

Chapter 8



Substitution and Elimination Reactions of Alkyl Halides

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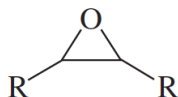
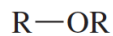
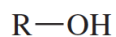
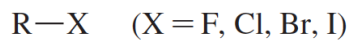
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The Families of Group III

Substitution Reactions of Alkyl Halides

Group III

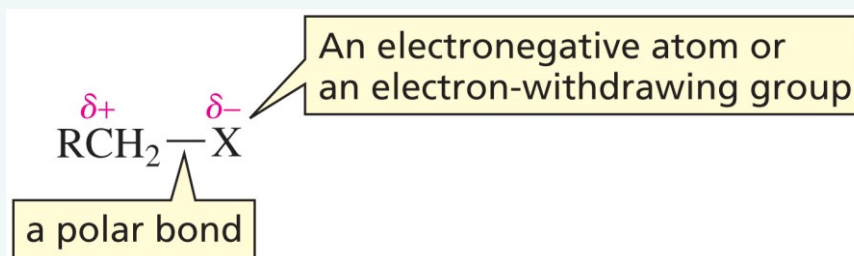


alkyl halides:
the first of the families
in Group III

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The Compounds in Group III are Electrophiles

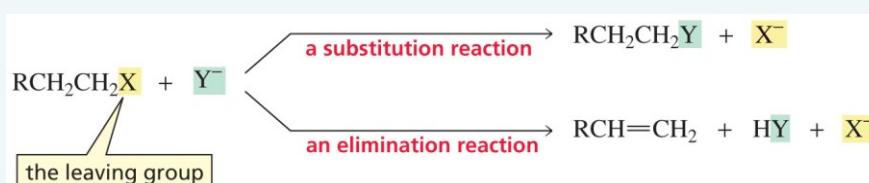


All the compounds in **Group III** have an **electron withdrawing atom or group** that is attached to an **sp³ carbon**.

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Because Group III Compounds are Electrophiles, they react with Nucleophiles



Substitution reaction—the electronegative group is **replaced** by another group.

Elimination reaction—the electronegative group is **eliminated** along with a hydrogen.

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Alkyl Halides

alkyl halides



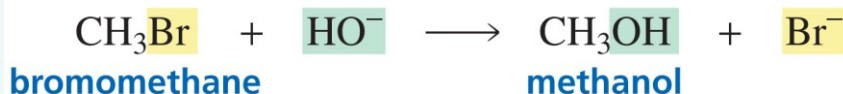
This chapter discusses the **substitution reactions** of **alkyl halides**.

Alkyl halides have **good leaving groups**.

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The S_N2 Reaction



The substitution reaction is more precisely called a **nucleophilic substitution reaction** because the atom replacing the halogen is a **nucleophile**.

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What is the Mechanism of the Reaction?

The **kinetics** of a reaction—the **factors that affect the rate of the reaction**—can help determine the mechanism.

$$\text{rate} \propto [\text{alkyl halide}][\text{nucleophile}]$$

$$\text{rate} = k [\text{alkyl halide}][\text{nucleophile}]$$

the rate constant

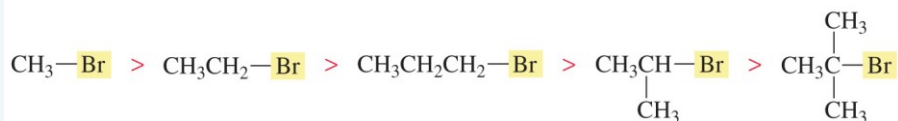
an S_N2 reaction

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Relative Rates of an S_N2 Reaction

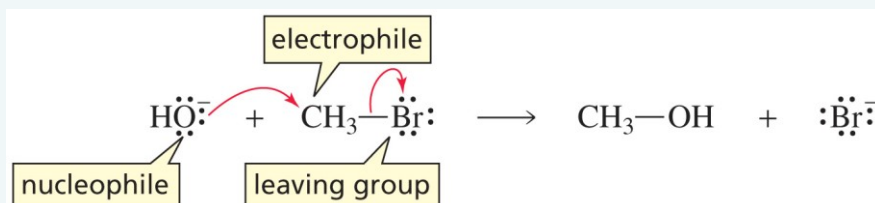
Relative Rates of an S_N2 Reaction



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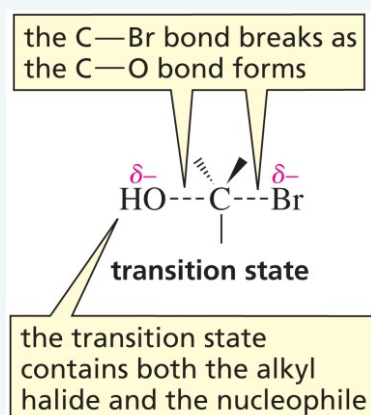
Mechanism for the S_N2 Reaction of an Alkyl Halide



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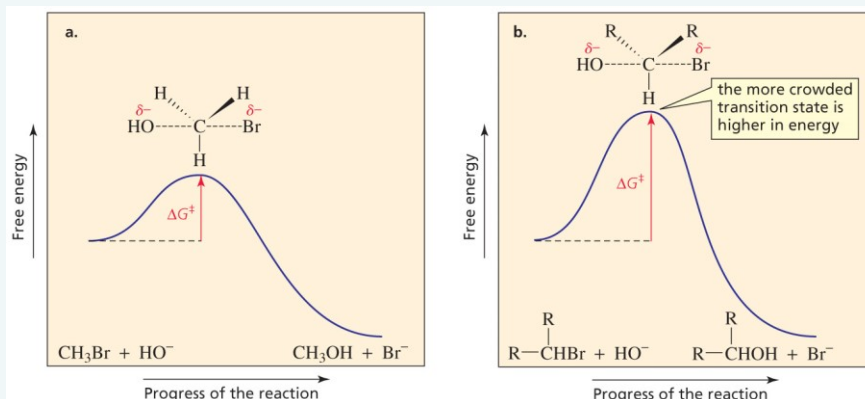
Why Bimolecular?



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Why Steric Hindrance decreases the rate



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Both are Primary Alkyl Halides but they react at different rates

more reactive in an S_N2 reaction

CH₃CH₂Br
an ethyl group

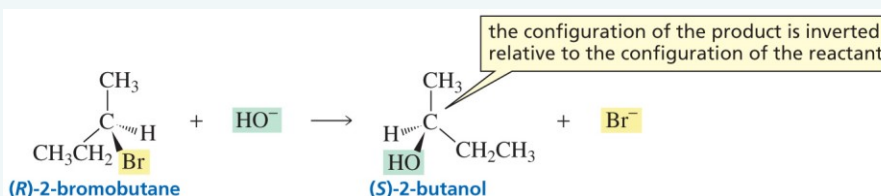
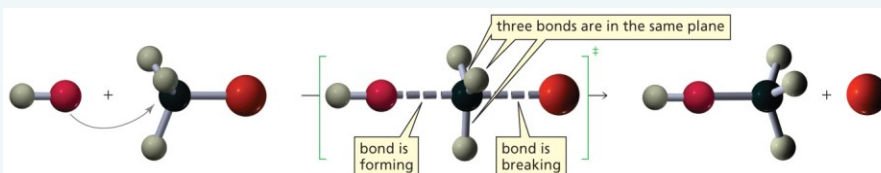
less reactive in an S_N2 reaction

CH₃CH₂CH₂Br
a propyl group

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Why the configuration of the Product is Inverted



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The rate of an S_N2 Reaction is affected by the Leaving Group

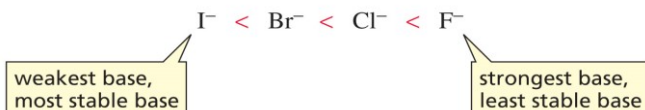
	relative rates of reaction	pK _a values of HX
HO ⁻ + RCH ₂ I → RCH ₂ OH + I ⁻	30,000	-10
HO ⁻ + RCH ₂ Br → RCH ₂ OH + Br ⁻	10,000	-9
HO ⁻ + RCH ₂ Cl → RCH ₂ OH + Cl ⁻	200	-7
HO ⁻ + RCH ₂ F → RCH ₂ OH + F ⁻	1	3.2

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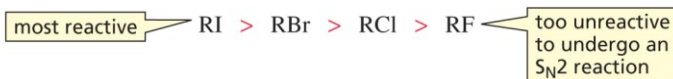
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The Weakest Base is the best Leaving Group

relative basicities of the halide ions



relative reactivities of alkyl halides in an $\text{S}_{\text{N}}2$ reaction

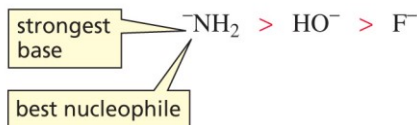


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Base Strength and Nucleophile Strength

relative base strengths and relative nucleophilicities



If atoms are in the same row, the **strongest base** is the **best nucleophile**.

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Base Strength and Nucleophile Strength

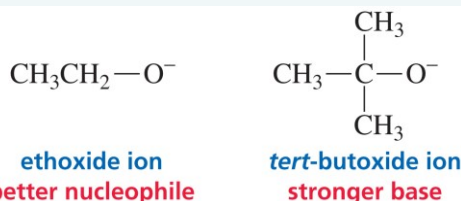
stronger base, better nucleophile		weaker base, poorer nucleophile
HO^-	>	H_2O
CH_3O^-	>	CH_3OH
$^-\text{NH}_2$	>	NH_3
$\text{CH}_3\text{CH}_2\text{NH}^-$	>	$\text{CH}_3\text{CH}_2\text{NH}_2$

A **negatively charged** atom is a **stronger base** and a **better nucleophile** than the same atom that is **neutral**.

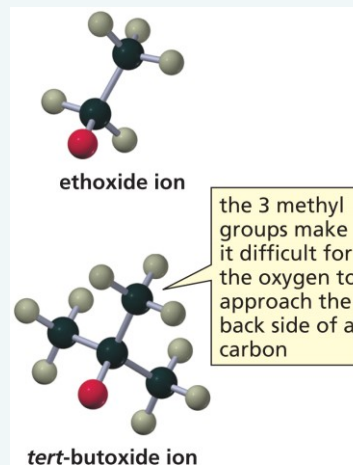
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Steric Hindrance Decreases Nucleophilicity



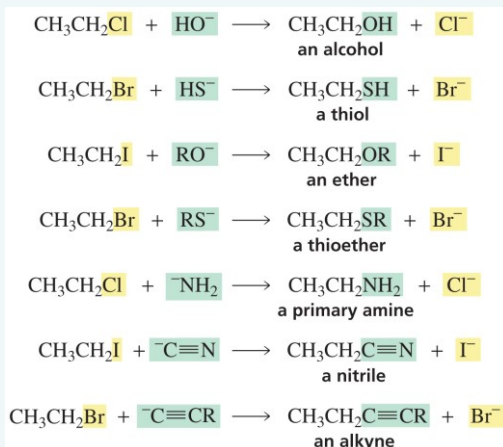
Even though the *tert*-butoxide ion is a **stronger base**, it is a **poorer nucleophile** because nucleophilic attack is more sterically hindered than proton removal.



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S_N2 Reactions can be used to make a variety of Compounds

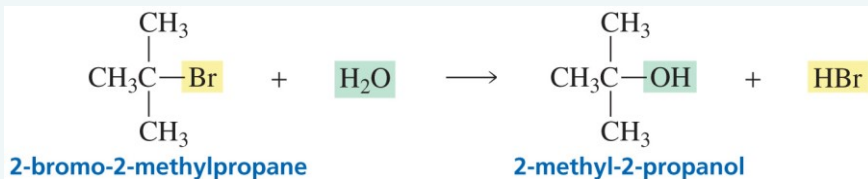


The reactions are irreversible because a **strong base** displaces a **weak base**.

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The S_N1 Reaction



A **tertiary alkyl halide** and a **poor nucleophile**

The reaction is surprisingly fast, so it must be taking place by a **different mechanism**.

Most S_N1 reactions are **solvolysis reactions**: the **solvent** is the **nucleophile**.

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The rate of an S_N1 Reaction depends only on the Alkyl Halide Concentration

$$\text{rate} = k [\text{alkyl halide}]$$

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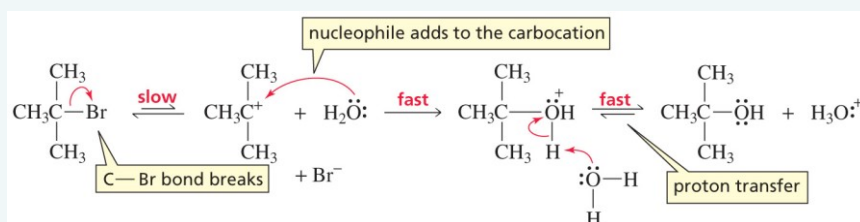
Summary of the Experimental Evidence for the Mechanism of an S_N1 Reaction

1. The rate of the reaction depends only on the concentration of the alkyl halide.
1. Tertiary alkyl halides react the fastest.
1. If the halogen is attached to an asymmetric center the product will be a pair of enantiomers.

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The Mechanism

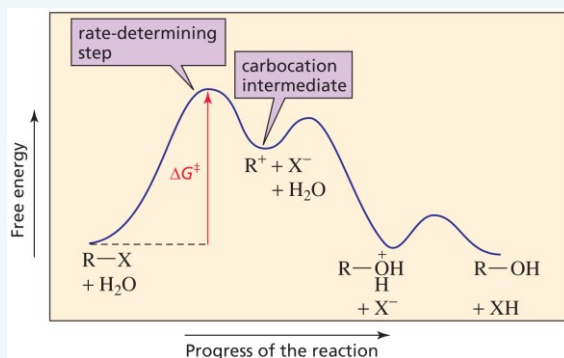


The leaving group departs **before** the nucleophile approaches.

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The Slow Step is Formation of the Carbocation



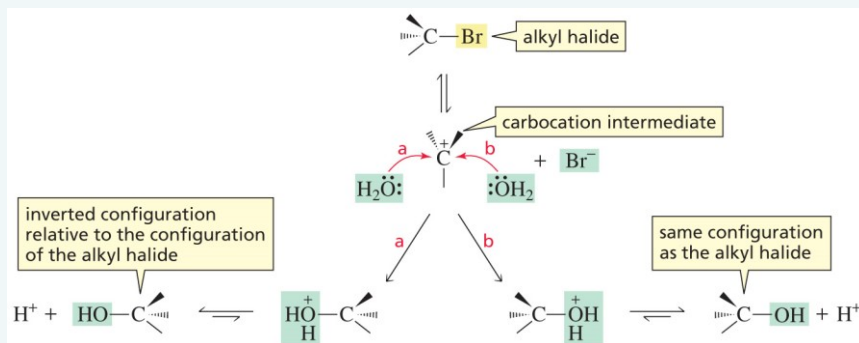
Tertiary alkyl halides react the fastest; they form the most stable carbocations.

Secondary and primary alkyl halides do **not** undergo $\text{S}_{\text{N}}1$ reactions.

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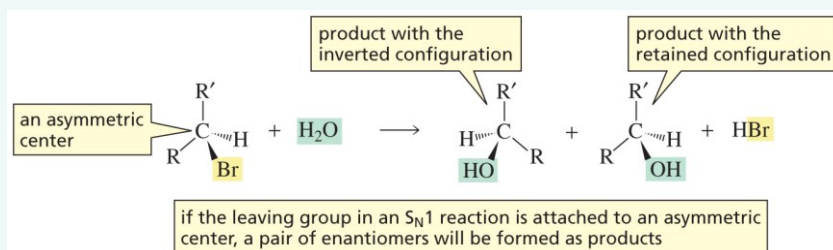
The Product is a Pair of Enantiomers



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The Product is a Pair of Enantiomers



If the halogen is bonded to an asymmetric center, the product will be a **pair of enantiomers**.

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The Leaving Group in an S_N1 Reaction

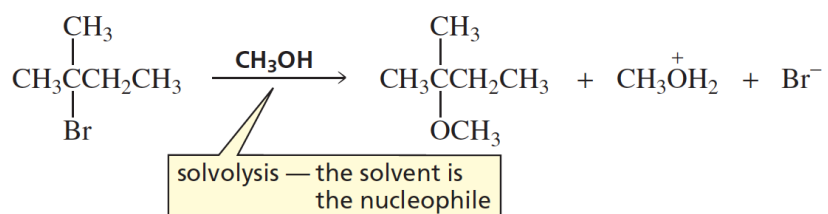
relative reactivities of alkyl halides in an S_N1 reaction



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The Nucleophile in an S_N1 Reaction



Most S_N1 reactions are **solvolysis** reactions; the **nucleophile** is the **solvent**.

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Comparing S_N2 and S_N1 Reaction

S_N2

- a one-step mechanism
- a bimolecular rate-determining step
- the rate is controlled by steric hindrance
- product has the inverted configuration relative to that of the reactant
- the leaving group: $\text{I}^- > \text{Br}^- > \text{Cl}^- > \text{F}^-$
- the better the nucleophile, the faster the rate of the reaction

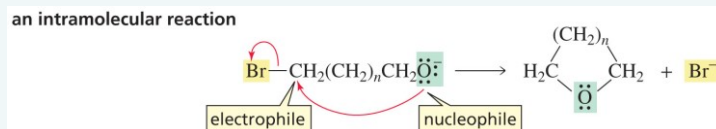
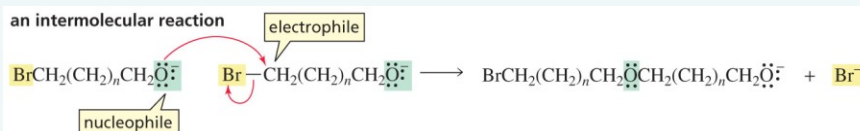
S_N1

- a two-step mechanism with a carbocation intermediate
- a unimolecular rate-determining step
- the rate is controlled by stability of the carbocation
- products have both the retained and inverted configurations relative to that of the reactant
- the leaving group: $\text{I}^- > \text{Br}^- > \text{Cl}^- > \text{F}^-$
- the strength of the nucleophile does not affect the rate of the reaction

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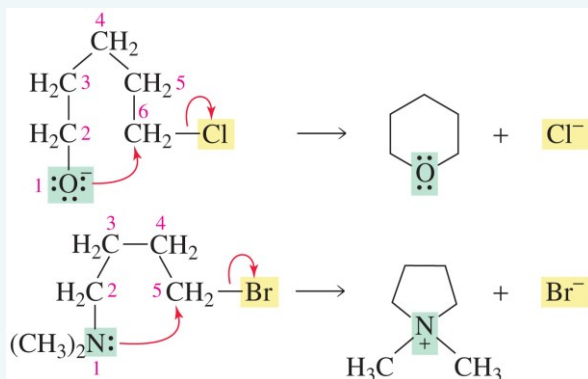
Intermolecular *versus* Intramolecular S_N2 Reactions



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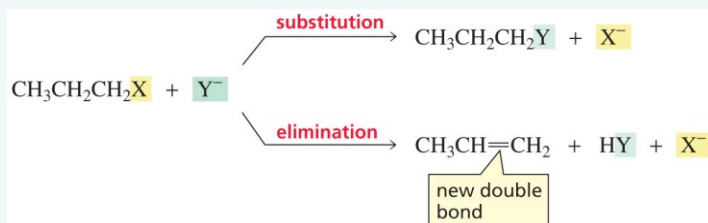
The Intramolecular Reaction is favored when a Five- or Six-Membered Ring can be formed



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Alkyl Halides undergo Substitution and Elimination Reactions



In an **elimination reaction**:

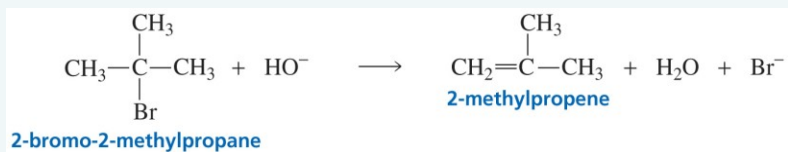
a **halogen** is removed from one carbon and
a **hydrogen** is removed from an adjacent carbon.

a **double bond** is formed between the two carbons
from which the atoms were removed.

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An E2 Reaction

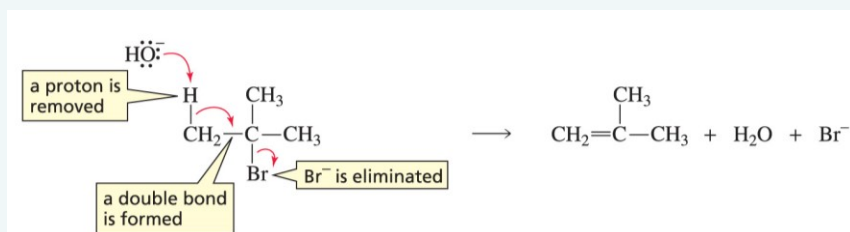


$$\text{rate} = k[\text{alkyl halide}][\text{base}]$$

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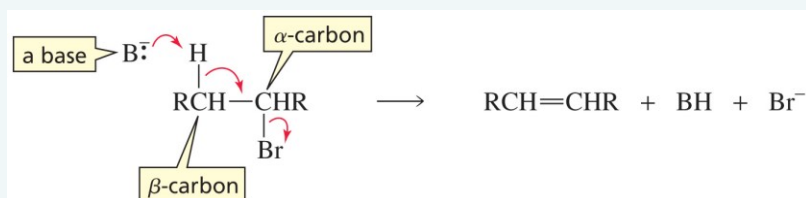
Mechanism for an E2 Reaction



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The Halogen comes off the Alpha Carbon;
the Hydrogen comes off the Beta Carbon

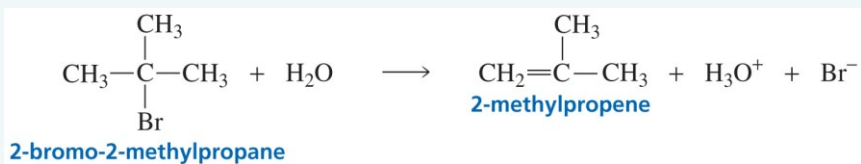


dehydrohalogenation

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An E1 Reaction

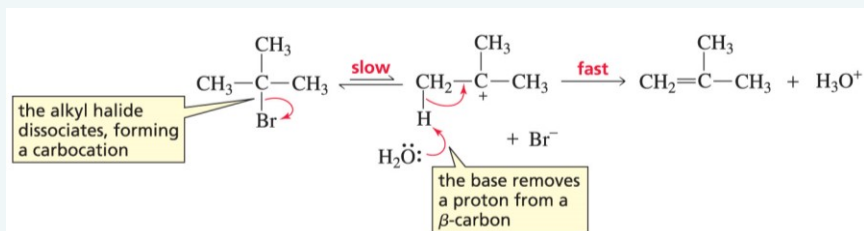


$$\text{rate} = k[\text{alkyl halide}]$$

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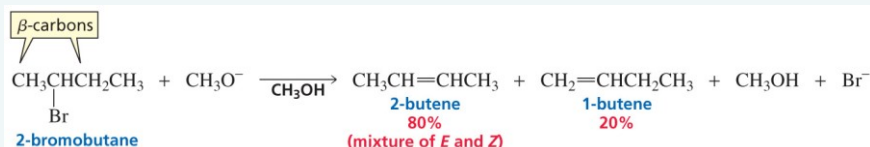
The Mechanism for an E1 Reaction



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An E2 Reaction is Regioselective



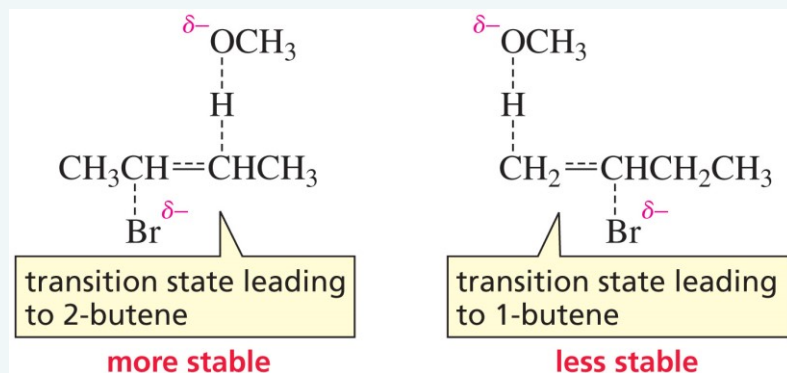
The **major product** is the **most stable** alkene.

The **most stable alkene** is (generally) obtained by removing a hydrogen from the beta carbon that is bonded to the **fewest hydrogens**.

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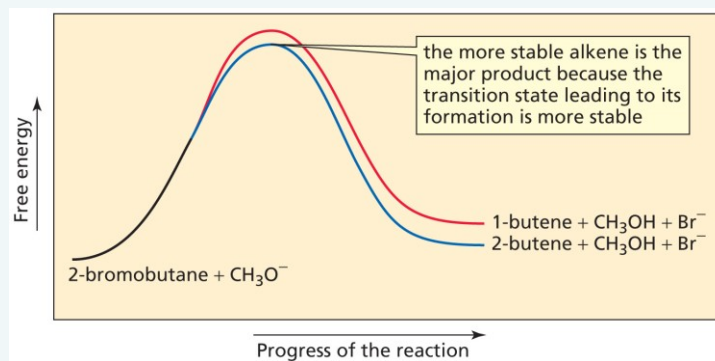
Alkene-Like Transition State



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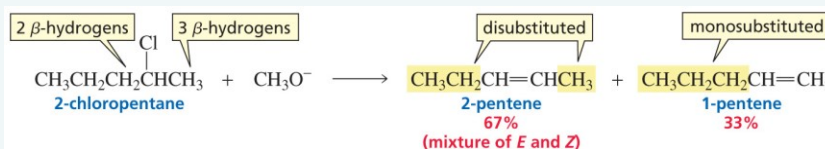
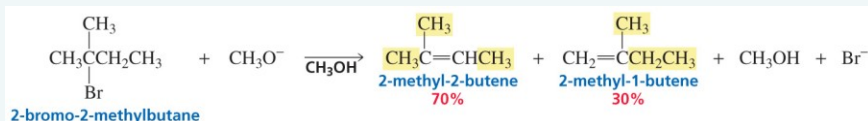
The more stable Alkene has the more stable Transition State



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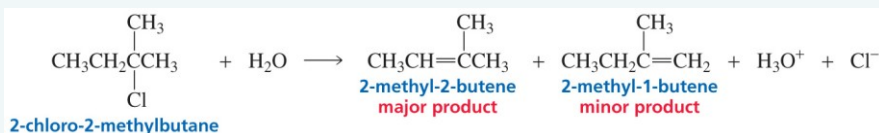
More Regioselective E2 Reactions



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An E1 Reaction is Regioselective



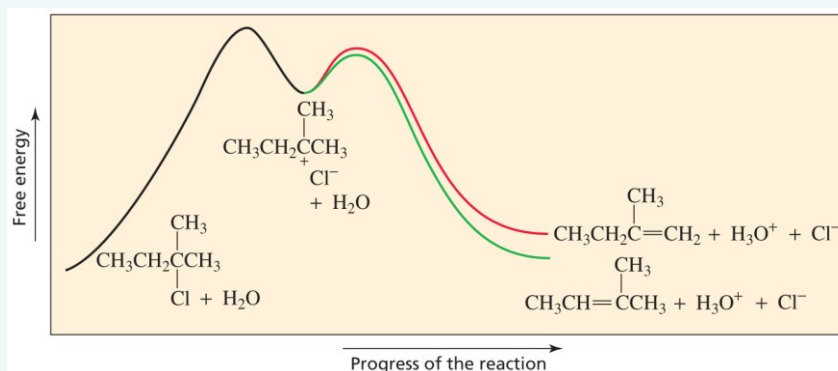
The **major product** is the **more stable alkene**.

The **most stable alkene** is obtained by removing a hydrogen from the beta carbon that is bonded to the **fewest hydrogens**.

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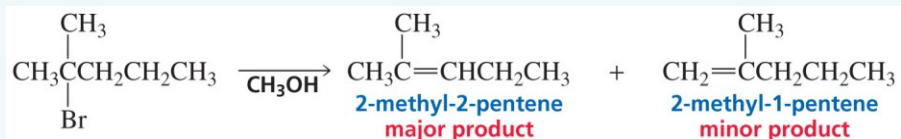
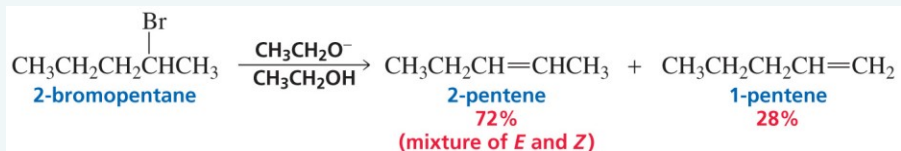
The more stable Alkene is the major product



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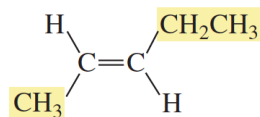
Both E2 and E1 Reactions are Regioselective



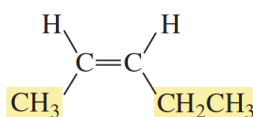
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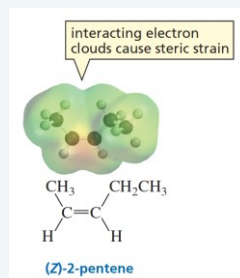
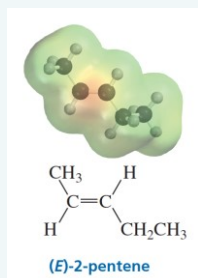
E2 and E1 Reactions are Stereoselective



(E)-2-pentene
major product



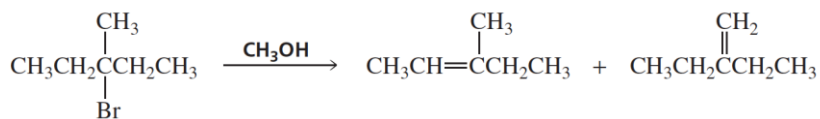
(Z)-2-pentene
minor product



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E2 and E1 Reactions are Stereoselective



3-bromo-3-methylpentane

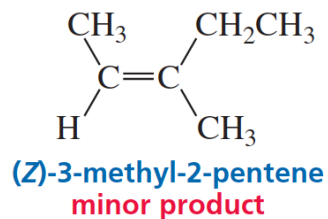
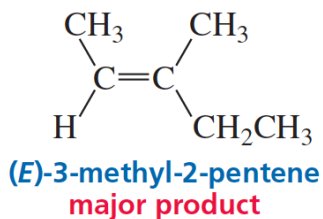
3-methyl-2-pentene
major product

2-ethyl-1-butene
minor product

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E2 and E1 Reactions are Stereoselective



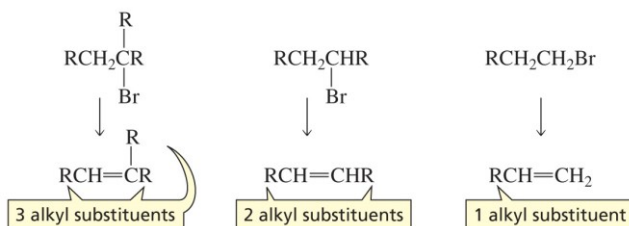
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Relative Reactivities of Alkyl Halides in an E2 Reaction

relative reactivities of alkyl halides in an E2 reaction

tertiary alkyl halide > secondary alkyl halide > primary alkyl halide



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Does a Tertiary Alkyl Halide undergo $S_N2/E2$ or $S_N1/E1$ Reactions?

$$\text{rate} = k_1[\text{alkyl halide}] + k_2[\text{alkyl halide}][\text{nucleophile}] + k_3[\text{alkyl halide}] + k_4[\text{alkyl halide}][\text{base}]$$

contribution to the rate by an S_N1 reaction

contribution to the rate by an S_N2 reaction

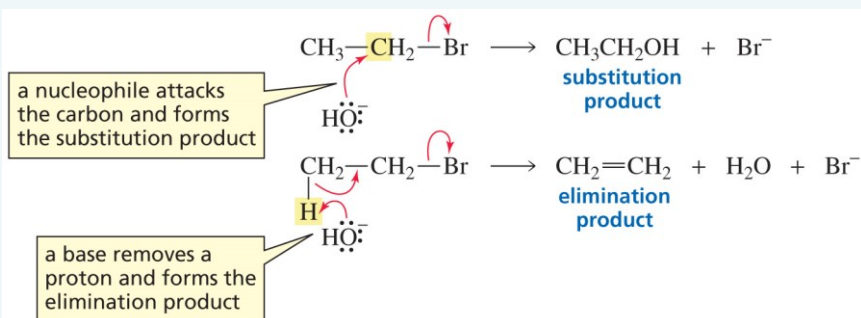
contribution to the rate by an E1 reaction

contribution to the rate by an E2 reaction

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Competition between Substitution and Elimination

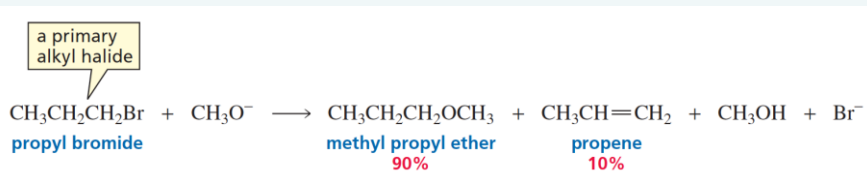


In an S_N2 reaction: $1^\circ > 2^\circ > 3^\circ$ In an E2 reaction: $3^\circ > 2^\circ > 1^\circ$

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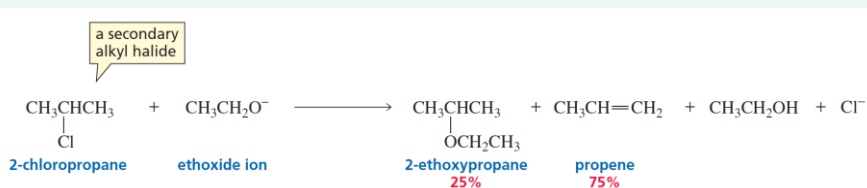
Under S_N2/E2 conditions, a Primary Alkyl Halide forms primarily a Substitution Product



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Under S_N2/E2 conditions, a Secondary Alkyl Halide forms Substitution and Elimination Products



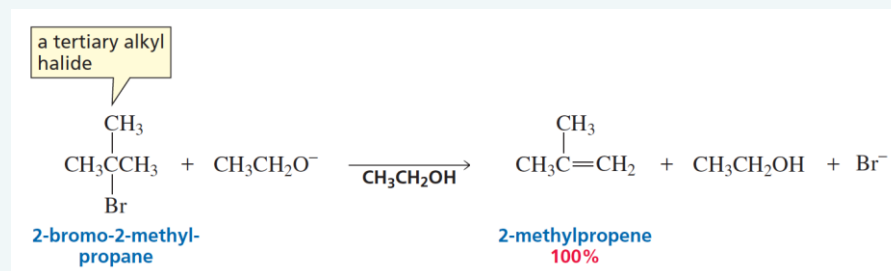
Substitution is favored by a weak base.

Elimination is favored by a strong base.

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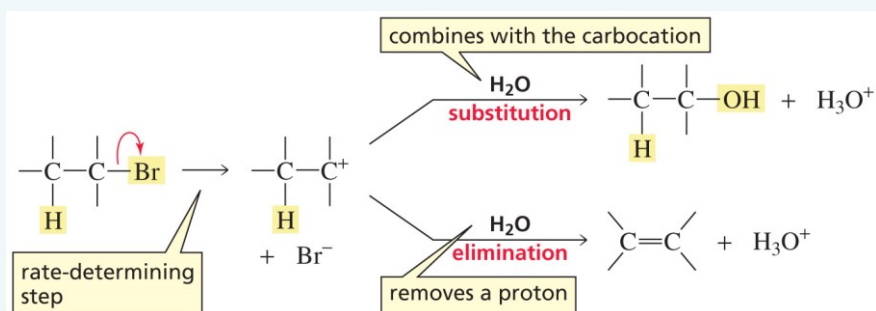
Under S_N2/E2 conditions, a Tertiary Alkyl Halide forms only an Elimination Product



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Under S_N1/E1 conditions, a Tertiary Alkyl Halide forms Substitution and Elimination Products

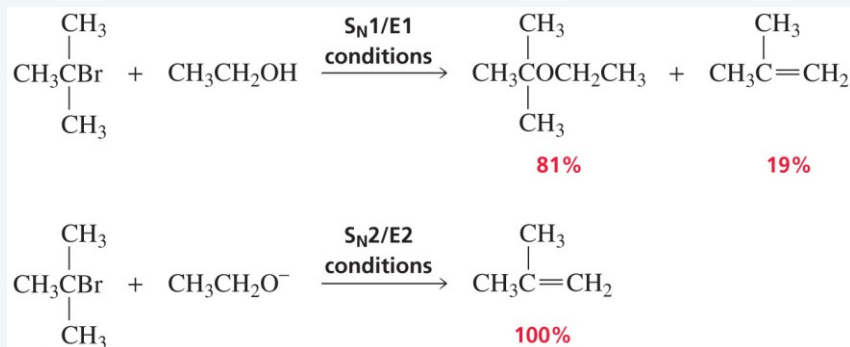


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Tertiary ($S_N1/E1$): Substitution is Favored

Tertiary ($S_N2/E2$): Only Elimination



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Summary of the products obtained from Substitution and Elimination Reactions

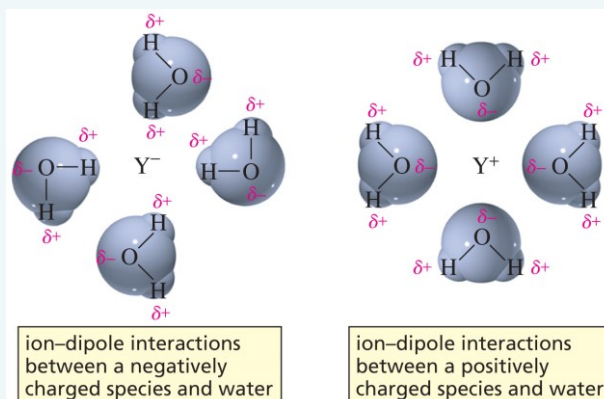
Table 8.2 Summary of the Products Expected in Substitution and Elimination Reactions

Class of alkyl halide	S_N2 versus $E2$	S_N1 versus $E1$
Primary alkyl halide	primarily substitution	cannot undergo $S_N1/E1$ solvolysis reactions
Secondary alkyl halide	substitution and elimination	cannot undergo $S_N1/E1$ solvolysis reactions
Tertiary alkyl halide	only elimination	substitution and elimination with substitution favored

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Solvation

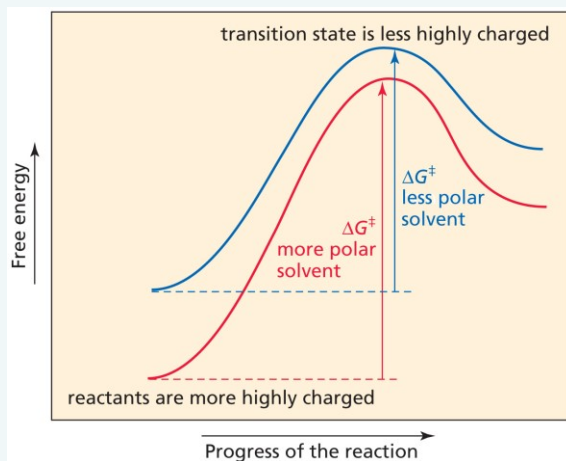


Polar solvents **stabilize charged species.**

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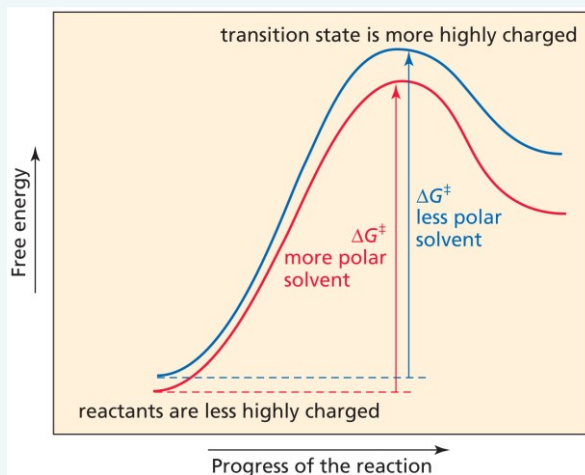
If a reactant is charged, increasing the polarity of the solvent decreases the Rate



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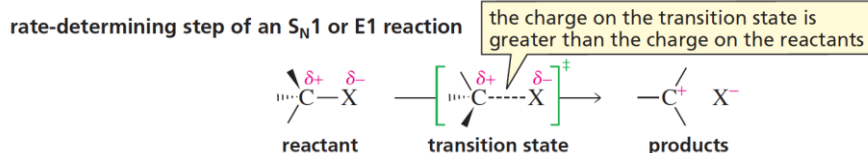
If the reactants are neutral, increasing the polarity of the solvent increases the Rate



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How a solvent affects the Rate of a Reaction that does not have a Charged Reactant



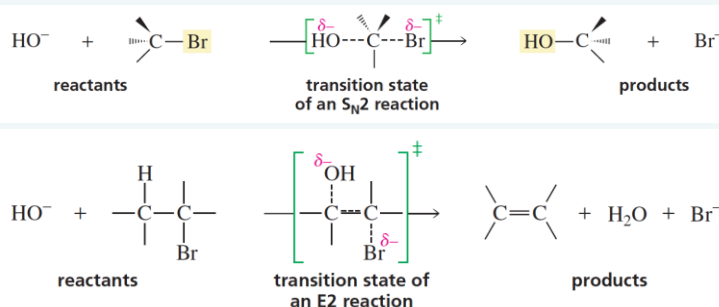
If a **reactant** is not **charged**:
 the **charge on the reactants** will be **greater** than the charge on the **transition state**.

Increasing the **polarity** of the solvent will **increase** the **rate** of the reaction.

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How a solvent affects the Rate of a Reaction that has a Charged Reactant



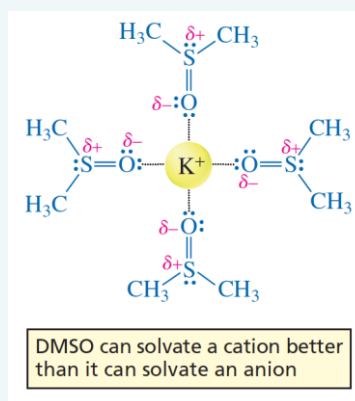
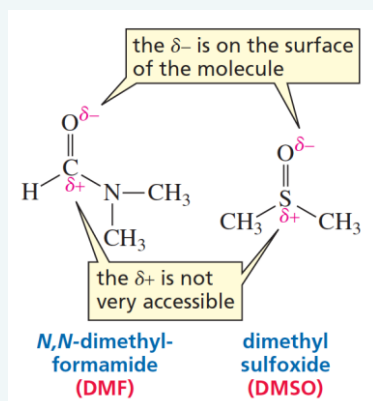
If a **reactant** is **charged**:
the **charge on the reactants** will be **greater** than the charge on the **transition state**.

Increasing the **polarity** of the solvent will **decrease** the **rate** of the reaction.

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DMF and DMSO



These solvents can solvate cations (+) better than they can solvate anions (-).

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William Ether Synthesis: an S_N2 Reaction

Williamson ether synthesis



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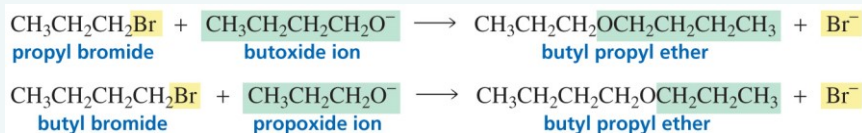
Forming an Alkoxide Ion



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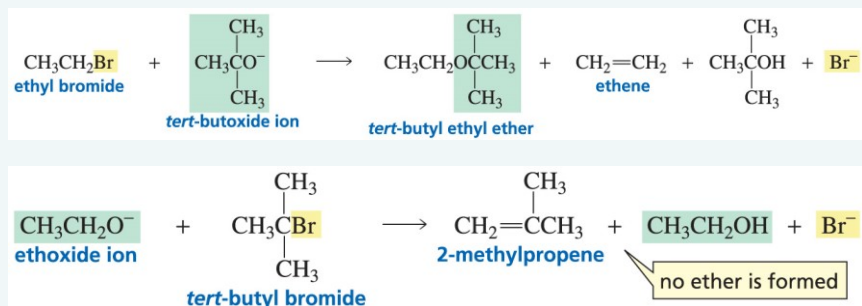
Synthesizing Butyl Propyl Ether



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Synthesizing *tert*-Butyl Ethyl Ether



The **less hindered** group should be provided by the **alkyl halide**.

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