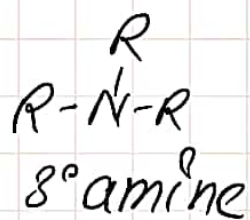
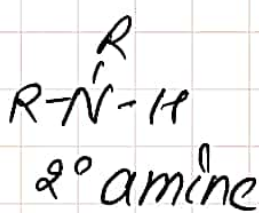
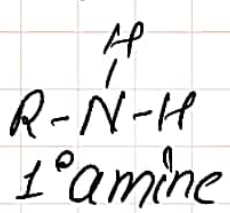
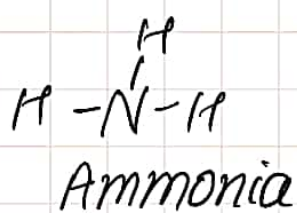


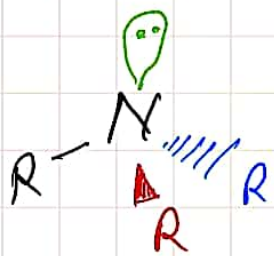
Amines

By Bharat Panchal Sir.

These are alkyl or aryl derivatives of Ammonia



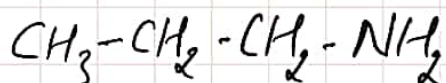
Structure of Amines



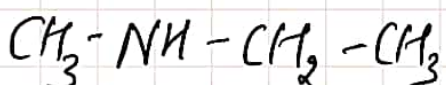
Structure - trigonal pyramidal

3 Bond Pair + 1 Lone Pair
Hybrid. $\rightarrow \text{sp}^3$

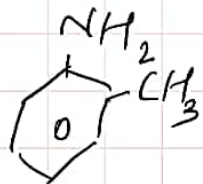
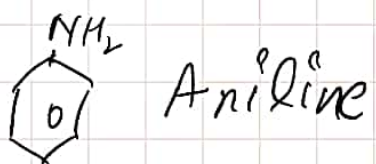
Nomenclature \Rightarrow



Propan-1-amine



N-Methyl ethanamine

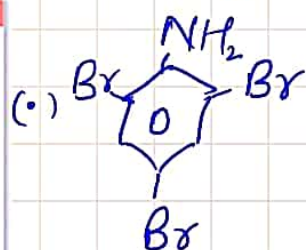


NCERT Ques.

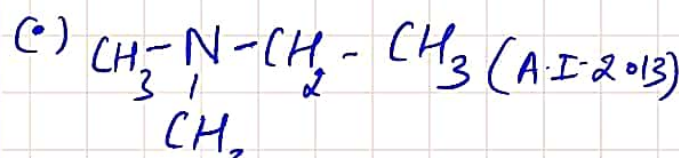
Amine	Common name	IUPAC name
$\text{CH}_3\text{-CH}_2\text{-NH}_2$	Ethylamine	Ethanamine
$\text{CH}_3\text{-CH}_2\text{-CH}_2\text{-NH}_2$	n-Propylamine	Propan-1-amine
$\begin{array}{c} \text{CH}_3\text{-CH}-\text{CH}_3 \\ \\ \text{NH}_2 \end{array}$	Isopropylamine	Propan-2-amine
$\begin{array}{c} \text{CH}_3-\text{N}-\text{CH}_2-\text{CH}_3 \\ \\ \text{H} \end{array}$	Ethylmethylamine	N-Methylethanamine
$\begin{array}{c} \text{CH}_3-\text{N}-\text{CH}_3 \\ \\ \text{CH}_3 \end{array}$	Trimethylamine	N,N-Dimethylmethanamine
$\text{C}_2\text{H}_5-\text{N}-\text{CH}_2-\overset{1}{\text{CH}_2}-\overset{2}{\text{CH}_2}-\overset{3}{\text{CH}_2}-\overset{4}{\text{CH}_3}$	N,N -Diethylbutylamine	N,N-Diethylbutan-1-amine
$\begin{array}{c} \text{NH}_2-\overset{1}{\text{CH}_2}-\overset{2}{\text{CH}}=\overset{3}{\text{CH}_3} \\ \\ \text{NH}_2 \end{array}$	Allylamine	Prop-2-en-1-amine
$\text{NH}_2-(\text{CH}_2)_6-\text{NH}_2$	Hexamethylenediamine	Hexane-1,6-diamine
$\begin{array}{c} \text{NH}_2 \\ \\ \text{C}_6\text{H}_5 \end{array}$	Aniline	Aniline or Benzenamine
$\begin{array}{c} \text{NH}_2 \\ \\ \text{C}_6\text{H}_4-\text{CH}_3 \end{array}$	<i>o</i> -Toluidine	2-Methylaniline
$\begin{array}{c} \text{NH}_2 \\ \\ \text{C}_6\text{H}_4-\text{Br} \\ \\ \text{N}(\text{CH}_3)_2 \end{array}$	<i>p</i> -Bromoaniline	4-Bromobenzenamine or 4-Bromoaniline
$\begin{array}{c} \text{NH}_2 \\ \\ \text{C}_6\text{H}_5-\text{N}(\text{CH}_3)_2 \end{array}$	N,N -Dimethylaniline	N,N -Dimethylbenzenamine

P.Y.Q

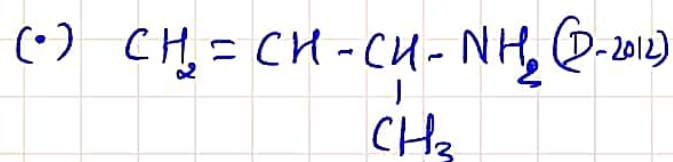
(D-2016)



2,4,6 - Tribromoaniline



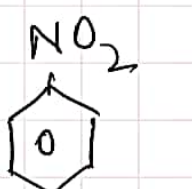
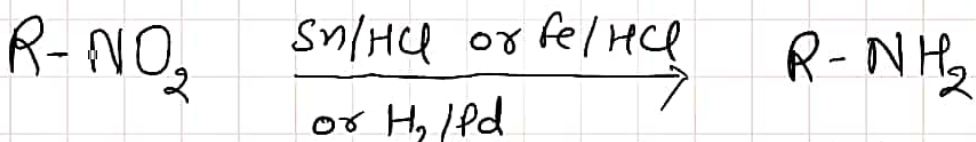
N,N -Dimethyl ethanamine



But-3-en-2-amine.

Preparation of Amines \rightarrow

- Reduction of Nitro Compounds -

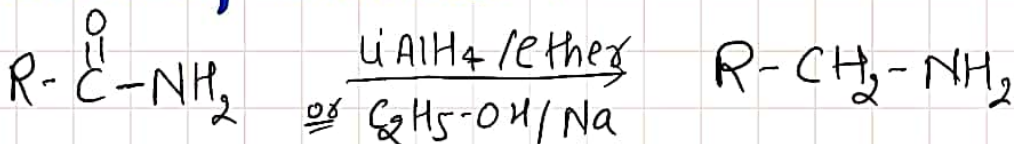


Nitrobenzene

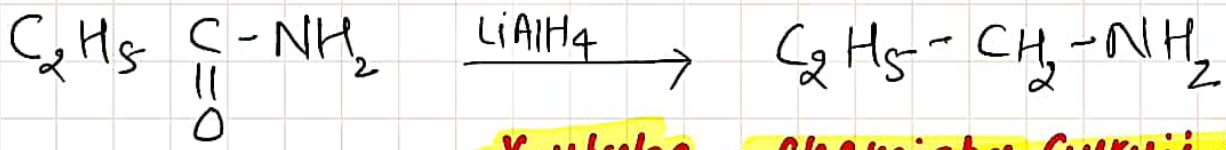


Aniline

- Reduction of Amides \rightarrow



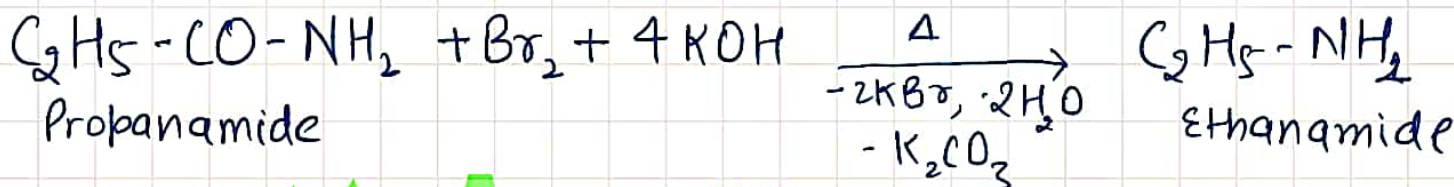
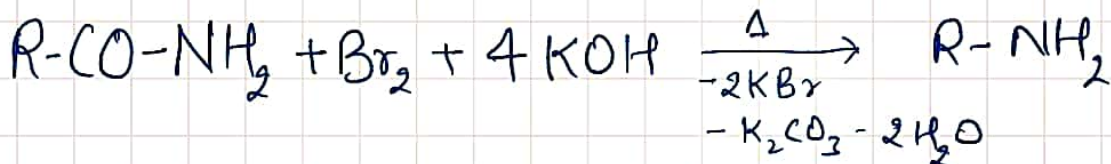
By Bharat Panchal
SST



Youtube - Chemistry Guruji 2.0

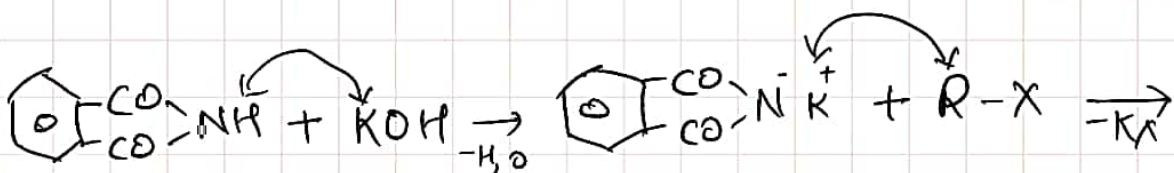
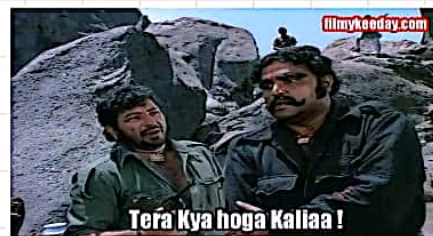
Hoffmann Bromamide Degradation

TRICK - Baby cut Base ፳፻፲፭ ይ!



Gabriel phthalimide synthesis :-

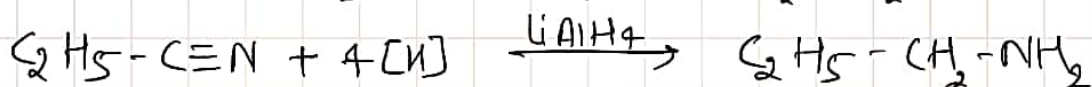
Trick - गतवर Basement में R-X को H₂O घिला।



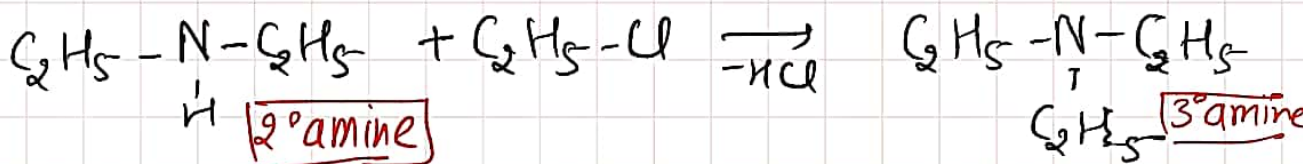
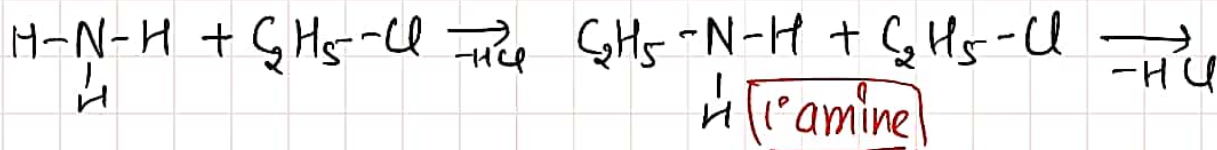
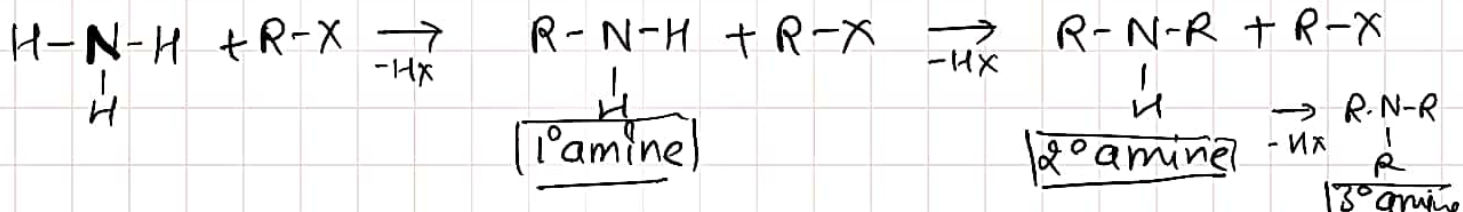
P.T.R

Aromatic 1° amines can't be prepared by this method because aryl halides do not undergo nucleophilic sub. with the anion formed by phthalimide.

• Reduction Of Nitriles:-



Hoffmann Ammonolysis of Alkyl Halide



Physical Properties of Amines:-

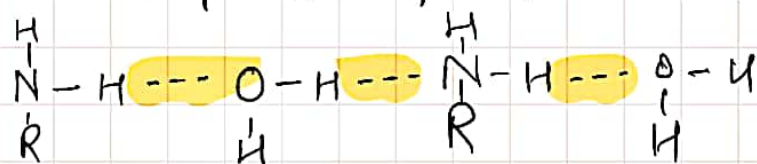
Physical State:-

Lower amines are gases and liquids but higher amines are solids. The lower aliphatic amines are gases with fishy odour.

Aryl amines are usually colourless but get coloured on storage due to atmospheric oxidation.

Solubility:-

Lower aliphatic amines are soluble in water because they can form H-Bond with water



Primary and Secondary amines are soluble in water due to H-bonding while 3° amines are insoluble in water.

- The Solubility decrease with increase in size of hydrophobic alkyl part.

P.T.R

The solubility of amines is less than that of alcohol of comparable molecular mass because alcohols are more polar than amines and form stronger H-Bond.

By - Bharat Panchal Sir

• Boiling Point \Rightarrow

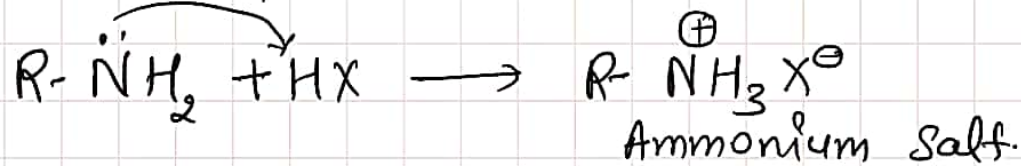
The order of b.pt isomeric amines
 1° amines $>$ 2° amines $>$ 3° amines

- ① 3° amines do not have intermolecular H-Bonding because no H- is attached to N-atom.
- ② 1° amines have maximum amount of H-Bonding because two H-atoms are attached to N-atom.

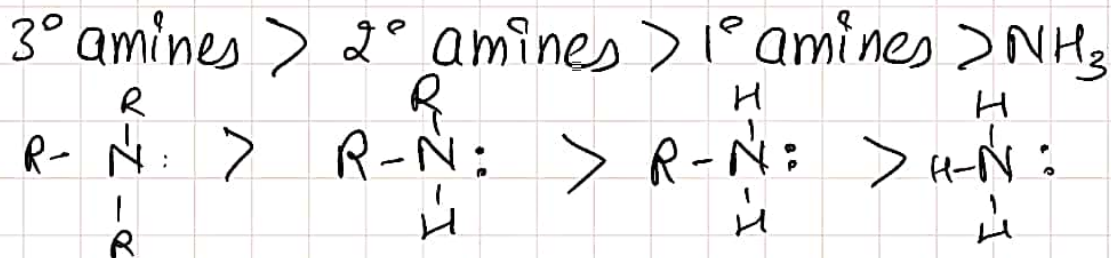
Basic Character of Amines

- Amines are basic in nature due to the presence of lone pair of e^- on nitrogen atom.
- Aliphatic amines are stronger bases than ammonia due to +I effect of Alkyl group.
- Aromatic amines are weaker bases than ammonia due to -I effect or Aryl group.
- Besides inductive effect, effects like steric effect, solvation effect, resonance effect also affect the basic strength of amines.

Amines are basic in nature and reacts with acids to form salts.

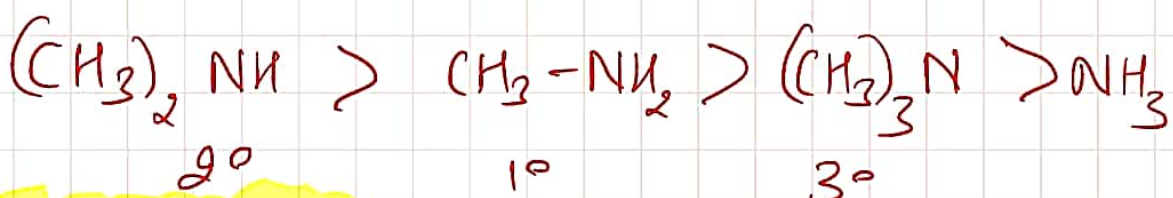
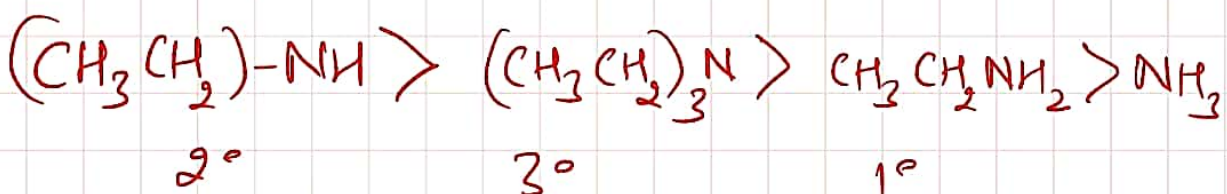


→ Order of basic character of amines in gaseous phase
(Acc. to +I effect)

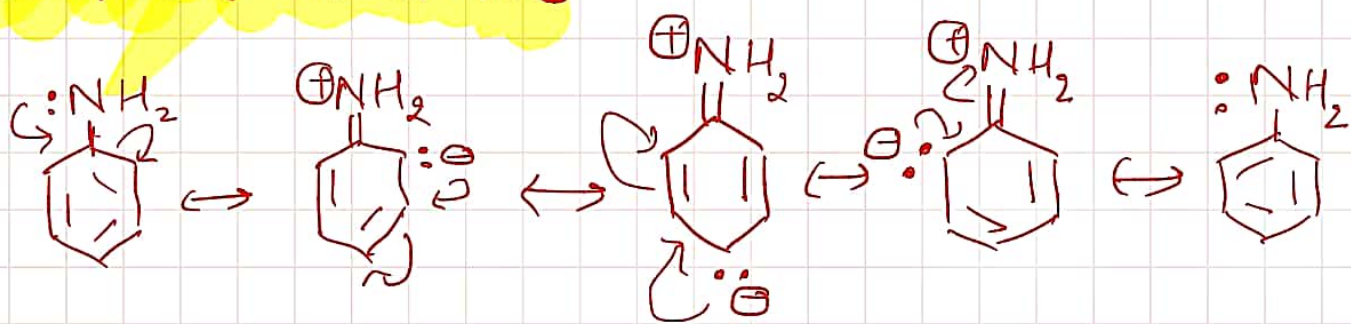


However in aqueous phase :-

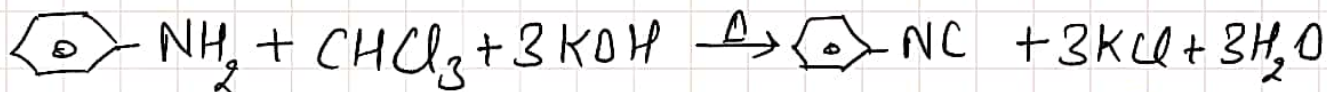
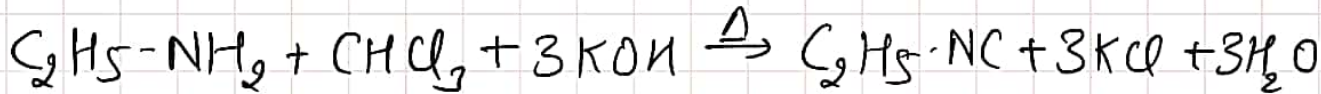
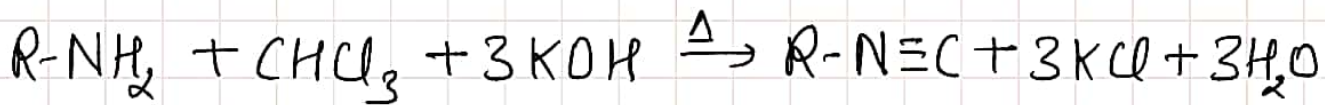
There is subtle interplay of Inductive effect, steric effect and solvation effect



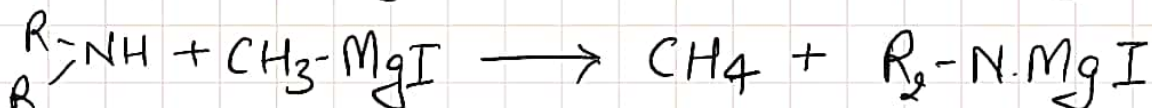
Resonance in Aniline



Carbylamine Reaction



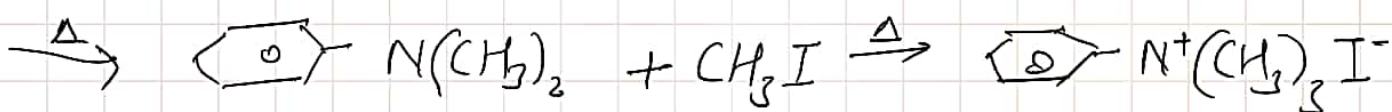
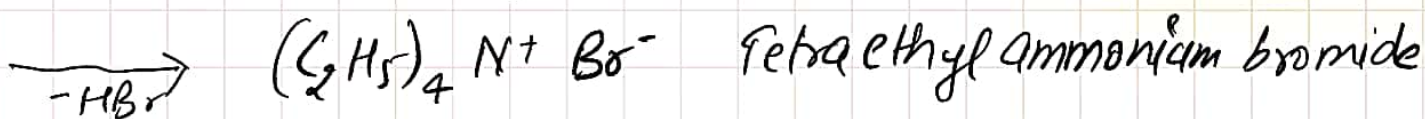
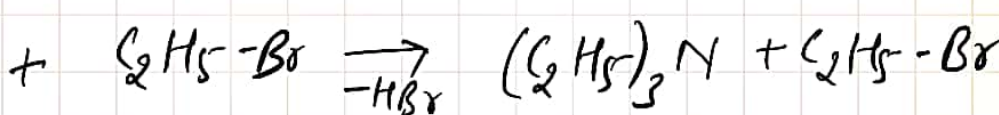
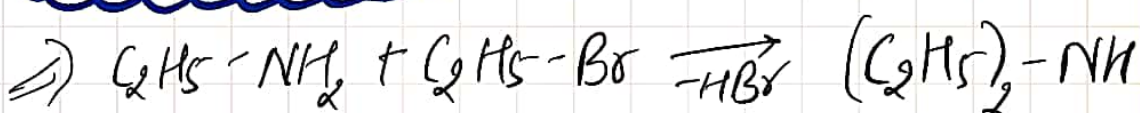
Reaction with Grignard Reagent By Bhagat Panchal



P.T.R

Tertiary amines do not react with Grignard reagents as they do not contain active H-atoms.

ALKYLATION

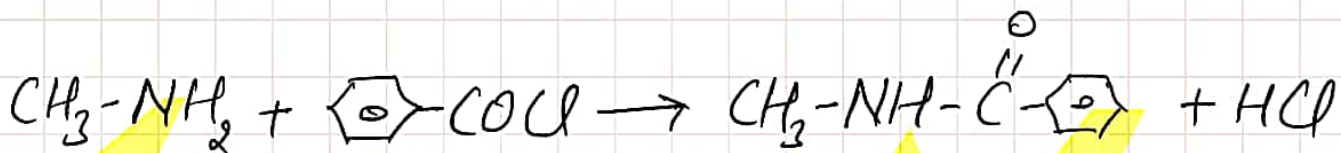
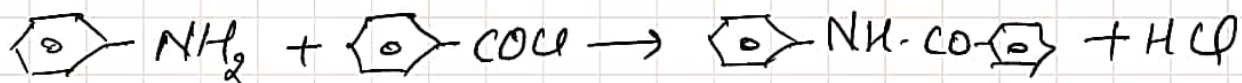


WHEN YOUR TEACHER SAYS



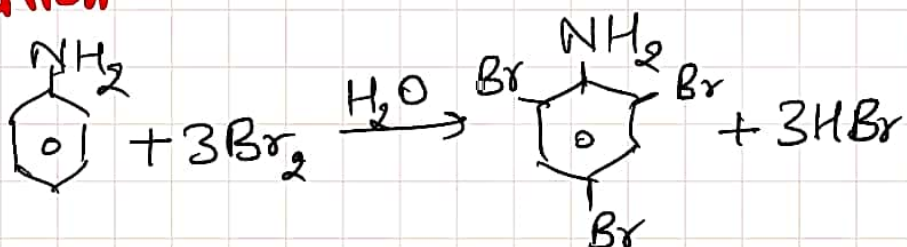
"HIGHLIGHT ONLY THE
IMPORTANT PARTS."

Acylation



Electrophilic Substitution Reaction \Rightarrow

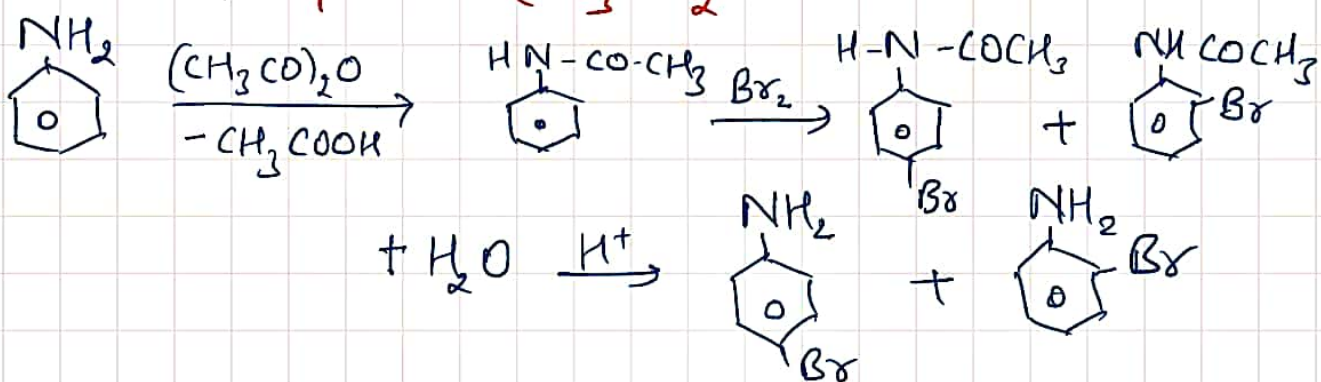
(i) Bromination -



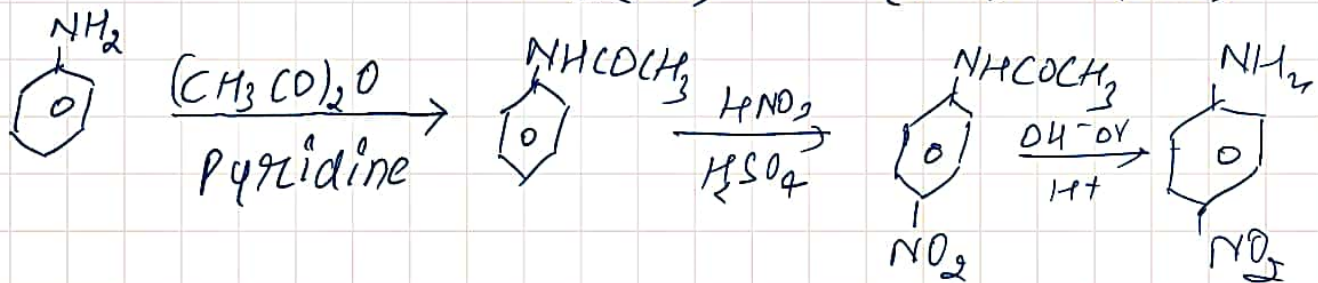
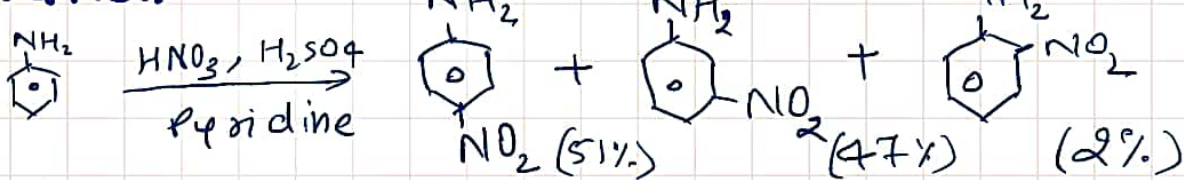
chemistry
Guruji
2.0

P.T.R

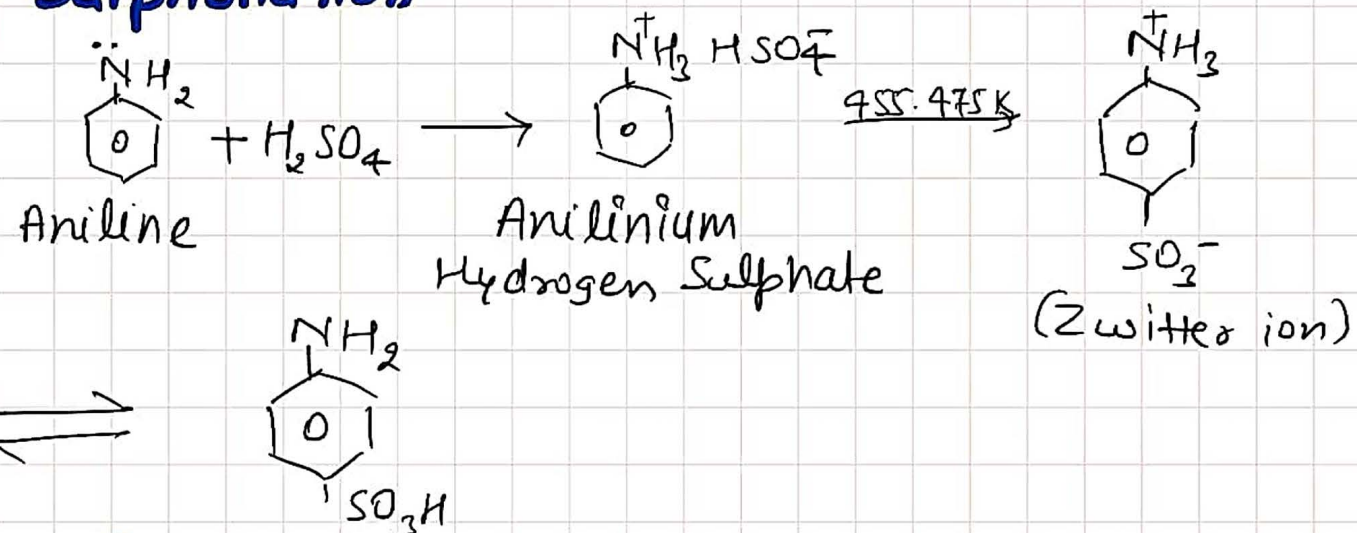
For monobromination, treat aniline with acetic anhydride $(\text{CH}_3\text{CO})_2\text{O}$



(ii) Nitration -

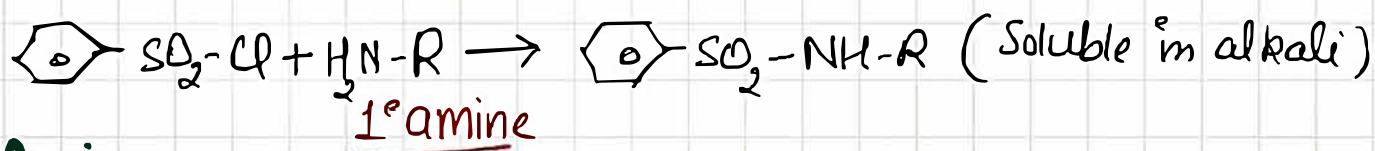


Sulphonation

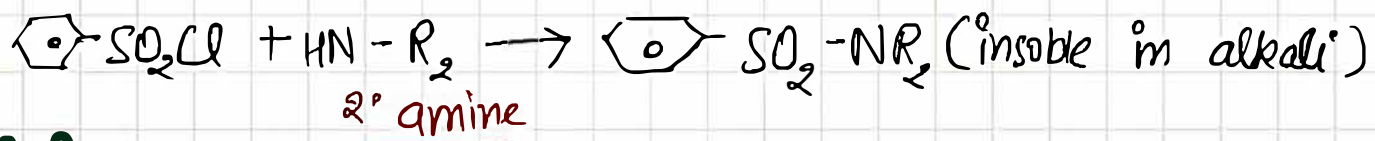


Hinsberg Test (benzene sulphonyl chloride)

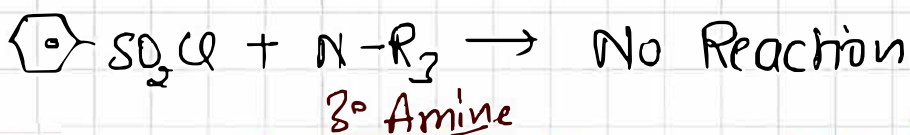
1° Amines



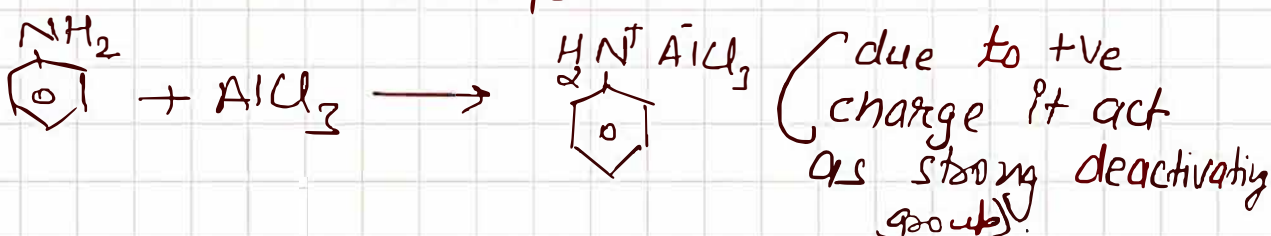
2° Amines



3° Amines



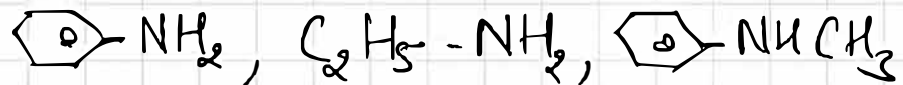
P.T.R Friedel craft Rxn are not possible for NH_2 because it form salt with anhy. AlCl_3 that is used as catalyst in the reaction.





Basic Character $\propto K_b$
Basic Character $\propto \frac{1}{P_{K_b}}$

- Arrange the following in the increasing order of P_{K_b} values:
(C.B.S.E - 2018)



Solution :- $\text{C}_2\text{H}_5\text{-NH}_2 < \text{C}_6\text{H}_5\text{NH-CH}_3 < \text{C}_6\text{H}_5\text{NH}_2$

(*) Give reasons : $(\text{CH}_3)_2\text{NH}$ is more basic than $(\text{CH}_3)_3\text{N}$ in an aqueous solution (C.B.S.E - 2018)

Ans: $(\text{CH}_3)_2\text{NH}$ is more basic than $(\text{CH}_3)_3\text{N}$ in an aqueous solution due to less steric hindrance.

≡ -