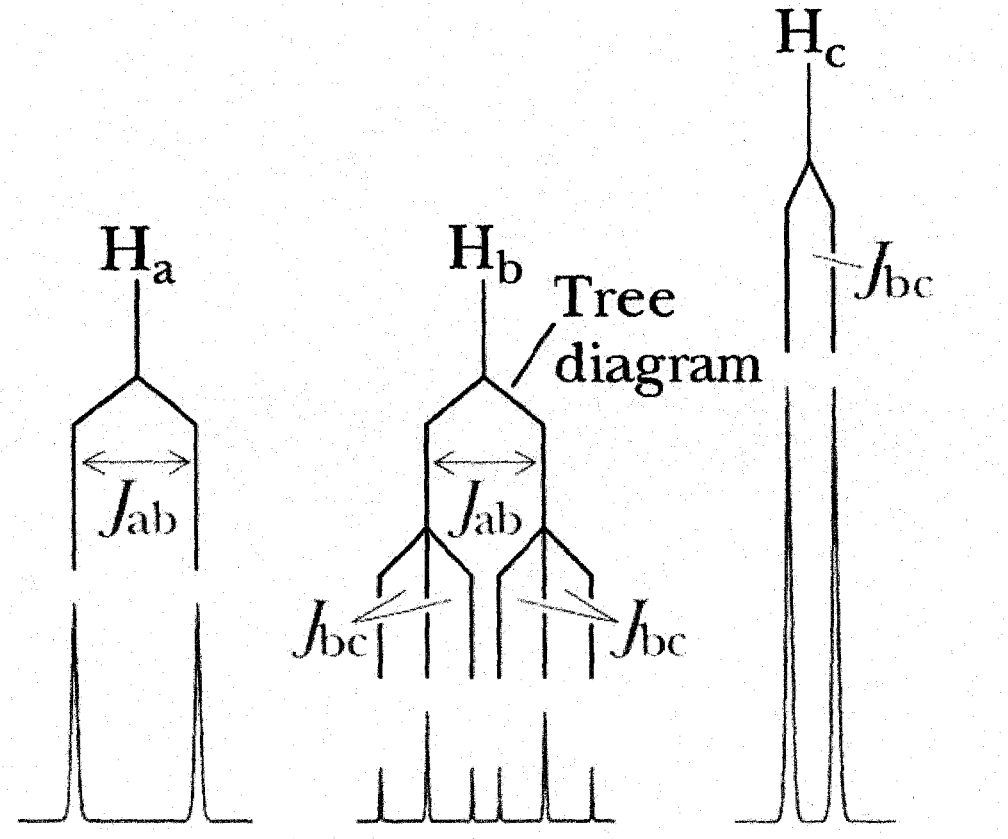
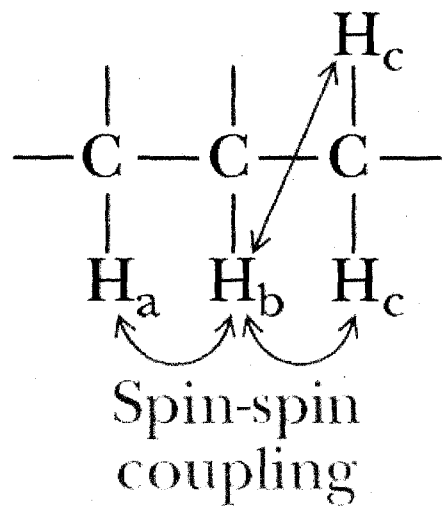


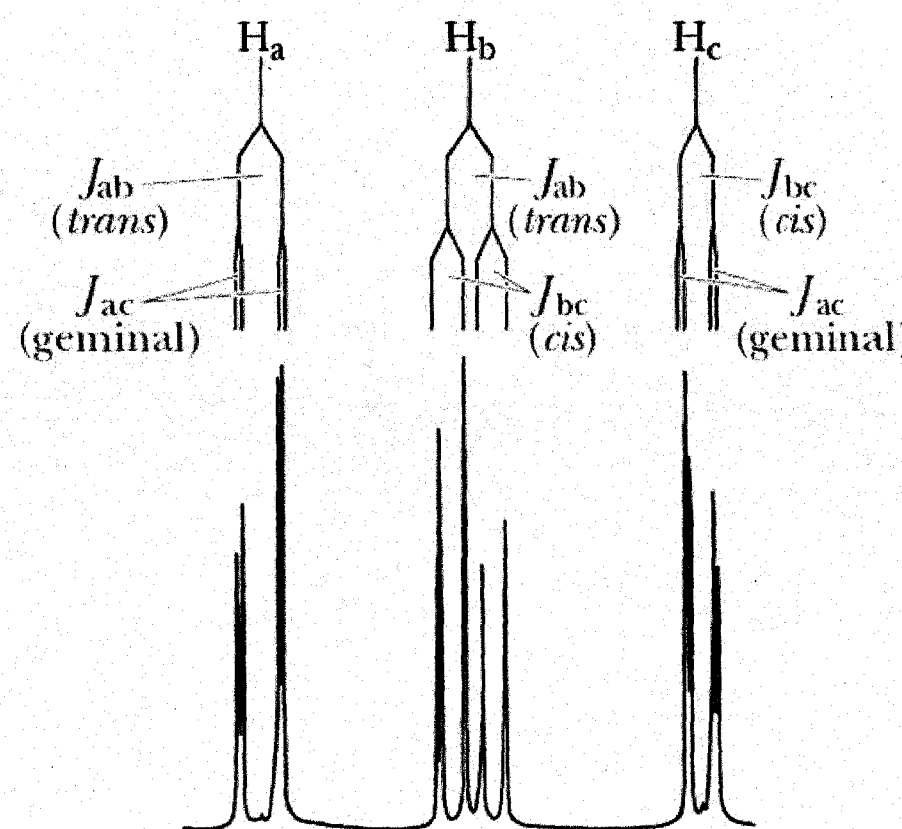
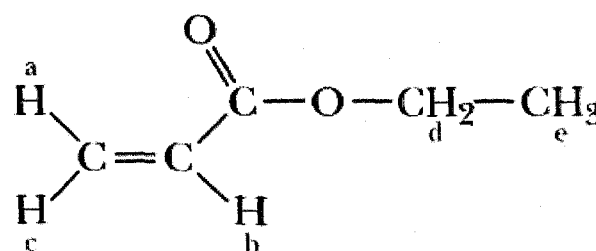
More Complex Splitting Patterns

- if H_c is a set of two equivalent H, then the observed splitting is a doublet of triplets



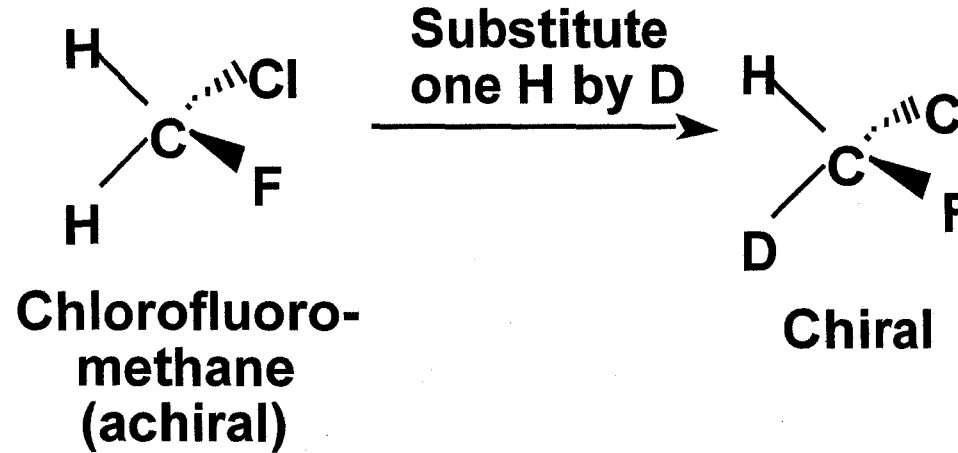
More Complex Splitting Patterns

- a tree diagram for the complex coupling of the three vinylic hydrogens in ethyl propenoate



Stereochemistry & Topicity

◆ Enantiotopic groups



Substitution produces a stereocenter; therefore, hydrogens are enantiotopic. Both hydrogens are prochiral; one is pro-R-chiral, the other is pro-S-chiral.

- enantiotopic atoms or groups have identical chemical shifts in achiral environments
- they have different chemical shifts in chiral environments



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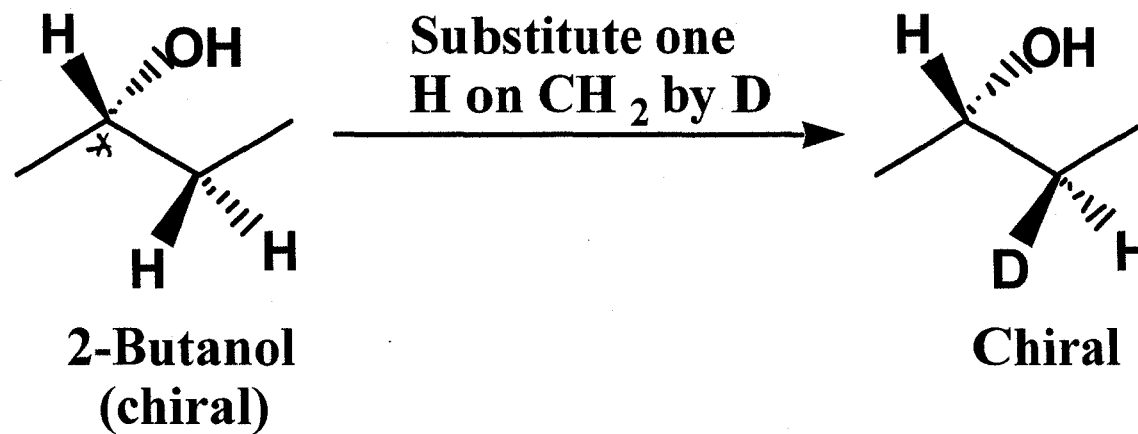


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Stereochemistry & Topicity

◆ Diastereotopic groups

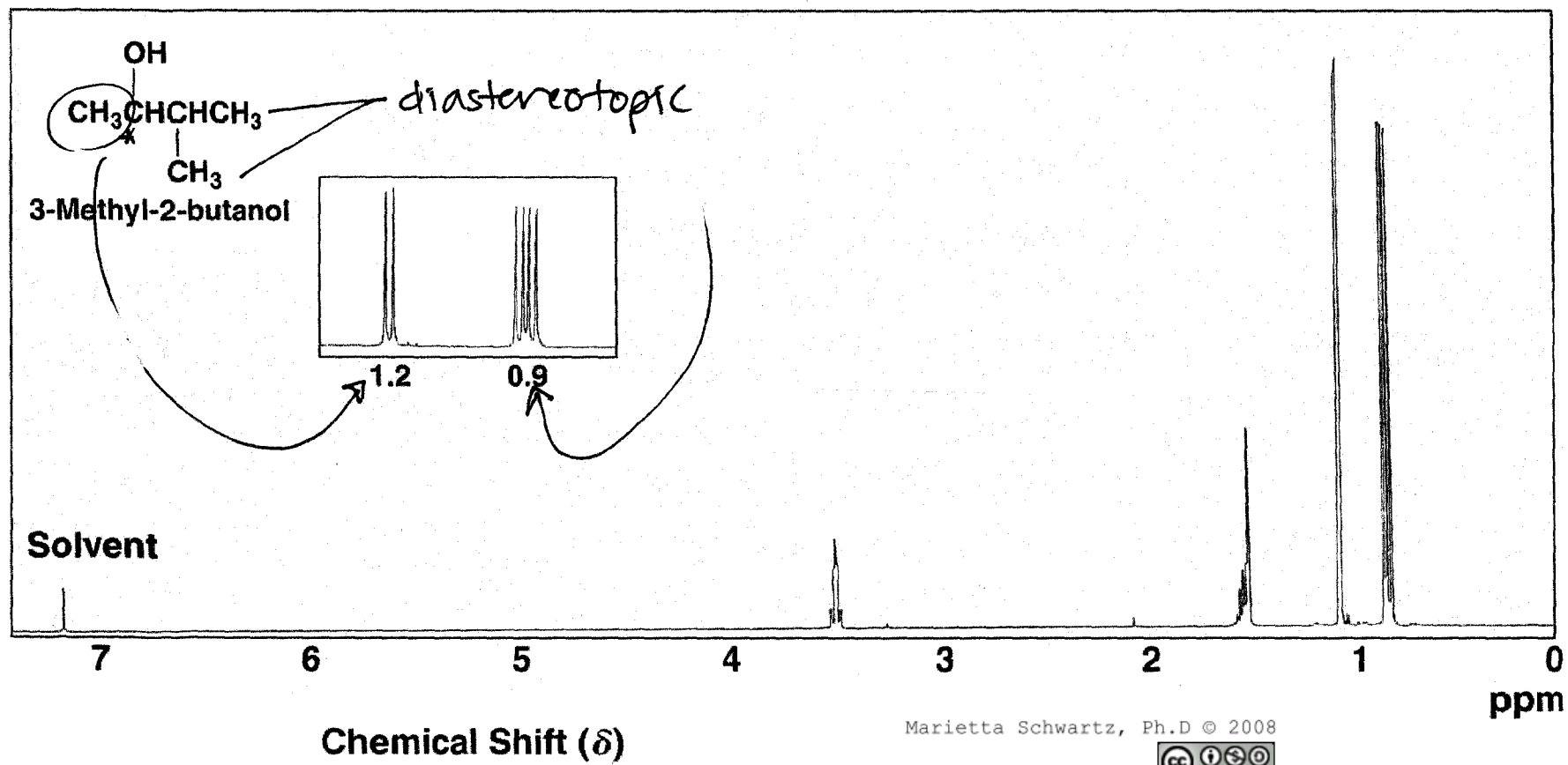
- H atoms on C-3 of 2-butanol are diastereotopic
- substitution by deuterium creates a chiral center
- because there is already a chiral center in the molecule, diastereomers are now possible



- diastereotopic hydrogens have different chemical shifts under all conditions

Stereochemistry and Topicity

- ◆ $^1\text{H-NMR}$ spectrum of 3-methyl-2-butanol
 - the methyl groups on carbon 3 are diastereotopic and appear as two doublets



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Ch. 10 Free Radicals

- a reactive intermediate w/ an unpaired e'
- formed by homolytic bond cleavage - (not heterolytic)

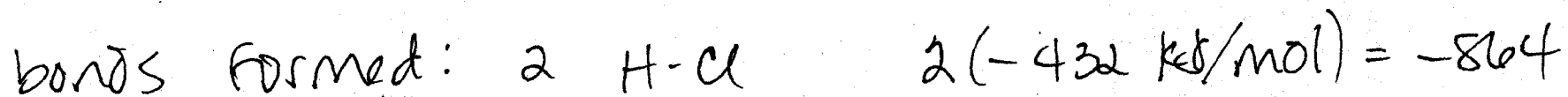
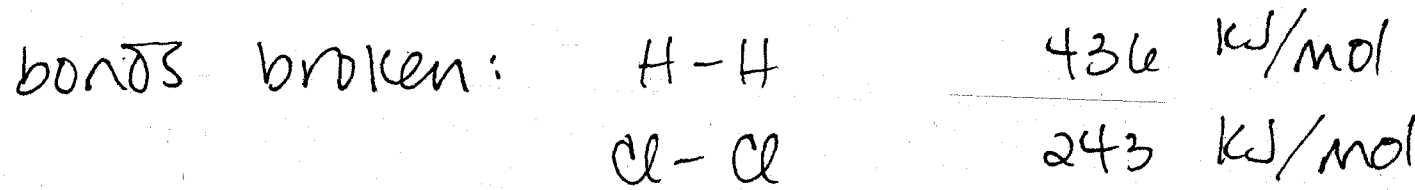
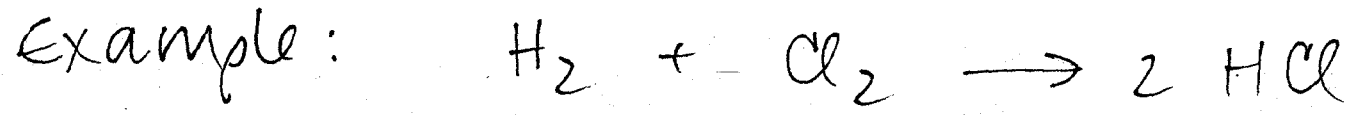
ΔH° - amt of energy required to break a bond
homolytically - table 10.1 p.430

numbers are all \oplus - require energy
(endothermic)

if you are going to make a bond - it's
the same number, just change the sign.

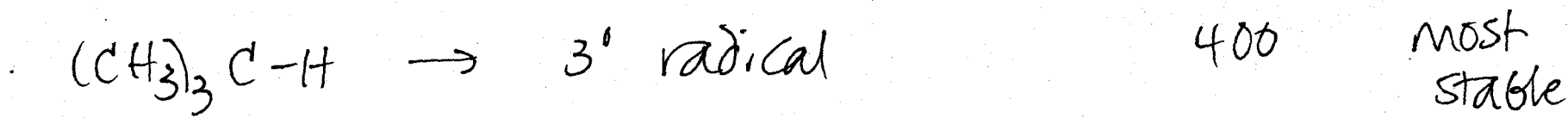
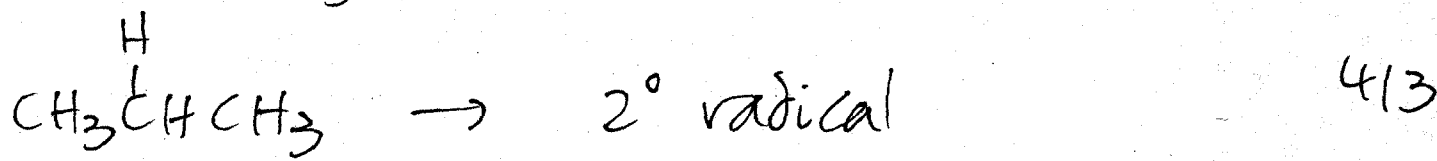
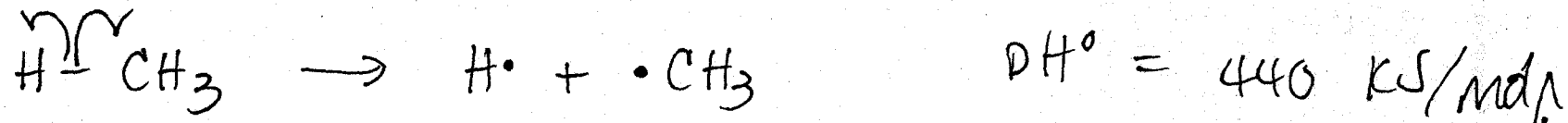
generally \ominus - energy is ~~required~~
(exothermic) produced

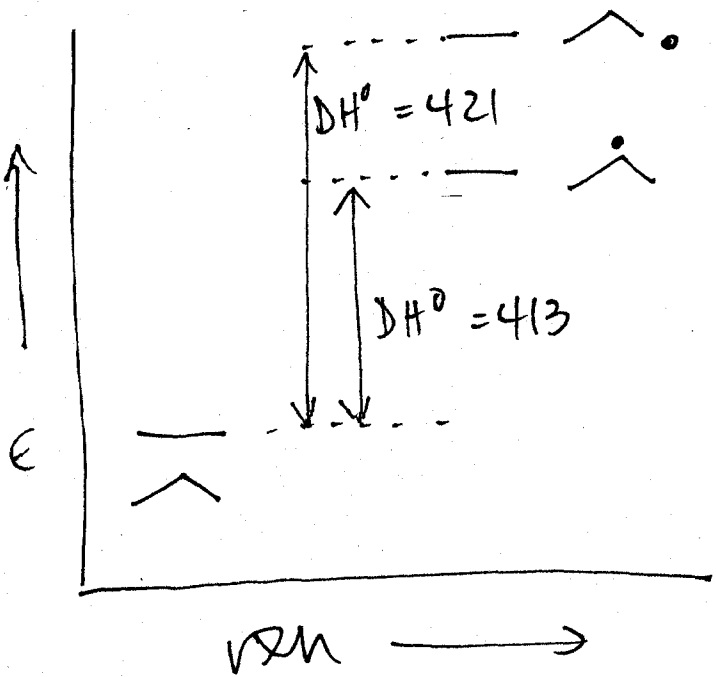
can use these ΔH° values to calculate ΔH°
for the rxn.



$$\Delta H^\circ = -185 \text{ kJ/mol}$$

These #s can help us predict relative stabilities of carbon-based free radicals.

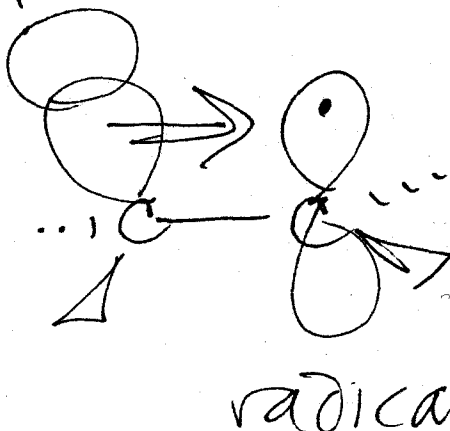
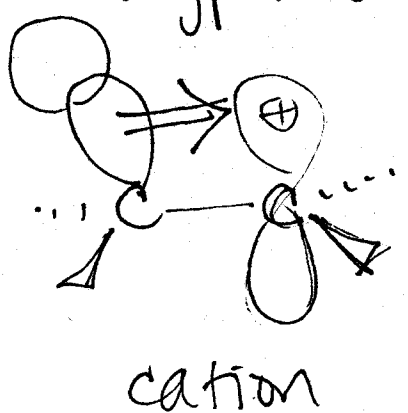




the lower the amount of energy required to break the bond, the more stable the radical (relatively speaking)

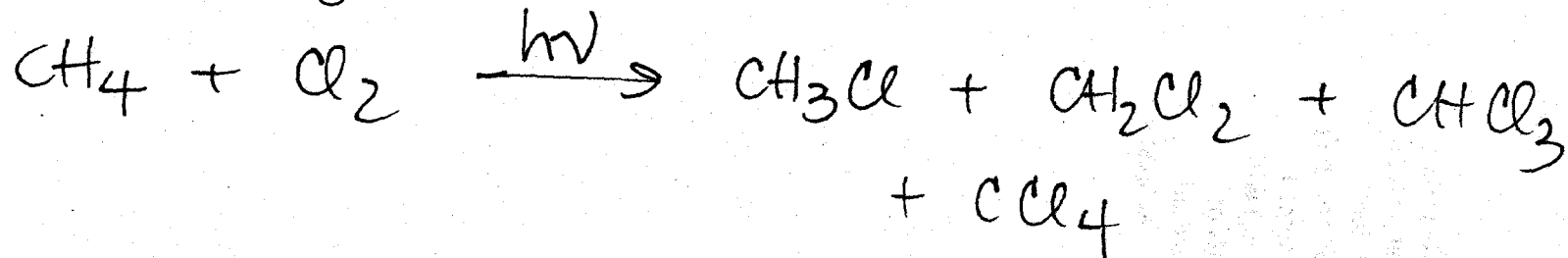
The order of stability for radicals is the same as for cations - rationale is the same.

- stabilized by alkyl substitution
- hyperconjugation



radicals are planar
 sp^2
 (get racemic products)

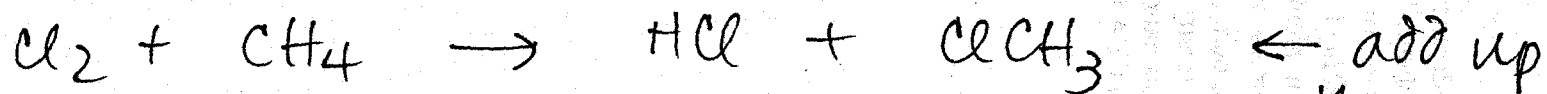
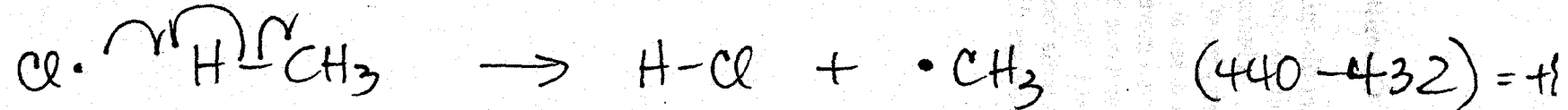
Radical Halogenation of Alkanes



1. Initiation - radicals are formed as products from non-radical reactants.



2. Propagation - radicals on both sides.

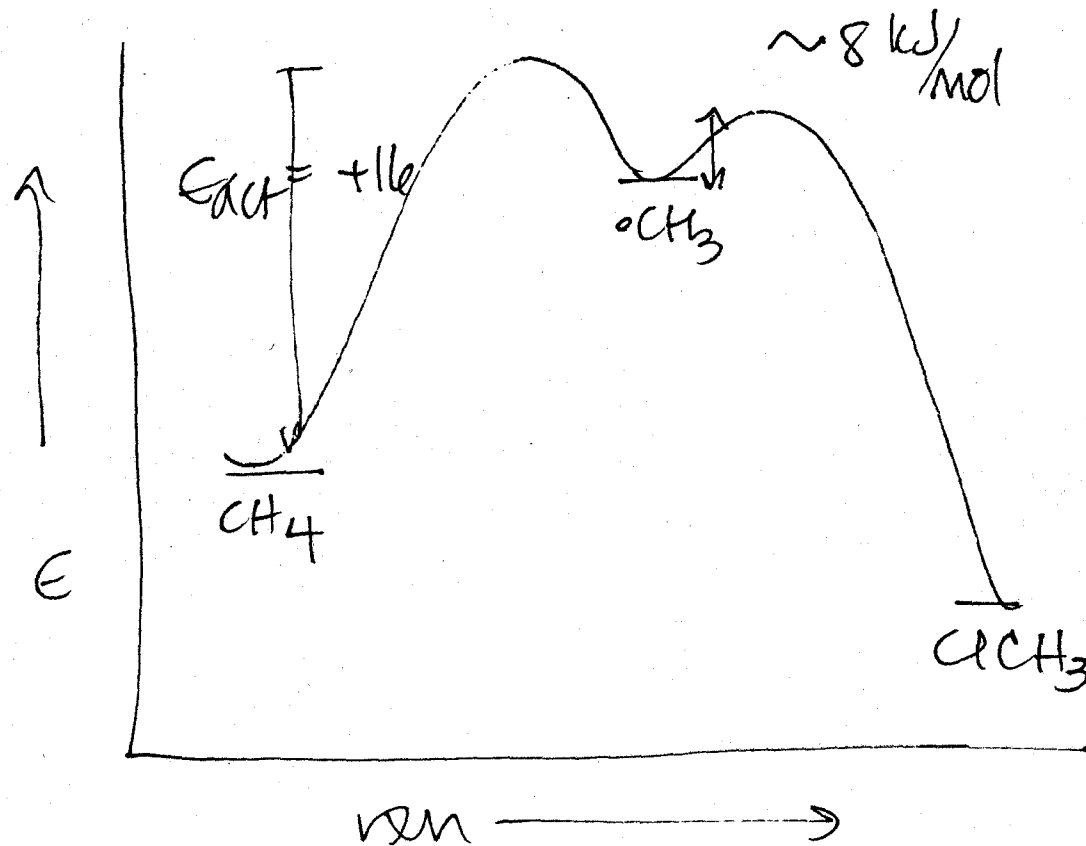
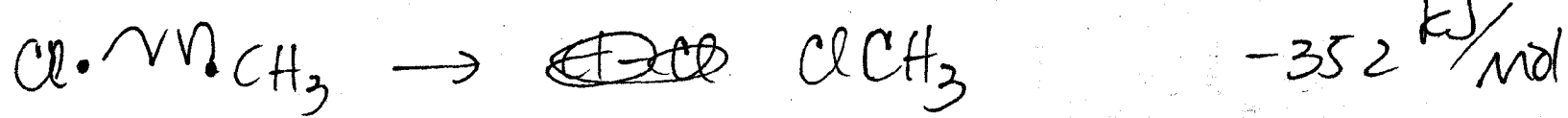


$$+8 - 109$$

$$= -101 \text{ kJ/mol}$$

rxn

3. Termination - radicals as reactants but not products - lots of possibilities



Also
need

