

## Midterm Results

Histogram of Midterm \#1


## Midterm 1

1. How many seconds are in 65 years?
a) $3.3 \times 10^{7} \mathrm{sec}$
b) $3.4 \times 10^{7} \mathrm{sec}$
c) $2.0 \times 10^{9} \mathrm{sec}$
d) $2.1 \times 10^{9} \mathrm{sec}$
2. The molecular formula of hydroquinone is $\mathrm{C}_{6} \mathrm{H}_{6} \mathrm{O}_{2}$. What is the empirical formula of hydroquinone?
a) $\mathrm{C}_{9} \mathrm{H}_{9} \mathrm{O}_{3}$
b) $\mathrm{C}_{6} \mathrm{H}_{6} \mathrm{O}_{2}$
c) $\mathrm{C}_{3} \mathrm{H}_{3} \mathrm{O}$
d) CHO

## Midterm 1

3. For the following set of four quantum numbers ( $n, I, m_{1,} m_{s}$ ) identify the set that is not valid:
a) $\{9,5,-3,1 / 2\}$
b) $\{3,3,3,1 / 2\}$
c) $\{2,0,0,1 / 2\}$
d) $\{4,2,-2,1 / 2\}$
4. How many electrons are in principle quantum number $n=3$ ?
a) 3
b) 6
c) 9
d) 18

## Midterm 1

5. How many radial nodes are in a 3d orbital?
a) 0
b) 1
c) 2
d) 3
6. Which of the following is isoelectronic with $\mathrm{N}^{3-}$
a) $F$
b) $\mathrm{Cl}^{-1}$
c) $\mathrm{Mg}^{2+}$
d) Ar

## Midterm 1

7. Which of the compounds below has bonds with more covalent character?
a) CaO
b) $\mathrm{Li}_{2} \mathrm{O}$
c) MgO
d) MgS
8. Which of the following is the most electronegative?
a) B
b) $\quad \mathrm{ln}$
c) Te
d) O

## Midterm 1

9. Which of the following has the largest atomic radius?
a) Be
b) S
c) Te
d) Sr

## Midterm 1

10. 1,3-Benzodioxol-5-ol, otherwise known as sesamol, is a natural organic compound found in sesame seed oil. The combustion analysis of a 100 mg sample of 1,3-benzodioxol-5-ol shows its composition is: $60.87 \%$ C, $4.38 \% \mathrm{H}$, and $34.75 \%$ O.
a) What is the empirical formula of 1,3-benzodioxol-5-ol?
b) If the molecular weight of 1,3-benzodioxol-5-ol is $138.12 \mathrm{~g} / \mathrm{mol}$, what is the molecular formula?

## Midterm 1

11. Sodium metal reacts readily with water to produce hydrogen gas and sodium hydroxide in solution.
a) Write a balanced reaction.
b) If $3.00 \times 10^{-1}$ grams of sodium react with 0.500 L of water (density $=1 \mathrm{~g} / \mathrm{mL}$ ), what is the theoretical yield of hydrogen gas?
c) If exactly $1.20 \times 10^{-2}$ grams of hydrogen gas is collected, what is the percent yield?
d) Knowing the percent yield of the reaction, what is the predicted concentration of NaOH in the solution assuming the volume of water does not change?

## Midterm 1

12. Write the noble gas electron configuration for the following atoms/ ions. How many unpaired electrons does each have?
a) $P$
b) $\mathrm{Ag}^{+1}$
13. Identify the $M^{3+}$ ion for each ground state electron configuration
a) $[\mathrm{Ar}] 3 d^{7}$
b) $\quad[\mathrm{Ar}] 3 \mathrm{~d}^{10}$
14. Briefly explain why oxygen has a lower first ionization energy than nitrogen.

## Midterm 1

15. Calculate the wavelength of the radiation emitted by a hydrogen atom (one electron atom) when an electron makes a transition from $n=5$ to $n=2$.
16. The work function for chromium metal is 4.37 eV or $7.00 \times 10^{-19} \mathrm{~J}$. Light with a wavelength of 11.0 nm is shined on the metal. What is the velocity of the ejected electron?

## Midterm 1

17. Draw the boundary surface for the three $3 p$ orbitals, making sure to name and label each axis correctly. How many radial nodes does a 3 p orbital have?

## Midterm 1

$\square$ Provide the Lewis dot structure for the following compounds. Make sure every electron is drawn. Indicate atoms with formal charge. If resonance structures can be drawn, draw all valid structures. Make sure to indicate formal charge when necessary.
a) $\mathrm{CH}_{3} \mathrm{CH}_{2} \mathrm{CH}_{2} \mathrm{OH}$
b) $\mathrm{CN}^{-}$
c) $\mathrm{HNO}_{3}$
d) $\mathrm{POCl}_{3}$

## Valence shell electron pair repulsion theory (VSEPR)

$\square$ Lewis Theory

- Connectivity, electron tracking
$\square$ VSEPR Theory
- 3-D Structure around an atom
$\square$ Valence Bond Theory
$\square$ Molecular Orbital Theory


## Review - VSEPR Theory

## Valence-Shell Electron-Pair Repulsion (VSEPR)

$\square$ Accounts for 3D shapes of molecules
$\square$ Based on electron-electron repulsion
$\square$ Determine bond angles $\rightarrow$ shape
Rules are based on experimental observation:

1. Areas of electron concentration (bonds and lone pairs) around the central atom repel each other.
2. Bonds and lone pairs stay as far away from each other as possible (without changing distance)


## Review - Polar Molecules

A polar covalent bond has a non-zero dipole moment


Polar molecules - molecules with non-zero dipole moment: $\mathrm{m} \neq 0 \mathrm{D}$

Examples:
HCl HF HBr
A non-polar molecule has no dipole moment: $m=0 D$
Examples: $\quad \begin{array}{llll} & \mathrm{O}_{2} & \mathrm{CH}_{4} & \mathrm{SF}_{6}\end{array}$

## Polar Molecules

Polar bonds, Non-polar molecule?


## $\mathrm{CO}_{2}$

If the dipole moments cancel out (Vector sum = 0):
Each C-O bond is polar
Two equally polar bonds, with opposing dipole moments


## Polar Molecules

## $\mathrm{H}_{2} \mathrm{O}$

Each O-H bond is polar
Overall molecule is ??????

## Polar Molecules



## Polar Molecules


non-polar
$\mathrm{CHCl}_{3}$

polar

## Valence Bond (VB) Theory

$\square$ Lewis Theory

- Connectivity, electron tracking
$\square$ VSEPR Theory
- 3-D Structure around an atom
$\square$ Valence Bond Theory
- Extended 3-D Structure Information
- Delocalization in Molecules
- Illustrates Multiple Bonding
- Prediction of Reactivity
$\square$ Molecular Orbital Theory


## Answers to $\mathrm{H} \Psi=\mathrm{E} \Psi$



## Answers to $\mathrm{H} \Psi=\mathrm{E} \Psi$



## Bonding With Orbitals



## 1s $1 s$

$1 s$
H

## Bonding with Orbitals



## Valence-Bond Theory

So far, we've been thinking of molecules using Lewis' Theory:
Bonding electrons are located in between bonded atoms - electrons are localized


But....
We learned in Ch. 1 that we learned to think of electrons as wavefunctions, which are described by atomic orbitals

Valence Bond Theory - quantum mechanical view of bonding

## ‘Types’ of Bonds $-\sigma$ bond

Lets start with $\mathrm{H}_{2}$, the simplest molecule:

Ground state H has one 1 s electron
When the 2 H atoms bond, the atomic orbitals merge, forming a $\sigma$-bond
o-bond (sigma bond) - along bond axis.

We say the atomic orbitals overlap
More overlap $=$ Stronger bond
All single covalent bonds are $\sigma$-bonds

## Atomic orbitals


$\sigma$-bond

## ‘Types’ of Bonds $-\sigma$ bond

All single covalent bonds are s-bonds


Can have $\sigma$-bonds between any types of orbitals:

Two s orbitals
Two p orbitals
An s and a p orbital
Etc....
Example: s-bond between 1s orbital of H and $2 p_{z}$ orbital of F


## ‘Types’ of Bonds - Multiple bonds

What happens in $\mathrm{N}_{2}$ ?
Remember the bond in $\mathrm{N}_{2}$ is a triple bond...


Each N atom has 3 unpaired 2 p electrons
Lets look at the atomic orbitals of N :


$\mathbf{P}_{\mathrm{x}}$

$P_{y}$

$\mathbf{P}_{\mathbf{z}}$

## ‘Types’ of Bonds - Multiple bonds

What happens in $\mathrm{N}_{2}$ ?
Remember the bond in $\mathrm{N}_{2}$ is a triple bond...

triple bond.
Each N atom has 3 unpaired 2 p electrons
How do the atomic orbitals bond?


Between the $2 p_{z}$ and $2 p_{z}$ orbitals, we have bonding on the bond axis

This is a $\sigma$-bond!

$$
\begin{array}{ll}
\mathbf{p}_{z} & \mathbf{p}_{z}
\end{array}
$$

## 'Types' of Bonds - Multiple bonds

What happens in $\mathrm{N}_{2}$ ?
Remember the bond in $\mathrm{N}_{2}$ is a triple bond...

Each N atom has 3 unpaired 2p electrons


How do the atomic orbitals bond?

$p_{x} \quad p_{x}$

Between $2 p_{x}$ and $2 p_{x}$ orbitals, the bonding is not on the bond axis

A different type of bond!
t-bond -nodal plane along bond axis
Bonding occurs above and below the bond axis

## Bonding of $\mathrm{N}_{2}$



## Bonding of $\mathrm{N}_{2}$



## Hybridization of Orbitals

What about methane $\left(\mathrm{CH}_{4}\right)$ ?
According to Valence Bond Theory:
C should only make 2 bonds!
But we know that $C$ can make 4 bonds


Carbon, [He] $2 s^{2} 2 p_{x}{ }^{1} 2 p_{y}{ }^{1}$

And $\mathrm{CH}_{4}$ has a tetrahedral shape according to VSEPR:
How do get a tetrahedral shape from the 2 s and 2 p orbitals?


## Hybridization of Orbitals

We solve the 4 bond problem by promoting an electron:


Carbon, [He] $2 s^{1} 2 p_{x}{ }^{1} 2 p_{y}{ }^{1} 2 p_{z}{ }^{1}$
It takes energy to promote an electron to a higher E orbital
But, overall the energy is lower if $C$ can make 4 bonds instead of 2.
Ok, what about the geometry problem?

## Hybridization of Orbitals

By promoting an electron, we can now make 4 bonds
But the geometry of the $p_{x}, p_{y^{\prime}}$ and $p_{z}$ orbitals don't match the tetrahedral shape of $\mathrm{CH}_{4}$
$s$ and $p$ orbitals are described by a wave-like model of the $e$
If we think of the orbitals as interfering with each other, we can define new hybrid orbitals:


$$
\begin{aligned}
& h_{1}=s+p_{x}+p_{y}+p_{z} \\
& h_{2}=s-p_{x}-p_{y}+p_{z} \\
& h_{3}=s-p_{x}+p_{y}-p_{z} \\
& h_{4}=s+p_{x}-p_{y}-p_{z}
\end{aligned}
$$



## sp ${ }^{3}$ Hybridization

C in $\mathrm{CH}_{4}$ uses 4 hybrid orbitals:


## $s p^{3}$ hybridized carbon

We took 1 s orbital and 3 p orbitals to make $4 \mathrm{sp}^{3}$ orbitals.
The sum of the atomic orbitals $=$ the sum of the hybrid orbitals
The $4 \mathrm{sp}^{3}$ orbitals point in the 4 directions of the tetrahedral bonds


## $s p^{3}$ Hybridization

Back to $\mathrm{CH}_{4}$ :
The $42 \mathrm{sp}^{3}$ hybrid orbitals on carbon make s-bonds with the 1 s orbitals of the 4 H atoms
$\mathrm{NH}_{3}$ also uses $\mathrm{sp}^{3}$ hybrid orbitals:


Ammonia, $\mathrm{NH}_{3}$
Whenever an atom has a tetrahedral structure, it is $\mathrm{sp}^{3}$ hybridized


## $s p^{3}$ Hybridization

Whenever an atom has a tetrahedral structure, we say it is $s p^{3}$ hybridized

This includes molecules with multiple central atoms:


## Atomic Orbitals $\rightarrow$ Hybrid Atomic Orbitals

$\mathrm{sp}^{3}$ bonding works for tetrahedral-shaped molecules
What about the other VSEPR shapes?

Linear bonding can be described by sp hybridization:

$$
\begin{aligned}
& \mathrm{h}_{1}=\mathrm{s}+\mathrm{p} \\
& \mathrm{~h}_{2}=\mathrm{s}-\mathrm{p}
\end{aligned}
$$

2 of the p orbitals remain as they were
Unused p orbitals are available for $p$ bonding

## Other Types of Hybridization

$\mathrm{sp}^{3}$ bonding works for tetrahedrally shaped molecules
What about the other VSEPR shapes?
Trigonal planar can be described by $\boldsymbol{s p}^{2}$ hybridization:

$$
\begin{gathered}
\mathrm{h}_{1}=\mathrm{s}+\sqrt{2} \mathrm{p}_{\mathrm{y}} \\
\mathrm{~h}_{2}=\mathrm{s}+\sqrt{\frac{3}{2}} \mathrm{p}_{\mathrm{x}}-\sqrt{\frac{1}{2}} \mathrm{p}_{\mathrm{y}} \\
\mathrm{~h}_{3}=\mathrm{s}-\sqrt{\frac{3}{2}} \mathrm{p}_{\mathrm{x}}-\sqrt{\frac{1}{2}} \mathrm{p}_{\mathrm{y}}
\end{gathered}
$$



The $p_{z}$ orbital is not used and remains as it was
$s p^{2}$

## Hybrid Orbitals



Overlapping sp hybrid orbitals

Simplified view of hybrid orbitals


## Hybrid Orbital Shapes



Look Familiar?

$\rightarrow+\infty$



## Other Types of Hybridization

N atomic orbitals always produce N hybrid orbitals
TABLE 3.2 Hybridization and Molecular Shape*

| Electron <br> arrangement | Number of <br> atomic orbitals | Hybridization of <br> the central atom | Number of <br> hybrid orbitals |
| :--- | :---: | :---: | :---: |
| linear | 2 | $s p$ | 2 |
| trigonal planar | 3 | $s p^{2}$ | 3 |
| tetrahedral | 4 | $s p^{3}$ | 4 |
| trigonal bipyramidal | 5 | $s p^{3} d$ | 5 |
| octahedral | 6 | $s p^{3} d^{2}$ | 6 |

Spectroscopic data suggests terminal atoms use hybrid orbitals as well
A terminal Cl uses $s p^{3}$ hybridization in the arrangement of its lone pairs?

## Examples

$\square$

## $1 s \downarrow^{1 s}$ $\mathrm{H}_{2}$



## m Bond

## Ethylene

## Ethylene



## Characteristics of Multiple Bonds

double bond $=1 \sigma$-bond $+1 \pi$-bond
triple bond $=1 \sigma$-bond $+2 \pi$-bonds
$\sigma$-bonds result from head-on overlap of orbitals $\pi$-bonds results from side-by-side overlap

Atoms in a single bond can rotate freely, whereas atoms in a double bond are much less likely to:


## Characteristics of Multiple Bonds

The shape of ethylene $\left(\mathrm{C}_{2} \mathrm{H}_{4}\right)$
Experimental evidence:
All six atoms lie in the same plane with $120^{\circ}$ bond angles
Suggests trigonal planar structure $\rightarrow \mathrm{sp}^{2}$ hybridization

$$
\text { Each C: } s p^{2}+p
$$

s-bonds with the sp ${ }^{2}$ orbitals

$$
\begin{gathered}
\mathrm{C}-\mathrm{C} \\
4 \mathrm{C}-\mathrm{H}
\end{gathered}
$$

p-bond with the leftover p orbital

$$
\mathrm{C}-\mathrm{C}
$$

## Characteristics of Multiple Bonds

The shape of acetylene $\left(\mathrm{C}_{2} \mathrm{H}_{2}\right)$ is linear $\rightarrow$ sp hybridization


## Hybridization and Benzene



## Molecular Orbital Theory

- Lewis Theory
- Connectivity, electron tracking
$\square$ VSEPR Theory
- 3-D Structure around an atom
$\square$ Valence Bond Theory
- Extended 3-D Structure Information
- Delocalization in Molecules (LIMITED)
- Illustrates Multiple Bonding
- Prediction of Reactivity
- Molecular Orbital Theory
- Orbitals as a function of the whole molecule
- Delocalization much more thorough
- Anti-bonding and Non-bonding electrons
- Behavior in a magnetic field
- Spectral Data


## Paramagnetic / Diamagnetic



## Constructive and Destructive Interference



## Molecular Orbitals

MO Theory - electrons occupy molecular orbitals spread over entire molecule

- All valence electrons are delocalized
- Molecular orbitals are built by adding together atomic orbitals
- Linear combination of atomic orbitals (LCAO)
$\mathrm{H}-\mathrm{H}$ bond in $\mathrm{H}_{2}$ with two 1s electrons:

$$
y=y_{A 1 s}+y_{B 1 s}
$$



- LCAO-MO shows constructive interference
- $\mathrm{E}_{\text {LCAO-mo }}<\mathrm{E}_{\mathrm{Ao}}$
- LCAO-MO are bigger than AO


## Molecular Orbitals

"Conservation of orbitals"
$\mathrm{H}_{2}$ bonds with two 1s orbitals - it must have 2 LCAO-MOs
$2^{\text {nd }}$ LCAO-MO - destructive interference, higher PE than atomic orbitals

Nodal plane! - much less e-density
"internuclearly"
This is an antibonding orbital

$1^{\text {st }}$ LCAO-MO is lower in energy than Atomic orbitals

This is called a bonding orbital

$$
y=y_{A 1 s}+y_{B 1 s}
$$

## Sigma Bonds



## Molecular Orbital Energy Diagram



## Electron Configurations of Diatomic Molecules

Shows ALL valence electrons using the Building-Up Principle:


Electron configuration of $\mathrm{H}_{2} \rightarrow \mathrm{~s}_{1 \mathrm{~s}}{ }^{2}$
Based on MO theory - a single electron can hold a bond together

## Bond Order



## $2^{\text {nd }}$ row Diatomic Molecules

Must include both $2 s$ and $2 p$ orbitals in making our MOs
Only orbitals that are close in energy will form MOs
We have $2(2 s)$ orbitals and $6(2 p)$ orbitals $\rightarrow 8$ atomic orbitals
2 s orbitals overlap in the same way as the 1 s orbitals of $\mathrm{H}_{2}$
$2 p$ orbitals overlap in same orientations as in Valence bond theory:


$\pi_{2 p}{ }^{*}$

$\pi_{2 p}$
$\sigma_{2 p}$

## $2^{\text {nd }}$ row Diatomic Molecules

We have $2(2 s)$ orbitals and $6(2 p)$ orbitals $\rightarrow 8$ atomic orbitals
$2 s$ orbitals overlap in the same way as the 1 s orbitals of $\mathrm{H}_{2}$
$2 p$ orbitals overlap in same orientations as in Valence bond theory:


## Pi Bonds



## $2^{\text {nd }}$ row Homonuclear Diatomic Molecules



## $2^{\text {nd }}$ row Homonuclear Diatomic Molecules


$\square$ Homonuclear Diatomic MO Energy diagram for all elements right of (and including) oxygen
$\mathrm{O}_{2}$
$F_{2}$
' $\mathrm{Ne}_{2}{ }^{\prime}$

And their combinations

## $2^{\text {nd }}$ row Homonuclear Diatomic Molecules

$\square$ Homonuclear Diatomic MO Energy diagram for all elements left of oxygen
$\begin{array}{llllll}\mathrm{Li}_{2} & \mathrm{Be}_{2} & \mathrm{~B}_{2} & \mathrm{C}_{2} & \mathrm{~N}_{2}\end{array}$

## Relative MO Energy Levels



## MO Diagram for $\mathrm{Li}_{2}$



## MO Diagram for $\mathrm{Be}_{2}$

Be
$\mathrm{Be}_{2}$
Be

$$
\begin{aligned}
& \text { 1s 倝 }
\end{aligned}
$$

## MO Diagram for $\mathrm{N}_{2}$



## MO Diagram for $\mathrm{O}_{2}$



## MO Diagram for $\mathrm{F}_{2}$




## Relative AO Energy Levels

$$
\begin{aligned}
& \underset{-2 s_{---} 2 p}{\text { H }} \text { Li } \quad \text { Be } \quad \text { B } \quad \text { C } \quad \text { N } \quad \text { O } \quad \text { F } \\
& -1 s-_{-}^{2} s_{---2 p}^{2 p}
\end{aligned}
$$

$$
\begin{aligned}
& -2 s \\
& -1 s \quad \quad-2 s \\
& \text { _ } 2 s \\
& \text { _ } 1 s \\
& \text { _ } 1 s \\
& \text { _ } 1 s
\end{aligned}
$$

## Heteronuclear Diatomic Molecules



Account for $\triangle \mathrm{EN}$ between AO's

More electronegative atoms will have lower energy atomic orbitals

Contributions are dependent on energy level

## MO diagram for HF



## Orbitals in Polyatomic Molecules

An electron pair in a bonding orbital helps to bind together the whole molecule - can become very complex

Occasionally some MOs are neither bonding nor antibonding -these are called nonbonding orbitals


## MO Diagram for Carbon Monoxide



## Organic Molecules

Chemists commonly mix VB and MO theories when discussing organic molecules:
VB theory is good for talking about $\sigma$ bonds MO theory is better for considering $\pi$ bonds

Consider Benzene ( $\mathrm{C}_{6} \mathrm{H}_{6}$ ):
VB perspective - each C is $\mathrm{sp}^{2}$ hybridized


## $\pi$ Orbitals of Benzene

## 84



## Coordination Compounds



FIGURE 16.17 When potassium cyanide is added to a solution of iron(II) sulfate, the cyanide ions replace the $\mathrm{H}_{2} \mathrm{O}$ ligands of the $\left[\mathrm{Fe}\left(\mathrm{OH}_{2}\right)_{6}\right]^{2+}$ complex (left) and produce a new complex, the hexacyanoferrate(II) ion, $\left[\mathrm{Fe}(\mathrm{CN})_{6}\right]^{4-}$ (right). The blue color is due to the polymeric compound called Prussian blue, which forms from the cyanoferrate ion.

## Colors of Coordination Compounds

Aqueous solutions of $\left[\mathrm{Fe}(\mathrm{SCN})\left(\mathrm{OH}_{2}\right)_{5}\right]^{2+}$, $\left[\mathrm{Co}(\mathrm{SCN})_{4}\left(\mathrm{OH}_{2}\right)_{2}\right]^{2-},\left[\mathrm{Cu}\left(\mathrm{NH}_{3}\right)_{4}\left(\mathrm{OH}_{2}\right)_{2}\right]^{2+}$, and $\left[\mathrm{CuBr}_{4}\right]^{2-}$


## Biological Importance



## Coordination Compounds

$\square$ Metals in the d-block form coordination compounds
$\square$ Coordination compound electrically neutral compound in which at least on of the ions present is a complex
$\square$ Ligands - ions or molecules that are attached to the central metal atom or ion

$\square$ Coordination number - number of ligands attached to the central metal

## Shapes of Complexes



3 A square-planar complex
2 A tetrahedral complex
1 An octahedral complex

## Shapes of Complexes



4 Dimethylmercury (0)


## Nomenclature

$\square\left[\mathrm{FeCl}\left(\mathrm{OH}_{2}\right)_{5}\right]^{+}$pentaqquachloridoiron(II)
$\square$ Overall complex charge $=+1$
$\square$ Ligand charge $=\mathrm{Cl}^{-1}$ and $\mathrm{OH}_{2}$ (water)
$\square$ Metal charge + ligand charge $=$ overall complex charge

- $\mathrm{Fe}+(-1)=+1$
- $\mathrm{Fe}=+2$
$\square\left[\mathrm{Co}\left(\mathrm{NH}_{3}\right)_{3}\left(\mathrm{OH}_{2}\right)_{3}\right]_{2}\left(\mathrm{SO}_{4}\right)_{3}$ triamminetriaquacobalt(III) sulfate
$\square$ Overall complex charge $=+3$
$\square$ Ligand charge $=0$
- $\mathrm{Co}=+3$


## Common Ligands

| Formula* | Name |
| :--- | :--- |
| Neutral ligands |  |
| $\mathrm{OH}_{2}$ | aqua |
| $\mathrm{NH}_{3}$ | ammine |
| NO | nitrosyl |
| CO | carbonyl |
| $\mathrm{NH}_{2} \mathrm{CH}_{2} \mathrm{CH}_{2} \mathrm{NH}_{2}$ | ethylenediamine (en) |
| $\mathrm{NH}_{2} \mathrm{CH}_{2} \mathrm{CH}_{2} \mathrm{NHCH}_{2} \mathrm{CH}_{2} \mathrm{NH}_{2}$ | diethylenetriamine (dien) |

## Common Ligands

```
Anionic ligands
F-
Cl-
Br
I-
OH-
O-
CN-
CN+
NCS
NCS
NO}\mp@subsup{}{2}{-}\mathrm{ as O-NO-
NO
CO}\mp@subsup{3}{}{2-}\mathrm{ as O-CO}\mp@subsup{2}{}{2-
C2O}\mp@subsup{\textrm{O}}{4}{}\mp@subsup{}{}{2-}\mathrm{ as }\mp@subsup{}{}{-}\mp@subsup{\textrm{O}}{2}{}\mp@subsup{\textrm{CCO}}{2}{-
```



```
\(\mathrm{SO}_{4}{ }^{2-}\) as \(\mathrm{OSO}_{3}{ }^{2-}\)
```

fluorido
chlorido
bromido
iodido
hydroxido
oxido
cyanido-кС
isocyanido, cyanido-к $N$
thiocyanato-к $N$
isothiocyanato, thiocyanato-кS
nitrito-к $O$
nitro, nitrito-к $N$
carbonato-к $O$
oxalato (ox) ${ }^{\dagger}$
ethylenediaminetetraacetato (edta) ${ }^{\text {S }}$
sulfato

## Prefixes

$\square 1$ mono
$\square 2$ di (bis)
$\square 3$ tri (tris)
$\square 4$ tetra (tetrakis)
$\square 5$ penta (pentakis)
$\square 6$ hexa (hexakis)
$\square 7$ hepta
$\square 8$ octa

Prefixes in parentheses are used when the ligand already contains a Greek prefix or is polydentate

Example: ethylenediamine oxalato (bidentate)

## More Nomenclature

$\square \mathrm{Na}_{2}\left[\mathrm{PtCl}_{2}(\mathrm{ox})_{2}\right]$ sodium dichloridobis(oxalato)platinate(IV)
$\square$ Overall complex charge $=-2$
$\square$ Ligand charge $=\mathrm{Cl}^{-1}$ and oxalato $\mathrm{C}_{2} \mathrm{O}_{4}{ }^{2-}$
$\square$ Metal charge + ligand charge $=$ overall complex charge

- $\mathrm{Pt}+(-6)=-2$
- $\mathrm{Pt}=+4$
$\square\left[\mathrm{CoBr}\left(\mathrm{NH}_{3}\right)_{5}\right] \mathrm{SO}_{4}$ pentaamminebromidocobalt(III) sulfate $\square$ Overall complex charge $=-2$
$\square$ Ligand charge $=\mathrm{Br}^{-1}$ and $\mathrm{NH}_{3}$
$\square$ Metal charge + ligand charge $=$ overall complex charge
- Co $+(-1)=+2$
- $\mathrm{Co}=+3$


## Polydentate Ligands

$\square$ Some ligands attach to the metal more than once


8 Ethylenediamine, $\mathrm{NH}_{2} \mathrm{CH}_{2} \mathrm{CH}_{2} \mathrm{NH}_{2}$

Bidentate ligand

## Polydentate Ligands

$\square$ Some ligands attach to the metal more than once


10 Ethylenediaminetetraacetic acid Hexadentate ligand


11 An edta complex

## Coordination Isomers


(a) trans- $\left[\mathrm{CoCl}_{2}\left(\mathrm{NH}_{3}\right)_{4}\right]^{+}$
(b) cis- $\left[\mathrm{CoCl}_{2}\left(\mathrm{NH}_{3}\right)_{4}\right]^{+}$

## Coordination Isomers


(a) trans- $\left[\mathrm{CoCl}_{2}(\mathrm{en})_{2}\right]^{+}$
(b) $c i s-\left[\mathrm{CoCl}_{2}(\mathrm{en})_{2}\right]^{+}$
(c) cis-[ $\left.\mathrm{CoCl}_{2}(\mathrm{en})_{2}\right]^{+}$

