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**RESEARCH ARTICLE** 

# $\begin{array}{c} \text{IO} \rightarrow \text{C} \\ \text{IN} & \text{RG} \\ \text{CHEMICAL KINETICS} \\ \text{OH} \rightarrow \text{IN} \\ \text{OH} \end{array} \\ \end{array}$

# Role of methyldioxy radical chemistry in high-pressure methane combustion in $CO_2$

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#### Abstract

The combustion chambers of direct-fired supercritical CO<sub>2</sub> power plants operate at pressures of approximately 300 bar and CO<sub>2</sub> dilutions of up to 96%. The rate coefficients used in existing chemical kinetic mechanisms are validated for much lower pressures and much smaller concentrations of CO<sub>2</sub>. Recently, the UoS sCO2 1.0 and UoS sCO2 2.0 mechanisms have been developed to better predict ignition delay time (IDT) data from shock tube studies at pressures from 1 to 260 bar in various CO<sub>2</sub>-containing bath gas compositions. The chemistry of the methyldioxy radical (CH<sub>3</sub>O<sub>2</sub>) has been identified as an essential combustion intermediate for methane combustion above 100 bar, where mechanisms missing this species begin to vastly overpredict the IDT. The current literature available on CH<sub>3</sub>O<sub>2</sub> is very limited and often concerned with atmospheric chemistry and low-pressure, low-temperature combustion. This means that the rate coefficients used in UoS sCO<sub>2</sub> 2.0 are commonly determined at sub-atmospheric pressures and temperatures below 1000 K with some rate coefficients being over 30 years old. In this work, the rate coefficients of new potential CH<sub>3</sub>O<sub>2</sub> reactions are added to the current mechanism to create UoS  $sCO_2$  2.1 It is shown that the influence of CH<sub>3</sub>O<sub>2</sub> on the IDT is greatest at high pressures and low temperatures. It has also been demonstrated that  $CO_2$  has very little effect on the chemistry of  $CH_3O_2$ at 300 bar meaning that CH<sub>3</sub>O<sub>2</sub> rate coefficients can be determined in other bath gases, reducing the impact of non-ideal effects such as bifurcation when studying in a CO<sub>2</sub> bath gas. The updated UoS sCO<sub>2</sub> 2.1 mechanism is then compared to high-pressure IDT data and the most important reactions which require reinvestigation have been identified as the essential next steps in understanding high-pressure methane combustion.

#### KEYWORDS

CH<sub>3</sub>O<sub>2</sub>, chemical kinetics, methane, supercritical CO<sub>2</sub>

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## 1 | INTRODUCTION

In recent years, the energy trilemma has become increasingly challenging on all three fronts. On the fuel security and equity fronts, the recent war in Ukraine has interrupted the supply of Russian gas to European countries and inflated oil and gas prices, forcing major European countries to find alternate suppliers, namely the United States and the Middle East.<sup>1,2</sup> Additionally, on the environmental security front, the 2019 "The Truth Behind the Climate Pledges" report found that 75% of the 184 report countries that signed up to the Paris Agreement had taken insufficient action to curb global temperature rises, with some still increasing their emissions.<sup>3</sup> Bringing together all these challenges identifies a need for energy technologies that provide for emissionsfree energy generation from the combustion of fossil fuels.

Direct-fired supercritical  $CO_2$  (s $CO_2$ ) power cycles such as the Allam-Fetvedt cycle<sup>4</sup> and the Supercritical Transformational Electric Power (STEP) project<sup>5</sup> have been proposed as methods of emissions-free power generation from fossil fuels. The Allam-Fetvedt cycle, the most developed of the two, burns a mixture of methane and oxygen in a 96% CO<sub>2</sub> dilution at pressures up to 300 bar.<sup>4</sup> Due to the combustion taking place in pure oxygen and  $CO_2$ , the only major products are  $CO_2$  and water (H<sub>2</sub>O), which can be easily separated to create a high-purity stream of  $sCO_2$ , ideal for transportation and subsequent storage or utilization. This means that Allam-Fetvedt cycle power plants have 100% carbon capture, at a price competitive with existing fossil-fueled power plants without carbon capture and storage. This is due to the greater power density of sCO<sub>2</sub> compared to traditional working fluids, and the high pressure reduces the overall plant footprint.<sup>4</sup> The deployment of the Allam-Fetvedt cycle worldwide has begun with the recent announcement of two new power plants in development in the United States.<sup>6</sup> and one in the United Kingdom.<sup>7</sup>

Most chemical kinetic combustion mechanisms have been developed using experimental data recorded at pressures and  $CO_2$  dilutions much smaller than those of the Allam-Fetvedt cycle.<sup>8</sup> Recently, Harman-Thomas et al.<sup>9</sup> studied the performance of four existing chemical kinetic mechanisms for modeling ignition delay time (IDT) of 52 datasets of methane, hydrogen, and syngas combustion at a range of equivalence ratios, pressures, temperatures, and fuel ratios in various concentrations of  $CO_2$ . From this work and Harman-Thomas et al.,<sup>10,11</sup> the University of Sheffield (UoS) s $CO_2$  2.0 mechanism was created and verified for methane, hydrogen, and syngas IDTs. Karimi et al.<sup>12</sup> first noted the importance of  $CH_3O_2$  chemistry in  $CO_2$  at 100 bar and 200 bar and this chemistry was subsequently found to the essential to the creation of UoS s $CO_2$  1.0. The rate coefficients for the reactions of  $CH_3O_2$  are often poorly understood with rate coefficients from outdated literature sources that have large uncertainty values, often not having been measured directly, and are based on estimations of similar reactions.

## 2 | METHYLDIOXY RADICAL CHEMISTRY

A literature review of CH<sub>3</sub>O<sub>2</sub> chemistry will bring up research focusing on its relevance to the oxidation of atmospheric hydrocarbons.<sup>13</sup> In atmospheric chemistry, CH<sub>3</sub>O<sub>2</sub> is an important intermediate in the low-pressure, lowtemperature oxidation of hydrocarbons, predominantly reacting with NO, HO<sub>2</sub>, and OH.<sup>14</sup> Similarly, CH<sub>3</sub>O<sub>2</sub> chemistry has also been discussed in its relevance to lowtemperature combustion where the third-body reaction to form CH<sub>3</sub>O<sub>2</sub> via Reaction 1 is favored, due to the complexes increased stability at low temperatures.<sup>15,16</sup> This means that most rate coefficients available for CH<sub>3</sub>O<sub>2</sub> chemistry currently used in chemical kinetic mechanisms are based on experimental data or theoretical calculations for temperatures and pressures well below that of the Allam-Fetvedt cycle. The importance of CH<sub>3</sub>O<sub>2</sub> chemistry to the combustion of the Allam-Fetvedt cycle has been discussed by Karimi et al.<sup>12</sup> and Harman-Thomas et al.<sup>9</sup> and the role of CH<sub>3</sub>O<sub>2</sub> in the high-pressure oxidation of methane in Hashemi et al.<sup>17</sup> where it was noted that  $CH_3O_2$  was pivotal for simulating  $CO_2$  formation. It was observed that the USC II18 and GRI 3.019 chemical kinetic mechanisms overpredicted IDTs above 100 bar due to their lack of CH<sub>3</sub>O<sub>2</sub> chemistry. The subsequent addition of the CH<sub>3</sub>O<sub>2</sub> and CH<sub>3</sub>O<sub>2</sub>H species and 14 of their reactions significantly improve these mechanisms' ability to model high-pressure methane IDTs.<sup>9</sup> Similarly, Shao et al.<sup>20</sup> found that FFCM-1,<sup>21</sup> a mechanism which also does not contain CH<sub>3</sub>O<sub>2</sub> chemistry, also showed poor agreement with the high-pressure methane IDTs. The present study seeks to improve upon UoS sCO<sub>2</sub> 2.0 by updating the mechanism with the most accurate rate coefficients from more recent theoretical, experimental, and mechanistic studies as listed in Table 1.

In the present study, the  $CH_3O_2$  chemistry in UoS sCO<sub>2</sub> 2.0 was expanded through the addition of new reactions  $CH_3O_2$ , updating rate coefficients, and the addition of  $CH_2O_2H$ . The importance of the addition of each of these reactions is then discussed and the mechanism created from the rate coefficients listed in Table 1 is denoted as UoS  $sCO_2$  2.1.

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 $\label{eq:constraint} \textbf{TABLE 1} \quad \text{Rate coefficients of } CH_3O_2 \text{ reactions in the UoS s} CO_2 \text{ 2.1 mechanism.}$ 

Reaction	A ( $cm^3 mol s$ )	n	Ea (cal/mol)	Ref.
Reaction 1 (DUP) $(1 + 0) + (1 + 0)$	$5.0 \times 10^{22}$	-3.85	2000	17
$CH_3 + O_2 (+M) \rightleftharpoons CH_3O_2 (+M)^a$	$5.0 \times 10^{22}$	-3.85	2000	22
Reaction 2 $CH_4 + CH_3O_2 \rightleftharpoons CH_3 + CH_3O_2H$	$1.81 \times 10^{11}$	0.00	18,480	22
Reaction 3 $CH_3O_2 + O \rightleftharpoons CH_3O + O_2$	$2.9 \times 10^{10}$	1.00	-724	17
Reaction 4 $CH_3O_2 + H \rightleftharpoons CH_3O + OH$	$9.6 \times 10^{13}$	0.00	0	23
Reaction 5 $CH_3O_2 + OH \rightleftharpoons CH_3OH + O_2$	$1.7 \times 10^{14}$	0.00	0	24
Reaction 6 $CH_3O_2 + HO_2 \rightleftharpoons CH_3O_2H + O_2$	$2.47 \times 10^{11}$	0.00	-1570	25
Reaction 7 $CH_3O_2H + HO_2 \rightleftharpoons CH_3O_2 + H_2O_2$	$4.01 \times 104$	2.50	10,206	26
Reaction 8 $CH_3O_2 + CH_3 \rightleftharpoons CH_3O + CH_3O$	$5.08 \times 10^{12}$	0.00	-1411	27
Reaction 9 $CH_3O_2 + CH_3O_2 \rightleftharpoons CH_2O + CH_3OH + O_2$	$2.0 \times 10^{11}$	-0.55	-1600	28
Reaction 10 (DUP)	$1.1 \times 10^{18}$	-2.4	1800	28
$CH_3O_2 + CH_3O_2 \rightleftharpoons O_2 + CH_3O + CH_3O$	$7.0  imes 10^{10}$	0	8000	
Reaction 11 $H_2 + CH_3O_2 \rightleftharpoons H + CH_3O_2H$	$1.5 \times 10^{15}$	0.00	26,030	29
Reaction 12 $CH_3OH + CH_3O_2 \Rightarrow CH_2OH + CH_3O_2H$	$2.06 \times 10^{-9}$	6.20	7128	22
Reaction 13 $CH_2O + CH_3O_2 \rightleftharpoons HCO + CH_3O_2H$	$1.981 \times 10^{9}$	1.111	12,499.5	30
Reaction 14 $CH_3O_2H(+M) \rightleftharpoons CH_3O + OH(+M)^a$	$4.1 \times 10^{19}$	-1.153	44,226	31
Reaction 15 $C_2H_6 + CH_3O_2 \rightleftharpoons C_2H_5 + CH_3O_2 H$	$4.04 \times 10^{1}$	3.55	16,900	32
Reaction 16 $C_2H_4 + CH_3O_2 \rightleftharpoons C_2H_3 + CH_3O_2H$	$3.83 \times 10^{3}$	3.06	20,799	33,34
Reaction 17 CO + CH <sub>3</sub> O <sub>2</sub> $\rightleftharpoons$ CH <sub>3</sub> O + CO <sub>2</sub>	$4.22 \times 10^{6}$	0.00	0	35
Reaction 18 $CH_3O_2 + H \rightleftharpoons CH_4 + O_2$	$7.05 \times 10^{12}$	1.02	16,599	36
Reaction 19 $CH_3O_2 + OH \rightleftharpoons CH_3O + HO_2$	$4.54 \times 10^{13}$	-0.22	-363	37
Reaction 20 $CH_3O_2 + HO_2 \rightleftharpoons CH_2O + H_2O + O_2$	$9.64 \times 10^{8}$	0.00	3437	38
Reaction 21 $CH_3OH + CH_3O_2 \Rightarrow CH_3O + CH_3O_2H$	$7.937 \times 10^{8}$	4.71	13,560	39
Reaction 22 $CH_3O_2H + H \Rightarrow CH_2O_2H + H_2$	$5.4 \times 10^{10}$	0	1860	40
Reaction 23 $CH_3O_2H + H \Rightarrow CH_3O + H_2O$	$1.2 \times 10^{10}$	0	1860	40
Reaction 24 $CH_3O_2H + O \rightleftharpoons CH_2O_2H + OH$	$1.6 \times 10^{13}$	0	4750	28
Reaction 25 $CH_3O_2H + O \rightleftharpoons CH_3O_2 + OH$	$8.7 \times 10^{12}$	0	4750	28

(Continues)

Reaction	A ( $cm^3 mol s$ )	n	Ea (cal/mol)	Ref.
Reaction 26 $CH_3O_2H + OH \rightleftharpoons CH_3O_2 + H_2O$	$1.1 \times 10^{12}$	0	-437	28
Reaction 27 $CH_3O_2H + OH \Rightarrow CH_2O_2H + H_2O$	$7.2 \times 10^{11}$	0	-258	28
Reaction 28 $CH_2O_2H(+M) \rightleftharpoons CH_2O + OH(+M)^a$	$2.4 \times 10^{12}$	-0.925	1567	41

<sup>a</sup>Pressure dependence.

TABLE 1 (Continued)

Preliminary reaction pathway analysis (RPA) revealed ethane  $(C_2H_6)$  formation via the recombination of methyl radicals  $(CH_3)$  in Reaction 29. Similarly, ethane  $(C_2H_4)$ and acetaldehyde (CH<sub>3</sub>CHO) are stable intermediates in methane combustion, so these reactions may be influential in the CH<sub>3</sub>O<sub>2</sub> chemistry. Reaction 17 was added as Harman-Thomas et al.<sup>11</sup> showed that CO formation via the reverse of Reaction 30 was appreciable when using a  $CO_2$  bath gas, due to the large concentration of  $CO_2$ making this reaction more favorable. This coupled with CO being a stable intermediate of methane combustion means that even though this rate coefficient is small, at large concentrations of CO it may be important, especially at the highest values of uncertainty. Bogdanchikov et al.<sup>36</sup> showed that the reverse of Reaction 18 was negligible compared to the accepted Reaction 31 at normal combustion conditions due to the large activation energy. The reaction was included because it may become more important at higher pressures due to the larger mole fraction of CH<sub>3</sub>O<sub>2</sub> at 300 bar relative to typical combustion pressures. Zhao et al.<sup>39</sup> studied the two competing reactions Reaction 12 and Reaction 21 at 1 atm from 300-1500 K and found that Reaction 21 became increasingly important as the temperature increased, becoming the dominant pathway at 1500 K.

 $CH_3 + CH_3 (+M) = C_2H_6 (+M)$  (Reaction 29)

$$CO + OH = CO_2 + H$$
 (Reaction 30)

$$CH_4 + O_2 = CH_3 + HO_2 \qquad (Reaction 31)$$

#### 3 | MODELLING PROCEDURE

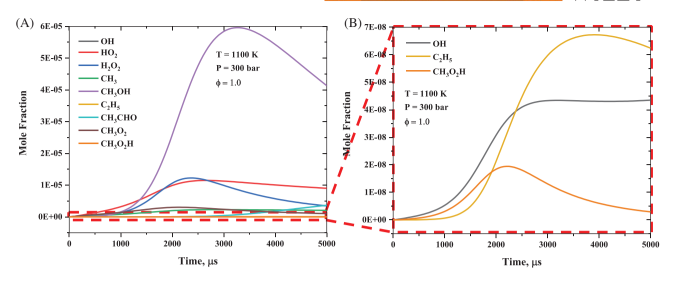
The modeling work in this work was performed using ANSYS Chemkin-Pro 2019 R3. To focus on the chemical kinetics a closed homogeneous batch reactor was used with the "constrain the volume and solve energy equation" problem type. The base conditions were set to resemble those of the Allam-Fetvedt cycle to focus on the role of  $CH_3O_2$  at large pressures. A pressure of 300 bar and a  $CO_2$  dilution of 96% was used, while the temperature was assumed to be 1100 K.<sup>42</sup> As the exact fuel-air concentrations have not been published, a stoichiometric ( $\phi = 1.0$ ) methane/oxygen mixture was used in a 96% dilution of  $CO_2$ . The UoS sCO<sub>2</sub> 2.1 mechanism was used to evaluate the need for the updated rate coefficients as well as including the important base Chemistry of UoS sCO<sub>2</sub> 2.0, which was shown to better model high-pressure IDTs of methane in a  $CO_2$  bath gas.<sup>12,20</sup>

The chemistry of  $CH_3O_2$  was investigated using sensitivity analysis, reaction pathway analysis and rate of production (ROP) analysis. Combining these analyses shows the contribution of  $CH_3O_2$  chemistry to the combustion of methane. The newly created UoS sCO<sub>2</sub> 2.1 mechanism was then compared to existing high-pressure literature IDT data to see how the changes made in the mechanism affected the simulations. Furthermore, the role of  $CO_2$  as a bath gas was investigated in  $CH_3O_2$  chemistry, by repeating the experimental run with a 96% N<sub>2</sub> and Ar bath gas. The IDT was determined as the time of maximum gradient in the temperature profile of the reaction.

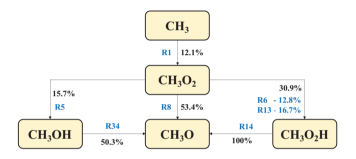
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Following the modelling procedure outlined, the chemistry of  $CH_3O_2$  was modeled at conditions relevant to the Allam-Fetvedt cycle. Figure 1A shows the mole fraction of nine selected radical species over the first 5 ms of the reaction including  $CH_3O_2$  and  $CH_3O_2H$  with Figure 1B focuses in on the three species of smallest concentration. The mole fraction of  $CH_3O_2$  in Figure 1 is comparable to  $CH_3$  under these conditions, 100% of methane will form  $CH_3$ , and as the mole fraction of  $CH_3O_2$  exceeds this. This suggests that the species has a reactivity smaller than  $CH_3$ , allowing an appreciable concentration to build up. Furthermore, the  $CH_3O_2$  mole fraction is greatest at the start of the reaction, before the ignition, suggesting that it plays a key role in the onset of ignition, explaining its importance to

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**FIGURE 1** Mole fraction of selected radical and combustion intermediates at 1100 K and 300 bar ( $CH_4:O_2:CO_2 = 1.333:2.667:96$ ) simulated by UoS sCO<sub>2</sub> 2.1.



**FIGURE 2**  $CH_3O_2$  reaction pathway at the point of ignition at 1100 K and 300 bar ( $CH_4:O_2:CO_2 = 1.333:2.667:96$ ) simulated by UoS  $sCO_2$  2.1.

high-pressure methane IDTs. However, the mole fraction of  $CH_3O_2H$  is much smaller and more closely resembles that of OH at the point of ignition despite Figure 2 showing that 30.9% of  $CH_3O_2$  goes on to form  $CH_3O_2H$ . This suggests that  $CH_3O_2H$  dissociates almost instantaneously via Reaction 14 to form  $CH_3O$  and OH under these conditions. Interestingly, Figure 1 shows that the OH concentration continues to increase after the ignition event due to the slower dissociation of  $H_2O_2$ .  $CH_3O$  an important product from the reactions of  $CH_3O_2$  is omitted from Figure 1 as the species reacted meaning a meaningful concentration is not achieved.

Figure 2 shows the key reactions and intermediates in the  $CH_3O_2$  reaction pathway. The formation of  $CH_3O_2$  via Reaction 1 competes with Reaction 32 and Reaction 33. At 300 bar, Reaction 1 and Reaction 33 are the major reaction pathways of the  $CH_3 + O_2$  reaction. These two reactions collectively account for almost 25.5% of  $CH_3$  consumption. At 30 bar, Reaction 1 is not a viable reaction path and therefore the  $CH_3O_2$  reaction pathway is negligible as almost all  $CH_3 + O_2$  proceeds via Reaction 33. It must also be noted that Karimi et al.<sup>12</sup> highlighted the greater importance of Reaction 2 in their datasets, most likely due to the greater fuel concentrations and thus increased methane concentration.

 $CH_3 + O_2 \rightleftharpoons O + CH_3O$  (Reaction 32)

$$CH_3 + O_2 \rightleftharpoons OH + CH_2O$$
 (Reaction 33)

As seen in Figure 2, 84.3% of  $CH_3O_2$  formed reacts to form  $CH_3O$ , either directly through Reaction 8 or via  $CH_3O_2H$ . The remaining  $CH_3O_2$  reacts with OH in Reaction 5 to form  $CH_3OH$ , and approximately half of  $CH_3OH$ subsequently reacts to form  $CH_3O$  under these conditions. As shown in Figure 3 the majority of  $CH_3O$  reacts to form  $CH_2O$  either via dissociation in Reaction 35 or by reacting with  $O_2$  in Reaction 35. Furthermore, despite Figure 3 showing that the majority of  $CH_3O$  is formed through Reaction 36, the next three biggest producers of  $CH_3O$  are via the  $CH_3O_2$  reaction pathway and cumulatively account for approximately one-third of the total production.

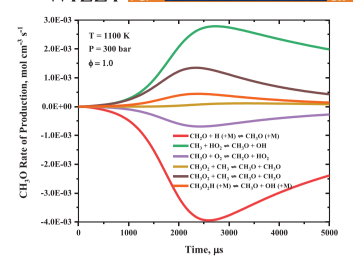
$$CH_3O(+M) \rightleftharpoons CH_2O + H(+M)$$
 (Reaction 34)

$$CH_3O + O_2 \rightleftharpoons CH_2O + HO_2$$
 (Reaction 35)

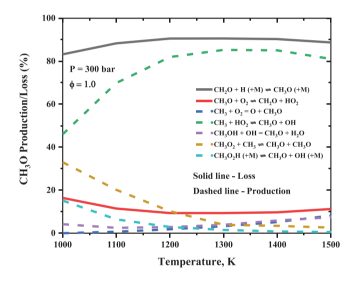
$$CH_3 + HO_2 = CH_3O + OH$$
 (Reaction 36)

Figure 4 shows the percentage of  $CH_3O$  production (dashed) and loss (solid) from 1000–1500 K across the completion reaction. Across the full temperature range, Reactions 34 and 35 remain similar in terms of  $CH_3O$  loss

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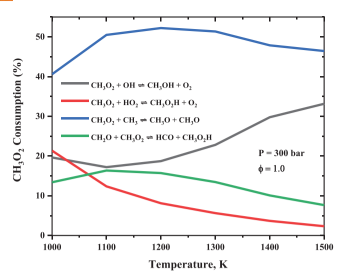
**FIGURE 3** Rate of  $CH_3O$  production at 1100 K and 300 bar simulated by UoS sCO<sub>2</sub> 2.1.



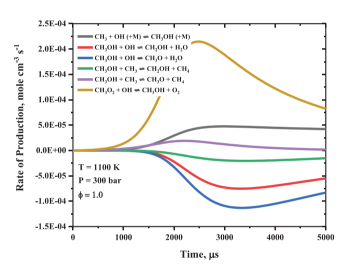
**FIGURE 4** Percentage of CH<sub>3</sub>O production (dashed) and loss (solid) simulated by UoS sCO<sub>2</sub> 2.1.

at approximately 85% and 15% respectively. Reaction 36 is the prominent route of  $CH_3O$  formation at 1000 K and becomes increasingly important as the temperature increases. The  $CH_3O_2$  reaction pathway contributes to approximately 50%  $CH_3O$  formation at 1000 K via Reactions 8 and 14 but this contribution is negligible at 1500 K, indicating a strong influence of temperature on  $CH_3O_2$  formation.

Figure 5 shows the percentage consumption of  $CH_3O_2$  at 300 bar between 1000 and 1500 K to see the influence of temperature on the reactivity since many of the rate coefficients listed in Table 1 have large temperature coefficients. Of the four most important reactions, Reaction 8 contributes to almost 45% of  $CH_3O_2$  consumption during the reaction. Reactions 6 and 13 become decreasingly important with temperature whilst Reaction 5 increases



**FIGURE 5** Effect of temperature on the consumption of  $CH_3O_2$  and the maximum net production rate of  $CH_3O_2$  simulated by UoS sCO<sub>2</sub> 2.1.



**FIGURE 6**  $CH_3OH$  rate of production at 1100 K and 300 bar simulated by UoS sCO<sub>2</sub> 2.1.

to compensate for this change. All other reactions listed in Table 1 are negligible under these conditions but could become more important at different pressure and different fuel/oxidizer combinations.

The other reactant,  $CH_3OH$ , must be considered as Figure 2 shows that 15.7% of  $CH_3O_2$  goes on to form  $CH_3OH$  via Reaction 5. Figure 6 shows the rate of  $CH_3OH$ production/loss at 1100 K and 300 bar.  $CH_3OH$  is formed primarily through Reaction 5 with Reactions 37 and 38 playing a much smaller role.  $CH_3OH$  is then consumed by reactions with OH via Reactions 39 and 40. Therefore, under these conditions, it can be considered that  $CH_3O_2$ formation dominates  $CH_3OH$  formation which subsequently reacts with OH to form  $H_2O$  and  $CH_3O$  or  $CH_2OH$ . However, it is important to note that there an alternate rate coefficient for Reaction 5 from Zhang and Huang<sup>37</sup> would significantly deemphasize the role of CH<sub>3</sub>O<sub>2</sub> on CH<sub>3</sub>OH formation. Both the rate coefficients of Bossolasco et al.<sup>24</sup> and Zhang and Huang<sup>37</sup> are validated for atmospheric chemistry and thus extrapolated well-beyond the conditions they were determined for when simulating the Allam-Fetvedt cvcle. Bossolasco et al.<sup>24</sup> was selected in this work due to it being determined using experimentally as well as being previously validated for a high-pressure methane combustion mechanism<sup>17</sup> and the negative effect Zhang and Huang<sup>37</sup> had on simulating the experimental data of Karimi et al.<sup>12</sup> However, given the large difference in the importance of CH<sub>3</sub>O<sub>2</sub> in CH<sub>3</sub>OH formation at high-pressures, it is essential that this rate coefficient is investigated under these conditions as a matter of research priority.

 $CH_3 + OH(+M) \rightleftharpoons CH_3OH(+M)$  (Reaction 37)

 $CH_3OH + CH_3 \rightleftharpoons CH_3O + CH_4$  (Reaction 38)

$$CH_3OH + OH = CH_2OH + H_2O$$
 (Reaction 39)

$$CH_3OH + OH = CH_3O + H_2O$$
 (Reaction 40)

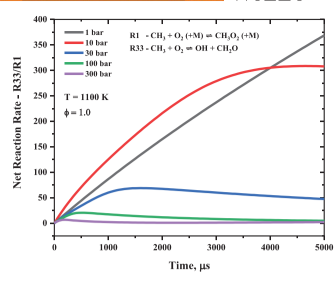
## 4.1 | Branching of CH<sub>3</sub> + O<sub>2</sub> reactions

The formation of  $CH_3O_2$  is completely due to the reaction of  $CH_3$  and  $O_2$  which makes the branching ratio of these reactions extremely important to the overall chemistry. As discussed, the rate of Reaction 32 is negligible at 300 bar. Therefore, only the rates of Reaction 1 and Reaction 33 need to be considered. Figure 7 shows the ratio of the net reaction rates Reaction 33/Reaction 1 at pressures ranging from 1 to 300 bar across the first 5000 ms of the reaction. As the pressure increases, the ratio decreases significantly from Reaction 1 which is negligible at 1 bar but becomes much more important at 100 bar and even the dominant reaction after ignition at 300 bar.

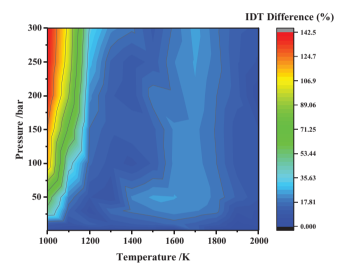
#### 5 | EFFECT OF TEMPERATURE AND PRESSURE ON IGNITION DELAY TIME

It was first noted by Karimi et al.<sup>12</sup> that  $CH_3O_2$  had an important influence on the IDT at pressures above 100 bar. It is also well-established that  $CH_3O_2$  is an important intermediate in low-temperature combustion.<sup>43</sup> The influence of  $CH_3O_2$  on the IDT was investigated over a pressure

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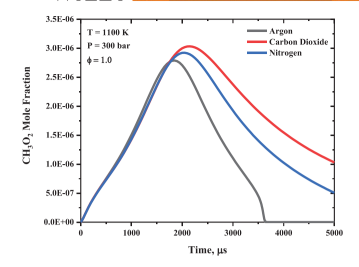
**FIGURE 7** The ratio of the Net Reaction Rate of Reaction 33/Reaction 1 at pressures from 1 to 300 bar.



**FIGURE 8** Contour plot of the difference in ignition delay time at pressures from 1–300 bar and 1000–2000 K with  $CH_3O_2$  chemistry included in the UoS s $CO_2$  2.1 mechanism.

and temperature range of 1–300 bar and 1000–2000 K, respectively. The IDT was determined as the time of the maximum gradient on the dT/dt curve. The IDT difference (%) was determined by finding the IDT using the UoS sCO<sub>2</sub> 2.1 mechanism and a modified version of the UoS sCO<sub>2</sub> 2.1 mechanism which had all CH<sub>3</sub>O<sub>2</sub> chemistry removed. The percentage difference was then determined which is shown in a contour plot in Figure 8. The importance of CH<sub>3</sub>O<sub>2</sub> to the IDT is greatest at 300 bar and 1000 K, which is to be expected based on previous research. The difference between the simulated IDTs here exceeds 140%, with the mechanism not containing CH<sub>3</sub>O<sub>2</sub> predicting the slower IDT. Furthermore, at only 50 bar and 1000 K removing the CH<sub>3</sub>O<sub>2</sub> chemistry leads to an increase in IDT by

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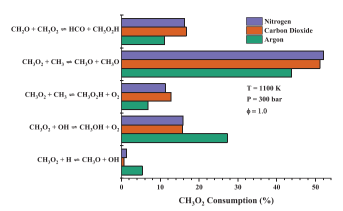


**FIGURE 9**  $CH_3O_2$  mole fraction for stoichiometric  $CH_4$  combustion in 96% carbon dioxide, nitrogen, and argon diluent at 1100 K and 300 bar.

approximately 50%. This shows a significant role of  $CH_3O_2$ at even lower pressures than discussed in Harman-Thomas et al.,<sup>9</sup> highlights the importance of using a mechanism which contains CH<sub>3</sub>O<sub>2</sub> chemistry to model methane combustion in CO<sub>2</sub> at pressures above 50 bar. Mechanisms such as FFCM-1,<sup>21</sup> USC II<sup>18</sup> and GRI 3.0<sup>19</sup> should be restricted to pressures below 50 bar when studying combustion at less than 1200 K. It should also be noted the deep blue may still indicate a pressure difference of almost 20%. Considering the experimental error in shock tubes typically varies from 15% to 25%,<sup>10,11</sup> this could still lead to large errors in chemical kinetic models. Furthermore, there is also a need to populate the current gap in methane IDT data which exists between 40 and 80 bar, to compare the effect of CH<sub>3</sub>O<sub>2</sub> on the IDT across this range where it is predicted to become an important factor. Utilizing rapid compression machines (RCMs) would allow the study of high-pressure combustion of CO<sub>2</sub> at much lower temperatures in shock tubes due to the longer achievable test times.

## $6 \quad | \quad EFFECT \ OF \ CO_2 \ ON \ CH_3O_2$ CHEMISTRY

As autoignition is rarely studied at pressures greater than 100 bar, the importance of  $CH_3O_2$  in the combustion of methane in Argon,  $N_2$  and  $CO_2$  has only been discussed briefly in existing literature.<sup>12,17,44,45</sup> Figure 9 shows the mole fraction of  $CH_3O_2$  for stoichiometric methane combustion at 1100 K and 300 bar in 96%  $CO_2$ ,  $N_2$ , and Argon. Despite the different temperatures achieved, which affect the time of maximum  $CH_3O_2$  mole fraction, the actual maximum mole fraction is consistent between the



**FIGURE 10** Percentage of  $CH_3O_2$  consumption for the five critical removal reactions for three different bath gas (carbon dioxide, nitrogen, and argon) at 1100 K and 300 bar.

three gases. As  $CO_2$  is a more efficient absorber of heat than nitrogen, and by extension argon, it absorbs more heat from the reaction and the maximum temperature achieved is less, meaning that slightly more  $CH_3O_2$  is formed as recombination reactions are favored at lower temperatures.

To investigate the influence of CO<sub>2</sub> on the actual chemistry of CH<sub>3</sub>O<sub>2</sub> the ROP of CH<sub>3</sub>O<sub>2</sub> was explored for the three different gases. As all CH<sub>3</sub>O<sub>2</sub> is formed through Reaction 1, and the CH<sub>3</sub>O<sub>2</sub> mole fractions are similar, the rate of production is similar for all three bath gases. Figure 10 shows the percentage of CH<sub>3</sub>O<sub>2</sub> consumption for the five key reactions. As shown, the chemistry of CH<sub>3</sub>O<sub>2</sub> consumption is largely unvaried between CO<sub>2</sub> and N<sub>2</sub>. For Ar, the chemistry is slightly different, the most significant change being the greater importance of Reactions 4 and 5 due to the greater temperature achieved due to Ar absorbing the heat of combustion less efficiently. The similarity of the CH<sub>3</sub>O<sub>2</sub> chemistry in the different bath gases is an essential observation for two reasons. First, the importance of CH<sub>3</sub>O<sub>2</sub> to high-pressure combustion expands beyond the application of direct-fired supercritical CO<sub>2</sub> power cycles. Other applications such as the Space-X Raptor engine which operates using a methane fuel in air at over 300 bar,<sup>46</sup> will also be heavily reliant on CH<sub>3</sub>O<sub>2</sub> chemistry and need to have accurate rate coefficients for these reactions to ensure accurate modelling of their combustion. Second, experimental techniques for the high-pressure investigation of CO<sub>2</sub>, such as shock tubes, are often susceptible to non-ideal effects such as bifurcation<sup>10,47</sup> which reduces the available test time, increases the uncertainty, and complicates the analysis. Therefore, by understanding that CH<sub>3</sub>O<sub>2</sub> chemistry is independent of bath gas, any future experimental work can be performed using argon, simplifying the analysis, and reducing the experimental error, if CO<sub>2</sub> would not have a chemical effect on any of the reactions, except for Reaction 1. It is important to consider however that the rate coefficient used for Reaction 1 has no third-body efficiencies for different gases as the rate coefficient uses a PLOG system to determine the effect of pressure on the reaction.

## 7 COMPARISON TO EXPERIMENTAL DATA

The original UoS sCO<sub>2</sub> 2.1 mechanism was developed based on three IDT datasets recorded at pressures over 200 bar from Karimi et al.<sup>12</sup> and Shao et al.<sup>20</sup> The mechanism originally showed a good agreement with the data over this pressure range. Figure 11 of NUIGMech1.1, DTU,<sup>48</sup> UoS sCO<sub>2</sub> 1.0 and UoS sCO<sub>2</sub> 2.1. Interestingly, the updating of the CH<sub>3</sub>O<sub>2</sub> rate coefficients in UoS sCO<sub>2</sub> 2.1 to those most recently published has slightly worsened the mechanism performance, although b) and Figure 11C) both fall within experimental error. One further improvement required is that Reactions 6 and 8 which were identified as important in Figure 2 are from outdated literature sources and have not been studied more recently to allow these rate coefficients to be updated. Especially given Reaction 8 is responsible for 50% of CH<sub>3</sub>O<sub>2</sub> consumption this should be reinvestigated as a research priority. The slight discrepancy from the experimental data could be due to the difference in the chemical kinetic mechanisms used outside of the CH<sub>3</sub>O<sub>2</sub> chemistry. However, the significant difference in UoS sCO<sub>2</sub> 1.0 and 2.1 shows that these rate coefficients do have a large influence on IDT under these conditions.

#### **RECOMMENDATIONS FOR FUTURE** 8 RESEARCH

This present study expands upon the work done by Harman-Thomas et al.<sup>9</sup> and Karimi et al.<sup>12</sup> further divulges the importance of the chemistry of  $CH_3O_2$  to high-pressure combustion in CO<sub>2</sub>. New investigations into the important rate coefficients are required for the most important reactions introduced in this work using experimental and theoretical investigations.

#### Important rate coefficients 8.1

Previous research into the rate coefficients of the reactions of CH<sub>3</sub>O<sub>2</sub> has been performed using high-pressure flow reactors and a pump, probe technique to monitor  $CH_3O_2$ radicals at 210, 224,49 and 240 nm.50 The most important reactions of the CH<sub>3</sub>O<sub>2</sub> reaction pathway have been dis-

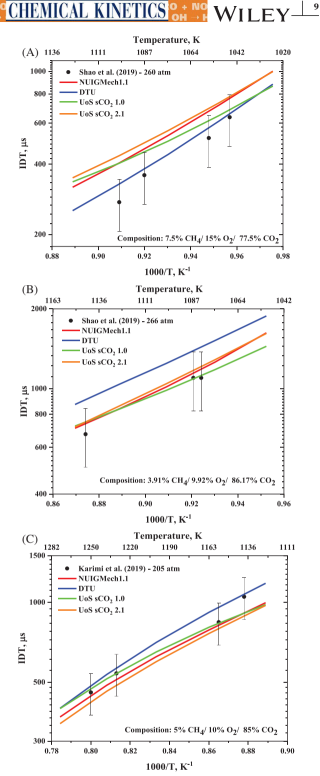


FIGURE 11 Comparison of four chemical kinetic mechanisms to IDT data from Shao et al.<sup>20</sup> and Karimi et al.<sup>12</sup>

cussed in this work and some should be revisited using modern experimental techniques as many rate coefficients currently utilized are based on limited experimental data and are almost 40 years old. As combustion at high pressures is so pertinent to modern applications, assuming full trust in these outdated rate coefficients at conditions well

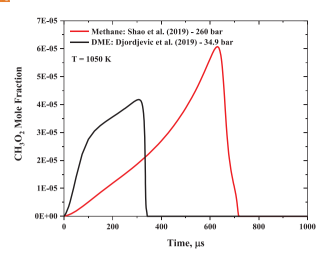
beyond those for which they were validated is no longer adequate.

Additionally, the development of new rate coefficients which deviate from that which is currently accepted must be carefully considered, regarding their validity and effect on the overall chemistry of combustion. For example, Xu et al.<sup>51</sup> recently determined a rate coefficient of hydrogen abstraction of methane by  $CH_3O_2$  (Reaction 2) which is over 100 times faster than that currently used in UoS sCO<sub>2</sub> 2.0<sup>33,34</sup> above 1000 K. This would significantly increase the impact of Reaction 2 on  $CH_3O_2$  formation from the 2.3% shown in Figure 2. However, as UoS sCO<sub>2</sub> 2.0 is looking at modeling combustion a specific environment of large  $CO_2$  dilutions and high-pressures, each new rate coefficient must be carefully evaluated based on its influence on the mechanisms ability to simulate experimentally determined combustion characteristics such as IDT.

The two most important rate coefficients for reinvestigation are Reactions 6 and 8 as identified by the RPA. Reaction 8 is responsible for 50% of CH<sub>3</sub>O<sub>2</sub> consumption and over 30% of CH<sub>3</sub>O formation at 1000 K (see Figure 4). The rate coefficient used in all of the mechanisms investigated in Section 7 is from a 1988 publication by Keiffer et al.<sup>27</sup> This work utilized a laser photolysis technique to study CH<sub>3</sub> decay curves which were subsequently fitted using a global technique to determine the rate coefficient of Reaction 1 and Reaction 8 at 1 atm over a narrow temperature range (293-530 K). Due to the importance of this reaction, it should be investigated as a priority for future research into the development of a mechanism for modelling high-pressure methane mechanism. Similarly, Reaction 6 was last studied by Lightfoot et al.<sup>23</sup> in 1991 between 600 and 719 K at atmospheric in flash photolysis study and RRKM calculations. Studying these two reactions under high-pressure conditions using modern experimental and theoretical techniques over a much broader temperature range is the next research hurdle to properly understanding the high-pressure reaction pathway of  $CH_3O_2$ .

## 8.2 | Ignition delay time studies

Furthermore, more experimental IDT data is required for the combustion of methane at pressures between 50 and 300 bar in  $CO_2$ ,  $N_2$ , and Ar to ratify the currently accepted  $CH_3O_2$  chemistry in terms of mechanism performance in these conditions. This would add important data for further mechanism refinement of the  $CH_3O_2$  chemistry and is essential to justifying some of the rate coefficients used in current mechanisms which are based on estimates or limited experimental and theoretical data. This is essential to allow accurate modeling of combustion in the Allam-



**FIGURE 12** CH<sub>3</sub>O<sub>2</sub> mole fraction for DME mixture (6.4% DME/19.2% O<sub>2</sub>/74.4% CO<sub>2</sub>) at 34.9 bar from Djordjevic et al.<sup>55</sup> and methane mixture (7.5% CH<sub>4</sub>/15% O<sub>2</sub>/77.5% CO<sub>2</sub>) at 260 bar from Shao et al.<sup>20</sup> simulated by NUIGMech1.1.

Fetvedt cycle and therefore used to increase performance and efficiency. Most shock tube facilities can only operate at pressures below 100 bar due to safety constraints in having sufficient design pressure to withstand detonation. This limits the potential to attain more methane IDT datasets under conditions where the chemistry of  $CH_3O_2$  is relevant, as the temperature must also be below 1200 K, which means longer test times are required. This can be achieved by studying methane ignition in rapid compression machines.

It is possible to validate the rate coefficients of  $CH_3O_2$ using different fuels with form CH<sub>3</sub>O<sub>2</sub> at lower pressures and with short ignitions at lower temperatures. The two obvious candidates are dimethyl ether (DME) and diethyl ether (DEE). Chemkin simulations (see Figure 12) using the NUIGMech1.1<sup>52-54</sup> chemical kinetic mechanism shows that the maximum mole fraction of CH<sub>3</sub>O<sub>2</sub> from a 38.4 atm (6.4% DME/19.2% O<sub>2</sub>/74.4% CO<sub>2</sub>) mixture at 900 K from Djordjevic et al.<sup>55</sup> is the same as that produced from methane ignition at 260 atm (75%  $CH_4/15\% O_2/77.5\% CO_2$ ) from Shao et al.<sup>20</sup> at 1050 K. NUIGMech1.1 was used as UoS sCO<sub>2</sub> 2.1 does not contain any DME chemistry. Furthermore, Issayev et al.<sup>56</sup> found that CH<sub>3</sub>O<sub>2</sub> played an important role in the ignition of DEE and DEE/ethanol blends over temperature and pressure ranges of 500-1000 K and 20-40 bar, respectively. Thus, the rate coefficients of the key reactions discussed in this work can be further validated using the DME and DEE experimental data.

#### 9 | CONCLUSION

This work takes this further by investigating the importance of  $CH_3O_2$  chemistry to high-pressure combustion in CO<sub>2</sub>. Identifying its importance to methane combustion above 50 bar at temperatures below 1200 K, where the absence of CH<sub>3</sub>O<sub>2</sub> chemistry can lead to an increase in the IDT by up to 140%. As CH<sub>3</sub>O<sub>2</sub> is formed in its entirety through the recombination of CH<sub>3</sub> and O<sub>2</sub>, the branching ratio compared to Reaction 33 is essential. It has been demonstrated that at 1 bar, CH<sub>3</sub>O<sub>2</sub> formation is negligible, but becomes the dominant CH<sub>3</sub>O<sub>2</sub> branching pathway at points during the reaction at 300 bar. Furthermore, the UoS  $sCO_2$  2.1 mechanism was created by updating the reactions of CH<sub>3</sub>O<sub>2</sub> to the most up-to-date rate coefficients as shown in Table 1. It was identified that Reaction 6 ( $CH_3O_2 + CH_3 = CH_3O + CH_3O$ ) and Reaction 8 (CH<sub>3</sub>O<sub>2</sub> + HO<sub>2</sub> = CH<sub>3</sub>O<sub>2</sub>H + OH) are essential to highpressure methane combustion and are a priority to restudy using experimental and theoretical techniques. Furthermore, it was noted that the selection of rate coefficient for Reaction 5 had a large impact on the importance of the methanol reaction pathway and thus this reaction should be studied at pressures relevant to the Allam-Fetvedt cycle. It was identified that DME and DEE produced similar maximum mole fractions of CH<sub>3</sub>O<sub>2</sub> at lower pressures and temperatures than methane, which are more achievable in experimental studies. Therefore, the chemistry of DME and DEE should be added to UoS sCO<sub>2</sub> 2.1 and validated using existing IDT data to further validate the mechanism. Furthermore, rapid compression machines should be used to generate methane ignition data at lower temperatures due to the longer test times achievable.

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#### DATA AVAILABILITY STATEMENT None.

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#### REFERENCES

- 1. Oxford A. Ukraine war threatens European climate goals. Emerald Expert Briefings. 2022. no. oxan-db.
- Liadze I, Macchiarelli C, Sanchez Juanino P, Mortimer-Lee P. The economic costs of the Russia-Ukraine conflict.NIESR Policy Paper, No.32. 2022. https://www.niesr.ac.uk/publications/ economic-costs-russia-ukraine-conflict?type=policy-papers
- 3. Watson R, McCarthy J, Canziani P, Nakicenovic N, Hisas L. *The truth behind the climate pledges*. FEU-US; 2019.

INTERNATIONAL JOURNAL OF

- Allam RJ, Palmer MR, Brown GW, et al. High efficiency and low cost of electricity generation from fossil fuels while eliminating atmospheric emissions, including carbon dioxide. *Energy Procedia*. 2013;37:1135-1149. doi:10.1016/j.egypro.2013. 05.211
- Lariviere B, Marion J, Macadam S, et al. "sCO<sub>2</sub> power cycle development and step demo pilot project". 4th European sCO<sub>2</sub> Conference for Energy Systems: March 23-24, 2021, Online Conference, 2021:352-362.
- Rathi A. U.S. startup plans to build first zero-emission gas power plants. Bloomberg Green. Accessed April 15, 2021. https://www.bloomberg.com/news/articles/2021-04-15/u-sstartup-plans-to-build-first-zero-emission-gas-power-plants
- Kelsall G. 8 Rivers Capital and Sembcorp Energy UK's first zero emissions power plant. IFRF. Accessed August 2, 2021. https://ifrf.net/ifrf-blog/8-rivers-capital-and-sembcorpenergy-uk-to-develop-uks-first-net-zero-emissions-powerplant/
- Coogan S, Gao X, McClung A, Sun W. Evaluation of kinetic mechanisms for direct fired supercritical oxy-combustion of natural gas. Presented at the Conference: ASME Turbo Expo 2016: Turbo-machinery Technical Conference and Exposition, Seoul, South Korea; 2016. [Online]. Available: doi:10.1115/GT2016-56658
- Harman-Thomas JM, Hughes KJ, Pourkashanian M. The development of a chemical kinetic mechanism for combustion in supercritical carbon dioxide. *Energy*. 2022:255:124490. doi:10. 1016/j.energy.2022.124490
- Harman-Thomas JM, Kashif TA, Hughes KJ, Pourkashanian M, Farooq A. Experimental and modelling study of hydrogen ignition in CO2 bath gas. *Fuel.* 2023;334:126664. doi:10.1016/j.fuel. 2022.126664
- Harman-Thomas JM, Kashif TA, Hughes KJ, Pourkashanian M, Farooq A. Experimental and modelling study of syngas combustion in CO2 bath gas. *Fuel.* 2023;342:127865. doi:10.1016/j.fuel. 2023.127865
- Karimi M, Ochs B, Liu Z, Ranjan D, Sun W. Measurement of methane autoignition delays in carbon dioxide and argon diluents at high pressure conditions. *Combust Flame*. 2019;204:304-319. doi:10.1016/j.combustflame.2019.03.020
- Assaf E, Sheps L, Whalley L, et al. The reaction between CH<sub>3</sub>O<sub>2</sub> and OH radicals: product yields and atmospheric implications. *Environ Sci Technol.* 2017;51(4):2170-2177. doi:10.1021/ acs.est.6b06265
- Fittschen C, Whalley LK, Heard DE. The reaction of CH<sub>3</sub>O<sub>2</sub> radicals with OH radicals: a neglected sink for CH<sub>3</sub>O<sub>2</sub> in the remote atmosphere. *Environ Sci Technol.* 2014;48(14):7700-7701. doi:10.1021/es502481q
- Villenave E, Lesclaux R. Kinetics of the cross reactions of CH<sub>3</sub>O<sub>2</sub> and C<sub>2</sub>H<sub>5</sub>O<sub>2</sub> radicals with selected peroxy radicals. *J Phys Chem.* 1996;100(34):14372-14382. doi:10.1021/jp960765m
- Jafri JA, Phillips DH. Ground and lower excited states of methylperoxy radical, CH<sub>3</sub>O<sub>2</sub>: a computational investigation. J Am Chem Soc. 1990;112(7):2586-2590. doi:10.1021/ja00163a017
- Hashemi H, Christensen JM, Gersen S, Levinsky H, Klippenstein SJ, Glarborg P. High-pressure oxidation of methane. *Combust Flame*. 2016;172:349-364. doi:10.1016/j.combustflame.2016.07.016

# <sup>12</sup> WILEY WILEY OF CHEMICAL KINETICS OF NO

- Wang H, You X, Joshi AV, et al. High-temperature combustion reaction model of H<sub>2</sub>. High-temperature Combustion Reaction Model of H<sub>2</sub>/CO/C1-C4 Compounds. 2007.
- Smith GP, Golden DM, Frenklach M, et al. GRI Mechanism 3.0. University of California, Berkeley. Accessed December 16, 2019. http://combustion.berkeley.edu/gri-mech/version30/ text30.html
- Shao J, Choudhary R, Davidson DF, Hanson RK, Barak S, Vasu S. Ignition delay times of methane and hydrogen highly diluted in carbon dioxide at high pressures up to 300 atm. *Proc Combust Inst.* 2019;37(4):4555-4562. doi:10.1016/j.proci.2018.08.002
- Smith GP, Tao Y, Wang H. Foundational Fuel Chemistry Model Version 1.0 (FFCM-1), http://nanoenergy.stanford.edu/ffcm1, 2016.
- 22. Wang Z, Zhao H, Yan C, et al. Methanol oxidation up to 100 atm in a supercritical pressure jet-stirred reactor. *Proceedings of the Combustion Institute*, 2022. doi:10.1016/j.proci.2022.07.068
- 23. Lightfoot PD, Roussel P, Caralp FO, Lesclaux R. Flashphotolysis study of the  $CH_3O_2 + CH_3O_2$  and  $CH_3O_2 + HO_2$ reactions between 600 and 719 K – unimolecular decomposition of methylhydroperoxide. *J Chem Soc, Faraday Trans.* 1991;87(19):3213-3220. doi:10.1039/ft9918703213
- Bossolasco A, Faragó EP, Schoemaecker C, Fittschen C. Rate constant of the reaction between CH<sub>3</sub>O<sub>2</sub> and OH radicals. *Chem Phys Lett.* 2014;593:7-13. doi:10.1016/j.cplett.2013.12.052
- 25. Lightfoot PD, Roussel P, Caralp FO, Lesclaux R. Flash-photolysis study of the  $CH_3O_2 + CH_3O_2$  and  $CH_3O_2 + HO_2$  reactions between 60 and 719 K Unimolecular decomposition of methylhydroperoxide. *J Chem Soc, Faraday Trans.* 1991;87(19):3213-3220. doi:10.1039/ft9918703213
- Rasmussen CL, Wassard KH, Dam-Johansen K, Glarborg P. Methanol oxidation in a flow reactor: implications for the branching ratio of the CH<sub>3</sub>OH+OH reaction. *Int J Chem Kinet*. 2008;40(7):423-441. doi:10.1002/kin.20323
- Keiffer M, Miscampbell AJ, Pilling MJ. A global technique for analysing multiple decay curves. Application to the CH<sub>3</sub> + O<sub>2</sub> system. *J Chem Soc, Faraday Trans 2*. 1988;84(5):505-514.
- Baulch DL, Bowman CT, Cobos CJ, et al. Evaluated kinetic data for combustion modeling: supplement II. *J Phys Chem Ref Data*. 2005;34(3):757-1397. doi:10.1063/1.1748524
- Tsang W, Hampson RF. Chemical kinetic data base for combustion chemistry. Part I. Methane and related compounds. *J Phys Chem Ref Data*. 1986;15(3):1087-1279. doi:10.1063/1.555759
- 30. Labbe NJ, Sivaramakrishnan R, Goldsmith CF, Georgievskii Y, Miller JA, Klippenstein SJ. Weakly bound free radicals in combustion: "Prompt" dissociation of formyl radicals and its effect on laminar flame speeds. *J Phys Chem Lett.* 2016;7(1):85-89. doi:10.1021/acs.jpclett.5b02418
- Jasper AW, Klippenstein SJ, Harding LB. Theoretical rate coefficients for the reaction of methyl radical with hydroperoxyl radical and for methylhydroperoxide decomposition. *Proc Combust Inst.* 2009;32(1):279-286. doi:10.1016/j.proci.2008.05.036
- 32. Carstensen H-H, Dean AM, Deutschmann O. Rate constants for the H abstraction from alkanes (R-H) by R'O2 radicals: a systematic study on the impact of R and R'. *Proc Combust Inst.* 2007;31(1):149-157. doi:10.1016/j.proci.2006. 08.091
- 33. Wu Y, Panigrahy S, Sahu AB, et al. Understanding the antagonistic effect of methanol as a component in surrogate fuel models:

a case study of methanol/n-heptane mixtures. *Combust Flame*. 2021;226:229-242.

- 34. Baigmohammadi M, Patel V, Nagaraja S, et al. Comprehensive experimental and simulation study of the ignition delay time characteristics of binary blended methane, ethane, and ethylene over a wide range of temperature, pressure, equivalence ratio, and dilution. *Energy Fuels.* 2020;34(7):8808-8823. doi:10. 1021/acs.energyfuels.0c00960
- 35. Sander SP, Watson RT. Kinetics studies of the reactions of  $CH_3O_2$  with NO,  $NO_2$ , and  $CH_3O_2$  at 298 K. J Phys Chem. 1980;84(13):1664-1674.
- 36. Bogdanchikov GA, Baklanov AV, Parker DH. The substitution reactions RH+O<sub>2</sub>→RO<sub>2</sub>+H: transition state theory calculations based on the ab initio and DFT potential energy surface. *Chem Phys Lett.* 2004;385(5):486-490. doi:10.1016/j.cplett.2004.01.015
- Zhang F, Huang C. Pressure-dependent kinetics of the reaction between CH<sub>3</sub>OO and OH focusing on the product yield of methyltrioxide (CH<sub>3</sub>OOOH). *J Phys Chem Lett.* 2019;10(13):3598-3603. doi:10.1021/acs.jpclett.9b00781
- Elrod MJ, Ranschaert DL, Schneider NJ. Direct kinetics study of the temperature dependence of the CH<sub>2</sub>O branching channel for the CH<sub>3</sub>O<sub>2</sub> + HO<sub>2</sub> reaction. *Int J Chem Kinet*. 2001;33(6):363-376. doi:10.1002/kin.1030
- Zhao Z, Song J, Su B, Wang X, Li Z. Mechanistic study of the reactions of methyl peroxy radical with methanol or hydroxyl methyl radical. *J Phys Chem A*. 2018;122(23):5078-5088. doi:10.1021/acs. jpca.7b09988
- Slemr F, Warneck P. Kinetics of the reaction of atomic hydrogen with methyl hydroperoxide. *Int J Chem Kinet*. 1977;9(2):267-282. doi:10.1002/kin.550090210
- Ing W-C, Sheng CY, Bozzelli JW. Development of a detailed high-pressure reaction model for methane/methanol mixtures under pyrolytic and oxidative conditions and comparison with experimental data. *Fuel Process Technol.* 2003;83(1):111-145. doi:10.1016/S0378-3820(03)00062-6
- 42. Allam RJ, Fetvedt JE, Forrest BA, Freed DA. The oxy-fuel, supercritical CO<sub>2</sub> Allam cycle: new cycle developments to produce even lower-cost electricity from fossil fuels without atmospheric emissions. *Proceedings of the Asme Turbo Expo: Turbine Technical Conference and Exposition.* 3b, 2014, Art no. V03bt36a016.
- Yan C, Kocevska S, Krasnoperov LN. Kinetics of the reaction of CH<sub>3</sub>O<sub>2</sub> radicals with OH studied over the 292–526 K temperature range. *J Phys Chem A*. 2016;120(31):6111-6121.
- Huang J, Hill PG, Bushe WK, Munshi SR. Shock-tube study of methane ignition under engine-relevant conditions: experiments and modeling. *Combust Flame*. 2004;136(1):25-42. doi:10. 1016/j.combustflame.2003.09.002
- Petersen E, Davidson D, Hanson R. Kinetics modeling of shock-induced ignition in low-dilution CH<sub>4</sub>/O<sub>2</sub> mixtures at high pressures and intermediate temperatures. *Combust Flame*. 1999;117(1):272-290. doi:10.1016/S0010-2180(98)00111-4
- O'Callaghan J. The wild physics of Elon Musk's methaneguzzling super-rocket. Wired. Accessed February 11, 2021. https://www.wired.co.uk/article/spacex-raptor-engine-starship
- Hargis JW, Petersen EL. Methane ignition in a shock tube with high levels of CO<sub>2</sub> dilution: consideration of the reflected-shock bifurcation. *Energy Fuels*. 2015;29(11):7712-7726. doi:10.1021/acs. energyfuels.5b01760

- Hashemi H, Christensen JM, Glarborg P. High-pressure pyrolysis and oxidation of DME and DME/CH<sub>4</sub>. *Combust Flame*. 2019;205:80-92. doi:10.1016/j.combustflame.2019.03.028
- Yan C, Krasnoperov LN. Pressure-dependent kinetics of the reaction between CH<sub>3</sub>O<sub>2</sub> and OH: TRIOX formation. J Phys Chem A. 2019;123(39):8349-8357. doi:10.1021/acs.jpca. 9b03861
- 50. Fernandes RX, Luther K, Troe J. Falloff curves for the reaction  $CH_3 + O_2 (+M) \rightarrow CH_3O_2 (+M)$  in the pressure range 2–1000 bar and the temperature range 300–700 K. *J Phys Chem A*. 2006;110(13):4442-4449. doi:10.1021/jp0568500
- Xu S, Liang J, Cao S, He R, Yin G, Wang Q-D. A hierarchical theoretical study of the hydrogen abstraction reactions of H2/C1–C4 molecules by the methyl peroxy radical and implications for kinetic modeling. *ACS Omega*. 2022;7(10):8675-8685. doi:10.1021/acsomega.1c06683
- Metcalfe WK, Burke SM, Ahmed SS, Curran HJ. A hierarchical and comparative kinetic modeling study of C1 – C2 hydrocarbon and oxygenated fuels. *Int J Chem Kinet*. 2013;45(10):638-675. doi:10.1002/kin.20802
- 53. Burke U, Metcalfe WK, Burke SM, Heufer KA, Dagaut P, Curran HJ. A detailed chemical kinetic modeling, ignition delay time

and jet-stirred reactor study of methanol oxidation. *Combust Flame*. 2016;165:125-136. doi:10.1016/j.combustflame.2015.11.004

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 Kéromnès A, Metcalfe WK, Heufer KA, et al. An experimental and detailed chemical kinetic modeling study of hydrogen and syngas mixture oxidation at elevated pressures. *Combust Flame*. 2013;160(6):995-1011. doi:10.1016/j.combustflame.2013.01.001

CHEMICAL KINETICS

- Djordjevic N, Rekus M, Vinkeloe J, Zander L. Shock tube and kinetic study on the effects of CO<sub>2</sub> on dimethyl ether autoignition at high pressures. *Energy Fuels*. 2019;33(10):10197-10208. doi:10.1021/acs.energyfuels.9b01575
- Issayev G, Mani Sarathy S, Farooq A. Autoignition of diethyl ether and a diethyl ether/ethanol blend. *Fuel.* 2020;279:118553. doi:10.1016/j.fuel.2020.118553

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