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Experimental Study on Stoichiometric Laminar Flame Velocities and Markstein Lengths of Methane and PRF95 Dual Fuels

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Abstract

Natural gas is one of the most promising alternative fuels. The main constituent of natural gas is methane. The slow burning velocity of methane poses significant challenges for its utilization in future energy efficient combustion applications. Methane-gasoline dual fuelling has the potential to improve methane’s combustion. The fundamental combustion characteristics of a methane-gasoline Dual Fuel (DF) blend needs further investigation. In the current experimental study, the relationship between laminar flame velocity and Markstein length, with the ratio of gas to liquid in a DF blend has been investigated using spherical flames in a constant volume combustion vessel. A binary blend of primary reference fuels (PRF95) was used as the liquid fuel. Methane was added to PRF95 in three different energy ratios 25%, 50% and 75%. Values of the stoichiometric laminar flame velocities and Markstein lengths are measured at pressures of 2.5, 5, 10 Bar and a temperature of 373 K. It has been found that with a 25% increase in the DF ratio, the Markstein length is reduced by 15%, 21%, 32% at a pressure of 2.5, 5 and 10 Bar respectively whereas at the same pressures the laminar flame velocity is reduced by 2%, 3% and 5%. The flame evolution at the early stages of combustion is found to be faster with an increase in the DF ratio, and gradually as the flame develops it becomes slower.

1) Introduction

Alternative fuels have a central contribution towards compliance with future emission legislations. Attributed mainly to its low carbon content and abundance reserves, methane can be classified as one of the most
promising alternative fuels. Historically, the slow burning velocity of methane has been a major concern for its utilisation in real energy efficient combustion applications. As emphasized in literature on experimental studies in SI engines [1,2], the addition of gasoline to methane (Methane-gasoline dual fuelling) has the potential to improve methane’s combustion, leading to an enhanced initial establishment of burning velocity even compared to that of gasoline.

Practical combustion phenomena, including burning velocity in SI engines, are governed by the fundamental laminar flame velocity ($S_f^0$) of the fuel-oxidizer mixture. Since all realistic flames are curved and/or travel through a strained flow field, another fundamental mixture parameter known as the Markstein length ($L_b$), which quantifies the response of the flame velocity to stretch rate, is also necessary to characterise flame behaviour more completely [3].

Substantial efforts have been devoted for improving the understanding on methane as well as gasoline combustion. Typical refinery gasoline consists of hundreds of hydrocarbons. Iso-octane as well as binary blends of primary reference fuels have been widely adopted as convenient gasoline surrogates. Studies reporting values of laminar flame velocities at elevated pressures have been conducted for gasoline [4,5] and its surrogates [5,6,7,8] as well as methane [9,10,11]. In all the above studies the reported laminar flame velocity of methane is consistently lower compared to that of gasoline and its surrogates when tested at similar conditions. The stretch sensitivity of iso-octane and methane air mixtures characterised by the Markstein length has been also reported in literature [6,9,10]. A part of the study of Gu et al. [9] compared the Markstein length of iso-octane and methane air mixtures at stoichiometric and lean conditions. As emphasized, these two fuels responded to flame stretch differently, both with respect to equivalence ratio as well as pressure.

As stated by Brequigny et al. [12], the flame stretch sensitivity observed in the laminar regime directly impacts the combustion process in an SI engine. The study of Petrakides et al. [13] quantifies the response of mass burning rate with methane addition to PRF95 in a constant volume combustion vessel and natural gas addition to gasoline in an SI engine. A comparison of burning rates between the two experimental environments reveal very similar qualitative trends supporting the comments of Brequigny et al. that phenomena of flame velocity and stretch interactions observed in the laminar regime are still applicable in the engine environment.

The flame stretch sensitivity characterised by the Markstein length is mainly governed by the thermo-diffusive properties, the so-called Lewis number effect [14-16]. The Lewis number is defined as the ratio of thermal to mass diffusivity of the combustible mixture. It has been reported in literature that the phasing of 5%, 10% and 50% mass fraction burned in an SI engine is linearly linked to the Lewis number and therefore to the Markstein Length [12,17] of the fuel-air mixture. It has been also reported by the same research group that the burning rate of high stretch sensitive fuels such as iso-octane, slow down when high levels of flame stretch is induced on the
flame through an increase of engine speed [18]. In the particular studies [12,17,18], the considered fuel-air mixtures in the SI engine were examined at different equivalence ratios to present similar laminar flame velocities at ignition timing, and therefore allow for the effect of the fuel's stretch sensitivity on the burning velocity to be investigated. Methane being the least sensitive fuel has shown the fastest combustion, in contrast to iso-octane being the most stretch sensitive fuel shown the slowest. The interactions of burning velocity with flame stretch in SI engines have been also investigated by the study of Aleifarris et al [19]. The study reports that fuels with low stretch sensitivity have the tendency to produce faster burning velocities in the early stages of combustion.

A comprehensive understanding of these two fundamental mixture parameters, laminar flame velocity and Markstein length, is essential for the development of energy efficient combustion applications. The laminar flame velocity and Markstein length of a methane-gasoline dual fuel blend needs further investigation. It is the aim of this study to experimentally investigate the relationship between laminar flame velocity and Markstein length, with the ratio of gas to liquid in a dual fuel blend. In the current experimental work a binary blend of primary reference fuels commonly known as PRF95 (95%vol\textsubscript{liq} of iso-octane and 5%vol\textsubscript{liq} n-heptane) was used as the liquid fuel. Methane was used as the gaseous fuel. Values of stoichiometric laminar flame velocities and Markstein lengths are measured at pressures of 2.5, 5, 10 Bar and a temperature of 373 K.

2) Experimental Technique

2.1) System Integration

A 100mm inner diameter cylindrical combustion vessel with a volume of 2.2L was employed for the experimental study. The experimental setup is illustrated in Figure 1. Optical access was possible through two opposing 80mm circular windows attached near the side of the vessel. The entire vessel was preheated uniformly by a set of electrical heating elements totaling 3.2-kW. One of the heaters was fully inserted inside the vessel to induce a transient temperature difference only during the filling process. The temperature difference evoked natural convection to stir the mixture enhancing the mixing of fuel and air. Similar technique has been used by Jerzembeck et al. [5]. The interior air temperature was controlled within 3 K using a closed-loop feedback controller set to 373K. The temperature could also be observed manually from a second temperature sensor mounted on the top of the vessel. The pressure rise during the combustion process was obtained using a Kistler 6113B pressure transducer. The mixture was ignited using a slightly modified standard ignition plug with extended electrodes of 1.35 mm in diameter. The ignition system generated a spark with duration of 0.7 ms. For safety reasons, a 6 MPa pressure release valve was installed on the combustion vessel.
The flame progress recorded at 6000 frames per second with a resolution of 512X512 pixels by high speed Schlieren photography arranged in a Z configuration. A 245W halogen lamp was used as the light source. The light was focused onto a slit using a focusing lens in order to generate the spotlight for the Schlieren technique. Passing through a group of mirrors, the light path was then cut by a knife-edge which is essential for the Schlieren method [4]. Two different high speed cameras have been used for the current experimental work. A Photron Fastcam SA5 was used for the experimental work at a pressure of 5 Bar, instead of a Photron Fastcam SA-X2 that was used at 2.5 and 10 Bar. The high speed cameras were synchronized with the spark timing and the interior pressure rise recording.

2.2) Dual Fuel Mixture Preparation

As the liquid fuel, PRF95 (95% volliq iso-octane and 5% volliq n-heptane) was used. High purity (99.9%) methane was used as the gaseous fuel. The dual fuel blends consist of methane and PRF95 in three different energy ratios (25%, 50%, 75%). A blend with 25% of its energy contributing from methane as defined in Eq. 1 was labelled as DF25, with 50% DF50, and for 75% DF75.

\[
DF_{Ratio} = \frac{M_{\text{CH}_4} \times LHV_{\text{CH}_4}}{M_{\text{PRF95}} \times LHV_{\text{PRF95}} + M_{\text{CH}_4} \times LHV_{\text{CH}_4}}
\]

The air to fuel ratio was set to stoichiometric throughout the study for all investigated conditions. The stoichiometric air to fuel ratio was calculated using the method of chemical balance and assuming products of
complete combustion. High purity technical air was used with an oxidizer concentration $[\text{O}_2/(\text{O}_2+\text{N}_2)]$ of 0.2 ± 0.01.

In every experimental condition, the air to fuel ratio was prepared inside the vessel using the partial pressure method. Initially the vessel was heated up to the desired temperature (373 K). Whilst the heater mounted inside the vessel was turned on, the liquid fuel was injected into the combustion vessel using a multi-hole gasoline direct injector with an injection pressure of 12 MPa. The targeted fuel mass was supplied inside the combustion vessel by individual injections using pre-calibrated data. The pre-calibration process involves the determination of the mass of liquid per single injection. After the injections were completed, two minutes were given to allow for the complete evaporation of the liquid fuel. Considering the correct increase in pressure inside the vessel caused by the evaporation of the liquid fuel compared to the thermodynamic ideal-gas law calculations, methane and air were fed in slowly using a fine needle valve and a pressure transducer to control the filling process. The technical air was heated by an external heater before flowing into the combustion vessel to better approximate an isothermal filling process. After the filling process was completed the interior heater was turned off, and three minutes of quiescence were given to minimize any flow structures and/or temperature stratifications inside the vessel. The quiescence time also promotes the homogeneous mixing of fuel and air.

For each test condition, the described experimental procedure that allowed the evaluation of the fundamental laminar flame velocity as well as burned gas Markstein length was carried out at a minimum of three times. The average values are reported as well as error bars evaluated based on standard error.
2.3) Flame Theory

A common approach for measuring burning velocity and Markstein length in a combustion vessel has been the constant pressure outwardly propagating spherical flame method [4-10]. The method is suitable for extrapolation of measured stretched burning velocities to their fundamental non-stretched values and the associate Markstein lengths due to the well-defined stretch rates of an outwardly spherical flame. The constant pressure outwardly propagating spherical flame method in combination with the relation given by Strehlow and Savage [20] have been used by most of the studies in literature [3,4,5,7,8]. The relation of Strehlow and Savage derived on the assumption that the burned gas is coming to rest after crossing an infinitesimally thin flame such as:

\[ S_u^0 = \frac{1}{\sigma} S_b^0 \]  

Where \( S_u^0 \) is the fundamental laminar flame velocity, \( S_b^0 \) is the unstretched burning velocity, and \( \sigma \) is the thermal expansion factor defined as the ratio of unburned to burned gas density.

The fundamental laminar flame velocity is defined as the velocity at which a one-dimensional planar, adiabatic flame travels through a quiescent unburned gas mixture. The flame stretch rate can collectively describe the various influences due to flow nonuniformity, flame curvature, and flow/flame unsteadiness on the surface of an outwardly propagating spherical flame [21]. It is defined as:

\[ \alpha = \frac{2}{R_f} S_b \]  

Where \( R_f \) is the instantaneous flame radius, and \( S_b \) the stretched burning velocity corresponding to the flame radii over time, measured by an in house flame processing code.

The method developed by Markstein [22] relates the stretched burning velocity with its corresponding stretch rate. Through a linear extrapolation of \( S_b \) back to zero stretch using relation 4, the value of the unstretched burning velocity \( (S_b^0) \) and the associate burned gas Markstein length \( (L_b) \) can be obtained:

\[ S_b^0 = S_b + L_b \alpha \]  

For the Markstein theory to be satisfied exactly, it requires an unwrinkled, spherical, infinitesimally thin, weakly stretched, adiabatic, quasi-steady flame with a constant expansion factor in a zero gravity, unconfined environment [3]. These assumptions are not satisfied in practical applications, even in well-controlled experiments.

The validity of the linear relation starts to be questionable when the Lewis number of a mixture significantly deviates from unity. As reported by Kelley and Law [23], a nonlinear extrapolation between stretched burning
velocity and stretch rate should be used for mixtures with Lewis numbers appreciably different from unity. According to Halter et al. [24], the use of a nonlinear methodology is only required when the burned gas Markstein length ($L_b$) reaches or surpasses the unity value (in mm). As will be illustrated in section 3.3 the maximum value of $L_b$ measured in the current experimental study corresponds to 0.67 mm. Following the correlation derived by Halter et al. [24] for evaluating the relative percentage difference between linear and not linear extrapolation methodology, the maximum difference in the current experimental study is lower than 1.3 %. Therefore, it was concluded that in the current study a linear extrapolation methodology can still be used with confidence.

Despite its limitations, the extrapolation of a spherical outwardly propagating flame to its zero stretch using the Markstein method is widely accepted and used in literature [3-10]. This method has been applied in the present study in order to allow a comparison of the measured values of $S_c^0$ and $L_b$ with the existing related literature information.

The required expansion factors have been computed using the model for a freely propagating flame in the Cantera software package [25]. The numerical model was integrated with the reduced kinetic scheme of Jurzemberck et al [5].

2.4) Non-symmetrical Flame Restriction

In the present experimental work, the use of a cylindrical combustion vessel instead of a spherical one imposes non-symmetrical confinement on the outwardly flame evolution. According to Burke et al. [3], at flame radii ($R_f$) larger than 30% of the vessel's radius ($R_w$), the cylindrical vessel geometry excessively disrupts the induced flow field from the unconfined case, causing the motion of burned gases within the burned zone. As a result, significant departures can be experienced from the commonly employed spherical flame theory described in the previous section.

To help the reader visualize the mentioned phenomena, a symbolic illustration is presented in Figure 2. The figure presents indicative flame surface contours as experienced during the current experimental work (solid lines), in comparison to artificially symbolic circular contours that would correspond to an unconfined flame evolution (dotted lines). At the early stages (i.e a,b), the burned gas is motionless and the flame shape remains similar to that of the unconfined case. However, in contrast to the unconfined case, as the flame develops (i.e c-d-e), the burned gas deviates from its motionless state causing a non-similar flame propagation velocity along the X and Y direction. Following the work of Burke et al. [3], flames were analyzed up to a maximum radius of 15 mm ($R_f/R_w = 0.3$) to avoid any excessive motion of the burned gas that will cause departures from the applied flame theory.
As the flame propagates, the increase of pressure inside the combustion vessel is another constraint that needs to be addressed. An increase in pressure will reduce the flame velocity. As proposed in literature, the direct pressure effect on the flame velocity can be reasonably neglected when the ratio of burned gas volume to the vessel volume is less than 0.125 [3]. Within the present experimental work, at a maximum flame radius of 15 mm, the ratio of burned gas volume to the vessel’s volume is considerably lower (0.00642) than the limiting value, due to the large volume of the vessel. Therefore, the effects on the flame velocity from an increase in pressure were neglected.

![Symbolic illustration of flame surface contours for an unconfined (dotted lines) and cylindrically confined (solid lines) flame propagation process.](image)

**Figure 2.** Symbolic illustration of flame surface contours for an unconfined (dotted lines) and cylindrically confined (solid lines) flame propagation process.

### 2.5) Image Processing and Radius Definition

The flame surface was tracked with an in-house image processing code specifically developed for the current experimental setup to track flame front radii over time. Despite not being the same as the cold flame radius [26], the Schlieren image radius is commonly used in literature for flame velocity calculations [4,5,8]. The chronological change in flame radius allows for the calculation of the stretched burning velocity.

The developed technique for measuring the chronological flame radius is based on the geometrical fact that a circle can be calculated knowing at least three points on its periphery. The technique is illustrated in Figure 3. For all the experimental conditions, the technique was consistently applied from the fourth frame following the initiation of spark where the flame could be clearly observed for all test conditions. In order to avoid the effects from the electrodes, the left part of the flame’s periphery is used for the analysis. The white dots represent the points identified by the edge detection technique on the periphery of the flame, with points A and C corresponding to the upper and lower boundaries. For each particular image, points A and C are taken as the two out of three needed for the calculation of a circle. Starting from point A and moving along the flame’s periphery towards C, each single point detected is used as the additional one needed for the calculation of a circle. All of
the calculated circles are presented in Figure 3 with a green color. The average radius within one standard
deviation of all the calculated circles has been used as the equivalent flame radius at each frame. The burning
velocity \( (S_b) \) was determined from the gradient of a first-order least squares fit through four radii adjacent to each
point under consideration [5,6].

![Figure 3. Illustration of the flame detection technique.](image)

3) Results – Discussion

3.1) Flame Morphology and Evolution

3.1.1) Flame Morphology

A set of raw flame images of three different Dual Fuel (DF) ratios at a pressure of 5 Bar is presented in Figure 4.
A DF ratio of 0% corresponds to the pure liquid fuel (PRF95) whereas 100% corresponds to the gaseous fuel
(CH4). The time elapsed from the point of spark is shown. The presentation is limited at 7.93 ms as the DF 50
flame had reached the maximum allowed radius at that time. There are no signs of flame wrinkling or any
indication of cellular structures up to the maximum radius of analysis. Minor cracking can be observed on the
flame surface due to spark perturbation for all fuels. The shape of the flames appears smooth and therefore
stable independently of the fuel. As far as flame morphology is concerned, flames at a pressure of 2.5 Bar
shown consisted behaviour as in 5 Bar.

Another set of raw images, this time at a pressure of 10 Bar is presented in Figure 5. The morphology of the
flames at a randomly selected radius of about 10 mm can be observed for all the DF ratios. Flame stability at 10
Bar appears to be affected by the DF ratio. As can be clearly observed from Figure 5, the wrinkling on the flame surface is increased by moving from pure liquid (PRF95) having the largest Markstein Length, to pure gas (CH4) having the lowest. This is in contrast to the observations of flame stability at 5 Bar. As reported in literature, mixtures with low Markstein lengths have an increased propensity to instabilities [6 11 14]. Similar behaviour has been observed in the current study. The same conclusions can be drawn if a different radius is selected as a point of reference for the comparison of flame morphology of all test fuels.
For all fuels, wrinkles are triggered by the spark and remain similar in morphology as the flame expands. As proposed by Rozenchan et al. [10] and supported by L.Qiao et al. [27] at the absence of cell cracking to smaller
scales (cellularity) the linear relationship between velocity and stretch still holds. Even though the Markstein
theory can still be applied, the uncertainty in applying the theory is increased as the value of Markstein Length is
decreased.

DF ratio (%)

<table>
<thead>
<tr>
<th></th>
<th>0</th>
<th>25</th>
<th>50</th>
<th>75</th>
<th>100</th>
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Figure 5. Schlieren flame images of all DF ratios at 10 Bar. Targeted radius: 10mm

At a pressure of 10 Bar, the burning velocity of DF75 resulted to be considerably higher compared to the rest of
the fuels. The response was consistent for all of its repeats. The higher burning velocity of DF75 is thought to be
caused by phenomena of flame instability. The effects of developed instability on the flame evolution are out of
the scope of this study.

3.1.2) Flame Evolution

The evolution of a stable flame is governed by the laminar flame velocity of a fuel-oxidizer mixture, and the
sensitivity of that flame to stretch characterised by the Markstein length. At a pressure of 5 Bar, the average radii
calculated from the different repeats of each fuel are presented in the upper plot of Figure 6. For each fuel, due to
the slightly different burning velocities at respective repeats, the average radius has been calculated only up to
the time where a minimum of three radii exist (one for each repeat). The presented times are consistent with
those of Figure 4 to allow for the visualization of flame evolution of the three selected fuels CH4, DF50 and
PRF95. At respective time steps up to 6.83 ms, the percentage difference of the flame's radius of each fuel in
comparison to that of the pure liquid fuel (PRF95) has been calculated and presented in the lower plot of Figure
6. The change in flame's radius among the different DFs can be clearly observed at each time step.
At 0.83 ms after spark, it has been found that with the addition of methane to PRF95 in a dual fuel blend the flame radius is increased. Moving to 1.83 ms, DF75 is having the largest radius and PRF the smallest. The radius of methane’s flame is smaller than those of DF50 and DF75 whereas is marginally larger than that of DF25. From 1.83 to 7.93 ms, the flame evolution of the DF50 blend forms a medium between all of the test fuels and is the first to reach the maximum allowed radius at a time of 7.93 ms. The flame evolution of DF25 and PRF95 are converging towards DF50 in contrast to DF75 and CH4 that are diverging.

The studies of Brequigny et al. [17,18] present the flame evolution of methane and iso-octane flames in an SI engine. Similar qualitative trends have been found in comparison to the base fuels of the current study. During the initial stages of flame evolution, methane has been found to have a larger flame radius as compared to iso-octane, and gradually as the flame develops, the flame radius of iso-octane to converge to the radius of methane.
In the current study, similar overall trends in flame evolution could be observed at a pressure of 2.5 and 10 Bar. It has to be noted that at a pressure of 10 Bar the flame radius at the early stages of combustion was considerably higher as the dual fuel ratio was increased.

The flame evolution of the different fuels at a pressure of 5 bar is complemented with plots of burning velocity versus time and versus radius presented in the subplot shown in Figure 7. To allow for the maximum amount of data points to be presented especially in the initial period of the flame evolution, the burning velocity for each fuel has been calculated using successive radius differences, and smoothed with a second order polynomial filter only for the presentation purposes of Figure 7. The burning velocity of all fuels is initially increasing attributed to the effect of a decreasing stretch for a mixture of a positive Markstein length. PRF95 is found to give the largest increase in speed whereas methane the lowest. From 0.83 to 1.83 ms after spark corresponding to a radius of 3 mm, methane is found to be faster than PRF95 although it was slower than all DFs. Initially, the fastest burning fuel is DF50 whereas at about 2 ms after spark corresponding to 5mm in radius, the burning velocity of DF25 reaches and eventually crosses that of DF50. From a radius of 8 mm onwards, PRF95 and DF50 have comparable burning velocities whereas the velocity of DF75 is lower.

As already discussed, methane has the largest flame radius at 0.83 ms after spark. It seems that methane exhibits the fastest burning velocity only for radii below 2 mm where flames have not been analysed, as they could not be clearly observed and therefore precisely tracked by the image processing code.
The experimental study of Aleifraris et al. [19] reported the stretched burning velocity versus radius as acquired in an SI engine during the early stages of combustion for stoichiometric methane, gasoline and iso-octane air mixtures. The mass fraction burned versus time is also presented for the mentioned fuels for the whole combustion process. It has been reported that up to a radius of 15 mm the burning velocity of methane is higher than the velocity of gasoline and to a larger extent that of iso-octane. However, from a flame radius of about 10 mm and onwards the burning velocity of gasoline and iso-octane gradually converges to that of methane and eventually becomes faster as can be concluded from the available plot of mass fraction burned versus time. As it was acknowledge in the study [19], the stretch rate experienced by the flames in the engine environment is considerably higher than in constant volume laminar combustion experiments. Thus, the flame stretch sensitivity is expected to have a greater influence on the burning velocity.

With the addition of methane to PRF95 is evident that flame evolution is altered. As the dual fuel ratio increases, the flame is expanding faster at the early stages of combustion in contrast to the later stages of combustion whereas the flame is expanding slower. Similar phenomena with regards to the base fuels of the current study are also observed in real combustion applications [17-19]. In the present study, the evaluation of laminar flame velocity and Markstein length will enhance the understanding behind the mechanism of flame evolution. For the
three different test pressures, the effects of methane addition to PRF95 on both fundamental combustion parameters will be quantified and discussed.

3.2) Extrapolation of \( S_b \) to zero Stretch

3.2.1) Definition of Spark Affected Regime

At the early stages of flame evolution, the ignition energy can affect the measured value of burning velocity. As suggested by Bradley et al. [6], the sharp fall in burning velocity \( (S_b) \) with the stretch rate indicates that in this regime a fully developed flame is not yet established. Presented in Figure 8 is a selection of experimental data showing the variation of burning velocity with flame stretch rate for different fuels at a pressure of 5 Bar, as well as a single fuel (DF50) at all tested pressures. It has to be clarified that for all test conditions due to the differentiation method and the fact that the image processing code is initially applied at the 4\(^{th}\) frame after the initiation of spark, the first point of \( S_b \) in Figure 8 corresponds to the burning velocity at the 9\(^{th}\) frame after spark.

![Figure 8. Stretched burning velocity versus stretch rate for three selected fuel-air mixtures.](image)

Considering the plot of stretch burning velocity versus stretch rate, two distinct regimes can be identified; the spark affected regime and the developed flame regime. As can be found in literature [5,6,11] different researches suggest that the ignition energy effect diminishes at flame radii between 5 -10 mm. In the present study, the radius that corresponds to the upper boundary of the spark affected regime was found to be depended on the test pressure as well as fuel. When the pressure is increased the radius is decreased. At each investigated pressure, PRF95 resulted to have the largest radius, in contrast to methane that had the lowest. Thus, for each investigated pressure, burning velocities associate with flame radii less than the radius at the upper boundary of
the PRF95 flame have been excluded from further analysis. Data have been excluded for radii below 7mm at 2.5
Bar, 6mm at 5Bar, and 5mm at 10 Bar.

3.2.2) Extrapolation procedure

The unstretched flame velocity \( S_b^0 \) and the corresponding Markstein length can be determined using a linear
extrapolation through the largest possible range of radii where there is no spark influence, and where the curve of
stretch burning velocity versus stretch rate is reasonably linear [6]. The intersection of the extrapolated line back
to zero stretch corresponds to the value of the unstretched flame velocity. The gradient of the extrapolated line
corresponds to the value of the Markstein length.

Historically, the choice of data range has been somewhat arbitrary. Different researchers made different choices
without giving quantitative justification [28]. In an effort to derive the values of the unstretched burning velocities
and Markstein lengths with a consistent approach, a sensitivity analysis has been performed through the selected
reasonably linear range of radii. The overall methodology is depicted in Figure 9, where the axes have been
magnified to point out the oscillatory trend of \( S_b \) purely for presentation purposes. The observed oscillations of \( S_b \)
are induced by the unavoidable acoustic disturbances inside the vessel [5,6]. The lower boundary of the
sensitivity analysis is defined as the first point of the selected reasonably linear range. An extrapolated line is
fitted starting from the lower boundary and moving with increments of 0.5 mm in radius towards the upper
boundary. The upper boundary is defined as the point at which the value of Markstein length changes sign
compared to its initial sign at the lower boundary. Each extrapolated line within the range of sensitivity analysis is
giving a value of the unstretched burning velocity. The selected unstretched burning velocity is defined as the
average within one standard deviation of all the resulted values. The extrapolated line with its intersection giving
the closest value to that of the selected unstretched burning velocity (dashed red-blue) is used to define the value
of Markstein length. As is illustrated in Figure 8, the values of the Markstein length are defined as the slope of
the selected extrapolated lines.
Figure 9. Definition of the sensitivity analysis applied at each test condition.

For each investigated pressure, five different fuels have been tested with a minimum of three repeats per fuel. A sensitivity analysis has been performed to determine the value of $S_0^0$ at each investigated repeat. At each pressure, the average standard deviation of the unstretched burning velocities for all the tested repeats is calculated and presented in Figure 10. Also for an immediate interpretation the coefficient of variation of $S_0^0$ is also shown at each pressure. It can be clearly observed that the uncertainty in the extrapolation procedure indicated by the standard deviation of $S_0^0$ appears to increase with the decrease of pressure. This trend is attributed to the fact that the available data points within the developed flame regime and therefore the selected linear range are reduced with a decrease in pressure due to a faster flame. Summarising the current analysis, it is suggested that the reasonably linear range should be as large as possible to minimize the uncertainties from the extrapolation procedure.

Figure 10. Averaged standard deviation of the unstretched burning velocity ($S_0^0$) at each investigated pressure.
3.3) Stretch Effects - Markstein Length

The influence of stretch rate on the burning velocity is characterised by the value of Markstein length ($L_b$). For all the presented conditions in Figure 8, as stretch rate increases $S_b$ is reduced. Therefore, stretch rate has an adverse effect on the burning velocity which is indicative of a positive $L_b$. On the other hand, a negative $L_b$ indicates the increase of $S_b$ with stretch rate. Inspection of Figure 8 reveals that the difference in $S_b$ between methane and PRF95 increases as the stretch rate is reduced. That's attributed to the different values of $L_b$ between the two fuels. Up to a stretch rate of about 750 s$^{-1}$, DF50 has a higher $S_b$ even compared to that of PRF95.

The effects on $L_b$ both with respect to the DF ratio as well as pressure are depicted in Figure 11. At each investigated point, error bars are evaluated based on the standard error of all the repeated tests. The uncertainty of the extrapolation procedure and the repeatability of the tests at each investigated point are contributing to the extent of the error bars. Available literature data are also presented in Figure 11 for the base fuels. For presentation purposes, the literature data are slightly shifted on the x-axis. It appears that there is no prior work reporting values of the $L_b$ for different ratios of methane addition to PRF95 at elevated pressures. At each test pressure, the data are correlated with a straight line fit (dotted lines) aiming to present the overall trend of $L_b$ relative to the DF ratio. The equations of the fitted lines are also presented.

**Fuel effect on $L_b$:** Considering the uncertainty of the experimental results, it has been found that as the DF ratio increases, $L_b$ is decreased following a fairly linear trend. The reduction of $L_b$ with the increase in DF ratio is consistent at each tested pressure. However, at a pressure of 2.5 Bar the absolute reduction in $L_b$ is higher (larger slope) than at 5 and 10 Bar where the reduction of $L_b$ with DF ratio is similar. With a 25% increase in the DF ratio, the value of $L_b$ is linearly reduced by 0.1, 0.063, 0.056 mm at pressure of 2.5, 5 and 10 Bar respectively. As percentages the above reductions correspond to 15%, 21%, and 32%, indicating that the burning velocity becomes less sensitive to stretch as DF ratio increases. As pressure increases, the percentage difference in stretch sensitivity with the increase of the DF ratio is larger. The responses have been calculated based on the slopes of the fitted lines.
Figure 11. Burned gas Markstein lengths for all test conditions, and comparison with literature data reported by Bradley et al. (x-markers) [6], Rozenchan et al. (stars) [10], and Gu et al. (crosses) [9].

Pressure effect on $L_b$: The value of $L_b$ is not only affected by a change in fuel but is also affected by a change in pressure. As pressure increases the value of $L_b$ is reduced for all fuels as can be clearly observed in Figure 11. The reduction of $L_b$ with pressure is following a non-linear trend. The absolute reduction of $L_b$ from 2.5 to 5 Bar is larger than from 5 to 10 Bar for all fuels. For the same increase in pressure, the percentage reduction in $L_b$ is larger with the increase of the DF ratio.

Available literature data are also presented in Figure 11. Bradley et al. [6] reported values of $L_b$ for an iso-octane-air mixture at different pressures, temperatures and equivalence ratios. Appropriate values from that study are illustrated with x-markers for a comparison to the values of PRF95 measured in the current study. For methane, the reported values of $L_b$ from the experimental studies of Rozenchan et al. [10] (stars), and Gu et al. [9] (crosses), are presented. Considering the reported discrepancies of the measured Markstein lengths by different researchers [29] that can even be larger than 300%, it can be concluded that the reported values of $L_b$ from the current experimental work are in satisfactory quantitative and qualitative agreement with the selected values from literature.

### 3.4) Unstretched Burning Velocity – $S_b^0$

With the evaluation of $L_b$, the values of the unstretched burning velocity ($S_b^0$) of all fuels can now be presented. Values of $S_b^0$ are presented in Figure 12. At each investigated pressure, derived values of $S_b^0$ for all tested fuels are correlated with a straight line fit as shown by the dotted lines. The equations of the fitted lines are also presented. At a pressure of 10 Bar the value of $S_b^0$ for the DF75 blend is considerably higher compared to the rest of the fuels. As discussed in section 3.1.1, DF75 is thought to be affected by phenomena of flame instability at 10 Bar. Therefore the $S_b^0$ of DF75 is not taken into consideration for the linear fit correlation at a pressure of 10 Bar.
Fuel effects on $S_{b0}$: At a pressure of 2.5 and 5 Bar, the values of $S_{b0}$ are converging for all dual fuel ratios with a distinct difference from the values of methane. This behaviour is not evident at a pressure of 10 Bar. As an overall trend, it appears that as DF ratio increases, the value of $S_{b0}$ is decreased. The response is the same for all the investigated pressures with the exception of DF75 blend at a pressure of 10 Bar. Following the slope of the fitted lines, a 25% increase in the DF ratio, will decrease the value of $S_{b0}$ by 0.12, 0.11, 0.1 m/s at pressure of 2.5, 5 and 10 Bar respectively. As percentages these differences correspond to 4% at 2.5 Bar, 5% at 5 Bar, and 6.5% at 10 Bar.

Pressure effects on $S_{b0}$: As pressure increases the value of $S_{b0}$ is decreased for all test fuels. For an increase in pressure between 2.5 and 5 Bar, the absolute reduction in $S_{b0}$ is smaller as DF ratio is increased. At a pressure of 5 and 10 Bar the slope of the fitted lines appears to be comparable. Therefore, for an increase in pressure from 5 to 10 Bar, the absolute difference in $S_{b0}$ is similar for all fuels apart from DF75. On average (evaluated based on the difference of PRF95 and methane), the absolute reduction in $S_{b0}$ from 2.5 to 5 Bar corresponds to 0.8 m/s and 0.56 m/s from 5 to 10 Bar. The adverse effect of pressure on $S_{b0}$ is reduced as pressure is increased for all fuels.

3.5) Fundamental Laminar Flame Velocity – $S_{u0}$

The fundamental laminar flame velocity ($S_{u0}$) can be derived by dividing the already reported values of $S_{b0}$ with the appropriate expansion factors. The required expansion factors are depended both on the fuel as well as on the test pressure. At each investigated condition the computed expansion factors are presented in Figure 13. It can be observed that with the increase of the DF ratio, the expansion factor is reduced in a fairly linear manner at all three test pressures. This behaviour is mainly attributed to the different molecular weight of each fuel, with
PRF95 being the heaviest hydrocarbon under examination and methane the lightest. As far as the effect of pressure is concerned, the value of the expansion factors at 2.5 Bar is on average 2% lower as compared to the values at 10 Bar. The difference is attributed to the effect of pressure on the equilibrium state of the burned gas.

Figure 13. Computed Expansion Factors for all test Conditions

The resulted values of $S_b^0$ with their corresponding error bars are presented in Figure 14 for all the investigated conditions. At each investigated pressure, the resulted values of $S_b^0$ are well correlated with a straight line fit (dotted lines) similar to the data of $S_b^0$. The equations of the fitted lines are also presented in the figure. The considerably higher value of DF75 at 10 Bar is not taken into consideration for the fitting process. Available literature data are also included in the plot. For presentation purposes, the literature data are slightly shifted on the x-axis. For methane, data are taken from the work of Gu et al. (stars) [9]. For PRF95 data are taken from the work of Bradley et al. (x-markers) [6], Jurzembeck et al. (cross) [5], and Beeckmann et al. (triangle) [8]. It appears that there is no prior literature study reporting values of laminar flame velocities of methane-PRF95 dual fuel blends at elevated pressures.

**Fuel effects on $S_b^0$**: Considering the slope of the fitted lines as presented in Figure 14, it can be concluded that as the pressure increases, the percentage reduction in $S_b^0$ is larger with the increase of the DF ratio. With a 25% increase in the DF ratio, the value of $S_b^0$ is reduced by 2%, 3% and 5% at pressure of 2.5, 5 and 10 Bar respectively. These percentage differences are on average 2% lower as compared to those derived for $S_b^0$, attributed to the unequal expansion factors of each fuel.

**Pressure effects on $S_b^0$**: As is clearly presented in Figure 14, with the increase of pressure, $S_b^0$ is reduced. However, the reduction of $S_b^0$ is larger for an increase in pressure between 2.5 and 5Bar in comparison to an increase in pressure from 5 to 10 Bar. The adverse effect of pressure on $S_b^0$ is reduced as pressure is increased for all fuels. For the methane flame, the percentage reduction in $S_b^0$ is 2% and 5% higher than that of PRF95, with an increase of the pressure from 2.5 to 5 Bar, and from 5 to 10Bar respectively. It can be concluded that the
$S_0^0$ of methane is more sensitive in pressure than that of PRF95. This response is consisted with literature [9]. For all DFs, the percentage reduction with an increase in pressure is between the values corresponding to the pure liquid fuel (PRF95) and the gaseous fuel (CH₄).

As illustrated in Figure 14, at a pressure of 2.5 Bar the experimental values of $S_0^0$ obtained in this work are on average 11% higher compared to those reported in literature. This trend does not show on the other two investigated pressures. There is a maximum deviation of 15% between the values of $S_0^0$ obtained in this work as compared to the ones reported in literature. The maximum deviation corresponds to the value of PRF95 at a pressure of 10 Bar.

With the evaluation of both fundamental combustion parameters $L_b$ and $S_0^0$, the mechanism behind the flame evolution as discussed in section 3.1.2 can now be explained. At a pressure of 5 Bar, with a 25% increase in the DF ratio, the values of $S_0^0$ and $L_b$ are reduced by 3% and 21% respectively. As already discussed, at the early stages of combustion the flame radius is increased with DF ratio. It is clear that the mechanism behind this phenomenon is attributed to the decrease of $L_b$ as the dual fuel ratio is increased. As the flame develops and flame radius is increasing, stretch rate is reduced. This implies that the effect of $L_b$ on the flame velocity is decaying. Therefore $S_0^0$ will start to dominate the flame evolution. As a result, an increase in the DF ratio will slow down the flame evolution. Indeed, the flame evolution of PRF95 becomes gradually faster than that of methane as the combustion process progress.

4) Conclusions

The effects of methane addition to PRF95 on the fundamental combustion parameters, laminar flame velocity ($S_0^0$) and Markstein length ($L_b$), were experimentally investigated at a stoichiometric air to fuel ratio, different
pressures (2.5, 5, 10 Bar) and a constant temperature of 373 K. A Dual Fuel (DF) blend was formed by adding
methane to PRF95 in three different energy ratios 25%, 50% and 75%. Spherically expanding flames were used
to measure burning velocities, from which the corresponding \( L_b \) and \( S_{u0} \) were derived. Where applicable, values
obtained from this work were compared with reported data in literature. It appears that there is no prior work
reporting values of either \( L_b \) or \( S_{u0} \) for different DF ratios at elevated pressures.

As far as \( L_b \) is concerned, it has been found that with a 25% increase in the DF ratio, the value of \( L_b \) is reduced
by 15%, 21%, 32% at a pressure of 2.5, 5 and 10 Bar respectively. As pressure increases, \( L_b \) is reduced for all
fuels. The absolute reduction of \( L_b \) from 2.5 to 5 Bar is larger than from 5 to 10 Bar. For the same increase in
pressure, the percentage reduction in \( L_b \) is larger with the increase of the DF ratio. A satisfactory qualitative and
quantitative agreement with the appropriate values from literature was obtained.

As far as \( S_{u0} \) is concerned, it has been found that with a 25% increase in the DF ratio, the value of \( S_{u0} \) is reduced
by 2%, 3% and 5% at pressure of 2.5, 5 and 10 Bar respectively. As pressure increases, \( S_{u0} \) is reduced for all
fuels. For the same increase in pressure, the percentage reduction in \( S_{u0} \) is larger with the increase of the DF
ratio. There is a maximum deviation of 15% between the values of \( S_{u0} \) obtained in this work and those reported in
literature.

At the early stages of combustion, the flame evolution is found to be faster with the increase in the DF ratio, and
gradually as the flame develops it becomes slower. At the early stages of combustion \( L_b \) has a dominant effect on
the flame evolution. As the flame develops, stretch rate is reduced, and \( S_{u0} \) becomes the governed parameter for
the flame evolution.

5) References

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