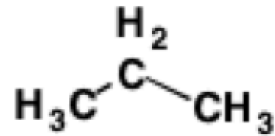


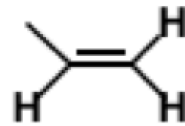


# Chemistry and Introduction to Biochemistry

## Functional Groups



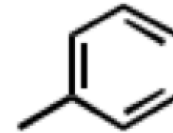
alkane



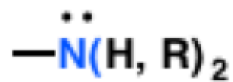
alkene



alkyne



benzene ring  
(phenyl)



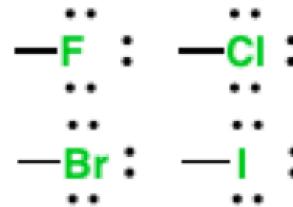
amine



alcohol



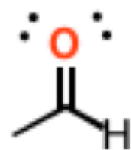
ether



alkyl halide



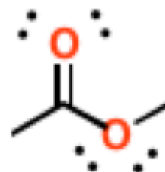
thiol



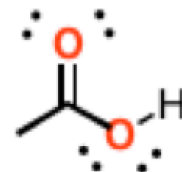
aldehyde



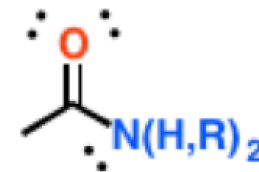
ketone



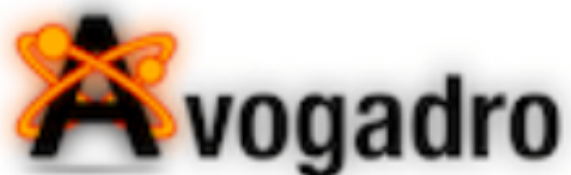
ester



carboxylic  
acid

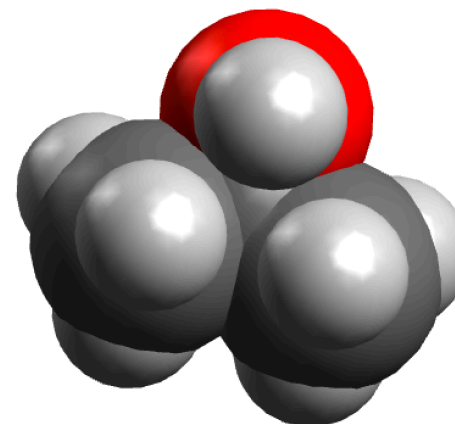


amide



Avogadro is an **open source** **advanced molecule editor and visualizer** designed for cross-platform use in *computational chemistry, molecular modeling, bioinformatics, materials science*, and related areas. It offers flexible high quality rendering and a powerful plugin architecture.

download it <http://avogadro.cc/>



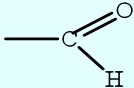
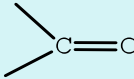
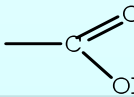


# Chemistry and Introduction to Biochemistry

## Functional Groups

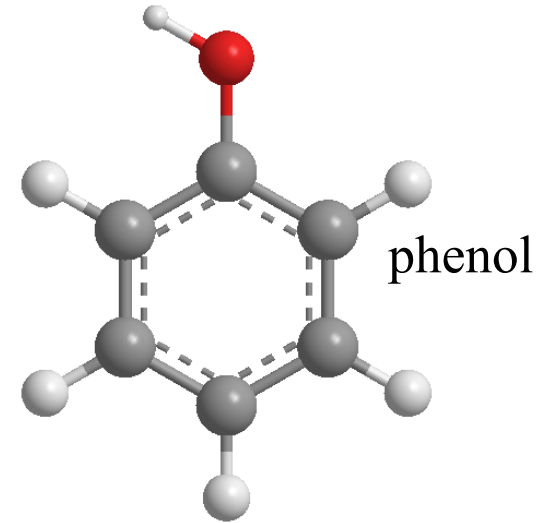
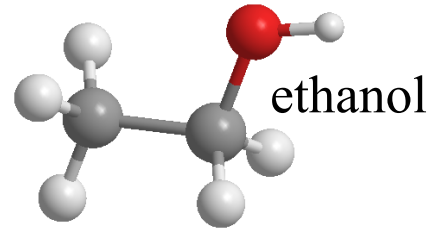
A functional group is defined as a **part of the structure** of a molecule characterized by **specific elements**, which gives the compound a **typical reactivity** similar to that of other compounds containing the same group.

In other words, the functional group constitutes the **center of the chemical reactivity of the molecule**.

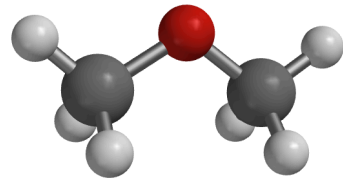
functional group		class of compounds	notes
$C=C$	carbon-carbon double bond	alkenes	
$C\equiv C$	carbon-carbon triple bond	alkynes	
$\text{---}X$	any halogen	<ul style="list-style-type: none"> <li>haloalkanes</li> <li>acyl halides</li> </ul>	<ul style="list-style-type: none"> <li>when it substitutes an H in an alkane</li> <li>when it substitutes an OH in a carboxylic acid</li> </ul>
$\text{---}OH$	hydroxy	<ul style="list-style-type: none"> <li>alcohols</li> <li>enols</li> <li>phenols</li> </ul>	<ul style="list-style-type: none"> <li>when bound to <math>sp^3</math> carbon</li> <li>when bound to <math>sp^2</math> carbon</li> <li>when bound to an aromatic ring</li> </ul>
$\text{---}SH$	sufyhydryl	thiols	when bound to $sp^3$ carbon
	aldehyde	aldehydes	
	carbonyl	ketones	
	carboxylic	carboxylic acids	
$\text{---}NH_2$	amine	<ul style="list-style-type: none"> <li>amines</li> <li>aromatic amines</li> <li>ammides</li> </ul>	<ul style="list-style-type: none"> <li>when it substitutes an H in an alkane</li> <li>when bound to an aromatic ring</li> <li>when it substitutes an OH in a carboxylic acid</li> </ul>

# Functional groups of oxygen

## 1) alcohols & phenols

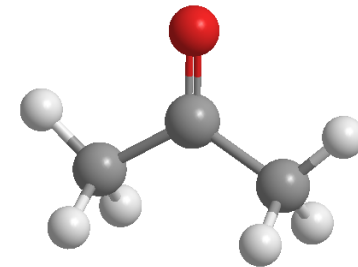
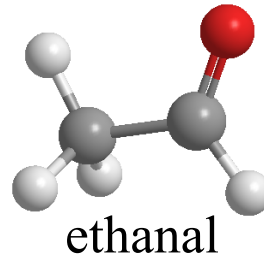


## 2) ethers



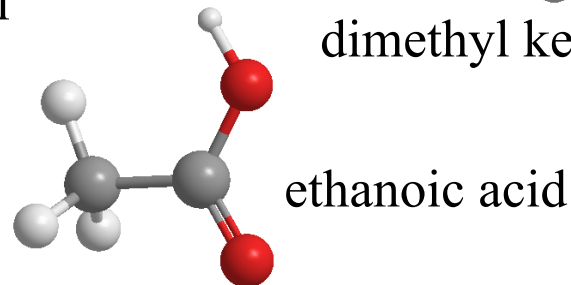
dimethyl ether

## 3) aldehydes & ketones

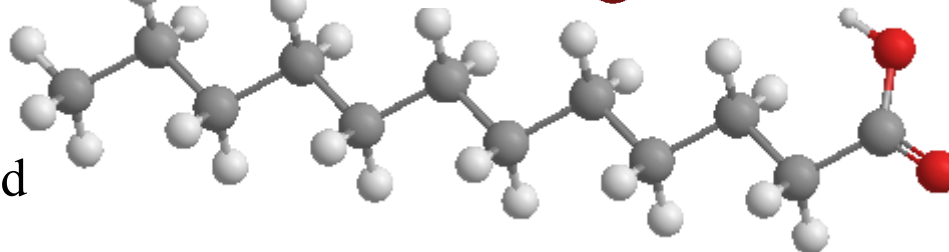


dimethyl ketone

## 4) carboxylic acids



dodecanoic acid



# Alcohols and phenols

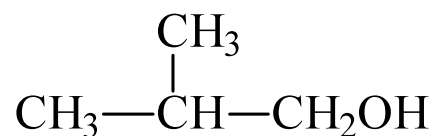
The functional groups of oxygen are numerous and important.

The simplest is the hydroxyl group:

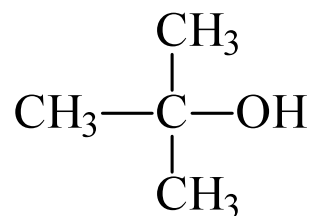
–OH present in alcohols and phenols

**Nomenclature:** the name of an alcohol is given by the name of the corresponding hydrocarbon, to which the ending **-ol** is added.

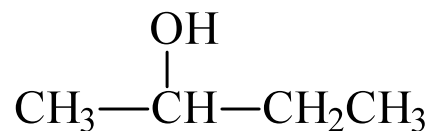
The longest hydrocarbon chain is identified, and is numbered by giving the lowest number to the C to which the OH is bound.



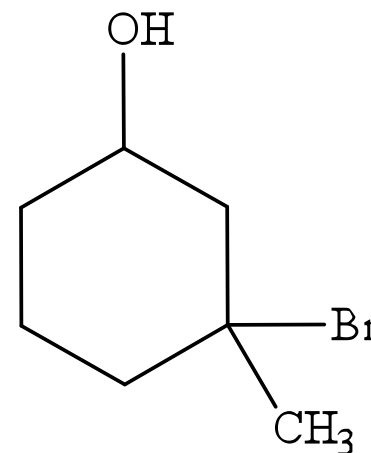
2-methyl-1-propanol



2-methyl-2-propanol



2-butanol

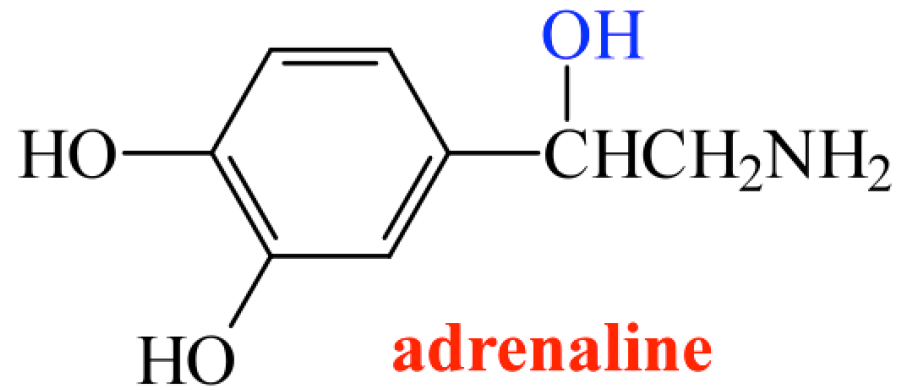


3-bromo-3-methylcyclohexanol

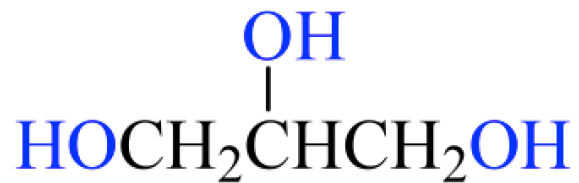
# Some alcohols



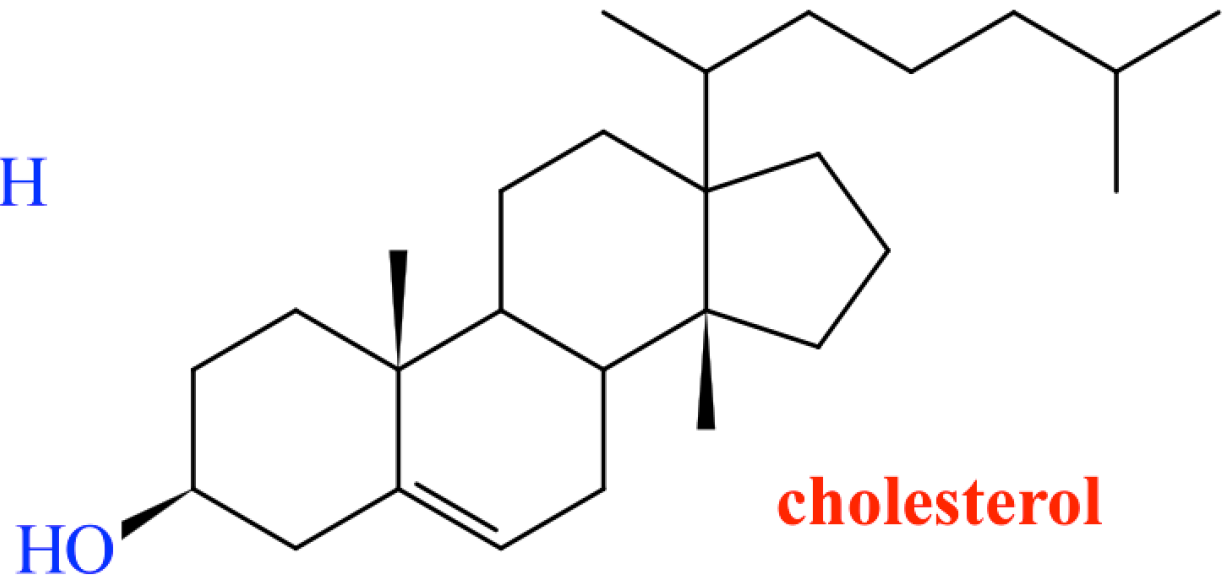
**ethanol**



**adrenaline**

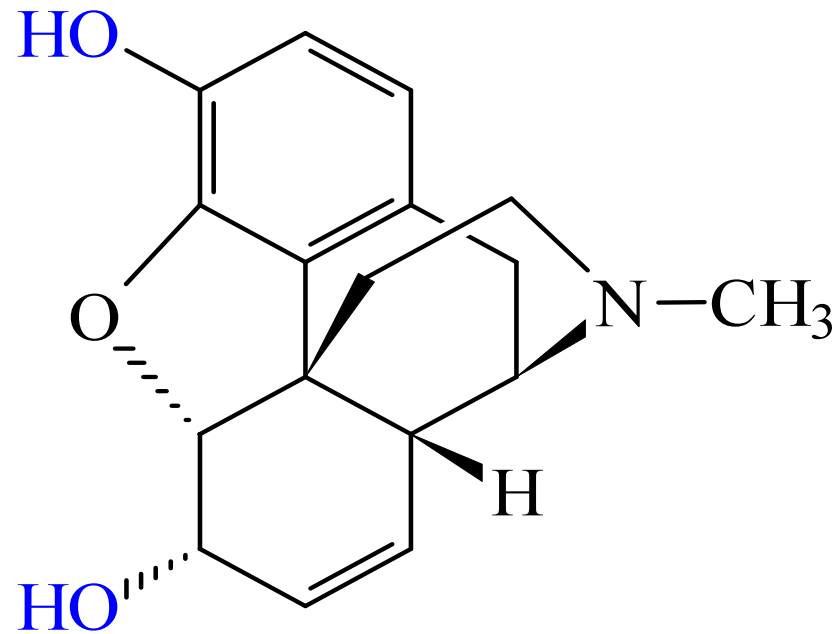


**glycerol**



**cholesterol**

## Alcohols are Found in Many Natural Products



**Morphine**

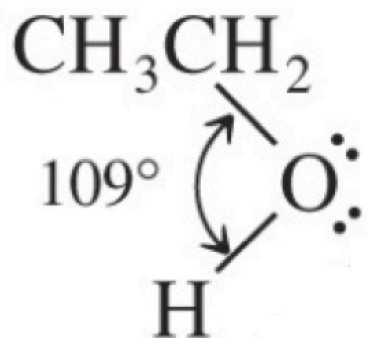
most abundant of opium's alkaloid



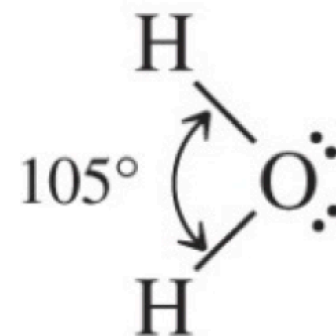
An alcohol can be viewed as either a hydroxyl derivative of an alkane or an alkyl derivative of water



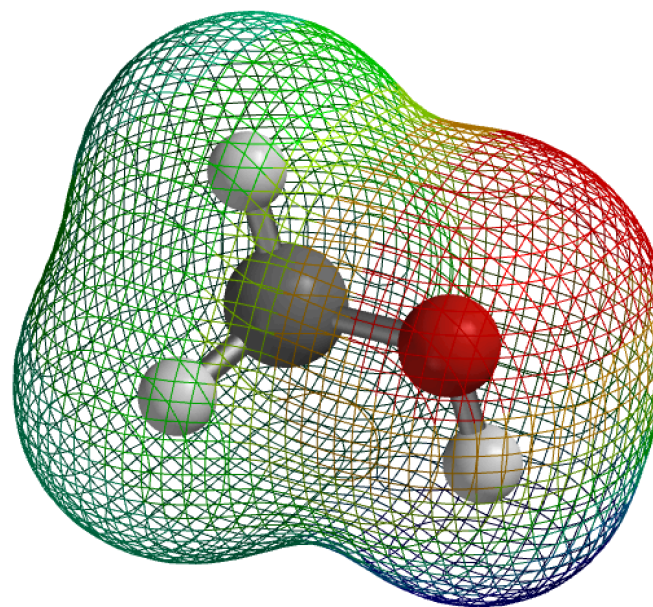
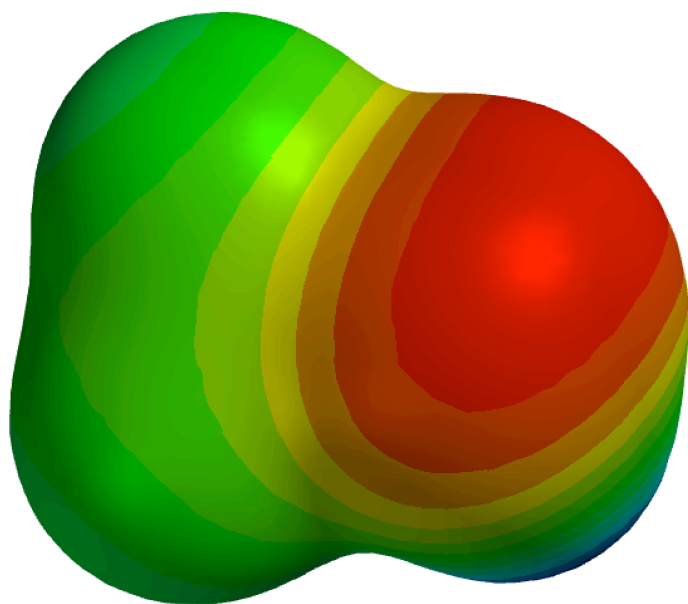
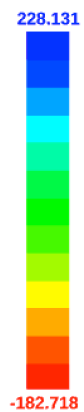
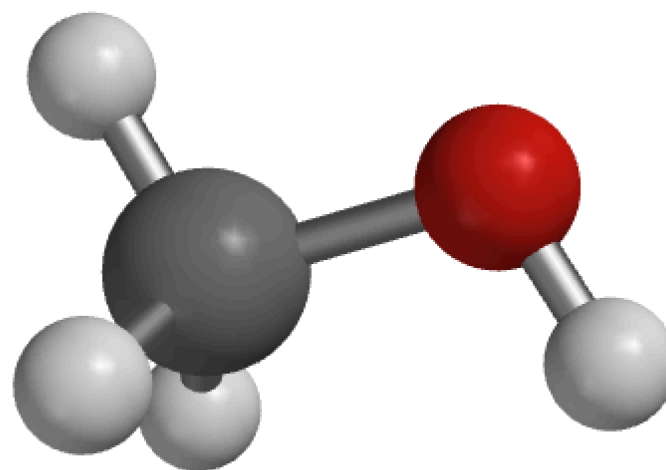
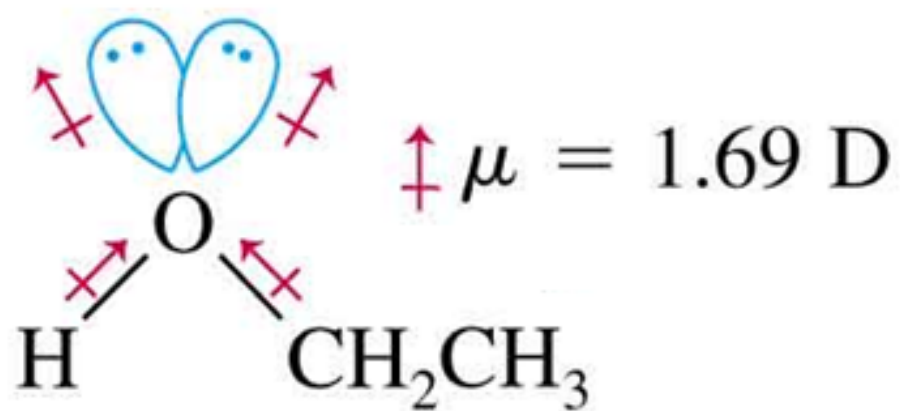
ethane

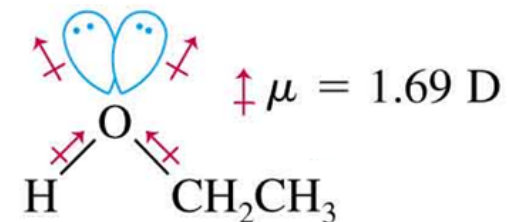
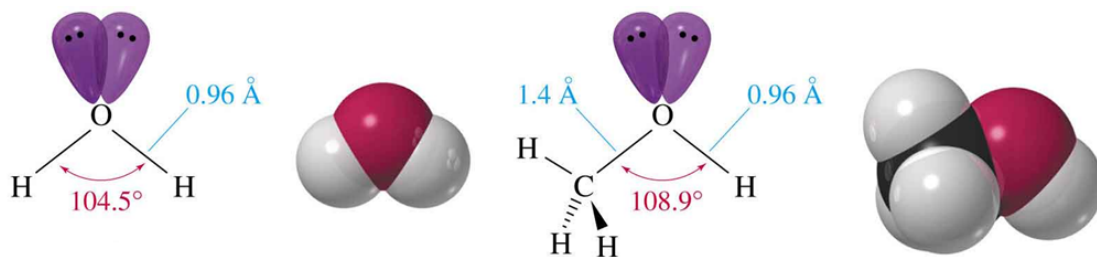


ethyl alcohol  
(ethanol)



water



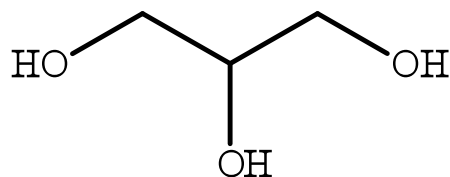
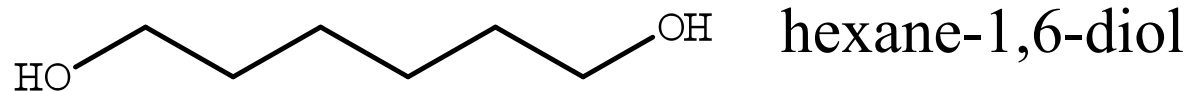


The presence of the OH group makes alcohols polar, therefore soluble in water, and it raises the boiling point with respect to the corresponding alkanes.

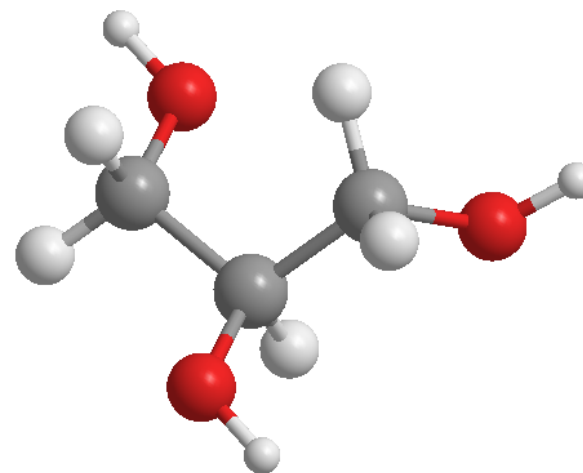
alkane	$T_{\text{eb}}$ (°C)	alcohol	$T_{\text{eb}}$ (°C)
CH <sub>4</sub>	-161.6	CH <sub>3</sub> OH	64.8
CH <sub>3</sub> CH <sub>3</sub>	-89.0	CH <sub>3</sub> CH <sub>2</sub> OH	78.3
CH <sub>3</sub> CH <sub>2</sub> CH <sub>3</sub>	-42.1	CH <sub>3</sub> CH <sub>2</sub> CH <sub>2</sub> OH	97.1

alcohol	solubility in H <sub>2</sub> O at 25 °C
methanol	miscible
ethanol	miscible
1-propanol	miscible
2-methyl-2-propanol	miscible
2-methyl-1-propanol	10%
1-butanol	9.1%
1-pentanol	2.7%
cyclohexanol	3.6%
1-hexanol	0.6%
phenol	9.3%

Compounds containing 2 (3) OH groups: diols (triols)

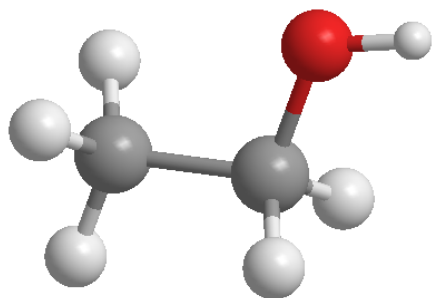
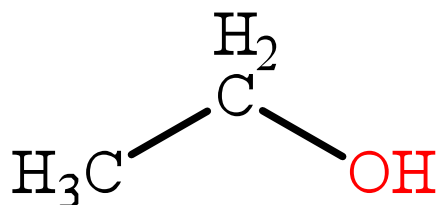


propane-1,2,3-triol (glycerol) → lipids

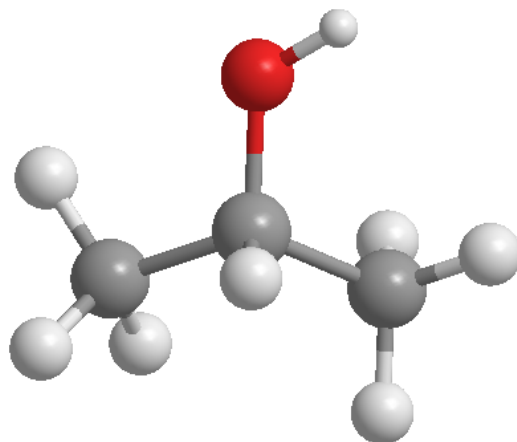
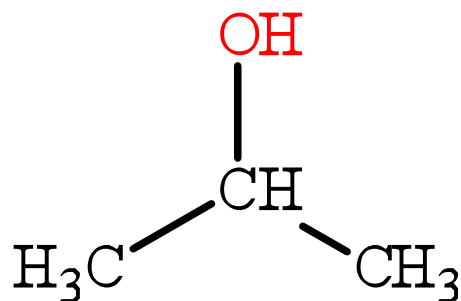


Alcohols are also classified according to the carbon the hydroxyl is directly attached to:

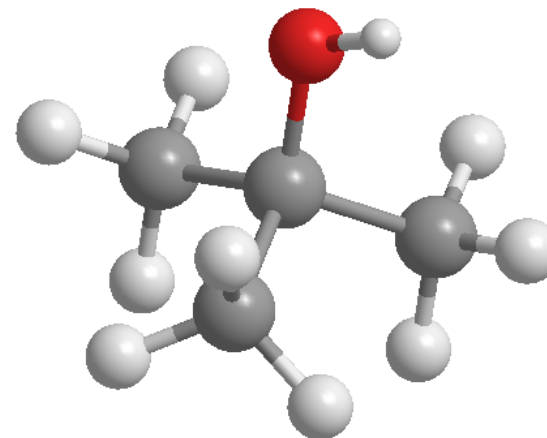
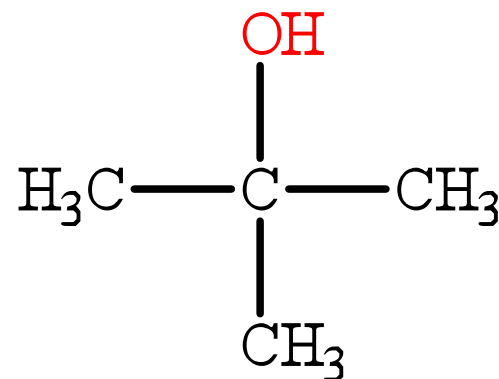
primary



secondary

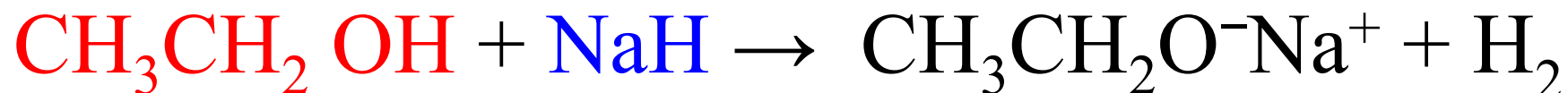
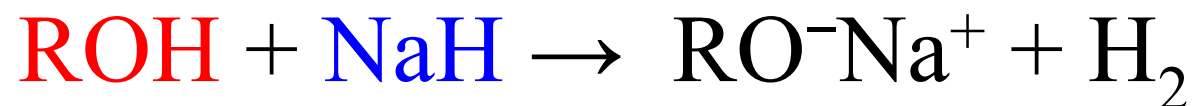


tertiary



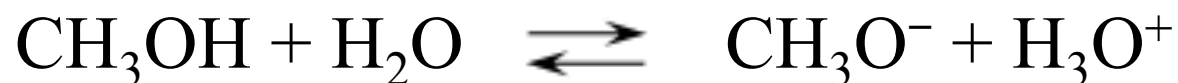
## Acid-base reactions

Alcohols have only **weak acidic character** (methanol and ethanol about as water, others with longer chain even less) and do not react with weak bases. It is therefore necessary to use very strong bases (e.g. sodium hydride)

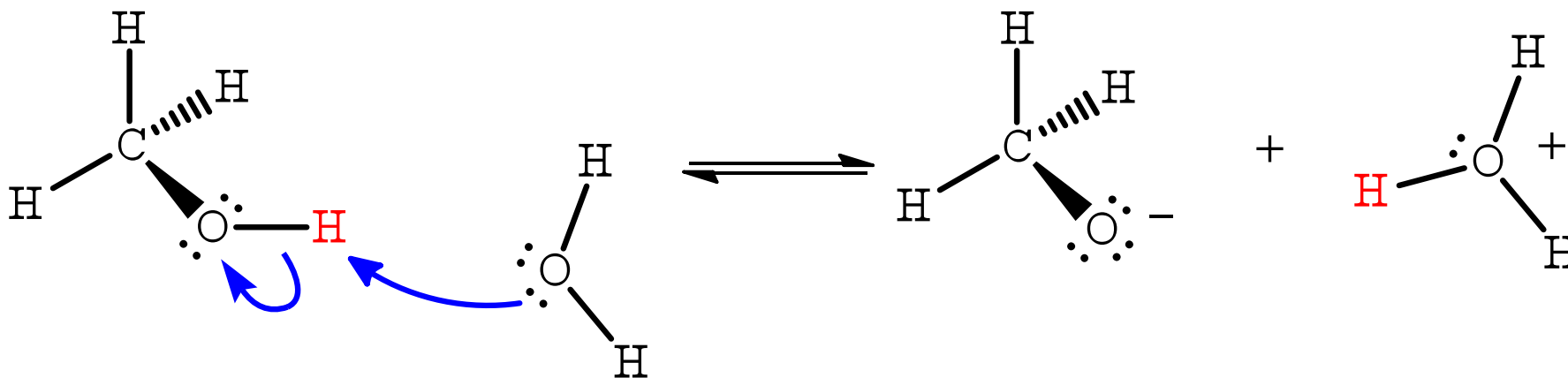


# Alcohols are very weak acids

In the absence of electron-withdrawing groups (which increase their acidity) and electron-repelling groups (which reduce their acidity) the ionization constants are ca.  $K_A = 1 \cdot 10^{-15} \text{ M}$

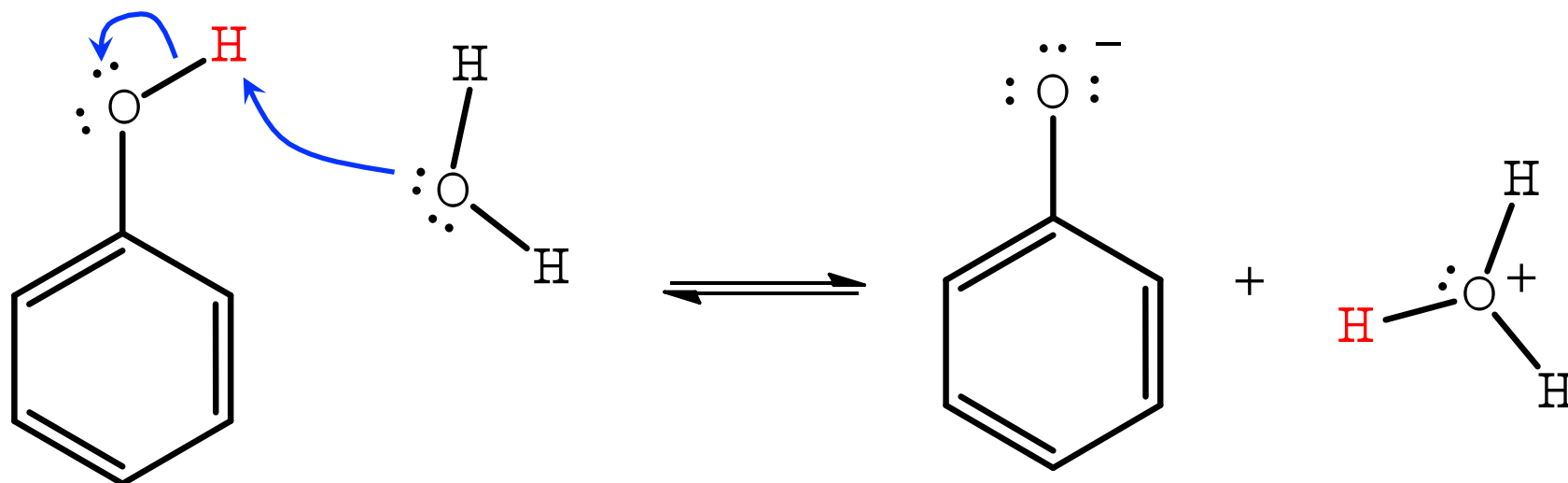


$$K_A = \frac{[\text{CH}_3\text{O}^-] \cdot [\text{H}_3\text{O}^+]}{[\text{CH}_3\text{OH}]} \cong 3.2 \cdot 10^{-16} \text{ M} \quad \text{p}K_A = 15.5$$



Although they have a hydroxyl group (-OH), the phenols do not behave like alcohols.

Phenol has a weak acidic character:  $pK_A = 9.95$

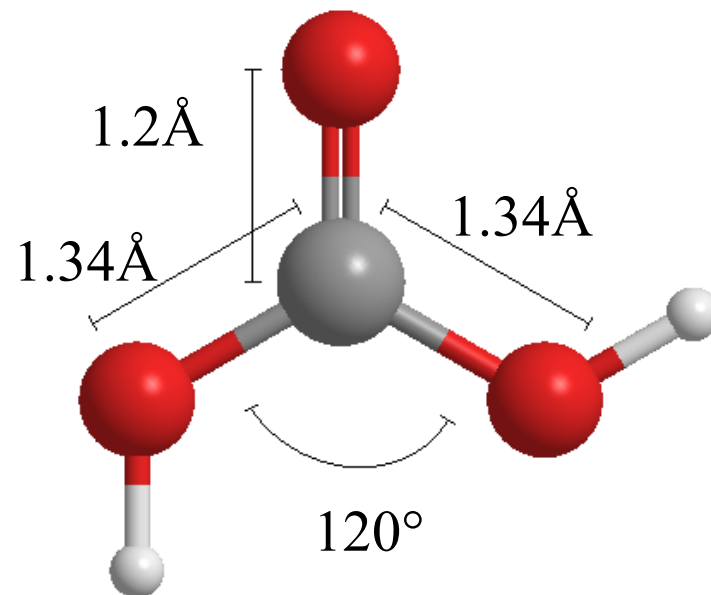


This happens because the negative charge of the anion that is formed by the dissociation of the hydrogen of the hydroxyl group (phenate ion) is stabilized by resonance on the whole aromatic ring.



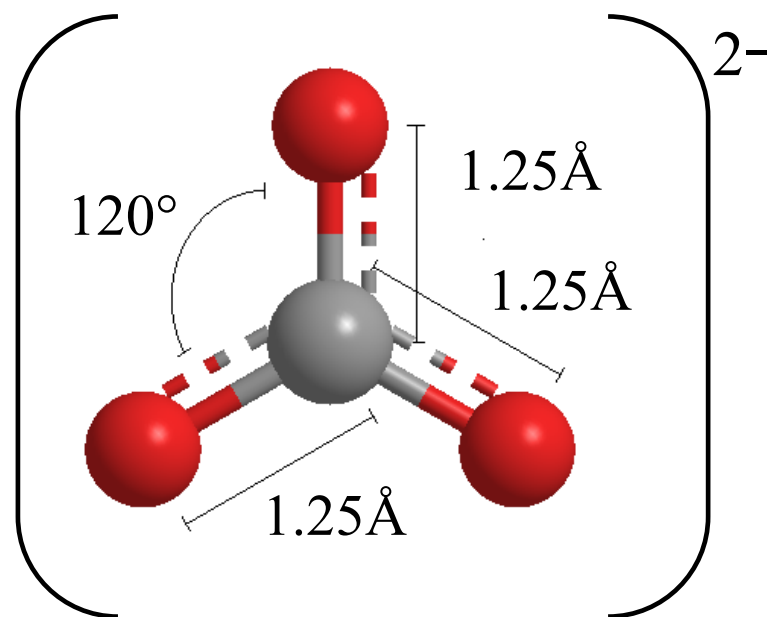
Resonance limit forms

carbonic acid  $\text{H}_2\text{CO}_3$

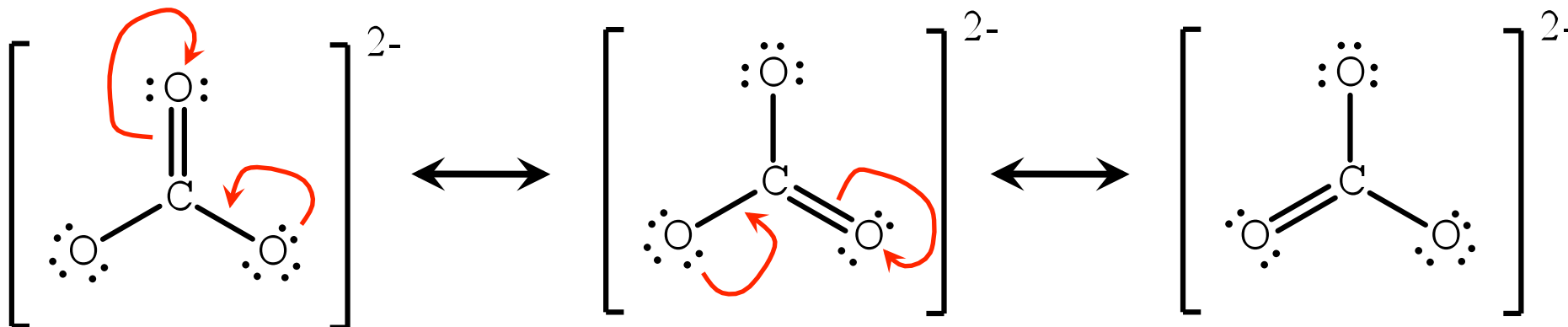


carbonate dianion  $\text{CO}_3^{2-}$

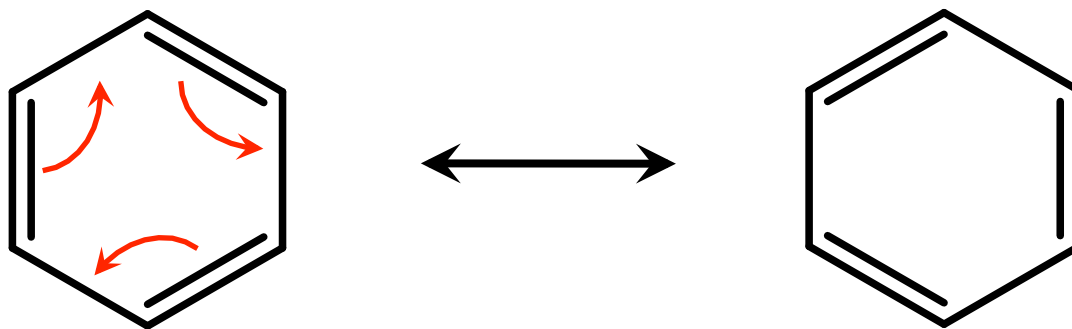
3 resonance limit forms



# Resonance limit forms

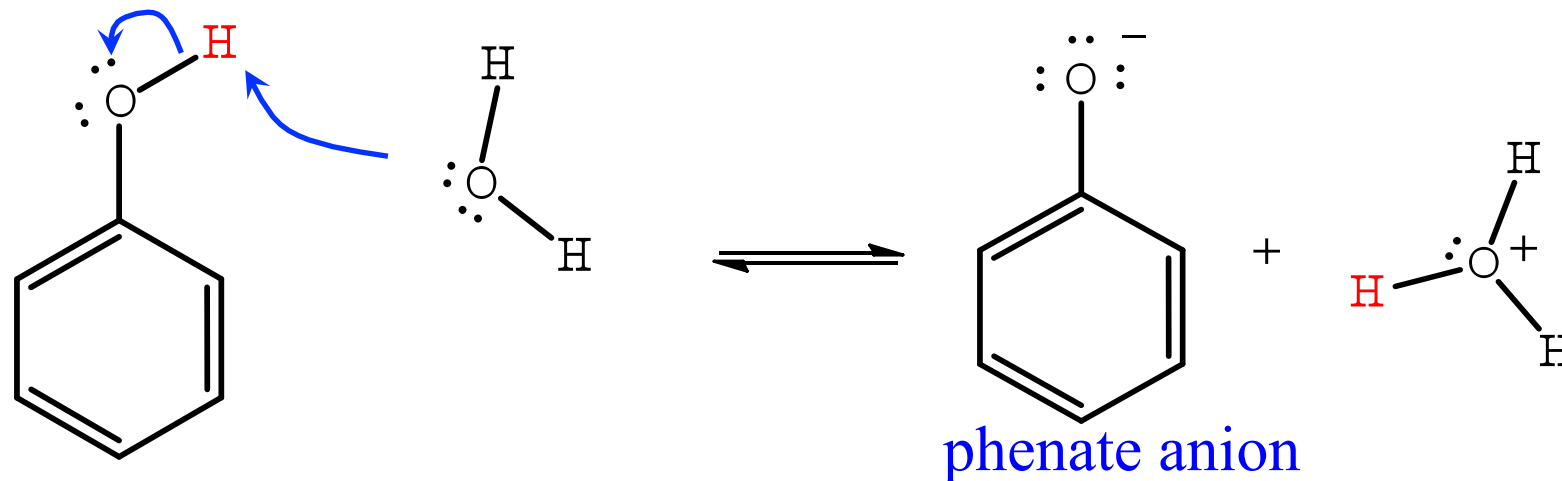


carbonate dianion

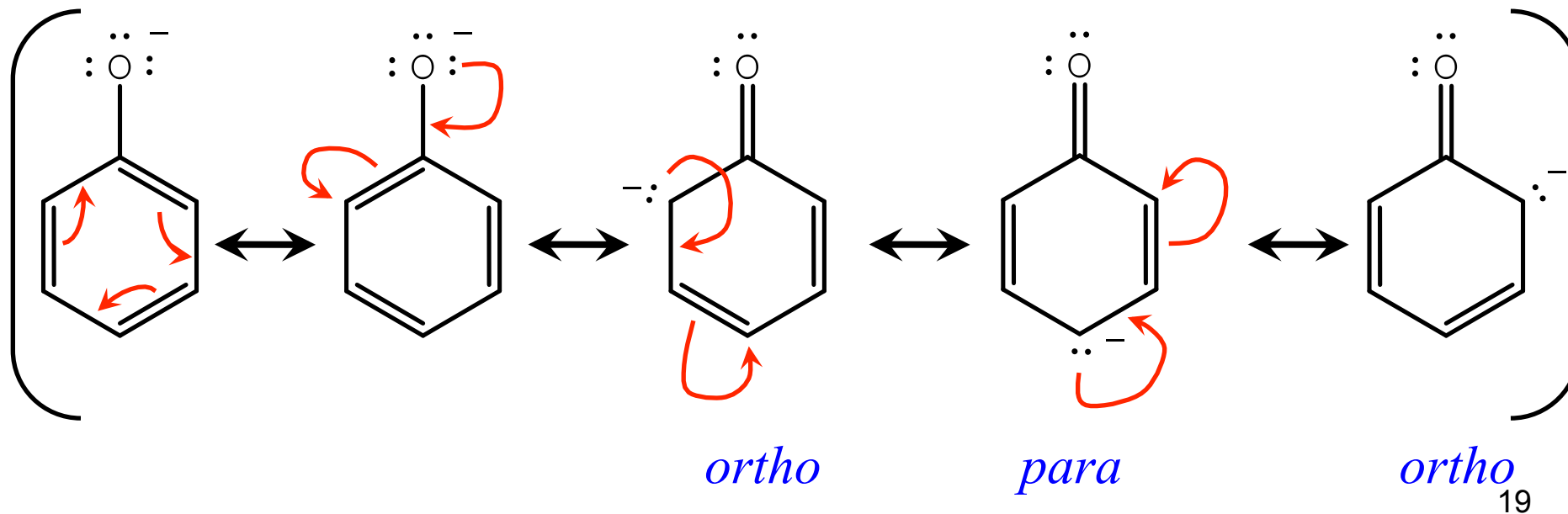


benzene

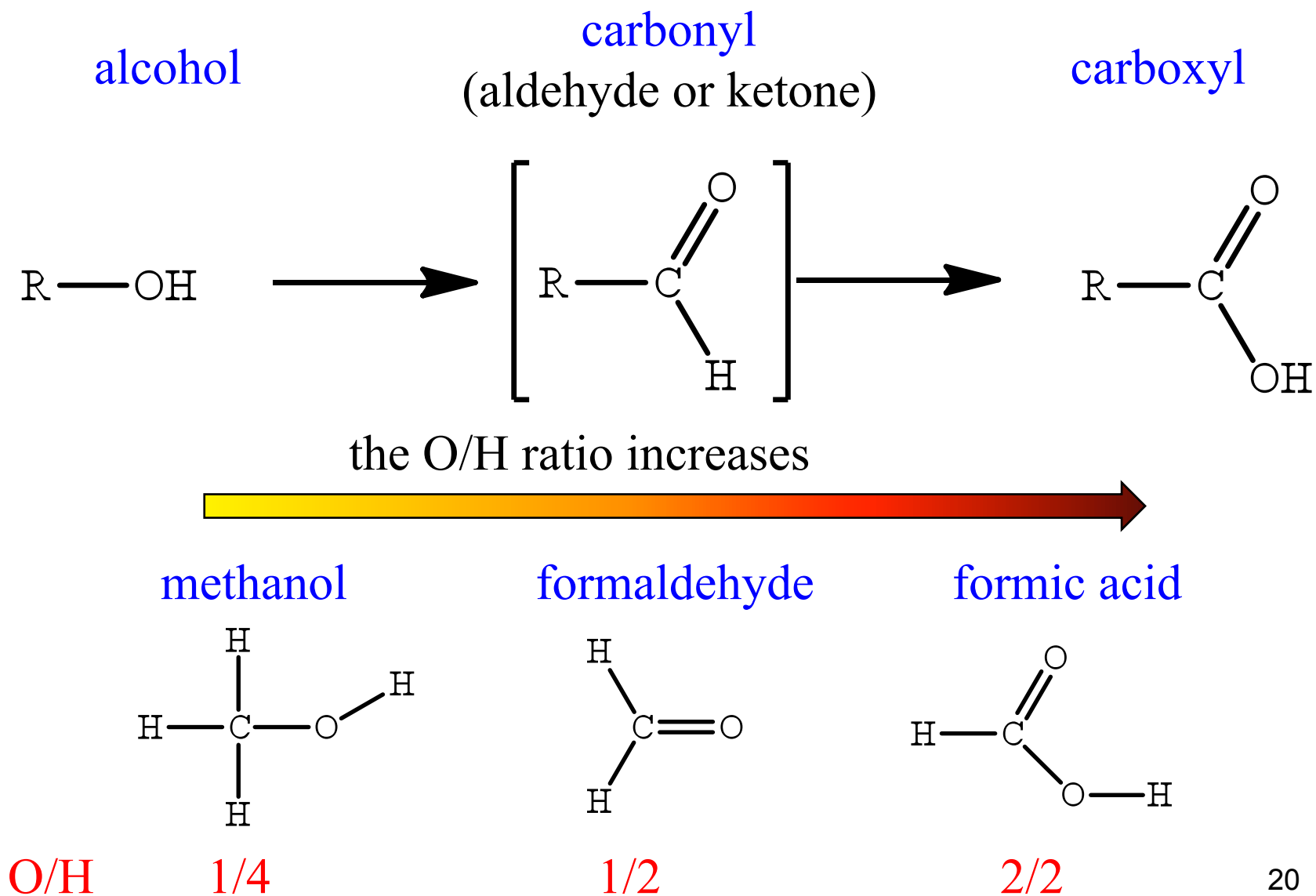
The phenate anion is stabilized by resonance among several limit forms. Phenols are stronger acids than alcohols.



The charge is delocalized on the ring and the conjugate base is stabilized

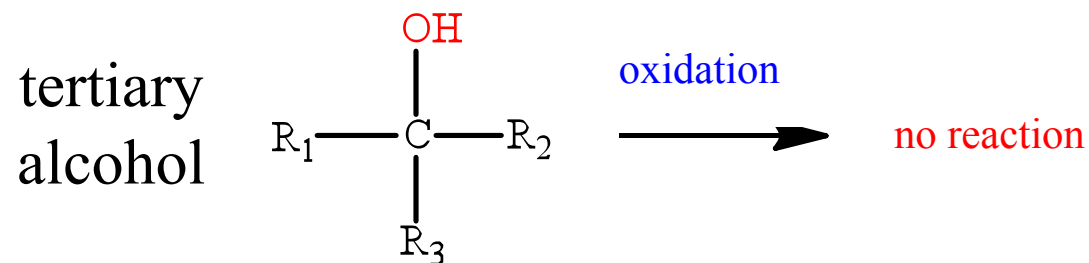
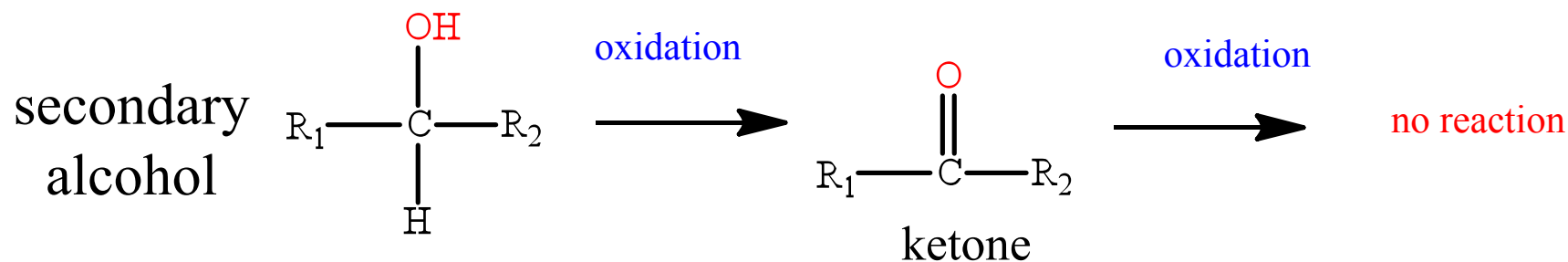
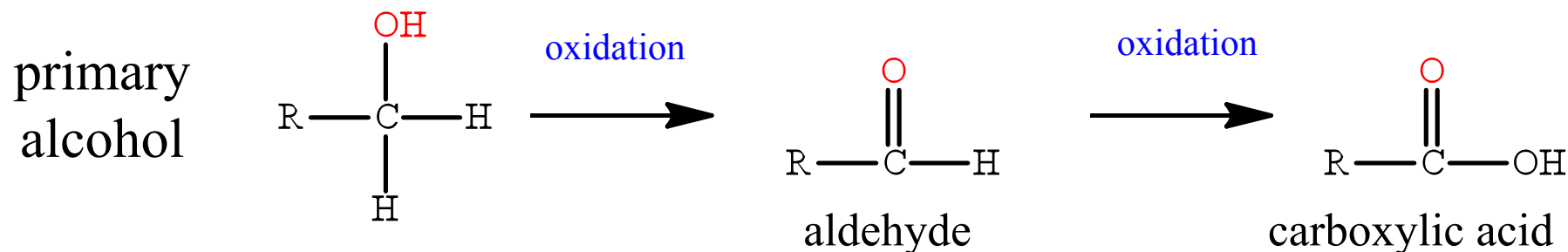


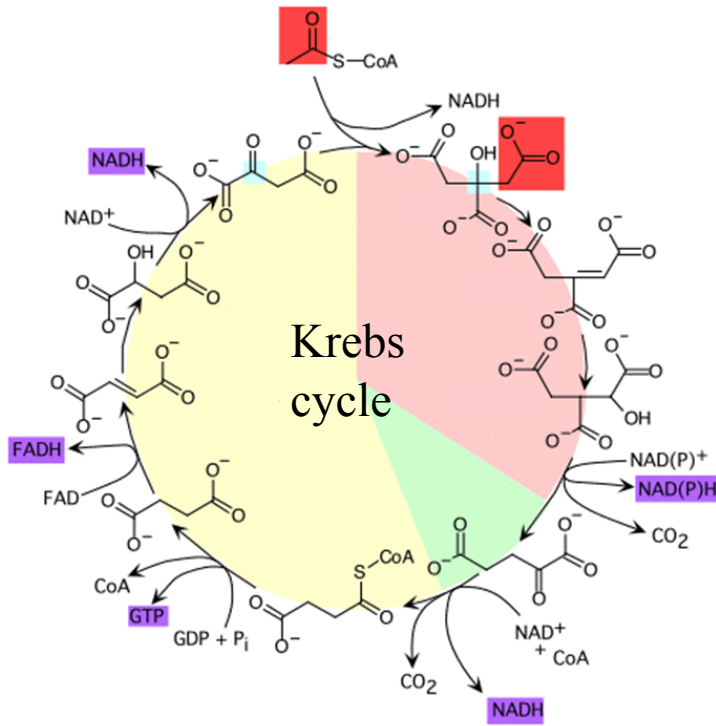
# Alcohol oxidation reactions



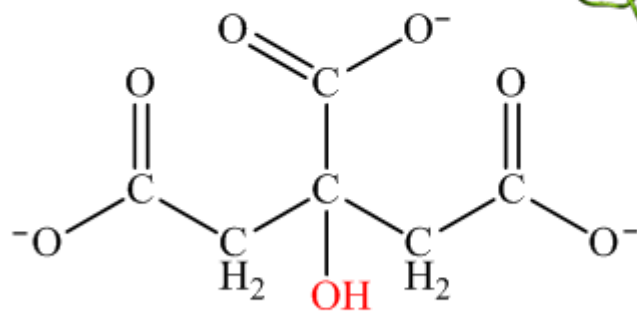
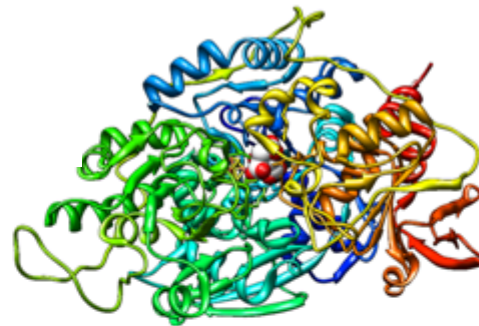
# Alcohol oxidation reactions

The result of alcohol oxidation depends on the nature of the alcoholic C substituents and the presence of at least 1 H atom

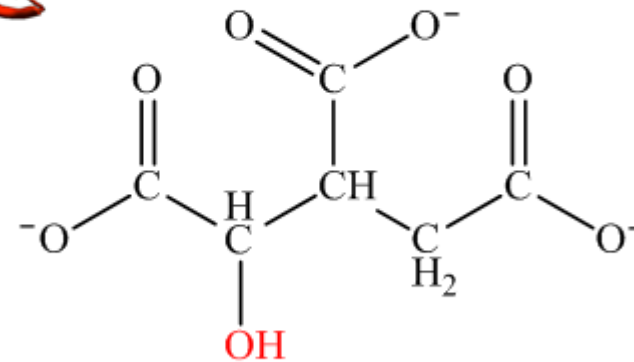




In the Krebs cycle, citrate (a tertiary alcohol) is isomerized to isocitrate (a secondary alcohol) by aconitase. This alcohol will be then oxidized to a ketone.

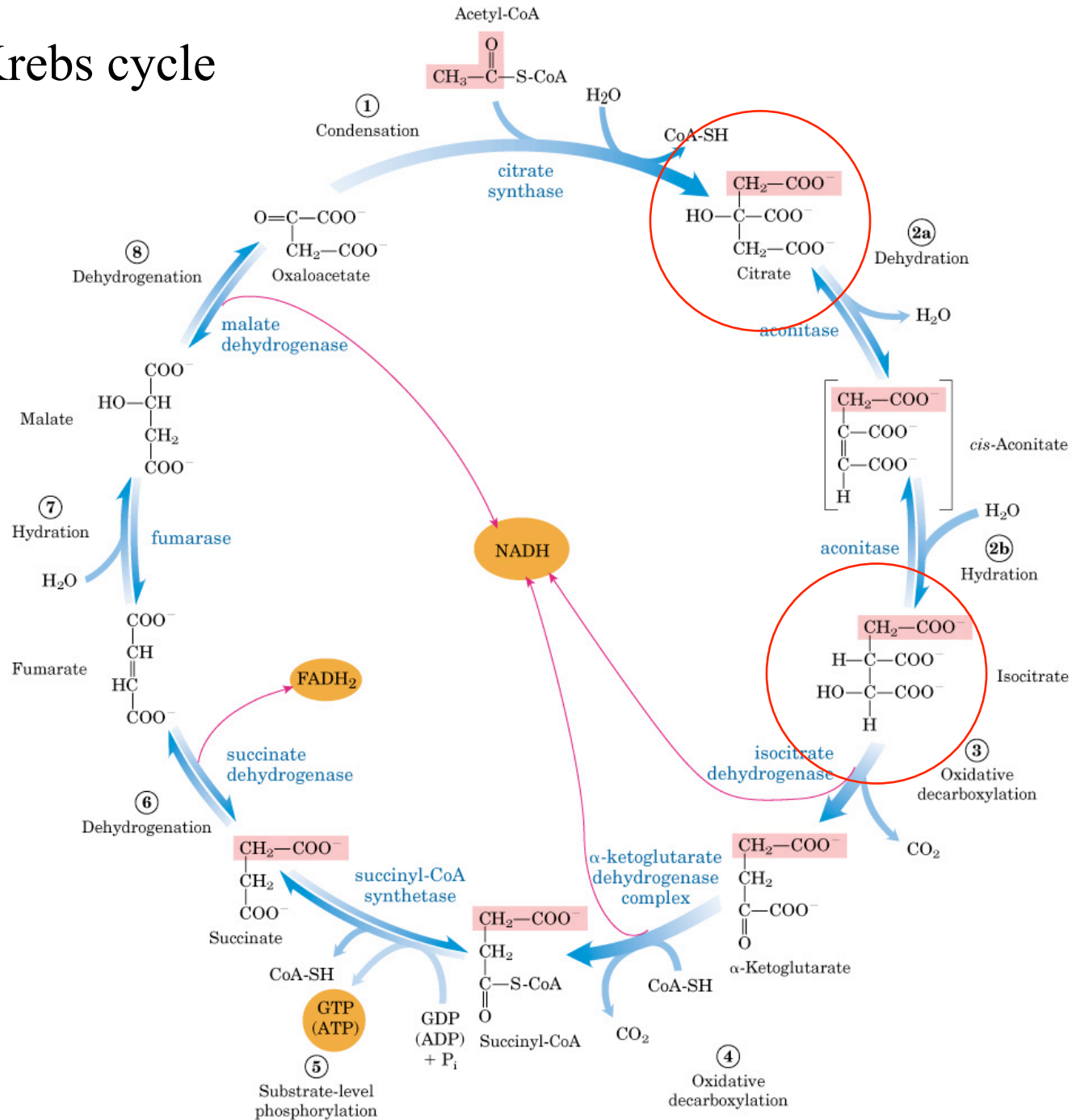


citrate



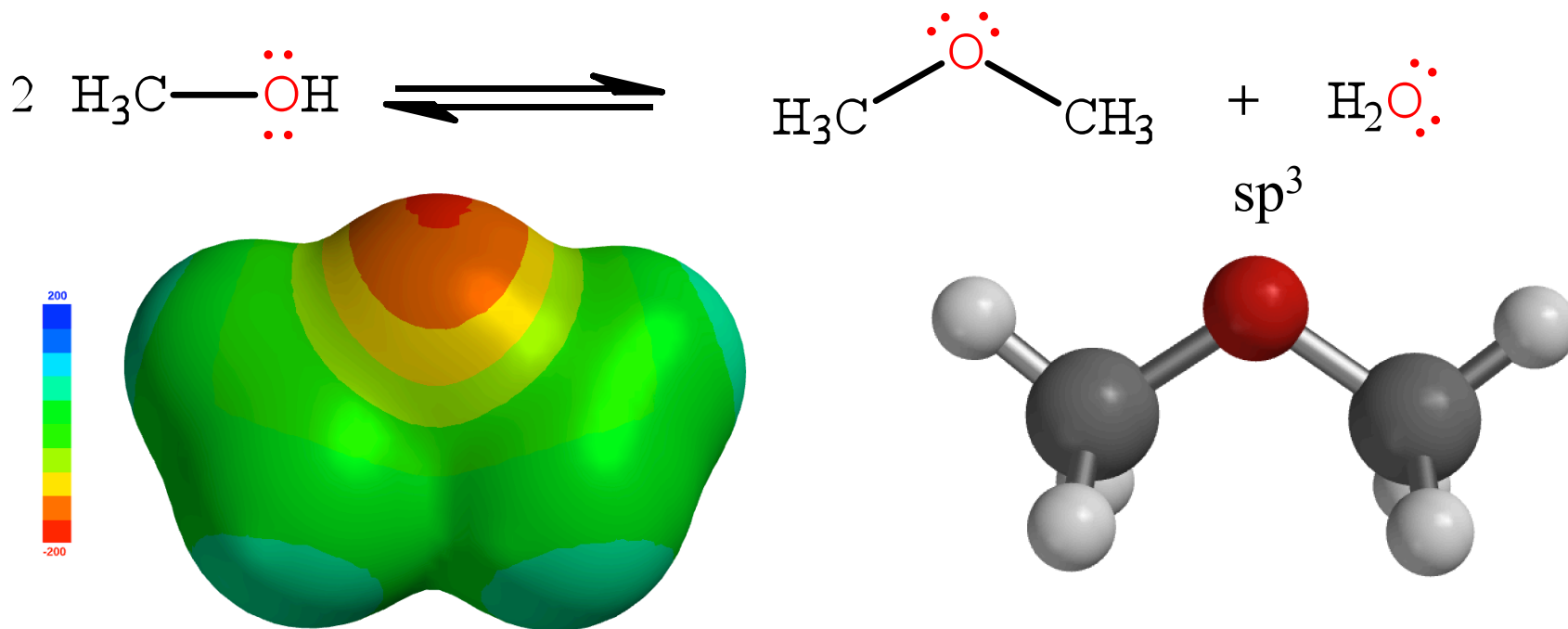
isocitrate

# The Krebs cycle



# Ethers

Ether is the functional group deriving from the combination of two alcohols, with the elimination of a water molecule. These can be considered organic derivatives of water in which both hydrogens are replaced by organic groups. The bond angle at the oxygen is close to the tetrahedral angle



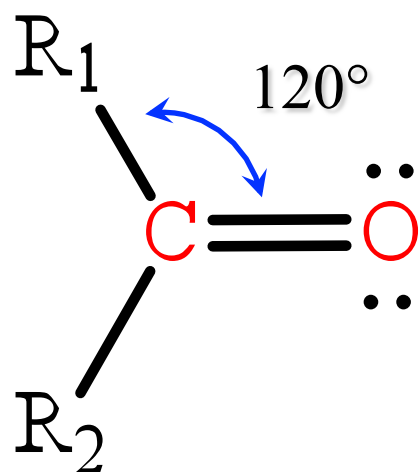
Ethers are molecules that behave like aprotic polar solvents, they lack hydrogen atoms available to form H-bonds. They are very volatile and have anesthetic properties.



# Aldehydes and ketones

Can be considered oxidation products of primary and secondary alcohols.  
Consistently, the reduction of an aldehyde returns a primary alcohol and that of secondary alcohol a ketone.

They contain the **carbonyl functional group** (C = O)

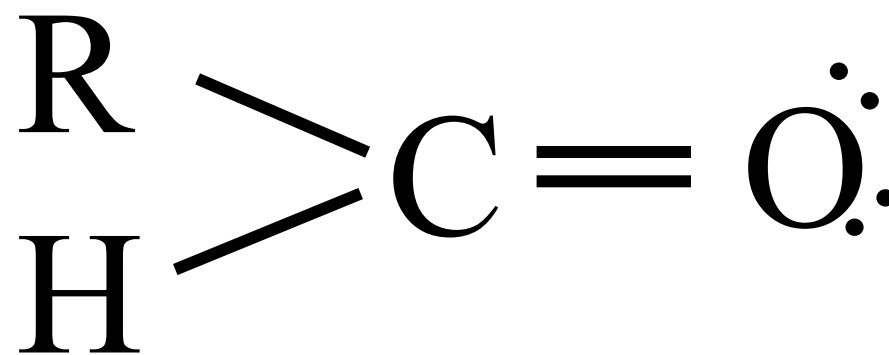
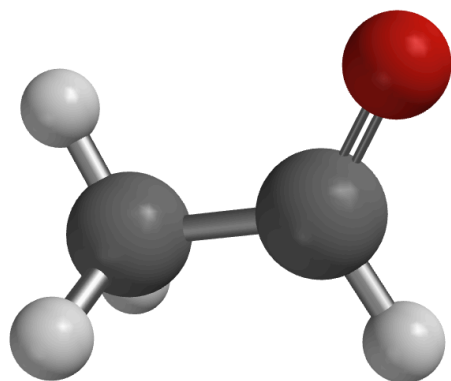
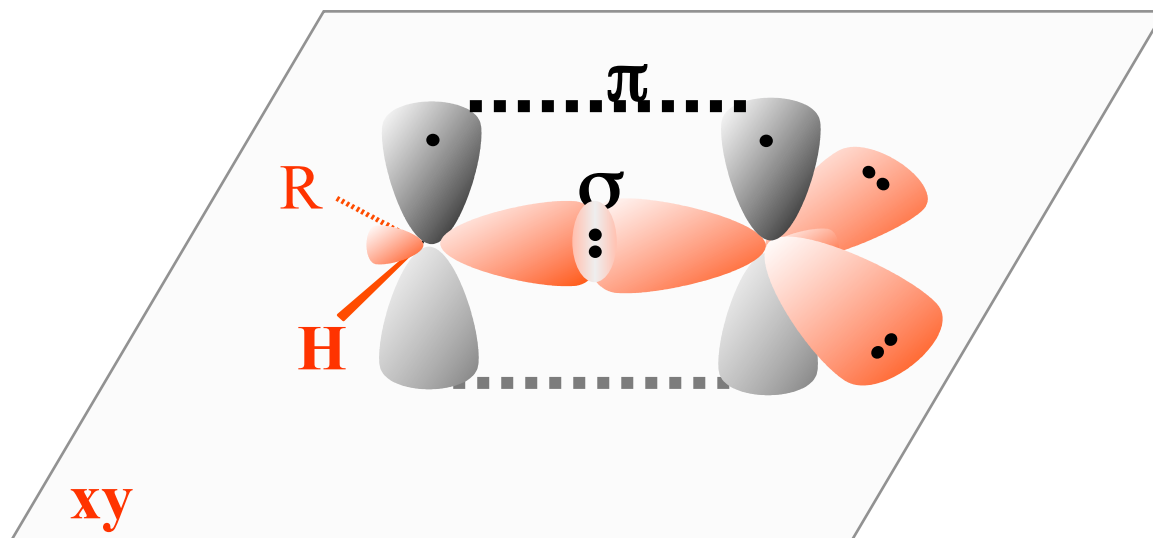
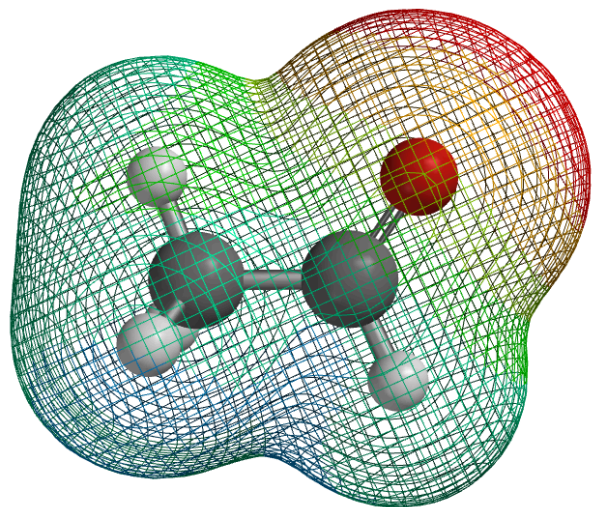


in the double bond between C and O we find a  $\sigma$  and  $\pi$  bond

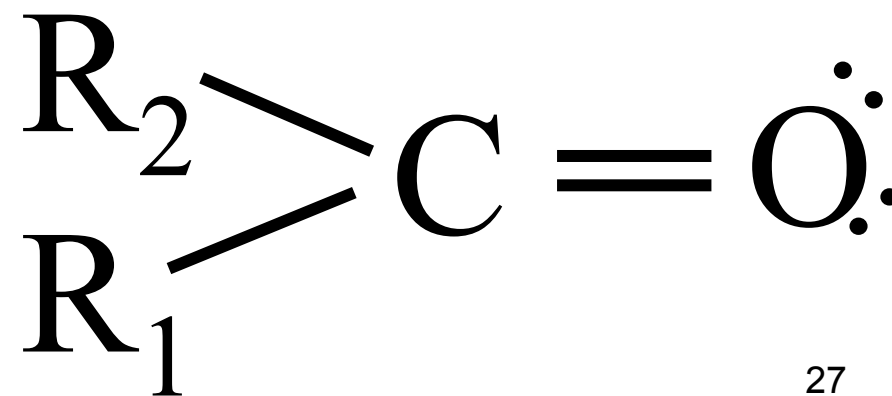
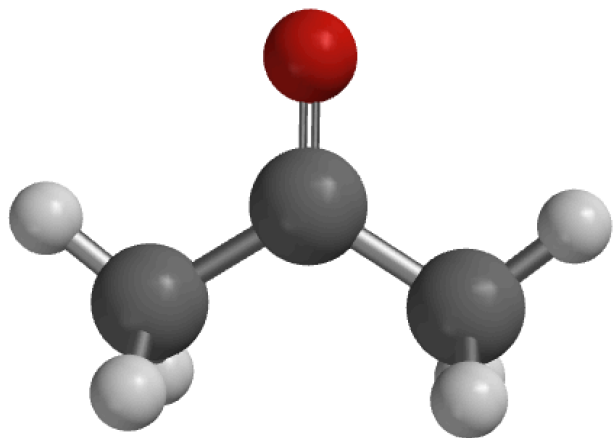
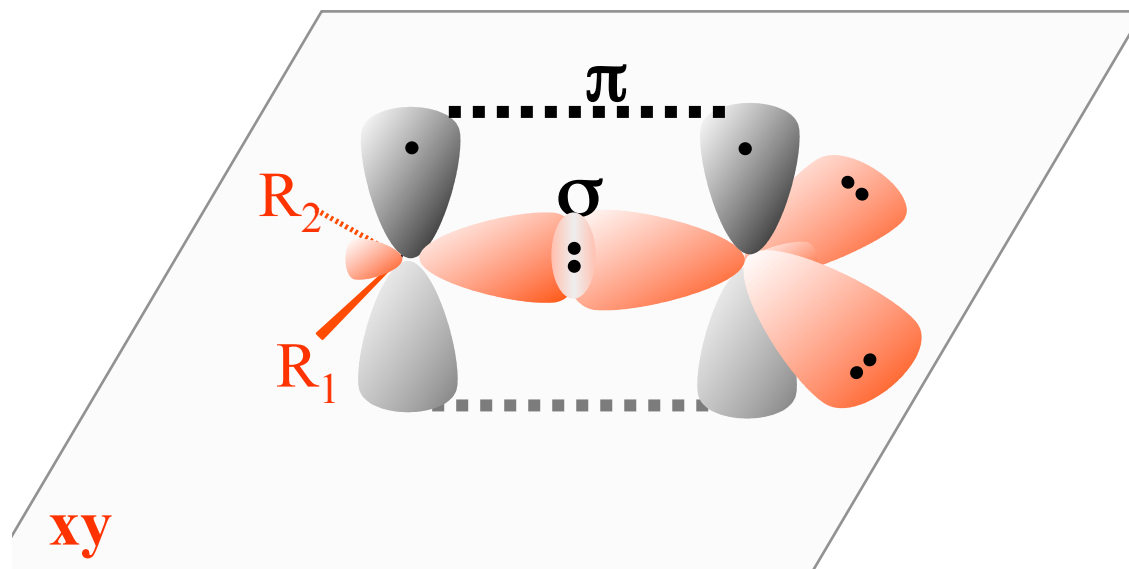
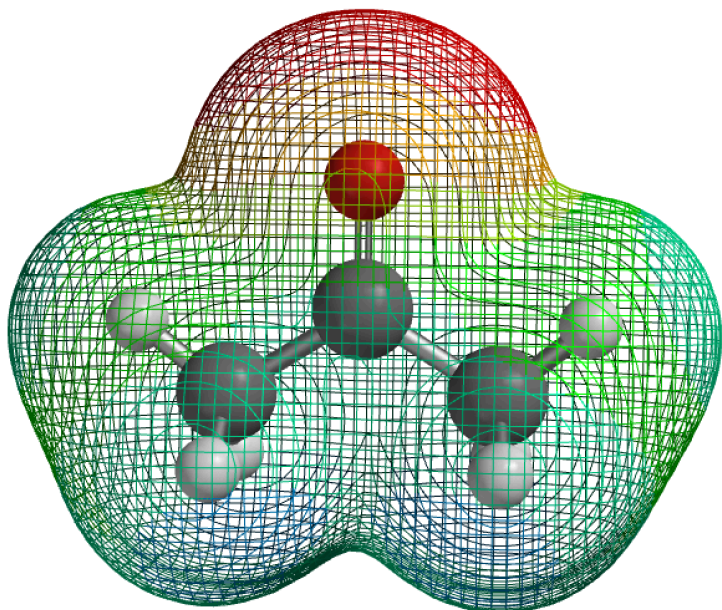
C and O have  $sp^2$  hybridization and the carbonyl group is planar with angles of  $120^\circ$

R<sub>1</sub> and R<sub>2</sub> can be two different groups and give rise to various functional groups containing the carbonyl group

Aldehydes contain a carbonyl group



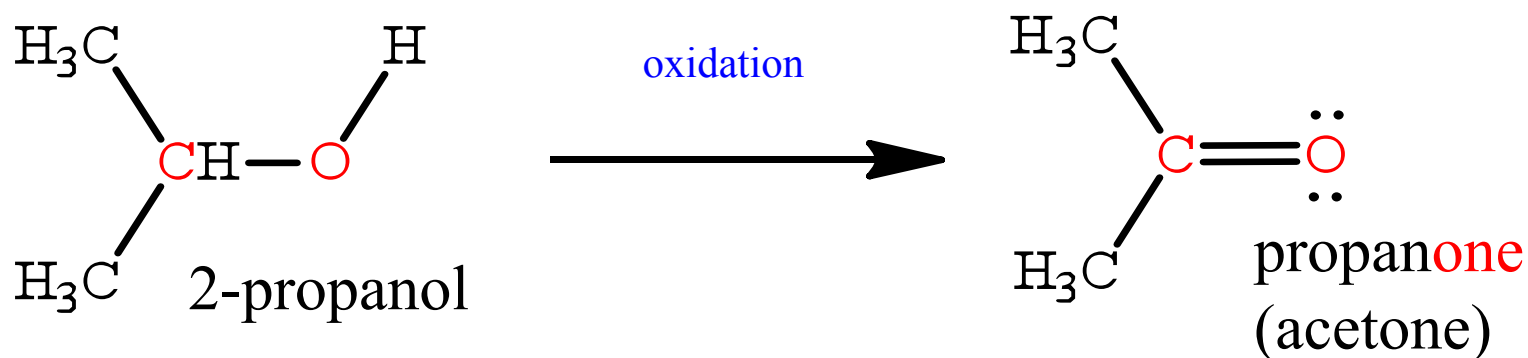
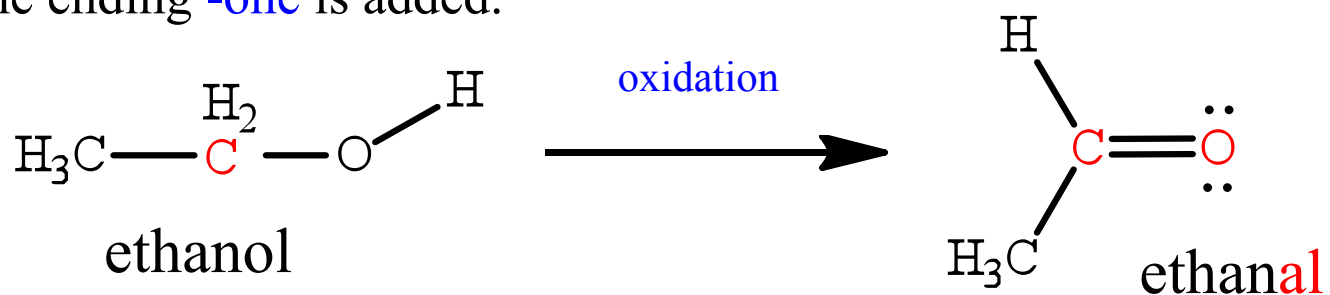
Ketones also contain a carbonyl group



## Nomenclature

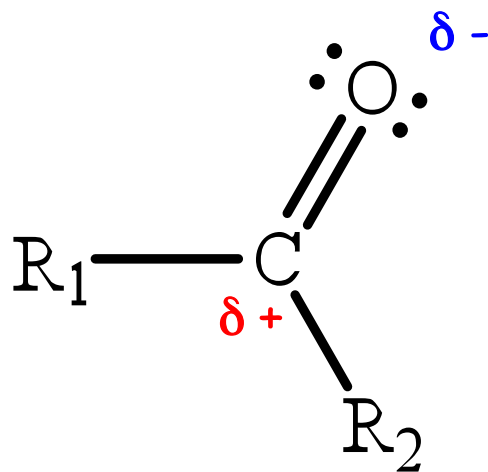
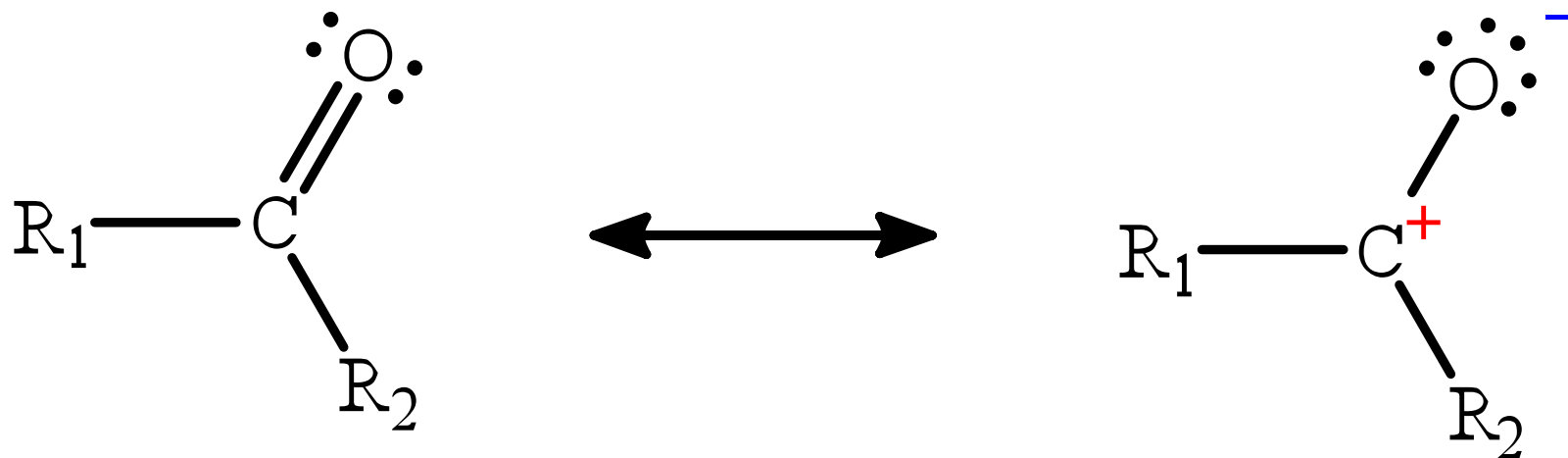
The name of aldehydes is given by the name of the corresponding hydrocarbon to which the ending **-al** is added

The name of ketones is given by the name of the corresponding hydrocarbon to which the ending **-one** is added.



The  $\text{C}=\text{O}$  group presents a planar trigonal geometry, therefore in the oxidation from alcohol to aldehyde or from alcohol to ketone, there is also a change of the corresponding hybridization of carbon from  $\text{sp}^3$  to  $\text{sp}^2$ .

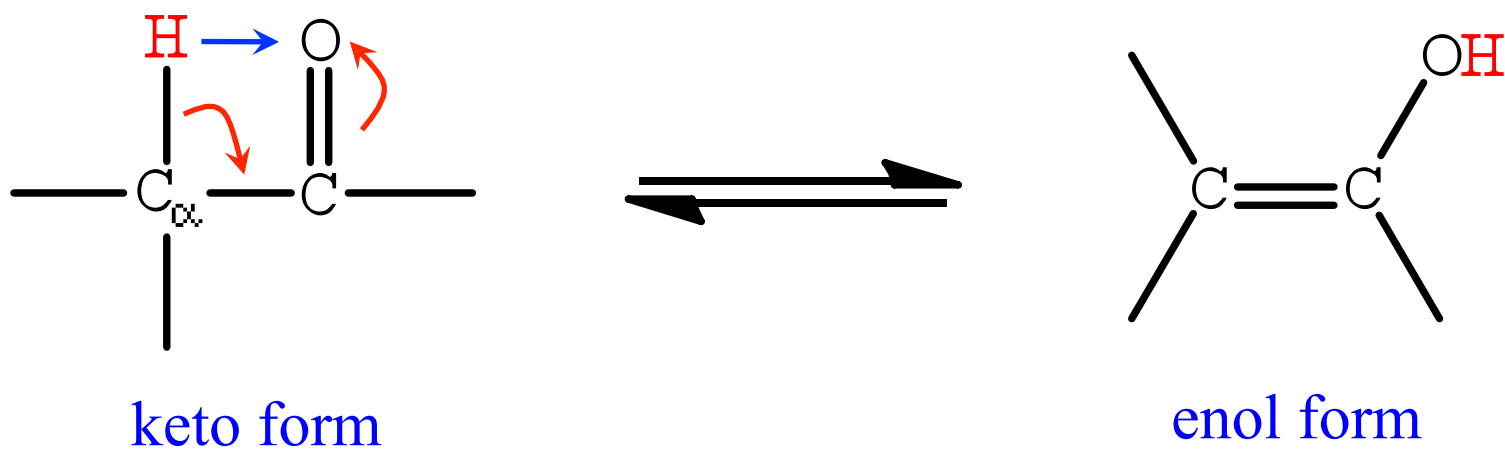
## Two resonant limit structures



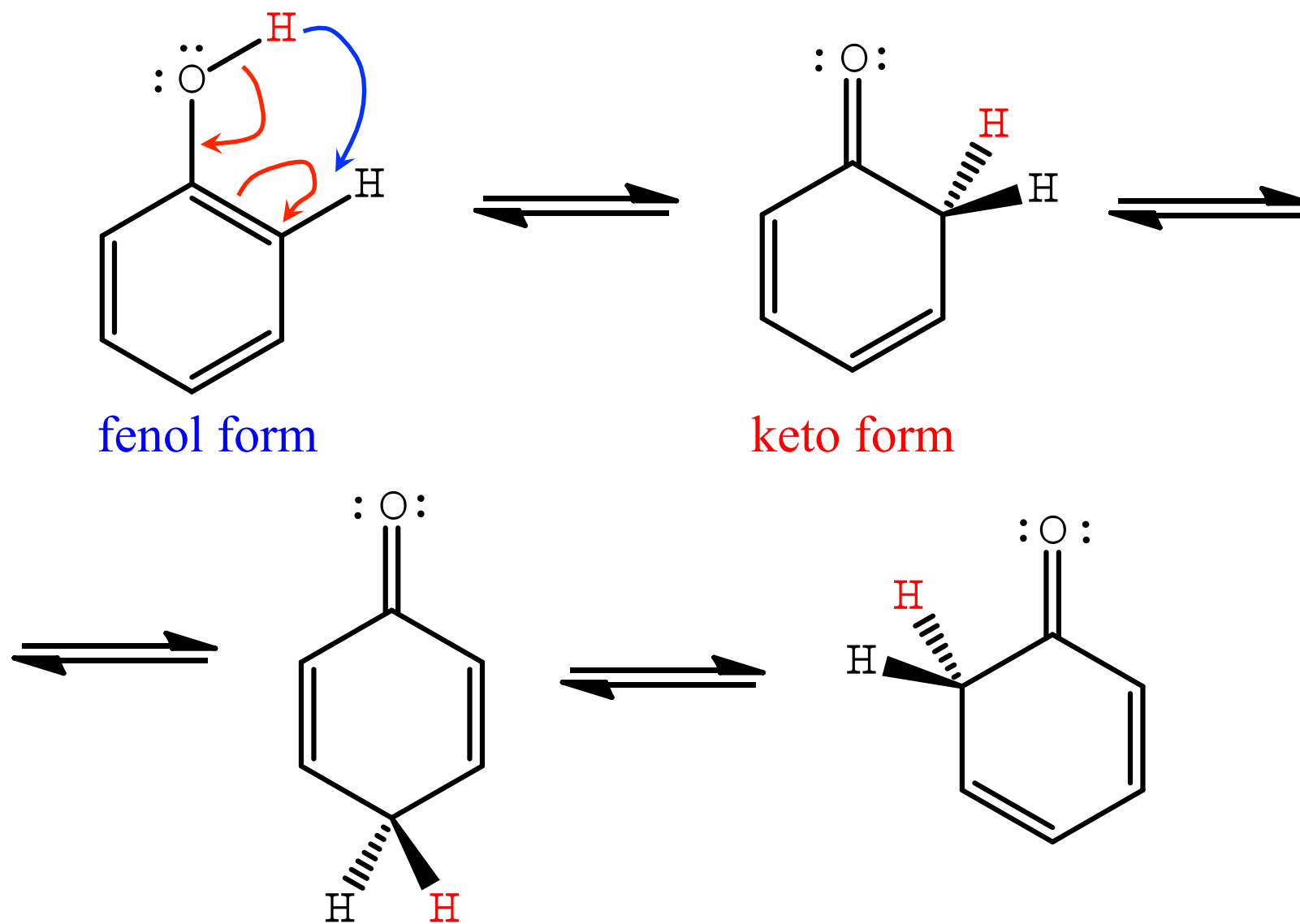
The carbonyl CO bond is polarized and this determines its reactivity

# Keto-enol tautomerism

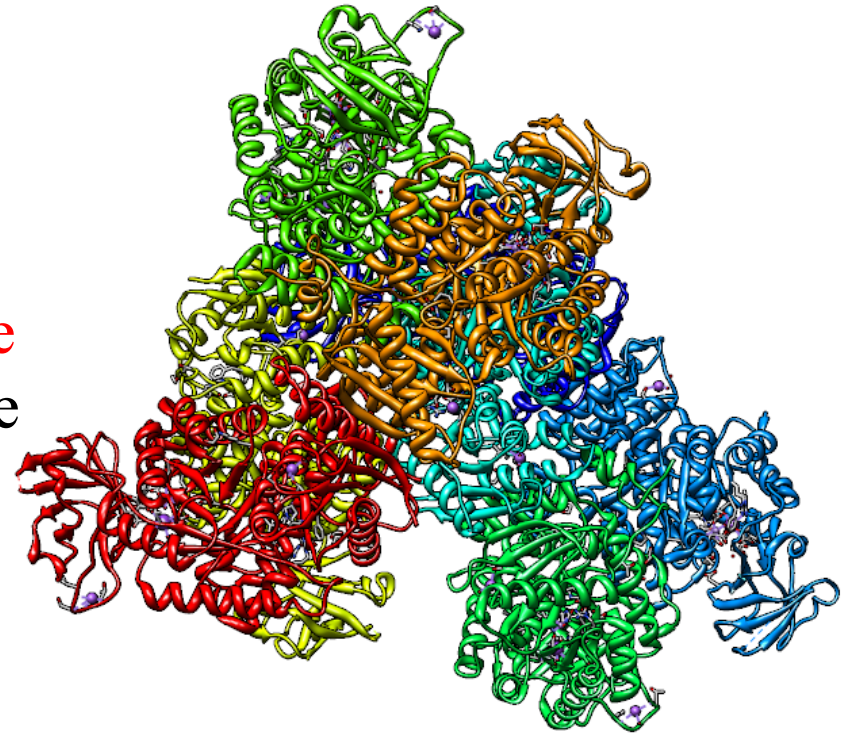
- Aldehydes and ketones can exist at equilibrium in the two keto and enol forms, which differ in the position of a proton and a double bond.
- This isomerism is called tautomerism, the forms are called tautomers and are two particular isomers that are in equilibrium.
- A carbonyl compound, in order to exist in the enol form, must have a hydrogen atom linked to the carbon atom adjacent to the carbonyl ( $\alpha$  carbon).



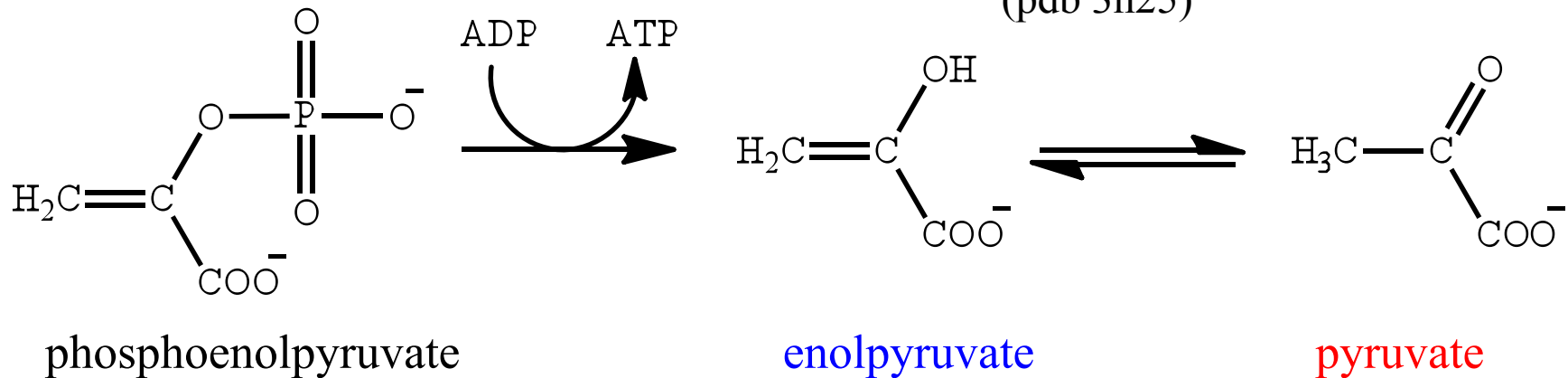
# Keto-enol tautomerism in phenol



In the last stage of glycolysis, **pyruvate kinase** catalyzes the phosphorylation of ADP with the formation of **enolpyruvate** which spontaneously tautomerizes to the more stable keto form, pyruvate.



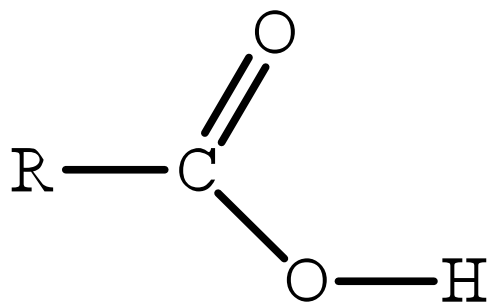
rabbit muscle pyruvate kinase  
(pdb 3n25)





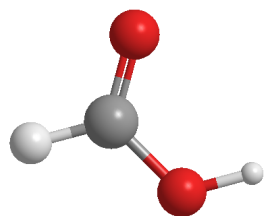
# Carboxylic acids

functional group: **carboxyl**

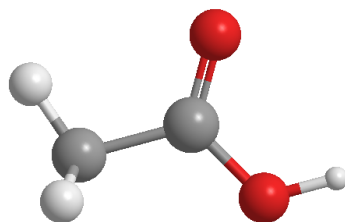


Nomenclature:

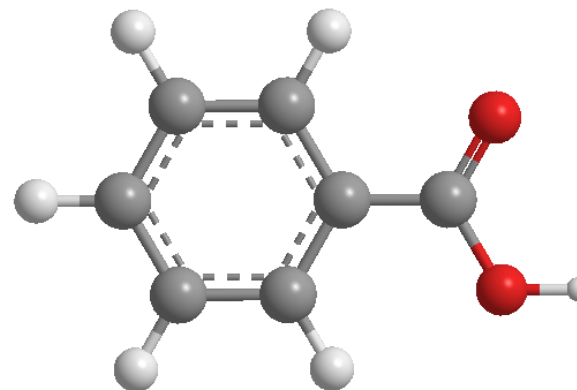
carboxylic acids are named by adding the **-oic** ending to the theme of the corresponding hydrocarbon name



H-COOH  
methanoic acid  
**formic acid**

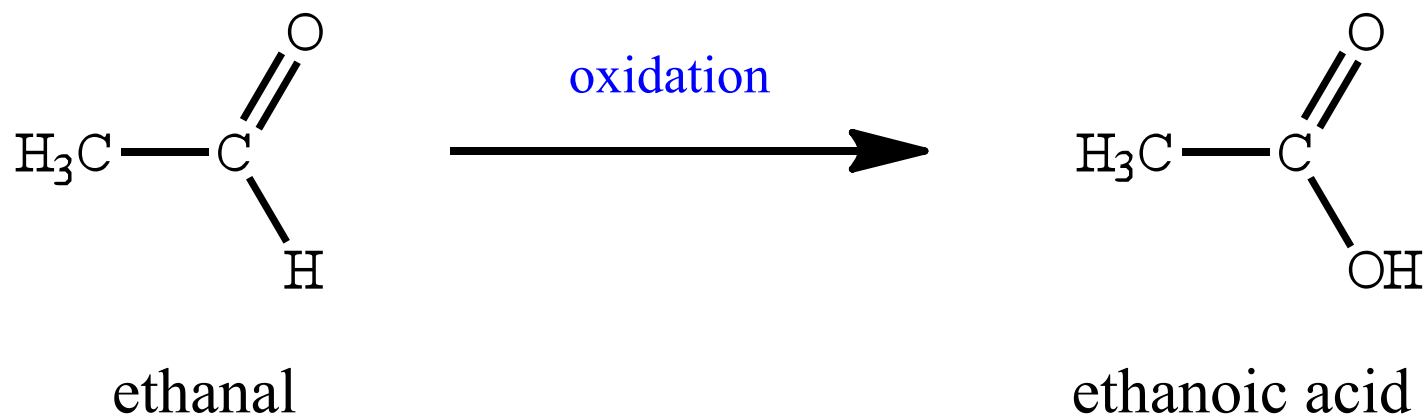


CH<sub>3</sub>-COOH  
ethanoic acid  
**acetic acid**

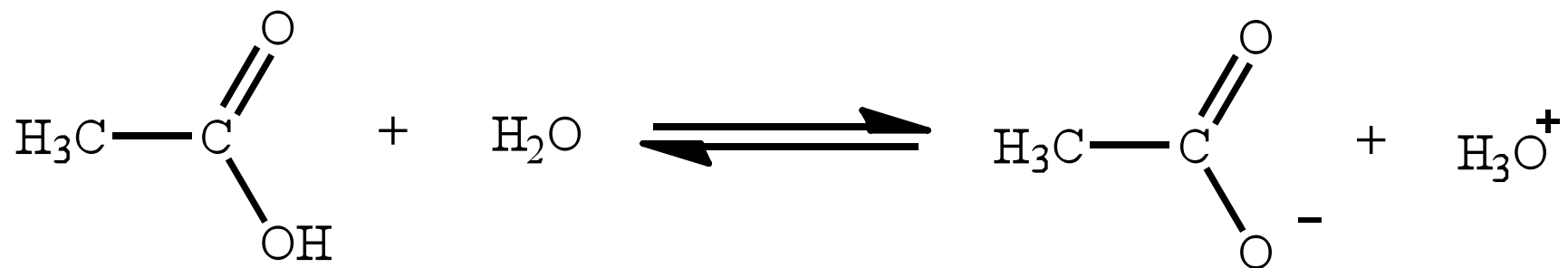


C<sub>6</sub>H<sub>5</sub>-COOH  
benzoic acid

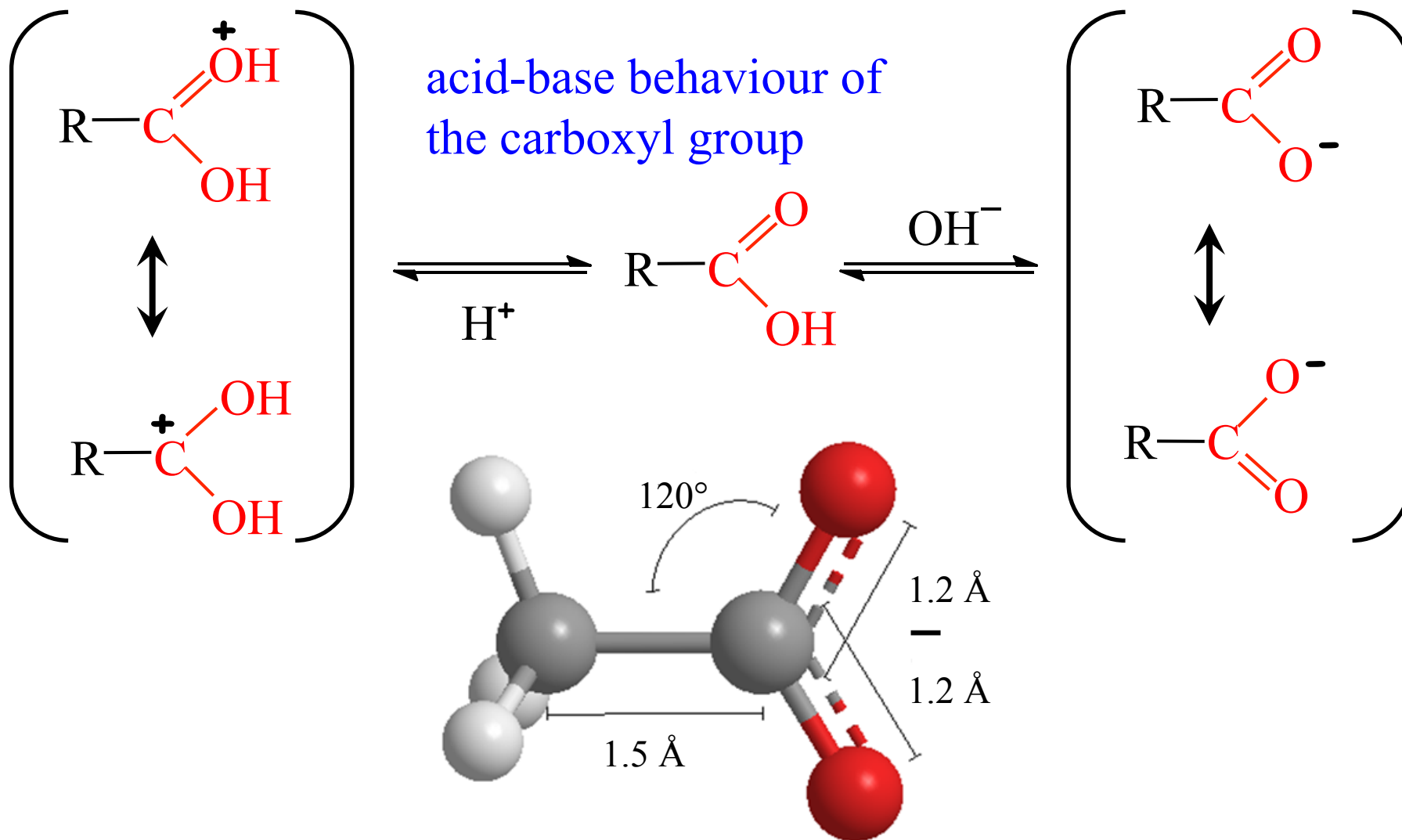
Carboxylic acids are produced from the corresponding aldehydes:



Carboxylic acids are weak acids, with  $\text{pK}_A$  values between 3 & 6

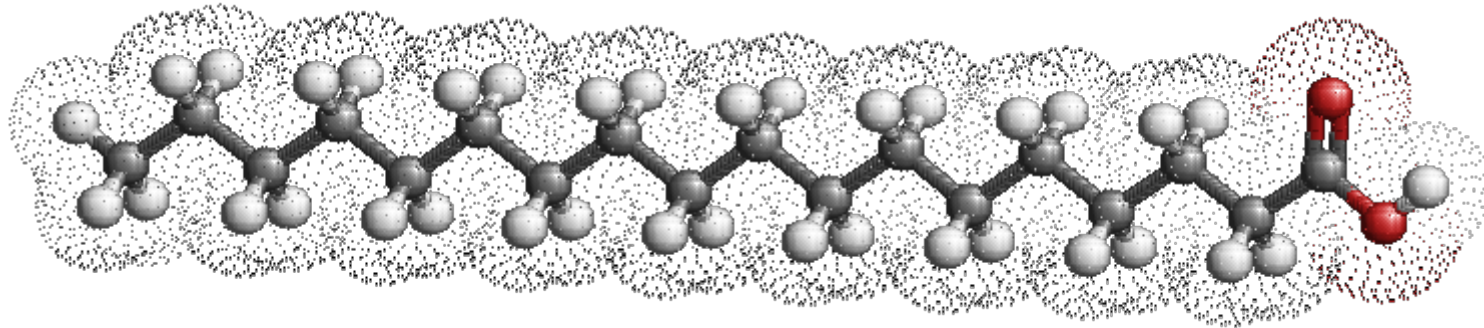


# Resonance increases the acidity of the carboxyl group

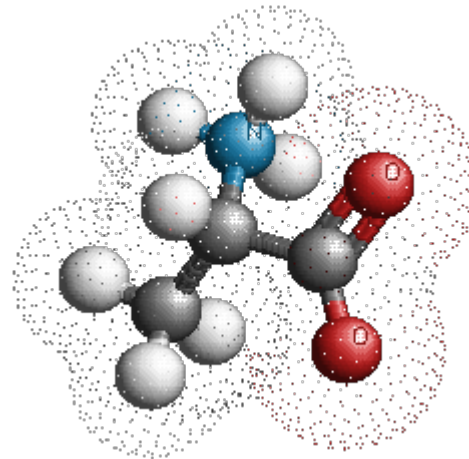


Carboxylic groups are present in (and characterize the chemical properties of) many important bio-molecules

palmitic acid (fatty acid)

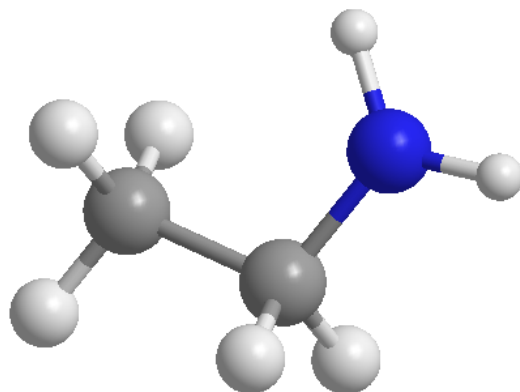


alanine (aminoacid)

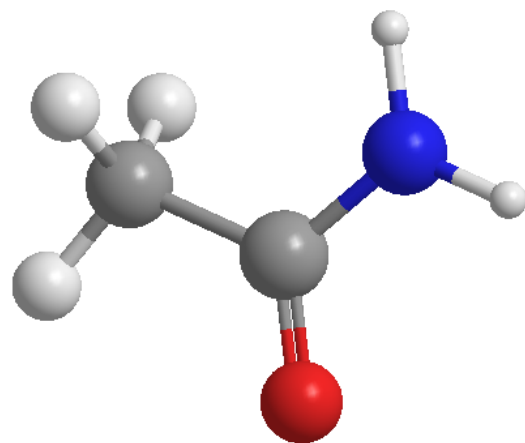


# Functional groups of nitrogen

## Amines

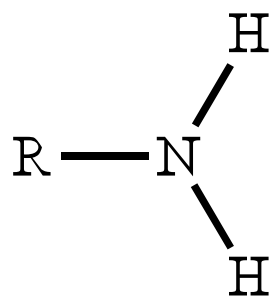


## Amides

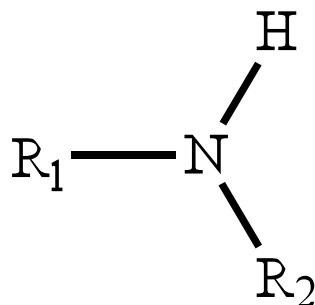


# Amines

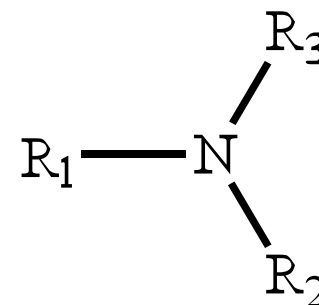
The amines having only one alkyl or aromatic group linked to nitrogen are called primary. The amines having the nitrogen atom bound to 2, 3 and 4 aromatic, aliphatic or alicyclic groups are termed secondary, tertiary and quaternary, respectively



primary



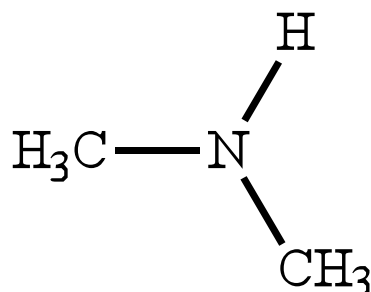
secondary



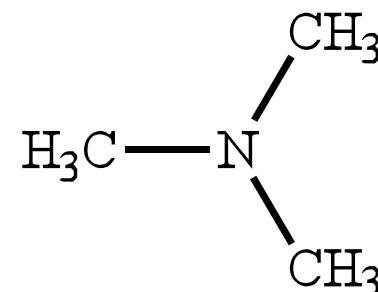
tertiary



methylamine

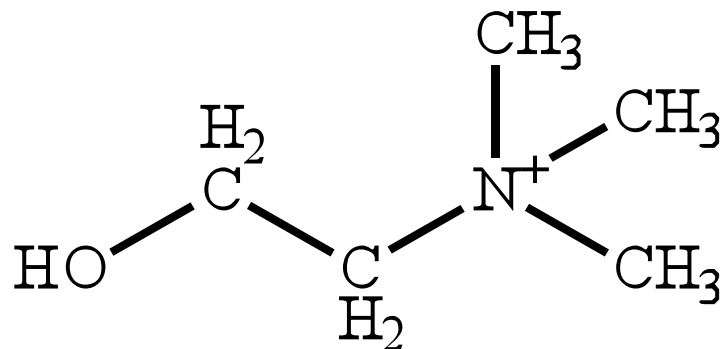


dimethylamine

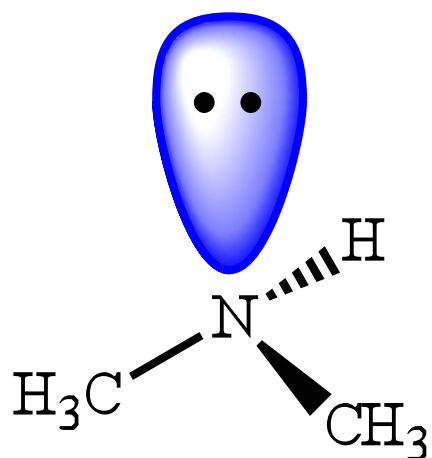


trimethylamine

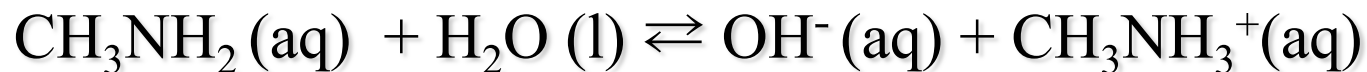
The **quaternary amines** contain an **ammonium ion** bound to four organic compounds. A very important quaternary amine for the physiology is the **choline** (trimethyl-ethanol-ammonium ion)



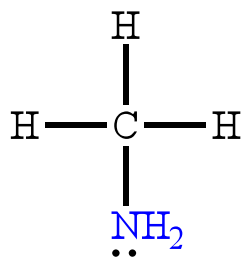
Amines are derivatives of ammonia with nitrogen having  $sp^3$  hybridization and tetrahedral geometry, as in ammonia.



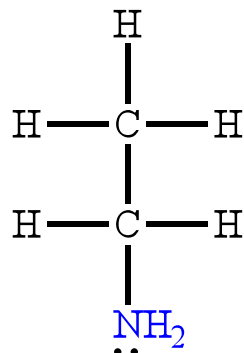
The lone pair in one of the 4 orbitals confers a basic character to the amines. All the amines are indeed weak bases with  $pK_B$  values between 4 and 6.



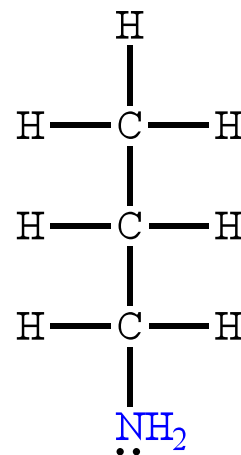
# Aliphatic amines



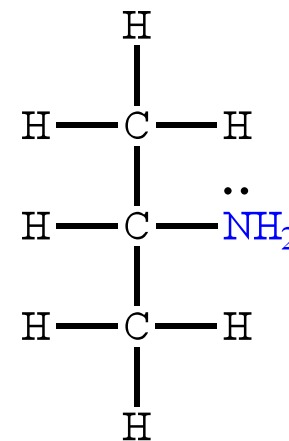
aminomethane  
methylamine



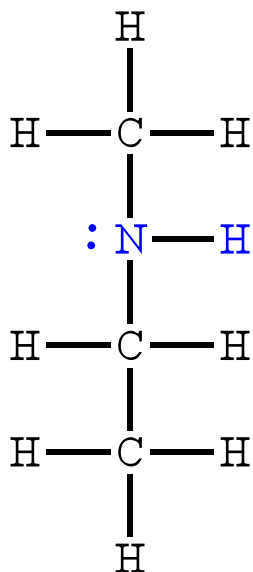
aminoethane  
etilamina



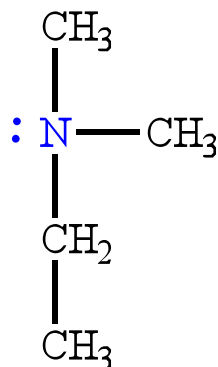
1-aminopropane  
1-propylamine



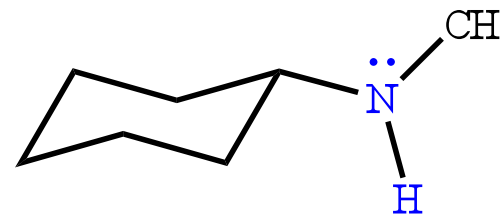
2-aminopropane  
2-propylamine



N-methyl-aminoethane  
methyl ethyl amine



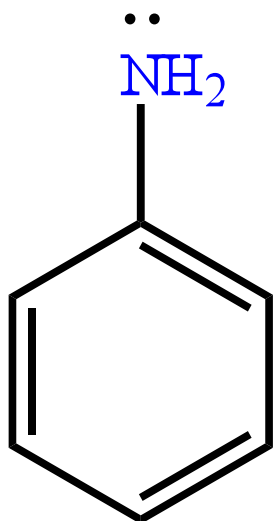
N,N-dimethyl-aminoethane  
dimethyl-ethyl-amine



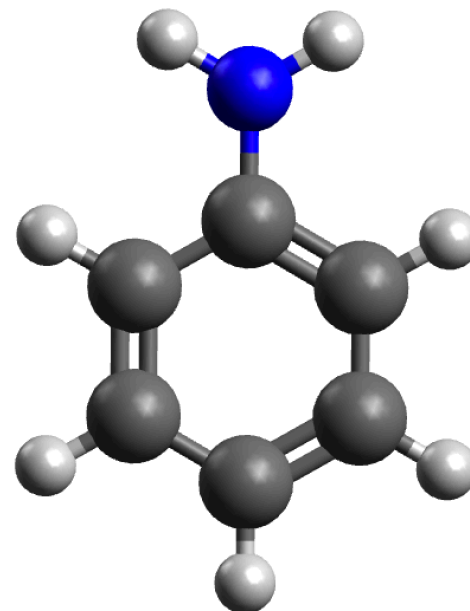
N-methyl-aminocyclohexane  
methyl-cyclohexyl-amine



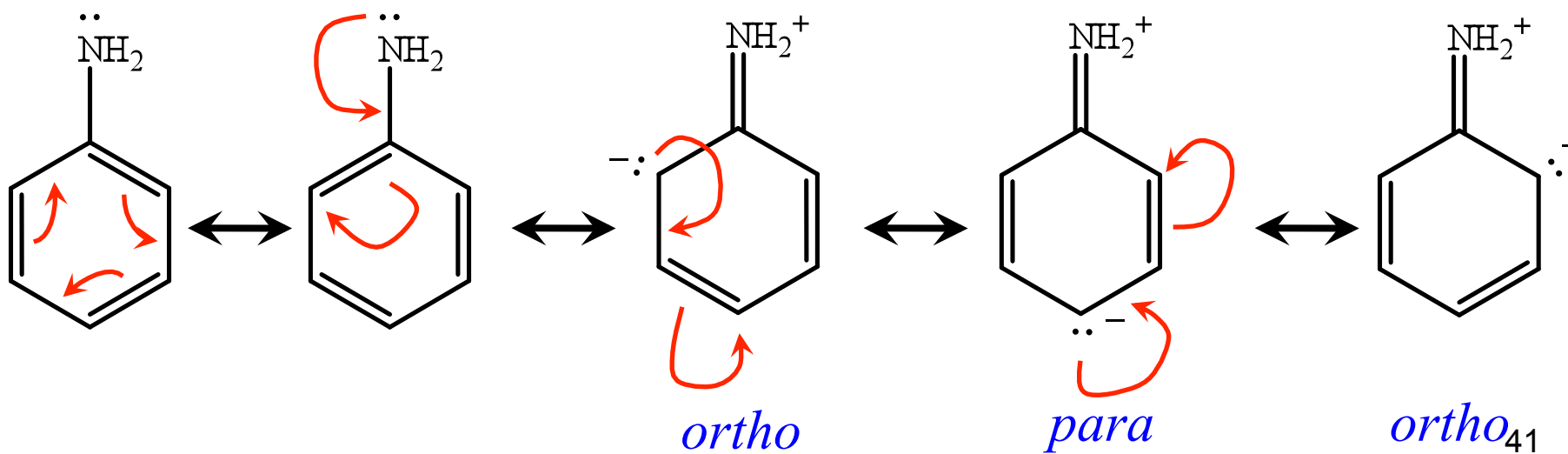
# Aromatic amines



benzylamine  
aminobenzene  
phenylamine  
aniline

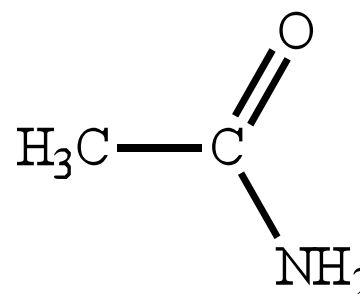
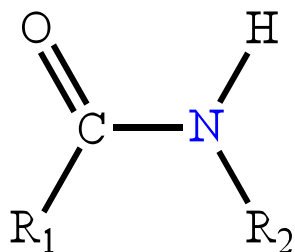
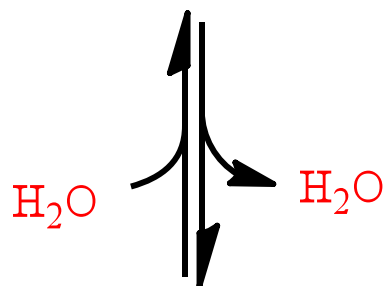
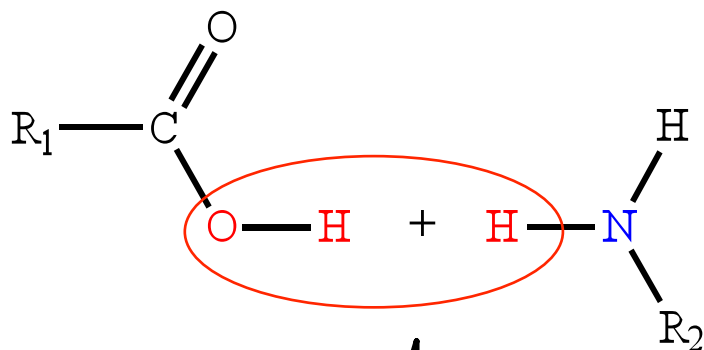
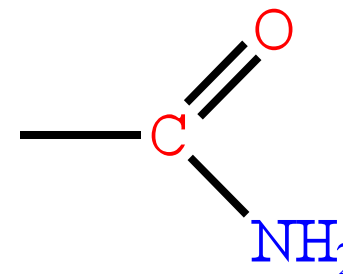


Aniline resonance  
limit structures

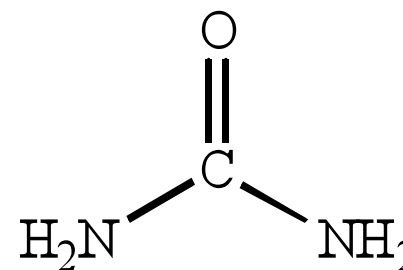


# Amides

The amide functional group has formula  $-\text{CONH}_2$  and is obtained by reaction of a carboxylic acid with an amine

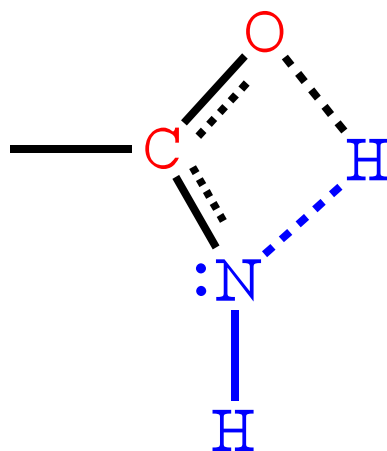
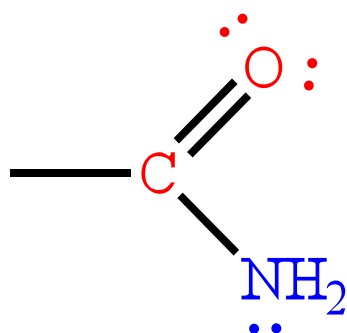


acetamide  
(acetic amide)

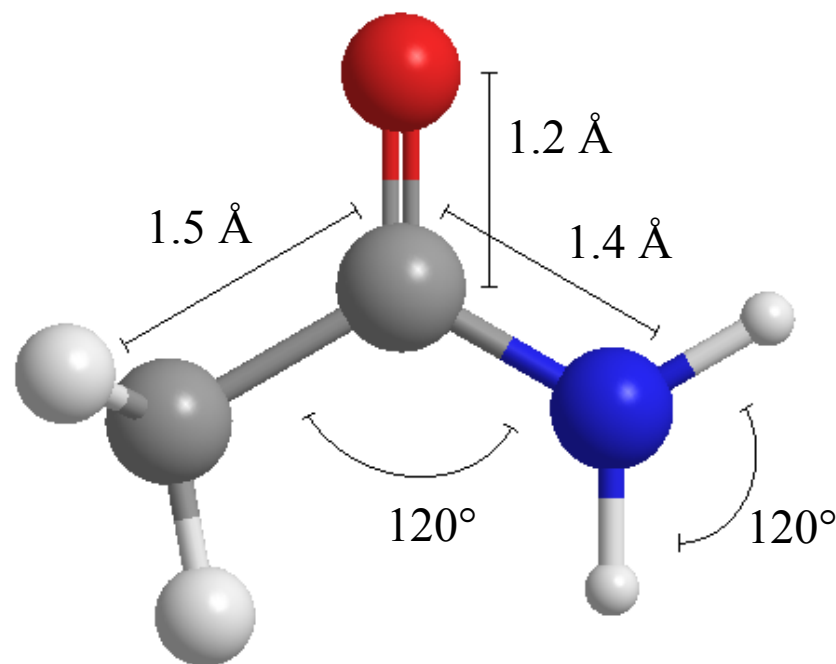


urea  
(a diamide of carbonic acid)

The structure of the amide group allows the delocalization of the  $\pi$  binding orbital on three atoms C, O and N and this imposes the  $sp^2$  geometry for the nitrogen in the delocalized O-C-N system: as a consequence **amides have no basic character**. To represent this structure, you can write the formula like this:

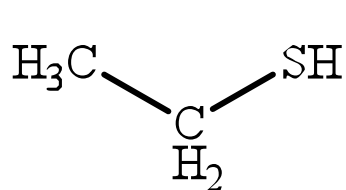


In this structure one of the nitrogen-linked hydrogen atoms is shared with the oxygen

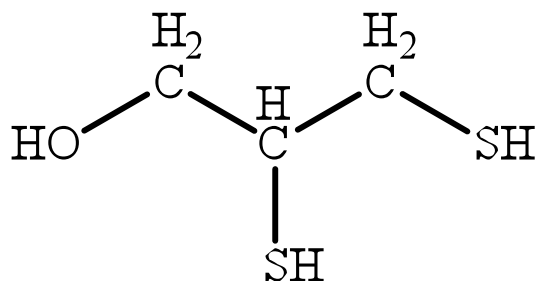


# Functional groups of sulfur

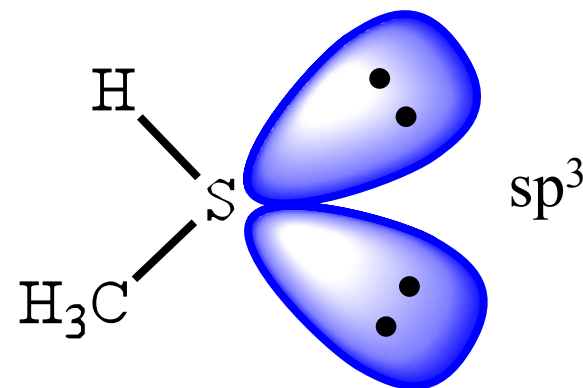
In practice the only important functional group of sulfur are **thioalcohols** or **mercaptans** (which resemble alcohols), which have the formula -SH



ethanethiol



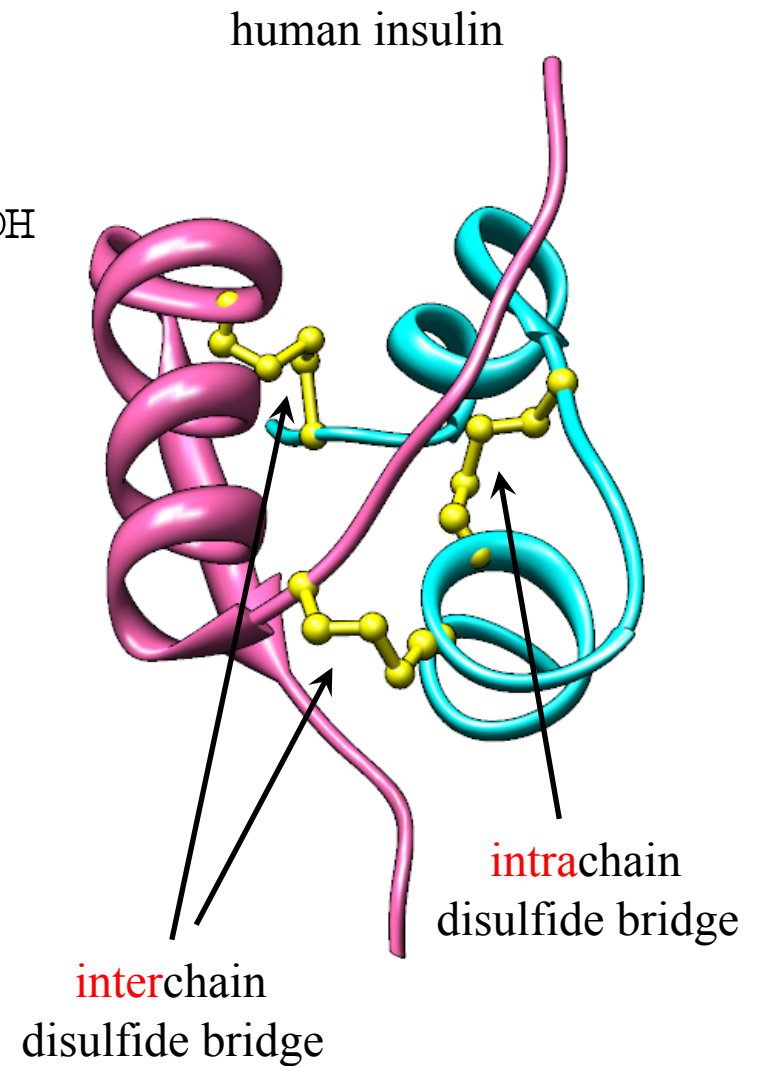
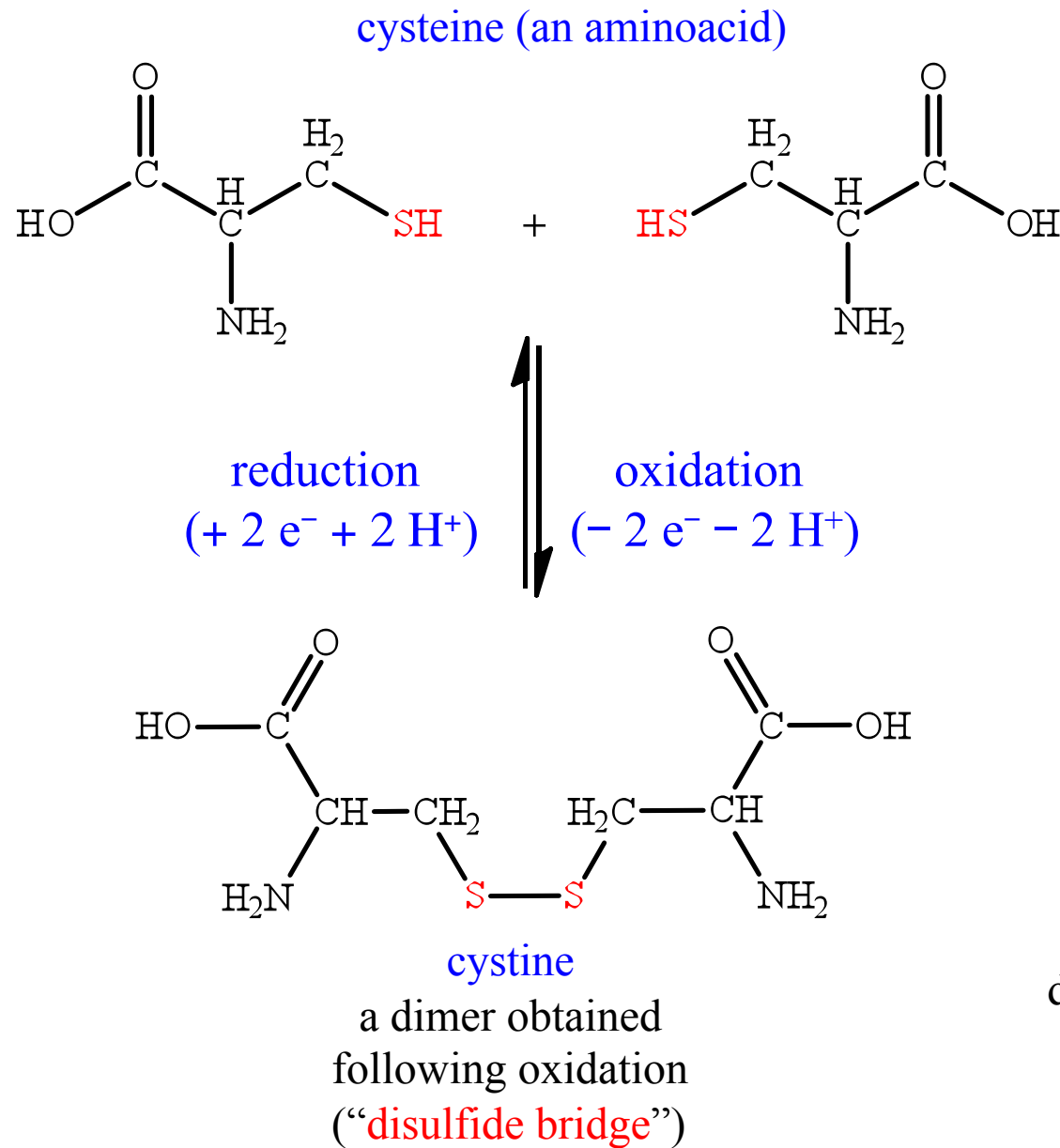
dimercaptopropanol



Sulfur is the homolog of oxygen and a thioalcohol has properties and geometry similar (not so really) to that of the corresponding alcohol, **but unlike alcohols it behaves like weak acid with pK<sub>A</sub> of about 9**



# Thiols: oxidation reactions



# The perm



## Aminoacid composition of some fibrous proteins (mol %)

Aminoacid	$\alpha$ -keratin (wool)	fibroin (silk)	collagen (bovine tendon)	Elastine (pig aorta)
Gly	8.1	44.6	32.7	32.3
Ala	5.0	29.4	12.0	23.0
Ser	10.2	12.2	3.4	1.3
Glu+Gln	12.1	1.0	7.7	2.1
Cys	11.2	0	0	0
Pro	7.5	0.3	22.1 <sup>a</sup>	10.7 <sup>c</sup>
Arg	7.2	0.5	5.0	0.6
Leu	6.9	0.5	2.1	5.1
Thr	6.5	0.9	1.6	1.6
Asp+Asn	6.0	1.3	4.5	0.9
Val	5.1	2.2	1.8	12.1
Tyr	4.2	5.2	0.4	1.7
Ile	2.8	0.7	0.9	1.9
Phe	2.5	0.5	1.2	3.2
Lys	2.3	0.3	3.7 <sup>b</sup>	3.6 <sup>d</sup>
Trp	1.2	0.2	0	0
His	0.7	0.2	0.3	0
Met	0.5	0	0.7	0

<sup>a</sup> ~ 39 % = hyd-Pro; <sup>b</sup> ~ 14 % = hyd-Lys; <sup>c</sup> ~ 13 % = hyd-Pro; <sup>d</sup> ~ 80 % = involved in crosslinks

# Keratin

most durable and less reactive protein of vertebrates.

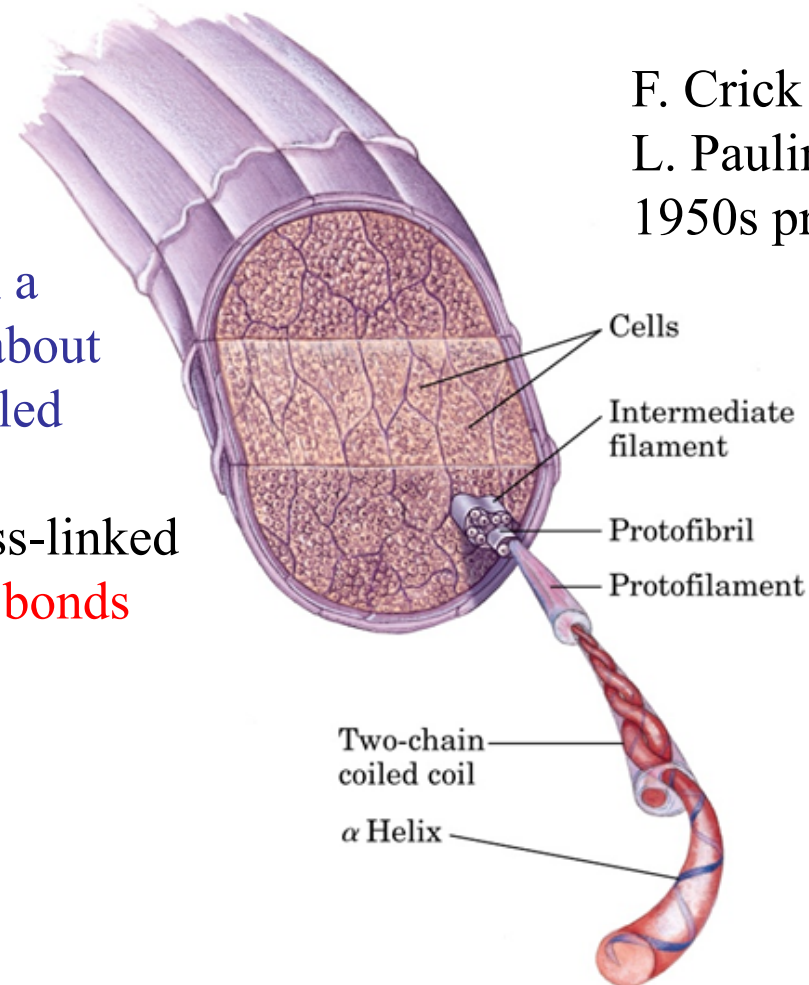
- mammals:  $\alpha$ -keratins
- birds & reptiles:  $\beta$ -keratins

Cross-section of a hair

F. Crick and  
L. Pauling  
1950s prediction

Keratin filaments are found in hair. In a single wool fiber with a diameter of about 20  $\mu\text{m}$ , millions of filaments are bundled together within dead cells.

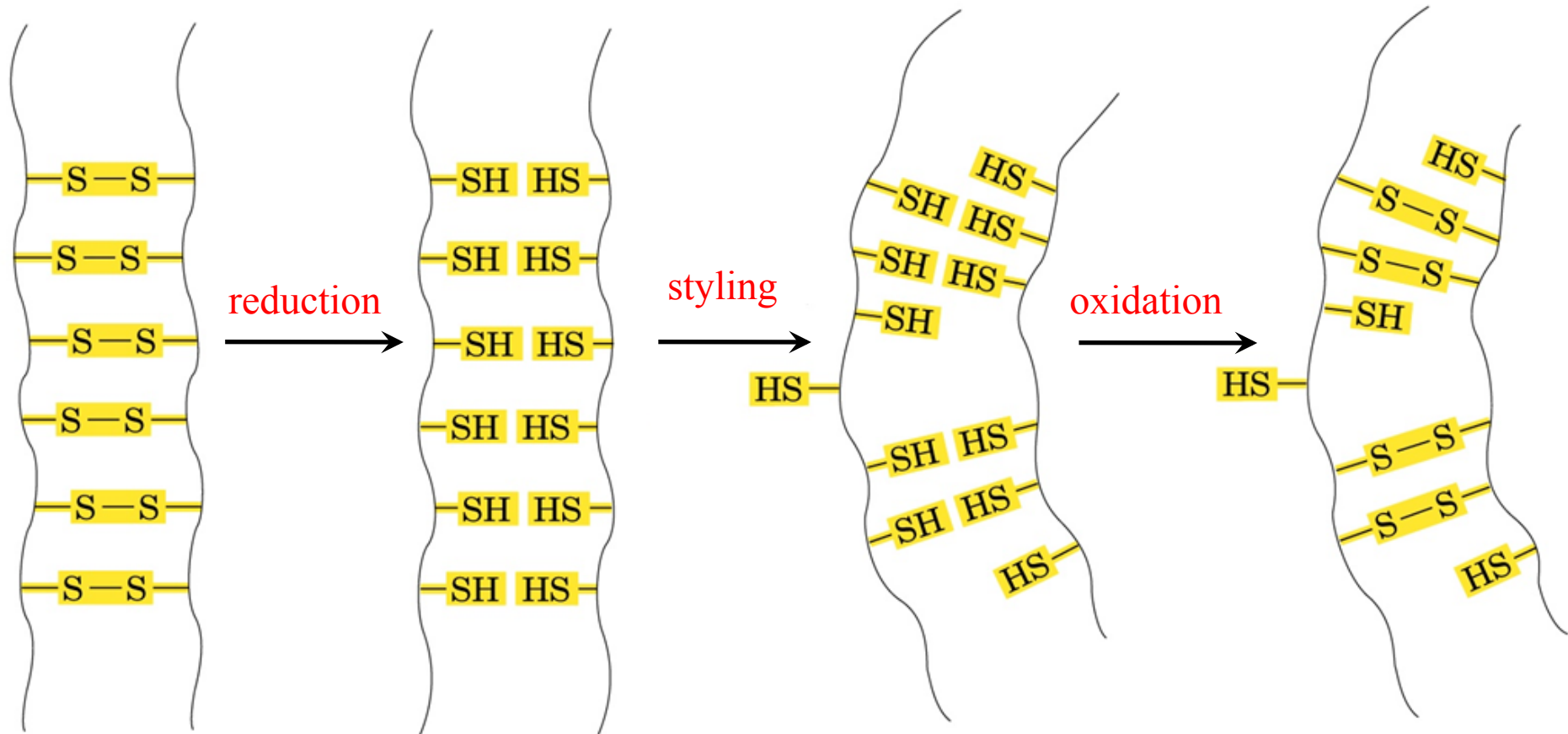
The individual keratin helices are cross-linked and stabilized by numerous **disulfide bonds**





The perm is a biochemical engineering operation

keratin strands



## Functional groups deriving from the combination of other groups

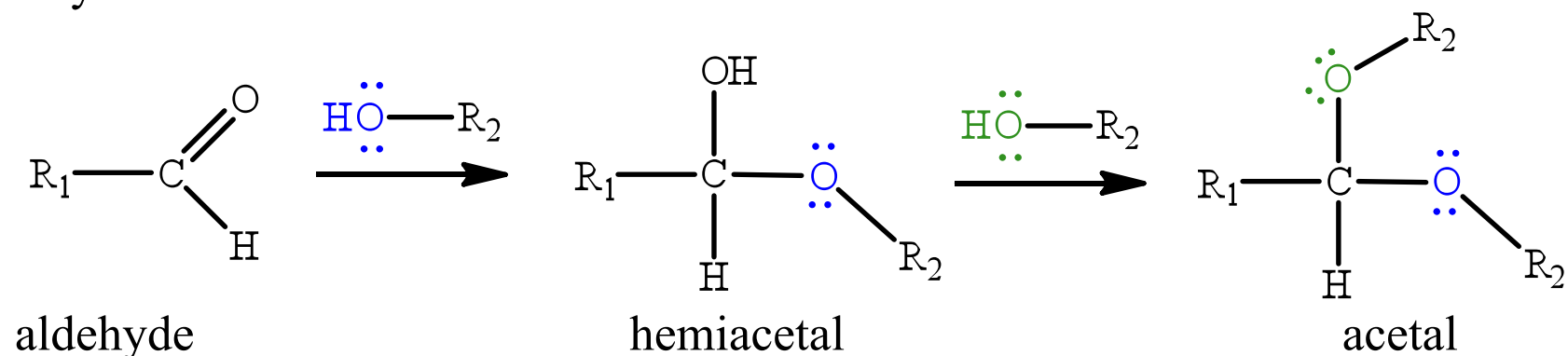
The functional groups studied up to this point, account for more or less the most frequent compounds, but do not consider **those compounds that derive from the reactions between groups**.

The most reactive groups are those that contain oxygen

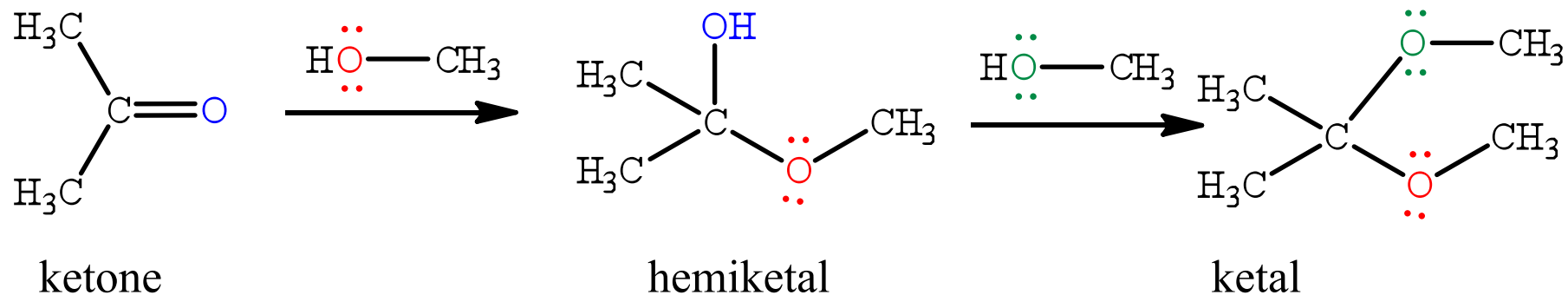
- anhydrides
- acetals and ketals
- esters

# Acetals and ketals

The **hemiacetals** and **acetals** derive from the reaction of **one** or **two** alcohols with an aldehyde

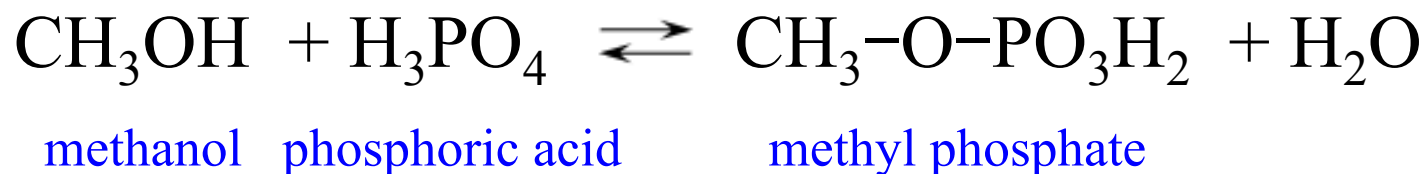
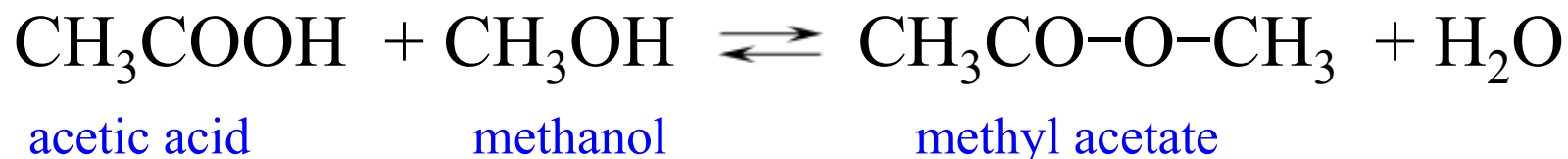
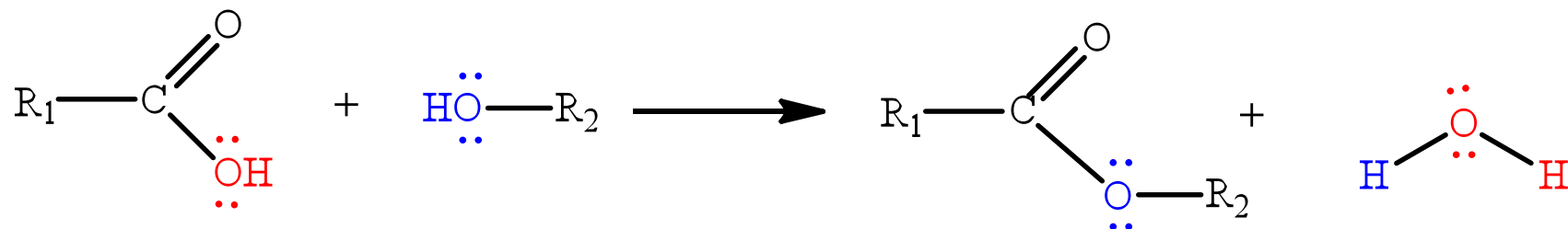


The **hemiketals** and **ketals** derive from the reaction of **one** or **two** alcohols with a ketone

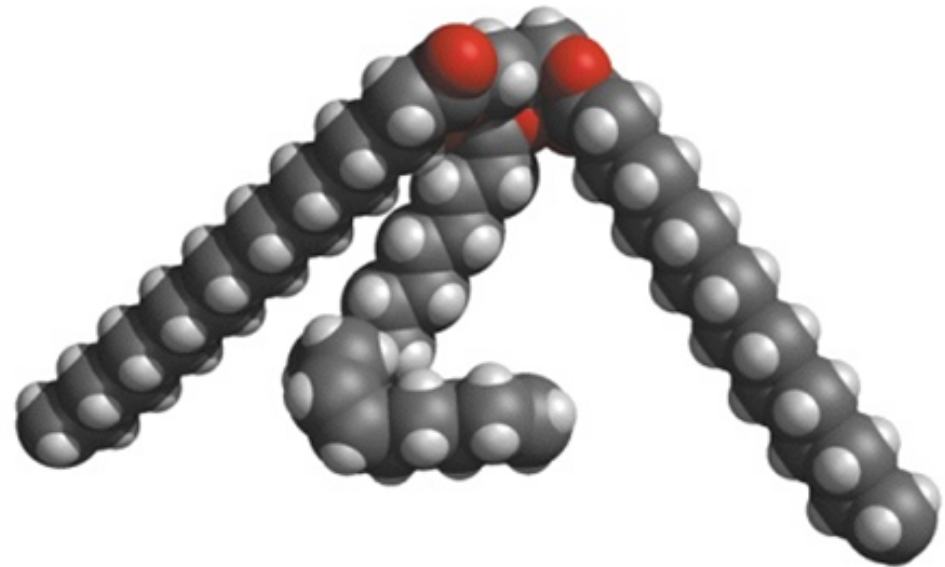
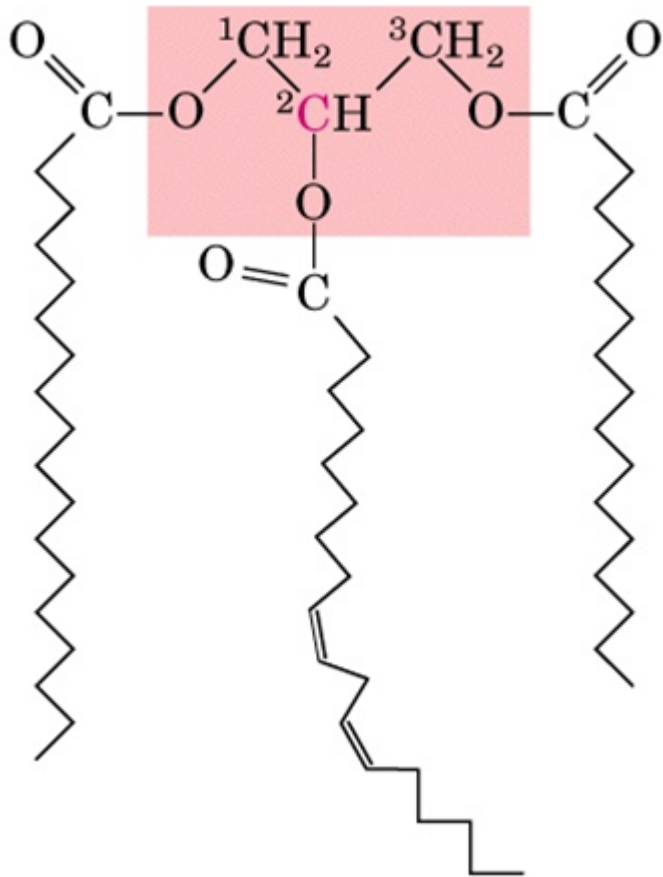
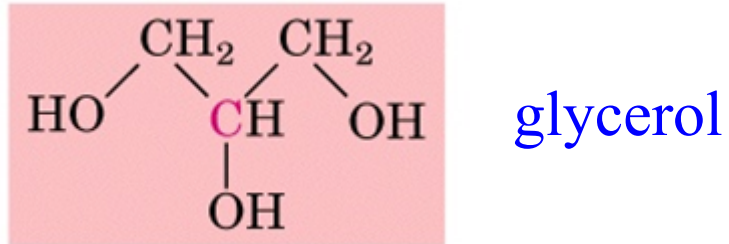


# Esters

The **ester** is the reaction product of an **alcohol** with an **acid**, with removal of a molecule of water



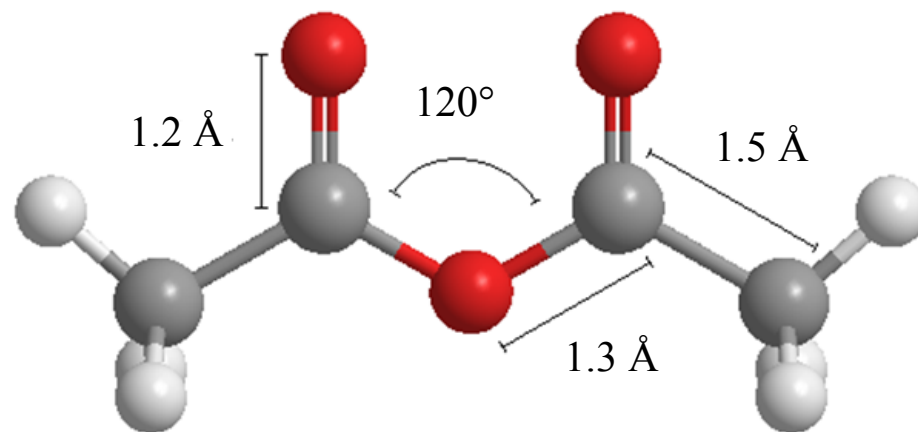
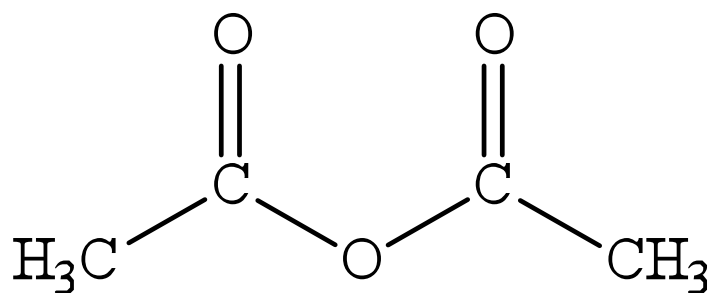
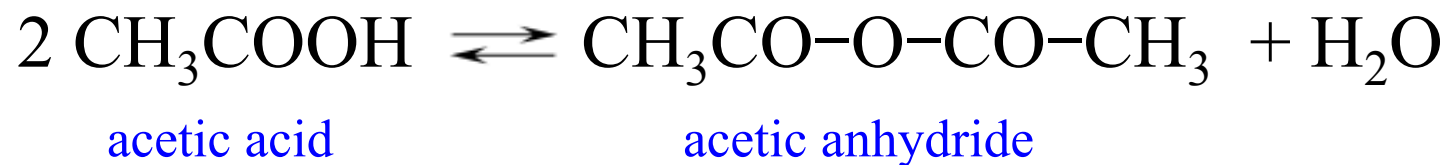
Triglycerides: glycerol esters with three fatty acids (lipid).



1-stearoyl-2-lineoyl-3-palmitoyl-glycerol

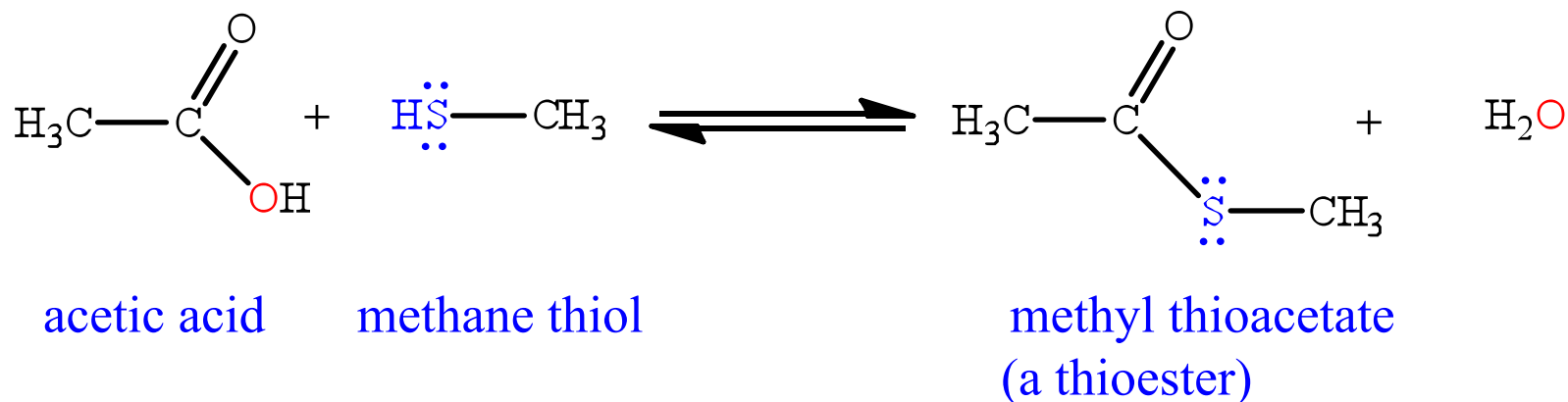
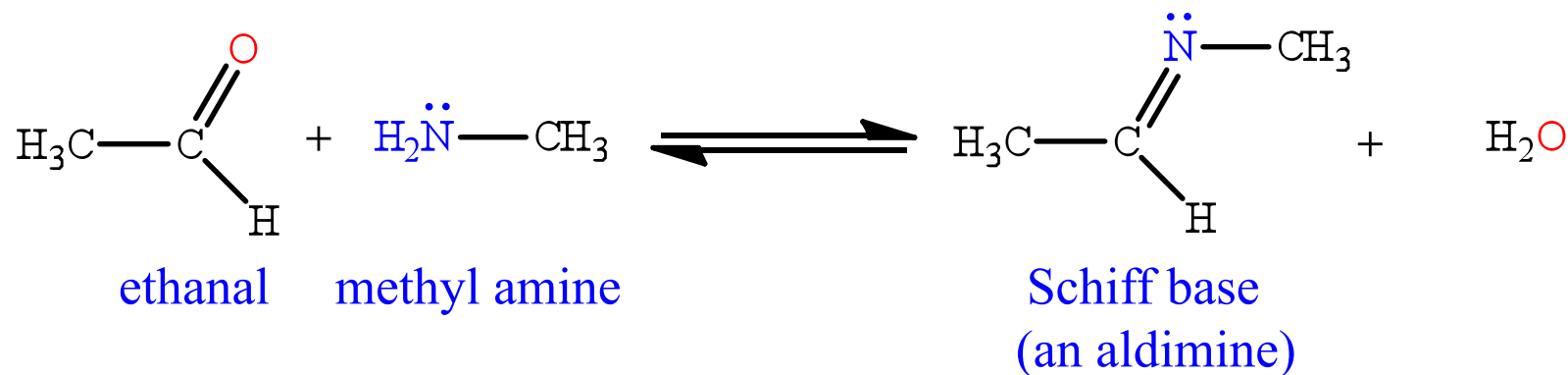
# Anhydrides

an **anhydride** is the reaction product between **two acids**, with the elimination of a water molecule

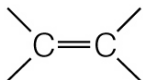
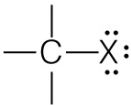
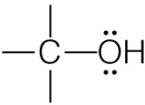
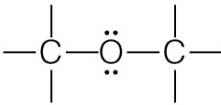
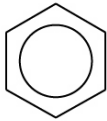


## Other important reactions and compounds

Among the characteristic reactions of functional groups, the reactions between **aldehydes** and **amines** and those between **carboxylic acids** and **thioalcohols** are quite important



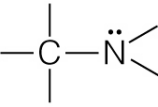
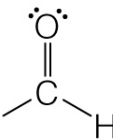
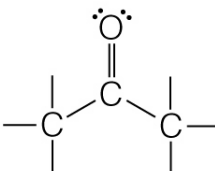
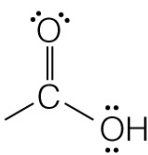
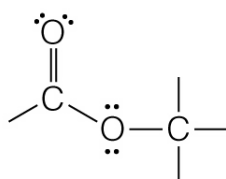
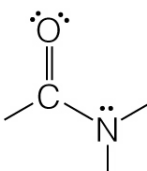
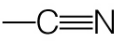


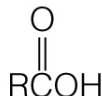

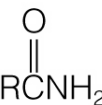
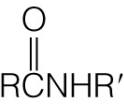
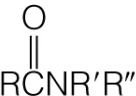

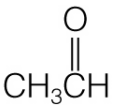
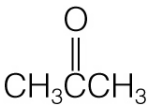
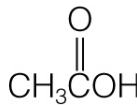
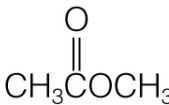
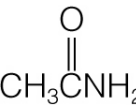
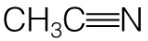
# Summary of Important Families of Organic Compounds

	Family						
	Alkane	Alkene	Alkyne	Aromatic	Haloalkane	Alcohol	Ether
Functional group	C—H and C—C bonds		$\text{—C}\equiv\text{C—}$	Aromatic ring			
General formula	RH	RCH=CH <sub>2</sub> RCH=CHR R <sub>2</sub> C=CHR R <sub>2</sub> C=CR <sub>2</sub>	RC≡CH RC≡CR	ArH	RX	ROH	ROR
Specific example	CH <sub>3</sub> CH <sub>3</sub>	CH <sub>2</sub> =CH <sub>2</sub>	HC≡CH		CH <sub>3</sub> CH <sub>2</sub> Cl	CH <sub>3</sub> CH <sub>2</sub> OH	CH <sub>3</sub> OCH <sub>3</sub>
IUPAC name	Ethane	Ethene	Ethyne	Benzene	Chloroethane	Ethanol	Methoxymethane
Common name <sup>a</sup>	Ethane	Ethylene	Acetylene	Benzene	Ethyl chloride	Ethyl alcohol	Dimethyl ether

<sup>a</sup>These names are also accepted by the IUPAC.

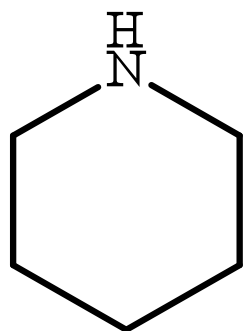


## Summary (cont.)

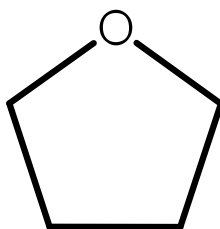
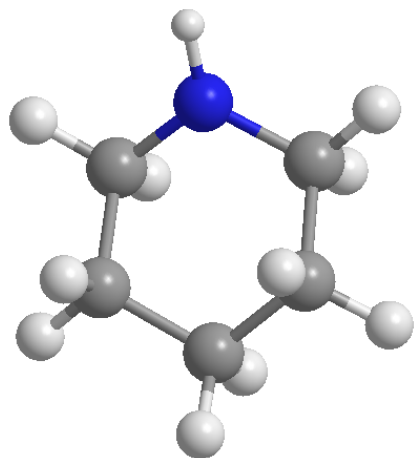
Family						
Amine	Aldehyde	Ketone	Carboxylic Acid	Ester	Amide	Nitrile
						
RNH <sub>2</sub> R <sub>2</sub> NH R <sub>3</sub> N					  	
CH <sub>3</sub> NH <sub>2</sub>						
Methanamine	Ethanal	Propanone	Ethanoic acid	Methyl ethanoate	Ethanamide	Ethanenitrile
Methylamine	Acetaldehyde	Acetone	Acetic acid	Methyl acetate	Acetamide	Acetonitrile

# Heterocyclic compounds

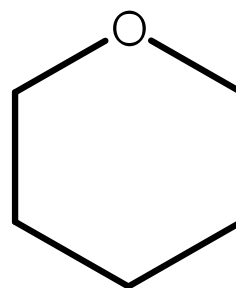
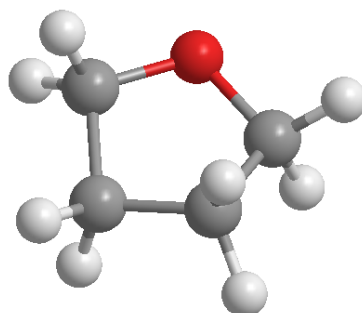
Heterocycles are cyclic, aromatic or aliphatic compounds, containing atoms other than carbon. Among the important aliphatic heterocyclics:



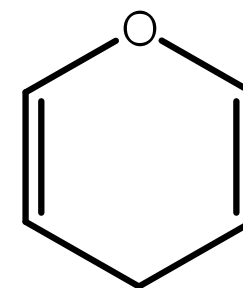
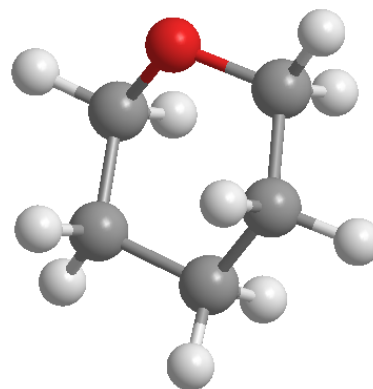
piperidine



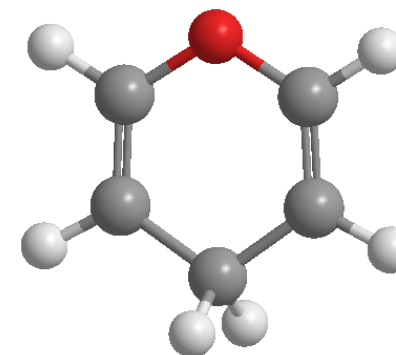
tetrahydrofuran



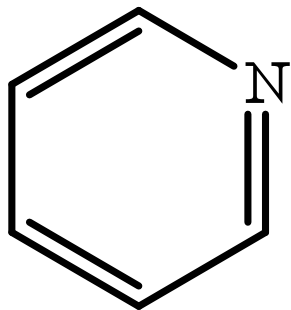
tetrahydropyran



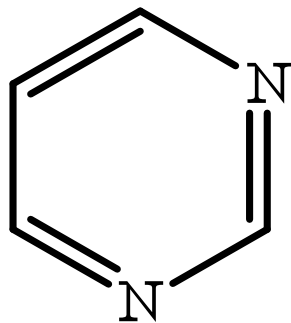
pyran



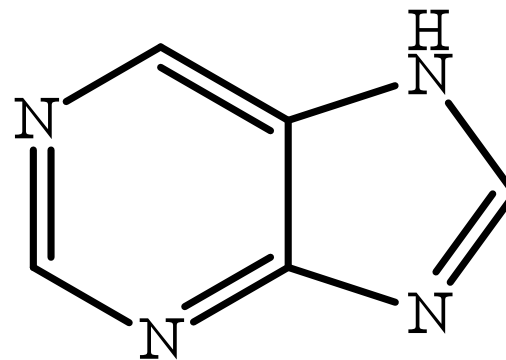
# Important aromatic heterocycles :



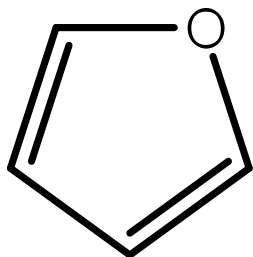
pyridine



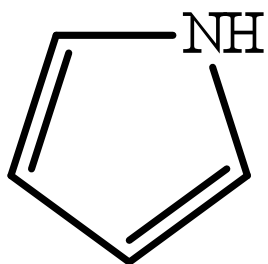
pyrimidine



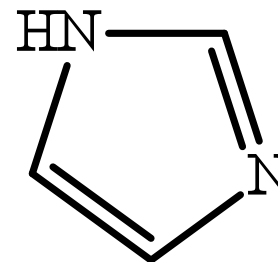
purine



furan



pyrrole



imidazole