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Co-oxidation kinetic model for the thermal oxidation of polyethylene-unsaturated substrate systems

Emmanuel Richaud a,*, Bruno Fayolle a, Jacques Verdu a, Jozef Rychlý b

a Arts et Metiers ParisTech, CNRS, PIMM UMR 8006, 151 bd de l’Hôpital, 75013 Paris, France
b Polymer Institute, Slovak Academy of Sciences, Dúbravská cesta 11, 842 36 Bratislava, Slovak Republic

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Methyl esters of unsaturated fatty acids
Co-oxidation
Kinetic modeling
Chemiluminescence

Abstract
The thermal oxidation of polyethylene (PE) impregnated by the methyl esters of unsaturated fatty acids (UFEs) was studied using chemiluminescence, and infra-red spectrophotometry. It was shown that the presence of UFEs accelerates the PE aging process. This can be interpreted as a co-oxidation phenomenon. In this study, the previously established models for PE and UFEs self-oxidation have been coupled in order to develop a co-oxidation model. Using the existing rate constants for the PE and UFEs self-oxidations, this model can simulate the complex shape of the kinetic curves of PE-UFE co-oxidation.

1. Introduction

In a first approach, the oxidation mechanism at relatively moderate temperatures (\(< 150^\circ C\)) of a hydrocarbon substrate with a single reactive site, when there is an oxygen excess (that is when there is no contribution of alkyl radicals \(P\) to the termination reactions) can be depicted by the following scheme [1,2]:

\[
\begin{align*}
(I) & \quad ROOH + ROOH \rightarrow R^+ + ROO^- + \text{carbonyls} + \text{scissions} & k_{1b} \\
(II) & \quad R^+ + O_2 \rightarrow ROO^- & k_2 \\
(III) & \quad ROO^- + RH \rightarrow ROOH + R^- & k_3 \\
(VI) & \quad ROO^- + ROO^- \rightarrow \text{inactive products} & k_6
\end{align*}
\]

Over the years, this model as well as several completed versions has been developed with a view to describing the oxidation of substrates under specific reaction conditions, namely, when it can be assumed that there is only one kind of reactive site which corresponds to the weakest C–H bond and the highest propagation rate constant \(k_3\) [3]. Some examples are: tertiary carbons in polypropylene [4], secondary carbons in polyethylene [5] and allylic ones in butadiene rubber [6].

It is well documented that oxidation can be favored by metallic impurities playing a catalytic (redox) role [7] or by species such as carboxylic acids capable of establishing strong hydrogen bonds with the polymeric ROOH groups [8]. These cases of assisted oxidation differ from the co-oxidation phenomena described in this current study. Here, the simultaneous in chain oxidation of two different kinds of reactive sites corresponding to two categories of C–H groups will be studied.

In such a case of co-oxidation, Decker et al. [9] showed that the oxidizability of ethylene propylene copolymer varies with the ethylene molar fraction \(e\) according to a pseudo-hyperbolic curve which can be approximated by the following function:

\[\frac{k_3}{k_6} = 10^{-4} \times \frac{2 - 1.63 \times e}{1 + 3.13 \times e}\]

Here \(k_3\) and \(k_6\) can be defined as the rate constants characteristic of a virtual homopolymer which would exhibit the same kinetic behavior as the copolymer under study. In fact, an approach in which the kinetic behavior of the copolymer could be predicted from the characteristics of the corresponding homopolymers would be far more satisfactory. In the case of propagation, the presence of two reactive sites needs to take into account four elementary reactions:

* Corresponding author. Tel.: +33 1 44246316.
E-mail address: emmanuel.richaud@ensam.eu (E. Richaud).
With this in mind, the coupled degradation of a PE-UFE series was studied using chemiluminescence at 150 °C. Another solid state (80 °C) series was also studied in an effort to separate the PE and UFEs oxidation by-products.

2. Experimental

2.1. Materials

Polyethylene was supplied as an antioxidant free powder. This was verified by observing the absence of an induction period at 200 °C. DSC analysis revealed a melting temperature of 132 °C with a melting enthalpy close to 150 J g⁻¹. This corresponds to a crystallinity ratio of about 50%.

According to Demirbaş et al. [15], vegetable oils are mainly composed of the methyl esters of fatty acids with 16 or 18 carbons in their linear hydrocarbon chains. Two important compounds are methyl oleate and methyl linolenate which have different oxidative stabilities. Methyl oleate (ref 554470-478) and methyl linolenate (ref 62210) were supplied by Sigma Aldrich.

The impregnation of the methyl esters into the PE at various temperatures has been previously described [16]. To summarize the initial state of the materials used were: ① pure PE, ② PE impregnated with 5% weight methyl oleate, ③ PE impregnated with 10% weight methyl oleate and ④ PE impregnated with 5% methyl linolenate.

2.2. Exposure conditions

Impregnated films were submitted to one of the two following conditions: ① thermal aging in ovens at 80 °C monitored by FTIR, ② thermal aging at 150 °C under 0.1 MPa O₂ monitored in situ by CL.

2.3. Characterization

2.3.1. Chemiluminescence

Chemiluminescence experiments were performed using a Lumipol 3 apparatus designed in-house at the Polymer Institute in the Slovak Academy of Sciences [7]. Films impregnated by methyl oleate and linolenate were placed into aluminum pans and heated under nitrogen to 150 °C. They were then maintained at this temperature under oxygen at 0.1 MPa pressure.

2.3.2. Extraction of the methyl esters of the unsaturated fatty acids

After thermal oxidation of the PE impregnated by UFE, both the UFE and its oxidation by-products were extracted using CH₂Cl₂ overnight at room temperature. An example of the observed differences in the FTIR spectra between an impregnated PE sample before and after extraction are shown in the spectra in Fig. 1 at t = 0. These show the total UFE extraction. Those in Fig. 3 represent the samples after several exposure durations.

2.3.3. FTIR

FTIR spectra of the PE films were recorded in transmittance mode using a Spectrum 100 spectrophotometer (Perkin Elmer), with 4 scans at 4 cm⁻¹ resolution. Both the virgin and oxidized methyl esters of the unsaturated fatty acids display an absorption peak at ca 1740 cm⁻¹. After removing the UFEs and its soluble by-products from PE, only a peak at 1720 cm⁻¹ remains. This is attributed to carbonyl compounds resulting from the PE oxidation. The carbonyl absorbance before and after extraction were converted into concentrations using the molar absorptivity of 300 l mol⁻¹ cm⁻¹ [17].
3. Results

3.1. Chemiluminescence study

Impregnated films were submitted to a thermo-oxidative aging procedure monitored in situ in a CL apparatus. TGA results confirmed that only a minor part of the absorbed UFE was lost by evaporation in the timescale under consideration.

Typical CL results for oxidation at 150 °C under 0.1 MPa O2 are shown in Fig. 2. A close examination of the results leads to the following observations:

① For pure PE, the curve has the classical sigmoidal shape. In addition, a small shoulder at time ~10000 s can be seen. According to Broska [18], this could be attributed to the existence of structural irregularities in PE (such as vinyl chain ends). This interpretation is, as will be confirmed later, in good agreement with the proposed kinetic model for co-oxidation involving the role of double bonds.

② Next, for the PE + methyl oleate, the CL kinetic curve shifts towards shorter times. This clearly suggests that the PE matrix is oxidized faster in the presence than in the absence of an UFE.

③ The trends observed for the PE + methyl oleate are strongly exaggerated for the PE + methyl linolenate system where an intense peak was seen to develop in the early days of exposure and also where the light emission in the first 5000 s is considerably stronger than for the other samples.

3.2. FTIR study

The absorbance of the impregnated samples is shifted during the first hours of exposure from 1745 to 1735 cm⁻¹ and then towards 1720 cm⁻¹. The former shift is a result of the UFE oxidation, whereas the PE oxidation leads to the second shift. After extraction by CH₂Cl₂, it is reasonable to assume that residual carbonyl peak centered at ca 1720 cm⁻¹ is linked to the oxidation level of PE matrix.

The results presented in Figs. 4 and 5 show a comparison between oxidation of a pure PE film and of an UFE-impregnated one. The kinetics curves which give the carbonyl concentration in the PE matrix with an UFE after aging are shifted towards shorter times compared to the kinetic curve for pure PE. Despite a noticeable scatter, these results clearly confirm that unsaturated fatty esters accelerate the polyethylene matrix oxidation. This effect is increased with the UFE concentration in PE (Fig. 4). It can also be seen that methyl linolenate has a stronger influence than methyl oleate (Fig. 5).

4. Discussion

4.1. Proposal of a mechanistic scheme

According to Figs. 2, 4 and 5, a co-oxidation mechanism, similar to that proposed for ethylene propylene copolymers EPR [9] can be expected to occur. This would be comparable to the mechanism also proposed in the case of PP in which the participation of both secondary and tertiary carbons was demonstrated as a result of the analysis of the hydroperoxides and alcohols [19,20].

The inherent nature of such a combined mechanism leads to a very complex mechanistic scheme with a greater number of kinetic parameters needing to be determined [12]. It is now pertinent to elaborate a model describing the most striking experimental features with the minimum number of adjustable kinetic parameters, that is starting from the PE oxidation model [4] and the UFE one [14] together with some cross reactions and retaining the previously determined rate constant values. The following mechanistic scheme is a valid first approach:

\[
\begin{align*}
(1u2) & \quad \text{PEOOH} \rightarrow 2\text{PE}^+ + \gamma_{\text{CO}}\text{PE} = \text{O} + \gamma_5 s \\
(2b2) & \quad \text{PEOOH} + \text{PEOOH} \rightarrow \text{PE}^+ + \text{PEOO}^+ + \gamma_{\text{CO}}\text{PE} = \text{O} + \gamma_5 s \\
(1b11) & \quad \text{UFEOOH} + \text{UFEOOH} \rightarrow \text{UFEO}^+ + \text{UFEOO}^+ + \text{UFE} = \text{O} \\
(1b12) & \quad \text{UFEOOH} + \text{PEOOH} \rightarrow \text{PE}^+ + \text{UFEOO}^+ + \text{PE} = \text{O}
\end{align*}
\]
The cross-initiation reaction (Ib12) between PEOOH and UFEOOH is theoretically the balance between the two pathways shown below:

\[
\begin{align*}
(\text{Ib12}) \quad & \text{UFEOOH} + \text{PEOOH} / \text{PE} / \text{C14} + \text{UFEOO} / \text{C14} + \text{PE} / \text{C14} \\
& \text{Ok b12}
\end{align*}
\]

On the basis of the thermochemical considerations shown in Table 1, the \(\Delta G_R\) is lower for pathway 2. It can be assumed, therefore, that the reaction between PEOOH and UFEOOH can be written as a single balance equation:

\[
\begin{align*}
(\text{Ib12}) \quad & \text{UFEOOH} + \text{PEOOH} \rightarrow \text{PE}^* + \text{UFEOO}^* + \text{PE} = \text{O} \\
& k_{b12}
\end{align*}
\]

4.2. Estimation of kinetic parameters for the co-oxidation model

Oxygen solubility in impregnated polyethylene is supposed unmodified by the presence of a few % by weight of UFE. Its value will therefore be taken equal to the value reported by Van Krevelen [21] in his monograph: \(sO_2 = 1.8 \times 10^{-4} \text{ mol L}^{-1} \text{ Pa}^{-1}\).

All the kinetic parameters for polyethylene (\(k_{62}, k_{622}, k_{32}, k_{322}\), and associated activation energies have been determined previously [5] hence will be used without change. They are shown in Table 2. For PE, \(k_{60}, k_{61}, k_{62}\) and \(k_{63}\) are combined to give an apparent rate constant \(k_{622}\) equal to:
Calculation of free enthalpy changes for cross bimolecular hydroperoxide decomposition.

<table>
<thead>
<tr>
<th>Macroradical</th>
<th>Model compound</th>
<th>( \Delta H (\text{kJ mol}^{-1}) )</th>
<th>( \Delta S (\text{J mol}^{-1} K^{-1}) )</th>
<th>( \Delta H_4(298 \text{K})(\text{kJ mol}^{-1}) )</th>
<th>( \Delta H_{353}(\text{kJ mol}^{-1}) )</th>
</tr>
</thead>
<tbody>
<tr>
<td>(-CH_2=CH-CH_2-)</td>
<td>iso-C_2H_5-O^-</td>
<td>-75.3</td>
<td>355.3</td>
<td>-225.6</td>
<td>-202.8</td>
</tr>
<tr>
<td>(-CH=CH=CH-)</td>
<td>H_2C=CH-CH_2-OO^-</td>
<td>96.3</td>
<td>350.3</td>
<td>-51.6</td>
<td>-27.4</td>
</tr>
<tr>
<td>(-CH=CH-CH=CH-)</td>
<td>iso-C_3H_7-OO^-</td>
<td>-46</td>
<td>345.3</td>
<td>-192.1</td>
<td>-168</td>
</tr>
<tr>
<td>(-CH=CH=CH=C)</td>
<td>(\equiv)CH-O^-</td>
<td>104.6</td>
<td>364.1</td>
<td>-49.4</td>
<td>-24</td>
</tr>
</tbody>
</table>

Fig. 5. Kinetic curves at 80 °C for carbonyl buildup for pure PE ( ), total carbonyl concentration in PE impregnated by methyl linolenate ( ), residual carbonyl in PE after removing by CHCl_2 of methyl linolenate and its oxidation by-products removing ( ) together with the corresponding curves generated by the kinetic model with parameters given in Tables 2 and 3.

\[ k_{622} = \frac{k_{60}}{1 + \frac{k_{60}}{k_{621} + k_{622}}} \left( \frac{1 + \frac{k_{63}}{k_{61} + k_{62}}} {2 \left( k_{61} + k_{62} \right)} \right) \]

The kinetic parameters for methyl oleate and linolenate were tentatively determined and reported in a recent work based on a study of the oxidation of pure compounds monitored by CL [14]. It is now necessary to consider the cross reaction rate constants. According to Korcek’s law [3], the rate constant for \( \text{ROO}^+ + \text{PH} \rightarrow \text{ROOH} + \text{P} \) depends only on the value of the bond dissociation energy. Hence it is possible to suppose:

\[ k_{312} = k_{322} \]

\[ k_{321} = k_{311} \]

In addition, for the initiation and termination rate constants, it will be assumed:

\[ k_{b12}^2 = k_{b11} \times k_{b22} \]

\[ k_{312} = k_{311} \times k_{322} \]

4.3. A comparison between model output data and chemiluminescence results

There is a wide range of species possibly present in PE in low concentrations that may be eventually responsible for the initial oxidation steps. In addition to others these are peroxides, and unreacted polymerization catalysts. It is warranted to consider that initial PE contains only hydroperoxides in an initial concentration \([\text{POOH}]_0\), the decomposition of \([\text{POOH}]_0\) yielding to the same quantity of radicals than the overall unstable species initially present. Since POOH groups accumulate during oxidation while other eventually unstable species are consumed, the POOH decomposition rapidly becomes the largely predominant initiation process and this in fact justifies the mechanism scheme chosen [22]. It was assumed that \([\text{PEOOH}]_0\) is equal to \(10^{-6} \text{ mol l}^{-1}\) which is consistent with our other papers dealing with PE oxidation modeling. In addition, \([\text{UFEOOH}]_0\) was fixed at a value 100 times higher because of the possible unsaturated fatty ester pre-oxidation occurring during the impregnation phase. It has been shown previously that such values should be employed for simulating the CL curves of pure UFEs oxidation [14].

The system of differential equations established from the kinetic scheme is solved using the following initial conditions:

At \( t = 0 \):

\[ [\text{UFE}^+]'_0 = [\text{UFEOOH}^+]_0 = [\text{PE}^+]_0 = [\text{POOOH}]_0 = 0 \]

\[ [\text{PEOOH}]_0 = 10^{-4} \text{ mol l}^{-1} \]

For the sake of simplicity, the densities of the PE amorphous phase and the fatty esters are assumed to be the same so that:

\[ [\text{PE}]_0 = \frac{\rho_{\text{PE}} M_{\text{CH}_2}} {1 - x_C} \times (1 - w_{\text{UFE}}^e) \]

\[ [\text{UFE}]_0 = \frac{1} {1 - x_C} \cdot \frac{\rho_{\text{PE}} M_{\text{UFE}}} {1 - w_{\text{UFE}}^e} \]

here:

- \( \rho_{\text{PE}} \) is the density of PE amorphous phase.
- \( M_{\text{CH}_2} \) is the molar mass of the methylene group since the PE repetitive structural unit holds an abstractable hydrogen.

Table 2

<table>
<thead>
<tr>
<th>Code</th>
<th>Rate constant</th>
<th>Pre-exponential factor</th>
<th>Activation energies</th>
</tr>
</thead>
<tbody>
<tr>
<td>1u</td>
<td>( k_{b11} )</td>
<td>( 8.0 \times 10^{12} \text{ s}^{-1} )</td>
<td>140 \text{ kJ mol}^{-1}</td>
</tr>
<tr>
<td>1b</td>
<td>( k_{b10} )</td>
<td>( 2.8 \times 10^{14} \text{ mol}^{-1} \text{ l} \text{ s}^{-1} )</td>
<td>105 \text{ kJ mol}^{-1}</td>
</tr>
<tr>
<td>2</td>
<td>( k_2 )</td>
<td>( 1.0 \times 10^{11} \text{ mol}^{-1} \text{ l} \text{ s}^{-1} )</td>
<td>0 \text{ kJ mol}^{-1}</td>
</tr>
<tr>
<td>3</td>
<td>( k_3 )</td>
<td>( 1.5 \times 10^{18} \text{ mol}^{-1} \text{ l} \text{ s}^{-1} )</td>
<td>73 \text{ kJ mol}^{-1}</td>
</tr>
<tr>
<td>60</td>
<td>( k_{60} )</td>
<td>( 6.9 \times 10^{19} \text{ mol}^{-1} \text{ l} \text{ s}^{-1} )</td>
<td>80 \text{ kJ mol}^{-1}</td>
</tr>
<tr>
<td>61</td>
<td>( k_{61} )</td>
<td>( 2.0 \times 10^{6} \text{ s}^{-1} )</td>
<td>0 \text{ kJ mol}^{-1}</td>
</tr>
<tr>
<td>62</td>
<td>( k_{62} )</td>
<td>( 1.0 \times 10^{18} \text{ mol}^{-1} \text{ l} \text{ s}^{-1} )</td>
<td>5 \text{ kJ mol}^{-1}</td>
</tr>
<tr>
<td>63</td>
<td>( k_{63} )</td>
<td>( 8.0 \times 10^{12} \text{ s}^{-1} )</td>
<td>50 \text{ kJ mol}^{-1}</td>
</tr>
</tbody>
</table>
The numerical resolution of the kinetic model gives rise to the kinetic functions \([UFE^1]\), \([UFEOO^1]\), \([UFEOOH^1]\), \([PE^1]\), \([PEOO^1]\), \([PEOOH^1]\) which permits comparisons with the experimental results reported above.

Let us first consider CL emission. Several mechanisms have been proposed for CL emission. In the first, emission is from a \(P=O^*\) generated from a Russel-type termination, which is in fact questionable because this reaction is not expected to occur in PE for which the CL intensity is, however, very high [23]. In a second mechanism, emission is from \(P=O^*\) generated by \(\beta\)-scission of alkoy: \(-\text{CH}_2\text{C}^1\text{O}^*\rightarrow\text{CH}_2\text{C}^0\) [24]. Emission from \(O^*_2\) has also been proposed [25] as well as emission from a bimolecular POOH combination [26,27].

The difference between all these mechanisms consists in the fact that the excited specie responsible for CL emission comes from either the initiation or termination reaction. It can be shown that the initiation rate varies pseudo linearly with the oxygen pressure while the termination rate varies linearly with the oxygen pressure [28]. This confirms that CL comes from the bimolecular decomposition of POOH [28,29]. This conclusion is in good agreement with the majority of literature reports as well as being based on other reasoning [26,27]. Here, it can be assumed that:

\[
\text{ICL} = \phi \times k_0 [\text{POOH}]^2
\]

It should be noted that it is easy to verify that the shape of the simulated curve remains almost the same by making the assumption that CL comes from POOH + POOH \(\rightarrow\) \(\cdots\) or from POO\(^*\) + POO \(\rightarrow\) \(\cdots\) ...

In the absence of any other useful information, it will be assumed that \(\phi\) is independent of the substrate. The chemiluminescence intensity can therefore be expressed as:

\[
\text{ICL} = \phi \times \left( k_{b11} [\text{UFEOOH}]^2 + k_{b12} [\text{UFEOOH}] [\text{POOH}] + k_{b22} [\text{PEOOH}]^2 \right)
\]

Next, we will consider the carbonyl buildup. Carbonyl concentration is calculated as a post treatment of model by:

\[
\frac{d[\text{PE} = \text{O}]}{dt} = (1 - x_C) \gamma_{CO} \left( k_{b2} [\text{PEOOH}] + k_{b22} [\text{PEOOH}]^2 + k_{b22} [\text{PEOO}] [\text{PEOOH}] + k_{b22} [\text{POOO}]^2 \right)
\]

\[
\frac{d[\text{UFEOH} = \text{O}]}{dt} = (1 - x_C) k_{b11} [\text{POOH}]^2
\]

Table 3

<table>
<thead>
<tr>
<th>Temperature (°C)</th>
<th>k₂₁</th>
<th>k₂₁</th>
<th>k₂₁</th>
<th>k₂₁</th>
<th>k₂₁</th>
</tr>
</thead>
<tbody>
<tr>
<td>150</td>
<td>2 \times 10^{-5}</td>
<td>10²</td>
<td>131</td>
<td>10⁵</td>
<td>111</td>
</tr>
<tr>
<td>LNO</td>
<td>1.25 \times 10^{-2}</td>
<td>10⁷</td>
<td>525</td>
<td>4.5 \times 10⁸</td>
<td>0.25</td>
</tr>
<tr>
<td>80</td>
<td>5 \times 10^{-6}</td>
<td>10⁷</td>
<td>92.5</td>
<td>2 \times 10⁷</td>
<td>0.24</td>
</tr>
<tr>
<td>LNO</td>
<td>4 \times 10^{-5}</td>
<td>10⁹</td>
<td>92</td>
<td>2 \times 10⁸</td>
<td>0.12</td>
</tr>
</tbody>
</table>

It should also be noted that some carbonyls can also be generated as a result of termination in an UFE self-oxidation process. But in the absence of quantitative information, this process will be neglected. By neglecting some minor terms, the following can be written:

\[
\frac{d[\text{PE} = \text{O}]}{dt} + \frac{d[\text{UFEOH} = \text{O}]}{dt} = \left\{ k_{b11} [\text{UFEOOH}]^2 + k_{b22} [\text{PEOOH}]^2 + k_{b12} [\text{UFEOOH}] [\text{PEOOH}] \right\}
\]

Thus:

\[
\frac{d\text{carbonyls}}{dt} = \lambda_{ICL}
\]

It is not surprising, therefore, to discover that the rate of carbonyl buildup varies in the same way with time as the CL intensity. Using the rate constants given in Table 3, the modeling runs gave rise to the following data:

A fair simulation of CL curve shape was found as shown in Fig. 6. The model permits a better understanding of the course of the PE oxidation as shown in Fig. 7. Here, the shift of the induction period duration relative to PE is shown when PE is oxidized in presence of UFE.

Acceptable simulations for carbonyl build-up curves are seen from the results shown in Figs. 4 and 5. Their shape is in reasonable agreement with the experimental data. Taking into consideration the relative complexity for analyzing FTIR spectra in the carbonyl region and the corresponding kinetic curves, this agreement cannot be a simple coincidence. The simulations of the oxidation state of the matrix (i.e. the concentration in carbonyl hold by PE chains) are acceptable (Figs. 4 and 5). Of particular note is the fact that methyl ester accelerates the course of the PE oxidation as shown in Fig. 7.
oxidation compared to when PE is oxidized alone. The model also describes the influence on the PE oxidation kinetics of increasing the methyl ester concentration in the PE amorphous phase. These results are shown in Fig. 4.

Finally, the rate constants for methyl esters are found to be relatively very close or even identical to those determined from the corresponding self-oxidation studies [14]. One minor discrepancy is observed for one termination constant (Table 3). It could actually be envisaged that the rate constant $k_{611}$ is equal to the one determined in the case of methyl ester self-oxidation. But the rate constants for cross termination $k_{612}$ would have a different value, actually lower than the geometric average ($k_{611}k_{622})^{1/2}$. In fact, given all the possible sources of error such as the differences in the methyl ester purity between this work and [14], or differences in polarity between the pure UFE medium the PE matrix, the numerous simplifying hypotheses used for the rate constant determination [14], then there is really rather a good agreement between the constants used here for simulating the co-oxidation curves and those used for simulating the CL curves for the oxidation of pure methyl esters. Even if the hypothesis of geometric means is considered questionable, it seems that it can still offer a rough estimation of the unknown rate constants for some cross reactions.

5. Conclusions

The kinetics of the thermal oxidation of PE combined with different unsaturated fatty esters was studied using chemiluminescence in PE molten state and by FTIR in the PE solid state. It was observed that these unsaturated molecules oxidize faster than PE and hence their addition to PE accelerates its degradation. This effect is attributed to the transfer of radical sites from UFE to the PE. A kinetic model for co-oxidation has been shown to simulate the main experimental trends. The model used kinetics parameters employed for modeling PE and UFE self-oxidation. Simulations can successfully be compared with the chemiluminescence results obtained for both oxidation in molten state (150 °C) as well as in the solid state (80 °C). Finally, both reactant oxidation by-products were successfully quantified.

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