Topic I

Review of VSEPR Theory

I.1 Module 1: Course Logistics and History

- 1/11: Homework questions will be similar to exam questions, so although you probably *can* find answers online, you shouldn't.
 - Submit Psets to chem201hw@gmail.com.
 - Watch modules before office hours and bring questions.
 - If you have a question outside of office hours, post it on Slack.
 - It's a difficult class, but he is open to and welcomes our feedback (via Slack, again).
 - You do need to read from Miessler et al. (2014), too; his class is not a replacement for this textbook.
 - He is a big fan of Cotton (1990).
 - There is an extra, new textbook to look for!
 - Convince yourself not to be afraid of time-independent quantum mechanics (we won't go too deep, but know wave functions and the like).
 - Exams will probably be open book/open note.
 - Inorganic chemistry contains too much information to rationalize empirically, so we need a system (the development of this system will be the focus of this course).
 - Reviews history of chemistry from Miessler et al. (2014) Chapter 1.
 - What is Nickel's electron configuration?
 - When Nickel is a free atom, the $[Ar]4s^23d^8$ electron configuration is the lowest energy.
 - When Nickel is chemically bound, the $[Ar]3d^{10}$ electron configuration is the lowest energy because it is energetically unfavorable to have a large 4s orbital pushing the bounds of the atom.
 - What is a **term symbol**?
 - Homework: Refresh Chapter 2 in Miessler et al. (2014).
 - Covalent bond: The sharing of pairs of electrons...?
 - G. N. Lewis predicts in 1916 (before Rutherford) that the atom has a positive **kernel** surrounded by a shell containing up to 8 electrons.
 - Also orbital penetration.
 - He recommends that we read the full paper: Lewis (1916).

I.2 Module 2: Molecular Geometries and VSEPR

- The easiest way to approach a new Lewis structure:
 - 1. Draw a valid Lewis structure for a molecule.
 - 2. Place electron pairs in the valence shell as far away from each other as possible. Use the σ -bond framework first.
 - 3. Add π -bonds to complete the molecule.
- Through the VSEPR approach, think of a molecule as arranged around a central atom A by m atoms or groups of atoms X and n lone electron pairs E.
- Steric number: The sum n+m of groups and electron pairs around the central atom.
- Steric numbers correspond to geometries.
- VSEPR is ok but it doesn't capture reality too well.
- Consider trimethyl boron (BMe₃).
 - Trigonal planar (D_{3h}) .
- Octahedral: O_h .
- Bent: C_{2v} .
- Order of the repulsive forces: lone pair lone pair lone pair bonding pair bonding pair bonding pair.
- In SF₄ (see-saw), will the lone pair be axial or equatorial?
 - Equatorial $2 \times 120^{\circ}$ and $2 \times 90^{\circ}$ vs. $3 \times 90^{\circ}$.
- In $BrF_3...$

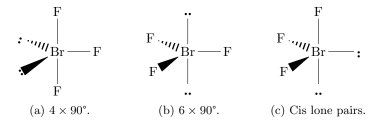


Figure I.1: VSEPR structure of BrF₃.

- T-shaped $4 \times 90^{\circ}$ vs. $6 \times 90^{\circ}$ or lone pairs in cis-position.
- In ions such as ICl_4^- , we get square planar (D_{4h}) .
- With mixed substituents (such as PF₂Cl₃)...
 - We need **Bent's rule**, which tells us that atoms share electrons from p- or d-orbitals to a greater extent than they do from s-orbitals.
 - Thus, when phosphorous excites $3s^23p^3$ to $3s^13p_x^13p_y^1ep_z^13d_{z^2}^2$ and then rehybridizes to create three sp^2 orbitals (each composed of $s+p_x+p_y$) and two "pd" hybrid orbitals (each composed of $p_z+d_{z^2}$), the equatorial sp^2 orbitals bond to the more **electropositive** chlorines and the axial "pd" hybrid orbitals bond to the remaining more electronegative fluorines.
- Bent's rule: Atomic s-character concentrates in orbitals directed toward electropositive substituents.

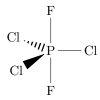


Figure I.2: Lewis structure of PF₂Cl₃.

- Electropositive (species): A species that has relatively lower electronegativity than another.
- For molecules with multiple bonds, ignore π -bonds.
- Linear: $D_{\infty h}$.
- Problems with VSEPR:
 - XeF₆ with 14 bonding electrons (7 pairs) is supposed to be pentagonal bipyramidal, but is actually octahedral (a known problem for 14 e[−] systems).
 - Heavy main group elements with no hybridization.
 - $H-C\equiv C-H$ is linear, but $H-Si\equiv Si-H$ is not.
 - No σ -bond exists in the latter species it's all π -bonding interactions.
- You maybe don't have to watch the modules and textbook and attend class.

I.3 Chapter 3: Simple Bonding Theory

From Miessler et al. (2014).

I.3.1 Notes

1/14:

- **Hypervalent** (central atom): A central atom that has an electron count greater than the atom's usual requirement.
 - There are rarely more than 18 electrons around a central atom (2 for s, 6 for p, 10 for d). Even heavier atoms with energetically accessible f orbitals usually don't have more surrounding electrons because of crowding.
 - With BeF₂, instead of getting the predicted double-bonded lewis structure, it forms a complex network with Be having coordination number 4.
 - BeCl₂ dimerizes to a 3-coordinate structure in the vapor phase.
 - Boron trihalides exhibit partial double bond character.
 - It is also possible that the high polarity of B-X bonds and the ligand-close packing (LCP) model account for the observed shorter bond length.
 - Boron trihalides also act as Lewis acids.

Steric Number	Geometry	Examples	Calculated Bond Angles	
2	Linear	CO_2	180°	o = c = o
3	Trigonal (triangular)	SO_3	120°	
4	Tetrahedral	CH_4	109.5°	H H
5	Trigonal bipyrimidal	PCl_5	120°, 90°	
6	Octahedral	${ m SF}_6$	90°	F F F
7	Pentagonal bipyrimidal	IF_7	72°, 90°	F F F
8	Square antiprismatic	$[{ m TaF_8}]^{3-}$	70.5°, 9.6°, 109.5°	F F F

Table I.1: VSEPR predictions.