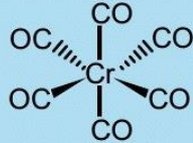
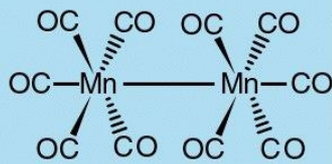
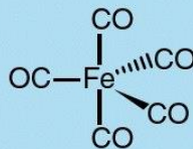
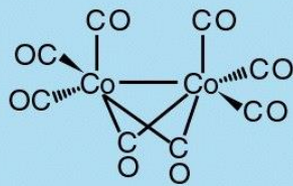
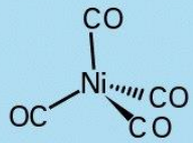
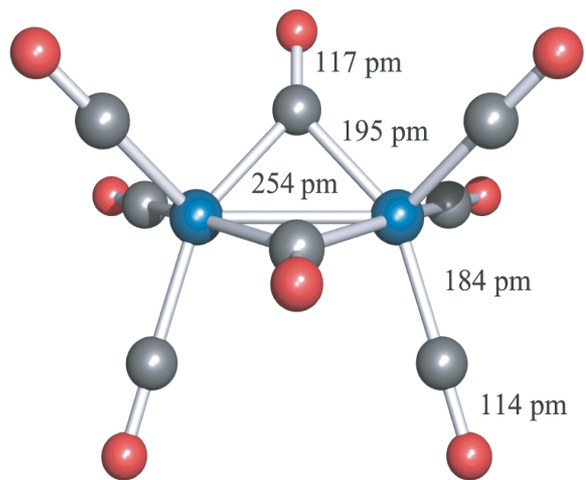
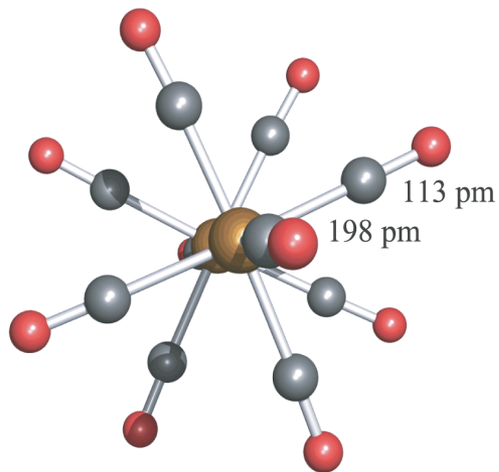


# Carbonili omoleptici

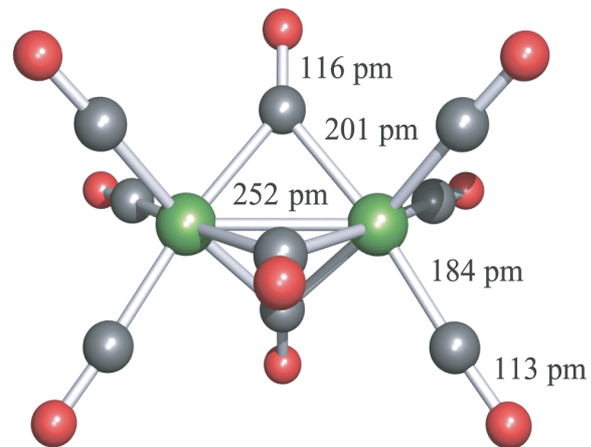
Table 21.5 Formulas and electron count for some 3d-series carbonyls				
Group	Formula	Valence electrons		Structure
6	$\text{Cr}(\text{CO})_6$	Cr	6	
		6(CO)	$\frac{12}{18}$	
7	$\text{Mn}_2(\text{CO})_{10}$	Mn	7	
		5(CO)	10	
		M—M	$\frac{1}{18}$	
8	$\text{Fe}(\text{CO})_5$	Fe	8	
		5(CO)	$\frac{10}{18}$	
9	$\text{Co}_2(\text{CO})_8$	Co	9	
		4(CO)	8	
		M—M	$\frac{1}{18}$	
10	$\text{Ni}(\text{CO})_4$	Ni	10	
		4(CO)	$\frac{8}{18}$	



(d)  
 $\text{Co}_2(\text{CO})_8$



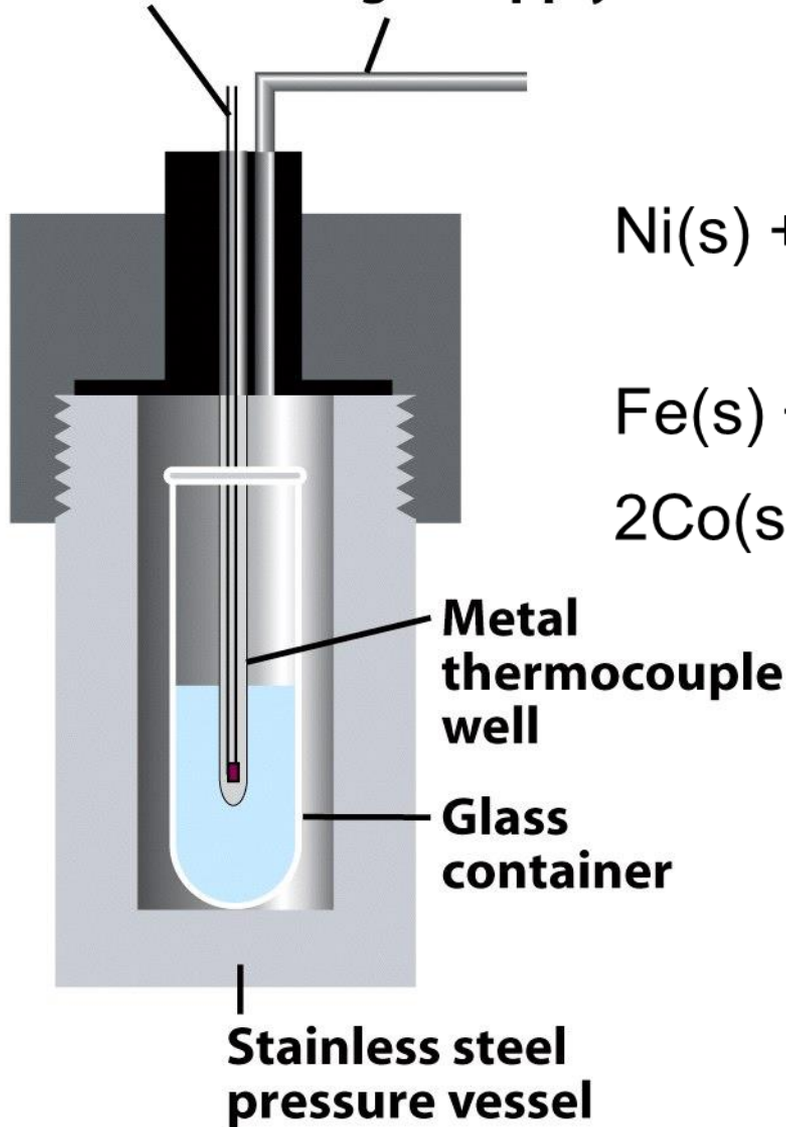
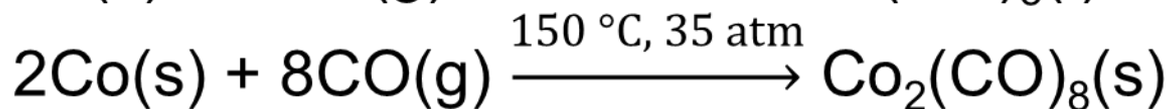
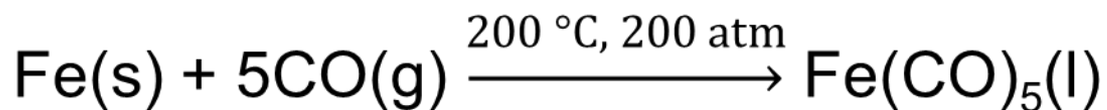
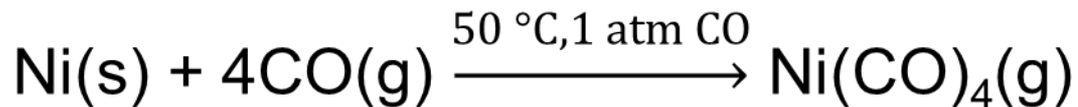
(b)  
 $\text{Re}_2(\text{CO})_{10}$



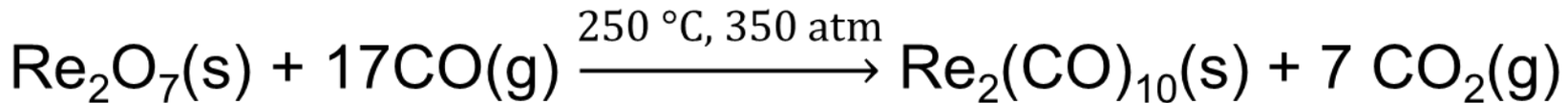
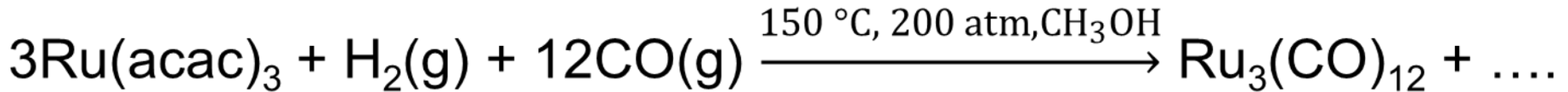
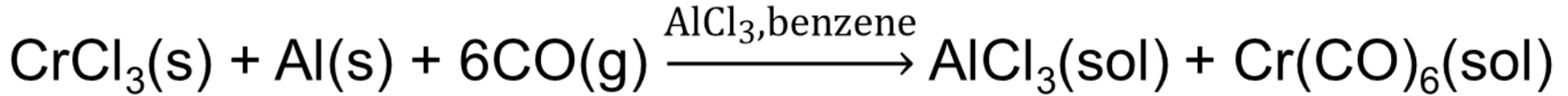
(c)  
 $\text{Fe}_2(\text{CO})_9$

Thermocouple  
High-pressure  
gas supply

# Reazione diretta

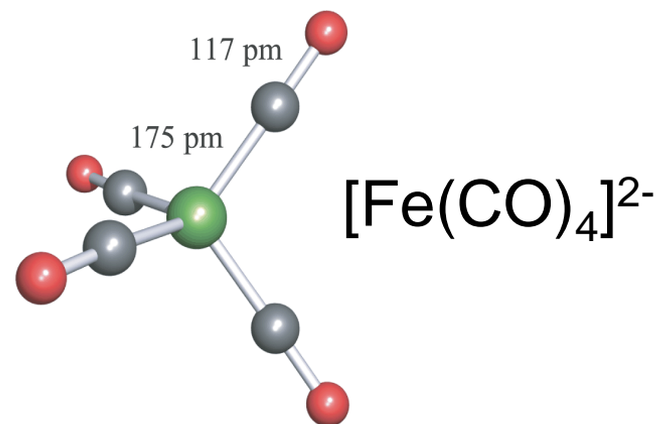


# Carbonilazione riduttiva

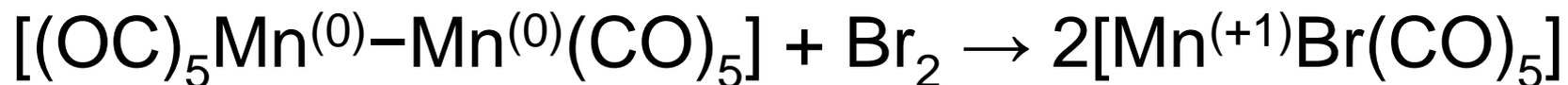


Group number	5	6	7	8	9	10	
First row metals	<b>V(CO)<sub>6</sub></b> Dark blue solid; paramagnetic; dec. 343 K	<b>Cr(CO)<sub>6</sub></b> White solid; sublimes <i>in vacuo</i> ; dec. 403 K	<b>Mn<sub>2</sub>(CO)<sub>10</sub></b> Yellow solid; mp 427 K	<b>Fe(CO)<sub>5</sub></b> Yellow liquid; mp 253 K; bp 376 K		<b>Co<sub>2</sub>(CO)<sub>8</sub></b> Air-sensitive, orange-red solid; mp 324 K	<b>Ni(CO)<sub>4</sub></b> Colourless, volatile liquid; highly toxic vapour; bp 316 K
Second row metals		<b>Mo(CO)<sub>6</sub></b> White solid; sublimes <i>in vacuo</i>	<b>Tc<sub>2</sub>(CO)<sub>10</sub></b> White solid; slowly dec. in air; mp 433 K	<b>Fe<sub>2</sub>(CO)<sub>9</sub></b> Golden crystals; mp 373 K (dec.)	<b>Fe<sub>3</sub>(CO)<sub>12</sub></b> Dark green solid; dec. 413 K	<b>Co<sub>4</sub>(CO)<sub>12</sub></b> Air-sensitive, black solid	
				<b>Ru(CO)<sub>5</sub></b> Colourless liquid; mp 251 K; dec. in air at 298 K to Ru <sub>3</sub> (CO) <sub>12</sub> + CO		<b>Co<sub>6</sub>(CO)<sub>16</sub></b> Black solid; slowly dec. in air	
				<b>Ru<sub>3</sub>(CO)<sub>12</sub></b> Orange solid; mp 427 K; sublimes <i>in vacuo</i>		<b>Rh<sub>4</sub>(CO)<sub>12</sub></b> Red solid; >403 K dec. to Rh <sub>6</sub> (CO) <sub>16</sub>	
Third row metals		<b>W(CO)<sub>6</sub></b> White solid; sublimes <i>in vacuo</i>	<b>Re<sub>2</sub>(CO)<sub>10</sub></b> White solid; mp 450 K	<b>Os(CO)<sub>5</sub></b> Yellow liquid; mp 275 K		<b>Rh<sub>6</sub>(CO)<sub>16</sub></b> Black solid; dec. >573 K	
				<b>Os<sub>3</sub>(CO)<sub>12</sub></b> Yellow solid; mp 497 K		<b>Ir<sub>4</sub>(CO)<sub>12</sub></b> Slightly air-sensitive yellow solid; mp 443 K	
						<b>Ir<sub>6</sub>(CO)<sub>16</sub></b> Red solid	

# Riduzione dei metallocarbonili a metallocarbonilati



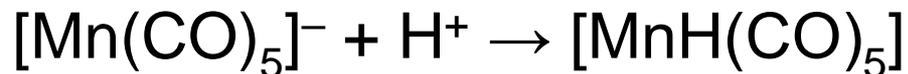
# Ossidazione (rottura ossidativa) di un legame M–M in metallo-carbonili



**Table 21.6** Acidity constants of *d*-metal hydrides in acetonitrile at 25°C

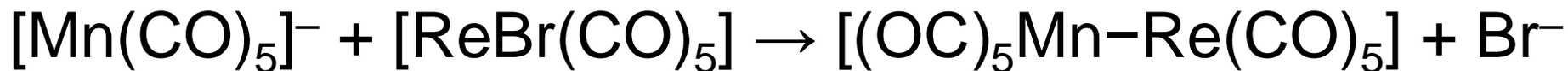
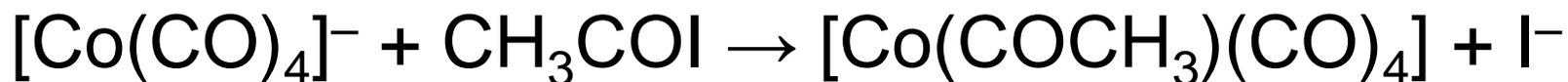
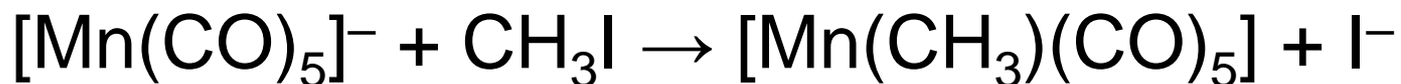
Hydride	pK <sub>a</sub>
[CoH(CO) <sub>4</sub> ]	8.3
[CoH(CO) <sub>3</sub> P(OPh) <sub>3</sub> ]	11.3
[Fe(H) <sub>2</sub> (CO) <sub>4</sub> ]	11.4
[CrH(Cp)(CO) <sub>3</sub> ]	13.3
[MoH(Cp)(CO) <sub>3</sub> ]	13.9
[MnH(CO) <sub>5</sub> ]	15.1
[CoH(CO) <sub>3</sub> PPh <sub>3</sub> ]	15.4
[WH(Cp)(CO) <sub>3</sub> ]	16.1
[MoH(Cp*)(CO) <sub>3</sub> ]	17.1
[Ru(H) <sub>2</sub> (CO) <sub>4</sub> ]	18.7
[FeH(Cp)(CO) <sub>2</sub> ]	19.4
[RuH(Cp)(CO) <sub>2</sub> ]	20.2
[Os(H) <sub>2</sub> (CO) <sub>4</sub> ]	20.8
[ReH(CO) <sub>5</sub> ]	21.1
[FeH(Cp*)(CO) <sub>2</sub> ]	26.3
[WH(Cp)(CO) <sub>2</sub> PMe <sub>3</sub> ]	26.6

## Protonazione di metallocarbonilati



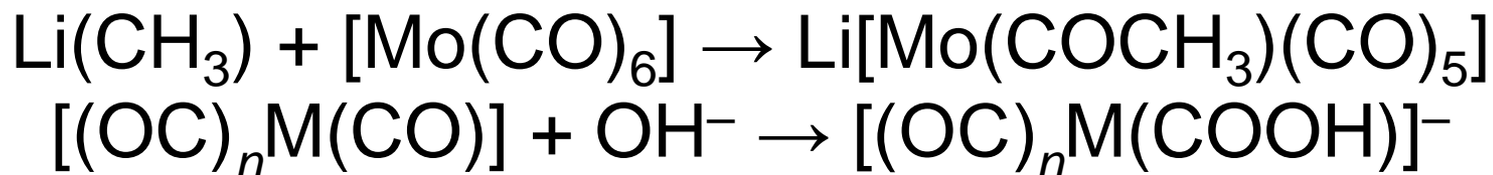
# Basicità dei metallocarbonilati

## Attacco nucleofilo

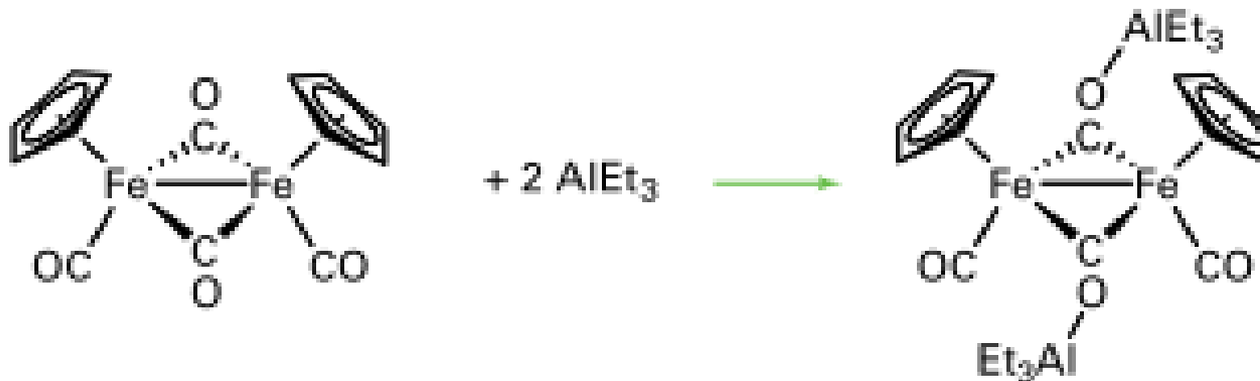


# Attacchi nucleofili ed elettrofili a carbonili

Metallo elettrone-povero: attacco nucleofilo su C



Metallo elettrone-ricco: attacco elettrofilo su O



**Table 21.7** Relation between the structure of a carbonyl complex and the number of CO stretching bands in its IR spectrum

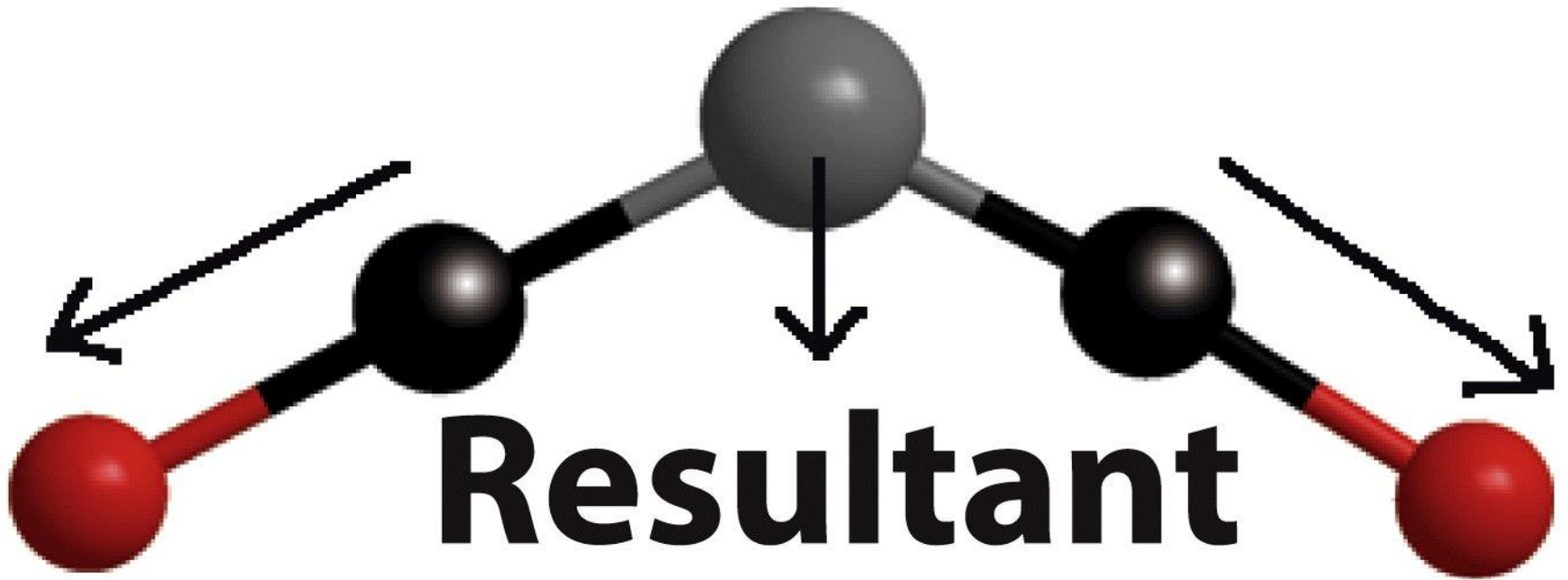
Complex	Isomer	Structure	Point group	Number of bands*	Complex	Isomer	Structure	Point group	Number of bands*
$M(CO)_6$			$O_h$	1	$M(CO)_5$			$D_{3h}$	2
$M(CO)_5L$			$C_{4v}$	3 <sup>†</sup>	$M(CO)_5$	<i>ax</i>		$C_{3v}$	3 <sup>§</sup>
$M(CO)_4L_2$	<i>trans</i>		$D_{4h}$	1	$M(CO)_4L$	<i>eq</i>		$C_{2v}$	4
$M(CO)_4L_2$	<i>cis</i>		$C_{2v}$	4 <sup>‡</sup>	$M(CO)_3L_2$	<i>trans</i>		$D_{3h}$	1
$M(CO)_3L_3$	<i>mer</i>		$C_{2v}$	3 <sup>‡</sup>	$M(CO)_3L_2$	<i>cis</i>		$C_s$	3
$M(CO)_3L_3$	<i>fac</i>		$C_{3v}$	2	$M(CO)_4$			$T_d$	1

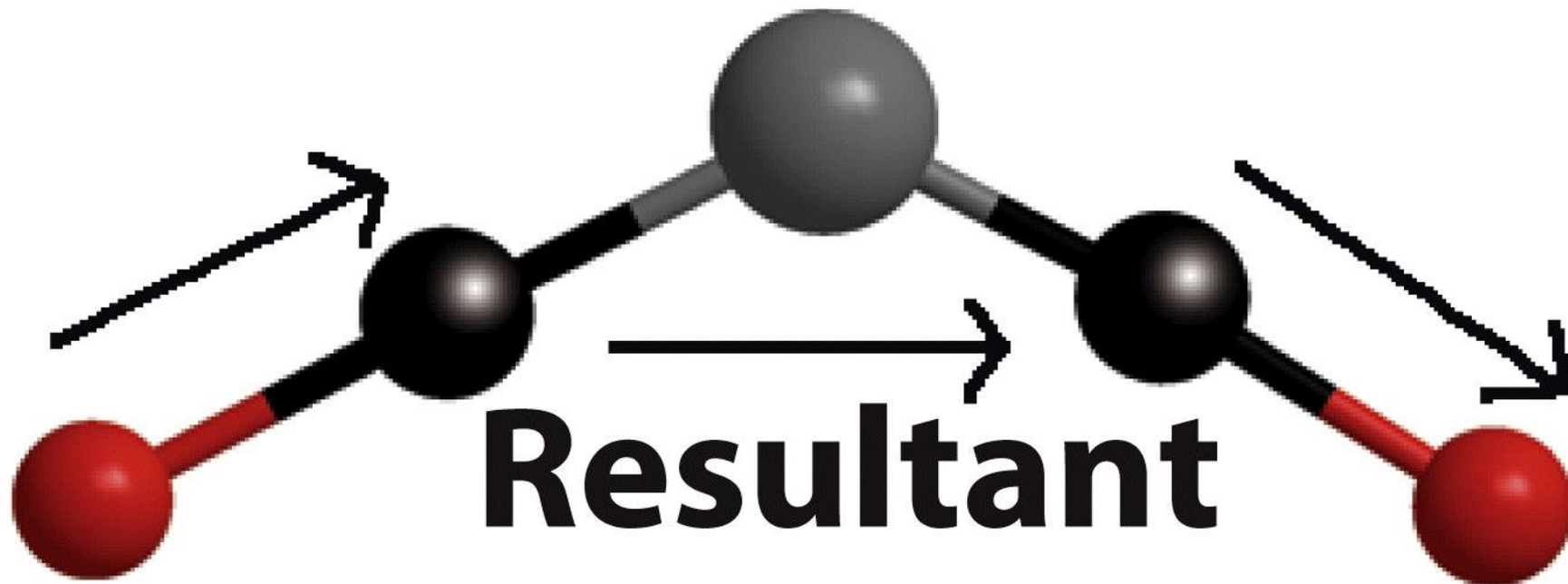
\* The number of IR bands expected in the CO stretching region is based on formal selection rules, and in some cases fewer bands are observed, as explained below.

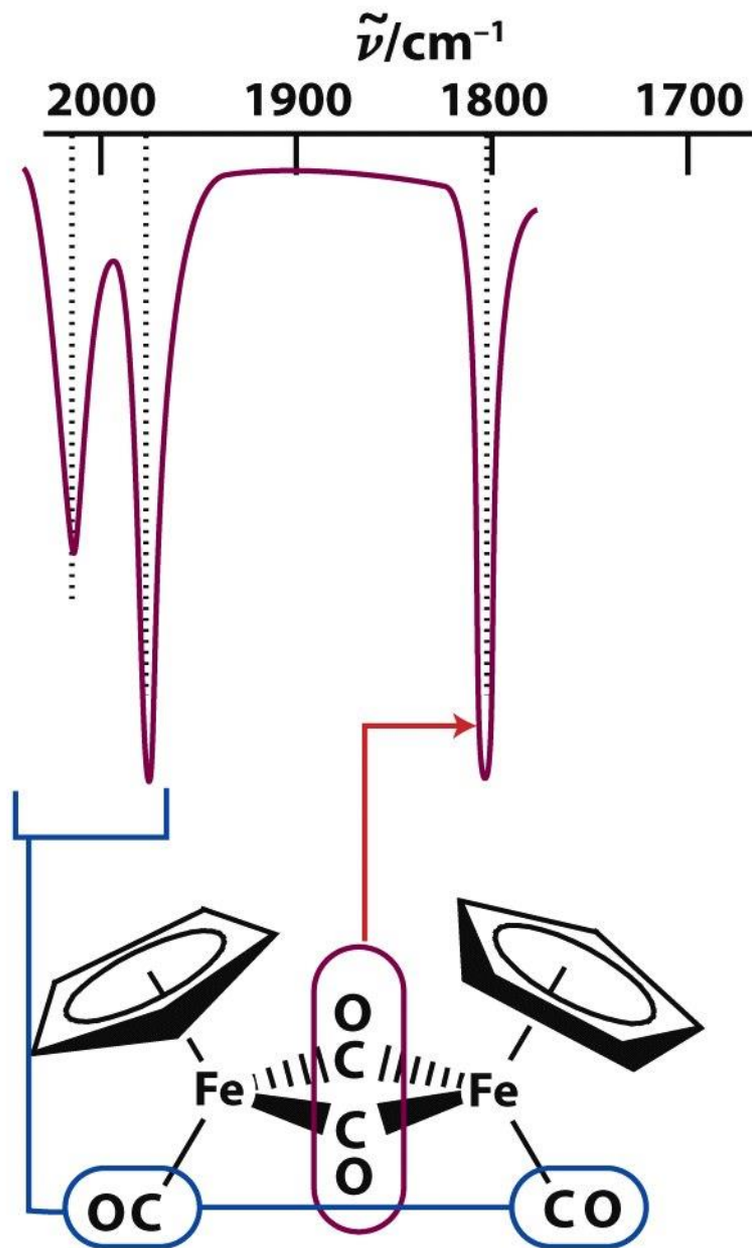
† If the fourfold array of CO ligands lies in the same plane as the metal atom, two bands will be observed.

‡ If the *trans* CO ligands are nearly collinear, one fewer band will be observed.

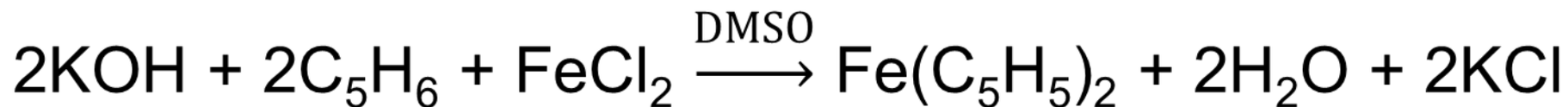
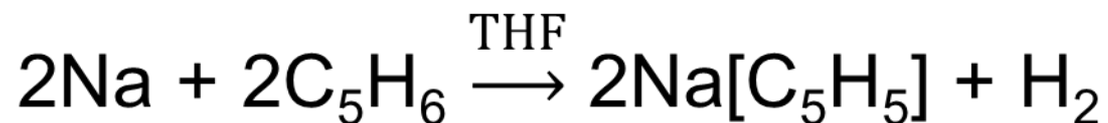
§ If the threefold array of CO ligands is nearly planar, only two bands will be observed.

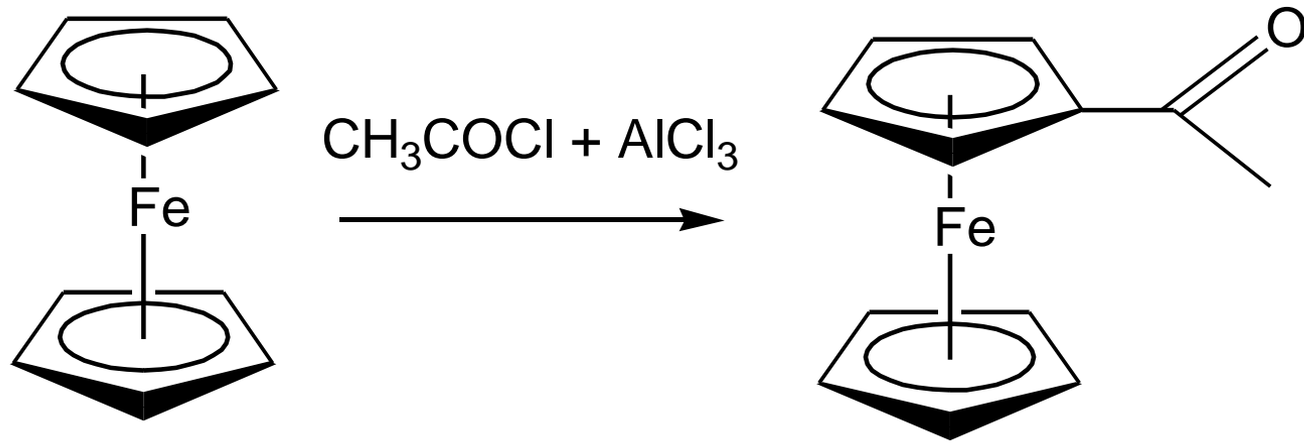




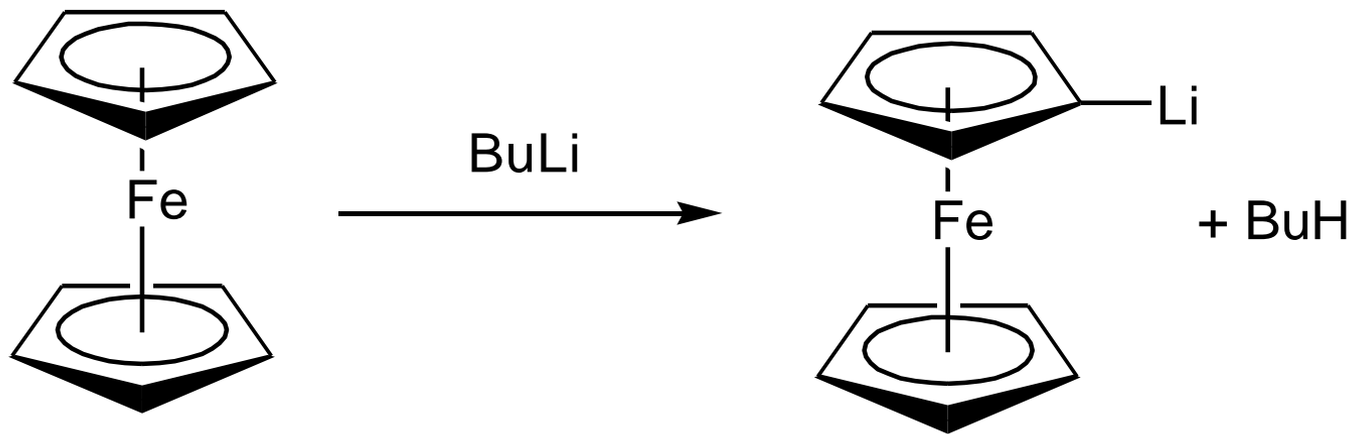


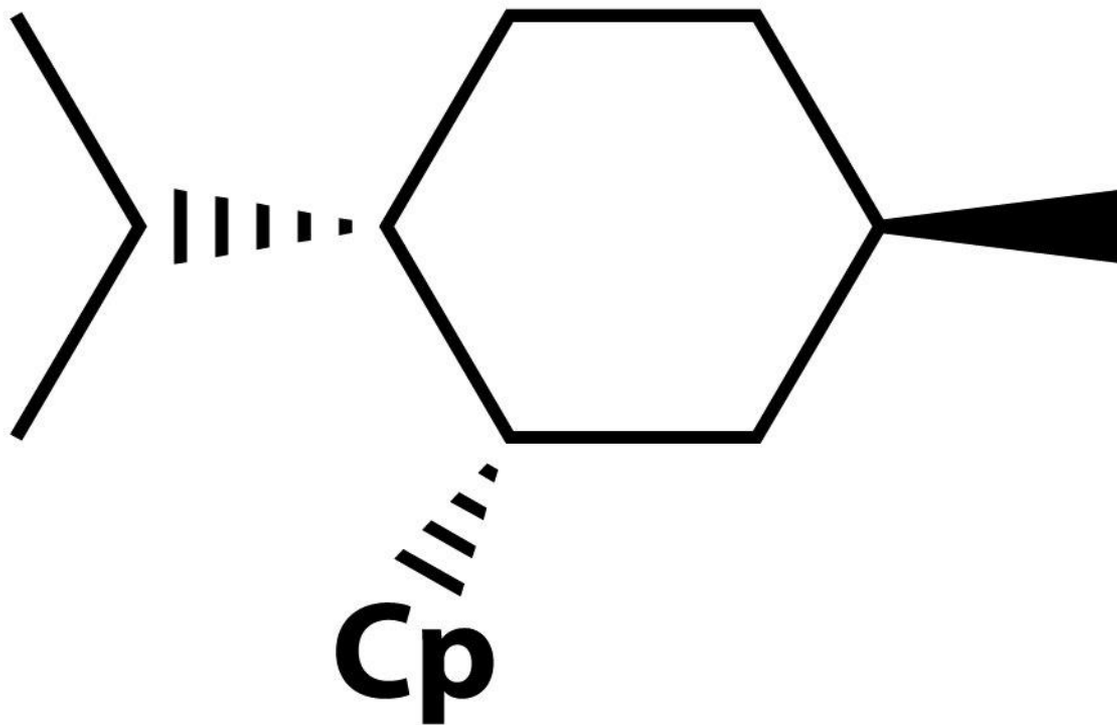
# Metalloceni



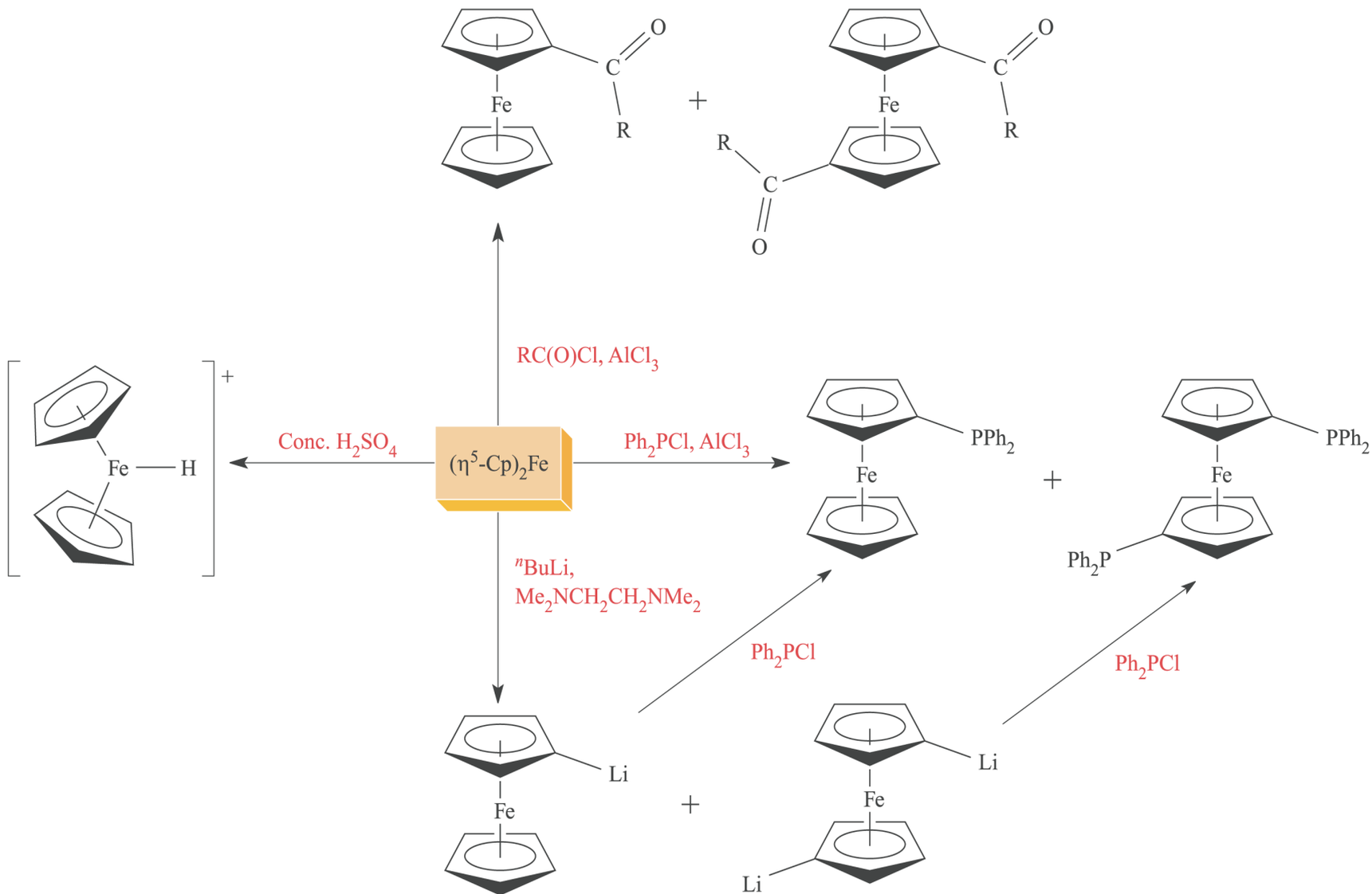


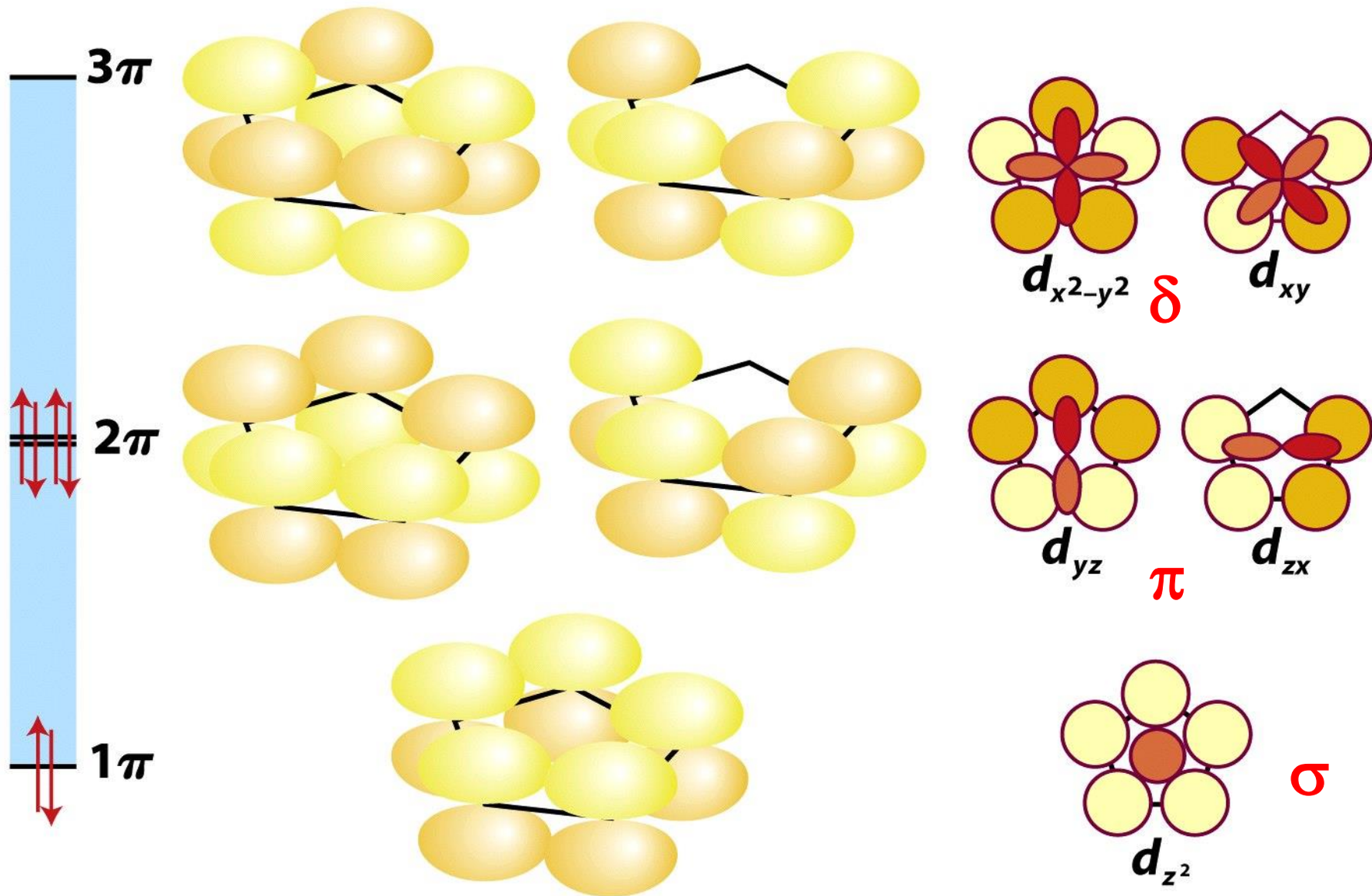
acilazione di Friedel – Crafts

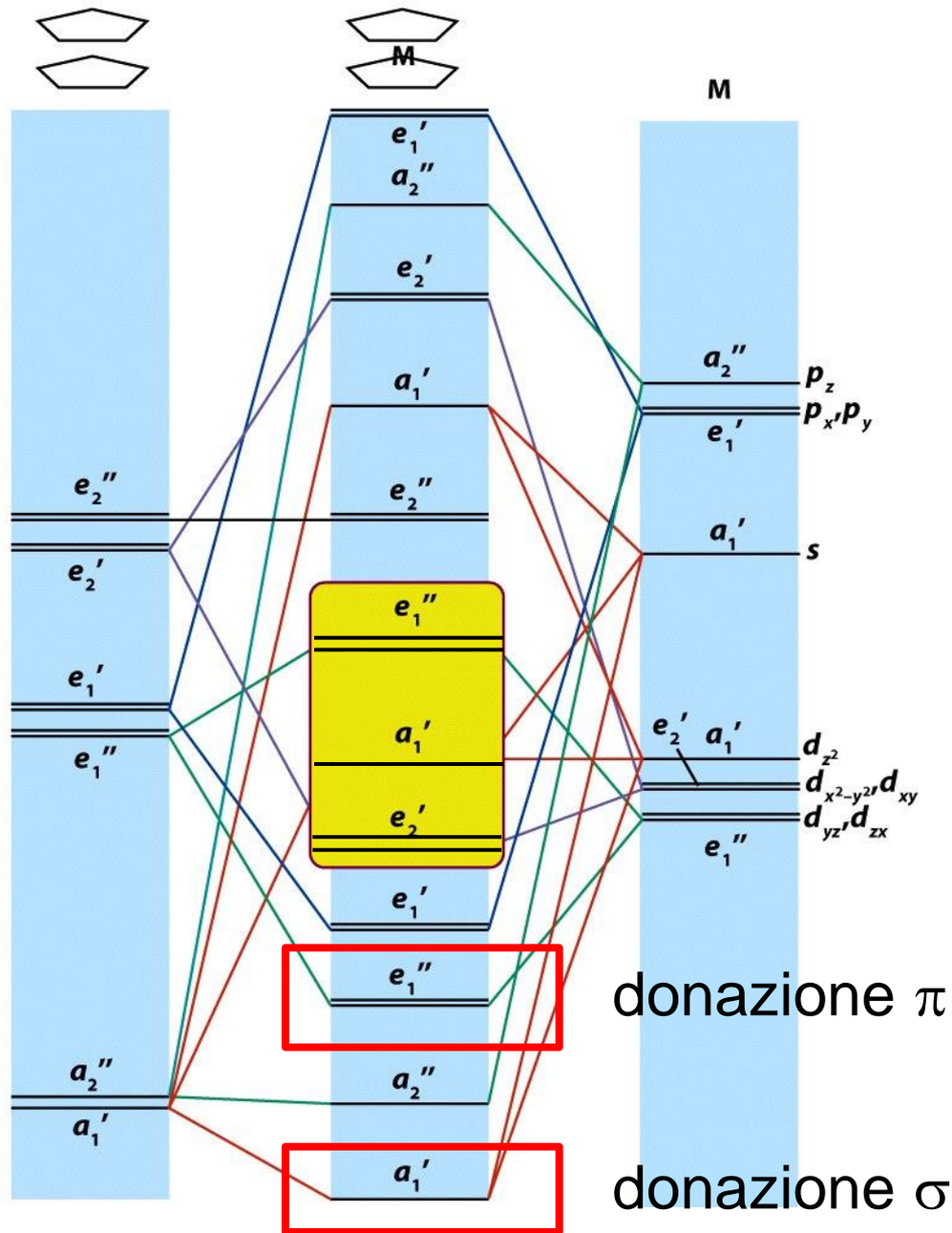


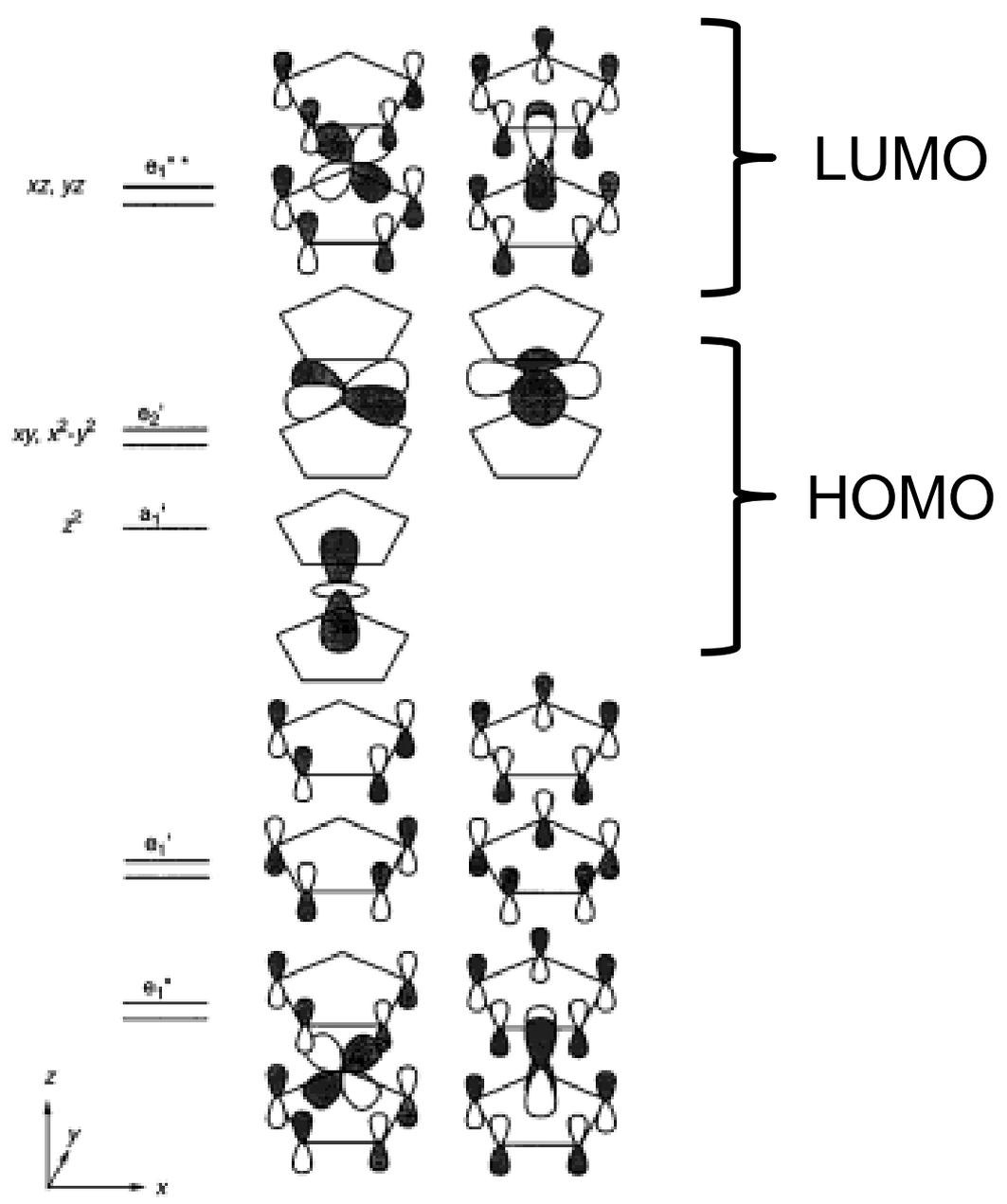


***neo-Menthylcyclopentadienyl***





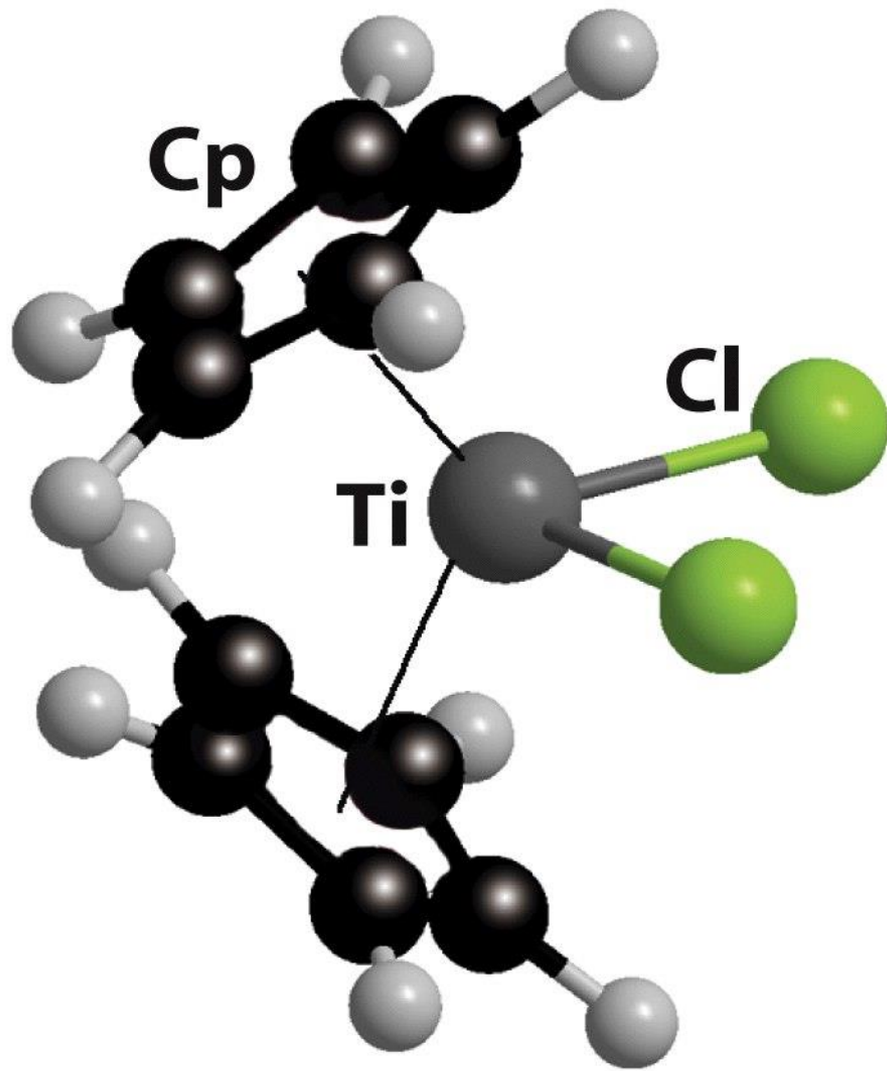




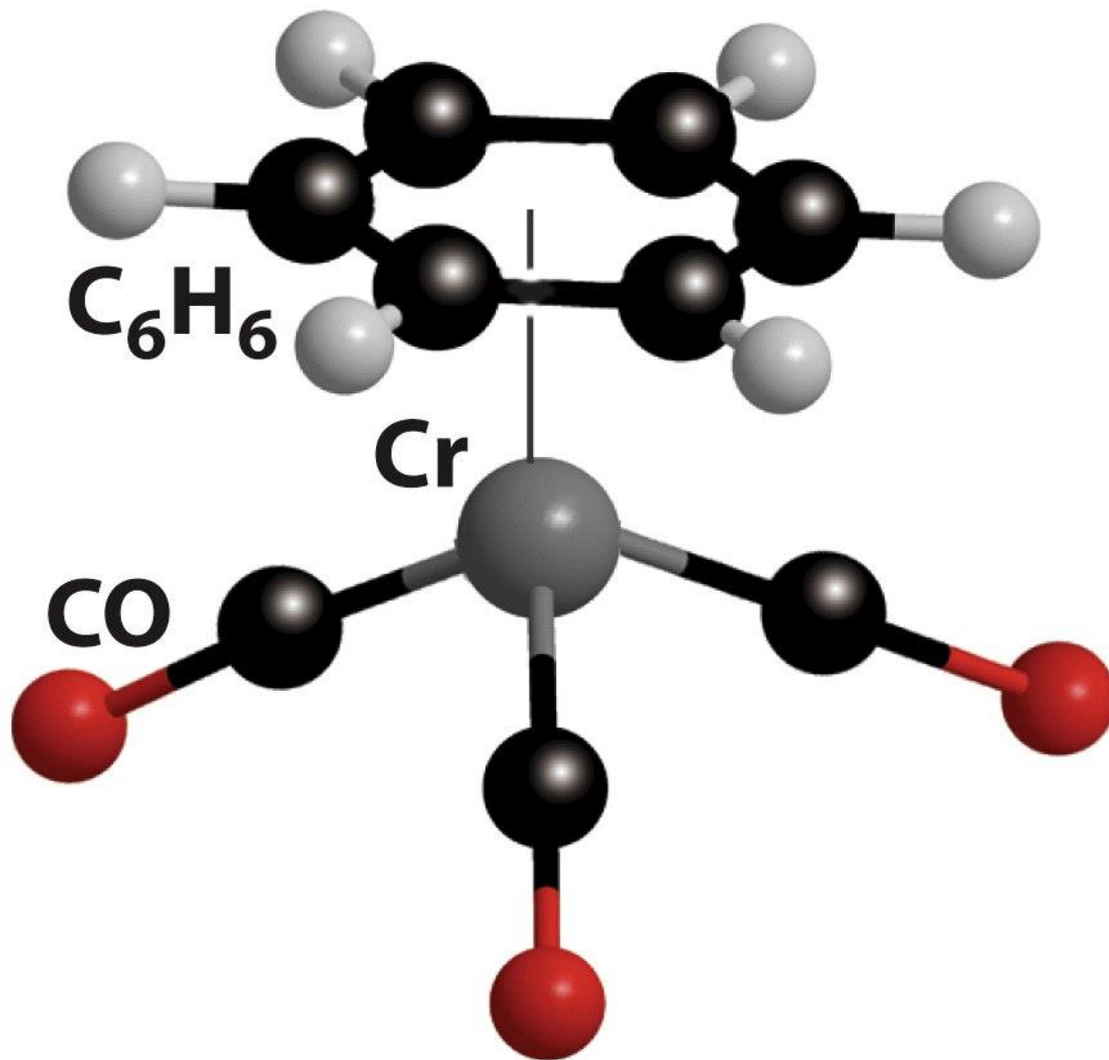
**Table 21.8** Electronic configuration and M—C bond length in  $[M(\eta^5\text{-Cp})_2]$  complexes

Complex	Valence electrons	Electron configuration	M—C bond length/pm
$[V(\eta^5\text{-Cp})_2]$	15	$e_2'^2 a_1'^1$	228
$[\text{Cr}(\eta^5\text{-Cp})_2]$	16	$e_2'^3 a_1'^1$	217
$[\text{Mn}(\eta^5\text{-Me-C}_5\text{H}_4)_2]^*$	17	$e_2'^3 a_1'^2$	211
$[\text{Fe}(\eta^5\text{-Cp})_2]$	18	$e_2'^4 a_1'^2$	206
$[\text{Co}(\eta^5\text{-Cp})_2]$	19	$e_2'^4 e_1''^1 a_1'^2$	212
$[\text{Ni}(\eta^5\text{-Cp})_2]$	20	$e_2'^4 e_1''^2 a_1'^2$	220

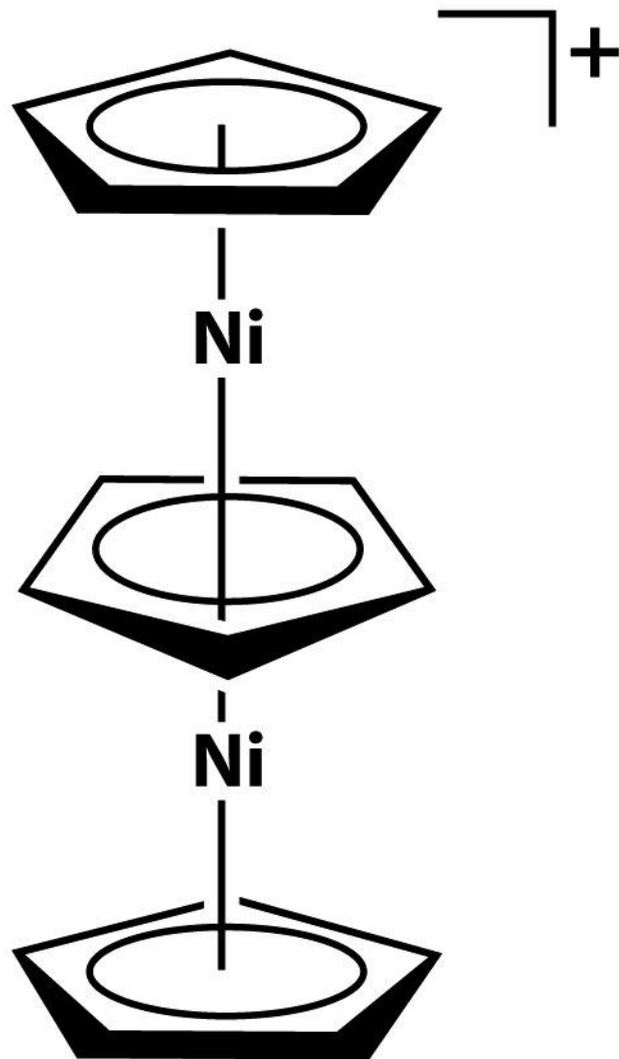
\*Data are quoted for this complex because  $[\text{Mn}(\eta^5\text{-Cp})_2]$  has a high-spin configuration and hence an anomalously long M—C bond (238 pm).



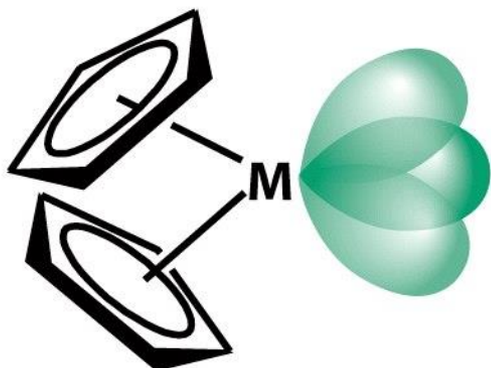
*Bent sandwich*



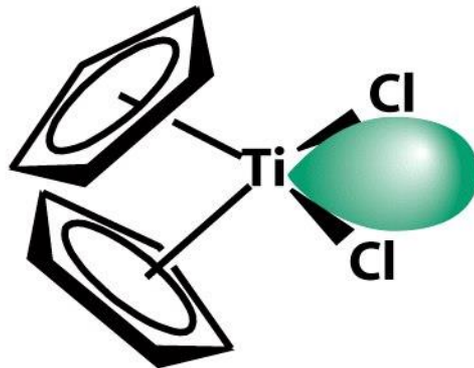
*Half sandwich o Piano stool*



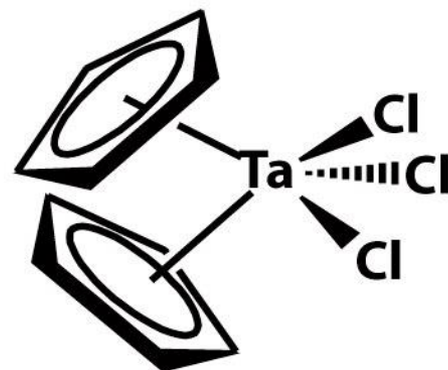
*Triple decker o doppio sandwich*



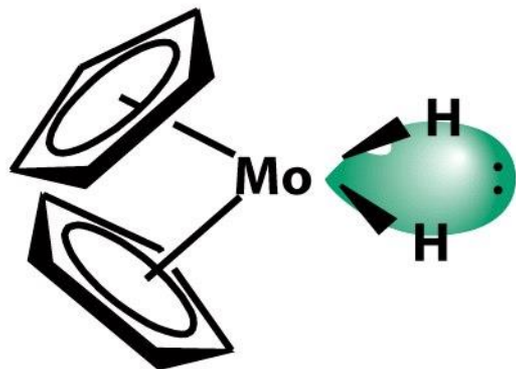
**General form**



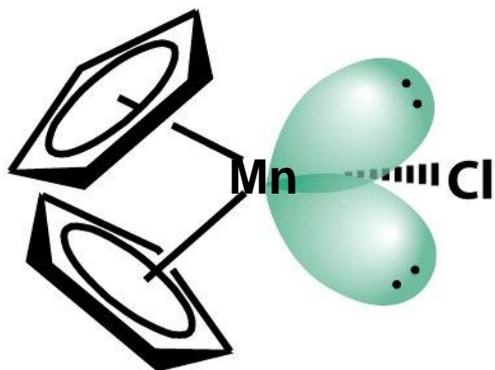
**16-electron**



**18-electron**



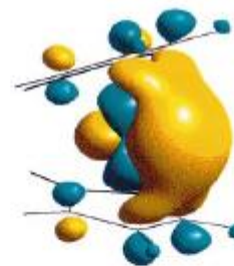
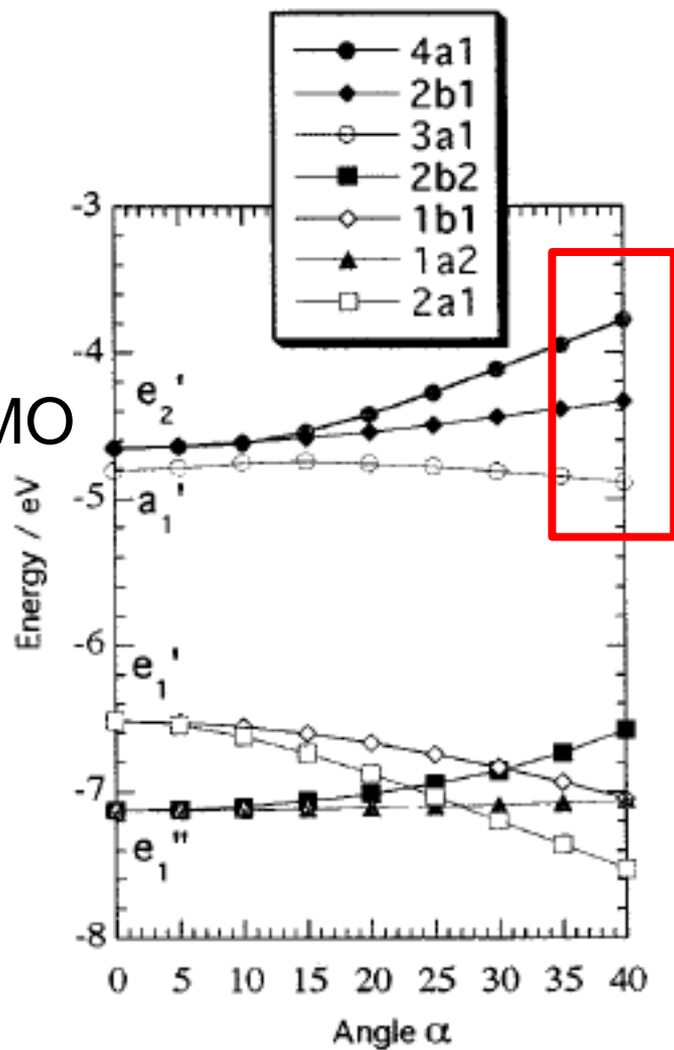
**18-electron**



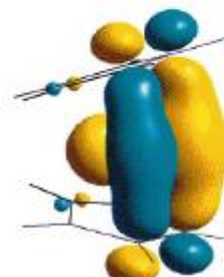
**18-electron**

# Diagramma di Walsh

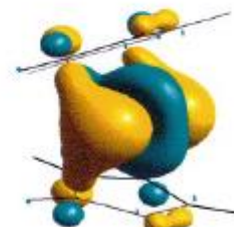
HOMO



4a1



2b1



3a1

# $\text{Cp}_2\text{ZrCl}_2$ : catalizzatore tipo Ziegler-Natta per la polimerizzazione di olefine

