# Molecular Orbitals in Inorganic Chemistry

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### **Octahedral Complexes**

- very large and important class of transition metal complexes
- **⊚** TM complexes act as catalysts

to control and improve catalysts we need to understand the underlying electronic structure

magnetic properties, colour!

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Image from p16 of Metal Ligand Bonding by R. Janes and E. Moore RSC, Cambridge, 2004.

#### **Outline**

- octahedral complexes
- **⊚** forming the MO diagram for O<sub>h</sub>
- **©** colour, selection rules
- $\bigcirc$   $\Delta$ <sub>oct</sub>, spectrochemical series

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#### **MO** checklist

**Steps to construct a MO diagram** 

determine the molecular shape and identify the point group of the molecule

define the axial system and find all of the symmetry operations on the molecule

identify the chemical fragments, and put them along the bottom of the diagram

determine the energy levels and symmetry labels of the fragment orbitals (use H1s as a reference)

combine fragment orbitals of the same symmetry, estimate the splitting energy and draw in the MO energy levels and MOs (in pencil!)

determine the number of electrons in each fragment and hence the central MO region, add them to the diagram

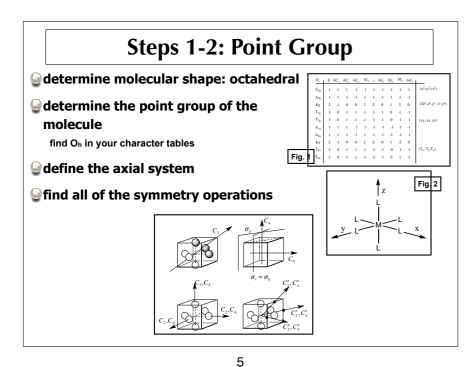
identify if any MO mixing occurs, determine the mixed orbitals and redraw the MO diagram with shifted energy levels and the mixed MOs

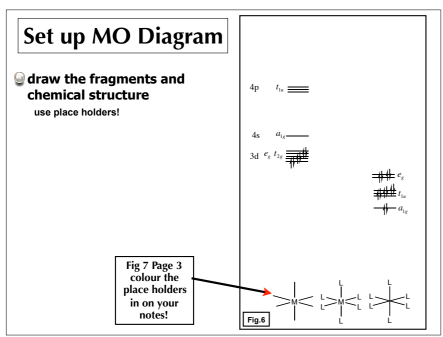
annotate your diagram

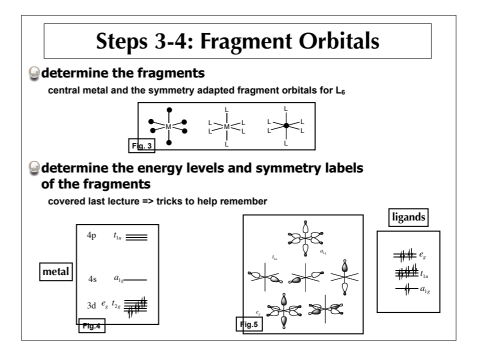
use the MO diagram to understand the structure, bonding and chemistry of the molecule

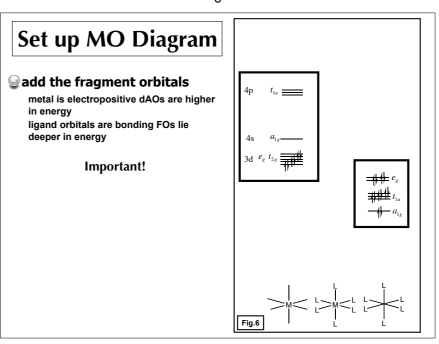
work through for an octahedral molecule

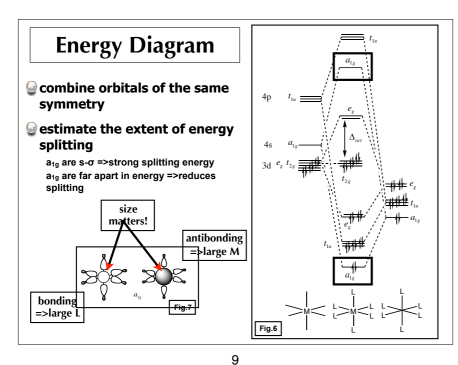
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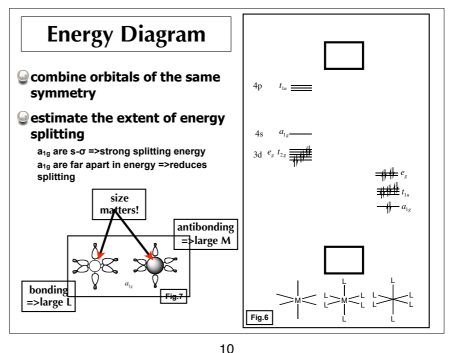


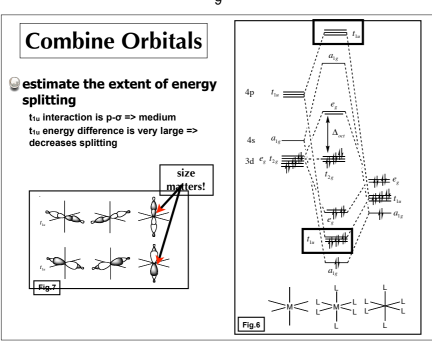


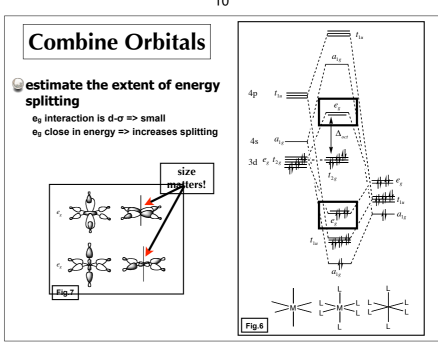








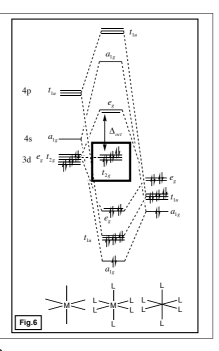




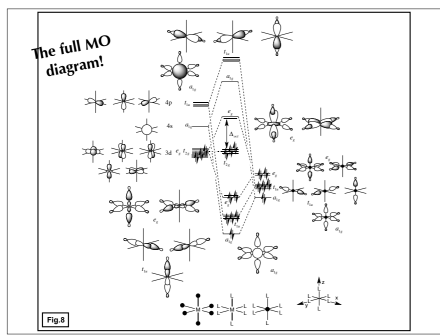
### **Combine Orbitals**

estimate the extent of energy splitting

metal  $t_{2g}$  remain non-bonding as there are no ligand orbitals of this symmetry



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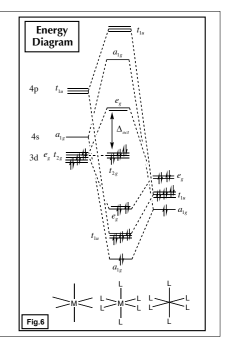
**Steps: 6-8** 

electronic configuration

six 2e donor ligands =12e number d electrons depends on the metal and oxidation state I've used six here

edetermine if there is mixing no mixing occurs

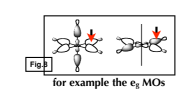
**■** use MO diagram checklist!



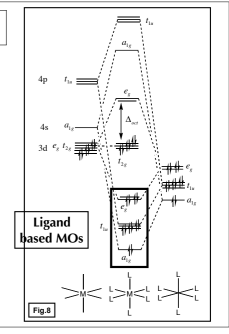
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three lowest energy sets of orbitals are ligand based

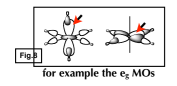


Important!

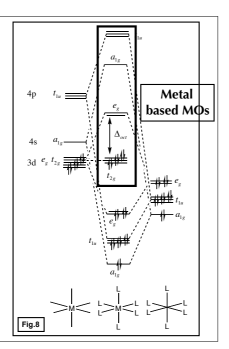




- three lowest energy sets of orbitals are ligand based
- higher energy MOs are metal based



Important!



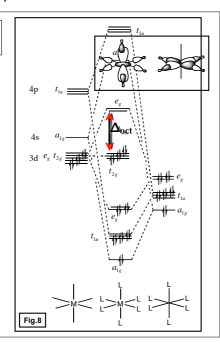
### $\Delta_{oct}$ parameter

- $\bigcirc$   $\sigma$ -ligands

t<sub>2g</sub> is non-bonding eg is antibonding!

Δ<sub>oct</sub> depends on strength of interaction:

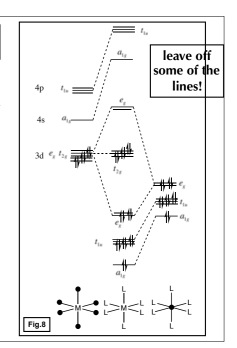
 $\Delta\epsilon$  energy between M and L FOs  $S_{ij}$  strength/density of overlap of FOs  $H_{ij}$  which is hard to estimate! total charge on complex oxidation state of the M (charge on M)



#### Note!

- leave off some of the "lines" for the metal dominated (s & p FOs) and the ligand dominated (low energy) FOs
- keep all the lines for the dAOs!

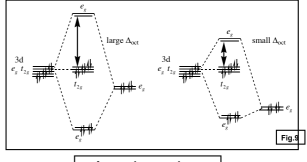
Important!



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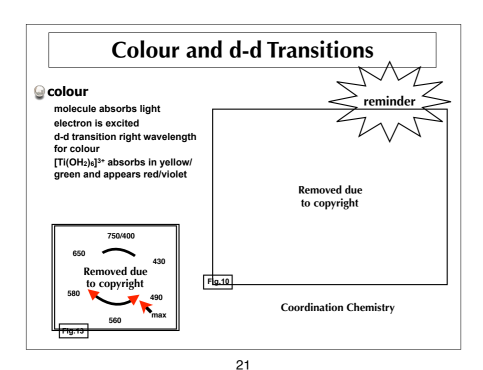
### $\Delta_{oct}$ parameter

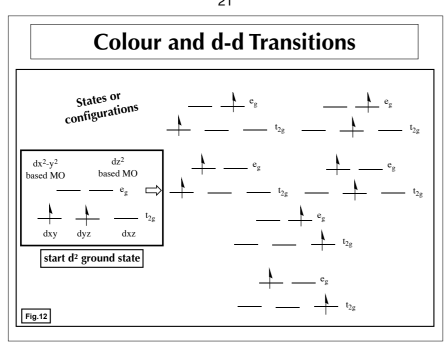
**y** higher in energy the ligand FOs the better the interaction

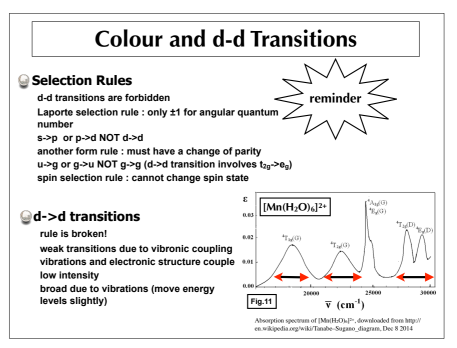


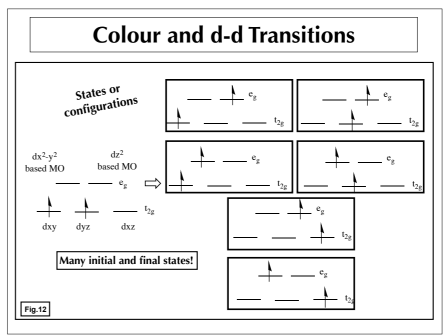
=> larger interaction => higher e<sub>g</sub> orbital => a larger Δ<sub>oct</sub>

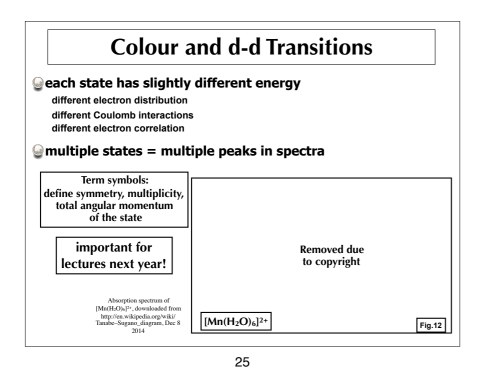
Important!











Spectrochemical Series

© replace NiL<sub>6</sub> ligands from stronger field to weaker field change Δ<sub>oct</sub> and the colour

 $[Ni(H_2O)_6]^{2+}$  has low energy, small  $\Delta_{\text{oct}}$  absorbs in the red and is green

example!

Fig.13

Colour of various Ni(II) complexes in aqueous solution.

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[Ni(NH<sub>3</sub>)<sub>6</sub>]<sup>2+</sup>, [Ni(en)<sub>3</sub>]<sup>2+</sup>, [NiCl<sub>4</sub>]<sup>2-</sup>, [Ni(H<sub>2</sub>O)<sub>6</sub>]<sup>2+</sup>

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Spectrochemical series

I- <Br- <Cl- <F-<OH- <O<sup>2</sup>- <H<sub>2</sub>O <NH<sub>3</sub> < en < NO<sub>2</sub>- <CH<sub>3</sub>-<C<sub>6</sub>H<sub>5</sub>- <CN- <CO

<C6Π5- <CN- <CC

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Image from http://chemwiki.ucdavis.edu/ Inorganic\_Chemistry/Crystal\_Field\_Theory/ Colors\_of\_Coordination\_Complexes, downloaded 1 Dec 2014

Image from http://en.wikipedia.org/wiki/ Nickel(II)\_chloride, LHcheM, downloaded 1 Dec 2014

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Info-graphics from http://www.compoundchem.com/

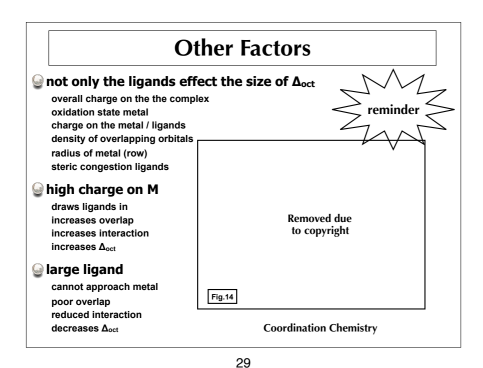
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CN- < CO < NO+ strong field

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In-Class Activity

determine new symmetry labels for metal AOs

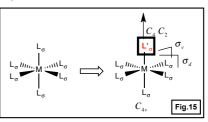
determine the new symmetry labels for the ligand FOs

VERY Important!

use short-cuts and your character table for  $C_{4v}$   $C_{4v}$ 

### **Reduce the Symmetry**

- **⊚** loss of symmetry O<sub>h</sub> -> C<sub>4v</sub>



**process** simple:

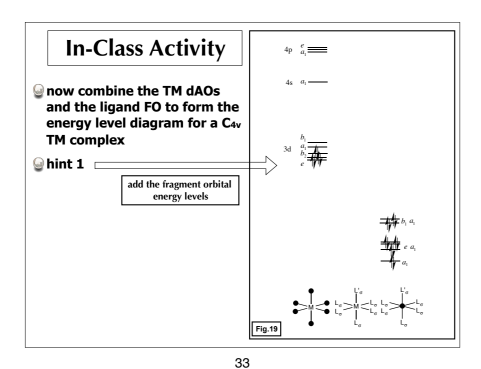
MOs remain the same just need to work out reduced symmetry labels! watch for degeneracy breaking

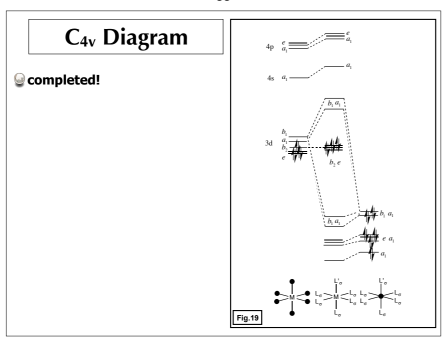
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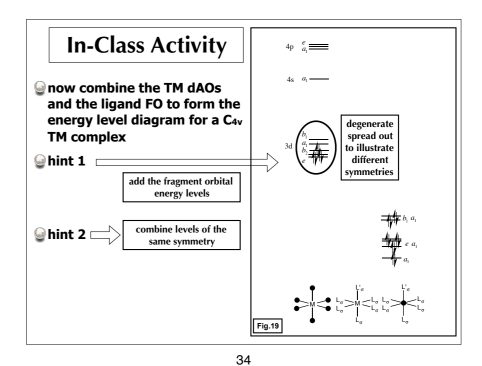
#### **In-Class Activity**

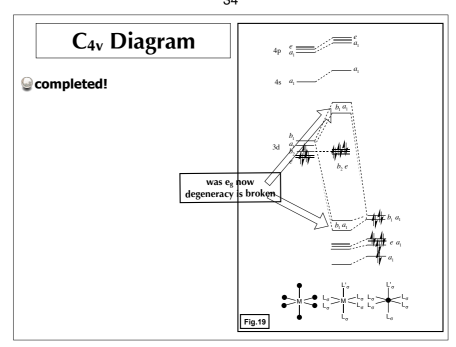
**⊚** now combine the TM dAOs and the ligand FO to form the energy level diagram for a C<sub>4v</sub> TM complex

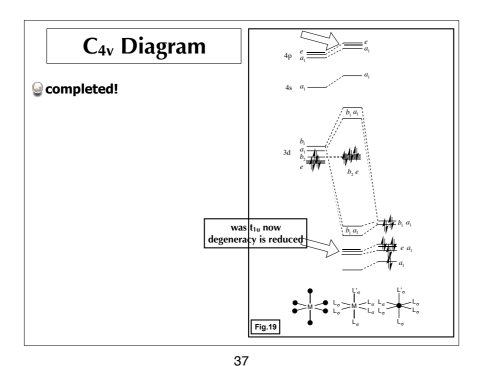
Fig.18, page 10 of your notes





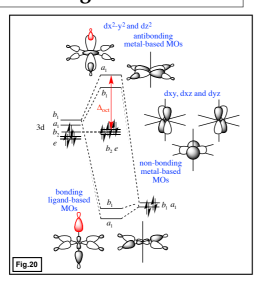






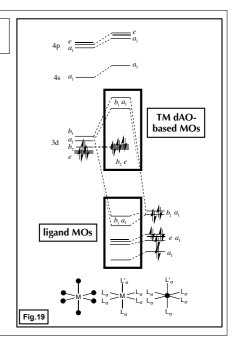
C<sub>4v</sub> TM Diagram

- primary effect of the reduction in symmetry => split the e<sub>g</sub> energy levels
- ordering of a<sub>1</sub> and b<sub>1</sub> will depend on the ligands
- ω Δ<sub>oct</sub> is splitting of dAO manifold (group of orbitals!)



C<sub>4v</sub> Diagram

**⊚** completed!



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#### **Key Points**

- $\ensuremath{\mbox{\ensuremath{\mbox{\sc be}}}}$  be able to draw energy level diagrams for octahedral TM complexes with  $L_\sigma$
- be able to draw and describe the important MOs
- be able to discuss key features of the diagrams, especially features relating to the character of the MOs
- be able to draw the energy diagram for a lower symmetry TM complex with σ-bonding ligands, including square planar

## **Finally**

#### http://www.huntresearchgroup.org.uk/

#### See my web-site

notes AND slides
link to panopto when it becomes available
optional background support for beginners
optional material to take you a little further
links to interesting people and web-sites
links to relevant research papers on MOs
model answers!!

