

Final projects

- I will be at your presentations next week!
- If you want to consult me about your slides, feel free to email them to me and I can give you comments over Thanksgiving



Final Exam

- ~50% new material
- Questions (not comprehensive):
- New material:
 - 1) Ligand substitution in coordination complexes
 - 2) Hard soft acid base theory
 - 3) 18 electron rule
 - 4) Analyzing molecular vibrations in organometallic complexes
 - 5) Predicting products/reactants in organometallic reactions
 - 6) Given a catalytic cycle, identify the type of reaction at each step, or predict the next intermediate given the reaction type, count electrons at each step, determine metal oxidation state

YOU WILL NOT BE TESTED ON REDOX REACTIONS, YOU WILL NOT HAVE TO MEMORIZE CATALYTIC CYCLES, JUST ANALYZE THEM



Final Exam

- ~50% new material
- Questions (not comprehensive):
- Old material:
 - 1) Identify point groups
 - 2) MO Theory/orbital interactions (you will not have to derive SALCs)
 - 3) Spectrochemical series, d orbital splitting
 - 4) Orbitals



ORGANOMETALLIC REACTIONS AND CATALYSIS

MFT Ch 14
CH 431



Organometallic complexes and catalysis

- Organometallic complexes, much like other inorganic complexes, can undergo a variety of different reactions
- Some organometallic complexes are used widely for industrial reactions as well as in common reactions performed in synthetic labs
- This lecture will cover the basic kinds of reactions organometallic complexes can undergo
- Lectures after this will then cover important complexes used in catalytic cycles



Two main categories of organometallic reactions

1. Reactions involving a gain or loss of ligands
2. Reactions involving modification of ligands



Reactions involving gain or loss of ligands

- Ligand dissociation
- Oxidative addition
- Reductive elimination
- Sigma bond metathesis



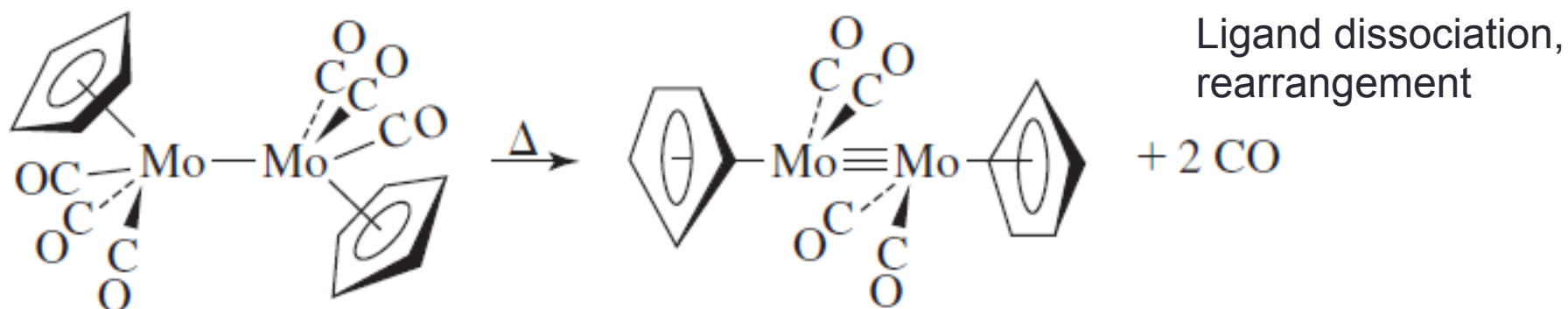
Ligand dissociation reactions

- Ligand dissociation reactions either involve the loss of a ligand or a ligand substitution reaction
- In reactions classified as ligand dissociation, metal oxidation state(s) do not change, however complexes may rearrange following ligand dissociation
- The same principles apply from what we have already learned about ligand substitution reactions
- If a reaction involves simply the gain of a ligand in the metal coordination sphere, this is called ligand association



Ligand dissociation: CO dissociation

- CO can be lost from a metal complex either thermally (with heat) or photochemically (with light)
- The reaction will either cause the metal complex to rearrange or for CO to be replaced by another ligand



ligand
substitution



Ligand dissociation: phosphines

- In some ligand dissociation reactions, steric bulk plays an important role, with bulkier ligands dissociating more quickly to relieve steric strain in the metal complex
- This can be observed in some phosphine dissociation reactions, where phosphines with bigger cone angles have faster ligand dissociation rates

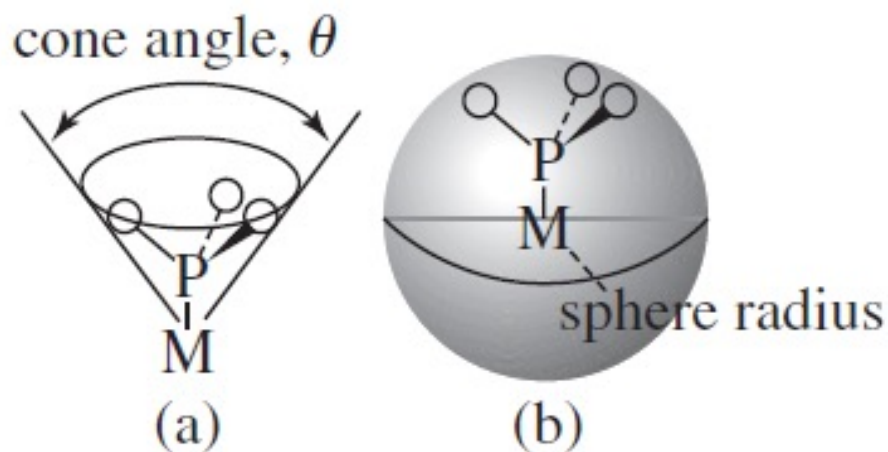
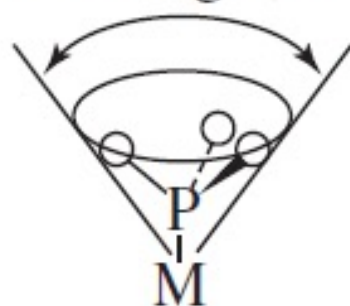


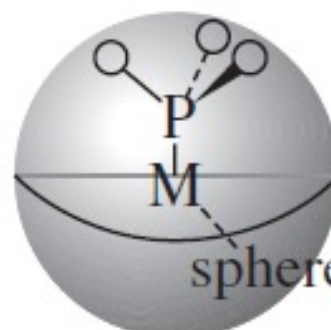
TABLE 14.1 Ligand Cone Angles and % V_{bur} for Tertiary Phosphines

Ligand	Cone Angle θ	% V_{bur}	Ligand	Cone Angle θ	% V_{bur}
PH_3	87°		$\text{P}(\text{CH}_3)(\text{C}_6\text{H}_5)_2$	136°	
PF_3	104°		$\text{P}(\text{CF}_3)_3$	137°	
$\text{P}(\text{OCH}_3)_3$	107° (128°) ^a	26.4 ^b	$\text{P}(\text{C}_6\text{H}_5)_3$	145°	29.6
$\text{P}(\text{OC}_2\text{H}_5)_3$	109°		$\text{P}(\text{cyclo-C}_6\text{H}_{11})_3$	170°	31.8
$\text{P}(\text{CH}_3)_3$	118°	22.2	$\text{P}(t\text{-C}_4\text{H}_9)_3$	182°	36.7
PCl_3	124°		$\text{P}(\text{C}_6\text{F}_5)_3$	184°	37.3
$\text{P}(\text{OC}_6\text{H}_5)_3$	128° (155°) ^a	30.7	$\text{P}(o\text{-C}_6\text{H}_4\text{CH}_3)_3$	194°	41.4
$\text{P}(\text{C}_2\text{H}_5)_3$	132°	27.8	$\text{P}\left(\text{Me-C}_6\text{H}_3\text{Me}\right)_3$	212°	47.6

cone angle, θ

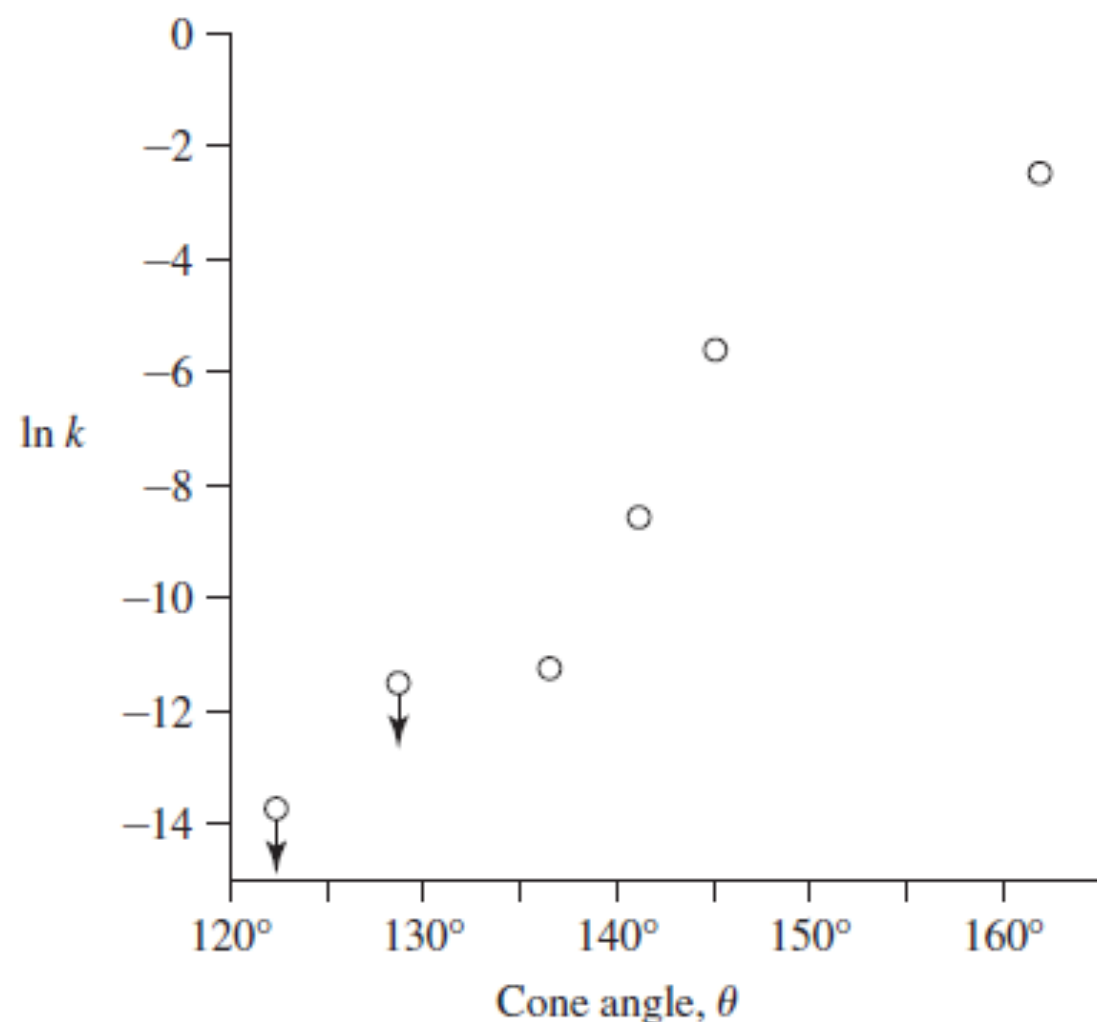
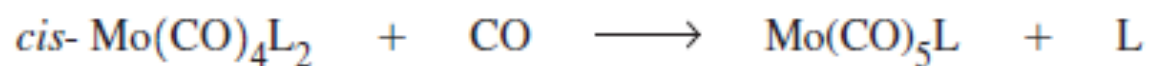


(a)



(b)



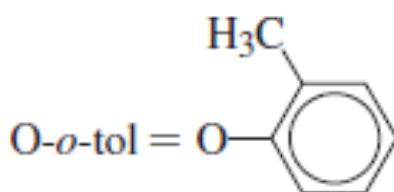


L^a	θ	$k \text{ (s}^{-1}\text{)}$
PPh ₂ Cy	162°	6.40×10^{-2}
PPh ₃	145°	3.16×10^{-3}
P(O- <i>o</i> -tol) ₃	141°	1.60×10^{-4}
PMePh ₂	136°	1.33×10^{-5}
P(OPh) ₃	128°	$<1.0 \times 10^{-5}$
PMe ₂ Ph	122°	$<1.0 \times 10^{-6}$

^a Cy = cyclohexyl

Ph = phenyl

Me = methyl



Oxidative addition and reductive elimination

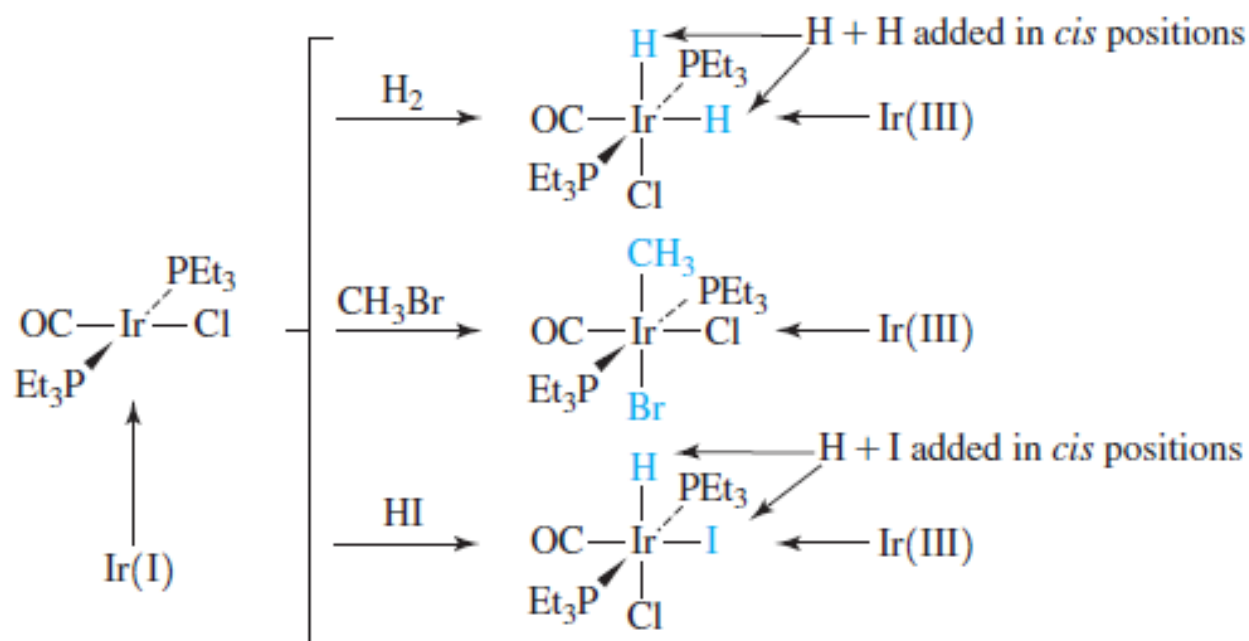
- **Oxidative addition (OA)** involves both *oxidation* of the metal center and *addition* of ligands to the metal coordination sphere
- The reverse reaction is called **reductive elimination (RE)** which involves both *reduction* of the metal center *elimination* of ligands from the metal coordination sphere



Type of Reaction	Change in Coordination Number	Change in Formal Oxidation State of Metal	Change in Electron Count
Oxidative addition	Increase by 2	Increase by 2	Increase by 2
Reductive elimination	Decrease by 2	Decrease by 2	Decrease by 2

Oxidative addition reactions

- OA reactions are very common with square planar complexes (why?)
- Example: OA reactions of *trans*-Ir(CO)Cl(PEt₃)₂

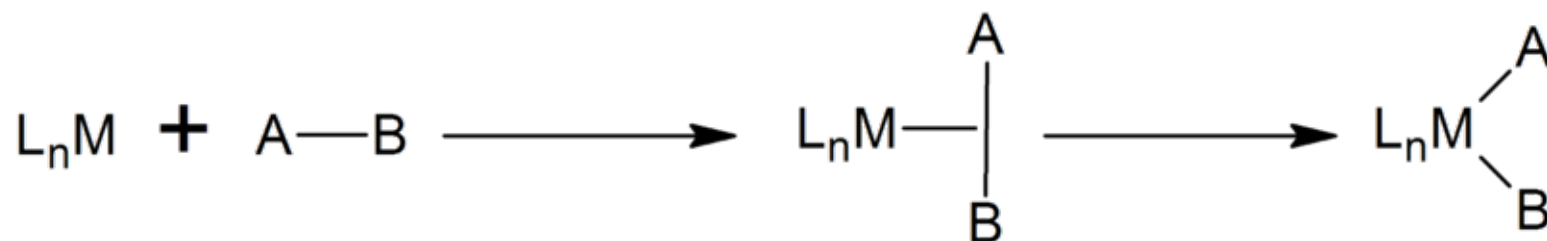


Depending on reactants, either *trans* or *cis* OA products can form

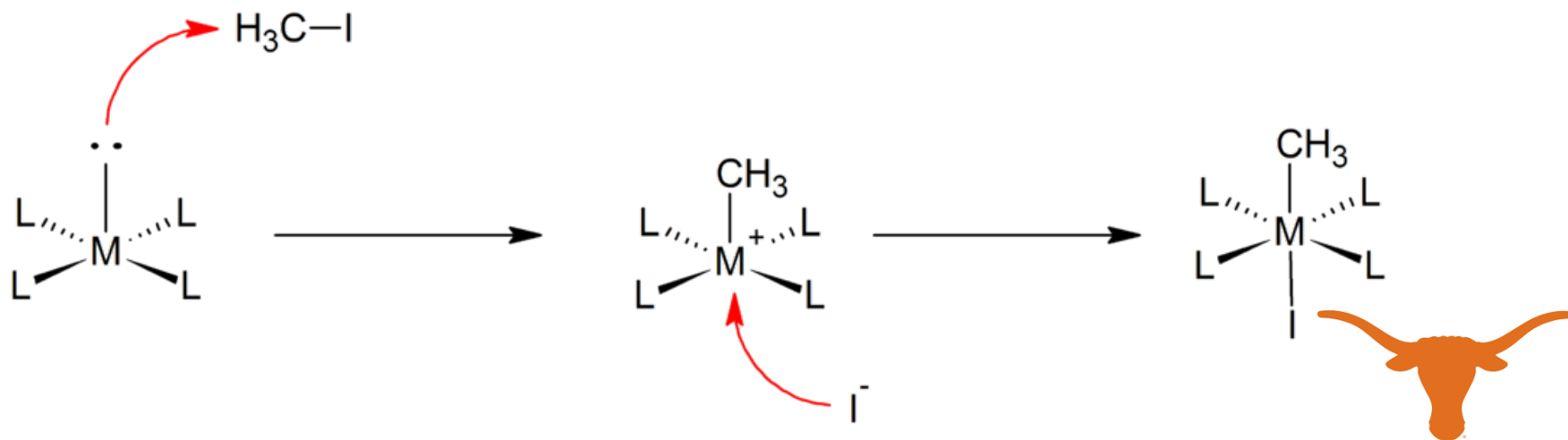


Oxidative addition reactions

- Different mechanisms possible including:
 - **Concerted pathway** (leads to cis addition)

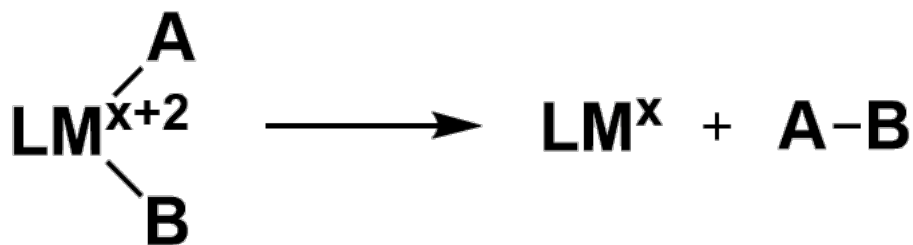


- **S_N²-type** (leads to trans addition)



Reductive elimination reactions

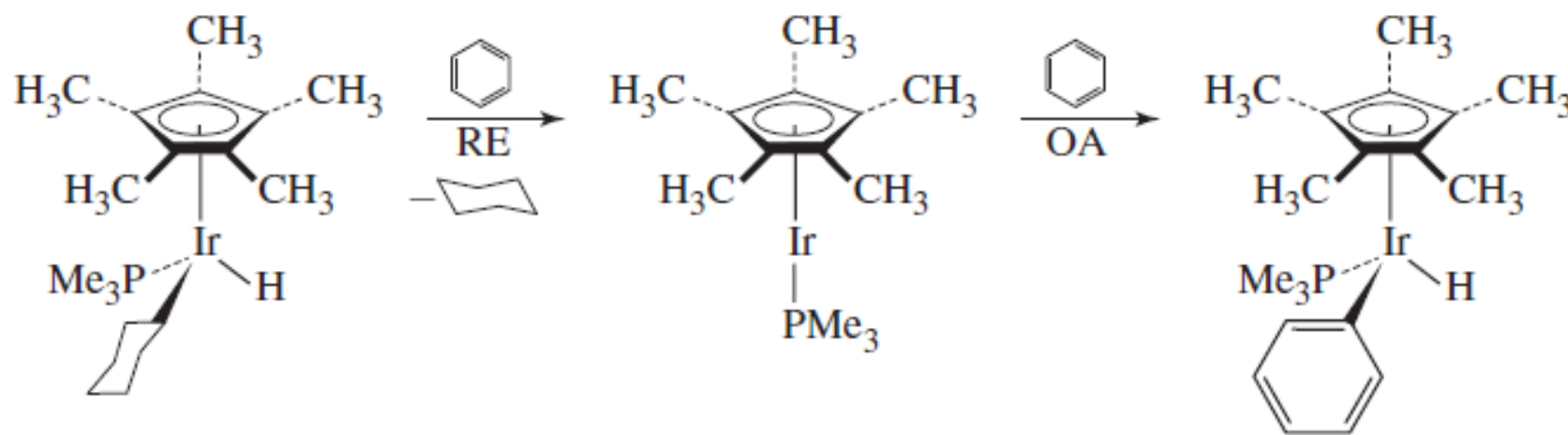
- RE is reverse of OA
- Will go through similar reaction mechanisms to OA (*principle of microscopic reversibility*)
- RE can result in the formation of a variety of bonds including H-H, C-H, C-C, and C-X (X = halide, amide, alkoxide, thiolate, phosphide)



OA and RE in C-H bond activation

- C-H bond activation is a reaction in which a transition metal catalyst cleaves a strong C-H bond

Activation of C-H bond in benzene (R.Bergman, UC-Berkeley)

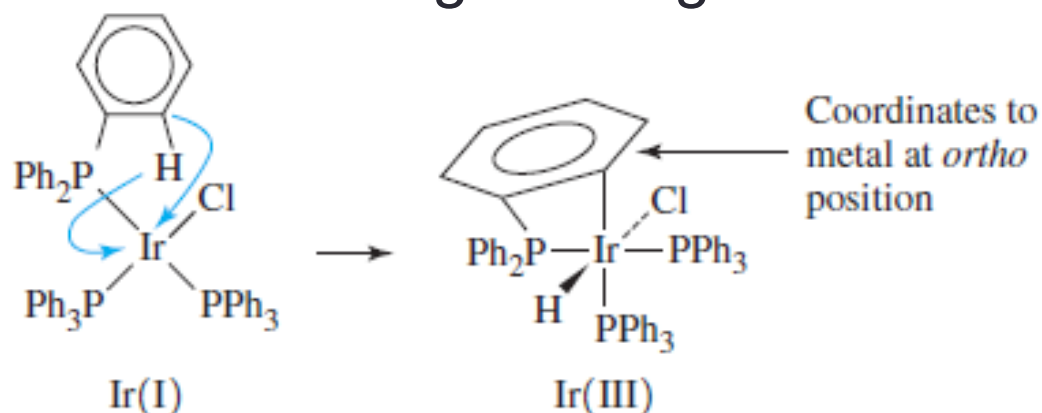


Potential way to make benzene derivatives

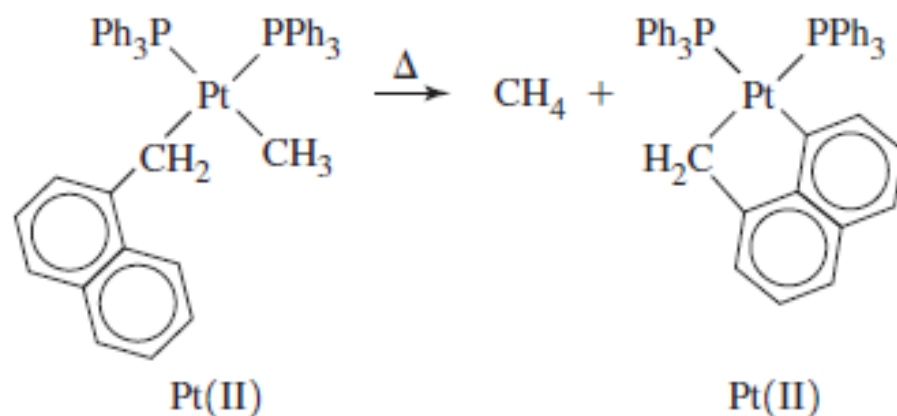


OA and RE in C-H bond activation

- Cyclometallation: C-H bond activations that incorporate metals into organic rings



OA forms a 6-coordinate cyclometallated species

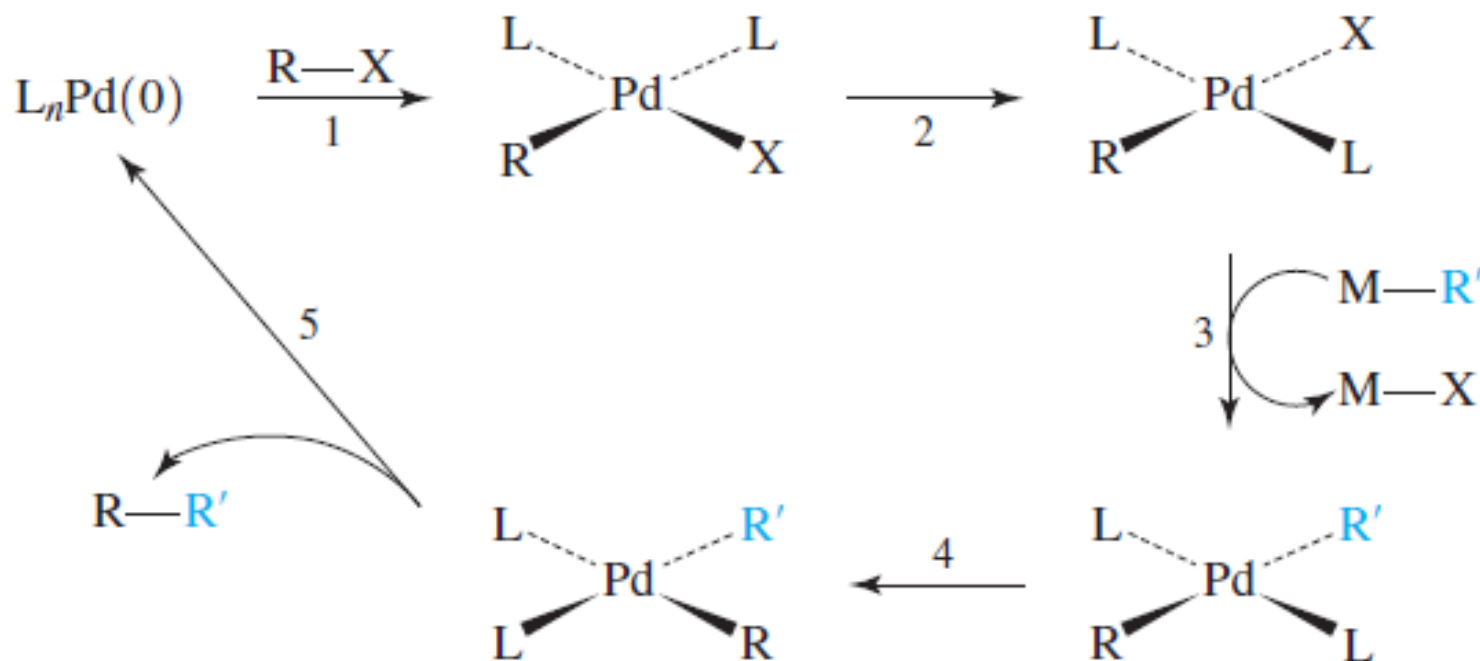


OA followed by RE of CH₄ forms a 4-coordinate cyclometallated species



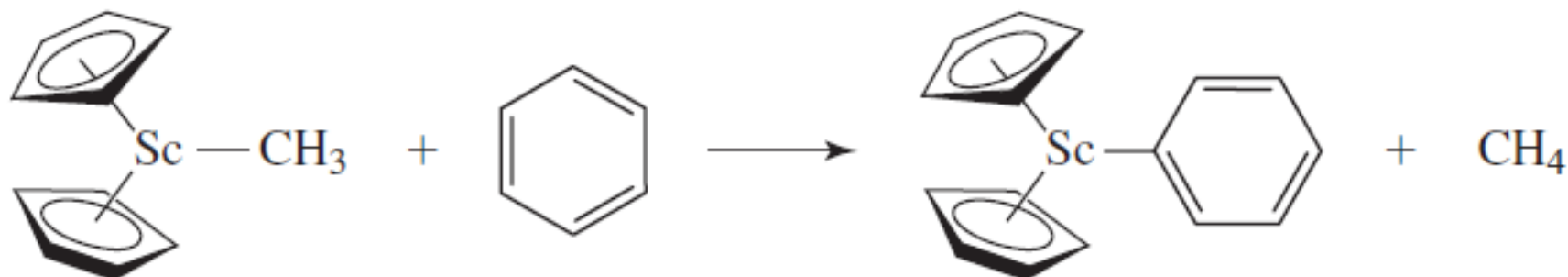
OA and RE in Pd-cross coupling

- Pd cross coupling reactions are widely used in organic chemistry, OA and RE are key steps
- Include important named reactions: Heck, Sonogashira, Negishi, Stille, Suzuki



Sigma bond metathesis

- In this type of reaction, two sigma bonded ligands are exchanged on a metal center without a change in metal oxidation state



- In this reaction, C-H bond of methane is made and C-H bond of benzene is cleaved



Reactions involving modification of ligands

- Insertion
- Migration
- Hydride elimination
- Abstraction



Insertion reactions

- In insertion reactions, an incoming molecule inserts into a metal-ligand bond (this is not a ligand substitution as a new covalent bond is formed)



(a)

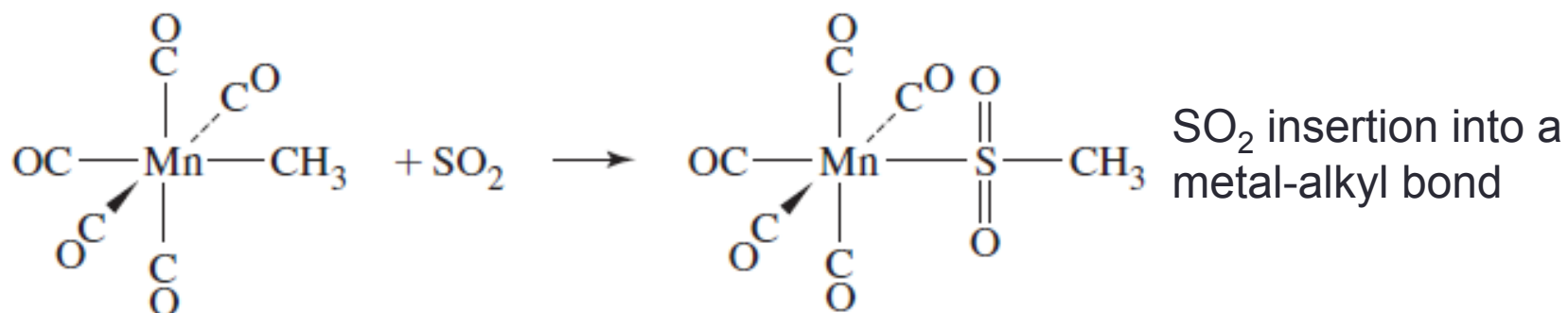
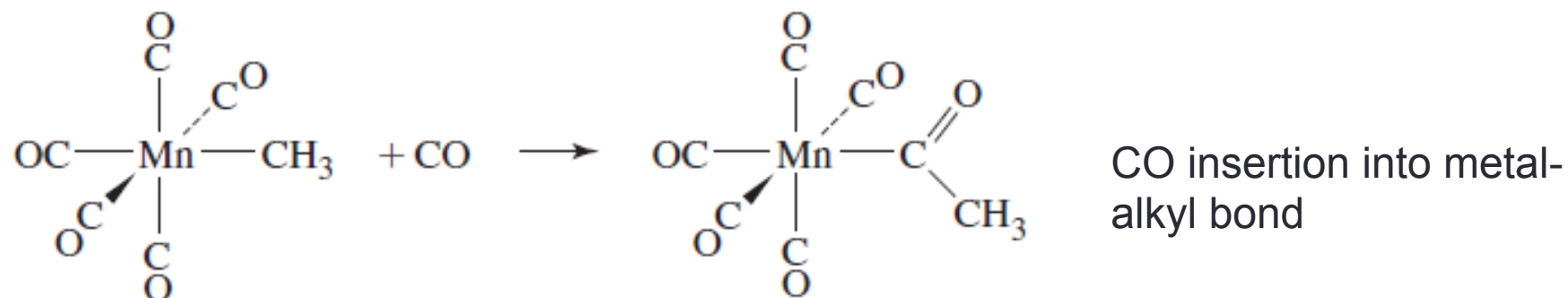


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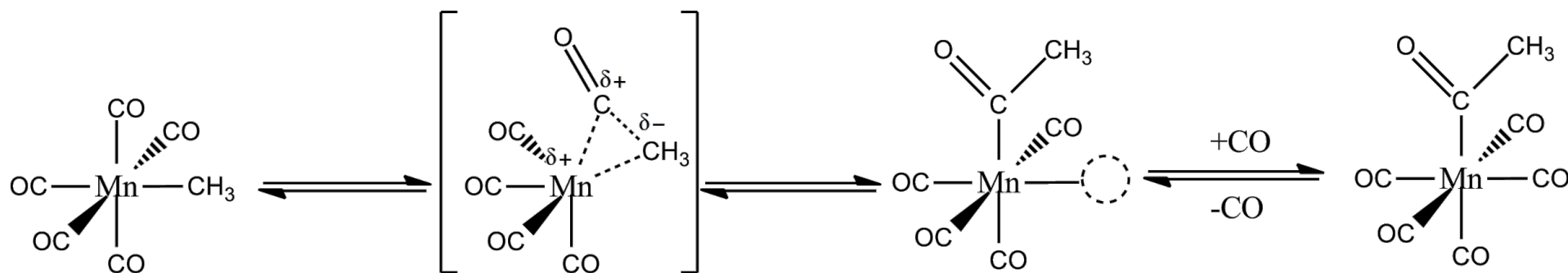
1,1 insertions

- Both new bonds formed are to the same atom



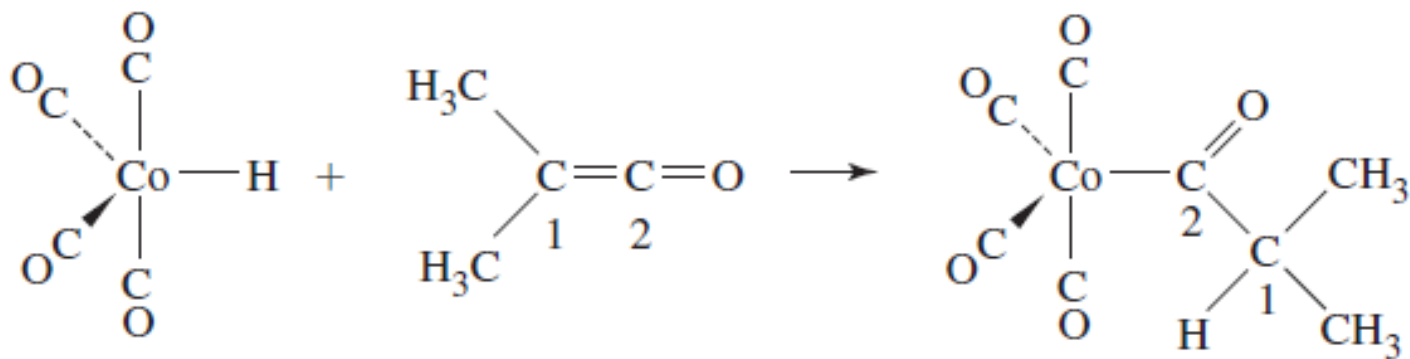
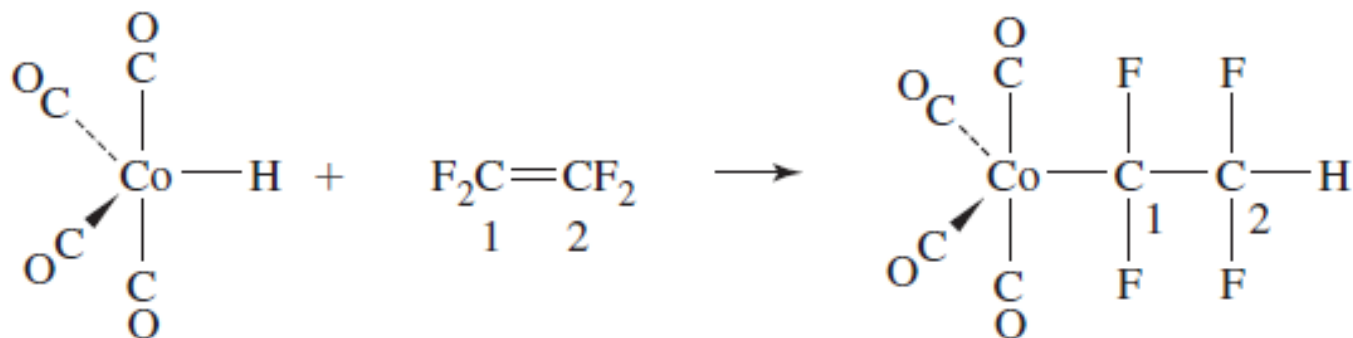
Migratory insertion

- Sometimes, inserted molecule is derived from another ligand on the metal center, this is called a migratory insertion (migratory 1,1-insertion in this case)



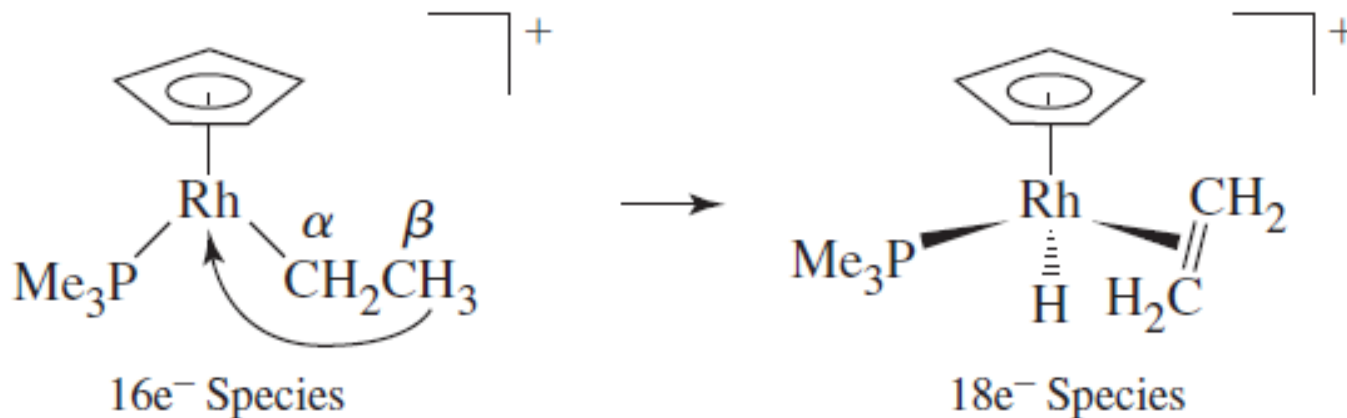
1,2 insertions

- New bonds are made to adjacent (1,2) atoms



Hydride elimination

- In hydride eliminations, a hydrogen atom is transferred from a ligand to a metal
- The most common is **Beta hydride elimination**

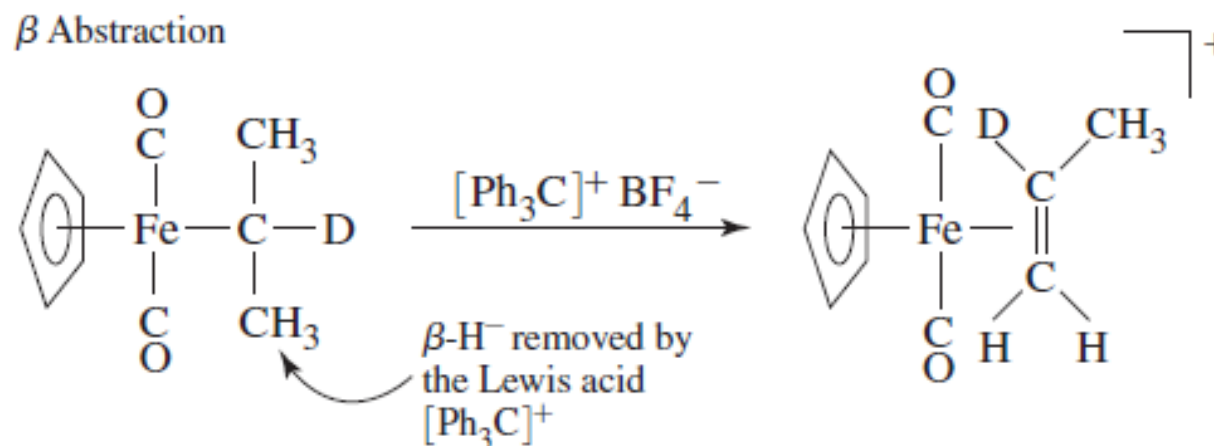
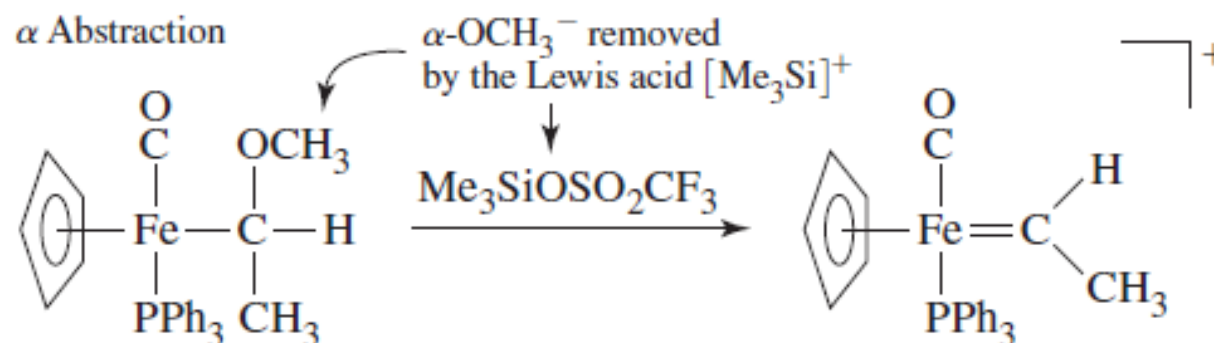


- Metal-alkyl complexes without beta hydrogens are more stable because they cannot undergo beta hydride elimination
- Alpha, gamma, and other hydride eliminations are also possible



Abstraction reactions

- These are elimination reactions in which the coordination number of the metal does not change
- Mediated by an external reagent like a Lewis acid



Organometallic catalysis

- Many of these single steps can be combined to form a catalytic cycle to convert one molecule into another
- This is of interest to industry: how to convert inexpensive feedstocks (e.g. petroleum) into \$\$\$ molecules?
- One of the simplest examples is catalytic deuteration, a process catalyzed by H_3TaCp_2 : this process is a series of RE and OA reactions

