**Abstract**

This paper reports computational results, to complement experimental observations, on the turbulence–chemistry interaction of nonpremixed jet flames issuing into a heated and highly diluted oxidant stream. It is found experimentally that large-scale vortices and flame stretch can lead to spatial thinning and a decrease in OH concentration. This reduction in OH is described as a weakening of the reaction zone. Accompanying reaction zone weakening is also an increase in H₂CO levels. The reduction in reaction rates is most noticeable at low oxidant stream O₂ levels. The heated and low oxygen oxidant conditions typify those of Moderate or Intense Low oxygen Dilution (MILD) combustion. The computational results indicate that the effects of the low oxygen levels of MILD combustion leads to both a reduction in reaction rates and an increase in transport of O₂ across the reaction zone. The relationship between the reaction rate and level of O₂ permeation suggests that a form of partial premixing can occur under MILD combustion conditions. This partial premixing leads to the formation of flame intermediates which contribute to the stabilisation of the flames. The permeation effects are most pronounced at high strain rates, which are commonly encountered in practical MILD combustors.

**Key words:** Reaction zone weakening, MILD combustion, JHC burner

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1 Introduction

Moderate or Intense Low oxygen Dilution (MILD) combustion is a particular combustion regime which involves the strong recirculation of exhaust gases back into the reaction zone (Cavaliere and de Joannon, 2004, Choi and Katuski, 2001, Wünning and Wünning, 1997). The subsequent reduction in localised \( O_2 \) concentration leads to a distributed reaction zone, reducing pollutant emissions (notably \( NO_x \)) and an increase in nett radiation flux (Cavaliere and de Joannon, 2004, Weber et al., 2000). Application of MILD combustion has been successfully applied in numerous applications, additionally incorporating the use of biomass fuel (Hi-TACG, 2008). Despite much progress in the application of MILD combustion to practical systems, there remain unresolved issues on the fundamental stabilisation, auto-ignition, and structure of the reaction zone near the jet exit under the hot and diluted conditions (de Joannon et al., 2005, Maruta et al., 2000).

To achieve the very high recirculation rates required for MILD combustion high velocity jets are often used (Cavaliere and de Joannon, 2004). The high strain rate conditions lead to a reduction in turbulence time-scales, coupled with larger chemical time-scales as a result of the low oxygen conditions, these flames are associated with low Damköhler numbers (Katsuki and Hasegawa, 1998). As an extension to this, in a furnace environment MILD combustion has been likened to a well-stirred reactor (WSR) (Plessing et al., 1998, Weber et al., 1999). Based on the concept of attempting to infer MILD combustion from a well-stirred reactor, de Joannon et al. (2000) has attempted to model a WSR with MILD combustion conditions. However as identified in that work the authors acknowledge that a well-stirred reactor is unfeasible for a practical combustor. Furthermore, the interaction between turbulence and chemistry in describing the nature of MILD combustion is expected to
play a significant role (Katsuki and Hasegawa, 1998). The combustion is controlled
by both the kinetics and the mixing (Milani and Saponaro, 2001), and so the anal-
yogy to a WSR may not always be applicable to practical systems. The importance
of both chemical and turbulent timescales has been shown by Galletti et al. (2007)
who reported Damköhler numbers near unity in MILD conditions.

To advance the fundamental understanding of practical MILD combustion furnaces
beyond WSR theory, there have been efforts to study MILD combustion furnaces
on an experimental level (e.g. (Plessing et al., 1998, Szegő et al., 2008)). Through a
series of subsequent studies (Coelho and Peters, 2001, Dally et al., 2004, Özdemir
and Peters, 2001) using the furnace of Plessing et al. (1998), it has been reported
that computational studies qualitatively match the experimental findings, except
in the flow-field near the jet and the NO formation. A similar observation was
made between the numerical work of Mancini et al. (2002) for the furnace of
Weber et al. (2000). The comparisons between the numerical and experimental
results from these furnaces highlights that there are aspects of the stabilisation
and turbulence–chemistry interaction under MILD combustion conditions which
remain unresolved.

To bridge the knowledge gap between the overly simplified WSR theory, and the
complex interactions within a furnace environment (even on an experimental level),
Dally et al. (2002a,b) reported on the structure of turbulent nonpremixed jet flames
issuing into a heated and diluted coflow. This burner configuration, referred to as
a jet in hot coflow (JHC) burner, emulates MILD combustion under simplified and
well-controlled conditions. These authors used single-point Raman-Rayleigh-LIF
diagnostic techniques to simultaneously measure temperature, major and minor
species at different locations in these flames. It was found that major changes in the
flame structure occur when reducing the oxygen concentration and that, at higher
jet Reynolds number and low oxygen concentration, oxygen leakage from the surroundings is related to local extinction of the flame. Medwell et al. (2007) extended that work by simultaneous imaging of the hydroxyl radical (OH), formaldehyde (H$_2$CO) and temperature in the same burner, finding evidence of partial premixing in these flames, and localised extinction in the presence of surrounding air.

It has been identified that the flame stabilisation mechanism appears fundamentally different under MILD conditions. Different stabilisation characteristics have been noted in a MILD combustion furnace (Szegő et al., 2007). Under the hot and diluted coflow conditions, in a JHC burner, pre-ignition reactions have been identified upstream of what appear to be lifted flames (Medwell et al., 2008). The presence of flame intermediates (viz. OH and H$_2$CO) in the apparently lifted region of these flames lead to the term transitional flames being adopted to describe this phenomenon. In both transitional and attached flames, H$_2$CO has been detected along the jet centreline (at $x/D=7.6$), suggesting diffusion of oxygen atoms well into the reaction zone. Further differences in the stabilisation of MILD combustion is evident by noting that the apparent liftoff height decreases with an increase in jet velocity (Medwell et al., 2008). This finding is consistent with the observation in a furnace environment that increasing the jet velocity helped stabilise the jet flame through enhanced mixing (Dally et al., 2004). Furthermore, Mancini et al. (2007) concluded a similar effect of “preconditioning” of the fuel whereby significant mixing occurs prior to the consumption of the fuel.

Using a JHC burner (Dally et al., 2002a,b, Medwell et al., 2007, 2008), this study aims to examine the structure of the reaction zone of a jet in a heated and diluted coflow. Temperature, the hydroxyl radical and formaldehyde are measured instantaneously and simultaneously using planar laser imaging techniques. The hydroxyl radical (OH) is used as a flame marker while the formaldehyde (H$_2$CO) interme-
The product of $[\text{OH}]$ and $[\text{H}_2\text{CO}]$ has also been suggested as an indicator of the formyl (HCO) radical, which is closely related to the heat release rate (Najm et al., 1998).

In this paper we report on the effect of turbulence–chemistry interaction on the reaction zone structure under hot and diluted oxidant stream conditions. Using the same experimental conditions presented in a companion paper (Medwell et al., 2007), a JHC burner is used to emulate MILD combustion. The fuel used is a mixture of natural gas & hydrogen, and issues into a coflow with a 3% $\text{O}_2$ level. Laminar flame calculations are used to complement, and provide additional insight into, the reaction zone structure effects which were observed experimentally.

2 Experimental Details

The MILD combustion burner used in this study is the jet in hot coflow (JHC) burner used previously (Medwell et al., 2007, 2008), and shown in Figure 1. It consists of a central insulated fuel jet ($\varnothing 4.6\text{mm}$) within an annular coflow ($\varnothing 82\text{mm}$) of hot exhaust products from a premixed secondary burner mounted upstream of the jet exit plane. The coflow $\text{O}_2$ level is 3% (volumetric), with an exit temperature of 1100K. The fuel used in the jet is natural gas ($92\% \text{CH}_4$) diluted with hydrogen ($\text{H}_2$) in an equal volumetric ratio. The addition of hydrogen reduces the levels of soot ( advantageous for laser techniques) and improves flame stability. Addition of $\text{H}_2$ also has implications for the potential use of hydrogen as a supplemental fuel additive. The jet Reynolds number for the experimental data presented in this paper is 15,000.
Laser induced fluorescence (LIF) is used to image OH and H$_2$CO, and temperature is inferred from Rayleigh scattering measurements. The laser pulses are fired sequentially to reduce interferences on the other systems, with the entire sequence occurring in 300ns to ensure the flow field is effectively frozen with respect to the fluid time scales. The in-plane resolution of all three ICCD cameras is 160µm, after spatial matching. The laser sheet heights were all ∼12mm, of which the central 8mm portion is presented herein. All images are corrected for laser power and profile variations shot-to-shot based on the signal from a laminar slot burner. Description of the experimental details is described in-depth in a previous publication by the authors (Medwell et al., 2007).

3 Laminar Flame Calculations

Laminar flame calculations have been performed to extend the study beyond the available measurements, and shed more light on the structure of the reaction zone and molecular transport. The OPPDIF routine of the Chemkin package is used to compute temperature and species concentration for opposed-flow diffusion flames. For all calculations the GRI-Mech 3.0 mechanism is used. Previous studies have shown the GRI mechanism to provide agreement with experimental measurements of the JHC burner used in this study under similar conditions (Christo and Dally, 2005, Dally et al., 2002a, Kim et al., 2005).

The opposed-flow laminar diffusion flame configuration represents a one-dimensional flame, analogous to the traverse across the well-defined reaction zone from fuel to oxidant. By increasing the velocity of the flow from two facing nozzles, the strain rate imposed on the flame front can be varied. The strain rate quoted throughout most of this paper is the average normal strain rate reported in the OP-
PDIF post-processor output. The use of a laminar, one-dimensional, configuration provided by the OPPDIF models is a well established methodology to enable the role of strain alone to be de-coupled from the more complex turbulent interactions that are observed experimentally.

Three different oxidant stream temperatures and compositions are used for the calculations, as listed in Table 1. The major species concentrations for the experimental conditions used in the previous publication (Medwell et al., 2007) are found from equilibrium calculations based on the coflow inlet flowrates. The oxidant stream temperature is 1100K (to match the measurement from the JHC burner). For comparison, standard air conditions are included as well. The jet composition is an equal molar ratio of CH₄ & H₂ (i.e. the natural gas is assumed pure methane).

Since the coflow oxidant stream consists of combustion products (H₂O and CO₂), the standard definition of mixture fraction is not appropriately defined for calculations based on the mass fraction of H & C (hydrogen & carbon) atoms. A normalised mixture fraction, \( \xi^* = (\xi - \xi_{oxi}) / (\xi_{fuel} - \xi_{oxi}) \) is used instead, where \( \xi_{fuel} \) & \( \xi_{oxi} \) refer to the standard definition of mixture fraction at the fuel and oxidant stream boundaries, respectively.

4 Results and Discussion

4.1 Motivation

Figure 2 shows a selection of instantaneous image triplets of OH, H₂CO, and temperature from the JHC burner (repeated from Medwell et al. (2007)). The jet Reynolds number is 15,000 and the coflow O₂ concentration 3% (volumetric). The
images are centred at 35mm downstream of the jet exit plane. The corresponding size of each image is 8mm in height and 30mm wide. The jet centreline is marked by the vertical dashed line.

Typically, near the jet exit the images do not show evidence of large-scale vortices. The images suggest that the reaction zone appears unconvoluted, as shown in Figure 2a. Nonetheless, a proportion of the images do show signs of large-scale vortices, as indicated by convolution of the temperature and/or flame species. As the flow becomes more convoluted and stretched by vortices, the OH images can show a localised decrease in concentration and a spatial thinning. The phenomenon relating to reductions in the OH layer is described as a “weakening” of the flame front, and forms the basis of this paper. An example of a weakened reaction zone is seen in Figure 2b.

The interaction of the vortices with the reaction zone intuitively suggests that flame stretch leads to the observed reduction in OH concentration. Over a range of strain rates up to extinction, the laminar flame calculations presented in Figure 3 suggest that the peak OH number density does have a dependence on the strain rate. Similarly, the peak temperature is also dependent on the strain rate. The strain rate presented in Figures 3 & 4 is defined as the maximum absolute normal strain rate across the reaction zone. Apparent from Figures 3 & 4 is that the flames in the heated and diluted oxidant conditions can sustain a much higher strain rate before extinction. It is also noted that the OH number density measured is of a similar magnitude to that predicted by the laminar flame calculations presented in Figure 3.

For comparison of the strain rate values to turbulent conditions, the scalar dissipation ($\chi$) is determined from the gradient of the mixture fraction ($\xi$);
\[ \chi = 2\mathcal{D} (\nabla \xi \cdot \nabla \xi) \]  

(1)

The diffusion coefficient \( \mathcal{D} \) is determined from the mole-fraction weighted mean of the mixture averaged diffusion of the fuel \( k = \text{CH}_4 \) and \( \text{H}_2 \) components, defined as:

\[ D_{km} = \frac{1 - Y_k}{\sum_{j \neq k} X_j / D_{jk}} \]  

(2)

The binary diffusion coefficients \( D_{jk} \) are determined based on polynomial fit with temperature provided by the TRANSPORT library of CHEMKIN.

For the two heated oxidant streams the approximate maximum scalar dissipation rate prior to extinction is \( \sim 1200 \text{ s}^{-1} \) for the 3% \( \text{O}_2 \) and \( \sim 6000 \text{ s}^{-1} \) for the 9% \( \text{O}_2 \) oxidant. At standard oxidant conditions the peak scalar dissipation of \( \sim 150 \text{s}^{-1} \) was estimated. These results confirm that MILD combustion can sustain much higher strain/scalar dissipation than under conventional oxidant stream conditions.

Experimental measurements of \( \text{H}_2\text{CO} \) show an increase with Reynolds number (Medwell et al., 2007). An increase in \( \text{H}_2\text{CO} \) with strain is also noted in the laminar flame calculations presented in Figure 5. \( \text{H}_2\text{CO} \) has also been shown to increase with the extent of partial premixing (Mc Enally and Pfefferle, 2000, Medwell et al., 2007). To investigate the possibility of a connection between the increase in \( \text{H}_2\text{CO} \) with strain and partial premixing, the \( \text{O}_2 \) concentration at the location of peak \( \text{H}_2\text{CO} \) is plotted against strain in Figure 6.

Figure 6 suggests a relationship between strain and partial premixing is plausible. Since \( \text{H}_2\text{CO} \) is formed on the fuel-rich side of the reaction zone, the \( \text{O}_2 \) concentra-
tion at the location of peak \( \text{H}_2\text{CO} \) is indicative of the amount of \( \text{O}_2 \) which has been transported across the flame front. The molecular transport of \( \text{O}_2 \) to the fuel-rich side is suggestive of a form of partial premixing, and is seen to increase with the strain rate. The presence of oxygen on the fuel-rich side of the reaction zone has been noted previously in MILD combustion conditions (Kim et al., 2005).

4.2 \textit{Formaldehyde Production}

The \( \text{H}_2\text{CO} \) production rates have been analysed from laminar flame calculations. For each oxidant stream composition, two strain rate conditions have been chosen to differentiate between low and high strain cases, \( \text{viz.} \ 30 \text{ s}^{-1} \) and \( 200 \text{ s}^{-1} \) (averaged normal strain rate, corresponding to maximum absolute normal strain rates of \( \sim 50 \text{ s}^{-1} \) and \( \sim 500 \text{ s}^{-1} \)). Previous studies have shown the validity of laminar flame calculations to model these flames when using moderate strain rates, of the order of \( 30 \text{ s}^{-1} \) to \( 200 \text{ s}^{-1} \) (Dally et al., 2002a).

Five main reactions have been found to be responsible for the majority of \( \text{H}_2\text{CO} \) formation and consumption. The key reactions identified are (numbered in accordance with GRI);

\[
R10: \quad \text{O+CH}_3 \rightarrow \text{H+H}_2\text{CO} \\
R56: \quad \text{H+H}_2\text{CO} (+M) \rightarrow \text{CH}_2\text{OH} (+M) \\
R58: \quad \text{H+H}_2\text{CO} \rightarrow \text{HCO+H}_2 \\
R101: \quad \text{OH+H}_2\text{CO} \rightarrow \text{HCO+H}_2\text{O} \\
R127: \quad \text{CH+H}_2\text{O} \rightarrow \text{H+H}_2\text{CO}
\]
For the 3% O\textsubscript{2} oxidant stream case, Figures 7a & 7b show the production rate of H\textsubscript{2}CO plotted against mixture fraction for the five selected reactions at two different strain rates. Reactions R10 and R127 are seen to be the main contributors to H\textsubscript{2}CO production. H\textsubscript{2}CO production via R10 increases by a factor of ten between the two strain rates considered. In comparison, R127 increases by a factor of three. The increase of H\textsubscript{2}CO production as the strain rate is increased is consistent with Figure 5. Accompanying the increase in H\textsubscript{2}CO production is also an increase in the consumption rate. Reactions rates for R58 and R101 increase by a factor of five and three, respectively, between the two strain rate cases. Nevertheless, the large increase in the reaction rate of R10 with strain over-compensates for the differences in consumption, leading to the higher H\textsubscript{2}CO concentration at the higher strain condition.

The rate of H\textsubscript{2}CO production via reaction R10 for the two strain cases under consideration is shown in Figures 8a & 8b. Also shown in these figures are the species relevant to H\textsubscript{2}CO production via reaction R10 (\textit{viz.} O and CH\textsubscript{3}). A vertical line is included as a reference point at the location of the peak nett H\textsubscript{2}CO production rate. At this mixture fraction, it is noted that the O concentration is very low for the low strain case, but is significantly (five fold) higher at the high strain case. This difference in O concentration between the two strain cases, coupled with a two fold increase in CH\textsubscript{3}, seems to explain the relative reaction rate of R10.

An increase in the concentration of O at the location of peak H\textsubscript{2}CO production seems to be one of the main contributors to the increase of H\textsubscript{2}CO with strain. Investigation of reaction rates reveals that the major production of O is from O\textsubscript{2}, via reaction \textit{R38}: H + O\textsubscript{2} $\rightleftharpoons$ O + OH. The production rate of O via reaction R38, and H & O\textsubscript{2} mole fractions, are also shown in Figures 8a & 8b. It is apparent that the production rate of O at the location of peak nett H\textsubscript{2}CO is ten fold greater at
the higher strain rate. This is predominately attributed to a five fold increase in O₂ concentration at the location of peak nett H₂CO between the two strain cases.

It is acknowledged that there are many concurrent processes that occur as the strain rate is increased. Nevertheless, the most significant increase in H₂CO with strain seems to occur via the following path;

\[
\text{O}_2 \xrightarrow{R38} + \text{H} \xrightarrow{R10} + \text{CH}_3 \rightarrow \text{H}_2\text{CO}
\]

The identified pathway relates the increase in H₂CO with strain to an increase in O₂ concentration at the location of peak nett H₂CO production. For these laminar nonpremixed calculations, any O₂ at the location of peak H₂CO must have been transported from the oxidant stream. This transport of O₂ across the reaction zone is likened to partial premixing. The possibility of O₂ existing on the fuel rich side of the reaction zone is not unheard of; in conventional nonpremixed flames O₂ has been measured along the centreline towards the base of an attached flame (McE-nally and Pfefferle, 1999).

It is worth noting that both the experimental measurements (Figure 2) and the laminar flame calculations (Figure 8) show a broad H₂CO distribution on the fuel-rich side of the reaction zone. Figure 7 indicates a cessation of significant H₂CO production/consumption above \(\xi^* \gtrsim 0.015\), yet high H₂CO concentrations are seen much further into the fuel-rich side (Figure 8). This is strongly indicative of diffusion of H₂CO towards the fuel-rich boundary, and is consistent with previous studies that have shown broad H₂CO distribution is a result of diffusive and convective transport effects (Fotache et al., 1997a,b).
4.3 Transport Effects

As the strain rate increases, it has been demonstrated in this paper that the O\textsubscript{2} concentration on the fuel-rich side of the reaction zone increases. The O\textsubscript{2} level on the fuel-rich side of the reaction zone is controlled by the competition between transport from the oxidant stream and consumption. The nett O\textsubscript{2} consumption rate (i.e. the opposite of production rate) for the two strain cases is shown in Figure 9. For reference purposes, key species mole fractions are also included on this plot. To aid in visualisation, since transport is a physical process, Figure 9 is plotted against distance, instead of mixture fraction. The distance origin is defined as the location of peak HCO\textsubscript{3} with the fuel-rich side denoted as negative, and the fuel-lean side positive. Although constant between the two plots, the range extends only over the region of interest, and not to the fuel and oxidiser boundaries. As expected, as the strain rate is increased it is apparent there is a spatial compression of the reaction zone. The width of the reaction zone (defined as the distance between the peak OH and H\textsubscript{2}CO, as markers of the fuel-lean and -rich peaks, respectively) reduces \(\sim2\)-fold with the increase in strain.

From Figure 9, the peak O\textsubscript{2} consumption rate is seen to be \(\sim5\) times higher at 200\textsuperscript{s\textsuperscript{-1}} compared to 30\textsuperscript{s\textsuperscript{-1}}. The increase in peak consumption rate is due in part to the spatial compression of the physical distances at the higher strain and therefore higher concentrations of the radicals involved. Nevertheless, the integrated O\textsubscript{2} consumption is \(\sim3\) times higher at the higher strain rate case.

Despite an increase in the consumption of O\textsubscript{2} with increased strain rate, it has been seen that more O\textsubscript{2} permeates the reaction zone. To compare the differences in transport, the velocity of O\textsubscript{2} for the two strain cases is also included in Figure 9.
The O\textsubscript{2} velocity ($v_{O_2}$) is defined as the combination of the bulk velocity ($u$) and the diffusion velocity ($V_{O_2}$) (Yamada et al., 2003). Using the O\textsubscript{2} velocity, the mass flux transport of O\textsubscript{2} ($j_k = \rho_k Y_k v_k$, with $k = O_2$ (Reaction Design, 1999)) has also been added to Figure 9. It is apparent that the transport flux is significantly higher as the strain rate is increased. Coupled with the spatial compression of the reaction zone with increased strain rate, there is a significant increase in the transport of O\textsubscript{2}.

It is therefore apparent that the increased O\textsubscript{2} transport, and reduced spatial scales, associated with the increase in strain rate over-compensate for the increased reaction consumption rates. The result is that more O\textsubscript{2} is capable of penetrating the reaction zone. This observation is consistent with the instantaneous images showing increases in H\textsubscript{2}CO near the location of local weakening of the flame front (Figure 2b).

4.4 Comparison between oxidant stream conditions

In the conditions analogous to MILD combustion presented so far, an increase in transport of O\textsubscript{2} across the reaction zone has been observed. To compare these heated and diluted conditions to more conventional flame conditions, Figure 10 presents the same details as shown in Figure 9, but now for a 21% O\textsubscript{2} & 300K oxidant stream. The permeation effects under consideration are primarily prevalent at high strain rates, and so data is presented only for the $a \approx 200s^{-1}$ condition. The scaling for each of the species is the same between the plots, but the vertical axis range is different to accommodate the differences in the relative intensities. Despite the seven-fold increase in oxidant O\textsubscript{2} concentration, it is apparent that under the conventional oxidant stream conditions there is significantly lower O\textsubscript{2} levels on the fuel-rich side of the reaction zone compared to MILD conditions. In order for
the higher O\textsubscript{2} level in the oxidant stream to be reflected on the fuel-rich side, the transport must also increase compared to the increase in consumption. The peak O\textsubscript{2} consumption rate is 7 times higher at the 21% O\textsubscript{2} level, whereas the O\textsubscript{2} mass flux only increases approximately 4-fold in comparison to the 3% O\textsubscript{2} & 1100K condition. In combination, the increase in consumption relative to transport counteracts the higher oxidant stream O\textsubscript{2} concentration in conventional conditions. In effect, the reaction zone becomes more permeable to O\textsubscript{2} under MILD combustion conditions. The increase in O\textsubscript{2} concentration on the fuel-rich side with a reduction in oxidant O\textsubscript{2} level is consistent with Figure 6.

To bridge the gap between the 3% O\textsubscript{2} & 1100K and 21% O\textsubscript{2} & 300K oxidant, Figure 11 shows the same details again, but for a 9% O\textsubscript{2} & 1100K oxidant stream. As expected (from Figure 6), it is seen in Figure 11 that the O\textsubscript{2} level on the fuel-rich side is significantly reduced at 9% O\textsubscript{2} as compared to the 3% O\textsubscript{2} oxidant stream. The three-fold difference in the O\textsubscript{2} level is accompanied by a three-fold increase in O\textsubscript{2} consumption rate. The increased O\textsubscript{2} level at 9% as compared to 3% leads only to a two-fold increase in O\textsubscript{2} flux. Again, the increase in consumption is greater than the increase in flux, such that the O\textsubscript{2} permeation to the fuel-rich side is less.

It is apparent that the lower the O\textsubscript{2} concentration in the oxidant stream the higher the degree of permeation of O\textsubscript{2} across the reaction zone. The lower reaction rates as a result of the low O\textsubscript{2} conditions associated with MILD combustion therefore lead to a form of partial premixing. These findings suggest that molecular transport and finite rate chemistry effects are essential in order to capture the stability and structure of these flames. The challenge remains, however, to capture the transition from a MILD regime to conventional flames (Christo and Dally, 2005).
4.5 Stabilisation

In the low oxygen & high temperature oxidant stream conditions presented, it has been seen that transport of $\text{O}_2$ across the reaction zone is responsible for an increase in the formation of key flame intermediates (notably $\text{H}_2\text{CO}$). As the strain rate is increased, these intermediates continue to be generated. The production of such flame intermediates is responsible for the stabilisation of the reaction.

For flames similar to those presented in the current paper, Medwell *et al.* Medwell *et al.* (2008) identified flame intermediates and pre-ignition reactions upstream of an apparent liftoff height. Due to the presence of intermediates in what is generally considered the liftoff region, the term “transitional” flames was adopted to describe these unique conditions.

As the velocity (hence strain rate) approaches lift-off, rather than the transitional flames becoming completely lifted, the formation of intermediates continues. It is proposed that in the “lift-off” region a pool of intermediates is produced. These precursors initiate the combustion reaction, and are believe to greatly enhance the stabilisation of the flame. The build-up of a pool of precursors (notably $\text{H}_2\text{CO}$) in jet flames in a similar high temperature coflow has been suggested by Gordon *et al.* (2007).

5 Conclusion

MILD combustion is a promising combustion regime offering simultaneous reductions in emissions and an increase in thermal efficiency. Laminar flame calculations in the low oxygen and high temperature conditions of MILD combustion have been
shown to lead to simultaneous reduction in reaction rates and increased transport across the reaction zone. Enhancing these effects, experimentally it is shown that interaction with large-scale vortices can lead to a spatial thinning of the reaction zone. The turbulence–chemistry interaction under the heated and highly diluted oxidant stream is referred to as reaction zone weakening. The effects of strain is particularly important for MILD combustion since practical combustors typically rely on high velocity jets to achieve the highly diluted conditions.

As the flame-front is stretched at the low oxygen (3% O$_2$) and high temperature ($T_{\text{ini}}=1100\text{K}$) conditions, reaction zone weakening manifests itself in two ways; the strain effect leads to an increase in H$_2$CO, and large-scale vortices lead to a reduction in OH concentration. The low O$_2$ conditions lead to a reduction in reaction rate, that is, a weaker reaction zone (as compared to higher O$_2$ oxidant stream conditions). When the strain rate is increased, the effects of weakening, combined with a spatial compression of the reaction zone, contribute to a greater degree of O$_2$ transport across the reaction zone. Therefore, in MILD combustion a form of partial premixing occurs. Large-scale turbulent mixing due to large-scale vortices can lead to a weakening of the flame front. This observation demonstrates the importance of turbulent mixing in the establishment of the MILD combustion regime.

Reaction zone weakening, which leads to a form of partial premixing, may contribute to the stabilisation of the MILD combustion reaction zone. Previous studies have shown that an increase in jet velocity helps stabilise flames under MILD conditions (Dally et al., 2004, Medwell et al., 2008). The partial premixing effects that have been identified provide an increase in the formation of flame intermediates, and thus an improvement in the stabilisation with jet velocity.
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Table 1
Oxidant stream temperature and composition (molar basis) used for laminar flame calculations.

<table>
<thead>
<tr>
<th>Oxidant</th>
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<th>9% O₂</th>
<th>21% O₂</th>
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</tbody>
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