

# Molecular Orbitals in Inorganic Chemistry

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Rm 110F (MSRH)

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## Outline

- octahedral complexes
- forming the MO diagram for  $O_h$
- colour, selection rules
- $\Delta_{oct}$ , spectrochemical series
- forming the MO diagram for  $C_{4v}$

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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. J. E. and E.  
Moore RSC, Cambridge, 2004.

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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

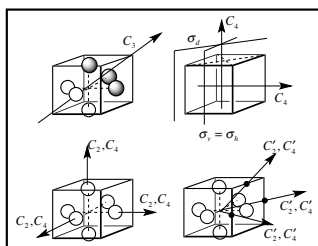
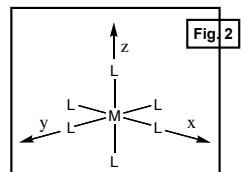
work through for  
an octahedral  
molecule

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**Steps 1-2: Point Group**

-  **determine molecular shape: octahedral**
  -  **determine the point group of the molecule**
    - find  $O_h$  in your character tables
  -  **define the axial system**
  -  **find all of the symmetry operations**
- Fig

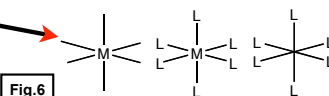
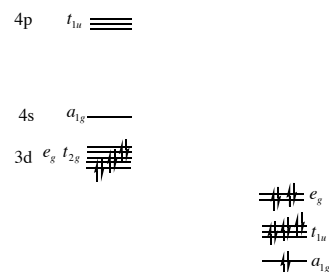
$O_i$	E	8C <sub>3</sub>	6C <sub>2</sub>	3C <sub>2</sub>	i	6S <sub>6</sub>	8C <sub>6</sub>	6σ <sub>v</sub>	
A <sub>1g</sub>	1	1	1	1	1	1	1	1	(x <sup>2</sup> y <sup>2</sup> , y <sup>2</sup> z <sup>2</sup> )
A <sub>2g</sub>	1	1	-1	-1	1	-1	1	-1	
E <sub>g</sub>	2	-1	0	0	2	2	0	-1	(2x <sup>2</sup> -y <sup>2</sup> -z <sup>2</sup> , x <sup>2</sup> -y <sup>2</sup> )
T <sub>1g</sub>	3	0	-1	-1	-1	3	0	-1	
T <sub>2g</sub>	3	0	1	-1	-1	3	-1	0	(xy, xz, yz)
A <sub>1u</sub>	1	1	1	1	-1	-1	-1	-1	
A <sub>2u</sub>	1	1	-1	-1	-1	-1	-1	1	
E <sub>u</sub>	2	-1	0	0	-2	0	-1	1	
T <sub>1u</sub>	3	0	-1	-1	-1	-3	0	1	(T <sub>x</sub> , T <sub>y</sub> , T <sub>z</sub> )
T <sub>2u</sub>	3	0	1	-1	-1	-3	1	0	-1



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## Set up MO Diagram

-  **draw the fragments and chemical structure**  
use place holders!

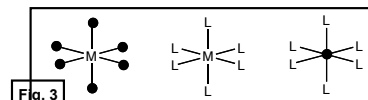


**Fig 7 Page 3**  
**colour the**  
**place holders**  
**in on your**  
**notes!**

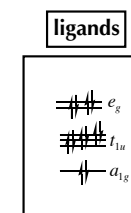
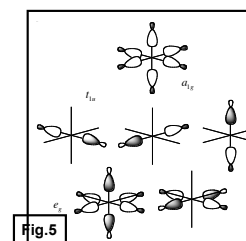
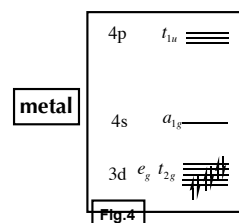
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### Steps 3-4: Fragment Orbitals

- determine the fragments**  
central metal and the symmetry adapted fragment orbitals for  $L_6$




- determine the energy levels and symmetry labels of the fragments**  
covered last lecture => tricks to help remember

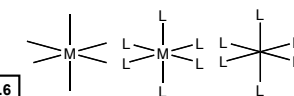
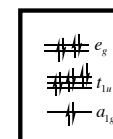
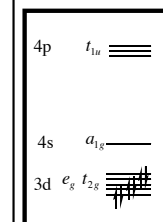


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## Set up MO Diagram

-  **add the fragment orbitals**
- metal is electropositive dAOs are higher in energy
  - ligand orbitals are bonding FOs lie deeper in energy

### Important!

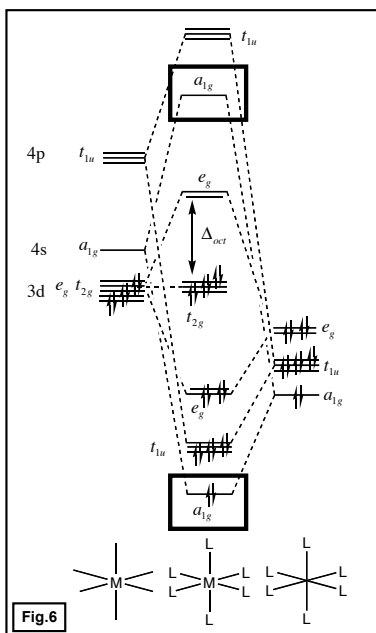
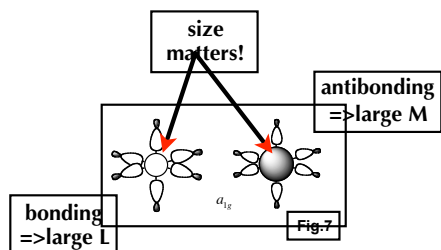


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## Energy Diagram

- combine orbitals of the same symmetry
- estimate the extent of energy splitting

$a_{1g}$  are s- $\sigma$  => strong splitting energy  
 $a_{1g}$  are far apart in energy => reduces splitting

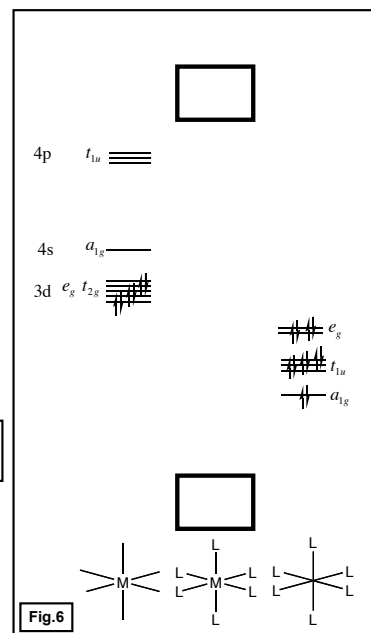
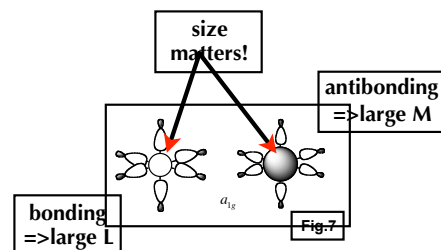


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## Energy Diagram

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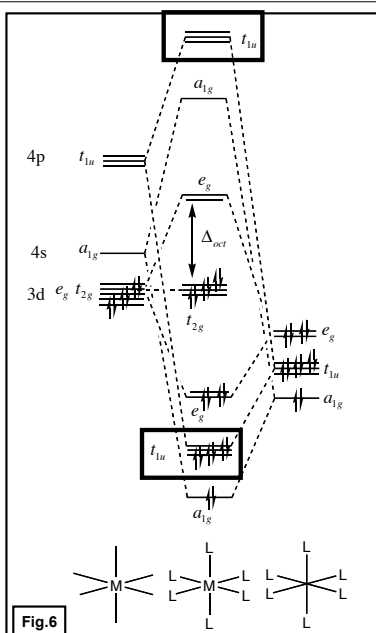
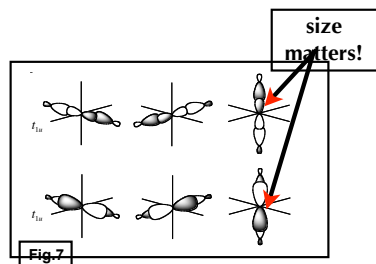


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## Combine Orbitals

- estimate the extent of energy splitting

$t_{1u}$  interaction is p- $\sigma$  => medium  
 $t_{1u}$  energy difference is very large => decreases splitting

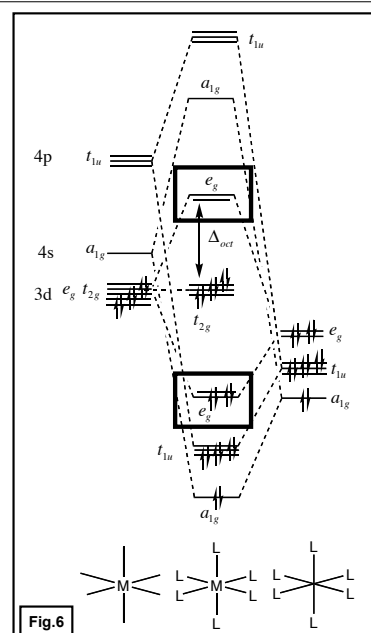
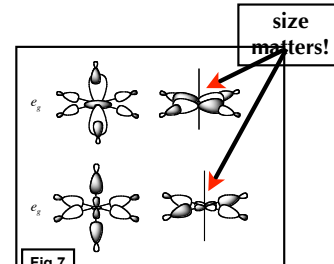


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## Combine Orbitals

- estimate the extent of energy splitting

$e_g$  interaction is d- $\sigma$  => small  
 $e_g$  close in energy => increases splitting

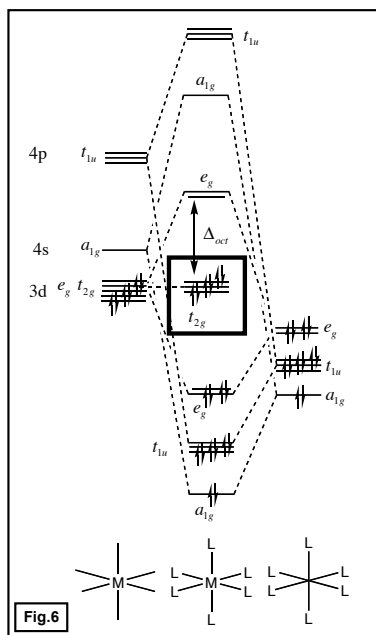


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## Combine Orbitals

### 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

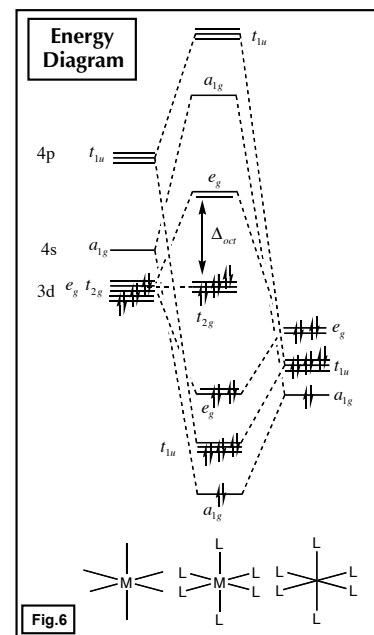
### electronic configuration

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

### determine if there is mixing

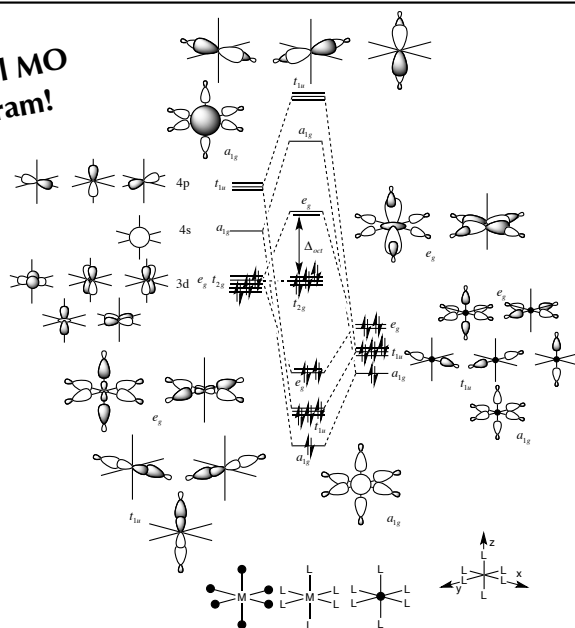
no mixing occurs

### use MO diagram checklist!



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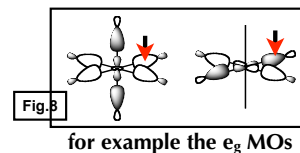
## The full MO diagram!



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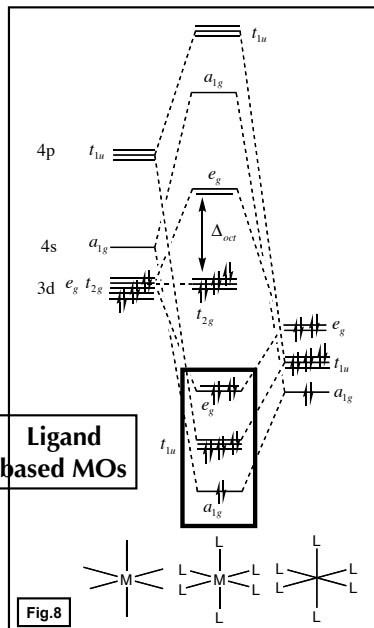
## Step 9: Analysis

### three lowest energy sets of orbitals are ligand based



for example the  $e_g$  MOs

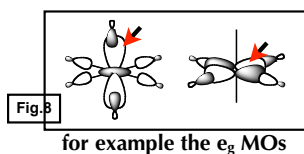
Important!



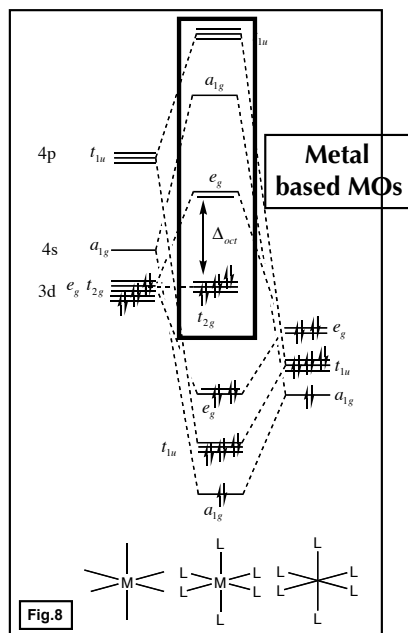
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## Step 9: Analysis

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



Important!

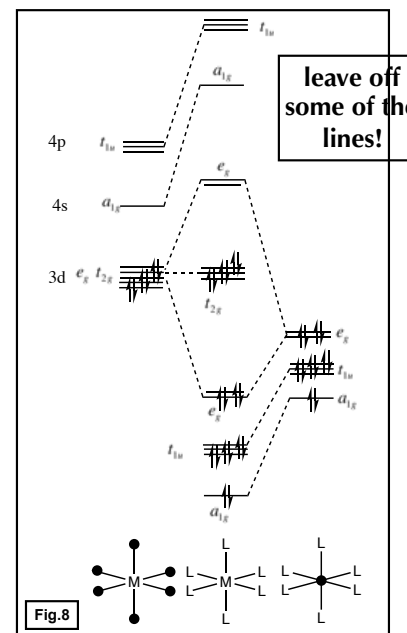


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## Note!

- 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

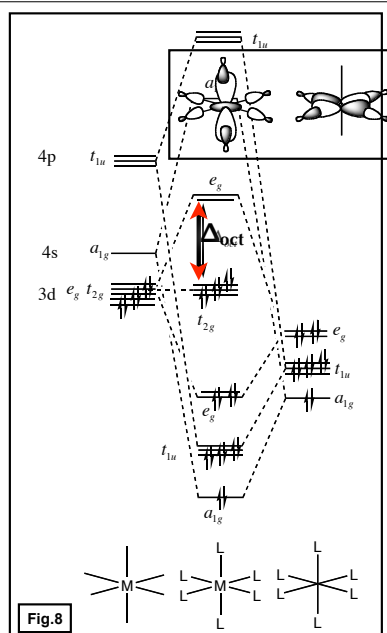
- $\Delta_{oct}$  energy span of the dAO dominated MOs

### $\sigma$ -ligands

$t_{2g}$  is non-bonding  
 $e_g$  is antibonding!

- $\Delta_{oct}$  depends on strength of interaction:

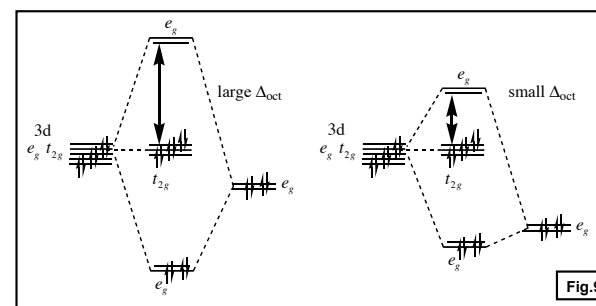
$\Delta \epsilon$  energy between M and L FOs  
 $S_{ij}$  strength/density of overlap of FOs  
 $H_{ij}$  which is hard to estimate!  
 total charge on complex  
 oxidation state of the M (charge on M)



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## $\Delta_{oct}$ parameter

- higher in energy the ligand FOs the better the interaction



=> larger interaction  
 => higher  $e_g$  orbital  
 => a larger  $\Delta_{oct}$

Important!

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## Colour and d-d Transitions

### colour

molecule absorbs light  
electron is excited  
d-d transition right wavelength  
for colour  
[Ti(OH<sub>2</sub>)<sub>6</sub>]<sup>3+</sup> absorbs in yellow/  
green and appears red/violet

reminder

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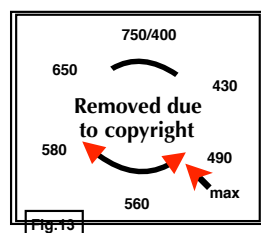


Fig.10

Coordination Chemistry

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## Colour and d-d Transitions

### Selection Rules

d-d transitions are forbidden  
Laporte selection rule : only  $\pm 1$  for angular quantum  
number  
s->p or p->d NOT d->d  
another form rule : must have a change of parity  
u->g or g->u NOT g->g (d->d transition involves  $t_{2g} \rightarrow e_g$ )  
spin selection rule : cannot change spin state

reminder

### d->d transitions

rule is broken!  
weak transitions due to vibronic coupling  
vibrations and electronic structure couple  
low intensity  
broad due to vibrations (move energy  
levels slightly)

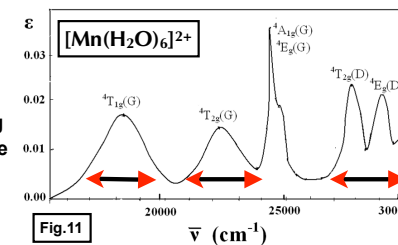


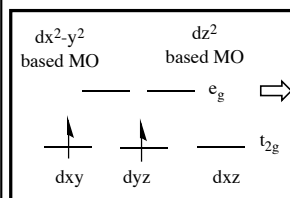
Fig.11

Absorption spectrum of [Mn(H<sub>2</sub>O)<sub>6</sub>]<sup>2+</sup>, downloaded from [http://en.wikipedia.org/wiki/Tanabe-Sugano\\_diagram](http://en.wikipedia.org/wiki/Tanabe-Sugano_diagram), Dec 8 2014

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## Colour and d-d Transitions

States or  
configurations



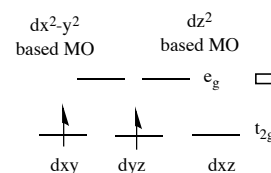
start d<sup>2</sup> ground state

Fig.12

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## Colour and d-d Transitions

States or  
configurations



Many initial and final states!

Fig.12

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## Colour and d-d Transitions

### each state has slightly different energy

different electron distribution  
different Coulomb interactions  
different electron correlation

### multiple states = multiple peaks in spectra

Term symbols:  
define symmetry, multiplicity,  
total angular momentum  
of the state

important for  
lectures next year!

Absorption spectrum of  
[Mn(H<sub>2</sub>O)<sub>6</sub>]<sup>2+</sup>, downloaded from  
[http://en.wikipedia.org/wiki/Tanabe-Sugano\\_diagram](http://en.wikipedia.org/wiki/Tanabe-Sugano_diagram), Dec 8  
2014

[Mn(H<sub>2</sub>O)<sub>6</sub>]<sup>2+</sup>

Fig.12

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## Spectrochemical Series

### uv-Vis experimentally measures $\Delta_{\text{oct}}$

### order ligands according to size $\Delta_{\text{oct}}$

### strong field ligand large $\Delta_{\text{oct}}$

for example CO

### weak field ligand small $\Delta_{\text{oct}}$

for example Cl<sup>-</sup>

Spectrochemical series

weak field

I<sup>-</sup> < Br<sup>-</sup> < Cl<sup>-</sup> < F<sup>-</sup> <

OH<sup>-</sup> < O<sup>2-</sup> < H<sub>2</sub>O <

py < NR<sub>3</sub> < NH<sub>3</sub> < en < NO<sub>2</sub><sup>-</sup>

< CH<sub>3</sub><sup>-</sup> < C<sub>6</sub>H<sub>5</sub><sup>-</sup> <

CN<sup>-</sup> < CO < NO<sup>+</sup>

strong field



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## Spectrochemical Series

### example!

### replace NiL<sub>6</sub> ligands from stronger field to weaker field change $\Delta_{\text{oct}}$ and the colour

[Ni(H<sub>2</sub>O)<sub>6</sub>]<sup>2+</sup> has low energy, small  $\Delta_{\text{oct}}$   
absorbs in the red and is green

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

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Fig.13

[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>

Spectrochemical series

I<sup>-</sup> < Br<sup>-</sup> < Cl<sup>-</sup> < F<sup>-</sup>

< OH<sup>-</sup> < O<sup>2-</sup> < H<sub>2</sub>O

< NH<sub>3</sub> < en < NO<sub>2</sub><sup>-</sup> < CH<sub>3</sub><sup>-</sup>

< C<sub>6</sub>H<sub>5</sub><sup>-</sup> < CN<sup>-</sup> < CO

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Fig.13

Image from [http://chemwiki.ucdavis.edu/Inorganic\\_Chemistry/Crystal\\_Field\\_Theory/Colors\\_of\\_Coordination\\_Complexes](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](http://en.wikipedia.org/wiki/Nickel(II)_chloride), LHCheM, downloaded 1  
Dec 2014

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## TM-Colour!

### Info-graphics from <http://www.compoundchem.com/>

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## Other Factors

### not only the ligands effect the size of $\Delta_{oct}$

overall charge on the the complex  
oxidation state metal  
charge on the metal / ligands  
density of overlapping orbitals  
radius of metal (row)  
steric congestion ligands

reminder

### high charge on M

draws ligands in  
increases overlap  
increases interaction  
increases  $\Delta_{oct}$

### large ligand

cannot approach metal  
poor overlap  
reduced interaction  
decreases  $\Delta_{oct}$

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Fig.14

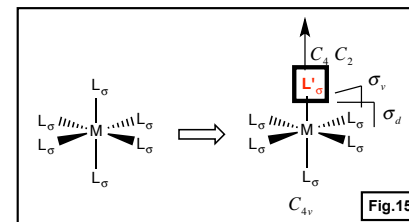
Coordination Chemistry

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## Reduce the Symmetry

### replace 1 ligand with a different $\sigma$ -ligand

### loss of symmetry $O_h \rightarrow C_{4v}$



### process simple:

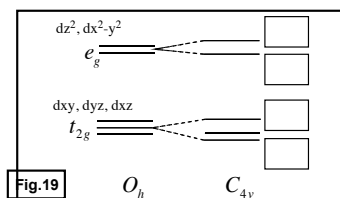
MOs remain the same  
just need to work out reduced symmetry labels!  
watch for degeneracy breaking

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## In-Class Activity

### determine new symmetry labels for metal AOs

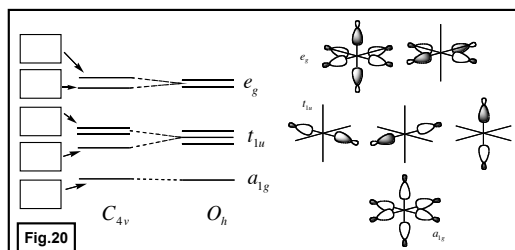
### determine the new symmetry labels for the ligand FOs



VERY Important!

degenerate levels remain degenerate, I've split them for ease of interpretation

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



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## In-Class Activity

### now combine the TM dAOs and the ligand FO to form the energy level diagram for a $C_{4v}$ TM complex

Fig.18, page 10 of your notes

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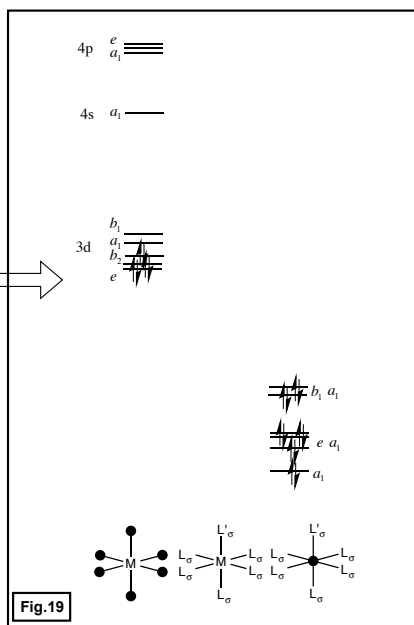


## In-Class Activity

now combine the TM dAOs and the ligand FO to form the energy level diagram for a  $C_{4v}$  TM complex

hint 1

add the fragment orbital energy levels



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## In-Class Activity

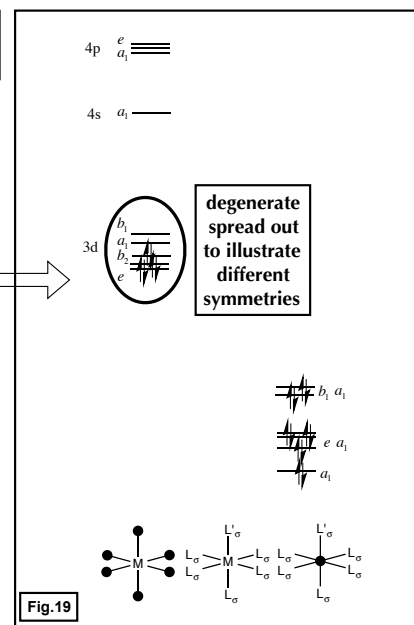
now combine the TM dAOs and the ligand FO to form the energy level diagram for a  $C_{4v}$  TM complex

hint 1

add the fragment orbital energy levels

hint 2

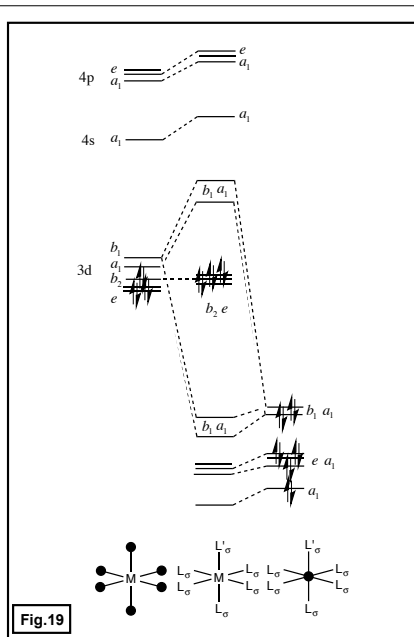
combine levels of the same symmetry



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## $C_{4v}$ Diagram

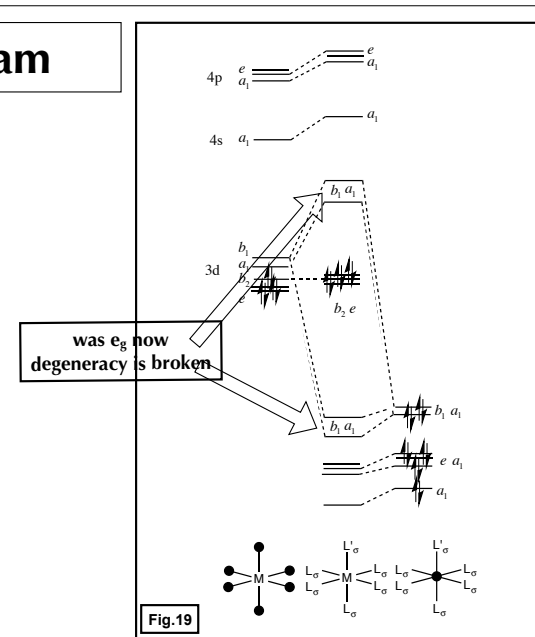
completed!



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## $C_{4v}$ Diagram

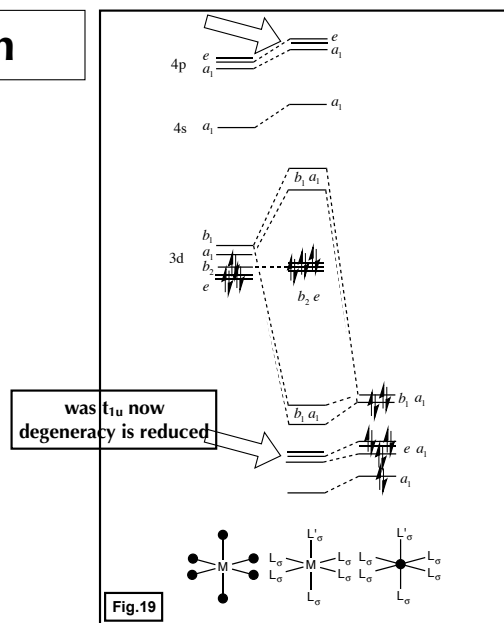
completed!



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

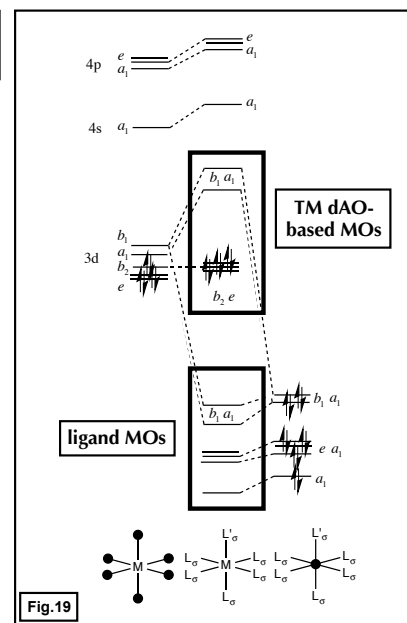
completed!



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

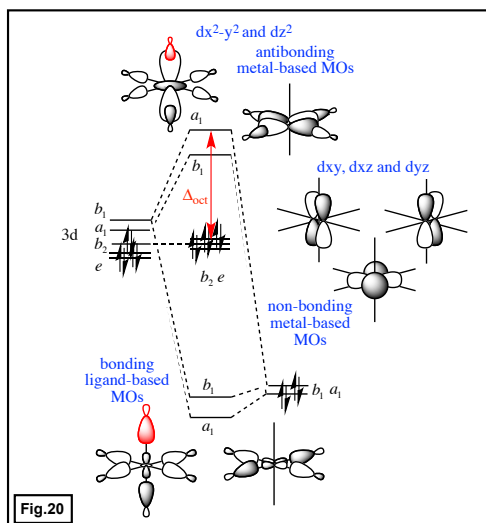
completed!



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

- primary effect of the reduction in symmetry => split the  $e_g$  energy levels
- ordering of  $a_1$  and  $b_1$  will depend on the ligands
- $\Delta_{oct}$  is splitting of dAO manifold (group of orbitals!)



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## Key Points

- 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 define the octahedral splitting parameter and be able to discuss key properties that impact on or effect  $\Delta_{oct}$
- be able to draw the energy diagram for a lower symmetry TM complex with  $\sigma$ -bonding ligands, including square planar

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# 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!!

