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OLEFIN OXIDATION

By L. BATEMAN

(The British Rubber Producers' Research Association, Welwyn Garden City, Herts.)

THE interaction of olefins with molecular oxygen is not only a subject of widespread industrial importance, but is one of the most thoroughly understood chemical processes. This Review attempts to give a broad picture of the main mechanistic features. An earlier article ¹ has reviewed the historical background and has given details of the method of approach and of the earlier kinetic data which were largely responsible for opening up this field.

In Section (1) we present the generally accepted chain mechanism and the simpler rate expressions which are often obeyed. Certain quantitative comparisons of olefinic reactivity derivable on this basis are then discussed. Our main concern, however, is to show how comparatively small changes in certain mechanistic details can give rise to substantial differences in the observed kinetics—so much so that a profound change in mechanism might be imagined. Section (2) deals with rate measurements under nonstationary state conditions designed to determine the propagation- and termination-rate constants separately, and emphasises the inherent limitations to accuracy which oxidation systems present in this respect.

Sections (3), (4), and (5) are concerned with the initiation of the oxidation chain, and the part played in this by the hydroperoxide which is the primary reaction product. This behaviour of the hydroperoxide is responsible for the autocatalytic character of the oxidations, and the complexity and environmental sensitivity of its decomposition serves to complicate the kinetics of the oxidations as a whole. Attention is drawn to circumstances where the fraction of hydroperoxide undergoing decomposition is large compared with that being formed, so that the character and kinetics of the process are greatly altered despite the same fundamental reactions being involved. Section (6) describes efforts to analyse the initiation process quantitatively in order that the number of oxidation chains being started under given conditions can be specified.

Under mild conditions of oxidation, the chain is long and the fraction of the hydroperoxide which decomposes to initiate fresh chains is very small. The overall yield of hydroperoxide should thus be nearly quantitative. In Section (7) serious discrepancies are interpreted in terms of the dual reactivity of peroxy-radicals towards olefins, the consequence being that the measured rate constants are composite quantities relating to both hydrogen extraction and double-bond addition.

The allylic radicals formed on removal of an α -methylenic hydrogen atom from an olefin are mesomeric and hence the derived product may

¹ Bolland, Quart. Reviews, 1949, **3**, 1. 147

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consist of allylic isomerides. The behaviour of mono-olefins and 1:4 diolefins in this respect is discussed in Section (8).

1. General Kinetic Behaviour

The following reaction scheme, where RH represents the olefin with $a_{\rm h}$ α -methylenic hydrogen atom H, $r_{\rm i}$ is the rate of chain initiation, and $t_{\rm he}$ k's are the velocity coefficients of the reactions indicated, accounts for $t_{\rm he}$ reaction characteristics with remarkable comprehensiveness.

Initiation :	Production of	f R \cdot or RO ₂ \cdot radicals	r_1
Propagation :		\rightarrow RO ₂ ·	k_2
	$RO_2 + RH$	$\rightarrow \mathrm{RO}_{2}\mathrm{H} + \mathrm{R}$	k_3
Termination :		\rightarrow) Non-initiating or	k_4
		\rightarrow -propagating	k_5^-
	$2 RO_2$ ·	\rightarrow products	k_{e}

The more obvious of these are : (i) high yields of the hydroperoxide RO_2H (cf. p. 162); (ii) catalysis by light and by free-radical producing substances, indicating the free-radical nature of the reaction; ² (iii) quantum, yields greater than one and a proportionality between rate and the square root of the light intensity in photo-oxidations, indicating a chain reaction with mutual destruction of two chain carriers in the termination step;³ (iv) a parallelism between oxidisability and the relative ease of rupture of the C-H bond in RH, indicating the importance of a hydrogen-exchange reaction such as (3) (cf. p. 149); (v) the formation of conjugated-diene hydroperoxides from 1:4-dienes, in agreement with the generation of mesomeric R-radicals as in (2) (cf. p. 164); and (vi) the marked retardation in rate produced by phenolic compounds (among others), which interfere with the propagation process by providing an alternative and easier reaction for the RO₄ radicals that does not liberate a radical equivalent to R-4

It being assumed that $k_5^2 = k_4 k_6$, the above mechanism yields the rate equation (for long chains)⁵

$$r = r_{\rm i}^{\frac{1}{2}} k_3 k_6^{-\frac{1}{2}} [\rm RH] \frac{k_2 k_6^{-\frac{1}{2}} [\rm O_2]}{k_3 k_4^{-\frac{1}{2}} [\rm RH] + k_2 k_6^{-\frac{1}{2}} [\rm O_2]} \qquad . \qquad (1)$$

where r is the overall rate of oxidation and the square brackets signify concentration terms.

Oxidisability at "High" Oxygen Pressures.—When reaction (2) is so much faster than (3) that $[R] \ll [RO_2]$, termination can be assumed to occur entirely by reaction (6), and equation (1) simplifies to

$$r_{\infty} = r_{1}^{\frac{1}{2}} k_{3} k_{6}^{-\frac{1}{2}} [\text{RH}]$$
 . . . (2)

Equation (2) accurately expresses the observed kinetics for most olefins at

² Bateman and Bolland, Proc. XIth International Congress of Pure and Applied Chem., 1947.

³ Bateman and Gee, Proc. Roy. Soc., 1948, A, 195, 376.

⁴ Bolland and ten Have, Trans. Faraday Soc., 1947, 43, 201; Discuss. Faraday Soc., 1947, 2, 252.

⁵ Bolland, Proc. Roy. Soc., 1946, A, 186, 218.

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 $_{0xygen}$ pressures greater than 100 mm. (the "high" pressure region)—an interesting exception being discussed later. At constant r_i , estimates of $k_{g}k_{6}^{-1}$ are thus obtained from $r_{\infty}/[\mathrm{RH}]$ and these measure the relative reactivities of different olefins in reaction (3), since k_6 is not very sensitive W to changes in R (see p. 153). Bolland 6 has in this way developed a correlation between olefinic structure and oxidisability. Referring to propene, he CH₃·CH:CH₂, at 45°, he concludes that :

(i) Replacement of one or two hydrogen atoms at (a) and/or (c) by (a) alkyl groups increases k_3 by $3 \cdot 3^n$, where n is the total number of substituents; similar replacement at (b) is without effect.

(ii) Replacement of a hydrogen atom at (a) by a phenyl group increases k3 23-fold.

(iii) Replacement of a hydrogen atom at (a) by an alk-l-enyl group increases k_3 107-fold.

(iv) The value of k_3 appropriate to an α -methylenic group contained in a cyclic structure is 1.7 times that of the group contained in an analogous in un acyclic structure.

These rules relate to broad variations, as implied by the assumed lan is equivalency of different alkyl groups. The assumptions that k_6 is invariable and that benzoyl peroxide (used as a standard initiator) initiates throughout eq with equal efficiency also introduce second-order uncertainties. More ng serious discrepancies occur in special cases. Thus 2:4-dimethylpent-2-ene ien is at least 10 times less reactive than would be predicted," presumably 1 fbecause of steric hindrance at C3-behaviour simulated in a saturated tio

TABLE	1	

Olefin, RH ^a	$E_3^{\ b}$	$\Delta H_{\rm 3}=E_{\rm 3}/0.4$	$\Delta (\Delta H_{a})$
$CH_2:CH \cdot CH_3 \cdot \cdot \cdot \cdot \cdot \cdot \cdot$	13·5	34	0
$CH_2:CH \cdot CH_2Alk$	$11 \cdot 5$	29	5
CHAlk:CH·CH2Alk	$10 \cdot {}_{5}$	26	8
$CAlk_2:CH \cdot CH_2Alk \dots$	9	23	11
CH2;CH·CH2Ph	10	25	9
$CHAlk:CH \cdot CH_2 \cdot CH:CHAlk$.	6	15	19
$\begin{array}{c} \text{CH:CH} \overset{\bullet}{\underset{\text{Alk}}{}} \\ \text{Alk} \\ \text{Alk} \\ \end{array}$	9.5	24	10

Reactive α -methylenic group indicated by an asterisk. ^bCalculated from $E_i = E_0 - \frac{1}{2}E_i + \frac{1}{2}E_a$, where E_0 designates the overall activation energy of oxidations stalysed by benzoyl peroxide. E_6 is taken as zero and E_i as 30 kcal./mole (Bolland's rublished values are based on $E_i = 31$ kcal./mole).

⁶ Bolland, Trans. Faraday Soc., 1950, 46, 358. ⁷ Morris, unpublished result.

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