Free-Ion Terms to Ligand-field Terms

- Orbital term symbols for free atoms and ions are identical to symbols for irreducible representations in R_3 .
 - The irreducible representations of R_3 include all possible degeneracies.
 - There are no inherent symmetry restrictions on possible orbital degeneracies in R_3 .
- In octahedral and tetrahedral crystal fields (O_h and T_d) the highest dimension irreducible representations are three-fold degenerate.
 - For O_h and T_d complexes, free-ion terms with orbital degeneracies greater than three (*D*, *F*, *G*, ...) must split into new terms, each of which can have no higher than three-fold degeneracy.
 - In crystal fields of lesser symmetry (e.g., D_{4h} , D_3) free-ion orbital multiplicity terms with (2L + 1) > 2 must split as a result of the descent in symmetry from R_3 to the finite point group of the complex.
 - Ligand-field terms can have no higher orbital degeneracy than allowed by the highest dimension irreducible representation of the complex's point group.
 - In any crystal field all the term symbols, including those that are not split, are redefined and newly designated with the appropriate Mulliken symbols of their corresponding irreducible representations in the point group of the complex.

Why Terms Split in a Ligand-field

- Lifting the degeneracy among the *d* orbitals can destroy the equivalences among microstates that give rise to a particular free-ion term.
 - Orbital assignments that were energetically equivalent in the free ion may now be quite distinct in the environment of the complex.
 - These differences result in new collections of equivalent microstates, each of which gives rise to a distinct *ligand-field term*.
 - The total number of microstates for the configuration, as represented by D_t , remains the same.

Example: Splitting of d^1 Terms in an O_h Field

- The 10 microstates for the free-ion configuration d^1 give rise to a 2D term.
- In an octahedral field, the electron may have either the configuration t_{2g}^{1} or e_{g}^{1} :



d¹ O_h Ground State Term



- In the ground state, the electron can be in any of the three t_{2g} orbitals with either spin orientation $(m_s = \pm 1/2)$.
 - This makes six equivalent microstates.
 - There are three equivalent orbital assignments, so the overall orbital degeneracy (orbital multiplicity) is three.

$$(2L+1)=3$$

• There are only two overall spin orientations $(M_s = \pm \frac{1}{2})$, so the spin degeneracy (spin multiplicity) is two.

$$(2S + 1) = 2$$

The resulting term is ${}^{2}T_{2g}$, in which the Mulliken symbol for the orbital term is appropriately three-fold degenerate.

$d^{1}O_{h}$ Excited State Term

$$e_g$$
 1
 t_{2g} Excited State

- In the excited state configuration e_g^{-1} the electron can be in either of the two e_g orbitals with either spin orientation ($m_s = \pm \frac{1}{2}$).
 - This makes four equivalent microstates.
 - There are two equivalent orbital assignments, so the overall orbital degeneracy (orbital multiplicity) is two.

$$(2L+1)=2$$

• There are only two overall spin orientations $(M_s = \pm \frac{1}{2})$, so the spin degeneracy (spin multiplicity) is two.

$$(2S + 1) = 2$$

The associated term is ${}^{2}E_{g}$, in which the Mulliken symbol for the orbital term is two-fold degenerate.

Total Degeneracy, *D_t*, Remains Unchanged

• Note that the total degeneracy of each ligand-field term, equivalent to the number of microstates giving rise to it, is the product of its spin degeneracy times its orbital degeneracy.

$$D_t$$
 (term) = $(2L + 1)(2S + 1)$

• For ${}^{2}T_{2g}$ we have

$$D_t ({}^2T_{2g}) = (2)(3) = 6$$

• For ${}^{2}E_{g}$ we have

$$D_t(E_g) = (2)(2) = 4$$

The sum of total degeneracies of the ligand-field terms is equivalent to D_t for the free-ion configuration d^1 .

$$D_t(d^1) = D_t(^2T_{2g}) + D_t(^2E_g) = 6 + 4 = 10$$

Determining Ligand-Field Terms from Free-ion Terms

• The fate of any free-ion term in the point group of a complex can be determined by applying equations by which the characters for an irreducible representation in *R*₃ can be calculated:

$$\chi(E)=2j+1$$

$$\chi[C(\theta)] = \frac{\sin(j + 1/2)\theta}{\sin\theta/2}$$

$$\chi(i) = \pm(2j+1)$$

$$\chi[S(\theta)] = \pm \frac{\sin(j + 1/2)(\theta + \pi)}{\sin(\theta + \pi)/2}$$

$$\chi(\sigma) = \pm \sin(j + 1/2)\pi$$

Making Free-ion Terms the Basis for a Representation

- It is possible to apply these equations to both the spin and orbital terms (*S* and *L* states), but the field does not interact directly on the electron spin in a chemical environment such as a complex ion.
 - The new ligand-field terms will retain the original spin multiplicities of the free-ion terms from which they originate.
 - We only apply these equations to the *L* state of a free-ion term to determine the identities of the terms that result from splitting in the ligand field.
- In the last three equations with variable sign (±), the positive sign is used with *gerade* functions and the negative sign is used with *ungerade* functions.
 - We will be concerned solely with terms arising from configurations of *d* electrons, which are inherently *gerade*. Therefore we will choose the positive expression in all cases.
 - Nonetheless, in noncentrosymmetric point groups (e.g., T_d , D_{3h}) the resulting Mulliken symbol for the new state will not have a g subscript notation, which would be inappropriate in such groups.

{If terms arising from p or f configurations are to be considered, use the negative sign equations, because states arising from these are inherently *ungerade*.}

- *S* state, for which L = 0, is nondegenerate.
 - As with an *s* orbital, it has no angular dependence and no orientation in space.
 - Without using the equations, we conclude that *in any point group* an *S* term will not be split and will bear the Mulliken symbol for the totally symmetric representation.

$$\square In O_h, S \to A_{1g}$$

- *P* state, for which L = 1, is triply degenerate.
 - Substituting L = 1 into the equations for the operations of O_h gives the following representation.

$$O_h$$
 E
 $8C_3$
 $6C_2$
 $6C_4$
 $3C_2$
 i
 $6S_4$
 $8S_6$
 $3\sigma_h$
 $6\sigma_d$
 Γ_P
 3
 0
 -1
 1
 -1
 3
 1
 0
 -1
 -1

• Inspection of the character table shows that $\Gamma_P \equiv T_{1g}$.

$$\mathbb{R} In O_h, P \to T_{1g}$$

• A *P* term is not split, but becomes a triply degenerate T_{1g} term.

{Recall that the three-fold degenerate p <u>orbitals</u> transform as T_{1u} in O_h , but as we now see a P <u>state</u> transforms as T_{1g} . The transformations are different because the p orbitals are inherently *ungerade*, but the P state arising from a d configuration is inherently *gerade*.}

- **D** state, for which L = 2, is five-fold degenerate.
 - Substituting L = 2 into the equations for the operations of O_h gives the following representation.

O_h	E	8 <i>C</i> ₃	$6C_{2}$	$6C_4$	$3C_{2}$	i	$6S_4$	8 <i>S</i> ₆	$3\sigma_h$	$6\sigma_d$
Γ_{D}	5	-1	1	-1	1	5	-1	-1	1	1

 \circ This is identical to the reducible representation we obtained for *d* orbitals.

$$\mathbb{R} In O_h, D \to E_g + T_{2g}$$

• The five-fold degeneracy of the *D* free-ion term is lifted to become a doubly degenerate term and a triply degenerate term because of the restrictions on maximum degeneracy in O_h .

- *F* state, for which L = 3, is seven-fold degenerate.
 - Substituting L = 3 into the equations for the operations of O_h gives the following representation.

• This reduces as $\Gamma_F = A_{2g} + T_{1g} + T_{2g}$.

$$\square O_h, F \to A_{2g} + T_{1g} + T_{2g}$$

• The splitting of other states (*G*, *H*, *I*, etc.) can be determined in similar manner, giving the following results:

Free-ion	
Term	Terms in O_h
S	A_{1g}
Р	T_{1g}
D	$E_g + T_{2g}$
F	$A_{2g} + T_{1g} + T_{2g}$
G	$A_{1g} + E_g + T_{1g} + T_{2g}$
H	$E_{g} + 2T_{1g} + T_{2g}$
Ι	$A_{1g} + A_{2g} + E_g + T_{1g} + 2T_{2g}$

Example: Splitting of d^2 Free-Ion Terms in O_h

• The free-ion terms for the configuration *nd*², in order of increasing energy are

$${}^{3}F < {}^{1}D < {}^{3}P < {}^{1}G < {}^{1}S$$

• Each of these terms will split into the ligand filed terms we have just identified.

Free-ion terms	^{3}F	^{1}D	^{3}P	${}^{1}G$	^{1}S
Octahedral terms	${}^{3}A_{2g}$	${}^{1}E_{g}$	${}^{3}T_{1g}$	${}^{1}\!A_{1g}$	${}^{1}A_{1g}$
	${}^{3}T_{1g}$	${}^{1}T_{2g}$		${}^{1}E_{g}$	
	${}^{3}T_{2g}$			${}^{1}T_{1g}$	
				${}^{1}T_{2g}$	
Microstates	21	5	9	9	1

$$D_t = 21 + 5 + 9 + 9 + 1 = 45$$

Ligand-Field Terms in Other Fields

- The splittings of free-ion terms and the Mulliken symbols for the ligand-field terms in other point groups can be obtained in similar manner by making them bases for representations in the appropriate point group.
 - It is usually more efficient to use the correlation tables
 - For example, inspection of the correlation table for O_h and T_d shows that the splittings are identical in both groups, except for the omission of the subscript g for the tetrahedral states.
 - Correlations with other groups (e.g., D_{4h} , D_3 , D_{2d}) are not as trivial, but are equally straightforward.

Correlation Diagrams for Ligand Field Splitting

- What is the energy order of the ligand field terms?
- How will the energies of the terms change with changing Δ_0 ?
- Group theory alone, cannot provide quantitative answers.
- It is possible to address the problem at least qualitatively with a *correlation diagram*, which shows how the energies of terms change as a function of the ligand field strength, measured as Δ_o.
- To construct the correlation diagram, we look at two extremes:
 - Left side: A *weak field*, just strong enough to lift the R_3 free-ion term degeneracies.
 - On the left side of the diagram we show the energies of the free-ion terms and the Mulliken symbols for the terms into which they are split in a weak octahedral field.

• Right side: A hypothetical extremely strong field.

- At the limit of an extremely large Δ_0 separation between t_{2g} and e_g orbitals, we assume that interactions between electrons in separate orbitals are negligible.
- At this limit we can assess the energy order of the possible electronic configurations for the ground state and all excited states.
- We can then identify the terms that will emerge from each of these configurations in a slightly less strong field, where electronic interactions begin to be felt.



Layout of a Term Splitting Correlation Diagram

Constructing A Correlation Diagram for d^2

- The job of constructing the diagram amounts to determining the correlations between terms in the weak field and the terms in the strong field.
- General Approach The Method of Descending Symmetry (Bethe).
 - Rigorous, generally applicable, but tedious.
- The *Noncrossing Rule* is observed: Correlation lines for states of the same symmetry and same multiplicity do not cross, but rather repel one another, thereby increasing their relative energy separation beyond a certain minimum as field strength increases.