Comparative Shock-Tube Study of Autoignition and Plasma-Assisted Ignition of C₂H₂, C₂H₄, C₂H₆ and C₂H₅OH

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Abstract
The kinetics of ignition in lean and stoichiometric C₂H₂:O₂:Ar, C₂H₄:O₂:Ar, C₂H₆:O₂:Ar and C₂H₅OH:O₂:Ar mixtures was analyzed in experiments using a shock tube with a discharge cell. Ignition delay time was measured by detecting CH radiation behind a reflected shock wave after a high-voltage nanosecond discharge and in its absence. Initiation of the discharge was shown to lead to a noticeable reduction in ignition delay time. Numerical simulation of discharge and ignition phases was used to show the main mechanisms that lead to a diversity of ignition properties of C₂-hydrocarbons.

Introduction
In recent decades particular attention has been drawn to applications of non–equilibrium plasma for plasma–assisted ignition and plasma–assisted combustion (PAI/PAC) [1-3]. It was shown experimentally and numerically that nanosecond discharge plasmas have profound effect on ignition delay reduction and flame stabilization in various combustible mixtures. As a result, fuel could be ignited under conditions when ignition did not occur in the absence of discharge plasma. Recently, plasma-assisted ignition and combustion had been considered as a promising application of low-temperature non-equilibrium plasmas. In particular, ignition by non-equilibrium gas discharges seems to be perspective for a number of applications under conditions of high speed flows and under conditions similar to automotive engines.

It is interesting to compare ignition of hydrocarbons under the same conditions when hydrocarbon molecules contain the same number of C atoms. This has been done for oxidation of C₂H₆, C₂H₄ and C₂H₂ by measuring the induction period in shock waves in thermally-equilibrium regime [4]. In this work, using observations in a shock tube with a discharge cell, measured ignition delay times were compared for stoichiometric and lean C₂H₆-, C₂H₄-, C₂H₂- and C₂H₅OH-containing mixtures both in autoignition regime and ignition with non-equilibrium plasma. Ignition delay was studied after a high-voltage nanosecond discharge and in its absence. We measured ignition delay time in a lean C₂H₆:O₂:Ar mixture and in lean and stoichiometric C₂H₄:O₂:Ar mixtures. The obtained results were compared with our previous studies of ignition of C₂H₆ [5], C₂H₄ [6] and C₂H₅OH [7]. Numerical simulation was used to analyze ignition properties of C₂-hydrocarbons with and without non-equilibrium discharge plasma.

Experiment
Shock tube technique has been widely used to measure autoignition delay times in combustible mixtures [8]. In this work, to show the effect of discharge plasma, ignition in C₂-hydrocarbon–containing mixtures was studied in a shock tube after the discharge and in its absence. Non-equilibrium plasma was created by a high-voltage nanosecond discharge. Gas mixtures under study were heated by a reflected shock wave just before discharge initiation. In this experiment, the discharge processes occurred on a nanosecond scale, whereas ignition developed at times longer than tens microseconds. Chemically active particles were formed in the discharge plasma to favor the ignition of C₂-containing mixtures at gas temperatures above self-ignition threshold.

A detailed description of the experimental setup and methods used to study PAI has been given elsewhere [9-11]. Observations were made in the shock tube (25×25-mm² square cross section) with a 1.6 m long low-pressure channel and a 60 cm long high-pressure chamber. The stainless-steel channel had two pairs of windows for optical diagnostics. In addition, the Plexiglas discharge section of the shock tube had eight optical windows made of quartz or MgF₂. The metal end plate of the tube served as a high – voltage electrode, whereas another electrode was the opposite grounded steel section of the tube.

Ignition was studied in a lean (ϕ = 0.5) C₂H₅OH mixture and in stoichiometric (ϕ = 1) and lean (ϕ = 0.5) C₂H₄O₂ mixtures (10%) diluted with Ar (90%). Ignition delay time was measured behind a reflected shock wave. In this region, the gas temperature was between 1100 and 1700 K and the corresponding pressure was between 0.3 and 0.7 atm. The values of gas pressure and temperature behind a reflected shock wave were obtained from the analysis of the data on initial mixture composition, initial pressure, and measured velocity of the incident shock wave. Thermodynamic data for pure gases required for this analysis were taken from [12].

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Shock wave velocity was measured with three laser Schlieren systems installed along the shock tube. Ignition delay time was defined as the time interval between the Schlieren signal on the reflected shock wave arrival at the diagnostic cross section near the endplate and the onset of the leading front of the CH emission at 431 nm. The emission onset was determined by extrapolating the tangent line to zero emission intensity.

To study the effect of discharge plasma on the ignition, a high-voltage discharge was initiated at the instant when the reflected shock wave reached the diagnostic cross section. The main discharge impulse was followed by one or two additional impulses. These impulses were induced by a partial reflection of the electromagnetic wave from the discharge cell and the high-voltage generator. The total time of plasma generation and decay was much shorter than the delay times for autoignition and PAI. Indeed, the composition of the dominant active species produced in the discharge reached their quasi-steady values in the discharge afterglow in ~ 0.5 µs, whereas the ignition delay time under the conditions studied was in the range 10 - 300 µs. This allowed a separate consideration of excitation kinetics and ignition kinetics.

Fig. 1 compares the new measured data on delay times for autoignition and plasma-assisted ignition in \( \text{C}_2\text{H}_6 \) and \( \text{C}_2\text{H}_4 \)-containing mixtures with our previous measurements for ignition of \( \text{C}_2\text{H}_6 \), \( \text{C}_2\text{H}_5\text{OH} \) and \( \text{C}_2\text{H}_2 \). Additional nonequilibrium excitation leads to a strong reduction in the ignition delay time. The effect is more profound for higher gas temperatures at which the specific deposited discharge energy is higher (see Fig. 2).

![Fig. 1. Comparison of measured induction delay times for \( \text{C}_2 \)-hydrocarbons.](image)

The data shown in figure 1 were obtained using the same setup and for the same range of gas parameters. Therefore, this comparison shows peculiarities for autoignition and plasma-assisted ignition in the \( \text{C}_2 \)-containing mixtures under study. The data obtained for
C$_2$H$_6$, C$_2$H$_4$ and C$_2$H$_5$OH are close to each other, whereas the delay time for autoignition and plasma-assisted ignition in C$_2$H$_2$ that is highly reactive are much shorter. Figure 3 shows the effect of equivalence ratio on the measured ignition delay times. The autoignition delay time is independent of the equivalence ratio in the C$_2$H$_6$- and C$_2$H$_4$-containing mixtures, whereas this time in the C$_2$H$_5$OH- and C$_2$H$_2$-containing mixtures is shorter for the stoichiometric mixtures than that for the lean mixtures. In plasma-assisted ignition, the delay time is approximately the same for $\phi = 1$ and 0.5 in the C$_2$H$_4$ and C$_2$H$_2$ mixtures; in contrast, the induction times in the lean C$_2$H$_6$- and C$_2$H$_5$OH-containing mixtures are shorter than those in the stoichiometric mixtures.

Numerical simulation
To show the main mechanisms that lead to a diversity of autoignition and plasma-assisted ignition times for C$_2$-hydrocarbons, ignition in the mixtures under consideration was numerically simulated with and without the additional excitation by the discharge. We studied the temporal evolution of the mole fractions of chemically active species in the discharge, in its afterglow and in the ignition phase. The method and kinetic schemes used to simulate the discharge phase and discharge afterglow have been described in detail elsewhere [5,11]. Ignition processes were simulated on the basis of the kinetic scheme for autoignition [13]. Ignition delay time was calculated from the analysis of the time evolution of CH mole fraction in the ignition phase. This approach allowed good agreement between our previous measurements and calculations for autoignition and plasma-assisted ignition of CH$_4$-C$_3$H$_8$ [5,11], C$_2$H$_4$ [6] and C$_2$H$_5$OH [7].

Fig. 4 compares ignition delay times calculated in this work for autoignition and plasma-assisted ignition with the measured data in C$_2$H$_4$-containing mixtures. Agreement between the calculated and measured results for autoignition and plasma-assisted ignition is good; it is better for the lean mixture and for lower gas temperatures when the specific deposited energy per one neutral particle is lower.

To demonstrate the difference in dominant ignition processes for various C$_2$-hydrocarbons, we made a sensitivity analysis for the autoignition and plasma-assisted ignition in the stoichiometric C$_2$H$_4$:O$_2$:Ar, C$_2$H$_2$:O$_2$:Ar, C$_2$H$_6$:O$_2$:Ar and C$_2$H$_5$OH:O$_2$:Ar mixtures at the same values of initial gas temperature and pressure. For this purpose the rate coefficients of every pair of forward and reverse reactions were in turn increased by 50% and the ignition delay time was calculated in each case. The sensitivity coefficients, $(\tau_i - \tau_0)/\tau_0$, were again calculated, where $\tau_0$ is the ignition delay time calculated with unchanged rate constants and $\tau_i$ is the ignition delay time calculated with the increased rate constants for the i-th pair of forward and reverse reactions.

We analyzed the calculated results for the reactions when the magnitude of the ratio $(\tau_i - \tau_0)/\tau_0$ was higher than 0.05 at least for autoignition or ignition by the discharge. The sensitivity analysis showed the most limiting reactions and led to the following conclusions.
For all C₂-hydrocarbons, the H + O₂ ↔ O + OH reaction always exhibits the greatest sensitivity (20-40% decrease in the ignition delay time) both for autoignition and for ignition after the discharge. This chain-branching reaction leads to an increase in the amount of radicals and is generally important in ignition of different fuels at T > 1100 K.

Ignition of C₂H₂ and C₂H₆ was controlled only by three and six chemical reactions, respectively, that significantly affected the ignition delay. The number of reactions that greatly affect ignition of C₂H₂ and C₂H₆OH is several times higher. All dominant reactions for the ignition of C₂H₂ lead to a reduction in the ignition delay time when increasing their rates. For other C₂-hydrocarbons under study, many reactions lead to a reduction in the ignition delay time when increasing their rates, whereas some other reactions have an opposite effect on ignition.

Conclusions

Using a shock tube with a discharge cell, ignition delay time was measured in a lean (φ = 0.5) C₂H₆:O₂:Ar mixture and in lean (φ = 0.5) and stoichiometric C₂H₅OH:Ar mixtures with a high-voltage nanosecond discharge and without it. The measured results were compared with the measurements made previously with the same setup for C₂H₅-, C₂H₅OH- and C₂H₂-containing mixtures. It was shown that the effect of plasma on ignition is almost the same for C₂H₆, C₂H₅, and C₂H₅OH. The reduction in time is smaller for C₂H₅OH, the fuel that is well ignited even without the discharge. Autoignition delay time was independent of the stoichiometric ratio for C₂H₂ and C₂H₆, whereas this time in stoichiometric C₂H₂- and C₂H₅OH-containing mixtures was noticeably shorter than that in the lean mixtures. Ignition after the discharge was not affected by a change in the stoichiometric ratio for C₂H₂ and C₂H₆, whereas the plasma-assisted ignition delay time for C₂H₆ and C₂H₅OH decreased as the equivalence ratio changed from 1 to 0.5.

Ignition delay time was calculated in C₂-hydrocarbon-containing mixtures under study by simulating separately discharge and ignition processes. Good agreement was obtained between new measurements and calculated ignition delay times. Sensitivity analysis was used to show the peculiarities of ignition with and without the discharge for C₂-hydrocarbons under study.

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References

