

Carbonyl Alpha Chem. II

Alkylation Reactions

- malonic ester synthesis
- acetoacetic ester synthesis
- “direct alkylation”

Ref (17: 7C; 19: 3 - 6)

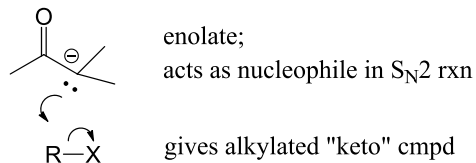
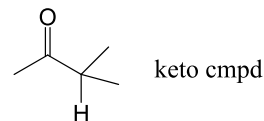
Prob 19: 8, 12, 15

Adv Rdg 17: 4 – 7, 9b; 19: 9

α Alkylation

(replace H by alkyl group at α posⁿ of “Keto”)

General:



$R-X$ gives alkylated "keto" cmpd

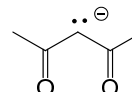
where $R-X$ = 1° halide, tosylate, ...

(incl. allylic, benzylic)

“keto” works best (easiest)

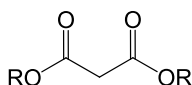
if α C is doubly activated

i.e., β – dicarbonyl cmpds, such as

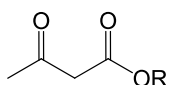


α Alkylation

esp.



malonic ester



acetoacetic ester

can form enolates easily,

with “weak bases”, such as EtO^-

(consult pKa Tables, if needed)

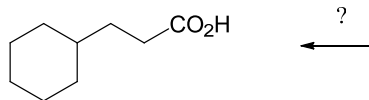
1.) Malonic Ester Synthesis

Ex. Mech.:

malonic ...

N.B. Adds $-\text{CH}_2\text{CO}_2\text{H}$ (2 C's)
to "1° substrate"

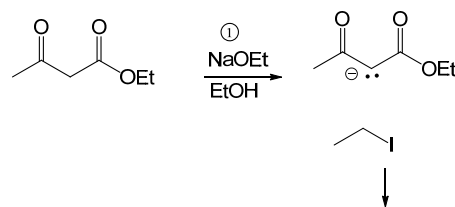
Practice: Retrosynthetic Analysis



Ans.

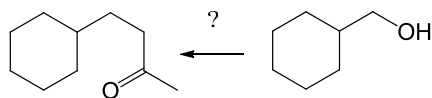
Do details as HMWK !

2.) Acetoacetic Ester Synthesis



N.B. 3 C's added to substrate

Practice



Ans.

Do details as HMWK!

3.) "Direct Alkylation"

(modern development)"1° substrate" + enolate of monoketo cmpd

→ "alkylated keto cmpd"

Ex.

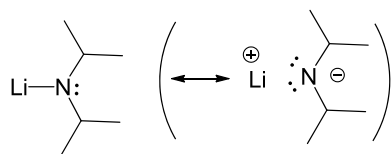
Problems

- need strong base (OR^- too weak) to make enolate
- side rxns (e.g. aldol) possible
- amide (e.g., NH_2^-) strong enough, but could attack carbonyl C (imine formⁿ)
- in ketones, 2 different α positions exist and could react

direct ...

Solution

Use LDA, lithium diisopropylamide, as base



- is very strong base (pKa of amine ≈ 40)
- “bulky”: does not form imine/enamine,
forms enolate at less substituted α C

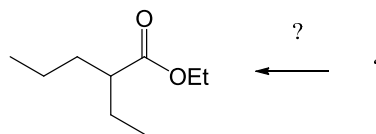
Suitable Conditions

1.) treat “keto cmpd” at -78°C

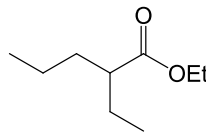
w/ 1 equiv. of LDA / THF

(makes enolate quantitatively, irreversibly, selectively
at less substituted C)

2.) Add substrate (“halide”) and allow $T \uparrow$ to r.t.

Practice

Ans.: α substituted ester; try "direct alkylation"



Potential Alternative:

