Characteristics of Ammonia/Hydrogen Premixed Combustion in a Novel Linear Engine Generator †

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Abstract: In order to support the development of a novel linear engine generator (LEG), the characteristics of ammonia/hydrogen premixed combustion are studied by using a detailed chemical kinetics mechanism. The ammonia combustion mechanism is identified among several mechanisms and validated with published experimental data. A parametric analysis is carried out under LEG typical working conditions to study the effects of equivalence ratio (0.80 – 1.60), hydrogen blending ratio (0.0 – 0.6), initial temperature (300 – 700 K) and initial pressure (1 – 20 bar) on premixed laminar flame speed, ignition delay and key flame species concentrations. It is shown that an equivalence ratio of around 1.10 – 1.20 is beneficial to both ammonia flame stability and lower NOx emission. Ignition delay is reduced with the increase in hydrogen blending ratio, initial temperature and initial pressure. At a certain initial temperature and initial pressure, the effects of hydrogen blending ratio can be negligible for over 50% hydrogen in the fuel. Under higher pressure (>10 bar), the initial pressure has a minor influence on the ignition delay reduction. It is also found that the high-pressure high-temperature environment contributes to reducing NOx emission considerably in ammonia/hydrogen combustion, which implies the potential of a low NOx LEG fuelled by ammonia/hydrogen.

Keywords: ammonia/hydrogen premixed combustion; linear engine generator; chemical kinetics model

1. Introduction

In order to reduce CO2 emission from the transportation industry, various alternative powertrain designs are proposed and developed in addition to the improvement of conventional powertrains such as the internal combustion engine. The concept of the linear engine generator (LEG) is one of the most promising approaches. In an LEG, a free piston linear engine converts chemical energy of fuel into mechanical energy, which will then be converted into electric power through an integrated linear alternator. Compared to conventional engines with a crankshaft system, free piston engines have a simpler and more compact structure, lower friction loss, more control freedom and wider fuel adaptability [1,2]. The integration between the linear engine and the alternator provides an attractive hybrid powertrain solution, which enables the engine to be operated under optimized working conditions and, therefore, achieve high system efficiency. The German Aerospace Centre (DLR) have been developing a free piston linear generator with either single or dual opposed piston configuration and a functioning demonstration model was put into operation in 2013 [3]. A prototype
engine–linear alternator (FPLA) system was developed at Sandia National Laboratories [4]. High efficiency was obtained by utilizing homogeneous charge compression ignition (HCCI) combustion concept with high compression ratio (~30:1) and low equivalence ratio (~0.35). Toyota developed a prototype two-stroke engine linear generator (FPEG) and claimed that 42% efficiency could be achieved [5,6]. Researchers from Newcastle University [7,8] raised the linear Joule engine generator (LJEG) concept, and a working prototype has been developed aiming for potential application in micro-scale combined heat and power (CHP) systems.

Zero-carbon-emission alternative fuel is another important approach to reducing CO₂ emission. In particular, extensive efforts have been devoted to the application of hydrogen and hydrogen carriers in the powertrain system. Ammonia is considered as a potential hydrogen carrier as well as a fuel without hydrogen extraction [9–11]. It is reported that ammonia utilization is vital for the future low-carbon energy system in Japan and the ammonia supply chain is under development and is scheduled to be completed by 2030 [11,12]. The attractiveness of ammonia mainly lies in the following aspects: ammonia has a high gravimetric and volumetric hydrogen density compared with various hydrogen carriers [13]; the industry of ammonia production, storage, transportation and utilization is well-established [9,11]; sustainable and renewable ammonia production technologies are available and under development and improvement [14].

The concept of ammonia-fuelled LEG is promising, as it combines the merits of LEG as an efficient powertrain and the merits of ammonia as a carbon-free hydrogen carrier. Inspired by this idea, an ammonia-fuelled LEG is under development by the authors’ team. Although reports on the application of ammonia in conventional spark-ignition or compression-ignition internal combustion engines [15,16] and gas turbines [17,18] are readily available, research on an ammonia-fuelled linear engine or LEG has not been reported to the knowledge of the authors.

Despite the merits of ammonia as a zero-carbon-emission alternative fuel, challenges should be overcome before practical application of ammonia in an LEG is realized. Firstly, ammonia has higher minimum ignition energy and slower burning velocity compared with conventional hydrocarbon fuel and hydrogen, which causes difficulty in ensuring reliable ignition and maintaining stable combustion within the combustor [19,20]. The addition of hydrocarbon or hydrogen [17,21] and increased oxygen content within the oxidizer [22] were reported as effective methods to improve combustion robustness. Another challenge is to effectively reduce NOx emission from ammonia combustion, as fuel-nitrogen plays a crucial role in NOx formation in addition to the thermal-NO mechanism [23], which is dominant for hydrocarbon fuel combustion. A deep understanding of the characteristics of ammonia/hydrogen combustion is of great importance for developing a robust and low-NOx emission combustion system of an ammonia-fuelled LEG.

The Gas Turbine Research Centre (GTRC) of Cardiff University carried out a premixed (50:50 vol%) ammonia/hydrogen burner experiment at the inlet temperature of 288 K and atmospheric pressure. It was found that the flame was stabilized in a relatively narrow range of equivalence ratio (0.43 – 0.52) in a swirl combustor due to the high diffusivity and reactivity of hydrogen, which may potentially lead to boundary layer flashback [21]. The burning velocity of premixed ammonia-hydrogen combustion was measured by Li et al. at normal temperature and pressure (NTP) and the emission characteristics were numerically analysed in relation to the hydrogen addition in the fuel of 40.0% – 66.7% (mole fraction) and the equivalence ratio of 1.00, 1.10 and 1.25 [24]. The results suggested that the stoichiometric conditions should be avoided since NOx emissions reached peak and unburnt hydrogen was also detected at the equivalence ratio of unity. Ichikawa et al. [25] performed the ammonia/hydrogen experiment up to 0.5 MPa for the first time at the 298 K initial temperature and stoichiometric condition and the effects of hydrogen addition (0.0 – 1.0) and initial pressure (0.1, 0.3, 0.5 MPa) on unstretched laminar burning velocity and burned Markstein length were also investigated with simulations using five different chemical kinetic mechanisms. Nozari et al. numerically explored the premixed laminar flame speed and species profiles of NH₃/H₂ at a pressure level of 17 bar and an initial temperature of 673 K under lean equivalence ratio (0.5), which is suitable for a general gas turbine combustor [26]. The evidence suggested that fuel composition is a key factor of NOx reduction and the authors deduced
that fuel rich conditions could reduce NOx emission considerably. Recently, a numerical study on 70:30 mol% NH₃/H₂ fuelled swirler combustor using Reynolds-averaged Navier–Stokes (RANS) models was presented at the rich equivalence ratio (1.2) and medium swirl (0.8) [27]. The 3D emission profiles were obtained under high pressures and high inlet temperatures. As these works contribute a lot to the understanding of ammonia/hydrogen combustion, more detailed and extensive investigation under engine-operation conditions is still required to better support the combustion system of the target LEG prototype.

In this work, a modelling study is conducted to investigate ammonia/hydrogen premixed combustion characteristics under the target LEG’s potential operating conditions. An ammonia combustion mechanism is identified among representative mechanisms based on extensive validation against published experimental data. Then, a parametric analysis in terms of laminar flame speed, ignition delay and flame species concentration is performed to study the effects of equivalence ratio (0.8−1.6), hydrogen blending ratio (0.0−0.6), initial temperature (300−700 K) and initial pressure (1−20 bar). The findings from this study will support the development of the combustor for the target LEG prototype.

2. Methodology

A comprehensive review of ammonia combustion mechanism development is conducted, and three representative mechanisms are identified, which are validated against experimental data under various conditions, respectively. Then, the three identified mechanisms are compared against the same sets of literature experimental data, which cover flame speed, ignition delay and flame species concentration. The mechanism which shows best agreement with the test results is employed for the subsequent parametric study.

2.1. Ammonia Combustion Mechanism Review

Miller and Bowman [28] developed a detailed kinetic nitrogen reaction mechanism and explained NOₓ formation in ammonia oxidation. Based on Miller and Bowman’s work, Klippenstein et al. [29] studied the role of NNH on NO formation (NNH + O, NNH + O₂, NH₂ + O₂). To investigate ammonia combustion under high temperature (450–925 K) and pressure (30 and 100 bar), an experiment of ammonia oxidation was carried out and a modified mechanism was proposed by Song et al. [30]. However, the concentrations of NO and NO₂ were not measured due to the limitations of the experimental equipment. Followed by Song’s mechanism, an updated kinetic reaction mechanism including NH₂, HNO and N₂H₂ relevant reactions was developed by Otomo et al. [31]. A detailed mechanism including 127 species and over 1200 reactions was proposed by Konnov [32,33]. This model can be used to investigate the oxidation process of the blends of ammonia with small hydrocarbons and hydrogen. Duynslaegher et al. [34,35] examined several mechanisms, including GRI [36], San Diego [37], Lindstedt [38], Bian [39] and Konnov [32,33], and found that Konnov’s mechanism has good agreement on main species except N₂O and NH₂ under low pressure conditions. To improve the modelling accuracy, a revised mechanism was developed and further reduced by Duynslaegher et al. [34]. Recently, on the basis of Miller and Bowman’s mechanism, Mathieu and Petersen’s mechanism [40], Konnov’s mechanism and Bugler’s updated more accurate database of the thermochemical parameters [41], Nakamura et al. [42] proposed an ammonia oxidation mechanism, which showed good agreement with previous measured results. In the present work, Otomo’s mechanism [31], Duynslaegher’s mechanism [34] and Nakamura’s mechanism [42] are compared against the same sets of literature experimental data to provide the basis for subsequent parametric analysis.

2.2. Parametric Study Conditions

2.2.1. Premixed Laminar Flame Speed Modelling

Premixed laminar flame speed is one of the most important combustion parameters. It is widely used for combustion kinetic model validation and has a decisive influence on turbulent flame speed,
which is a key parameter for practical combustion system development. As ammonia is reported to have a much lower flame speed compared with hydrocarbon fuel and hydrogen, it is important to promote ammonia oxidation based on the knowledge of the influence of various parameters on ammonia flame speed. Laminar flame speed calculation is conducted based on the 1D premixed freely propagating flame sub-model of ANSYS CHEMKIN PRO [43]. The influence of hydrogen blending ratio \((0.0 - 0.6)\), fuel-air equivalence ratio \((\phi)\ \((0.8 - 1.6)\), initial temperature \((300 - 700 \, K)\) and initial pressure \((1 - 20 \, bar)\) on laminar flame speed is investigated. The hydrogen blending ratio, \(x\%H_2\), which is defined as the mole fraction of hydrogen in the fuel mixture, is determined by:

\[
x\%H_2 = \frac{x(H_2)}{x(H_2) + x(NH_3)}
\]

where \(x(H_2)\) and \(x(NH_3)\) denote the mole fraction of hydrogen and ammonia, respectively.

2.2.2. Ignition Delay Time Modelling

Ignition delay, which is defined as the time for a mixture to reach autoignition, is another critical parameter for premixed combustion. It is important for both theoretical study and dealing with either normal or abnormal ignition behaviour within a practical combustion system. The ignition delay time modelling is carried out by using the 0D closed homogeneous reactor sub-model of ANSYS CHEMKIN PRO, where the ignition delay time can be determined as the time during which a certain species concentration reaches its maximum or the inflection point of temperature appears [43]. In the present study, the ignition delay time is defined as the time to reach the maximum concentration of \(OH\).

2.2.3. Burner-Stabilized Flame Structure Modelling

Minimizing \(NO_x\) emissions is a key target in the development of an ammonia-fuelled engine, where the fuel-bound \(NO_x\) emission is expected to play an important role. In order to gain a deeper insight into \(NO_x\) formation and the consumption mechanism of ammonia/hydrogen premixed combustion, the burner-stabilized flame structure is investigated by employing ANSYS CHEMKIN PRO software. The mole fraction profiles of fuels \((NH_3\ and \ H_2)\), \(NO_x\ (NO, \ N_2O\ and \ NO_2)\) and key radicals \((OH, \ H\ and \ O)\) are derived and analysed with an initial temperature of \(700 \, K\) and initial pressure of \(20 \, bar\), which is typical for the target LEG working condition.

3. Mechanism Selection for Parametric Study

In order to determine the mechanism to be used for parametric study, a cross-comparison of the three mechanisms identified in the previous section is conducted against the same sets of test data, which cover flame speed, ignition delay and flame species concentration.

Kumar et al. [44] conducted an experimental study on ammonia/hydrogen blends premixed flame speed under standard temperature and pressure (STP), and the reported test data are used for comparison of the three mechanisms identified previously in terms of flame speed prediction. As shown in Figure 1, the modelling results based on Nakamura’s mechanism agree better with the measurement results than the other two, particularly when using ammonia addition by energy at \(80\%\).

The measured ignition delay time of \(NH_3/O_2/Ar\) mixtures at \(1.4, 11, 30 \, atm\) are obtained from the shock-tube experiments reported by Mathieu [40], where the ignition delay time was defined as the time between the passage of the reflected shock wave and the intersection of the slope of hydroxyl radical \((OH^+)\) and the zero-concentration horizontal line. Figure 2 indicates that Nakamura’s mechanism shows the best agreement with the experimental results, while Otomo’s mechanism slightly overestimates the results, and apparent underestimation is found in the modelling results based on Duynslaegher’s reduced mechanism.

In Figure 3, the mole fraction profiles of the calculated flame species using three mechanisms are compared with the measured data of a low pressure \((0.05 \, bar)\) \(NH_3/H_2/O_2/Ar\) premixed burner-
stabilized flame (\(0.21NH_3/0.13H_2/0.21O_2/0.45Ar\)) reported by Duynslaegher [35]. The mole fraction profiles of \(NH_3\), \(H_2\) and \(N_2\) are satisfactorily predicted by all three mechanisms. Duynslaegher’s reduced mechanism can better capture the measure profiles of \(NO\), \(N_2O\) and \(NH_3\), which are key species for \(NO_x\) formation, although the peak mole fractions of \(NO\) and \(N_2O\) are still underestimated.

![Figure 1](image1.png)

**Figure 1.** Comparison between experimental and computational premixed \(NH_3/H_2\) laminar flame speed when \(E%NH_3\) is (a) 20%, (b) 50% and (c) 80%. Experimental results by Kumar et al. [44] plotted in conjunction with computational results using Nakamura’s mechanism [42], Otomo’s mechanism [31] and Duynslaegher’s reduced mechanism [34].

![Figure 2](image2.png)

**Figure 2.** Comparison between experimental and computational ignition delay time for the case of \(0.01143NH_3/0.00857O_2/0.98Ar\) (\(\phi = 1.0\)) at (a) 1.4 atm, (b) 11 atm and (c) 30 atm. Experimental results by Mathieu et al. [40] plotted in conjunction with computational results using Nakamura’s mechanism, Otomo’s mechanism and Duynslaegher’s reduced mechanism.

![Figure 3](image3.png)

**Figure 3.** Comparison between experimental and computational concentration profiles of (a) \(NO\), \(NH_3\), \(N_2O\) and (b) \(NH_3\), \(H_2\), \(N_2\) for the case of \(0.21NH_3/0.13H_2/0.21O_2/0.45Ar\) at 0.05 bar. Experimental results by Duynslaegher et al. [35] plotted in conjunction with computational results using Nakamura’s mechanism, Otomo’s mechanism and Duynslaegher’s reduced mechanism.
Based on the comparison above, Nakamura’s and Otomo’s mechanisms have better agreement with the experimental data in the study of premixed $\text{NH}_3/\text{H}_2$ laminar flame speed and ignition delay. Although Duynslaegher’s reduced mechanism can better predict flame species concentration, apparent deviations from the measured results are found in ignition delay. By comparing the computational results of NO concentration, the model with Nakamura’s mechanism predicts better than that of Otomo’s mechanism. Therefore, Nakamura’s detailed reaction kinetic mechanism is chosen to conduct a parametric analysis of premixed $\text{NH}_3/\text{H}_2$ combustion characteristics under LEG operational conditions.

4. Results and Discussion

4.1. Premixed Laminar Flame Speed

Figure 4 illustrates the effects of hydrogen blending ratio and equivalence ratio on the premixed flame speed of $\text{NH}_3/\text{H}_2$ blends. The flame speed increases significantly and non-linearly as more hydrogen is introduced in the blends. This behaviour can be attributed to the much higher reactivity and higher flame speed of hydrogen than ammonia. As ammonia is gradually replaced by hydrogen, the latter takes the dominant role in the oxidation reaction. However, the high hydrogen concentration in $\text{NH}_3/\text{H}_2$ mixtures should be avoided, as high reactivity of hydrogen could result in boundary layer flashback [21]. Moreover, it can be observed that the flame speed increases steadily at fuel-lean conditions, reaches the peak when the equivalence ratio is around $1.1 - 1.2$, and then decreases at fuel-rich conditions. The premixed flame speed reaches approximately $34 \text{ cm/s}$ when equivalence ratio and hydrogen blending ratio are $1.1$ and $0.4$, respectively. Under the same conditions, the premixed flame speed of pure methane is $\sim 37 \text{ cm/s}$ [45] while the flame speed of pure hydrogen is over $200 \text{ cm/s}$ [25]. This suggests that it is practical to apply a $\text{NH}_3/\text{H}_2$ blend with an equivalence ratio of $1.1 - 1.2$ to realize stable combustion within the combustor.

![Figure 4](image)

Figure 4. Flame speed as a function of equivalence ratio in various hydrogen fractions at initial temperature of $300 \text{ K}$ and pressure of $1 \text{ bar}$.

Figure 5 demonstrates the effects of initial temperature and initial pressure on the premixed flame speed of the $\text{NH}_3/\text{Air}$ and $\text{NH}_3/\text{H}_2/\text{Air}$ ($x%\text{H}_2 = 0.3$) mixtures. As temperature has dominant effects on chemical reaction rate, flame speed increases rapidly with the increase in initial temperature, and the growth rate is steeper under a higher initial temperature. As the initial temperature rises from $300 \text{ K}$ to $700 \text{ K}$, flame speed is increased by around $40 \text{ cm/s}$ under atmospheric pressure. With the presence of $30\%$ hydrogen in fuel, the flame speed is increased by around $90 \text{ cm/s}$. On the contrary, flame speed is reduced as initial pressure goes up. At higher pressures, the thickness of the flame increases and the burning speed slows down. It is also notable in $\text{NH}_3/\text{H}_2/\text{Air}$ combustion that when the initial pressure increases from $1$ to $5 \text{ bar}$, the flame speed decreases by almost the same as the initial pressure increases from $5$ to $20 \text{ bar}$. This indicates that the flame speed is less sensitive to initial pressure under high pressure conditions, which is more typical under engine working conditions.

As reported, the reactions of $\text{NH}_3 + H = \text{NH}_2 + H_2$, $\text{NH}_3 + OH = \text{NH}_2 + H_2O$, $\text{NH}_3 + O = \text{NH}_2 + OH$ dominate during the beginning of ammonia oxidation [44]. Mole fraction profiles of $H$, $O$, $OH$
radicals in the flames with an equivalence ratio of 1.1 are presented in Figure 6. With the increase in hydrogen blending ratio, the concentrations of $H$, $O$, $OH$ are all increased—in particular, the radical of $H$ and $OH$. This means that the $OH$ radical has a greater contribution to increasing the flame speed of the $NH_3/H_2$ mixture. Similar behaviour is also found for fuel-lean and stoichiometric conditions [26,44].

**Figure 5.** Premixed $NH_3/H_2/Air$ flame speed as a function of initial temperature under 1, 5, 10, 15, 20 bar ($\phi = 1.10$).

**Figure 6.** Radicals profiles of $O$, $H$, $OH$ with hydrogen blending ratio at (a) 0.0, (b) 0.2, (c) 0.4 and (d) 0.6 ($\phi = 1.10$).

### 4.2. Ignition Delay Time

In Figure 7, the ignition delay time is plotted as a function of the hydrogen concentration in fuel mixtures with equivalence ratio at 0.5, 1.0, 1.7 and 2.0. For a certain equivalence ratio, ignition delay time is shortened considerably with hydrogen addition. For instance, the ignition delay time drops from $\sim 4 \times 10^{-3} s$ to $\sim 0.6 \times 10^{-3} s$ as the hydrogen blending ratio reaches a value of 0.2 at stoichiometric conditions. However, there is a minor difference that can be negligible as the hydrogen blending ratio is over 0.5. This indicates that by adding a higher reactive fuel component to ammonia, the ignition delay can be effectively shortened, which is beneficial to realize robust ignition. Similar behaviour was reported in $NH_3/CH_4$ combustion [46]. Furthermore, it can be found that the
influence of equivalence ratio on the ignition delay becomes weaker as more hydrogen is added to the fuel, which is also reported by Mathieu et al. [40], based on experiments of ammonia oxidation diluted by Ar.

As shown in Figure 8, the influence of the initial temperature (1040 − 1620 K) and initial pressure (1 − 20 bar) is discussed at the hydrogen blending ratio of 0.1 and 0.3, respectively. The ignition delay is prolonged significantly with the decreasing initial temperature. It also can be seen that a higher initial pressure can also reduce the ignition delay time, but the reduction rate decreases with the increasing initial pressure. Take the hydrogen blending ratio of 0.3, for instance: when the initial temperature is 1250 K (1/T = 0.8), a factor ($\tau_{1bar}/\tau_{5bar}$) of $\sim$3.13 is found between the ignition delay time under 1 and 5 bar, while a smaller factor ($\tau_{5bar}/\tau_{10bar}$) of $\sim$1.40 is found between 5 and 10 bar. When the initial pressure is over 10 bar, the ignition delay time is almost the same and the influence of initial pressure on ignition delay is negligible.

![Figure 7](image)

**Figure 7.** Ignition delay time as a function of hydrogen blending ratio for various equivalence ratios at initial temperature of 1400 K and pressure of 1 bar.

![Figure 8](image)

**Figure 8.** Ignition delay time as a function of $1000/\tau$ under the initial pressure of 1 − 20 bar at the hydrogen blending ratio of (a) 0.1 and (b) 0.3 ($\phi = 1.10$).

4.3. Mole Fractions of the Flame Species

As ammonia and hydrogen are zero-carbon fuels, carbon-related emissions will not be generated from ammonia/hydrogen blend combustion. However, NOx emissions become the most concerning issue as they can be produced not only through thermal NOx or Zel’dovich mechanism ($N_2 + O = NO + N$, $N + O_2 = NO + O$, $N + OH = NO + H$), but also through fuel-bound NOx mechanism. Further insight into the NOx formation mechanism within ammonia/hydrogen combustion can be derived from premixed flame structure study based on detailed kinetic modelling, which will provide valuable information for the development of a low NOx combustion system.

In Figures 9 and 10, the effects of equivalence ratio on flame species concentration are demonstrated with a hydrogen blending ratio of 0.1 and 0.3, respectively. It can be found that NO is the dominant component of NOx emissions, while the concentrations of NO2 and N2O are nearly zero in the post-flame zone. As the equivalence ratio increases, NOx emission is reduced apparently. This trend is even more explicit when the equivalence ratio is close to unity. For instance, with 10%
hydrogen in fuel, NO concentration decreases by a factor of 1.75 as ϕ increases from 1.05 to 1.10. Additionally, it is observed that NO concentration increases rapidly before reaching a steady state under a certain equivalence ratio (1.10 for 10% hydrogen in fuel and 1.15 for 30% hydrogen in fuel), while it firstly peaks and then drops above the equivalence ratio. Besides this, the apparent increase in H₂ in the post-flame zone is noted as the flame becomes richer. Thus, an optimized equivalence ratio exists considering high NOx emissions in fuel-lean flame and high H₂ emission in fuel-rich flame.

To better understand the observed phenomena, rate of production (ROP) analysis is carried out for lean (ϕ = 0.90) and rich (ϕ = 1.20) flames with 30% hydrogen in the fuel. As indicated in Figure 11, in the lean flame, NO is produced primarily through the reactions between HNO and H/OH radicals, and through the thermal-NO (Zel’dovich) mechanism reactions (N + O₂ = NO + O and N + OH = NO + H) to a lesser extent; the consumption of NO is mainly through NH₂ + NO reactions, and through the reverse reaction of the rate-limiting step of thermal-NO mechanism (N₂ + O = NO + N) to a lesser extent. In the rich flame, the contribution from HNO + OH to NO production reduces, while that from N + O₂ increases; the dominant role of the reverse thermal-NO mechanism (N₂ + O = NO + N) in NO reduction is noted and should be relevant to the drop in NO concentration in rich flames. As reported by Kobayashi [9], when the flame becomes richer, concentrations of O/H/OH radicals drop, which tends to inhibit NO formation from NH₂ oxidation via HNO. Meanwhile, the relative concentration of H in radical pool rises in rich flame, which promotes the reactions between NH₂ and H. As a result, abundant H₂ and N will be produced. The existence of more nitrogen atoms will promote the reactions of the aforementioned thermal-NO (Zel’dovich) mechanism, which aligns with the findings of ROP analysis.

Figure 9. Species profiles of NH₃, H₂, NO, NO₂, N₂O at (a) ϕ = 0.90, (b) ϕ = 1.00, (c) ϕ = 1.05, (d) ϕ = 1.10, (e) ϕ = 1.15 and (f) ϕ = 1.20 under initial temperature of 500 K and pressure of 1 bar for hydrogen blending ratio of 0.1.
Figure 10. Species profiles of \( \text{NH}_3, \text{H}_2, \text{NO}, \text{NO}_2, \text{N}_2\text{O} \) at (a) \( \phi = 0.90 \), (b) \( \phi = 1.00 \), (c) \( \phi = 1.05 \), (d) \( \phi = 1.10 \), (e) \( \phi = 1.15 \) and (f) \( \phi = 1.20 \) under initial temperature of 500 \( K \) and pressure of 1 bar for hydrogen blending ratio of 0.3.

Figure 11. Rate of production of \( \text{NO} \) at (a) \( \phi = 0.90 \) and (b) \( \phi = 1.20 \) (\%\( \text{H}_2 = 0.3 \)).

Figure 12 shows the flame structure under various hydrogen blending ratios in fuel, with equivalence ratio, initial temperature and pressure kept constant. With more hydrogen addition, \( \text{NO} \) concentration in the post-flame zone only shows a marginal increase. Despite the higher initial \( \text{H}_2 \) concentration in the high-hydrogen-ratio flame, the steady state \( \text{H}_2 \) concentration almost remains unchanged. Although the total rate of \( \text{NO} \) production should share similar behaviour to \( \text{NO} \) mole fraction, the contribution from certain reactions changes with the \( \text{H}_2 \) blending ratio. As shown in Figure 13, both the thermal \( \text{NO} \) mechanism and fuel-bound \( \text{NO} \) mechanism are active for \( \text{NO} \) production and reduction. However, the contribution from the thermal \( \text{NO} \) mechanism (\( \text{N} + \text{OH} \) for \( \text{NO} \) production and \( \text{N} + \text{NO} \) for \( \text{NO} \) reduction) increases as more \( \text{H}_2 \) is added into the fuel. The unchanged \( \text{H}_2 \) concentration in the post-flame zone can be attributed to the promoted \( \text{H}_2 \) consumption rate through the reaction between \( \text{H}_2 \) and \( \text{OH} \) radical when \( \text{H}_2 \) is added into the fuel, as indicated by Figure 14.
The effects of the initial temperature range of $300 - 700 \, K$ and the initial pressure range of $1 - 20 \, bar$ on flame structure are demonstrated in Figures 15 and 16, respectively. The hydrogen concentration in the fuel mixtures is set to 0.3, and the equivalence ratio is set to 1.10. The effect of the initial temperature is investigated at atmospheric pressure. Figure 15 indicates that $NO$ concentration in the post-flame zone decreases from $9.6 \times 10^{-8}$ to $4.61 \times 10^{-8}$ as the initial temperature rises from $300$ to $700 \, K$, although the increase in initial temperature is usually expected to promote the thermal $NO_\alpha$ formation. In a modelling study of a stoichiometric $NH_3 \ (60\%) / CH_4 \ (40\%)$ premixed flame by Xiao [46], $NO_\alpha$ emissions were found to rise with the increase in initial temperature. The unexpected reduction of $NO$ under high initial temperature condition can be explained based on the ROP analysis shown in Figure 17. In a lower inlet temperature, fuel-bound $NO$ is dominant in the production of nitric oxide, while thermal-$NO$ contributes less; the reverse thermal-$NO$ reaction is dominant in the consumption of nitric oxide, while the combination with $NH_3$ contributes less. The same behavior is also found in the case with a higher inlet temperature, but the rate of both production and consumption is decreased, which could be the reason for the reduction in $NO$ emission.

Figure 12. Species profiles of $NH_3$, $H_2$, $NO$, $NO_2$, $N_2O$ at the hydrogen blending ratio of (a) 0.0, (b) 0.04, (c) 0.1, (d) 0.2, (e) 0.5 and (f) 0.8 under initial temperature of $500 \, K$ and pressure of 1 bar for equivalence ratio of 1.10.

Figure 13. Rate of production of $NO$ at (a) $x\%H_2 = 0.0$, (b) $x\%H_2 = 0.2$ and (c) $x\%H_2 = 0.5$. 
Figure 14. Rate of production of $H_2$ at (a) $x\%H_2 = 0.0$, (b) $x\%H_2 = 0.2$ and (c) $x\%H_2 = 0.5$.

Figure 15. Species profiles of $NH_3$, $H_2$, $NO$, $NO_2$, $N_2O$ under the initial temperature of (a) 300 K, (b) 400 K, (c) 600 K and (d) 700 K at atmospheric pressure ($\phi = 1.10$).
Figure 16. Species profiles of \( NH_3, H_2, NO, NO_2, N_2O \) under (a) 1 bar, (b) 5 bar, (c) 10 bar, (d) 15 bar and (e) 20 bar at the initial temperature of 600 K (\( \phi = 1.10 \)).

Figure 17. Rate of production of \( NO \) at the inlet temperature of (a) 300 K and (b) 700 K.

The influence of initial pressure on \( NH_3/H_2 \) flames is studied, with the equivalence ratio, initial temperature and hydrogen blending ratio kept at 1.10, 600 K and 0.3, respectively. As shown in Figure 16, \( NO \) is reduced significantly as pressure increases, which agrees with the phenomena reported by other works [9]. This trend can be attributed to the consumption of \( NH_3 \) species via the promoted radical combination reactions under high pressure conditions [9]. It suggests that low \( NO_x \) emission is possible under high pressure conditions, which is practical for engine application. In addition, it is noticeable that the peak in \( NO_2 \) concentration in the reaction zone is increased at higher initial pressure.

5. Conclusions

To support the development of a novel linear engine generator (LEG), the characteristics of ammonia/hydrogen premixed combustion are studied by using a detailed chemical kinetics mechanism, which is selected among several mechanisms based on validation against the same sets of experimental data.

Extensive parametric analysis is carried out under LEG typical working conditions to study the effects of equivalence ratio (0.80 – 1.60), hydrogen blending ratio (0.0 – 0.6), initial temperature (300 – 700 K) and initial pressure (1 – 20 bar) on premixed laminar flame speed, ignition delay and key flame species concentrations.

It is shown that an optimized equivalence ratio exists around 1.10 – 1.20 for \( NH_3/H_2 \) premixed flames to realize stable and efficient combustion as well as low \( NO_x \) emission. Ignition delay is reduced with the increase in hydrogen blending ratio, initial temperature and initial pressure. At a certain initial temperature and initial pressure, the effects of hydrogen blending ratio can be negligible for over 50% hydrogen in the fuel. Under higher pressure (> 10 bar), the initial pressure has a minor influence on the ignition delay reduction. It is also found that the high-pressure, high-
temperature environment is favourable for low NO\textsubscript{x} emission from ammonia/hydrogen combustion, which implies the potential for a low NO\textsubscript{x} LEG fuelled by ammonia/hydrogen.

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