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INTRODUCTION

Atmospheric oxygen is the most prevalent as well as the most economically important of the oxidising agents for the fats and fatty acide. In some cases the action of oxygen is highly desirable, as in the production of the blown paint oils, while in other cases its action is to be avoided if it is at all possible. For example, the process of film formation is one involving oxidation, but the failure of these same films is due to prolonged exposure to atmospheric oxygen.

Both the favorable and the unfavorable results of the oxidative process follow the same type of reaction between oxygen and the unsaturated centers of the fatty acide. Because of this, an enorance amount of work has been done in an attempt to determine the mechanism involved in autoxidation so that these reactions may be better controlled. Theoriee regarding the course of the reaction are many and varied, and considerable experimental evidence has been offered in support of the hypothness.

Modern theories conserving the exidation process are generally considered to date from the works of Bach (2) and Engler (15). Prior to their publications it was believed generally that melecular coxygen was partially dissociated into atomic oxygen and that this was the agent responsible for the slow extention observed in organic compounds. Bach and Ragler believed that the exygen involved was molecular and that the exygen added

at the site of the double bond to give a compound of the general formula ROOR. This compound could then in turn oxidise some other oxidisable compound.

Another investigator, Fehrion (14), assumed that the oxidation occurred in the fatty sold to form a cyclic percent which then rearranged to a dihydroxy ethylenic or hydroxyketo configuration in the following manners

Fahrion also assumed that the cyclic peroxides could give rise to products having a more stable configuration, such as

A theory of tautomeric ketchydroxy-dihydroxy rearrangement was proposed by Ellis (12) and was based on the detection of characteristic groups in linsed oil films. According to Ellis, autoxidation cocurs as a result of the addition of a molecule of oxygen at the double bond to form a dihydroxy compound which then rearranges to a ketchydroxy compound.

Still another scheme was that proposed by Fokin (80) in which the first step of the reaction was the addition of oxygen at the double bond to form an ethylene oxide ring.

This configuration is known to be formed in the oxidation of monochemoid eaids with perbensoic and peracetic eaids. Fuldence in support of Fokin's proposal is less extensive than that
for some of the other hypotheses. Aside from the work of Fokin,
soms support for the theory is found in the work of Saent-Gyorgyi
(40) who investigated the autoxidation of limbelenic acid in the
presence of sulfnydryl groups as estalyst and found that the
molecular weight of the product, 995, indicated one atom of oxygen per mole of limbelnic acid oxidized.

Standinger (87) has proposed a theory in which the peroxido reaction of Bach and Engler is the second rather than the first step in the autoxidation of a double bond. This was based on a study of asym-diphenylethylene which yielded a stable peroxide. The primary product of oxidation was termed a moloxide and could not be isolated. It was assumed that the oxygen added at the double bond to form the moloxide which then rearranged to form the peroxido compound.

Until recently this theory was widely accepted by the majority of the workers in this field. The evidence for the existence of such a structure has all been obtained indirectly on the basis of sertain analytical data. Unfortunately, all the analytical methods involved are not definitely quantitative and it is a well known fact that the exidation of a fatty acid leads to a

wide variety of products, making the proper interpretation of such analytical data quite complicated. It also should be noted that a product having such a structure has not been isolated and characterized as such from the oxidation products obtained. Those which have been characterized have all possessed structures quite different from the assumed heterocyclic peroxidic structure.

In 1986, Criegee (10) suggested that cyclohexene autoxidized to form a hydroperoxide having the structure

Following this, Ricoho (35) suggested that unasturated fats and oils probably behaved in a similar manner. According to Ricoho the autoxidation of mono- or polyumesturated compounds could occur through the formation of oxygen-activated methylane groups.

$$-CE = CHCE_2CE = CH- + O_2 \longrightarrow -CE = CHCHCH = CH-$$

The hydropercxide mature of certain types of clefinic compounds has been substantiated by the works of Orieges (10) and Stephens (38) on cyclohexene and also by the work of Orieges, Fils and Flygare (11) and of Hook and Susemini (28) on tetralin.

The development of the hydroperoxide theory of autoxidation has been credited generally to the works of Farmer (15,16) and Farmer et al. (17,18,19). Farmer (15) originally postulated that the autoxidation of practically all unconjugated ciefinic compounds proceeded by the addition of a molecule of oxygen to the carbon atom in the O-position with respect to the double bond forming a hydroperoxide and leaving the double bond intact.

Farmer and Sutton (10), in studying the oxidation of methyl oleate, found that a mixture of mono- and di-hydroperoxides was produced, the hydroperoxide group being attached at 0g, 01, or at both. Each of these hydroperoxides contained either one atom of active oxygen and one atom of active hydrogen or two atoms of active oxygen and an intest double bond. The peroxidised methyl cleate was estimated to not exceed 70 percent purity but refractionation gave a product which was estimated to be mearly 100 percent pure methyl peroxido cleate.

This work of Farmer and Sutton was repeated by Swift, et al.
(39) who reported that the product obtained possessed the properties and responded to the reactions described by Farmer and Sutton. The identity of the product was further substantiated by the findings of Atherton and Hilditch (1) who subjected the peroxidised methyl cleats to further oxidation with powdered potassium permangante. Among the products they identified were four acide, suberic, aselaie, octanoic and nonanoic, which were predicted by Farmer as some of the solssion products of hydroperoxido cleic acid.

Parmer et al., (17,18) believed that the reaction proceeded

by way of a free radical mechanism in which an active methylene group is somehow dehydrogenated to form a free radical. This free radical then absorbs oxygen, forming a new free radical which then adds a hydrogen atom to form the hydroperoxids.

$$\circ$$
CH $_{2}$ CH = CH $_{-}$ \longrightarrow \circ CHCH \longrightarrow \bigcirc CHCH \longrightarrow \circ CHCH \longrightarrow \bigcirc CHCH

According to Farmer, Eoch and Sutton (18), the first step in the autoxidation of an clefinic system of the methyleme-interrupted type, such as is characteristic of the drying oil acide, was the severance of a thormally or photochemically activated C-H bond in the Q-position with respect to the double bond leaving an olefinic free radical. This free radical could exhibit resonance between the two forms.

so that the addition of oxygen would most likely occur at C₁ or at C₂ and the double bond would then either remain at its original position or appear at the adjacent carbon atom. The most active methylens group was postulated to be the one flanked on either side by a double bond, such as the CH₂ group at C₁₁ in limbaic acid.

In 1946 Farmer (16) modified his original hypothesis. In view of the fact that approximately 80 keal of energy are required for the severance of a C-H bond and because of the apparent sase with which the oxidation occurs, the original hypothesis did not seem to be the most plausible. Experiments had shown

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that in the case of conjugated compounds, even though the senters of unsaturation were flanked on either side by active methylene groupe, autoxidation cocurred additively leaving the methylene group intact. Because of this and remembering that the monomeric peroxides of simple elefins and unconjugated polyeletine are mainly questhylenic hydroperoxides, there seemed to be justification for postulating universal initiation by addition at the double bond and a continuation of the attack substitutively by means of a chain reaction. The high expenditure of energy required for severance of the C-H bond would thus be reduced. By this scheme the actual extent of addition would be insignificant but would be sufficient to start the necessary chain reactions.

It should be noted that intramolecular reaction of a type permitting the first formed di-radical, -GH₂GHGH(00)-, to become stabilized by the detachment of a hydrogen atom from the adjacent G-methylene group to form -GH = GHGH(00H)- is highly unlikely since it would not propagate reaction chains. There was also an indication of spontaneously occurring chain scission reactions which could be interpreted as indicating two kinds of exidative attack occurring simultaneously.

Since, under proper conditions it is possible to obtain hydropercoridee in mearly quantitative rields, Solland and Gee (6,7) have made a study of the amtoridation mechanism kinetically and thermochemically. The studies were carried out on ethyl linoleste, ethyl linoleste, methyl cleate and equalene. Absorption spectra in the quarts ultra-violet region indicated that the oxidation was accompanied by a displacement of the double bonds. This displacement was confined to oxidised molecules and, in the case of the linoleste ester, the primary product of oxidation was a confugated hydropercoxide. In each case the process was identified as a chain reaction in which hydrocarbon radicals were formed, absorbed caygen, and then reacted with a second molecule of olefin to form a hydropercoxide and a new free redical.

From their experiments, Bolland and Gee determined the energy of activation from the temperature dependence of the velocity constante obtained from kinetic data. They also have estimated the heate of reaction for various reaction paths, using the formal strengthe of bonds broken and formed and making corrections for the estimated resonance energies of the reactante and products. These were then used, along with the activation energies, in an attempt to determine the most likely path of the reaction. Two possible points of attack were considered, attack at the double bond and at an active methylene group. For a mono-olefin these reactions may be formulated as follows:

In the case of a 1:5 di-elefin, attack at one end of the double bond would be assisted by the development of allyl resonance while the other path, (b), would be assisted by a somewhat greater gain of resonance energy. For a 1:4 di-elefin, attack at the double bond would be equivalent to that in a mono-elefin, but attack at an active methylene group would be easier because of 11 kilocalories resonance energy in the radical. The values of resonance energies employed in the calculations were

0 = 0 = 0 0 = 0 = 0 0 = 0 = 0 = 0 0 = 0 = 0 = 0 0 = 0 = 0 0 = 0 = 0 0 = 0 = 0 0 0 = 0 0 0 = 0 0 0 = 0 0 0 0 =

and the bond strengths used, though not identical with those of Pauling (54), were

C-C 81 keal C = C 145 keal C-H 90 keal C = C 174 keal C-O 87 keal C = C (Ogmolecule) 118 keal

Using these values, the estimated heats of reaction were tabulated as follows for the various elefins considered.

0-0 (peroxide)

66 koal

| Hono- | 115 | 114 |
5 Ha | 14 kcal | 2 kcal | 14 kcal |
5 Hb | 7 kcal | 3 kcal | -4 kcal |

0-H 110 kenl

These reactions have beets lying below the experimentally determined energies of activation, 28 keal, so that none of them can be ruled out on this basis. The order of reactivity of double bond attack was then determined as 1:8>mono- or 1:6; while for \(\alpha\) enthylenic reactivity the order was 1:6\1:5\mono-. The two reaction paths were thus postulated not to be mutually archainty and would be expected to proceed simultaneously.

The double bond displacement which cocurred during oxidation was explained by Bolland and Gee on the basis that one of the chain carriers was the radical ROH = CHPGCH = CHR!, since this radical will possess two other resonance hybrid configurations giving rise to a large resonance stabilization and permitting the formation of isomeric peroxide radicals on the attack of oxygen. The chain cycle was believed to be

The estimated heats of reaction for reactions (c) and (d) for the olafins considered were

Because of the exothermic character of these reactions, they would be highly probable steps in a chain reaction and were consistent with the kinetically deduced energies of activation, It has not been possible in all cases to isolate the hydroperoxide materials of the autoxidation, for, as pointed out by Farmer (16), chain scission reactions apparently set in unpreventably at the outset of the reaction, possibly by a direct mechanism of scission or occurring simultaneously with a less direct mechanism. Little study has been given to the course of the decomposition or chain scission reactions but experience has shown that the course is greatly affected by experimental conditions; i.e., the presence of catalysts, the acidity or alkalinity of the medium, the temperature at which the reaction is carried out, and incident radiation. For example, clefin hydroperoxides decomposed in acid media have been found to give rise to triols, probably derived from the corresponding epoxides, while an alkaline media leads to the formation of hydroxylated scission products, yielding aldebydes, ketones, and acids.

A polymeric tendency is also of rather general occurrence as a secondary reaction of the autoxidized material, for even in the most favorable examples of hydroperoxide formation, there usually has been an appreciable polymer fraction which may partially consist of dialry peroxides.

ROO- + RH --- ROOR + H-

Bolland and Gee (7), have pointed out that polymers sould be formed by the radicals formed in the direct attack of oxygen at the double bond and an unoxidized molecule of olefin.

The free radical ends thus produced could then react further with oxygen to again produce peroxide radicals which could then add olefin and thus build up a polymeric chain.

There is very little direct experimental evidence in support of any of the proposed polymeric reactions. Whatever the mechanism of the reaction, certain factors must be taken into consideration: (a) the autoxidation mechanism must be considered, (b) unoxidised earbon to carbon double bonds must not be involved, and (c) the general requirements of the general polymerisation theory must be met.

In recent years, infrared spectroscopy has found wide application as an aid in the solution of various problems of both theoretical and practical importance and has been recently reviewed by Barnes et al., (4). The applications of infrared spectroscopy can be divided into two general eategories, qualitative and quantitative. Qualitatively it has been used in the identification of compounds, in the recognition of specific chemical bonds, linkages or groups, and also in the fields of deutero-chemistry, polymerization, and isomerism. The qualitative analysis of mixtures is also facilitated by the study of the absorption spectrum, for the absorption spectrum of a misture, except in the case of strong molecular interaction, is equivalent to a superposition of the spectra of the various components of the mixture. Quantitatively, infrared epectroecopy is quite useful in the analysis of mixtures, for, if the nature of the components is known, it is usually possible to determine a frequency at which only one of these will absorb strongly, facilitating the determination of its concentration by comparteon with standards. This ease principle also has been applied in the study of reaction rates as a function of temperature, pressure, or catalyst. Gertain thermodynamic constants also may be calculated from a knowledge of the infrared absorption frequencies.

When light of various wave-lengths is passed into a group of molecules, some of this light may be absorbed and some will page through. Absorption can occur only if the molecule is capable of a rearrangement of ite possible motions in such a way as to take up the energy associated with the incident radiation. A molecule is capable of three types of internal motion: motion of the electrons surrounding the mucleus, motions of the muclei with respect to one another, and a rotation of the molecule as a whole. The energies associated with the three types of motion are of entirely different magnitudes, making it possible to correlate one of these types of motion with a particular region of the electromagnetic spectrum. The ultraviolet spectra are said to be due to electronic motions, the near infrared are due to muclear vibrations, and the far infrared are due to molecular rotations. These regions are not sharply defined and there is considerable overlapping. The effect of electronic motions on

vibrations of the nuclei may be neglected for these are associated with relatively high energies.

Not all molecules have far infrared spectra and some few have no spectra even in the near infrared. Radiation may be absorbed or emitted only by a system whose electrical configuration is changing relative to a point in space, meaning that a vibration can result in absorption only if the electric moment of the molecule changes in the course of that vibration, and similarly for rotational absorption. According to the quantum theory, the frequency of vibration and rotation are limited to certain discrete values determined by the nature of the molecule so that there are only certain frequencies at which absorption can occur. Thus by irradiating a sample with a series of monochromatic bands of infrared radiation and plotting the percent of the radiation absorbed or transmitted as a function of either wave-length or frequency, the resulting graph may be interpreted in terms of the intramolecular motion of the molecule and is known as the infrared spectrum of the substance under investigation.

A molecule containing a stome will have Se degrees of freedom. Of these, three are associated with translation of the molecule as a whole and three are associated with rotation, so that the molecule will them possess Se-6 internal degrees of freedom which will express themselves as Se-6 normal modes of wibration provided that the molecule is mon-linear. For a linear molecule there will be Se-6 normal modes of wibration since there are only two degrees of freedom associated with the rotation of a linear molecule. A normal vibration is defined as a vibration in which the center of gravity of the molecule does not move, and in which all the atoms move with the same frequency and in phase. Bormal modes of vibration need not be independent, for frequently several degenerate modes are found, depending on the symmetry of the system.

The atoms of any molecule not at absolute sero are constantly oscillating about their equilibrium positions. These oscillations are of extremely small amplitude and high frequency. The frequencies are of the same order of magnitude as are those of infrared radiation so that some relationship might be expected to exist between the motions of the atoms and their effects upon incident infrared radiation. It also might be expected that a mathematical calculation of the normal modes of vibration is possible, which should permit a unique determination of the structure of the molecule. The correct structure of the molecule would then be that for which the calculated frequencies correspond exactly to those observed experimentally. These calculations can be carried out if all the interatomic forces are known, the complexity of the calculations depending on the number of atoms and their symmetry. This has been done for a few molecules in which the number of atoms is small or which possess a high degree of symmetry. However, since most molecules of intorest to the organic chemist are relatively complex, some other method must be utilized. Considerable success in this direction

has been made by the purely empirical approach of Barnes etal.,

As has been pointed out by Hersberg (24), it is the bond force constant and not the frequency which is characteristic of a specific bond. If the restoring force between two atoms is the same in one molecule as in another, the conclusion would be that the electronic structures are the same in the two cases. And conversely, if the structures are the same then the force constants should also be the same. This actually has been found to be the case. For example, the bond-stretching force constant in HCN, ClCN, BrCH and (CH), is approximately 17 x 10 dynes per on throughout the series. Similar results have been obtained for other bonds. However, this principle holds strictly only if the bonds are in similar surroundings, since the force constants will be affected by such factors as the proximity of unsaturation and the degree of such unsaturation. This same principle holds for the bond-bending force constants and in this case the condition of similar surroundings is even more critical. It is not sufficient that the same bond type be adjacent to the bond under consideration, the atoms at the other end of the adjacent bond also must be the same.

Grawford and Brinkely (9) have shown for a number of molceules, that, by taking over not only the bond-stretching and bond-bending force constants but also the interaction constants from other molecules with the same groups, some or all of the mormal frequencies may be predicted to within 1 or 2 percent of the observed values. Such a procedure is of great importance since in this way it is possible to determine the correct assignment of the observed frequencies in the more complex molecules or even to obtain the approximate values without actual observation. With increasing accuracy of observation and calculation, slight differences in the bond force constants are to be expected since the surroundings of a given group are not the same in difference because.

The application of these principles to a correct assignment of the observed vibrational frequencies is greatly simplified by a corollary of the principle of invariance of bond force constants, namely the constancy of bond or group frequencies in different molecules. This was first established by observation and later explained theoretically on the basis of the constancy of bond force constants.

The observation of characteristic frequencies had led Necke (\$1,38,38) to the introduction of the concept of valence and deformation vibrations; that is, the idea that to every bond in a molecule there corresponds a vibration in which this bond is stretched and another vibration of smaller frequency in which it is bent. An explanation of this can be given on the backs of the mechanical molecular models of Kettering, Shutts and Andrews (37) or simply by a consideration of the atoms taking part in the vibration. If the 0-H, 0-H, and N-H vibrations are considered, it is seen that the masses of the other atoms are so much greater than the mass of the hydrogen nucleus that the smalltude

of vibration of the hydrogen mucleus will be much larger than those of the muclei of greater mass. Therefore, to a first approximation, the hydrogen muslaus may be considered as oscillating against an infinitely large mass so that the vibration frequency depends practically only on the force by which the hydrogen atom is bound to the remainder of the molecule and will be nearly the same for different molecules with the same C-H. O-H or N-H force constants. Since the hydrogen atom is always an end atom, it can move only in the line of the bond or perpendicular to it; that is, it can undergo only stratching or handing vibrations of frequencies corresponding to the respective force constants. If two such groups are present, or more than two, these will vibrate simultaneously and there will then he several normal frequencies which will be of nearly the same marnitude as for a single group, differing only by a small amount which is an indication of the strength of interaction of the equivalent groups. Apart from the end atoms thus far discussed. experiments have shown that in many cases, bonds involving heavier atoms have characteristic frequencies even if none of these are end atoms.

The degree of definition of the characteristic frequencies also depends on the angle between successive bonds in the chain of atoms. It has been shown Partholome and Teller (5) that if the angle is 90 degrees there will be no transfer of vibrational energy from one bond to the next so that the characteristic frequencies will then be best defined. In the case of the bond bending frequencies, the chance that in a molecule there is another frequency, not necessarily a bending frequency, of similar magnitude is much larger than for the bond-stretching frequencies. Thus, the bending frequencies are often not very characteristic.

In case a chain-like molecule has two or more equivalent bonds, there will be a resonance which leads to a splitting of the characteristic frequency and the stronger the coupling, the greater will be the degree of splitting. Also the greater the number of equivalent bonds, the greater will be the splitting into a correspondingly larger number of frequencies which will deviate increasingly from the characteristic frequency of a single bond of the same type.

Using these principles it is possible to obtain approximate values for some of the frequencies of a molecule, particularly an organic molecule, without actual observation, if the war-acteristic stretching and bending frequencies of the bonds in the molecule are known. Such predictions, even though only approximate, are often quite helpful in the analysis of observed vibrational spectra.

As shown by Hersberg (28) the vibrations may be considered to obey Hocks's law. Therefore, any equation set up to describe these motions must be dimensionally similar to that which holds for simple harmonic vibrations, $\hat{\nu} = (1/2\pi e)/\frac{1}{k}$, where $\bar{\nu}$ is the requesty in m^{-1} , e is the velocity of light, μ is the reduced mass of the vibrating atoms and k is the force constant existing

between the two atoms. The reduced mass is given by the equation $\frac{1}{h} = \frac{1}{h_0} + \frac{1}{h_0}$ where m_1 and m_2 are the masses of the vibrating atoms. The constant k is related to the vibration by the equation $v = \frac{1}{h}(x^2)$, where v is the potential energy and x is the displacement of the atoms from their equilibrium positions. Substitution of the proper constants reduces the dimensional equation to $\vec{j} = 1307 \sqrt{k_0}$ cm⁻¹, where k is expressed in units of 10^5 dynes on and μ is expressed in atomic mass units,

The value of k for practically all single bonds lies between 4 and 6 x 105 dynes per cm. The corresponding value for double bonds lies between 8 and 12 x 105 dynes per cm. while those for triple bonds generally lie between 12 and 18 x 105 dwnas per cm. These values, along with the proper values of \mu may be used to calculate the approximate vibration frequencies for a few twoisal atom pairs. as shown by Barnes et al. (4). For the C-H of methane the value of k has been found to be almost exectly 5 x 105 dynes per cm. The carbon atoms, because of its large mass relative to that of hydrogen, remains essentially at rest, and μ is close to 1. Substitution of these values gives $\bar{\nu} = 1307 \sqrt{5/1} =$ 2920 cm-1. Similarly, for the C-O of methyl alcohol, the value of k is still very close to 5 x 105 dynes per cm. but the value of M is now 6.85 so that the frequency will be lower than in the case of the C-H bond, $\nu = 1507 \sqrt{5/6.85} = 1110 \text{ cm}^{-1}$. In the infrared spectra of these compounds there are strong absorption bands at 2915 and 1054 cm-1. Also, a study of a series of compounds containing these linkages shows the characteristic bands

at approximately the calculated frequencies. Thus it is possible in many cases to use the exact frequency empirically as a
means of telling whether or not the particular atomic group is
present and also its relationship to the rest of the molecule.

And conversely, once the absorption frequency has been definitely attributed to a particular atomic group, this frequency may
be used to calculate the value of k, the bond force constant.

This method of extracting a portion of the molecule and treating
it mathematically is, however, only an approximation and should
not be carried to extraces. The method is applicable to only a
few compounds. Barnes and his associates (4) have used an empirical approach and examined the spectra of several hundred
organic compounds. Only rarely have they failed to find correlations between bond frequencies and structural relationships.

It can thus be seen that infrured spectroscopy offers promise of improved methods in blochemical and medical research but very little work has as yet been reported. Mention has been made by Jones (86) to investigations dealing with the application to analysis of sterole and fatty acids. The infrured spectra of limoleic and limolenic acids have been determined, the only noticeable difference being at 0.50 microns (1168 cm⁻¹). Fromination of these acids gave a much greater difference in the spectra, as would be expected. Jones also states that preliminary measurements had been made on some of the epoxy and hydroxy acids obtained by a controlled caridation of limoleic acid and had indicated a possibility of following the successive

stages of oxidation of this soid.

McOutcheon et al. (30) have determined the infrared absorption spectra of the ethyl esters of claic, elizido, limcleic, limcleidic and limclenic acids. They observed strong absorption in the region of 1667 cm⁻¹ in the case of the esters of the claic, limcleic and limclenic acids, while elizidic and limcle-laidic acids exhibited relatively wider and weaker bands in this region. For ethyl limclenate there was a secondary absorption in the region of 1006 cm⁻¹. From theoretical considerations ois-acids abould exhibit a strong absorption characteristic of the double bond in the 1667 cm⁻¹ region. Since the ethyl elaidate and limclelaidate did not absorb strongly in this region, the results were interpreted as proof of the cis-cis and trans-trans configurations of these two esters respectively.

Gemble and Barmett (31) have examined the infrared absorption spectra of the methyl, glycol and glyceryl esters of Oleic, linolesic, linolesic and elseosteric soids. The glyceryl esters all showed absorption in the region of 1607 cm⁻¹, characteristic of the double bond. In addition to this, the elseostearate showed an absorption at 1000 cm⁻¹ which the authors attributed to conjugation. These same authors determined the spectra of the methyl esters of cleic and elseostearia saids before and after exposure to ultraviolet light. A strong absorption band at 2940 cm⁻¹ was found to have shifted to 3520 cm⁻¹. Absorption at 3400 cm⁻¹ is generally attributed to the presence of an -CE group. The observed shift was thus attributed to the intro-

duction of -OH groups. More probably, however, the shift was due to the introduction of hydroperoxide groups, -OOH, which are readily formed in oleic acid when irradiated with ultraviolet light in the presence of oxymen.

It was because of the diversity of opinion as to the autoxidation mechanism and the inexessing use of infrared spectroscopy as an aid in such problems that the present investigation was undertaken in an attempt to add epectroscopic evidence to the chemical evidence supporting the hydroperoxide mechanism of confidence.

EXPERIMENTAL

Preparation of Materials

Unless otherwise stated, the methyl lineleate used in these experiments was obtained by the debromination of tetrahromostearic acid by the Rollett method (56).

Frequention of the Fatty Acid. Three hundred g of potassium hydroxide were placed in a five liter round bottom flask, 1200 ml of ethyl alcohol and 100 ml of water were added and the mixture heated to near boiling on a steam bath. One thousand g of cottonseed oil were then added, a reflux condenser attached, and the mixture allowed to reflux for 50 minutes. The mixture was then ecoled to about 40° 0. In a stream of tap water after which 1500 ml of cold distilled water were added. Five hundred fifty ml of concentrated hydrochloric sold were then added in small portions with cooling and agitation. The mixture was then placed

in a large separatory funnel and shaken to insure decomposition of the scape. The fatty acids were washed trice with about 1 liter of cold distilled water, being careful to avoid emilsification. After drawing off the water, the fatty acids were dried over anhydrous sodium sulfate. One liter of redistilled petrolum ether was then added and the solution allowed to stand overnight at -6° C. The asturated acids and the sodium sulfate were then filtered off and 1 liter of petroleum ether added to the filtrate. The filtrate was then ready for bremination.

Preparation of Crystalline Tetrabromostearic Acid. A five liter flask containing the solution of the fatty acids in petroleum ether was clamped firmly in an ice-salt bath, care being taken that the flask was far enough from the bottom of the bath to insure proper cooling. Bromine was added from a separatory funnel at such a rate that the temperature of the reaction mixture at no time exceeded 100 C. and stirred continuously with a mechanical stirrer. Saturation of the fatty saids was indicated by the persistence of the bromine color. The flask was then tightly stoppered and allowed to stand overnight at -50 C. The orude grystalline tetrabromostearic acid was then filtered off using a Buchner furmel, washed with redistilled petroleum ether and 1.5 liters of other then added and the mixture heated on a steam bath to effect solution. Twenty g of Morit were then added and heating continued for a few minutes. The hot solution was then filtered through a warm Buchner funnel and the filtrate allowed to again stand overnight at -5° C. The crystalline produet was then filtered and washed with redistilled petroleum ether and the crystals dried at room temperature. The melting point was then determined (114-118°) and if low, the product was recrystallised as before from petroleum sther before proceeding.

Preparation of Nethyl Lincleate. Two hundred g of tetrabromostesric acid and 200 g of granulated sine were placed in a dry, ground-nack, round bottom flask. A reflux condenser was attached and 200 ml of methyl alcohol added. The soid wad dissolved by carefully warming on a steam bath, conling by a streem of tap water to control the initial reaction if necessary. The mixture was then allowed to reflux for two hours, after which the reaction mixture was cooled and poured into a separatory funnel containing 500 ml of distilled water. A small amount of hydrochloric acid was added to decompose any gine soaps and then shaken vigorously. The mixture was then allowed to stand until the ester had separated completely. The aqueous phase was then drawn off and extracted twice with 200 ml of netroleum ether. The ester and the ether solution were added and washed twice with 500 ml of 2 percent sodium carbonate solution and twine with 100 ml of cold distilled water. The ether solution was then dried over anhydrous sodium sulfats, filtered, and the solvent removed by distillation under reduced pressure. The ester was distilled in an all-glass vacuum distillation apparatus and the fraction collected that boiled at 135-1400. 0.1 mm measure.

<u>Proparation of a^{10,12} Linoleic Acid</u>. The conjugated linoleic acid was prepared by the method of von Wikusch (41).

Sixty g of solid potassium hydroxide were dissolved in 840 ml of 05 percent ethyl alcohol and 80 ml of distilled water. Two hundred g of dehydrated easter oil were then added to this solution and the mixture allowed to reflux for l hour on a steem bath. The reaction mixture was then cooled in a stream of tap water and 880 ml of distilled water added. The mixture was then acidified to litms with concentrated hydrochloric acid, placed in a large separatory funnel and chaken to insure decomposition of the coaps. The aqueous phase was drawn off and extracted with Skellysolve B and the Skellysolve phase added to the acids. The solution was then dried over anhydrous sodium sulfate, filtered, and the solvent removed by distillation under reduced pressure.

The saids thus obtained were then isomerized by adding them slowly to a solution of 400 g of potassium hydroxide in 1100 ml of ethylene glycol at 180° C. in an oil bath and contiming the heating for 30 minutes. The heating was carried out in a large Erlemeyer flack and stirred slowly to prevent local heating. Following the heating, the hot reaction mixture was poured into a beaker containing 1000 g of chipped ice, acidified to litems with concentrated hydrochloric acid, transferred to a large experatory funnel and shaken vigorously. The acids were then extracted with Skellysolve F, washed with distilled water, dried over ambydrous codium sulfate, filtered, and the solvent removed by distillation under roduced pressure. The ethylene glycol phase was extracted with Skellysolve F and the Skellysolve

fraction treated as before. The two fractions were then combined and refrigerated at 3° C. for 48 hours to allow crystallisation of the solid scids. The solid scids thus formed were filtered on a Buchner funnel and recrystallized twice from each of the following solvents: Skellysolve F, 95 percent ethyl alcohol and diethyl ether at -17° C. This yielded δ^{10} , 12 octadesediencie acid of shout 90 percent purity.

Preparation of Dissonsthame. A mixture of 40 ml of ether and 12 ml of 40 percent aqueous potassium hydroxide were placed in a 250 ml Briemmeyer flask and cooled to 3-5° C. in a nicesalt bath. This mixture was continuously stirred mechanically as 5.45 g of N-nitrosomethylures were added in small portions as rapidly as the crystals dissolved. The yellow ethereal solution of discomethane was separated from the aqueous fraction by means of a separatory furmel and the ethereal solution dried by allowing it to stand overnight over snhydrous potassium hydroxide polices in the cold (4-6° C).

<u>Frenaration of the Methyl Ester of 2^{10,12} Octadecadiomolo Acid</u>. Approximately one-half of the solution of diazomethane in other was added from a burrette to a solution of 8,7 g of the conjugated lincleic acid in diethyl ether. Hitrogen gas was evolved and the methyl ester was obtained by removing the ether by distillation under reduced pressure.

Determination of Constants

Peroxide Value. The determination of the peroxide value was carried out by the method of Wheeler (42). Approximately 100 mg of the sample was weighed into a 125 ml Erlemmeyer flask and dissolved in 5 ml of a mixture of glacial acetic acid and chloroform (2:1 by volume). One all of a saturated solution of potassium iodide was then added and the flask rotated slowly for one mimute. Fire all of water were then added and the liberated iodine titrated with 0.01 H sodium thiceulfate solution, using starch solution as the indicator. The peroxide value in millimolee of peroxide caygen per kilogram of sample was then calculated by use of the formula $P = \frac{1 \times N}{2} \times \frac{1}{2} \times 1000$ where V is the volume of sodium thiceulfate expressed in ml, and of normality N, used in the titration of weight W, in grams, of the sample.

Spactral Daterminations

Spectrophotometric Analysis. The ultraviolet absorption of the samples was determined by means of a Beckmann D. U. Quartz Spectrophotometer. Samples of approximately 100 mg were weighed into a 100 ml volumetric flask, discolved in absolute sthyl alcohol and the flask filled to the mark. The absorption at 8380 A was then determined, diluting if necessary to keep the absorption in the region of maximum instrumental accuracy. The scale readings were then calculated in terms of the specific absorption, dy

by the method of Kraybill (28), $(\log I_0/I)/cI = \alpha$ where C is the concentration of the sample expressed in grams per liter and 1 is the length of the cell, in this case 1 cm.

Infrared Absorption Spectra. The infrared absorption spectra of the samples was determined by means of a Perkin-Klaer Infrared Spectrometer, model 185, using rock selt optics and equipped with an automatic wave-length drive and a Leeds and Northrup Specdomax recorder. The cell thicknesses used were 0.08 mm for the $A^{10,10}$ methyl limoleate and 0.08 mm in the case of the $A^{9,10}$ methyl limoleate. These particular cell thicknesses were chosen because they apparently gave the best absorption on the uncidized samples.

Experimental Procedure. The exidation was carried out by bubbling dried air through the compound contained in a small test tube suspended in a thermostatically controlled water bath maintained at 50° G. At various recorded time intervals samples were withdrawn for a determination of the infrared absorption spectra and the corresponding perceide values in the manner described. Two determinations were made for each eample, the first with the cell in place in the instrument and the second with no cell to eliminate errors in the absorption due to atmospheric carbon dioxide and water vapor.

For comparative purposes the absorption spectra of systohaxens and tetralin were determined both before and after oxidation in the described manner. In addition, wixtures of the two methyl linolestes were made up for a determination of the infrared absorption and the ultraviolet absorption at 2580 A. The region of 2580 A is known to be the region in which comjugated double bonds absorb strongly. The percent conjugation
ould thus be determined by multiplying the epecific absorption
by the factor 1,095. This factor was determined on the basis of
a theoretical absorption of 115 for the pure acid and the relationship between the molecular weights of the said and ester.
These were then used to determine the degree of conjugation from
the corresponding infrared absorption band for the various semples withdrawn during the oxidation.

RESULTS

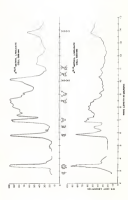
The absorption spectra of the two methyl limoleates preceeding exidation are shown in Fig. 1. These are seen to have the absorption bands which would be expected of compounds having their respective structures. The essential difference in the two spectra is in the absorption band at $10.15 \, \mu$ where the $4 \, ^{10} \, ^{18}$ methyl limoleate absorbe strongly due to the conjugation. This same band is also present in the unconjugated compound but to a much lesser extent, showing that a small smount of conjugated material is present. The other apparent differences in the absorption are due to the differences in the thickness of the cells used.

The preliminary experiments with cyclohexene and tetralin and also with the conjugated and unconjugated esters indicated a gradual change in the amount of absorption in the region of $8.9\,\mu$, the region in which absorption is generally attributed to bonded OH groups, in all cases except that of the conjugated ester. The absorption spectra of the esters of the conjugated and unconjugated acids gave the same result, as shown in Fig. 2, the lower of the curves represents the unoxidized material.

As the oxidation time increased, there was also un increase in the amount of absorption at 10.15 μ in the case of the uneonjugated methyl ester (Fig. 3). The band remained constant for the conjugated ester. As a result of this increase, mixtures of the two esters were made up and the percent of conjugated material present determined by the ultraviolet absorption at 2800 A. The infrared absorption spectra of these semples showed an increase in the degree of absorption at 10.18 μ with increasing conjugation as shown in Fig. 4.

Still another change was noted in the oxidation of the unenjugated ester which was not found in the conjugated material. This was a gradual decrease in the amount of absorption at $18.68~\mu$ with increasing exidation time as shown in Fig. 5. This change is not directly commested with the increase in the amount of conjugation as the mixtures previously described showed no such change, indicating therefore, that this is a result of the oxidation.

The results of the determinations of percentic values made during the oxidation are shown in Fig. 7. The percentic value of the unconjugated methyl limoleate is seen to rice rather rapidly with oxidation time while that for the unconjugated ester rises only very slowly.



The absorption spectra of the unoxidized esters.



The increase in absorption at 2.9 \$\mu\$ region during autoxidation of the unconjugated ester. Fig. 2.



The increase in absorption at 10.15 autoxidation of the unconjugated es Fig. 3.



The increase in absorption at 10.15 increasing conjugation of prepared n



Fig. 5. The decrease in absorption at 13.82 μ during autoxidation of the unconjugated ester.

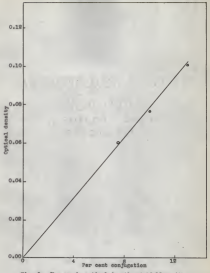


Fig. 6. Changes in optical density at 10.15 μ with increasing conjugation of prepared mixtures.

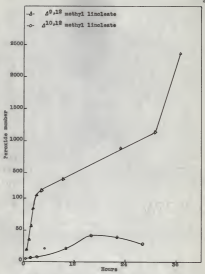


Fig. 7. Peroxide numbers during autoxidation.

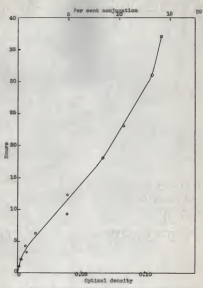


Fig. 8. Changes in optical density and per cent conjugation at 10.15 μ with autoxidation time.

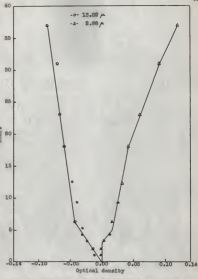


Fig. 9. Changes in optical density at 2.98 μ and 13.82 μ with autoxidation time.

DISCUSSION

There are four functionally distinct groups containing the OH linkage which presumably could be formed as a result of the autoxidation process and thus account for the increase in the absorption at 2.9 µ. These groups are the hydroperoxide group, the CH of water, the simple alcoholic CH and the carboxylic CH. However, since the reaction conditions were very mild, the likelihood of formation of a carboxylic OH is very slight, at least in the early stages of the reaction. The presence of peroxide oxygen leads to the theory of the formation of a hydroperoxide group as one of the primary steps in the autoxidation process rather than the alcoholic OH since this group would not contain peroxidic oxygen. Alcohols are possible products of secondary reactions as shown by Parmer (15) so that this possibility cannot be entirely discredited. Later reaction stages may result in the release of water by a condensation of the hydroxy compounds to form polymers so that all of these groups may be accounted for by the autoxidation mechanisms advanced. On the basis of this, it therefore seems most likely that the peroxides formed in the initial stages of the autoxidation reaction are essentially all hydroperoxides which is in agreement with the theory advanced by Farmer (16). There is no evidence to indicate that the initial step is the addition of oxygen to the double bond although, as Farmer pointed out (16), the actual amount of addition necessary to start the reaction chains would

be insignificant and probably could not be detected.

On the other hand, there is no increase in the amount of absorption at $8.9\,\mu$ over the 48.5 hour exidation period of the conjugated methyl limoleate. The percentle value does increase slightly with increasing exidation time, but comparison with the unconjugated ester indicates that the amount of percent coryen present is probably so small that it could not be detected in the absorption spectra. The mechanisms involved in the autoridation of these two esters is evidently of a different nature.

The increase in absorption at 10.15 µ in the spectra of the unconjugated ester indicates an increase in conjugation occurring as a result of the autoxidation reaction. This band also did not change in the autoxidation of the confugated methyl linolegte. From the prepared mixtures it was calculated that the actual amount of conjugation occurring in the unconjugated ester during autoxidation increased to a value of about 17 persent after background corrections were applied by conversion of the absorption to optical densities. This increase in conjugation is in agreement with the hydroperoxide theory of autoxidation in which diolefins of the methylene-interrupted type form a free radical capable of resonance between two confugated isomeric structures leading to the formation of isomeric hydroperoxides. This free radical is formed in both the theory which involves an initial addition of oxygen at the double bond and that of initial detachment of hydrogen in the Q-position with respect to the double bond.

The third change in the absorption spectra was a decrease in the absorption at 15.82 µ and this again was found in the unconjugated ester but not in the conjugated compound. It was at first believed that this was in some way connected with the absorption at 10.15 m since both of these involve carbon to carbon linkages. The prepared mixtures in which the amount of conjugated material was varied exhibited a constant absorption in this region indicating that the decrease in absorption is a result of the autoxidation process. This absorption band generally is attributed to straight chain carbon compounds in which the chain length is greater than four. Since the number of carbon to carbon linkages or the type of these linkages is not altered merely by a conversion of an unconjugated to a conjugated diene, this decrease is due apparently to chain scission reactions. It has been stated (15) that the formation of hydroperoxides may lead to scission between an ethylenic bond and an q-carbon atom, and also that such seission might occur during the original autoxidation by momentary formation and immediate decomposition of cyclic peroxides. Due to the evidence in support of the hydroperoxide theory and its incompatability with the theory of cyclic peroxide formation in the early reaction stages, it seems more likely that the degreese noted is due to ssission reactions of the former type.

It seems, therefore, in view of the results obtained, that the reaction of the unconjugated methyl linoleate with oxygen results in the formation of a free radical which may then become conjugated and lead to the formation of isomeric hydroperoxides. A second free redical may be formed during this process, thus accounting for the chain reaction. The hydroperoxide formed may undergo secondary reactions leading to a wide variety of end products. The mechanism by which the conjugated and unconjugated esters undergo autoxidation apparently is different as evidenced by the changes in the absorption spectrum of the unconjugated ester and the constancy of the spectrum of the conjugated ester during the autoxidation periods of this investigation. It seems possible that the oxidation of the conjugated ester may result in a polymerisation of this empound by a more direct mechanism than is the case with the unconjugated ester.

STIMARY

The autoxidation of the conjugated and unconjugated esters of limoleic acid was studied by means of infrared absorption spectra and a correlation with the theories of autoxidation attempted.

The unconjugated material is shown to undergo autoxidation in a different manner than that of the conjugated material as evidenced by a constancy of absorption during the autoxidation of the conjugated ester and enhances at 2.9, 10,15, and 13,58 \(\times \) in the case of the unconjugated ester. These changes may be explained on the basis of the hydroperoxide theory of autoxidation very matinifectority.

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LITERATURE CITED

- Atherton, D. and T. P. Hilditch.
 The union of gaseous oxygen with methyl cleate at 20° and 120°. Chem. Soc. Jour., 1945; 105-108.
- (2) Bach, A. Function of peroxides in phenomena of slow exidation. Compt. rend., 124: 951-954. 1897.
- (5) Barnes, R. B. Infrared spectra and organic chemistry. Rev. Sci. Instr., 7: 265-271. 1936.
- (4) Barnes, R. B., Urner Liddel and V. Z. Williams. Infrared spectroscopy. Industrial applications. Indus. and Engin. Chem., Anal. Ed., 15: 659-709. 1045.
- (5) Bartholome, E. and E. Teller. Calculation of characteristic frequencies of organic chain-molecules by means of models. Z. Physik. Chem. B., 19: 366. 1932.
- (6) Bolland, J. L. and Geoffrey Gee. Kinetic studies in the chemistry of rubber and related materials. II. The kinetics of oxidation of unconjugated clefins. Faraday Soc. Trans., 42: 236-245. 1946.
- (7) Bolland, J. L. and Geoffrey Gee. Kinetic studies in the chemistry of rubber and related materials. III. Thermochemistry and mechanism of olefin oxidation. Faraday Soc. Trans., 42: 244. 1946.
- (8) Bolland, J. L. and E. P. Koeh. The cause of autoridation reactions in polyisoprenes and allied compounds. IX. The primary thermal oxidation product of ethyl linelests. Chem. Soc. Jour., 1945; 445-447.
- (9) Grawford, B. L., Jr. and S. R. Brinkley, Jr. Force constants in some organic molecules. Chem. Phys. Jour., 9: 69-75. 1941.
- (10) Griegee, R. Ognic acid esters as intermediate products in oxidation. Annalen, 522: 75-95. 1936.

- (11) Griegee, R., H. Pils and E. Flygare.
 Olefin peroxides. Ber., 72: 1799-1804. 1959.
- (12) Ellis, G. W. The chemistry of drying oils. Soc. Chem. Ind. Jour., 44: 401-406T. 465-468T. 469-472T. 486T. 1925.
- (18) Engler, C.
 The rendering active of oxygen. Ber., 35: 1090-1111.
 1900.
- (14) Fahrion, Wilhelm. Drying process of linseed oil. Chem. 2tg., 28: 1196-1200. 1904.
- (15) Farmer, E. H. Methylenic reactivity in clefin and polyclefinic systems. Faraday Soc. Trans., 38: 340-548. 1942.
- (16) Farmor, E. H.
 Peroxidation in relation to elefin structure. Faraday
 Soc. Trans., 42: 228-236, 1946.
- (17) Farmer, E. H., O. F. Bloomfield, A. Sundralingum and D. A. Subbon. The course and mechanism of autoxidation reactions in olefinio and polyclefinio subtances, including rubber. Faraday Soc. Trans., 38: 348-356. 1942.
- (18) Farmer, E. R., H. P. Eoch and D. A. Sutton. Course of autoxidation resetions in polysisoprenes and allied compounds. VII. Rearrengement of double bonds during autoxidation. Chem. Soc. Jour., 1945; 54: 547.
- (19) Farmer, E. H. and D. A. Sutton. Course of autoxidation reactions in polyisoprenes and allied compounds. IV. The isolation and constitution of photochemically formed methyl cleate peroxide. Chem. Soc. Jour., 1945: 129-122.
- (20) Fokin, S., Catalytic exidation and reduction reactions of organic compounds. Z. Angew. Chem., 22: 1451-1459, 1492-1502. 1909.
- (21) Gamble, D. L. and C. E. Barnett. Infrared absorption spectra of drying cils. Indus. and Engin. Chem., 32: 375-378. 1040.

- (32) Gunstone, F. D. and T. F. Hilditch.

 The union of gasecus oxygen with methyl cleate,
 linoleste, and linolenate. Chem. Soc. Jour., 1945:
 836-841.
- (25) Hersberg, Gerhard.

 Molecular spectra and molecular structure. I. Diatomic molecules. Now York: Prentice-Hall, p. 69-80.
- (24) Herzberg, Gerhard. Infrared and Raman spectra. Hew York: D. Van Hostrand and Co. p. 192-201. 1945.
- (25) Hock, E. and W. Susemihl.
 Autoxidation of hydrocarbons. I. A tetrahydronaphthalene peroxide obtained by autoxidation. Ber., 668: 61-68. 1955.
- (26) Jones, R. Horman. Infrared spectroscopy in organic chemistry. Canad. Chem. and Process Ind., 30: 85. 1946.
- (27) Kettering, C. P., L. W. Shutts and D. H. Andrews. A representation of the dynamic properties of molecules by mechanical models. Phys. Rev., 36: 531, 1930.
- (28) Erapbill, H. R., J. H. Witchell, Jr. and P. P. Sacheile. Ultraviolet absorption spectra of lineed oil. Determination of bodied-in-wacuo and blown linsed oil in mixtures with raw linsed oil. Indus. and Engin. Chem., Analyt. Ed., 13: 705-705. 193.
- (29) Markley, K. S. Fatty acids, their chemistry and physical properties. New York: Interscience, p. 137, 324, 431. 1947.
- (50) McOutcheon, J. W., M. F. Crawford and H. L. Welsh. Isomeric structure of the G. unsaturated acids from their Raman and infrared spectra. Oil and Soap, 18: 9-11. 1341.
- (31) Mecks, R.

 Valence and deformation vibrations of simple molecules. I. General theory. Z. Physik. Chem., B., 16: 409-420. 1932.
- (32) Meske, R.

 Valence and deformation vibrations of simple molecules. II. Triatomic molecules. Z. Physik. Chem.,
 B, 16: 421-437. 1932.

- (33) Meake, R.
 - Valence and deformation vibrations of polyatomic molecules. III. Methane, acetylene, ethylene, and halogen derivatives. Z. Physik. Chem., B, 17: 1-20. 1982.
- (34) Pauling, Linus.

 The nature of the chemical bond. New York: Cornell University Press, p. 53, 131. 1945.
- (35) Rieche, A. The oxidation of organic compounds with air oxygen. Angew. Chem., 50: 520-524. 1937.
- (36) Rollett, A. Lincleic acid. Z. fur Physiol. Chem., 62; 410-421. 1909.
- (37) Staudinger, H. Autoxidation of organic compounds. III. Autoxidation of asym-diphonylethylene. Ber., 58: 1075-1079. 1925.
- (38) Stephens, H. H. Autoxidation. I. Cyclohexene peroxide. Amer. Chem. Sec. Jour., 50: 568-571. 1928.
- (59) Swift, C. E., F. G. Dollear and R. T. O'Connor. The oxidation of methyl oleate. I. The preparation, properties, and reactions of methyl peroxido cleate. Oil and Soap, 23: 356-359. 1946.
- (40) Ssent-Gyorgyi, A. v. Studies on biological exidation. I. The absorption of exygen by the system limelenic acid-SH groups. Z. Biochem., 146: 245-255. 1924. II. The mechanism and significance of -SH catalysis. Z. Biochem., 146: 254-255. 1924.
- (41) von Mikusch, F. D. Solid 10,12-octadecadiencic acid-1. A new conjugated lincleic acid melting at 57°. Amer. Chem. Soc. Jour., 54: 1580. 1942.
- (42) Wheeler, D. H. Peroxide formation as a measure of autoxidative deterioration. Oil and Scap, 9: 89-97. 1982.