Organic Chemistry Aromatic Compounds

#### Arenes:

compounds containing both aliphatic and aromatic parts.

- Alkylbenzenes
- Alkenylbenzenes
- Alkynylbenzenes
- Etc.

Emphasis on the effect that one part has on the chemistry of the other half.

**Reactivity & orientation** 

### **Aromatic Hydrocarbons**



Aliphatic compounds: open-chain compounds and ring compounds that are chemically similar to open-chain compounds. Alkanes, alkenes, alkynes, dienes, alicyclics, etc.

Aromatic compounds: unsaturated ring compounds that are far more stable than they should be and resist the addition reactions typical of unsaturated aliphatic compounds. Benzene and related compounds.

### Nomenclature – common names



### Nomenclature – common names



# Systematic Nomenclature

- Monosubstituted benzenes
- Hydrocarbon with *benzene as parent*
- $C_6H_5Br = bromobenzene$
- $C_6H_5NO_2 =$  nitrobenzene
- $C_6H_5CH_2CH_2CH_3 = propylbenzene$



#### others named as "alkylbenzenes":







o-diethylbenzene

n-butylbenzene

# The Phenyl Group

- When a benzene ring is a substituent, the term phenyl is used (for C<sub>6</sub>H<sub>5</sub><sup>±</sup>)
- You may also see "Ph" or " $\phi$ " in place of "C<sub>6</sub>H<sub>5</sub>"
- "**Benzyl**" refers to " $C_6 H_5 C H_2^{\pm}$



#### Use of phenyl $C_6H_5 = "phenyl"$





2-methyl-3-phenylheptane

1,2-diphenylethane

#### do not confuse phenyl ( $C_6H_5$ -) with benzyl ( $C_6H_5CH_5$ -)

### Nomenclature: Side Chains

If side chain has < 6 carbons</li>
 Alkyl benzene

If side chain has > 6 carbons
 – Phenyl alkane

#### Alkenylbenzenes, nomenclature:

Special name



Rest are named as substituted alkenes



3-phenylpropene (allylbenzene)



#### Alkynylbenzenes, nomenclature:





phenylacetylene

phenylethyne

5-phenyl-2-hexyne

#### Alcohols, etc., nomenclature:



1-phenylethanol

a-phenylethyl alcohol



#### benzyl alcohol

CH<sub>2</sub>CH<sub>2</sub>-Cl



1-chloro-2-phenylethane

cyclohexylbenzene

phenylcyclohexane

### Nomenclature Disubstituted Benzene

Relative positions on a benzene ring *ortho- (o)* on adjacent carbons (1,2) *meta- (m)* separated by one carbon (1,3) *para- (p)* separated by two carbons (1,4)

Describes reaction patterns ("occurs at the



### Nomenclature More Than Two Substituents

- Choose numbers to get lowest possible values
- List substituents alphabetically with hyphenated numbers
- Common names, such as "toluene" can serve as root name (as in TNT)







- Three double bonds
- Unreactive towards normal reagents (compare to alkenes)
- Very stable
- Why?
- How can we get benzene to react?
- Can we control these reactions?

### **Observations: Reactions of Benzene**

- Benzene reacts slowly with Br<sub>2</sub>
- Product is bromobenzene
- Substitution Product
- Addition products are not observed.



# Stability of Benzene

- KMnO<sub>4</sub>
  - Reacts with alkenes
  - No reaction with benzene
- HCl
  - Reacts with alkenes
  - No reaction with benzene
- HBr
  - Reacts with alkenes
  - No reaction with benzene

# Stability of Benzene

### Heat of Hydrogenation data



### C-C bond length



- Electrostatic potential
- Electron density at C is the same



### planar

### • August Kekule proposed:



1,3,5-cyclohexatriene structureExplained single monobromo product

### • Dibromobenzene



### Issue was resolved by Kekule



- Explains the observed products
- Does not explain
  - Unreactive nature of benzene
  - Observation of only substitution products
- A triene
  - As reactive as any alkene
  - Would give addition products
  - Not expected to be more stable

Resonance Hybrid



- Not
- Never
- -6.023 X 10<sup>23</sup> points



# Stability of Benzene

- MO Description
- 6 p atomic orbitals combine in cyclic manner
- Generate 6 molecular orbitals



### Key Ideas on Benzene

- Unusually stable
- heat of hydrogenation 150 kJ/mol lower than a cyclic triene
- Planar hexagon:
- bond angles are 120°
- carbon–carbon bond lengths 139 pm
- Undergoes substitution not addition
- Resonance hybrid
- One more important factor is the number of electrons in the cyclic orbital

# Aromaticity

#### • E Huckel (1931)

- Aromaticity is a property of certain molecules
- Chemistry would be similar to benzene
- Meet the following criteria
  - Planar
  - Mono cyclic system
  - Conjugated pi system
  - Contains 4n + 2 п electrons
- Can apply rules to variety of compounds and determine aromatic nature.
- Led to wild chase to make compounds
  - Met the rules
  - Violated the rules

# Aromaticity and the 4n + 2 Rule

- Huckel's rule, based on calculations a planar cyclic molecule with alternating double and single bonds has aromatic stability if it has *4n+ 2 π electrons (n is 0,1,2,3,4*)
- For n=1: 4n+2 = 6

**benzene** is stable and the electrons are delocalized



Compounds With 4n п Electrons Are Not Aromatic (May be Anti-aromatic)

- Planar, cyclic molecules with *4 п п* electrons are much *less* stable than expected (anti-aromatic)
- They will distort out of plane and behave like ordinary alkenes
- 4- and 8-electron compounds are not delocalized
- Alternating single and double bonds

# Cyclobutadiene

 Cyclobutadiene is so unstable that it dimerizes by a self-Diels-Alder reaction at low temperature



# Cyclooctatetraene

- Cyclooctatetraene has four double bonds
- Behaves as if it were 4 separate alkenes
- It reacts with Br<sub>2</sub>, KMnO<sub>4</sub>, and HCl
- Non-planar structure





### **Aromatic Heterocycles**

- Heterocyclic compounds contain elements other than carbon in a ring, such as N,S,O,P
- There are many heterocyclic aromatic compounds
- Cyclic compounds that contain only carbon are called carbocycles
- Nomenclature is specialized
- Four are important in biological chemistry

# Pyridine

- A six-membered heterocycle with a nitrogen atom in its ring
- п electron structure resembles benzene (6 electrons)
- The nitrogen lone pair electrons are not part of the aromatic system (perpendicular orbital)
- Pyridine is a relatively weak base compared to normal amines but protonation does not affect aromaticity



# Pyrrole

- A five-membered heterocycle with one nitrogen
- Four *sp*<sup>2</sup>-hybridized carbons with 4 *p* orbitals perpendicular to the ring and 4 p electrons
- Nitrogen atom is *sp*<sup>2</sup>-hybridized, and lone pair of electrons occupies a *p* orbital (6 п electrons)
- Since lone pair electrons are in the aromatic ring, protonation destroys aromaticity, making pyrrole a very weak


# Pyrimidine

- Similar to benzene
- 3 pi bonds
- 4n + 2 pi electrons
- aromatic



### Imidazole

- Similar to pyrrole
- Pair of non-bonding electrons on N used
- 4n + 2 pi electrons



### Thiophene and Furan

- Non-bonding electrons are used
- 4n + 2 pi electrons





#### Substitution Reactions of Benzene

- Benzene is aromatic: a cyclic conjugated compound with 6 п electrons
- Reaction with E<sup>+</sup> Leads to Substitution
- Aromaticity of Benzene is retained



#### **Aromatic Substitutions**

- The proposed mechanism for the reaction of benzene with electrophiles involves a cationic intermediate
- first proposed by G. W. Wheland of the University of Chicago
- Often called the Wheland intermediate



# Chemistry of the Intermediate

- Loss of a proton leads to rearomatization and substitution
- Loss of E<sup>+</sup> returns to starting material



# Halogenation

- Add Cl, Br, and I
- Must use Lewis acid catalyst
- F is too reactive and gives very low yields



# **Biological Halogenation**

- Accomplished during biosynthesis of
- thyroxine



#### **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



#### Nitrobenzenes: Precursors to Anilines

- Nitric acid destroys alkenes through [O]
- In sulfuric acid reacts with benzene giving nitrobenzene
- Nitrobenzene may be reduced to aniline
- Aniline useful precursors to many industrially important organic compounds

#### **Important Anilines**





### Aromatic Dyes



- William Henry Perkin
- Age 17 (1856)
- Undergraduate student in medicine
- Reacted aniline with potassium dichromate
- Tarry mess

# Aromatic Dyes



- Mauve a purple color
- Dyed white cloth
- Patented material and process
- First chemical company

#### Mauveines -> 1994 !





# Some Aniline Chemistry

 Anilines readily react with nitrous acid



- Diazonium salts
  Coupling reaction
  - giving an azo compound
- Dyes and sulfa drugs

# **Aniline Chemistry**



#### How do we make sulfuric acid?

- H<sub>2</sub>SO<sub>4</sub> least expensive manufactured chemical
- S (mined pure) +  $O_2$  SO<sub>3</sub> • SO<sub>3</sub> + H<sub>2</sub>O H<sub>2</sub>SO<sub>4</sub>
- Continue adding SO<sub>3</sub> gives

Fuming sulfuric acid: H<sub>2</sub>SO<sub>4</sub>/SO<sub>3</sub>

#### **Aromatic Sulfonation**

- Substitution of H by SO<sub>3</sub> (sulfonation)
- Reaction with a mixture of sulfuric acid and SO<sub>3</sub>
- Reactive species is sulfur trioxide or its conjugate acid
- Reaction occurs via Wheland intermediate and is reversible



#### **Benzene Sulfonic Acid**

- Manufacture of Ion Exchange Resins
  - Water softening
  - Water purification
  - Environmental restoration (removal of toxic metal ions)

#### Benzene Sulfonic Acid

- Starting material for Sulfa Drugs
- First useful antibiotics



# Hydroxylation

- Direct hydroxylation is difficult in lab
- Indirect method uses sulfonic acid



# **Biological Hydroxylation**

- Frequently conducted
- Example,



#### Coenzyme necessary

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

- Aromatic substitution of a R<sup>+</sup> for H
- Aluminum chloride promotes the formation of the carbocation
- Wheland intermediate forms



#### Limitations of the Friedel-Crafts Alkylation

- Only *alkyl* halides can be used (F, Cl, I, Br)
- Aryl 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



#### Limitations

Multiple alkylations occur because the first alkyl group activates the ring



#### polyalkylation



The alkyl group activates the ring making the products more reactive that the reactants leading to polyalkylation. Use of excess aromatic compound minimizes polyalkylation in the lab.

#### Limitations

- Carbocation Rearrangements During Alkylation
- Similar to those that occur during electrophilic additions to alkenes
- Can involve H or alkyl shifts





#### **Related Reactions**

#### Chloromethylation



#### **Related Reaction**

- Acylation of Aromatic Rings
- Reaction of an acid chloride (RCOCl) with an aromatic ring in the presence of AlCl<sub>3</sub> introduces the **acyl group**,
- \_COR
- Benzene with acetyl chloride yields acetophenone
- Acyl group deactivates ring
- Reaction stops after one group is added



# **Biological Alkylations**

- Common reaction
- No AlCl<sub>3</sub> present
- Utilizes an organodiphosphate
- Dissociation is facilitated by Mg<sup>+2</sup>
- Important reaction in biosynthesis of Vitamin K<sub>1</sub>



### **Ring Substitution Effects**

- Activation and deactivation of ring
  - Alkyl activates the ring
  - Acyl deactivates the ring
- Activating Groups
  - group promotes substitution faster than benzene
- Deactivating Groups

group promotes substitution slower than benzene

#### Activating and Deactivating Groups

- Activating groups
  - electron donating groups
  - stabilizes the carbocation intermediate
  - activates through induction or resonance
- Deactivating groups
  - electron withdrawing groups
  - destabilizes the carbocation intermediate
  - deactivates through induction or resonance

**Common substituent groups and their effect on EAS:** 



#### Activating and Deactivating Groups



# **Origins of Substituent Effects**

- Inductive effect withdrawal or donation of electrons through a  $\sigma$  bond
- 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  $\sigma$  bond connected to ring
- Alkyl groups *donate* electrons through  $\sigma$  bond





Alkyl group; inductively electron-donating

#### (X = F, Cl, Br, I)

The groups attached to the aromatic rings are inductively electronwithdrawing because of the polarity of their bonds. © 2004 Thomson/Brooks Cole
#### **Resonance Effects: Electron Withdrawal**

- C=O, CN, NO<sub>2</sub> substituents withdraw electrons from the aromatic ring by resonance
- п electrons flow from the rings toward the substituent



Rings substituted by a group with an electron-withdrawing resonance effect have this general structure. © 2004 Thomson/Brooks Cole

#### **Resonance Effects: Electron Donation**

 Halogen, OH, alkoxyl (OR), and amino substituents *donate* electrons through resonance

OH

OR

п electrons flow from into the ring



x = Halogen

Rings substituted by a group with an electron-donating resonance effect have this general structure.

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NHa

# Consider the following data



## Analysis of Data

- Methoxy and Methyl
- Activating
- Ortho and para products
- Nitro and Carbomethoxy
- Deactivating
- Meta product
- Bromine
- Deactivating
- Ortho and para products

# **Ring Effects - Conclusions**

- Activating groups
- Substitution is faster than for benzene
- Groups direct substitution to o/p positions
- Deactivating Groups
- Substitution is slower than for benzene
- Groups direct substitution to m position
- Halogens
- Deactivate ring
- Substitution is slower than for benzene
- Groups direct substitution to o/p positions

## **Ring Effects – The Explanation**

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



# Important

#### You need to know this:



# **Oxidation of Benzene**

- Toluene is readily oxidized by reagents
- Benzene is inert to oxidizing agents
  - Benzene is toxic to humans
  - Benzene is a suspected carcinogen
- Cytochrom P
  - strong oxidant in Liver
  - Primary detoxification process used

# **Proposed Chemistry**



## **Biological Oxidations of Side Chains**

#### Biosynthesis of norepinephrine



enzyme = dopamine-beta-monooxygenase

### **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> and Na<sub>2</sub>Cr<sub>2</sub>O<sub>7</sub> if they have a C-H next to the ring
- Converts an alkylbenzene into a benzoic acid, Ar  $-R \rightarrow Ar -CO_2H$



#### Bromination of Alkylbenzene Side Chains

 Reaction of an alkylbenzene with N-bromo-succinimide (NBS) and benzoyl peroxide (radical initiator) introduces Br into the side chain



## **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)



## **Reduction of Aromatic Compounds**

 Aromatic Rings can be reduced using Li or Na metal dissolved in liquid ammonia



# Reduction of Aryl Alkyl Ketones

- Aromatic ring activates neighboring carbonyl group toward reduction
- Ketone is converted into an alkylbenzene by catalytic hydrogenation over Pd catalyst

