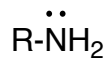


Amines (McM chapt 24)

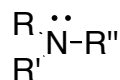
(R: alkyl, aryl)



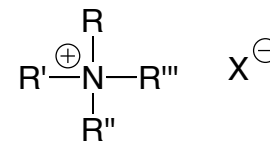
Primary amine



Secondary amine

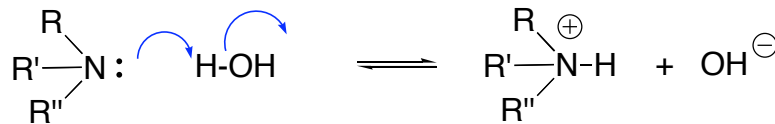


Tertiary amine



Quaternary ammonium salts

Basic compounds



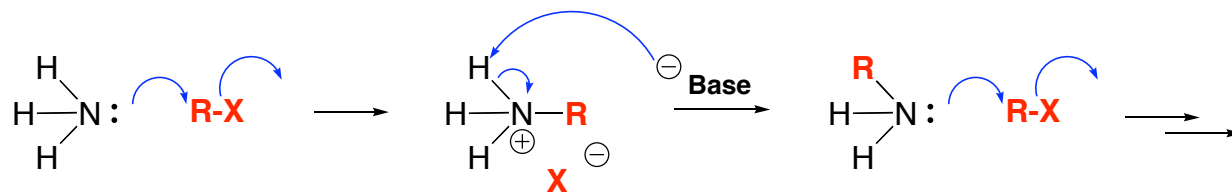
(R: H, alkyl, aryl)

pKa Alkylamines: ca 9-11
Arylamines: ca 4-5
(anilines)

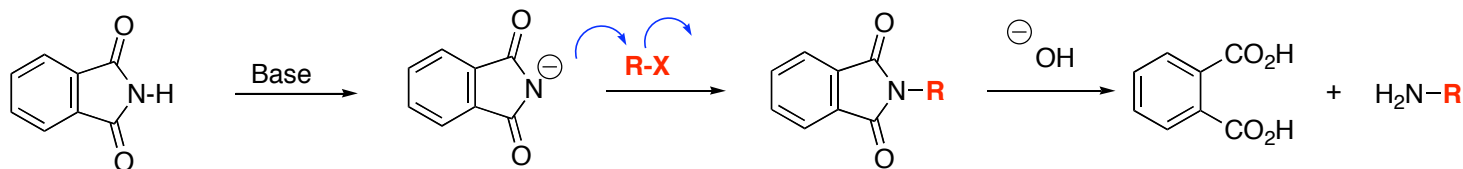
Synthesis

Known react. from KJM10xx

Alkylation (ammonia or amine, phthalimide)

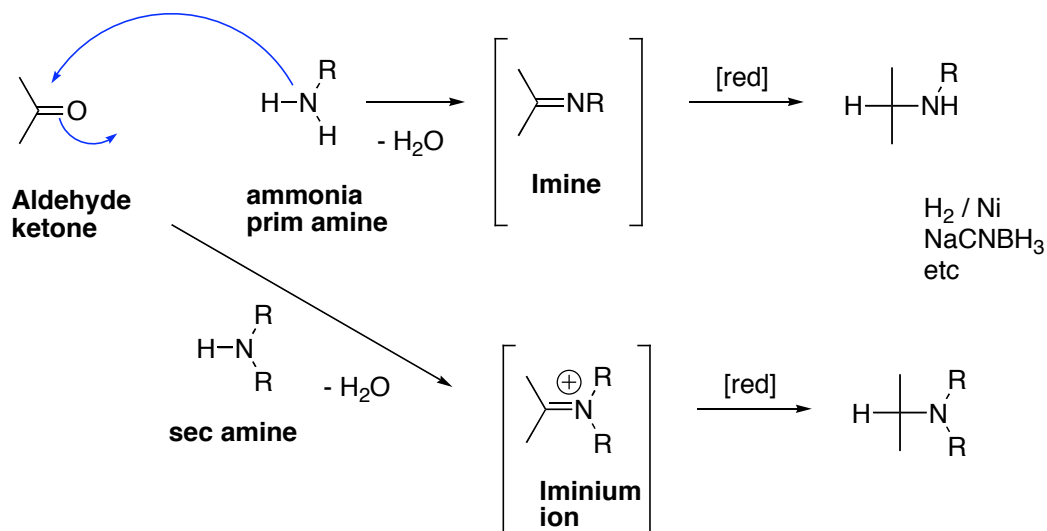


(R: H, alkyl, aryl)



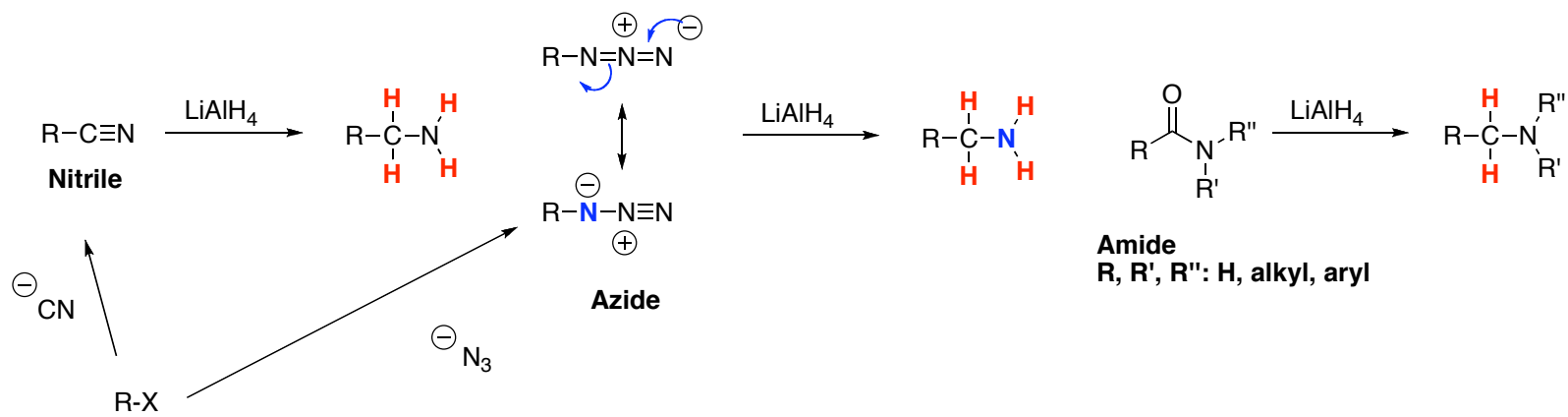
Acidity
c.f. 1,3-dicarbonyls

Reductive amination

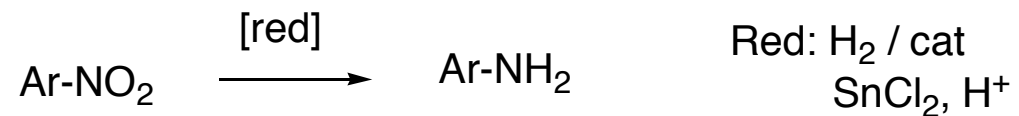


Reductions

- Nitriles
- Azides
- Amides

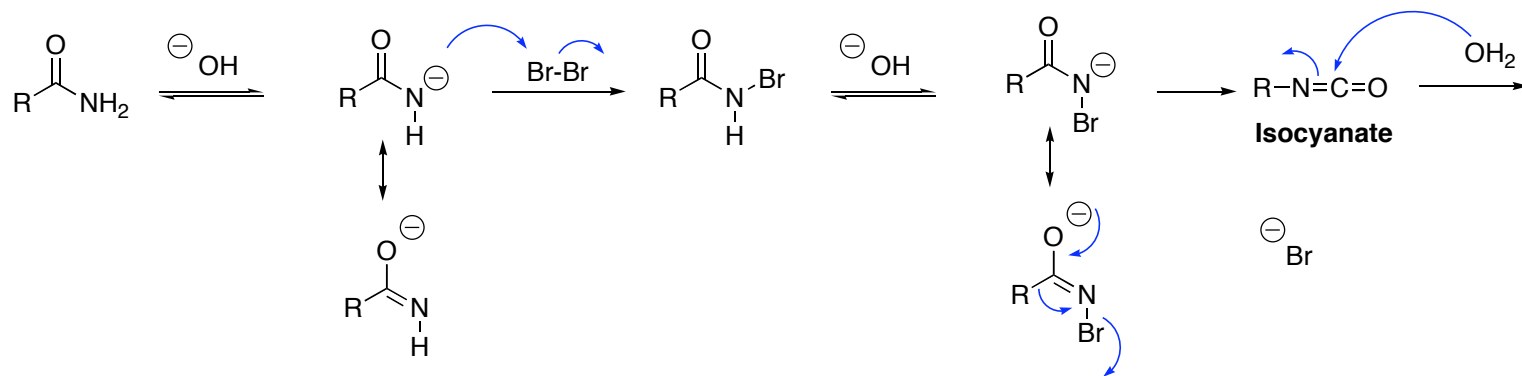
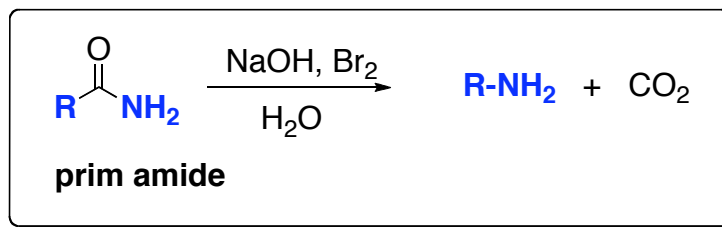


•Aromatic nitro compounds

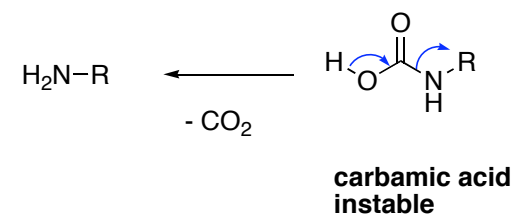
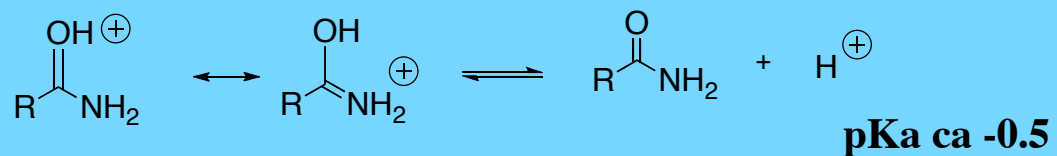
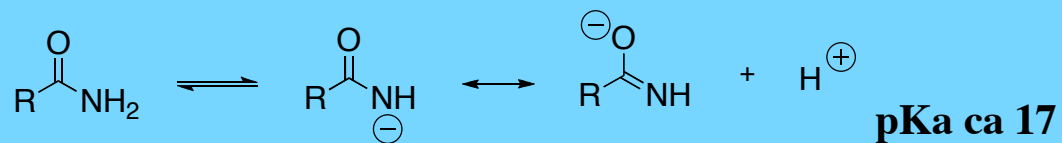


Synthesis: "New" reactions

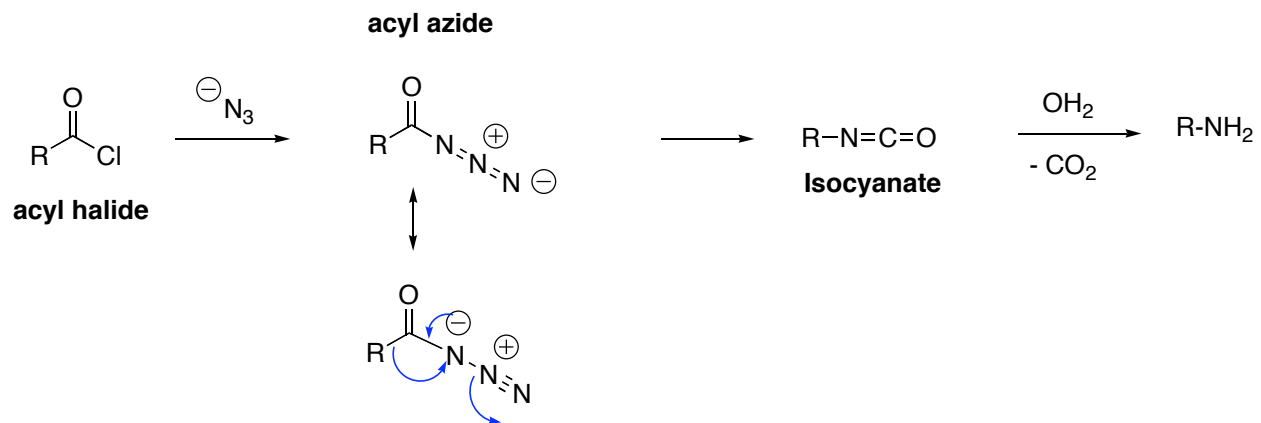
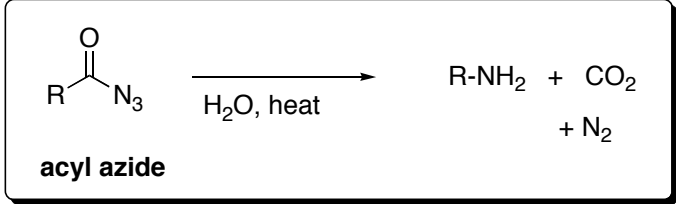
Hofmann rearrangement



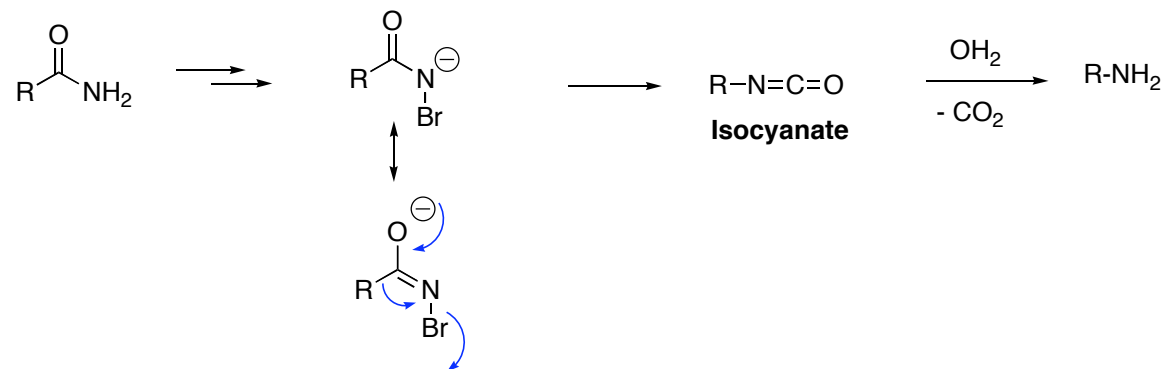
Acid / base properties amides



Curtius rearrangement



Mechanistic. related to Hofmann rearrang.



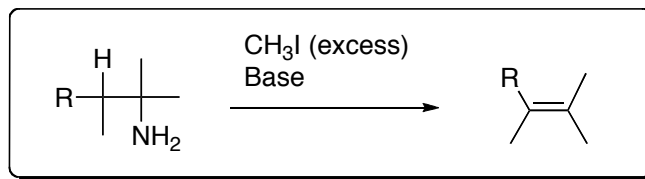
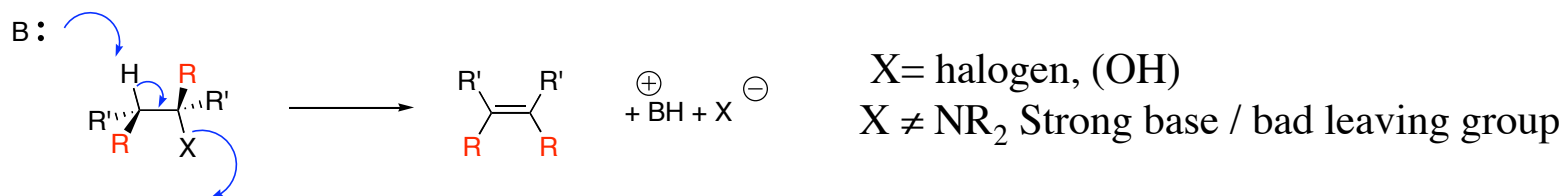
Reactions of amines (Alkylamines)

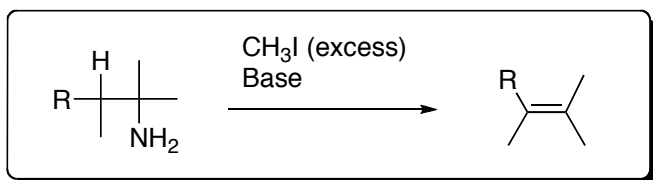
Alkylamines:

- Alkylation
- Acylation / synth of amide
- **Hofmann elimination** (\neq Hofmann rearrangement)

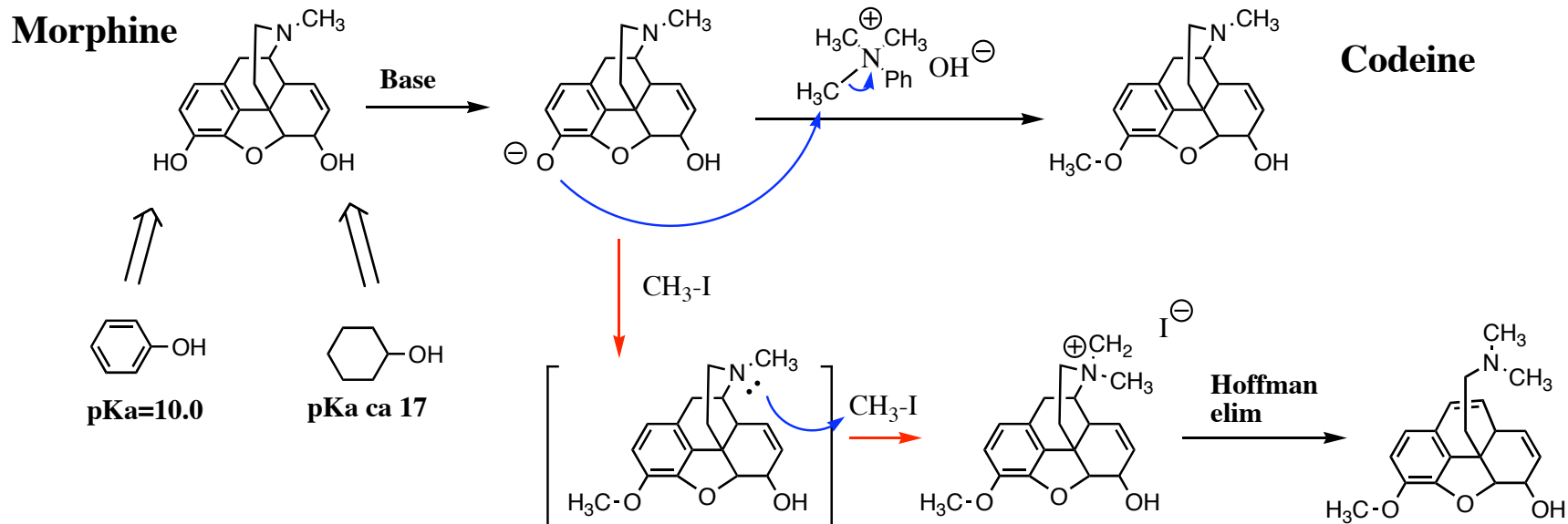
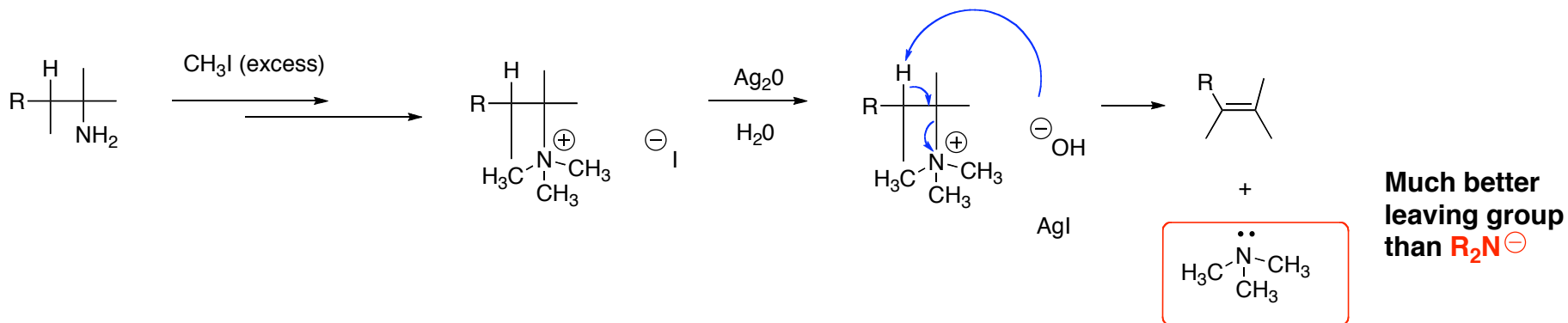
E2 elimination to form alkene

E2: mechanism

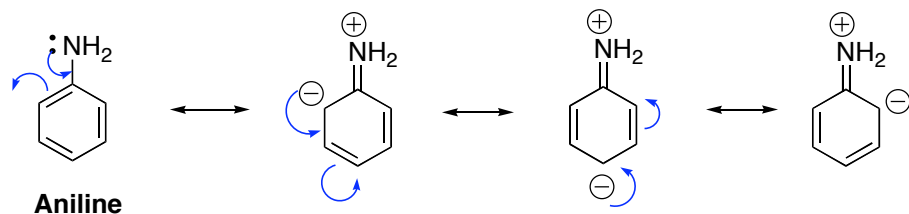




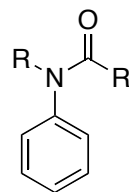
Removal of less sterically hindered H
Not necessarily most stable alkene formed



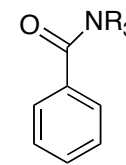
Reactions of Arylamines (aniline derivatives)



- Weak base (pKa ca 4.6)
- Highly Activated for E-fil Ar Subst (o/p)
- Protect. as amide: Less activated, still o/p

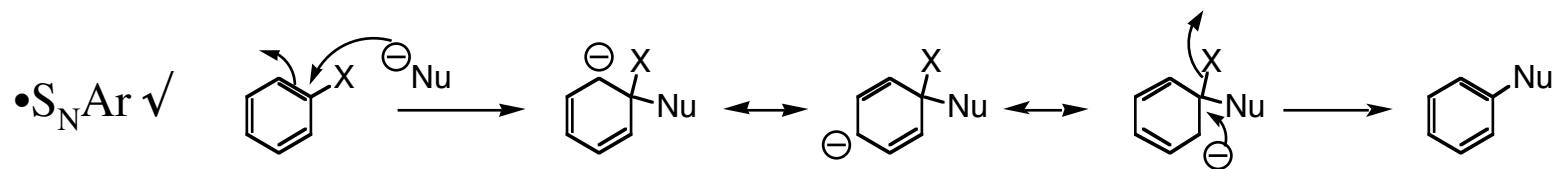


o/p directing



m-directing

Nucleophilic Aromatic Substitution - Mechanisms



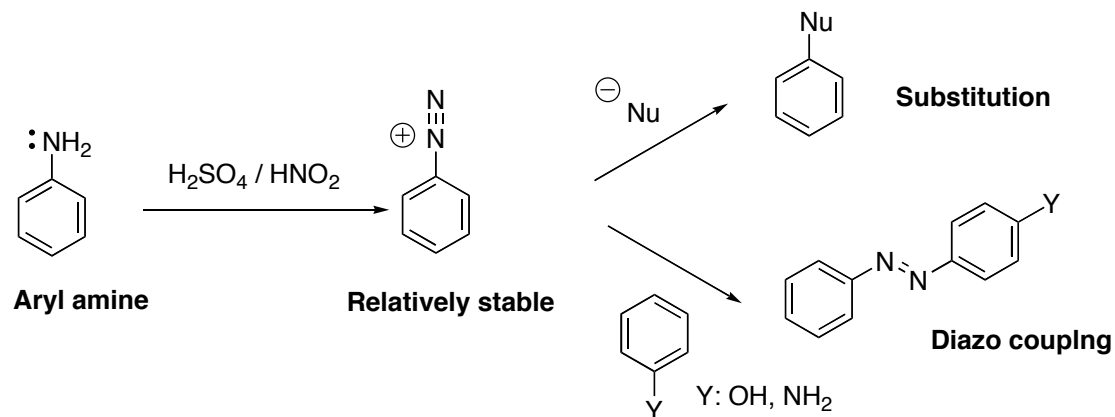
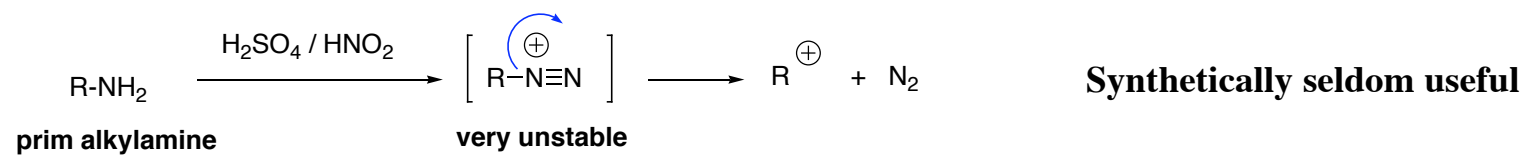
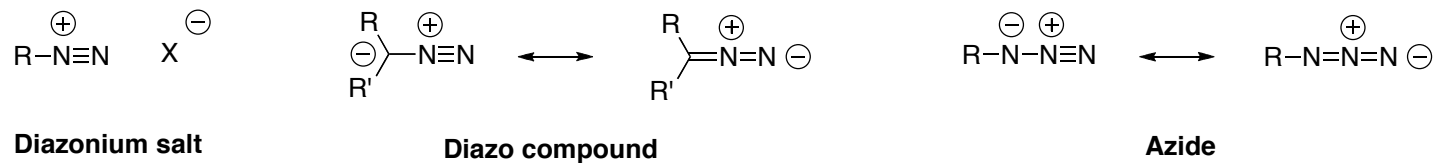
• S_N1

• Benzyne ✓

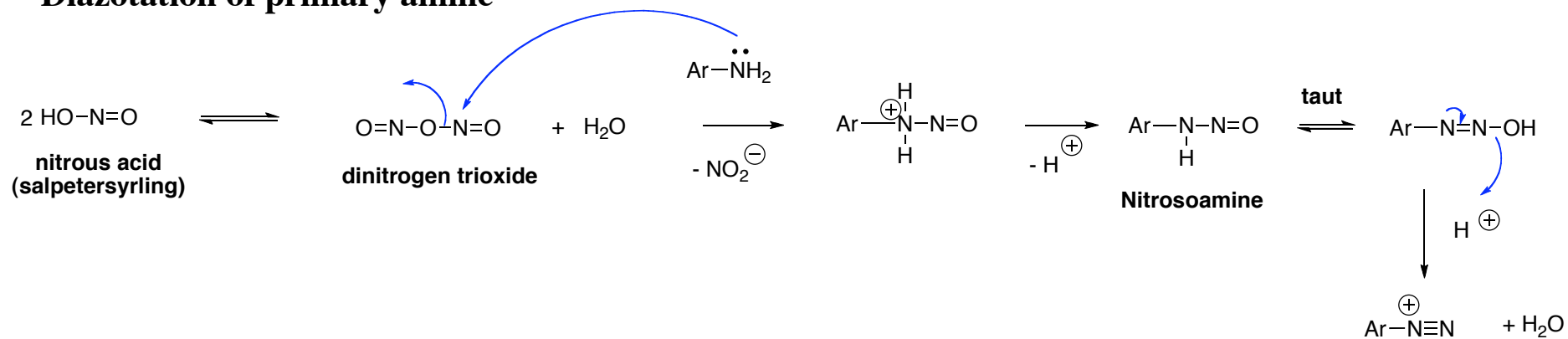
• **SRN1: Involves radicals**

• (VNS: Vicarious Nucl. Subst.)

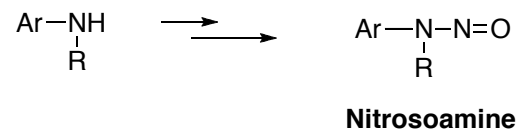
Formation of Diazonium Salts and the Sandmeyer Reaction



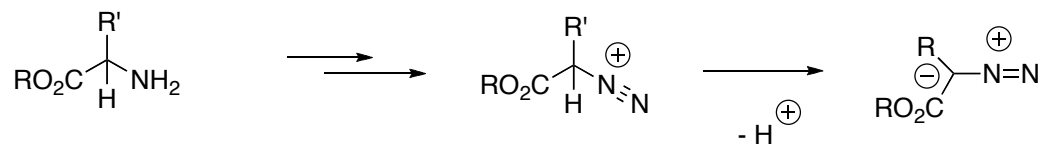
Diazotation of primary amine



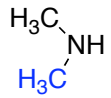
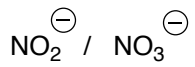
Sec amine gives nitroso compound



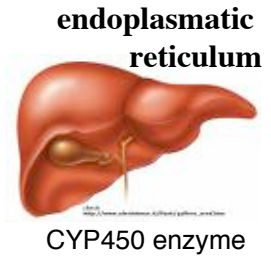
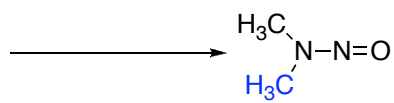
Amines with acidic α -H may give diazo compounds



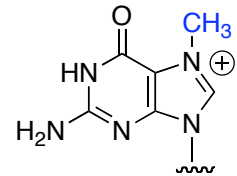
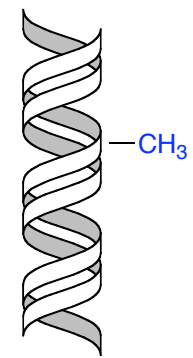
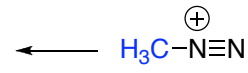
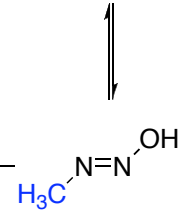
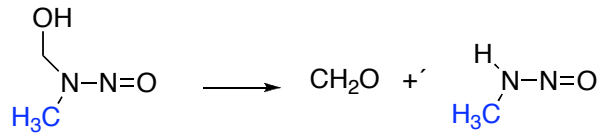
Toxicity nitroso compounds (not in McM) - Alkylation of biomolecules



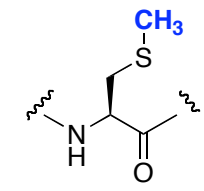
sec alkylamine



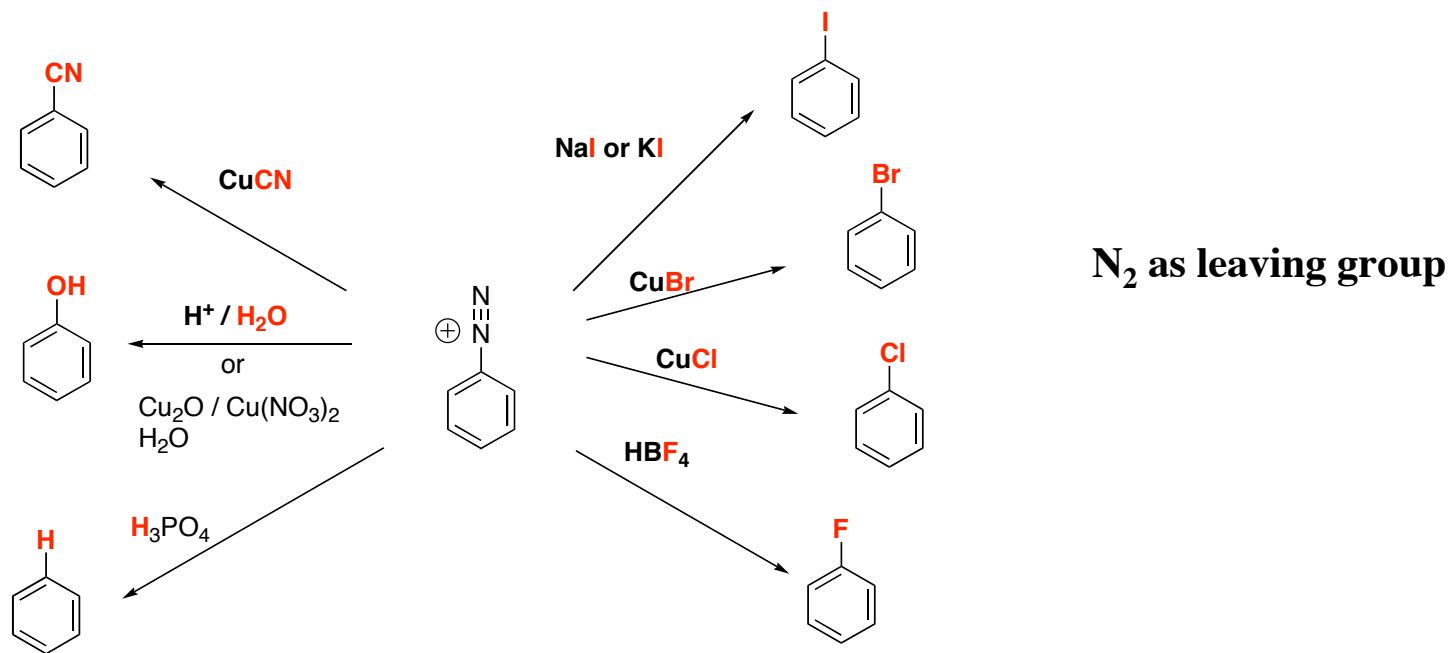
CYP450 enzyme



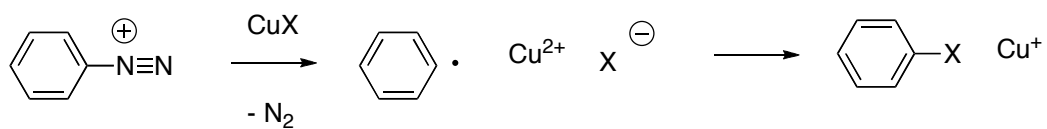
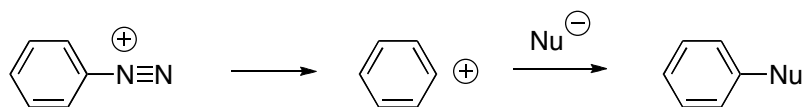
N-7 alkylation of guanine in DNA



S-methylated cysteine in a peptide/protein

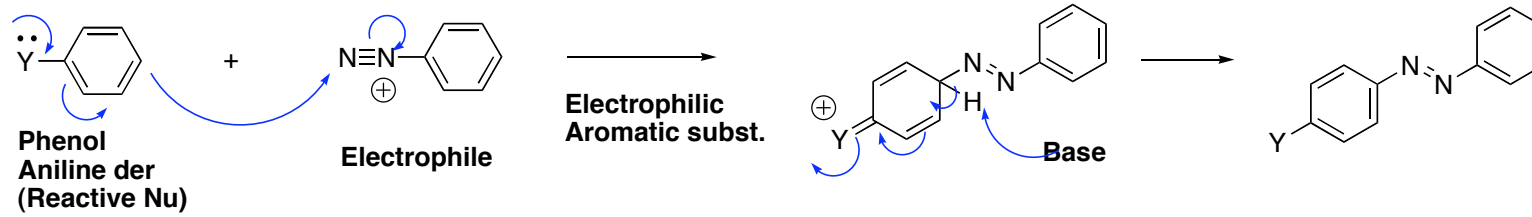


SN1 like mechanism or radical mechanism

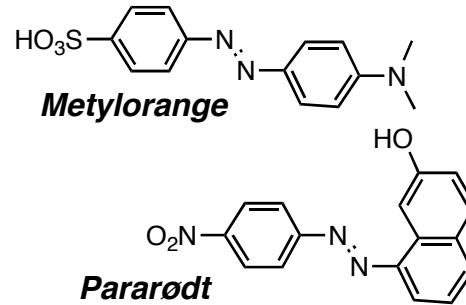


**Cu-salt mediated react.
Sandmeyer react.
(radical mech.)**

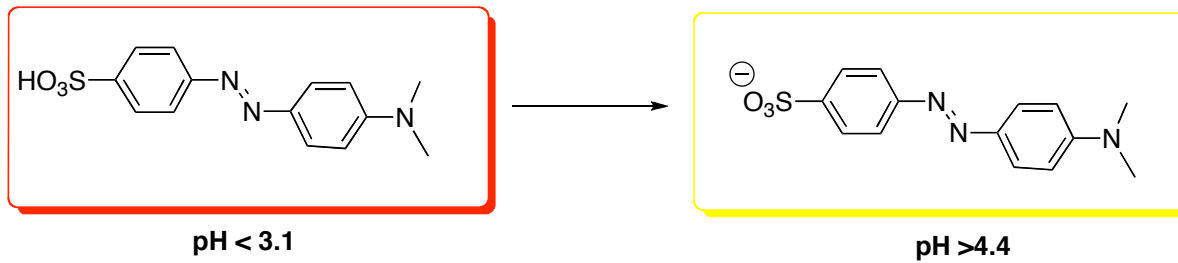
Diazo coupling



Azo dyes
Bayer etc
Late 1800-century, ex.



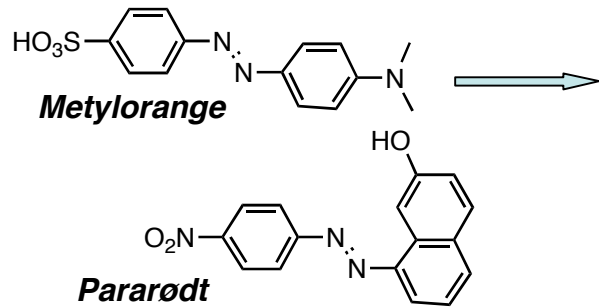
Metylorange



Azo dyes

Bayer etc

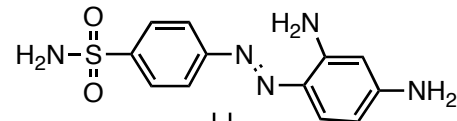
Late 1800-century, ex.



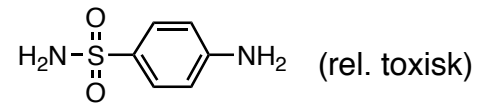
Screening of dyes as antibacterials



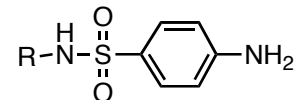
1932: **Prontocil** active against Streptococcus infection
no activity on bacterial cultures



1935: Prontocil metabolized (azoreductase) to **Sulfanilamid** *in vivo*



Modern sulfa drugs



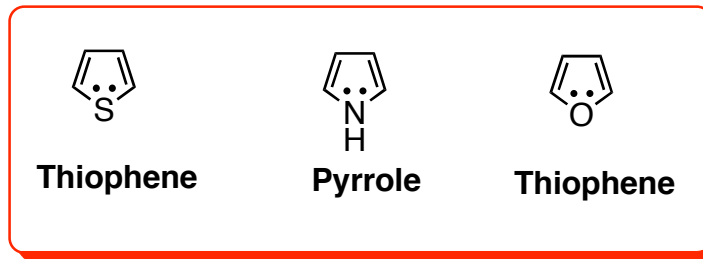
R: Aryl or hetroaryl

Antibacterial sulfonamides

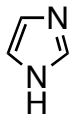
Heterocycles (McM chapt 24)

- Monocyclic or fused rings
- Cont. one or more ring atom \neq C (normally O; N; S)
- Aromatic, partly saturated or saturated ring(s)

5-Membered rings (Heteroatom N, O, S)



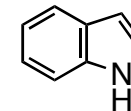
Other examples



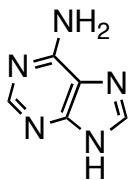
Imidazole



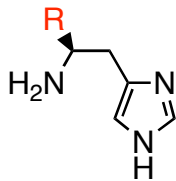
Thiazole



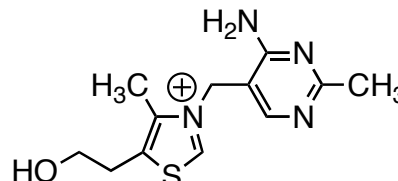
Indole



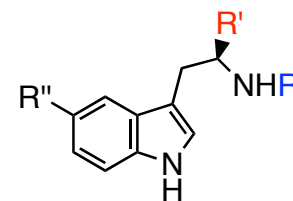
Adenine
(purine der.)



R=CO₂H: Histidine
R=H: Histamine



Thiamin
(Vit B1)



R=H
R'=H
R''=OH } Serotonin

R=H
R'=CO₂H
R''=H } Tryptophane

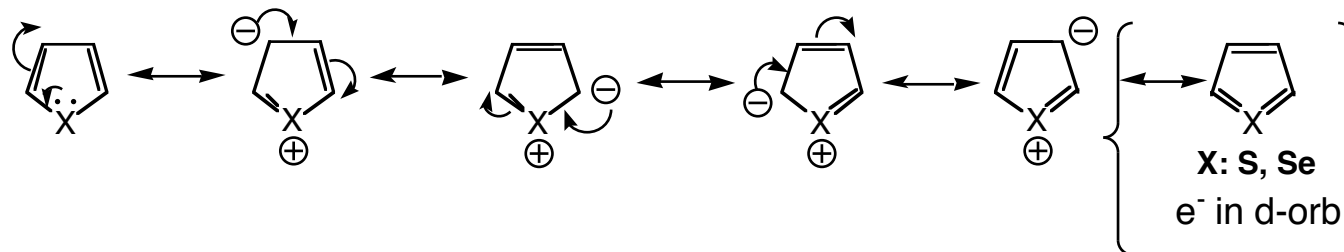
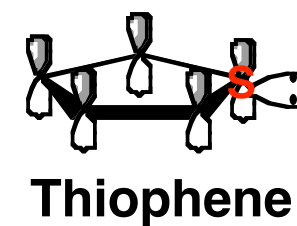
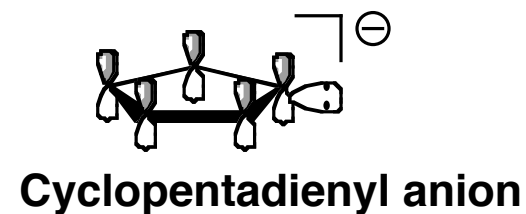
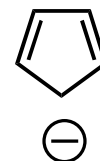
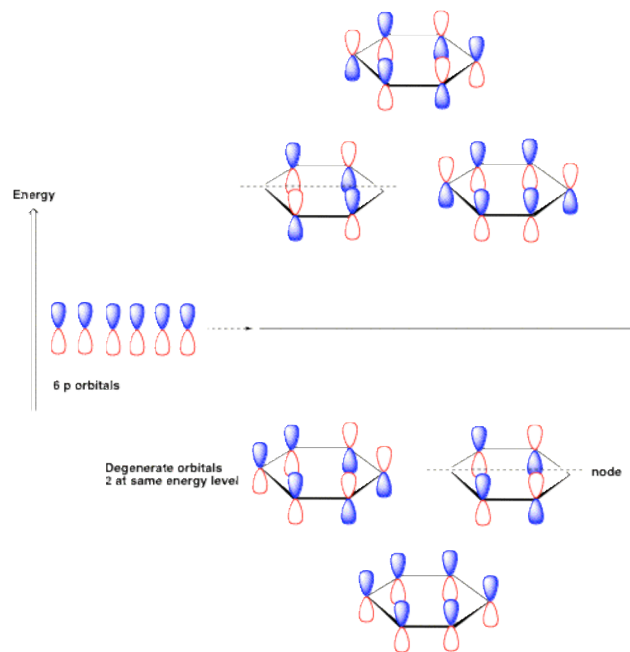
R=Ac
R'=H
R''=OCH₃ } Melatonin

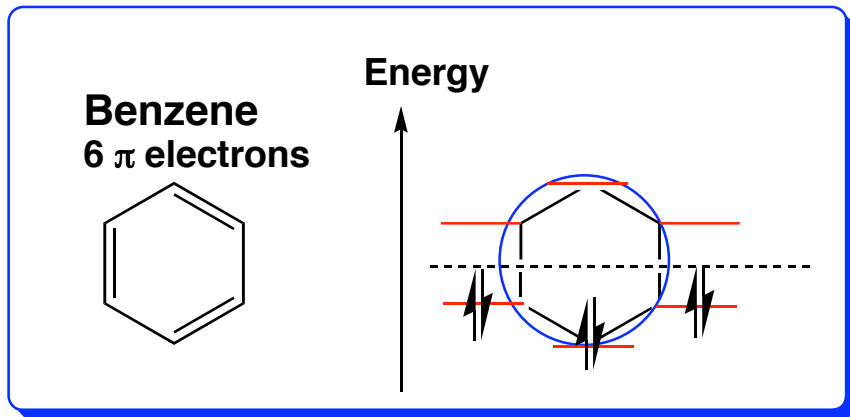
Criteria for Aromaticity (Hückel)

(Monocyclic) ring

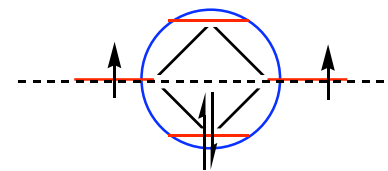
Planar

No of π -electrons in conjugation $4n+2$ ($n: 0, 1, 2, \dots$)



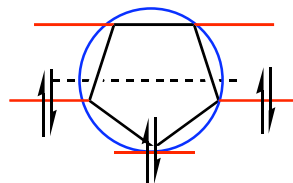
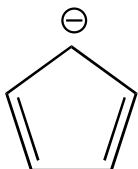


Cyclobutadiene
4 π electrons



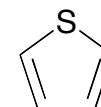
Diradical

Cyclopentadienyl anion
6 π -electrons

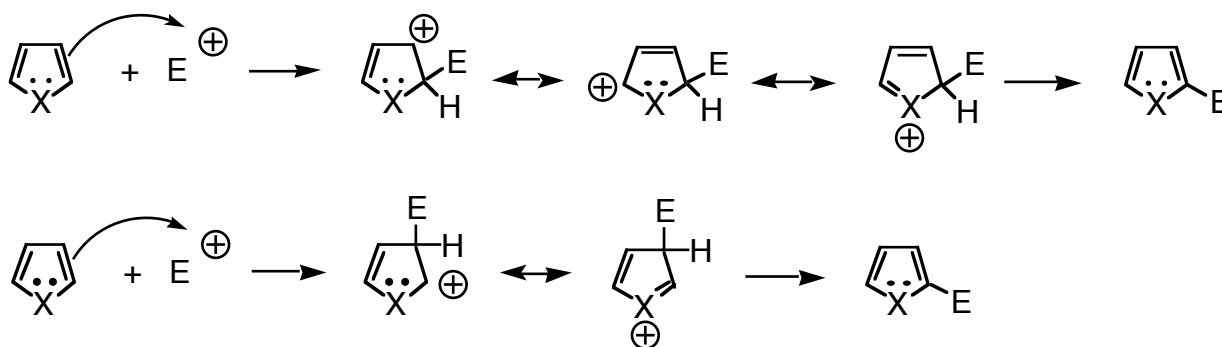
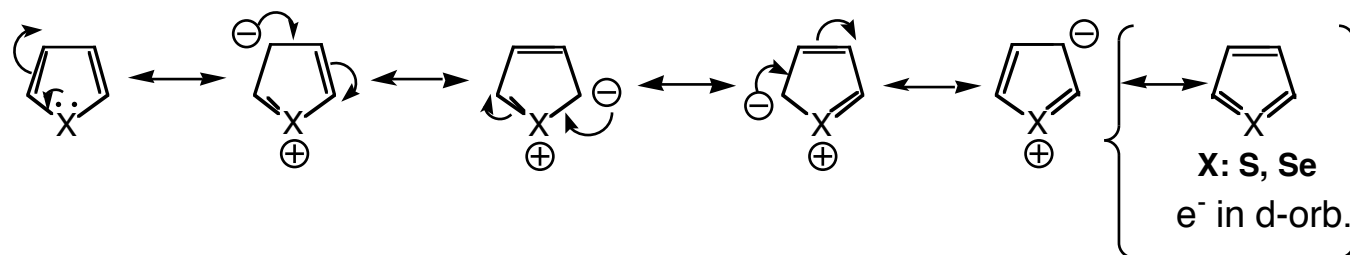


**All π electrons in
the bonding MO**

Thiophene
6 π -electrons



5-membered rings - electron rich on C - reactive i E-fil. Ar subst.

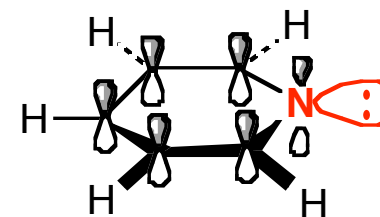
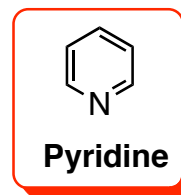


React. in α -position generally preferred

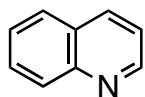
Selectivity not always good

React.: Pyrrole > thiophene > furan

6-Membered rings (Heteroatom N)



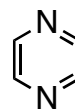
Other examples



Quinoline



Pyrimidine

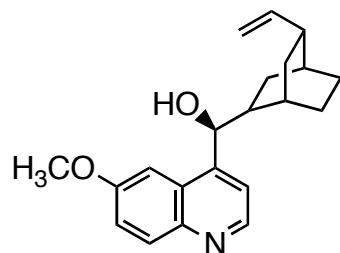


Pyrazine

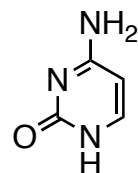


Pyridazine

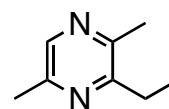
Rare in nature



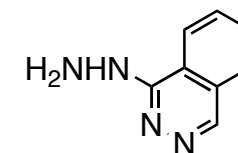
Quinine



Cytosine

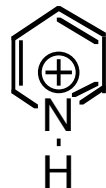
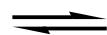
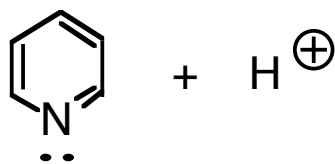


Ant pheremone

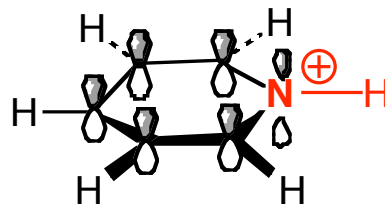
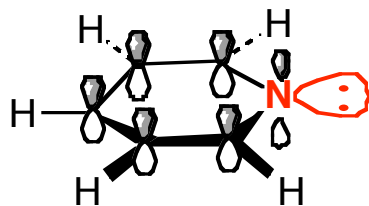


Hydralazine
Antihypertensive drug

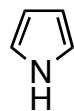
Pyridine as a base



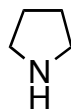
pKa: 5.2



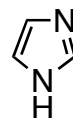
sp₂ N less basic than sp₃



pKa 0.4



pKa 11.3



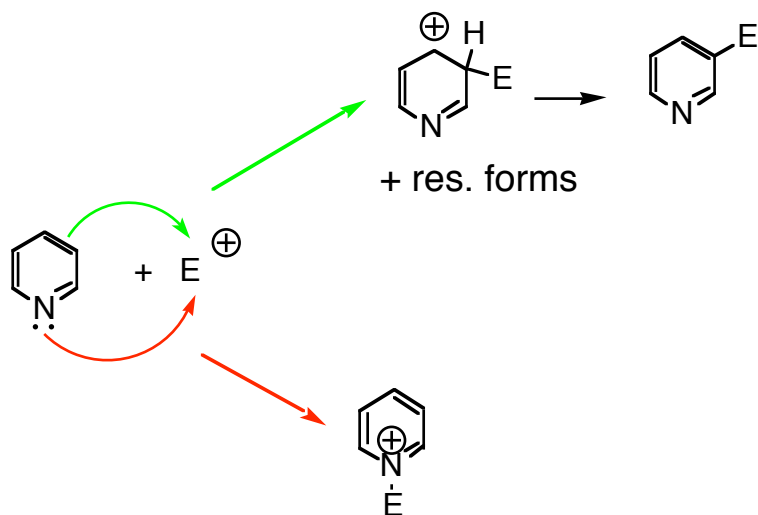
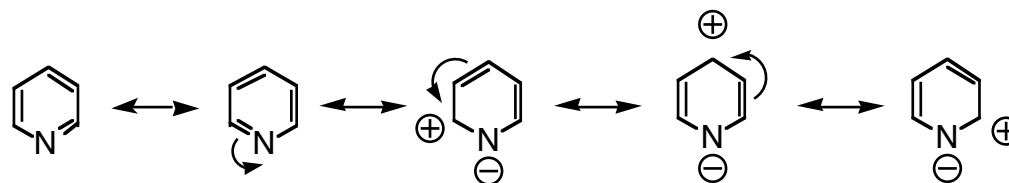
**pKa 7.1
(≈amidine)**



pKa 2.5

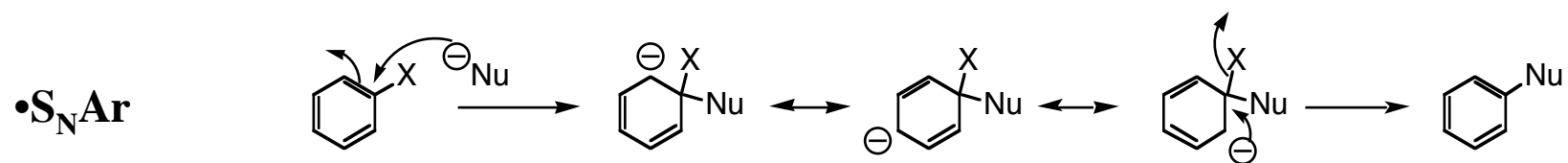
Electrophilic Reaction on Carbon: E-phil. Ar. Subs.

6-membered rings - electron deficient on C - ↓ reactivity



- Both C and N may react
- 3/5 pos. most reactive C
- Diazines less reactive
- Sulfonation, Nitration, halogenation
- Not FC react.

Nucleophilic Aromatic Substitution



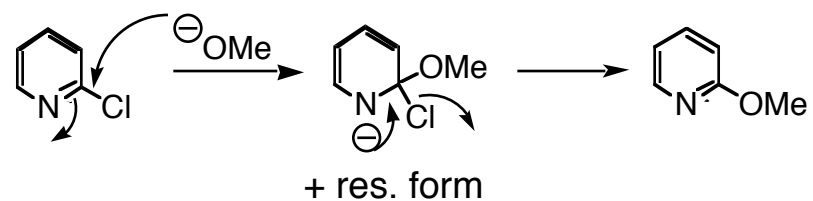
• **S_N1: Via aryl cation**

• **Benzyne**

• **SRN1: Involves radicals**

• **VNS: Vicarious nucl. Subst.**

6-membered rings - electron deficient - reactive in Nu-fil. Ar subst.



2 / 4 Pos. reactive; electron def. C, neg. charge partly on N in intermed
3 / 5-Pos. much less reactive (benzenoid pos.)