Fuel-lean ammonia/biogas combustion characteristics under the reacting swirl flow conditions

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Abstract

Ammonia has been identified as a viable energy vector for power generation. Using dual-fuel operation that mixes the ammonia with higher reactivity gaseous fuel can be vital in enhancing ammonia combustion. This study examined the fundamental swirl combustion characteristics of fuel-lean premixed ammonia/biogas via a numerical approach. The flame was established at an input thermal power of 7 kW and a global equivalence ratio of 0.8. The numerical model was verified and validated with biogas emissions data acquired through experimental work. At the 20 mm burner downstream, increased carbon dioxide mass fraction in the biogas lowered the peak flame temperature up to ~400 K. Moreover, the deformation of flame temperature radial profiles was also found aggravated as carbon dioxide concentration in the biogas elevated from 0% to 40%. The reduction in premixed reactant mixture reactivity not only initiated flame temperature profile deformation but also reduced the peak Damköhler number significantly. The peak Damköhler number was lowered by a factor of ~1.5 when carbon dioxide dilution in the biogas elevated by 40%. The premixed combustion was directed into the thin reaction flamelets zone with elevated carbon dioxide mass fraction, owing to the intensified flow fluctuation. This, in turn, gave rise to the average flow velocities, turbulent kinetic energy, and normalised turbulent flame speed, notwithstanding that heat release rate and laminar flame propagation declined. In all, the presence of carbon dioxide has been shown to lower the ammonia/methane mixture reactivity whilst escalating the reacting flow fluctuation.

Keywords: Ammonia; biogas; swirl combustion; premixed combustion; dual fuel

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1.0 Introduction

Carbon-free molecules like hydrogen (H₂) and ammonia (NH₃) are identified as potential substitutes for traditional carbon-based energy sources to mitigate global carbon emissions [1]. H₂ is an appealing carbon-free molecule, but its storage is challenging and pricey. On the contrary, the energy density of NH₃ is inherently greater than that of H₂. Furthermore, storage and transportation cost for NH₃ is also substantially more economic than for H₂ [2]. During the energy crises of the 1960s and 1970s, ammonia was temporarily used as transportation fuel [3]. Since the previous decade, initiatives to accelerate global carbon reduction have once more prioritised NH₃ as a viable alternative fuel, extending its use in direct-combustion-based power production and transportation fuel [2]. However, challenges are encountered when fuelling power generation machines with ammonia.

Due to ammonia's lower flammability than fossil fuels, it was shown that considerably greater ignition energy was needed to ignite the chemical [4]. At operation near the stoichiometric conditions, the minimum ignition energy for the NH₃/air mixture was ~21.5 times greater than the premixed propane (C₃H₈)/air mixture [5]. However, it was also observed that disassociated ammonia into hydrogen at ~28% could enhance the ammonia/air combustion, thus the minimum ignition energy was lowered by approximately two orders of magnitude [5]. Ammonia's flammability limit is likewise substantially narrower than that of H₂ and methane (CH₄). Methane and hydrogen can operate in a broader equivalence ratio (φ) range of 0.5-1.7 and 0.1-7.1, respectively. NH₃, however, can only operate within an equivalence ratio of 0.63-1.40 [6]. Owing to ammonia exhibiting lower flame stability than propane, a broader quenching distance is required for ammonia flame propagation at a global equivalence ratio of 0.8, a value that is around 50 mm broader than that for C₃H₈/air flame propagation [6]. To expedite ammonia as a greener fuel for upcoming power generation, elevating ammonia combustion quality along with minimal toxic emissions are crucial.
To enhance the combustion of neat ammonia, CH₄ was suggested as one of the potential solutions. Valera-Medina et al. [7] used a common swirl burner to study premixed NH₃/CH₄ combustion at various global equivalence ratios and CH₄ mass fractions. Global $\phi > 1.25$ caused flame instability because the core recirculation zone was weaker. When $\phi > 1.1$, the NO emission was very low (~20 ppm), mostly because amidogen radicals (NH₂⁺) consumed most of the NO. On the other hand, carbon monoxide (CO) production was ~900 ppm when the global flame equivalence ratio exceeds 1.1. But in the fuel-lean combustion, reactions among the nitrogen atoms (N⁺), imidogen (NH⁺), and oxygen atoms (O⁺), altogether with nitroxyl (HNO) + H (hydrogen atom) $\rightarrow$ NO + H₂, were identified as the main sources of NO production.

In another study, blending the NH₃ with CH₄ was reported to reduce NO production in the excess air operation to ~70 ppm/kW [7], which is substantially lower than the premixed combustion of ammonia/air where NO emission was ~151.5 ppm/kW [8]. When compared to the emissions performance of biodiesel, diesel, and natural gas, NH₃/CH₄ is still quite high [9,10]. Khateeb et al. [11] recently reported that when mixture input velocity and thermal power increased, the ammonia component in the fuel blend need to be reduced to ensure flame stability.

The non-premixed NH₃/CH₄ combustion under swirling flow and elevated atmospheric pressure conditions was studied by Somarathne et al. [12]. When the energy proportion of ammonia raised by 40%, The nitric oxide generation from NH₃/CH₄ non-premixed combustion surged by about three orders of magnitude. In another study [13], it was found that excess air non-premixed NH₃/CH₄ combustion resulted in encouraging NO mitigation. In contrast to premixed ammonia/air combustion, premixed ammonia/methane/air combustion lifted nitric oxide production by ~1.3 times. As opposed to non-premixed ammonia/air combustion, non-premixed ammonia/methane/air combustion lowered nitric oxide by a factor of ~2 [13]. Additionally, it was also unveiled that premixed ammonia/methane/air combustion produced
much greater nitrogen dioxide (NO₂) and nitrous oxide (N₂O) emissions than non-premixed ammonia/methane/air combustion. It was presumed that the main mechanism for NO generation during NH₃/CH₄ burning is due to fuel-bound nitrogen [13].

In a more recent study [14], the emission of premixed ammonia/methane/air swirl flames was examined numerically by elevating the NH₃ mole fraction up to 60%. The NO transport equation computed throughout the simulation is found to improve the NO emission. The study showed that residence period, temperature, and nitrogenous components are crucial determinants of NO concentration profile. For CH₄/NH₃ flames, the inner recirculation residence time determines the NO level, and locally high NO concentration is mostly driven by prolonged local residence time. The formation of OH⁺ and NO were tied to the high temperature [14]. The investigation of the radical components unveils that HNO is the primarily responsible for the important reaction route with regards to NO formation. The H, O, and O₂ concentrations declined as the NH₃ fraction was raised (leading to elevated HNO concentration). When the NH₃ mass fraction was increased to 40%, the NO production is the highest among all cases examined [14].

Zhang et al. [15] examined the blow-off features of NH₃/air and NH₃/CH₄/air swirl flames experimentally and numerically. The findings demonstrated that the ammonia swirl flame exhibited weak lean flame stability that may be significantly enhanced by incorporating CH₄ into the fuel. Furthermore, the lean blow-off limit for the NH₃ swirl flame does not appear to be affected by the elevated swirl number. On the other hand, the swirl number for the 50 wt.% NH₃ flame results in an obvious extension of the lean blow-off limit. Four distinct flame macrostructures responsible for eventual flame blow-off were identified when lowering the NH₃/air and NH₃/CH₄/air flame global equivalence ratio. In all, the NH₃/air flame extinguished much quicker than the NH₃/CH₄/air flame, which is mostly due to the excessive stretch that results in local extinction during the blow-off process [15]. Despite studies that are dedicated
to examine and understand the fundamental combustion characteristics of NH$_3$/CH$_4$/air swirl flames, the effects of carbon dioxide (CO$_2$) addition into NH$_3$/CH$_4$/air flame have not been analysed yet. CH$_4$ and CO$_2$ are the major components in a typical biogas mixture that is commonly produced via the anaerobic digestion of organic matter. Anaerobic digestion is widely recognised as a more environmentally friendly way of producing renewable methane. Considering the scarce understanding of NH$_3$/biogas swirl combustion, the present study aims to examine it numerically. By using NH$_3$/CH$_4$ swirl combustion as baseline, the flame temperatures, velocities, turbulent kinetic energy, premixed combustion regime, turbulent Damkhöler number (Da), and turbulent flame speed for NH$_3$/biogas swirl flame are examined.
2.0 Research method

2.1 Construction of swirl burner

Figure 1a depicts the schematic diagram of a swirl burner with the integration of premixed gas to improve flame stabilisation. The plenum was designed for the premixing of fuel and air to attain a homogeneous gaseous mixture, then transfer towards the swirler to be combusted. The reference for the exit plane of the burner is \( h = 0 \text{ mm} \). The swirler hub has an internal diameter of 26 mm and an outer diameter of 40 mm. The axial swirler comprises six straight vanes with a thickness of 1.5 mm and aligned 45° angle from the axial centerline axis, resulting in a geometric swirl number of roughly \( S_N \approx 0.84 \). The introduction of the swirl is to create a central toroidal recirculation zone (CTRZ) that enhances the mixing of combustion products with high thermal energy and unburned premixed reactants. This consequently promotes cleaner combustion whilst improving flame stabilisation. A quartz tube was placed on the exit plane’s flange to visualise flames. The delivery system is depicted in Figure 1b. The dry air was delivered and controlled with a minimal deviation of 1.5% through a mass flow controller (Sierra). To replicate the biogas composition, \( \text{CH}_4 \) and carbon dioxide (\( \text{CO}_2 \)) were delivered to the mixing chamber at a specific ratio. Mass flow controllers were employed to regulate the flow rate of \( \text{CH}_4 \) and \( \text{CO}_2 \) (Sierra, 1.5% full-scale accuracy). At the burner’s exit, the mixture of air and fuel was ignited to establish the swirl flame.

2.2 Test procedures and operating conditions

Exhaust emissions like nitric oxide (NO), oxygen (\( \text{O}_2 \)), and carbon dioxide (\( \text{CO}_2 \)) were quantified by using a sampling probe connected to a gas analyser (KANE Quintox 905), and the sampling probe was placed 13 mm upstream from the quartz tube. The gas analyser can quantify NO, carbon monoxide (CO), \( \text{O}_2 \), and \( \text{CO}_2 \) in the ranges of 0-1000 ppm, 0-10,000 ppm, 0-25 %, and 0-99.9 %, respectively. The emission data were collected at five points radially
across the quartz tube outlet. By using the area-weighted average velocity approach [16], the mean average of all emissions was computed from the five points readings collected. For this study, a biogas composition of CH$_4$ (70 wt.%) blended with CO$_2$ (30 wt.%) was adopted. The flow rates of CH$_4$, CO$_2$, and air were configured to establish the swirl flame at an input thermal power of 7 kW and a global equivalence ratio of 0.8. The data collected was used for model verification and validation.

Figure 1 (a) The model of swirl burner and (b) schematic representation of the gas transport system. (Reprinted from [16] with permission from Elsevier)
2.3 Numerical modelling

2.3.1 Flamelet generated manifold (FGM)

The flamelet is simulated using a comprehensive chemistry reaction scheme in the FGM approach. The present work employed the Gas Research Institute mechanism (GRIMECH 3.0 - GRI 3.0) scheme (53 species, 325 reactions) as a chemical kinetics reaction mechanism. The FGM technique employs the chemistry from a laminar flamelet database built from many 1D flamelet computations done with full chemical kinetics and transport equations. The type of flamelet (premixed or non-premixed) is decided based on the boundary conditions of each flamelet, which vary somewhat. The premixed swirl flame was modelled in this work with the software - ANSYS Fluent. A one-dimensional premixed flamelet was constructed to solve the flamelets in reaction progress space. The following equation defines the variable reaction progress:

\[ c = \frac{\sum_k \alpha_k (Y_k-u)}{\sum_k \alpha_k (Y_k^{eq}-Y_k)} \frac{Y_c}{Y_c^{eq}} \]  

(1)

The progress variable is defined as a normalised sum of the mass fraction of the product species across all species in the chemical process. From equation (1), \( Y_k \) represents the mass fraction of the \( k^{th} \) species, \( u \) represents the unburned reactant at the flame input, and \( eq \) represents chemical equilibrium at the flame exit. Accordingly, the coefficient \( \alpha_k \) is provided so that the flame's reaction rate, \( c \), grows monotonically. \( \alpha_k = 0 \) for all species, with the exception of \( \alpha_{CO_2} = \alpha_{CO} = 1 \) for hydrocarbon combustion and \( \alpha_{H_2O} = 1 \) for fuels containing no C element, such as H\(_2\). The adiabatic flamelet equations in one dimension can be translated from physical-space to reaction-progress space.

\[ \rho \frac{\partial Y_k}{\partial t} + \rho \frac{\partial Y_k}{\partial c} \dot{\omega}_c = \rho c \frac{\partial^2 Y_k}{\partial c^2} + \dot{\omega}_k \]  

(2)
\[
\rho \frac{\partial T}{\partial t} + \rho \frac{\partial T}{\partial c} \omega_c = \rho Y_c \frac{\partial^2 T}{\partial c^2} + \frac{1}{c_p} \sum_k h_k \omega_k + \rho Y_c \left( \frac{\partial c_p}{\partial c} + \sum_k c_{p,k} \frac{\partial Y_k}{\partial c} \right) \frac{\partial T}{\partial c} \quad (3)
\]

where \( Y_k \) is the \( k \)th species mass mass fraction, \( T \) is the temperature, \( \rho \) is the density of the fluid, \( t \) is time, \( \omega_k \) is the \( k \)th species mass fraction rate, \( h \) is the total enthalpy and \( c_{p,k} \) is the \( k \)th species specific heat at constant pressure. The definition of the scalar dissipation rate \( \chi_c \) is:

\[
X_c = \frac{\lambda}{\rho c_p} |\nabla c|^2 \quad (4)
\]

Where \( \lambda \) corresponds to heat conductivity. Variable with \( c \), the scalar dissipation \( \chi_c \) is an input to the equation set, where Equation 4 corresponds to:

\[
\rho \frac{\partial Y_k}{\partial t} + \rho \frac{\partial Y_k}{\partial c} \omega_c = \frac{\lambda}{c_p} |\nabla c|^2 \frac{\partial^2 Y_k}{\partial c^2} + \omega_k \quad (5)
\]

Other than the progress variable, the mixture fraction in FGM corresponds directly to the single equivalence ratio of the 1D premixed flamelet. The maximum scalar dissipation, \( \chi_{\text{max}} \), of a premixed flamelet at varying mixture fractions is distinct. Modelling the scalar dissipation \( \chi_c (f, c) \) at any mixture fraction \( f \) as

\[
\chi_c (f, c) = \chi_{\text{max}}^\text{STO} \exp \left( -2 \left( \text{erfc}^{-1} \left( \frac{f}{f_{\text{STO}}} \right) \right)^2 \right) \exp \left( -2 \left( \text{erfc}^{-1} (2c) \right)^2 \right) \quad (6)
\]

where \( \text{STO} \) represents the stoichiometric proportion of the mixture and \( \text{erfc}^{-1} \) is the inverse complementary error function. The only model input to the premixed flamelet generator in ANSYS Fluent is the scalar dissipation at stoichiometric mixture fraction, \( \chi_{\text{max}}^\text{sto} \). The solution of unstrained (freely propagating) physical space flamelets for rich, lean, and stoichiometric hydrocarbon and H\( _2 \) flames at standard temperature and pressure [16] often matches the default value of \( \chi_{\text{max}}^\text{sto} = 1000 / \text{s} \).
To achieve a reliable simulation result, the numerical grid setup is vital. Location with high temperature and possible species concentration requires high-quality components with a slow development rate. As depicted in Figure 2, the cut shell approach, which largely comprises of a structured hexahedron grid, was selected for this simulation investigation. The minimum and the maximum number of cells in the grid are 0.7 and 1 million, respectively. Aspect ratio and orthogonal quality determined the quality of the mesh. According to Zerrin et al. [17], a hexahedron grid with a maximum aspect ratio of 35 and a minimum orthogonal quality of 0.15 is of high quality. In this instance, the highest aspect ratio was 13.43 and the minimum orthogonal quality was 0.19, which falls within the range specified by Zerrin and colleagues [17]. Near the burner outlet, a denser grid of cells was formed, while the grid grew coarser as it approached the burner outlet. At the exit of the burner, a refined mesh is required to predict the occurrence of high velocity, species, and temperature gradient.

Figure 2 The simulation setting of boundary conditions and mesh setup. (Reprinted from [16] with permission from Elsevier)
2.3.3 Fuel compositions and boundary conditions

As stated in Table 1, several biogases/ammonia compositions were used as operational fuels in this simulation. Based on these compositions, a mass flow rate model setup was built at the inlet. Simulations were performed at an input thermal power of 7 kW and $\phi = 0.8$. The turbulence intensity and hydraulic diameter at the fuel input were set to 5% and 10 mm, respectively. The turbulence intensity value is derived from the value indicated by Krieger, et al. [18]. The combustor wall was assumed to be a no-slip boundary and species flux is absent. Flow outlet at the exit of the burner was regarded as the burner outlet condition. At the outlet boundary, the static pressure was set to atmospheric surrounding pressure.

<table>
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<tr>
<th>Biogas composition</th>
<th>NH$_3$ wt.% in biogas/NH$_3$ mixture</th>
<th>Acronym</th>
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<tr>
<td>CH$_4$ wt.% in biogas</td>
<td>CO$_2$ wt.% in biogas</td>
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2.3.4 Convergence criteria

The grid independence test was performed using various numbers of elements, as shown in Figure 3a. The NO emissions for biogas converges at 1.7 million elements, irrespective of the variation in global equivalence ratio from 0.9 to 0.75. The grid number of 1.7 million was deemed grid-independent since the results were nearly comparable to those of the >1.8 million scenario. Several criteria reported by earlier scholars were used to determine the convergence of a solution. Mayr et al. [19,20] claim that the simulation is deemed to converge if the fluctuations of maximum temperature is <5 K and species concentration is <0.001 mol fraction. The residuals for mixture fraction variance and mean mixture fraction should be reduced to less than $10^{-6}$, whereas the residuals for other equations, such as continuity, velocity, and k-epsilon are less than $10^{-3}$ [19,20]. In addition to residual, the number of iterations is also an indicator of the convergence of simulation process. As depicted in Figure 3b, the variations in NO emission have no obvious changes after 5000 iterations. As the NO value fluctuates minimally and enters a steady state, the NO$_x$ mean value is regarded to have converged. Figure 3c and Figure 3d show that flame shape obtained via numerical simulation resemble that captured during the experimental work. A notable feature in 2D swirl flame image is the formation of High Momentum Flow Region (HFMR) at burner outlet. Such feature is also shown in previous studies involved swirl flames [16,21].

2.3.5 Fluid flow modelling

The mass, momentum, energy, and heat transfer equations were solved in this investigation using ANSYS Fluent with finite volume. For steady-state conditions, a pressure-based solver was utilized. Solving the RANS (Reynold-averaged Navier Stokes) equations describes the fluid flow. To close the RANS equations, turbulence models were used. In the Computational Fluid Dynamics (CFD) code, several turbulence models are provided. The
standard k- model was used for the turbulence model in this simulation. The pressure-velocity coupling was calculated using a SIMPLE computational technique [19,20]. The governing equations for momentum, turbulent kinetic energy and dissipation rate, progress variable, and mixture fraction are discretised using a second-order upwind approach. The PRESTO! system is implemented for pressure. Mayr et al. [19,20] reported that the PRESTO! technique expedites simulation convergence.

Figure 3 Emission of NO from biogas at 7 kW input thermal power as a function of (a) grid or mesh element size, and (b) the number of iterations. Biogas swirl flame images (7 kW input thermal power, $\varphi = 0.8$) obtained via (c) experiment, and (d) numerical simulation.
Figure 4a compares the experimental data with the estimated NO emissions by the FGM models at various equivalence ratios. It has been noted that the FGM-estimated NO concentration exhibits a tendency that is comparable to that of experimental data. At lean regions, prediction by the FGM technique demonstrated good agreement with experimental results, particularly at equivalence ratios of 0.85 and below. However, when stoichiometry was approached, the concentration of NO was much over anticipated. A similar trend was also observed by [16]. Premixed FGM's divergence from experimental data is mostly attributable to the species' inability to disseminate in the direction of the Z gradient. Thus, one of the variables causing the variation in NO concentration is the absence of the diffusion effect in the premixed FGM model [22].

As can be shown in Figure 4b, the deviation of the CO\textsubscript{2} species predicted by the FGM model from the actual experimental value is often less than 10%. However, when the equivalency ratio becomes closer to stoichiometric, the FGM technique underestimates the CO\textsubscript{2} emissions. According to Najafi-Yazdi et al. [23], the mass fraction is decomposed before considerable heat release, hence the progress variable frequently produces erroneous findings for rich mixes. Figure 4c demonstrates that the FGM technique accurately predicts the O\textsubscript{2} species since the error percentage is less than 10% for all equivalence ratios. For quick chemical processes, the FGM technique primarily use quasi-steady state approximation, and this method is ideally suited to forecast free radical species like O\textsubscript{2} [24].
Figure 4 (a) NO, (b) CO\textsubscript{2}, and (c) O\textsubscript{2} emissions of biogas combustion at 7 kW input thermal power obtained from experiments and numerical simulations using FGM method as a function of equivalence ratio.
3.0 Result and discussion

Figure 5a shows that for the fuel mixture with 20 wt.% NH$_3$, the flame temperature reduces by averaging 150 K as CO$_2$ wt.% increases from 0% to 40% at 20 mm downstream. In addition to the flame temperature reduction, the peak temperature is also shifted radially outwards, and the dual temperature peak are apparent when CO$_2$ wt.% increases to 40%. A similar temperature profile deformity is advanced to a lower CO$_2$ wt.% when NH$_3$ wt.% in the reactant increases to 30% and 40%, as shown in Figure 5b and Figure 5c. Figure 6 illustrates the dual peak temperature by comparing flame temperature profiles with normalised OH$^+$ intensity. It is shown that the peak OH$^+$ is present at nearly the same location where the temperature peak is found (C0N40). With elevated CO$_2$ mass fraction, dual peak OH$^+$ is formed and aligns with the radial position of dual peak temperature.

For fuel mixture without CO$_2$ infusion (i.e., C0N20, C0N30, and C0N40), peak temperature takes place at a radial position ~35 mm for 20 mm downstream, regardless of the variation in NH$_3$ mass fraction. Thus, the displacement of peak temperature to an outer radial position at elevated CO$_2$ wt.% suggests that a portion of the premixed reactant mixture is not consumed at the inner branch of the reaction zone. Conversely, it is carried to an outer branch of the reaction zone by the positive radial velocity of the swirling flow. At 30 mm downstream (Figure 5d-f), the peak temperature for all fuel mixture is stretched radially by another ~5 mm. This is primarily due to the expansion of the swirling flow diameter farther downstream. Meanwhile, increased NH$_3$ mass fraction in the absence of CO$_2$ (i.e., C0N20, C0N30, and C0N40) does not result in dual peak temperature at 20 mm downstream. This denotes that CO$_2$ is the main contributor to the dual peak temperature formation, seemingly because of CO$_2$ decelerates the reaction rate that hinders the establishment of uniform reaction across the HFMR. At 30 mm downstream (Figure 5d-f), the dual peak temperature diminishes despite the radial displacement of peak temperature remains illustrious. The shifting of peak temperature
to the outer branch and the existence of multiple reaction zones due to the addition of secondary fuel were also observed in a study pertaining to n-heptane/natural gas dual fuel swirl flame [25]. The flame peak temperature at a 15 mm burner downstream was split up and shifted by ~5 mm radially outwards when natural gas was added to the n-heptane swirl flame. Moreover, the dual peak was also seen in the normalised OH$^+$ signal at 5 mm and 15 mm downstream, respectively. For neat n-heptane swirl flames, however, no double peak temperature was observed [25]. Premixing the CO$_2$ into the CH$_4$/NH$_3$ blend has resulted in notable change in flame temperature radial profiles, especially at near burner outlet. The addition of CO$_2$ tends to stretch the peak temperature radially outwards and demands an extended downstream distance to develop a more uniform reaction radially across the HMFR.
Figure 5 Flame temperature radial profiles for CH₄/NH₃/CO₂ premixed combustion (7 kW input thermal power, $\phi = 0.8$) at (a-c) 20 mm, and (d-f) 30 mm downstream from the burner outlet.
Figure 6 Normalised OH$^+$ and temperature radial profiles for CH$_4$/NH$_3$/CO$_2$ premixed combustion (7 kW input thermal power, $\phi = 0.8$) at 20 mm downstream from the burner outlet.
Figure 7 shows that peak velocities (axial, radial, and tangential) are concentrated within the HMFR. It can be observed also that the peak velocities for fuel mixture with 40 wt.% CO$_2$ is overall higher than 0% CO$_2$ by averaging 6 m/s, irrespective of the variation in NH$_3$ mass fraction. The introduction of CO$_2$ has also altered the size and turbulent kinetic energy of the structure as depicted in Figure 8, whereas the turbulent kinetic energy for the CH$_4$/NH$_3$ is notably lower than the CO$_2$-infused fuel mixtures. Increased CO$_2$ mass fraction not only elevates turbulent kinetic energy, but it also shifts peak turbulent kinetic energy and velocities radially outward by ~10 mm.

Figure 7 Velocities profiles of CH$_4$/NH$_3$/CO$_2$ premixed combustion (7 kW input thermal power, $\phi = 0.8$) at 20 mm downstream from the burner outlet.
Figure 8 Turbulent kinetic energy for CH$_4$/NH$_3$/CO$_2$ premixed combustion (7 kW input thermal power, $\phi = 0.8$) at 20 mm downstream from the burner outlet.
With reference to Figure 5, increased CO\textsubscript{2} wt.% leads to an averaging lower flame temperature, owing to the less released chemical energy. Thus, the elevation of gas velocities and turbulent kinetic energy due to the increased CO\textsubscript{2} wt.% are apparently not because of the increased heat release rate. As illustrated in Figure 9, increased CO\textsubscript{2} wt.% directs the premixed combustion into the thin reaction zone where the fluctuation is becoming more prominent than flame laminar propagating speed. Aggravated flame fluctuation leads to the persistent local flame quenching and reignition. The eddies of premixed reactant mixture that receive an influx of heat from active species are ignited and burned irregularly, depending on the eddy size and local mixture composition [26]. This promotes the formation of a combustion wave that consequently gives rise to the gas velocities and turbulent kinetic energy as indicated in Figure 7 and Figure 8, respectively. The O\textsubscript{2}/CO\textsubscript{2} volumetric ratio has consequential effects on flame fluctuation. As O\textsubscript{2}/CO\textsubscript{2} decreased from 35% to 31% (i.e., increased CO\textsubscript{2} volume fraction in the premixed reactant), the heat release fluctuation for a CH\textsubscript{4} swirl flame increased by ~90% [27]. Although lowering the O\textsubscript{2}/CO\textsubscript{2} to < 31% resulted in relative peak fluctuation amplitude reduction by ~40%, this was accompanied by a wider band peak with multiple peaks resided, denoting the flame started to fluctuate at a wider bandwidth prior to its extinction at 21% O\textsubscript{2}/CO\textsubscript{2} volumetric ratio [27]. Increased CO\textsubscript{2} wt.% in the premixed reactant mixture tends to provoke flame fluctuation, and this basically promotes the development of a combustion wave which in turn raises the gas velocities and turbulent kinetic energy.

Figure 7 also shows that radial velocity has reduced noticeably when landing on the left-hand side of the reaction zone. The radial velocity on the right is ~12 m/s, decreasing its value on the left to ~8 m/s, a reduction of ~33%. On the contrary, the tangential velocity at the left-hand side of the reaction zone is remarkably higher than that at the right, showing an energy tradeoff between radial and tangential velocities in the flow field whereby the radial flow momentum is transferred to raise tangential velocity that consequently set up the swirl flow.
motion to anchor the flame right at the top of the burner outlet. In other study [28], the axial velocity gradient increased notably right at the vicinity of Central Toroidal Recirculation Zone (CTRZ) and the HMFR on the right-hand side of the swirl flame, but its magnitude on the left-hand side of the swirl flame was ~50% lower than that of the right-hand side, signifying flow momentum exchange in the swirling flow field [28].

Figure 9 Regimes of CH₄/NH₃/CO₂ premixed turbulent combustion (7 kW input thermal power, $\phi = 0.8$) in the Borghi diagram.
Figure 10 shows that the turbulent Damköhler number for the methane/ammonia is considerably higher than the premixed reactant mixture with 30 wt.% and 40 wt.% CO$_2$ by ~40%. In all cases, notably high Da takes place at the central region of the reaction zone and declines radially outwards. The increase in length scales of the flame and the fall in turbulence intensity cause the Da to grow in the central region. Increased length scales of the flow are reflected in Figure 7 where flow velocities are minimal in the central region of the reaction zone. The length scales decline near the boundary of CTRZ and HMFR, owing to the elevated turbulent intensity in this region and this, in turn, lowers the Da [26]. O’Doherty and Gardner [28] study on swirling flow demonstrated that a wide range of eddies appeared in the vicinity of the CTRZ and the HMFR. These eddies promote rigour mixing and at the same time elevate turbulent intensity in this region [28].

Meanwhile, increased CO$_2$ wt.% reduces the Da drastically at the central region of the combustor, this is mainly because of the substantial reduction in laminar flame speed with increasing CO$_2$ mass fraction (Figure 11b). A similar tendency is also observed when the NH$_3$ mass fraction is raised, owing to the decline in the laminar flame speed and increased flame thickness. Owing to the lower NH$_3$/air burning rate, the thickness of the reaction zone for NH$_3$/air stoichiometric laminar flame under atmospheric conditions is wider than CH$_4$/air by a factor of approximately 12 [6]. A secondary peak of Da is observed at the near-wall area, despite it is considerably lower than Da at the central region. Like the central region, the Da for methane/ammonia is distinctively higher than the premixed reactant mixture with CO$_2$ infusion. The elevation of CO$_2$ mass fraction in the premixed methane/ammonia fuel mixture has inherently led to a significant reduction in Da, mainly due the decline in mixture reactivity.
Figure 10 Da radial profile for CH$_4$/NH$_3$/CO$_2$ premixed combustion (7 kW input thermal power, $\phi = 0.8$) at 20 mm downstream from burner outlet.
Figure 11a illustrates that normalised turbulent flame speed increases in a linear fashion with normalised turbulent intensity below 1.5. As normalised turbulent intensity goes beyond 1.5, the increase of normalised turbulent flame speed turns out to be somewhat steeper. Furthermore, CH₄/NH₃ mixture with a higher CO₂ mass fraction results in a noticeably higher normalised turbulent flame speed than a fuel mixture with a lower CO₂ mass fraction. Likewise, increased NH₃ mass fraction also leads to an increased normalised turbulent flame speed when the CO₂ mass fraction in the fuel mixture is fixed. Elevated normalized turbulent flame speed is postulated because of the magnified combustion pulsation when CO₂ and NH₃ mass fractions are escalated. As shown in Figure 9, the wrinkling flamelet is diminishing with higher levels of turbulent intensity. The idea of a coherent flame front would no longer be valid in this case, and the combustion is preserved almost exclusively by the reactions established at the interface of fresh mixture’ eddies and hot combustion active species. Depending on the composition of the mixture and the magnitude of the local turbulence intensity, the flow of heat and active species consume each eddy at an irregular rate. Intensified flow pulsations are produced in this way, elevating the gas flow velocities and turbulent flame speed as a result. The influence of small turbulent eddies are consequential, it was demonstrated that when the proposed numerical model considered small turbulent eddy effects on the flame front, the estimated turbulent flame speed was more closely matched to the measured values, indicating the significance of small-scale turbulent eddies that enter the preheating layer of the premixed flame front in affecting the normalised turbulent flame speed [29]. A fragmented flame front at higher turbulent intensity generates pressure pulsation that overall elevates the normalised turbulent flame speed.

Figure 11b, however, shows that the laminar flame speed declines with the rise of turbulent intensity. This ascertains that the increased $S_T/S_L$ in Figure 11a is predominantly because of the exaggerated flow pulsations but not because of the increased reactivity when turbulent intensity is amplified. The reduction in $S_L$, on the other hand, can be attributed to the
flame stretching as flow momentum increased especially in the fuel-lean operation. The flame surface for the fuel-lean CH$_4$/air premixed turbulent flame was found to expand by a factor of approximately 3 as Reynolds number (Re) was raised by an order of magnitude [30]. With inflated flame stretching, the heat loss turns out more prominent than heat generation, and this leads to the laminar flame speed reduction as a result.

Figure 11 (a) Normalised turbulent flame speed vs normalised turbulent intensity, and (b) Laminar flame speed vs turbulent intensity for CH$_4$/NH$_3$/CO$_2$ premixed combustion (7 kW input thermal power, $\phi = 0.8$).
4.0 Conclusion

The fundamentals swirl combustion characteristics of CH$_4$/NH$_3$ under elevated CO$_2$ addition and at a global equivalence ratio of 0.8 were examined. A commercial CFD package was employed to model the flow domain and simulate the physics of the reacting swirl flow. The numerical results were verified and validated using emissions data obtained via in-house experimental works. This study unveils that the introduction of CO$_2$ has profound effects on CH$_4$/NH$_3$ combustion characteristics. Increased CO$_2$ mass fraction to 40 wt.% in the premixed CH$_4$/NH$_3$ mixture lowers the peak flame temperature by 150-400 K on average as compared to the 0 wt.% CO$_2$ case. Moreover, the peak temperature is stretched by ~5 mm radially outwards and visible radial temperature profile deformation is observed particularly in the upstream region. These are mainly due to the reduction in reactivity of the premixed reactant mixture with elevated CO$_2$ wt.%. The reduction in mixture reactivity is also reflected in the peak Da reduction by a factor of ~1.5 when CO$_2$ wt.% increases from 0% to 40%.

Furthermore, increased CO$_2$ wt.% also directs the premixed combustion from corrugated flamelet into thin reaction flamelet, primarily because of the turbulent fluctuation has prevailed over the laminar flame propagation significantly. Increased flow pulsation due to the elevated CO$_2$ wt.% raises the flow velocities and turbulent kinetic energy visibly. Furthermore, aggravated flow pulsation also has a prominent effect on turbulent flame propagation. Despite a drastic reduction in laminar flame speed, the normalised turbulent flame speed has increased by a factor of 1.6 when CO$_2$ wt.% is raised from 0% to 40%, corresponding to an increase of $u'/S_L$ from 1 to 3. Overall, the escalation of normalised turbulent flame speed is mostly due to the exaggerated flow pulsation instead of the increased heat release rate at elevated CO$_2$ wt.%.

The use of biogas (CH$_4$/CO$_2$ mixture) to replace neat CH$_4$ for co-combustion with NH$_3$ is expected to incur intensified flow pulsation, the fluctuation intensity is likely to exacerbate as NH$_3$ wt.% in the biogas/NH$_3$ mixture is increased > 40%. There is a
lot of room for system optimisation going forward, especially with appropriate choice of swirl
number, preheating temperature, and elevated atmospheric pressure.
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