



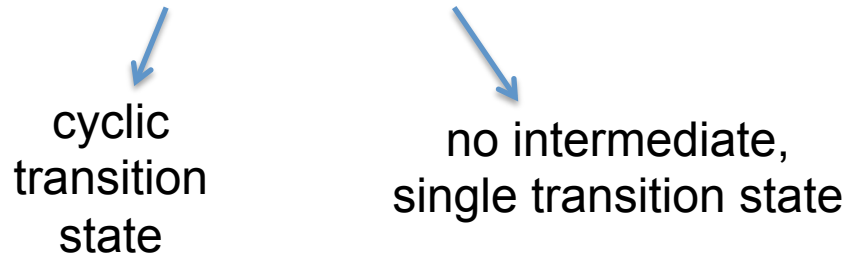
Chem 634

## Pericyclic Reactions

Reading:  
CS-B Chapter 6  
Grossman Chapter 4

## *Pericyclic Reactions*

Definition: Continuous, concerted reorganization of electrons



Bond breaking & bond making occur at the same time.

Can be synchronous (equal extent of breaking & making in TS) or asynchronous (unequal extent of breaking & making in TS).

Fukui & Hoffmann: Nobel Prize in Chemistry, 1981, "for... their theories, developed independently, concerning the course of chemical reactions" (Woodward dies in 1979)

## *5 Types*

1. Electrocyclic
2. Cycloadditions
3. Sigmatropic
4. Chelotropic
5. Group Transfer

## 3 Theories

All 3 theories are correct!

### 1. **Woodward–Hoffmann:** Conservation of Orbital Symmetry

- 1<sup>st</sup> historically
- Uses *correlation diagrams*

### 2. **Fukui:** Frontier Molecular Orbital Theory

- Easier than Woodward–Hoffmann (usually)
- Based on HOMO/LUMO interactions

### 3. **Dewar–Zimmerman:** Aromatic Transition State

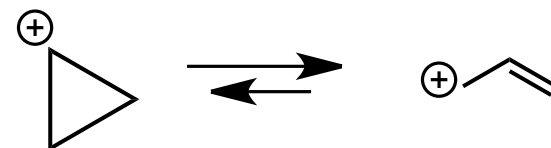
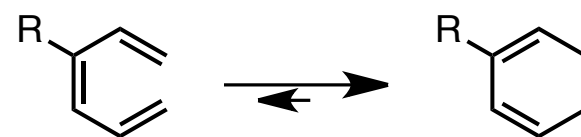
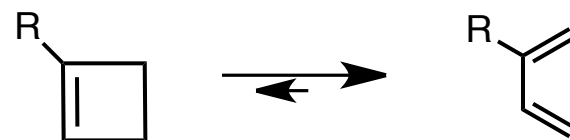
- Easiest to apply for all reaction types, but not intuitive to understand why it's valid

## *3 Things Matter*

1. Number of electrons involved
2. Stereospecificity
3. Conditions: heat ( $\Delta$ ) vs. light ( $h\nu$ )

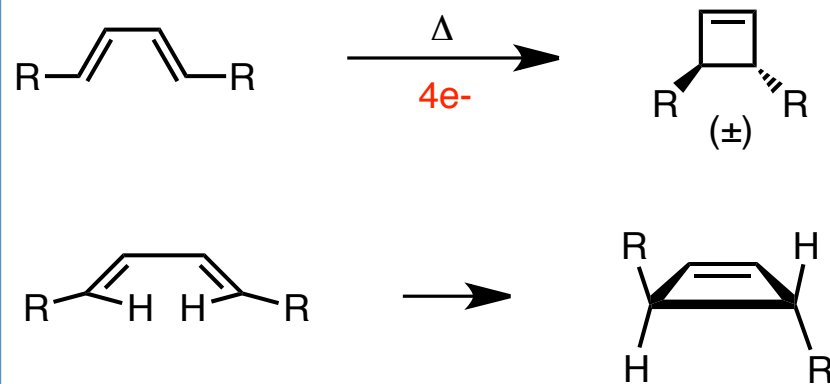
## Type 1: Electrocyclic Reactions

- Ring openings and closures
- Exchange  $\pi$ -bond for  $\sigma$ -bond
- Classified by number of electrons

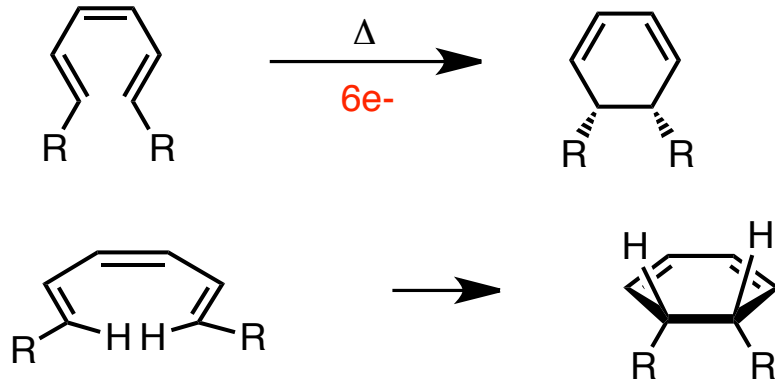


*Diastereoselectivity – Observations*

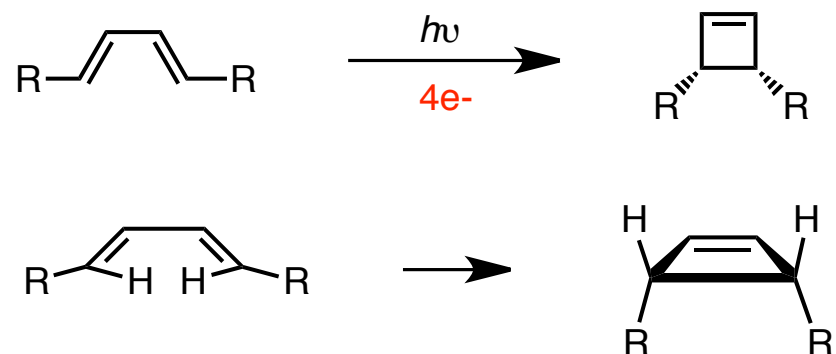
Case 1:



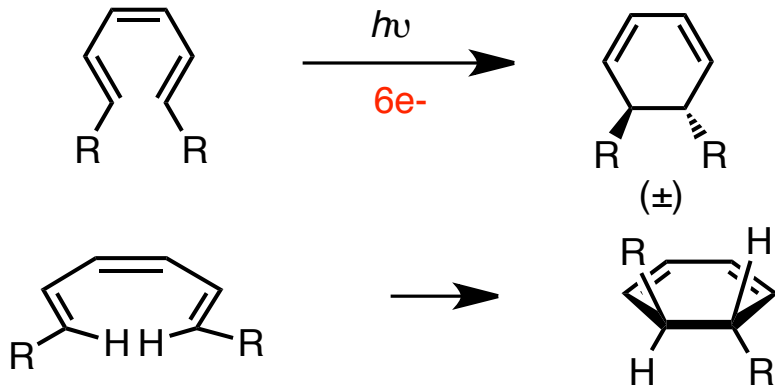
Case 3:



Case 2:



Case 4:



*General Phenomenon... Woodward–Hoffmann Rules*

Number of electrons	Thermal	Photochemical
$4n$	Con	Dis
$4n+2$	Dis	con

( $n = \text{integer}$ )

6 points for a touchdown  $\rightarrow$  6e-, thermal, disrotatory

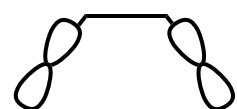
... *But why???*

## *Theory #1: Woodward–Hoffmann Correlation Diagrams*

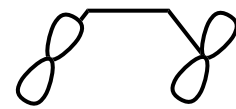
*Angew. Chem. Int. Ed.* **1969**, *8*, 781.

- Consider all molecular orbitals (MO's) involved
- Consider symmetry of MO's in starting material, product, *and transition state*.
- Orbitals of different symmetry can cross (orthogonal orbitals).
- Orbitals of same symmetry *cannot* cross (extreme energetic cost).
- We are about orbitals where electrons end up.

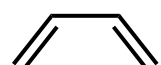
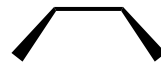
*Example of W–H Correlation Diagrams*



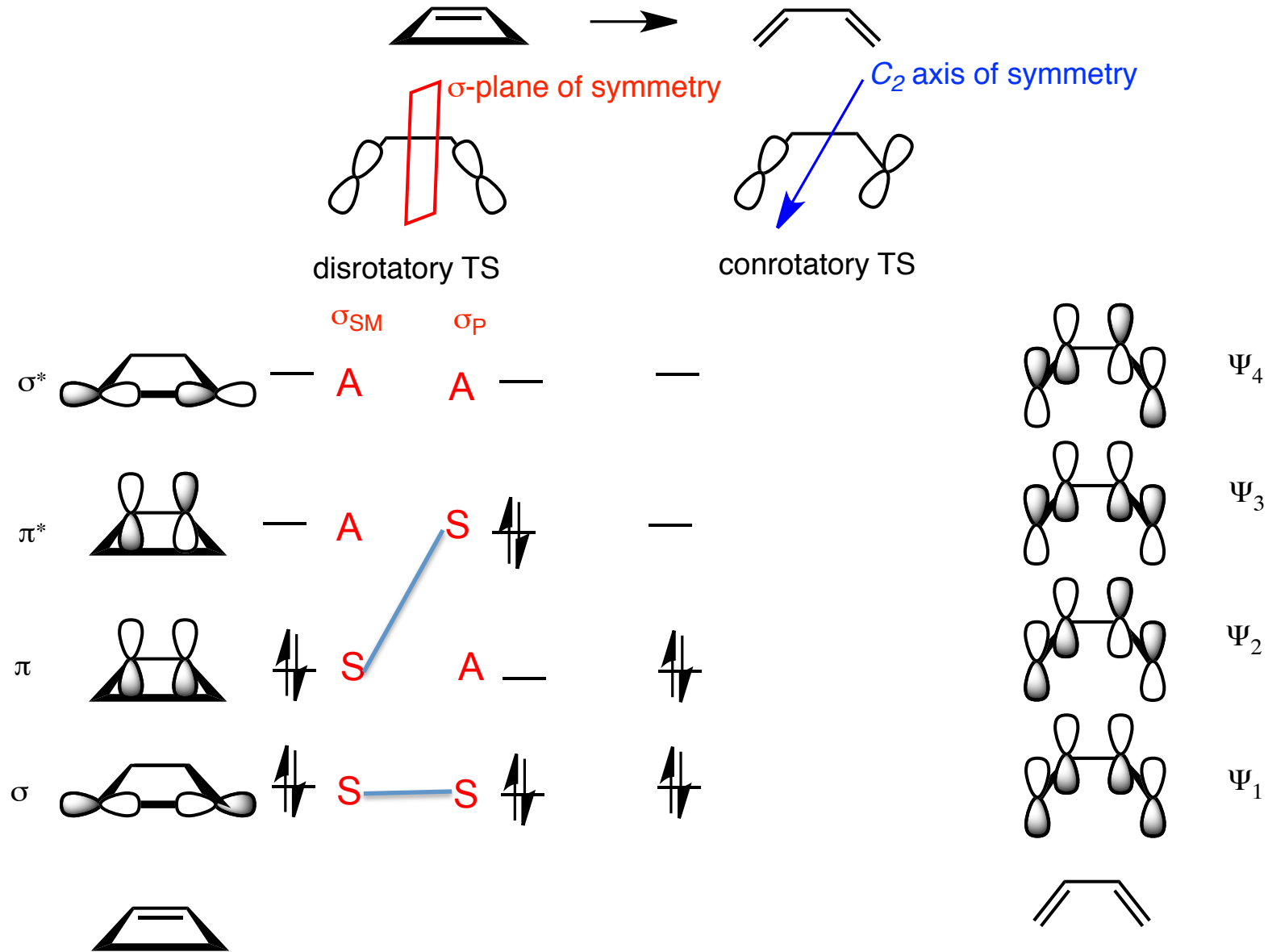
disrotatory TS



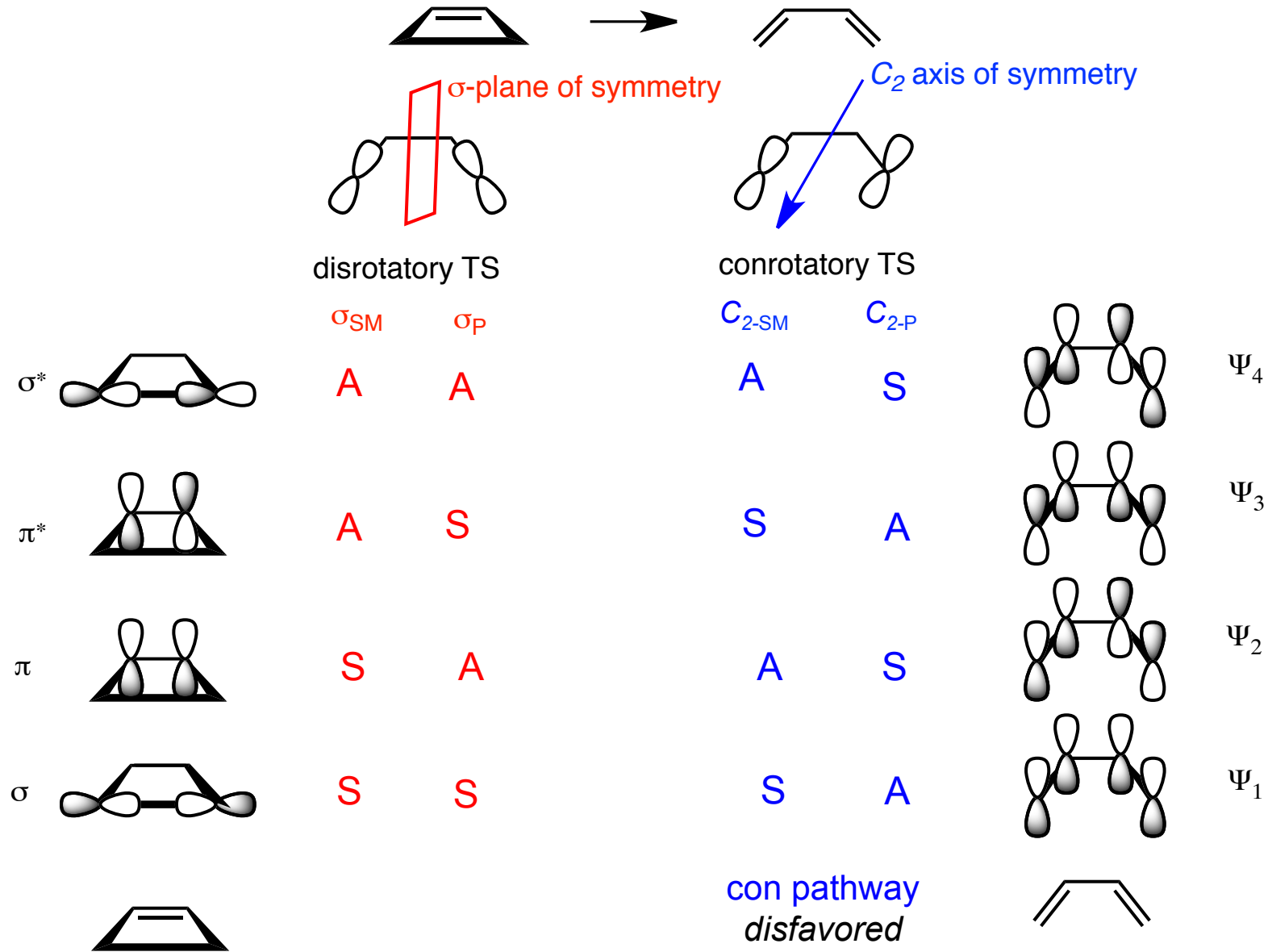
conrotatory TS

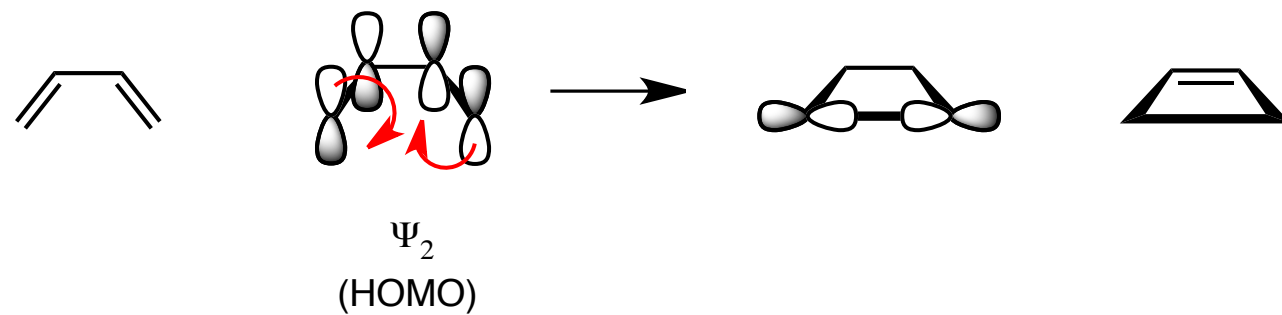
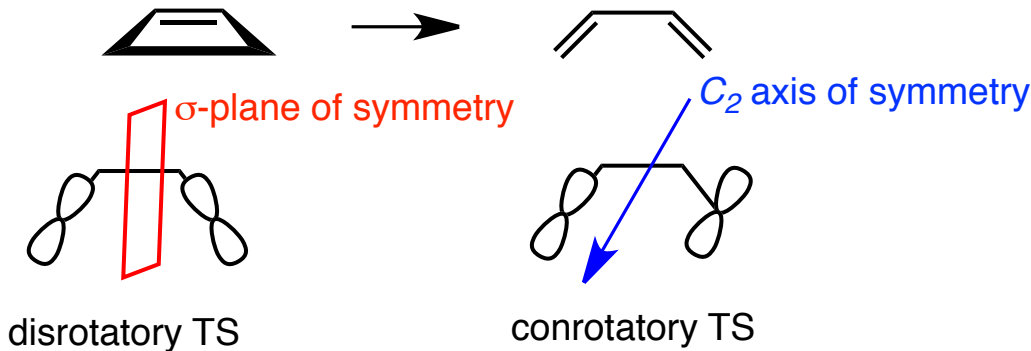


## Example of W–H Correlation Diagrams: Thermal Conditions



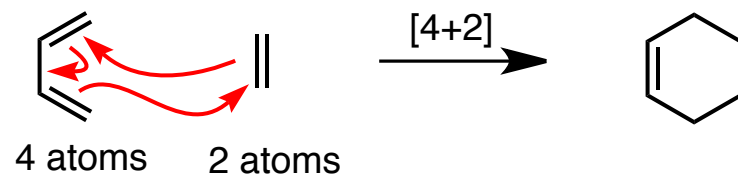
## Example of W–H Correlation Diagrams: Photochemical Conditions



*W–H Conservation of Orbital Symmetry Shortcut*

## Cycloadditions & Cycloreversions

- Union of 2  $\pi$ -systems
- Exchange  $\pi$ -bonds for  $\sigma$ -bonds
- Classified by  $[m+n]$ , m & n = # of conjugated atoms in each  $\pi$ -system



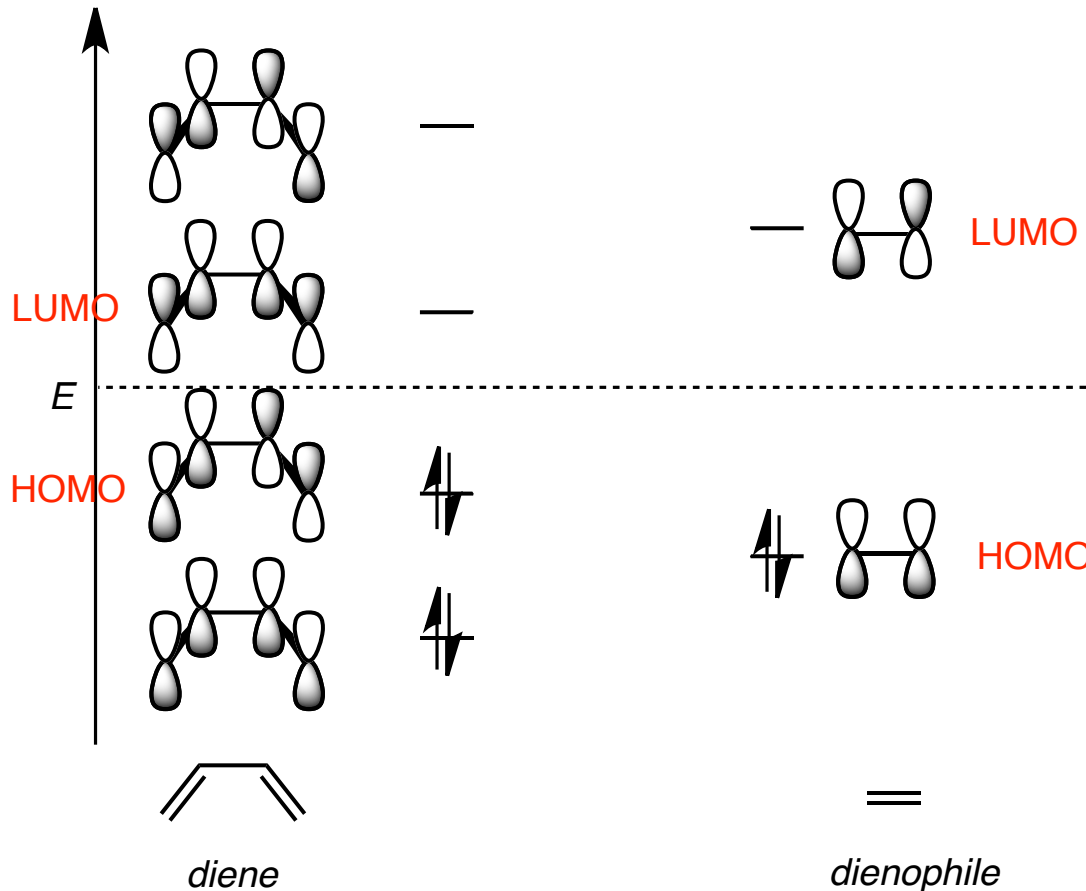
Diels–Alder Reaction!

Note: 6 e-

Great way to make cyclohexenes & cyclohexanes

## Fukui: Frontier Molecular Orbital (FMO) Theory

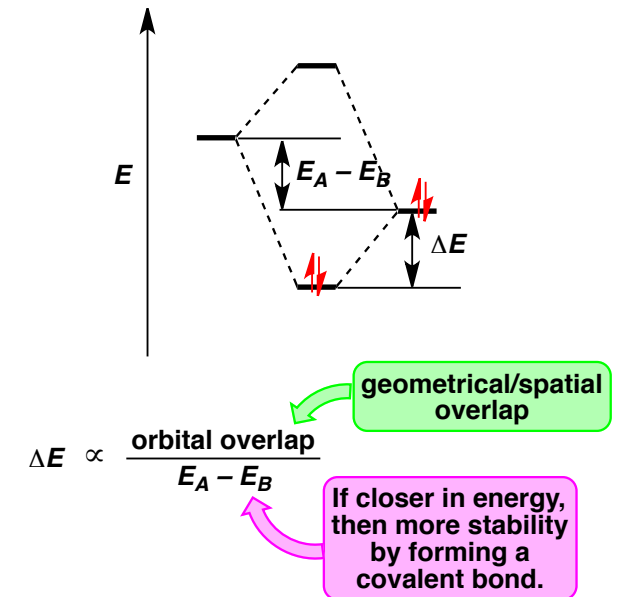
The idea: Use FMO's (HOMO + LUMO)



In this case, it doesn't matter... HOMO/LUMO gaps are the same.

*nonbonding level*

Which HOMO & LUMO?



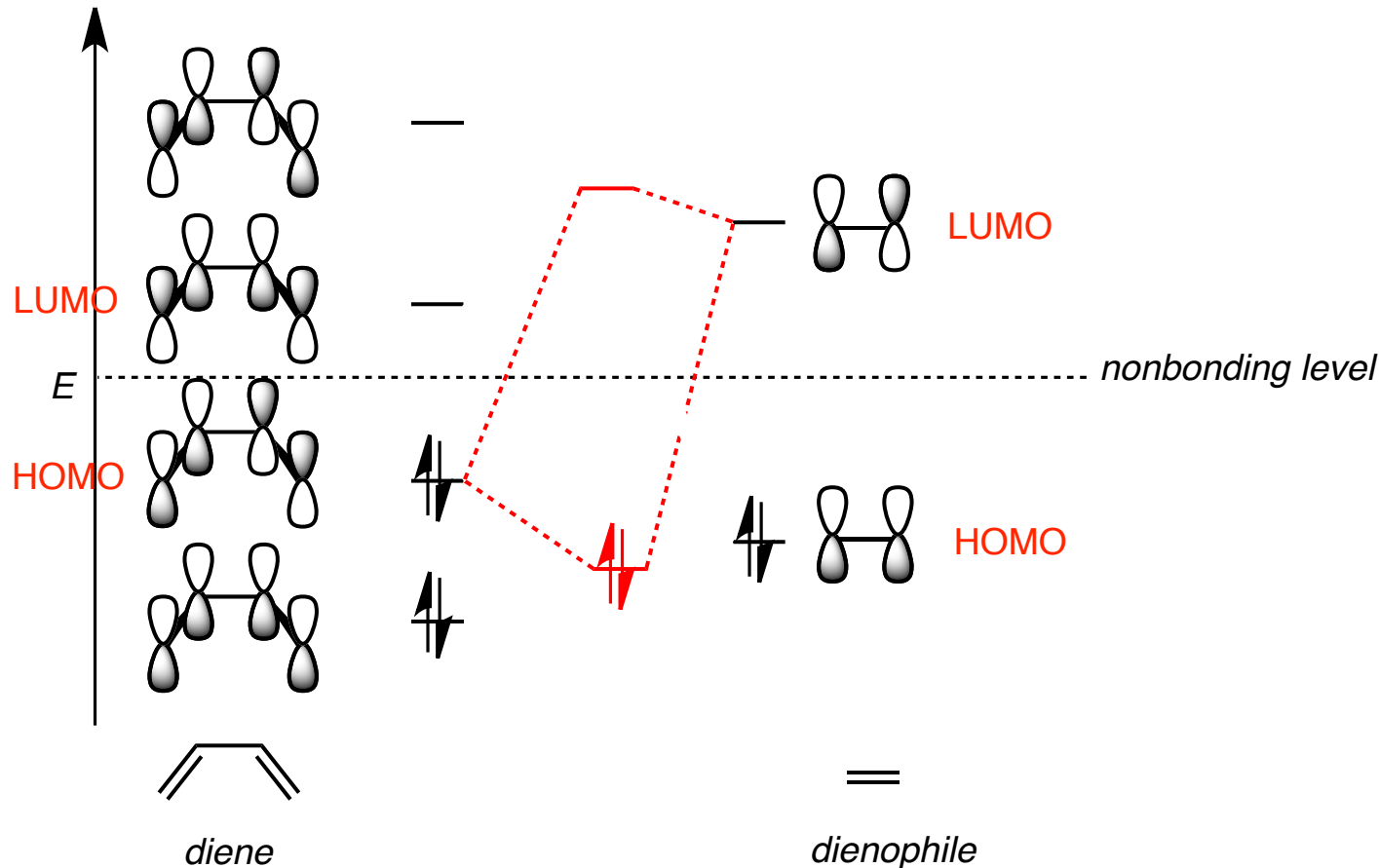
## *Types of Diels–Alder Reactions*

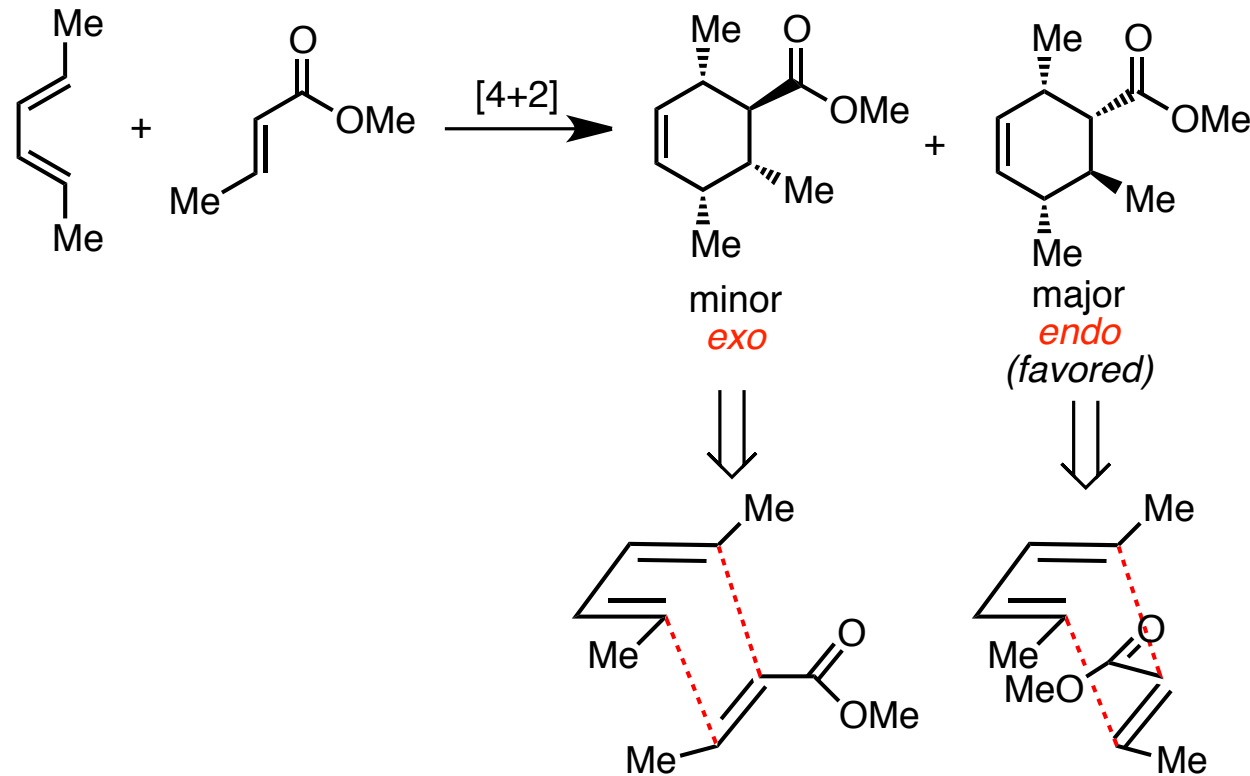
*Normal electron demand* = HOMO of diene + LUMO of dienophile

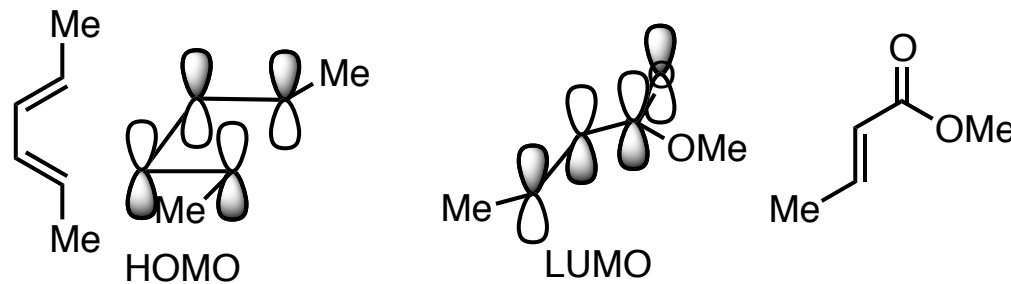
*Inverse electron demand* = HOMO of dienophile + LUMO of diene

## *Net Bonding Interaction?*

The idea: Use FMO's (HOMO + LUMO)



*Diastereoselectivity: Endo vs. Exo*

*Why? ... Secondary Orbital Interactions*

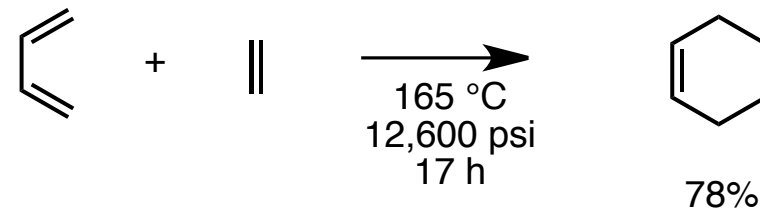
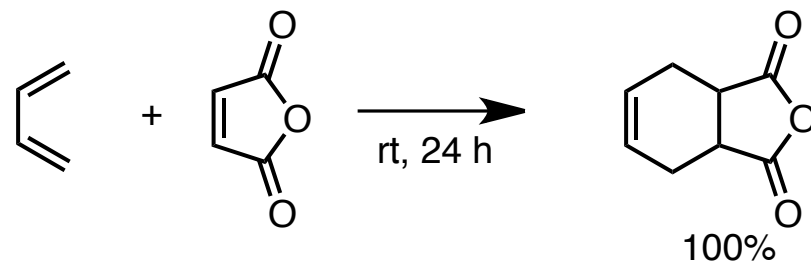
## Regioselectivity & Rates: Substituent Effects

Rates depend on HOMO/LUMO gap.

Perturbation	HOMO	LUMO
extra conjugation	↑	↓
electron-withdrawing group	↓	↓
electron-donating group	↑	↑

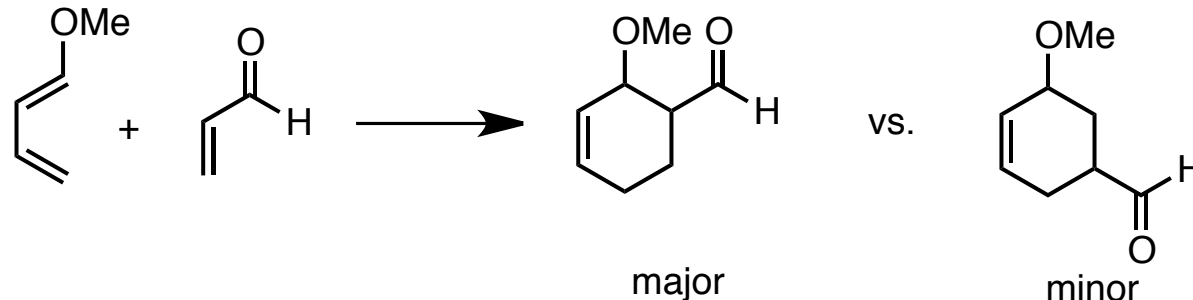
Effects apply to both dienes & dienophiles.  
Effect of substitution is biggest if on C1 of diene.



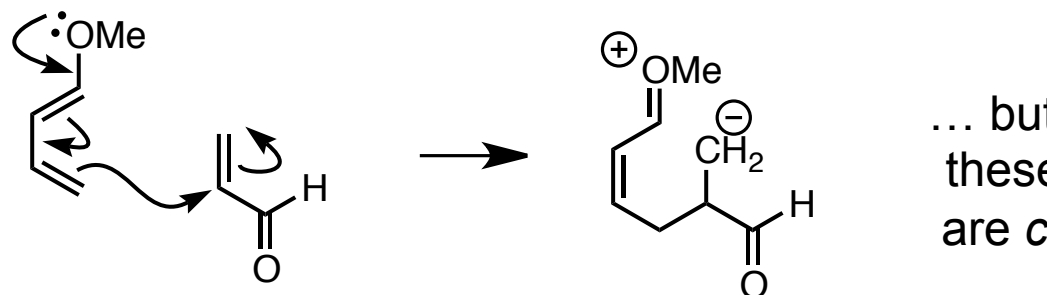
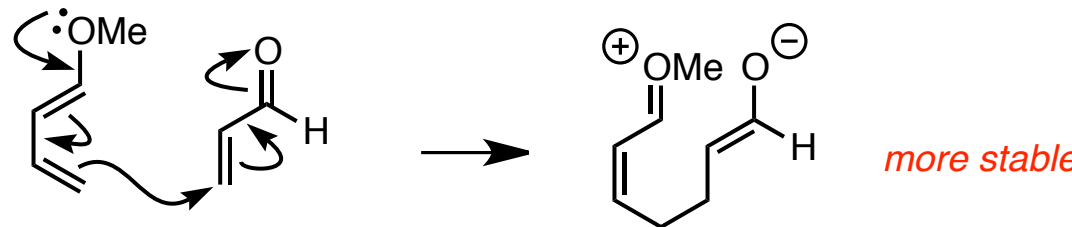
*Examples*

## Regioselectivity

Related to polarization of HOMO and/or LUMO

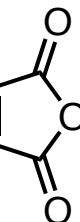
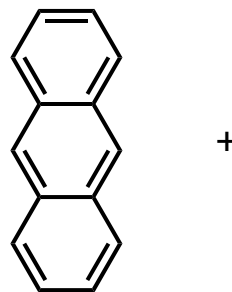


Quick prediction: “imaginary intermediate”  
(push arrows to get maximum effect of substituents)

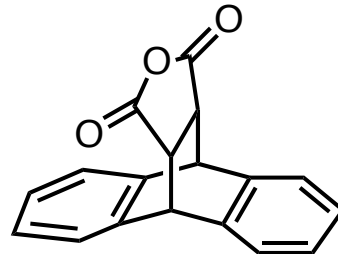


## Lewis Acid Effects

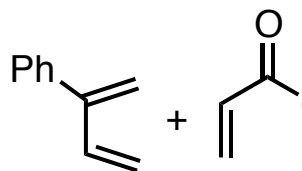
One of the first Lewis acid-accelerated organic transformations!



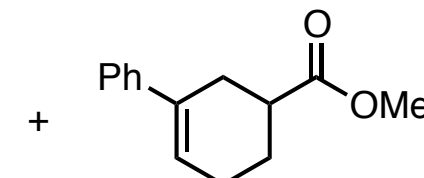
$\xrightarrow[\text{25 } ^\circ\text{C}]{\text{CH}_2\text{Cl}_2}$



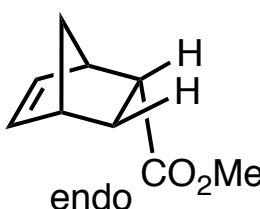
additive	$t_{1/2}$
none	2400 h
$\text{AlCl}_3$ (1 equiv)	< 1 min



1,4



additive	1,4 : 1,3
none	80 : 20
$\text{AlCl}_3$	97 : 3



1,3

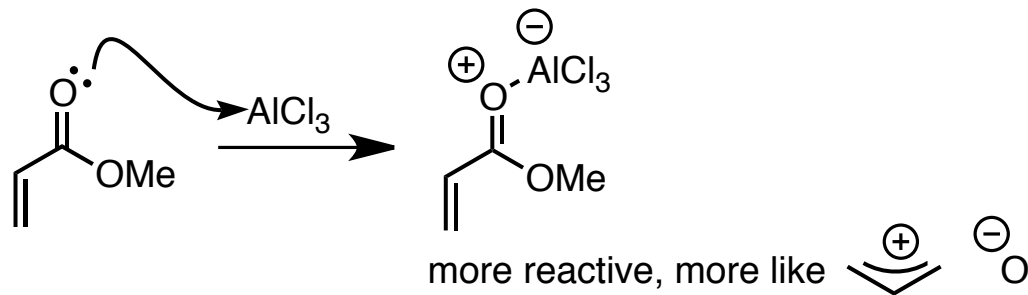
additive	endo:exo
none	82 : 18
$\text{AlCl}_3 \cdot \text{OEt}_2$ (1 equiv)	99 : 1

Lewis acid increases rate, endo/exo selectivity & regioselectivity!

Yates, Eaton. JACS **1960**, 82, 4436

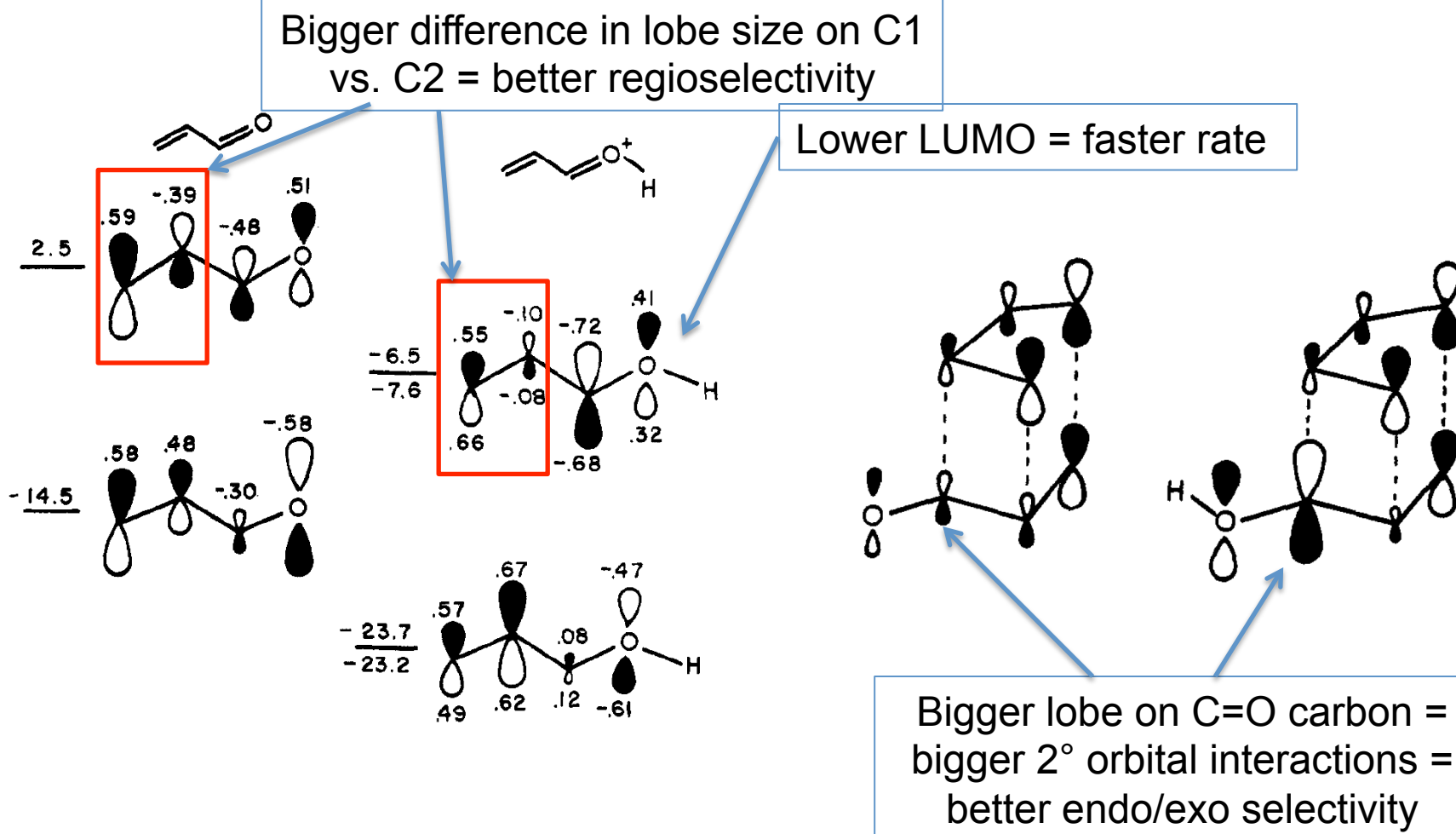
*Why???*

MO perturbation!

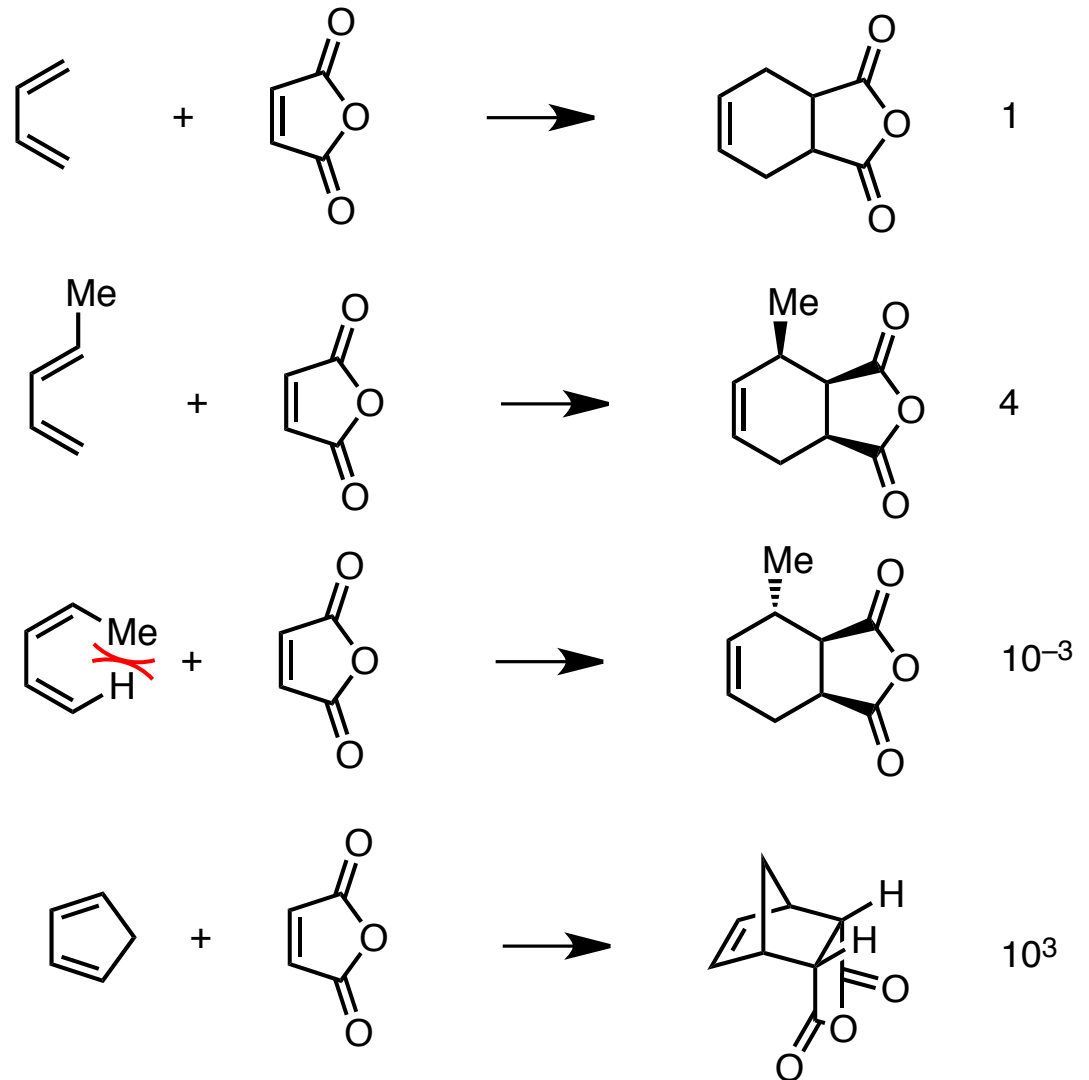
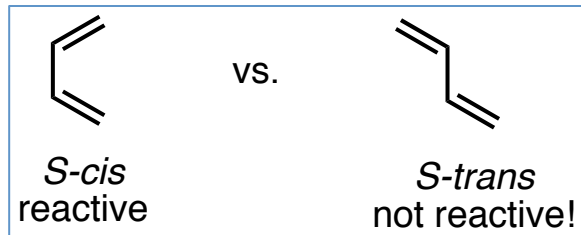


Explains rates, but what about selectivity issues???

*Why???*



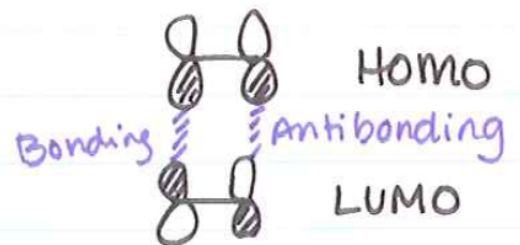
## One More Consideration: *S-cis* vs. *S-trans*



## [2+2] Cycloadditions



FMO Analysis:

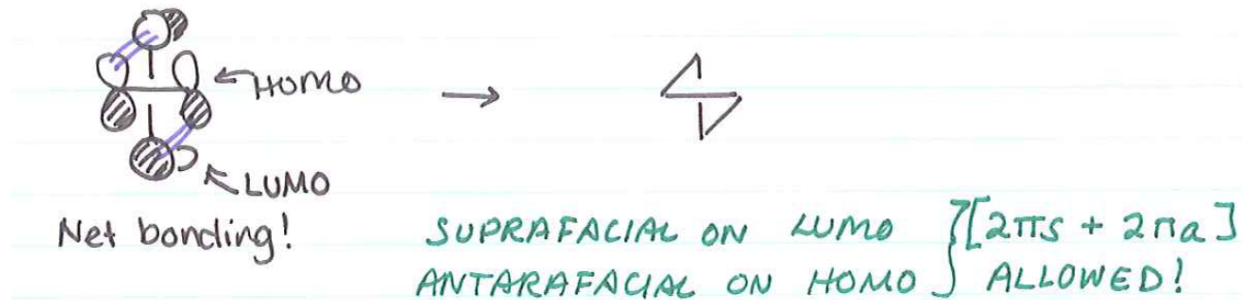


- No net bonding... “forbidden”
- This geometry is *suprafacial* on both  $\pi$  bonds  $\Rightarrow [2\pi_s + 2\pi_s]$

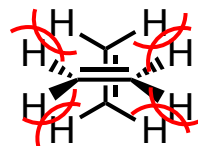
*Suprafacial* = same face of  $\pi$ -system

*Antarafacial* = opposite faces of  $\pi$ -system

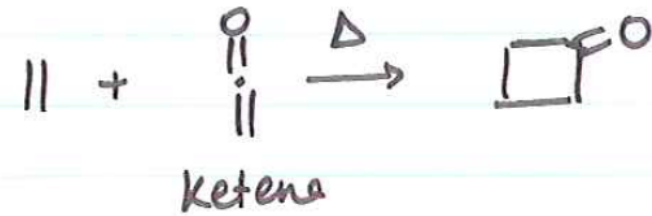
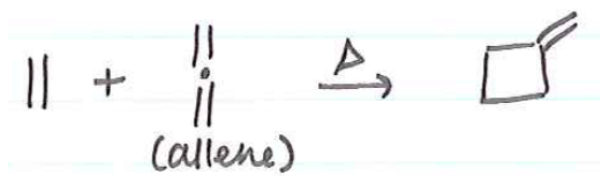
## *Alternative Transition State Geometry*



Problem: Steric Hindrance!



Solution: Remove steric hindrance!



## *Sigmatropic Reactions*

- Reorganization of  $\sigma$  and  $\pi$  bonds (migration of a  $\sigma$ -bond)
- Number of  $\sigma$  and  $\pi$  bonds remains constant
- Classify by  $[m,n]$ -rearrangement or  $[m,n]$ -shift ( $m, n$  = number of atoms in fragment)

## [1,3]-Sigmatropic Rearrangement



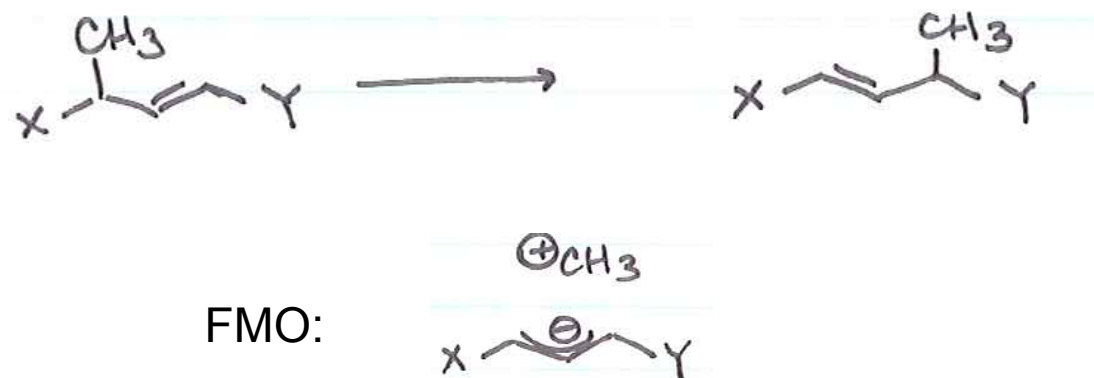
Does this rearrangement proceed under thermal conditions?

Supra- or antara-facial??

For FMO, break into HOMO and LUMO:

*[1,3]-Sigmatropic Rearrangements*

Alkyl Shift?



*[3,3]-Sigmatropic Rearrangements*

FMO Analysis :

2 Allyl radicals :

or

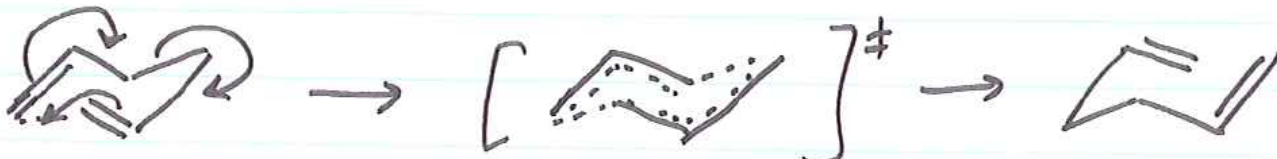
Allyl cation &amp; anion

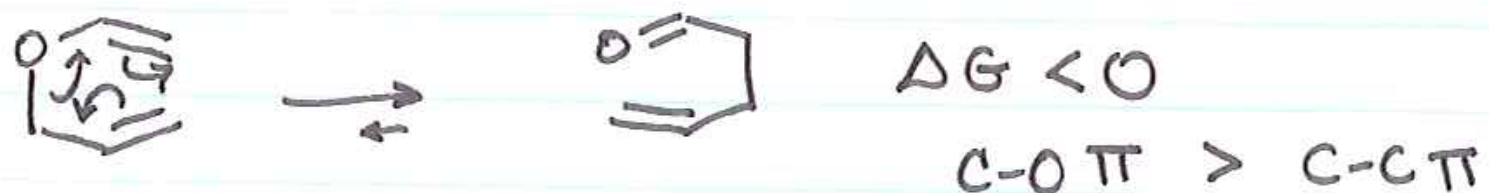


2 bonding interactions



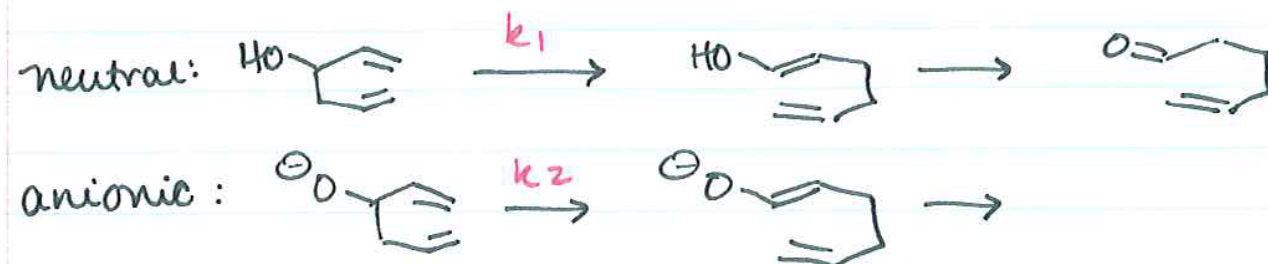
Suprafacial on both components!

Highly predictable TS  $\rightarrow$  "chair-like" (can predict stereochem)

*Claisen Rearrangement*

## Oxy-Cope

2) Tautomerization of Pdt: Oxy-Cope.

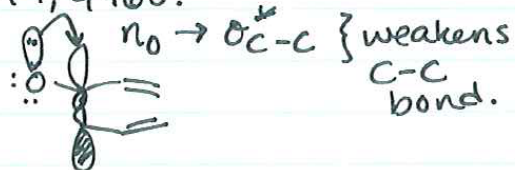
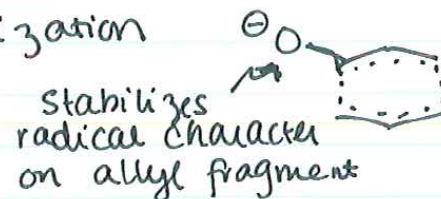


$$\frac{k_2}{k_1} = 10^{10} - 10^{17} !!$$

JACS 1975, 97, 4465.

Why? Ground-state destabilization

TS-stabilization



$\Rightarrow$  Asynchronous  
(bond breaks more  
than bond forms in  
TS)

## Theory #3: Dewar–Zimmerman: Aromatic Transition State

Steps:

1. Choose basis set of 2p AO's (or 1s for H atoms)
2. Assign phases (any phases)
3. Connect orbitals that interact in the starting material
4. Connect lobes that begin to interact in the reaction
5. Count the number of phase inversions
6. Identify topology
  1. Odd # of phase inversions = Möbius
  2. Even # of phase inversions = Hückel
7. Assign Transition State as *aromatic* (thermally allowed) or *antiaromatic* (photochemically allowed)

System/Topology	Aromatic	Antiaromatic
Huckel	$(4n+2) e^-$	$(4n) e^-$
Möbius	$(4n) e^-$	$(4n+2) e^-$

*Example of D–Z Theory*