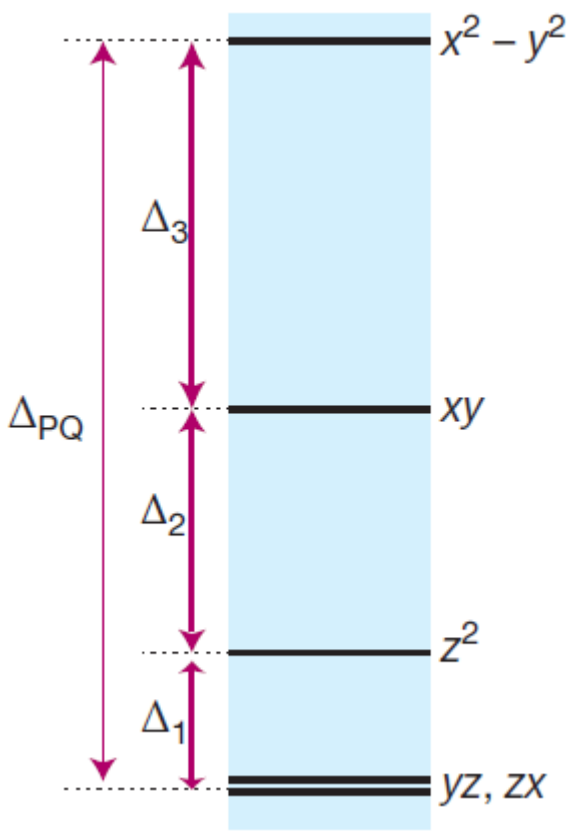
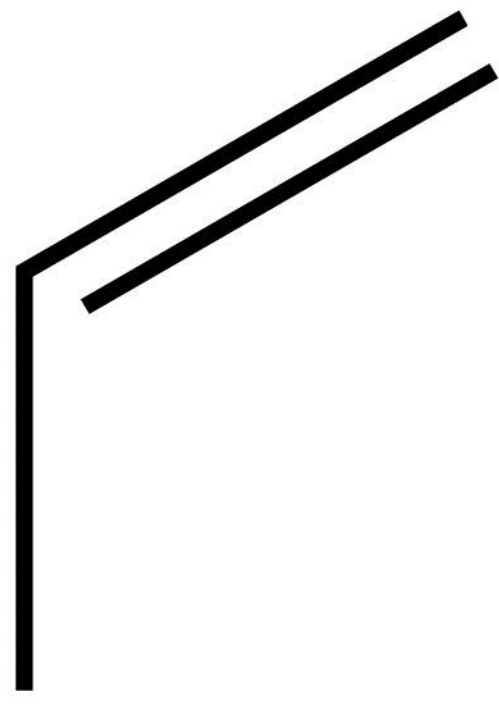


σ -donatore
+
 π -accettore

σ -donatore
+
 π -donatore





M

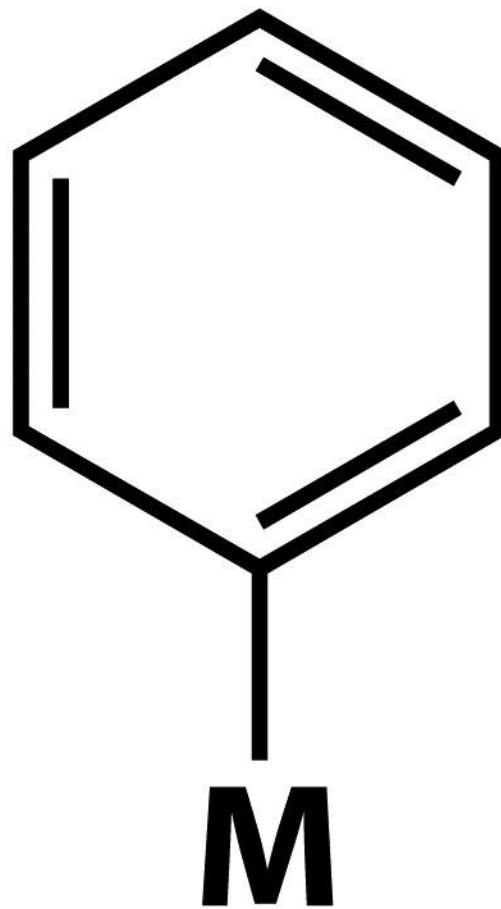
η^1 -alchenile

R



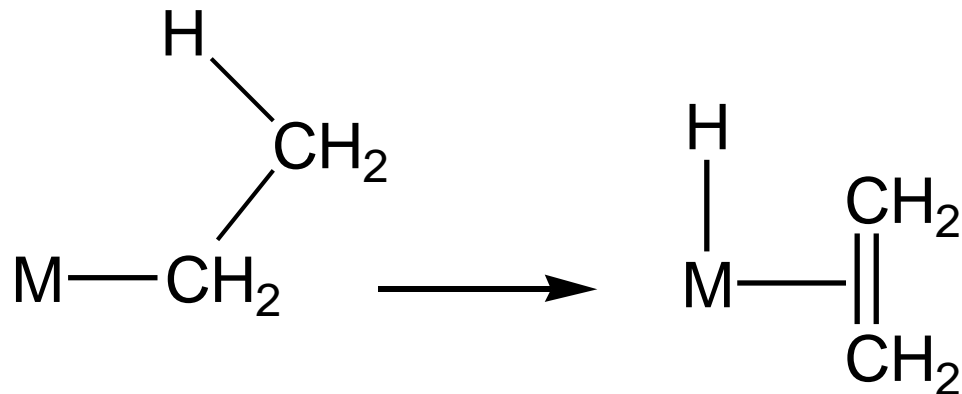
M

η^1 -alchinile



η^1 -arile

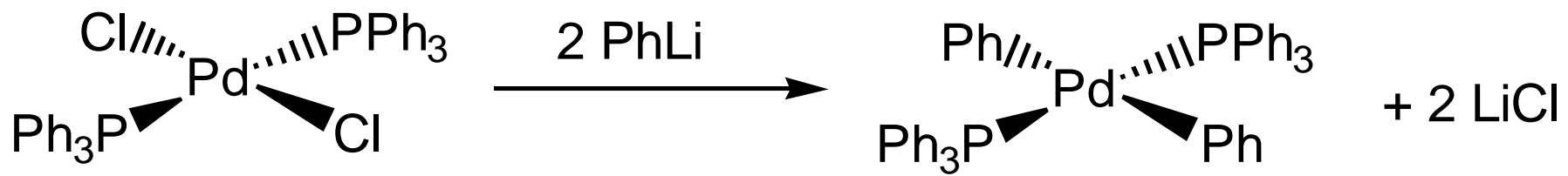
β -eliminazione di idruro



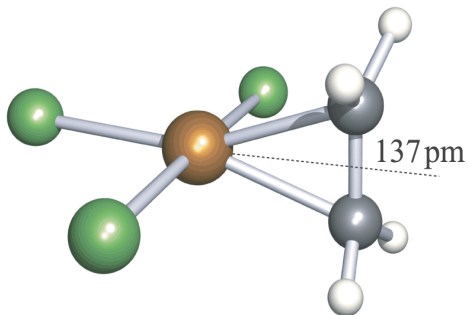
Gruppi stabili:

metile, benzile ($\text{CH}_3\text{C}_6\text{H}_5$), neopentile (CH_3CMe_3), e
trimetilsililmetile (CH_3SiMe_3)

Tipico processo di sintesi

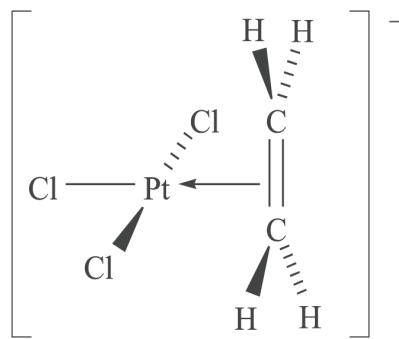


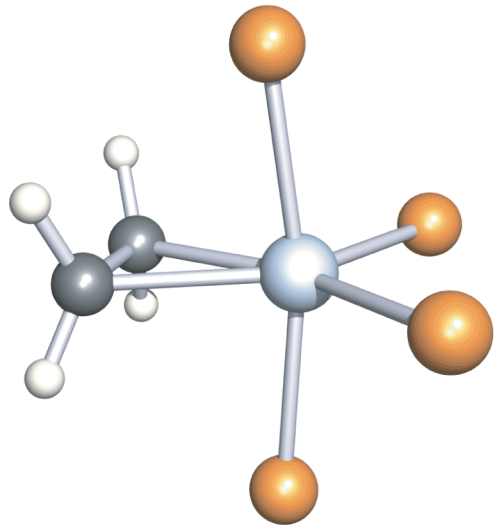
In alternativa si usano reattivi di Grignard



C–C = 137 pm vs 134 pm nell'etene

η^2 -alchene coordinazione *side-on*



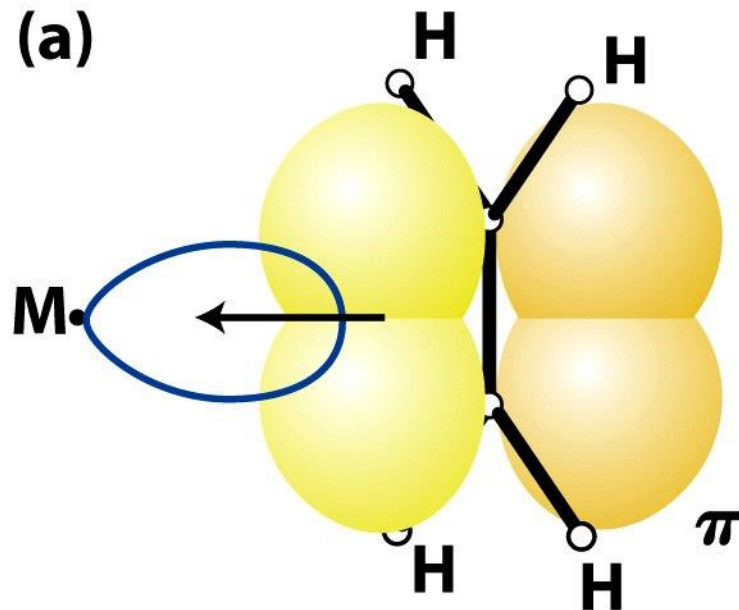


Struttura ai raggi-X di Ru(η^2 -C₂H₄)(PMe₃)

C-C = 144 pm vs 134 pm nell'etene

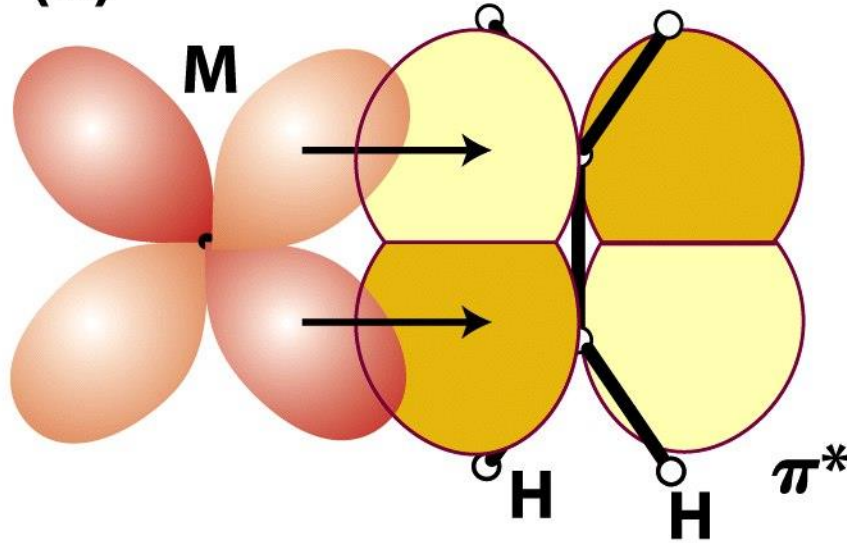
metallaciclopropano

(a)

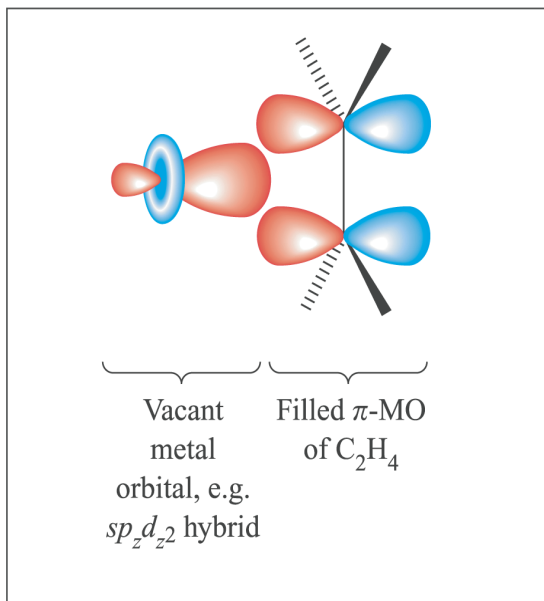
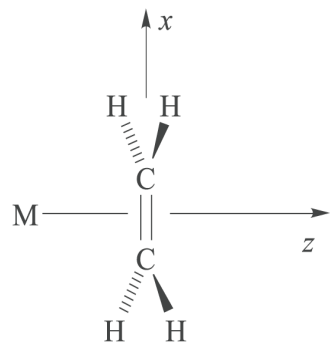


Orbitale di simmetria π per l'etene isolato, ma di simmetria σ quando l'etene è coordinato η^2 , cioè *side-on*

(b)

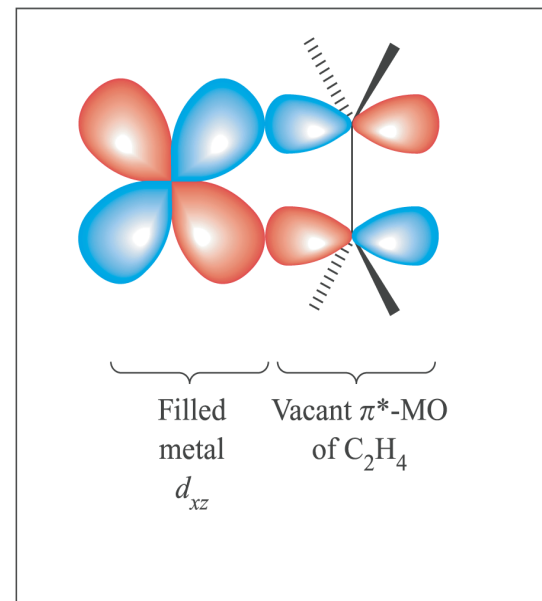


modello di Dewar – Chatt – Duncanson



Alkene-to-M donation

(a)

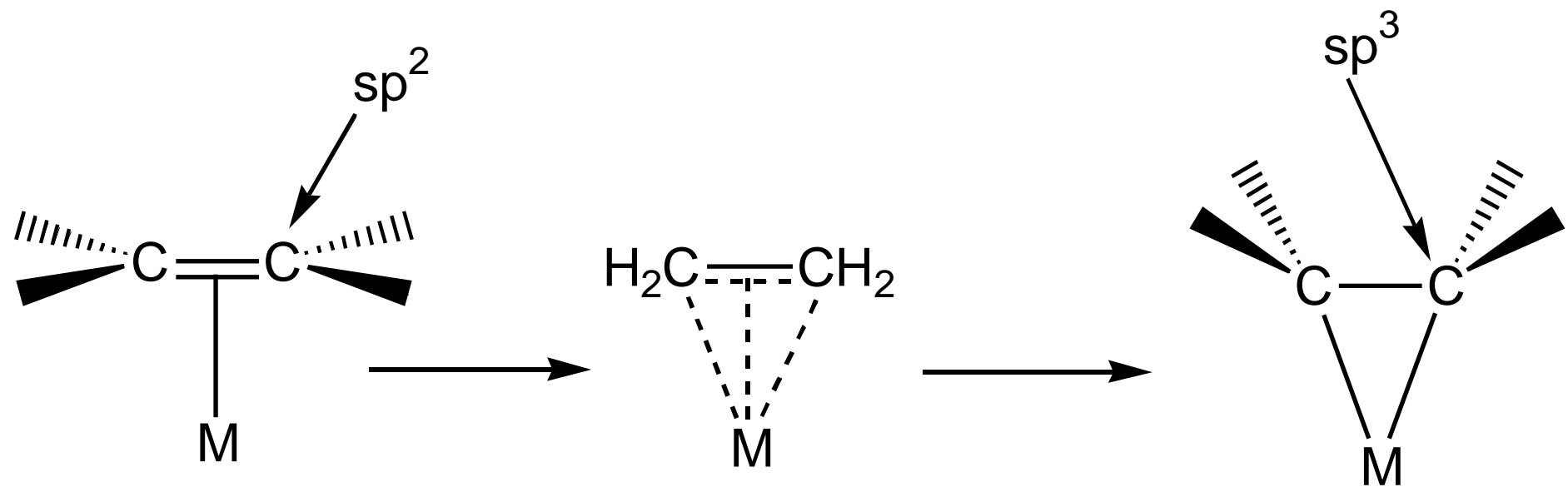


M-to-alkene back-donation

(b)

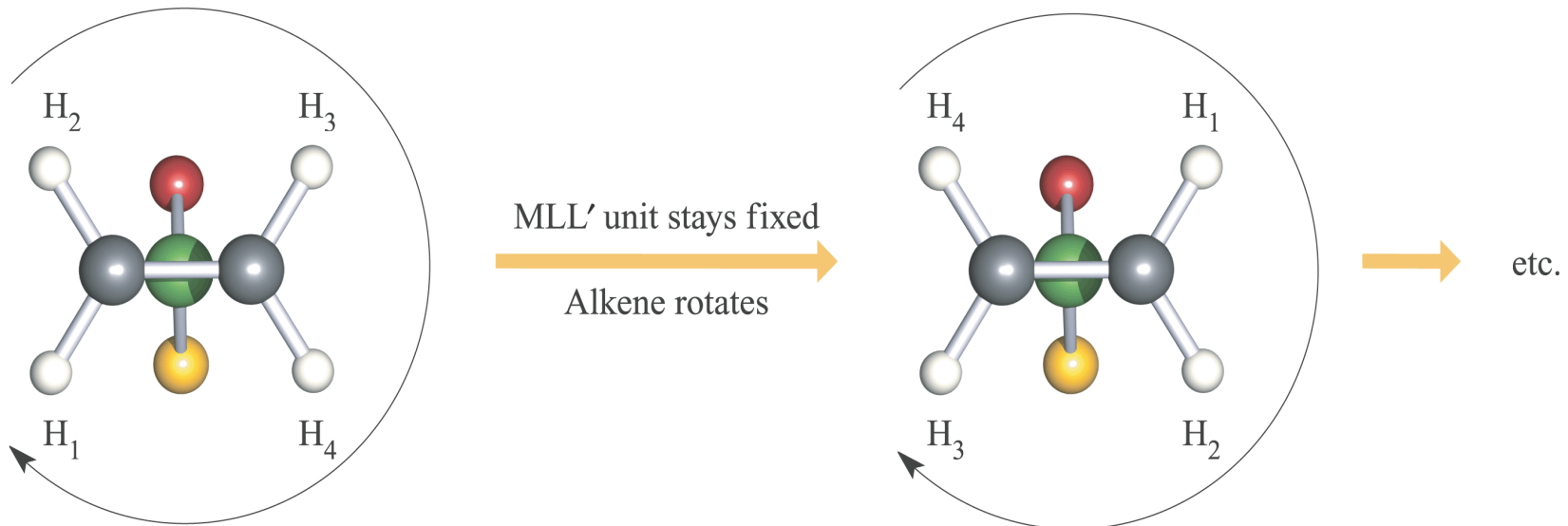
modello di Dewar – Chatt – Duncanson

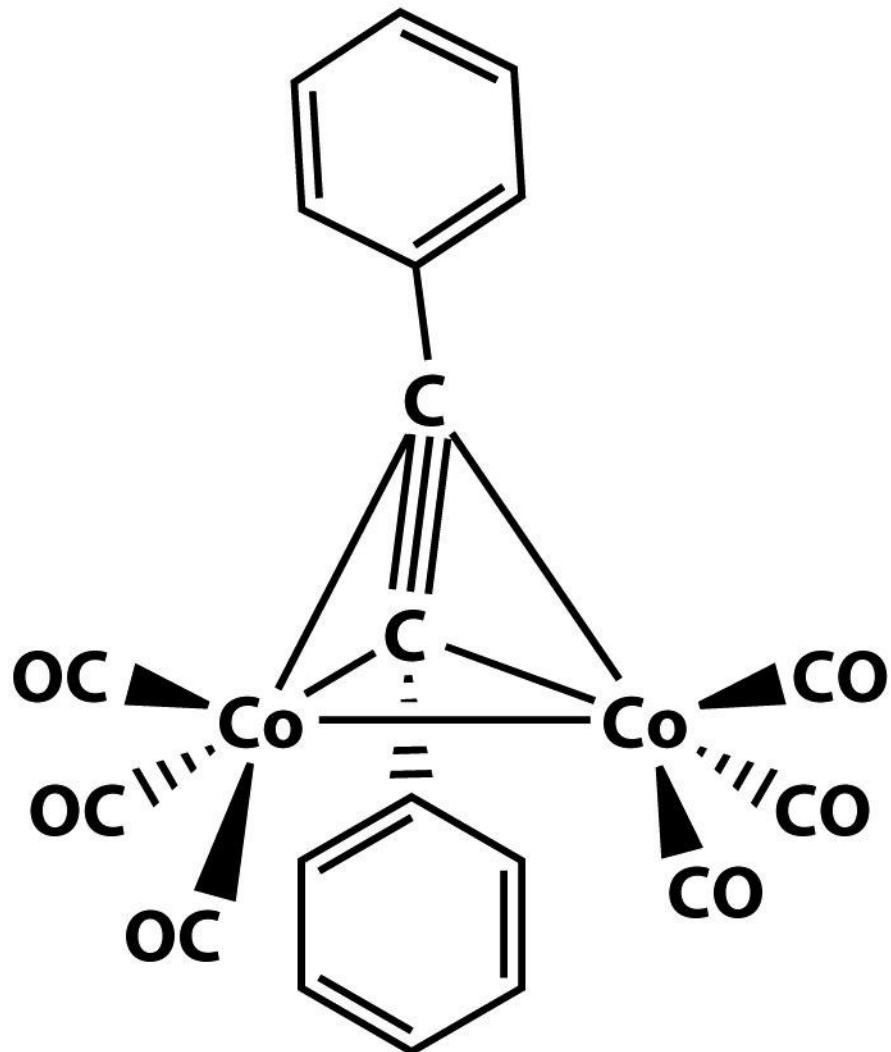
Retrodonazione π



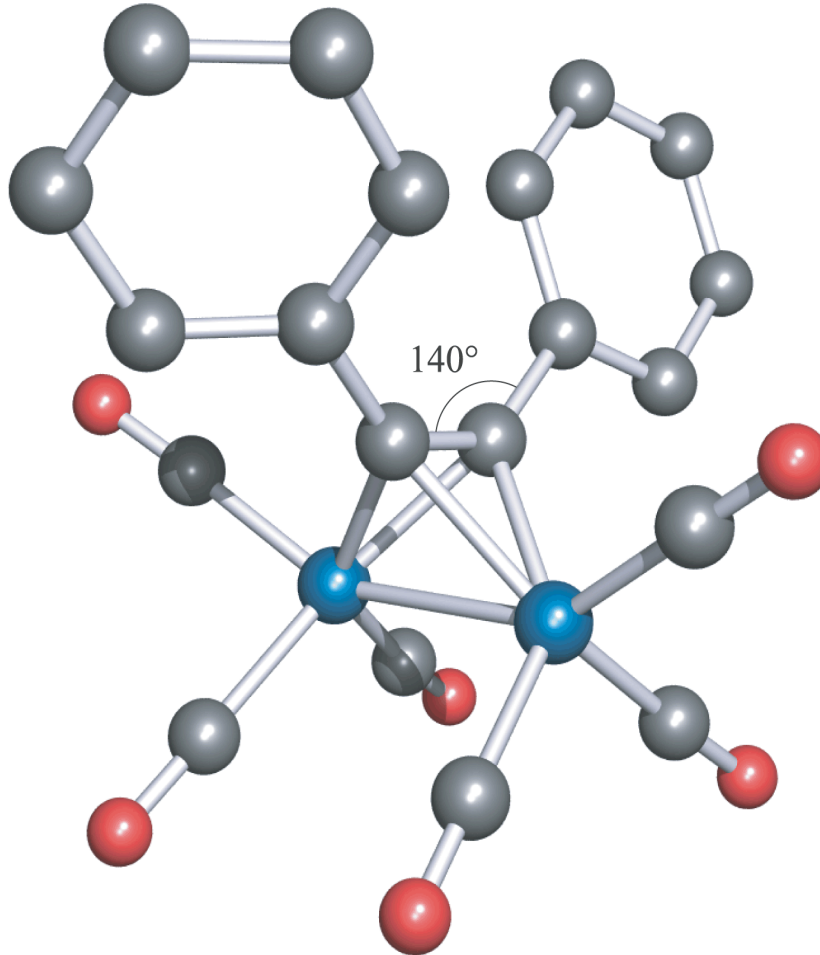
metallociclopropano

Flussionalità del legame η^2 -alchene





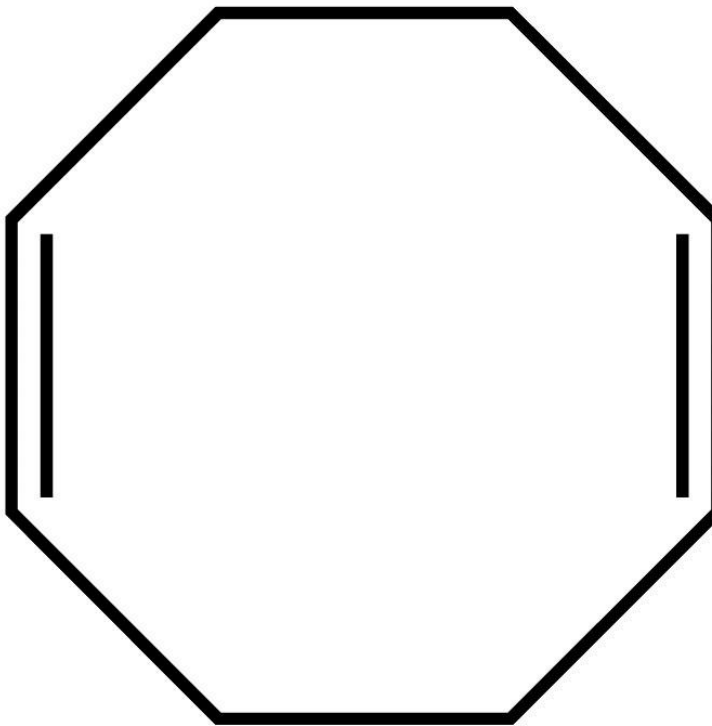
4-electron donor



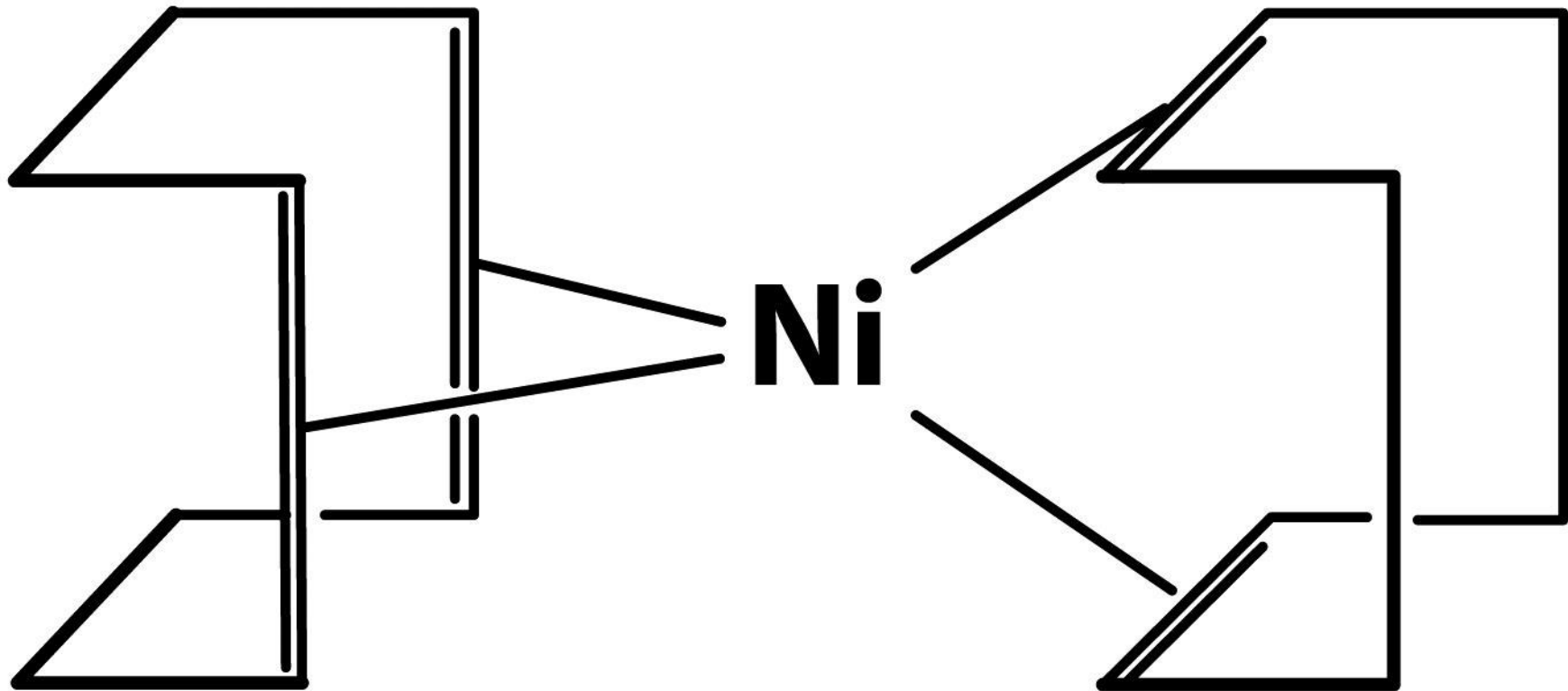
C—C in Co_2C_2 -unit = 136 pm vs 120 pm in C_2Ph_2

I due piani C_2Co , cioè i due legami η^2 , sono circa ortogonali

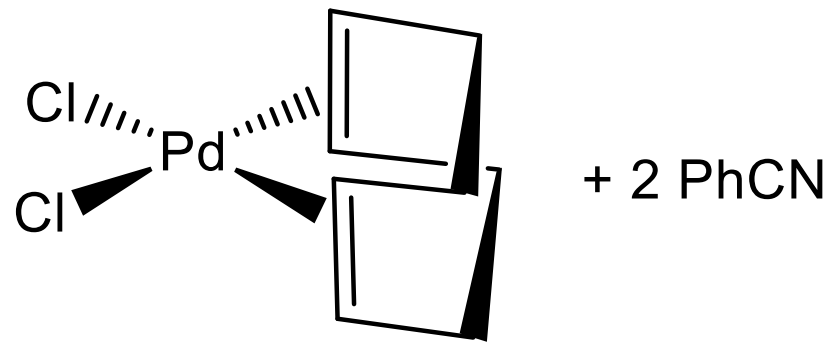
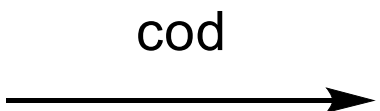
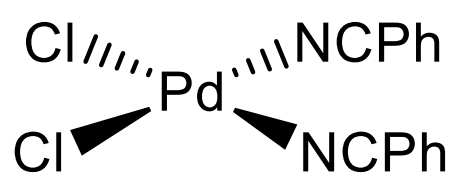
Dieni non-coniugati

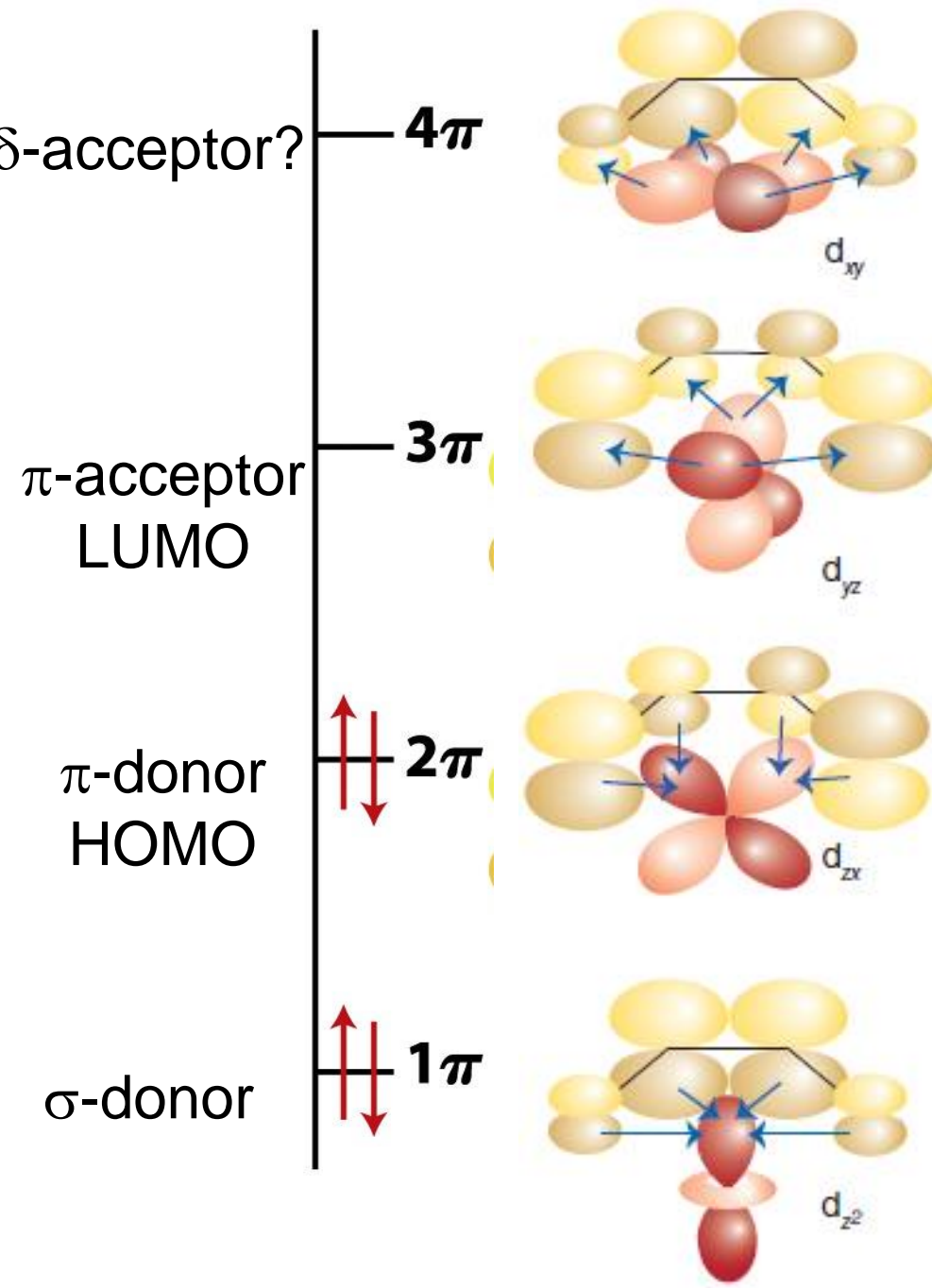


Cycloocta-1,5-diene, cod

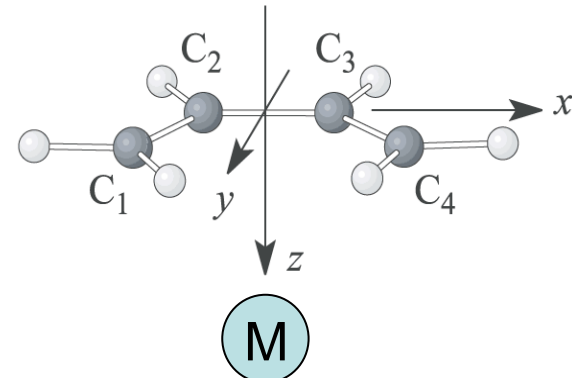


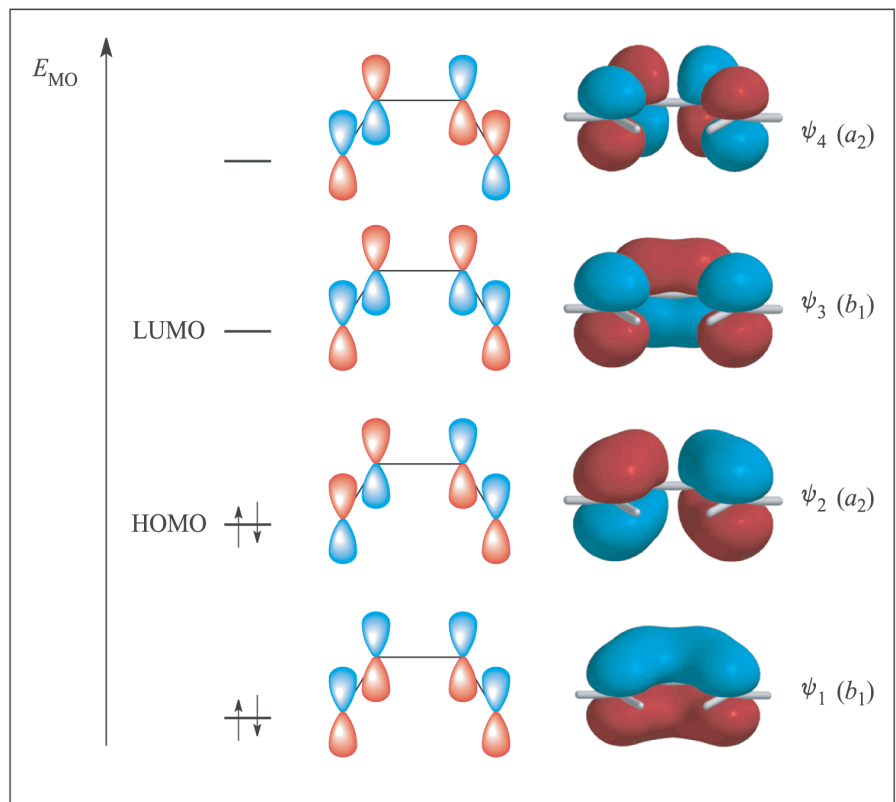
Ni(cod)₂



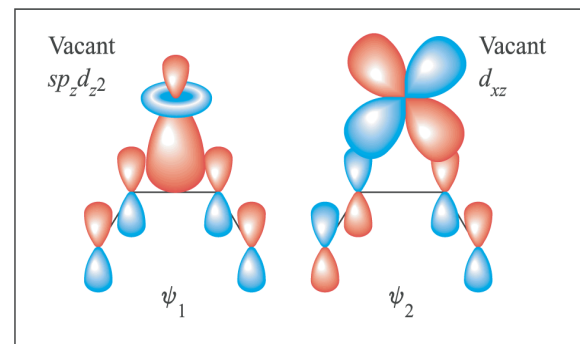
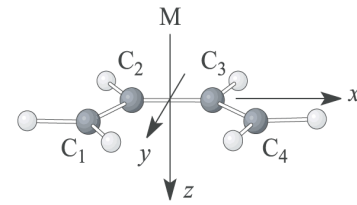


Butadiene -
giace nel piano xy ,
sopra al metallo

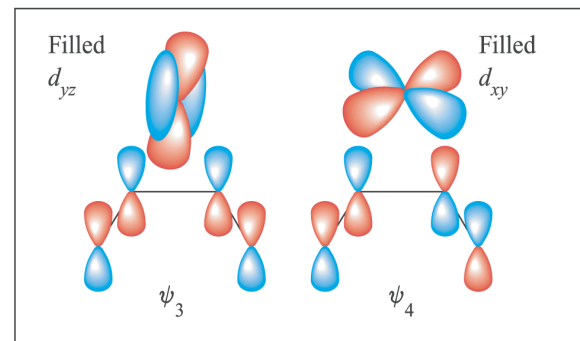


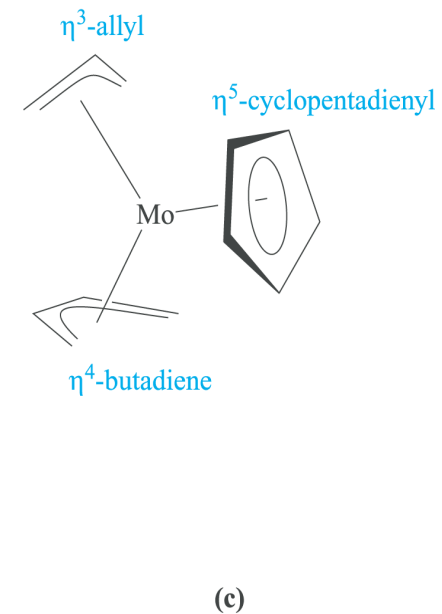
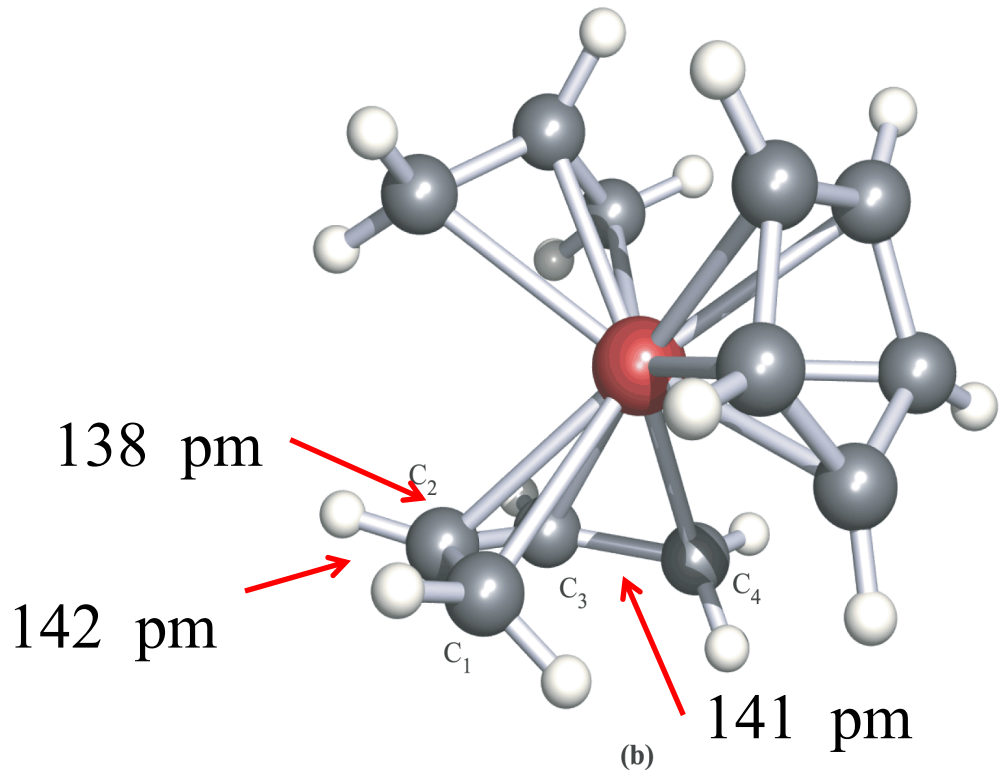


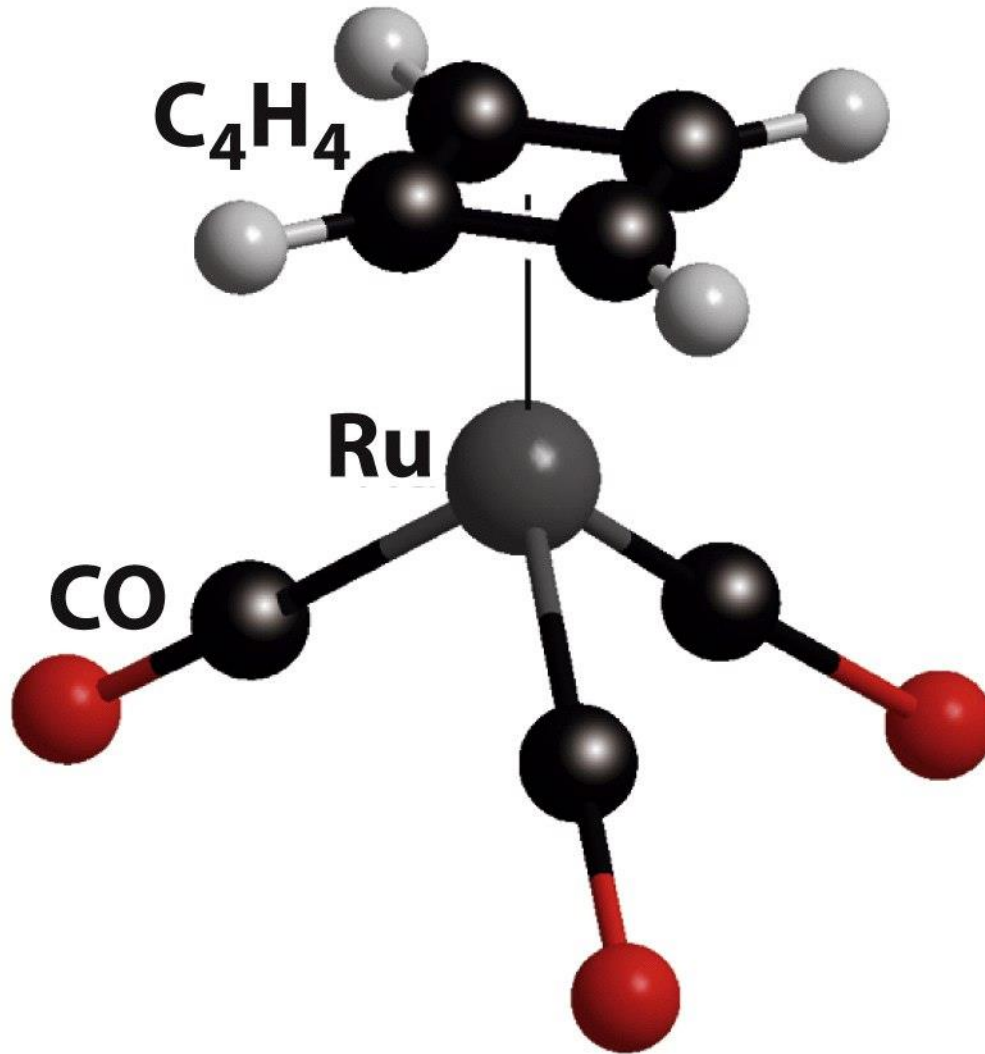
(a)



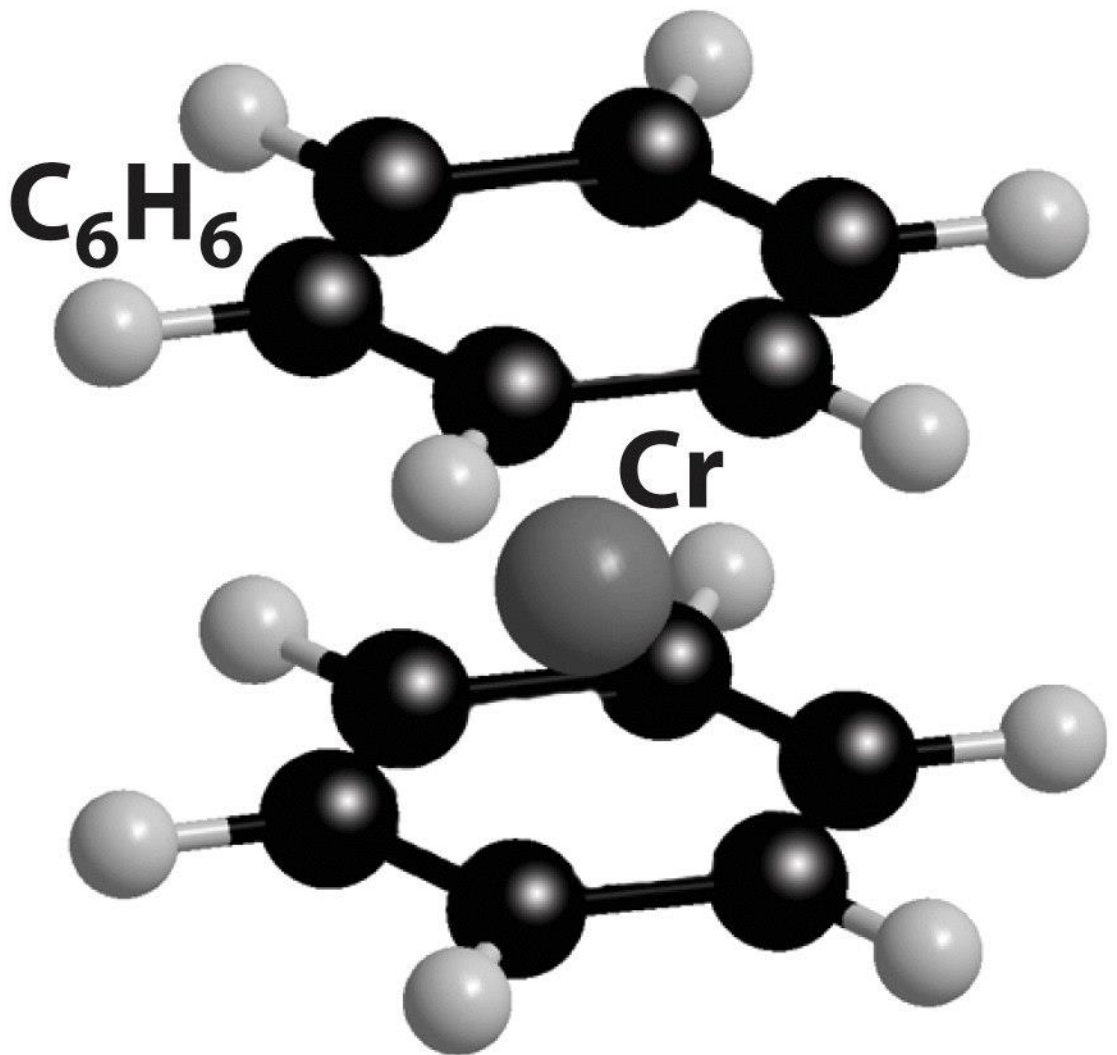
(b)

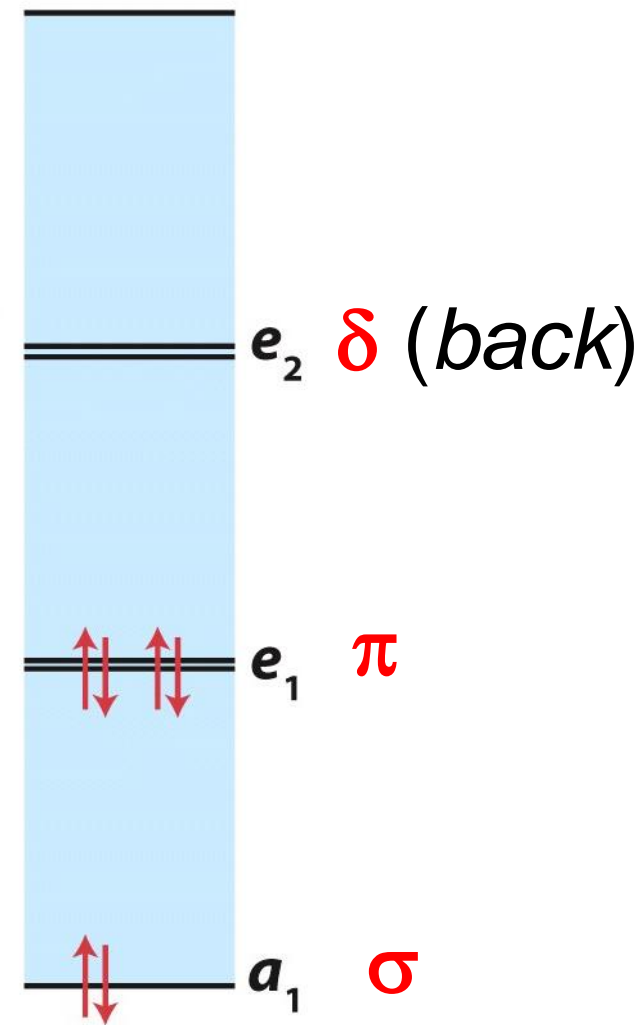
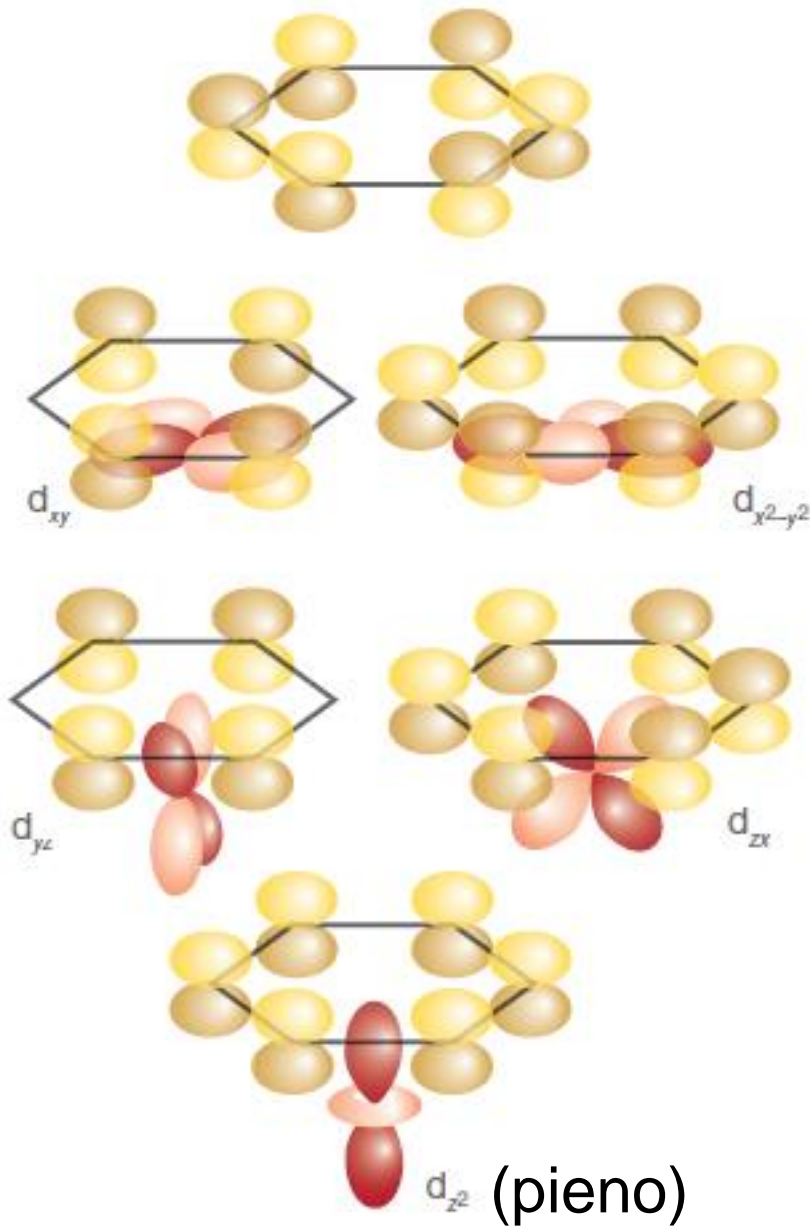


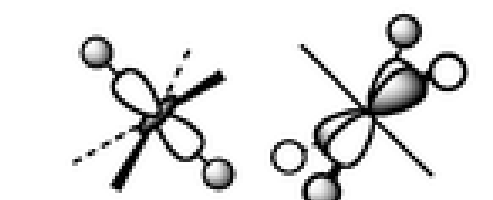
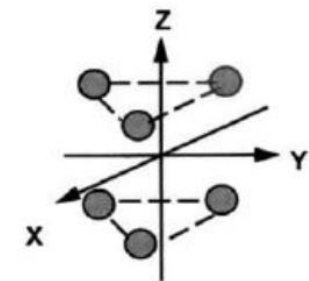
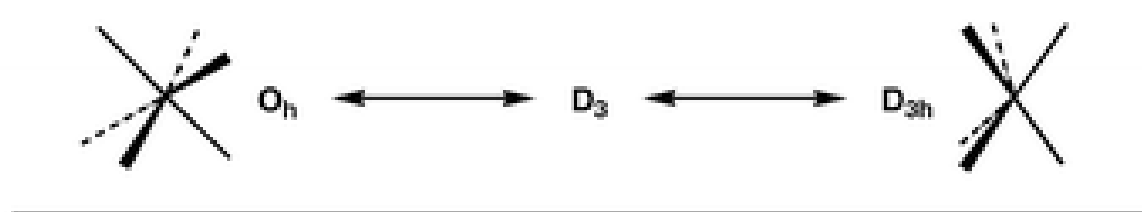




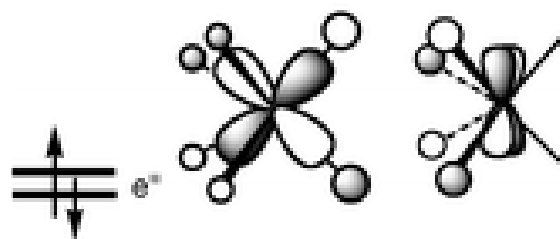
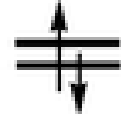
[Ru(C₄H₄)](CO)₃]



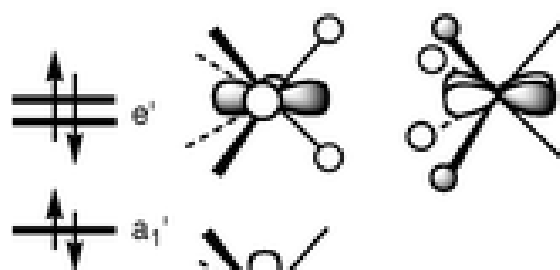




dz^2
 dx^2-y^2

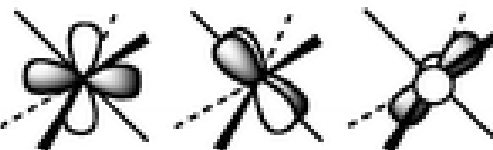


dxz
 dyz

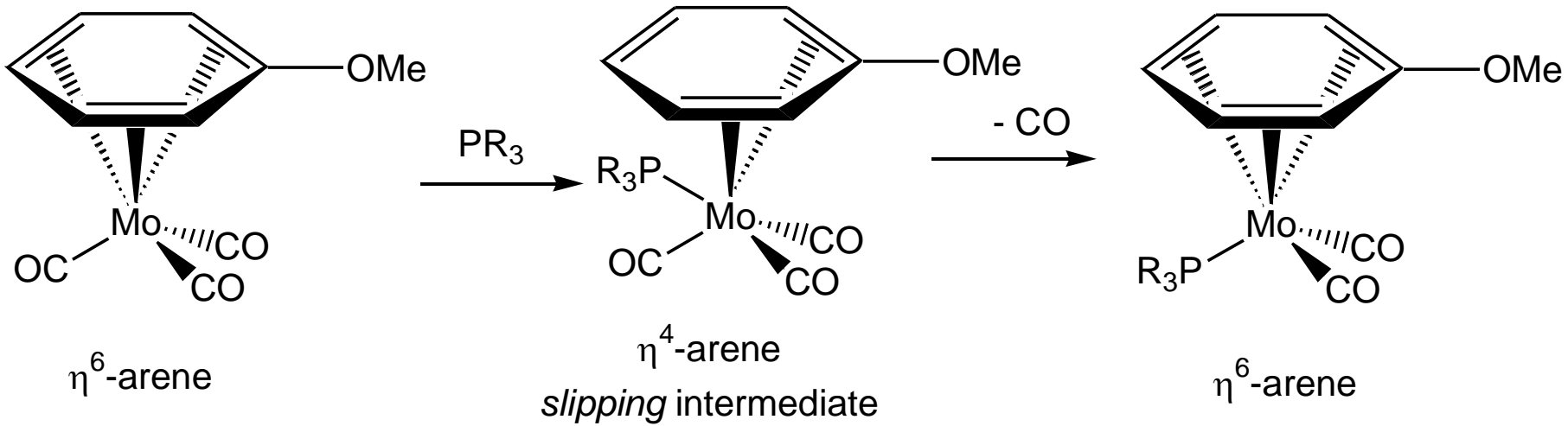
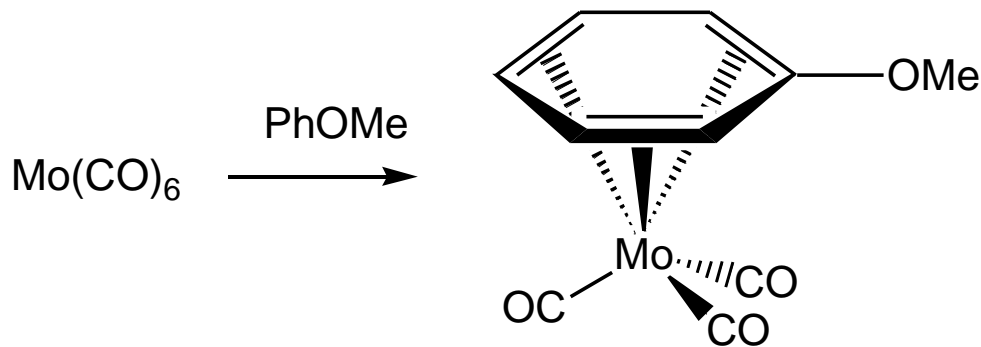


dxy ,
 dx^2-y^2

dxz
 dyz
 dxy

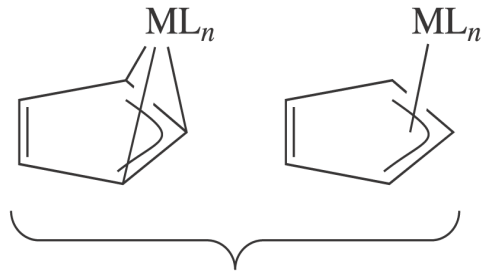


dz^2

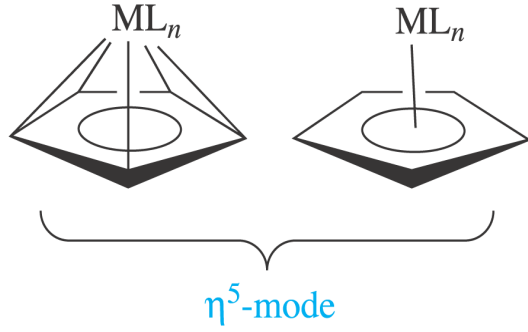




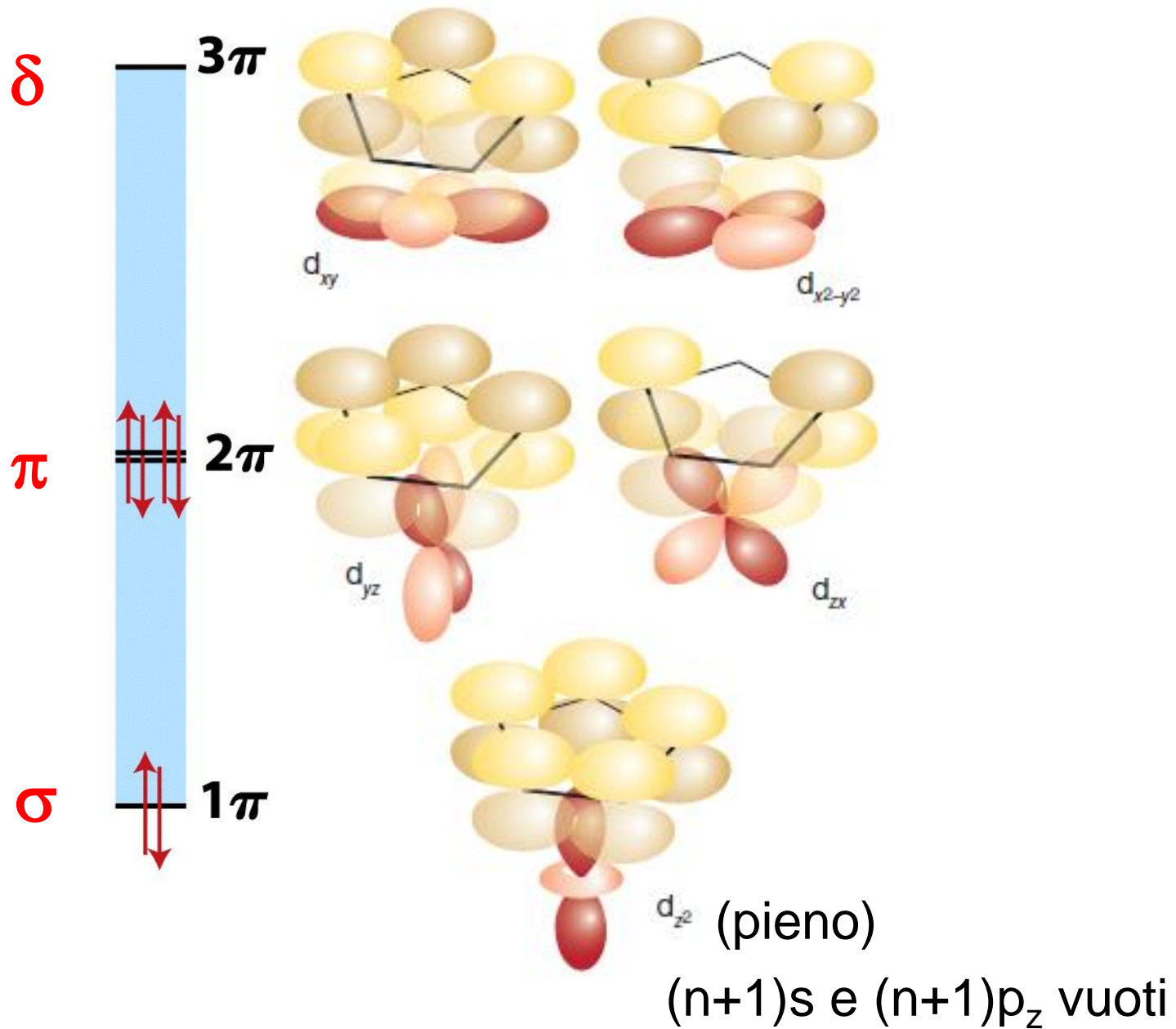
η^1 -mode

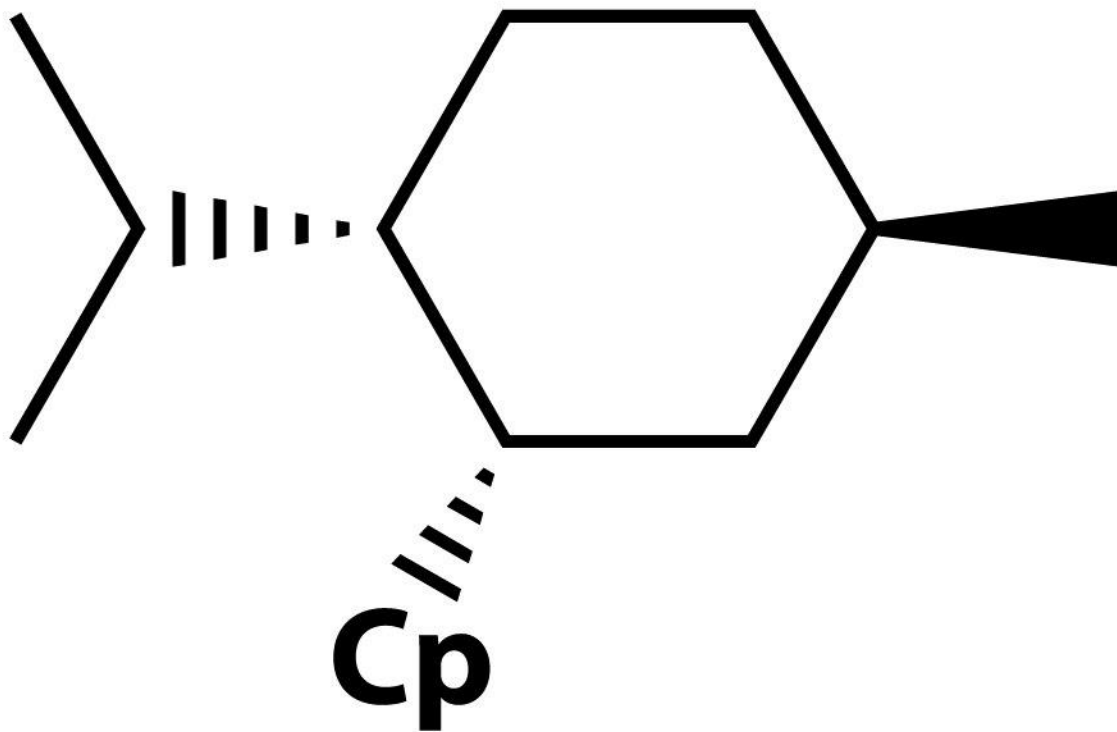


η^3 -mode



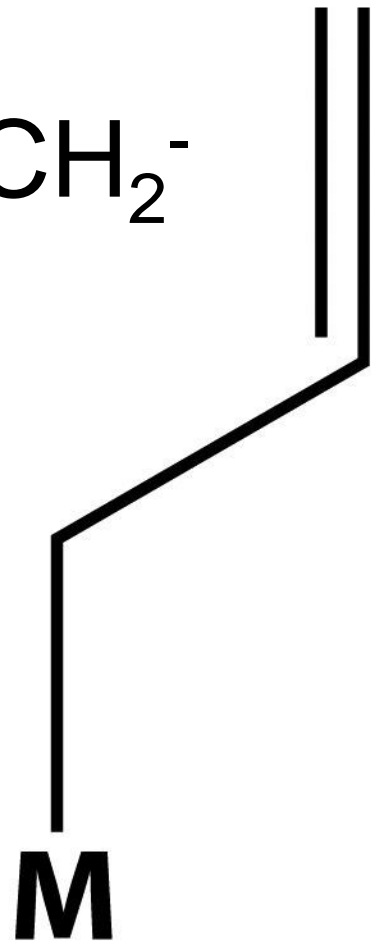
η^5 -mode



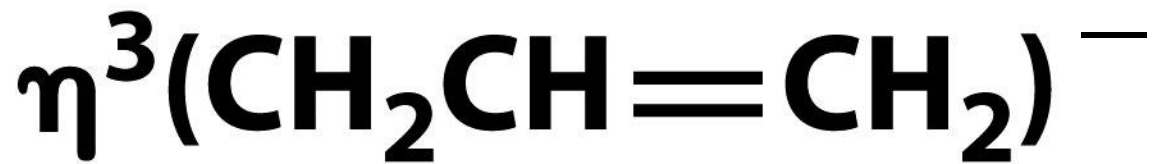
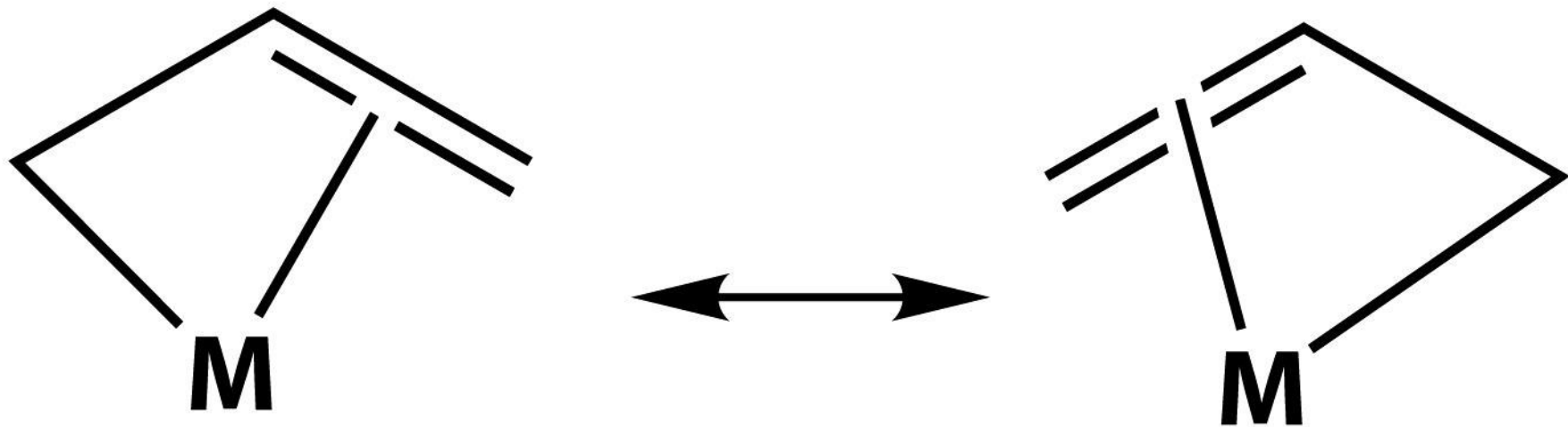


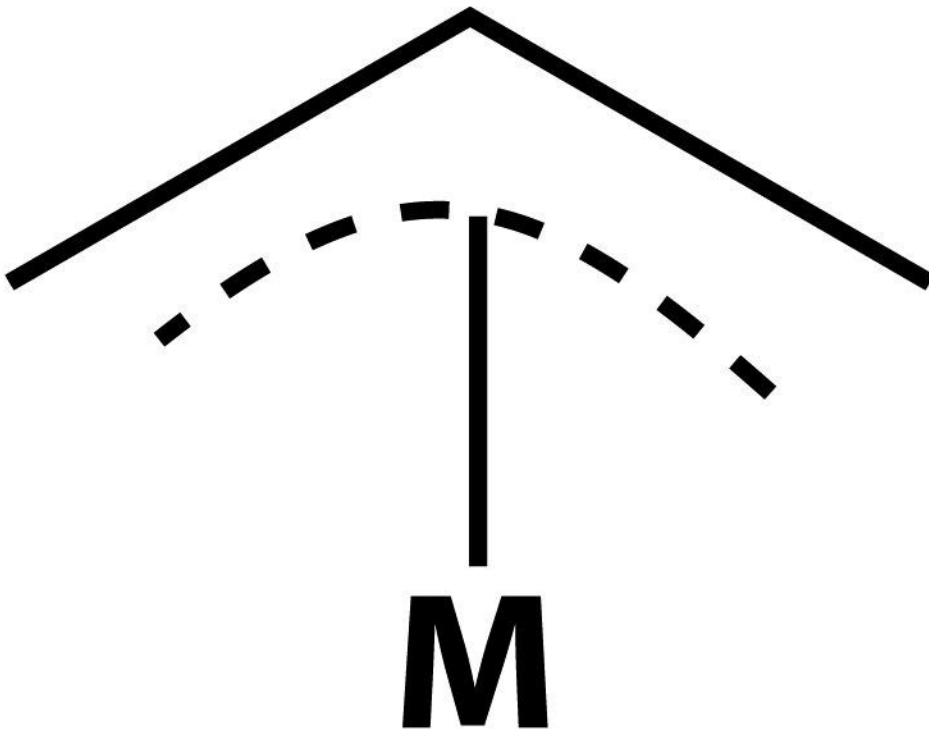
neo-Menthylcyclopentadienyl

Allile, $\text{CH}_2=\text{CH}-\text{CH}_2^-$



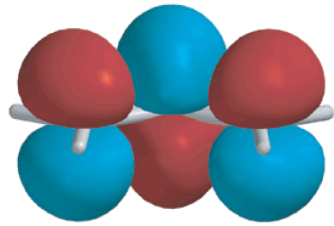
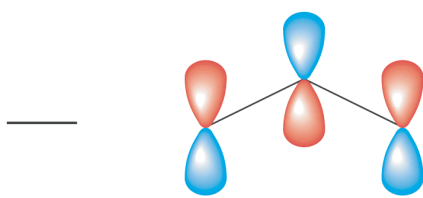
$\eta^1-(\text{CH}_2\text{CH}=\text{CH}_2)$



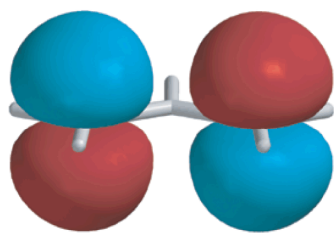
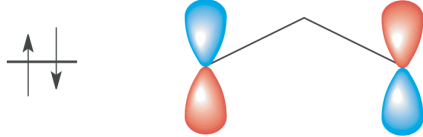


$\eta^3\text{-}(\text{CH}_2\text{CH}=\text{CH}_2)^-$

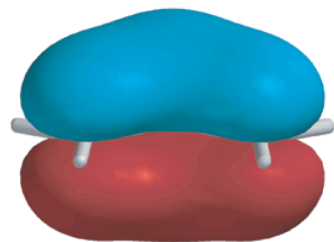
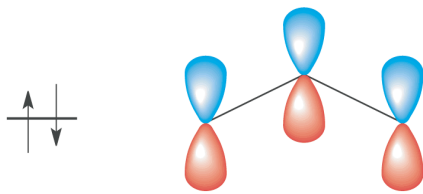
E_{MO}



$\psi_3 (b_1)$ Antibonding



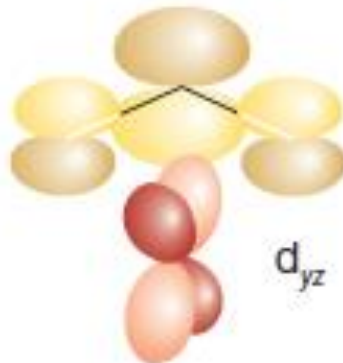
$\psi_2 (a_2)$ Non-bonding



$\psi_1 (b_1)$ Bonding

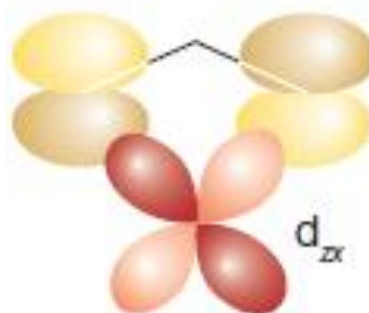
Accettore π

3π



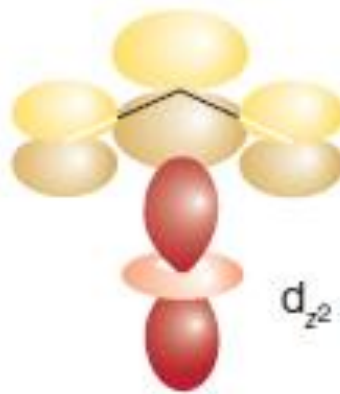
Donatore π

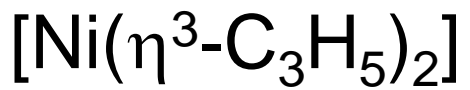
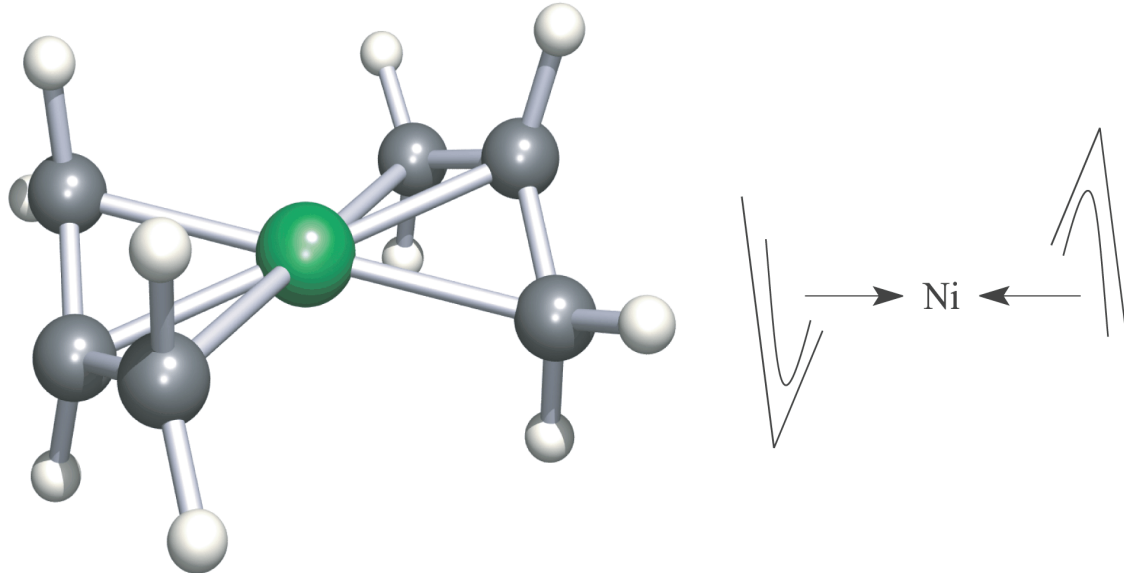
2π

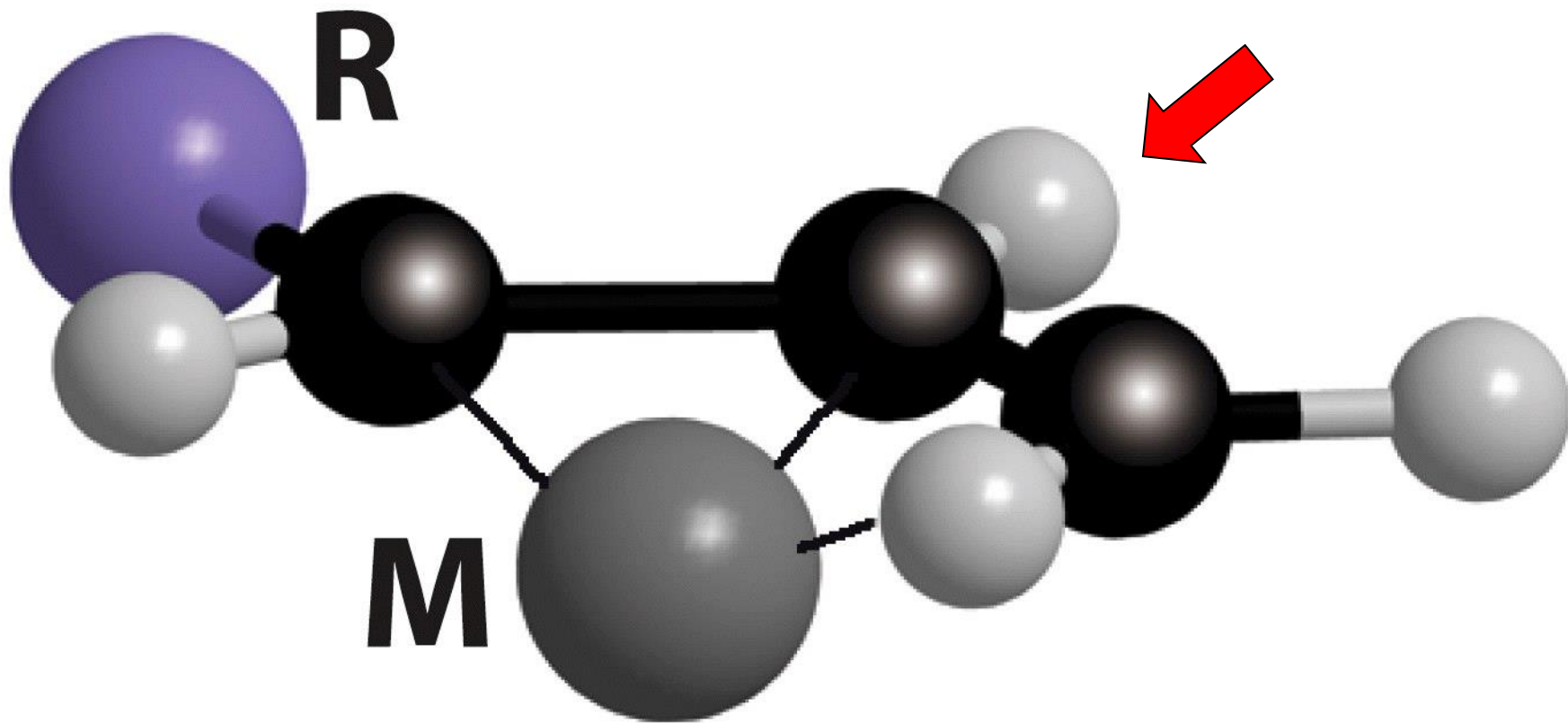


Donatore σ

1π

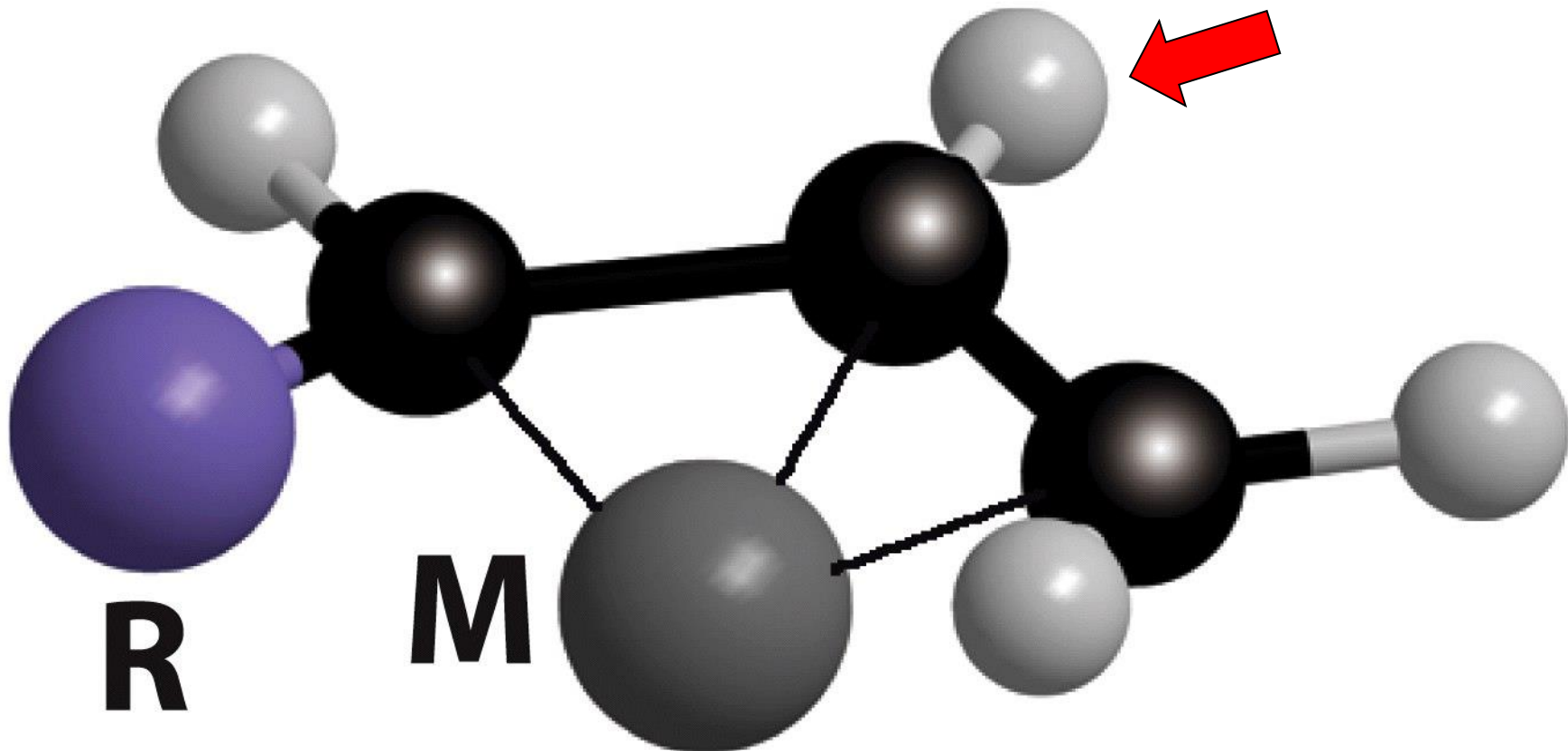






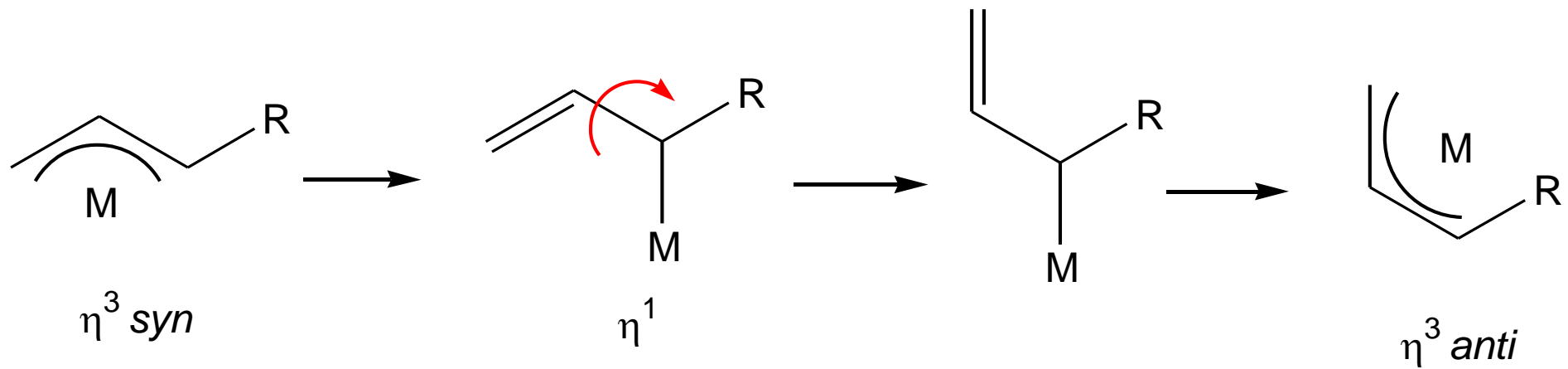
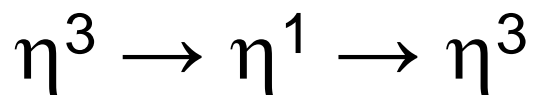
M

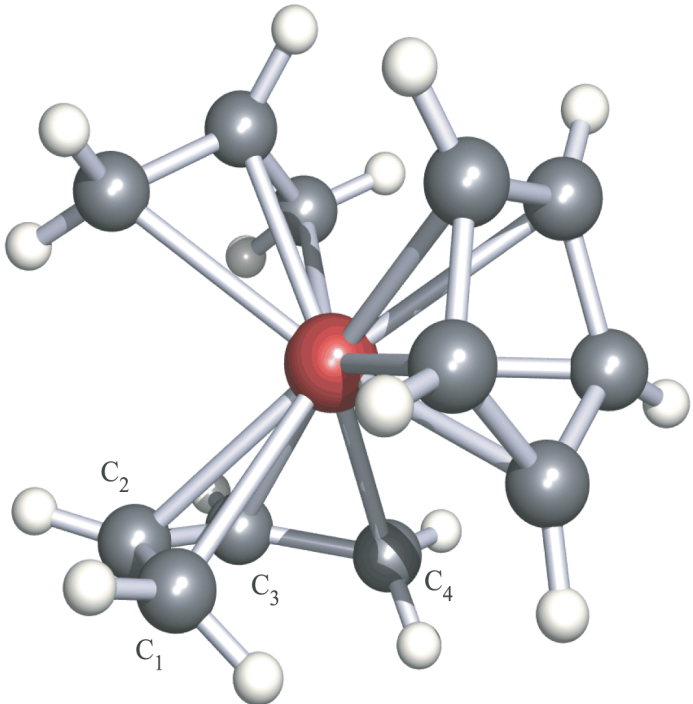
syn



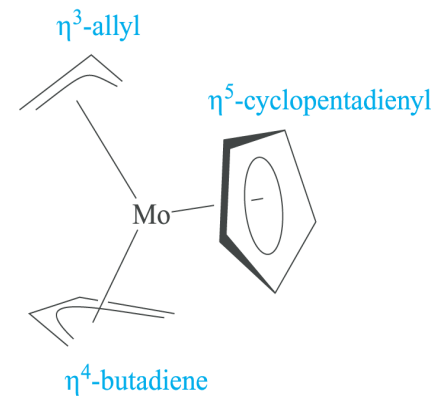
anti

Meccanismo di scambio *syn-anti*





(b)



(c)

