The Mechanism of Ozone-Alkene Reactions in the Gas Phase. A Mass Spectrometric Study of the Reactions of Eight Linear and Branched-Chain Alkenes

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Abstract: The stable products of the low-pressure (4-8 torr (1 torr = 133.33 Pa)) gas-phase reactions of ozone with ethene, propene, 2-methylpropene, cis-2-butene, trans-2-butene, trans-2-pentene, 2,3-dimethyl-2-butene, and 2-ethyl-1-butene have been identified by using a photoionization mass spectrometer coupled to a stirred-flow reactor. The products observed are characteristic of (i) a primary Criegee split to an oxoalkane (aldehyde or ketone) and a Criegee intermediate, (ii) reactions

of the Criegee intermediates such as unimolecular decomposition, secondary ozonide formation, etc., and (iii) secondary alkene chemistry involving OH and other free-radical products formed by the unimolecular decomposition of the Criegee intermediates. The secondary OH-alkene-O₂ reactions account for a significant fraction of the alkene (C_nH_{2n}) consumed and lead to characteristic products such as C_n dioxoalkanes [R'C(O)C(O)R''], at m/e $(C_nH_{2n} + 30)$, C_n acyloins [R'C(O)CH(OH)R''], at m/e $(C_nH_{2n} + 34)$. C_n oxoalkanes and C_n epoxyalkanes observed at m/e ($C_nH_{2n} + 16$) are probably formed primarily via epoxidation of the alkene by O_3 . A general mechanism has been proposed to account for the observations.

There is considerable evidence 1-10 that the gas-phase reactions of ozone with alkenes can be described, at least in part, by the Criegee mechanism shown in eq 1 and 2 for solution-phase ozo-

$$0_3 + > c = c < - > c - c < - > c - c < - > c - c < (1)$$

$$> c = c + > c - c - c < (2)$$

nolysis.^{11,12} In the gas phase, however, reaction 2 is not the sole reaction of the Criegee intermediate¹³ formed in reaction 1. In earlier work in our laboratory on the low-pressure (8 torr = 1.1 kPa) O₃-ethene reaction, 4 the secondary ozonide was too minor a product to be detected and the data were interpreted on the basis

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(11) R. Criegee, Angew. Chem., Int. Ed. Engl., 49, 13 (1977). (12) P. S. Bailey, "Ozonation in Organic Chemistry"; Academic Press: New York, 1978; Vol. 1.

(13) The question as to whether the Criegee intermediate is a diradical or Zwitterion is discussed in ref 6. For the purposes of this discussion the question is unimportant, and we shall use the term "Criegee intermediate" when referring to the dioxy form >C-O-O of the intermediate.

of a complex sequence of free-radical reactions induced by the unimolecular decomposition of the Criegee intermediate (eq 3a-d).

$$H_2C - O - O \rightarrow H_2 + CO_2 \tag{3a}$$

$$H_2C-O-O \to H_2O + CO$$
 (3b)

$$H_2C-O-O \to 2H + CO_2$$
 (3c)

$$H_2C-O-O \rightarrow HO + HCO$$
 (3d)

Subsequent work on the O₃-propene and O₃-2-methylpropene reactions confirmed this general low-pressure mechanism.⁵ In these studies it was apparent that some fraction of the alkene was being consumed in secondary hydroxyl radical reactions. Furthermore, many products have been identified in these reactions which cannot be accounted for by reactions 1 and 2 alone. 14,15 Detailed studies of the O₃-trans-2-butene reaction indicated that these anomalous products accounted for a significant fraction (\sim 20-30%) of the alkene consumed. 16

In this paper we report the mass spectra of low-pressure reacting mixtures of ozone with eight linear and branched-chain alkenes and propose a general mechanism for the O₃-alkene reaction systems to account for the major products formed in reaction 1 and in subsequent free-radical reactions.

⁽¹⁴⁾ Nor could such products be explained by the O'Neal-Blumstein mechanism for O_3 -alkene reactions. ¹⁵ There is now considerable evidence¹⁻¹⁰ against the general validity of the O'Neal-Blumstein mechanism, and, based on new kinetics information, there is general agreement (in reference 8, page 22} that the Criegee reaction pathway will be dominant under many reaction conditions

⁽¹⁵⁾ H. E. O'Neal and C. Blumstein, Int. J. Chem. Kinet., 5, 397 (1973). (16) R. I. Martinez, R. E. Huie, and J. T. Herron, unpublished data, measured with the same instrument as was used in this work. This instrument was a modified version of one described earlier. 4.17 The vacuum envelope, constructed from stainless-steel tubing, consisted of a 6-in. o.d. front section and a 10-in. o.d. rear section. The front section, which contained the reactor and was pumped by a 6-in. oil diffusion pump, was separated from the rear section by a plate which contained a skimmer with a 2-mm diameter hole to provide collimation of the molecular beam entering the rear section. The rear section contained the ionization region and mass filter and was pumped by a 6-in. mercury diffusion lamp. Both pumps used liquid-nitrogen-cooled traps. Sealed and gettered argon and krypton resonance lamps¹⁸ were attached via flanges. Each lamp was fitted with a light shield which had in its base a copper ring which served both as a collimator and as a photon detector to monitor lamp intensity.

Table I. Mass Spectrum of Ozone Plus Ethene Using Krypton and Argon Resonance Lamps^a

	rel int	ensity							
m/e	Kr	Ar	probable stable products	comments					
28 (R)	-1000	-2160		reactant ethene					
29	-10	-38							
30		940	H₂CO						
31		390		fragment ion, probably H_2COH^+ from m/e 48, 60, 62, and 64					
32		160	СН₃ОН	possible contribution from O_2 $(a^1 \Delta_g)$					
33 34	7	39		IP(HO ₂) = 11.5 eV (>Kr line); assignment is therefore uncertain					
34		13	H ₂ O ₂						
42 (R + 14)	54		$H_2C=C=O$	A					
43	11	14		fragment ion; probably H ₂ CCOH ⁺					
44 (R + 16)	21	28	сн _в сно, н ₂ с—сн ₂						
46		165	HC(O)OH						
47		57		fragment ion; probably CH ₂ OOH ⁺ from m/e 64					
48	15	$(8)^b$	$CH_3OOH, CH_2(OH)_2$						
			9 9						
58 (R + 30)	16	17	HC—CH						
59	$(2)^b$	1,	1 C G11						
	(-)		о он						
60 (D + 20)		••	ı I						
60 (R + 32)	15	20	HCCH ₂						
			OH OH						
62 (R + 34)	8		н ₂ с — Сн ₂						
64	12	49	(HO)CH ₂ (OOH)						
67	14	.,	(3)0112(0 011)						

 a $[O_{3}]_{0} = 1.2 \times 10^{-8}$ mol cm⁻³, $[C_{2}H_{4}]_{0} = 4.2 \times 10^{-9}$ mol cm⁻³, $\Delta [C_{2}H_{4}] = 1.8 \times 10^{-9}$ mol cm⁻³, reaction time = 53 s, and total pressure = 8.4 torr. b Values in parentheses may be in error by as much as a factor of 4 in some cases because of background problems at that m/e with lamp indicated and, therefore, are only qualitative.

Experimental Section

The reactions were studied by using a spherical, thermostatically controlled glass reactor which was coupled to a photoionization mass spectrometer through a 200-μm sampling orifice. The instrument was a modified version of one described earlier. I one from the mass analyzer were detected by using a Daly detector and standard pulse-counting techniques, as described earlier. Thus signals from the Daly detector were accumulated (multichannel scaling mode) under software control in a Tracor-Northern TN-11 system which consists of a PDP 11/05 mini-computer, a CRT display, dual cassette tape decks, and four-input scaler, all on the PDP-11 bus. The data, which were stored on the magnetic tape cassettes, were subsequently analyzed on the TN-11 system.

In a typical experiment, reactant flows were adjusted as desired in conjunction with the concomitant throttling of the gas flow to the pump in order to achieve the reaction time which would result in the consumption of $\sim 40-95\%$ of an alkene in the presence of excess ozone. [The reaction time was then estimated (by using known rate constants²¹) from the amount of alkene consumed in the presence of the excess ozone.] Mass spectra were then taken by repeatedly sweeping the mass filter and accumulating the resulting spectrum in the computer. The location of mass peaks was determined by using several known calibrants, and, on the basis of a least-squares regression analysis of mass as a function of channel number, a binomial interpolation routine was used to locate intervening mass peaks. Typically, 1-3 h of accumulation time were needed in order to completely separate minor peaks from the background [accumulation of a mass spectrum was terminated either when the most minor peak exceeded the background count rate n by more than $3n^{1/2}$ (i.e., S/N > 3) or at the end of 3 h of accumulation (a limitation imposed by the limited long-term stability of the instrumental components and the reactant flows)]. A typical difference mass spectrum, obtained by subtracting the mass spectrum of an unreacted mixture (with the ozonizer off) from the reaction mass spectrum (with the ozonizer on), is shown

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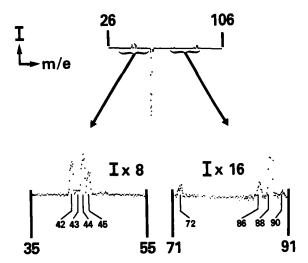


Figure 1. Difference mass spectrum (see text) for the reaction of ozone with trans-2-butene. Lower part of figure shows two horizontally expanded regions of interest for which the vertical display scale (ion intensity I) has also been expanded by factors of 8 and 16, respectively.

in Figure 1 for the O₃-trans-2-butene reaction. This shows products as positive peaks and reactants as negative peaks. Background peaks cancel.

Difference mass spectra, obtained by subtracting the mass spectrum with the ozonizer off from the mass spectrum with the ozonizer on, are given in Tables I–VIII for the reactions of ozone with ethene, propene, 2-methylpropene, cis-2-butene, trans-2-butene, trans-2-pentene, 2,3-dimethyl-2-butene, and 2-ethyl-1-butene. Reaction conditions are specified in each table. All experiments were done at 294 K. For the reaction with 2-ethyl-1-butene, measurements were made with an argon lamp 18 only. For the other alkenes, measurements were made with a krypton lamp (10.0- and 10.6-eV photons) and with an argon lamp (11.6- and 11.8-eV photons), neither of which is sufficiently energetic to ionize molecules such as $\rm H_2$, CO, CO₂, and $\rm H_2O$, which are known products of these reactions. The spectra were normalized by setting $\Delta(\rm alkene)$ (i.e., alkene consumed), as measured with the Kr lamp, equal to -1000.

For the reaction of ozone with *trans*-2-butene, under the conditions given in Table V, final products were analyzed by gas chromatography using a Porapak PS column at 100, 130, and 150 °C. The final products were drawn from the low-pressure *stopped-flow*^{4,5,17} reactor into an

⁽¹⁷⁾ R. I. Martinez and J. T. Herron, Int. J. Chem. Kinet., 10, 433 (1978).
(18) R. Gordon, Jr., R. E. Rebbert, and P. Ausloos, NBS Tech. Note
(IJS) No. 496 (1969)

⁽²⁰⁾ Certain commercial materials and equipment are identified in this paper in order to adequately specify the experimental procedure. In no case does such an identification imply recommendation or endorsement by the National Bureau of Standards, nor does it imply that the material or equipment identified are necessarily the best available for the purpose.

Table II. Mass Spectrum of Ozone Plus Propene Using Krypton and Argon Resonance Lamps^a

	rel in	tensity						
m/e	Kr	Ar	probable stable products	comments				
30 31 32 42 (R) 43	-1000 -17	154 23 16 -510 37	H ₂ CO CH ₃ OH H ₂ C=C=O	fragment ion; probably H_2COH^+ from m/e 48 and 74 possible contribution from O_2 ($a^1\Delta_g$) reactant propene and product ketene fragment ion; probably CH_3CO^+ from m/e 72 and 74.				
44 45	70 28	79 4 7	CH₃CHO	plus isotopic variant of m/e 42 fragment ion; probably CH ₃ CHOH ⁺ from m/e 62 and 74				
46 48	$(4)^b$	11 7	HC(O)OH, CH ₃ CH ₂ OH CH ₃ OOH, CH ₂ (OH) ₂	74				
58 (R + 16)	7	6	сн ₃ сн ₂ сно, (сн ₃) ₂ со, нс—сн, сн ₃ сносн ₂					
61 62	$(2)^{b}$	(4) ^b	CH ₃ CH(OH) ₂ , C ₂ H ₅ OOH	fragment ion				
72 (R + 30)	5		cH ₃ C—CH					
74 (R + 32)	31	16	O OH OH O CH3C—CH2, CH3CH—CH OH OH O—O	the different isomers give rise to different major fragments at m/e 31 and 45				
76 (R + 34)	7		СH ₃ CH—СH ₂ , H ₂ С СН ₂					
90 (R + 48)	$(2)^{b}$		0-0 CH ₃ CH CH ₂					

 $^{^{}a}$ [O₃]₀ = 6.3 × 10⁻⁹ mol cm⁻³, [C₃H₆]₀ = 2.8 × 10⁻⁹ mol cm⁻³, Δ [C₃H₆] = 1.2 × 10⁻⁹ mol cm⁻³, reaction time = 15 s, and total pressure = 4.0 torr. b Values in parentheses may be in error by as much as a factor of 4 in some cases because of background problems at that m/e with lamp indicated and, therefore, are only qualitative.

evacuated sampling coil of ~5-mm i.d. The sampling coil was then pressurized to 1 atm in order to gas dynamically compress, and thus concentrate the gas sample into the end of the sampling coil containing a rubber septum. The gas sample thus pressurized and concentrated was then sampled with a gas syringe and injected into the GC. Ethanal (acetaldehyde), oxirane (ethylene oxide), cis- and trans-2,3-epoxybutane (with a trans/cis ratio of about 6.7), and 3-hydroxy-2-butanone (acetoin) were identified. A broad peak centered about the retention times for 2-butanone and 2,3-butanedione (diacetyl) could not be separated, but, on the basis of the different "tailing" behavior of each compound and the occurrence of corresponding mass peaks at m/e 72 and 86, it was inferred that both were present and there was more 2-butanone than 2,3-buta-

Results and Discussion

In Tables I-VIII we attempt to assign each observed peak in the mass spectra to a stable reaction product or to a fragment ion arising from a product. The number of possible isomeric species or species of the same nominal mass corresponding to any given mass spectral peak makes identification on the basis of mass spectra alone very uncertain. The identification of products here is supported in part by GC analysis of the products of the O₃trans-2-butene reaction, microwave and infrared identification of products of the O₃-ethene reaction, ^{22,23} studies of the effects of added ethanal and nitrogen dioxide on the products of the O_3 -trans-2-butene reaction, 10 and the existing body of data on ozone chemistry.¹² In some cases probable product identities are based solely on mechanistic considerations. In the absence of either confirmatory evidence or plausible mechanistic schemes, however, the mass spectral peaks are left unidentified. Wherever possible we have attempted to identify peaks which do not correspond to a stable product but rather are fragment ions formed by dissociative ionization of a parent product, since, even with low-energy photoionization, fragmentation is a very important process, particularly with large oxygen-containing organic molecules. For

example, in Table IX we give the mass spectra of a group of selected reaction products particularly relevant to the O₃-2-butene reaction. It can be seen that in general the krypton lamp leads to the least ambiguous mass spectra, whereas with the argon lamp fragmentation becomes much more important. For some minor products, a fragment ion but not a parent ion might be found, thus the absence of a parent ion does not rule out the presence of the corresponding product molecule. Highly speculative, but plausible, mass peak assignments have been enclosed in brackets.

A second important consideration with respect to the mass spectra is that the ion intensities do not quantitatively reflect the partial pressures of the corresponding neutral species since ion intensity is a function of both partial pressure and mass spectral sensitivity. The latter can vary widely, which explains why a very minor product such as ketene, which has a very high sensitivity, is so prominent in the spectra. This caveat is stressed in order to prevent misinterpretation of the data in Tables I-VIII.

In section A we provide a general overview of the types of reaction products that were observed and indicate their probable origins. In sections B through F we discuss in some detail the reaction mechanisms enumerated in section A. In every case, the discussion is referenced to our observations for the O₃-trans-2butene reaction, since the latter has been studied over the widest range of conditions and is the one for which we have supporting GC product identification data.

A. General Information. The most important product of the ozonation reaction is in every case the oxoalkane (aldehyde or ketone) expected on the basis of the Criegee mechanism shown in eq 4a,b. Approximately 0.9-1.0 mole of oxoalkane is produced

⁽²²⁾ H. Kühne, S. Vaccani, A. Bauder, and Hs. H. Günthard, Chem. Phys., 28, 11 (1978)

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Table III. Mass Spectrum of Ozone Plus 2-Methylpropene Using Krypton and Argon Resonance Lampsa

		tensity						
m/e	Kr	Аr	probable stable products	comments				
30	0.6	129	H ₂ CO	5				
31 32	1	8 4	CH ₃ OH	fragment ion; probably CH ₃ O ⁺ from m/e 48 or 62 or 74 possible contribution from O ₂ ($a^1\Delta_g$)				
34		i	H_2O_2	possible contribution from O ₂ (a Mg)				
40		-17	* *					
41	-1	-94		4.41				
42 43	27 32	3	$H_2C=C=O$	probable contribution from fragment ion $(CH_3)_2C^+$ from m/e 56				
44	32 16	83 32	CH ₃ CHO	fragment ion; probably CH_3CO^+ from m/e 58 and 74				
45	4	8	C113C110	fragment ion; possibly CH ₃ CHOH ⁺ from m/e 60				
46		10	HC(O)OH	, , , , , , , , , , , , , , , , , , ,				
47	0.5	3		fragment ion; possibly $CH_3O_2^+$ from m/e 48 or CH_2OOH^+ from m/e 6				
48	1.5	4	$CH_3OOH, CH_2(OH)_2$					
53 54	0.5	-2						
55		-47						
56 (R)	-1000	-360		reactant 2-methylpropene				
57	-40	-6						
58	262	78	(CH ₃) ₂ CO					
59	51	65	(011) (011(011)	fragment ion; probably $(CH_3)_2COH^+$ from m/e 76, 88, and 90				
60	2.6	5	(CH ₃) ₂ CH(OH),					
61	0.7	1.7	CH₃C(O)OH	fragment ion; probably $CH_3C(OH)_2^+$ from m/e 76				
62	0.5	1.7		riaginent ion, probably $Cri_3C(O11)_2$ from m/e 70				
64	0.5	0.7	(HO)CH ₂ (OOH)					
66	0.5		•					
			н₃с О					
70 (R + 14)	2.7	3.2	(CH ₃) ₂ C=C=0, H ₂ C=C-C-H					
71		1.3	52					
/ 1		1.3	0.0					
			кн _э) ₂ снсно, кн ₃) ₂ с—сн ₂ , сн ₃ с—сн					
72 (R + 16)	31	12	(СН ₃) ₂ СНСНО, (СН ₃) ₂ С́—СН ₂ , СН ₃ С́—СН					
73	7 13		fragment ion; probably $CH_3C(OH)CHO^+$ from m/e 88					
			Q.					
74	8	7	 CH3COCH3					
75	10	25	•	fragment ion; probably CH ₃ C(OH)CH ₂ OH ⁺ from m/e 90				
	- •		0-0					
7.0	2		ŢŢ					
76	2	1.6	H ₂ C СH ₂ , (СН ₃) ₂ С(ОН) ₂					
0.1		0.6	ğ					
81 82	$(0.5)^{b}$	0.6 1.6						
84	$(0.3)^{b}$	1.3						
85	(0.7)	1						
86 (R + 30)	2							
87	0.7	1						
			OH O					
88 (R + 32)	4	1.4	(СН ³) ² С — СН					
89	1	1.4	V L					
37	1	1.7	OH OH					
00 (D + 24)	•	4						
90 (R + 34)	3	1	(CH3)2C-CH2					
95	0.0	1						
96 97	0.9	1.3 0.7						
98	$(0.5)^{b}$	1						
101	(0.0)	1.3						
102		$\frac{1.3}{(0.5)^b}$						
103	0.7							
			0-0					
104 (R + 48)	1		(CH3)2C CH2					
			\forall					
105	0.7	0.6						
117		0.6						
121	$(0.5)^b \\ (0.5)^b$							
124								

 $^{^{}a}$ $[O_{3}]_{o} = 9.7 \times 10^{-9}$ mol cm⁻³, $[C_{4}H_{8}]_{o} = 1.1 \times 10^{-9}$ mol cm⁻³, $\Delta [C_{4}H_{8}] = 7.5 \times 10^{-10}$ mol cm⁻³, reaction time = 14 s, and total pressure = 4.1 torr. b Values in parentheses may be in error by as much as a factor of 4 in some cases because of background problems at that m/e with lamp indicated and, therefore, are only qualitative.

Table IV. Mass Spectrum of Ozone Plus cis-2-Butene Using Krypton and Argon Resonance Lampsa

	rel intensity						
m/e	Kr	Ar	probable stable products	comments			
30		54	H ₂ CO				
40		-28					
41		-174					
42	69		$H_2C=C=O$				
43	12	48	0	fragment ion; probably $\mathrm{CH_3CO^+}$ from m/e 72, 86, and 88			
44	66	194	сн _з сно, н _г с—сн _г				
45	33	89		fragment ion; probably CH ₃ CHOH ⁺ from m/e 88 and 90			
46		11	HC(O)OH,				
			CH ₃ CH ₂ OH				
47	[4] ^b [4] ^b		3 - 2	fragment ion; possibly $CH_3O_2^+$ from m/e 48			
48	[4] ^b		CH ₃ OOH				
55		-85	,				
56 (R)	-1000	-574		reactant cis-2-butene			
57	-43						
			0 0				
58	20						
36	20		HCCH				
			Ĭ				
70 (R + 14)	[6] ^b	13	сн₂ — сн—ё—сн₃				
			8 R				
72 (R + 16)	12		CH-CH-CHCH- CH-CH-CCH-				
			2 13 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2				
82		17					
			9 9				
86 (R + 30)	[13] ^b						
00 (R 50)	[13]		ongo-roong				
			C OH				
88 (R + 32)	33	15	сн _з с—ёнсч _з				
	1 h						
89	[6] ^b						
			OH OH				
90 (R + 34)	[9] ^b		CH3CH—CHCH3				
,	[-1		0-0				
104 (0) + 40)	141 h		ŢŢ				
104 (R + 48)	[4] ^b		CH3CH CHCH3				
			8				
110	[2] ^b						

 a $[O_{3}]_{0} = 3.3 \times 10^{-9}$ mol cm⁻³, $[C_{4}H_{8}]_{0} = 7.3 \times 10^{-11}$ mol cm⁻³, $\Delta [C_{4}H_{8}] = 4.6 \times 10^{-11}$ mol cm⁻³, reaction time = 4.1 s, and total pressure = 4.1 torr. b Taken under different experimental conditions: $[O_{3}]_{0} = 3.2 \times 10^{-9}$ mol cm⁻³, $[C_{4}H_{8}]_{0} = 4.7 \times 10^{-10}$ mol cm⁻³, Δ [C₄H₈] = 2.3 × 10⁻¹⁰ mol cm⁻³, reaction time = 3.1 s, and total pressure = 4.0 torr.

per mole of alkene consumed. We also find a large number of other products which, for the most part, arise from the subsequent chemistry of the Criegee intermediate.

Examination of Tables I-VIII reveals certain common features in the distribution of the products. In particular, we find products at masses corresponding to the alkene reactant (R) plus 16, 30, 32, 34, and in some cases 14 and 48 mass units [identified in the tables as (R + 16), (R + 30), etc.]. In the case of 2,3-dimethyl-2-butene, only the R + 16 product was observed. In only four cases do we detect the R + 48 peak of the secondary ozonide which can be formed in reaction 2. This presumably is a consequence of competing loss processes (e.g., unimolecular decomposition) for the initially formed Criegee intermediate²⁴ and/or low instrumental sensitivity. Titration of the Criegee intermediate by addition of aldehyde leads to enhancement of the secondary ozonide signal as a consequence of reaction 2.10,24

Methanal (formaldehyde) is observed at m/e 30 as a reaction product for all ozone-alkene reactions studied, even when it is not a product of the primary Criegee split (reaction 1), in which case it is probably formed by reactions of free radicals such as CH₃· and CH₃O· with O₂ and/or O₃ and probably also by reaction with O₂ of the •CH₂(OH) radical which for terminal alkenes may

(24) J. T. Herron, R. I. Martinez, and R. E. Huie, submitted for publication.

be formed via OH-alkene reactions, as is discussed in sections B and E.

An m/e 44 product peak is also observed for every O₃-alkene system studied. It is probably due mainly to ethanal, but at least in the case of O_3 + ethene and of O_3 + trans-2-butene, it also includes a contribution from oxirane. In the O₃-ethene reaction, m/e 44 represents an R + 16 product as is discussed in section C. In the case of the reaction of O₃ with 2-butene and with some of the other alkenes (e.g., propene, trans-2-pentene, etc.), m/e44 arises from both the primary Criegee split (reaction 1) and via OH-alkene reactions, as is discussed in section B.

On the basis of its kinetic behavior, 16 the product peak at m/e42 is almost certainly due to ketene. Although the ketene peak is a prominent feature in all the mass spectra because of the high mass spectral sensitivity of ketene, the actual amount formed is small. In the case of *trans*-2-butene, the ketene yield is only about 3% of that of ethanal. Moreover, ketene has been characterized as an R + 14 product in the case of O_3 + ethene, while, in the case of other O₃-alkene reactions, ketene could be a Criegee rearrangement product (i.e., it is formed by an intramolecular elmination which occurs subsequent to one of the possible intramolecular rearrangements of a Criegee intermediate). This is discussed in section D with reference to both C_n and C_m ketenes from C_nH_{2n} alkenes (where m < n). In all subsequent discussions of C_m and C_n products, we will always have m < n.

Secondary OH-alkene reactions, which account for a significant fraction (\sim 13-22%) of the alkene consumed, lead to the char-

Table V. Mass Spectrum of Ozone Plus trans-2-Butene Using Krypton and Argon Resonance Lamps^a

	rel in	ntensity		
m/e	Kr	Ar	probable stable products	comments
30		167	H ₂ CO	
31		11	CT. CT.	fragment ion; probably CH_3O^+ from m/e 48
32 33		18 3	CH₃OH	possible contribution from O_2 ($a^1 \Delta_g$)
34		6	H_2O_2	
40		-42	2 - 2	
41		-320		
42	62 7	14	$H_2C=C=O$	fragment ion metable CH COt from m/s 72 96 and 99 and C H Ot
43	/	117		fragment ion; probably CH_3CO^+ from m/e 72, 86, and 88, and $C_2H_3O^+$ from m/e 44
44	76	432	сн _з сно, н ₂ сссн ₂	
45	41	253	HO(O)OH OH OH OH	fragment ion; probably CH_3CHOH^+ from m/e 62, 88, and 90
46 47	5	19 19	HC(O)OH, CH ₃ CH ₂ OH	fragment ion; probably CH ₃ O ₂ + from m/e 48 or CH(OH) ₂ + from m/e 62
48	4	23	CH ₃ OOH	magnification, productly crising from my control of crising from my control
55		-129	•	
56 (R)	-1000	-1100		reactant trans-2-butene
57	-41	-38		
••	_			
58	7	20	HC—CH	
60 61		12 14	CH₃C(O)OH	
62		$(3)^b$	CH ₃ CH(OH) ₂	
68		$(3)^b$		
			0	
70 (R + 14)		10	сн₂ = сн−с−сн₃	
			0 9	
72 (R + 16)	10	11	CH3CH-CHCH3, CH3CH2CCH3	
73				
75		$(3)^b$		
82		8 (3) ^b 6		
86 (R + 30)	15	6	сн ₃ С—С—сн ₃	
			о он II I	
88 (R + 32)	49	37	сн ₃ С—снсн ₃	
89	5	6		
			он он	
90 (R + 34)	8	4	CH3CH—CHCH3	

 $a = [O_3]_0 = 2.5 \times 10^{-9} \text{ mol cm}^{-3}, [C_4H_8]_0 = 4.4 \times 10^{-10} \text{ mol cm}^{-3}, \Delta [C_4H_8] = 3.1 \times 10^{-10} \text{ mol cm}^{-3}, \text{ reaction time} = 4.0 \text{ s, and total pressure} = 4.1 \text{ torr.}$ b Values in parentheses may be in error by as much as a factor of 4 in some cases because of background problems at that m/e with lamp indicated and, therefore, are only qualitative.

acteristic products R + 30, R + 32, and R + 34 which are discussed in section B and, with the exception of ethene, propene, and 2,3-dimethyl-2-butene, may also lead to the R + 14 products which are discussed in section D with reference to C_n conjugated oxoalkenes. While the R + 30 products derived from OH-alkene- O_2 reactions are C_n dioxoalkanes, the C_m dioxoalkanes discussed in section D are Criegee rearrangement products.

Finally, in section E we discuss the origins of both C_m and C_n acids and acid derivatives, while in section F we discuss some anomalous products.

Throughout our discussion we shall attempt to emphasize patterns among the products, whether they be characteristic or apparently anomalous for the ozone-alkene system studied.

(OH)R"] which are probably formed via a complex sequence of reactions initiated by reaction of hydroxyl radicals with the alkene. 10 This is discussed next with reference to 2-butene.

The peak at m/e 86 is probably due to 2,3-butanedione and accounts for about 2-3% of the butene consumed. As noted above, an unresolved peak from the gas chromatograph corresponded to both 2-butanone and 2,3-butanedione. From the mass spectral sensitivities of the epoxybutanes, 2-butanone, and 2,3-butanedione, the relative peak heights of m/e 72 and 86, and the relative gas-chromatographic peak areas for the epoxybutanes and the unseparated butanone and butanedione, we are able to estimate that there is about twice as much 2-butanone formed as 2,3-epoxybutane. These latter two products together account for about 2-3% of the butene consumed.

The peak at m/e 88 is due to 3-hydroxy-2-butanone (acetoin) and was confirmed by GC. It accounts for about 10–18% of the butene consumed. The corresponding product of the O_3 -ethene reaction, CH(OH)CHO, has been confirmed by microwave and infrared spectroscopy.²²

The small peak observed at m/e 90 is probably due to 2,3-butanediol.

Since with the argon lamp the magnitudes of the peaks at m/e 86, 88, and 90 decreased relative to the peak heights observed with

Table VI. Mass Spectrum of Ozone Plus trans-2-Pentene Using Krypton and Argon Resonance Lamps^a

	rel in	tensity		
m/e	Kr	Ar	probable stable products	comments
28	3	1.7	C ₂ H ₄	probably includes contribution of CH ₃ CH ⁺ fragment ion
30		12	H₂CO	
32		5	CH₃OH	possible contribution from O_2 ($a^1 \Delta_g$)
34		1	H_2O_2	
42	20	-43	$H_2^*C=C=O$	probably includes contribution of C ₂ H ₅ CH ⁺ fragment ion from m/e 70 and 86
43	10	8		fragment ion; probably CH_3CO^+ from m/e 100 and 102
44	39	17	CH ₃ CHO	
45	10	6	WG(0)0H G H OH	fragment ion; probably CH_3CHOH^+ from m/e 62, 102, and 10-
46	2	2	HC(O)OH, C₂H₅OH	
48	2 -3	2	CH ₃ OOH	
55	-3	-71		
			Ħ	
56	4		он _з c=c=0	
57	9	7		fragment ion; probably $C_2H_5CO^+$ from m/e 58, 74, 86, 100 and 102
			9 9	
58	196	26	 С ₂ н₅сно, нс—сн	
59	20	6		fragment ion; probably C ₂ H ₅ CHOH ⁺ from m/e 102 and 104
60	1.4	U	CH ₃ C(O)OH, CH ₃ CH ₂ CH ₂ OH	raginent ion, probably C ₂ 11 ₅ CHOH Hom m/e 102 and 104
61		3	CH ₃ C(O)OH, CH ₃ CH ₂ CH ₂ OH	fragment ion
62	1.6 6	0.9	CH ₃ CH(OH) ₂	Tagnient ton
70 (R)		-5 4	CH ₃ CH(OH) ₂	reactant trans-2-pentene
	-1000 -55	-34 -2.4		reactant trans-2-pentene
71	-33	-2.4	0.0	
72	5	$(0.8)^{b}$	Ĭ ch₌c—ch	
		(0.0)	3	
74	1.6		$C_2H_5C(O)OH$, $HC(O)OC_2H_5$	
75	$(1)^{b}$			
			9	
84 (R + 14)	$(1)^{b}$			
0.(20, 1.)	(1)		52 go 5 or g. or gon — or or or or or	
			, i i	
86 (R + 16)	12		c_2H_5 ĆH $-$ ČHC H_3 , c_2H_5 $-$ Č $-$ С $_2H_5$, c_3H_7 $-$ Č $-$ С H_3	
			ү—	
90	1.4		CH ₂ CH CH ₂	
	1.1			
0.4	4		5	
94	1			
			î î	
00 (R + 30)	18	$(0.1)^{b}$	C ₂ H ₅ C—CCH ₃	
		. ,		
01	$(2)^{b}$			
			о он оно	
02 (R + 32)	11		о он он о 	
			ĬŢŢŢŢŢŢ	
04 (R + 34)	2		C2H3CH—CHCH3. CH3CH CHCH3	
			X	
			γ—γ	
10 (0 + 40)	1.4		C5H²€¢H ¢HCH³	
18 (R + 48)	1.4		Comach Chicha	

 $^{^{}a}$ [O₃]₀ = 3.1 × 10⁻⁹ mol cm⁻³, [C₅H₁₀]₀ = 6.8 × 10⁻¹⁰ mol cm⁻³, Δ [C₅H₁₀] = 6.1 × 10⁻¹⁰ mol cm⁻³, reaction time = 5.4 s, and total pressure = 4.0 torr. b Values in parentheses may be in error by as much as a factor of 4 in some cases because of background problems at that m/e with lamp indicated and, therefore, are only qualitative.

the krypton lamp, we concluded that some of the new peaks which appeared when the argon lamp was used were probably due to dissociative ionization of these larger molecules. Examination of the photoionization mass spectra of 3-hydroxy-2-butanone, 2,3-butanedione, trans-2,3-epoxybutane, and 2-butanone given in Table IX confirmed this hypothesis. In addition, the spectra of Table IX demonstrate that dissociative ionization can be important even when a krypton lamp is used. This may help to explain the observation by Atkinson et al.²⁵ of mass spectral peaks at masses corresponding to free radicals in their study of O₃-alkene reactions. These mass peaks were probably due to ion fragments produced by dissociative ionization. Their work was possibly further complicated in the particular instrument they used (which was described in ref 26) by electron-impact ionization by photoelectrons produced by scattered photons.26

In a study involving the addition of radical scavengers to the ozone-trans-2-butene system, 10 we were able to infer that these products at m/e 86, 88, and 90 were due to secondary radical reactions (eq 6a-c and 7) arising from the reaction of the hydroxyl

⁽²⁵⁾ R. Atkinson, B. J. Finlayson, and J. N. Pitts, Jr., J. Am. Chem. Soc., 95, 7592 (1973).

⁽²⁶⁾ R. I. Martinez, thesis, University of California, Los Angeles, 1976; Diss. Abstr. Int. B, 36, 6193-B (1976).

Table VII. Mass Spectrum of Ozone Plus 2,3-Dimethyl-2-butene Using Krypton and Argon Resonance Lamps^{a,b}

	rel inter	nsity					
m/e	Kr	Ar	probable stable products	comments			
30	-10	47	H₂CO ^b				
41		3					
42	11		H ₂ C=C=O				
43 44	50	112	-				
44		7	CH₃CHO				
45		6	-				
48		6	CH₃OOH				
56	- 2	-35					
58	731	178	(CH ₃) ₂ CO				
59	91	62		fragment ion; probably (CH ₃) ₂ COH ⁺			
60		7	$CH_3C(O)OH^b$				
69	-36	-430					
70		-25					
71		-2					
			0 0				
72	24	10	 CH₃C—CH				
			о <i>р</i> фон б				
74	9	9	CH ₃ C—OCH ₃ . (CH ₃ C=CH ₂) — CH ₃ C—CH ₂ OH				
81		6					
82		6					
84 (R)	-1000	-270		reactant 2,3-dimethyl-2-butene			
85	-74	-17		reactant 2,5 dimethyr-2-butche			
100 (B + 16)	(e)C		Å				
100 (R + 16)	$(5)^c$		(CH ₃) ₂ CC(CH ₃) ₂				
101		6					

 a [O₃]₀ = 1.6 × 10⁻⁹ mol cm⁻³, [C₆H₁₂]₀ = 6.2 × 10⁻¹¹ mol cm⁻³, Δ [C₆H₁₂] = 5.8 × 10⁻¹¹ mol cm⁻³, reaction time = 2.8 s, and total pressure = 4.1 torr. b Although not observed here, CH₃OH and HC(O)OH have been reported as products by Smith and Eastman.⁵⁰ They also identified as products H₂CO, CH₃C(O)OH, and CH₃C(O)OCH₃. c Values in parentheses may be in error by as much as a factor of 4 in some cases because of background problems at that m/e with lamp indicated and, therefore, are only qualitative.

radical with butene (eq 5), thus leading to 3-hydroxy-2-butanone and 2,3-butanediol.

OH OO*

OH C4H8
$$\frac{O_2}{C}$$
 CH3C—CCH3 (5)

OH OO*

CH3CH—CHCH3 + RCH2O2* — CH3CH—CHCH3 + RCH2O* + O2 (6a)

OH OH

CH3CH—CCH3 + RCH2OH + O2 (6b)

OH OH

CH3CH—CHCH3 + RCH0 + O2 (6c)

OH O*

CH3CH—CHCH3 + O2 — CH3CH—CCH3 + HO2* (7)

2,3-Butanedione probably arises from similar reactions (eq 8a–c and eq 9) The source of the $CH_3C(O)C(CH_3)(H)OO$ · radical

is not known, although its presence, along with the radical products of reactions 5, 6a, and 8a, has been inferred from scavenging experiments by using NO_2 .¹⁰

In addition to the ethanal produced via the Criegee split, significant amounts of ethanal probably also result from the OH-trans-2-butene-O₂ reaction system²⁷ via such reactions as eq 10-12. (See, however, comments and discussion on this type of reaction in ref 8.)

$$CH_3CH + O_2 \longrightarrow CH_3CHO + HO_2$$
 (11)

$$\begin{array}{c|cccc}
0 & 0 & & & \\
& & & & & \\
CH_2C - CHCH_2 & - & CH_2CHO + CH_2CO & & \\
\end{array}$$
(12)

The types of reactions shown above can be written for all the alkenes considered here, except for 2,3-dimethyl-2-butene. For the linear, nonsymmetric alkenes, isomeric hydroxy-substituted oxo compounds are possible, which, although they have the same mass, will have different ion fragmentation patterns. Fragments corresponding to the different isomers can be found (see, for example, Table II).

C. Products of Alkene Epoxidation Reactions. For the O_3 -trans-2-butene reaction, the peak at m/e 72 is due to 2,3-epoxybutane and 2-butanone. These two products together account for about 2-3% of the butene consumed, and as was indicated above, there is about twice as much 2-butanone as there is 2,3-epoxybutane. Both cis- and trans-epoxybutanes were found, with a trans-to-cis ratio of about 6.7. Epoxides and oxoalkanes arising from the parent alkene are known to be products of the solution phase ozonation of alkenes, but there remains some uncertainty as to the mechanism of their formation. In his recent review, Bailey¹² favored a mechanism, in eq 13a,b, in which ozone adds

⁽²⁷⁾ H. Niki, P. D. Maker, C. M. Savage, and L. P. Breitenbach, J. Phys. Chem., 82, 135 (1978).

Table VIII. Mass Spectrum of Ozone Plus 2-Ethyl-1-butene Using an Argon Resonance $Lamp^{a,b}$

m/e	rel intensity Ar	probable stable products	comments
28 30	20 380	C ₂ H ₄ H ₂ CO	probably includes contribution of CH ₃ CH ⁺ fragment ion
31	23	_	fragment ion; probably $\rm H_2COH^+$ and/or $\rm CH_3O^+$ from m/e 48
34 41	-16	H_2O_2	
42 44 45	-172 140 42	СН₃СНО	fragment ion; probably $C_2H_5O^+$ from m/e 62 and m/e 120
			[HOCH ₂ CH ₂ C(OH) ₂ C ₂ \hat{H}_s] and/or m/e 134 [HOCH ₂ CH ₂ C(OH)-(C ₂ H _s)CH ₂ OH] and/or COOH ⁺ from m/e 46 and/or 74
46 47	35 8	HC(O)OH, C₂H₅OH	fragment ion; probably $CH_3O_2^+$ from m/e 48
48 54	11 -46	CH ₃ OOH, CH ₂ (OH) ₂	•
55 56	-262 -737		
57 59	514 19		fragment ion; probably $C_2H_5CO^+$ from m/e 86 and/or 100 fragment ion; probably $C_2H_5CHOH^+$ from m/e 88 or CH_2COOH^+ from m/e 74
62	12	C ₂ H ₅ OOH	nom me 14
69 70 71	-2362 -112 55		fragment ion; probably $(C_2H_5)_2CH^+$ from $m/e~100$ and/or 88
70	24	0	
72 73	34 10	c ₂ H₅ [©] CH₃	fragment ion; probably (C ₂ H ₅)CH(OH)CH ₂ + from m/e 88
74	19	$C_2H_5C(O)OH$	
76	3	H ₂ C CH ₂	
82	14	· ·	
83 84 (R)	0.8 1000		reactant 2-ethyl-1-butene
85 86	-50 390	$(C_2H_5)_2CO$	
87	189	372	fragment ion; probably $(C_2H_5)_2COH^+$ from m/e 118 and/or 116 $[(C_2H_5)_2C(OH)CHO]$ and $C_2H_5C(OH)CHO^+$ from m/e 116
88 89	31 44	$(C_2H_5)_2CH(OH)$	fragment ion; probably $C_2H_5C(OH)_2CH_2^+$ from m/e 118 and/or 120
07	7.4		[C ₂ H ₅ C(OH) ₂ CH ₂ CH ₂ OH] and/or HOCH ₂ C(OH)(C ₂ H ₅)* from m/e 118 and 132 [HC(O)CH ₂ C(OH)(C ₂ H ₅)CH ₂ OH] and/or 134 [HOCH ₂ CH ₂ C(OH)(C ₂ H ₅)CH ₂ OH]
95	6		
96	2	Ο ς ₂ H ₅	
98 (R + 14)	14	HC—C=CHCH ₃ , (C ₂ H ₅) ₂ C=C=O	
99	14		fragment ion; probably $(C_2H_5)_2C(CHO)^+$ from m/e 116 $[(C_2H_5)_2C(OH)CHO]$
100 (R + 16)	104	(C.H.), (C.H.), (C.H.), CHCHO, C.H.CC, H., CH, CH, CH, CH.	3
101	16		fragment ion; probably $(C_2H_5)_2C(CH_2OH)^+$ from m/e 118 and/or $HC(O)CH_2C(OH)(C_2H_5)^+$ from m/e 132 [$HC(O)CH_2C(OH)-(C_2H_5)CH_2OH$]
103	50		fragment ion; probably HOCH ₂ CH ₂ C(OH)C ₂ H ₅ * from m/e 120 [HOCH ₂ CH ₂ C(OH) ₂ C ₂ H ₅] and/or 134 [HOCH ₂ CH ₂ C(OH)-
112	3		(C ₂ H _s)CH ₂ OH]
113 114 (R + 30)	5 8		
115	6		fragment ion; possibly $HC(O)CH_2C(C_2H_5)CH_2OH^+$ from m/e 132 [$HC(O)CH_2C(OH)(C_2H_5)CH_2OH$]
117	7		fragment ion; possibly HOCH ₂ CH ₂ C(C ₂ H ₅)CH ₂ OH ⁺ from m/e 134 [HOCH ₂ CH ₂ C(OH)(C ₂ H ₅)CH ₂ OH]
118 (R + 34)	4	0 	
121 123	6 4		
	7		

 $a [O_3]_0 = 9.6 \times 10^{-9} \text{ mol cm}^{-3}$, $[C_6H_{12}]_0 = 6.5 \times 10^{-10} \text{ mol cm}^{-3}$, $\Delta [C_6H_{12}] = 4.9 \times 10^{-10} \text{ mol cm}^{-3}$, reaction time = 14 s, and total pressure = 4.1 torr. b Quantitative observations were not possible beyond m/e 130 due to background noise problems.

directly to the alkene. A different mechanism involving epoxidation by the Criegee intermediate has been given very strong support by Hinrichs et al.28 in a study in which the Criegee intermediate was produced in the absence of ozone. Possibly, either mechanism could be important, depending upon the reactant and the solvent. However, the addition of a 20-fold excess of ethanal to the low-pressure ozone-trans-2-butene reaction system led to a doubling of the yield (per mole of butene consumed) of m/e 72.²⁹ This suggests that these products do not arise from reactions of a stabilized Criegee intermediate under these conditions. The apparent yield increase is a consequence of the fact that in the presence of added ethanal the butene is no longer being consumed by hydroxyl radicals. Thus, it rules out the involvement of hydroxyl radicals in the formation of products at m/e 72 via reactions such as those proposed by Morris et al. 30 for OH-alkene reactions in the absence of added O₂. The work of Niki et al.²⁷ provides additional support for this conclusion. Therefore, we conclude that under our experimental conditions, these epoxides and oxoalkanes arise from a primary ozonation reaction, as discussed by Bailey. 12,31 Oxirane and ethanal, which are known products of the O₃-C₂H₄ reaction²² and which correspond to the observed m/e 44 peak in Table I, are probably also formed via reactions analogous to (13a) and (13b).

D. Dioxoalkanes, Ketenes, and Conjugated Oxoalkenes. (i) C_m Dioxoalkanes. For the O_3 -trans-2-butene reaction, the peak

(28) T. A. Hinrichs, V. Ramachandran, and R. W. Murray, J. Am. Chem. Soc., 101, 1282 (1979).

(29) R. I. Martinez, J. T. Herron, and R. E. Huie, unpublished data. Under our experimental conditions, with $[trans-2-C_4H_8]_0 \simeq (1.1 \pm 0.07) \times 10^{-9}$ mol cm⁻³, the fractions of trans-2-buttene (T2B) consumed in the presence and absence of added CH₃CHO $\{\pm A\}$ were respectively $F_{T2B}^{\Lambda} = \Delta [T2B]^{\Lambda}$ [T2B]₀ = 0.27 and $F_{T2B}^{\pi\circ\Lambda} = \Delta [T2B]^{\pi\circ\Lambda}/[T2B]_0 = 0.4_2$, and $\alpha_{T2B} = F_{T2B}^{\pi\circ\Lambda}/F_{T2B}^{\Lambda} = 1.6$. In the presence of added CH₃CHO, with [CH₃CHO]₀/[0₃]₀/[T2B]₀ $\simeq 20.8:1$, more than 75% of any OH produced by secondary reactions should have been scavenged by the CH₃CHO instead of reacting with the T2B. Hence, for any product X, by comparing its yield per mole of T2B consumed, $Y_X \{=\Delta[X]/\Delta[T2B]\}$, in the presence of added excess CH₃CHO, $Y_X^{\pi\circ\Lambda}$, relative to its product yield in the absence of added cxcess CH₃CHO, $Y_X^{\pi\circ\Lambda}$, one can infer whether or not X is a product of secondary reactions initiated by OH-alkene reactions. Thus, by making the simplifying assumption that T2B is consumed primarily by O₃ and OH, we concluded, on the basis of the measured $R_X = Y_X^{\Lambda}/Y_X^{\pi\circ\Lambda}$, that while diacetyl, acctoin, and 2,3-butanediol certainly arise from OH-T2B-initiated reactions ($R_X \approx 0.4$). Also, for Me = 58 [(HCO)₂], $R_X \approx 1$. This suggests that (HCO)₂ does not arise via OH-T2B chemistry but probably from O₃-T2B-initiated reactions. However, the relatively low R_X value also suggests that the (HCO)₂ precursor may be partially consumed by the added CH₃CHO. This is suggestive of a precursor such as the excited Criegee intermediate, 24 which is effectively titrated (i.e., quenching plus scavenging of the excited Criegee intermediate) by added CH₃CHO in competition with its rearrangement to glyoxal.

(30) E. D. Morris, Jr., D. H. Stedman, and H. Niki, J. Am. Chem. Soc., 93, 3570 (1971).

(31) An alternative path for epoxyalkane formation, suggested by Harding and Goddard⁶ with reference to oxirane formation in the gas-phase $O_3-C_2H_4$ reaction, is the intramolecular elimination of O_2 , (a¹ Δg) from the primary

ozonide. Under our experimental conditions we cannot distinguish between reactions (13a) and (RN1) as the source of epoxyalkane.

at m/e 58 is probably due to dioxoethane (ethanedial, glyoxal) since its phosphorescence has been identified in the O₃-2-butene systems. 32,33 (For the other alkenes studied, the m/e 58 peak may also correspond to other products, as indicated in Tables I-VIII.) The source of the glyoxal is not known. We have observed, however, that the addition of a 20-fold excess of ethanal to the reacting system resulted in no change in the yield of dioxoethane per mole of butene consumed.²⁹ If dioxoethane was to arise from hydroxyl radical initiated reactions or from the stabilized Criegee intermediate, the yield would have been expected to decrease by a factor of about 2. Further, in their studies of the chemiluminescence from ozone-alkene systems, Schurath et al.33 reported that dioxoethane phosphorescence is observed only for alkenes such as R'CH₂C(H)=CR"R" and R'CH₂C(H)= C(H)CH₂R", while 1,2-dioxopropane (methylglyoxal) phosphorescence is observed only for alkenes such as (CH₃)₂C=CR'R" and $(CH_3)_2C = C(CH_3)_2$ (where R', R", R" = H or alkyl). Also, this phosphorescence was independent of added O₂.

These observations can be rationalized by assuming that the dioxoalkane phosphorescence is a consequence of a rearrangement of the excited Criegee intermediate,²⁴ a primary product of the ozone-alkene reaction. If the phosphorescent dioxoalkane was to arise from a direct ozone-alkene interaction, as do the epoxyalkanes, then its yield would be expected to increase upon addition of ethanal. Hence, the constant dioxoalkane yield probably reflects a competition between the rearrangement process, and quenching and scavenging of the excited Criegee intermediate by the ethanal.²⁴ Thus, consistent with the above observations, we propose the chemiexcitation mechanism of Scheme I for methylglyoxal from 2-methylpropene, 2-methyl-2-butene, and 2,3-dimethyl-2-butene [all of which produce methylglyoxal

⁽³²⁾ B. J. Finlayson, J. N. Pitts, Jr., and R. Atkinson, J. Am. Chem. Soc., 96, 5356 (1974).

⁽³³⁾ U. Schurath, H. Güsten, and R. D. Penzhorn, *J. Photochem.*, **5**, 33 (1976).

Table IX. Mass Spectra of Selected Reaction Products of Ozone-Alkene Reactions Using Krypton and Argon Resonance Lamps^a

	СН	0 ₃ сн	CH ₃ C	CCH ₃	C₂⊦) И _Б ССН ₃	снзсн–	снсн _з	сн _з с—	•о <i>-</i> сн _з	СН ₃ С) 	li l	он -снсн _з
m/e	Kr	Ar	Kr	Ar	Kr	Ar	Kr	Ar	Kr	Ar	Kr	Ar	Kr	Ar
28							0.9	5						
29 31 32 39 41 42		3						7		_				
31										9 8				
32										8				0.0
39 41														0.8 1
42			1	1		0.4	4	5						1
43		306	10	68	4	44	867	685	3	1400	132	156	6	23
44	1000	1890	10 0.4	68 2	·	2	471	1470	•	38	5	7	·	3
45	23	45	• • •	_		0.3	13	52		15	-	0.8	410	388
46	23 2	4					2 39 8	4						10
57					10 2	23 0.9	39	40						
58			1000	190	2	0.9	8	3			5			
59			38 3	6 0.7		0.1								2 2
45 46 57 58 59 60 71 72 73 74			3	0.7										2
71								8			_			
72					1000	148	1000	137			3	0.8	23	33
73					49	7	46	6	1000	1500				
74					3	0.6	3		1000	1560				
13 76									32 6	55 7				
96									U	,	1000	133	32	5
87											51	7	11	3
75 76 86 87 88 89 100											6	0.9	1000	100
89													45	6
100											2 ^b	0.3 ^b		
114													100 b	13 b

^a Normalized to parent peak = 1000 using the krypton lamp. ^b Impurity peaks.

phosphorescence³³ and yield the $(CH_3)_2COO$ Criegee intermediate], while we propose Scheme II for glyoxal from alkenes which yield RCH₂CHOO Criegee intermediates,³³ where $R = H \cdot$ for propene and *cis*- and *trans*-2-butene, $CH_3 \cdot$ for 1-butene, and $HC(O)CH_2CH_2CH_2 \cdot$ for cyclohexene.³³ These reactions are discussed next.

Reaction 14 is important in the solution-phase chemistry of O₃ + 2,3-dimethyl-2-butene, 12 and reaction 18 is its analogue. Reaction 17 was proposed by Wadt and Goddard³ as a source of hydrogen atoms and H₂COO in the gas-phase reaction of O₃ with 2,3-dimethyl-2-butene. [These products will be discussed in section F with reference to OH Meinel band chemiluminescence and H₂CO (¹A") production in ozone-alkene systems.] By analogy with reaction 13b, we propose reactions 15 and 19 as the sources of excited glyoxal and methylglyoxal. In the case of $O_3 + cis$ -2-butene, the measured specific rate of phosphorescent emission from glyoxal³² suggests that reaction 20 accounts for less than 10⁻³% of the butene consumed, while the measured specific rates of emission from OH and H₂CO³² suggest that (21) accounts for less than 10⁻⁵% of the butene consumed. Clearly, reactions such as (19)-(21), and by analogy probably also (15)-(17), are very minor processes in O₃-alkene systems.

The chemiexcitation mechanism proposed above would explain why glyoxal phosphorescence is not observed in the O_3 -ethene system, even though glyoxal is a known product of the reaction (see Table I and section B with reference to OH-alkene reactions as sources of C_n dioxoalkanes). Clearly, reactions 14 and 18 are not possible for H_2COO (e.g., from $O_3 + C_2H_4$). Furthermore, according to the proposed mechanism, the formation of excited 2,3-butanedione would not be expected for any of the alkenes discussed, and therefore in the absence of other excitation mechanisms, diacetyl phosphorescence should not be observed in any of the O_3 -alkene systems discussed. Such was found to be the case. 32,33 Diacetyl chemiexcitation by the proposed mechanism might occur for an alkene such as

where various R' and R" substituents would have to be tested, including H and CH_3 . However, this too would not necessarily provide positive evidence for the proposed mechanism because the dynamics of the intramolecular rearrangement reactions (14) and (18) may preferentially involve intramigration of the H atoms of the C^{α} , resulting in formation of $(CH_3)(R')(R'')C-C(O)-C-(O)H$ instead of diacetyl.

It is interesting to note that glyoxal (m/e 58) is produced in both the O_3 -cis-2-butene and O_3 -trans-2-butene systems (see Tables IV and V). This suggests that the syn-anti isomers of the Criegee intermediate which is involved in reaction 18 do readily interconvert in the gas phase, as has been proposed by Harding and Goddard.⁶ The calculated barrier to syn-anti interconversion is sufficiently high so that in solution the isomers will not interconvert, and, on the basis of conformational analysis of primary ozonides, trans-alkenes produce only syn isomers while cis-alkenes produce only anti isomers.^{6,12} Hence, if the syn-anti isomers did not interconvert in the gas phase, glyoxal would be produced only by the syn isomer and thus only in the O_3 -trans-2-butene reaction and not in the O_3 -cis-2-butene reaction.

Examination of Tables I-VIII shows that the presence or absence of product peaks at m/e 58 (glyoxal) and m/e 72 (methylglyoxal) is in general agreement with the proposed chemiexcitation mechanism, reactions 15 and 19. However, for several of the alkenes, these peaks also correspond to other products or to these same products but from other sources such as the OHalkene reactions discussed in section B. For example, it is especially noteworthy that the same dioxoalkane, glyoxal, can be either a C_m or C_n product, depending on its origin. Thus, while glyoxal is a C_n product of the OH-ethene reaction, it is a C_m product of the O₃-2-butene reaction, produced by the proposed chemiexcitation mechanism. Similarly, while methylglyoxal is a C_n product of the OH-propene reaction, it is a C_m product of the $O_3-2,3$ dimethyl-2-butene reaction, produced by the proposed chemiexcitation mechanism. This was not immediately obvious and probably delayed the characterization of the various mechanisms responsible for the same functional-class compounds.

An alternate route to C_m dioxoalkanes is suggested as a possibility by the proposal of Wadt and Goddard³ that, in addition to undergoing reaction 17 [and by analogy (21)], the hydroper-

oxide formed in reaction 14 [and by analogy in (18)] might also decompose by losing an OH group. Then, in the presence of O_2

where the hydroperoxide R'C(OOH)CHR" is formed by rearrangement of an initially formed Criegee intermediate in a reaction analogous to (14) and (18). The O_3 -1-butene and/or O_3 -trans-2-pentene reactions in the presence and absence of O_2 would provide a unique test of the possible validity of reaction 22. The observation of m/e 72 from O_3 -trans-2-pentene (see Table VI; presumed to be methylglyoxal based on reaction 22, with R' = H, R" = CH₃) provides some support for this hypothesis, although GC identification is still necessary.

(ii) C_m Ketenes. For the C_nH_{2n} alkenes studied, examination of Tables I-VIII also reveals product peaks which correspond to C_m ketenes (m < n); e.g., ketene from $O_3 + 2$ -butene in Tables IV and V, and both ketene $(m/e \ 42)$ and methylketene $(m/e \ 56)$ from $O_3 + trans$ -2-pentene in Table VI. In the case of $O_3 + trans$ -2-butene, the yield of ketene was about 0.03-0.04 mol/mole of butene consumed. One can readily account for all the observed C_m ketenes if the hydroperoxide R'C(OOH)CHR", which is formed by rearrangement of an initially formed Criegee intermediate in a reaction analogous to (14) and (18), undergoes the intramolecular elimination processes indicated in Scheme III. Reaction 27 is the well-known Wolff rearrangement.³⁴

(iii) C_n Ketenes and Conjugated C_n Oxoalkenes. Examination of Tables I-VIII indicates that for most C_nH_{2n} alkenes there are product peaks designated in the tables as R+14 which correspond to C_n -substituted ketenes (e.g., in Table I, $O_3+C_2H_4 \rightarrow$ ketene²²) and/or C_n conjugated oxoalkenes (e.g., in Tables IV and V, $O_3+2-C_4H_8 \rightarrow 3$ -buten-2-one³²). Ketene has been confirmed as a product of the $O_3-C_2H_4$ reaction by microwave spectroscopy. Its production in the $O_3-C_2H_4$ reaction obviously cannot be explained on the basis of an isomerized Criegee intermediate. 3-3-Buten-2-one has been confirmed as a product of the O_3-2 -butene reaction by gas chromatography; it was observed only in the presence of added O_2 . This suggests that the 3-buten-2-one is a product of secondary radical reactions, probably initiated by OH addition to the butene.

Scheme III

(34) J. March, "Advanced Organic Chemistry: Reactions, Mechanisms, and Structure"; McGraw-Hill: New York, 1968.

(35) Ketene was observed as a product of the O_3 -ethanal reaction (unpublished data). Ethanal is a known product of the O_3 -ethene reaction. Hence the ketene observed in the O_3 -ethene reaction may have been produced by addition of O_3 to the product ethanal. However, because of their relative specific rates, the secondary O_3 -CH $_3$ CHO reaction would become important relative to an O_3 -alkene reaction only under conditions of excess O_3 after most of the alkene would have been consumed. The observation of ketene from O_3 + CH $_3$ CHO led to an examination of the recent work of several investigators. This indicated that the currently accepted O_3 -aldehyde reaction mechanism was incomplete. An alternative mechanism has been proposed O_3 -to explain the observations.

(36) R. I. Martinez, Int. J. Chem. Kinet., in press.

Scheme IV

$$R_2$$
 R_1
 R_2
 R_3
 R_4
 R_5
 R_5
 R_5
 R_5

E. C_m and C_n Alkanoic Acids and Acid Derivatives. A puzzling feature of the low-pressure stopped-flow studies of the O_3 -ethene⁴ and O_3 -propene⁵ reactions was the induction times found for methanoic (formic) and ethanoic (acetic) acid formation. In addition, alkanoic acid yields from ozone-alkene reactions ($\sim 0.05-0.08$ mol acid/mol of alkene consumed) appear to be substantially independent of the total pressure.^{4,7,23} These observations seem to rule out simple quenching of an isomerized Criegee intermediate as a significant source of alkanoic acids and acid derivatives (Scheme IV).²⁴ Also, the induction times would rule out the *direct* production of the acids from the reactions of OH or HO_2 radicals with oxo compounds (aldehydes and ketones).

A possible explanation for the observed induction times is that the alkanoic acids arise from the thermal decomposition of compounds such as (HOO)CR'R"(OH), possibly produced by secondary reactions involving HO₂ addition to oxo compounds. For example, the thermal and photochemical decomposition of (HOO)CH₂(OH) has been implicated as one source of methanoic acid in the photooxidation of methanal.^{37,38} Their suggested mechanism is shown in eq 30–32. The possibility that reaction 32 is

$$HO_{2^{\bullet}} + CH_{2}O \rightleftharpoons (HO_{2}CH_{2}O \cdot) \rightleftharpoons (HO)CH_{2}OO \cdot (30)$$

$$HO_{2'} + (HO)CH_2OO \rightarrow (HO)CH_2(OOH) + O_2$$
 (31)

$$(HO)CH2(OOH) \xrightarrow{h\nu} HC(O)OH + H2O (32)$$

the source of methanoic acid in the O_3 -ethene reaction is supported by the observation of a product peak at m/e 64 (see Table I), a fairly unique mass, corresponding to (HOO)CH₂(OH). However, it seems unlikely that the rate of reaction 30 is fast enough to produce (HOO)CH₂(OH) in our system.³⁹ An alternate path (eq 33) involves addition of H₂O to the stabilized Criegee in-

$$H_2O + H_2COO - H_2C$$
 H_2C $O-OH$ OH OH

termediate. 12,24 This would provide an atmospherically important loss process for the Criegee intermediate 24 but cannot explain (HOO)CH₂(OH) formation under our low-pressure conditions. The reaction sequence (33) + (32) is discussed in detail elsewhere 24 with reference to its postulated role in O_3 -alkene- SO_2 systems.

Scheme V

(37) F. Su, J. G. Calvert, J. H. Shaw, H. Niki, P. D. Maker, C. M. Savage, and L. D. Breitenbach. Chem. Phys. Lett., 65, 221 (1979).

and L. D. Breitenbach, Chem. Phys. Lett., 65, 221 (1979).
(38) F. Su, J. G. Calvert, and J. H. Shaw, J. Phys. Chem., 83, 3185 (1979).

(39) Parent product peaks corresponding to the homologous (HOO)-CR'R"(OH), where R'R"C=O are the oxo compounds produced via reactions 4, were not observed for any oxo compound other than H_2 CO[(HOO)-CR'R"(OH) product peak at m/e 64] and then only in the O_3 -ethene and O_3 -2-methylpropene reactions, even though H_2 CO is a significant product of every O_3 -alkene reaction studied.

$$\cdot CH_2OH + O_2 \rightarrow H_2CO + HO_2 \cdot \tag{39}$$

as has been confirmed recently by Radford.⁴⁰ Moreover, the homologous (HOO)CR'R"(OH), which would have been expected from analogous OH-alkene-O₂ reactions, were not observed.³⁹

An additional source of alkanoic acids was suggested by the study of Horowitz et al.41 and by our observation that when a large excess of ethanal is added to the O₃-trans-2-butene system, the yield of methanoic acid drops, while that of ethanoic acid increases. This suggested suppression of reaction 40c ($k_{40a} >> k_{40b}$ and k_{40c} $\simeq 2k_{40b}$) due to scavenging of HO radicals by ethanal with the

$$HO + H_2CO \rightarrow H_2O + HCO$$
 (40a)

$$\rightarrow$$
 H₂O + HCO* (40b)

$$\rightarrow$$
 HC(O)OH + H (40c)

consequent generation of ethanoic acid via (41b). However (41a)

$$HO + CH_3CHO \rightarrow HC(O)OH + CH_3$$
 (41a)

$$\rightarrow$$
 CH₃C(O)OH + H (41b)

probably predominates over (41b), because of the relative bond strengths of the C-C and C-H bonds involved in reaction 41. Hence, (41) probably cannot explain our observation.

Finally, we found no evidence for the formation of methanoic acid anhydride in the O₃-ethene reaction, as has been reported.^{22,23} However, examination of reaction conditions suggests that the anhydride is a product of a pressure-dependent process whose rate is probably dependent on $[O_3]^p[C_2H_4]^q[M]'$. Thus, Kühne et al.²² found the anhydride for $[O_3] = [C_2H_4] = 1-2$ torr (total pressure $P_{\rm t} = 2-4 \; {\rm torr}, \; T = 240-300 \; {\rm K}, \; {\rm reaction \; time} \; \tau \simeq 20-40 \; {\rm s}) \; {\rm but}$ not for $[O_3] = [C_2H_4] = 10-50$ mtorr $(P_1 = 20-100$ mtorr, T = 10-100220-300 K, $\tau \simeq 2-100$ s). On the other hand, Su et al.²³ observed the anhydride with $[O_3] \simeq [C_2H_4] \simeq 7$ mtorr ($P_t = 700$ torr, $T = 291-299 \text{ K}, \tau = 0-65 \text{ min}$), while we observed no anhydride for $[O_3] \simeq 200$ mtorr, $[C_2H_4] \simeq 80$ mtorr $(P_t \approx 8 \text{ torr}, T \approx 300)$ K, $\tau \approx 50$ s). Furthermore, when observed, Kühne et al.²² found an anhydride yield of only $\sim 0.8\%$ on the basis of $\Delta[C_2H_4]$, while Su et al.²³ found ~18-29% yield on the basis of $\Delta[C_2H_4]$ (when CO, H₂CO, and SO₂ were not included as additives).

F. "Anomalous" Products. (i) Chemiluminescent Species. The discussion of section D on C_m dioxoalkanes in terms of the reactions of the primary Criegee intermediates provides a basis for discussion of the observations by Finlayson et al.32 of OH and H₂CO chemiluminescence from O₃-alkene reactions in the presence and absence of O₂. Thus reaction 14 followed by reaction 17 was proposed by Wadt and Goddard³ as a source of H atoms in order to rationalize the OH $(X^2II_i)_{\nu'\leq 9}$ Meinel bands (generated by H + O₃ \rightarrow OH[†] + O₂) observed in the O₃-2,3-dimethyl-2butene reaction.³² The intent of their proposal was to demonstrate how other alkenes might give rise to some of the same chemiluminescent species as are observed in the O₃-ethene reaction. For example, chemiluminescence from H₂CO (¹A") was observed in all the O₃-alkene systems studied by Finlayson et al.,³² and it was not affected by the presence of O₂. Wadt and Goddard³ argued that the most likely mechanism for H₂CO (¹A") production in the ozone—ethene system is the attack by O_3 on the planar ${}^1A'(4\pi)$ state of H₂COO.

$$H_2$$
COO [¹A′(4π)] + O₃ →
 H_2 CO (¹A″) + 2O₂ ΔH ≈ -113 kJ mol⁻¹ (42)

The reaction is sufficiently exothermic so that the $H_2CO(^1A'')$ is probably produced with at least 29 kJ mol-1 of vibrational excitation, in which case it can dissociate to H + HCO, thereby providing a source of H atoms and, therefore, a source of OH Meinel bands via $H + O_3 \rightarrow OH^{\dagger} + O_2$. The isomerization and subsequent decomposition of H₂COO also provides a source of H atoms and therefore OH[†].

It is clear that for the alkenes studied by Finlayson et al., 32 the sequence of reactions 14 and 17, and by analogy (18) and (21), do provide a source of H_2COO and hence of H_2CO ($^1A''$) and OH $(X^2\Pi_i)_{\nu \le 9}$. Thus, while ethene, propene, 1-butene, and 2-methylpropene can provide H₂COO directly via the primary Criegee split (reaction 4), the alkenes 2-methylpropene, 2methyl-2-butene, and 2,3-dimethyl-2-butene can provide H₂COO via reaction 17, and the alkenes propene and cis- and trans-2butene can provide H₂COO via reaction 21. Consequently, the observations by Finlayson et al.³² of OH and H₂CO chemiluminescence from the reaction of O3 with these alkenes in the presence and absence of O₂ can be explained by reactions of the primary Criegee intermediates, as outlined above. And, as was discussed in section D, such reactions [(15)-(17) and (19)-(21)] represent very minor processes in the O₃-alkene systems.

(ii) C_m Alkene Products. C_m alkenes have been reported as products of the reaction of O_3 with a variety of C_n alkenes. 12,43 However, this discussion will be restricted to the production of C_2H_4 (~0.02–0.04 mol/mol of alkene consumed) by the reaction of O_3 with 1-butene^{12,43} or 2-pentene^{12,43} (also see Table VI) or 2-ethyl-1-butene (see Table VIII). One possible source of C₂H₄ is via its unimolecular elimination from the corresponding primary Criegee intermediate, subsequent to its rearrangement to an excited alkanoic acid derivative 24 (reactions 43-46), where R' = H

for 1-butene and 2-pentene and C₂H₅ for 2-ethyl-1-butene. Peaks corresponding to the expected acid derivatives were observed at m/e 46 for 2-pentene (see Table VI) and m/e 74 for 2-ethyl-1butene (see Table VIII). Additional support for this plausible mechanism comes from the work of Ausloos⁴⁴ and Wijnen⁴⁵ on the gas-phase photolysis of alkyl esters. They demonstrated that C₂H₄ was a primary product arising from the intramolecular rearrangement and decomposition of both ethyl propanoate^{44,45} and ethyl methanoate,44 which correspond respectively to the Criegee intermediates expected from 2-ethyl-1-butene and 2pentene (or 1-butene).

The above mechanism [(43)-(46)] necessitates the migration of bulky alkyl groups to form an actual "hot" acid intermediate, which although possible, seems less probable than a radical fragmentation process whereby the migrating bulky alkyl group falls off during its attempted rearrangement, thus forming a radical pair such as $(R'C(=O)-O + C_2H_5)$ or $(R'C(=O) + C_2H_5O)$. Consequently, we propose the reaction sequence (43), (47), and (48) as the most probable source of the molecular products R'C(O)OH and C_2H_4 .

Vrbaski and Cvetanovic⁴³ proposed instead an H-atom migration within the primary Criegee intermediate followed by

⁽⁴⁰⁾ H. E. Radford, Chem. Phys. Lett., 71, 195 (1980). (41) A. Horowitz, F. Su, and J. G. Calvert, Int. J. Chem. Kinet., 10, 1099 (1978)

⁽⁴²⁾ The arguments presented by Finlayson et al. (ref 32, p 5366) to rationalize the significant increase in the intensity of the Meinel bands in the absence of O₂ are not affected by our considerations since the isomerization and decomposition of the primary Criegee intermediate proceeds unaltered (43) T. Vrbaski and R. J. Cvetanovic, Can. J. Chem., 38, 1063 (1960). (44) P. Ausloos, Can. J. Chem., 36, 383 (1958). (45) M. H. J. Wijnen, J. Am. Chem. Soc., 82, 3034 (1960).

elimination of CH₃CH and the corresponding Criegee fragment.

However, the newly formed Criegee intermediate R'C(H)OO-would be expected to rearrange and decompose, without significant stabilization to the corresponding acid product.²⁴

It is clear that the mechanisms involved in the formation of "anomalous" products such as C_m alkenes are still poorly understood but do suggest the involvement of intramolecular rearrangement processes such as those outlined here. For example, alkenes could also be formed from the decomposition of the hot epoxyalkane⁴⁶ formed in the ozone-alkene reaction.

Conclusions

The products of gas-phase ozone-alkene reactions are characteristic of (i) a primary Criegee split (reaction 4) to a C_m oxoalkane and a Criegee intermediate, (ii) competing loss processes for the Criegee intermediates, viz., unimolecular decomposition vs. reactions such as secondary ozonide formation etc., and (iii) secondary alkene chemistry involving OH and the other freeradical products which are formed by the unimolecular decomposition of the Criegee intermediates. The alkanoic acids appear to be products of a variety of slow reactions involving oxoalkanes which are produced in the O_3 -alkene systems. The C_n oxoalkanes and C_n epoxyalkanes appear to be formed via epoxidation of the C_nH_{2n} alkene primarily by O_3 . While the production of C_n ketenes and C_n oxoalkenes is probably initiated by OH addition to the alkene, the C_m ketenes appear to be Criegee rearrangement products. Thus, for different O₃-alkene reactions, the same compound or classes of compounds (e.g., ketenes) may arise from very different reaction mechanisms, thereby complicating the unraveling of the overall gas-phase O₃-alkene reaction mechanism.

The secondary OH-alkene-O₂ reactions account for a significant fraction of the alkene consumed and lead to characteristic products such as dioxoalkanes, acyloins, and alkanediols. The OH radicals which initiate these reactions can be produced directly from the unimolecular decomposition of a Criegee intermediate in reactions such as (3d) or its homologue, reaction 50. Sub-

$$CH_3\dot{C}HOO \rightarrow [CH_3C(O)OH]^{\dagger} \rightarrow OH + CH_3C(O)$$
 (50)

$$\rightarrow$$
 CH₃ + COOH (51)

sequent reactions of hydrogen atoms, alkyl radicals, and other primary free-radical decomposition products of Criegee intermediates [e.g., CH₃ from (51)] also can lead to OH or HO₂. Reactions of radicals with ozone may be an additional OH source, e.g.

$$CH_3 + O_3 \rightarrow OH + H_2COO$$
 (52)

This reaction is of considerable interest as a possible new source of the Criegee intermediate. Reactions such as (52), however, can only be important under experimental conditions such as used here (\gtrsim 4% O_3 in O_2 , total pressure \lesssim 4-8 torr), where the consumption of CH_3 by reaction with O_3 is competitive with the consumption of CH_3 by reaction with O_2 . Thus, for example, on the basis of a recent determination of the rate constant for the $CH_3 + O_3$ reaction, the rates of reactions $CH_3 + O_3$ and $CH_3 + O_2$ are about equal under our reaction conditions. Under atmospheric conditions, however, the $CH_3 + O_3$ reaction would be unimportant.

As we discuss elsewhere, ²⁴ relative rates of reactions participating in O₃-alkene systems are very significantly altered by reaction conditions, thereby influencing observed product distributions and assumed mechanisms. For example, high aldehyde concentrations will tend to stabilize the initially formed excited Criegee intermediate,²⁴ thereby reducing the free-radical activity of the system while encouraging secondary ozonide formation. Increasing the total pressure will also affect the product distribution because of third-body effects. However, these will not only alter the secondary radical chemistry (e.g., $H + O_2 \stackrel{M}{\longrightarrow} HO_2$ followed by $HO_2 + O_3 \rightarrow OH + 2O_2$) but also affect to varying degrees the stabilization of the initially formed excited Criegee intermediate and consequently the free-radical activity of the O₃-alkene system. Hence, in order to unravel the complete gas-phase reaction mechanism, one must determine the product distributions as a function of total pressure and reactant concentrations.

The extremely complex secondary chemistry found in ozone-alkene reactions is a consequence of the unimolecular decomposition of Criegee intermediates. The chemistry of the Criegee intermediate remains the single most important problem area in understanding the gas-phase mechanism of ozone-alkene reactions. It is interesting to note, therefore, that it has been suggested that the Criegee intermediate may also participate in combustion⁴⁸ and in the photooxidation of aldehydes.⁴⁹

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⁽⁴⁸⁾ P. S. Nangia and S. W. Benson, Int. J. Chem. Kinet., 12, 43 (1980).

⁽⁴⁹⁾ B. M. Morrison, Jr., and J. Heicklen, J. Photochem., 11, 183 (1979).

⁽⁵⁰⁾ H. E. Smith and R. H. Eastman, J. Am. Chem. Soc., 83, 4274 (1961).