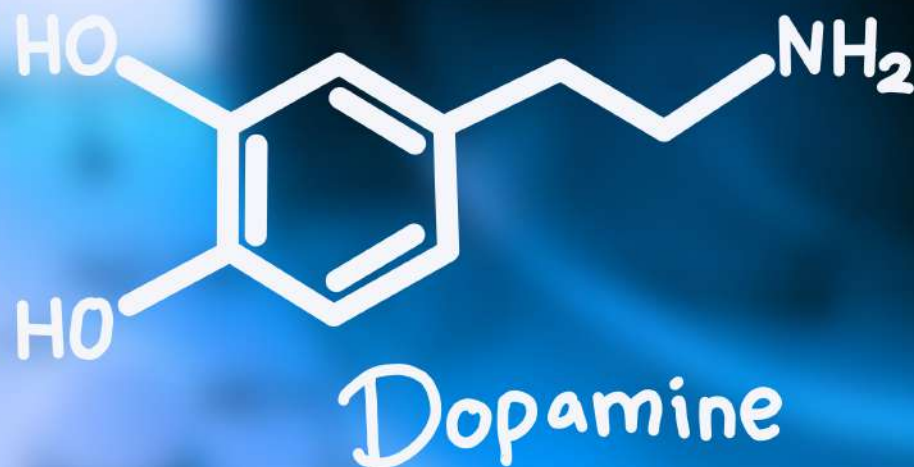


Organic 1



Subject: Alcohols,
Phenols and Ethers



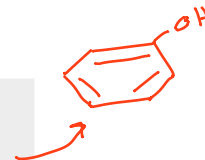
Pharmaceutical Organic Chemistry-1

Chapter-4: Alcohols, Phenols and Ethers

Alcohols, Phenols and Ethers

- Alcohols, ethers and phenols have a common functional group, the **hydroxyl group, -OH**.

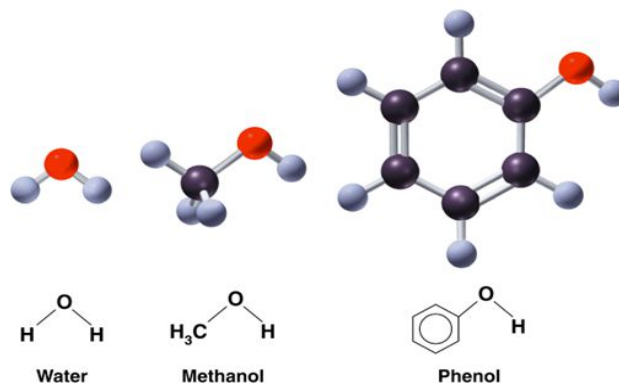
H-O-H Water	R-OH Alcohol	R-O-R Ethers	Ph-O-H Phenol
-----------------------	------------------------	------------------------	-------------------------



- Alcohols** are compounds whose molecules have a **hydroxyl group** attached to a **saturated** carbon atom. (لازم تكون الرابطة بين OH و R أحادية)
- Phenols** are compounds that have a **hydroxyl group** attached directly to a **benzene ring**.
- Ethers** are compounds whose molecules have an **oxygen** atom bonded to **two** carbon atom.

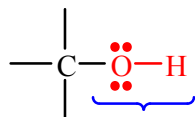
Alcohols and Phenols

- o **Alcohols and phenols** may be viewed as organic *derivatives of water*. (رنا ان الڪحول و الفينول اصلهم من الماء)



- o **Alcohols** have the general formula $R-OH$, and structurally similar to water, but with one of the hydrogens replaced by an alkyl group.
- o **Phenols** have a hydroxyl group attached directly to an aromatic ring.

Alcohols



This is the functional group of an alcohol

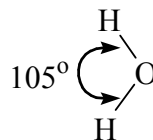
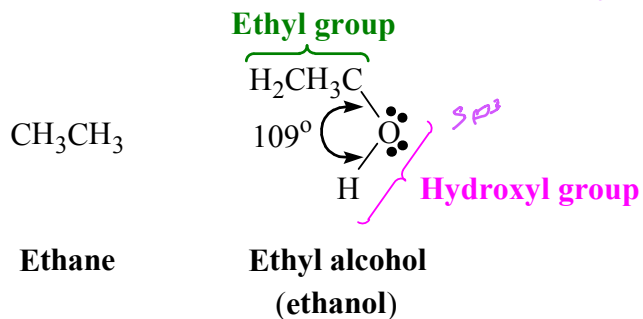
o Alcohols can be viewed in two ways structurally:

(1) as **hydroxyl derivatives** of alkanes

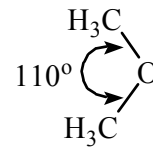
لما ابدل ذرة هيدروجين في الألكان بمجموعة OH يحصل على كحول

and (2) as **alkyl derivatives** of water.

لما ابدل ذرة H في الماء بمجموعة alkyl يحصل على كحول



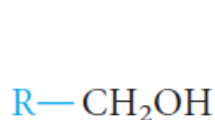
Water



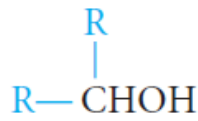
Dimethyl ether
(a typical ether)

Classification of Alcohols

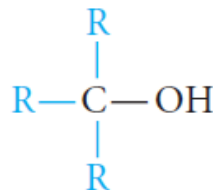
- o **Alcohols** are classified as **primary** (1°), **secondary** (2°), or **tertiary** (3°), depending on whether one, two, or three organic groups are connected to the hydroxyl-bearing carbon atom.



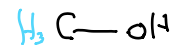
primary (1°)



secondary (2°)



tertiary (3°)

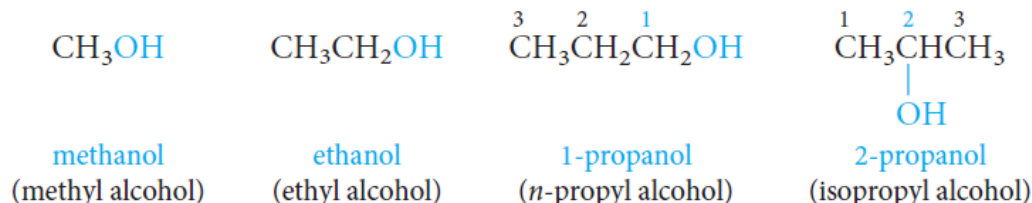


methyl Alcohol

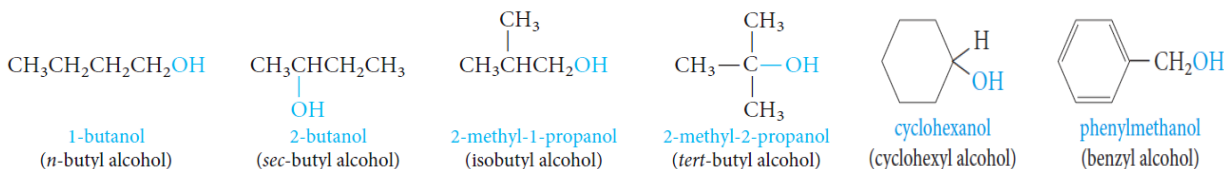
- o **Methyl alcohol**, which is not strictly covered by this classification, is usually grouped with the **primary** alcohols.

Nomenclature of Alcohols

- o The **common names** for the simplest alcohols consist of **alkyl group** attached to the hydroxyl function followed by the word alcohol: *Alkyl alcohol*.
- o In the **IUPAC system**, alcohols are named according to the following rules.
 1. Select the **longest continuous carbon chain** that *contains the -OH group*.
Drop the *-e* ending of the parent alkane and replace it by the suffix *-ol*: *Alkanol*
 2. When isomers are possible, the chain is numbered so as to give the functional group (-OH) the *lowest possible number*.

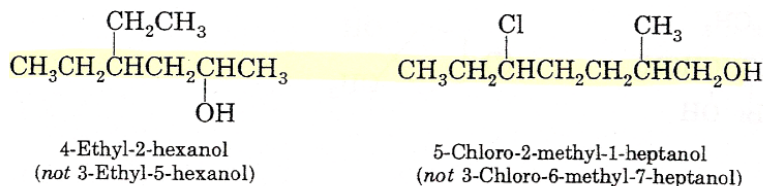


Nomenclature of Alcohols



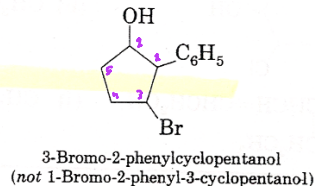
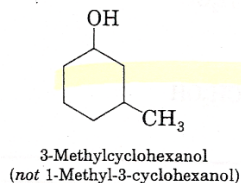
3. When alkyl side chains or other groups are present, they are named alphabetically and their positions are indicated by a number.

*The position of the functional group (-OH) is always given the **lowest possible number** at the end of the name.*



Nomenclature of Alcohols

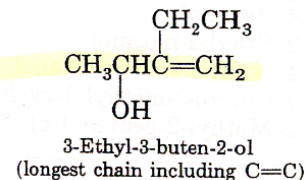
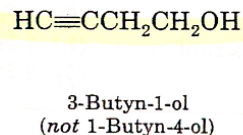
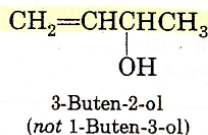
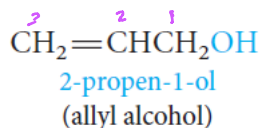
For cyclic alcohols, numbering always starts from the carbon bearing the -OH group.



مع باصلة يكون دائماً
ترقيم OH (1)

4. With Unsaturated Alcohols; If a molecule contains both an -OH group and a C=C or C-C triple bond, the -OH group takes **preference** before the double or triple bonds in getting the lower number.

The name should include (if possible) both the hydroxyl and the unsaturated groups, even if this does not make the longest chain the parent hydrocarbon.



← الأولوية بالترقيم اعطي OH اقل رقم →

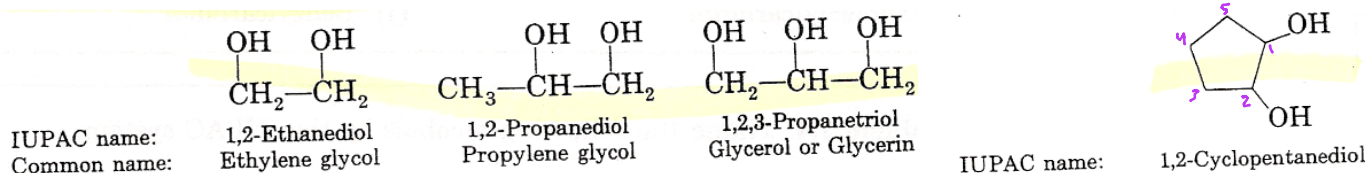
Nomenclature of Alcohols

o Alcohols with More Than One Hydroxyl Group

- Compounds with two adjacent alcohol groups are called **glycols**.

The most important example is ethylene glycol. Common name

- Compounds with more than two hydroxyl groups are also known, and several, such as glycerol and sorbitol, are important commercial chemicals.



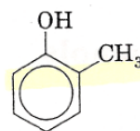
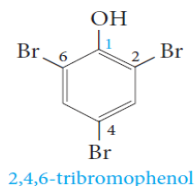
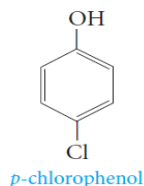
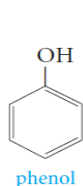
- **Ethylene glycol** is used as the “permanent” antifreeze in automobile radiators and as a raw material in the manufacture of Dacron. (بوليمستر)
- **Ethylene glycol** is completely miscible with water.
- **Glycerol** is a syrupy, colorless, water-soluble, high-boiling liquid with a distinctly sweet taste. Its soothing qualities make it useful in shaving and toilet soaps and in cough drops and syrups. العلاقة

سائل كثافة عالية نوع الصمغ

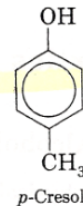
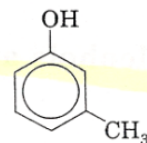
Nomenclature of Phenols

- o **Phenols** are usually named as derivatives of the parent compounds.

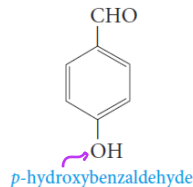
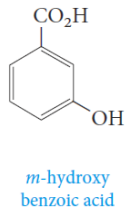
الترقيم لا يكون
(3) وجعدين الأولوية
للسار الي بعضي
الفرقات اقل ترقيم



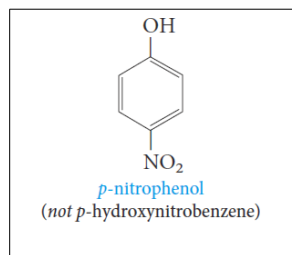
عبارة عن CH₃
و OH مع بعض



- o The hydroxyl group is named as a substituent when it occurs in the same molecule with carboxylic acid, aldehyde, or ketone functionalities, which have **priority in naming**.



but



ترتيب الأولويات :



Physical Properties of

Alcohols

Physical

State The simplest alcohol, methanol, is a **liquid** at room temperature.

In contrast, alkanes from methane to butane (C1-C4) are **gases**.

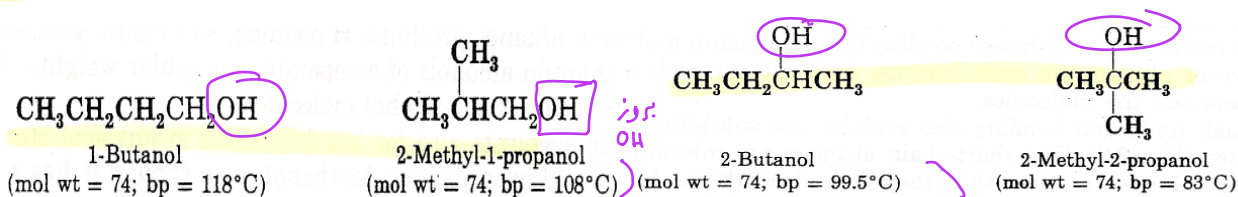
Solubilit

- The **lower alcohols** are completely miscible with water. ذائب
- As the number of **carbons in the alcohol increases**, the **solubility in water decreases**.

Boiling

Points Series of normal alcohols; The boiling points increase with increase in molecular weights.

- A comparison of boiling points among isomeric alcohols; The boiling points **decrease** as the number of alkyl branches from the **carbinol group increases**.



↓
 ار OH في هذا الموضع
 أكثر بروزا
 ↓
 أعلى bp

عند تساوي
 عدد C
 بين المركبات

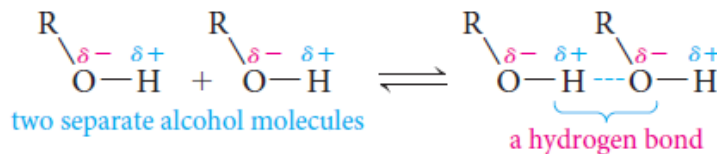
Hydrogen Bonding in

- o The **boiling points** (bp,s) of **Alcohols** much higher than those of **ethers** or **hydrocarbons** with similar molecular weights.

	<chem>CH3CH2OH</chem>	<chem>CH3OCH3</chem>	<chem>CH3CH2CH3</chem>
mol wt	46	46	44
bp	+78.5°C	-24°C	-42°C

Why? Because alcohols form hydrogen bonds with one another.

The O-H bond is polarized by the high electronegativity of the oxygen atom and places a partial positive charge on the hydrogen atom and a partial negative charge on the oxygen atom.



Two or more alcohol molecules thus become loosely bonded to one another through hydrogen bonds.

کمازاد عدد روابط ه
زاد pb وزادت solubility الکن > ایشتر > کحول

Hydrogen Bonding in Alcohols

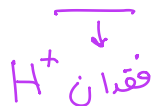
- o Consequently, alcohols have relatively **high boiling points** because they must supply enough heat to break the hydrogen bonds before each molecule.
- o **Hydrogen bonds** are weaker than ordinary covalent bonds.
- o **Water**, of course, is also a hydrogen-bonded liquid.
- o The **lower molecular-weight alcohols** can readily replace water molecules in the hydrogen bonded network.
- o This accounts for the complete **miscibility of the lower alcohols with water**.
- o However, as the organic **chain lengthens** and the alcohol becomes **relatively more hydrocarbon** like, its **water solubility decreases**.

Table 7.1 Boiling Point and Water Solubility of Some Alcohols

Name	Formula	bp, °C	Solubility in H ₂ O g/100 g at 20°C
methanol	CH ₃ OH	65	completely miscible
ethanol	CH ₃ CH ₂ OH	78.5	completely miscible
1-propanol	CH ₃ CH ₂ CH ₂ OH	97	completely miscible
1-butanol	CH ₃ CH ₂ CH ₂ CH ₂ OH	117.7	7.9
1-pentanol	CH ₃ CH ₂ CH ₂ CH ₂ CH ₂ OH	137.9	2.7
1-hexanol	CH ₃ CH ₂ CH ₂ CH ₂ CH ₂ CH ₂ OH	155.8	0.59

Physical Properties of Phenols

- o **Phenol** is a colorless, crystalline, low-melting solid, with a high boiling point, that is **moderately** soluble in water.
- o Most other phenols also **are solids**, with **slight solubility** in water and high boiling points.
- o The most significant physical property that **distinguishes alcohols from phenols** is the **acidity of phenols**.



يعبر عنها بواسطة
 K_a / pK_a

↓
كما كانت
إعادة قدرة على
منح H^+ على
كانت acidity
أعلى و K_a أعلى

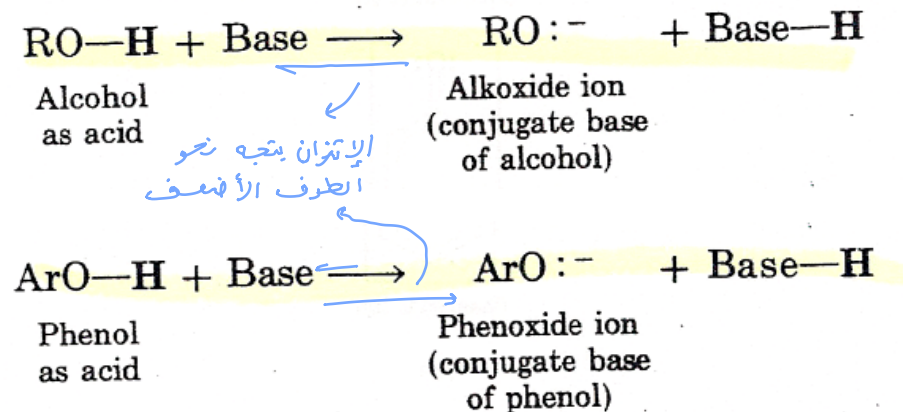
تفكك

↓ أعلى dissociation و pK_a أقل
كما كان conjugate base أكثر استقرارا

The Acidity of Alcohols and Phenols

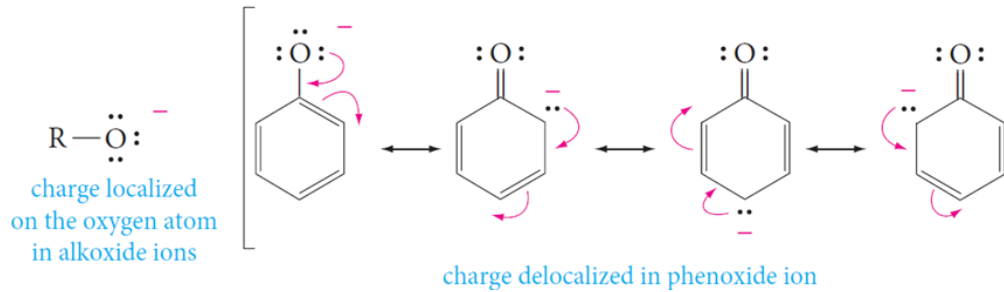
- o **Like water, alcohols and phenols are weak acids.**

The hydroxyl group can act as a proton donor, and dissociation occurs in a manner similar to that for water



The Acidity of Alcohols and Phenols

- o **Phenols are stronger acids than alcohols** mainly because the corresponding **phenoxide ions** are stabilized by resonance.



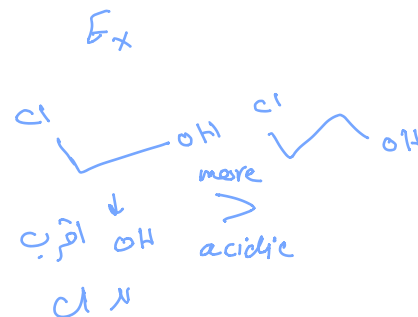
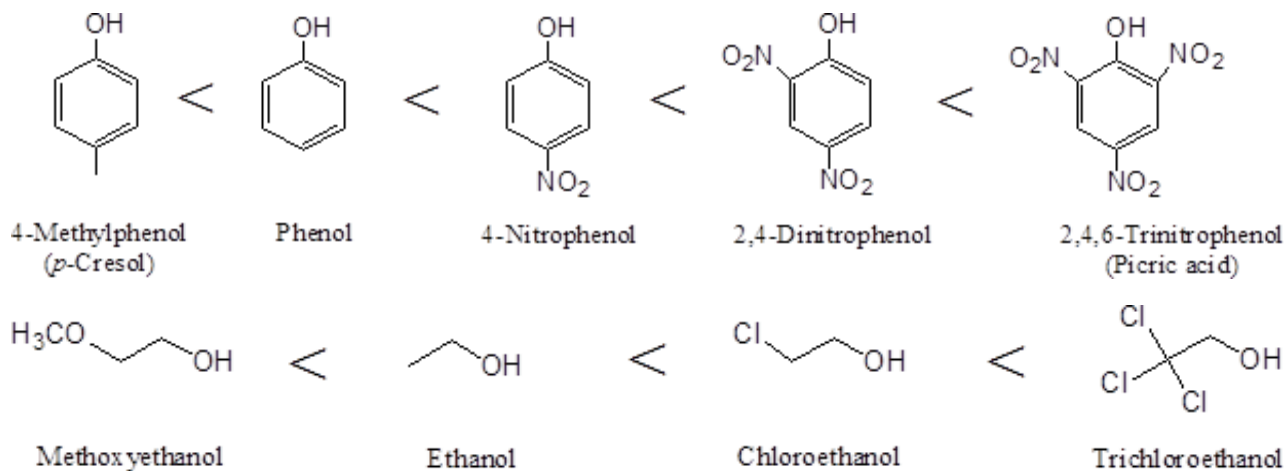
The negative charge of an **alkoxide ion** is concentrated on the oxygen atom, but the negative charge on a phenoxide ion can be delocalized to the ortho and para ring positions through resonance.

Because **phenoxide ions** are stabilized in this way, the equilibrium for their formation is more favorable than that for alkoxide ions

يوجد Resonance بالphenols لكي
 لا يكون باركول ← Resonance لكي كانت
 Stability للقاعدة اعرفقة أكبر .
 1. Resonance اعرفقة من :
 2. Electron-withdrawing group (EWG)
 3. Electron-donating group (EDG)
 اقل pK_a → اقل K_a → اقل acidity → اقل stability → اقل acidity
 اقل pK_a → اقل K_a → اقل acidity → اقل stability → اقل acidity

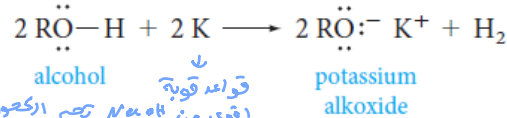
The Acidity of Alcohols and Phenols

- All **electron-withdrawing groups increase acidity** by stabilizing the conjugate base. **Electron-donating groups decrease acidity** because they **destabilize** the conjugate base.



The Acidity of Alcohols and Phenols

- o **Alkoxides**, the conjugate bases of alcohols, can be prepared by the reaction of an alcohol with sodium or potassium metal.

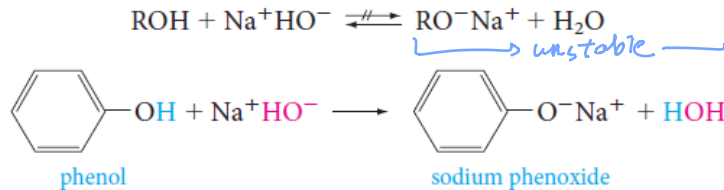


- o Treatment of alcohols with sodium hydroxide does not convert them to their alkoxides.

This is because alkoxides are stronger bases than hydroxide ion, so the reaction goes in the reverse direction.

Since alcohols are weaker acids than water, it is not possible to form the salt of an alcohol in aqueous alkaline solutions.

- o Treatment of phenols with sodium hydroxide converts them to phenoxide ions.



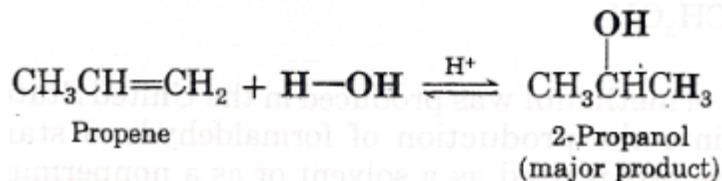
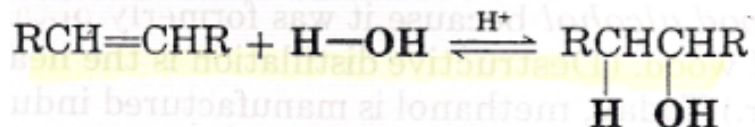
لذلك لا
يجد هذا
التفاعل لأن
القواعد أقوى من
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القواعد أقوى من
الماء

Preparation of Alcohols

o *From*

A. Hydration of Alkenes

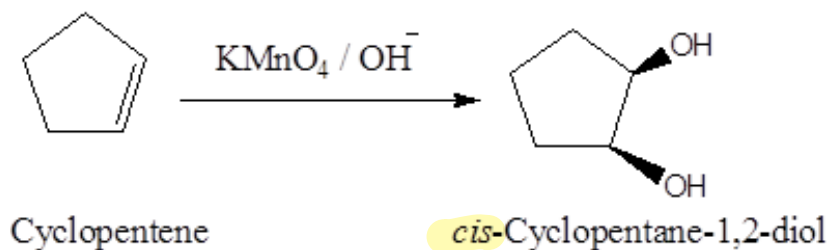
1. Addition of water to a double bond in the presence of an *acid catalyst, H⁺*.
2. The addition *follows Markovnikov's rule*.
3. *It is not possible to prepare primary alcohols* except Ethanol.



o *From Alkenes*

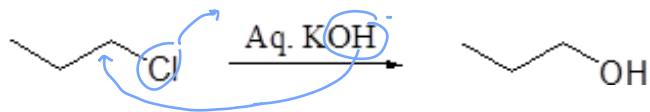
B. Oxidation of Cycloalkenes

Alkenes react with alkaline potassium permanganate to form **glycols** (compounds with two adjacent hydroxyl groups).



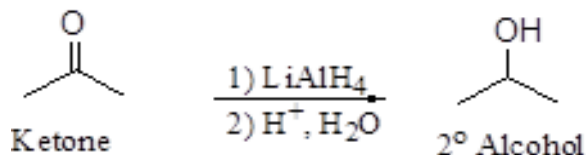
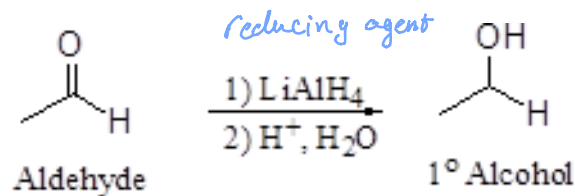
Preparation of Alcohols

o *Nucleophilic Substitution of Alkyl Halide*



o *Reduction of Ketones, and Aldehydes*

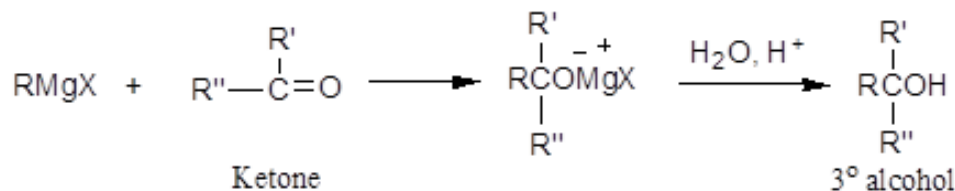
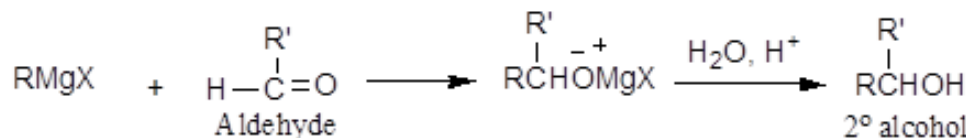
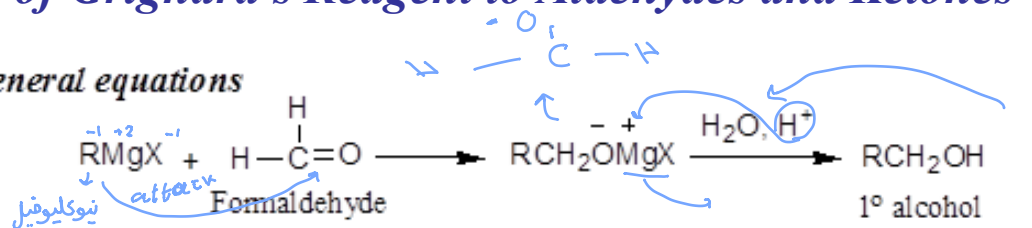
Aldehydes and ketones are easily reduced to primary and secondary alcohols, respectively.



Preparation of Alcohols

o *Addition of Grignard's Reagent to Aldehydes and Ketones*

General equations



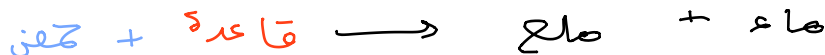
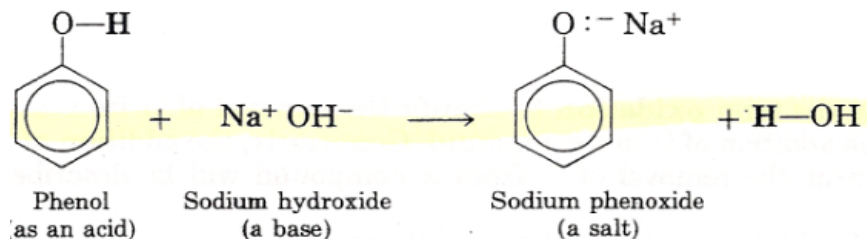
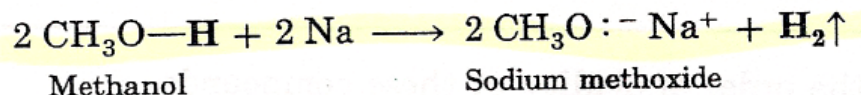
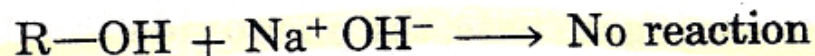
Reactions of Alcohols and Phenols

- o **Alcohols** undergo two kinds of reactions:
 - Those that involve the **breaking of the oxygen-hydrogen** bond (CO-H).
 - Those that involve the **rupture of the carbon-oxygen** bond (C-OH).
- o **Phenols** do not participate in reactions where the C-OH bond is broken.

Reactions of Alcohols

A) Those that involve the breaking of the oxygen-hydrogen bond (CO-H).

1) Reactions of Alcohols and Phenols as Acids: Salt Formation.

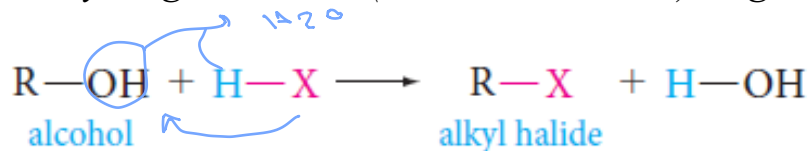


Reactions of Alcohols

B) Those that involve the rupture of the carbon-oxygen bond (C-OH).

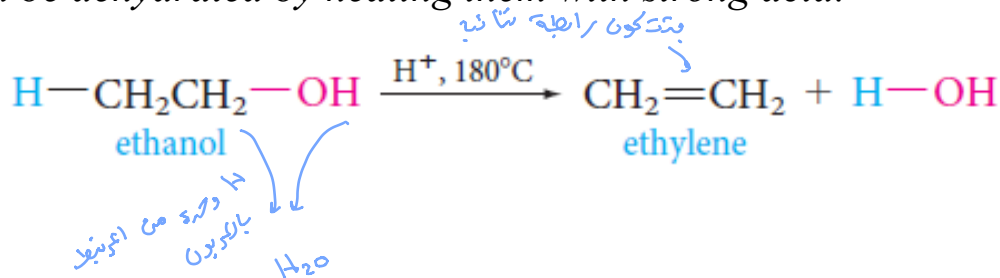
1) The Reaction of Alcohols with Hydrogen Halides: Alkyl Halides

Alcohols react with hydrogen halides (HCl, HBr and HI) to give alkyl halides.



2) Dehydration of Alcohols: Formation of Alkenes

Alcohols can be dehydrated by heating them with strong acid.



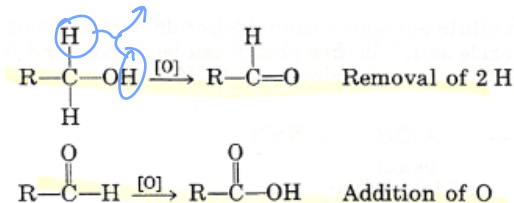
$3^\circ > 2^\circ > 1^\circ$

عندما تخرج مجموعة OH ينسأ على ذرة C شحنة موجبة والتي يفضل ان تكون على كربونة ثلاثية ثم ثانوية ثم أولية

Reactions of Alcohols

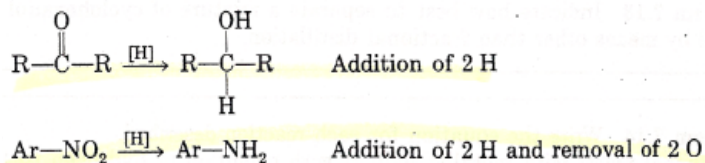
C) Oxidation Reactions

- **Oxidation** is the **removal of H** from a compound and/or the **addition of O** to a compound.



An oxidizing agent is the chemical reagent that does the oxidation.

- **Reduction** is the **addition of H** to a compound and/or the **removal of O** from a compound.

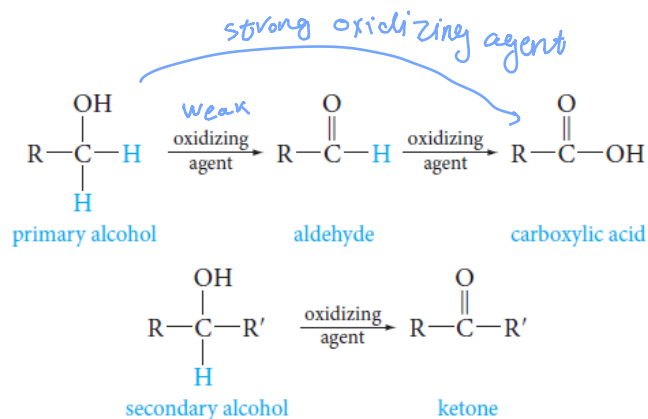


A reducing agent is a substance that does the reduction.

Reactions of Alcohols

C) Oxidation Reactions

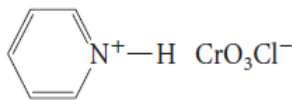
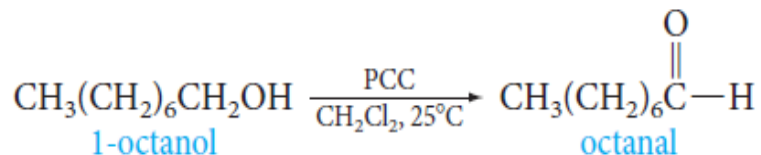
- o Alcohols with at least one hydrogen attached to the hydroxyl-bearing carbon can be oxidized to carbonyl compounds.
- **Primary alcohols** give **aldehydes**, which may be further oxidized to **carboxylic acids**.
 - **Secondary alcohols** give **ketones**.
 - **Tertiary alcohols**, having no hydrogen atom on hydroxyl-bearing carbon, do not undergo oxidation.



Reactions of Alcohols

C) Oxidation Reactions

- o **Primary alcohols**, oxidation can be stopped at aldehyde stage by special reagents, such as “pyridinium chlorochromate (PCC)”.



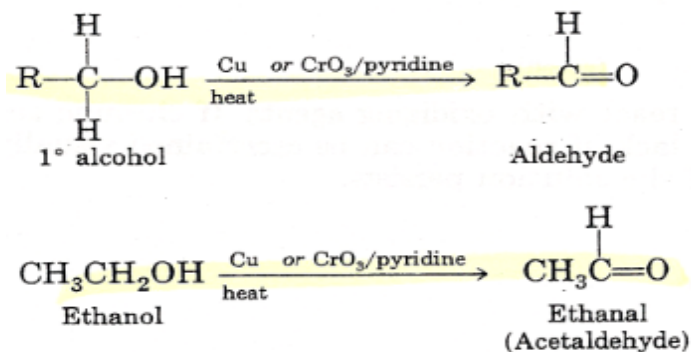
pyridinium chlorochromate
(PCC)

weak oxidizing agent

Reactions of Alcohols

C) Oxidation Reactions

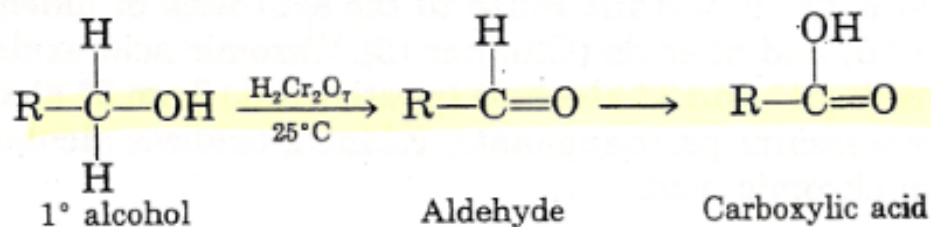
- o **Primary alcohols** yield **aldehydes** when treated with mild oxidizing agents such as hot metallic copper or CrO₃ in pyridine.



Reactions of Alcohols

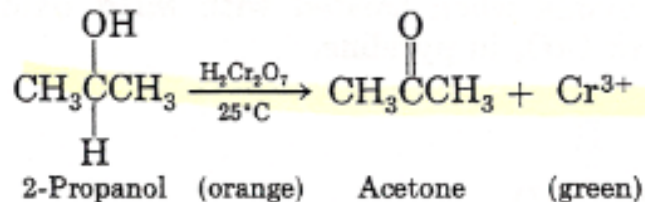
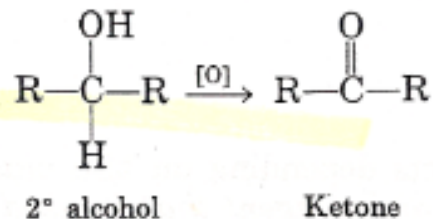
C) Oxidation Reactions

- o **Primary alcohols** yield **aldehydes** when treated with stronger oxidizing agents, such as **chromic acid, $\text{H}_2\text{Cr}_2\text{O}_7$** , or **neutral potassium permanganate, KMnO_4** , the **intermediate aldehydes** formed initially are oxidized further to **carboxylic acids**.

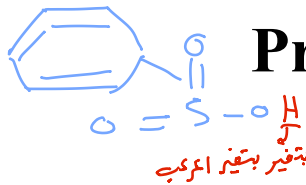


C) Oxidation Reactions

- o **Secondary alcohols**, when treated with *any of the oxidizing agents* mentioned previously, **yield ketones**.



sulfonate



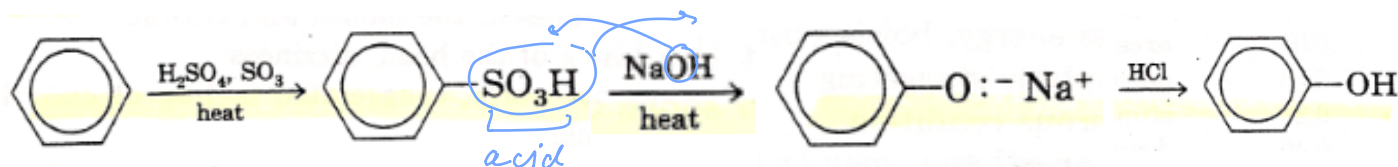
Preparation of Phenols

o

The Alkali Fusion of Sulfonates

The alkali fusion of sulfonates involves the following steps;

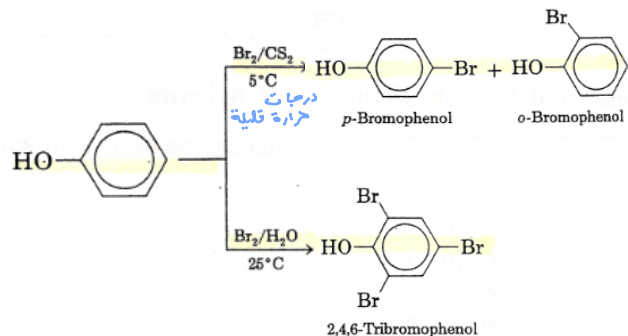
1. **Sulfonation** of an aromatic ring.
2. **Melting (fusion)** of the aromatic sulfonic acid with sodium hydroxide to give a **phenoxide salt**.
3. **Acidification** of the phenoxide with HCl to produce the **phenol**.



ليس من تفاعلات
حمضي - قاعدية

Reactions of Phenols

- Halogenation takes place *without catalyst*.



- The products depend on the solvent used.
- In *aprotic solvents* (solvents that do not release protons) (CCl_4 , CS_2)-bromination gives a mixture of *o*- and *p*-bromophenol.
 - In *protic solvents* (solvents that can release protons) (H_2O)-halogenation gives a trisubstituted phenol is produced.

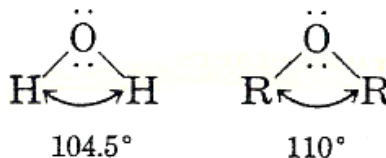
Ethers

Structure of Ethers

- o All **ethers** are compounds in which two organic groups are connected to a single oxygen atom. ممکن یکونوا نفس بعض ارمختلفات
- o The **general formula for an ether** is $R-O-R'$, where R and R' may be identical or different, and they may be alkyl or aryl groups



- o The geometry of simple ethers is similar to that of water.

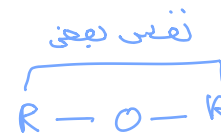


Structure of Ethers

o The ether is classified as

➤ Symmetrical ethers;

When the organic groups attached to the oxygen are identical.



➤ Unsymmetrical ethers (mixed ethers);

When the organic groups attached to the oxygen are different.

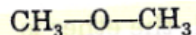


Nomenclature of Ethers

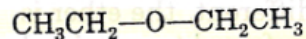
Common Names

Ethers are usually named by giving the name of each alkyl or aryl group, in alphabetical order, followed by the word *ether*.

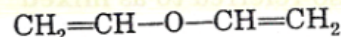
Methyl ether



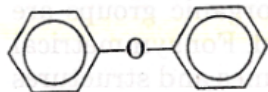
Ethyl ether



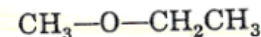
Vinyl ether



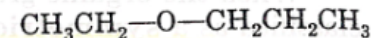
Phenyl ether



Ethyl methyl ether



Ethyl-*n*-propyl ether

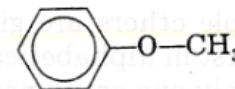


t-Butyl methyl ether



Methyl phenyl ether

(anisole)



benzil methyl
ether



فیه
فروق

Nomenclature of Ethers

IUPAC

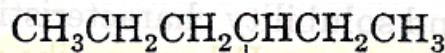
System

For ethers with more complex structures, it may be necessary to name the -OR group as an alkoxy group. In the IUPAC system, the smaller alkoxy group is named as a substituent.



بمائله كثره

2-Methoxypentane



3-Ethoxyhexane



2-Methoxyethanol

في الألوكة بالترقيم لإحاطة، R-O-R أقل رفق ممكن

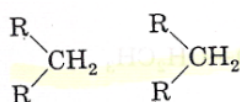
Physical Properties of Ethers

Physical

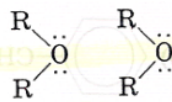
State

Boiling

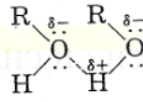
- **Points** They have *lower boiling points (bp,s) than alcohols* with an equal number of carbon atoms.
- In fact, an ether has nearly the **same bp as the corresponding hydrocarbon** in which a -CH₂- group replaces the ether's oxygen.
- *Because of their structures (no O-H bonds), ether molecules cannot form hydrogen bonds with one another.*



Alkanes: No hydrogen bonding between molecules; low boiling points



Ethers: No hydrogen bonding between molecules; low boiling points



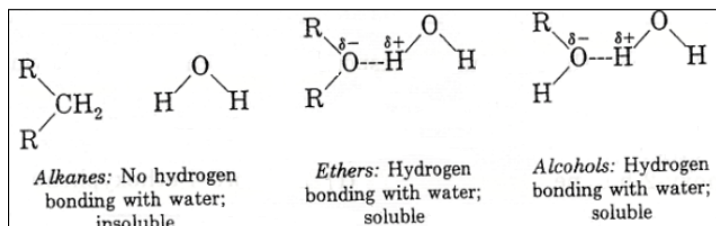
Alcohols: Hydrogen bonding between molecules; high boiling points

Compound	Formula	bp	mol wt	Water solubility (g/100 mL, 20°C)
1-butanol	CH ₃ CH ₂ CH ₂ CH ₂ OH	118°C	74	7.9
diethyl ether	CH ₃ CH ₂ -O-CH ₂ CH ₃	35°C	74	7.5
pentane	CH ₃ CH ₂ -CH ₂ -CH ₂ CH ₃	36°C	72	0.03

Physical Properties of Ethers

Solubility

- Low-molecular-weight ethers, such as dimethyl ether, are quite soluble in water.
- Ether molecules can form hydrogen bonds to water.*



Structure	Name	Mol.wt.	Bp (°C)	Solubility in H ₂ O At 20 °C
CH ₃ CH ₂ CH ₃	propane	44	−42	insoluble
CH ₃ OCH ₃	methyl ether	46	−24	soluble
CH ₃ CH ₂ OH	ethanol	46	78	soluble
CH ₃ CH ₂ CH ₂ CH ₃	<i>n</i> -butane	58	−0.5	insoluble
CH ₃ CH ₂ OCH ₃	ethyl methyl ether	60	8	soluble
CH ₃ CH ₂ CH ₂ OH	1-propanol	60	97	soluble
CH ₃ (CH ₂) ₃ CH ₃	<i>n</i> -pentane	72	35	insoluble
CH ₃ CH ₂ OCH ₂ CH ₃	ethyl ether	74	36	7.5 g/100 g
CH ₃ (CH ₂) ₂ CH ₂ OH	1-butanol	74	118	7.9 g/100 g
CH ₃ (CH ₂) ₅ CH ₃	<i>n</i> -heptane	100	98	insoluble
CH ₃ (CH ₂) ₂ O(CH ₂) ₂ CH ₃	<i>n</i> -propyl ether	102	91	0.2 g/100 g
CH ₃ (CH ₂) ₄ CH ₂ OH	1-hexanol	102	157	0.6 g/100 g

Preparation of Ethers

- o There are *two general methods* for synthesizing ethers.

1) Dehydration of alcohols

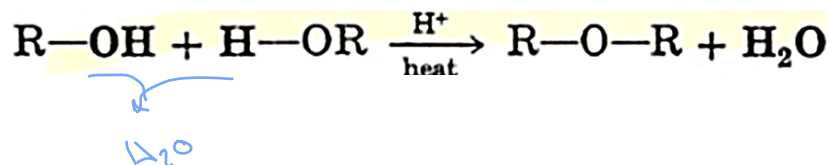
It is used **commercially** and in the **laboratory** to make certain **symmetrical ethers**.

2) Williamson synthesis

General **laboratory** method used to prepare all kinds of ethers, **symmetrical** and **unsymmetrical**.

1) Dehydration of Alcohols

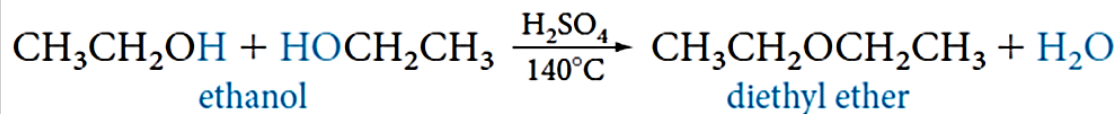
It takes place in the presence of acid catalysts (H_2SO_4 , H_3PO_4) (intermolecular reaction)



لازم يستخدم نفس نوع الـ R

Example;

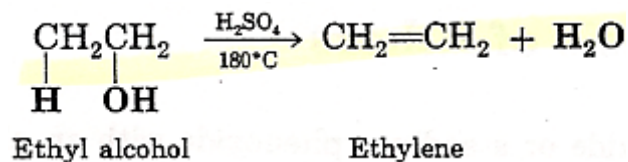
The most important commercial ether is diethyl ether. It is prepared from ethanol and sulfuric acid.



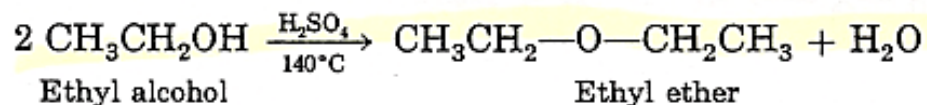
Scope	and
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Limitations

- o When ethyl alcohol is dehydrated by sulfuric acid at 180° C, the dominant product is **ethylene**.



- o **To prepare ethyl ether**
 - Dissolve ethyl alcohol in sulfuric acid at ambient temperature.
 - Heat the solution to 140°C while adding more alcohol.



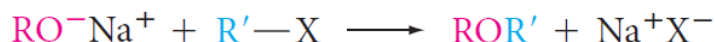
2)	Williamson
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Synthesis

- This method has two steps;
 - 1) An alcohol is converted to its alkoxide by treatment with a reactive metal (sodium or potassium).



- 2) Displacement is carried out between the alkoxide and an alkyl halide.



- To obtain the best yields of mixed dialkyl ethers, we select a 1° rather than a 2° or 3° alkyl halide and react it with a sodium alkoxide
- To prepare an alkyl aryl ether, we must be careful not to pick a combination in which one of the reagents has a halogen directly attached to an aromatic ring.

2) Williamson Synthesis

o Example 1; Preparation of *t*-butyl methyl ether, (CH₃)₃C-O-CH₃.

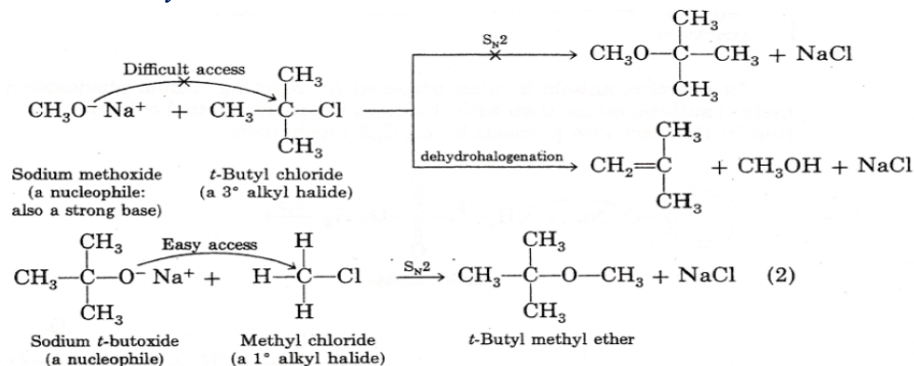
➤ In theory, this could be done by either of two reactions.

1. You could react sodium methoxide, CH₃O-Na⁺, with *t*-butyl chloride, (CH₃)₃C-Cl.

This combination leads to dehydrohalogenation to an alkene, an elimination reaction.

2. You could react sodium *t*-butoxide, (CH₃)₃C-O-Na⁺, with methyl chloride, CH₃Cl.

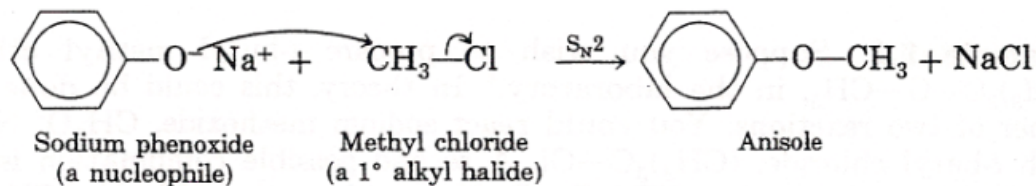
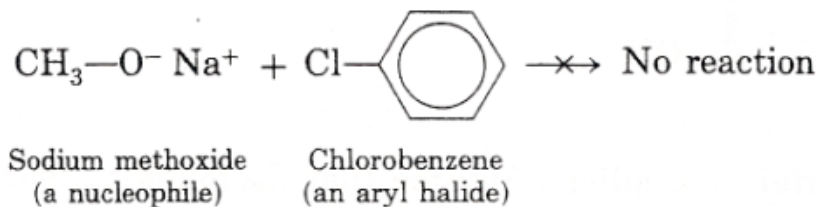
This route gives the desired ether by substitution.



2)Williamson Synthesis

Example 2; Assume you need to synthesize **methyl phenyl ether (anisole), CH₃-O-C₆H₅**, by the Williamson method.

➤ *In theory, you could obtain anisole in either of two ways.*

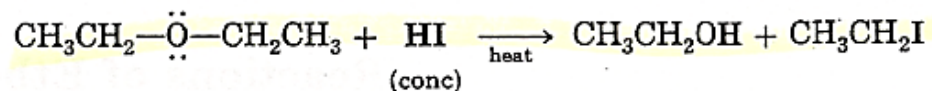


Reactions of Ethers

- o **Ethers** are quite stable compounds.
- o The **ether** linkage does not react with bases, reducing agents, oxidizing agents, or active metals.
- o ***Ethers** react only under strongly acidic conditions.*

Cleavage of Ethers by Hot Concentrated Acids

- When **ethers** are heated in concentrated acid solutions, the ether linkage is broken.



- The acids most often used in this reaction are HI, HBr, and HCl.
- If an *excess of acid* is present, the alcohol initially produced is converted into an alkyl halide by the reaction.



For example,

