SYNTHESIS OF ALKENES BY OXIDATIVE DECARBOXYLATION OF CARBOXYLIC ACIDS; MECHANISTIC VIEWS; THE ORGANIC CHEMISTRY NOTEBOOK SERIES, A DIDACTICAL APPROACH, Nº 6

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ABSTRACT

This is the sixth chapter in the series published by the same authors: “The Organic Chemistry Notebook Series, a Didactical Approach” [1-5].

Here we offer the mechanistic views of the synthesis of alkenes by oxidative decarboxylation of carboxylic acids. The aim of this series of studies is to help students to have a graphical view of organic synthesis reactions of diverse nature. The oxidative decarboxylation of carboxylic acids is a useful method for generating alkenes. Here we propose the mechanism and its discussion for the application of the method of decarboxylation of diacids lacking nearby double bonds. Also, the route is explained mechanistically for the preparation of Dewar benzene. The thermal or photolytic decomposition of di-t-butyl per-esters is described. The treatment of monocarboxylic acids to afford alkenes in the presence of lead tetraacetate and copperII acetate is briefly discussed. The alkylation-decarboxylation of aromatic acids is also explained.

The oxidative decarboxylation of carboxylic acids can eventually conduct to the obtaining of ketones instead of alkenes. We have used a series of reactions reviewed by W. Carruthers, and we have proposed didactical and mechanistic views for them. This latest approach is included in the synthetic methods reviewed by W. Carruthers with respect to the “Formation of carbon-carbon double bonds”. Spanish title: Síntesis de alquenos por descarboxilación de ácidos carboxílicos; vistas mecanísticas; De la serie: El cuaderno de notas de química orgánica, un enfoque didáctico, Nº6.

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MECHANISTIC PROPOSALS AND DISCUSSION

Decarboxylation of vicinal diacids lacking nearby double bonds

As academics we are concerned with the didactical importance of covering the needs of debutant students in organic synthesis. This is the sixth study in: “The Organic Chemistry Notebook Series, a Didactical Approach” [1-5]. The method of decarboxylating carboxylic acids has proven to be useful [6]. Variations on the basic steps have been introduced essentially for the improvement of the yields [6]. The aim is the generation of double C-C bonds starting with vicinal dicarboxylic acids and subsequent decarboxylation, as schematically viewed in Fig. 1 [6,7].

Figure 1. Vicinal dicarboxylic acid as starting material for generating double C-C bonds; reviewed by W. Carruthers [5]
The treatment with lead tetraacetate in boiling benzene can be improved with the presence of oxygen becoming thus applicable for vicinal dicarboxylic acids lacking nearby double bonds [6] (Fig. 2).

**Figure 2.** Synthesis of 1,4-cyclohexadiene by decarboxylation 4,5-ciclohexendicarboxylic acid, reviewed by W. Carruthers [6]

The mechanism is seen in Fig. 3. In step 1, lead acetate pulls off a proton from one of the carboxylic acid residues of the substrate to form a carboxylate residue. What follows is the immediate excision of the bonding of the substituent carboxylate. This electronic movement evokes somehow the Diels Alder mechanism with respect to the formation of the new double bond. The contrast is the rupture of both sigma bonds instead of the transformation of two double into single bonds in Diels Alder. However, indeed, this is not a pericyclic reaction. Besides cyclohexenone, CO$_2$ and carbanionic carboxylic acid evolve from this fragmentation. The acid-base interaction between the acetic acid formed from the lead tetraacetate and the carbanionic carboxylic acid gives formic acid and tetraacetate is regenerated. In step 2, tetraacetate pulls off a proton from formic acid provoking decarboxylation (generation of CO$_2$) and acetic acid plus 4 hydrides. In step 2, two of the four acetic acid equivalents ionize and generate two acetate and two proton equivalents. These two proton equivalents plus two from the four hydride equivalents of step 2 generate 4 protons and 4 e$^-$ due to oxidation of hydride in step 3. In step 4, Pb$^{IV}$ (Pb$^{4+}$) is reduced to Pb$^0$ thanks to the previous oxidation step that provided four e$^-$. In step 5, two of the four proton equiv generated in step 3 interact with two acetate residues to generate 2 acetic acid equiv. Step 6 is the covalence formation between 2H$^+$ and 2H$^-$ generating H$_2$.

**Figure 3.** Synthesis of 1,4-cyclohexadiene by oxidative decarboxylation of 4,5-ciclohexendicarboxylic acid; mechanistic views

An interesting procedure with good yields is the obtaining of dienes by electrolysis of the dicarboxylic acid in pyridine solution with trimethylamine added. The reaction goes on in mild conditions, and the method is appropriate “for preparing highly strained unsaturated small and bridged ring compounds” [6]. As an example let’s mention the preparation of Dewar benzene from bicyclo[2,2,0]hex-2-en-5,6-dicarboxylic acid, Fig. 4 [6].

**Figure 4.** Dewar benzene obtained by electrolysis in pyridine solution with trimethylamine from bicyclo[2,2,0]hex-2-en-5,6-dicarboxylic acid. Reviewed by W. Carruthers [6]
In the first step an acid reaction occurs with the liberation of a proton from one of the acid residues of bicyclo[2,2,0]hex-2-en-5,6-dicarboxylic acid. Triethylamine serves as temporary base to accommodate this proton but soon after the proton is partially reduced by 1e\(^-\) to H. Also, decarboxylation occurs with the first carboxylate residue to form the alkene and expel the second carboxylic acid residue under the form of carbanion. In the second step, the carbanionic carboxylic acid frees its proton which neutralizes the carbanion to afford formiate. Formiate expels a hydride and generates an equivalent of carbon dioxide. In the third step the hydride oxidizes to give a proton and two electrons, one of which generates a second H species. The H species overlap each other and form gaseous hydrogen. This step regenerates the electron employed in the electrolytic method under review. See Fig. 5 for mechanistic proposal.

$$\begin{align*}
\text{NiC}_2\text{H}_5\quad & \quad \text{O} \quad \text{O} \quad \text{H} \\
\text{H} & \quad \text{O} \quad \text{O} \quad \text{H} + \text{H}^+ & \quad \text{H}^+ + \text{NiC}_2\text{H}_5 \\
\text{H} + \text{H}^+ & \quad \text{O} \quad \text{O} \quad \text{H} + \text{H}^+ & \quad \text{H}^+ + \text{H}^+ + \text{H}_2 \\
\text{H}^+ + 3\text{e}^- & \quad \text{H} & \quad 3\text{e}^- + \text{H}_2 \\
\text{H} & \quad \text{O} \quad \text{O} \quad \text{H} + \text{e}^- & \quad 2\text{CO}_2 + \text{H}_2 + 3\text{e}^- \\
\text{H} & \quad \text{O} \quad \text{O} \quad \text{H} & \quad 2\text{CO}_2 + \text{H}_2
\end{align*}$$

**Final balanced equation:**

$$\begin{align*}
\text{H}_2\text{C} = \text{C} & \quad \text{O} \quad \text{O} \\
\text{H}_2\text{C} = \text{C} & \quad \text{O} \quad \text{O} + 2\text{CO}_2 + \text{H}_2 + 3\text{e}^-
\end{align*}$$

**Figure 5.** Dewar benzene obtained by electrolysis in pyridine solution with trimethylamine from bicyclo[2,2,0]hex-2-en-5,6-dicarboxylic acid. Mechanism

An easy and useful method, done under mild conditions, and activated under thermal or radiation conditions to provoke photolytic decomposition, uses t-butyl peresters, obtained at its turn from diacid chlorides and t-butyl hydroperoxide, Fig. 6 [6].

This photolysis reaction is applicable for synthesizing thermally unstable alkenes [6]. The vicinal dicarboxylic acids employed in this reaction can be obtained by Diels Alder reaction [6]. An alternative is the photosensitised addition of maleic anhydride to dienes or alkenes [6].

**Figure 6.** Photolytic decomposition of t-butyl peresters under thermal or radiation conditions to afford unstable alkenes. Reviewed by W. Carruthers [6]

The obtaining of the t-butyl peresters is effected by nucleophilic substitution using diacid chloride as starting substrate and t-butylhydroperoxide as nucleophile. The outer oxygen of one equivalent of the two t-butylhydroperoxide equivalents acts as nucleophile over one of the acid chloride function of the bicyclo-compound. This forms a mono-t-butylperester derivative protonated. The second step is the substitution on the second acid chloride extreme of the bicyclo-compound to afford a di-t-butylperester. What follows is the irradiation of the di-t-butylperester which provokes homolytic rupture of the weakest bonding in the substrate, namely the peroxide bridge. This fact generates two single radicals (t-butoxy) and a double radical (di-carboxylate radical). This extremely reactive species (the double radical), collapses into another double radical (proto-alkene) and two carbonic anhydride residues. The two single radicals (t-butoxy) prefer to couple each other to generate a t-butoxyperoxide molecule. See Fig. 7 for this mechanism.
The oxidative decarboxylation of mono-carboxylic acids is a related reaction to the previously described reactions, and this is effected under treatment with lead-tetraacetate [6]. The development of the reaction depends on the structure of the acid giving poor yields and mixture of products, this, under normal conditions [6]. However, with the addition of catalytic amounts of copperII acetate the increasing of the yields of alkenes synthesized is significant [6]. The reaction is applied to primary and secondary carboxylic acids, either under photolysis, or under thermolysis, in benzene. Under the review, the initial oxidation of the acid is followed by reduction of PbII to PbI, etc.

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Pb$^{III}$ with the electron provided soon after by the oxidation of Cu$^{I}$ to Cu$^{II}$. The acetate and the Pb$^{III}$ they form an adduct; this adduct absorbs radiation ($h\nu$) and gives rise to R', CO$_2$ and Pb$^{II}$. The alkyl R' becomes an alkene and an H radical. This H radical is oxidized to H$^+$ to provide the electron to reduce Cu$^{II}$ into Cu$^{I}$. The complementary part of the chain propagation is the oxidation of Pb$^{II}$ into Pb$^{IV}$ and Cu$^{I}$ into Cu$^{II}$ and the generation of H$_2$. After this, the cycle reinitiates at the stage of the dissociation of the carboxylic acid. This mechanistic proposal is highly limited and restrained by certain lack of information which is unavailable in the review [6], and could be improved after proper consultations with the correct bibliographic source. Let us remark with respect to Fig. 8 and 9, that those reactions inside the blue and red square in Fig. 8, are mechanistically explained in squares of the corresponding color in Fig. 9.

\[
\begin{align*}
\text{Pb}(\text{AcO})_4 & \rightarrow \text{Pb}^{IV} + 4\text{AcO}^- \\
\text{Cu}(\text{AcO})_2 & \rightarrow \text{Cu}^{II} + 2\text{AcO}^-
\end{align*}
\]

Initiation step

\[
\begin{align*}
h\nu & \rightarrow \text{Pb}^{IV} + \text{H}^+ \\
1e^- & \rightarrow \text{Cu}^{I} + \text{H}^+
\end{align*}
\]

\[
\begin{align*}
\text{Pb}^{IV} + 4\text{AcO}^- & \rightarrow \text{Pb}^{IV} + \text{Pb}(\text{AcO})_4 \\
\text{Cu}^{II} + 2\text{AcO}^- & \rightarrow \text{Cu}^{II} + \text{Cu}(\text{AcO})_2
\end{align*}
\]

\[
\begin{align*}
\text{Pb}^{IV} & \rightarrow \text{Pb}^{IV} + 2\text{AcO}^- \\
\text{Cu}^{II} & \rightarrow \text{Cu}^{II} + 1e^- \\
\text{H}^+ & \rightarrow \text{H}_2
\end{align*}
\]

\[
\begin{align*}
\text{Cu}^{II} + 1e^- & \rightarrow \text{Cu}^{I} + \text{H}^+ \\
\text{H}^+ + 1e^- & \rightarrow \text{H}_2
\end{align*}
\]

Figure 9. Decarboxylation of mono-carboxylic acids with lead tetraacetate; free radicals mechanism. Mechanistic views

As an example of the applications of this reaction let’s cite the alkylation-decarboxylation of aromatic acids. Fig. 10.

Figure 10. Alkylation-decarboxylation of aromatic acids. Reviewed by W. Carruthers [6]
The lithium and ammonia employed provoke somehow delocalization of one of the aromatic π systems until locating the negative charge over the carbonyl oxygen of the aromatic acid. The direct alkylation occurs when the carbanion attacks the alkyl bromide in a SN2 manner [6]. The furan derivative, rosefuran can be obtained following the oxidative radical process of Fig. 10 and 11 [6].

**Figure 11. Alkylation-decarboxylation of aromatic acids. Mechanistic views.**

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These two last mechanisms (Fig. 9 and 11) although presenting some species stoichiometrically unjustified (H\(^+\) and AcO\(^-\)), give at least a global idea of the movement of species as well as the oxidation-reducing processes.

A carboxylic acid can be converted into a ketone instead of an alkene following oxidative decarboxylation. Carbon loss is verified during the process [6]. The acid converted in a dianion suffers sulphenylation under dimethyl disulphide [6]. The sulphenylated acid reacts with \(N\)-chlorosuccinimide and gives a diacetal with loss of carbon anhydride adding sodium bicarbonate [6]. The diacetal goes a hydrolysis to afford the ketone at 0\(^\circ\)C [6]. The acid can be synthesized from cyclohexadiene and acrylic acid; this last is equivalent to the ketene in this homologue of the Diels Alder reaction [6]. See Fig. 12.

Another example is shown in Fig. 12 with the decarboxylation of bicyclo[2.2.2]hept-5-ene-2-carboxylic acid into \((1R,4R)\)-bicyclo[2.2.2]hept-5-en-2-one [6]. The mechanistic views corresponding to Fig. 12 are exposed in Fig. 13 and 14.

![Diagram](http://www.bolivianchemistryjournal.org)
negative charge (two electrons) over the carbanion, delocalize to form the alkene’s double bond and thus to dispatch the mobile \(\pi\)electrons of carbonyl to the carbonyl’s oxygen to generate an alcoxy group. The allylalcoxy (stabilized by the cation Li\(^+\)) moves its \(\pi\)electrons as nucleophile over dimethyl disulphide breaking the inter-sulfur linkage, all provoked by the reformation of the carbonyl group by localization of the negative excessive charge on oxygen. The \(\alpha\)-methyl thioether carboxylic acid, after the introduction of the S-group in \(\alpha\) presents now acidity enough to interact with the N-chlorosuccinimide. The N-chlorosuccinimide protonated frees a chloride (Cl\(^-\)) by means of an nucleophilic attack from the sulfur from the \(\alpha\)-methyl thioether carboxylic acid. The chlorinated thioether carboxylate residue frees a chloride by decarboxylation and formation of the double bond between carbon and sulfur. A double nucleophilic addition of ethanol with configuration inversion occurs with this substrate to generate an ethyl-acetal which by means of protonation of the first ethanol residue and subsequent separation as alcohol by hydrolysis and elimination of the second protonated ethanol, finally conducts to the ketone product. Figure 14 is a similar mechanism to figure 13.

\[ \text{Figure 14. Conversion of bicyclo[2.2.2]hept-5-ene-2-carboxylic acid into (1R,4R)-bicyclo[2.2.2]hept-5-en-2-one instead of an alkene following oxidative decarboxylation. Mechanistic views.} \]

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