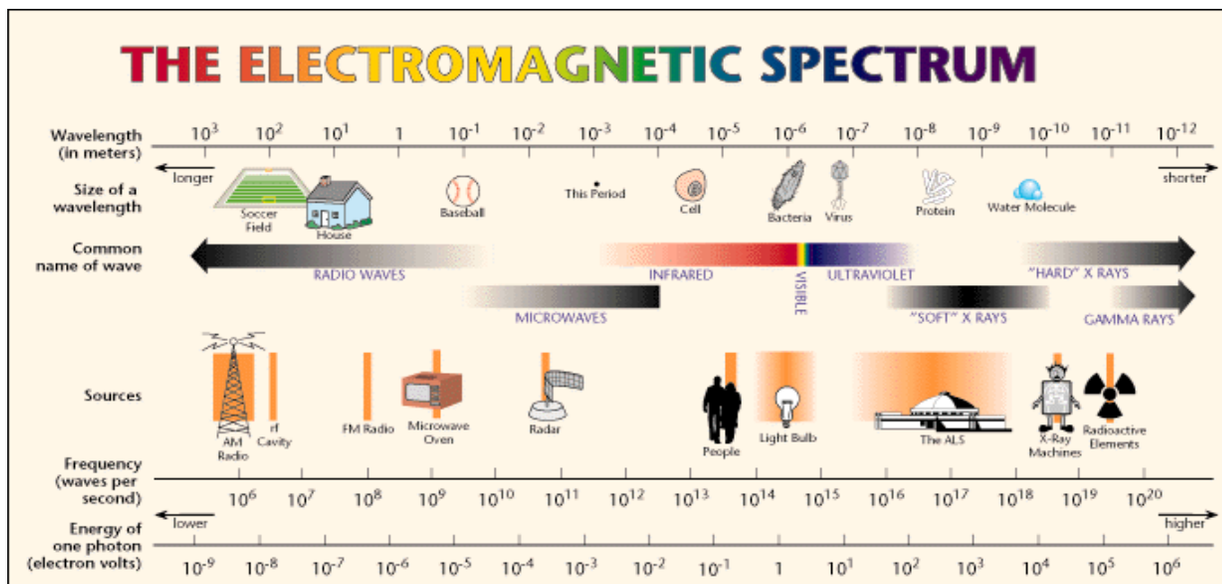
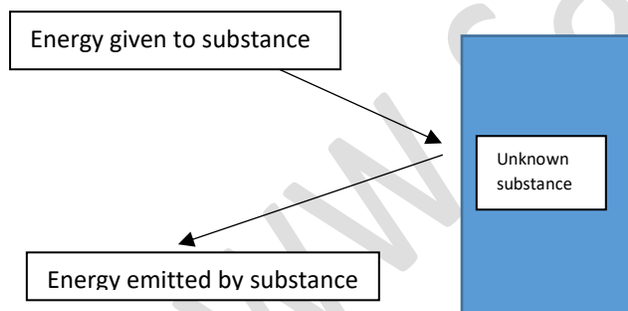


Chapter 7 – Introduction to Spectroscopy

Spectroscopy is the study of the interaction of radiant energy and matter. The electromagnetic spectrum diagram is shown below to use as a reference.



When radiation is shone on a sample of an element or a molecule, the radiation might just bounce off the particles. But in many cases, some of the radiation might be absorbed by the element or molecule. And the remainder of the energy of the radiation is emitted. **The wavelength at which energy is absorbed gives us information about the particle.**



Example 1: When **x-ray radiation** is shone at a substance, it has so much energy that it will remove electrons. The particular wavelength of x-ray used to remove each electron within the substance gives scientists an idea of the number of electrons in shells and how tightly they are held to the nucleus and thus, the identity of an element. This is known as **photoelectron spectroscopy**.

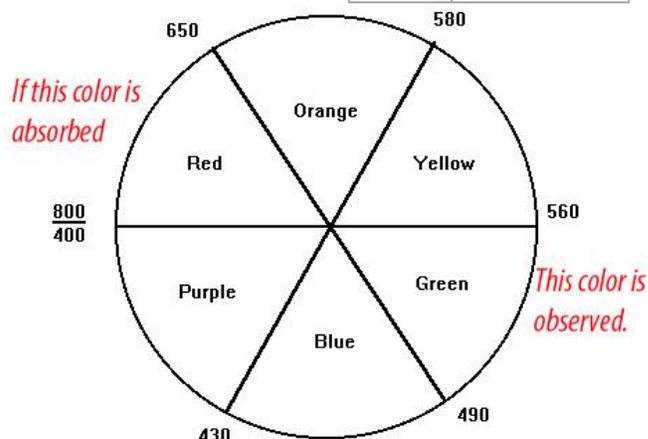
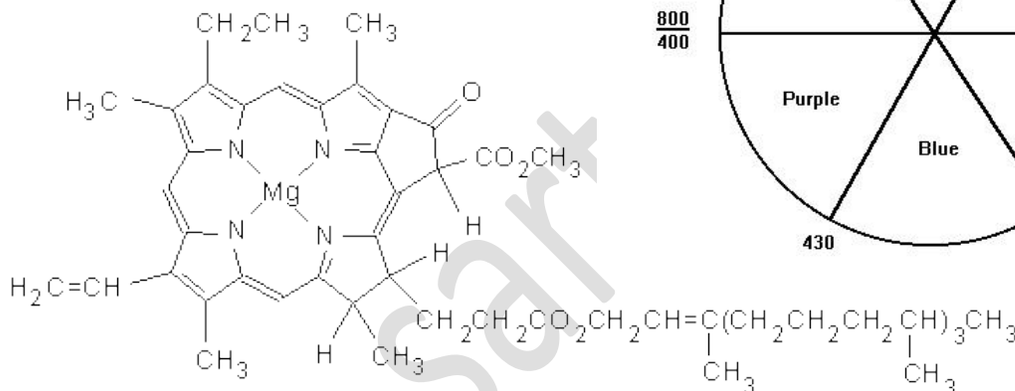
Example 2: When **white visible light** is shone on certain colorful molecules or transition metal ions, electrons in the bonds (or in the d sublevels of the transition metal ions) can be excited. Since visible light is lower energy than x-rays, the electrons are not removed from the atom, but rather the valence electrons only are excited to a different state. Since, electrons in atoms AND in molecules have quantized states, only certain wavelengths of light will be absorbed. The other wavelengths will bounce off giving a certain "color" to the molecule or ion. Molecules with extensive alternating double and single bonds (aka delocalized electrons/conjugation/resonance), often absorb visible light. Molecules without this extensive delocalization tend to absorb wavelengths at much higher energies and therefore absorb light in the ultraviolet region, but reflect all visible light and, therefore, appear colorless to us. Using the wavelength of light absorbed within the UV or visible region of the electromagnetic spectrum to identify a molecule is known as **UV-Visible spectroscopy**.



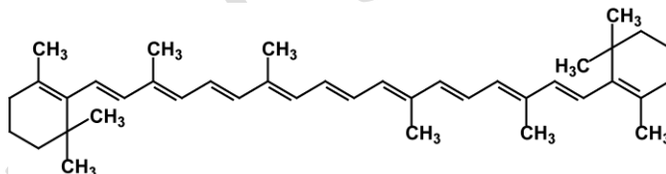
violet	380–450 nm
blue	450–495 nm
green	495–570 nm
yellow	570–590 nm
orange	590–620 nm
red	620–750 nm

When white light is observed, what is actually being seen is all the colors of light combined. When this light passes through a substance, certain energies (or colors) of light are absorbed while the other color(s) are allowed to pass through or are reflected. This is why some substances appear colored. The color we see is the combination of the energies of visible light that are not absorbed by the sample. If the substance does not absorb any light, it appears white or colorless. A solution appears a certain color due to the absorbance and transmittance of visible light. For example, an orange solution appears orange because it is absorbing all of the colors except orange. A sample may also appear orange if all colors of light except blue are transmitted. This is because blue and orange are complimentary colors. (see figure below right)

Chlorophyll, absorbs extensively at blue (~450 nm) and orange-red (~650 nm) wavelengths and appears green to our eyes



Beta-carotene, found in carrots, absorbs many wavelengths, especially at blue (~454 nm) and appears orange to our eyes.



Example 3: In the less energetic regions of the electromagnetic spectrum (infrared radiation and microwave radiation), the radiation doesn't have enough energy to excite electrons and certainly not enough to remove electrons. Instead, energy at certain wavelengths of infrared or microwave can be absorbed and cause a particular bond in a molecule to have a different vibrational state. Microwave ovens cook by causing water molecules to rotate. However, since the wavelength is on the order of millimeters, the water must be in objects at least that size. Ants and rice are largely unaffected by microwaves because they're too small.



We tend to think of bonds as being rigid, but in reality, the atoms and electrons have a certain amount of kinetic energy. The atoms on either side of a bond can bounce closer and further away from each other. If there are several bonds near each other in a molecule, the atoms can even scissor. See this link for an animation of the above:

http://chemwiki.ucdavis.edu/Physical_Chemistry/Spectroscopy/Vibrational_Spectroscopy

Or another: https://www.youtube.com/watch?v=S8R30EdcIT4&feature=player_embedded

Each of these vibrations has a specific energy. Therefore, if the molecule switches from one vibrational state to another state, a specific amount of energy is absorbed or released, and this is usually in the infrared region of the spectrum. Further, the amount of energy absorbed depends on the order of the bonds and the atoms in the bonds. So, chemists can measure at what wavelengths energy is absorbed and thus the region of the electromagnetic spectrum to determine what types of bonds or functional groups are present. Carbon dioxide, absorbs energy at 4 micrometers and 15 micrometers (in the infrared part of the spectrum), and does not allow these wavelengths to escape into space, effectively heating up our atmosphere. Other greenhouse gases absorb different, specific wavelengths of infrared depending on their particular molecular structure. Measuring the wavelengths that are absorbed within the infrared spectrum can help identify a compound and/or determine its molecular structure and is known as **infrared spectroscopy**.

Summary: Types of Spectroscopy

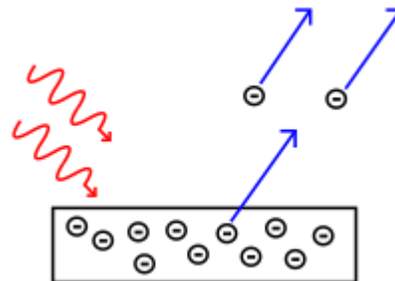
Type of Radiation Used	Frequency	Relative Energy	What it does to the atom/molecule	What it tells us about the atom/molecule	Name of Spectroscopy
X-ray	$10^{16} - 10^{19}$ Hz	Quite High	Removes core electrons	How tightly the electrons are held by the nucleus Measures their binding energy; the identity of an element	Photoelectron spectroscopy (PES)
Ultraviolet	$10^{16} - 10^{19}$ Hz	High	Excites valence electrons	The identity of a molecule or element	UV-Visible Spectroscopy
Visible Light	$10^{16} - 10^{19}$ Hz	Medium	Excites valence electrons	The identity of concentration of a molecule	UV-Visible Spectroscopy
Infrared	$10^{16} - 10^{19}$ Hz	Low	Changes the vibrations in the bonds	The types of bonds/atoms/functional groups within a molecule	IR (vibrational) spectroscopy
Microwave	$10^{16} - 10^{19}$ Hz	Quite Low	Changes the rotations of the atoms in the bonds	The locations of hydrogen atoms within a molecule	Microwave (rotational) spectroscopy

Ideas to consider:

1. Our eyes are spectrophotometers. The colors we see can help us to understand the structures of the molecules we are looking at. The colors we see are opposite on the color wheel to the colors being absorbed by the electrons.
2. Not many molecules absorb in the visible region of the electromagnetic spectrum, because it is such a narrow region of the spectrum.
3. If we could “see” wavelengths outside of the visible spectrum, we would be able to “see” color from some molecules that currently appear colorless to us. Indeed, some insects and birds can “see” different wavelengths of the ultraviolet region and some reptiles can see wavelengths in the infrared region. This is because their eye structures contain different molecules that are sensitive to different wavelengths. These animals do not “see” the world as we do!
4. By shining light of all types of wavelength on molecules, we can measure not just the visible light wavelengths they absorb but also the wavelengths of other types of radiation we can't see to figure out the structure of the molecule.
5. Why is the sky blue? O_2 and N_2 have a vibrational absorbance that is blue. The white light from the sun hits these molecules, and blue light in particular is absorbed but then re-emitted/scattered. The other colors of the light have too low of energy to interact with the O_2 and N_2 and aren't scattered as much.

Photoelectron Spectroscopy (PES)

The Photoelectric Effect is the observation that photons short-wave (visible or ultraviolet) light can cause it to emit electrons. This is an example of ionization, with the photons providing the ionization energy. When visible or ultraviolet light is shone on a substance, the **energy from the photons of light excites** electrons in the substance. If the energy exceeds the ionization energy of an atom in the substance, the electron is emitted. The emitted electrons are called photoelectrons.



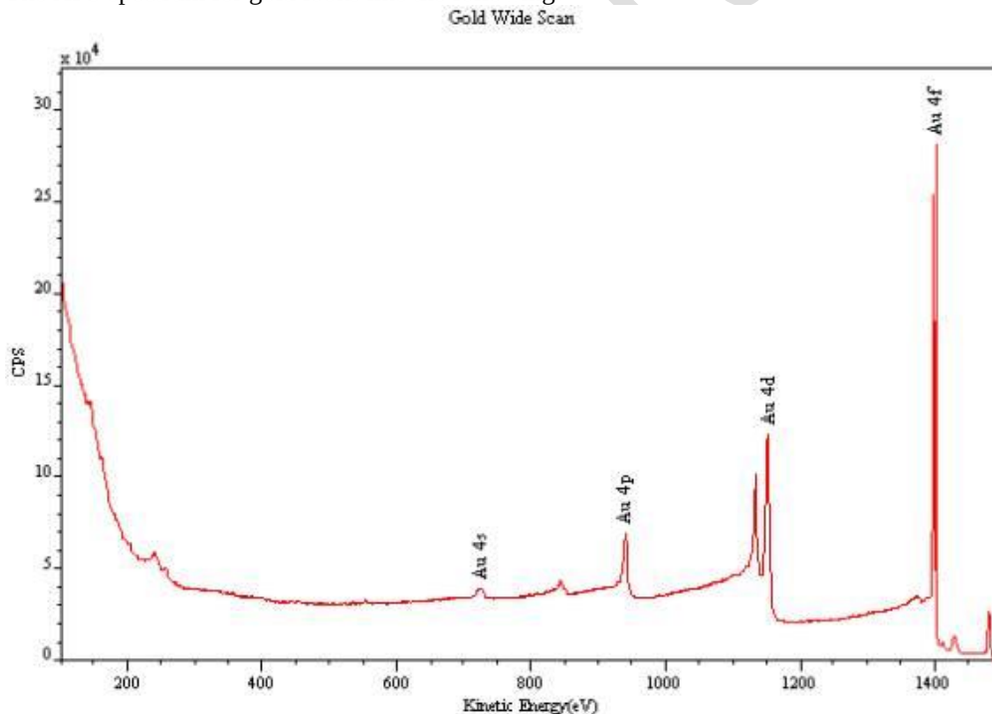
Photoelectron Spectroscopy or photoemission spectroscopy (PES) involves using the energy from electrons emitted via the photoelectric effect to gain information about the electronic structure of a substance. The term “photoelectron spectroscopy” is generally used for the technique when applied to gases, and “photoemission spectroscopy” is generally used for electrons emitted from solid surfaces. In photoelectron (or photoemission) spectroscopy, a substance is bombarded with photons, which have a given amount of energy based on their frequency: $E_{\text{photon}} = h\nu$

By measuring the kinetic energy of the emitted electron, E_k , and predetermining the “work function”, W_0 , of the substance (the amount of additional energy it takes to move the delocalized electron to the surface of the material—zero for gases, but nonzero for solids), we can calculate the binding energy, E_B , of the electron from Einstein’s equation for the photoelectric effect: $E_k = h\nu - E_B - W_0$

The photons used for photoelectron spectroscopy range from ultraviolet light to X-rays. Ultraviolet photoelectron spectroscopy (UPS) and extreme ultraviolet photoelectron spectroscopy (EUPS) are used to study valence electrons and the electrons that participate in chemical bonding. X-ray photoelectron spectroscopy (XPS) is most often used to study core electrons, particularly in solids.

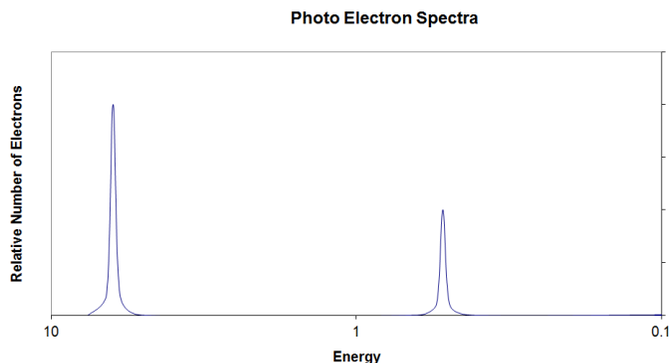
Interpreting Photoelectron Spectra

The XPS spectrum for gold looks like the following:



Notice that the peaks for 4s, 4p, 4d, and 4f are different heights. The height of the peak is proportional to the number of electrons in a given sublevel.

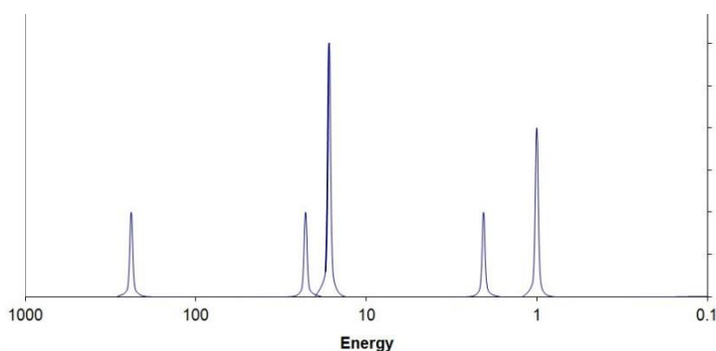
Analysis of UV photoelectron spectra for a single element with a relatively low atomic number is straightforward. For example, the following is an idealized plot of the photoelectron spectrum for lithium: $1s^2 2s^1$



The x -axis has units of binding energy of the electron, usually electron-volts (eV). Recall that electrons in higher sublevels have less binding energy, which makes them easier to remove. This means the peak at the right corresponds with the easiest-to-remove electrons (for lithium, this is the $2s$ electron). The peak at the left corresponds with the electrons that are hardest to remove (for lithium, these are the two $1s$ electrons).

The y -axis indicates the number of photons emitted with that energy. This means the height of each peak is proportional to the number of electrons in the corresponding sub-level. Notice that the peak at 6.26 eV (the $1s$ electrons) is twice as high as the peak at 0.52 eV (the $2s$ electron). This means there are twice as many electrons in the $1s$ sublevel as in the $2s$ sublevel. The only element that matches this spectrum is lithium ($1s^2 2s^1$).

You try: 1. Below is the PES of sulfur, and a table of successive ionization energies. The energy scale for the PES is in MJ/mol (1 MJ = 1000 kJ). There is exactly one peak on this graph whose energy corresponds to an ionization energy in table 7.2. What is it? Why doesn't anything else match up?



In PES, all energies are for removing an electron from a neutral atom. Thus, for the $3p$ electrons of S, there is a single peak because it takes the same energy to remove any $3p$ electron. This is different from successive ionization energies, where the 2nd electron is removed from the S^+ ion. That's much harder than removing the first electron, so the second ionization energy is much greater. The point of overlap (or consistency) is the 1st ionization energy and, from PES, the binding energy of the most easily removed electron. These

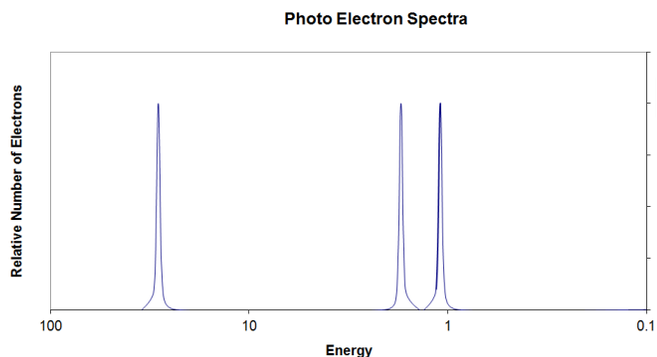
represent the same event, and indeed have the same value. For S, the 1st IE is 1000 kJ/mol, and the binding energy of the $3p$ peak in the S PES is 1.0 MJ/mol (1000 kJ/mol). The same consistency will be true of other elements.

TABLE 7.2 • Successive Values of Ionization Energies, I , for the Elements Sodium through Argon (kJ/mol)

Element	I_1	I_2	I_3	I_4	I_5	I_6	I_7
Na	496	4562					
Mg	738	1451	7733				
Al	578	1817	2745	11,577			
Si	786	1577	3232	4356	16,091		
P	1012	1907	2914	4964	6274	21,267	
S	1000	2252	3357	4556	7004	8496	27,107
Cl	1251	2298	3822	5159	6542	9362	11,018
Ar	1521	2666	3931	5771	7238	8781	11,995

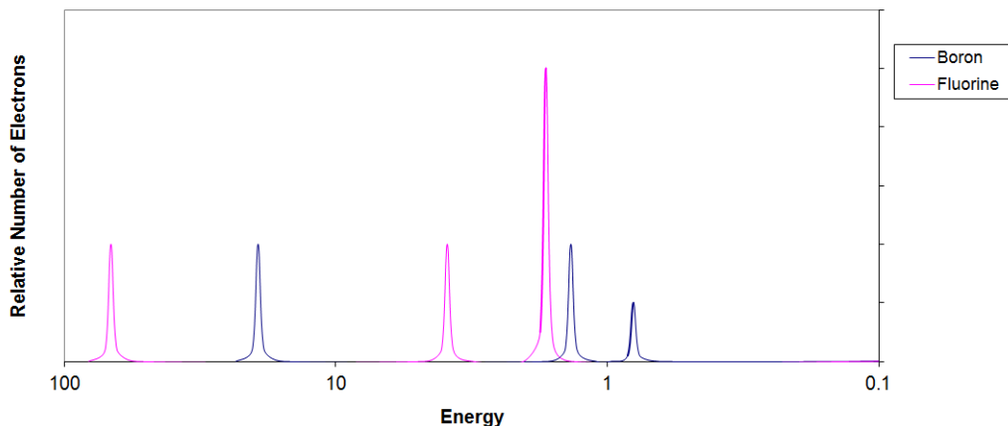
(From Brown et al., Chemistry the Central Science 12 ed., copyright Pearson Prentice Hall)

2. Identify the element represented by the PES spectrum shown to the right.



3. Here is a PES spectrum of boron ($Z=5$; blue) superimposed on that of fluorine ($Z=9$; pink)

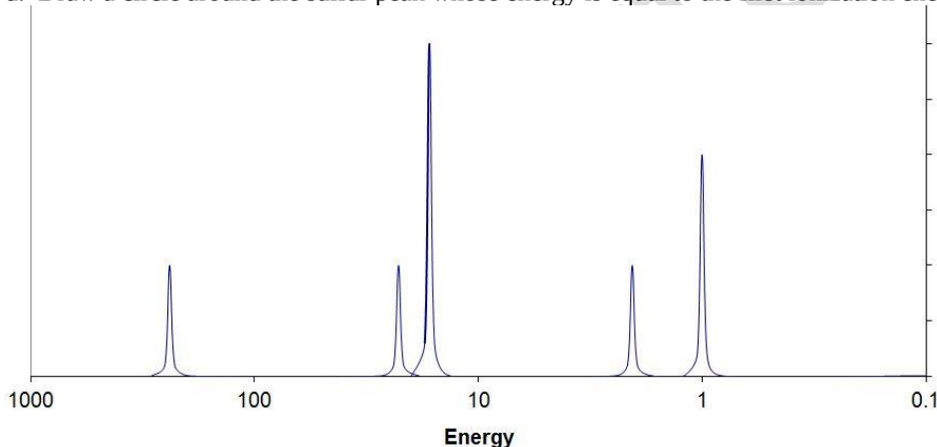
Photo Electron Spectra



- Why are the fluorine peaks to the left of the boron peaks?
- Why is there one peak in fluorine that is so much taller than all the others?

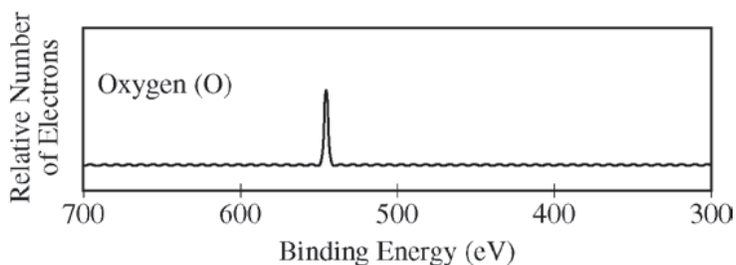
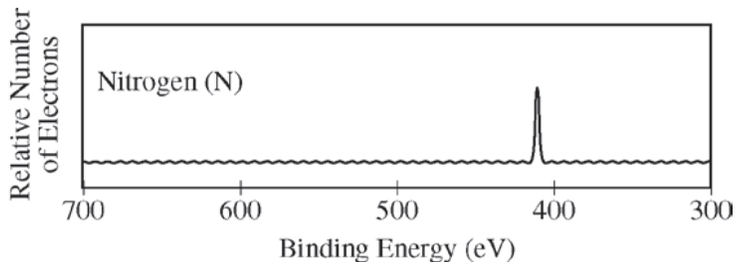
4. Below is shown the PES spectrum of sulfur (atomic number = 16).

- Write the full electron configuration of sulfur.
- Label each peak in the spectrum to show which subshell it represents (i.e., 1s, 2s, etc.)
- On the spectrum, sketch in the relative locations and correct peak heights for the spectrum of aluminum (atomic number = 13). By relative location, I mean correctly to the left or right of the same subshell peak in the sulfur spectrum.
- Draw a circle around the sulfur peak whose energy is equal to the first ionization energy of sulfur.



5. The photoelectron spectra above show the energy required to remove a 1s electron from a nitrogen atom and from an oxygen atom. Which of the following statements best accounts for the peak in the upper spectrum being to the right of the peak in the lower spectrum?

- Nitrogen atoms have a half-filled p subshell.
- There are more electron-electron repulsions in oxygen atoms than in nitrogen atoms.
- Electrons in the p subshell of oxygen atoms provide more shielding than electrons in the p subshell of nitrogen atoms.
- Nitrogen atoms have a smaller nuclear charge than oxygen atoms.
- Oxygen atoms are smaller than nitrogen atoms.



Answer 1: The first ionization energy of sulfur matches the furthest-right peak on the PES. The different peaks in PES represent the effort needed to evict an electron in different subshells of a neutral sulfur atom. In contrast, the ionization energy chart represents the energy needed to evict the most easily removed electron from a neutral sulfur atom (IE1), an S^+ ion (IE2), an S^{2+} ion (IE3), etc. These two data sets have only one point of overlap, which is IE1 on the chart and the lowest-energy peak on the PES. None of the peaks in the PES relate to removing an electron from a sulfur cation, and, likewise, none of the successive ionization energies concern removing an electron from other than the highest subshell.

Answer 2: Carbon

Answer 3: a. Why are the fluorine peaks to the left of the boron peaks?

Answer: Fluorine has a greater nuclear charge, causing the electrons in all the subshells to be held more tightly.

b. Why is there one peak in fluorine that is so much taller than all the others?

Fluorine has 4 more electrons than boron; all the additional electrons are in the 2p subshell, giving it 5 in all, causing that peak to be much higher than 1s or 2s, which each have 2 electrons.

Answer 4: Below is shown the PES spectrum of sulfur (atomic number = 16).

a. Write the full electron configuration of sulfur. $1s^2 2s^2 2p^6 3s^2 3p^4$

b. Label each peak in the spectrum to show which subshell it represents (i.e., 1s, 2s, etc.) **They go in order from left to right as written in the electron configuration above.**

c. On the spectrum, sketch in the relative locations and correct peak heights for the spectrum of aluminum (atomic number = 13). By relative location, I mean correctly to the left or right of the same subshell peak in the sulfur spectrum. **All the aluminum peaks should be shifted to the right relative to the same peak in sulfur because aluminum has a smaller nuclear charge. The heights should all be the same as in sulfur, except the 3p peak, which should be half the height of the 1s, 2s, and 3s peaks.**

d. Draw a circle around the sulfur peak whose energy is equal to the first ionization energy of sulfur. **It's the 3p peak.**

Answer 5: D Nitrogen atoms have a smaller nuclear charge than oxygen atoms.