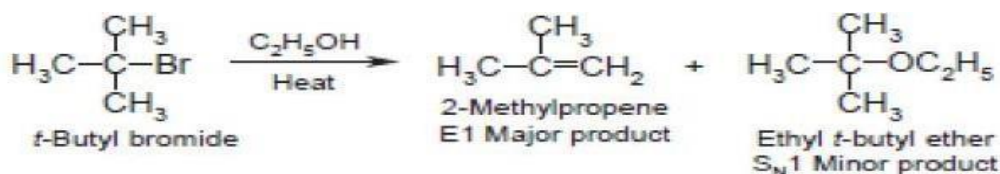


BP 202T - PHARMACEUTICAL ORGANIC CHEMISTRY - I**UNIT: 2 ALKANES, ALKENE & CONJUGATED DIENES****CLASS: 3****TOPIC: E1 & E2 REACTIONS-KINETICS, E1 VS E2 REACTIONS,
FACTORS AFFECTING E1 & E2****ELIMINATION REACTION**

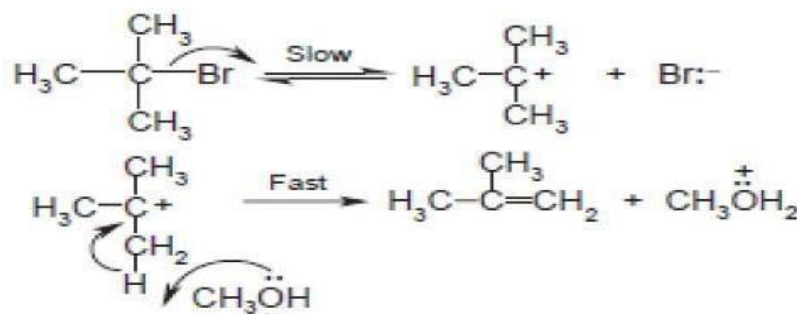
- Elimination reaction is a type of organic reaction in which two substituent's are removed from a molecule in either a one or two-step mechanism.
- The one-step mechanism is known as the E2 reaction, and the two-step mechanism is known as the E1 reaction.
- In most organic elimination reactions, at least one hydrogen is lost to form the double bond: the unsaturation of the molecule increases.
- It is also possible that a molecule undergoes reductive elimination, by which the valence of an atom in the molecule decreases by two, though this is more common in inorganic chemistry.
- There are three fundamental events in these elimination reactions:
 - removal of a proton
 - formation of the CC π bond
 - breaking of the bond to the leaving group

E₁ REACTION

- E1 reaction or first order elimination results from the loss of a leaving group to form a carbocation intermediate, followed by the removal of a proton to form the C=C bond.
- This reaction is most common with good leaving groups, **stable carbocations** and **weak bases** (strong acids)



Mechanism.



- In the E1 mechanism which is also **known as unimolecular elimination**, there are usually two steps involved – ionization and deprotonation.
- During ionization, there is a formation of **carbocation as an intermediate**. In deprotonation, a proton is lost by the carbocation.
- This happens in the presence of a base which further leads to the formation of a pi-bond in the molecule.
- In E1, the reaction rate is also proportional to the concentration of the substance to be transformed.
- It exhibits **first-order kinetics**.
- **For example**, ter-butyl bromide (3° alkyl halide) reacts with water to form 2-methylpropene, following an E1 mechanism.

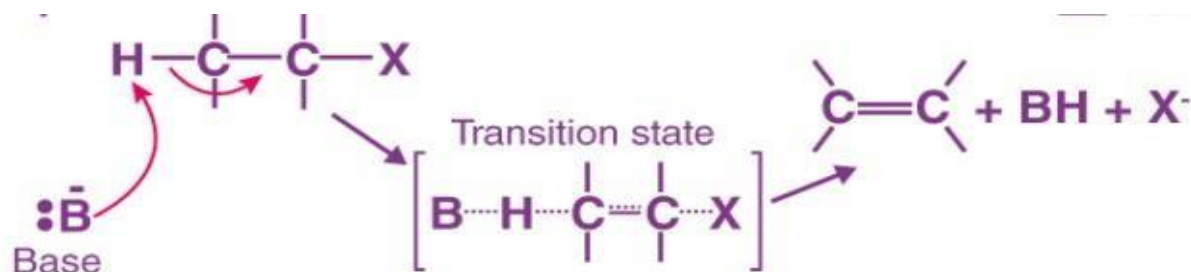
Characteristics of an E₁ Reaction

- Kinetics – First order
- Mechanism – Two step process
- Identity of R group – More substituted halides react faster
 - Rate: $\text{R}_3\text{CX} > \text{R}_2\text{CHX} > \text{RCH}_2\text{X}$
 - Strength of the base – Favored by weaker bases such as H_2O and ROH
 - Leaving group – Better leaving group leads to faster reaction rates. The rate determining step involves the C—X bond cleavage
 - Type of solvent – Favored by **polar Protic solvents**, which can stabilize the ionic

intermediates

E₂ REACTION

- Kinetics – Second order
 - Mechanism – Single step
 - Identity of R group – More substituted halides react faster
 - Rate: $R_3CX > R_2CHX > RCH_2X$
 - Strength of the base – Stronger bases favor the reaction
 - Leaving group – Better leaving group leads to faster reaction rates
 - Type of solvent – Favored by polar aprotic solvents
 - E₂ reactions are stereo selective, resulting in the formation of trans-double bonds preferably.
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- Dehydrohalogenation of sec- and ter- alkyl halides undergo both E₁ and E₂ reactions. However, prim-halides undergo only E₂ reactions.
 - They cannot undergo E₁ reaction because of the difficulty of forming primary carbocations.
 - E₂ elimination is stereospecific,
 - The E₂ reaction is the most effective for the synthesis of alkenes from primary alkyl halides.
 - In an E₂ mechanism which refers to bimolecular elimination is basically a one-step mechanism.
 - Here, the carbon-hydrogen and carbon-halogen bonds mostly break off to form a new double bond.
 - However, in the E₂ mechanism, a base is part of the rate-determining step and it has a huge influence on the mechanism.
 - The reaction rate is mostly proportional to the concentrations of both the eliminating agent and the substrate.
 - It exhibits second-order kinetics.
 - The E₂ mechanism can generally be represented as below. In the below-mentioned representation, B stands for base and X stands for halogen.

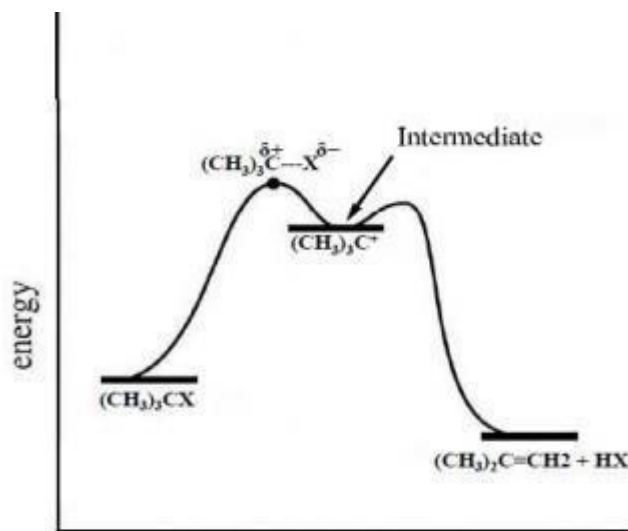
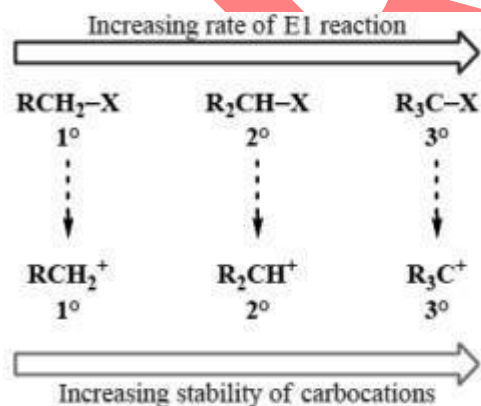


E1 Reaction influences the reaction pathway:

- E1 mechanistic pathway is most common with:
 - ✓ Good leaving groups
 - ✓ Stable carbocations
 - ✓ Weak bases.
- E1 Reactions is Non-stereospecific- follows Zaitsev (Saytseff) Rule -Does NOT occur with primary alkyl halides (leaving groups)

1. Effect of R-

- Reactivity order: $(\text{CH}_3)_3\text{C}^- > (\text{CH}_3)_2\text{CH}^- > \text{CH}_3\text{CH}_2^- > \text{CH}_3^-$
- In an E1 reaction, the rate determining step is the loss of the leaving group to form the intermediate carbocation.
- The more stable the carbocation is, the easier it is to form, and the faster the E1 reaction.
- The rate of an E1 reaction increases as the number of R groups on the carbon with the leaving group increases.



Energy Profile for an E₁ Reaction

2. Leaving Group (LG)

- The only event in the rate determining step of the E1 is breaking the C- LG bond. Therefore, there is a very strong dependence on the nature of the leaving group, the better the leaving group, the faster the E1 reaction will be.
- In the acid catalysed reactions of alcohols, the -OH is protonated first to give an oxonium ion, providing the much better leaving group, a water molecule.

3. Base (B)

- Since the base is not involved in the rate determining step, the nature of the base is unimportant in an E1 reaction.
- Favored by weaker bases such as H₂O and ROH.

4. Type of Solvent

- Favored by polar protic solvents, which can stabilize the ionic intermediates.

E2 Reaction influence the reaction pathway:

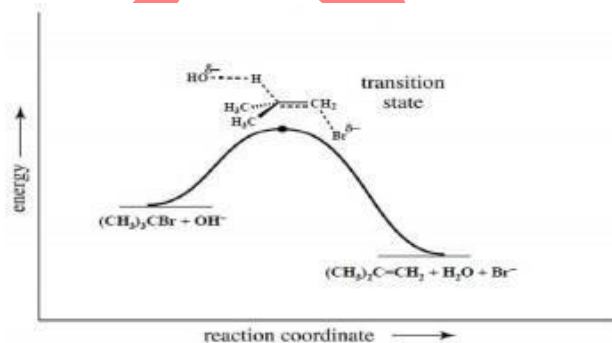
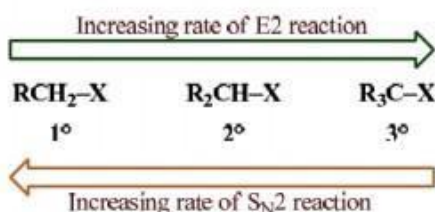
- Kinetics – Second order
- Mechanism – Single step
- Stereospecific (Anti-periplanar geometry preferred, Syn-periplanar geometry possible)
- Concerted - all bonds form and break at same time.
- Bimolecular - rate depends on concentration of both base and substrate - Favoured by strong bases.
- Favored by polar aprotic solvents.
- Better leaving group leads to faster reaction rates.

Identity of R group –

More substituted halides react faster Rate: R₃CX > R₂CHX > RCH₂X

(i) Effects of R-

- ✓ Reactivity order: $(\text{CH}_3)_3\text{C}- > (\text{CH}_3)_2\text{CH}- > \text{CH}_3\text{CH}_2- > \text{CH}_3-$
- ✓ In an E2 reaction, the reaction transforms 2 sp^3 C atoms into sp^2 C atoms. This moves the substituents further apart decreasing any steric interactions.
- ✓ So more highly substituted systems undergo E2 eliminations more rapidly. This is the same reactivity trend as seen in E1 reactions.
- ✓ As the number of R groups on the carbon with the leaving group increases, the rate of the E2 reaction increases.

**Energy Profile for an E₂ Reaction****(ii) Leaving Group (LG)**

The C-LG bond is broken during the rate determining step, so the rate does depend on the nature of the leaving group.

However, if a leaving group is too good, then an E1 reaction may result. -Rate of reaction follows the order Rate of reaction follows the order, **R-I**

**(iii) Base (B)**

Stronger bases favor the reaction. Since the base is involved in the rate determining step, the nature of the base is very important in an E2 reaction.

More reactive bases will favor an E2 reaction.

Sl.No	Characteristics of E1 reaction	Characteristics of E2 reaction
1.	Unimolecular reaction	Bimolecular reaction
2.	Two step reaction	Single step reaction
3.	Carbocation intermediate formed	Hydrogen removes from beta carbon.
4.	Reactivity order of RX is $3^\circ > 2^\circ > 1^\circ$	Trans elimination because low energy Consumption.
5.	No stereospecific.	Anti periplanar attack
6.	Follow ziatsev rule	Polar aprotic solvent best
7.	Polar protic solvent good because stabilized ionic intermediate.	Phenyl group influence elimination because product alkene further stabilised by resonance.
8.	Rate of reaction increases when concentration of substrate increases.	Reactivity order $3^\circ > 2^\circ > 1^\circ$. No steric effect
9.	Rearrangement may take place.	Strong nucleophile influence elimination
10.		No intermediate formed.

5 MARKS QUESTIONS

1. Explain E1 & E2 reactions with mechanism?
2. Write the factors effecting E1 & E2 reactions?
3. Write the differences between E1 & E2 reactions?

10 MARKS QUESTION

1. Explain E1 & E2 reactions with mechanism and explain the factors effecting E1 & E2 reactions?
2. Explain E1 & E2 reactions with mechanism and write the differences between E1 & E2 reactions?