

# CHAPTER-10

# ELECTROPHILIC AROMATIC SUBSTITUTION

BY,

G.DEEPA.

## Electrophilic Aromatic Substitution (Aromatic compounds)

Ar-H = aromatic compound

### 1. Nitration



### 2. Sulfonation



### 3. Halogenation

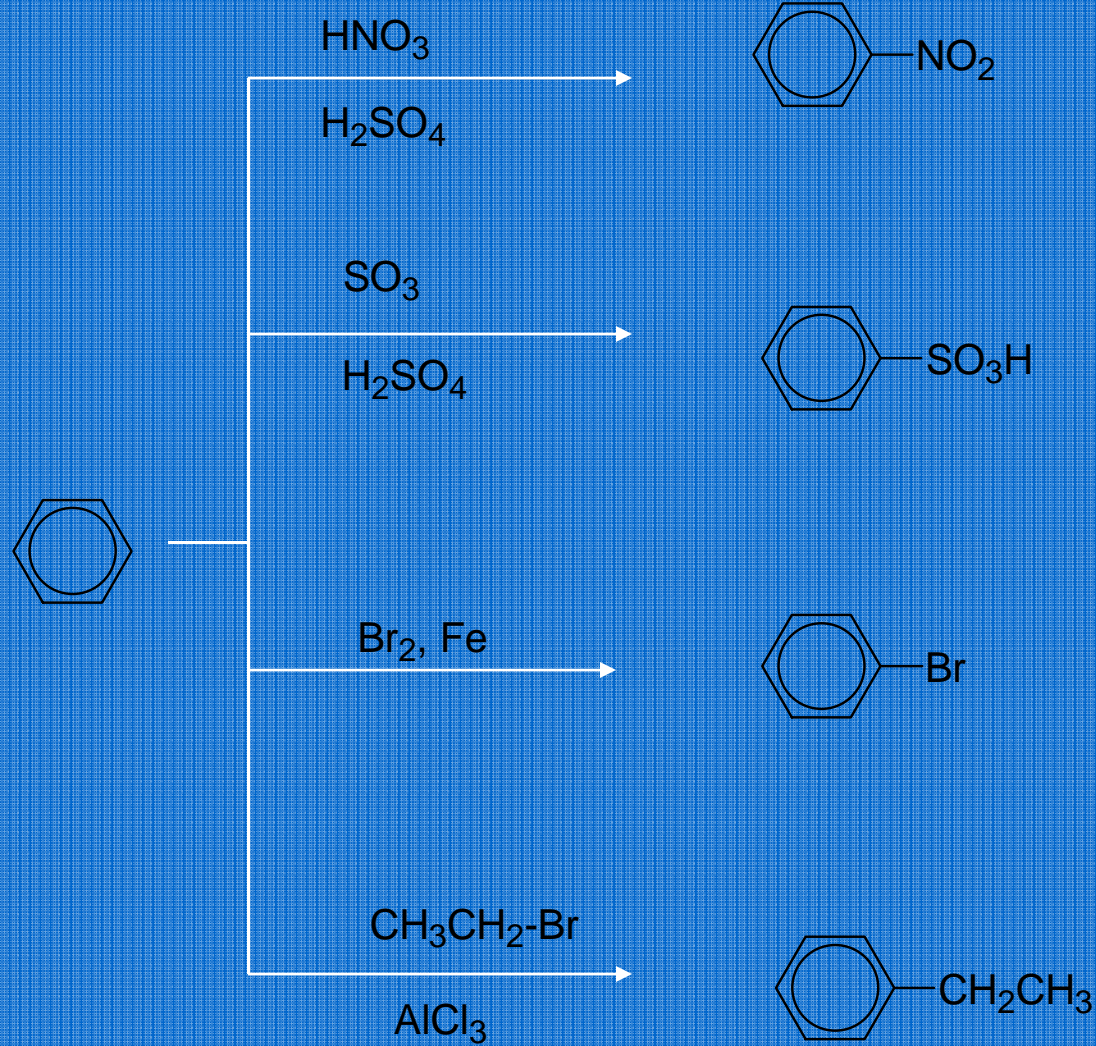


### 4. Friedel-Crafts alkylation

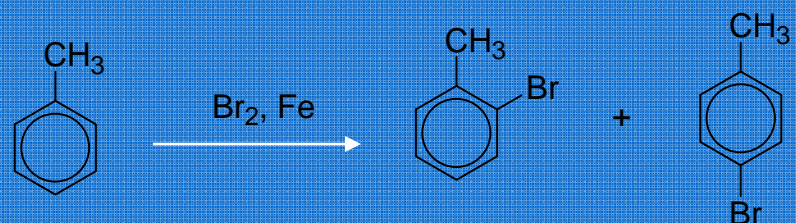
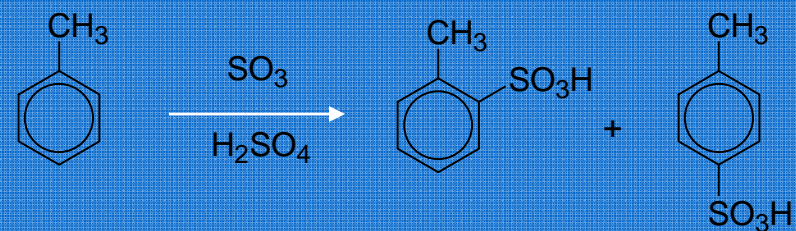
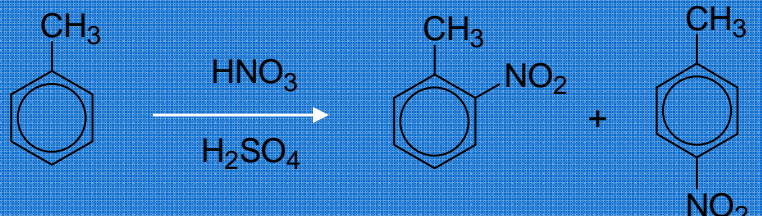


## Friedel-Crafts alkylation (variations)



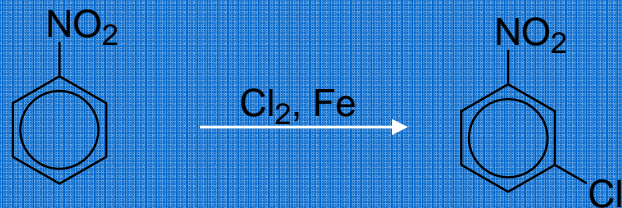
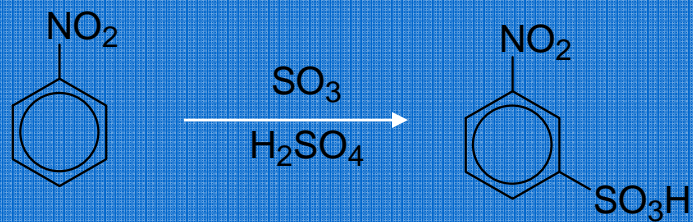
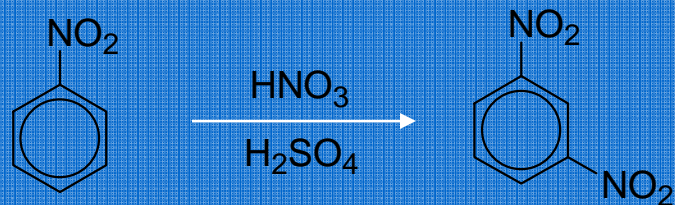


## Toluene



**faster** than the same reactions with benzene

## Nitrobenzene



slower than the same reactions with benzene

Substituent groups on a benzene ring affect electrophilic aromatic substitution reactions in two ways:

**1) reactivity**

activate (faster than benzene)

or deactivate (slower than benzene)

**2) orientation**

*ortho*- + *para*- direction

or *meta*- direction

-CH<sub>3</sub>

activates the benzene ring towards EAS

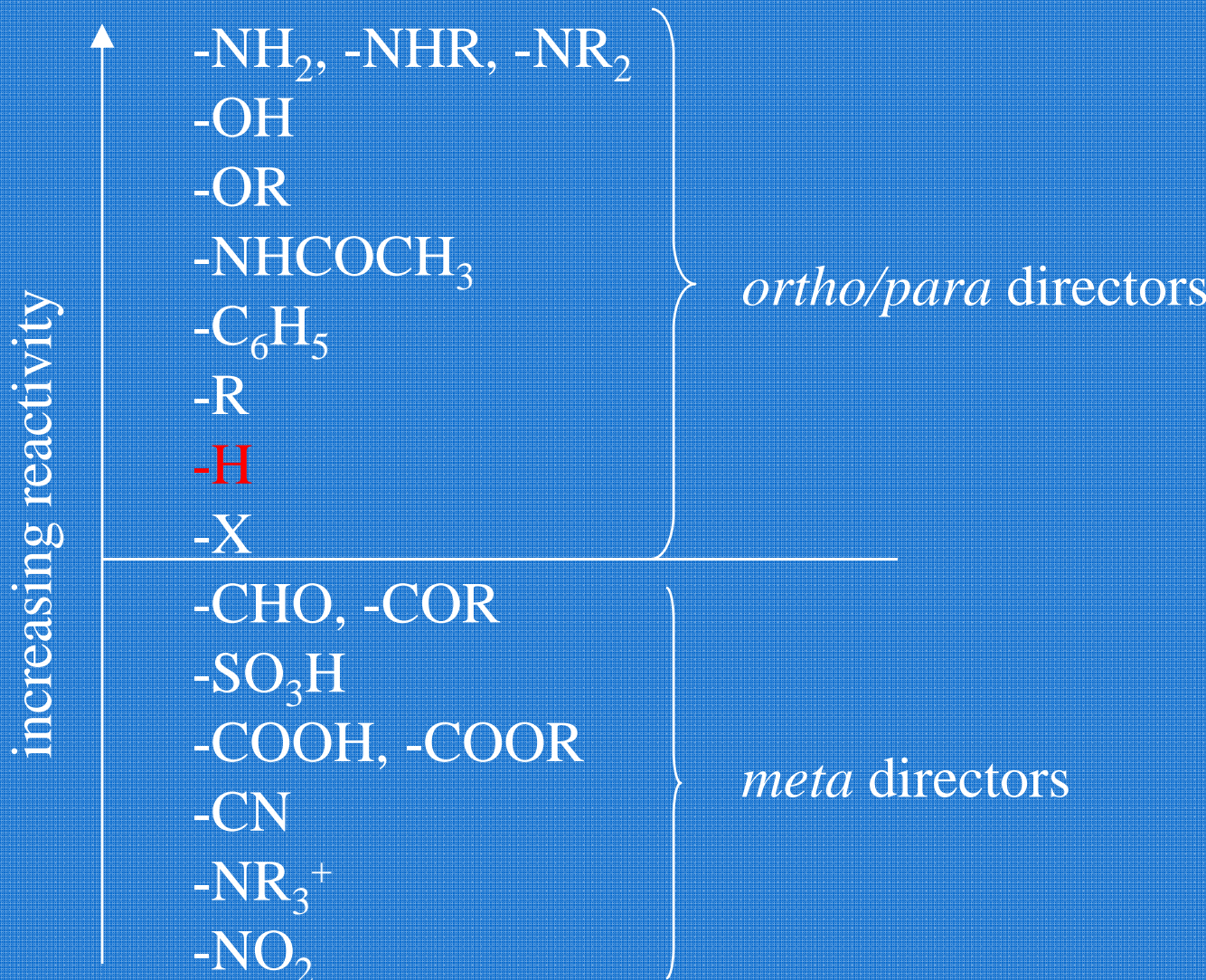
directs substitution to the *ortho*- & *para*- positions

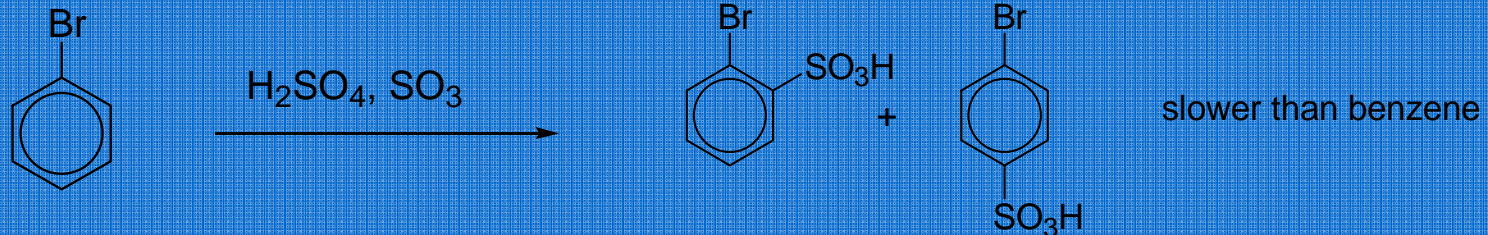
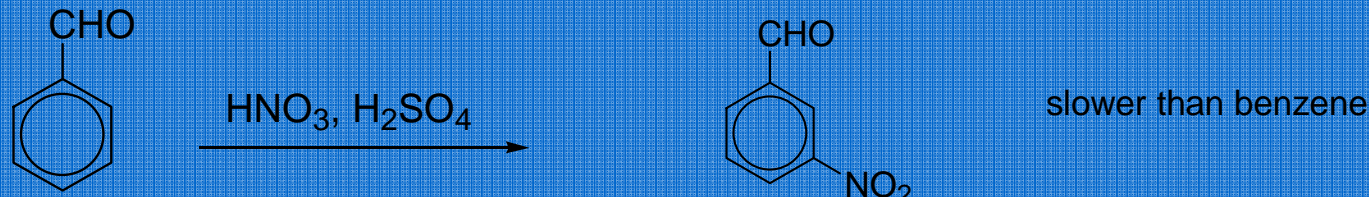
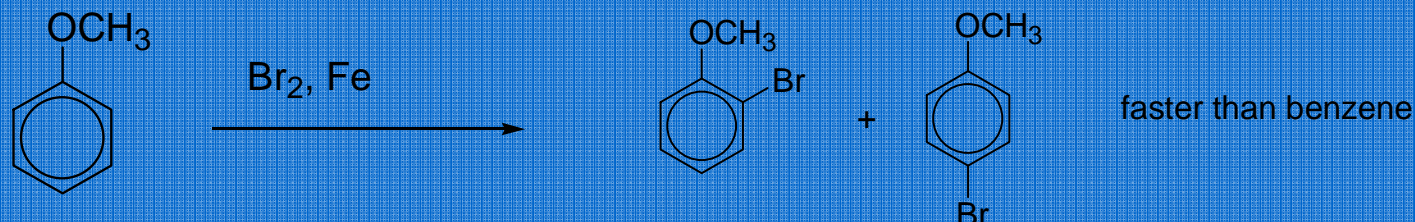
-NO<sub>2</sub>

deactivates the benzene ring towards EAS

directs substitution to the *meta*- position

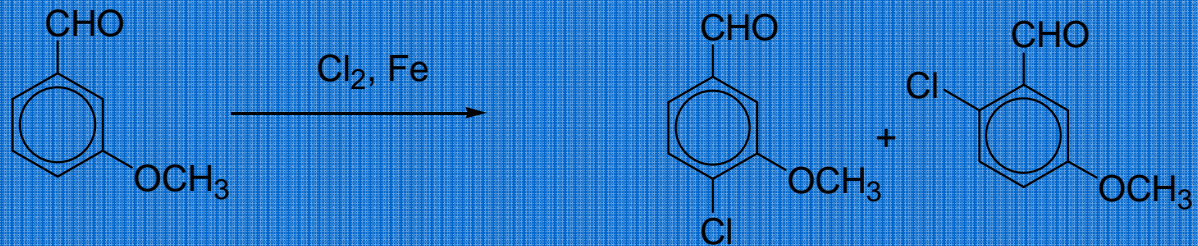
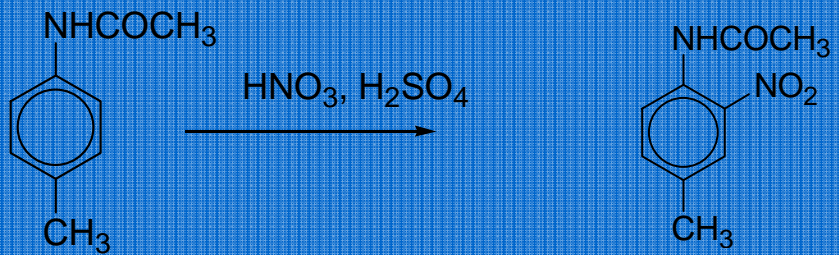
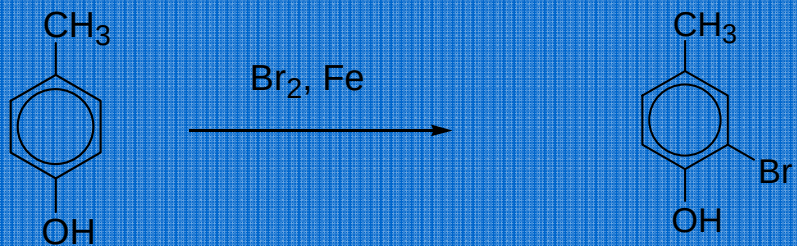
# Common substituent groups and their effect on EAS:





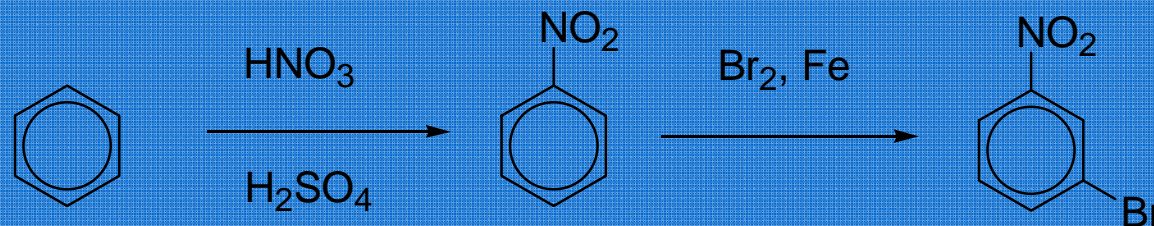
If there is more than one group on the benzene ring:

1. The group that is more activating (higher on “the list”) will direct the next substitution.
2. You will get little or no substitution between groups that are *meta*- to each other.

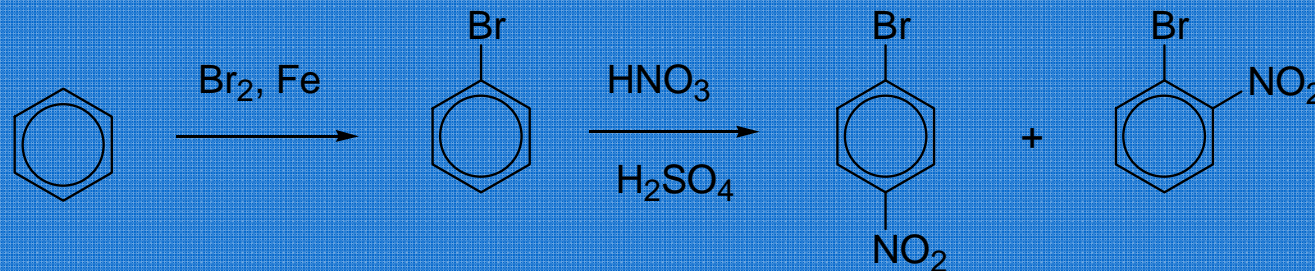


## Orientation and synthesis. Order is important!

synthesis of *m*-bromonitrobenzene from benzene:



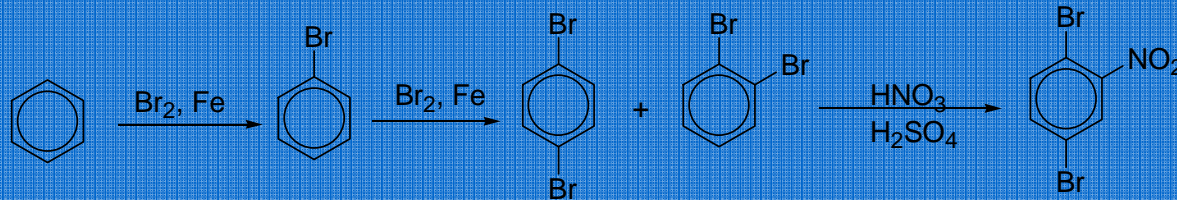
synthesis of *p*-bromonitrobenzene from benzene:



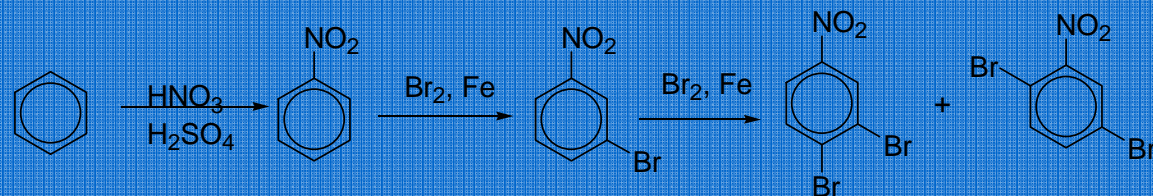
You may assume that you can separate a pure *para*-isomer from an *ortho-/para*- mixture.

note: the assumption that you can separate a pure para isomer from an ortho/para mixture does not apply to any other mixtures.

synthesis of 1,4-dibromo-2-nitrobenzene from benzene

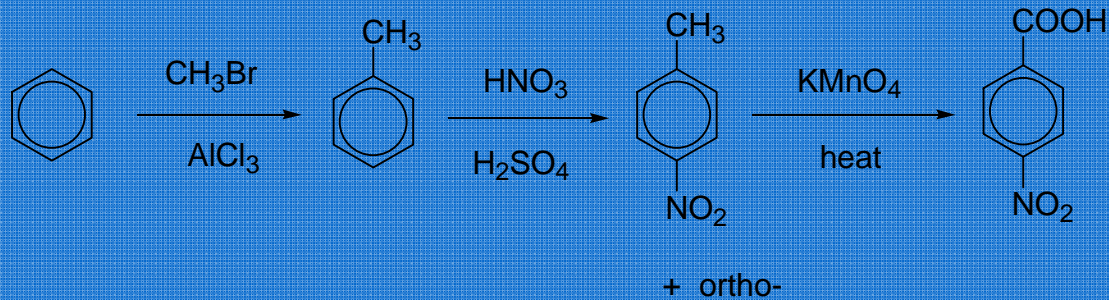
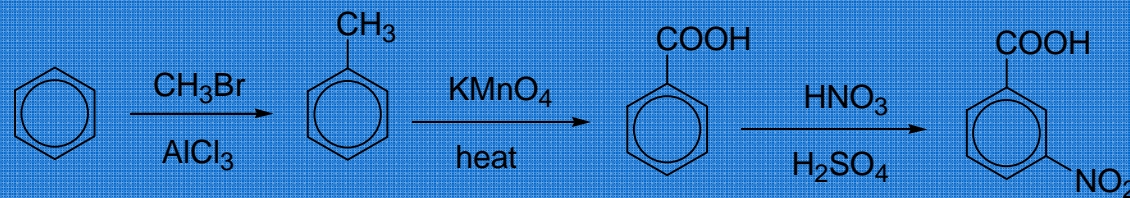
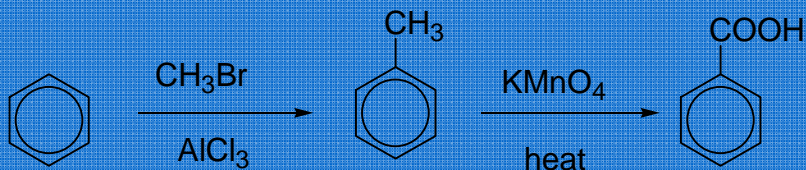


separate pure para isomer from ortho/para mixture

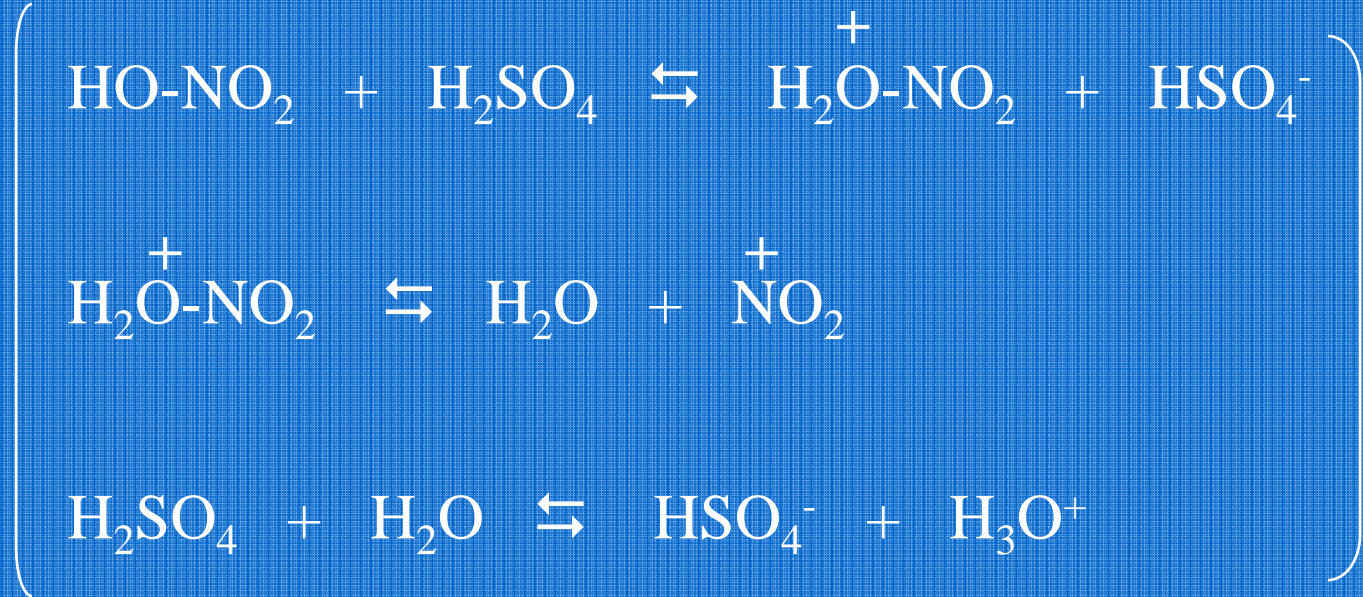


cannot assume that these can be separated!

## synthesis of benzoic acids by oxidation of $-\text{CH}_3$



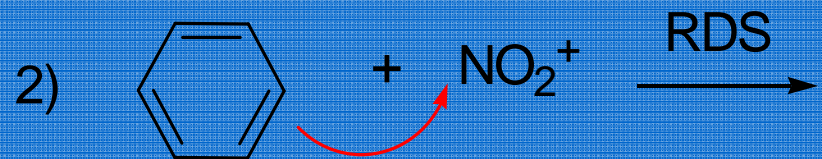
## nitration



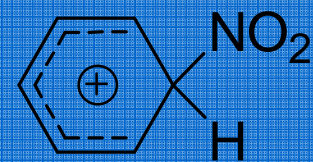
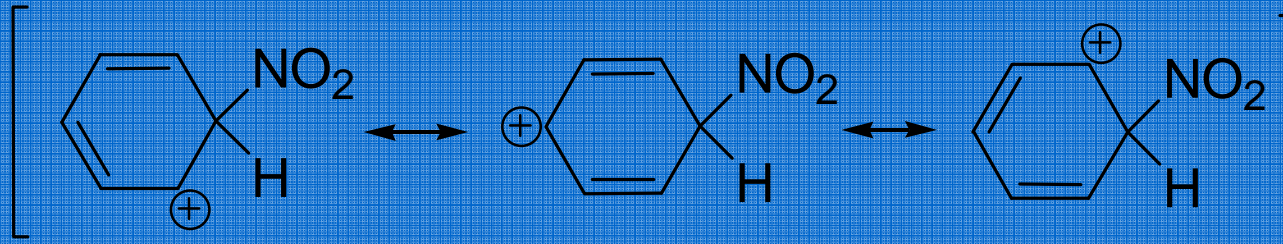
## Nitration



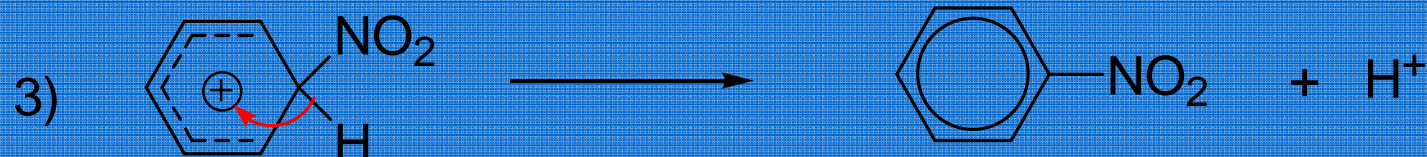
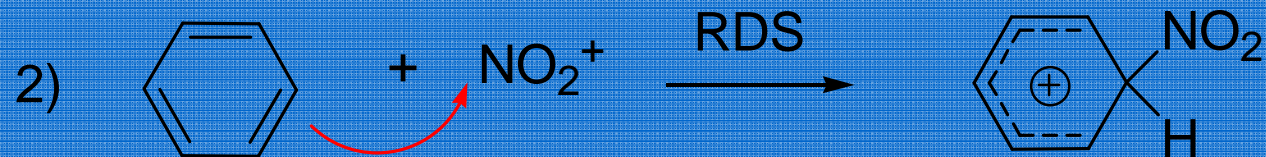
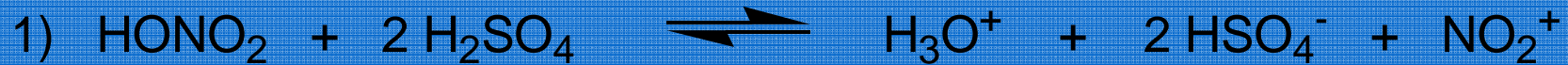
electrophile



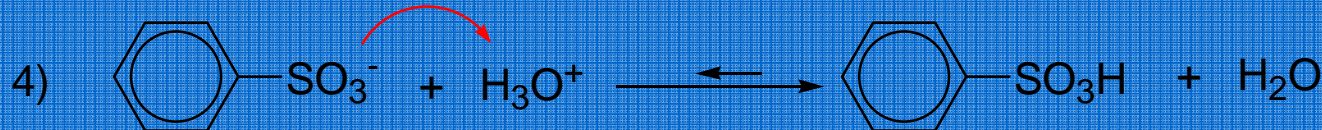
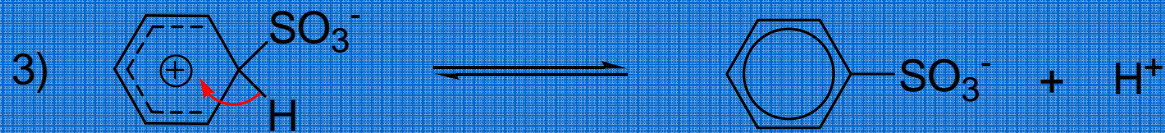
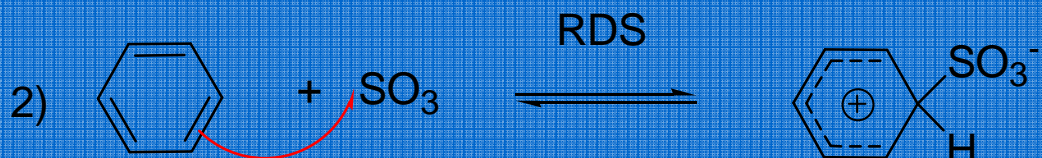
# Resonance



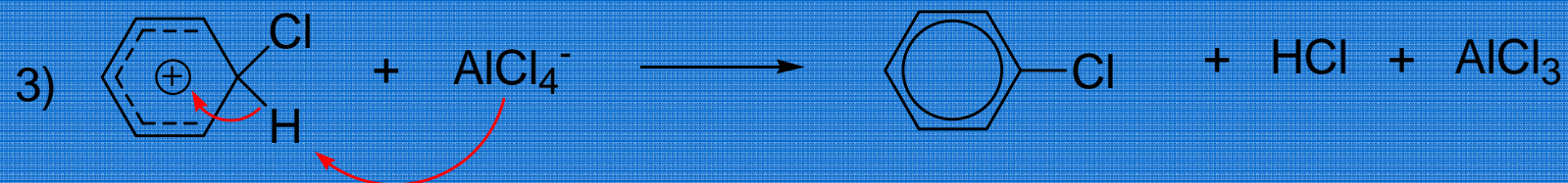
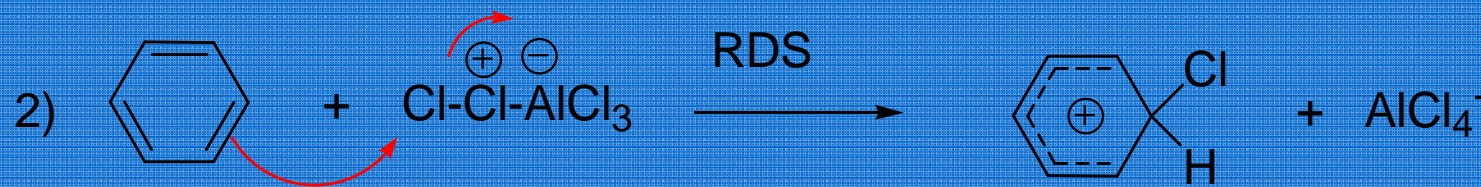
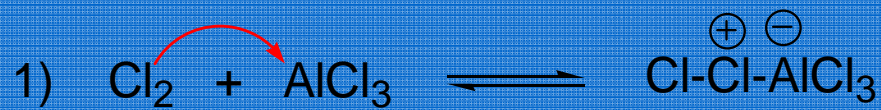
## Mechanism for nitration:



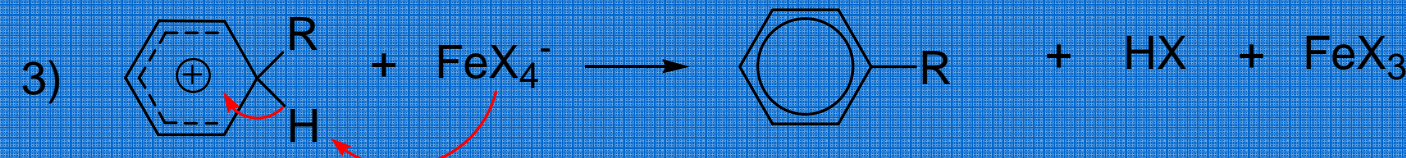
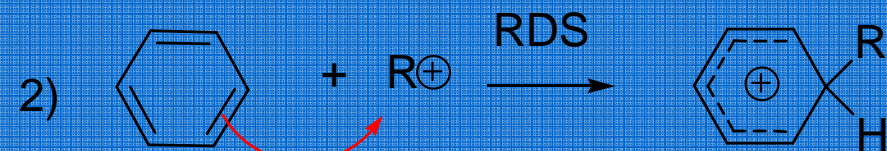
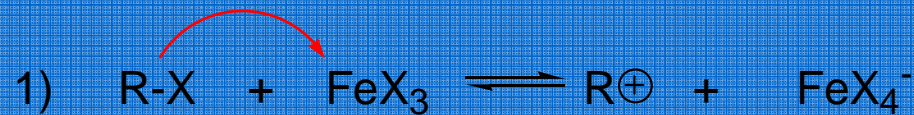
## Mechanism for sulfonation:



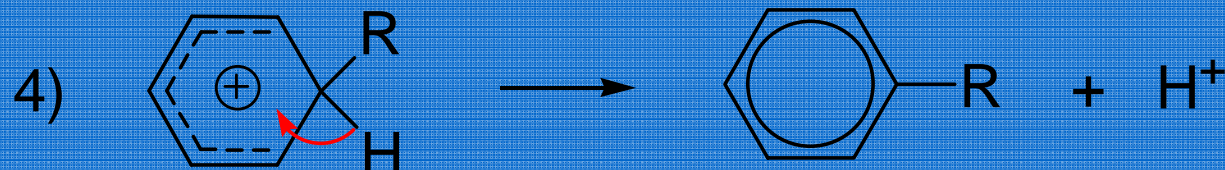
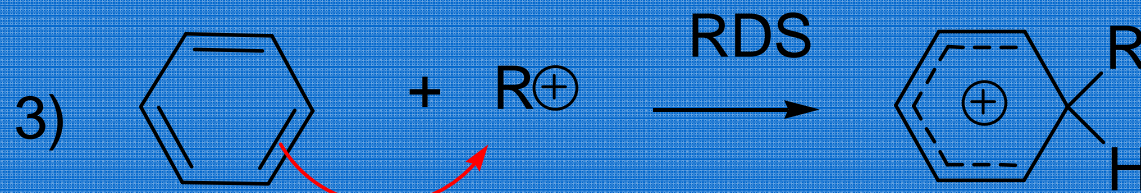
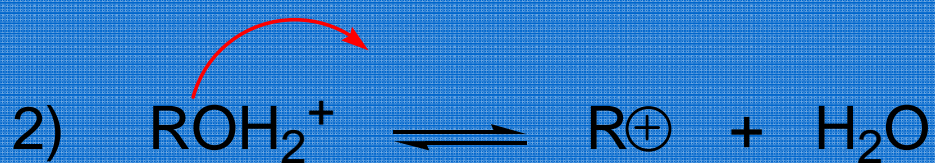
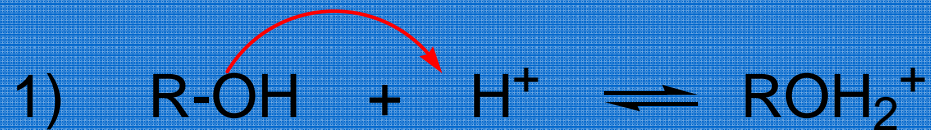
## Mechanism for halogenation:



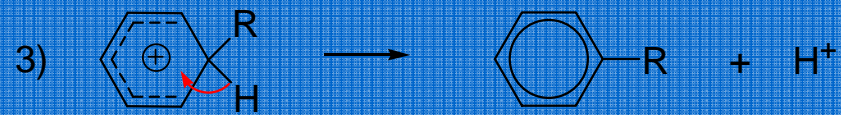
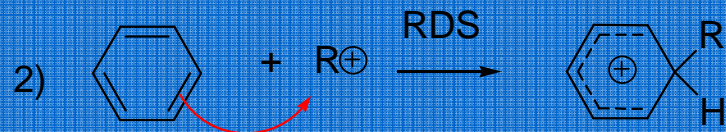
## Mechanism for Friedel-Crafts alkylation:



## Mechanism for Friedel-Crafts with an alcohol & acid

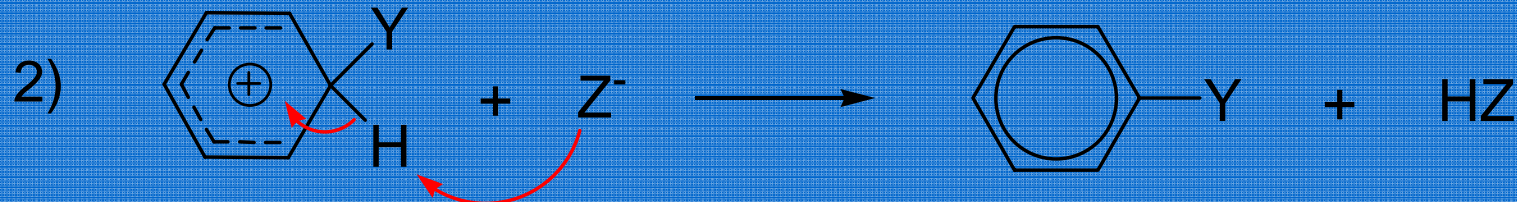
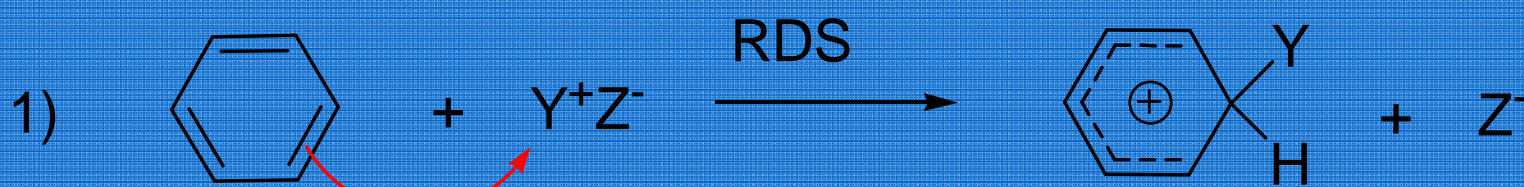


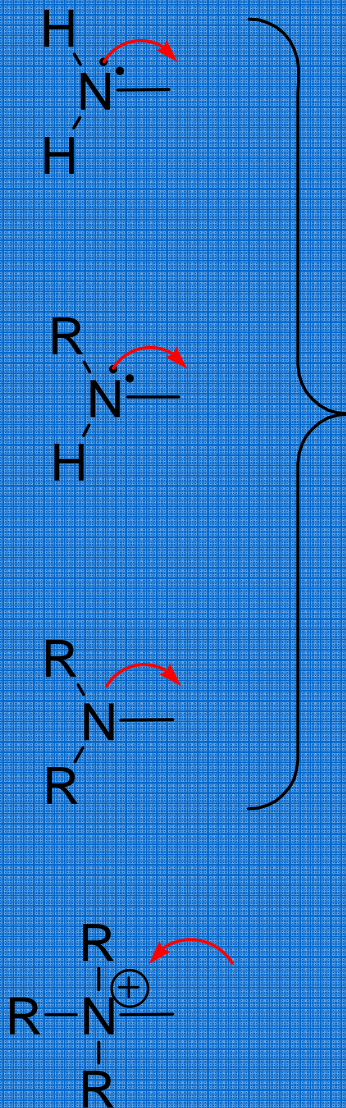
## Mechanism for Friedel-Crafts with alkene & acid:



electrophile in Friedel-Crafts alkylation = carbocation

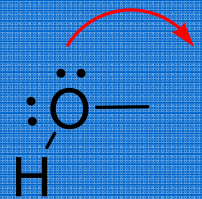
## “Generic” Electrophilic Aromatic Substitution mechanism:



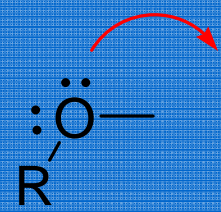


unshared pair of electrons on the nitrogen  
**resonance donating groups**  
(weaker inductive withdrawal)

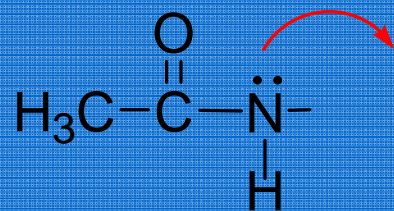
**strong inductive withdrawal**  
(no unshared pair of electrons on the  
nitrogen & no resonance possible)



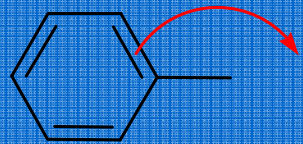
**resonance donation**  
(weaker inductive withdrawal)



**resonance donation**  
(weaker inductive withdrawal)



**resonance donation**  
(weaker inductive withdrawal)



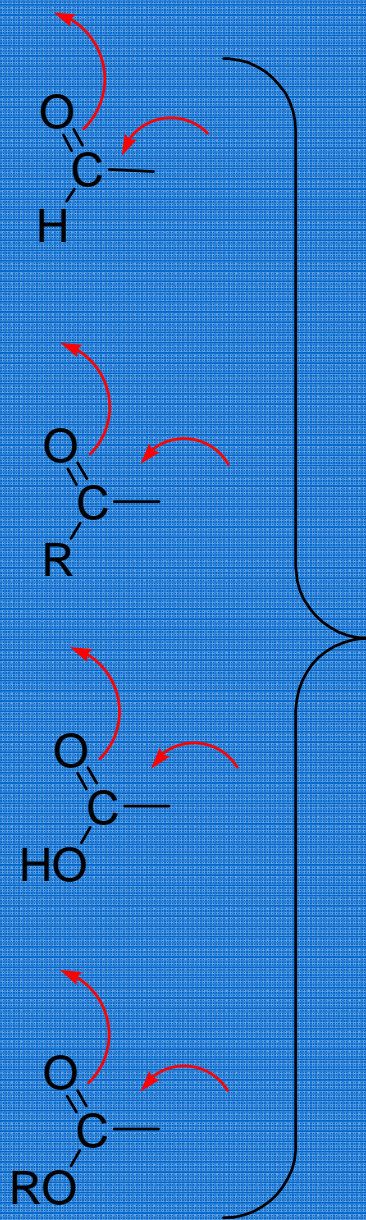
resonance **donation**



inductive **donation**  
sp<sup>3</sup>      sp<sup>2</sup> ring carbon



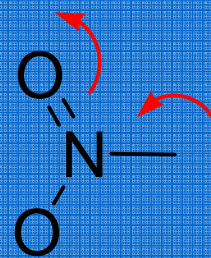
inductive **withdrawal**



**resoance withdrawal and  
inductive withdrawal**

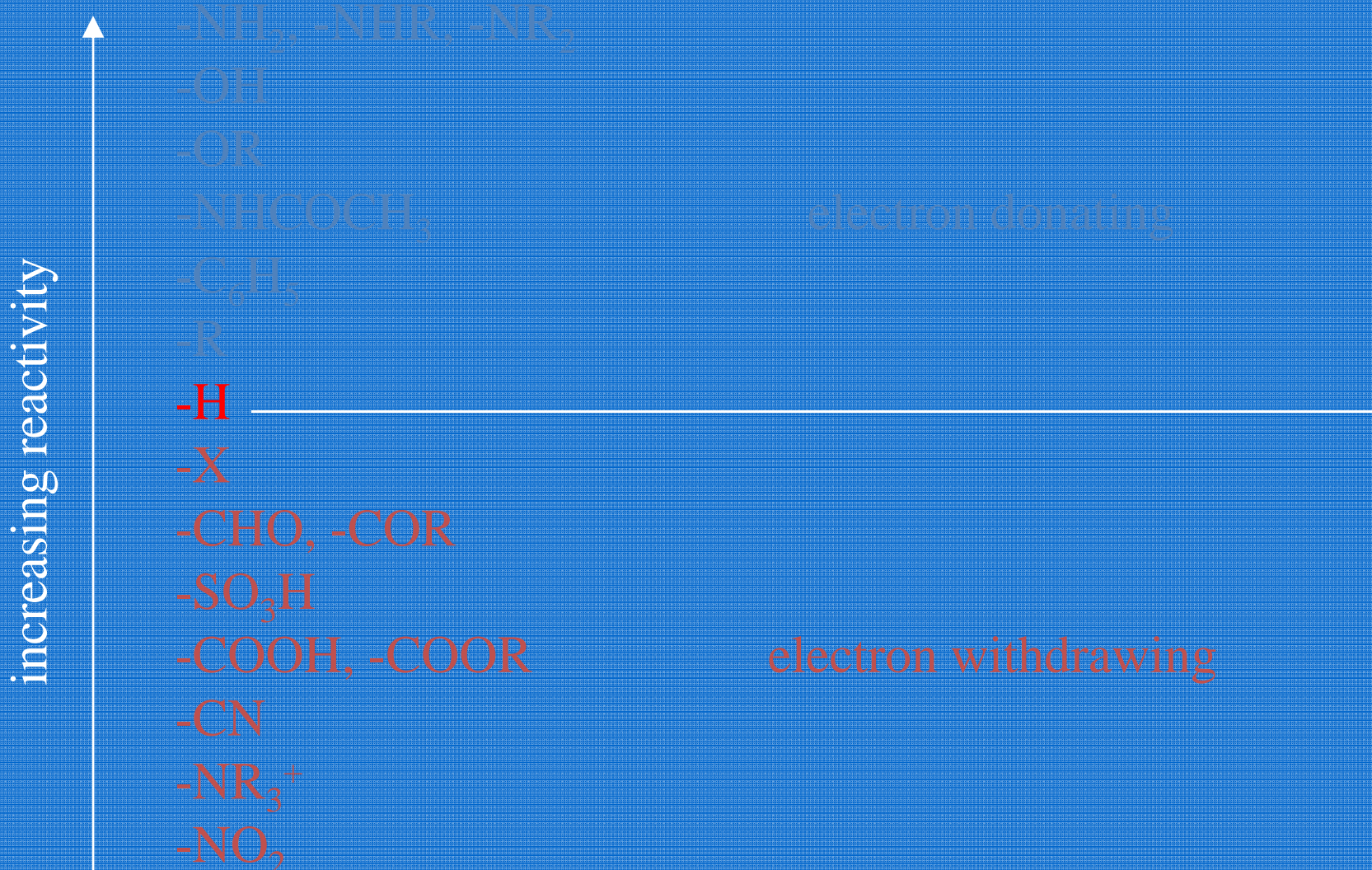


resonance and  
inductive **withdrawal**



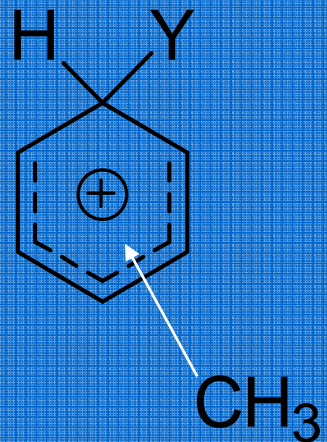
resonance and  
inductive **withdrawal**

# Common substituent groups and their effect on **reactivity** in EAS:



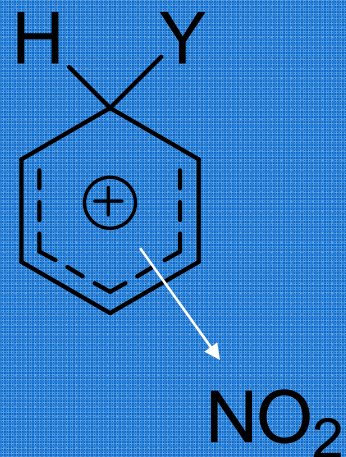
Electron donating groups activate the benzene ring to electrophilic aromatic substitution.

1. electron donating groups increase the electron density in the ring and make it more reactive with electrophiles.
2. electron donation stabilizes the intermediate carbocation, lowers the Eact and increases the rate.



Electron withdrawing groups deactivate the benzene ring to electrophilic aromatic substitution.

1. electron withdrawing groups decrease the electron density in the ring and make it less reactive with electrophiles.
2. electron withdrawal destabilizes the intermediate carbocation, raising the Eact and slowing the rate.

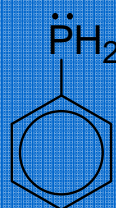




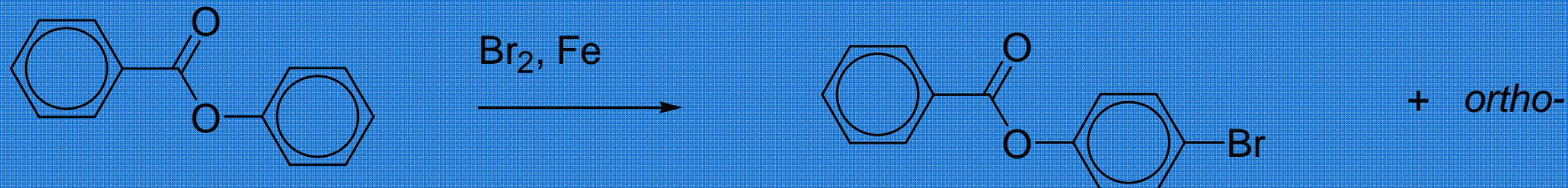
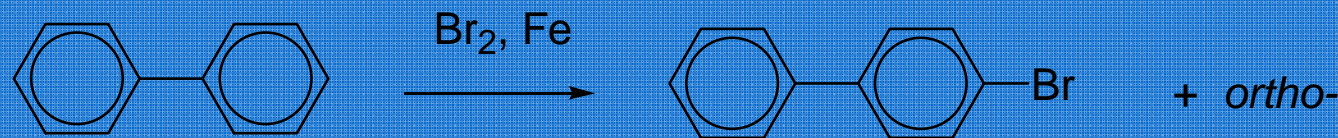
electron withdrawing = deactivating & *meta*-director



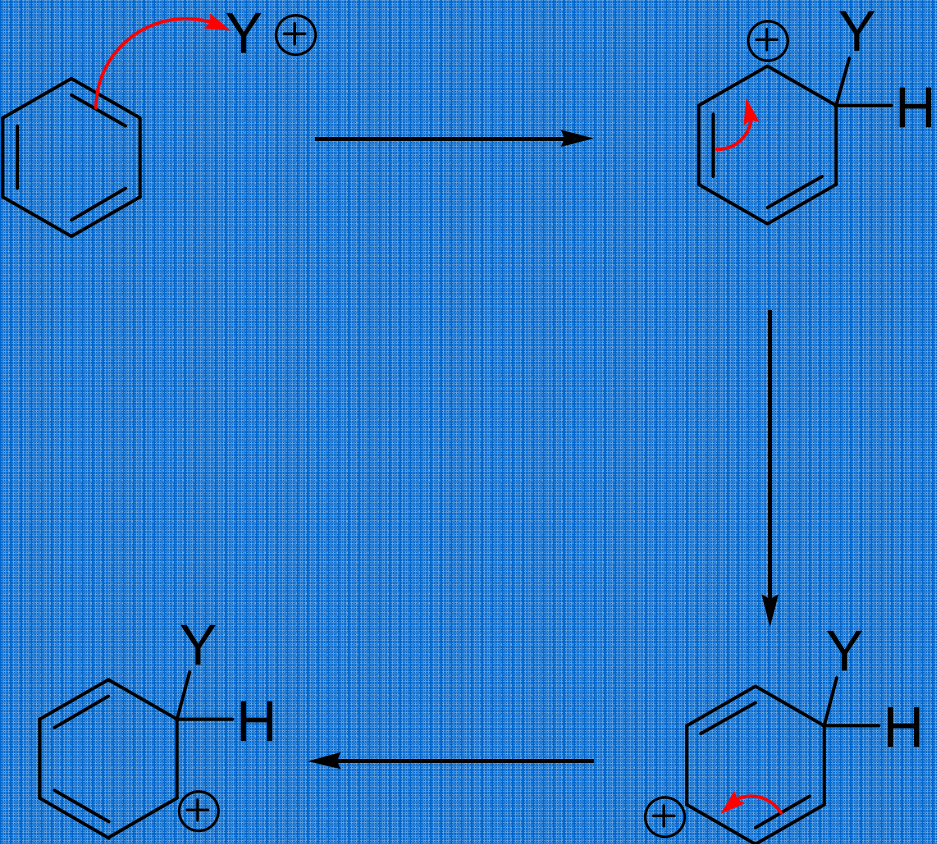
electron withdrawing = deactivating & *meta*-director

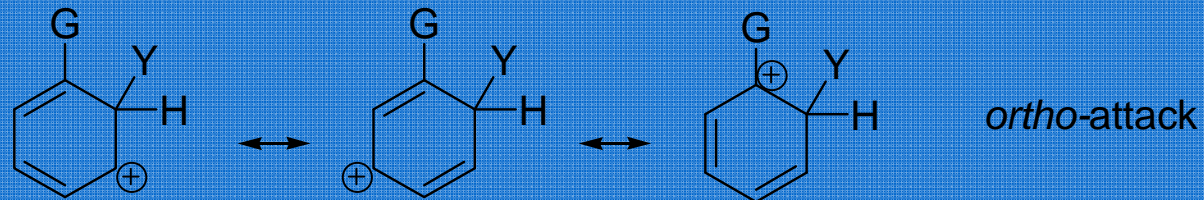


electron donating = activating & *ortho-/para*-director

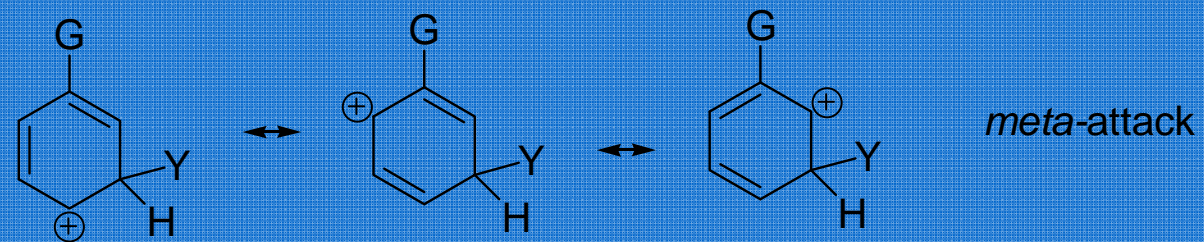


## How to draw resonance structures for EAS

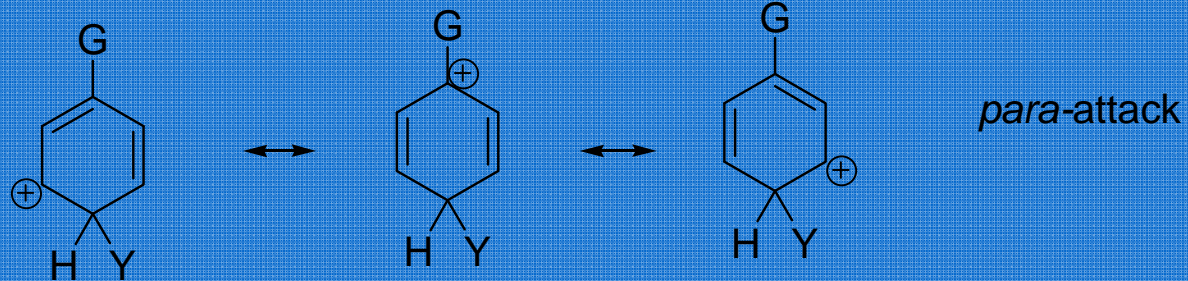




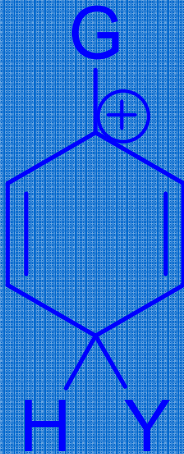
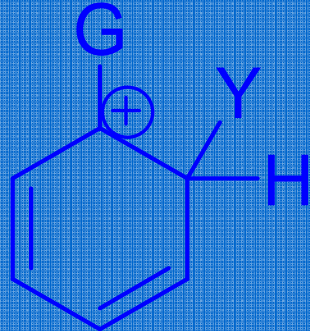
*ortho-attack*



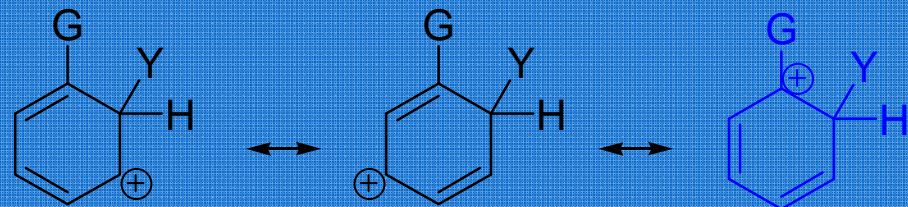
*meta-attack*



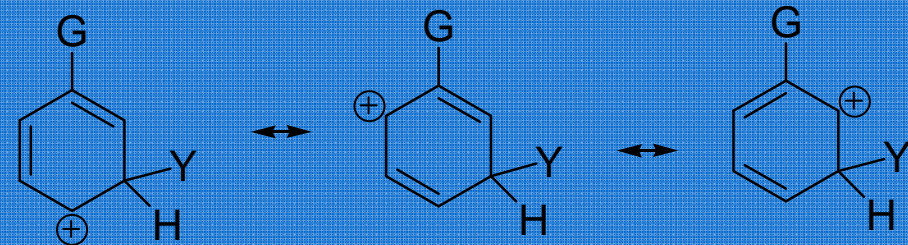
*para-attack*



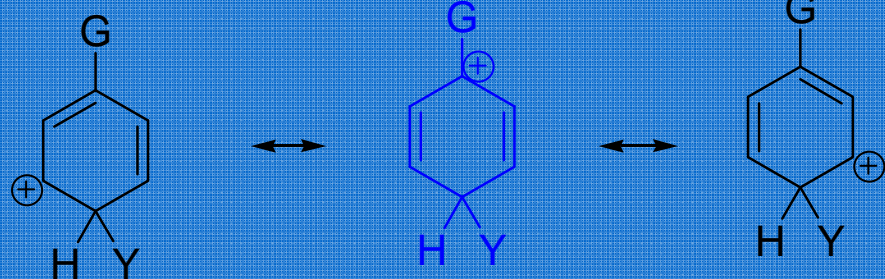
If G is an electron donating group, these structures are especially **stable**.



*ortho-attack*



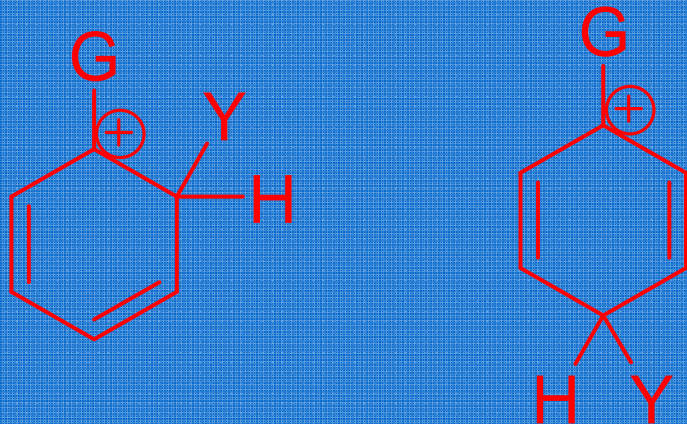
*meta-attack*



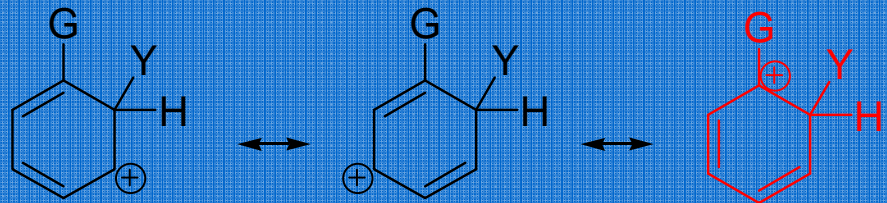
*para-attack*

Electron donating groups stabilize the intermediate carbocations for *ortho*- and *para*- in EAS more than for *meta*-. The Eact's for *ortho-/para*- are lower and the rates are faster.

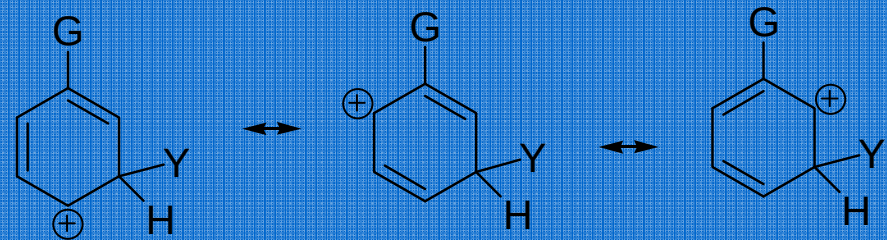
**Electron donating groups direct *ortho-/para*- in EAS**



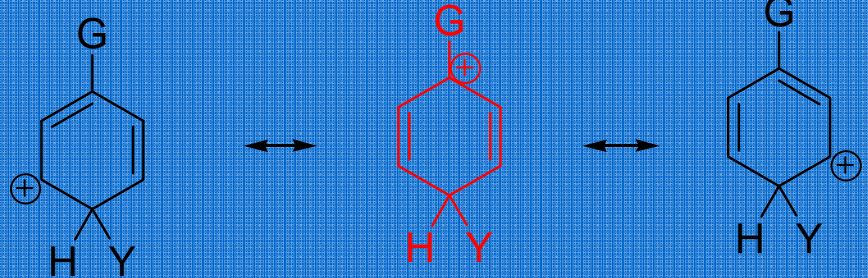
If G is an electron withdrawing group, these structures are especially **unstable**.



*ortho-attack*



*meta-attack*



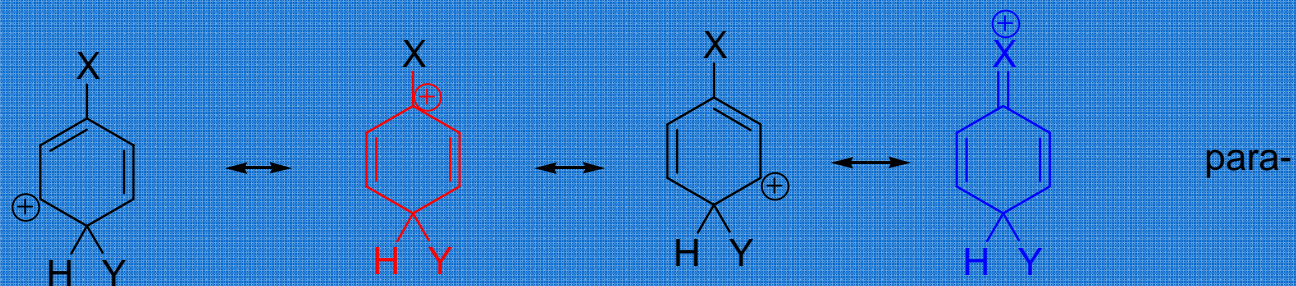
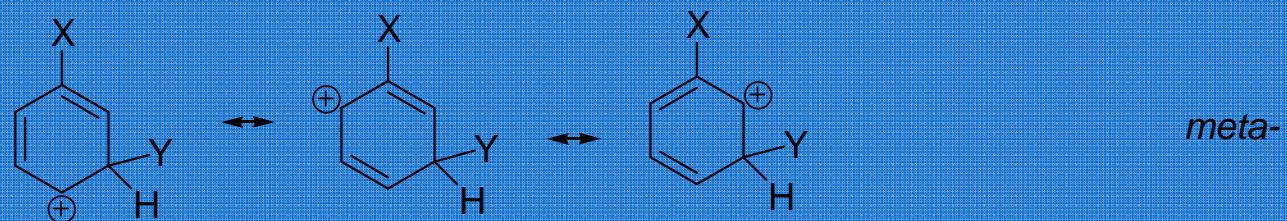
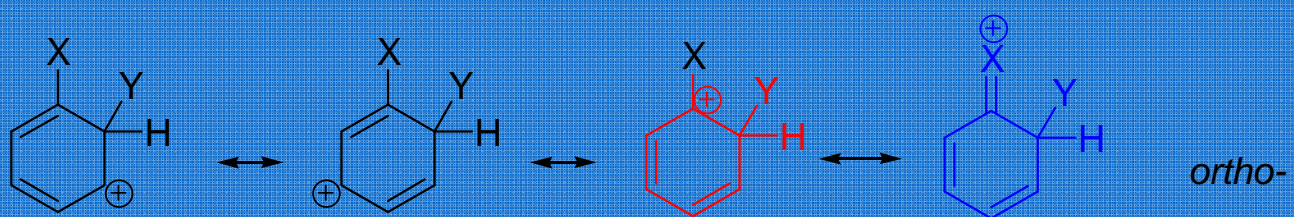
*para-attack*

Electron withdrawing groups destabilize the intermediate carbocations for *ortho*- and *para*- in EAS more than for *meta*-. The Eact's for *ortho-/para*- are higher and the rates are slower.

**Electron withdrawing groups direct *meta*- in EAS**

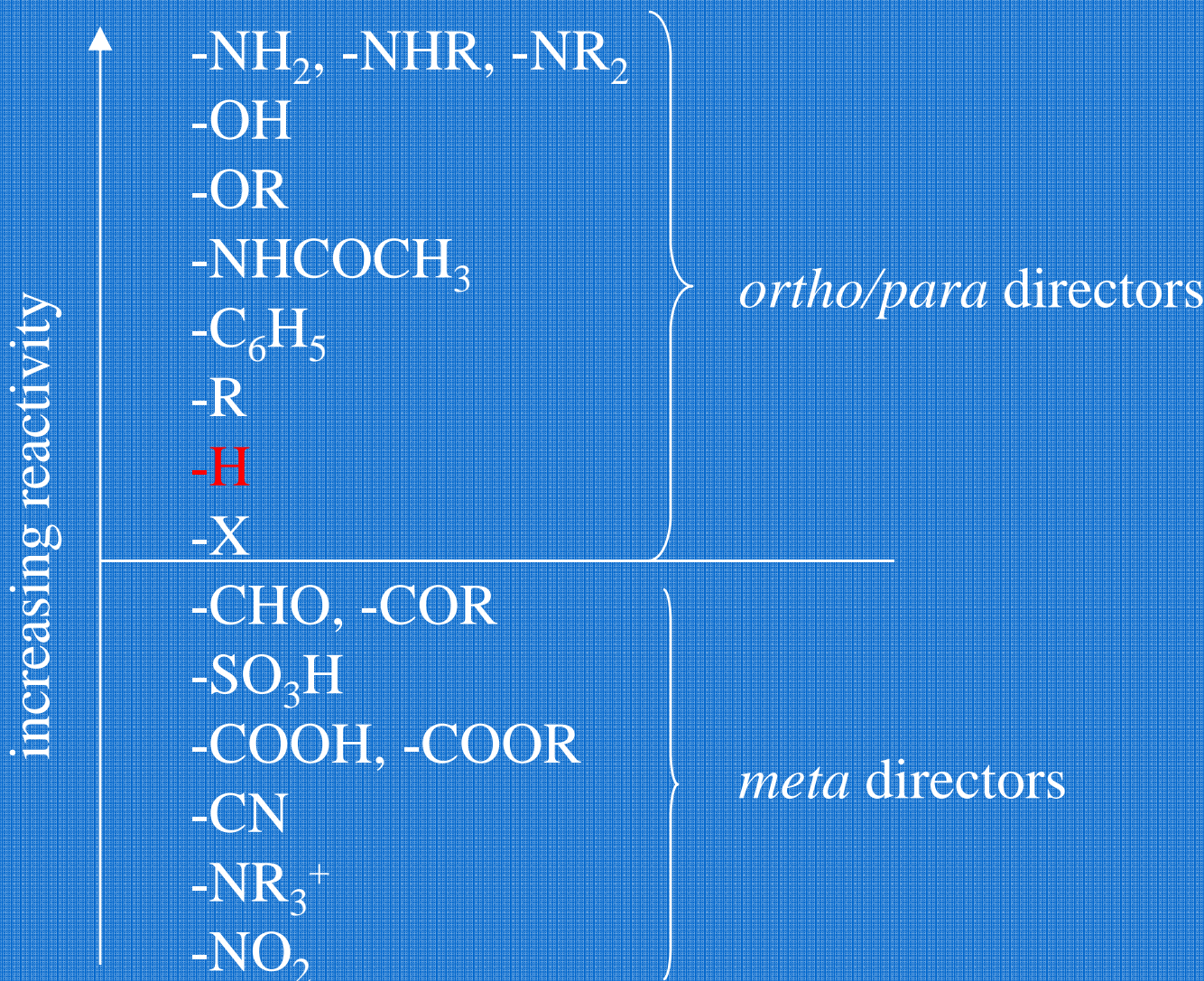
Halogens are electron withdrawing but are ortho/para directing in EAS.

The halogen atom is unusual in that it is highly electronegative but also has unshared pairs of electrons that can be resonance donated to the carbocation.



halogens are deactivating in EAS but direct *ortho* and *para*

# Common substituent groups and their effect on EAS:



**THANK YOU**