For alkylation reactions of enolate anions to be useful, these intermediates must be generated in high concentration in the absence of other strong nucleophiles and bases. The aqueous base conditions used for the aldol condensation are not suitable because the enolate anions of simple carbonyl compounds are formed in very low concentration, and hydroxide or alkoxide bases induce competing S_N2 and E2 reactions of alkyl halides. It is necessary, therefore, to achieve complete conversion of aldehyde or ketone reactants to their enolate conjugate bases by treatment with a very strong base (pK_a > 25) in a non-hydroxylic solvent before any alkyl halides are added to the reaction system. Some bases that have been used for enolate anion formation are: NaH (sodium hydride, pK_a > 45), NaNH_2 (sodium amide, pK_a = 34), and Li[N(CH(CH_3)_2)_2] (lithium diisopropylamide, LDA, pK_a 36). Ether solvents like tetrahydrofuran (THF) are commonly used for enolate anion formation. With the exception of sodium hydride and sodium amide, most of these bases are soluble in THF. Certain other strong bases, such as alkyl lithium and Grignard reagents, cannot be used to make enolate anions because they rapidly and irreversibly add to carbonyl groups. Nevertheless, these very strong bases are useful in making soluble amide bases. In the preparation of lithium diisopropylamide (LDA), for example, the only other product is the gaseous alkane butane.

Because of its solubility in THF, LDA is a widely used base for enolate anion formation. In this application, one equivalent of diisopropylamine is produced along with the lithium enolate, but this normally does not interfere with the enolate reactions and is easily removed from the products by washing with aqueous acid. Although the reaction of carbonyl compounds with sodium hydride is heterogeneous and slow, sodium enolates are formed with the loss of hydrogen, and no other organic compounds are produced.

The presence of these overlapping p orbitals gives \(\alpha\) hydrogens (Hydrogens on carbons adjacent to carbonyls) special properties. In particular, \(\alpha\) hydrogens are weakly acidic because the conjugate base, called an enolate, is stabilized though conjugation with the \(\pi\) orbitals of the carbonyl. The effect of the carbonyl is seen when comparing the pK_a for the \(\alpha\) hydrogens of aldehydes (~16-18), ketones (~19-21), and esters (~23-25) to the pK_a of an alkane (~50).

\[
\begin{align*}
\text{C}_2\text{H}_5\text{Li} & + [\text{C}_2\text{H}_5\text{CH}_2\text{NH}] & \rightarrow & [\text{C}_2\text{H}_5\text{CH}_2\text{NH}^\ominus] & + & \text{C}_3\text{H}_7
\end{align*}
\]

\[\text{Butyl lithium} \quad \text{Diisopropylamine} \quad \text{LDA} \]

\[\begin{align*}
\text{Li}^+ & \quad \text{Na}^+ \\
\text{H}_3\text{C} & \quad \text{H} \\
\text{C} & \quad \text{N} \\
\text{CH}_3 & \quad \text{H} \\
\text{CH}_3 & \quad \text{H}
\end{align*}\]

\[\text{LDA} \quad \text{Sodium Amide}\]
Of the two resonance structures of the enolate ion the one which places the negative charge on the oxygen is the most stable. This is because the negative charge will be better stabilized by the greater electronegativity of the oxygen.

### Acidity of α-Hydrogens in Some Activated Compounds

<table>
<thead>
<tr>
<th>Compound</th>
<th>RCH$_2$–NO$_2$</th>
<th>RCH$_2$–COR</th>
<th>RCH$_2$–C≡N</th>
<th>RCH$_2$–SO$_2$R</th>
</tr>
</thead>
<tbody>
<tr>
<td>pK$_a$</td>
<td>9</td>
<td>20</td>
<td>25</td>
<td>25</td>
</tr>
</tbody>
</table>

### Examples

\[
\text{H}_3\text{C} = \text{C} - \text{H} + \text{NaNH}_2 \rightarrow \text{Na}^{+} + \text{H}^{+} \text{NH}_2^{-}
\]
If the formed enolate is stabilized by more than one carbonyl it is possible to use a weaker base such as sodium ethoxide.

\[ \text{NaOCH}_2\text{CH}_3 = \text{Na}^+ \cdot \text{OCH}_2\text{CH}_3 = \text{NaOEt} \]

Because of the acidity of \( \alpha \) hydrogens, carbonyls undergo keto-enol tautomerism. Tautomers are rapidly interconverted constitutional isomers, usually distinguished by a different bonding location for a labile hydrogen atom and a differently located double bond. The equilibrium between tautomers is not only rapid under normal conditions, but it often strongly favors one of the isomers (acetone, for example, is 99.999% keto tautomer). Even in such one-sided equilibria, evidence for the presence of the minor tautomer comes from the chemical behavior of the compound. Tautomeric equilibria are catalyzed by traces of acids or bases that are generally present in most chemical samples.

### General Reaction of Enolates

The Ambident Character of Enolate Anions

Since the negative charge of an enolate anion is delocalized over the alpha-carbon and the oxygen, as shown earlier, electrophiles may bond to either atom. Reactants having two or more reactive sites are called **ambident**, so this term is properly applied to enolate anions. Modestly electrophilic reactants such as alkyl halides are not sufficiently reactive to combine with neutral enol tautomers, but the increased nucleophilicity of the enolate anion conjugate base permits such reactions to take place. Because alkylations are usually irreversible, their products should reflect the inherent (kinetic) reactivity of the different nucleophilic sites.
If an alkyl halide undergoes an S\textsubscript{N}2 reaction at the carbon atom of an enolate anion the product is an alkylated aldehyde or ketone. On the other hand, if the S\textsubscript{N}2 reaction occurs at oxygen the product is an ether derivative of the enol tautomer; such compounds are stable in the absence of acid and may be isolated and characterized. These alkylations (shown above) are irreversible under the conditions normally used for S\textsubscript{N}2 reactions, so the product composition should provide a measure of the relative rates of substitution at carbon versus oxygen. It has been found that this competition is sensitive to a number of factors, including negative charge density, solvation, cation coordination and product stability.

**Contributors**

- Prof. Steven Farmer (Sonoma State University)

**Enolate of Unsymmetrical Carbonyl Compounds**

Now let’s consider what happens when an unsymmetrical carbonyl is treated with a base. In the case displayed below there are two possible enolates which can form. The removal of the 2o hydrogen forms the kinetic enolate and is formed faster because it is less substituted and thereby less sterically hindered. The removal of the 3o hydrogen forms the thermodynamic enolate which is more stable because it is more substituted.

**Kinetic Enolates**

Kinetic enolates are formed when a strong bulky base like LDA is used. The bulky base finds the 2o hydrogen less sterically hindered and preferable removes it.

Low temperature are typically used when forming the kinetic enolate to prevent equilibration to the more stable
thermodynamic enolate. Typically a temperature of -78 °C is used.

**Thermodynamic Enolates**

The thermodynamic enolate is favored by conditions which allow for equilibration. The thermodynamic enolate is usually formed by using a strong base at room temperature. At equilibrium the lower energy of the thermodynamic enolate is preferred, so that the more stable, more substituted enolate is formed.

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