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Journal of Macromolecular Science, Part A Publication details, including instructions for authors and subscription information: http://www.informaworld.com/smpp/title~content=t713597274

Free-Radical Ring-Opening Polymerization

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To cite this Article Bailey, William J., Chen, Paul Y., Chen, Shuh-Chung, Chiao, Wen-Bin, Endo, Takeshi, Gapud, Benjamin, Lin, Yin-Nian, Ni, Zhende, Pan, Cai-Yuan, Shaffer, Scott E., Sidney, Luann, Wu, Shang-Ren, Yamamoto, Noboru, Yamazakl, Noboru and Yonezawa, Kazuya(1984) 'Free-Radical Ring-Opening Polymerization', Journal of Macromolecular Science, Part A, 21: 13, 1611 — 1639

To link to this Article: DOI: 10.1080/00222338408082081

URL: http://dx.doi.org/10.1080/00222338408082081

Science Part A:

Pure and Applied Chemistry

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Free-Radical Ring-Opening Polymerization

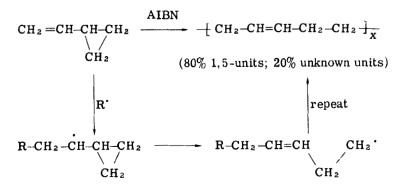
WILLIAM J. BAILEY, PAUL Y. CHEN, SHUH-CHUNG CHEN, WEN-BIN CHIAO, TAKESHI ENDO, BENJAMIN GAPUD, YIN-NIAN LIN, ZHENDE NI, CAI-YUAN PAN, SCOTT E. SHAFFER, LUANN SIDNEY, SHANG-REN WU, NOBORU YAMAMOTO, NOBORU YAMAZAKI, and KAZUYA YONEZAWA

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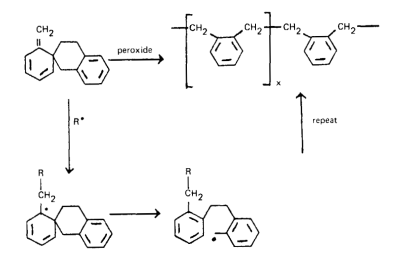
Although the ionic ring-opening polymerization of heterocyclic compounds, such as ethylene oxide, tetrahydrofuran, ethylenimine, β -propiolactone, and caprolactam, as well as the Ziegler-Natta ring-opening of cyclic olefins, such as cyclopentene and norbornene, are well known, free radical ring-opening polymerizations are rather rare. The few examples that are reported in the literature involve cyclopropane derivatives or highly strained bicyclic olefins. For example, Takahashi [1] studied the free radical polymerization of vinylcyclopropane and reported that the cyclopropane ring opened to give a polymer containing about 80% 1,5-units and about 20% of undetermined structural units.

Apparently the radical adds to the vinyl group to give the intermediate cyclopropylmethyl radical which opens at a rate faster than the addition to the double bond of another monomer. Somewhat similar results were obtained with the chloroderivatives. More recently, Cho and Ahn [2] studied the related malonic ester derivative, which underwent ring-opening free-radical polymerization to produce a high molecular weight polymer containing only the 1,5 units:

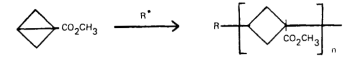
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Errede [3] found the dimer of o-xylylene would undergo free-radical ring-opening polymerization to give the corresponding poly-o-xylylene:

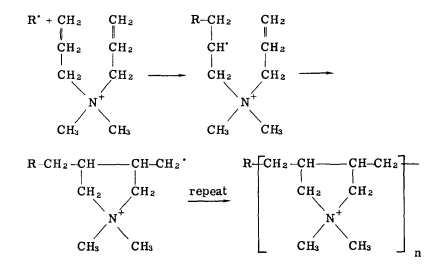


In this case the driving force for the ring-opening step is the formation of the aromatic ring. Hall and co-workers [4] showed that derivatives of bicyclo[1.1.0] butane would polymerize by free radicals by cleavage of the highly strained central bond:



Of course, the ring-opening polymerization of S_{θ} has been postulated to involve free radicals [5].

Unstrained five- and six-membered carbocyclic rings, on the other hand, are usually involved in ring-closing reactions rather than ring opening. For example, Butler and Angelo [6] in 1957 reported that when diallyldimethylammonium bromide was polymerized by a free radical mechanism, a soluble polymer containing five-membered rings was obtained by an inter-intramolecular polymerization.



Apparently the reaction is kinetically controlled to form the fivemembered ring rather than the thermodynamically favored sixmembered ring. The course of some of these ring-opening and ringclosing polymerizations can be explained by the recent data of Maillard, Forest, and Ingold [7], who studied the transformations in the cyclopropylmethyl and the cyclopentylmethyl series by electron spin resonance. In the case of the three-membered radical the reaction involves ring-opening since the energy is favorable and the rate of the reaction is very high. In the case of the five-membered ring system the reaction proceeds in the direction of ring-closure since the energetics of that reaction is favorable and the rate of the ring closure is also moderately high (Table 1).

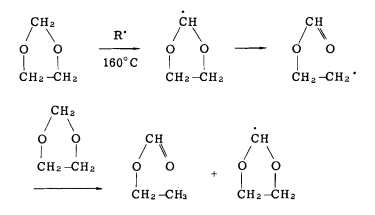
With the assumption that the carbon-oxygen double bond is at least 50-60 kcal/mol more stable than the carbon-carbon double bond [8], one can calculate that the introduction of an oxygen atom in place of the carbon atom in the cyclopentylmethyl radical would favor the reverse reaction or the ring opening. In other words, the ring-opening reaction would be favored by at least 40 kcal/mol by producing the more stable carbonyl double bond.

| | TABLE 1 | | |
|--|-------------------------|--------------|--------------------------|
| | k250 (S ⁻¹) | E (kcal/mol) | Log A (s ⁻¹) |
| $\begin{array}{c} CH_{2} \\ \\ CH_{2} \\ CH_{2} \\ CH_{2} \end{array} \xrightarrow{ CH_{2} \cdot CH_{2} \cdot CH_{2} \cdot CH_{2} \\ CH_{2} \\ CH_{2} \\ CH_{2} \end{array}$ | $1.3 	imes 10^{8}$ | 5.94 | 12,48 |
| $\begin{array}{cccc} CH_2 & CH_2 &$ | $1.0 	imes 10^{5}$ | 7.8 | 10.7 |

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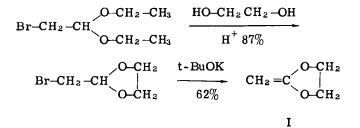
FREE-RADICAL RING-OPENING POLYMERIZATION

A search of the literature indeed revealed many ring systems containing an oxygen atom that would undergo a ring-opening reaction in the presence of free radical catalysts. One such case was the cyclic formal, ethylene formal. Maillard, Cazaux, and Lalande [9] found that when ethylene formal was heated at 160° C, it rearranged to ethyl formate:



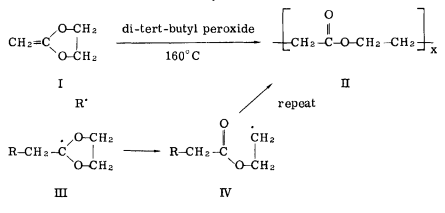
The reaction could be rationalized as indicated where the driving force for the ring-opening reaction in the chain reaction was the formation of the stable carbon-oxygen double bond in the final ester. With the knowledge that such a ring system would undergo cleavage, it seemed to be a fairly straightforward process to synthesize a monomer that would undergo ring-opening polymerization by introducing a double bond at the carbon atom flanked by the two oxygens.

The monomer desired for this ring-opening polymerization had indeed been prepared by McElvain and Curry [10] in 1948. Although Johnson, Barnes, and McElvain [11] had treated diethyl ketene acetal with peroxide and had reproted that there was no reaction, no such study was reported for the 2-methylene-1,3-dioxolane (I). A reinvestigation of the cyclic ketene acetal I was therefore undertaken. The ketene acetal I was found to be extremely sensitive to acid [12]. Ex-



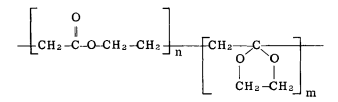
posure of I to the atmosphere would cause immediate polymerization to give a polymer with little or no ring opening. In order to handle this monomer conveniently, it was necessary to have some base, such as potassium tert-butoxide or a tertiary amine, always present; under these conditions no spontaneous polymerization would occur.

Treatment of this monomer I with benzoyl peroxide gave a high molecular weight polyester by a free radical ring-opening polymerization which can be rationalized by the scheme



The structure of the polyester was established by analysis as well as infrared and NMR spectroscopy. This polyester is difficult to synthesize with high molecular weight from the γ -hydroxybutyric acid because of the stability of the competing lactone.

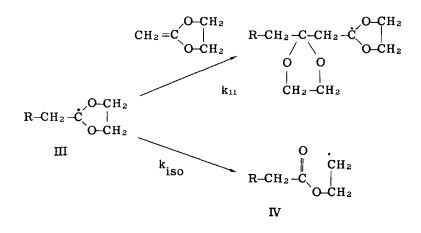
At lower temperatures the ring opening is not complete. Thus at 60° C only 50% of the rings are opened to give a random copolymer of the following structure [12]:



The structure of the polyester was established by analysis as well as infrared and NMR spectroscopy. An alternative method of analysis of the extent of ring-opening was the basic hydrolysis of copolymer II, which cleaved the ester groups but left the cyclic ketals intact.

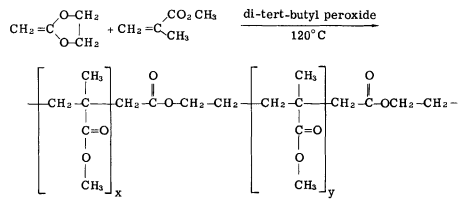
Even at 120°C only 87% of the rings are opened. Apparently the unopened radical III can add directly to monomer I in competition with the ring-opening process to form open chain radical IV. High dilution was found to favor the ring-opening process since the addition of III

to monomer I is a second-order reaction while the conversion of III to the open chain radical is first order. The extent of ring opening is kinetically controlled with a direct competition between the rate of direct addition, k_{11} , and the rate of ring opening, k_{150} .

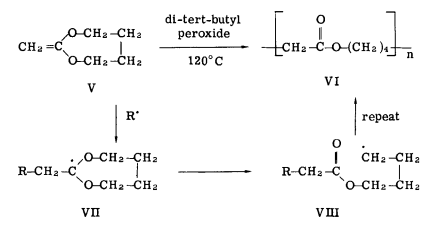


Any change in the system that would increase the steric hindrance of the cyclic radical would similarly increase the extent of ring opening by decreasing k_{11} . Similarly, any change in the system that would produce a more stable radical upon ring opening would be expected to favor the ring-opening process by increasing k_{150} .

One of the intriguing characteristics of the ethylene ketene acetal is its ability to copolymerize with a wide variety of common monomers, including styrene and methyl methacrylate. One should note that this process introduces an ester group into the backbone of an addition polymer. Although the copolymerization of oxygen would introduce a peroxide linkage into the backbone, this is the first time that a relatively stable but yet hydrolizable functional group has been introduced into the backbone of an addition polymer [12].

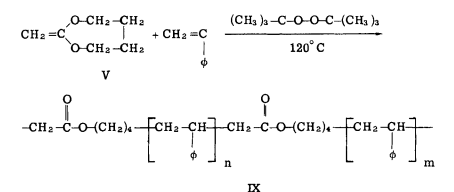


In a program to find other cyclic ketene acetals that would undergo quantitative ring-opening even at room temperature, we prepared the seven-membered ketene acetal, 2-methylene-1,3-dioxepane (V), which underwent essentially complete ring opening at room temperature [13]. This process makes possible the quantitative introduction of an ester group in the backbone of an addition polymer.



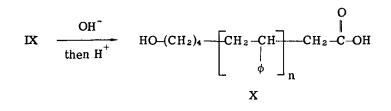
Apparently the seven-membered ring increases the steric hindrance in the intermediate free radical VII to eliminate practically all of the direct addition and also introduces a small amount of strain so that the ring opening to the radical VIII is accelerated.

When the seven-membered ketene acetal V was copolymerized with styrene, a styrene copolymer containing ester groups in the backbone was obtained ($r_1 = 0.021$ and $r_2 = 22.6$).



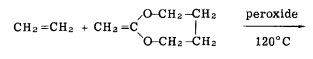
FREE-RADICAL RING-OPENING POLYMERIZATION

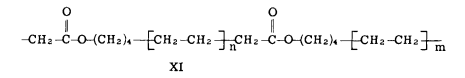
Hydrolysis of these copolymers makes possible the synthesis of oligomers of styrene capped with a hydroxyl on one end and a carboxylic acid group on the other [14].



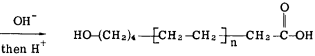
With a mixture containing about 80% V and 20% styrene, a copolymer containing 90 mol% styrene and 10 mol% ester-containing units was obtained. The hydrolysis of this copolymer gave an oligomer X of styrene containing an average of about nine styrene units end capped with the hydroxyl and carboxylic acid groups. Thus a very general method has been developed for the synthesis of a wide variety of oligomers with any desired molecular weight range. Of course, since the copolymers are random, the molecular weight distribution of the oligomers is quite broad. However, these oligomers should prove quite useful for the synthesis of polyurethanes and block polyesters.

Although most synthetic polymers are nonbiodegradable since they have not been on the earth long enough for microorganisms or enzyme systems to have evolved to utilize them as food, polyesters that are of relatively low molecular weight and rather low melting are biodegradable. This observation is related to the fact that poly- $(\beta$ -hydroxybutyric acid) occurs widely in nature and many microorganisms use this polyester to store energy in the same way that animals use fat. On the other hand, no synthetic addition polymer was known that was readily biodegradable. In an effort to produce a biodegradable addition polymer, the 2-methylene-1,3-dioxepane (V) and ethylene were copolymerized at 120°C for 30 min at a pressure of 1800 psi to give a low conversion of copolymers XI with estercontaining units varying from 2.1 to 10.4 mol%. The copolymers were, in fact, biodegradable, with the copolymers containing the high amount of ester groups being rapidly degraded and the copolymers containing only 2.1% comonomer only slowly degraded. Apparently there are enzymes in the microorganisms that are capable of hydrolyzing the ester linkages in the ethylene copolymer to produce the oligomers with terminal carboxylic acid groups; these oligomers are then degraded as analogs of fatty acids by the normal metabolic processes.





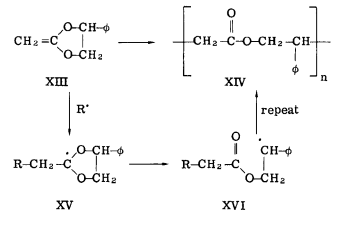
v



XII

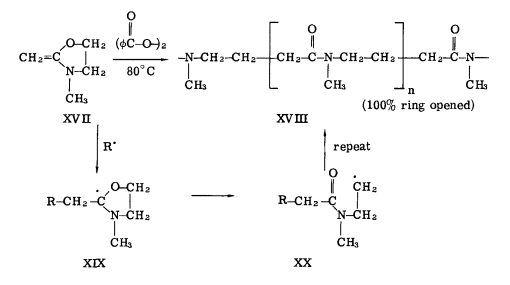
In a separate step the ethylene-2-methylene-1,3-dioxepane copolymer was hydrolyzed to give oligomers XII that were capped with a hydroxyl group at one end and a carboxylic acid group at the other. When the ester-containing unit was 2.1 mol%, the value of n was approximately 47, and when it was 10.4 mol% the value of n was approximately 9. The copolymers with 6 or less mol% of the ester-containing units had melting points in excess of 90° C.

Additional cyclic ketene acetals [17-19] that have been studied have included the 4-phenyl-2-methylene-1,3-dioxepane (XIII) which undergoes quantitative ring opening to give the polyester XIV. Apparently

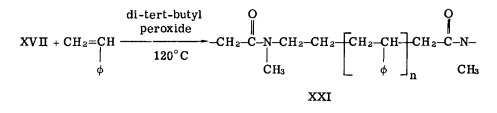


the ring-opening step from XV to XVI is greatly enhanced by the formation of the relatively stable benzyl radical in XVI even though XV is a five-membered ring analogous to the radical III.

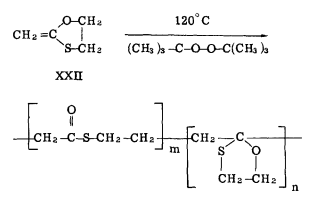
Since the nitrogen analogs of the cyclic ketene acetals were readily synthesized and would polymerize with essentially 100% ring opening, their copolymerization with a variety of monomers was undertaken [20]. Apparently the amide group is sufficiently more stable than the ester group that in contrast with the 2-methylene-1,3-dioxolane (I)



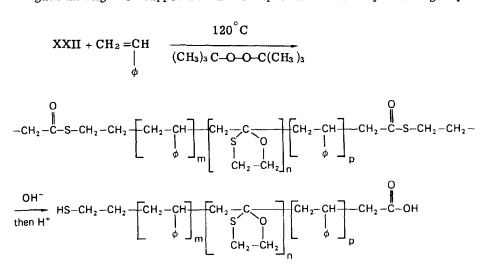
the nitrogen analog XVII undergoes essentially quantitative ring opening. Copolymerization of XVII with styrene give a copolymer XXI, which upon hydrolysis gives an oligomer capped with a methylamino group on one end and a carboxylic acid group on the other [21].



Although the sulfur analog of the cyclic ketene acetal I was prepared and polymerized, apparently the resulting thioester is of higher energy than the ordinary ester and therefore retards the extent of ring opening. Even at 120° C, only 45% of the rings were opened [22].

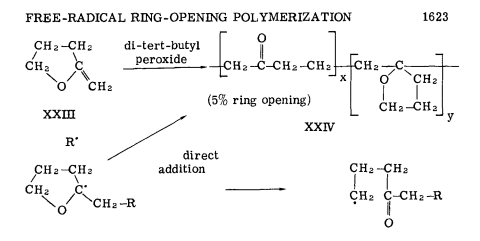


Nevertheless, copolymerization of XXII with styrene gave a copolymer containing some thioester groups, and hydrolysis of this copolymer gave an oligomer capped with a mercaptan and a carboxylic acid group.

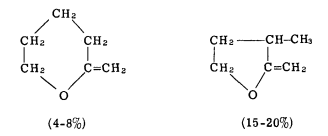


Since the nitrogen and sulfur analogs of the cyclic ketene acetal I gave interesting results, the study was extended to the carbon analog, the cyclic vinyl ethers. Thus, 2-methylenetetrahydrofuran (XXIII), when treated with di-tert-butyl peroxide at 120° C, gave a polymer XXIV in which only about 5% of the rings had opened [20]. Apparently the ketone group is sufficiently less stable than the ester group that the extent of ring opening decreases from 87% for I to 5% for XXIII. Even when the cyclic vinyl ether XXIII was diluted with an equal volume of benzene, the ring opening increased only to 7%.

Since the extent of ring opening of the cyclic vinyl ethers is less

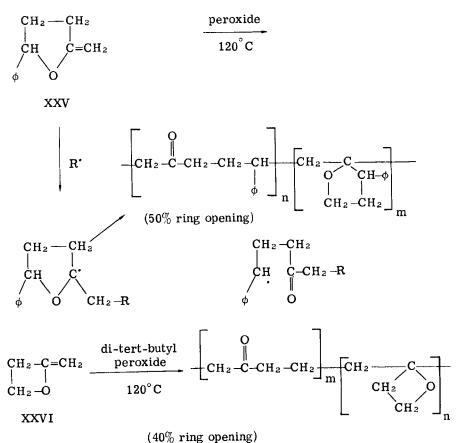


than that of the cyclic ketene acetals, this series appeared to an ideal system to study the effect of steric hindrance and the presence of radical-stabilizing substituents on the extent of ring opening. Thus the following monomers were prepared and polymerized at 120° C in the presence of di-tert-butyl peroxide. The number in parentheses indicate the extent of ring opening.



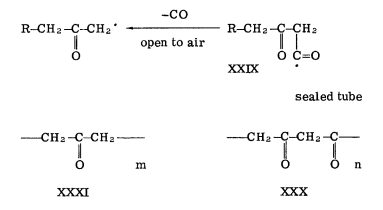
The highest extent of ring opening in this series was observed for the 4-phenyl-2-methylenetetrahydrofuran (XXV). In this case the formation of the relatively stable benzyl radical helps promote ring cleavage so that about one-half of the rings are opened [20].

Even ring strain will not produce quantitative ring opening in this series. Although the oxetane ring possesses considerable ring strain, the corresponding radical does appear to open at a rapid rate. For example, 2-oxetanyl gives no signals in the ESR spectrum for the ring-opened product, which is in contrast to the five- and six-membered acetals which give the ring-opened radical signals at room tempera-ture [23]. For example, the polymerization of 2-methyleneoxetane (XXVI) [24] at 120°C gave a copolymer in which only about 40% of the rings were opened [22]. Apparently the small size of the ring reduces the steric hindrance so that the direct addition can effectively compete with the sluggish ring-opening step.



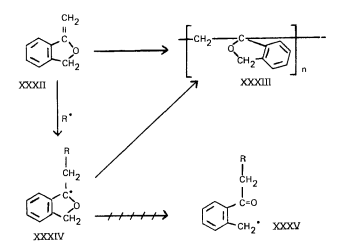
Ketene dimer (XXVII), which is an analog of the 2-methyleneoxetane (XXVI), will undergo a higher extent of ring opening on polymerization to give copolymers with different structures depending on whether the reaction is run in a sealed tube or open to the air.

 $\begin{array}{cccc} CH_2 = C - CH_2 & R' & R - CH_2 - C - CH_2 \\ | & | & & | & | \\ O - C = O & 120^{\circ}C & O - C = O \\ \\ XXVII & & XXVIII \\ & & & | \\ \end{array}$



The extent of ring opening is higher because the acyl radical XXIX is more stable than the primary radical that one would expect to obtain from the 2-methylene-1,3-oxetane (XXVI). In the sealed tube the acyl radical XXIX does not lose very much carbon monoxide to give a copolymer XXX which contains a large amount of the 1,3-diketone structure. On the other hand, the acyl radical XXIX, when the reaction is open to the air, can lose a substantial proportion of the acyl groups to give the copolymer XXXI is largely the 1,4-diketone.

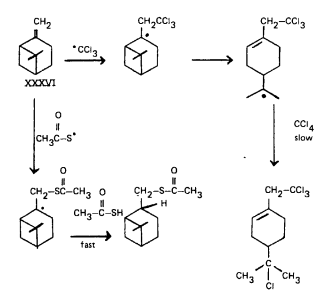
A surprising result was obtained from the polymerization of the 2-methylene-3, 4-benzotetrahydrofuran (XXXII); XXXII gives a polymer XXXIII with little or no ring opening. Apparently the intermediate radical XXXIV, which is a tertiary benzyl radical with addition stabilization from the oxygen, is sufficiently more stable than the

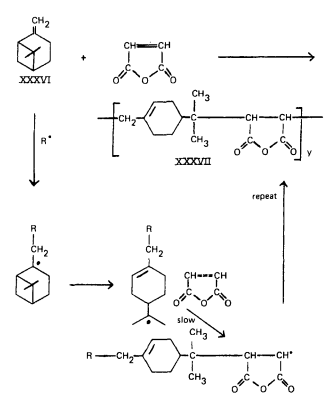


open chain primary benzyl radical XXXV that only direct addition takes place.

Copolymerization of several of these unsaturated cyclic ethers, particularly the 2-methylene-5-phenyltetrahydrofuran (XXV) with various monomers, is a convenient way to introduce a ketone group into the backbone of the addition polymer. Such polymers containing the keto groups can be expected to be photodegradable. The competive process involves the copolymerization of the monomer with carbon monoxide under high pressure [25].

In the previous examples discussed, the three-membered rings had sufficient ring strain to undergo ring opening even in carbocyclic systems but a five- or six-membered ring would open under radical conditions only if some other driving force was present to promote the ring opening. Since four-membered rings represent systems in between these two extremes, they appeared to be interesting cases for study. The ring opening of cyclobutylcarbinyl is exothermic (ΔH° = -4.0 kcal/mol compared to ΔH° of -5.1 kcal/mol for the ring opening of cyclopropylcarbinyl), and the E has been estimated at 13.0 kcal/ mol, corresponding to a k of about 4.5×10^3 s⁻¹ at 25°C, which is less than the values listed in Table 1 for cyclopropylcarbinyl ring opening. Finally, cyclobutylcarbinyl radicals can be observed in ESR at temperatures [26] at which cyclopropylcarbinyl radicals undergo ring opening. Studies with β -pinene show that ring opening can compete with relatively slow abstraction reaction but cannot compete with relatively fast abstraction reactions. For example, β -pinene will react with carbon tetrachloride to give a ring-opened product [27] but with thioacetic acid to give the nonring-opened product [28].

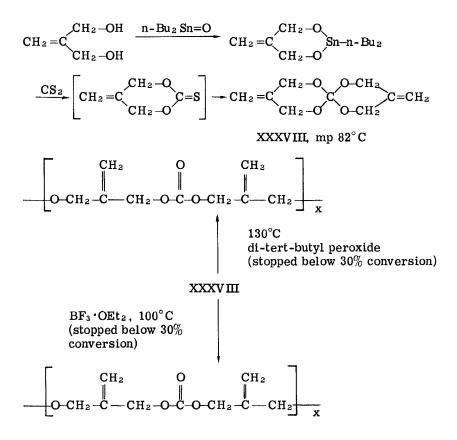




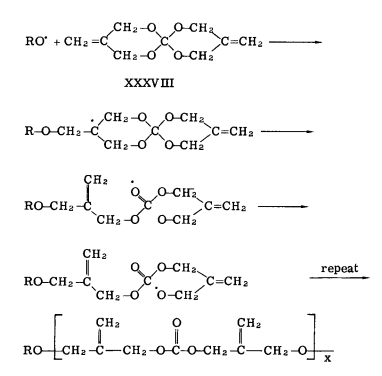
It was reasoned that if β -pinene was copolymerized with a monomer in which the addition step was relatively slow and involved some steric hindrance, there was a good chance that a copolymer containing ringopened units could be obtained. Thus maleic anhydride appeared to be an attractive choice since it does not have a high tendency to homopolymerize and is bulky enough to give considerable steric hindrance in the addition step. When equivalent quantities of β -pinene and maleic anhydride were heated at 130°C in the presence of di-tert-butyl peroxide, an essentially regular alternating copolymer containing the ringopened β -pinene units was obtained.

DOUBLE RING - OPENING POLYMERIZATION

In a program to develop monomers that expand upon polymerization, we had prepared a series of spiro ortho esters, spiro ortho carbonates, trioxabicyclooctanes, and ketal lactones that could be polymerized with ring opening by ionic expansion in volume [29]. Since a large proportion of industiral polymers are prepared by free-radical polymerization, it was desirable to have available a series of monomers that would polymerize by a free radical process with no change in volume or slight expansion. Such monomers could be added to common monomers to produce copolymers with reduced shrinkage or expansion in volume. It was reasoned that the introduction of unsaturation into a spiro ortho carbonate would permit double ring opening with expansion in volume. For that reason we undertook the synthesis of 3,9-dimethylene-1,5,7,11-tetraoxaspiro[5.5] undecane (XXXVIII) by the following set of reactions [30]:



It was found when this monomer was treated with di-tert-butyl peroxide at 130° C and the reaction stopped below 30% conversion, a soluble polymer was obtained having a structure of a polycarbonate with pendant methylene groups. The structure of the polymer was established by elemental analysis as well as infrared and NMR spectroscopy. A very similar polymer could be attained by treatment of the monomer with boron trifluoride etherate at low conversions. The mechanism of the polymerization appeared to involve a radical double ring-opening according to the following mechanism:



The driving force for the double ring-opening polymerization apparently is the relief of the strain at the central spiro atom as well as the formation of the stable carbonyl group [31].

At high conversion this monomer produced a highly cross-linked resin very similar in appearance to the material produced from the polymerization of diallyl carbonate. As indicated in Fig. 1, the volume change that occurred during homopolymerization was quite unusual. At room temperature a 4.3% expansion in volume occurred, while just below its melting point at 70° C, a 7% expansion in volume occurred; at 85° C a 2% expansion took place; and the expansion decreased until above 115° C a slight shrinkage occurred. It is obvious from these data that the large expansion in volume that occurs below the melting point involves not only the increase in volume due to the double ring opening but also a change in volume of 3-6% due to the process of going from a crystalline monomer to a liquid monomer. Since the monomer is a crystalline solid, it is difficult to find examples of homopolymerization in which the full 7% expansion in volume can be utilized. However, in copolymerizations it is possible

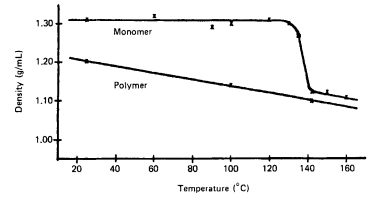
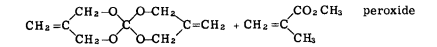
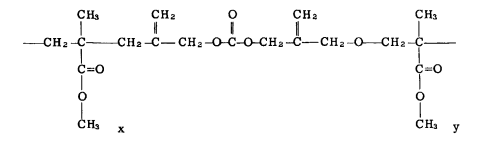


FIG. 1. Densities of the monomeric spiro ortho carbonate and related polyoxycarbonate versus temperature.

to use a slurry of the crystalline monomer in a liquid monomer so that as copolymerization progresses, the crystalline monomer dissolves with some expansion and also polymerizes with expansion [32-37].

The dimethylene spiro ortho carbonate XXXVIII was shown to copolymerize with a variety of monomers, including methyl methacrylate, hydroxyethyl methacrylate, styrene, and diallyl carbonate. With methyl methacrylate, XXXVIII was shown to be less reactive





with reactivity ratios of $r_1 = 0.87$ and $r_2 = 16.4$. Since $r_1 r_2$ is greater than 1, this fact was interpreted as involving association of XXXVIII in the monomer mixture rather than existing in a homogeneous solution. In this copolymerization, if the monomer mixture contains 10%

XXXVIII and 90% methyl methacrylate and the reaction is stopped at 69% conversion, the copolymer contained only 1% of the carbonate units.

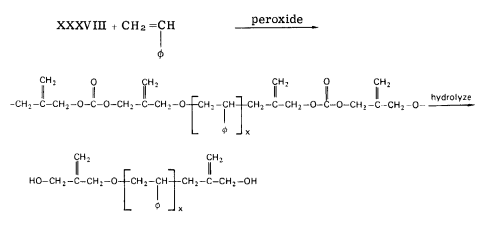
As discussed earlier, the free-radical ring-opening copolymerization of the dimethylene monomer XXXVIII with methyl methacrylate introduces a functional group into the backbone of an addition polymer. It was obvious that the introduction of even a small amount of this unit would greatly affect the chemical and physical properties of the base polymer. For example, polymethyl methacrylate is somewhat thermally unstable with the mechanism of decomposition involving a depolymerization process. It has been shown that, when a break occurs in the chain to form a radical above 225°C, reverse polymerization will take place to convert essentially the entire polymer molecule to monomers. Since the introduction of ethylene oxide units into polyformaldehyde produces increased thermal stability, it was reasoned that the introduction of the ring-opened units into polymethyl methacrylate would also result in increased thermal stability. Thus, if a break occurred in the polymer chain, depolymerization would proceed until the radical end reached a ring-opened unit. Since it is unlikely that the ring would reclose, and there is no other simple process for the removal of this end unit, the unzipping process will stop and further decomposition of that polymer molecule will cease. The fact that this prediction actually occurs is shown in Table 2.

The fact that the presence of 1% of the ring-opened unit in the copolymer reduces the weight loss to less than one-half that of the homopolymer is very encouraging. Presumably, any of the monomers that have been discussed previously that will undergo free radical ringopening copolymerization, including the 2-methylene-1,3-dioxepane (V), will also serve to stabilize polymers against reverse polymerization.

Since the carbonate group is a relatively easy group to hydrolyze, several other uses of these copolymers were suggested. A copolymer

| Amount of DSOC in copolymer (%) | Weight of polymer residue after 30 min at 225°C under N ₂ (%) |
|------------------------------------|--|
| 0 | 78.6 |
| 1 | 89.8 |
| 3 | 94,9 |
| 8 | 96.3 |

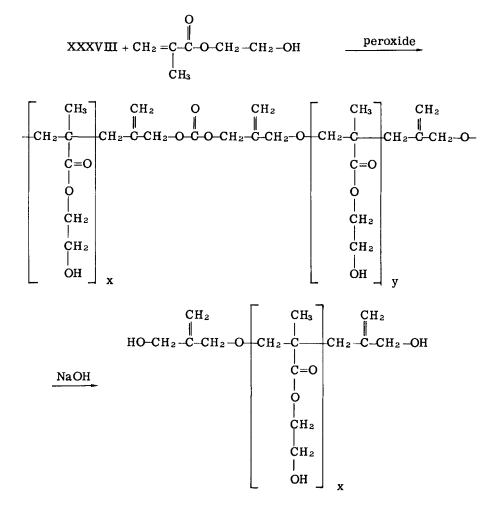
TABLE 2. Thermal Stability of Copolymers of Methyl Methacrylateand Dimethylene Spiro Ortho Carbonate



of the dimethylene monomer XXXVIII and styrene was hydrolyzed to produce a hydroxy-terminated polystyrene. The use of other monomers that polymerize by ring opening should permit the synthesis of a variety of addition polymers containing a wide assortment of functional endgroups.

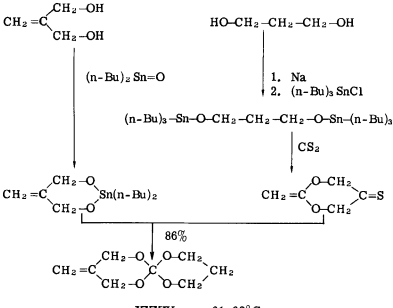
Still another use of the presence of a functional group within the backbone of an addition polymer is to render the copolymer biodegradable. When the dimethylene monomer XXXVIII and hydroxyethyl methacrylate (HEMA) were copolymerized to produce a copolymer containing 14 mol% of the ring-opened units, hydrolysis of the copolymer in an alcoholic solution containing 1% sodium hydroxide gave within 3 h at room temperature a copolymer with 1/5th of the original viscosity average molecular weight. Preliminary results indicate that this copolymer is biodegraded by microorganisms. For use in the human body, complete biodegradability may not be necessary since polymeric materials with molecular weights below 5000 can be eliminated from the body. Although some hydrophilic character is necessary for fast biodegradation, the introduction of a sufficient number of such functional groups in the backbone may render even hydrophobic polymers, such as polystyrene, slowly biodegradable.

A potential use of this monomer is in the area of dental fillings in which a slurry containing 20% of very fine crystals of the unsaturated spiro ortho carbonate XXXVIII and 60% of the adduct of methacrylic acid to bisphenol-A diglycidyl ether (Bis-GMA) plus 20% trimethylolpropane trimethacrylate produces a material with essentially no change in volume upon polymerization. An investigation of a bubble test on tooth enamel showed that this copolymer had nearly double the adhesion to the tooth structure that the base resin had without the addition of the unsaturated spiro ortho carbonate. The copolymer also had improved impact strength but yet essentially the same modulus, and filled composites appeared to have somewhat improved abrasion resistance [37].



Since the synthesis of the spiro ortho carbonates through the tin compounds could be modified to produce unsymmetrical materials, we undertook the synthesis of the unsymmetrical 2-methylene-1,5,7,11tetraoxaspiro[5.5] undecane XXXIX by the accompanying set of reactions.

The resulting monomer was a crystalline solid with a melting point of 61-62°C. When the polymerization was carried out in the presence of di-tert-butyl peroxide and the reaction was stopped at low conversion, a linear polycarbonate containing pendant methylene groups was obtained. The structure of the polymer was established by elemental analysis as well as infrared and NMR spectroscopy. The structure of this material was very similar to the polymer that could be obtained

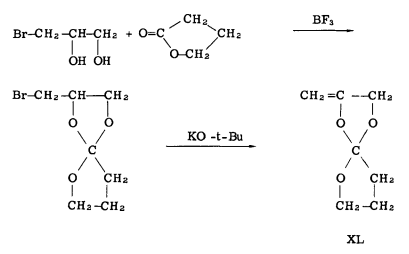


XXXIV, mp 61-62°C

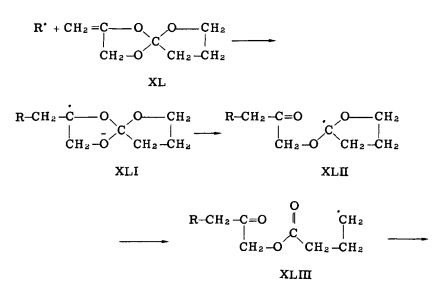
by the ionic polymerization of this same monomer at low conversions. Bulk polymerization of XXXIX with peroxide catalyst gave a material at 25°C with an expansion of 4.5% and at 60°C an expansion of 5.5%; above the melting point of XXXIX (61-62°C) the expansion decreased until at 111°C the density of the monomer and the density of the polymer were the same.

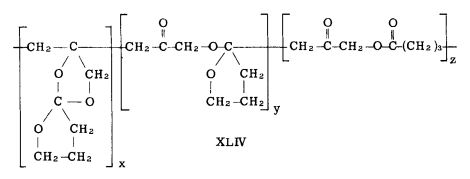
XXXIX $\frac{\begin{array}{c} \text{di-tert-butyl} \\ \text{peroxide} \end{array}}{130^{\circ}\text{C}} \quad \begin{bmatrix} \text{OCH}_2 - \text{C} - \text{CH}_2 - \text{O} - \text{C} + \text{O} - \text{CH}_2 - \text{CH}_2 - \text{CH}_2 \\ \parallel \\ \text{CH}_2 & \text{O} \end{array}} \\ \begin{bmatrix} \eta \end{bmatrix}_{\text{CHCl}_3}^{25^{\circ}} = 0.11 \end{bmatrix}$

Since a variety of unsaturated heterocyclic monomers will undergo ring-opening polymerization quite readily, it appeared undesirable to prepare a related spiro ortho ester for use as a monomer with either no change in volume or slight expansion. Thus an unsaturated spiro ortho ester was synthesized by the accompanying reaction. Polymerization of the 2-methylene-1, 4, 6-trioxaspiro[4, 4] nonane with benzoyl



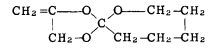
peroxide gave a polymer with a complex structure that indicated that at 120° C only about 10% ring opening had occurred. Apparently the radical adds to the double bond to give the intermediate XLI which can add to another monomer or undergo ring opening to give the semiopened radical XLII.





XLII, in turn, can either add to a monomer or undergo a second ring opening to give the double ring-opened radical XLIII. In fact the final polymer XLIV has about 90% unopened units and about equal quantities of the semiopened and double-opened units.

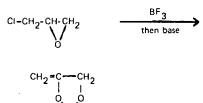
When the related unsaturated spiro ortho ester XLV derived from ϵ caprolactone was polymerized, it gave a similar structure containing slightly more of the double ring-opened units but still about 90% of the



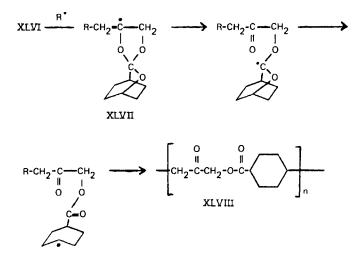
 $\mathbf{X}\mathbf{L}$

unopened units. However, when the strained unsaturated spiro ortho ester XLVI was polymerized, essentially complete double ring opening occurred.





XLVI



Apparently the intermediate radical XLVII has sufficient strain to promote the ring opening. The resulting polymer XLVIII is intriguing since it contains both ester groups and keto groups. Such a polymer should be both biodegradable and photodegradable.

Thus we have been able to demonstrate that free radical ringopening of unsaturated heterocyclic monomers is a very general phenomenon. This reaction has been shown to be, for the first time, a way to introduce functional groups such as esters, amides, thioesters, carbonates, and ketones into the backbone of an addition polymer. As a consequence, this copolymerization is a useful method for the synthesis of oligomers that can be capped on both ends with reactive groups such as carboxyl, amino, mercapto, and hydroxyl for the synthesis of biodegradable and photodegradable polymers, and for the stabilization of copolymers against decomposition by reverse polymerization. Free radical double ring-opening polymerization is useful to control the shrinkage of monomers during free radical polymerization.

ACKNOWLEDGMENTS

This work was supported in part by the Polymer Program of the National Science Foundation, the Frasch Foundation, and the Goodyear Tire and Rubber Company.

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