s^22p^63s^23p^64s^23d^{10}4p^4$	[Ar] $4s^2 3d^{10} 4p^4$			
35	Br	$\uparrow\downarrow$	$\uparrow\downarrow \uparrow\downarrow \uparrow\downarrow \uparrow\downarrow \uparrow\downarrow \uparrow\downarrow$	$\uparrow\downarrow\uparrow\downarrow\uparrow$	$1s^2 2s^2 2p^6 3s^2 3p^6 4s^2 3d^{10} 4p^5$	[Ar] $4s^2 3d^{10} 4p^5$			
36	Kr	$\uparrow\downarrow$	$\boxed{\uparrow\downarrow \uparrow\downarrow \uparrow\downarrow \uparrow\downarrow \uparrow\downarrow}$	$\uparrow\downarrow\uparrow\uparrow\downarrow\uparrow\downarrow$	$1s^2 2s^2 2p^6 3s^2 3p^6 4s^2 3d^{10} 4p^6$	[Ar] $4s^2 3d^{10} 4p^6$			

^{*}Colored type indicates the sublevel to which the last electron is added.





Figure 8.5

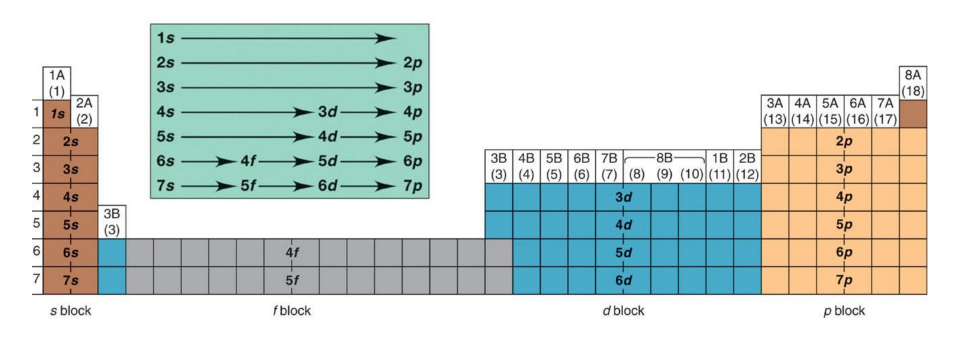
A periodic table of partial ground-state electron configurations.

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	Copyright © The McGraw-Hill Companies, Inc. Permission required for reproduction or display.																		
		Main-Group Elements (s block)								Main-Group Elements (<i>p</i> block)									
	1A (1)											8A (18)							
		ns ¹																	ns ² np ⁶
	1	1 H	2A (2)													5A (15)	6A (16)	7A (17)	2 He
		1s ¹	ns ²											ns ² np ¹	ns ² np ²	ns²np³	ns ² np ⁴	ns²np5	1s ²
<u></u>		3	4											5	6	7	8	9	10
Period number: highest occupied energy level	2	Li 2s ¹	Be 2s ²		Transition Elements (<i>d</i> block)										C 2s ² 2p ²	N 2s ² 2p ³	O 2s ² 2p ⁴	F 2s ² 2p ⁵	Ne 2s ² 2p ⁶
ene		11	12					(0.0)	ook)					13	14	15	16	17	18
peidr	3	Na 3s1	Mg 3s ²	3B (3)	4B (4)	5B (5)	6B (6)	7B (7)	(8)	— 8B — (9)	(10)	1B (11)	2B (12)	AI 3s ² 3p ¹	Si $3s^23p^2$	P 3s ² 3p ³	S 3s ² 3p ⁴	CI 3s ² 3p ⁵	Ar $3s^23p^6$
CC		19	20	21	22	23	24	25	26	27	28	29	30	31	32	33	34	35	36
ghest	4	K 4s ¹	Ca 4s ²	Sc 4s ² 3d ¹	Ti 4s ² 3d ²	V 4s ² 3d ³	Cr $4s^13d^5$	Mn 4s ² 3d ⁵	Fe $4s^23d^6$	Co $4s^23d^7$	Ni 4s ² 3d ⁸	Cu 4s ¹ 3d ¹⁰	Zn 4s ² 3d 10	Ga $4s^24p^1$	Ge $4s^24p^2$	As $4s^24p^3$	Se $4s^24p^4$	Br 4s ² 4p ⁵	Kr $4s^24p^6$
: hi		37	38	39	40	41	42	43	44	45	46	47	48	49	50	51	52	53	54
lbe	5	Rb	Sr	Υ	Zr	Nb	Мо	Тс	Ru	Rh	Pd	Ag	Cd	In	Sn	Sb	Те	- 1	Xe
nn		5s ¹	5s ²	5s ² 4d ¹	5s ² 4d ²	5s ¹ 4d ⁴	5s ¹ 4d ⁵	5s ² 4d ⁵	5s ¹ 4d ⁷	5s ¹ 4d ⁸	4d ¹⁰	2000 0000	5s ² 4d ¹⁰		5s ² 5p ²	$5s^25p^3$	5s ² 5p ⁴	5s ² 5p ⁵	5s ² 5p ⁶
jod		55	56	57	72	73	74	75	76	77	78	79	80	81	82	83	84	85	86
Per	6	Cs 6s1	Ba 6 <i>s</i> ²	La* 6 <i>s</i> 25 <i>d</i> 1	Hf 6s ² 5d ²	Ta 6s ² 5d ³	W 6s ² 5d ⁴	Re 6s ² 5d ⁵	Os 6s ² 5d ⁶	1r 6s ² 5d ⁷	Pt 6s ¹ 5d ⁹	Au 6s ¹ 5d ¹⁰	Hg 6s ² 5d ¹⁰	TI 6s ² 6p ¹	Pb 6s ² 6p ²	Bi 6s ² 6p ³	Po 6s ² 6p ⁴	At 6s ² 6p ⁵	Rn 6 <i>s</i> ² 6 <i>p</i> ⁶
	-	87	88	89	104	105	106	107	108	109	110	111	112	113	114	115	116	65-6p-	118
	7	Fr	Ra	Ac**	Rf	Db	Sg	Bh	Hs	Mt	Ds	Rg	Cn	113	114	113	110		110
	'	7s ¹	7s ²	7s ² 6d ¹	7s ² 6d ²			7s ² 6d ⁵	7s ² 6d ⁶					7s ² 7p ¹	7s ² 7p ²	7s ² 7p ³	7s ² 7p ⁴		7s ² 7p ⁶
	Inner Transition Elements (f block)											1	92.						
					59	60	61	62	63	64	65	66	67	68	69	70	71		
	6	*Lanth	anides	Се	Pr	Nd	Pm	Sm	Eu	Gd	Tb	Dy	Ho	Er	Tm	Yb	Lu		
	_			6s ² 4f ¹ 5d ¹	6s ² 4f ³	6s ² 4f ⁴	6s ² 4f ⁵	6s ² 4f ⁶	6s ² 4f ⁷	6s ² 4f ⁷ 5d ¹	6s ² 4f ⁹					6s ² 4f ¹⁴			
	7	** ^ ^	tinides	90 Th	91 Pa	92 U	93 No	94 Pu	95	96 Cm	97 Bk	98 Cf	99 Es	100 Fm	101 Md	102 No	103 Lr		
	1	AC	uillues	in	Pa	U	Np	Pu	Am	CIII	BK O	0 10	ES 11	rm			LI		

 $\left| 7s^26d^2 \right| \left| 7s^25f^26d^4 \right| \left| 7s^25f^36d^4 \right| \left| 7s^25f^46d^4 \right| \left| 7s^25f^46d^4 \right| \left| 7s^25f^6 \right| \left| 7s^25f^7 \right| \left| 7s^25f^76d^4 \right| \left| 7s^25f^9 \right| \left| 7s^25f^{10} \right| \left| 7s^25f^{11} \right| \left| 7s^25f^{12} \right| \left| 7s^25f^{13} \right| \left| 7s^25f^{14} \right| | \left| 7s^25f^{14} \right| \left| 7s^25f^{14} \right| | | \left| 7s^25f^{14} \right| | | \left| 7s^25f^{14} \right| | | | \left| 7s^25f^{14} \right| | | \left$

Figure 8.6 Orbital filling and the periodic table.



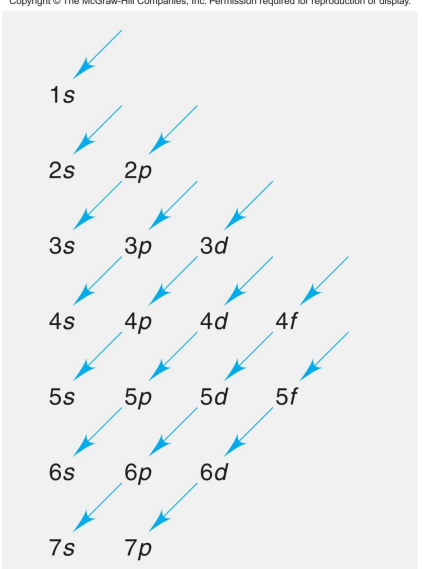
The order in which the orbitals are filled can be obtained directly from the periodic table.





Aid to memorizing sublevel filling order.





The n value is constant horizontally. The l value is constant vertically. n + l is constant diagonally.



Categories of Electrons

Inner (core) electrons are those an atom has in common with the pervious noble gas and any *completed* transition series.

Outer electrons are those in the *highest* energy level (highest *n* value).

Valence electrons are those involved in forming compounds.

For **main group** elements, the valence electrons **are** the outer electrons.

For **transition elements**, the valence electrons include the outer electrons and any (n-1)d electrons.





Determining Electron Configurations

PROBLEM: Using the periodic table on the inside cover of the text (not Figure 8.5 or Table 8.3), give the full and condensed electron configurations, partial orbital diagrams showing valence electrons only, and number of inner electrons for the following elements:

$$(K; Z = 19)$$

$$(K; Z = 19)$$
 $(Tc; Z = 43)$

(Pb;
$$Z = 82$$
)

PLAN: The atomic number gives the number of electrons, and the periodic table shows the order for filling orbitals. The partial orbital diagram includes all electrons added after the previous noble gas except those in filled inner sublevels.



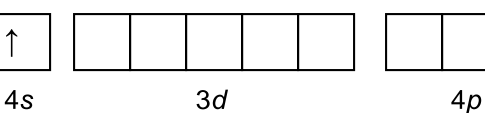
SOLUTION:

(a) For K (Z = 19)

full configuration $1s^22s^22p^63s^23p^64s^1$

condensed configuration [Ar] 4s¹

partial orbital diagram



There are 18 inner electrons.

SOLUTION:

(b) For Tc (Z = 43)

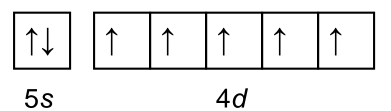
full configuration

 $1s^22s^22p^63s^23p^64s^23d^{10}4p^65s^24d^5$

condensed configuration

[Kr]5*s*²4*d*⁵

partial orbital diagram



5*p*

There are 36 inner electrons.



SOLUTION:

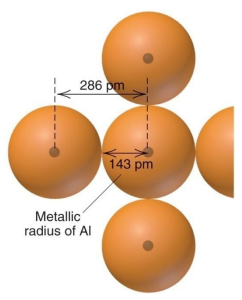
(a) For Pb (Z = 82)

full configuration $1s^22s^22p^63s^23p^64s^23d^{10}4p^65s^24d^{10}5p^66s^24f^{14}5d^{10}6p^2$ condensed configuration [Xe] $6s^24f^{14}5d^{10}6p^2$

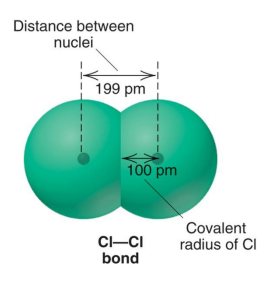
There are 78 inner electrons.



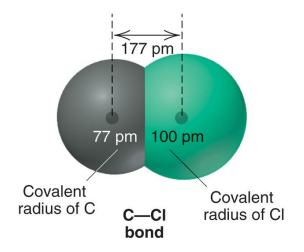
Figure 8.7 Defining atomic size.



A. The metallic radius of aluminum.



B. The covalent radius of chlorine.



C. Known covalent radii and distances between nuclei can be used to find unknown radii.



Trends in Atomic Size

Atomic size *increases* as the principal quantum number *n increases*.

As *n* increases, the probability that the outer electrons will be farther from the nucleus increases.

Atomic size **decreases** as the effective nuclear charge Z_{eff} increases.

As Z_{eff} increases, the outer electrons are pulled closer to the nucleus.

For *main group elements*:

atomic size *increases* down a group in the periodic table and *decreases* across a period.





Figure 8.8

Atomic radii of the maingroup and transition elements.

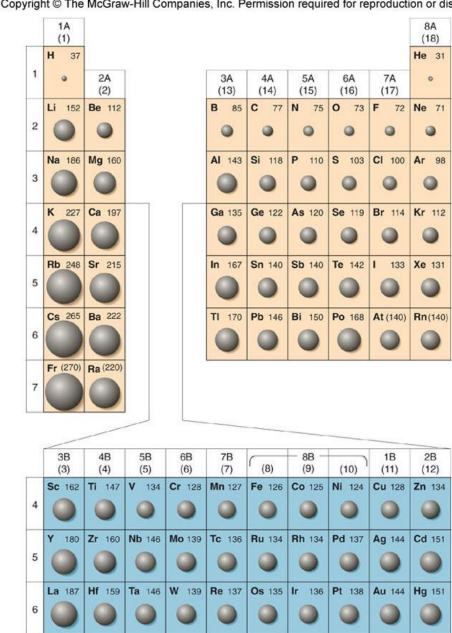
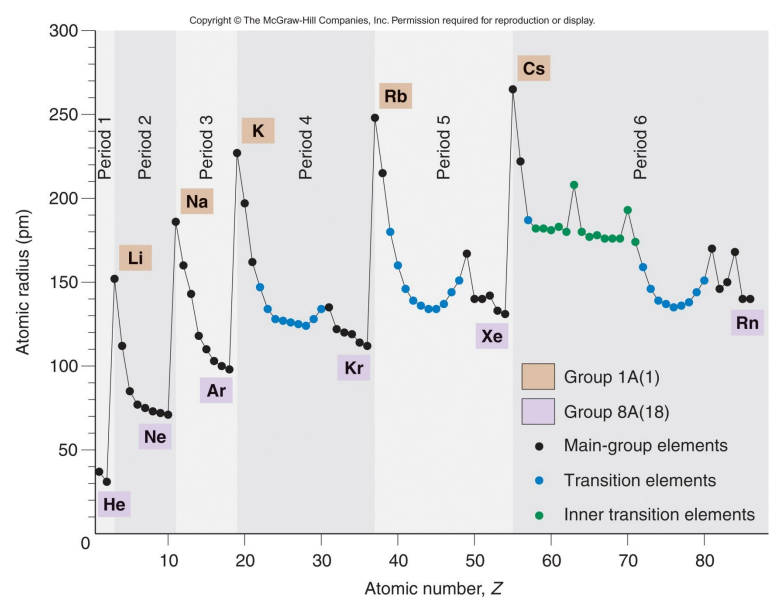


Figure 8.9 Periodicity of atomic radius.



Ranking Elements by Atomic Size

PROBLEM: Using only the periodic table (not Figure 8.8), rank each set of main-group elements in order of *decreasing* atomic size:

- **(a)** Ca, Mg, Sr **(b)** K, Ga, Ca
- (c) Br, Rb, Kr (d) Sr, Ca, Rb

PLAN: Locate each element on the periodic table. Main-group elements increase in size down a group and decrease in size across the period.



SOLUTION:

(a) Sr > Ca > Mg

Ca, Mg, and Sr are in Group 2A. Size increases down the group.

(b) K > Ca > Ga

K, Ga, and Ca are all in Period 4. Size decreases across the period.

(c) Rb > Br > Kr

Rb is the largest because it has one more energy level than the other elements. Kr is smaller than Br because Kr is further to the right in the same period.

(d) Rb > Sr > Ca

Ca is the smallest because it has one fewer energy level. Sr is smaller than Rb because it is further to the right in the same period.



Trends in Ionization Energy

Ionization energy (IE) is the energy required for the **complete removal** of 1 mol of electrons from 1 mol of gaseous atoms or ions.

Atoms with a *low IE* tend to form *cations*. Atoms with a *high IE* tend to form *anions* (except the noble gases).

Ionization energy tends to *decrease* down a group and *increase* across a period.

Figure 8.10 Periodicity of first ionization energy (IE₁).

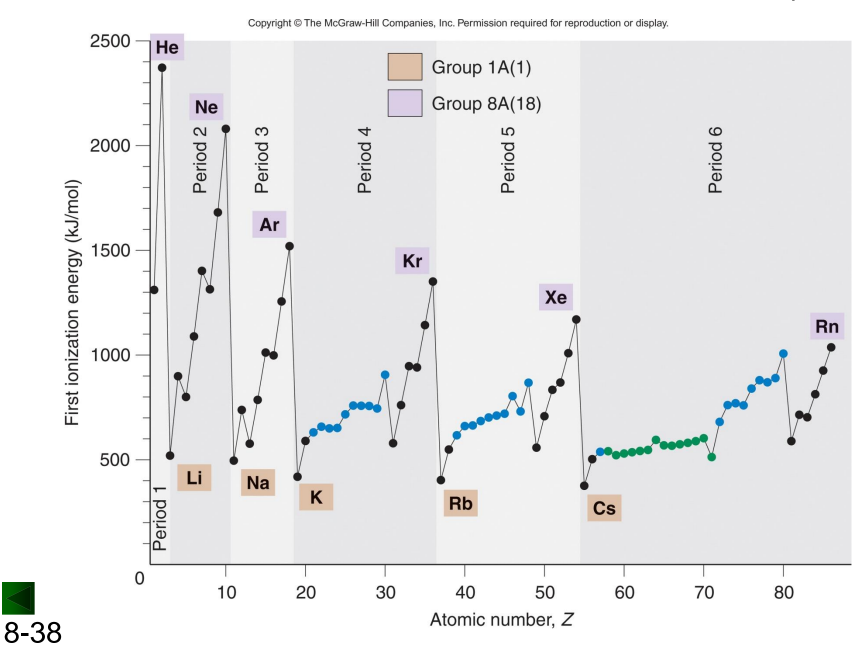
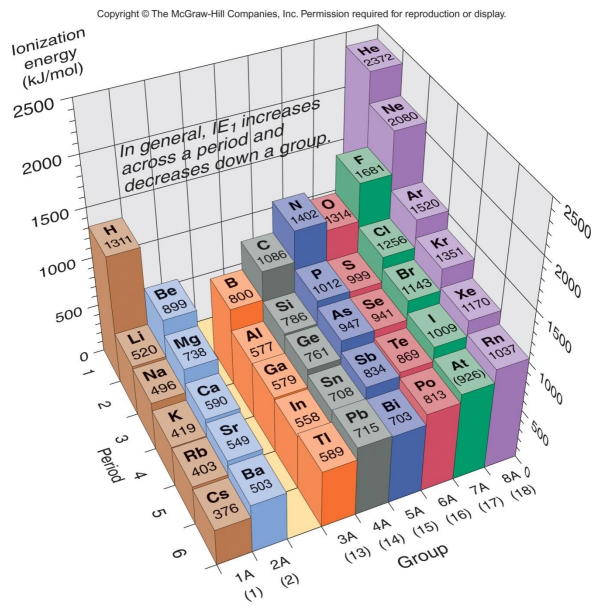


Figure 8.11 First ionization energies of the main-group elements.



Ranking Elements by First Ionization Energy

PROBLEM: Using the periodic table only, rank the elements in each of the following sets in order of *decreasing* IE₁:

- (a) Kr, He, Ar
- **(b)** Sb, Te, Sn
- **(c)** K, Ca, Rb
- (d) I, Xe, Cs

PLAN: Find each element on the periodic table. IE₁ generally decreases down a group and increases across a period.

SOLUTION:

(a) He > Ar > Kr

Kr, He, and Ar are in Group 8A. IE₁ decreases down the group.



SOLUTION:

(b) Te > Sb > Sn

Sb, Te, and Sn are in Period 5. IE₁ increases across a period.

(c) Ca > K > Rb

K has a higher IE₁ than Rb because K is higher up in Group 1A. Ca has a higher IE₁ than K because Ca is further to the right in Period 4.

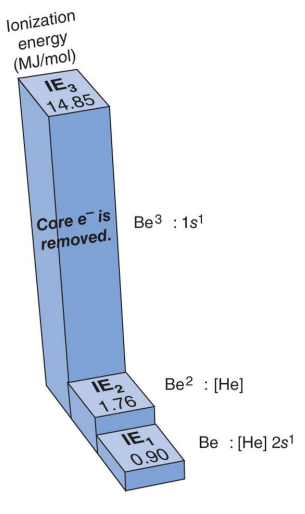
(d) Xe > I > Cs

Xe has a higher IE₁ than I because Xe is further to the right in the same period. Cs has a lower IE₁ than I because it is further to the left in a higher period.



Figure 8.12 The first three ionization energies of beryllium.

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Beryllium has 2 valence electrons, so IE₃ is much larger than IE₂.



Table 8.4 Successive Ionization Energies of the Elements Lithium Through Sodium

		Number		Ionization Energy (MJ/mol)*								
Z	Element	of Valence Electrons	IE ₁	IE ₂	IE ₃	IE ₄	IE ₅	IE ₆	IE ₇	IE ₈	IE ₉	IE ₁₀
3	Li	1	0.52	7.30	11.81							
4	Be	2	0.90	1.76	14.85	21.01			CORE	ELECTR	ONS	
5	В	3	0.80	2.43	3.66	25.02	32.82					
6	C	4	1.09	2.35	4.62	6.22	37.83	47.28				
7	N	5	1.40	2.86	4.58	7.48	9.44	53.27	64.36			
8	O	6	1.31	3.39	5.30	7.47	10.98	13.33	71.33	84.08		
9	F	7	1.68	3.37	6.05	8.41	11.02	15.16	17.87	92.04	106.43	
10	Ne	8	2.08	3.95	6.12	9.37	12.18	15.24	20.00	23.07	115.38	131.43
11	Na	1	0.50	4.56	6.91	9.54	13.35	16.61	20.11	25.49	28.93	141.37

^{*}MJ/mol, or megajoules per mole = 10^3 kJ/mol.

Identifying an Element from Its Ionization Energies

PROBLEM: Name the Period 3 element with the following ionization energies (in kJ/mol) and write its electron configuration:

IE ₁	IE_2	IE ₃	IE_4	IE ₅	IE ₆
1012	1903	2910	4956	6278	22,230

PLAN: Look for a large increase in IE, which occurs after all valence electrons have been removed.

SOLUTION:

The largest increase occurs after IE_5 , that is, after the 5th valence electron has been removed. The Period 3 element with 5 valence electrons is **phosphorus** (**P**; Z = 15).

The complete electron configuration is $1s^22s^22p^63s^23p^3$.



Trends in Electron Affinity

Electron Affinity (EA) is the energy change that occurs when 1 mol of electrons is **added** to 1 mol of gaseous atoms or ions.

Atoms with a *low EA* tend to form *cations*. Atoms with a *high EA* tend to form *anions*.

The trends in electron affinity are not as regular as those for atomic size or IE.

Figure 8.13 Electron affinities of the main-group elements (in kJ/mol).

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Сорупу	Copyright © The McGraw-Hill Companies, Inc. Permission required for reproduction or display.									
1A (1)								8A (18)		
H -72.8	2A (2)		3A (13)	4A (14)	5A (15)	6A (16)	7A (17)	He (0.0)		
Li -59.6	Be ≤0		B -26.7	C – 122	N +7	O -141	F -328	Ne (+29)		
Na – 52.9	Mg ≤0		AI -42.5	Si - 134	P -72.0	S -200	CI -349	Ar (+35)		
K -48.4	Ca -2.37		Ga -28.9	Ge - 119	As -78.2	Se -195	Br -325	Kr (+39)		
Rb - 46.9	Sr -5.03		In –28.9	Sn – 107	Sb -103	Te – 190	I -295	Xe (+41)		
Cs -45.5	Ba -13.95		TI –19.3	Pb -35.1	Bi -91.3	Po – 183	At -270	Rn (+41)		

Behavior Patterns for IE and EA

Reactive nonmetals have high IEs and highly negative EAs.

These elements attract electrons strongly and tend to form negative ions in ionic compounds.

Reactive metals have low IEs and slightly negative EAs. These elements lose electrons easily and tend to form positive ions in ionic compounds.

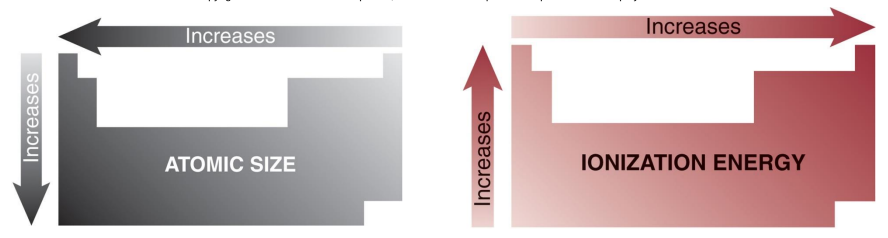
Noble gases have very high IEs and slightly positive EAs. These elements tend to neither lose nor gain electrons.

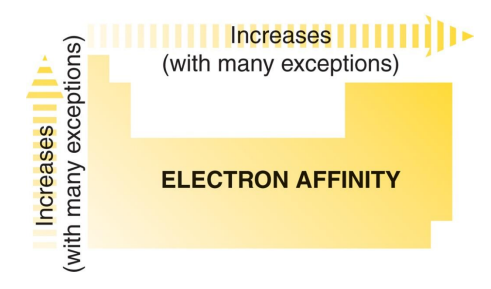




Figure 8.14 Trends in three atomic properties.

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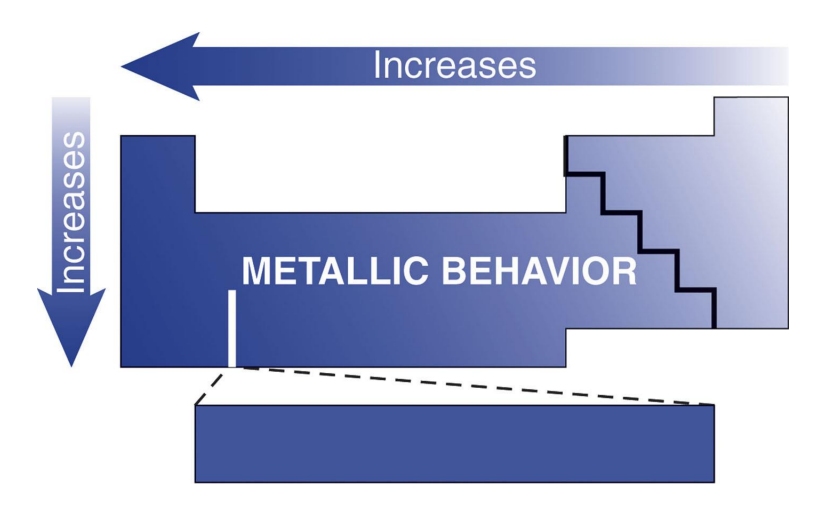


Metallic Behavior

- Metals are typically shiny solids with moderate to high melting points.
- Metals are good conductors of heat and electricity, and can easily be shaped.
- Metals tend to lose electrons and form cations, i.e., they are easily oxidized.
- Metals are generally strong reducing agents.
- Most metals form ionic oxides, which are basic in aqueous solution.



Figure 8.15 Trends in metallic behavior.



Acid-Base Behavior of Oxides

Main-group metals form *ionic oxides*, which are *basic* in aqueous solution.

Main-group nonmetals form *covalent oxides*, which are *acidic* in aqueous solution.

Some metals and metalloids form *amphoteric oxides*, which can act as acids or bases in water:

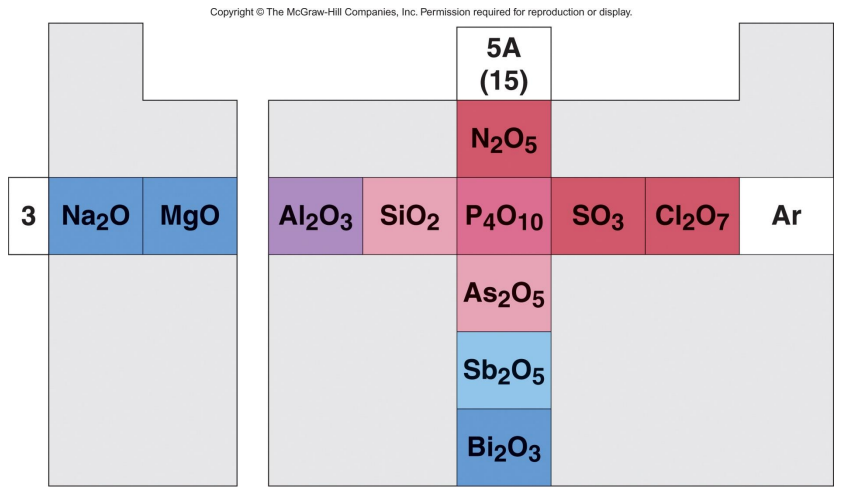
$$Al_2O_3(s) + 6HCI(aq) \rightarrow 2AICl_3(aq) + 3H_2O(l)$$

 $Al_2O_3(s) + 2NaOH(aq) + 3H_2O(l) \rightarrow 2NaAl(OH)_4(aq)$





Figure 8.16 Acid-base behavior of some element oxides.



Oxides become more basic down a group and more acidic across a period.





Electron configurations of Monatomic Ions

Elements at either end of a period gain or lose electrons to attain a filled outer level. The resulting ion will have a *noble gas electron configuration* and is said to be *isoelectronic* with that noble gas.

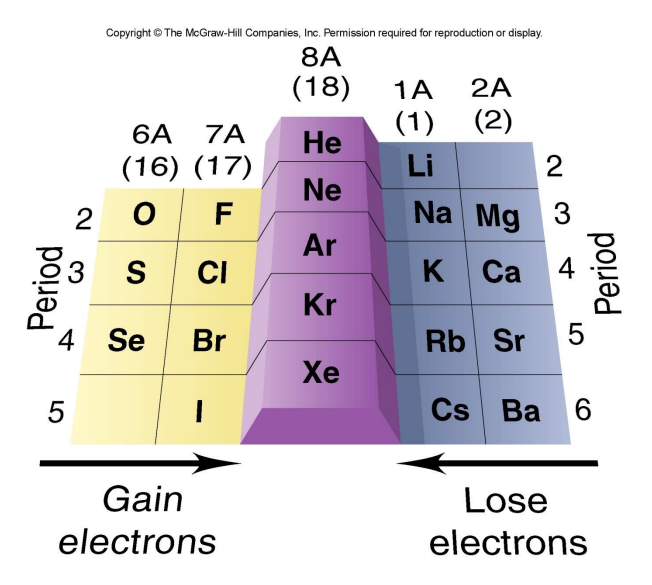
Na(1
$$s^22s^22p^63s^1$$
) \rightarrow e⁻ + Na⁺([He]2 s^22p^6)
[isoelectronic with Ne]

Br([Ar]
$$4s^23d^{10}4p^5$$
) + $e^- \rightarrow Br^-$ ([Ar] $4s^23d^{10}4p^6$) [isoelectronic with Kr]





Figure 8.17 Main-group elements whose ions have noble gas electron configurations.



Electron Configurations of Monatomic Ions

A *pseudo-noble gas configuration* is attained when a metal atom empties its highest energy level.

The ion attains the stability of empty ns and np sublevels and a filled (n-1)d sublevel.

$$Sn([Kr]5s^24d^{10}5p^2) \rightarrow 4e^- + Sn^{4+} ([Kr]4d^{10})$$

A metal may lose only the *np* electrons to attain an *inert pair configuration*.

The ion attains the stability of a filled ns and (n-1)d sublevels.

$$Sn([Kr]5s^24d^{10}5p^2) \rightarrow 2e^- + Sn^{2+} ([Kr]5s^24d^{10})$$





Writing Electron Configurations of Main-Group Ions

PROBLEM: Using condensed electron configurations, write reactions for the formation of the common ions of the following elements:

(a) Iodine (Z = 53) (b) Potassium (Z = 19) (c) Indium (Z = 49)

PLAN: Identify the position of each element on the periodic table and recall that:

- Ions of elements in Groups 1A(1), 2A(2), 6A(16), and 7A(17)
 are usually isoelectronic with the nearest noble gas.
- Metals in Groups 3A(13) to 5A(15) can lose the *ns* and *np* electrons or just the *np* electrons.

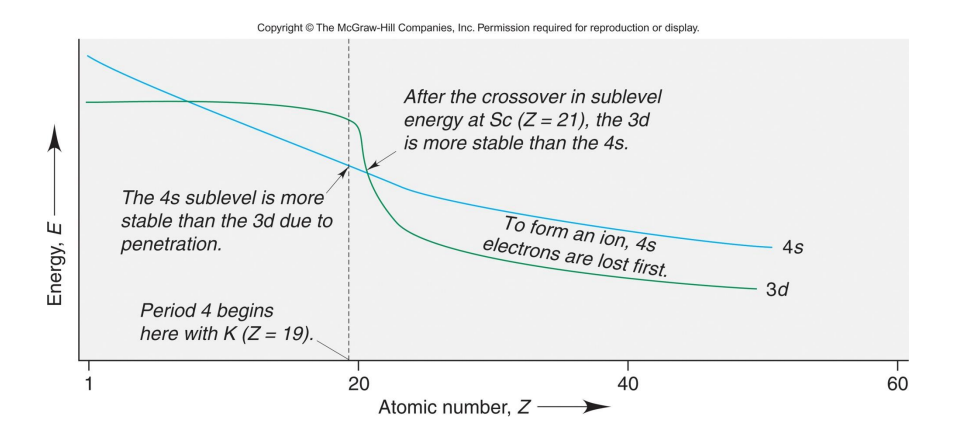


SOLUTION:

- (a) Iodine (Z = 53) is in Group 7A(17) and will gain one electron to be isoelectronic with Xe: I ([Kr] $5s^24d^{10}5p^5$) + e⁻ \rightarrow I⁻ ([Kr] $5s^24d^{10}5p^6$)
- **(b)** Potassium (Z = 19) is in Group 1A(1) and will lose one electron to be isoelectronic with Ar: K ([Ar] $4s^1$) \rightarrow K⁺ ([Ar]) + e⁻
- (c) Indium (Z = 49) is in Group 3A(13) and can lose either one electron or three electrons: In ([Kr] $5s^24d^{10}5p^1$) \rightarrow In⁺ ([Kr] $5s^24d^{10}$) + e⁻ In ([Kr] $5s^24d^{10}5p^1$) \rightarrow In³⁺([Kr] $4d^{10}$) + 3e⁻



Figure 8.18 The crossover of sublevel energies in Period 4.



Magnetic Properties of Transition Metal Ions

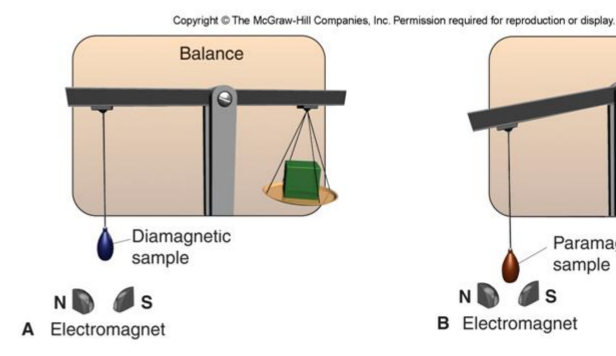
A species with one or more unpaired electrons exhibits *paramagnetism* – it is attracted by a magnetic field.

Ag
$$(Z=47)$$

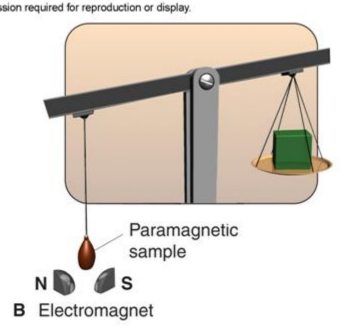
$$\uparrow \qquad \uparrow \downarrow \qquad \uparrow \downarrow \qquad \uparrow \downarrow \qquad \uparrow \downarrow$$
5s
$$4d$$
5p

A species with all its electrons paired exhibits diamagnetism – it is not attracted (and is slightly repelled) by a magnetic field.

Figure 8.19 Measuring the magnetic behavior of a sample.



The apparent mass of a diamagnetic substance is unaffected by the magnetic field.



The apparent mass of a paramagnetic substance increases as it is attracted by the magnetic field.



Magnetic Properties of Transition Metal ions

Magnetic behavior can provide evidence for the electron configuration of a given ion.

Ti²⁺ has 2 unpaired electrons and is paramagnetic, providing evidence that the 4*s* electrons are lost before the 3*d* electrons.

Writing Electron Configurations and **Predicting Magnetic Behavior of Transition Metal Ions**

PROBLEM:

Use condensed electron configurations to write the reaction for the formation of each transition metal ion, and predict whether the ion is paramagnetic or diamagnetic.

(a)
$$Mn^{2+}(Z = 25)$$

(b)
$$Cr^{3+}(Z = 24)$$
 (c) $Hg^{2+}(Z = 80)$

(c)
$$Hg^{2+}(Z = 80)$$

PLAN:

Write the condensed electron configuration for each atom, recalling the irregularity for Cr. Remove electrons, beginning with the *ns* electrons, and determine if there are any unpaired electrons.



SOLUTION:

- (a) $Mn^{2+}(Z = 25)$ $Mn ([Ar] 4s^23d^5) \rightarrow Mn^{2+}([Ar] 3d^5) + 2e^-$ Since there are 5 d electrons they are all unpaired. Mn^{2+} is **paramagnetic**.
- **(b)** $Cr^{3+}(Z = 24)$ $Cr([Ar] 4s^{1}3d^{5}) \rightarrow Cr^{3+}([Ar] 3d^{3}) + 3e^{-}$ Since there are 3 d electrons they are all unpaired. Cr^{3+} is **paramagnetic.**
- (c) $Hg^{2+}(Z = 80)$ Hg ([Xe] $6s^24f^{14}5d^{10}) \rightarrow Hg^{2+}$ ([Xe] $4f^{14}5d^{10}) + 2e^-$ The 4f and the 5s sublevels are filled, so there are no unpaired electrons. Hg^{2+} is **diamagnetic**.



Ionic Size vs. Atomic Size

Cations are *smaller* than their parent atoms while anions are *larger*.

Ionic radius *increases* down a group as *n* increases.

Cation size *decreases* as charge *increases*.

An *isoelectronic series* is a series of ions that have the same electron configuration. Within the series, ion size *decreases* with increasing nuclear charge.



Figure 8.20 Ionic radius.

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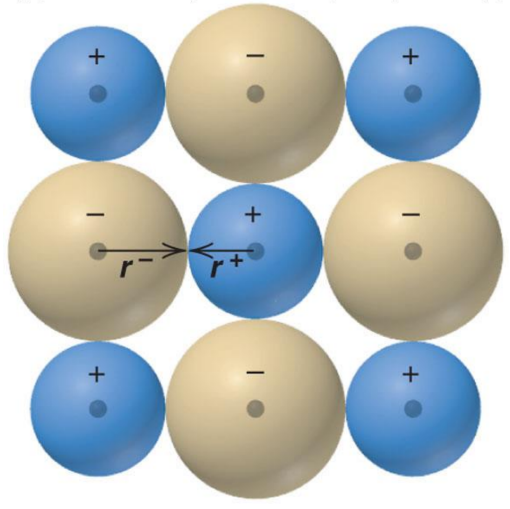
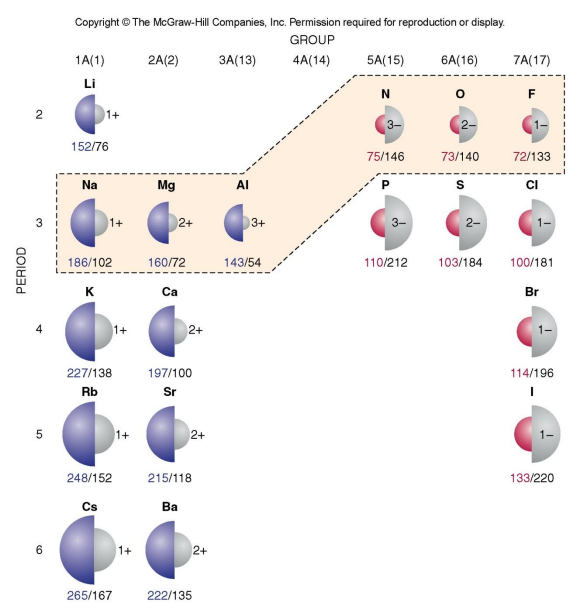


Figure 8.21

Ionic vs. atomic radii.



Ranking Ions by Size

PROBLEM: Rank each set of ions in order of *decreasing* size, and

explain your ranking:

(a) Ca^{2+} , Sr^{2+} , Mg^{2+} (b) K^+ , S^{2-} , Cl^- (c) Au^+ , Au^{3+}

PLAN: Find the position of each element on the periodic table and

apply the trends for ionic size.

SOLUTION:

(a) $Sr^{2+} > Ca^{2+} > Mg^{2+}$

All these ions are from Group 2A, so size increases down the group.

SOLUTION:

(b)
$$S^{2-} > CI^- > K^+$$

These ions are isoelectronic, so size decreases as nuclear charge increases.

(c)
$$Au^+ > Au^{3+}$$

Cation size decreases as charge increases.