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Detailed Product Analysis during Low- and Intermediate-Temperature Oxidation of Ethylcyclohexane

B. Husson†, O. Herbinet†, P. A. Glaude†, S. S. Ahmed‡, and F. Battin-Leclerc*†

† Laboratoire Réactions et Génie des Procédés, CNRS, Université de Lorraine, 1 Rue Grandville, BP 20451, 54001 NANCY Cedex, France
‡ Saudi-Aramco Research & Development Center, Dhahran 31311, Saudi Arabia

Abstract

An experimental study of the oxidation of ethylcyclohexane has been performed in a jet-stirred reactor with online gas chromatography, under quasi-atmospheric pressure (800 Torr), at temperatures ranging from 500 to 1100 K (low- and intermediate-temperature zone including the negative temperature coefficient area), at a residence time of 2 s, and for three equivalence ratios (0.25, 1, and 2). Ethylcyclohexane displays important low-temperature reactivity with a well-marked negative temperature coefficient behavior. In addition to 47 products with a mass lower than ethylcyclohexane which have been quantified, many species with a \( \text{C}_8\text{H}_{14}\text{O} \) formula (molecular weight of 126) were detected by GC-MS and 7 of them were quantified. These molecules are cyclic ethers, ketones, and aldehydes with the same carbon skeleton as the reactant. Experiments were also carried on under the same conditions for two other \( \text{C}_8 \) hydrocarbons, \( n \)-octane and 1-octene, showing that the reactivity of ethylcyclohexane is close to that of the alkene and lower than that of the alkane. Simulations using a detailed kinetic model of the literature allow a good prediction of the global reactivity and of the main hydrocarbon products for temperatures above 800 K. The main reaction channels leading to the observed reaction products at both low (below 800 K) and intermediate temperature (above 800 K) are discussed.

Corresponding Author:

Frédérique Battin-Leclerc
Laboratoire Réactions et Génie des Procédés
Ecole Nationale Supérieure des Industries Chimiques
BP 20451, 1 rue Grandville, 54000 Nancy, France
Tel: +33 (0)3 83 17 51 25
Fax: +33 (0)3 83 17 81 20
E-mail: Frederique.Battin-Leclerc@ensic.inpl-nancy.fr
Introduction

The \( n \)-alkylated cyclohexanes are a homologous series of hydrocarbon compounds that are commonly present in crude oil and refinery products such as diesel fuels (1). The components of diesel fuels have a number of atoms of carbon which ranges approximately from 10 to 22 with an average around 14 and can be classified into the following chemical families: \( n \)-alkanes, iso-alkanes, aromatics, and cycloalkanes (2). Cycloalkanes are usually present less than 5% in diesel fuel derived from conventional crude-oil sources but are much more abundant in alternative diesel fuel derived from oil–sand deposits. These species typically have one ring with multiple alkyl side chains, e.g., alkylcyclohexanes (3).

The literature is relatively rich in detailed kinetic models for combustion of alkanes (e.g., refs 4–7) and recently alkyl–aromatics (e.g., refs 8–10). However, there are fewer studies on the other classes of molecules including alkylcyclohexanes, and the mechanisms involved in their combustion are poorly known.

Table 1 summarizes previous experimental studies of the oxidation of cyclohexane and alkylcyclohexanes. Although a fair number of studies have already been performed on the oxidation of cyclohexane and methylocyclohexane in different experimental apparatuses, very little information is available on the oxidation of alkylcyclohexanes with a larger alkyl chain. Moreover, studies performed on cyclohexane and \( n \)-propylcyclohexane oxidation in a jet-stirred reactor enclosed no information about their low-temperature reactivity.

<table>
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<th>Compounds</th>
<th>Reactor( ^d )</th>
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<th>( P ) (bar)</th>
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<td>0.68</td>
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<td></td>
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<td>0.7-1.5</td>
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</table>

$^a$ Species profiles in a premixed laminar flame.
$^b$ Flame speed, temperature is then the unburned gases temperature.
$^c$ Species profiles in a nonpremixed laminar flame.

To our knowledge, only two studies were performed on kinetic measurements related to the oxidation of ethylcyclohexane: in 2009, Vanderover et al. (35) measured the ignition delay times of a mixture of ethylcyclohexane/air in a shock tube ($T = 881–1319\, K; P = 10.8–69.5\, \text{bar}; \Phi = 0.25–1$) but did not perform any attempt of modeling. In 2011, Ji et al. (27) measured the laminar flame speeds of ethylcyclohexane/air mixtures in the counterflow configuration at atmospheric pressure ($T = 353\, K; \Phi = 0.7–1.5$). A good simulation of these results was obtained using the JetSurF 1.1 mechanism (44).

The purpose of this paper is to present results on the low- and intermediate-temperature oxidation of ethylcyclohexane in a jet-stirred reactor, with careful identification and analysis of the products.

Experimental Method

This study of the oxidation of ethylcyclohexane has been performed in a jet-stirred reactor which can be modeled as a perfectly stirred reactor (45,46). This reactor has been used by our research team in Nancy for numerous gas-phase kinetic studies (e.g., refs 47–49). The influence of surface reactions has been shown to be very minor in this kind of reactor (50). Experiments were performed under a constant pressure of 1.06 bar (800 Torr), at a residence time of $2 \pm 0.03\, s$, temperatures ranging from 500 to 1100 $K$, for the three equivalence ratios of 0.25, 1, and 2. Helium was used as diluent, and the study was performed with a fuel mole fraction of 0.005. Ethylcyclohexane, helium, and oxygen were provided by Sigma-Aldrich (purity $\geq 99\%$) and Messer (purity of 99.999% for He and 99.995% for O$_2$), respectively. Concerning the determination of equivalence ratios and fuel mole fraction, note that the main error is due to the error in the mass flow measurements (using Bronkhorst High-Tech Mass Flow Controller) which leads to an error of about 1%.

The reactor, made of fused silica, consists of a quartz sphere (volume = 95 cm$^3$) into which diluted reactants enter through an injection cross located at its center. It is operated under constant temperature and pressure and preceded by an annular preheating zone in which the temperature of the gases is progressively increased up to the reactor temperature. Gas mixture residence time inside the annular preheater is very short compared to its residence time inside the reactor (a few percent). Both the spherical reactor and the annular preheating zone are heated by resistances rolled up around the walls.

The outlet products were analyzed online by gas chromatography. Online analysis of products liquid under standard conditions was allowed by a heated transfer line between the reactor outlet and the chromatograph sampling valves which were also heated. During the study the temperature of the
transfer line was set at 473 K. This temperature was high enough to keep all the reaction products in the gas phase.

Three gas chromatographs were used for quantification of the different species. The first gas chromatograph, equipped with a Carbosphere packed column, a thermal conductivity detector (TCD), and a flame ionization detector (FID), was used for quantification of O₂, CO, CO₂, methane, ethylene, acetylene, and ethane. Among the products which are known to be important for hydrocarbon oxidation, only hydrogen, water, and formaldehyde were not quantified, but the presence of the two last one is well shown in GC-MS analyses. The second one was fitted with a PlotQ capillary column and a FID and used for quantification of C₂–C₄ hydrocarbons and small oxygenated compounds. The third one was fitted with a HP-5 capillary column and a FID and used for quantification of hydrocarbons and oxygenated species with more than five heavy atoms (i.e., carbon and oxygen atoms). Calibration was performed by injecting known amounts of pure substances when available; otherwise, the method of the effective carbon number (51) was used (species having the same number of carbon atoms and the same functional groups were assumed to have the same response in the FID). Calculated uncertainties on the species quantifications were about ±5% with online analysis of oxygen and C₁–C₂ hydrocarbons and about ±10% for analysis of other species. The detection threshold is better than 1 ppm for hydrocarbons and 100 ppm for carbon oxides quantified by TCD.

Identification and calibration of light species (e.g., carbon oxides, C₁–C₄ hydrocarbons) were performed by injecting gaseous samples provided by Messer and Air Liquide. Identification of species which were not calibrated before was performed using a fourth gas chromatograph coupled with a mass spectrometer (GC-MS) operating under the same conditions as the gas chromatograph used for quantification, enabling direct comparison of both chromatograms. Identification of these species was not performed “online”. The reaction products were condensed in a trap located at the outlet of the reactor and maintained at the temperature of liquid nitrogen. The content of the trap was then analyzed by GC-MS. Mass spectra of all detected reaction products were included in the spectra database “NIST 08”, except those with a molecular weight of 126 apart from 7-oxabicyclo[4.3.0]nonane. Note that the software of library identification of the MS always provides an answer, but when the proposed identification looked strongly unlikely according to the possible chemical pathways, a careful deciphering of the mass spectrum was necessary. Their deciphering was performed based on the experience gained in analysis of the oxygenated compounds formed during oxidation of alkanes (52).

Experimental Results

A table giving the complete set of experimental results for ethylcyclohexane is given as Supporting Information. Table 2 presents the 47 reaction products with a molecular weight lower than that of ethylcyclohexane (112 g/mol), which have been quantified online by gas chromatography. Figures 1–7 display the evolution as a function of temperature of the mole fractions of the reactants (ethylcyclohexane and oxygen) and of these 47 reaction products.
Figure 1. Evolution of the mole fractions of the reactants and carbon oxides with temperature for 3 equivalence ratios: (●) $\phi = 0.25$; (+) $\phi = 1$; (□) $\phi = 2$. Symbols are experiments and lines simulations.
Figure 2. Evolution of the mole fractions of the C₁–C₄ hydrocarbons with temperature for 3 equivalence ratios: (●) $\phi = 0.25$; (+) $\phi = 1$; (□) $\phi = 2$. Symbols are experiments and lines simulations.
Figure 3. Evolution of the mole fractions of the C_2–C_3 oxygenated species with temperature for 3 equivalence ratios: (●) $\phi = 0.25$; (+) $\phi = 1$; (□) $\phi = 2$. 
Figure 4. Evolution of the mole fractions of the C4–C6 oxygenated species with temperature for 3 equivalence ratios: (●) $\phi = 0.25$; (+) $\phi = 1$; (□) $\phi = 2$. 
Figure 5. Evolution of the mole fractions of C₅–C₆ nonaromatic hydrocarbons with temperature for 3 equivalence ratios: (●) $\phi = 0.25$; (+) $\phi = 1$; (□) $\phi = 2$. Symbols are experiments and lines simulations.
Figure 6. Evolution of the mole fractions of C7–C8 nonaromatic hydrocarbons with the temperature for 3 equivalence ratios: (●) $\phi = 0.25$; (+) $\phi = 1$; (□) $\phi = 2$. Symbols are experiments and lines simulations.
Figure 7. Evolution of the mole fractions of aromatic hydrocarbons with temperature for 3 equivalence ratios: (●) $\phi = 0.25$; (+) $\phi = 1$; (□) $\phi = 2$. Symbols are experiments and lines simulations.

As shown in Figure 1 the mole fraction profile of ethylcyclohexane and oxygen exhibit a very marked negative temperature coefficient (NTC) behavior. While the equivalence ratio has no real impact on the temperature for which the reactivity starts, it has a significant effect on ethylcyclohexane conversion from 625 to 780 K. The low-temperature reactivity is the largest for an equivalence ratio of 0.25. The stoichiometric mixture is less reactive, and the rich mixture shows the lowest reactivity. In addition, the extent of the NTC zone and conversion near 625 K (the temperature of the maximum of reactivity at low temperature) decrease noticeably when the equivalence ratio increases. Above 800 K, the equivalence ratio has a much lower impact. The same conclusion can be derived from the profiles of carbon monoxide. Formation of carbon dioxide occurs mostly above 800 K and is favored in lean and stoichiometric mixtures. However, as shown by Figure 8, which displays a zoom on the 550–800 K reaction zone for some of the major low-temperature reaction products, formation of 500 ppm of $\text{CO}_2$ is encountered at 625 K in stoichiometric mixtures.
Table 2. Reaction products (with a mass lower than that of the reactant (112)) identified and quantified during oxidation of ethylcyclohexane \((T = 500–1100 \text{ K}; \phi = 0.25, 1, \text{ and } 2; P = 800 \text{ Torr}; \tau = 2 \text{ s})\)

| Saturated and unsaturated linear hydrocarbons | methane, ethane, ethylene, acetylene, propene, 1-butene, 1,3-butadiene, 2-butene (cis and trans), 1-butyne (traces), 1,3-pentadiene (cis & trans) |
| Branched hydrocarbons | 3-methyl-1-butene, 2-methyl-1,3-pentadiene [ethylbutadiene]<sup>a,b</sup> |
| Aldehydes | acetaldehyde, acrolein (2-propenal), propanal, butanal, 2-butenal, pentanal |
| Ketones | acetone, 2-butane, 2-hexanone, 1-penten-3-one |
| Cyclic oxygenated species | ethylene oxide, oxetane, cyclohexanone, 2-cyclohexen-1-one |
| Other oxygenated species | carbon oxides, ethanol, acetic acid |
| Aromatics | benzene, toluene, ethylbenzene, styrene |
| Cyclic hydrocarbons | cyclopentene, 1,3-cyclopentadiene, cyclohexene, 1,3-cyclohexadiene, methylcyclohexane, methylenecyclohexane [1-methylcyclohexene]<sup>a</sup>, 3-methylcyclohexene, 3-ethylcyclohexene, 4-ethylcyclohexene, 1-ethylcyclohexene<sup>b</sup>, vinylcyclohexane, 3-ethylidenecyclohexene<sup>b</sup>, 4-ethylidenecyclohexene<sup>b</sup> |

<sup>a</sup> When another compound is given in square brackets that means it has a close mass spectrum, making fully certain identification difficult.

<sup>b</sup> Cis and trans forms can exist, but only one peak was observed.

Figure 8. Zoom on the 550–800 K reaction zone for some of the major low-temperature reaction products for a stoichiometric mixture.
Figure 2 presents the evolution with temperature of the mole fractions of the C\textsubscript{1}–C\textsubscript{4} hydrocarbons. Below 800 K, the only species of this type which have been detected are ethylene, propene, and 1-butene. As shown in Figure 8, at 625 K in stoichiometric mixtures, 140 and 20 ppm of ethylene and propene are formed, respectively. Above 800 K, large concentrations of these species are formed especially in the rich mixture, the major ones being methane, ethylene, and propene. Noticeable amounts (up to 200 ppm in rich mixtures) of acetylene and 1,3-butadiene are also observed.

Figure 3 displays the evolution with temperature of the mole fractions of C\textsubscript{2}–C\textsubscript{3} oxygenated species. Large concentrations of these species are formed even below 800 K, demonstrating again the high reactivity of ethylcyclohexane at low temperature. The most abundant species of this type is acetaldehyde with a maximum around 600 K of about 300 and 400 ppm under stoichiometric and lean conditions, respectively. Note that as shown in Figure 8 the maximum of acetaldehyde occurs at lower temperature than that of carbon oxides and small alkenes. Note also that an important formation of acetic acid was also observed in a similar amount as acetaldehyde in lean mixtures and below 800 K but with a higher sensitivity to the change in equivalence ratios than that of the aldehyde. We have no evidence showing this behavior for other intermediates, except perhaps compounds including two carbonyl groups. This profile of acetic acid is not presented in Figure 3, because as shown by Herbinet et al. (49) in the case of n-heptane its formation is probably mostly due decomposition of ketohydroperoxides in the transfer line. Among other important products are ethylene oxide, acrolein (up to 300 ppm in lean mixtures), and propanal. Formation of these species is favored in lean mixtures.

Figure 4 presents the evolution with temperature of the mole fractions of C\textsubscript{4}–C\textsubscript{6} oxygenated species, which are also more easily formed in lean mixtures. Formation of these species is lower than that of most of the smaller oxygenated species and favored below 800 K, apart from 2-butenal. The most abundant of these species are the cyclic ones: cyclohexanone and cyclohexenone (with mole fraction up to 120 ppm at 625 K under lean conditions).

Figure 5 presents the evolution with temperature of the mole fractions of C\textsubscript{5}–C\textsubscript{6} nonaromatic hydrocarbons which are all mainly produced above 800 K. Cyclic C\textsubscript{6} species (1,3-cyclohexadiene and cyclohexene) are the most abundant hydrocarbons formed during this study apart from C\textsubscript{1}–C\textsubscript{3} species, with a maximum of 300 ppm. Note that 1,3-cyclohexadiene is formed in larger amounts in lean mixtures, while formation of all other produced C\textsubscript{1}–C\textsubscript{6} hydrocarbons is larger in rich mixtures. Cyclic C\textsubscript{5} species (1,3-cyclopentadiene and cyclopentene) are much less abundant with a maximum of 70 and 50 ppm, respectively. The most abundant of the acyclic species is 1,3-pentadiene (with a maximum mole fraction of 130 ppm). Two chromatographic peaks have been observed corresponding to the cis and trans isomers, but it has not been possible to attribute a peak to a given isomer.

Figure 6 presents the evolution with temperature of the mole fractions of C\textsubscript{7}–C\textsubscript{8} nonaromatic hydrocarbons. Methylene cyclohexane is the most abundant among C\textsubscript{7} products, with a maximum of 90 ppm at 875 K under rich conditions but also slight formation at lower temperature (up to 10 ppm at 625 K in lean mixtures). All C\textsubscript{8} products have similar mole fractions below and above 800 K with a marked NTC behavior in rich and stoichiometric mixtures. Under lean conditions formation of these products is significantly favored and the NTC behavior is much less pronounced. The most abundant of these species are ethylcyclohexene isomers with a maximum total formation of 270 ppm. It was not possible to distinguish between the 2- and the 3-ethylcyclohexenes. Formation of vinylcyclohexane is much lower (up to 25 ppm). Note that among unsaturated C\textsubscript{8} compounds vinylcyclohexane and ethylidenecyclohexenes have been observed but not ethylidenecyclohexane or vinylcyclohexenes. In addition, while their formation could have been expected, no ethylcyclohexadiene was detected in noticeable amounts. While isomers of heptene have been detected in flames of methycyclohexane by Skeen et al. (36), acyclic C\textsubscript{8} compounds of the same mass.
as the reactant were not observed in the present study, even in rich mixtures. Formation of isomers of octane could be obtained through an internal disproportionation of biradicals formed from the cyclic reactant by unimolecular initiations involving breaking of a C–C bond in the ring.

Figure 7 presents the evolution with temperature of the mole fractions of the four aromatic hydrocarbons analyzed. Benzene is the aromatic species formed in the largest amount (up to 260 ppm at 875 K). However, like toluene, it is only formed above 800 K. Ethylbenzene and styrene are formed in lower amounts (up to 70 ppm) but are produced also in a noticeable way below 800 K.

In addition to these products of molecular weight lower than that of ethylcyclohexane (molecular weight = 112), a lot of molecules having a molecular weight of 126 (C₈H₁₂O formula) were detected in a large amount. Figure 9 displays the evolution with temperature of the total mole fraction of these species, which are formed both below and above 800 K and are more abundant in lean mixtures, especially above 625 K (up to 1000 ppm for the sum of these isomers). The equivalence ratio has a larger impact on formation of these species from 650 to 850 K than at lower temperatures, which lead us to think that the low-temperature mechanism is certainly still important above 800 K in lean mixtures.

![Figure 9. Evolution of the total mole fraction of the species with a molecular weight of 126 with temperature for 3 equivalence ratios: (●) \( \phi = 0.25 \); (●) \( \phi = 1 \); (□) \( \phi = 2 \).](image)

Among these C₈H₁₂O products, interpretation of mass spectra has allowed us to propose a structure for the 8 compounds which are listed in Table 3. An example of a chromatogram related to these products is given in Figure 10. Mass spectra of the 7 compounds with a molecular weight of 126 which were identified in addition to 7-oxabicyclo[4.3.0]nonane are given as Supporting Information. As shown in Figure 10, the chromatograms also include many small peaks for which a reliable identification was not possible. For longer residence times other small peaks could correspond to compounds with a C₈H₁₂O₂ formula (a molecular weight of 140). These compounds could be species including two carbonyl groups in the same way as what has recently been observed during oxidation of n-butane (48) and n-heptane (49). Figure 11 presents the evolution with temperature of the mole fractions of the seven C₈H₁₄O species for which quantification was possible. It shows that octenone is the most abundant of that type of compound with a larger formation under lean conditions and a higher maximum at 825 K (up 700 ppm) than at 625 K (up 400 ppm). Formation of this unsaturated ketone is notably more favored in lean mixtures than that of all the other products displayed in Figure 11. The bicyclic ethers, with both cycles sharing at least a bond, are formed in similar amounts whatever the size of the additional ring: 3 or 5 members (maximum at 625 K of about 90 ppm). 3-Methyl,2-oxaspiro[1,5]octane and cyclohexylmethylketone are formed in lower amounts (maximum at 625 K below 50 ppm).
Figure 10. Example of chromatogram obtained with mass spectrometry detection for products with a molecular weight of 126.

Figure 11. Evolution with the temperature of the mole fractions of the species with a molecular weight of 126 which can be quantified: (●) $\phi = 0.25$; (+) $\phi = 1$; (□) $\phi = 2$. 
Table 3. Reaction products (with a mass with a molar mass of 126) identified and quantified during oxidation of ethylcyclohexane

<table>
<thead>
<tr>
<th>Name</th>
<th>Structure</th>
</tr>
</thead>
<tbody>
<tr>
<td>3-methyl,2-oxaspiro[1,5]octane</td>
<td><img src="image1" alt="structure" /></td>
</tr>
<tr>
<td>1-ethyl,7-oxabicyclo[4.1.0]heptane</td>
<td><img src="image2" alt="structure" /></td>
</tr>
<tr>
<td>oct-7-en-3-one</td>
<td><img src="image3" alt="structure" /></td>
</tr>
<tr>
<td>1-methyl,7-oxabicyclo[2.2.1]heptane</td>
<td><img src="image4" alt="structure" /></td>
</tr>
<tr>
<td>7-oxabicyclo[4.3.0]nonane</td>
<td><img src="image5" alt="structure" /></td>
</tr>
<tr>
<td>(2 peaks corresponding to 2 isomers)</td>
<td></td>
</tr>
<tr>
<td>cyclohexymethylketone</td>
<td><img src="image6" alt="structure" /></td>
</tr>
<tr>
<td>2-cyclohexylacetaldehyde</td>
<td><img src="image7" alt="structure" /></td>
</tr>
<tr>
<td>(2 peaks corresponding to 2 isomers, but one has the same retention time as that of 1-methyl,7-oxabicyclo[2.2.1]heptane)</td>
<td></td>
</tr>
<tr>
<td>Ethylcyclohexanone</td>
<td><img src="image8" alt="structure" /></td>
</tr>
<tr>
<td>(The peak obtained with FID detection is too small to allow quantification)</td>
<td></td>
</tr>
</tbody>
</table>

The carbon balance is generally better than 10%, apart from the case of the study of the leanest mixtures (\(\phi = 0.25\)) around 650 K for which heavy oxygenated compounds condensing directly at the reactor outlet are certainly formed.
Discussion

Comparison with Other C₈ Hydrocarbon Reactants

To compare the reactivity of alkycyclohexanes with that of other types of hydrocarbons we studied the oxidation of two other C₈ hydrocarbons, n-octane and 1-octene, under the same conditions for stoichiometric mixtures. The results are plotted in Figure 12 for the fuel mole fraction and for that of some representative reaction products (CO, acetaldehyde, C₂H₄, C₂H₂, benzene, and ethylene oxide). This figure shows that the reactivity of ethylcyclohexane is very close to that of 1-octene and much lower than that of n-octane below 800 K. This is also shown by the much larger formation of carbon monoxide, aldehydes (see acetaldehyde in Figure 12), or alkenes (see ethylene in Figure 12).

![Figure 12. Comparison of the evolution of the mole fractions of the reactants and several products obtained during oxidation of stoichiometric mixtures containing different C₈ reactants: (●) ethylcyclohexane; (○) 1-octene; (■) n-octane.](image-url)
In addition, the extent of the NTC zone is much larger for the linear alkane than for the two unsaturated compounds. The presence of the cycle, like that of a double bond, makes isomerization of peroxy radicals more difficult and then hinders the low-temperature reaction. Note however that formation of ethylene oxide is larger for 1-octene and ethylcyclohexane. Above 800 K, the reactivity of the three C₆ hydrocarbons is very similar (still slightly larger for n-octane) as can be shown by fuel conversion and formation of carbon monoxide and acetaldehyde. However, formation of ethylene and ethylene oxide is larger for the linear alkane than for the alkene and cyclic compounds. On the other hand, formation of acetylene is largest for ethylcyclohexane and 1-octene. It is worth noting that because of the presence of a C₆ ring in the molecule formation of benzene is considerably larger in the case of ethylcyclohexane than for the two acyclic C₆ hydrocarbons. Alkylcyclohexanes are then a favored source of aromatic compounds at intermediate temperatures, which are soot precursors.

**Comparison with Simulations Performed with a Mechanism Available in the Literature**

Simulations have been performed using the JetSurF 1.1 mechanism (44), which was already used with success to model laminar flame speeds of ethylcyclohexane/air mixture at atmospheric pressure (27). Since this mechanism has a short globalized low-temperature submechanism we tested it over the full temperature range studied. Comparison between experimental and predicted results is presented in Figures 1, 2, 5, 6, and 7. Figure 1 indicates that the model predicts well the reactivity above 770 K, as shown by the reactants and carbon oxide profiles. At lower temperatures, no reactivity is predicted while a significant experimental one was observed. Prediction of the formation of C₃–C₅ hydrocarbons is also relatively good, apart from that of acetylene and 1-butene which is significantly overestimated especially in rich mixture. As formation of oxygenated species by a low-temperature mechanism is not adequately taken into account, no comparison is presented for the species of Figures 3 and 4. Figure 5 shows a satisfactory prediction of formation of 1,3-cyclopentadiene, but the agreement is less good for other C₅–C₆ hydrocarbons. Figure 6 shows that while methylcyclohexane and vinylcyclohexane are included in the model, no formation of C₇–C₈ nonaromatic hydrocarbons is predicted. As displayed in Figure 7, formation of benzene is well predicted in rich and stoichiometric mixtures but overestimated in lean mixture. Formation of toluene is considerably underestimated, while that of ethylbenzene and styrene is not considered by the model. Note that the model also predicts noticeable amounts of H₂, H₂O, H₂O₂, CH₂O, and CH₃CO, which cannot be quantified by the experimental method used in the present study.

**Ways of Reactions Important To Explain Formation of the Main Products**

We tried below to identify the reactions which are of importance to explain formation of the main products observed in this study.

- **Ways of Reactions Important Above 800 K**

Figure 13 shows a reaction rate analysis for a temperature of 850 K under the conditions of Figures 1–7 (for stoichiometric mixtures) corresponding to 90% conversion of ethylcyclohexane. Under these conditions ethylcyclohexane is mainly consumed by H abstractions mostly with OH radicals to give the six possible ethylcyclohexyl radicals (RC1–RC6). RC3 radicals can be obtained by abstraction of one tertiary H atom (13% of ethylcyclohexane consumption, RC1 radicals by that of three primary H atoms; 13% consumption), RC2 (8% consumption) and RC6 (11% consumption) radicals by that of 2 secondary H atoms, and finally RC4 (28% consumption) and RCS (24% consumption) radicals can be produced by abstraction of 4 secondary H atoms.
Above 800 K, the most important type of reaction consuming ethylcyclohexyl radicals is by breaking of a C–C bond. \( \beta \)-Scissions from RC1 radicals lead to ethylene and cyclohexyl radicals, which produce by opening of the cycle ethyl radicals and 1,3-butadiene, a major product of this study as shown in Figure 2. RC2 radicals react by breaking a C–C bond with opening of the cycle, yielding resonance-stabilized radicals, which isomerize and then decompose to give propyl radicals, a source of propene, and 1,3-pentadiene (see Figure 5). \( \beta \)-Scissions from RC3 radicals produce methylenecyclohexane, which is an important product of our study as shown in Figure 6. \( \beta \)-Scissions from RC4 radicals lead to ethyl radicals and cyclohexene, which is a source of 1,3-cyclohexadiene and then of benzene. The rate constant of the \( \beta \)-scission from RC4 radicals yielding ethyl radicals and cyclohexene has been estimated in the JetSurf 1.1 mechanism (44) as that of 2-methyl-pent-4-yl radicals, while Sirjean et al. (53) has shown that the \( \beta \)-scission involving formation of a double bond inside the cycle have an activation energy about 7 kcal/mol larger than for acyclic radicals. That certainly explains the large overprediction observed for cyclohexene, as shown in Figure 5. 1,3-Cyclohexadiene is mostly obtained by H abstractions from cyclohexene followed by reaction with an oxygen molecule to give \( \text{HO}_2 \) radicals and this diene. Benzene was mostly obtained by H abstractions from 1,3-cyclohexadiene followed by C–H bond breaking. The rate constant used in the JetSurf 1.1 mechanism (44) for formation of 1,3-cyclohexadiene by reaction with \( \text{O}_2 \) \( (k = 1 \times 10^{-11} \text{ cm}^3\text{-mol}^{-1}\text{s}^{-1}) \) is about 400 times smaller than the rate constant of the \( \beta \)-scission from RC4 radicals.
faster at 800 K than the rate constant proposed by Touchard et al. (54) \((k = 3.2 \times 10^{12} \text{ cm}^3\text{mol}^{-1}s^{-1})\). That certainly explains why formation of 1,3-cyclohexadiene is correctly predicted, while cyclohexene is overpredicted, and the simulated formation of benzene is favored in lean mixture, while it is not the case experimentally (see Figure 7). Breaking of a C–C bond with opening of the cycle of RC5 radicals is a source of 1-butene, an important reaction product above 800 K. \(\beta\)-Scissions from RC6 radicals produce radicals which readily isomerize to give resonance-stabilized radicals, which decompose yielding 1,3-butadiene and 2-butyl radicals. Note that a significant part (about 20%) of RC6 radicals also isomerizes to RC2 radicals.

RC1 and RC2 radicals could also react with an oxygen molecule to produce HO_2 radicals and vinylcyclohexane, a minor product of this study (see Figure 6). RC3 and RC2 radicals could also react with an oxygen molecule to produce HO_2 radicals and ethylidenecyclohexane, which has not been observed but could be the source of the ethylidenecyclohexenes, the mole fraction profiles of which are displayed in Figure 6. RC3, RC4, RC5, and RC6 radicals could also react with an oxygen molecule to produce HO_2 radicals and the different isomers of ethylcyclohexenes. Via formation of ethylcyclohexadienes, these species are certainly a good source of ethylbenzene and then styrene. This series of reactions leading to aromatic compounds can be important even at low temperature, which explains why a noticeable formation of ethylbenzene and styrene has been observed below 800 K as shown in Figure 7. These reactions are shown with dotted arrows in Figure 14, because formation of cyclic C_6H_{14} species is not considered in the JetSurF 1.1 mechanism (44). Flow rate analysis shows that within the present model ethylcyclohexyl radicals also react with oxygen molecules to give a globalized radical which at 850 K mostly decomposes to give cyclohexane, ethylene, and HO_2 radicals. These reactions have an important flow rate in the case of RC1, RC5, and RC6 radicals.

- **Possible Ways of Reactions Important below 800 K**

According to Walker and Morley (55), cyclohexane reacts at low temperature following the same pathway as that proposed for alkanes and shown in Figure 14. Alkyl radicals R• are produced by H abstractions and add thereafter to O_2 and yield peroxy radicals ROO•. Some consecutive steps will then lead to formation of hydroperoxides, which are degenerate branching agents explaining the high reactivity of the alkanes at low temperature. When the temperature increases, the equilibrium R• + O_2 = ROO• begins to be displaced back to the reactants and formation of ROO• radicals is less favored. The larger concentration of R• radicals increases the rate of the strongly inhibiting oxidation reaction R• + O_2 = •HO_2 + conjugated olefin. Furthermore, unimolecular reaction channels from •QOOH radicals become important and lead to formation of cyclic ethers, ketones, or conjugated olefins.

Considering that the same scheme can be applied to alkylcyclohexanes, Figure 15 presents the ways of formation which can be envisaged to explain formation of the main products observed in this study below 800 K. At higher temperatures ethylcyclohexane is consumed by H abstractions to form the six cyclohexylethyl radicals previously described. In the lower part of the temperature range studied cyclohexylethyl radicals react mainly by addition to oxygen molecules to produce 6 cyclohexylethylperoxy radicals (shown in Figure 15 only for RC3 and RC4 radicals). These 6 cyclohexylethylperoxy radicals can lead to a large number of hydroperoxy radicals via isomerization by internal transfer of a H atom. Figure 15 presents only the hydroperoxy radicals leading to compounds which have been experimentally observed.
Figure 14. General scheme of low-temperature oxidation of an alkane (RH).

Note that the observed intermediates show that isomerizations by transfer of a tertiary H atom, which require a lower activation energy for the same ring strain energy in the bicyclic transition state, are favored. This type of isomerization occurs in the case the peroxy radicals deriving from RC6, RC5, RC4, and RC2 radicals, and decomposition of the obtained hydroperoxy radicals lead to OH radicals and 1-methyl,7-oxabicyclo[2.2.1]heptane, oct-7-en-3-one, 1-ethyl,7-oxabicyclo[4.1.0]heptane, and 3-methyl,2-oxaspiro[1,5]octane, respectively. Octenone is obtained by decomposition of a four-membered ring cyclic ether, according to what has been shown by Walker and Morley (55) in the case of cyclohexane: the bicyclic ether with an additional ring involving 4 members is very unstable and decomposes very quickly to 1-hexenal. Figure 16 presents this pericyclic reaction in the case of the cyclic ether, leading to formation of octenone. Note that branched unsaturated C₈ aldehyde, which has not been experimentally observed, could also be a possible product. Analysis of this reaction for all possible bicyclic ethers with an additional ring involving 4 members deriving from ethylcyclohexane shows that this channel is the only one leading to an acyclic ketone: all other ones lead either to C₈ aldehydes, to cyclohexanone and ethylene, or to smaller aldehydes and alkenes.
Figure 15. Ways of formation which can be envisaged to explain formation of the main products observed in this study below 800 K.

Other types of isomerisations which are favored are those abstracting a H atom from an atom of carbon also carrying a O atom. As four-membered ring isomerizations require a high strain energy, these isomerisations do not occur directly but certainly in two steps. The first step is formation of a hydroperoxy radical though abstraction a H atom from an atom of carbon not carrying any O atom. The second step is isomerization abstracting this time a H atom from the atom of carbon also carrying a O atom. This type of isomerization requires a lower activation energy for the same ring strain energy of the transition state and the same type of abstracted H atom. This is the case of peroxy radicals deriving from RC4, RC2, and RC1 radicals, with production of OH radicals and 2-ethyl-cyclohexanone, cyclohexymethylketone, and 2-cyclohexylacetaldehyde, respectively. The only product which does not derive from the two previously described types of isomerisations is 7-oxabicyclo[4.3.0]nonane. This product can be obtained from hydroperoxy radicals deriving from RC4...
and RC1 radicals. The channel deriving from RC1 radicals is certainly favored as it involves transfer of 4 secondary H atoms, instead of 3 primary ones in the case of the channel deriving from RC4 radicals. In both cases, the transition state involves formation of a new seven-membered ring, outside from the cyclohexyl cycle. In this case no additional strain energy is due to formation of the bicycle.

\[
\begin{align*}
\text{Figure 16. Mechanism of the pericyclic reaction leading to octenone.}
\end{align*}
\]

In addition to decomposition to cyclic ethers, aldehydes, or ketone, the hydroperoxy radicals can also add a second time to an oxygen molecule to form radicals which in their turn isomerize and decompose to form ketohydroperoxides. An example of formation of a ketohydroperoxide from RC1 is shown in Figure 15. Ketohydroperoxides, which are responsible for the high reactivity observed from 600 K, easily decompose and are certainly the source of the C₂–C₅ aldehydes and ketones observed.

Cyclohexylethylperoxy radicals can also react with HO₂ radicals to yield cyclohexylethylhydroperoxides. Such a channel is displayed for the peroxy radicals deriving from RC3 radicals. In this case decomposition of the obtained cyclohexylethylhydroperoxide can easily lead to formation of cyclohexanone, an important product of this study as shown in Figure 4.

**Conclusion**

A study of the oxidation of ethylcyclohexane has been performed in a jet-stirred reactor under quasi-atmospheric pressure, at a residence time of 2 s, at temperatures ranging from 500 to 1100 K, for three equivalence ratios (0.25, 1, and 2), and a fuel mole fraction of 0.005. A high reactivity has been observed at temperature below 650 K with a marked NTC behavior. The low-temperature reactivity is lower than that of n-octane but close to that of 1-octene. A wide range of products has been identified both below and above the NTC zone, showing a high complexity of the involved chemistry. Formation of all the observed products can be qualitatively explained using reactions pathways directly derived from those usually proposed for oxidation of acyclic alkanes (55). Reactions of all isomers of cyclohexylethyl radicals have to be considered carefully, with no obvious a priori simplification. A noticeable formation of aromatic species (ethylbenzene and styrene) can be observed at temperatures as low as 600 K.

Better knowledge of the chemistry of the oxidation of alkylcyclohexanes will require development of a complete low-temperature mechanism in order to model the present results but also to extend the experimental methodology described here to study compounds with a longer alkyl chain such as n-butylcyclohexane or n-hexylcyclohexane.
Supporting Information

Mass spectra of compounds of mass 126 are given as well as a table of full experimental data. This material is available free of charge via the Internet at http://pubs.acs.org.bases-doc.univ-lorraine.fr.

Acknowledgment

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References