Article

Study on the Combustion Process of Premixed Methane Flames with CO₂ Dilution at Elevated Pressures

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Abstract: The article presents the results of experimental and numerical investigation of turbulent premixed methane flames diluted by carbon dioxide (up to 30%) at atmospheric and elevated pressures (up to 0.5 MPa). The study included the influence of fuel properties and operation parameters on the emission of NOₓ and CO as well as flame properties. The investigation has been prepared for two combustion system configurations (axisymmetric flames and flames supported by a pilot flame) in a wide range of air/fuel equivalence ratios ($\phi = 0.42 \div 0.85$). It has been reported that reduction of NOₓ emission by CO₂ fuel dilution reached a level of up to 45% in atmospheric conditions and 30% at elevated pressure, decreasing with a drop in the equivalence ratio. The results have shown influence of pressure on NOₓ composition, where for pressurized tests, NO₂ was doubled compared to atmospheric tests. Carbon monoxide emission rises with CO₂ content in the fuel as a result of thermal dissociation, but this phenomenon is mitigated by a pressure increase. Planar laser induced fluorescence (PLIF) study has shown that flame length decreases with an increase in pressure and CO₂ content in the fuel. Fuel staging increased NOₓ emission, especially for richer flames ($\phi > 0.6$) at low pressure, while CO increased in the whole range of equivalence ratios.

Keywords: carbon dioxide dilution; nitric oxide emission; gas turbine; elevated pressure; fuel staging combustion; PLIF; flame length

1. Introduction

Reports published at the International Panel on Climate Change in Paris showed that human economic activity is responsible for global warming. Almost 200 countries committed to reduce their carbon dioxide emissions, considered to be the main cause of climate change. In the presented scenario, the lack of emissions reduction will lead to an increase in global average temperature of 3 °C. This would have catastrophic consequences for people and the natural environment. The aim of the agreement is to limit temperature growth by 2 °C, or by 1.5 °C if possible [1]. Carbon dioxide emissions should be reduced by almost half by 2030. Achieving these goals will require unprecedented transformation of the energy sector. In the 2050 perspective, it would be necessary to completely abandon combustion of hard coal and lignite. In addition, in the transitional period of abandoning fossil fuel consumption, it is necessary to develop methods of reducing carbon dioxide emissions and preventing their release into the atmosphere.

Due to imperfections of currently available renewable energy technologies (RE), network management methods, and energy storage systems [2,3], it is necessary to further develop combustion techniques that reduce carbon dioxide emissions. Currently, there are a number of methods allowing decarburization of the energy sector. These are solutions which are either in the research and development phase or already commercially available. They include carbon capture and storage
(CCS), the use of hydrogen in electricity generation, and natural gas (NG) combustion in place of coal in the transitional period.

There are many available methods allowing CO$_2$ capture related to CCS. Precombustion capture and oxyfuel combustion are the most effective; however, they are not applicable in many processes. Referring to Bounaceur et al. [4], the most cost-effective post-combustion CCS method is membrane separation. To properly function while maintaining profitability, this approach requires at least 30% CO$_2$ content in exhaust gas. For this reason, it is necessary to develop solutions allowing to increase the CO$_2$ content. Carbon dioxide containing natural gases and exhaust gases recirculation (EGR) are potential solutions to cope with CO$_2$ dilution obstacles. Combustion of gaseous fuels in gas turbines is a very difficult process because of high power concentration, elevated pressure, limitation of maximum temperature (turbine inlet temperature, TIT), and a high share of oxygen in flue gases (around 15 vol %). These parameters influence flame stability, turbine durability, cycle efficiency, and emission of pollutant compounds. Additional carbon dioxide delivered to the combustion chamber, e.g., from EGR, can affect combustion parameters such as flame temperature, laminar and turbulent flame speed, heat transfer, flame length, combustion efficiency, and NO and CO emission. A lot of research has been done on EGR application in micro-gas turbines (MGT) and gas turbines in the context of CCS [5–7]. Recently, Best et al. investigated the impact of CO$_2$-enriched air combustion in the micro-gas turbine Turbec T100 PH. They expected that 8% CO$_2$ by volume may be supplied to combustion air without blowoff (about 35% EGR). The influence of CO$_2$ on NO$_x$ reduction was in the range from 0.20 g/kWh to 1.01 g/kWh, although a minor decrease of efficiency was observed [5]. Elkady et al. [8] tested the general electric DLN combustor with EGR rate up to 40%. Their results showed significant reduction in NO$_x$ formation amounting to 47% and lower acoustic instability as well as almost constant combustion efficiency and carbon monoxide emission. A similar conclusion was conveyed by Lipardi et al. [9]; during numerical and experimental investigations of an opposed jet burner, they confirmed NO formation decrease. However, the condition for this is a sufficiently long post-flame residence time; otherwise, NO formation in the flame front is increased. Investigation of the biogas combustion process with CO$_2$ content of up to 40% in a swirl dual lean premixed combustor presented by Lee et al. [10] showed a decrease in flame temperature when the dilution rate is increased, resulting in a reduction of thermal NO$_x$ emissions. Furthermore, they showed that delivering more than 30% of a mixture of CH$_4$/CO$_2$ in the pilot mode increased NO$_x$ emission significantly, due to different temperature distribution in the combustion area. Lafay et al. [11] studied biogas combustion in a gas turbine configuration. They observed that CO$_2$-diluted methane/air flames become unstable in higher equivalence ratios compared to methane/air flames. Furthermore, the addition of CO$_2$ shifts reaction zone location and reaction intensity.

Perris et al. [12] experimentally and numerically investigated the methane/air swirl flames diluted by CO$_2$ at atmospheric pressure. They found CO$_2$ dilution limits equal to 12% and 30% for rich and lean conditions, respectively. The limits correspond to flame blowoff. Simulations performed by Giorgetti et al. showed that applying EGR to micro-gas turbines can reduce the mass flow rate of exhaust gases by as much as 50% while tripling CO$_2$ concentration compared to the traditional cycle. An additional share of carbon dioxide in the supplied fuel considered in this work can further increase the effectiveness of the MGT cycle [13].

Summarizing the previous studies, it can be noticed that fuel dilution by CO$_2$ results in:

- a decrease in flame temperature;
- a reduction of thermal NO$_x$ emissions;
- relatively slower heat release;
- suppression of combustion oscillations.

Another important issue related to the decarburization process is the wider utilization of nonstandard gases such as low-heating-value natural gases with CO$_2$ content, biogas, and landfill gas, as well as synthesis gases, especially from processes with CO$_2$ and liquid biofuels [14]. Significant
natural gas reservoirs containing high CO\textsubscript{2} concentrations are spread around the world. For example, in the Pannonian Basin region, CO\textsubscript{2} content ranges from 0.5 to 99.5% with an average value of 28\% [15]. The direct use of natural gases containing CO\textsubscript{2} increases its concentration in the flue gas and indirectly reduces carbon dioxide emissions during processing into carbon dioxide-free, high-methane gas. The aim of the performed research work focuses on understanding the CO\textsubscript{2} dilution effect on flames in the premixed methane/air mixture at elevated pressures. The obtained results can be helpful for utilization of nonstandard, high-CO\textsubscript{2}-content gases in conditions corresponding to micro-gas turbines and gas turbine operation, as well as in application to EGR for pressurized gas combustion systems.

2. Experimental Study

2.1. Test Rig

The experiment was performed in a high-pressure test rig with a maximum thermal power of 400 kW and a maximum operation pressure of 2 MPa. The combustion process of the air/fuel mixture took place in an optically accessible combustion chamber of a length of 320 mm and an internal diameter of 75 mm. The air/fuel mixture stream was delivered through a 25-mm nozzle located in the axis of a quartz tube, while the secondary fuel mixture was delivered through an array of five jets (a nozzle of a diameter of 1 mm) placed on the perimeter of the main burner nozzle (the case with a pilot flame). The secondary fuel jets were injected parallel to the main stream.

The air/fuel mixture was mixed and preheated to 630 K using an 8D electrical heater located upstream of the combustion chamber. The flame stability was provided by a sudden radial expansion by a factor of three between the inlet nozzle and combustion chamber walls. It provided external zone recirculation which stabilizes the premixed flame. The target elevated pressure was obtained using a fully automatically controlled valve located downstream of the combustion chamber. The flue gas composition was measured using a set of gas analyzers at atmospheric pressure (500 mm downstream of the pressure valve). The analyzers, manufactured by Siemens, measured carbon monoxide, carbon dioxide, and oxygen content, while for nitric oxides, the Eco-Physics analyzer was used. The main parameters of gas analyzers are presented in Table 1.

| Analyzer Name (Method) | Used Range | Accuracy |
|------------------------|------------|----------|
| Oxymat 5E O\textsubscript{2} (paramagnetic) | 0–25% | 1% of FS |
| Ultramat 6 CO\textsubscript{2} (infrared, IR) | 0–25% | 1% of FS |
| Ultramat 6 CO (infrared, IR) | 0–11% | 1% of FS |
| CLD 700EL ht NO\textsubscript{x} (chemiluminescence) | 0–0.01% | 1% of FS |

FS—full-scale operating range

The temperature inside the combustion chamber and the flue gas temperature were measured using S-type thermocouples within a range of up to 1800 K and an accuracy of 1%, while for temperature of the air/fuel mixture, a K-type thermocouple was used (a measuring range of up to 800 K at 1% accuracy). The scheme of the test rig is shown in Figure 1. All measured data collected in the experiments were registered online and recorded on a computer disk with a frequency of 1 Hz.
Aside from thermodynamic properties of the combustion process and emission of toxic compounds, the study also measured the reaction zone position in the combustion chamber as well as flame length using planar laser induced fluorescence (PLIF) techniques. In the PLIF diagnostic method, OH radicals were detected to assess the flame front position in the combustion chamber. The OH PLIF system was built on a dye laser system pumped by a solid-state Nd:YAG laser with a 9 mJ pulse at 285 nm, a pulse duration of 30 ns, and an ICCD camera with a repetition rate of 20 Hz. The 285-nm PLIF light sheet entered the reactor from the rear axial window of the tank. The signal from excited OH radicals was collected by an image intensifier camera equipped with a UV filter lens. The camera was located perpendicular to the axis of the combustion chamber. More information about the laser/camera configuration can be found in [13]. For each studied case, more than 1000 raw images were collected. The post-process adaptation of the methodology is proposed in [16], with the latest modifications described in [17].

2.2. Measurements

The experimental study was done for three fuels: methane as a comparative fuel and a mixture of methane and carbon dioxide with a volume fraction of 15 and 30 vol %. For all experiments, main fuel stream mass velocities were kept on a constant level equal to 40 m/s, while the secondary fuel velocity was changed depending on the test’s conditions. The composition and main parameters of fuels are presented in Table 2.

### Table 2. Composition and main parameters of fuels.

| No | Fuel Name | Fuel Composition | Low Heating Value (LHV) |
|----|-----------|------------------|------------------------|
|    |           | CH₄ (vol %)      | CO₂ (vol %)            | (MJ/m³)    |
| 1  | M100      | 100              | 0                      | 35.8       |
| 2  | M85       | 85               | 15                     | 30.4       |
| 3  | M70       | 70               | 30                     | 25.1       |

During the investigation, two combustion system setups were used. In the first one, the whole fuel dose was delivered to the combustion chamber by the main nozzle, while in the second case, the fuel was injected using both the main and secondary fuel nozzles at a 9 and 20 vol % share of the
volume (the pilot mode combustion, PMC), respectively. The PMC was used to increase flame stability, especially in the case of lean flammable mixtures.

The experiments were performed in a wide range of operational parameters: the air/fuel equivalence ratio ranged from 0.4 to 0.86, and absolute pressure of up to 0.5 MPa was used. The main parameters of the combustion process are presented in Table 3.

| Fuel   | M100 | M85  | M70  |
|--------|------|------|------|
| Primary air/fuel mixture temperature (K) | 633  | 628  | 631  |
| Primary air/fuel mixture bulk velocity (m/s) | 40   | 40.5 | 40   |
| Equivalence ratio | 0.42–0.84 | 0.41–0.84 | 0.44–0.86 |
| Combustion pressure (MPa) | 0.1; 0.25; 0.5 | 0.1; 0.25; 0.5 | 0.1 |
| Share of secondary fuel F2 (vol %) | 0; 10; 20 | 0; 10; 20 | 0; 10; 20 |
| Average temperature of combustion chamber (K) | 1620 | 1580 | 1611 |

Table 4. Numerical setup matrix.

| Parameter                              | Value                                      |
|----------------------------------------|--------------------------------------------|
| Fuel                                   | M100, M85, M70                            |
| Air/fuel mixture temperature (K)       | 630                                        |
| Air/fuel mixture bulk velocity (m/s)   | 40                                         |
| Investigated models                    | 1D FreeFlame, BurnerFlame                 |
| Domain size (m)                        | 0.5                                        |
| Equivalence ratio                      | 0.4–0.9                                    |
| Combustion pressure (MPa)              | 0.1; 0.25; 0.5; 1.0                       |
| Kinetic reaction mechanism             | SanDiego, Creck, GRI-Mech3.0, Tian        |

Numerical calculation results of the thermodynamic properties of laminar flames are presented in Tables 5 and 6 as a function of the pressure, equivalence ratio, and dilution rate. It was observed that
a decrease in the equivalence ratio leads to a decrease of $S_L$ and $T_A$. The same variations of laminar flame speed and adiabatic flame temperature were observed in the case of a growing ratio of CO$_2$ dilution. A more notable effect of CO$_2$ as an inert gas was observed for mixtures close to stoichiometric conditions as a result of a higher combustion process temperature. For given shares of CO$_2$ laminar flame speed and adiabatic flame temperature, values were growing with an increase in the equivalence ratio. The decrease in the value of $T_A$ as a result of thermal dissociation of CO$_2$ and H$_2$O is reduced by an increase in pressure. Since thermal dissociation occurs only at high temperatures (above 2000 K), the impact of pressure on $T_A$ is visible only for combustion of rich mixtures. There is a slight increase in temperature with a pressure increase. Based on the laminar flame speed definition proposed by Mallard and Le Chatelier [24], it can be noted that pressurization of the combustion system will lead to an increase in $S_L$ as a result of increasing density.

For each tested chemical reaction mechanism, trends were reproduced properly. However, some differences were observed, especially for laminar flame speed and emission results. The best quantitative results were achieved with SanDiego and Creck mechanisms, when the mechanism proposed by Tian underpredicted NO and $S_L$.

Radiation heat flux affected adiabatic flame temperature for atmospheric pressure from 15 K for lean methane flames up to 48 K for $\phi = 0.91$ when CO$_2$ dilution was used.

Figures 2 and 3 show a comparison of the numerical and experimental measurement results of nitrogen oxides as a function of the equivalence ratio. There is a good qualitative and quantitative agreement between the model predictions and the experimental results for the entire investigated parameter ranges, such as $\phi$, pressure, and the CO$_2$ dilution rate. Figure 4 presents calculated and measured results of CO emission. A significant discrepancy between the predicted and measured carbon monoxide was found. The trends are similar only for mixtures close to stoichiometric conditions, but there is a large quantitative discrepancy, which could be a result of the ideally premixed inlet composition and the laminar flow of the model. Moreover, a slight increase in CO emission for lean mixtures near the blowout point could not be reproduced without complex flow modeling. Both the experimental and numerical results show that CO emission increased with CO$_2$ dilution and decreased with pressurization of the combustion system.

| Table 5. Adiabatic flame temperature at elevated pressures and $T_{sub} = 630$ K (SanDiego). |
|---|---|---|---|---|---|---|---|
| Fuel | M100 | M85 | M70 |
| Pressure (MPa) | 0.1 | 0.25 | 0.5 | 1.0 | 0.1 | 0.25 | 0.5 | 1.0 | 0.1 | 0.25 | 0.5 | 1.0 |
| $\phi = 0.91$ | 2354 | 2379 | 2394 | 2408 | 2326 | 2349 | 2363 | 2375 | 2287 | 2287 | 2287 | 2287 |
| $\phi = 0.77$ | 2136 | 2219 | 2225 | 2229 | 2185 | 2193 | 2198 | 2202 | 2150 | 2158 | 2162 | 2166 |
| $\phi = 0.67$ | 1999 | 2068 | 2070 | 2071 | 2044 | 2046 | 2047 | 2049 | 2015 | 2017 | 2018 | 2019 |
| $\phi = 0.59$ | 1937 | 1938 | 1939 | 1940 | 1920 | 1920 | 1921 | 1921 | 1915 | 1915 | 1915 | 1915 |
| $\phi = 0.50$ | 1781 | 1781 | 1781 | 1782 | 1768 | 1767 | 1767 | 1768 | 1748 | 1748 | 1748 | 1748 |

| Table 6. Laminar flame speed at elevated pressures and $T_{sub} = 630$ K (SanDiego). |
|---|---|---|---|---|---|---|---|
| Fuel | M100 | M85 | M70 |
| Pressure (MPa) | 0.1 | 0.25 | 0.5 | 1.0 | 0.1 | 0.25 | 0.5 | 1.0 |
| $\phi = 0.91$ | 1.55 | 1.18 | 0.95 | 0.66 | 1.43 | 1.07 | 0.82 | 0.59 | 1.28 | 0.95 | 0.71 | 0.51 |
| $\phi = 0.77$ | 1.34 | 1.00 | 0.83 | 0.54 | 1.25 | 0.92 | 0.69 | 0.50 | 1.13 | 0.83 | 0.62 | 0.44 |
| $\phi = 0.67$ | 1.10 | 0.79 | 0.67 | 0.41 | 1.03 | 0.74 | 0.54 | 0.37 | 0.94 | 0.67 | 0.48 | 0.33 |
| $\phi = 0.59$ | 0.87 | 0.60 | 0.51 | 0.28 | 0.82 | 0.56 | 0.39 | 0.26 | 0.75 | 0.51 | 0.36 | 0.23 |
| $\phi = 0.50$ | 0.58 | 0.37 | 0.32 | 0.15 | 0.55 | 0.35 | 0.23 | 0.14 | 0.51 | 0.32 | 0.21 | 0.13 |
Both the experimental and numerical results show that CO emission increased with CO₂ dilution and decreased with pressurization of the combustion system.

**Figure 2.** Calculated and experimental NO emissions for the tested fuels at atmospheric and elevated combustion pressure (P = 0.5 MPa).

**Figure 3.** Calculated and experimental NO emissions for the tested fuels at elevated combustion pressures of 0.25 MPa and 1 MPa.
4. Experimental Results and Discussion

4.1. Emission of Pollutant Compounds

During measurements, the influence of the physicochemical properties of the gaseous fuels and operating parameters on the combustion characteristics was investigated. The first part of the measurements examined the influence of the equivalence ratio, fuel composition, and pressure in the combustion chamber on toxic compound emission. The tests were done for the first combustion chamber setup, in which the entire fuel dose was delivered by the main nozzle. The emission characteristics are presented in Figures 5, 6, and 7.

Figures 5 and 6 show the expected decrease of NO and NO\textsubscript{2} emissions as indicated by the flame cooling effect caused by addition of carbon dioxide. This effect is more significant for the tests performed at atmospheric pressure, where an increase of the CO\textsubscript{2} share in the fuel to 30 vol % caused the overall nitric oxide emissions to decrease by 45% (\(\phi = 0.83\)). It was observed that the effect of dilution on NO\textsubscript{x} emission became more significant with a decreasing equivalence ratio. Such a trend was observed for both atmospheric and pressurized tests.
Figure 5. Experimental NO and NO\textsubscript{2} emissions for the tested fuels for atmospheric combustion pressure (P = 0.1 MPa) versus equivalence ratio.

Figure 6. Experimental NO and NO\textsubscript{2} emissions for the tested fuels at elevated combustion pressure (P = 0.25 MPa) versus equivalence ratio.
The analysis of the combustion chamber pressure impact on NOx emission revealed a significant increase of NOx for both analyzed fuels. The emission rises almost four times in the rich mixture \((t = 0.85)\) when the combustion pressure changes from 0.1 to 0.5 MPa. It was observed that for pressurized tests, the composition of NOx varies and amount of nitric dioxide rises. This augmentation is caused mainly as a result of the reaction \(\text{NO} + \text{HO}_2 \rightarrow \text{NO}_2 + \text{OH}\), where \(\text{HO}_2\) is formed according to the third body reaction \(\text{H} + \text{O}_2 + \text{M} \rightarrow \text{HO}_2 + \text{M}\). The increase of combustion pressure could increase the probability of a three body reaction occurring once the mean free path of the molecules decreases [25]. It can be noticed that pressure increase above 0.5 MPa promoted \(\text{HO}_2\) formation from the OH \(\rightarrow \text{H}_2\text{O} \rightarrow \text{HO}_2 \rightarrow \text{OH}\) feedback loop; direct formation through H radicals is no longer visible.

Nitric dioxide increased in the whole range of investigated equivalence ratio values for both fuels (M100 and M85) and became more significant in richer mixture conditions, for which incomplete fuel combustion occurs during the process (see Figure 6). Numerical and experimental studies presented in [26] showed that trace amounts of hydrocarbons greatly promote nitric oxide conversion to \(\text{NO}_2\).

The influence of carbon dioxide content in the fuel as well as the impact of pressure on carbon monoxide emission are presented in Figure 7. When \(\text{CO}_2\) increases, \(\text{CO}\) is higher for the same value of the equivalence ratio. It is a result of thermal dissociation of carbon dioxide in the high-temperature region of the flame. The process of thermal dissociation increases in magnitude with an increase in the equivalence ratio (an increase in flame temperature) and decreases with pressure.

In the second part of the investigation, a new combustion setup was introduced (the pilot mode). Volume shares of 9\% and 20\% of gas were delivered as a secondary fuel through nozzles located on the perimeter of the main burner nozzle. Such a burner design was tested as a method for the increase of flame stability, especially for lean-burn. The influence of pilot mode combustion (PMC) on emission characteristics of \(\text{CO}_2\)-diluted premixed flames is presented in Figures 8–10.

**Figure 7.** Experimental CO emission at elevated combustion pressure \((P = 0.25 \text{ MPa})\) versus equivalence ratio.
In the second part of the investigation, a new combustion setup was introduced (the pilot mode). Volume shares of 9% and 20% of gas were delivered as a secondary fuel through nozzles located on the perimeter of the main burner nozzle. Such a burner design was tested as a method for the increase of flame stability, especially for lean-burn. The influence of pilot mode combustion (PMC) on emission characteristics of CO₂-diluted premixed flames is presented in Figures 8, 9, and 10.

**Figure 8.** Comparison of experimental NOₓ emissions for tested fuels for various configurations of the combustion process at atmospheric pressure (P = 0.1 MPa).

**Figure 9.** Comparison of experimental NOₓ emissions for tested fuels in various configurations of the combustion process at elevated pressure (P = 0.25 MPa).
It can be observed that introducing a portion of fuel by a pilot nozzle (a pilot flame) caused an increase in nitrogen oxides emission. The increase depends on the amount of secondary fuel and becomes significant for a volume flow rate of 20 vol%. The effect of PMC on NOx emission is stronger in the case of atmospheric pressure, especially in a richer combustion mixture ($\psi > 0.6$). This is because NOx emission is proportional to temperature, and the temperature is higher in a rich zone. The opposite effect was observed only for the fuel M70, where for richer mixtures, nitrogen oxide emission decreased (M70_PMC_20%). This may be a result of a greater radiation heat loss from the reaction zone and a decreased local temperature. It should be noted that for 9% (vol) of fuel delivered by the pilot nozzle, the influence on NOx emission is negligible for lean mixtures in both atmospheric and pressurized tests. Another measured toxic compound was carbon monoxide. Implementation of PMC caused a significant increase in CO emission in the whole range of investigated fuel mixtures and combustion pressures (Figure 10). This may be due to a weak mixing process of pilot fuel with oxidizer delivered by the main nozzle. In addition, the diffusion flame is located next to the cooled combustion chamber wall, which can result in extinguishing of the flame. The analysis of pollutant emissions from combustion systems (atmospheric and pressurized) showed that dilution of methane by CO2, from exhaust gas recirculation, for example, caused a decrease in nitrogen oxide emission, but an increase in carbon monoxide. The received values of NOx emission for lean mixtures were below the emission limits (81.4 ppmv [27]) for the gas turbine used. In addition, increasing the share of carbon dioxide above 20% (volume) in the fuel shifted emission limits by up to 195 ppmv and exceeded the limit of CO emission (280 ppmv for natural gas). It can be concluded that dilution of natural gas by CO2 provided a positive effect on emission characteristics of the investigated flames and provided better conditions for the decarbonization process of electricity production through implementing CCS in a gas turbine plant.
4.2. Flame Structure

The flame structure was studied by means of flame visualization using the method described in Section 2.1. The flame structure and the reaction zone position constitute very important information for understanding nitrogen oxide emission and flame stability. In addition, flame visualization allows calculating flame length and turbulent flame speed. Flame structures marked on the basis of the OH radical (OH PLIF) for the investigated fuels are presented in Figures 11 and 12. The presented results focus on describing the influence of fuel properties (content of CO₂) and operating parameters (pressure, φ) on the shape, reaction zone position, and flame length.

**Figure 11.** Examples of OH-PLIF images (scaled min–max) for M100 and M85 fuels, with PMC at 0.25 MPa and φ = 0.63.

**Figure 12.** Examples of OH-PLIF images (scaled min–max) for M100 and M85 fuels, with PMC at 0.5 MPa and φ = 0.63.

Figures 11 and 12 are examples of typical OH-PLIF images in lean mixture conditions (φ = 0.63) at a constant temperature and bulk velocity (T_u = 630 K, u_b = 40 m/s). The color scale represents the OH-PLIF signal, with the white color corresponding to the maximum signal. The constant color scale was used during the image analysis. It can be seen that CO₂ diluted in the fuel caused an elongation of the flame front, increasing the flame volume, and consequently leading to a more distributed heat
release. Aside from a temperature decrease through CO\textsubscript{2} dilution, a more distributed heat release weakens the process of nitrogen oxide formation. An increase in combustion pressure has an opposite effect to CO\textsubscript{2} fuel dilution. As shown, a significant reduction of flame length and volume was observed.

The flame structure changed when the combustion process was done in the PMC mode. When 9\% of fuel was delivered to the combustion system (M100\textsubscript{PMC}_9\%, M85\textsubscript{PMC}_9\%), the additional signal of OH radicals was collected next to the nozzle output. The higher intensity of OH formation appeared between the flame front and the external recirculation zone (ERZ). This suggests an increase in the reaction zone size, resulting in a more distributed heat release, slower combustion, and reduction of flame stability. When PMC reaches 20\% (M100\textsubscript{PMC}_20\%, M85\textsubscript{PMC}_20\%), the pilot burner starts to play an important role in the combustion process. Two reaction zones were observed: one in the center and the other next to the combustion chamber wall, where diffusion of the combustion process of the secondary fuel and oxidizer from the ERZ occurs. Increasing the CO\textsubscript{2} dilution rate at PMC values of 9 and 20\% did not significantly change the flame structure and did not reduce the signal intensity of the OH radicals.

For each testing point, an averaged OH-PLIF image was prepared and supplemented with two fitted curves. The first one was based on scaled OH signal strength along the burner symmetry axis, and the second presented a calculated progress variable. Flame length was determined as the most probable flame front axial position and marked as X\textsubscript{MP} for a point where the reaction progress variable was equal to 0.5. The influence of CO\textsubscript{2} dilution and the combustion setup on flame length is presented in Figure 13.

![Figure 13](image_url)

**Figure 13.** Flame length for the fuels M100 and M85 at elevated pressure (P = 0.25 MPa) versus equivalence ratio.

The flame length (X\textsubscript{MP}) increased with a raised share of CO\textsubscript{2} in the fuel, but a more significant change (elongation) can be observed when the equivalence ratio is lowered (X\textsubscript{MP} was doubled with a change in $\phi$ from 0.83 to 0.52). The use of the pilot mode combustion shortened the reaction zone for both investigated fuels.
5. Conclusions

The experimental and numerical studies of methane/CO$_2$/air flames were performed at elevated pressures. The impact of CO$_2$ dilution, pressure, mixture composition, and combustion setup configuration on the overall emission of toxic compounds and flame properties was shown. Based on the obtained results, the following conclusions can be provided:

- Experimental and numerical results showed that dilution of the methane/air mixture by carbon dioxide decreased the emission of nitrogen oxides at atmospheric and elevated pressures. The emissions drop by up to 45% in atmospheric conditions and 30% at elevated pressure. In the pressurized system, the composition of NO$_x$ was changing while the NO$_2$/NO$_x$ ratio increased.
- Introduction of CO$_2$ changed the flame structure, causing elongation of the reaction zone and increasing the flame volume, especially in lean mixture conditions. It provided a more distributed heat release, resulting in an increased flame stability.
- The combustion setup in the pilot mode combustion increased flame stability, but also increased NO$_x$ and CO emission. The effect of PMC on pollutant emission increased with an increasing share of the secondary fuel, but was mitigated by an increase in the CO$_2$ dilution rate.
- The presented model of numerical calculation of NO$_x$ emission ensured good quantitative and qualitative agreement with experimental data, indicating that it can be used as a tool for predicting NO$_x$ emission in other combustion systems.

The results of the presented study indicate the possibility of using nonstandard gases with a high content of CO$_2$ (landfill gas, biogas, low calorific natural gas), as well as introduction of an external recirculation system in pressurized combustion systems, such as micro- and gas turbines.

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**Nomenclature**

| Abbreviation | Description |
|--------------|-------------|
| CCS          | carbon capture and storage |
| EGR          | external gas recirculation |
| ERZ          | external recirculation zone |
| F1           | primary fuel |
| F2           | secondary fuel delivered in PMC |
| ICCD         | intensified digital camera |
| NG           | natural gas |
| OH           | hydroxyl radicals |
| PLIF         | planar laser induced fluorescence |
| PMC          | pilot mode combustion |
| RE           | renewable energy sources |
| TIT          | turbine inlet temperature |
| Q$_{load}$   | heat delivered to combustion chamber |
| Q$_{loss}$   | heat loss from combustion chamber |
| S$_L$        | laminar flame speed |
| T$_A$        | adiabatic flame temperature |
| T$_{sub}$    | substrates temperature |
| X$_{MP}$     | flame front position |
| u$_b$        | bulk velocity |
| $\phi$       | equivalence ratio |
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