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Experimental Investigation on flame stability of Ammonia-LPG Flames

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Abstract- NH₃ as an alternate fuel is gaining huge importance in the recent past owing to the urgency to mitigate global warming caused by conventional hydrocarbon fuels. However, NH₃ because of its low laminar flame speed and reactivity when compared to those of conventional hydrocarbon fuels, needs to be blended with conventional fuels for it to be a viable alternative. In this paper, NH₃ in various proportions is blended with LPG. Experiments are conducted using a single slot burner of 17.3 mm inner diameter for different combinations of parameters like (i) mixing ratios of NH₃ and LPG in fuel mixture, (ii) low power ratings (0.1 to 1.1 kW) and (iii) percentages of partial premixing with air (0% to 30%). Analysis of varying the aforementioned parameters is done using flame photographs and trends in flame length data. This is followed by a stability map for different configurations which divides the flames into different regimes of (A) steady flames, (B) stable flames with oscillating tip, (C) flames with vortex shedding, and (D) unstable flames with lift-off. An increase in partial premixing is observed to have a stabilising effect (making flames transition from the regime of vortex shedding to that with oscillating tips) whereas an increase in power rating has the opposite effect. It is found that increase in partial premixing shortens the average flame length whereas increase in power rating has the opposite effect. Stable and steady flames are observed with either lower percentage of partial premixing coupled with higher concentrations of NH₃ or higher percentage of partial premixing of NH₃. Additionally, the effect of increasing NH₃ volume fractions in the fuel mixture on adiabatic flame temperature, NO, CO and CO₂ production is also studied using equilibrium calculations and the results are reported.

Keywords: Ammonia, LPG, Diffusion, Partially Premixed, Flame photographs, Stability Maps, Average Visible Flame Height, NO, Adiabatic Flame Temperature.

1. Introduction

In the recent times, global warming has increased tremendously due to the contribution of greenhouse gases released from hydrocarbon combustion. Several research works focus in the direction of carbon-neutralisation in order to mitigate the release of greenhouse gases. Ammonia (NH₃) is considered to be one of the alternative fuels of interest [1] owing to its clean energy nature i.e., zero carbon emissions [2]. Being a zero-carbon fuel and an economic hydrogen carrier, its complete combustion products include only steam and nitrogen. However, NH₃ combustion comes with its own set of challenges such as lower burning velocity, flammability limit, calorific value and adiabatic flame temperature when compared to conventional hydrocarbon fuels [3]. Therefore, it is difficult to burn pure NH₃ in order to establish stable flames [4]. Additionally, ammonia combustion leads to high NOx emissions due to the presence of N atom [5].

To address the above issues, NH₃ is often blended with conventional fuels such as hydrogen and various hydrocarbon fuels like methane, gasoline and dimethyl ether (DME). The type of fuel blends and equivalence ratio plays a significant effect in determining the laminar flame speed of the fuel mixture. NH₃ blends including methane [3], syngas [6] and dimethyl ether [7] is found to enhance laminar burning velocity (LBV).

Lee et al. [8] observed the increase in LBVs with increasing hydrogen concentration in the fuel mixture at 0.1 MPa. The effect of adding H₂, CO and CH₄, on NH₃/air mixtures was investigated using a heat flux burner by Han et al. [9]. It was found that the maximum LBV shifts towards slightly richer conditions in comparison to pure ammonia, for up to 40% H₂ concentration. Lhuiller et al. [10] found an increment in the flame speed from 7 cm/s (0% H₂) to nearly 80 cm/s (60% H₂) at 298 K. Another study [11] shows that NH₃-H₂ blends can have LBVs comparable to that of CH₄ as H₂ ratio increases from

33.3% to 60%. This was also explained by Li et al. [12] using a numerical model. It was found that H₂ caused a reduction in overall chemical activation energy and increased transport effect such as mass and thermal diffusion.

Okafor et al. [13] tested the unstretched LBV of a premixed NH₃-CH₄ flame in a constant volume chamber for different equivalent ratio ranged from 0.8 to 1.3 and the NH₃ content (% volume) varied from 0 to 0.3. The findings demonstrated that as NH₃ content and equivalence ratio increases, the relation between flame velocity and flame stretching rate changed from linear to nonlinear. Ku et al. [14] investigated the potential of CH₄- NH₃ blends in expanding spherical premixed flames and measured Markstein numbers, flame structures, and LBVs. Their findings were in close agreement with those of Okafor et al. [13]. The range of these values was further extended to higher pressures, up to 5 bar, by Shu et al. [15]. They found a strong linear relation between LBV and CH₄ volume fraction for NH₃-CH₄ mixtures with the same equivalence ratio. Experimental and numerical study of Lubrano Lavadera et al. [16] investigated the NH₃-CH₄ flame using a heat flux burner and a spherical chamber. Results were found to differ from one another, especially at higher temperatures, equivalence ratios, and hydrocarbon concentrations where the burner's values were less than 5 cm/s and the chamber's values were greater. Their investigation revealed that, for all NH₃ mass fractions, the S_L/S_{L,0} ratio trended linearly, but the temperature behaved non-linearly. Additionally, Liu et al. [17] used CO₂ dilution (25–65%) in the oxidizer to study these patterns under oxygen fuel conditions. Across all equivalence ratio, they revealed that burning velocity increased with increasing CH₄ content.

Given its carbon-neutral characteristics and improved combustion performance, dimethyl ether (DME) is drawing interest as a possible NH_3 enhancer [18]. Cai and Zhao's work [7] assessed the laminar flame speed (S_L) of NH_3 -DME flames using a 1D freely propagating flame model. They found that the addition of DME considerably increased the ammonia laminar flame speed to levels that are comparable to those of hydrocarbon fuels. It is also seen that increasing the amount of oxygenated alternative fuels like ethanol [19, 20] and methanol [20] is essential to raising NH_3 's reactivity and combustion efficiency.

As it is clearly seen, blending conventional hydrocarbon fuels with NH_3 is a rising area of research and overcomes most of the shortcomings of pure ammonia flames. One such hydrocarbon fuel which requires attention in recent times is Liquefied Petroleum Gas (LPG). It finds a variety of uses in domestic and industrial areas owing to its efficiency, portability, versatility and clean burning characteristics compared to other hydrocarbon fuels. In industries, it is used for metal processing, firing of ceramic products, drying and steaming in textile industries, powering automotive vehicles, etc. In the domestic sphere, LPG is primarily used in cooking and in heating devices. In India, LPG is the most commonly used fuel for cooking. However, there are no studies related to flame stability of ammonia-LPG blended flames available in the literature. Therefore, this study initiates an experimental investigation on flame stability for different amounts (% by volume) of NH₃ (0, 50, 70, 75, 80%) blended with commercially available LPG. A newly designed and fabricated single slot Bunsen burner (of inner diameter 17.34 mm) is used to establish canonical laminar diffusion and partially premixed NH₃-LPG flames. A series of experiments are conducted using this burner by varying different parameters like (i) mixing ratios of NH₃ and LPG in fuel mixture, (ii) power ratings (0.1 to 1.1 kW) and (iii) percentages of partial premixing with air (0% to 30%). A qualitative analysis is done using the flame photographs in order to understand the effect of these parameters on the flame stability. The variation of the measured visible flame heights with respect to aforementioned parameters are also reported. Additionally, the effect of increasing NH₃ volume fractions in the fuel mixture on adiabatic flame temperature, NO, CO and CO₂ production is also studied using equilibrium calculations and the results are reported.

2. Experimental Methods

In the present study, the flame stability of ammonia and LPG blends is characterized by establishing a canonical diffusion and partially premixed flames in a newly designed and fabricated single slot Bunsen burner. The burner tube made up of stainless steel has internal diameter and length of 17.34 mm and 15 cm, respectively. The inner diameter of the burner is chosen such that the Reynolds' number for the flows inside the tube are well within the laminar regime. The tube length is selected such that the flow is fully developed as it comes out of the burner. A thickness of 1.83 mm is used to ensure optimal heat dissipation. The burner is provided with four inlets around the periphery of the settling chamber. The photographic image of the Bunsen burner and the schematic of the experimental setup are shown in the Figs. 1.1 and 1.2, respectively. The fuel cylinders of commercial LPG and ammonia (purity 99.99%) supplies fuel to the burner and the flow rates of each fuel are controlled using respective Mass Flow Controllers (MFC1 and MFC2). The outlets of the LPG and ammonia MFCs deliver the required fuel flow rates and are connected directly to a settling (or mixing) chamber through the flexible pneumatic pipes. The air used in the experiments are drawn from the main compressor line. The flow rate of air is controlled using MFC3 before it is allowed to mix with the fuel. All the flow rates of ammonia, LPG and air are regulated using Alicat MFC with 0.5% of reading accuracy. The flame photographs are captured with the help of a high-speed camera (EOS M50 Mark II).

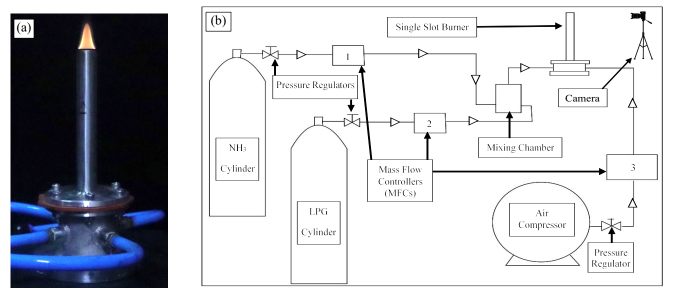


Fig. 1: (a) Photographic image of the singe slot burner and (b) Schematic of the experimental set up (1- MFC for ammonia, 2- MFC for LPG and 3- MFC for air)

The MFC used in measuring the flow rate of a multicomponent fuel such as LPG has some limitations in handling the different components. Therefore, analogous fuel compound has to be chosen based on the molecular weight, viscosity and the major component present in the actual fuel. The molar composition of LPG used in the experiments is measured using a gas chromatography (Model: Perkin Elmer Clarus 580; Asset ID: 8819). It consists of the following components by volume: 44.82% C₃H₈, 33.27% n-C₄H₁₀, 16.82% iso-C₄H₁₀, 1.75% iso-C₄H₈, 1.06% trans-C₄H₈, 0.33% C₃H₆, 0.25% iso-C₅H₁₂ and 0.082% CH₄. The equivalent molecular weight and (lower) calorific value of LPG thus calculated is 51.58 g/mole and 45895.41 kJ/kg, respectively. The viscosity of this multi-component fuel, i.e., LPG, calculated using Wilke's formula [21] is found to be 7.79 x 10⁻⁶ Pa-s. As the calculated molecular weight and viscosity of LPG ($\eta_{LPG}/\eta_{butane} = 1.06$) is closer to butane, LPG flow rate in MFC 2 is therefore controlled by setting the fuel as n-butane.

The relationship between the desired flow rate of LPG and the indicated flow rate of n-butane on the MFC display is given by the relationship (Eq. (1)) [22],

$$Q_{\text{butane}} = Q_{\text{LPG}} x (\eta_{\text{LPG}} / \eta_{\text{butane}}) (1)$$

where, Q = volume flow rate, and $\eta =$ viscosity of gas.

The gas set for the other two MFCs is the same as the fluid passing through them. The calorific value and viscosity of ammonia are 18601 kJ/kg [23] and 10.083 x 10^{-6} Pa-s [23], respectively. LPG and ammonia are blended together to form fuel mixtures of various compositions by volume as shown in Table 1. For each fuel mixture composition, the diffusion and partially premixed flames are established with the power ratings ranging from 0.1 to 1.1 kW.

Table 1: Experimental Configurations				
Burner Inlet &	Fuel Compositions		Power Rating	Flame
Ambient			(kW)	configurations
Conditions				
Temperature:	% vol.	% vol.	0.1, 0.2, 0.3,	(a) Diffusion
298 K	NH ₃	LPG	0.4, 0.6, 0.8,	
Pressure: 1 atm	0	100	1.1	(b) Partially
	50	50		Premixed
	70	30		(i) 10% (ii) 20% (iii)
	75	25		30% of
	80	20		stoichiometric air.

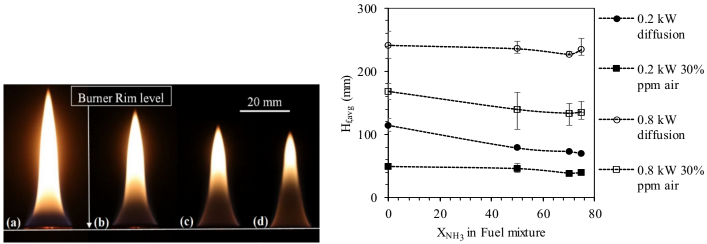
To set up a flame for each trial, LPG is passed initially through MFC 2 at the required flow rate and a flame is established. Then, ammonia and air (if required) are passed through MFC 1 and 3, respectively at the required flow rates. The photographic images are taken once the initial disturbances in the flame caused by the flow are removed. In each trial, photographic images of the flames are taken for a span of 120 seconds, with the help of a high-speed camera. In each second, 50 consecutive images were captured at regular intervals, leading to 6000 images. A developed MATLAB code is used to retrieve the average flame height from each of these frames in pixels. This flame length is then converted in the units of millimetres from a suitable reference image. A minimum of three such trials are performed for each case. Apart from flame height, the nature and stability of flames are also observed and categorised.

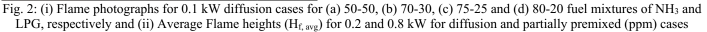
3. Results and Discussion

The effect of power rating, ammonia percentage in fuel mixture and percentage of partial premixing on the flame stability of NH_3 -LPG flames is analysed qualitatively with the help of photographic flame images. A flame stability map is then developed based on the collective observations obtained from the series of experiments to demarcate the various stability regimes.

3.1. Effect of Ammonia Percentages in Fuel mixture

The effect of varying ammonia volume percentage is studied by establishing a diffusion and partially premixed flames with constant power ratings of 0.1, 0.2 and 0.8 kW. The typical flame photographs of 0.1 kW diffusion flame and variation of flame height with respect to ammonia volume percentages are shown in the Figs. 2(i) and 2(ii), respectively. Two primary effects are observed as shown in Figure 2.(i),— (a) a reduction in flame height (~11% for 0.1 kW diffusion flames when 0.8 mole fraction of ammonia in fuel mixture as compared to pure LPG flames) and (b) a decrease in the sooty or luminous zone of the flame. The second observation is promising as it indicates production of less soot with increasing ammonia mole fraction in fuel mixture. A steady flame can be established on a single slot burner with ammonia volume percentage as high as 80% in the fuel mixture provides scope for future steady measurements of temperature and emissions of the flames. Also, the flames being diffusion flames ensure that they are closer to real life domestic applications. It is noted from Figure 2.(ii) that for lower power ratings like 0.2 kW, ammonia causes a higher decrease in flame height for diffusion cases (~39%) than partially premixed cases (~18.7% for 30% partially premixed). However, on increasing power rating (to 0.8 kW in this case), a larger decrease in flame height on ammonia addition is observed for partially premixed cases (~19.9% for 30% partially premixed) than compared to corresponding diffusion cases (~2.94%).





3.2. Effect of Power Rating

The effect of increasing power by keeping the fuel mixture composition constant at 50:50 by volume for diffusion cases is seen (Fig. 3.1). An increase in flame height by approximately 200% is observed as power rating is increased from 0.1 kW to 0.8 kW. However, the flame moves from a momentum driven flow regime to a buoyancy driven flow regime. As a result, the flame tip starts oscillating as we go from 0.1 to 0.2 kW power rating. Thus, the flame is no more steady. Similarly, a further increase in power rating to 0.4 kW takes the flame into a zone where vortex shedding is observed. This phenomenon is even more pronounced when 0.8 kW flames are observed.

20 mm

3.1

3.2 (a)

3.2 (b)

3.2 (c)3.2 (d)3.2 (e)Fig. 3.1: Instantaneous Flame photographs of diffusion cases for 50-50 fuel mixtures of NH3 and LPG (by volume) with power
ratings (kW), (a) 0.1, (b) 0.2, (c) 0.4 and (d) 0.8.Fig. 3.2: Time-averaged visible flame length (H_{f, avg}) of diffusion (0%) and partially premixed flames (~10%, ~20% and ~30%
air) for (a) 0-100 (NH3:LPG by volume) fuel mixture, (b) 50-50 fuel mixture (c) 70-30 fuel mixture (d) 75-25 fuel mixture and (e) 80-

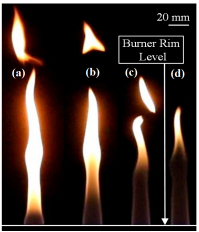
20 fuel mixture

The time averaged visible flame heights are experimentally measured using the help of a camera for different compositions of fuel mixture, power ratings and percentages of partial premixing. For a given fuel mixture composition, they are plotted (Fig. 3.2 (a)-(e)) against different power ratings and percentages of partial premixing. The average flame length increases with power rating when fuel mixture composition and percentage of partial premixing remains the same. It can also be noted that the increase in flame length with power rating is more pronounced for diffusion cases than partially premixed cases. For example, in Fig. 3.2(e), the flame height increases by approximately 300% for 10% partially premixed cases as compared to 240% for 30% partially premixed cases.

3.3. Effect of Partial Premixing with air

The effect of partial premixing with air on LPG-NH3 flames are observed at a constant power rating of 0.6 kW and composition of 50-50 fuel mixture. As the extent of partial premixing increases from 0% to 30% of stoichiometric air (Fig. 4.1 and Fig. 3.2 (b)), the average flame length decreases by approximately 39%. The increase in availability of air leads to burning of the fuel mixture over a shorter distance from the burner rim. There exist luminous regions on top of these flames due to soot formation, the extent of which reduces with increase in the percentage of premixed air (Fig. 4.1). Beneath these luminous sooty regions, there exist non-luminous regions which are associated with combustion closer to stoichiometry. The extent of these non-luminous zones increases despite the decrease in flame heights.

A transition in tendency of flames with vortex shedding to flames with oscillating tips with increase in air flow rate is also observed. Fig. 4.2 shows that for a given fuel composition, the transition of flames from regimes of vortex shedding to regimes of oscillating tip happen at lower extents of partial premixing when the power rating is low. These qualitative observations on increasing the extent of partial premixing hold true for flames with different power ratings and compositions of fuel mixtures.



250 0.3 kW • 0.4 kW 200 **-** 0.6 kW H_{favg} (mm) 150 100 С В 50 0 20 40 0 % of partial premixing air

Fig. 4.1: Instantaneous Flame photographs of 50:50 fuel mixtures of NH_3 and LPG by volume with power ratings 0.6 kW (kW) for (a) 0% (diffusion), (b) 10%, (c) 20% and (d) 30%

Fig. 4.2: Average Flame Heights of photographs of 50:50 fuel mixtures of NH₃ and LPG by volume with power ratings 0.3, 0.4 & 0.6 kW (kW) for 0% (diffusion), 10%, 20% & 30% of partial premixing with air. Zone C is for flames which show vortex shedding whereas Zone B demarcates flames with oscillating tip for the flames considered in the graph

3.4. Stability Maps

Different regimes (steady, oscillating tips and vortex shedding) of flames are observed for different combinations of power rating, fuel mixture composition and degrees of partial premixing with air. Thus, it is important to develop a stability map (shown in Fig. 5) which demarcates the stability regime in which the flame lies for a particular combination of the aforementioned parameters. From Fig. 5, it is observed that the increase in percentage of partial premixing has a stabilising effect on flames. For example, as shown in Fig. 5 (a), fuel mixture with 0.8 mole fraction of NH₃ exhibits lift off (regime D). But this flame transitions into a vortex shedding regime (C) on partial premixing. Also, for any given fuel composition and percentage of partial premixing with air, an increase in power rating increases the tendency of the flame to transition from the regime of oscillating tip to that of vortex shedding. It is interesting to note that for 10% and 20% partially premixed cases (Fig. 5(b)&(c)), flames with oscillating tip can be found for higher concentrations of NH₃ in fuel mixtures whereas for 30% partially premixed case, the same is found at lower concentrations of NH₃.

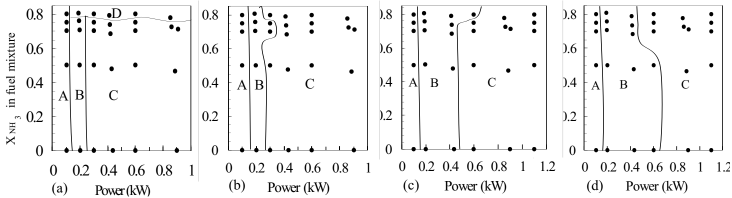


Fig. 5: Stability maps for different configurations. (a) Diffusion Case, (b) 10% partially premixed, (c) 20% partially premixed, (d) 30% partially premixed. (Regimes: A – Steady flames with no tip oscillations, B – Stable Flames with tip oscillations, C – Flames with vortex shedding, D – Unstable flames with lift-off)

3.5. Equilibrium Calculations

In this study, the stoichiometric equations for the combustion of the 5 different compositions (namely, 0:100, 50:50, 70:30, 75:25 and 80:20 ratios of NH₃ and LPG by volume) of fuel mixtures are estimated by equilibrium calculations using the software GasEQ [24]. A constant pressure adiabatic process was considered. The probable product species considered were N₂, O₂, CO, CO₂, H₂O, NO, H₂, H, OH, O and N. This was done on the basis of the concept of major and minor species [25]. However, the significant products turned out to be N₂, O₂, CO, CO₂, H₂O and NO.

It is seen that the number of moles of stoichiometric air required for the complete combustion of the fuel mixture decreases as the mole fraction of NH_3 increases in fuel mixture. As shown in Fig. 6.1, at constant pressure for one mole of these fuel mixtures, on increasing NH_3 mole fraction, a reduction in adiabatic flame temperature is observed. This is accompanied by a drop in CO and CO₂ mole fractions in the product mixture (Fig. 6.2) because of decreasing concentrations of hydrocarbon (LPG) in the fuel mixture. However, the mole fraction of NO (the most dominant contributor to NO_x emissions) remains the same even on increasing mole fraction of NH_3 in fuel mixture. This is interesting because although NO production is enhanced by NH_3 addition due to N atom contribution in the combustion zone, NO production is also found to be highly sensitive to temperature (thermal NO) which decreases on increasing NH_3 mole fraction in the fuel mixture.

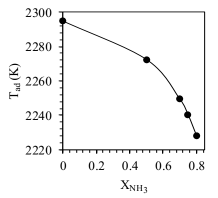


Fig. 6.1: Variation of Adiabatic Flame Temperature (K) T_{ad} with mole fraction of NH₃ in fuel mixture.

Fig. 6.2: Variation of NO, CO and CO₂ in product mixture with varying mole fraction of NH₃ in fuel mixture.

4. Conclusion

Ammonia is considered to be a good alternative for traditional hydrocarbon fuels in order to mitigate the global warming caused by greenhouse gases. However, for NH₃ to be a practical substitute, it must be blended with traditional fuels due to its lower laminar flame speed and reactivity when compared to those of conventional hydrocarbon fuels. In this work, an experimental investigation is carried out using NH₃-LPG blends in varying proportions for different power ratings (0.1-1.1 kW) by establishing both diffusion and partially premixed (10%, 20% and 30%) flames in a newly constructed single slot burner. The important findings from the study are as follows,

(i) Steady diffusion flames of lower power rating 0.1 kW are established with varying NH_3 mole fraction till as high as 0.8 in fuel mixture. It is observed that on increasing NH_3 in the fuel mixture for steady flames, there is a decrease in flame length, shortening of the luminous sooty zone and an increase in length of the non-luminous zone.

(iv) Stability maps are plotted for different configurations of fuel compositions, power ratings and percentages of partial premixing. This identifies four stability regimes among flames—(A) steady flames, (B) flames with oscillating tip, (C) flames with vortex shedding and, (D) unstable flames with lift-off. As power rating is increased, flames transition from being at regimes A to B and finally C. All diffusion flames except for 0.1 kW, with 80% ammonia (by volume) in the fuel mixture lie in regime D.

(v) From stability maps, it is inferred that there is a tendency of the flames to undergo transition from regime (C) to regime (B) as the extent of partial premixing increases.

(vi) Flames in regime B are found for larger concentrations of NH₃ in fuel mixtures for 10% and 20% partially premixed instances, whereas the same can be found for lower concentrations of NH₃ for 30% partially premixed case.

(vii) Flame lengths are found to increase with increase in power rating and decrease with increase in percentage of partial premixing with air.

(viii) At constant pressure for one mole of fuel mixture, the adiabatic flame temperature, mole fractions of CO and CO_2 in product mixture decreases with increase in mole fraction NH_3 in the fuel mixture.

(ix) NO is the primary contributor to NO_x emissions. However, for one mole of fuel mixture under constant pressure, an increase in NH₃ mole fraction in fuel does not produce drastic changes in NO mole fraction among products. This is because the effect of increasing the mole fraction of N atom containing NH₃ is nullified by the decrease in adiabatic flame temperature since NO formation is highly dependent on temperature (thermal NO).

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