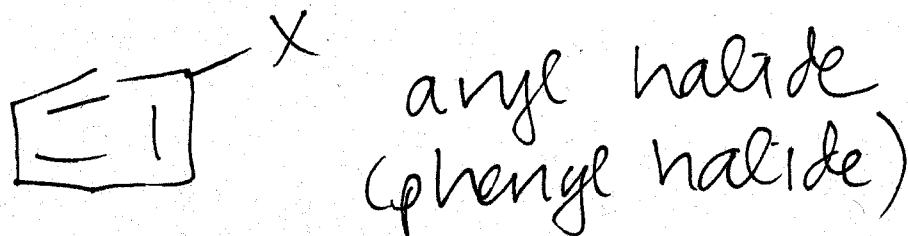
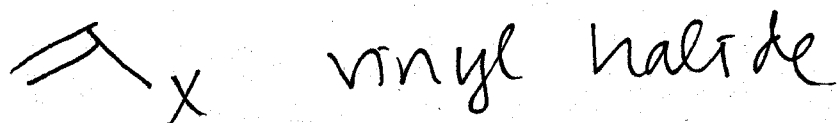


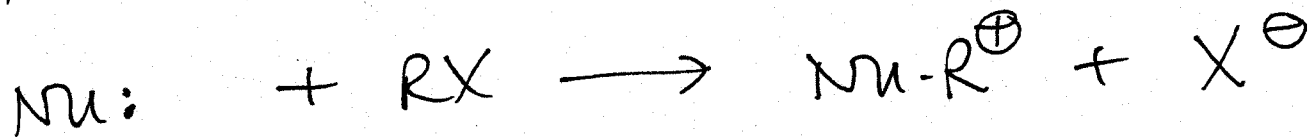
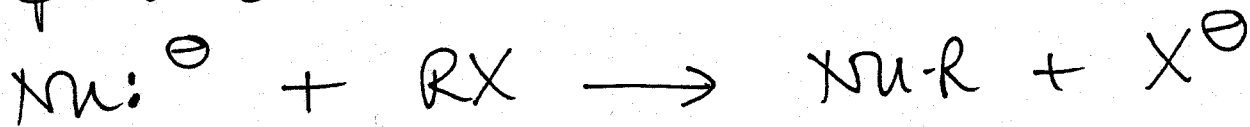
# Ch. 6 - Ionic RXNS - Alkyl Halides

an alkyl halide is a halogen on an  $sp^3$  carbon.



Not these.

Nucleophilic Substitution RXNS.



## Questions:

- \* What is the mechanism?
- \* Does the C-Nu bond form 1st, and then the C-X bond breaks? or is it simultaneous? Or the other way around?
- \* What makes a good nucleophile?
- \* Does the size/nature of R matter?
- \* Does X matter?
- \* Can we use something besides X?

## Definitions:

Nucleophile - nucleus loving - basically, anything with a lone pair. Can have  $\ominus$  charge but don't have to.

Electrophile - electron loving - e<sup>-</sup> deficient. Often cations; always at least partial  $\oplus$

Leaving Group - the conjugate base of a strong acid. Able to handle a neg. charge w/o difficulty. Halides are good L.G.'s - remember. HCl acid;  $\text{Cl}^-$  conj. base.

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sample Rxn:  $\text{CH}_3\text{Cl} + \text{NaOH} \rightarrow \text{CH}_3\text{OH} + \text{NaCl}$

<u>Experiment</u>	<u>[NaOH]</u>	<u>[CH<sub>3</sub>Cl]</u>	<u>Relative Rate</u>
1	1	1	1
2	1	2	2
3	2	1	2
4	2	2	4

$$\text{Rate} = k [\text{OH}^-] [\text{CH}_3\text{Cl}]$$

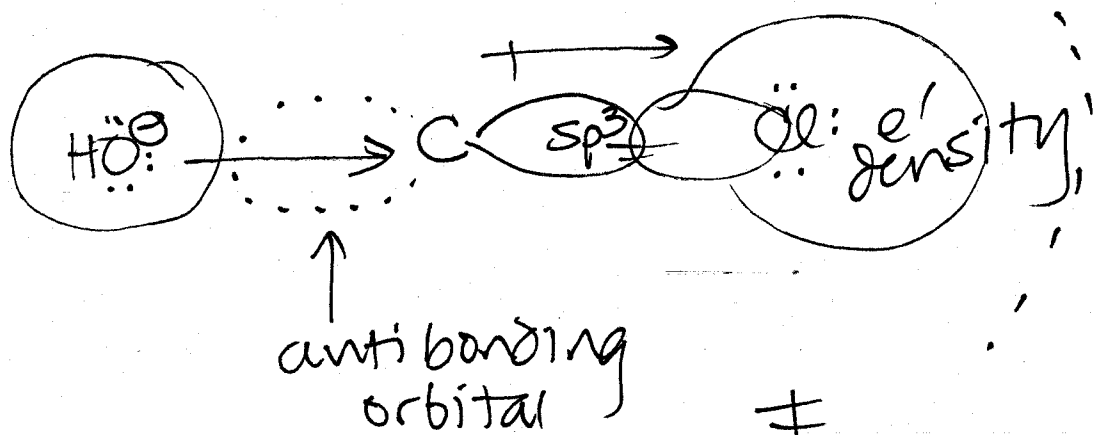
2<sup>nd</sup> order  
kinetics

2<sup>nd</sup> order rxn means both species are involved in rate-det. step. (bimolecular)

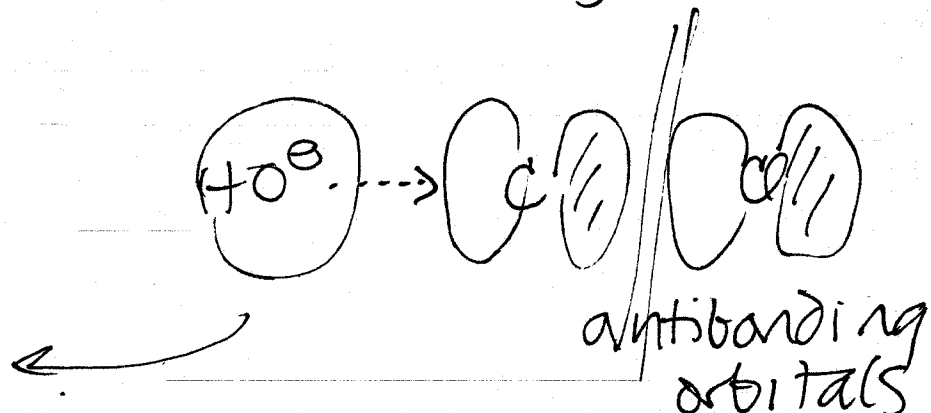
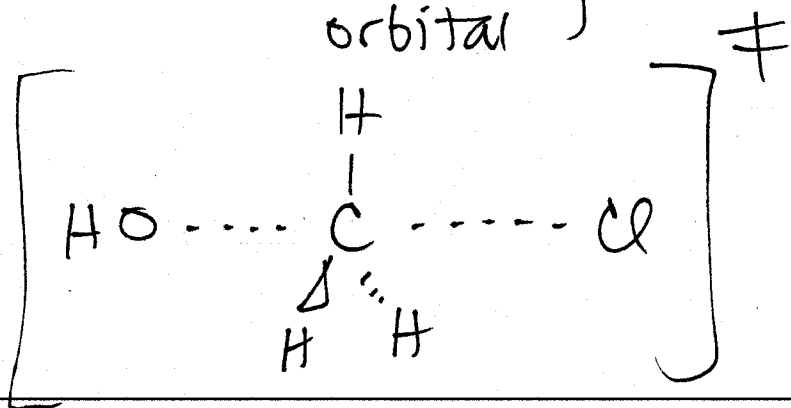
THIS IS AN S<sub>N</sub>2 RXN.

substitution, nucleophilic, bimolecular

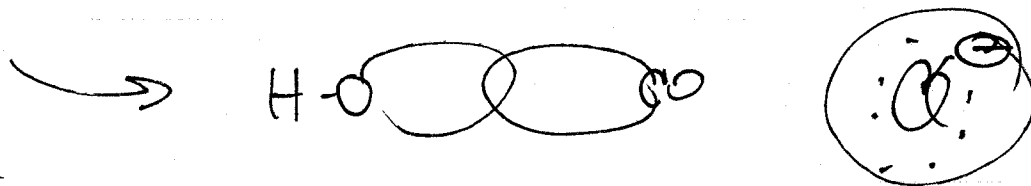
consider the orbitals on the carbon.



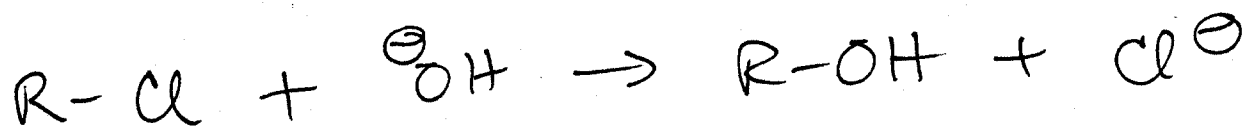
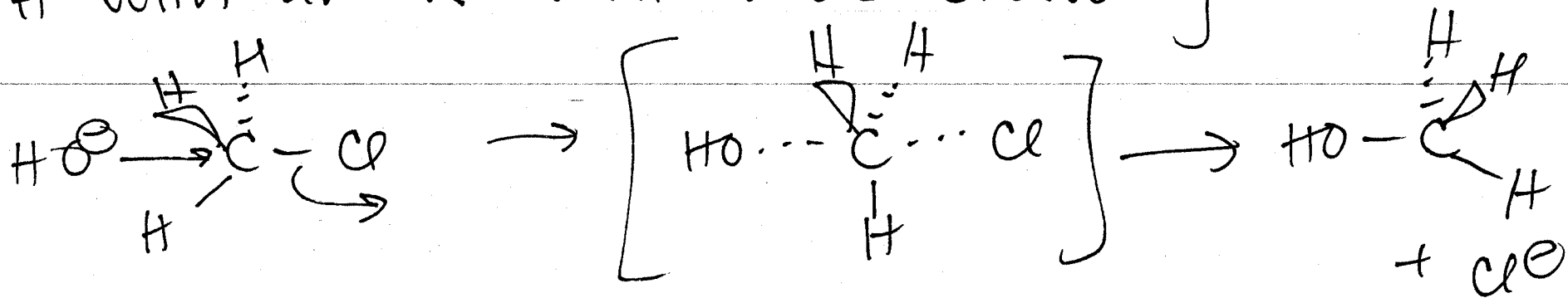
It won't attack from this side - too many e's repelling it.



penta coordinate transition state

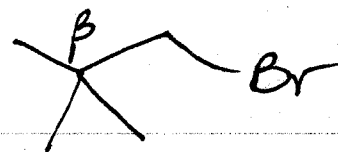
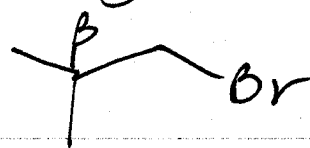
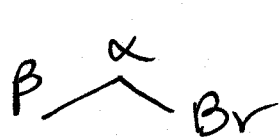


steric effects - look at TS. - replacing a H with an R will cause crowding.



R:	CH <sub>3</sub> -	Et-	Pr-	iPr-	tBu-
Nature of R:	methyl	1°	1°	2°	3°
Relative rate:	100	1.31	0.81	0.015	0.004 (essentially no reaction)

more sterics:  $\beta$ -branching



rate:

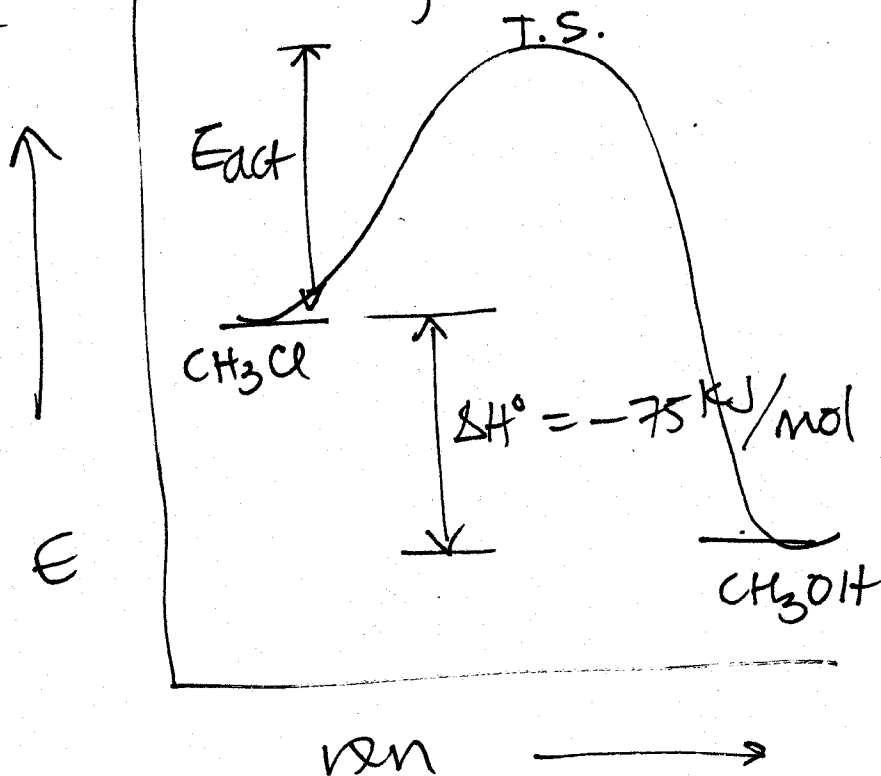
1

0.8

0.003

$1.3 \times 10^{-5}$   
(NRX)

(still looking at  $\text{CH}_3\text{Cl} + \text{NaOH}$ )



THIS rxn has  
 $\Delta G^\circ = -100 \text{ kcal/mol}$   
 calc.  $K_{eq} = 5 \times 10^{15}$

$E_{act}$  = activation energy

generally if  
 $E_{act} < 84 \text{ kJ/mol}$   
 the rxn will go.