

Chem 634

# Pericyclic Reactions

Reading: CS-B Chapter 6 Grossman Chapter 4

#### Pericyclic Reactions

Definition: Continuous, concerted reorganization of electrons

cyclic transition state

no intermediate, single transition state

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. Chelatropic
- 5. Group Transfer

#### 3 Theories

#### All 3 theories are correct!

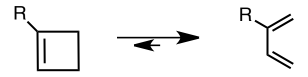
- 1. Woodward-Hoffmann: Conservation of Orbital Symmetry
  - 1<sup>st</sup> historically
  - Uses correlation diagrams
- 2. Fukui: Frontier Molecular Orbital Theory
  - Easier than Woodward–Hoffmann (usually)
  - Based on HOMO/LUMO interactions
- 3. Dewar–Zimmerman: Aromatic Transition State
  - Easiest 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 (hu)

### Type 1: Electrocyclic Reactions

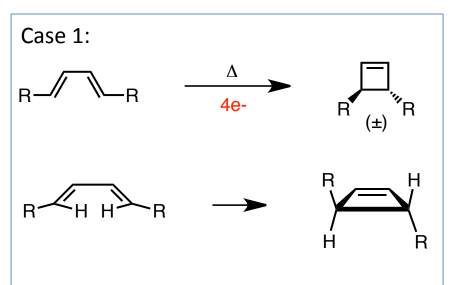
- Ring openings and closures
- Exchange  $\pi$ -bond for  $\sigma$ -bond
- Classified by number of electrons

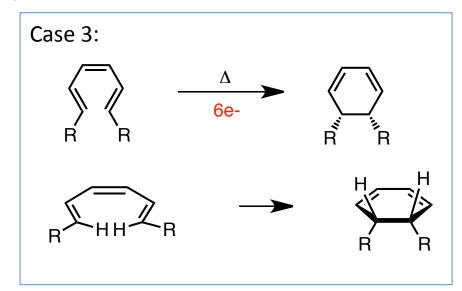


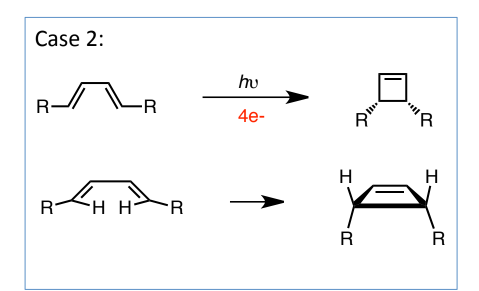
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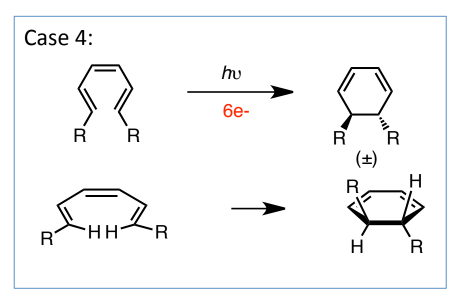
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### Diastereoselectivity – Observations









#### General Phenomenon... Woodward-Hoffmann Rules

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

(n = integer)

<u>6</u> points for a <u>t</u>ouch<u>d</u>own -> <u>6</u>e-, <u>t</u>hermal, <u>d</u>isrotatory

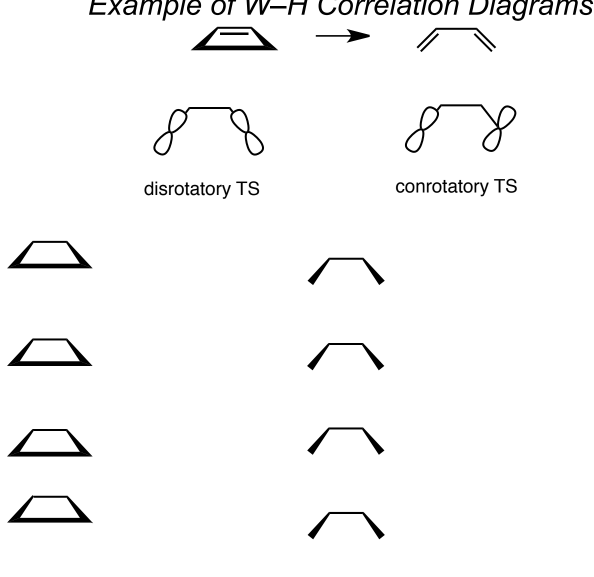
... But why???

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

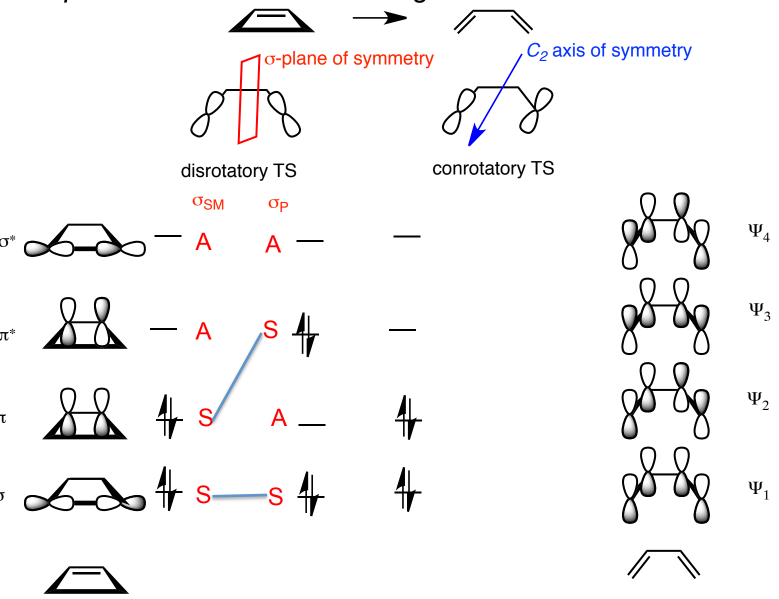
Angew. Chem. Int. Ed. 1969, 8, 781.

- 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.

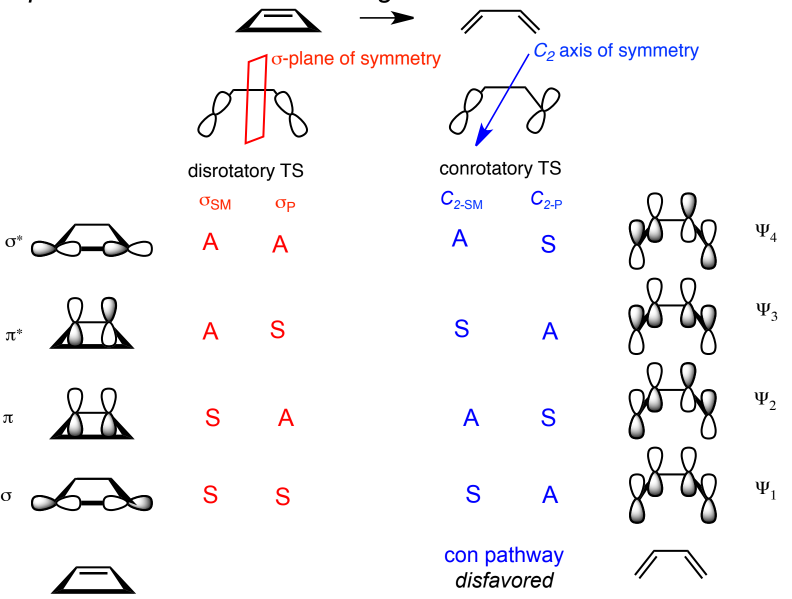
### Example of W–H Correlation Diagrams



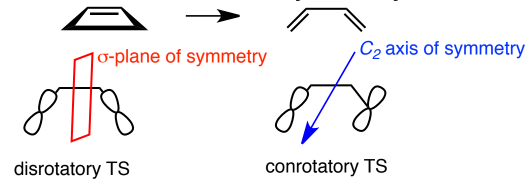
#### Example of W–H Correlation Diagrams: Thermal Conditions

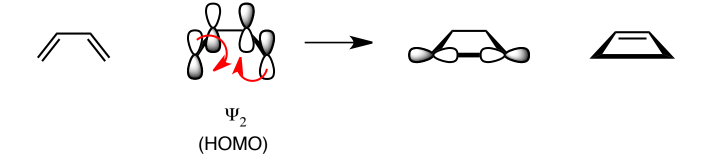


### Example of W–H Correlation Diagrams: Photochemical Conditions



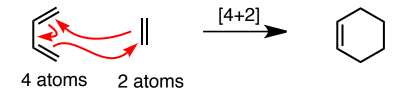
### W-H Conservation of Orbital Symmetry Shortcut





### Cycloadditions & Cycloreversions

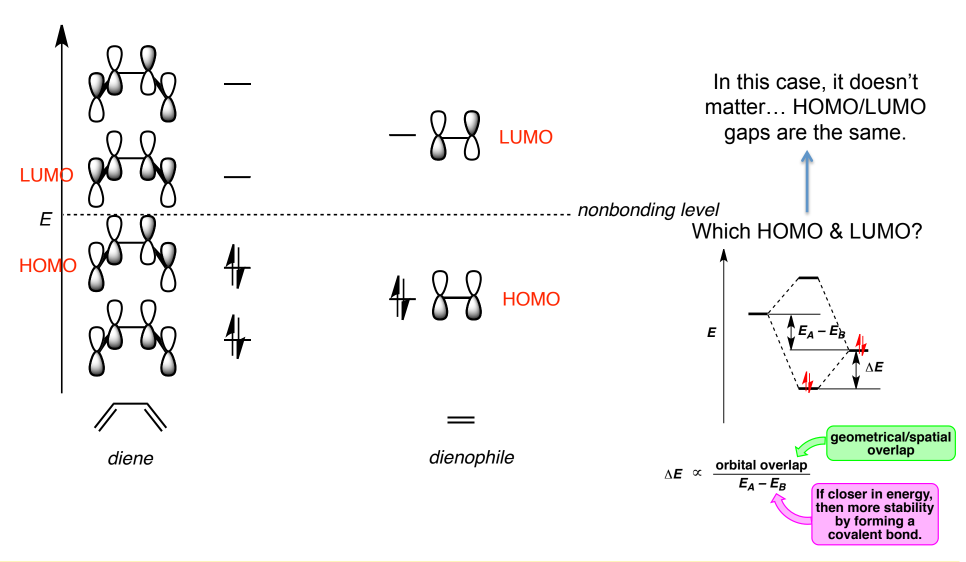
- Union of 2 π-systems
- Exchange  $\pi$ -bonds for  $\sigma$ -bonds
- Classified by [m+n], m & n = # of conjugated atoms in each  $\pi$ -system



Diels–Alder Reaction!
Note: 6 eGreat way to make cyclohexenes & cyclohexanes

#### Fukui: Frontier Molecular Orbital (FMO) Theory

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



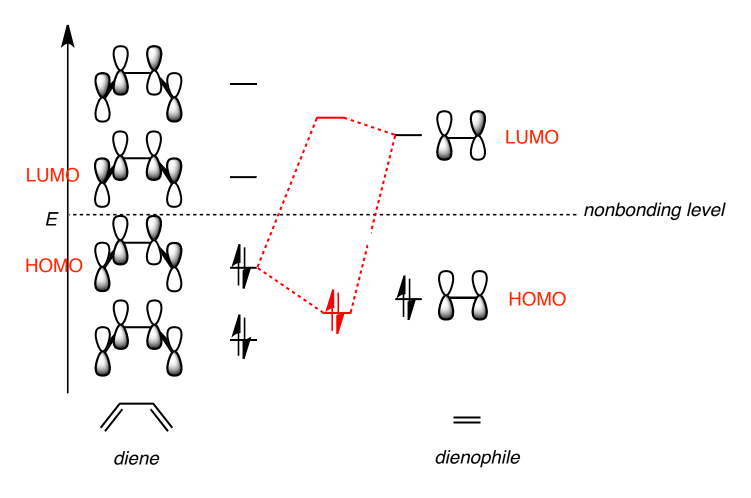
#### Types of Diels-Alder Reactions

Normal electron demand = HOMO of diene + LUMO of dienophile

Inverse electron demand = HOMO of dienophile + LUMO of diene

### Net Bonding Interaction?

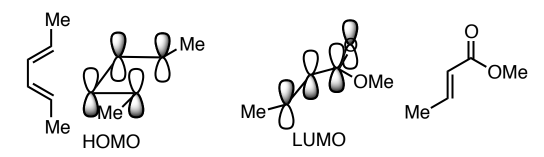
The idea: Use FMO's (HOMO + LUMO)



# Diastereoselectivity: Endo vs. Exo

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# Why? ... Secondary Orbital Interactions



#### Regioselectivity & Rates: Substituent Effects

Rates depend on HOMO/LUMO gap.

Perturbation	НОМО	LUMO
extra conjugation	<b>^</b>	•
electron-withdrawing group	•	•
electron-donating group	<b>^</b>	<b>^</b>

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!

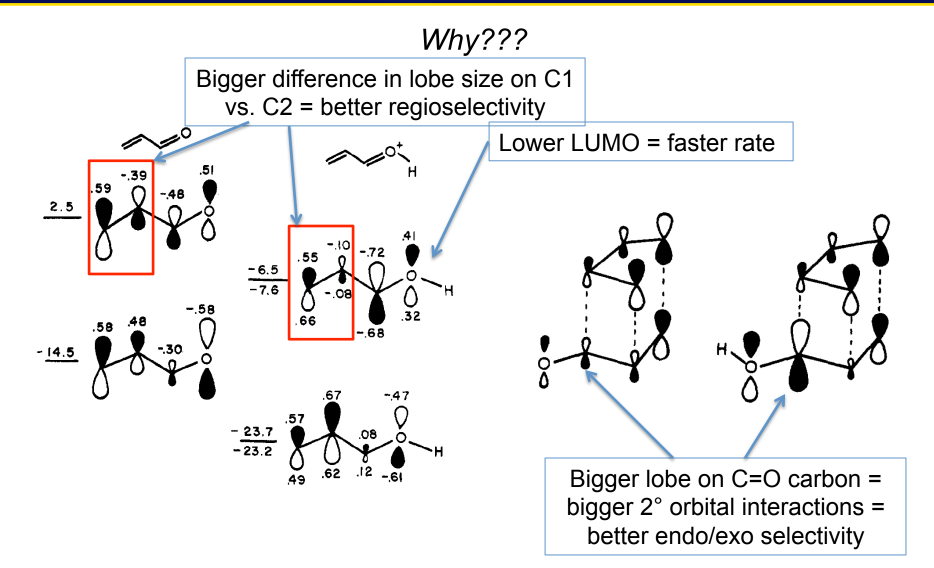
Lewis acid increases rate, endo/exo selectivity & regioselectivity!

Yates, Eaton. JACS 1960, 82, 4436

### *Why???*

#### MO perturbation!

Explains rates, but what about selectivity issues???



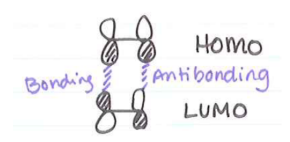
#### One More Consideration: S-cis vs. S-trans



#### [2+2] Cycloadditions



FMO Analysis:



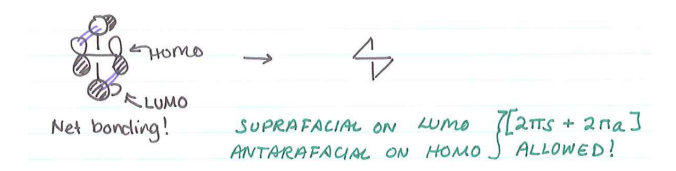
- No net bonding... "forbidden"
- This geometry is *suprafacial* on both  $\pi$  bonds =>  $[2\pi_s + 2\pi_s]$

Suprafacial = same face of  $\pi$ -system

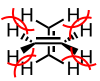
Antarafacial = opposite faces of  $\pi$ -system



#### 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:



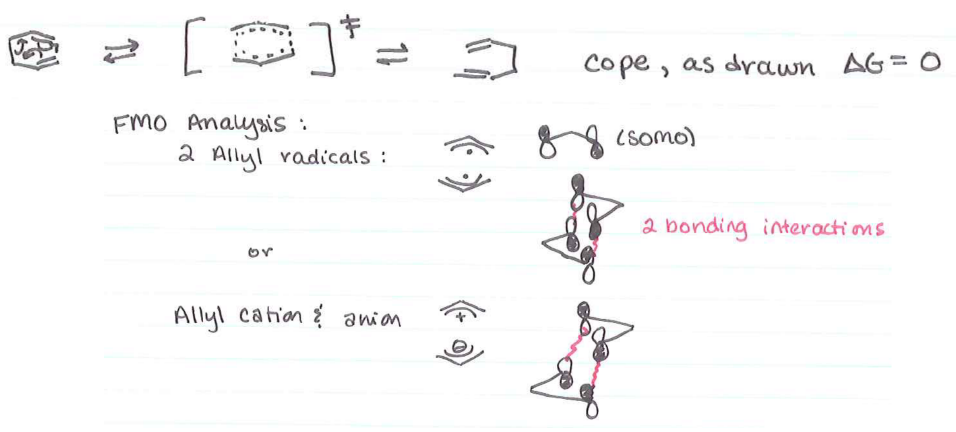
### [1,3]-Sigmatropic Rearrangements

Alkyl Shift?

FMO:

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### [3,3]-Sigmatropic Rearrangements



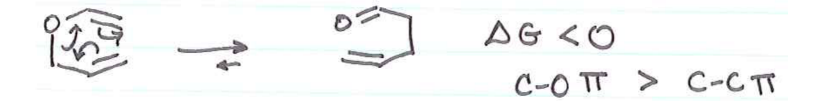
Suprafacial on both components!

Highly predictable TS -> "chair-like" (can predict stereochem)

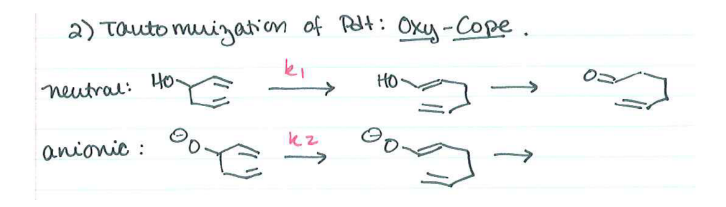




### Claisen Rearrangement



### Oxy-Cope



#### 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-
Mobius	(4n) e-	(4n+2) e-



# Example of D–Z Theory