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DOI:

[10.1016/j.ijhydene.2024.02.281](https://doi.org/10.1016/j.ijhydene.2024.02.281)

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*Document Version*

Publisher's PDF, also known as Version of record

*Citation for published version (Harvard):*

Zhang, F, Zhang, G, Wang, Z, Wu, D, Jangi, M & Xu, H 2024, 'Experimental investigation on combustion and emission characteristics of non-premixed ammonia/hydrogen flame', *International Journal of Hydrogen Energy*, vol. 61, pp. 25-38. <https://doi.org/10.1016/j.ijhydene.2024.02.281>

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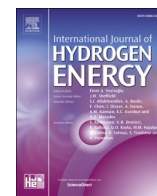
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# Experimental investigation on combustion and emission characteristics of non-premixed ammonia/hydrogen flame

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## ARTICLE INFO

Handling editor: J Ortiz

### Keywords:

Ammonia-hydrogen combustion  
Non-premixed flame  
Flame stability  
Emissions  
Chemiluminescence

## ABSTRACT

The use of ammonia and hydrogen as fuels to decarbonise the heavy transport industry has attracted worldwide attention. In this work, a swirl-enhanced combustion test rig has been built to investigate the combustion and emission characteristics of non-premixed ammonia/hydrogen flames. The blowoff limits were first examined to ensure that a range of stable flames could be achieved. Emissions containing nitric oxide (NO), nitrous oxide (NO<sub>2</sub>), ammonia slip (NH<sub>3</sub>) and unburnt hydrogen (H<sub>2</sub>) were then measured with a variety of global equivalence ratios, hydrogen blending ratios, inlet gas temperature, swirl numbers and combustion chamber insulation conditions. The structure of non-premixed ammonia/hydrogen flames and the relationship between excited hydroxyl radicals (OH\*) and NO emission were revealed using OH\* chemiluminescence profiles. The stable non-premixed ammonia/hydrogen flames were achieved when the hydrogen blending ratio was greater than 20%. The optimal emission performance was achieved under stoichiometric conditions, as determined by combustion efficiency. The insulation conditions of the combustor wall played a key role in the emission results, as significant growth of NO and NO<sub>2</sub> as well as a reduction of ammonia slip were found. Although a lower swirl number improved the flame stability range, the increases in NO and NO<sub>2</sub> emissions were observed.

## 1. Introduction

Electric vehicles introduce a new trend on the market of passenger cars and light-duty vehicles. However, the hard-to-decarbonise heavy transport sectors would not be possible to be solely driven by batteries due to their application backgrounds, such as heavy load and long distance. To decarbonise heavy transports, ammonia and hydrogen are considered as promising alternative fuels. Ammonia and hydrogen are both carbon-free, which avoids carbon emissions during the combustion of ammonia and hydrogen. As ammonia has been extensively applied as fertilizer and cleaning chemicals in agriculture, the production, storage and distribution processes of ammonia have been well established. Although the dominate way of producing ammonia in the current stage still relies on the conventional Haber-Bosch method, many projects on green ammonia production are ongoing [1–4]. If the production of green ammonia can be commercially introduced, the carbon footprint in the life cycle is expected to be significantly reduced. Owing to the physical properties similarity as propane, it is easy for ammonia to be stored as

liquefied ammonia under approximately  $-34\text{ }^{\circ}\text{C}$  at atmospheric pressure and compressed ammonia in an around 10 bar tank at room temperature. Given these advantages, ammonia, as one of the highly competitive hydrogen carriers, is worthy of in-depth research and discussion as a green alternative fuel.

Nevertheless, the direct adoption of ammonia as a fuel has encountered challenges. The narrow flammability (around 18% – 28% fuel mole fraction) [5], the high ignition temperature (903 K) [6] and the low laminar flame speed ( $\sim 0.07\text{ m/s}$  @NTP (normal temperature and pressure)) [7–9] of ammonia make it difficult to be ignited and maintain robust flame. Due to the high reactivity and high laminar flame speed of hydrogen, it is considered to be an ideal enhancer for ammonia flames [10–14]. Joo et al. [10] studied the flame stability of premixed ammonia/hydrogen flames in a tube type combustor at NTP. The findings concluded the stability limits increased as more hydrogen was blended with ammonia. The comparison of combustion limits between premixed ammonia/hydrogen flames and premixed ammonia/methane flames was conducted experimentally [11]. It was concluded that hydrogen

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<https://doi.org/10.1016/j.ijhydene.2024.02.281>

Received 7 November 2023; Received in revised form 9 January 2024; Accepted 21 February 2024

Available online 28 February 2024

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addition showed a wider combustion limit than methane addition into premixed ammonia flames. Furthermore, elevating the inlet temperature had a great impact on expanding the combustion limits of premixed ammonia/hydrogen flames and premixed ammonia/methane flames. Wei et al. [15] analysed the lean blowoff characteristics of premixed ammonia/hydrogen combustion in a gas turbine combustor. The flame structure and flow field results showed that there were less large-scale and more small-scale wrinkles in the flame front when 10% hydrogen was premixed with 90% ammonia. The promoted combustion intensity facilitated the flame stability. A generic swirl burner was used to test the stability limits of premixed ammonia/hydrogen flames up to 5 bar in KAUST (King Abdullah University of Science and Technology) [12]. It indicated that the stability range was broadened by increasing the pressure, however, the impact was inapparent. Moreover, the results showed that 40%  $\text{NH}_3$ /60%  $\text{H}_2$  flame was surprisingly stabilised even at very lean conditions (the equivalence ratio was at approximately 0.3). Zhen et al. [16] studied the flame stability of premixed ammonia/hydrogen Bunsen flames for impingement heating applications. It was found that there exists a critical value of hydrogen addition to obtain a stable premixed ammonia/hydrogen Bunsen flame for a given Reynolds number of the fuel/air mixture jet and equivalence ratio. It is evident that a majority of scholars have initiated research on the combustion characteristics of premixed ammonia and hydrogen, however, there are few studies on the flame stability limits of non-premixed ammonia/hydrogen flames so far.

Emissions are another concern in the combustion of ammonia and hydrogen. While carbon emissions are effectively reduced, high-temperature conditions, particularly when ammonia with nitrogen atoms is present, can lead to fuel-bound  $\text{NO}$  formation. Valera-Medina et al. [17] carried out a 50%  $\text{NH}_3$ /50%  $\text{H}_2$  premixed flame under fuel-lean conditions ( $\varphi$ : 0.41–0.56) in Gas Turbine Research Centre (GTRC) in Cardiff University under NTP. The  $\text{NO}_x$  emission was high in fuel-lean conditions and reached over 1000 ppmv when the equivalence ratio was 0.52 and 0.56. Later, the same research group [18] investigated premixed 70%  $\text{NH}_3$ /30%  $\text{H}_2$  and 60%  $\text{NH}_3$ /40%  $\text{H}_2$  combustion under fuel-rich conditions, but it was noticed that 60%  $\text{NH}_3$ /40%  $\text{H}_2$  flame tended to flashback. It was observed that the  $\text{NO}_x$  emission decreased when the equivalence ratio increased from 1.0 to 1.4. Experimental results of ammonia slip were not published in this work though.  $\text{NO}$  emission from premixed ammonia flames and premixed ammonia/hydrogen flames was investigated in a porous combustor by Chen et al. [19]. When the equivalence ratio increased from 1.0 to 1.2,  $\text{NO}$  emission decreased by 98.3% to 23.6 ppm. The calculated conversion rate of nitrogen atoms in the fuel to  $\text{NO}$  was very low under fuel-rich conditions for both premixed ammonia flames and premixed ammonia/hydrogen flames. Compared to pure ammonia combustion, the conversion rate of  $\text{NO}$  was only 0.42% higher when 30% hydrogen was added to the fuel. Recently, Tong et al. [20] analysed premixed ammonia/hydrogen/oxygen combustion by building a 3D meso-scale combustor model. The model was employed to investigate the  $\text{NO}$  emissions under different hydrogen additions (0–20%). The results presented that increasing hydrogen addition to 15% resulted in  $\text{NO}$  emission reduction at a given low inlet velocity (less than 0.4 m/s) under stoichiometric conditions. However, as the hydrogen addition increased to 20%,  $\text{NO}$  emission was increased. The relationship between  $\text{NO}$  concentration and  $\text{OH}^*$  in premixed ammonia/hydrogen flames was explored under fuel-lean conditions by Zhu et al. [21]. The local  $\text{NO}$  concentration and  $\text{NO}$  emission were found to align well with the local  $\text{OH}^*$  chemiluminescence intensity in the combustor when the equivalence ratio was at 0.40–0.85 and ammonia concentration in the fuel was greater than 0.25. It indicates  $\text{OH}^*$  chemiluminescence imaging technology can be employed to qualitatively visualise local  $\text{NO}$  concentration trends or  $\text{NO}$  emission trend in premixed ammonia/hydrogen flames which is a very cost-effective approach compared to local  $\text{NO}$  detection using  $\text{OH}$ -PLIF technology. Pugh et al. [22] investigated  $\text{OH}^*$ ,  $\text{NH}_2^*$ ,  $\text{NH}^*$  chemiluminescence in premixed and non-premixed

ammonia/air flame as well as partially premixed 70%  $\text{NH}_3$ /30%  $\text{H}_2$  flame with the same net thermal power at 25 kW. There were two types of configurations of partially premixed ammonia/hydrogen flame. One is premixed ammonia and air entering from outer swirling flow and hydrogen entering from the central flow, which was called  $\text{H}_{2\text{Diff}}$ . The other is premixed hydrogen and air entering from outer swirling flow and ammonia entering from the central flow, which is called  $\text{NH}_{3\text{Diff}}$ . By comparing the premixed ammonia flame,  $\text{H}_{2\text{Diff}}$  and  $\text{NH}_{3\text{Diff}}$ , it was noted that for the case of  $\text{NH}_{3\text{Diff}}$ ,  $\text{NH}_2^*$  chemiluminescence intensity was the greatest and  $\text{OH}^*$  and  $\text{NH}^*$  chemiluminescence intensities were the lowest among three. Meanwhile,  $\text{NO}$  emission produced in  $\text{NH}_{3\text{Diff}}$  showed the lowest value. When both the temperature and pressure were lifted, a similar relational phenomenon was observed, i.e.,  $\text{NH}_2^*$  intensity increased,  $\text{OH}^*$  intensity and  $\text{NO}$  emission decreased. Moreover, a significant reduction of  $\text{NO}$  was observed in non-premixed ammonia/air flame compared to premixed ammonia/air flame, with 200 ppmvd@15%  $\text{O}_2$  under stoichiometric condition, but with a high ammonia slip. Nonetheless, it is still unclear about the emission characteristics of non-premixed ammonia/hydrogen flames, especially with different hydrogen blending ratios and swirlers. Experimental data on  $\text{NO}_x$  emission has been reported in many studies, but it is equally important to have unburnt ammonia and hydrogen to provide solid evidence for chemical mechanism development and model validation.

It is essential to conduct a thorough investigation on the combustion and emission characteristics of non-premixed ammonia/hydrogen flames in a swirl-enhanced combustor. To the best of our knowledge, this area has not been extensively explored. In this paper, the stable range conditions of non-premixed ammonia/hydrogen flame were first identified by measuring the blowoff limits. Secondly, emissions including  $\text{NO}_x$  and unburnt ammonia and hydrogen at the exhaust were analysed with various global equivalence ratios, hydrogen blending ratios, inlet gas temperature, combustor wall conditions and swirl numbers.  $\text{OH}^*$  chemiluminescence images were then used to reveal the flame structure. For the quantitative analysis,  $\text{OH}^*$  intensity was calculated based on pixel data of the  $\text{OH}^*$  chemiluminescence images. The experimental results are anticipated to make a significant contribution to the validation of computational fluid dynamics models and to the development of ammonia combustion mechanisms. In turn, it provides guidance for the development of ammonia/hydrogen fuelled heat engines for use in the heavy transport sectors.

## 2. Methodology

### 2.1. Experimental test rig

All the experiments were conducted in the Future Engines & Fuels Laboratory at the University of Birmingham. The non-premixed combustion chamber is cylindrical with a detachable quartz glass window. The inner diameter of the combustion chamber is 150 mm with 310 mm in length. The optical window is in rectangular shape (249 mm  $\times$  53 mm). Fuel blend enters the combustion chamber through an 8-hole fuel nozzle while air enters through a swirler on the annulus of the fuel nozzle. The injection angle of the fuel nozzle is determined to be 45° in the present work as it was concluded by Okafor et al. [23] that 45° shows the best performance of mixing fuel and air near the inlet in ammonia combustor among four different injection angles (0°, 30°, 45°, 60°). Swirl-enhanced strategy is utilised for air injection to promote the ammonia flame robustness. Fig. 1 demonstrates the swirler diagram with a vane angle,  $\alpha$ . There are three different swirlers available for investigation with the vane angle at 40°, 45°, 50°. The technical data of the swirlers is illustrated in Table 1. Swirl number ( $S$ ) is used to represent the strength of the swirling flow. It is determined as the ratio of the axial flux of angular momentum to the axial flux of the axial momentum [24]. The swirl number is calculated using the equation [25]:

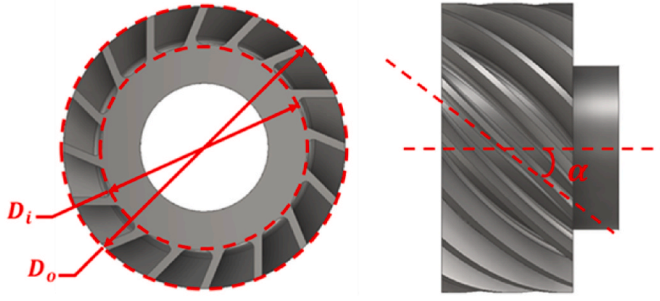


Fig. 1. Swirler.

Table 1  
Swirler details.

Swirler	$\alpha$ ( $^\circ$ )	$D_i$ (mm)	$D_o$ (mm)	Number of vanes (–)	$S$ (–)
$S(a)$	40	32	44	16	0.73
$S(b)$	45	32	44	16	0.87
$S(c)$	50	32	44	16	1.04

$$S = \frac{2}{3} \cdot \frac{1 - \left(\frac{D_i}{D_o}\right)^3}{1 - \left(\frac{D_i}{D_o}\right)^2} \cdot \tan \alpha \quad (1)$$

where  $D_i$ ,  $D_o$  and  $\alpha$  represent the inner diameter, the outer diameter of the swirler and the vane angle, respectively. In the present work, swirl number is changing between 0.73, 0.87 and 1.04, which is sufficient (above 0.6) to obtain a strong swirling flow [26–28].

Fig. 2 illustrates the schematic diagram of experiment test rig, which spans two test rooms and two control rooms. It is comprised primarily of gas cylinders, air compressor, mass flow controllers (MFCs) for each gas, flame arrestors, flame mixer, in-line heating pipes with their temperature controllers, combustion chamber, thermocouples, purge system, high speed COMS camera, heated sample lines, an emission multi-gas analyser for  $NO_x$  and  $O_2$  measurements, a gas chromatography for  $NH_3$  and  $H_2$  measurements, a data acquisition device and a computer.

Ammonia (99.98% purity) and hydrogen (99.99% purity) were supplied in pressurised gas cylinders from BOC company. The flow rate of ammonia and air were controlled by Aalborg™ MFC model GFC57S-BDN2 and GFC67S-VEN2, respectively, with the accuracy at  $\pm 3.0\%$  of set point for 0 – 20% of full-scale range scale and  $\pm 1.5\%$  of set point for 20 – 100% of full-scale flow. The MFC for hydrogen is model GFC47S-VDN6 with the accuracy within  $\pm 1.0\%$  of set point for full-scale flow. The ignition system is composed of a power supply, capacitor discharge ignition, a pulse generator, an ignition coil and two electrodes. Two electrodes are made of stainless steel with sharpened tip at the front. The negative electrode is mounted on the inlet plate of the combustor. The positive electrode is inserted from the bottom plate of the combustor. The electrodes gap is approximately 3 mm.

The total input thermal power ( $P$ ) was maintained at 8 kW throughout the test cases. The mass flow rate of fuel blends ( $\dot{m}_{fuel}$ ) was calculated based on the total input thermal power and LHVs (Lower Heating Values) of the individual fuels using the equation as follows:

$$\dot{m}_{fuel} = \frac{P}{LHV_{NH_3} \times x[NH_3] + LHV_{H_2} \times x[H_2]} \quad (2)$$

where  $LHV_{NH_3}$ ,  $LHV_{H_2}$  denote the LHVs of ammonia and hydrogen, which is 18.8 MJ/kg and 120 MJ/kg, respectively;  $x[NH_3]$  and  $x[H_2]$  represent the mole fraction of ammonia and hydrogen in the fuel blend. Hydrogen blending ratio ( $x\% H_2$ ) is calculated by dividing the concentration of hydrogen in the total fuel blend, shown as:

$$x\% H_2 = \frac{x[H_2]}{x[H_2] + x[NH_3]} \quad (3)$$

The global equivalence ratio ( $\phi_{global}$ ) is the ratio of the fuel-air ratio to the stoichiometric fuel-air ratio using the equation below and controlled by adjusting the mass flow rates of the fuel blend and air:

$$\phi_{global} = \frac{\dot{m}_{fuel}/\dot{m}_{air}}{(\dot{m}_{fuel}/\dot{m}_{air})_{st}} \quad (4)$$

where  $\dot{m}_{fuel}$  and  $\dot{m}_{air}$  represents the mass flow rate of fuel and air respectively;  $(\dot{m}_{fuel}/\dot{m}_{air})_{st}$  represents the fuel-air ratio under the stoichiometric condition.

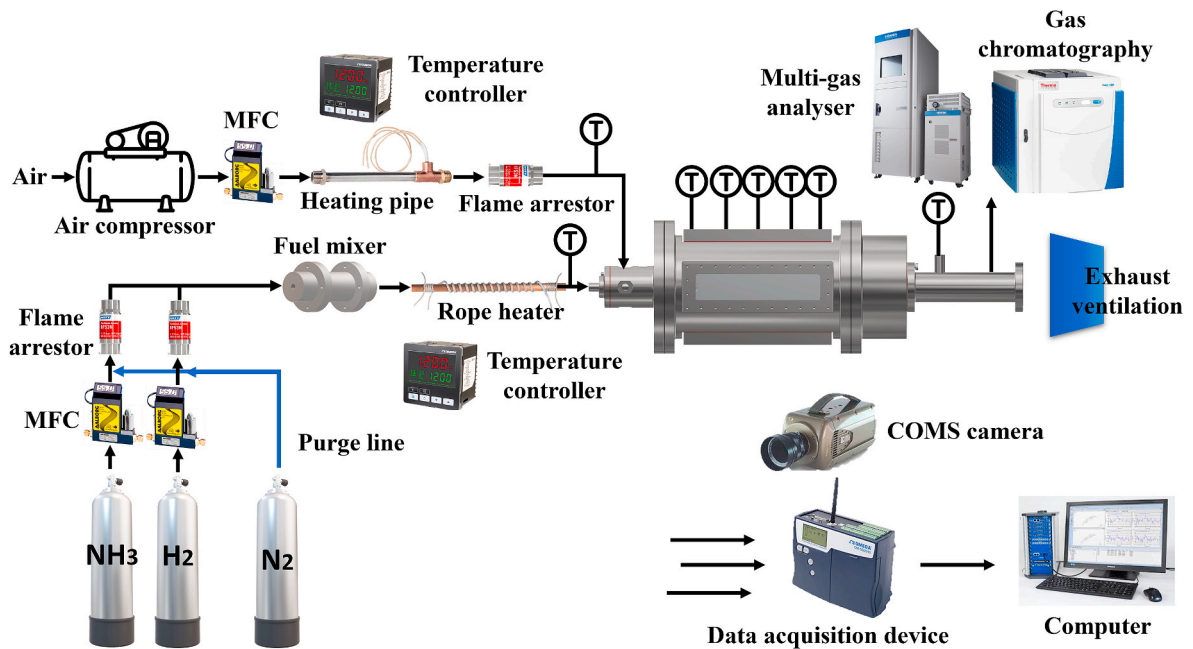


Fig. 2. Experiment test rig diagram.



## 2.2. Blowoff limits measurement

The blowoff limit is a critical parameter to ensure the range of a reliable flame. It is identified by gradually increasing the air flow rate by approximately 2% every 20 seconds while keeping the fuel flow rate unchanged until the flame is no longer sustained [29]. The global equivalence ratio at the blowoff limit is therefore calculated based on the measured air flow rate when the flame becomes stable and when the flame blows off. The blowoff limit of ammonia and hydrogen was investigated at different hydrogen blending ratios, inlet temperatures and swirl numbers so as to first gain the knowledge of the conditions of non-premixed ammonia/hydrogen flame stability. This would aid in the subsequent analysis on emission characteristics and  $OH^*$  chemiluminescence. The experimental conditions for examining the flame stability are shown in Table 2.

## 2.3. Emission measurement

Horiba multi-gas analyser (MEXA-7100DEGR) was performed to measure  $NO$  and  $NO_2$  emissions and oxygen at the exhaust. The repeatability is within  $\pm 0.5\%$  of the full scale for zero point or within  $\pm 0.5\%$  of the readings for span point. Thermo Fisher Trace 1300 gas chromatography with Pulse Discharge helium ionisation Detector (PDD) and Thermal Conductivity Detector (TCD) was utilised to obtain the concentration of unburnt ammonia and hydrogen at the exhaust. For both types of detectors, the repeatability of typical retention time is less than 0.0008 min and the repeatability of the typical peak area is less than 0.5% relative standard deviation. The sample gas was collected from the centreline of the exhaust duct through a heating pipe. The temperature of the gas in the heated sample pipe was kept at 193 °C for Horiba multi-gas analyser and 140 °C for gas chromatography to prevent the water condensation and aqueous dissolution of ammonia. The heated sample pipe was coated with functionalised hydrogenated amorphous silicon to avoid adsorption of ammonia and deliver a faster response in the measurement. The  $NO_x$  emissions from non-premixed ammonia/methane using the current test rig were validated with the single-staged results of premixed ammonia/methane flames in the literature [30]. The trends of  $NO$  and  $NO_2$  showed satisfactory agreements with the literature results. The effects of global equivalence ratios, hydrogen blending ratios, inlet temperature, swirl numbers and combustion chamber wall conditions on emissions of non-premixed ammonia/hydrogen combustion were conducted in the swirl-enhanced combustor. The experimental conditions are elaborated in Table 3. All data were collected while emission levels remained stable. As ammonia and hydrogen were burnt under different global equivalence ratios, oxygen normalisation was applied to compare the emissions fairly. According to the regulation in the UK [31,32], all the emission results were normalised to the standard oxygen concentration basis at 15% using the equation:

$$ppm_{norm} = ppm_{mea} \left[ \frac{20.95 - 15}{20.95 - O_{2mea}} \right] \quad (5)$$

where  $ppm_{norm}$  and  $ppm_{mea}$  indicate the normalised emission concentration and measured emission concentration, respectively;  $O_{2mea}$  indicate the measured oxygen concentration in the exhaust gases [31].

The error bar was calculated using the standard deviation equation

**Table 2**

The experimental conditions for blowoff limit test.

Factor	Range
Air flow rate	1.37 – 6.87 g/s
Hydrogen blending ratio	20%, 30%, 40%
Fuel inlet temperature	300, 400, 475 K
Air inlet temperature	300, 400, 435 K
Swirl number	0.73, 0.87, 1.04

**Table 3**

Experimental conditions for emission measurements/ $OH^*$  chemiluminescence imaging.

Factor	Range
Global equivalence ratio	0.6 – 1.2
Hydrogen blending ratio	20%, 30%, 40%
Fuel inlet temperature	300, 325, 350, 400 K
Air inlet temperature	300, 325, 350 K
Swirl number	0.73, 0.87, 1.04
Combustion chamber wall condition <sup>a</sup>	with and without silicate ceramic fibre blanket insulation

<sup>a</sup>  $OH^*$  chemiluminescence imaging was performed without silicate ceramic fibre blanket insulation.

based on the five measurements:

$$\sigma = \sqrt{\frac{1}{5} \sum_{i=1}^5 (x_i - \bar{x})^2} \quad (6)$$

where  $x_i$  and  $\bar{x}$  represent the average value for each measurement and average value of the five times measurements, respectively. It indicates the uncertainty of the emission measurement at each point.

## 2.4. $OH^*$ chemiluminescence imaging measurement and image post-processing

The use of chemiluminescence imaging for investigating radicals of interest is non-invasive, cost-effective and has been applied to a variety of combustion applications with small optical access [33–36]. Fig. 3 demonstrates the optical system for capture  $OH^*$  chemiluminescence images. A high-speed Phantom® v710 CMOS camera was utilised with resolution of 1280 × 800, sample rate at 100 pps and exposure time at 9900  $\mu$ s. Hamamatsu Photonics C10880 image intensifier was fitted on the CMOS camera through the relay lens to intensify the chemiluminescence at low-light level and facilitate the image visualisation. The image intensifier gain was constant at 999 in all the measurements. As the  $OH^*$  chemiluminescence primarily occurs in the wavelength between 280 – 350 nm [37], UV lens (UV-Nikkor 105 mm f/4.5 lens) and narrowband filter (centre: 310 nm) are employed. In the present work,  $OH^*$  chemiluminescence images have been measured and post-processed in relation to the global equivalence ratios, hydrogen blending ratios, inlet temperatures and swirl numbers. Table 3 also reveals the experimental conditions for  $OH^*$  chemiluminescence imaging measurement.

For each case, a thousand images were collected and post-processed with the in-house developed MATLAB code. The raw chemiluminescence images were averaged and then background corrected with averaged background images. Minimum value correction and 2D 3 × 3 median filter were applied to remove noise. Since the images collected in the laboratory were line-of-sight and the combustion chamber provided equidistant boundary for the flames with the axisymmetric fuel nozzle and swirler, the Abel inversion algorithm [38, 39] was implemented to obtain a representative spatially resolved image of the  $OH^*$  chemiluminescence distribution assuming that the flames were axisymmetric. Namely, the Abel inversion algorithm was utilised to deconvolute each row of images into integrals using an expansion of cosine functions on the principle of Fourier analysis [38,39]. Considering the computation time, the expansion value of cosine functions was assigned to five in the present study. The processed time-averaged spatially resolved  $OH^*$  chemiluminescence images were presented with false colour mapping, as shown in Fig. 4(b). To quantify the  $OH^*$  chemiluminescence,  $OH^*$  intensity ( $I_{OH^*}$ ) was determined on a pixel-by-pixel basis in the image before the application of the Abel inversion algorithm using the formula as follows [40]:

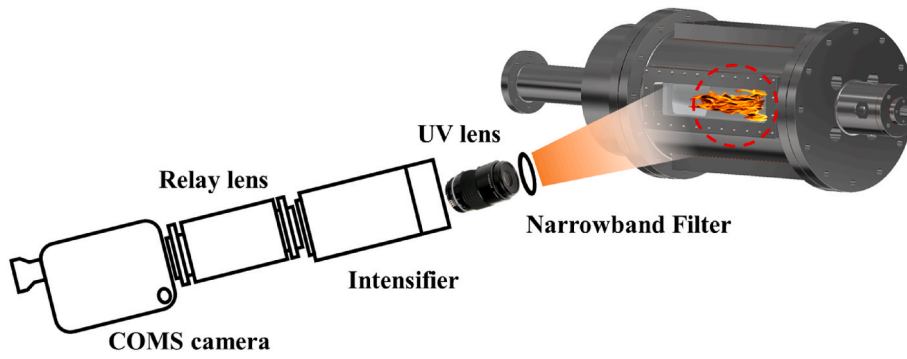
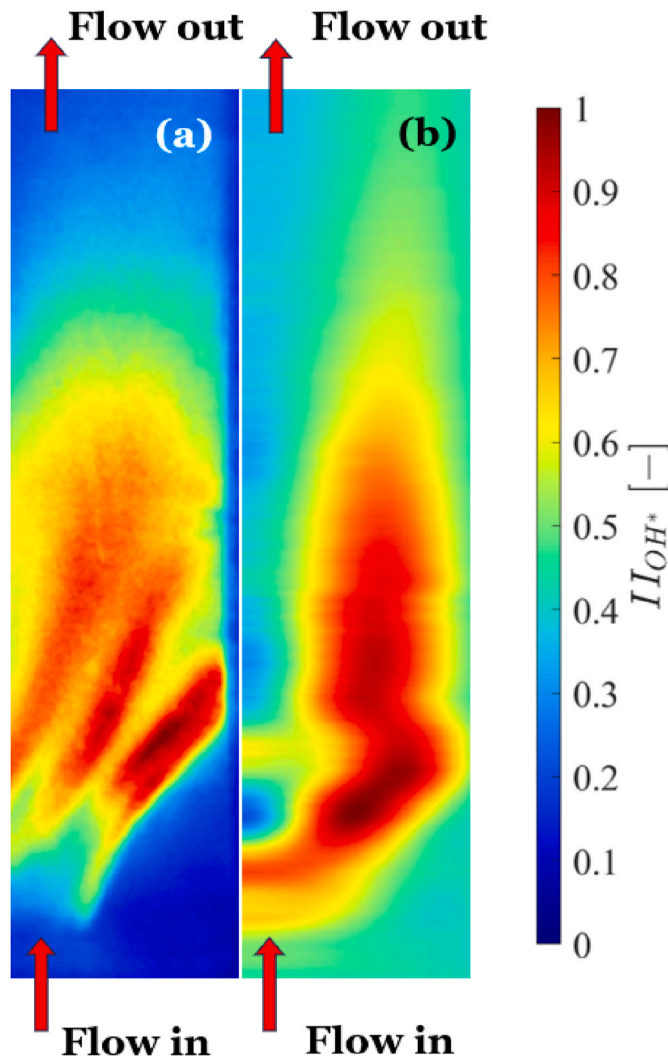


Fig. 3. Chemiluminescence visualisation setup.

Fig. 4. Time-averaged  $\text{OH}^*$  chemiluminescence distributions (a) before and (b) after Abel inversion algorithm.

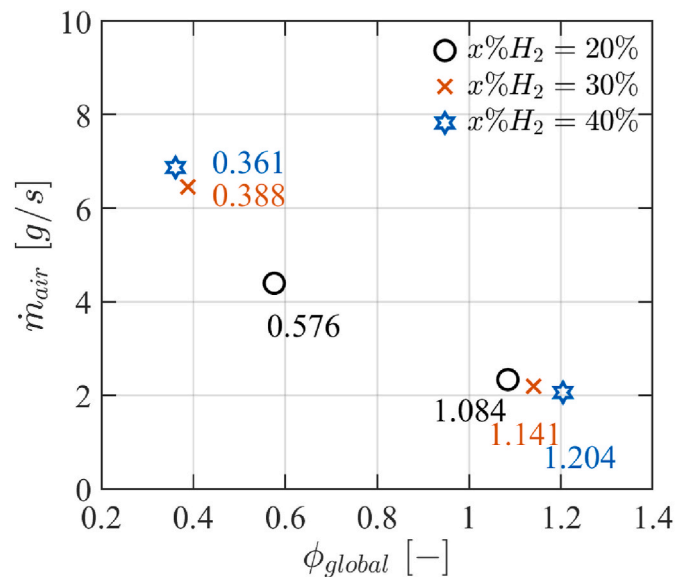
$$\Pi'_{\text{OH}^*} = \sum_{i=1}^{800} \sum_{j=1}^{1280} \bar{I}_{\text{OH}^*_{ij}} \quad (6)$$

### 3. Results and discussion

#### 3.1. Blowoff limits

It has been reported that ammonia flames have narrow flammability and low flame velocity, so it is important to determine the flame stability conditions when ammonia is used as the primary fuel and blended with a high flame velocity substance, hydrogen. Blowoff limits were examined with various hydrogen blending ratios and different swirlers at different gas inlet temperatures to ensure the flame stability condition and range (from lean blowoff limit to rich blowoff limit). The rich blowoff limit is defined as the minimum mass flow rate of air that can ignite and burn the fuel blends stably while the lean blowoff limit is defined as the maximum mass flow rate of air that keeps the flame stable before blowing away at a constant mass flow rate of fuel blends [41].

The effect of hydrogen blending ratios varying from 10% to 40% on the blowoff limits of non-premixed ammonia flames was conducted with swirler  $S(b)$  (swirl number: 0.87) at the inlet gas temperature of 300 K. It is observed that the flame was not stabilised when only 10% hydrogen is blended with ammonia. When the hydrogen blending ratio is above 20%, the flame became stable. The rich and lean blowoff limits of non-premixed ammonia flames with 20%, 30%, 40% hydrogen addition is

Fig. 5. Blowoff limits of non-premixed ammonia/hydrogen flames at the inlet temperature of 300 K ( $S = 0.87$ ).

plotted in Fig. 5. The region between the rich and lean blowoff limits is the region of stable non-premixed ammonia/hydrogen flame. It can be seen that hydrogen involvement expands the stable range of ammonia flames, and it has more pronounced effect on the lean blowoff limits than that on rich blowoff limits. For example, the stable range (global equivalence ratio) for 80%  $\text{NH}_3$ / 20%  $\text{H}_2$  flame is 0.576 – 1.084 while that for 70%  $\text{NH}_3$ / 30%  $\text{H}_2$  is 0.388 – 1.141. Due to the limitation of the maximum air flow rate, 60%  $\text{NH}_3$ / 40%  $\text{H}_2$  flame burns steadily at 6.870 g/s and its lean blowoff limit can be extended further. Increasing hydrogen addition is effective in igniting the fuel mixture, generating sufficient heat and maintain a stable flame, especially under lean-burn conditions due to better mixing with air; however, the effect is weakened under rich-burn conditions as less oxidant exits and combines with the fuel. The finding is comparable with the conclusions drawn from the lean blowoff limits of premixed ammonia/hydrogen flames [11]. The trend of lean blowoff limit in non-premixed ammonia/hydrogen flames is consistent with that of the premixed ammonia/hydrogen flames, i.e., the lean blowoff limit is gradually decreased as hydrogen blending ratio increases.

The effect of inlet temperature on the blowoff limits of non-premixed 60%  $\text{NH}_3$ / 40%  $\text{H}_2$  flame is presented in Fig. 6. It is noticed that the lean blowoff limits of non-premixed 60%  $\text{NH}_3$ / 40%  $\text{H}_2$  can be further extended because the flames under room temperature and elevated temperature are still robust with the maximum air mass flow rate available in the laboratory. Therefore, only the effect of inlet temperature on the rich blowoff limit can be observed. The ranges to achieve the stable flames are widened when rising the inlet temperature. For instance, the rich blowoff limit is extended from 1.204 to 1.290 when the inlet temperature rises from 300 K to 400 K and it extends further to 1.505 when the inlet air temperature rises to 435 K and the inlet fuel temperature rises to 475 K. A dimensionless number, Damköhler number ( $Da$ ), has to be greater than unity to obtain a stable flame [42]. Damköhler number is determined by the ratio of the characteristic mixing time to the characteristic chemical reaction time. An increase in the inlet temperature enhances the chemical reaction rate and burning rate, which reduces the chemical reaction time. As a result, Damköhler number is greater with a high inlet gas temperature and the stable range of the flame was extended.

Adding swirlers is generally considered as an effective way to increase the mixing ratio of fuel and oxidizer in the shear layer region. Therefore, in this work, three swirlers,  $S(a)$ ,  $S(b)$ ,  $S(c)$ , are employed to

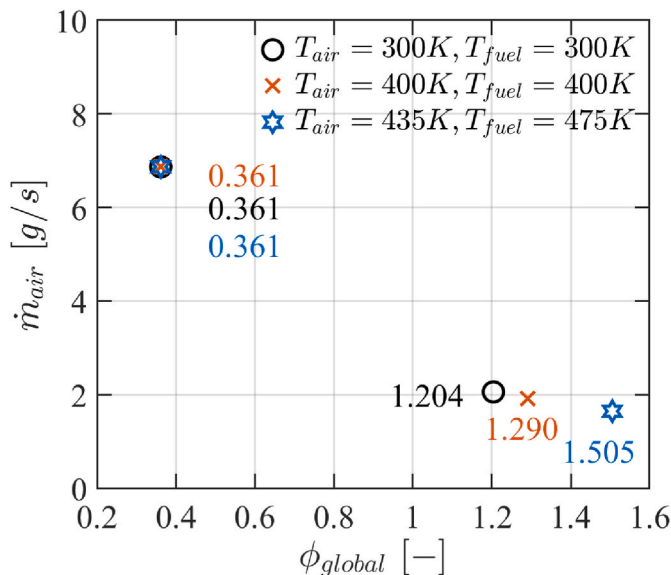


Fig. 6. Blowoff limits of non-premixed flames 60%  $\text{NH}_3$ / 40%  $\text{H}_2$  under different inlet gas temperature ( $S = 0.87$ ).

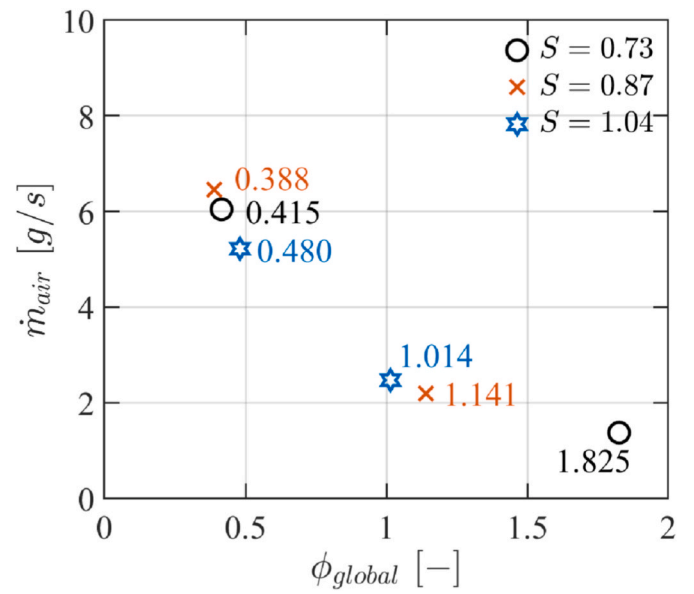


Fig. 7. Blowoff limits of non-premixed 70%  $\text{NH}_3$ / 30%  $\text{H}_2$  flames with different swirl numbers at the inlet temperature of 300 K.

investigate the effect of swirl numbers on the blowoff limits of non-premixed ammonia/hydrogen flames. Fig. 7 demonstrates the increase in the swirl number shortens the flame stability range considerably for non-premixed 70%  $\text{NH}_3$ / 30%  $\text{H}_2$  flames. The effect on the rich blowoff limits is more pronounced than the effect on the lean blowoff limits. The effect of swirl numbers on the blowoff limits of premixed ammonia/methane was also analysed by Zhang et al. [43]. It was reported that when the swirl number was changed between 0.42, 0.71, 1.27, the lean and rich blowoff limits almost stayed unchanged in premixed ammonia flames; however, an apparent extension on the lean blowoff limits was observed in premixed 50%  $\text{NH}_3$ / 50%  $\text{CH}_4$  flames.

### 3.2. Emissions

#### 3.2.1. Emission performance of different hydrogen blending ratios at different global equivalence ratios

Although no carbon emissions will be emitted from non-premixed ammonia/hydrogen combustion, nitric oxides, ammonia slip and unburnt hydrogen can be produced, all of which are harmful to the atmosphere. A clear knowledge of emission characteristics of non-premixed ammonia/hydrogen combustion is beneficial for emission control when designing the industrial applications powered by ammonia and hydrogen. Therefore, emissions of  $\text{NO}$ ,  $\text{NO}_2$ ,  $\text{NH}_3$ ,  $\text{H}_2$  are emphasised in this section and all the results have been normalised to the standard oxygen level, 15%(vol).

Fig. 8 depicts the effects of global equivalence ratios and hydrogen blending ratios on the emissions of non-premixed ammonia/hydrogen flames. When the hydrogen blending ratio is 20% and 30%, it can be seen that  $\text{NO}$ ,  $\text{NO}_2$  emissions increase when the global equivalence ratio rises from 0.6 – 0.8, reach the peak at the global equivalence ratio of 0.8 with about 100  $\text{ppmvd}@15\%\text{O}_2$  for  $\text{NO}$  emission and about 35  $\text{ppmvd}@15\%\text{O}_2$  for  $\text{NO}_2$  emission and decrease with further increase in the global equivalence ratio. A similar trend is also observed when the hydrogen blending ratio is 40%, but both  $\text{NO}$  and  $\text{NO}_2$  emissions are much higher at a low global equivalence ratio compared to the results with hydrogen blending ratio at 20% and 30%. The maximum  $\text{NO}$  and  $\text{NO}_2$  emissions reach approximately 350  $\text{ppmvd}@15\%\text{O}_2$  and 60  $\text{ppmvd}@15\%\text{O}_2$ , respectively. There are dramatic reductions for  $\text{NO}$  and  $\text{NO}_2$  emission when the global equivalence ratio is close to the unity.  $\text{NO}$  and  $\text{NO}_2$  emission were almost the same low value under fuel-rich conditions when hydrogen blending ratio was at 20%, 30% and 40%.

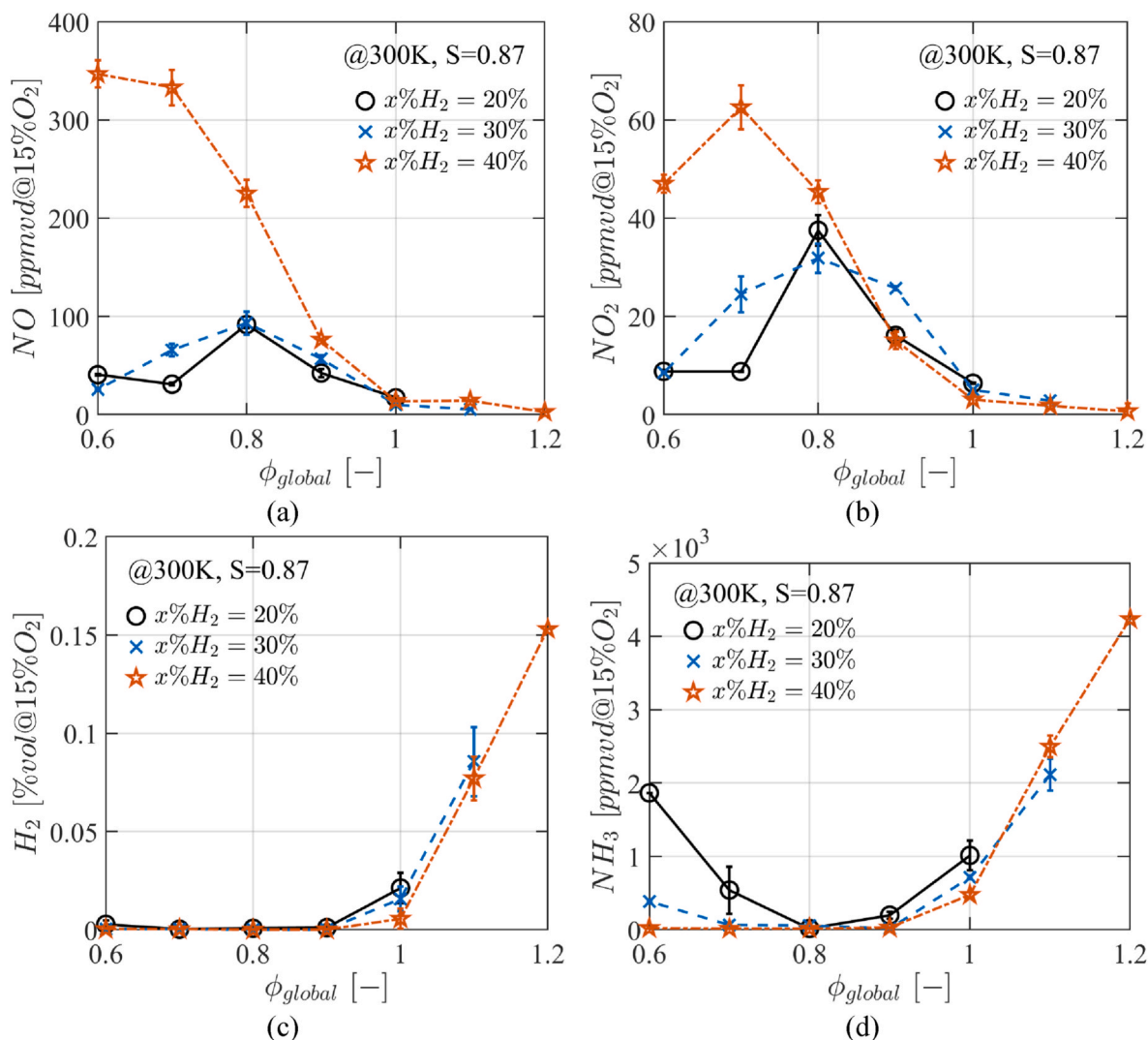


Fig. 8. Emissions of non-premixed ammonia/hydrogen flames versus various global equivalence ratios at different hydrogen blending ratios.

In ammonia combustion,  $NO$  and  $NO_2$  formations and consumptions are mainly associated with the flame temperature (thermal  $NO_x$  route) and concentrations of  $O/H/OH$  radicals in the flame (fuel-bound  $NO_x$  route). When the global equivalence ratio is far-lean, the flame temperature is very low which suppressed  $NO$  and  $NO_2$  production. As the global equivalence ratio increases but below the unity, fuel-bound  $NO_x$  ( $HNO + H = NO + H_2$ ,  $HNO + OH = NO + H_2O$ ) is the main pathway for  $NO_x$  formation. Meanwhile, the flame temperature is relatively higher which induced further  $NO_x$  emission caused by the thermal  $NO_x$  formation. This can be explained the reason of low emissions of  $NO$ ,  $NO_2$  found at a very low equivalence ratio and the increase of emissions with the global equivalence ratio increasing to one. Conversely, under fuel-rich conditions, the fuel-bound route for  $NO_x$  formation is inhibited as the concentrations of  $O/H/OH$  radicals are decreased. Instead, the thermal  $NO_x$  route for  $NO_x$  consumption is promoted and dominate to consume  $NO_x$ , resulting the reduction in  $NO$  and  $NO_2$  emissions when the global equivalence ratio is greater than one.

Although  $NO$  and  $NO_2$  emissions keep decreasing under fuel-rich conditions, the unburnt fuel including hydrogen and ammonia rises significantly. At the global equivalence ratio of 1.2, ammonia slip is over 4000 ppmvd@15% $O_2$  while unburnt hydrogen is over 0.15%vol@15% $O_2$  at the exhaust. Combustion efficiency is an indicator of the combustion completeness. The unburnt fuels and products in the exhaust gas are considered. Taking the combustion efficiency into account, the optimal

emission control occurred at stoichiometric condition. Relative low emissions including  $NO_x$  and unburnt fuel were also found at far-lean global equivalence ratio with hydrogen blending ratio at 20% and 30%, but the flame robustness of ammonia and hydrogen at far-lean global equivalence ratio should be considered and investigate further.

Surprisingly, changes in hydrogen blending ratios have no significant impact on the unburnt hydrogen over the wide range of the global equivalence ratios. However, due to the flame temperature elevated by more hydrogen involvement, thermal  $NO_x$  formation is greatly affected. The elevated flame temperature also improves the ammonia fuel consumption. Consequently, increased  $NO$  and  $NO_2$  as well as reduced ammonia slip at the exhaust were observed as more hydrogen was blended with ammonia. However, the effect is weakened considerably when the global equivalence ratio is above the unity. It indicates that more hydrogen blending ratios provide less penalty on emissions under fuel-rich conditions.

### 3.2.2. Emission performance at different inlet temperatures

In Fig. 9, variations of emissions for non-premixed 60%  $NH_3$ / 40%  $H_2$  flames are presented with the constant global equivalence ratio at 1.1 and swirl number at 0.87. The inlet fuel temperature varied from 300 to 400 K and the inlet air temperature varied from 300 to 350 K. It was well known that increasing the inlet temperature improves the flame temperature and chemical reaction rate. According to the Zel'dovich



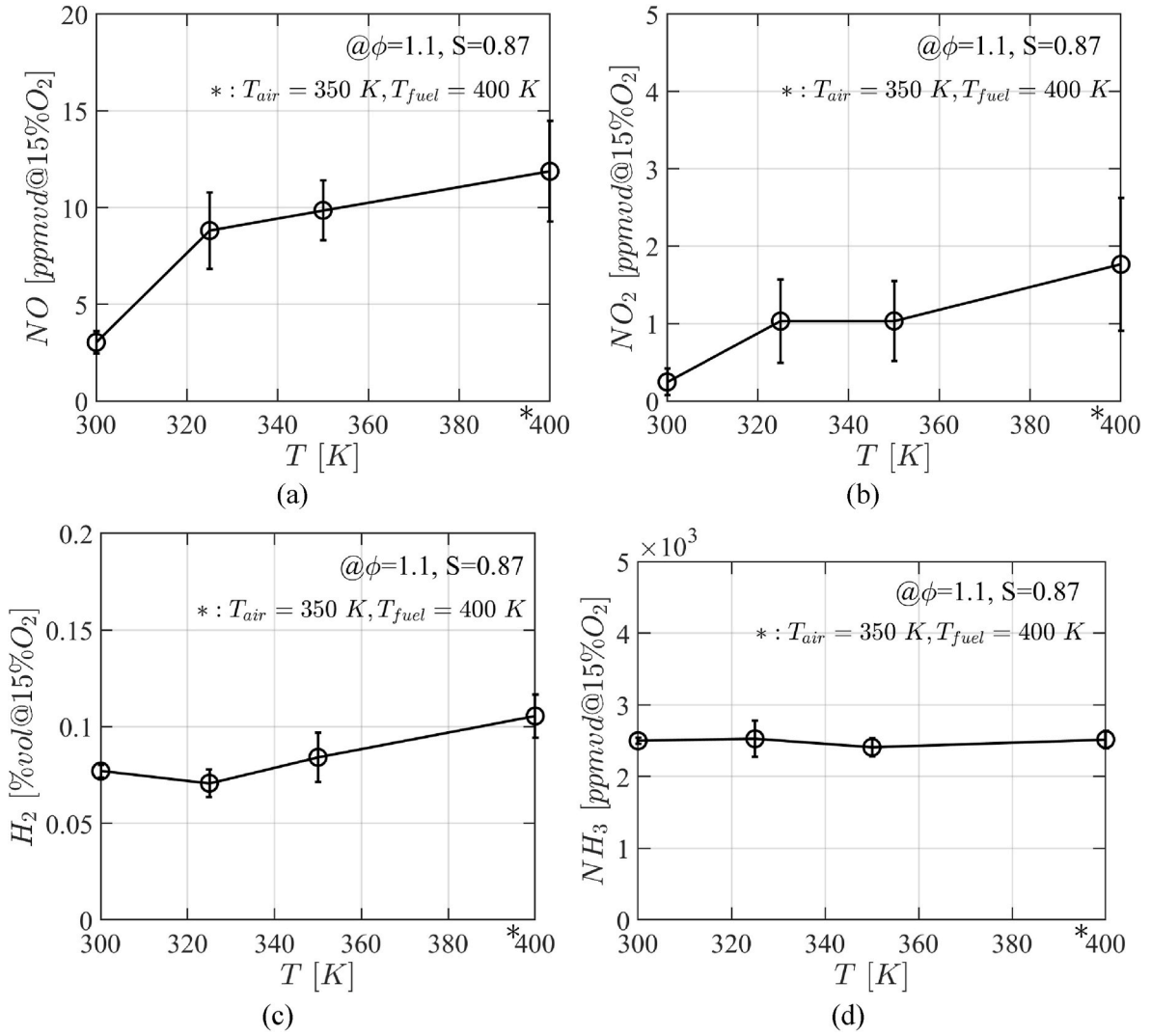


Fig. 9. Emissions of non-premixed 60%  $\text{NH}_3$ /40%  $\text{H}_2$  flames versus different inlet temperature.

mechanism, when the temperature is above 1800 K,  $\text{NO}_x$  formation through thermal  $\text{NO}_x$  route is promoted. Valera-Medina et al. [17] investigated premixed ammonia/hydrogen flames using 0D Gaseq analysis, finding  $\text{NO}_x$  concentration of flue gas was increased as the inlet temperature increase from 300 to 900 K at 10 bar. However, the experimental results in the present work show the lifted inlet temperature had no obvious impact on the emission at the global equivalence ratio of 1.1. The measured inner wall temperature was increased by approximately 46 K for every 25 K increase in the inlet temperature. The moderate effect of increasing the inlet gas temperature on the inner wall temperature could be owing to the heat loss from the sizeable optical window and the insufficient increase in the inlet temperature. This suggests that the increase in the flame temperature not be significant in showing the profound difference in emissions. The conjecture of flame temperature was verified in section 3.3  $\text{OH}^*$  chemiluminescence, Fig. 14.

### 3.2.3. Emission comparison under the wall insulated and non-insulated conditions

As temperature plays a key role in the flame and affects emissions considerably, the effect of wall condition on emissions of non-premixed ammonia/hydrogen combustion has been investigated. The combustion chamber was insulated with silicate ceramic fibre blankets to compare the emission results between insulated wall conditions and non-

insulated wall conditions. Fig. 10 illustrates the emissions for non-premixed 70%  $\text{NH}_3$ /30%  $\text{H}_2$  flames with the inlet gas temperature of 300 K under insulated wall condition and non-insulated wall condition. The results show that a substantial growth of up to 107% ( $\varphi_{\text{global}} = 0.8$ ) in  $\text{NO}$  emission and up to 92% ( $\varphi_{\text{global}} = 0.8$ ) in  $\text{NO}_2$  emission, alongside a notable reduction of up to 44% ( $\varphi_{\text{global}} = 1.1$ ) in unburnt ammonia at the exhaust, when the combustion chamber is converted from non-insulated to insulated. Meanwhile, unburnt hydrogen kept almost the same. It suggests a great heat loss from the optical window and combustion chamber. The great heat loss, especially in the region close to the wall, considerably suppresses the generation of  $\text{O}/\text{H}$  radicals which are mainly yielded by the chain branching reaction,  $\text{H} + \text{O}_2 \rightarrow \text{OH} + \text{O}$  [44]. Thus, thermal  $\text{NO}_x$  generation is suppressed. The substantial ammonia concentration at the exhaust suggests further low  $\text{NO}_x$  generation from ammonia through fuel-bound  $\text{NO}_x$  route. The emission trends are consistent with the study on non-premixed ammonia/air flames that have been examined experimentally with non-insulated quartz glass liner, non-insulated steel liner and insulated steel liner [45].

### 3.2.4. Emission performance at different swirl numbers

Three swirlers with different vane angles were tested. Fig. 11 demonstrates the emissions of non-premixed 70%  $\text{NH}_3$ /30%  $\text{H}_2$  flames with various swirl numbers. Due to the flame stability difference using different swirl numbers, flames measured with small swirl number

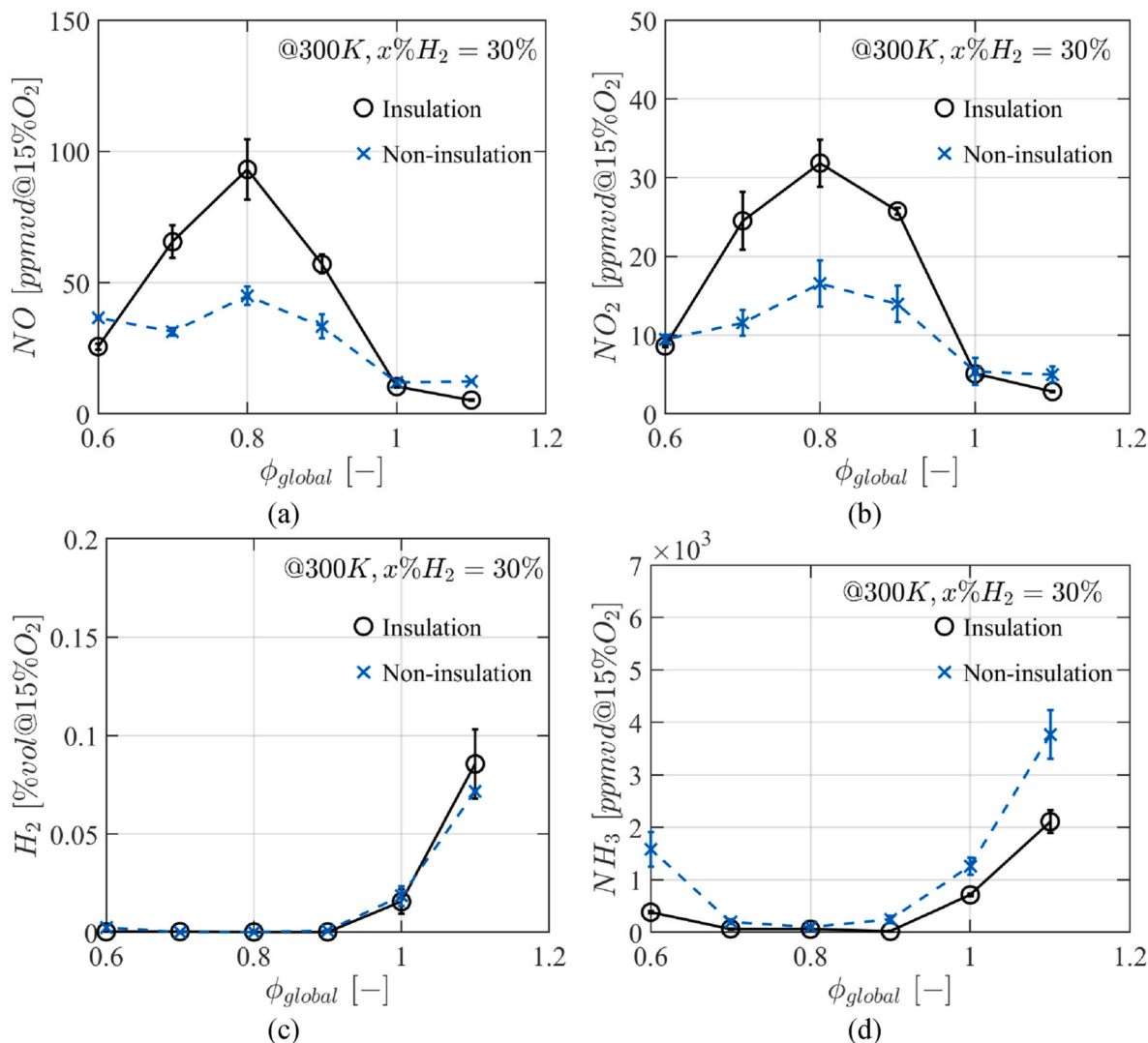


Fig. 10. Emissions of non-premixed 70%  $\text{NH}_3$ / 30%  $\text{H}_2$  flames versus various global equivalence ratios under insulated and non-insulated wall conditions ( $S = 0.87$ ).

provide a wider range of emission results. It is found that the change in the swirl number has a moderate effect on the  $\text{NO}_x$  emission, although a slight reduction of  $\text{NO}$  and  $\text{NO}_2$  concentration with swirler number at 0.87 and 1.04 was observed when the global equivalence ratio was greater than 0.8. Under far-lean conditions, unburnt hydrogen concentration at the exhaust is almost unaffected by swirl numbers while unburnt ammonia concentration at the exhaust is reduced considerably. For instance, at the global equivalence ratio of 0.6, the unburnt ammonia was 1880  $\text{ppmvd@15\%O}_2$  with swirl number at 0.73, 1580  $\text{ppmvd@15\%O}_2$  with swirl number at 0.87 and 887  $\text{ppmvd@15\%O}_2$  with swirl number at 1.04. In the premixed 30%  $\text{NH}_3$ / 70%  $\text{CH}_4$  flames, it was reported in the literature by conducting a large eddy simulation that a decreased  $\text{NO}$  emission was achieved by increasing the swirl number from 0.58 to 0.99, and when the swirl number was greater than 0.76, less effect was observed on  $\text{NO}$  reduction [46]. It suggests there is potentially an optimal swirl number for  $\text{NO}$  emission control in non-premixed ammonia/hydrogen flames. Under the fuel-rich conditions, both the concentrations of unburnt hydrogen and ammonia were increased dramatically with a higher swirl number. As the angle of fuel nozzle is  $45^\circ$ , the vane angle of swirler at  $40^\circ$  ( $S = 0.73$ ) enables the fuel and air to be mixed near the nozzle. The vane angle of swirler at  $50^\circ$  ( $S = 1.04$ ) provides a stronger swirling flow but the mixing flow is being pushed closer to the cold wall. The mixing flow is burnt near the nozzle but the mixing flow close to the cold wall is likely to be pushed to

the exhaust without reacting. The behaviour became stronger with more fuel dominate in the chamber, which led to less unburnt fuel (hydrogen and ammonia) with a low swirl number under fuel-rich condition. The effect of swirl numbers on unburnt fuel emission became greater when the global equivalence ratio was further increased.

### 3.3. $\text{OH}^*$ chemiluminescence

Chemiluminescence of a flame is the spontaneous emission of light that occurs when a chemical species descends from the excited to the ground state. Excited hydroxyl ( $\text{OH}^*$ ) chemiluminescence is recognised as an important parameter to understand physical properties without interfere with the actual flame, e.g. flame structure, relationship with  $\text{NO}$  concentration. In this section, a thousand of images of  $\text{OH}^*$  chemiluminescence were taken for each case to gain the knowledge qualitatively and quantitatively. For the distribution of  $\text{OH}^*$  chemiluminescence, normalisation was conducted to the image data by dividing by the maximum intensity of each case and rescaling to the same range shown in the colour bar. Therefore, it is noted that the value ( $\text{II}_{\text{OH}^*}$ ) does not represent the absolute value.

Fig. 12(a) shows the  $\text{OH}^*$  chemiluminescence profiles of the non-premixed 60%  $\text{NH}_3$ / 40%  $\text{H}_2$  flame with various global equivalence ratio in the range of 0.6 – 1.2 measured at the room temperature with the swirl number of 0.87. Owing to the axis symmetry of the image, only

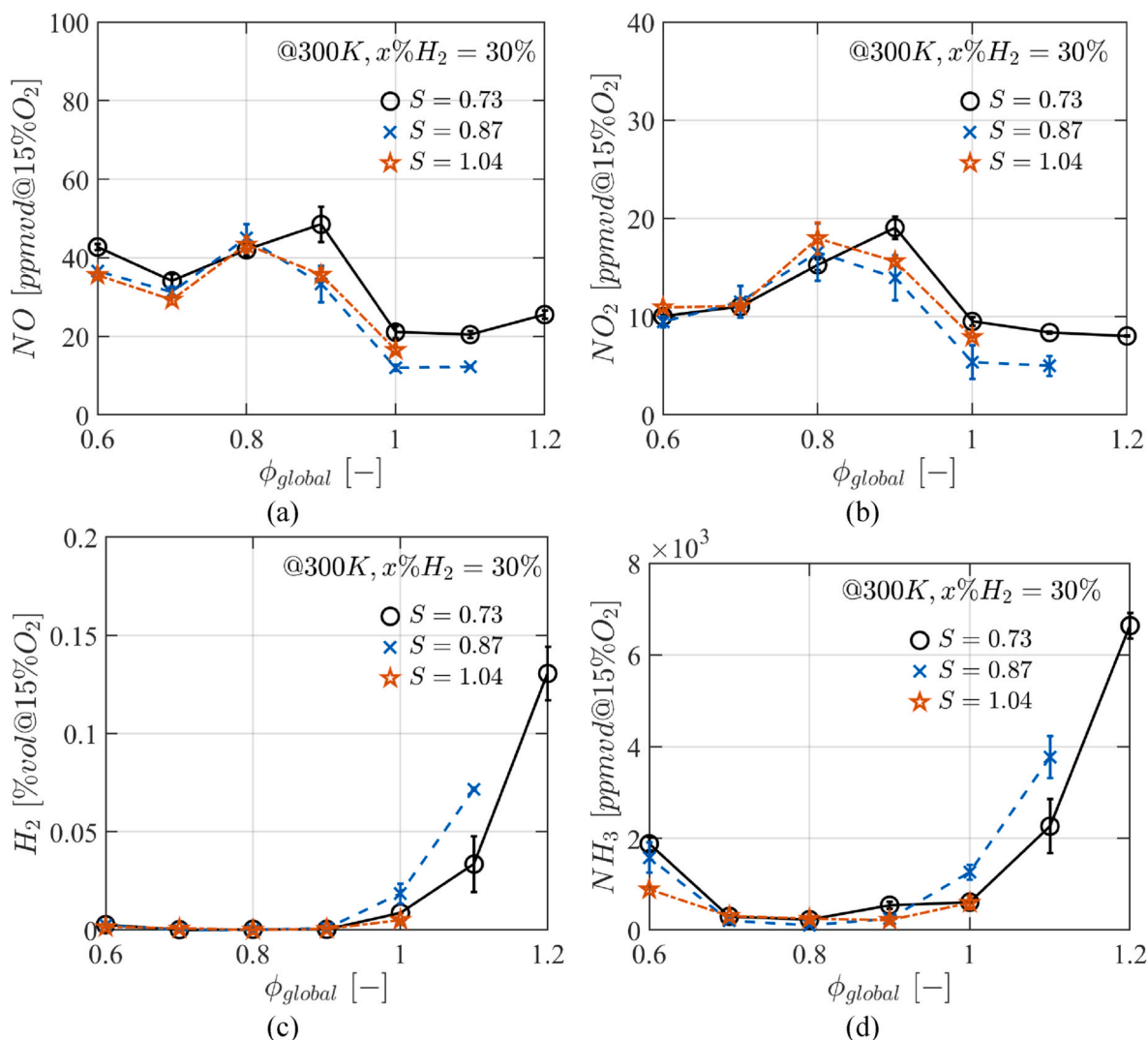


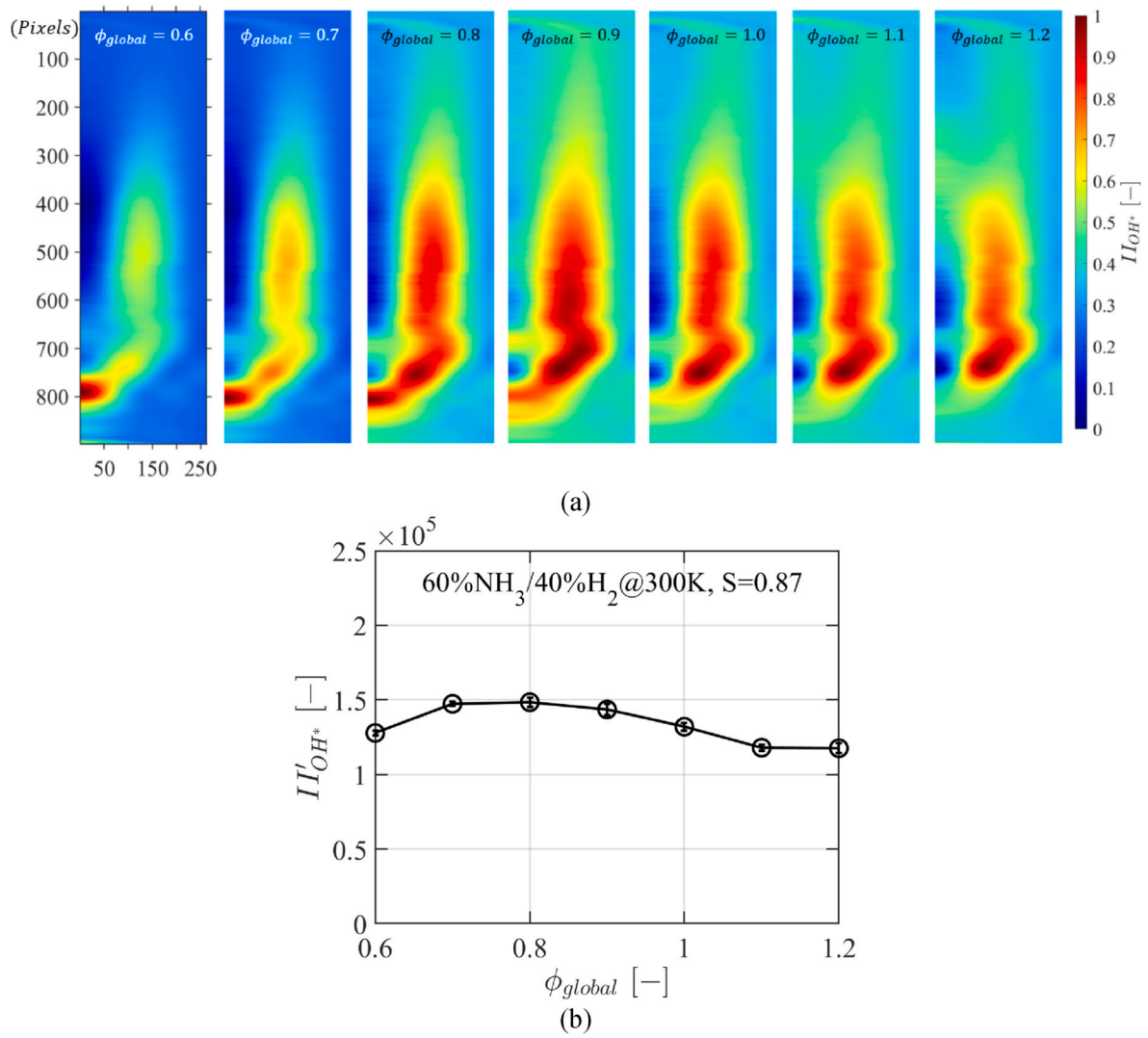
Fig. 11. Emissions of non-premixed 70%  $\text{NH}_3$ /30%  $\text{H}_2$  flames versus various global equivalence ratios with different swirl numbers.

the top half of each image is rotated for analysis. From the perspective of topology in each case, it is found that the  $\text{OH}^*$  concentration was generally located at the upstream close to the nozzle in all cases and it was moved further away from the nozzle when the global equivalence ratio increased. Furthermore,  $\text{OH}^*$  chemiluminescence distribution was expanded as the flame became richer. The phenomenon was consistent with the observation of flame shape. At the far-lean conditions, the flame height of ammonia/hydrogen was short and orangish. As less oxidizer was participated in the reaction, the flame height became longer and turned to yellow-ish. It demonstrates that the feature of  $\text{OH}^*$  chemiluminescence distribution that can be used to identify flame shape can also be extended to non-premixed ammonia and hydrogen flames. Fig. 12 (b) reveals the  $\text{OH}^*$  intensity results with the respect to the global equivalence ratios. It can be seen that the  $\text{OH}^*$  intensity trend is compatible with the trend in NO emission in Fig. 8 (a). Hydroxyl radical plays a critical role in fuel-bound NO formation ( $\text{HNO} + \text{OH} = \text{NO} + \text{H}_2\text{O}$ ) and fuel-bound NO formation dominates in the NO emission under fuel-lean conditions. Under fuel-rich conditions, fuel-bound NO formation route is inhibited due to the insufficient O/H/OH radicals. Therefore, hydroxyl radical and excited hydroxyl radical measurement reflects nitric oxide concentrations at the post-flame zone. The findings agree with the results observed with premixed ammonia/air flames and partially premixed ammonia/hydrogen flames in the literature [22].

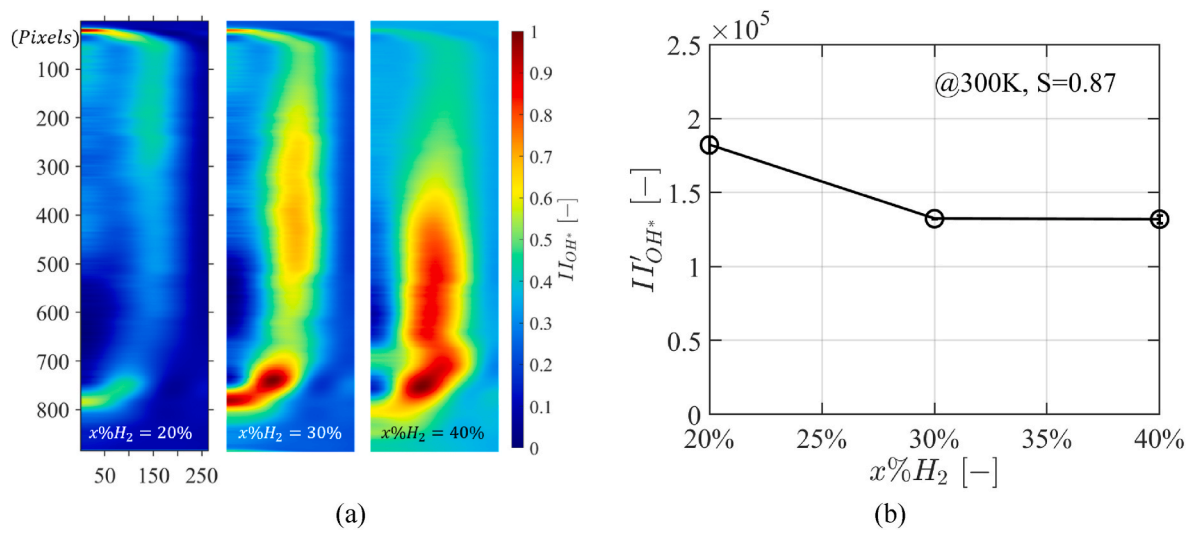
It is depicted in Fig. 13(a) that  $\text{OH}^*$  chemiluminescence images of the

non-premixed ammonia/hydrogen flames with hydrogen blending ratio at 20%, 30% and 40% at stoichiometric condition. The result shows that  $\text{OH}^*$  chemiluminescence near the nozzle region was evidently enlarged as more hydrogen was involved in the fuel blends. For the case of non-premixed 80%  $\text{NH}_3$ /20%  $\text{H}_2$  flame,  $\text{OH}^*$  is intense at the region near the nozzle and downstream, showing a weak flame behaviour. It agrees with the results in Fig. 5 that the rich blowoff limit for this case is slightly above the global equivalent ratio at 1.0. With the increase of the hydrogen blending ratio, the  $\text{OH}^*$  radicals became more concentrated near the nozzle and the flame became more robust. As illustrated in Fig. 13 (b), the  $\text{OH}^*$  intensity is about 37% higher at a hydrogen blending ratio of 20% than at a hydrogen blending ratio of 30% that is almost the same as at the hydrogen blending ratio of 40%. In comparison with results in Fig. 8, the change in  $\text{OH}^*$  intensity is consistent with the change in NO emission at the global equivalence ratio of 1.0. Hydrogen involvement lifted the flame temperature, an increase in NO emission via thermal NO production is expected. However, much higher  $\text{OH}^*$  concentration was found in 80%  $\text{NH}_3$ /20%  $\text{H}_2$ , indicating that the fuel-bound NO route still substantially contributes to NO production and it is slightly greater than the effect of increasing the hydrogen blending ratio up to 40% when the global equivalence ratio was at 1.0.

In Fig. 14(a), the distributions of  $\text{OH}^*$  chemiluminescence of non-premixed 60%  $\text{NH}_3$ /40%  $\text{H}_2$  flames are presented with different inlet temperature at the global equivalence ratio of 1.1. The flame shape

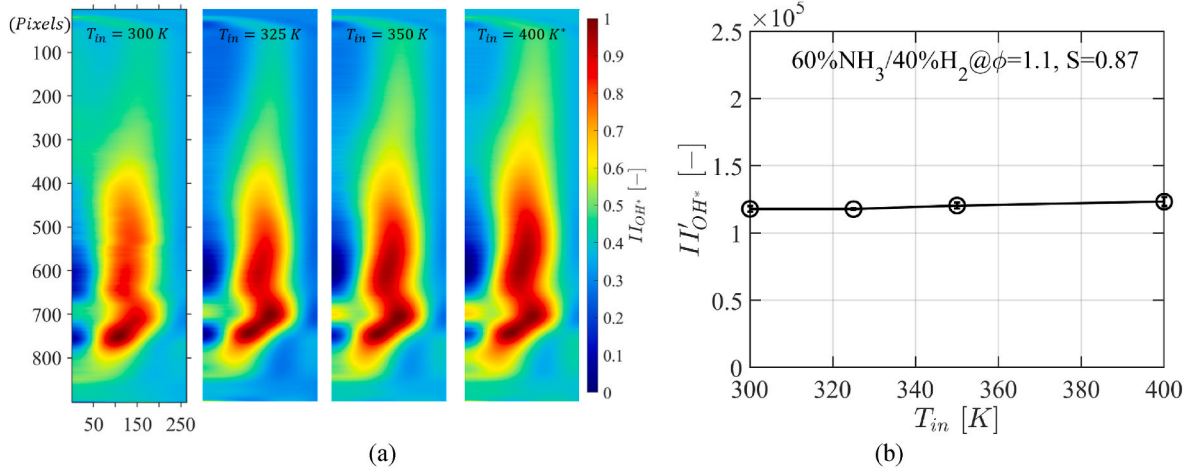


**Fig. 12.** (a)  $OH^*$  chemiluminescence profiles (normalised colormap applied) and (b)  $OH^*$  intensity for non-premixed  $60\% NH_3 / 40\% H_2$  flames with various global equivalence ratios ( $T = 300\text{ K}$ ,  $S = 0.87$ ).



**Fig. 13.** (a)  $OH^*$  chemiluminescence profiles (normalised colormap applied) and (b)  $OH^*$  intensity for non-premixed flames with various hydrogen blending ratios ( $\phi_{global} = 1$ ,  $T = 300\text{ K}$ ,  $S = 0.87$ ).





**Fig. 14.** (a)  $OH^*$  chemiluminescence profiles (normalised colormap applied) and (b)  $OH^*$  intensity for non-premixed 60%  $NH_3$ /40%  $H_2$  flames with various inlet temperature ( $\phi_{global} = 1.1, S = 0.87$ ).

remained almost unchanged, but the regions of the peak  $OH^*$  concentration (deep red) was expanded. Fig. 14(b) also reflects rising the inlet temperature has no significant effect on the  $OH^*$  intensity, with a slight increase observed. It indicates that there was no considerable change in the flame temperature in the current experiment as the inlet air temperature rising from room temperature to 350 K and inlet fuel temperature rising to 400 K. Therefore, it can explain the result of  $NO$  emission in Fig. 9 (a) which was found almost the constant.

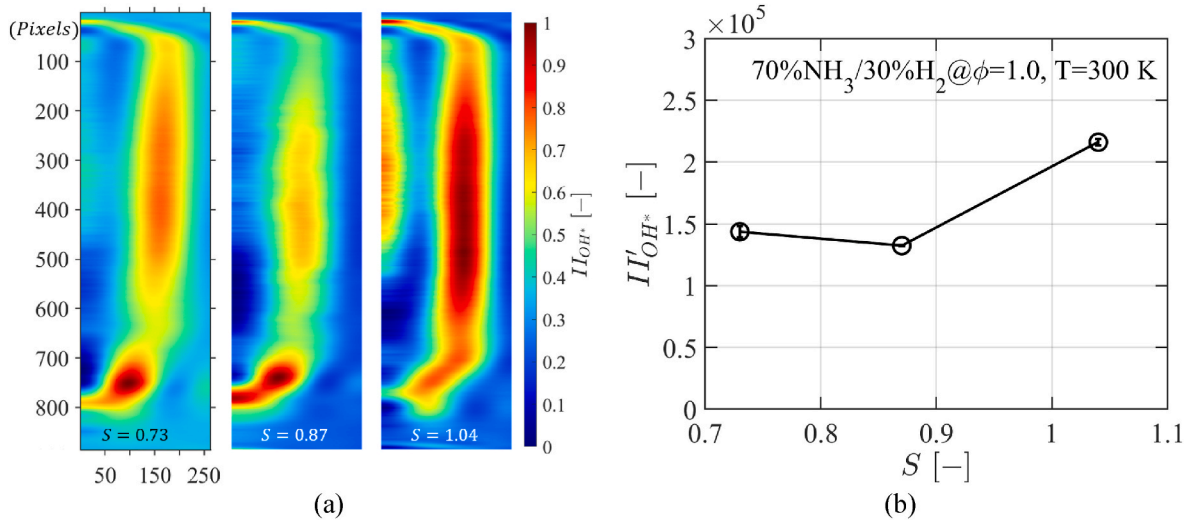
Three different swirlers were changed to create various flow recirculation in the combustor. The profiles of  $OH^*$  chemiluminescence of non-premixed 70%  $NH_3$ /30%  $H_2$  flames with different swirlers are demonstrated in Fig. 15(a) at the stoichiometric condition under the room temperature. As the vane angle of the swirler varied from  $40^\circ$  to  $50^\circ$ , the angle of the  $OH^*$  concentration upstream was deflected, which helped the circulation and mixing ratio of the fuel and air near the nozzle. Additionally, with a higher swirl number,  $OH^*$  concentration in the region near the nozzle and transition region expanded and showing a more strengthened flame. For instance, when the swirl number reached 1.04, the red marked  $OH^*$  chemiluminescence was from the nozzle all the way downstream. In Fig. 15 (b), it is notable that the  $OH^*$  intensity decreased and then increased significantly as the swirl number rose from 0.73 to 1.04. However, the emission results in Fig. 11 (a) show a higher

$NO$  emission was found at the swirl number of 0.73, which does not match the results of  $OH^*$  intensity under the same condition. It is necessary to conduct a further spatial and temporal information investigation into the pathways of  $NO$  formation to understand the reasons behind it.

#### 4. Conclusions

Combustion and emission characteristics of non-premixed ammonia/hydrogen flames were studied experimentally with a swirl-enhanced combustor test rig. The blowoff limits were identified firstly. The effects of global equivalence ratios (0.6–1.2), hydrogen blending ratios (20%–40%), inlet gas temperatures (300–400 K), swirl numbers (0.73, 0.87, 1.04) and combustion chamber wall insulation conditions on emissions ( $NO$ ,  $NO_2$ ,  $NH_3$ ,  $H_2$ ) were analysed. Excited hydroxyl radical ( $OH^*$ ) chemiluminescence imaging technology was unveiled. The main findings are summarised below.

- (1) Taking the combustion efficiency into account, the optimal emission control occurred at stoichiometric condition. At far-lean global equivalence ratios,  $NO_x$  and unburned fuel also showed



**Fig. 15.** (a)  $OH^*$  chemiluminescence profiles (normalised colormap applied) and (b)  $OH^*$  intensity for non-premixed 70%  $NH_3$ /30%  $H_2$  flames with various swirl numbers ( $\phi_{global} = 1.0, T = 300\text{ K}$ ).

relatively low emissions with hydrogen blending ratio at 20% and 30%; however, flame stability is of concern.

- (2) Robust flames were obtained at hydrogen blending ratio over 20%. More hydrogen involvement resulted in the increase of  $NO$  and  $NO_2$  emissions and the decreased ammonia slip. However, the effect was weakened under fuel-rich conditions.
- (3) With the global equivalence ratio of 1.1, the rising inlet gas temperature has a marginal effect on emissions due to the heat loss from the combustor wall and optical windows. After the non-insulation rig was converted to the wall insulated combustor,  $NO$  and  $NO_2$  emissions increased by a maximum of 107% and 92%, respectively, despite a reduction in ammonia slip of up to 44%.
- (4) In the current combustor setup, when the global equivalence ratio above 0.8,  $NO$  and  $NO_2$  emissions were lower with the swirl number of 0.87 and 1.04 than that with the swirl number of 0.73. This suggested there is an optimal swirl number for  $NO_x$  control and its value changes depending on the geometry and size of combustor.
- (5)  $OH^*$  chemiluminescence distribution can effectively identify the shape of non-premixed ammonia and hydrogen flames. The trend

of  $OH^*$  intensity was consistent with the results of  $NO$  emissions at the same swirl number.

## Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

## Acknowledgements

The authors are grateful for the financial support from EPSRC (Engineering and Physical Sciences Research Council, United Kingdom) through the project 'Powering Carbon-free Autonomous Shipping: Ammonia/Hydrogen dual-fuelled Linear Engine-Generator' (EP/S00193X/2) and the project 'Premixed Combustion Flame Instability Characteristics' (EP/W002299/1). The authors also thank for the project support from Merchant Marine College of Shanghai Maritime University.

## Nomenclature

NTP	Normal Temperature and Pressure
GTRC	Gas Turbine Research Centre
CRN	Chemical Reactor Network
MFCs	Mass Flow Controllers
LHVs	Lower Heating Values
PDD	Pulse Discharge helium ionisation Detector
TCD	Thermal Conductivity Detector
$S$	Swirl number (–)
$Da$	Damköhler number (–)
$D_i$	Inner diameter of swirler (mm)
$D_o$	Outer diameter of swirler (mm)
$\alpha$	Vane angle of swirler (°)
$P$	Total input thermal power (kW)
$\dot{m}_{fuel}$	Mass flow rate of fuel blends (g/s)
$\dot{m}_{air}$	Mass flow rate of air (g/s)
$x[NH_3]$	Mole fraction of ammonia in the fuel blends (–)
$x[H_2]$	Mole fraction of hydrogen in the fuel blends (–)
$\phi_{global}$	Global fuel-air equivalence ratio (–)
$x\%H_2$	Hydrogen blending ratio (–)
$ppm_{norm}$	Normalised emission concentration (ppmvd@15%O <sub>2</sub> )
$ppm_{mea}$	Measured emission concentration (ppmvd)
$O_{2mea}$	Measured oxygen concentration in the exhaust gases (ppmvd)
$II_{OH^*}$	Normalised $OH^*$ intensity (–)
$II'_{OH^*}$	$OH^*$ intensity before normalisation (–)
$\bar{I}_{OH^*}$	$OH^*$ intensity of the $i^{th}$ column and the $j^{th}$ row (–)
$T_{fuel}$	Inlet fuel temperature (K)
$T_{air}$	Inlet air temperature (K)

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