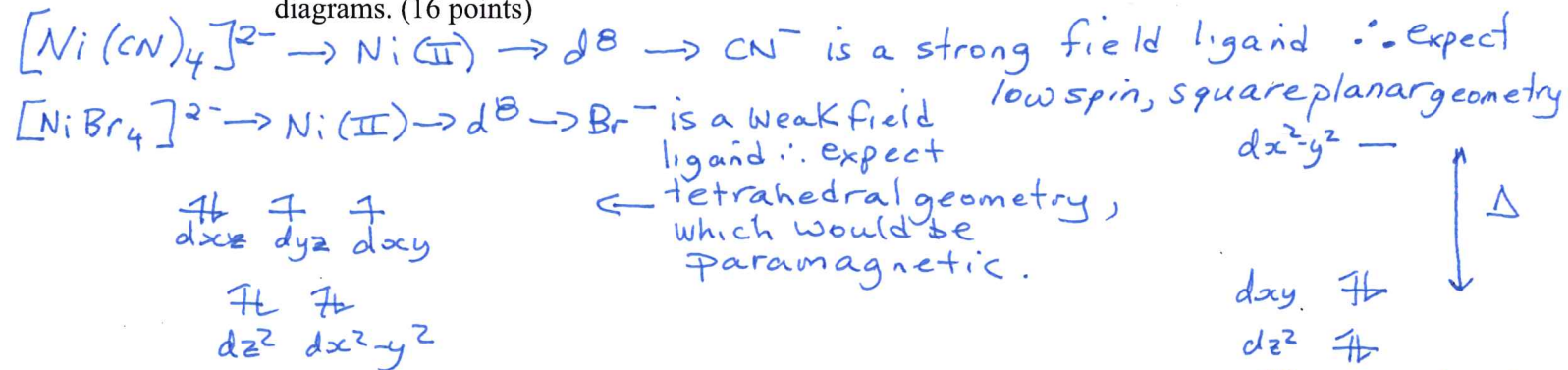


Chemistry 3840
Midterm Examination #2 (March 12, 2025)

1. The nickel complex $[\text{Ni}(\text{CN})_4]^{2-}$ is diamagnetic (all electrons are paired), while $[\text{NiBr}_4]^{2-}$ is paramagnetic (has unpaired electrons). Fully explain these observations. Include labelled diagrams. (16 points)



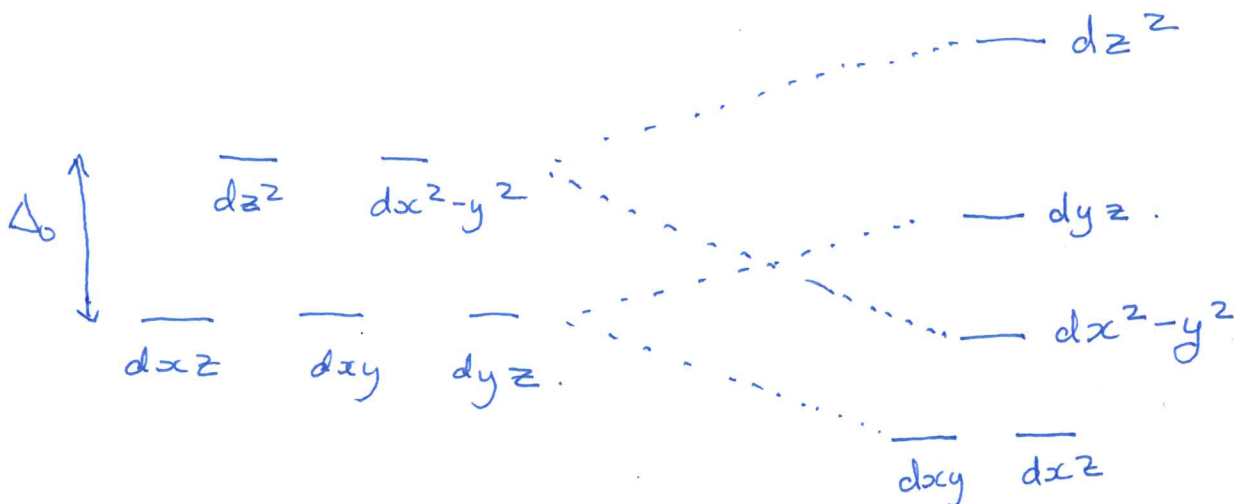
$\rightarrow \text{Ni}(\text{II})$ complexes can be either square planar or tetrahedral, depending upon the ligands. Since $[\text{NiBr}_4]^{2-}$ is paramagnetic, it must have tetrahedral geometry, which makes sense as Br^- is a weak field ligand. Conversely, $[\text{Ni}(\text{CN})_4]^{2-}$ is diamagnetic, meaning all e^- are paired. Therefore, it must be square planar which agrees with CN^- being a strong field ligand.

2. Give an example of an ionization isomer of $[\text{CoBr}(\text{H}_2\text{O})_5]\text{Cl}$. No explanation is required. (6 points)

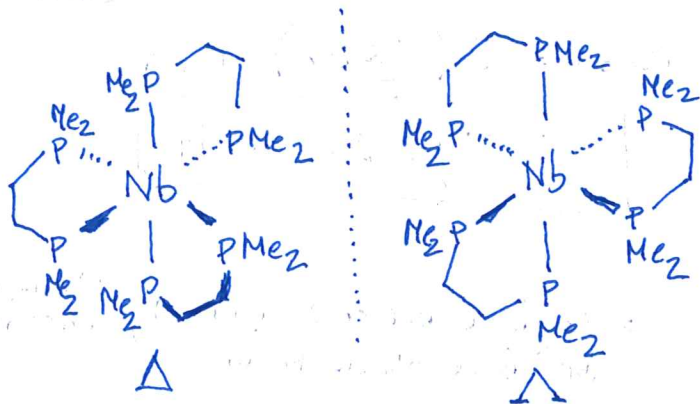
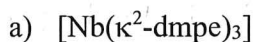


3. Starting from an octahedral orbital splitting diagram, use the principles of Crystal Field Theory (CFT) to indicate (and discuss) the orbital splitting diagram for a square planar metal complex (complete removal of the two ligands along the x-axis). Be sure to label all orbitals. (12 points)

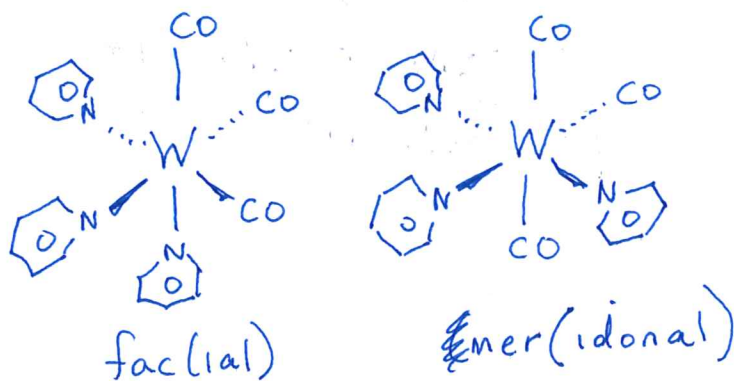
\rightarrow removal of ligands along the x-axis will decrease the energy of orbitals with x-character. Orbitals with y and z character will increase in energy.



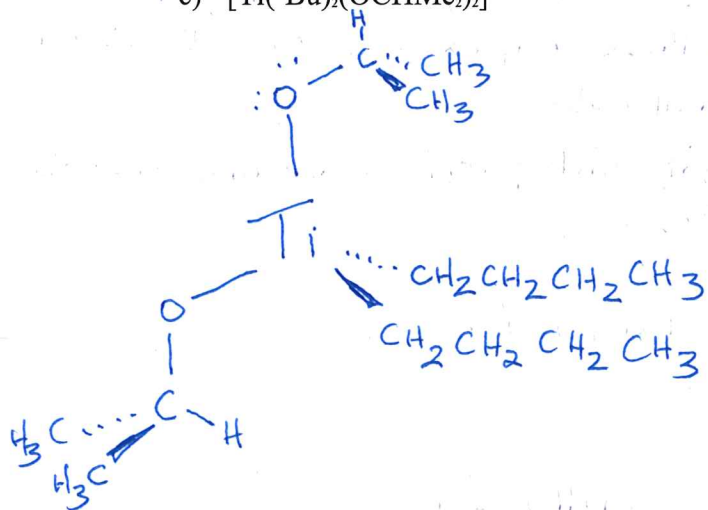
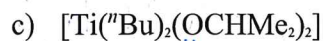
4. **Properly** draw the following complexes using **fully expanded** Lewis structures. Be sure to draw and label all *reasonable* isomers. (dmpe = $\text{Me}_2\text{PCH}_2\text{CH}_2\text{PMe}_2$; py = pyridine ($\text{C}_5\text{H}_5\text{N}$); ${}^n\text{Bu}$ = $\text{CH}_2\text{CH}_2\text{CH}_2\text{CH}_3$) (18 points)



→ $\text{Nb}(\text{CO}), d^5, 17e^-$
 → dmpe is a bidentate ligand ∴ Nb is 6 coordinate → octahedral geometry expected



→ $\text{W}(\text{CO}), d^6, 18e^-$
 → 6 coordinate
 ↳ expect octahedral geometry.

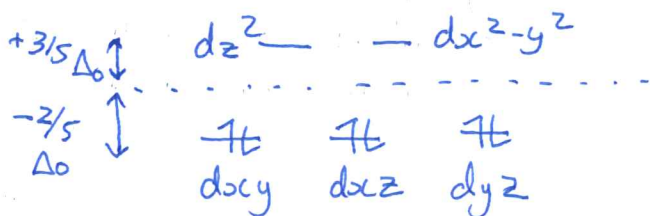


→ $\text{Ti}(\text{IV}), d^0, 8e^-$
 → 4 coordinate
 ↳ expect tetrahedral geometry.

→ No other isomers

5. Calculate the Crystal Field Stabilization Energy (CFSE) for one of the following two octahedral complexes. Show your work and explain (in 10 words or less) any assumptions you made. (12 points)

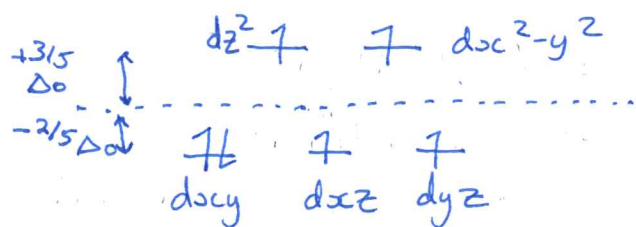
a) $[\text{W}(\text{CO})_6] \rightarrow \text{W}^0 \rightarrow d^6 \rightarrow \text{W}$ is a 3rd row element } \therefore strong field environment
 CO is a strong field ligand



$$\text{CFSE} = 6 \left(-\frac{2}{5} \Delta_o \right) + 3P$$

$$= -\frac{12}{5} \Delta_o + 3P$$

b) $[\text{CoBr}_6]^{3-} \rightarrow \text{Co}(\text{III}) \rightarrow d^6 \rightarrow \text{Co}$ is a 1st row element } weak field environment
 Br^- is a weak field



$$\text{CFSE} = 4 \left(-\frac{2}{5} \Delta_o \right) + 2 \left(\frac{3}{5} \Delta_o \right) + P$$

$$= -\frac{8}{5} \Delta_o + \frac{6}{5} \Delta_o + P$$

$$= -\frac{2}{5} \Delta_o + P$$

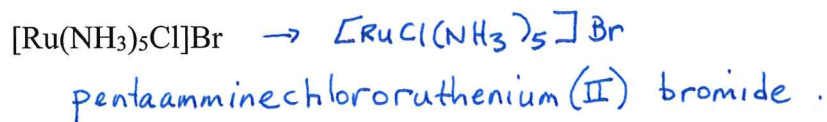
6. As a recent university graduate with a major in Chemistry, you are very excited about your job interview for a research chemist position at NOVA Chemicals. When you arrive, you are sent into a lab where you find two vials on a bench. One of the vials is filled with a purple powder and the other contains a yellow powder. Nearby you see two labels ($[\text{Co}(\text{NH}_3)_6]^{3+}$ and $[\text{CoI}(\text{NH}_3)_5]^{2+}$). If you place the labels on the correct vials you will get the job. Which label do you place on which vial? Why? Explain in four sentences or less. (18 points)

- Both complexes are $\text{Co}(\text{III}), d^6$
- I^- is a weak field ligand, while NH_3 is a stronger field ligand
- ↳ since the only difference between these complexes is one I^- vs one NH_3 , $[\text{CoI}(\text{NH}_3)_5]^{2+}$ would have a smaller Δ_o
- yellow compounds absorb blue/purple light, while purple compounds absorb green/yellow (lower energy \therefore smaller Δ_o).

$\therefore [\text{Co}(\text{NH}_3)_6]^{3+}$ is yellow (larger Δ_o)
 and

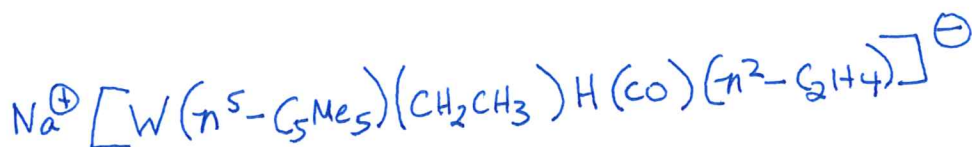
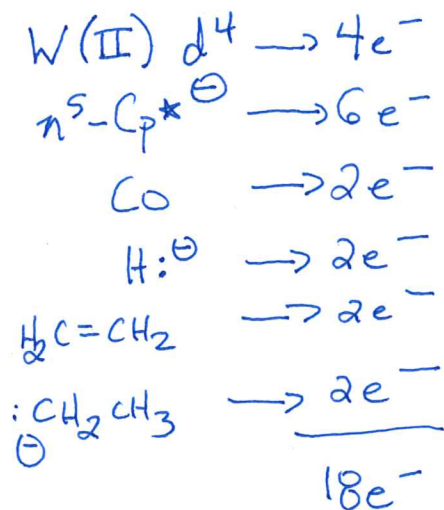
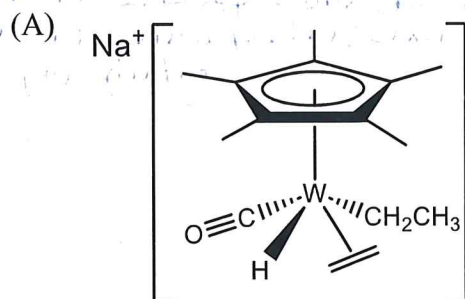
$[\text{CoI}(\text{NH}_3)_5]^{2+}$ is purple (smaller Δ_o).

7. Name the following complex according to IUPAC convention. If there are any errors in the given formulae, provide the correct chemical formulae. (6 points)



8. For the following molecule:

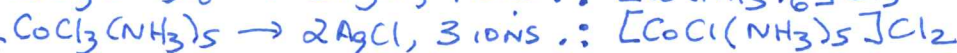
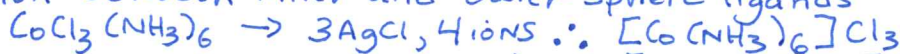
- (i) What is the metal oxidation state, the d-electronic configuration and the electron count at the metal centre? (8 points)
 (ii) Provide the correct IUPAC formula (4 points)



Bonus

Alfred Werner is often considered the "Father of Modern Coordination Chemistry" and was awarded the Nobel Prize in Chemistry in 1913 for his work that established the prevalence of octahedral geometry for late transition metal complexes. Briefly explain how conductivity experiments were essential for providing support for the structures he proposed for various cobalt complexes. Use examples. (4 points)

→ Werner proposed octahedral geometry for a series of Co complexes and by precipitating AgCl and conducting conductivity measurements he was able to distinguish between inner and outer sphere ligands



→ Werner also made chiral molecules using en, and ~~supported~~ separated them, and also created the first chiral complexes (and separated them), that did not contain any carbon atoms.

Additional Information

Spectrochemical series for ligands:

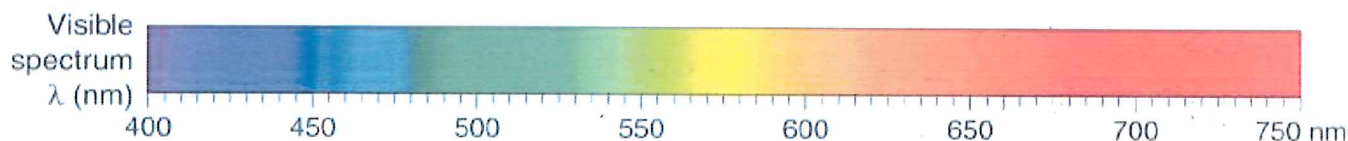
weak field ligands (small Δ_o)

strong field ligands (large Δ_o)



Spectrochemical series of metals:

[Note: not all metals included, use general trends to predict relative ordering]



Colour Wheel:

