Ignition Delay Characteristics of Kerosene with Decomposed Hydrogen Peroxide

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The ignition chemical delay of kerosene with decomposed hydrogen peroxide was simulated by using n-decane as a surrogate fuel for kerosene. The influences of H₂O₂ concentration, the H₂O₂ decomposition rate, and pressure on the chemical delay were investigated. The ignition delay time had the negative temperature coefficient phenomenon for a low H₂O₂ decomposition rate, and the increasing pressure weakened the negative temperature coefficient phenomenon, which nearly disappeared when the pressure was greater than 4 MPa. An R_H²O₂-dependent coefficient of the ignition delay time was almost not influenced by H₂O₂ concentration and pressure when the pressure was less than 4 MPa, and R_H²O₂ had a noticeable influence on the ignition delay time at R_H²O₂ < 80%. Increasing the H₂O₂ concentration effectively reduced the ignition delay time at R_H²O₂ > 70%, but it had a slight effect on the ignition delay time at R_H²O₂ > 70%.

Nomenclature

<table>
<thead>
<tr>
<th>Symbol</th>
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<tbody>
<tr>
<td>C_H²O₂</td>
<td>H₂O₂ concentration, %</td>
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<td>P</td>
<td>pressure, MPa</td>
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<td>R_H²O₂</td>
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<td>τ₁</td>
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<td>first-stage ignition delay time</td>
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<td>τ₂</td>
<td>second-stage ignition delay time</td>
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<td>φ</td>
<td>equivalence ratio</td>
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I. Introduction

In recent years, to enhance safety and reduce the toxicity of liquid rocket engines, the green propellants have received wide attention, and the combination of kerosene with H₂O₂ is promising due to its high-density specific impulse, long-term storage capability, and low toxicity [1,2]. Many studies are focusing on developing H₂O₂/kerosene bipropellant liquid engines [3].

Rocket-grade hydrogen peroxide (RGHP) is characterized by a high mass ratio of hydrogen peroxide to water (usually greater than 85%) [4]. In a rocket engine, RGHP is passed through a catalyst bed (silver screens or other catalytic material) and aft-mounted injector, and the catalyst bed initiates RGHP decomposition [5–7]. Assuming that 90% H₂O₂ is decomposed, the decomposition reaction for this concentration can be written as

$$\text{H}_2\text{O}_2(l) + 0.21\text{H}_2\text{O}(l) \Rightarrow 1.21\text{H}_2\text{O}(g) + 0.5\text{O}_2(g) \quad (1)$$

The RGHP decomposition mixture of high-temperature steam and oxygen leads to the automatic ignition of kerosene in a rocket engine, and the ignition delay is a critical kinetic parameter [3,7]. If the ignition does not occur on time, propellants can accumulate in the chamber and then spontaneously ignite, resulting in the chamber bursting and combustion instability, which leads to rocket engine failure [8]. The ignition delay of hypergolic propellants depends on the complex interactions between many chemical and physical factors, the process of ignition delay includes the physical delay (atomization, mixing, vaporization, and diffusion) and the chemical delay (gas phase reactions), and the characteristic times of each process combine to form a global ignition delay time [9–12]. The physical delay is greatly influenced by the injection technique, which varies with the injector configuration and injecting parameters, and the chemical delay is determined by the reaction potential of the two propellants (such as fuel structure, mixture composition, the fuel/air ratio, temperature, and pressure) [13].

There is relatively little information about the ignition delay characteristics of kerosene with the decomposed hydrogen peroxide. In this study, the ignition physical delay is not considered; the ignition chemical delay of kerosene with decomposed hydrogen peroxide is simulated by using n-decane as a surrogate fuel for kerosene; and the influences of C_H²O₂, R_H²O₂, and pressure on the chemical delay are investigated in a perfectly stirred reactor model.

II. Methodology

A. Reaction Mechanism for n-Decane

The commercial Jet-A1 is kerosene-type fuel, it is complex mixtures of a large number of different hydrocarbons, and the developing chemical kinetic models of kerosene combustion is of practical importance. Because of the very large chemical kinetic mechanisms and the associated computational resources required, the simulation of kerosene combustion is challenging [14]. Zeng et al. [15] and Dagaut and Cathonnet [16] used a surrogate fuel containing only n-decane to simulate kerosene combustion at 1–4 MPa. Chang et al. [17] developed a new skeletal mechanism including 40 species and 141 reactions for n-decane oxidation to simulate kerosene combustion, the mechanism for C₈–C₁₀ was used to model the oxidation of heavy hydrocarbons for the prediction of the ignition characteristics, and it was extensively validated by comparing it to the experimental ignition delay data of n-decane oxidation under the low-to-high temperature and high-pressure conditions [18]. In the study, the skeletal mechanism developed by Chang et al. is used to simulate the ignition delay characteristics of kerosene with the decomposed hydrogen peroxide.

B. Homogeneous Reactor

To understand the ignition delay mechanism of kerosene with H₂O₂ decomposition gas in a rocket engine, the homogeneous, isobaric, and adiabatic combustion reactor model is employed to predict the ignition time, and it can qualitatively discover the effects of operating parameters on the ignition delay process.

The chemical kinetic behavior of reactants is modeled by Sandia National Laboratory’s SENKIN program of CHEMKIN subroutines.
The program solves the conservation equations for mass and energy, it can calculate the temporal evolution of mole fractions of species for a homogeneous mixture in a closed reactor, and it can account for finite-rate elementary chemical reactions. Therefore, the SENKIN model is appropriate to investigate the ignition chemical delay of kerosene with H$_2$O$_2$ decomposition gas in a closed chamber. In this reactor model, the reactive system is defined by specifying initial values of volume, temperature, pressure, and chemical compositions. Based on a steepest-increase criterion of the temperature, $\tau_{ig}$ is extracted. The reaction pathway flux analysis is performed using MixMaster (a Python program that is part of the Cantera suite) based on a conserved scalar of reaction fluxes [18].

### C. Simulation Conditions

Using a thermodynamic equilibrium model based on the minimization of Gibbs free energy, the dominant products of H$_2$O$_2$ decomposition are H$_2$O and O$_2$, the mass concentration sum of other productions is less than $10^{-5}$ in the operating conditions of catalyst beds using silver screens, and thus it is considered that the decomposition products are only H$_2$O and O$_2$. The effect of pressure on the H$_2$O$_2$ decomposition temperature was reported by General Kinetics [21]. According to the real gas specific heat and standard formation enthalpy of H$_2$O, O$_2$, and H$_2$O, the decomposition temperature is calculated [21,22], and Ge et al. [22] provided the calculation method and formula. Figure 1 shows the decomposition temperature of 87–92% H$_2$O$_2$ at 1 MPa.

For a given C$_{H2O2}$, when H$_2$O$_2$ is completely decomposed in the catalyst bed, the mass fractions of O$_2$ and H$_2$O in the high-decomposition gas mixture are

$$f_{O2} = 0.47037C_{H2O2}/100$$  \hspace{1cm} (2)

$$f_{H2O} = 1 - 0.47037C_{H2O2}/100$$  \hspace{1cm} (3)

To avoid catalyst bed overtemperature or the instability in the practical operation of the rocket engine, H$_2$O$_2$ is partly decomposed in the catalyst bed, and the residual H$_2$O$_2$ in the supercritical liquid state is injected into combustion chamber, which has no sufficient time to participate in the ignition due to the delay of the vaporization and decomposition process. Thus, it is necessary to investigate the effect of R$_{H2O2}$ on the ignition delay of kerosene with the H$_2$O$_2$ decomposition gas. At given C$_{H2O2}$ and R$_{H2O2}$, the mass fractions of O$_2$, H$_2$O, and the residual H$_2$O$_2$ in the high-decomposition mixture are

$$f_{O2} = 0.47037 \times (C_{H2O2}/100) \times (R_{H2O2}/100)$$  \hspace{1cm} (4)

$$f_{H2O} = (1 - 0.52963 \times R_{H2O2}/100) \times C_{H2O2}/100$$  \hspace{1cm} (5)

$$f_{H2O2} = (1 - R_{H2O2}/100) \times C_{H2O2}/100$$  \hspace{1cm} (6)

In the ignition simulation of kerosene with decomposed hydrogen peroxide at a given C$_{H2O2}$, R$_{H2O2}$ influences the initial ignition temperature and $\varphi$, and $\varphi$ is defined as

$$\varphi = \frac{(m_{C10H22}/m_{H2O2})_{actual mixture}}{(m_{C10H22}/m_{H2O2})_{stoichiometric mixture}}$$  \hspace{1cm} (7)

where $m_{C10H22}$ and $m_{H2O2}$ are the mass contents of C$_{10}$H$_{22}$ and H$_2$O$_2$, respectively. In the study, m$_{C10H22}$ is kept at a constant, $\varphi$ is kept at a constant of 1.0 when H$_2$O$_2$ is completely decomposed (R$_{H2O2}$ = 100%), the actual amount of m$_{H2O2}$ participating in the ignition reactions is equal to the product of stoichiometric H$_2$O$_2$ mass and R$_{H2O2}/100$, and then $\varphi$ is derived from Eq. (7):

$$\varphi = 100/R_{H2O2}$$  \hspace{1cm} (8)

As known from Eq. (8), decreasing R$_{H2O2}$ means that the equivalence ratio decreases, and the ignition is under a fuel-rich atmosphere at $\varphi < 1.0$. In the study, the H$_2$O$_2$ concentration and decomposition rate vary from 87 to 92% and 60 to 100%, respectively.

### III. Results and Discussion

#### A. Ignition Process Analysis

In the study, a high-temperature gas stream of the decomposed hydrogen peroxide mixes with n-decane, and the autoignition can be achieved. Ignition is an oxidizing fuel and increasing temperature process. Figure 2 shows the temperature history of the mixture as it achieved. Ignition is an oxidizing fuel and increasing temperature process. Figure 2 shows the temperature variations in the ignition process at 1 MPa and 87% C$_{H2O2}$.

In the study, a high-temperature gas stream of the decomposed H$_2$O$_2$ is injected into the combustion chamber, which has no sufficient time to participate in the ignition due to the delay of the vaporization and decomposition process. Thus, it is necessary to investigate the effect of R$_{H2O2}$ on the ignition delay of kerosene with the high-temperature gas. At given C$_{H2O2}$ and R$_{H2O2}$, the mass fractions of O$_2$, H$_2$O, and the residual H$_2$O$_2$ in the high-decomposition mixture are

The behavior of two-stage ignition for low R$_{H2O2}$ in Fig. 2 is further demonstrated in Fig. 3. With the increase of R$_{H2O2}$, $\tau_1$ monotonically decreases, and $\tau_1$ is nearly equal to 0 ms at R$_{H2O2}$ > 88%; $\tau_2$ first increases at R$_{H2O2}$ < 88% and then decreases; it indicates that the ignition delay is two stages at R$_{H2O2}$ < 88% and a single stage at R$_{H2O2}$ > 88%; the total ignition delay time $\tau_1$ is the sum of $\tau_1$ and $\tau_2$, and the ignition delay mainly results from the first-stage delay at R$_{H2O2}$ < 68%.

#### Fig. 1 Effect of R$_{H2O2}$ on decomposition temperature at 1 MPa.

#### Fig. 2 Temperature variations in the ignition process at 1 MPa and 87% C$_{H2O2}$. 

\[
f_{H2O} = (1 - 0.52963 \times R_{H2O2}/100) \times C_{H2O2}/100
\]

\[
f_{H2O2} = (1 - R_{H2O2}/100) \times C_{H2O2}/100
\]

\[
\varphi = \frac{(m_{C10H22}/m_{H2O2})_{actual mixture}}{(m_{C10H22}/m_{H2O2})_{stoichiometric mixture}}
\]

\[
\varphi = 100/R_{H2O2}
\]
Figure 3, with the increase of $R_{H2O2}$, $\tau_i$ first decreases at $R_{H2O2} < 72\%$ and then increases at $R_{H2O2} = 72-88\%$. As is known from Fig. 1, the temperature of the $H_2O_2$ decomposition gas increases with the increase of $R_{H2O2}$, and thus there is a negative temperature coefficient (NTC) phenomenon in the C-D region where $\tau_i$ increases with the increase of the initial ignition temperature. Practically, the NTC phenomenon is of particular relevance to the intrinsically low-temperature phenomenon of ignition for the low-temperature oxidation of large hydrocarbons [23].

As is known from Figs. 2 and 3, the ignition process goes through two stages to achieve intense burning at $R_{H2O2} < 88\%$, but it goes through only one stage, controlled by low-temperature chemistry, at $R_{H2O2} > 88\%$. To understand the dominant chemical reaction mechanism of ignition at the points of A and B in Figs. 2 and 4 provides the species concentration variations with time in the ignition process for $C10H2O2 = 87\%$ and $R_{H2O2} = 64\%$ at 1 MPa. In Figs. 4a and 4b, with the increase of time, these species except $C10H22$ gradually increase, there is a single peak of point A (at 1941 $\mu$s) where hydrocarbon species sharply decrease (see Figs. 4a and 4b) and the temperature rapidly increases to 822 K at point A (see Fig. 2), and thus the first-stage ignition occurs at point A. In the first-stage ignition, $C10H22$ is decomposed and oxidized with the slow increase of the temperature, the temperature increases from 722 to 822 K, and thus the first-stage ignition is governed by the low-temperature chemistry. At point B in Fig. 2 (at 2606 $\mu$s), the concentration of hydrocarbon species dramatically decreases in Figs. 4c and 4d, the temperature sharply increases to 1828 K, and thus the secondary-stage ignition occurs at point B in Fig. 2, which is governed by the high-temperature chemistry.

To analyze the reaction mechanisms of the low-temperature chemistry and the high-temperature chemistry in the ignition process, the reaction paths are investigated at 1930 and 2548 $\mu$s, as shown in Fig. 5.

Figure 5 shows the low-temperature reaction path of the first-stage ignition at 1941 $\mu$s, and the low-temperature chain sequence is as follows. First, H is abstracted from $C10H22$ by OH and H radicals to form the heptyl radicals ($C10H21$):

$$C10H22 + H \rightarrow C10H21 + H_2$$ (9)

$$C10H22 + OH \rightarrow C10H21 + H_2O$$ (10)

H and OH radicals attack $C10H22$ to result in H abstraction, and the major consuming fuel paths are exothermic reactions.

Second, the heptyl radical ($C10H21$) combines with $O_2$ to form the heptylperoxy radical ($C10H22O2$):

$$C10H21 + O_2 \rightarrow C10H22O2$$ (11)

11 is one of the major heat release reactions in the low-temperature reaction of the first-stage ignition; 11 can process toward the inverse direction with the increase of temperature, and then it has an inverse temperature dependence.

Third, the isomerization of heptylperoxy ($C10H22O2$) forms the heptylhydroperoxy radicals ($C10H22OOH$):

$$C10H21OO \rightarrow C10H22OOH$$ (12)

12 is an endothermic reaction.

Fourth, the $C10H22OOH$ proceeds to take the chain branching reaction with the second $O_2$ addition to form $OOC10H22OOH$:

$$C10H22OOH + O_2 \rightarrow OOC10H22OOH$$ (13)

$OOC10H22OOH$ then forms ketohydroperoxide ($C10$ ket) by releasing $OH$:

$$OOC10H22OOH \rightarrow C10 ket + OH$$ (14)

Moreover, the ketohydroperoxide molecule undergoes decomposition to form $C3H7CO$, olefin, and formaldehyde in reaction:

$$C10 ket \rightarrow CH2O + C3H7CO + OH + C3H6$$ (15)

The ketones ($C10$ ket) decomposition reaction of 15 is the most important low-temperature endothermic chain branching reaction. The $C3H7CO$ continues to decompose into $C0 - C3$ species:

$$C3H7CO + O_2 \rightarrow C3H7 + C2H5 + CO + HO_2$$ (16)

Finally, the carbonyl decomposition reactions generate smaller $C1$ and $C2$ radicals, which subsequently undergo further reactions:

$$C2H + O_2 \rightarrow CH2O + HCO$$ (17)

Specifically, it is noted that the isomerization and decomposition reactions have the dominant effect on the first ignition stage.

In the previous reactions, 10, 11, 13, 14, and 17 are important exothermic reactions, the concentration of OH radical originating from 14 and 15 increases with the increase of time at 1800-2020 $\mu$s, and this accelerates the reaction rate of 17 and the rise of temperature until the temperature rises to a value at which the 16 reaction becomes the important reaction:

$$C10H22 + O_2 \rightarrow C10OH20 + HO_2$$ (18)

$H_2O_2$ is formed:

$$HO_2 + HO_2 \rightarrow H_2O_2 + O_2$$ (19)

In the first-stage ignition process, the OH radical is largely consumed in the range of 1940 to 1960 $\mu$s, the temperature noticeably rises (see Fig. 2), 11 shifts toward the decomposition of the $C10H22OO$ radical, the production rate of OH decreases, and then the low-temperature chemistry of the first-stage ignition is shut down [23].

Since the reaction rates of 18 and 19 increase with the increase of temperature in the first-stage ignition, 18 competes with the reactions of 11, 13, 14, and 17; this suppresses OH radical formation and results in the decrease of the 10 reaction rate, and then 18 reaction becomes important, terminating the first-stage ignition when the temperature rises to 822 K (see point A in Figs. 2, 4a, and 4b).

In the second-ignition stage, initially, 18 becomes a dominant path reaction involving slightly exothermic process, it suppresses the formation of the OH radical, OH concentration begins to decrease after the first-stage ignition, the 10 reaction rate decreases, this results in the decrease of the $C10H21$ concentration (see Figs. 4a and 4b), and the temperature slowly rises (see Fig. 2) until 20 becomes an important reaction:
H₂O₂(+)M → OH + OH(+)M \hspace{2cm} (20)

20 accelerates H abstraction from C₁₀H₂₂ by 10 and promotes reaction, and the temperature rapidly rises. At high temperature, the pyrolysis reaction becomes important:

C₁₀H₂₁ → 2C₃H₆ + C₂H₅ + C₂H₄ \hspace{2cm} (21)

As is known from Fig. 5 at 2606 \(\mu s\), C₁ – C₃ hydrocarbons are rapidly oxidized to CO₂ by O₂ and the free radicals of HO₂ and OH. With the acceleration of the oxidation reactions, the temperature sharply rises to 1828 K at 2606 \(\mu s\) (see point B in Fig. 2), and the second-stage ignition process is terminated.

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B. Effects of Pressure, H₂O₂ Decomposition Rate, and H₂O₂ Concentration on Ignition Delay

In the ignition simulation of H₂O₂/kerosene, at a given C₁₀H₂₂, R₃H₂O₂ influences the initial ignition temperature and \(\phi\). The temperature of the decomposition gas mixture increases with the increase of R₃H₂O₂, C₁₀H₂₂, and pressure \([21,22]\). As is known from Eq. (8), \(\phi\) decreases in the ignition stage with the increase of R₃H₂O₂.

Figure 6 shows the effects of pressure, C₁₀H₂₂, and R₃H₂O₂ on the ignition delay time. With the increase of pressure, the ignition delay time rapidly decreases in the range from 1 to 4 MPa (see Figs. 6a and 6c) and slowly decreases at \(P > 4\) MPa (see Figs. 6b and 6d), and this phenomenon is directly related to the oxidation of the alkyl radicals (see 11). With the increase of pressure, the equilibrium of 11 shifts toward the formation of the C₁₀H₂₂OO radical, the reaction releases a large amount of heat in the low-temperature reaction of the first-stage...
ignition, and this effectively reduces \( \tau_1 \) when the pressure increases from 1 to 4 MPa. Since the \( \text{H}_2\text{O}_2 \) decomposition temperature increases with pressure, the ignition time decreases with the further increase of pressure from 4 to 9 MPa, but \( \text{H}_2\text{O}_2 \) process toward the inverse direction with the increase of temperature, and thus the variation of the ignition delay time with the increase of pressure becomes slow (see Figs. 6b and 6d).

Since the low-temperature chain reactions produce a large amount of \( \text{H}_2\text{O}_2 \) by \( 19 \), \( \text{H}_2\text{O}_2 \) decomposition leads to the termination of the NTC process through \( 21 \), which is pressure-dependent reaction, and thus the \( \text{H}_2\text{O}_2 \) decomposition rate is greater at higher pressures. Therefore, increasing the pressure makes the NTC region shift to a higher \( R_{\text{H}_2\text{O}_2} \) range, and the corresponding NTC region becomes narrow, as shown in Fig. 7.

To investigate the influencing extent of \( \text{C}_\text{H}_2\text{O}_2 \) and pressure on the NCT region, the difference of \( \tau_{\text{min}} \) and \( \tau_{\text{max}} \) is defined as

\[
\Delta \tau = \tau_{\text{max}} - \tau_{\text{min}}
\]  

(22)

where \( \tau_{\text{min}} \) and \( \tau_{\text{max}} \) are, respectively, the minimal and maximal ignition delay times corresponding points C and D of the NCT region in Fig. 3. If the value of \( \Delta \tau \) is great, the variation extent of the NCT region is noticeable. Figure 8 shows the influencing extent of \( \text{C}_\text{H}_2\text{O}_2 \) and pressure on the NCT region corresponding to Fig. 7; the increase of pressure weakens the variation extent of the NTC. As is known from Fig. 6, at \( P > 4 \) MPa, the NTC phenomenon of ignition nearly disappears, and the ignition process goes through only one stage governed by low-temperature chemistry [23].

In the study, when \( \text{H}_2\text{O}_2 \) is completely decomposed by catalysis (\( R_{\text{H}_2\text{O}_2} = 1 \)), the equivalence ratio is kept at 1. If \( \text{H}_2\text{O}_2 \) is partly decomposed by catalysis, the equivalence ratio decreases with the decrease of \( R_{\text{H}_2\text{O}_2} \), and the decomposition temperature decreases. In Fig. 6, \( R_{\text{H}_2\text{O}_2} \) has complex effects on the ignition delay time: at low \( R_{\text{H}_2\text{O}_2} \), increasing \( R_{\text{H}_2\text{O}_2} \) makes \( \tau_{\text{ig}} \) rapidly decrease; in the NTC region, \( \tau_{\text{ig}} \) increases with \( R_{\text{H}_2\text{O}_2} \); and at high pressure, \( \tau_{\text{ig}} \) monotonically decreases with the increase of \( R_{\text{H}_2\text{O}_2} \).

To further investigate the effect of \( R_{\text{H}_2\text{O}_2} \) on \( \tau_{\text{ig}} \), the \( R_{\text{H}_2\text{O}_2} \) dependent coefficient of the ignition delay time is defined as

\[
\zeta = \frac{(\tau_{i+1} - \tau_i)/\tau_i}{\Delta R/R_1}
\]  

(23)
where $R_1$ and $\tau_1$, respectively, denote 60% $R_{\text{H}_2\text{O}_2}$ and the corresponding ignition delay time; $\Delta R$ is the increment interval of $R_{\text{H}_2\text{O}_2}$; and $\tau_i$ and $\tau_{i-1}$ are, respectively, the ignition delay times at $R_{\text{H}_2\text{O}_2} = (60 + i\Delta R)$ and $R_{\text{H}_2\text{O}_2} = 60 + ((i + 1)\Delta R)$. As is known from Eq. 23, if $\zeta$ is negative, this means that the ignition delay time decreases with the increase of $R_{\text{H}_2\text{O}_2}$.

Figure 9 presents $R_{\text{H}_2\text{O}_2}$-dependent coefficients of the ignition delay time at different $R_{\text{H}_2\text{O}_2}$; the coefficients are almost similar when pressure is less than 4 MPa, and thus it can be considered that $\zeta$ is almost only dependent on $R_{\text{H}_2\text{O}_2}$ but independent of $C_{\text{H}_2\text{O}_2}$ and pressure. At $R_{\text{H}_2\text{O}_2} < 80\%$, $\zeta$ rapidly varies with $R_{\text{H}_2\text{O}_2}$, and it indicates $R_{\text{H}_2\text{O}_2}$ has a great influence on $\tau_{i-1}$. At $R_{\text{H}_2\text{O}_2} > 80\%$, $\zeta$ slowly varies with $R_{\text{H}_2\text{O}_2}$, and it indicates that the influence of $R_{\text{H}_2\text{O}_2}$ on $\tau_{i-1}$ decreases. Since the initial ignition temperature and the equivalence ratio are low at $R_{\text{H}_2\text{O}_2} < 80\%$, increasing $R_{\text{H}_2\text{O}_2}$ can effectively reduce the first-stage ignition delay time (see Fig. 4), the ignition delay time rapidly decreases with the increase of $R_{\text{H}_2\text{O}_2}$, and the $R_{\text{H}_2\text{O}_2}$-dependent coefficient of the ignition delay time rapidly increases (see Fig. 9).

In Fig. 9, the range of positive value $\zeta$ is the corresponding NTC region. Since the temperature of the $\text{H}_2\text{O}_2$ decomposition gas increases with $R_{\text{H}_2\text{O}_2}$, the initial ignition temperature also rises with $R_{\text{H}_2\text{O}_2}$ (see Fig. 1); this can effectively reduce the first-stage ignition time, but the released heat at the end of the first-ignition stage decreases, which leads to a longer delay time for the second-stage ignition, and consequently the entire ignition delay time becomes longer with $R_{\text{H}_2\text{O}_2}$ in the NTC region, as shown in Fig. 3.
Figure 10 shows the effect of $\text{H}_2\text{O}_2$ concentration on the ignition delay time, and it indicates that at $R_{\text{H}_2\text{O}_2} < 70\%$ the increase of the $\text{H}_2\text{O}_2$ concentration can effectively reduce ignition delay time and at $R_{\text{H}_2\text{O}_2} > 70\%$ the $\text{H}_2\text{O}_2$ concentration has a slight effect on the ignition delay time. In the study, the increasing $\text{H}_2\text{O}_2$ concentration means the increase of the initial ignition temperature and $\text{O}_2$ concentration in the $\text{H}_2\text{O}_2$ decomposition mixture gas, and thus the ignition delay rapidly decreases with the $C_{\text{H}_2\text{O}_2}$ at $R_{\text{H}_2\text{O}_2} < 70\%$.

**IV. Conclusions**

In this study, fuel $n$-decane is used as a surrogate fuel for kerosene; the effects of $C_{\text{H}_2\text{O}_2}$, $R_{\text{H}_2\text{O}_2}$, and pressure on the ignition chemical delay are investigated; and the reaction paths of the ignition process are provided.

The ignition delay time has the negative temperature coefficient (NTC) phenomenon for a low $\text{H}_2\text{O}_2$ decomposition rate, $\tau_{ig}$ increases with $R_{\text{H}_2\text{O}_2}$ in the NCT region, the increasing pressure makes the NTC region shift to a higher $R_{\text{H}_2\text{O}_2}$ range and weakens the variation extent of the NTC, the corresponding NTC region becomes narrow, and the NTC phenomenon of ignition delay nearly disappears when pressure is greater than 4 MPa. The $R_{\text{H}_2\text{O}_2}$-dependent coefficient of the ignition delay time is almost independent of $C_{\text{H}_2\text{O}_2}$ and the pressure when the pressure is less than 4 MPa, and $R_{\text{H}_2\text{O}_2}$ has a great influence on $\tau_{ig}$ at $R_{\text{H}_2\text{O}_2} < 80\%$. Increasing the $\text{H}_2\text{O}_2$ concentration can effectively reduce the ignition delay time at $R_{\text{H}_2\text{O}_2} < 70\%$, and the $\text{H}_2\text{O}_2$ concentration has a slight effect on the ignition delay time at $R_{\text{H}_2\text{O}_2} > 70\%$.

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**References**
