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To cite this version:
Marion Blocquet, Coralie Schoemaecker, Damien Amedro, Olivier Herbinet, Frédérique Battin-Leclerc, et al.. Quantification of OH and HO2 radicals during the low-temperature oxidation of hydrocarbons by Fluorescence Assay by Gas Expansion technique. Proceedings of the National Academy of Sciences of the United States of America, National Academy of Sciences, 2013, 110 (50), pp.20014-20017. 10.1073/pnas.1314968110. hal-00923973

HAL Id: hal-00923973
https://hal.archives-ouvertes.fr/hal-00923973
Submitted on 6 Jan 2014

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Quantification of OH and HO₂ radicals during the low-temperature oxidation of hydrocarbons by Fluorescence Assay by Gas Expansion technique

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Abstract

•OH and •HO₂ radicals are known to be the key species in the development of ignition. A direct measurement of these radicals under low-temperature oxidation conditions (T = 550 – 1000 K) has been achieved by coupling a technique named fluorescence assay by gas expansion, an experimental technique designed for the quantification of these radicals in the free atmosphere, to a jet-stirred reactor, an experimental device designed for the study of low-temperature combustion chemistry. Calibration allows conversion of relative fluorescence signals to absolute mole fractions. Such radical mole fraction profiles will serve as a benchmark for testing chemical models developed to improve the understanding of combustion processes.

Keywords: HO₂ radicals; autoignition

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In the context of a needed significant reduction of the emission of greenhouse gases, understanding the autoignition chemistry of hydrocarbons and biofuels is of critical importance to allow the development of new combustion strategies for clean and efficient internal combustion engines (1). Advanced combustion concepts that rely on compression self-ignition (1,2), as well as the improvement of safety in oxidation processes (3), demand an improved understanding of the reaction kinetics governing the detailed mechanisms of organic compounds, particularly in the low-temperature oxidation regime. The chemistry of oxidation and autoignition is governed by species, with unpaired electrons, called free radicals, which are present in trace amounts. Among free radicals important in oxidation are H atoms, alkyl radicals, and hydroxyl (•OH) or hydroperoxy (•HO$_2$) radicals, the last two being the most important. It is then of critical importance for a better understanding of this chemistry to develop methods to measure radicals under all possible reaction conditions governing autoignition, including low-temperature oxidation (below 1000 K).

Because •OH radicals are present in relatively high concentrations at high temperatures, they have been measured for a long time in flames by molecular beam mass spectrometry (4) and by laser-based methods such as laser-induced fluorescence (LIF) or absorption spectroscopy (5,6). The detection of •HO$_2$ radicals has recently been achieved in a flame using a photo-fragmentation laser-induced fluorescence technique (7) as well as during the oxidation of dimethyl ether in a laminar flow reactor using midinfrared Faraday rotation spectroscopy (8). A method to measure the ratio between •HO$_2$ and •RO$_2$ radicals based on electron spin resonance spectroscopy was proposed by Carlier and Sochet (9) but has never been applied to real hydrocarbons and low-temperature oxidation.

H atoms (10) or •OH radicals (11) were also followed under high-temperature (T > 1200 K) shock tube conditions. Finally, planar LIF using a laser sheet has been used in optical access engines to image OH radical concentrations to map the hot combustion zones (12). This technique does not directly give access to a real quantitative measurement of the radical concentration due to differences in quenching and line broadening caused by local gradients of fuel/air equivalence ratio and temperature (13).

In this work, we present the coupling of an optical technique named Fluorescence Assay by Gas Expansion (FAGE), initially designed for the measurement of absolute concentrations of •OH and •HO$_2$ radicals in the atmosphere (14), to a combustion device called a jet-stirred reactor (JSR), with the goal of quantifying both radicals under reaction conditions similar to those found before development of autoignition (2). A JSR allows studying gas-phase reactions at temperatures up to 1000 K without external radical production. Such measurements will be particularly valuable for improving knowledge of the chemistry and detailed kinetic models in the temperature zone (650 – 900 K), in which reaction mechanisms are the most complex and the kinetic data are the least accurate (15).

The schematic reaction mechanism currently accepted for the low-temperature combustion of many typical hydrocarbon fuel molecules is shown in Figure 1, in which the key reactions are numbered to facilitate the following discussion. The starting point of the chain reaction leading to ignition is the reaction of a hydroxyl radical (•OH), the main chain carrier, with the fuel molecule, named hereafter RH. Abstraction of an H atom by the OH radical (reaction 1) leads to formation of a radical •. At low temperatures (below around 750 K), • rapidly yields a peroxy radical (ROO•) after a barrierless reaction with an oxygen molecule (reaction 2). This ROO• radical leads then to the formation of hydroperoxides and an •OH radical, mainly by a complex mechanism involving two isomerizations and addition of a second oxygen molecule, not described here. The obtained hydroperoxides can easily decompose giving two radicals, including a second •OH radical. This degenerate chain-branching reaction explains the high reactivity of hydrocarbon/oxygen mixtures at low temperatures.

An increase in temperature enhances the reversibility of reaction 2, thus hindering the formation of peroxy radicals and consequently of •OH radicals. At moderate temperatures (between around 750 and 950 K), the radical • reacts with oxygen, yielding an unsaturated hydrocarbon (named alkenes in Fig. 1) and an •HO$_2$ radical (reaction 3), or, if temperature is high enough,
decomposes into a small alkene and another radical 'R•' (reaction 4). •HO₂ radicals are much less reactive than •OH radicals and mainly react in a termination step leading to hydrogen peroxide (H₂O₂) (reaction 5). This reduced importance of degenerate chain-branching reactions in favor of chain-terminating reactions leads to a slowdown of the reactivity and explains the occurrence of a feature specific to mixtures of organic compounds with oxygen (16): the commonly called “negative temperature coefficient” (NTC) zone; i.e., a zone (usually around 650 K under atmospheric pressure) where the reactivity decreases with increasing temperature.

With further increased temperature, the decomposition of H₂O₂ leading to two •OH radicals becomes faster and is the dominant degenerate chain-branching reaction, again promoting high reactivity. Above around 950 K, small hydrocarbon radicals such as obtained by reaction 4 can in turn decompose, leading to alkenes and •H atoms. The reaction of an •H atom with oxygen at high temperatures leads to an •OH radical and to an •O• atom (reaction 6), which in turn regenerates an •OH radical by H abstraction from the fuel. Reaction 6 is a true branching step, ensuring the full development of ignition and complete combustion. However, the pressure-dependent reaction 7 can compete with reaction 6, depending on temperature and pressure, and leads to the formation of the less reactive •HO₂ radicals.

This scheme shows clearly the critical importance of •OH and •HO₂ radicals in all stages linked to the development of ignition. The purpose of this study is therefore to present an experimental device allowing the quantification of both radicals under conditions observed during the reaction period before the autoignition of hydrocarbons. n-Butane has been studied as it is the smallest hydrocarbon structure that exhibits an oxidation behavior representative to species commonly present in gasoline and diesel fuels.

The goal of this work was to reproduce the chemistry leading to this phenomenon rather than to observe unsteady phenomena directly such as autoignition. For this purpose, a heated JSR has been used, a device that has already been used many times for studying the chemistry of organic compounds before ignition (15). Note that this reactor has been recently used to follow the formation of alkylhydroperoxides (17) and H₂O₂ (18) through its coupling with a reflectron time-of-flight mass spectrometer and cw-Cavity Ring Down Spectroscopy (CRDS) cell, respectively. CRDS is known to be a very sensitive technique to measure •HO₂ radicals, but surprisingly these radicals

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**Figure 1:** Simplified scheme of the mechanism of oxidation of hydrocarbons (dotted arrows represent a succession of several elementary steps).
could not be detected in these experiments, even though concentrations much above the detection limit of this technique (19,20) were expected (18).

**Results and Discussion**

FAGE is a highly sensitive method for quantifying •OH and •HO$_2$ radicals, originally designed for the quantification of these radicals in the free atmosphere. Mole fractions of •OH and •HO$_2$ radicals found in the atmosphere are typically 0.2 and 8 ppt, respectively (21). Today eight research groups in the world have operational FAGE systems (22–28), but all of them are used in research solely linked to atmospheric chemistry. To our knowledge, no other work has been published where the advantages of radical detection by FAGE technique are exploited in the field of combustion research.

FAGE is based on LIF after expansion of the gas mixture (in general, the ambient atmosphere) through a small orifice (few 100 µm to 1 mm) to pressures of around 0.5 torr. •HO$_2$ radicals, which do not fluoresce, are converted to •OH radicals by their fast reaction with NO. Indirect detection of •HO$_2$ after conversion to •OH radicals through addition of NO to the reaction system is a common technique (29), leading possibly to complications due to the reaction of NO with other intermediates. In the case of FAGE, however, NO is added only after the expansion of the gas mixture into the FAGE cell; i.e., when the reaction system is already at low temperature and pressure. Recent experiments have shown that, even under these conditions, •RO$_2$ radicals might also be converted to •OH radicals in a two-step reaction sequence with the conversion efficiency depending strongly on the NO concentration (30). In the present experiments, the NO concentration was kept very low (see supplementary material) and no interference has been detected.

![Diagram](image)

**Figure 2:** Simplified scheme of the experimental device (not to scale). The upper part (JSR) is at atmospheric pressure and heated at high temperatures, and the lower part (FAGE) is at low pressure (< 1 torr) and ambient temperature.

The FAGE system of Lille has been described in detail several times (31–33), so only its coupling with the JSR will be described here. The spherical quartz JSR (diameter around 5.5 cm), in which a gas mixture is continuously flowing, was operated at atmospheric pressure and was electrically heated to a constant temperature for each data point of a temperature profile (see Figures 3 and 4). This type of reactor, which can be heated up to 1,200 K, is well adapted for mechanistic studies (see supplementary material for more details): the gas phase within the reactor is well stirred, meaning that concentrations and temperatures are close to homogenous.
As schematically shown in Figure 2, the JSR was adapted for the coupling with the FAGE system by transforming the gas exhaust from the usual tube into an orifice with a diameter of 7 mm. Only a small fringe of roughly 2 mm at the rim of the opening was kept to maintain the heating coils. The inlet of the FAGE was made from a stainless-steel cone with an orifice of 400 μm: this cone was placed in the center of the exhaust hole of the JSR. The exhaust of the JSR was not connected leak tight to the FAGE inlet to avoid perturbing the chemistry within the JSR; the air intake into the FAGE is higher than the output of the JSR. For calibration purposes, it has been estimated from flow rate measurements that the gas mixture of the JSR is diluted by a factor of 2.6–5.2 during the intake into the FAGE (see supplementary material). This contamination with ambient laboratory air has no impact on the chemistry, because ambient indoor air contains only extremely low radical concentrations. A loss of •OH radicals during the short transition period from the JSR to the FAGE inlet due to reaction with trace gases possibly present in the laboratory air can safely be neglected compared with the concentration of n-butane and its oxidation products. The JSR is surrounded by a Plexiglas cylinder to avoid air drafts: for a few experiments, this cylinder has been flooded with N2 (Alpha 3.0), and no differences have been observed in the LIF signal.

Figure 3 shows the •OH and •HO2 profiles obtained during the oxidation of a mixture of 2.3% n-butane/13% O2 in helium at a residence time of 6 s within the JSR. Each data point of such a profile is the result of an individual experiment: the electrical heating of the JSR is set to obtain the desired temperature, the gas flows are adapted to keep the residence time constant (the change in gas density with temperature needs to be taken into account), and once the chemical system has stabilized to the new conditions, the steady-state concentrations of the different species are measured at the exit of the JSR. Figure 3 also presents the n-butane mole fraction profile measured by gas chromatography under the same conditions (18). The reaction starts when the JSR temperature reaches around 600 K, visible through a sharp increase in •OH and •HO2 LIF signal following the evolution of the n-butane consumption. The LIF signal intensity reaches a maximum for both radicals roughly at the same JSR temperature at which the n-butane concentration reaches a minimum. When further increasing the temperature of the JSR, the NTC zone is clearly visible above 650 K, the LIF intensity decreases to a minimum for both radicals, but nonzero LIF signal is still present near 700 K, a temperature for which no conversion of n-butane is observed. Further increase of the JSR temperature leads to a second increase in reactivity with an initially steeper rise in the •HO2 signal compared with that of the •OH signal. Above 900 K, the •HO2 signal decreases again, whereas the •OH signal sharply rises.

Because FAGE is based on LIF, it is only a relative method, and a calibration is needed for obtaining absolute concentrations. Different calibration techniques have been developed for the quantification of radicals in the atmosphere (34). One of these methods, based on the photolysis of H2O2, has been set up for the FAGE system in Lille. However, it cannot be applied straightforward for the calibration of the measurements presented in this work because typical radical concentrations in atmospheric chemistry and in combustion systems differ by many orders of magnitude, especially for HO2 radicals (parts per trillion in the atmosphere, parts per million in combustion systems). Thus, the FAGE detection system, if directly used in the “atmospheric” configuration, would saturate when analyzing combustion exhaust gases and the sensitivity of the instrument has been attenuated (see supplementary material).

However, the calibration system is designed for generating typical atmospheric concentrations, and therefore the signals would be very weak if the “atmospheric” calibration system would be used in the attenuated “combustion” configuration. An attempt has nevertheless been made to convert relative LIF signals into absolute concentrations and then into mole fractions for both, •OH and •HO2 radicals. The profiles are shown in Fig. 4, together with model predictions. For this purpose, several corrections have been applied to the calibration factors such as obtained by the atmospheric calibration system (for details, see supplementary material). Note that the mole fractions shown for both radicals in Fig. 4 are probably only lower limits, because (i) the dilution factor (see supplementary material) has been calculated by considering that the JSR reactor flow is completely sampled into the FAGE, which is probably not true, and (ii) loss of radicals can be
expected between the exit of the JSR and the expansion into the FAGE. •OH radicals are much more reactive than •HO₂ radicals and therefore their concentration will probably decrease faster. Therefore, the agreement between the model and the measurement is excellent for the maximal mole fractions of •HO₂ radicals at both temperatures with the model predicting somewhat higher •HO₂ concentrations. As for the concentration of •OH radicals, the model underpredicts the concentration at nearly all temperatures; only at temperatures above 900 K is the measured •OH concentration lower than the predicted.

Figure 3: •OH (−) and •HO₂ (○) radical fluorescence intensity signals (in arbitrary units; Left y axis) as a function of temperature, as well as experimental (●) and predicted (black line; see supplementary material for details) n-butane mole fraction profiles (18) (Right y axis).

Figure 4: Estimated •OH (−) and •HO₂ (○) radical mole fractions obtained from calibrated LIF signals compared with predictions using a detailed kinetic model (18).

Summary

We have presented here direct measurements of •OH and •HO₂ radical profiles under combustion conditions combining the advantages of a radical detection by FAGE to a chemical reactor well adapted for studying low-temperature combustion phenomena, a JSR. Although the absolute concentrations are somewhat uncertain due to calibration and sampling issues, the relative temperature profiles are well defined. Such measurements will certainly lead to a big step forward in refining current models of combustion chemistry, as these radical profiles will be a powerful benchmark for testing models that have so far been developed mostly on the basis of concentration profiles for stable species only.
Acknowledgments

PhysicoChimie des Processus de Combustion et de l’Atmosphère participates in the Institut de Recherche en Environnement Industriel financed by the Région Nord Pas-de-Calais, the Ministère de l’Enseignement Supérieur et de la Recherche, the Centre National de la Recherche Scientifique, and the European Regional Development Fund.

This project was supported by the French Agence Nationale de la Recherche under Contract ANR-11-LabEx-0005-01 “Chemical and Physical Properties of the Atmosphere,” by the European Commission through the “Clean ICE” Advanced Research Grant of the European Research Council, and by European Cooperation in Science and Technology Action CM0901.

Reference