

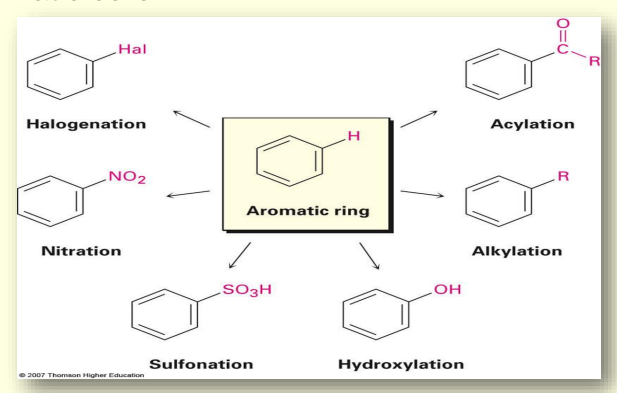
# Chemistry of Benzene Electrophilic Aromatic Substitution

Based on McMurry's *Organic Chemistry*, 7<sup>th</sup> edition Assistant Lecturer. Jalal Hasan Mohammed



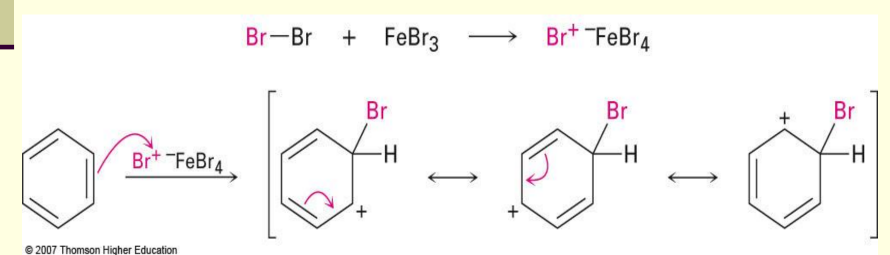
# Substitution Reactions of Benzene and Its Derivatives

- Benzene is aromatic: a cyclic conjugated compound with 6 π electrons
- Reactions of benzene lead to the retention of the aromatic core.



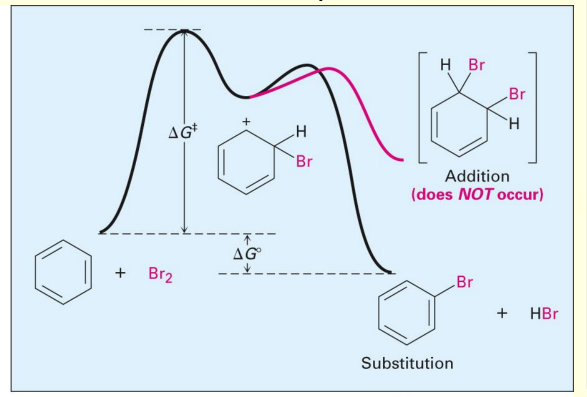
### Electrophilic Aromatic Bromination

- Benzene's  $\pi$  electrons participate as a Lewis base in reactions with Lewis acids
- The product is formed by loss of a proton, which is replaced by bromine
- FeBr<sub>3</sub> is added as a catalyst to polarize the bromine reagent
- In the first step the  $\pi$  electrons act as a nucleophile toward  $Br_2$  (in a complex with  $FeBr_3$ )
- This forms a cationic addition intermediate from benzene and a bromine cation
- The intermediate is not aromatic and therefore high in energy

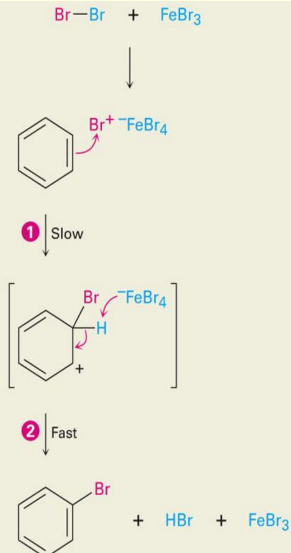


### Formation of Product from Intermediate

- The cationic addition intermediate transfers a proton to FeBr<sub>4</sub>- (from Brand FeBr<sub>3</sub>)
  - This restores aromaticity (in contrast with addition in alkenes)



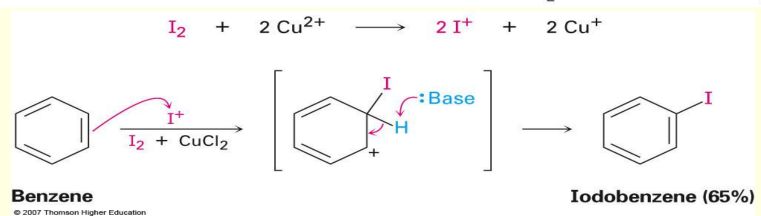
Reaction progress



Energy

### Other Aromatic Halogenations

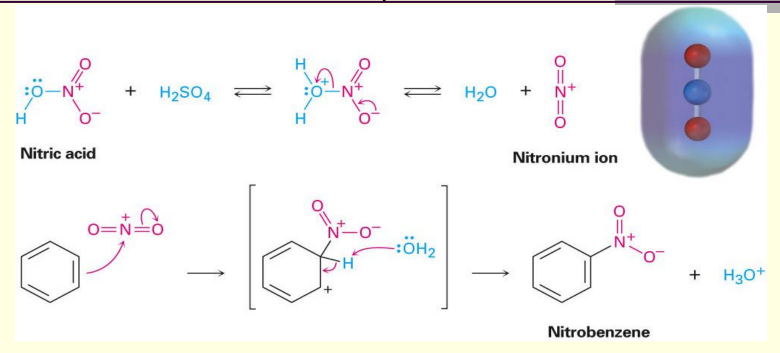
- Chlorine and iodine (but not fluorine, which is too reactive) can produce aromatic substitution with the addition of other reagents to promote the reaction
- Chlorination requires FeCl<sub>3</sub>
- Iodine must be oxidized to form a more powerful I+ species (with Cu<sup>2+</sup> from CuCl<sub>2</sub>)



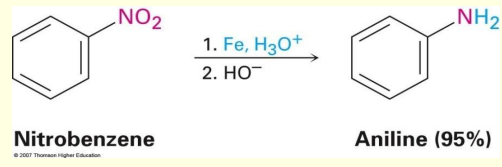
5

### **Aromatic Nitration**

- The combination of nitric acid and sulfuric acid produces NO<sub>2</sub><sup>+</sup> (nitronium ion)
- The reaction with benzene produces nitrobenzene

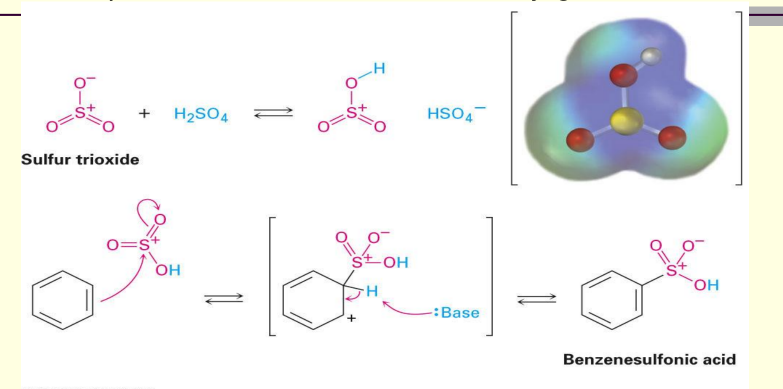


The Nitro group can be reduced to an Amino group if needed



# Aromatic Sulfonation Substitution of H by SO<sub>3</sub> (sulfonation)

- Reaction with a mixture of sulfuric acid and SO<sub>3</sub> ("Fuming H<sub>2</sub>SO<sub>4</sub>)
  - Reactive species is sulfur trioxide or its conjugate acid

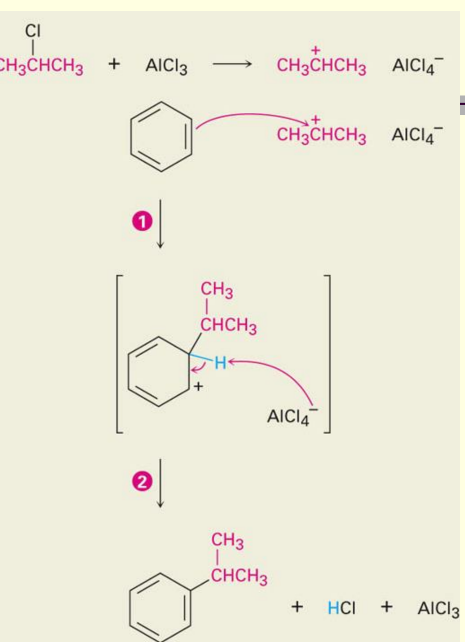


Sulfonamides are "sulfa drug" antibiotics

### Alkylation of Aromatic Rings: The Friedel–Crafts

#### Reaction

- Alkylation among most useful electrophilic aromatic substitution reactions
- Aromatic substitution of R+ for H+
- Aluminum chloride promotes the formation of the carbocation



### Limitations of the Friedel-Crafts Alkylation

- Only alkyl halides can be used (F, Cl, I, Br)
- Ary/ halides and vinylic halides do not react (their carbocations are too hard to form)
- Will not work with rings containing an amino group substituent or a strongly electron-withdrawing group

+ R-X 
$$\xrightarrow{AICI_3}$$
 **NO reaction** where Y =  $-\stackrel{+}{NR_3}$ ,  $-NO_2$ ,  $-CN$ ,  $-SO_3H$ ,  $-CHO$ ,  $-COCH_3$ ,  $-CO_2H$ ,  $-CO_2CH_3$  ( $-NH_2$ ,  $-NHR$ ,  $-NR_2$ )

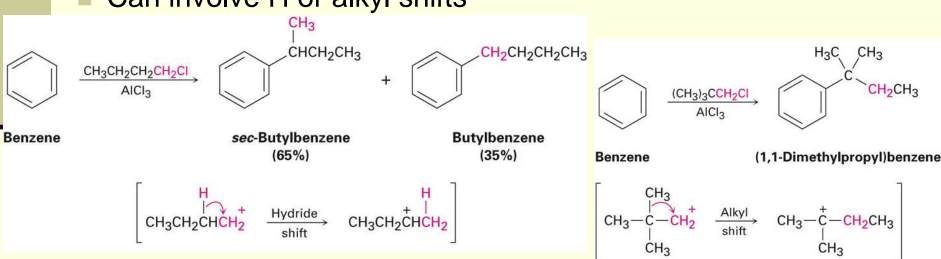
9

### Other Problems with Alkylation

Multiple alkylations can occur because the first alkylation is

activating

- Carbocation Rearrangements Occur During Alkylation
  - Similar to those occurring during electrophilic additions to alkene
  - Can involve H or alkyl shifts



### Acylation of Aromatic Rings

- Reaction of an acid chloride (RCOCI) and an aromatic ring in the presence of AICI<sub>3</sub> introduces **acyl group**, —COR
  - Benzene with acetyl chloride yields acetophenone

- Avoids many of the problems of alkylation
  - Only substitutes once, because acyl group is deactivating
  - No rearrangement because of resonance stabilized cation

### Mechanism of Friedel-Crafts Acylation

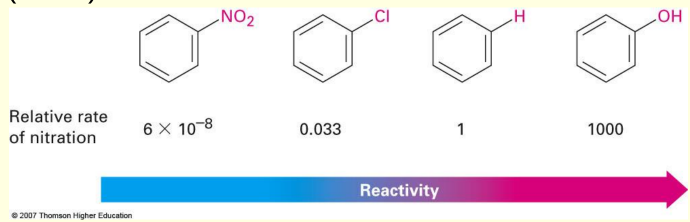
- Similar to alkylation
- Reactive electrophile: resonance-stabilized acyl cation
- An acyl cation does not rearrange

Can reduce carbonyl to get alkyl product

$$\begin{array}{c|c} C & H & H \\ \hline C & CH_2CH_3 & \underline{H_2/Pd} & C & CH_2CH_3 \end{array}$$

### Substituent Effects in Aromatic Rings

 Substituents can cause a compound to be (much) more or (much) less reactive than benzene



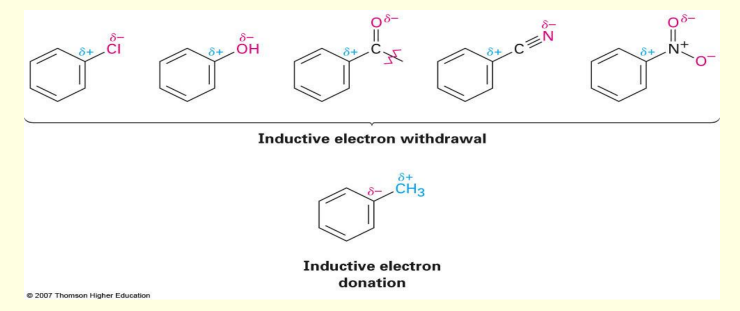
- Substituents affect the orientation of the reaction the positional relationship is controlled
- ortho- and para-directing activators, ortho- and paradirecting deactivators, and meta-directing deactivators.

### Origins of Substituent Effects

- An interplay of inductive effects and resonance effects
- Inductive effect withdrawal or donation of electrons
   through a σ bond = Polar Covalent Bonds
- Resonance effect withdrawal or donation of electrons through a π bond due to the overlap of a p orbital on the substituent with a p orbital on the aromatic ring

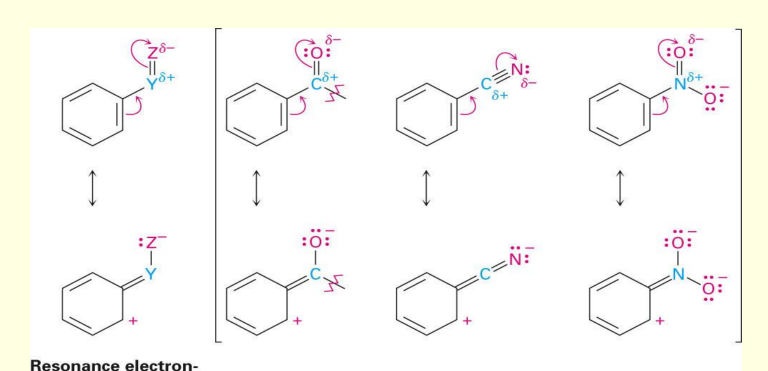
### Inductive Effects

- Controlled by electronegativity and the polarity of bonds in functional groups
- Halogens, C=O, CN, and NO<sub>2</sub> withdraw electrons through σ bond connected to ring
- Alkyl groups donate electrons



#### Resonance Effects – Electron Withdrawal

- C=O, CN, NO<sub>2</sub> substituents withdraw electrons from the aromatic ring by resonance
- $\blacksquare$   $\pi$  electrons flow from the rings to the substituents
- Look for a double (or triple) bond connected to the ring by a single bond

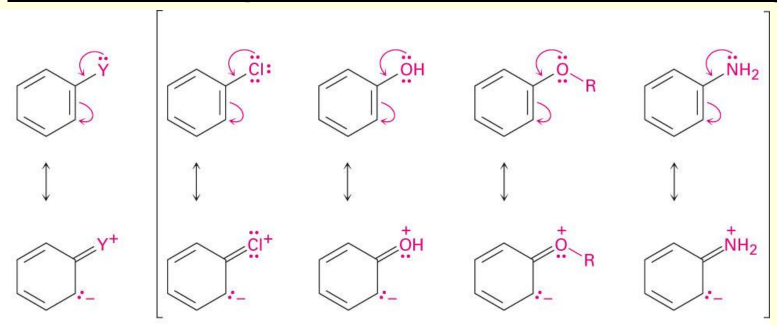


withdrawing group
© 2007 Thomson Higher Education

16

### Resonance Effects – Electron Donation

- Halogen, OH, alkoxyl (OR), and amino substituents donate electrons
- $\pi$  electrons flow from the substituents to the ring
- Effect is greatest at ortho and para positions
- Look for a lone pair on an atom attached to the ring

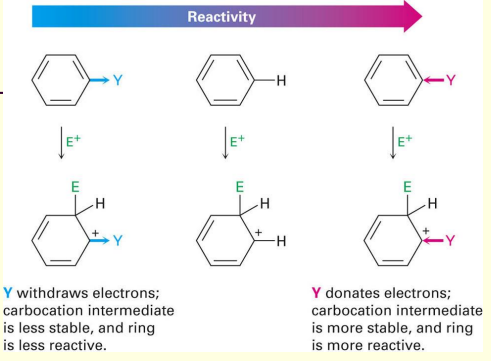


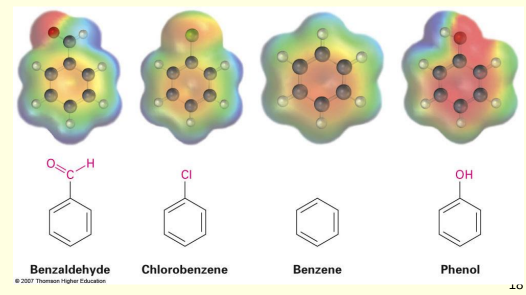
Resonance electrondonating group

© 2007 Thomson Higher Education

### An Explanation of Substituent Effects

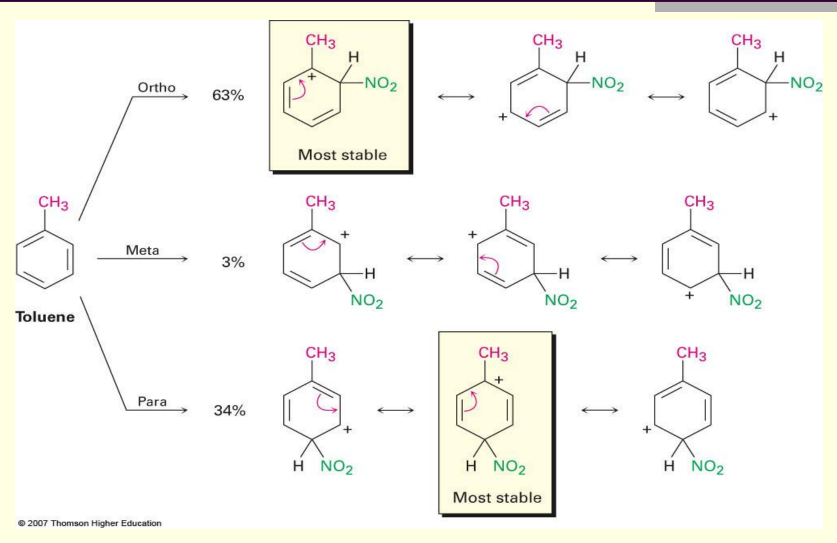
- Activating groups donate electrons to the ring, stabilizing the carbocation intermediate
- Deactivating groups withdraw electrons from the ring, destabilizing carbocation intermediate





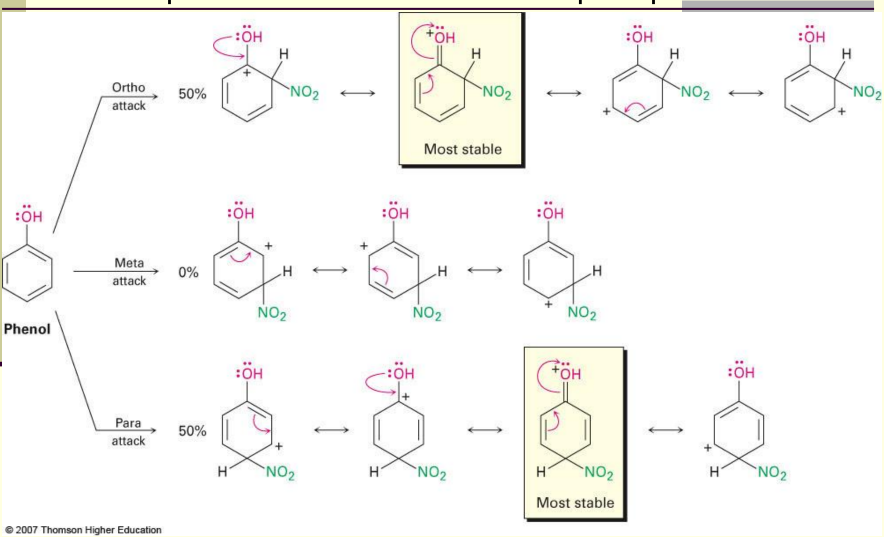
# Ortho/Para-Directing Activators: Alkyl Groups Alkyl groups activate by induction: direct further substitution to positions ortho and para to themselves

Alkyl group has most effect on the ortho and para positions



### Ortho/Para-Directing Activators: OH and NH<sub>2</sub> Alkoxyl, and amino groups have a strong, electron-

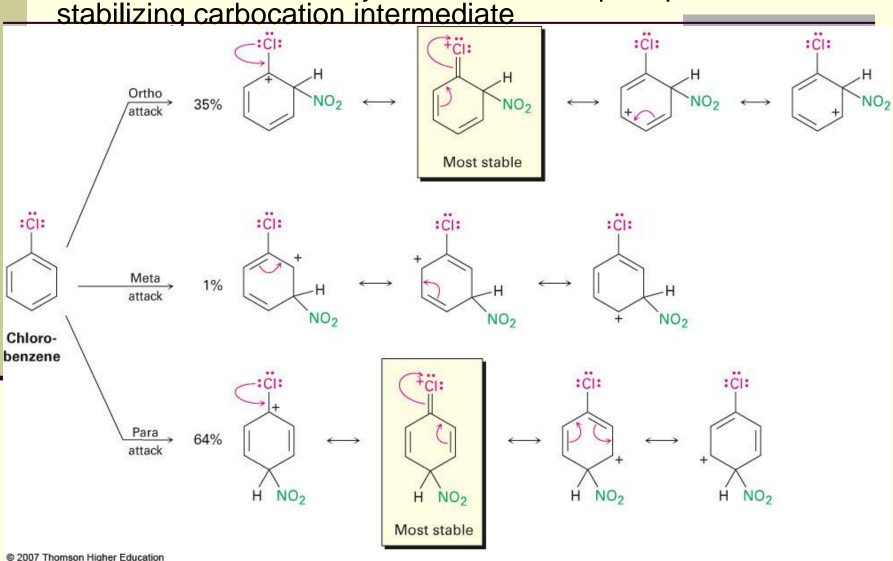
- donating resonance effect
- Most pronounced at the ortho and para positions



### Ortho/Para-Directing Deactivators: Halogens Electron-withdrawing inductive effect outweighs weaker electron-

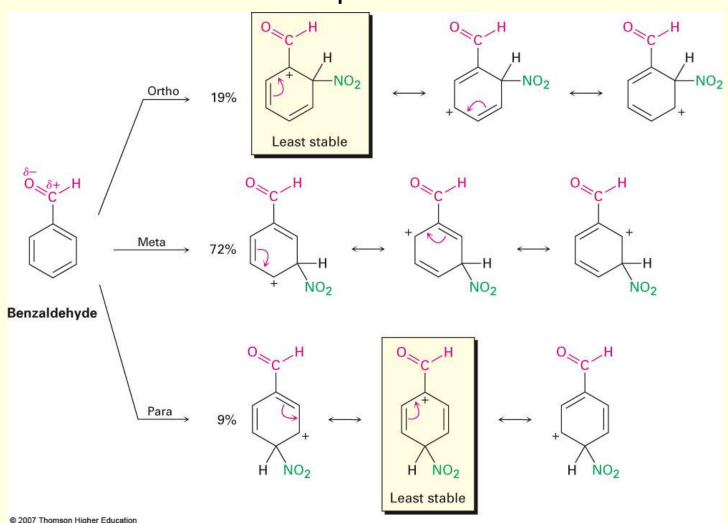
donating resonance effect

Resonance effect is only at the ortho and para positions,



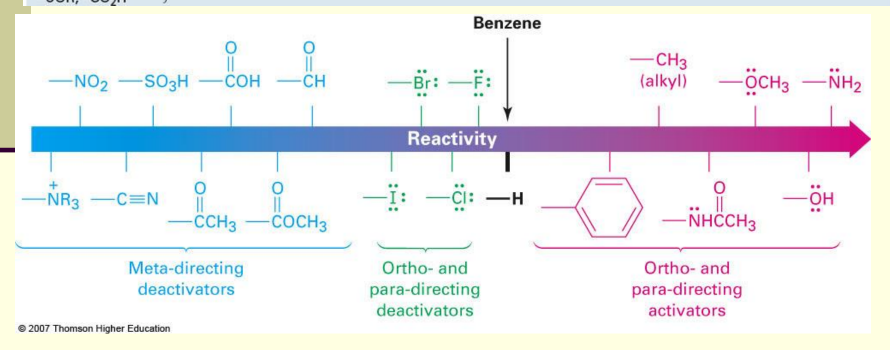
## Meta-Directing Deactivators Inductive and resonance effects reinforce each other

- Ortho and para intermediates destabilized by deactivation of carbocation intermediate
- Resonance cannot produce stabilization



# Summary Table: Effect of Substituents in Aromatic Substitution

	Table 16.2 Substituent Effects in Electrophilic Aromatic Substitution					
_	Substituent		Reactivity	Orienting effect	Inductive effect	Resonance effect
	-CH <sub>3</sub>		Activating	Ortho, para	Weak donating	, <del>, , ,</del>
	-OH, -NH <sub>2</sub>	ļ.	Activating	Ortho, para	Weak withdrawing	Strong donating
	–F, –Cl –Br, –I	}	Deactivating	Ortho, para	Strong withdrawing	Weak donating
	-NO <sub>2</sub> , -CN -CHO, -CO -COR, -CO	D <sub>2</sub> R	Deactivating	Meta	Strong withdrawing	Strong withdrawing



### Trisubstituted Benzenes: Additivity of Effects

If the directing effects of the two groups are the same, the result is additive

If the directing effects of two groups oppose each other, the more powerful activating group decides the principal outcome

Usually gives mixtures of products

24

### Meta-Disubstituted Compounds

- The reaction site is too hindered
- To make aromatic rings with three adjacent substituents, it is best to start with an orthodisubstituted compound

But:

CH<sub>3</sub>

CH<sub>3</sub>

CH<sub>3</sub>

CH<sub>3</sub>

CH<sub>3</sub>

CH<sub>3</sub>

CH<sub>3</sub>

CH<sub>3</sub>

CH<sub>3</sub>

PeCl<sub>3</sub>

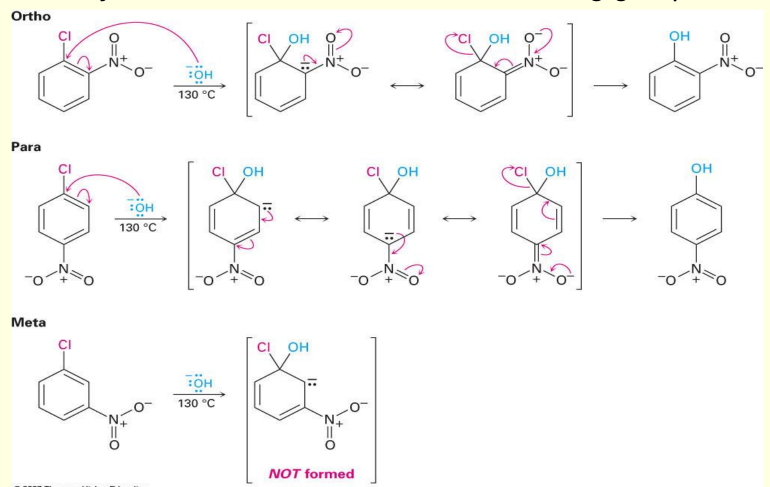
CH<sub>3</sub>

C

© 2007 Thomson Higher Education

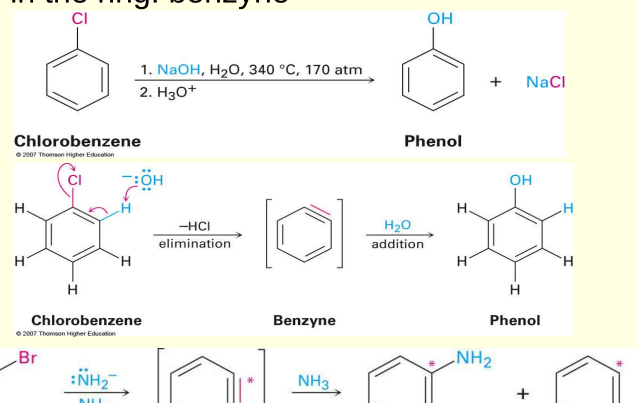
## Nucleophilic Aromatic Substitution Aryl halides with electron-withdrawing substituents ortho and para

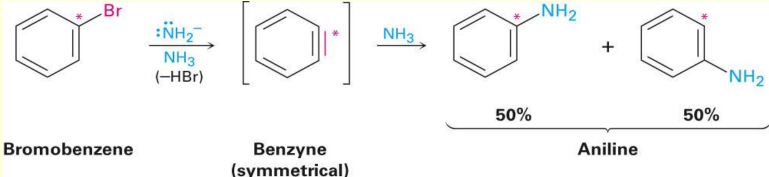
- Aryl halides with electron-withdrawing substituents ortho and para react with nucleophiles (electron withdrawing needed to accept electrons from the nucleophile)
- Form addition intermediate (Meisenheimer complex) that is stabilized by electron-withdrawal. Halide is leaving group.



### Benzyne: Substitution of Unactivated Aromatics Phenol is prepared industrially by treatment of chlorobenzene

- with dilute aqueous NaOH at 340°C under high pressure
- The reaction involves an elimination reaction that gives a triple bond in the ring: benzyne



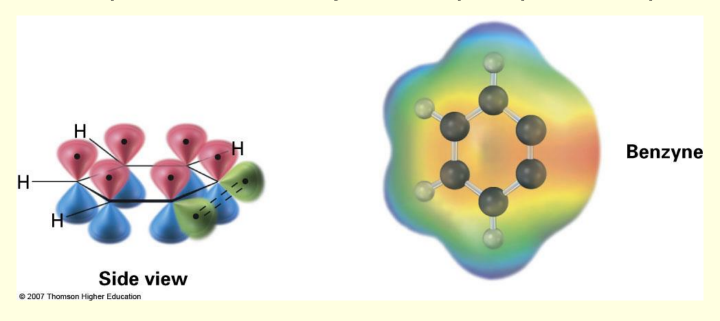


@ 2007 Thomson Higher Education

27

### Structure of Benzyne

- Benzyne is a highly distorted alkyne
- The triple bond uses  $sp^2$ -hybridized carbons, not the usual sp
- The triple bond has one  $\pi$  bond formed by p-p overlap and another by weak  $sp^2-sp^2$  overlap



### Oxidation of Aromatic Compounds

- Alkyl side chains can be oxidized to —CO<sub>2</sub>H by strong reagents such as KMnO<sub>4</sub> if they have a C-H next to the ring
- Converts an alkylbenzene into a benzoic acid, Ar—R → Ar—CO<sub>2</sub>H
- A benzylic C-H bond is required, or no reaction takes place

$$\begin{array}{c|c} & H_3C & CH_3 \\ \hline & C & CH_3 & \hline & KMnO_4 \\ \hline & H_2O & No \ reaction \\ \hline \\ & & & & \\ \hline \\ & &$$

# Reduction of Aromatic Compounds Aromatic rings are inert to catalytic hydrogenation under

- conditions that reduce alkene double bonds
- Can selectively reduce an alkene double bond in the presence of an aromatic ring
- Reduction of an aromatic ring requires more powerful reducing conditions (high pressure or rhodium catalysts)

© 2007 Thomson Higher Education

© 2007 Thomson Higher Education

30

#### Synthesis of Trisubstituted Benzenes

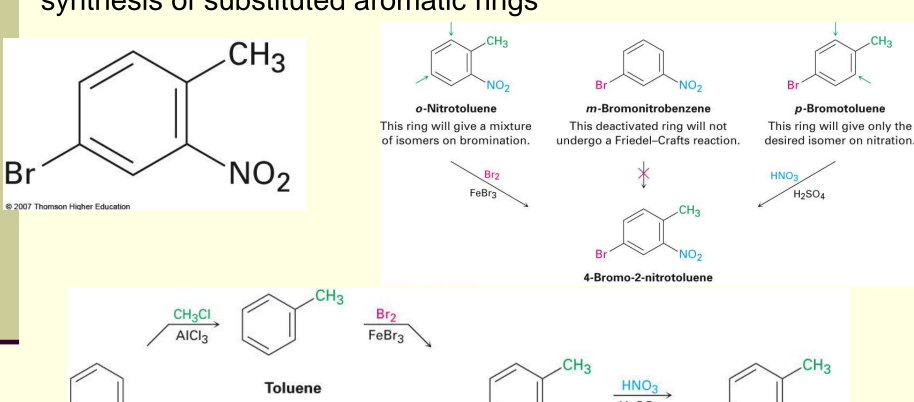
Benzene

© 2007 Thomson Higher Education

FeBr<sub>3</sub>

Bromobenzene

- These syntheses require planning and consideration of alternative routes
- Ability to plan a sequence of reactions in right order is valuable to synthesis of substituted aromatic rings



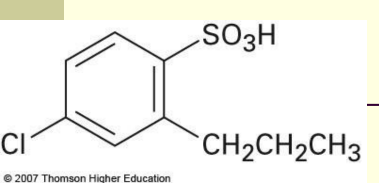
p-Bromotoluene

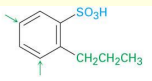
CH<sub>3</sub>CI

AICI<sub>3</sub>

31

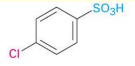
4-Bromo-2-nitrotoluene





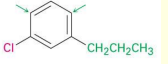
#### o-Propylbenzenesulfonic acid

—This ring will give the wrong isomer on chlorination.



#### p-Chlorobenzenesulfonic acid

This deactivated ring will not undergo a Friedel–Crafts reaction.



#### m-Chloropropylbenzene

This ring will give the desired product on sulfonation.

4-Chloro-2-propylbenzenesulfonic acid

$$\begin{array}{c}
O \\
\parallel \\
CH_3CH_2CCI \\
\hline
AICI_3
\end{array}$$

© 2007 Thomson Higher Education

#### Benzene

#### **Propiophenone**

#### *m*-Chloropropiophenone

m-Chloropropylbenzene