CHARACTERISING HYDROGEN MICROMIX FLAMES: COMBUSTION MODEL CALIBRATION AND EVALUATION

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ABSTRACT

Hydrogen micromix combustion is a promising concept to reduce the environmental impact of both aero and land-based gas turbines by delivering carbon-free and ultra-low-NOₓ combustion without the risk of autoignition or flashback. The ENABLEH2 project aims to demonstrate the feasibility of such a switch to hydrogen for civil aviation, within which the micromix combustion, as a key enabling technology, will be matured to TRL3. The micromix combustor comprises thousands of small diffusion flames for which air and fuel are mixed in a cross-flow pattern. This technology is based on the idea of minimizing the scale of mixing to maximize mixing intensity. The high-reactivity and wide flammability limits of hydrogen in a micromix combustor can produce short and low-temperature small diffusion flames in lean overall equivalence ratios.

For hydrogen-air mixtures there is a need to further characterize the physical importance and calibration process of the laminar Schmidt (Sc), Lewis (Le) and Prandtl (Pr) and turbulent Schmidt (Sc) numbers. In addition, there is limited numerical and experimental data about flame characteristics and emissions of hydrogen micromix combustor at high pressure and temperature conditions.

In this paper, the CFD software STAR-CCM+ was used with the FGM (Kinetic Rate) combustion model to simulate and calibrate hydrogen micromix flames. The research was divided into two parts. In the first part, the values of laminar Schmidt (Sc), Lewis (Le) and Prandtl (Pr) and turbulent Schmidt (Sc) numbers were obtained from the flamelet tables. A Turbulent Schmidt number was then obtained by comparing RANS and LES simulations of a single injector. If Sc$t$>0.2, the predicted NOₓ production of RANS simulations approaches that of LES; while Sc$t$<0.2 provides similar overall flame structure between RANS and LES. It is concluded that, for the current simulations, Sc$t$=0.2 is a good compromise between flame structure and emissions prediction.

Flame characteristics and NOₓ emissions given by Thickened Flame and FGM Kinetic Rate models in a single injector geometry were also compared.

Keywords: Hydrogen; Micromix; Combustion; Calibration; ENABLEH2; STAR-CCM+; FGM Kinetic rate; Thickened Flame.

NOMENCLATURE

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<thead>
<tr>
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<td>α</td>
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**INTRODUCTION**

The aerospace sector, as well as the automotive sector, is under the most restrictive anti-emissions law in history and these low-emission requirements are going to be further increased as time goes by. For the aerospace industry, CO2 and NOx are mainly the emissions that must be mitigated. In order to make it possible, the European programs Vision 2020 and Flightpath 2050 aim to reduce these emissions by setting goals and funding projects to achieve them. The goals target a reduction of CO2 and NOx emissions by 50% and 80% in 2020 and by 75% and 90% in 2050 respectively relative to 2000 levels [1, 2]. In contrast, according to ICAO, with the current growing trend of the civil aerospace sector, CO2 emissions are expected to increase [3]. This is clearly a fact that goes against EU objectives and must be addressed urgently since the aviation sector is responsible for roughly the 5% of man-made global warming [4].

At this point, a clear conclusion can be extracted: even with more electric propulsion, EU objectives are very challenging and might not be fulfilled. In this situation, alternative fuels such as hydrogen may be part of the solution.

Hydrogen is proposed as fuel not only because its combustion does not produce carbon emissions but also because of its high reactivity. This property has mainly two consequences that tend to reduce NOx production. Hydrogen flames are characterised by lower residence time therefore lower NOx. Hydrogen is able to burn at much leaner equivalence ratio than kerosene [5, 6], which potentially reduces local flame temperatures. However, if fuel-to-air mixing is not adequate, its high calorific value leads to higher flame temperature at stoichiometry, increasing NOx production.

Since H2 properties are very different from hydrocarbon fuel (mainly higher reactivity, diffusivity and calorific power [7]), a conventional combustion system is not suitable for air-to-hydrogen mixing. If it is used in a conventional combustion chamber, relatively high equivalence ratio regions are produced hence slightly higher NOx emissions than kerosene combustion are generated [6].

Micromix combustion system allows to make use of the full potential of hydrogen combustion by reducing significantly NOx emissions. This technology was developed based on the idea of minimizing the scale of mixing to maximize the mixing intensity [6]. In order to do that, a micromix combustor is composed of thousands of small cross-flow injectors. By this way, for a given point of operation, the amount of fuel required is distributed among numerous orifices ensuring lean diffusion combustion in short flames, thus reducing high local temperatures and residence time hence NOx production.

Recently, ENABLEH2 project [9] was launched in order to develop this technology to meet emission targets for 2050. The aim of this project is to mature LH2-based propulsion technologies to TRL4 to enable free-CO2 and ultra-low-NOx engines in the future. This paper is part of the research carried out by Cranfield University in ENABLEH2 project.

Developing technology to TRL4 means to create a system/subsystem model or prototype demonstrator in a relevant environment [10]. Fortunately, nowadays CFD software tools are very powerful and are used extensively in industry. In order to raise the TRL of a technology whose physics have been understood but not accurately predictable in any operating condition, validation of CFD models is a necessary step for future optimization of the technology. In this paper, preliminary analyses were carried out to establish best practices for CFD modelling of H2/air cross-flow combustion. Validation of CFD simulations will be performed in the following years with the experimental data that will be generated by Cranfield University as part of ENABLEH2 project.

The high diffusivity of H2 makes the mixing characteristics of any H2/air mixture different from those of any hydrocarbon/air mixture. Therefore, the molecular properties of the mixture are different as well. These fluid characteristics are usually expressed by means of the Lewis (Le), Schmidt (Sc) and Prandtl (Pr) numbers. For CFD simulations, knowing the value of these numbers is critical to implement the molecular properties of the fluid in the simulations. However, in the literature there is a lack of information about the value of these parameters at different temperatures, pressures and equivalence ratios. In this paper, the basic guidelines to obtain these numbers from correlations and their values for a wide range of temperatures, pressures and equivalence ratios are provided.

In RANS simulations, the mixing produced by turbulence is modelled through the turbulent Schmidt number (Scₜ). This parameter only depends on the flow characteristic and has been estimated as 0.2 in simulations for non-reactive cross-flow patterns [11]. However, the changes in the flow pattern produced by combustion and their subsequent effect on Scₜ have not been assessed. The value of this number was re-evaluated for H2-micromix flames.

The flame regime defines the turbulence-chemistry interactions for any flame. In any simulation, knowing the flame regime is critical in order to choose the CFD combustion models since they define the chemistry and the turbulence-chemistry interactions. In the literature, the Borghi diagram has been extensively used for almost any type of flame regardless of the fuel. However, using it with H2-micromix diffusion flames is not justified because it implicitly assumes Sc = Le = Pr = 1 and should only be used for pre-mixed flames. Since Sc, Le and Pr for H2-micromix flames differ from 1 and they are diffusion...
flames, a new flame diagram was proposed to establish the flame regime and preliminarily choose the CFD combustion models.

In order to model combustion, Flamelet Generated Manifold (FGM) and Thickened Flame models seem to be promising. FGM model has been used during recent decades providing good predicting capabilities for solving hydrocarbon and hydrogen combustion [12, 13, 14, 15, 16]. The flamelet model generates beforehand tabulated values for the combustion. Then these flamelets are integrated in the actual simulation by using the progress variable and the mixture fraction parameters, computed from their corresponding transport equation. Tabulated values calculated prior to the simulation decouples chemistry with flow simulations hence reduce considerably the computational cost of solving the combustion phenomenon [12]. Thickened flame model, used only for LES simulations, has been recently used as a promising model to calculate flame physics at sub-grid scale with relatively coarse mesh. This is achieved by artificially thickening the flame front to solve it in detail while keeping constant laminar flame speed [17, 18]. The results provided using these two models were compared to provide a deeper insight on the different possibilities to model this type of flames.

Along this paper, a know-how about the calibration and model-selection process for H_2-micromix combustion simulation is provided.

1 Physics of H_2/air mixtures

The first step before running any numerical study is to identify the physics behind it in the range of p and T that are going to be simulated. This could be done by estimating the molecular and turbulent properties of the working fluid and the expected value for the interest variables. For this it is necessary to conduct a thorough literature review to get correlations and mixing models that can describe the fluid and flow field properties. This task is especially important in hydrogen-fueled cross-flow combustion since most of the work done for combustion is usually related to hydrocarbon fuels in other types of flow field such as the toroidal re-creation vortex in conventional combustion chambers.

Regarding the properties of the fluid, the usual way of characterizing them is through the Schmidt (Sc), Lewis (Le) and Prandt (Pr) numbers:

$$Sc = \frac{v}{D} ; \quad Le = \frac{a}{D} ; \quad Pr = \frac{v}{\alpha} ;$$

(1)

Traditionally, the value of these parameters was assumed 1 for hydrocarbon-fueled combustion. However, this is not correct when the fuel is hydrogen, since its high molecular diffusivity implies lower-than-1 Le and Sc numbers. Due to the relevance of the molecular and thermal diffusivities in the mixing process and flame structure of any combustion system, these non-dimensional numbers must be known for all the operating conditions.

The molecular properties that are required to obtain these parameters are the dynamic viscosity (\(\nu\)), the density (\(\rho\)), the compressibility factor (\(z\)), the thermal conductivity (\(k\)) and the isobaric specific heat (\(C_p\)). In this section the correlations that were used to calculate these properties are referenced and the values of the Sc, Le and Pr numbers for a wide range of pressures presented.

1.1 Dynamic viscosity

Lemmon et al. [19] developed a correlation to calculate the dynamic viscosity of air and validated it with REFPROP 7.0 database. All the predicted values fall within ±5% of error for T \(\in\) [100,1000] K and p \(\in\) [1,1200] bar.

Yusibani et al. [20] developed a similar correlation to predict the dynamic viscosity of H_2 valid in the range of T \(\in\) [100,1000] K and p \(\in\) [1,2200] bar with a maximum error of ±4% validated with experimental results.

In both correlations the dynamic viscosity is calculated as 
\(\mu = \mu_0 + \mu_r\), where \(\mu_0\) is the dilute dynamic viscosity (independent of pressure) and \(\mu_r\) the residual fluid viscosity (representing the effect of pressure).

1.2 Density and compressibility factor

These two parameters are calculated together. First, correlations and data from tables are used to calculate the compressibility factor. Finally, this parameter is used in the ideal gas law to calculate density (1).

$$\rho = \frac{p}{zRT}$$

(2)

The compressibility factor of air is obtained from linear interpolation in temperature and pressure from experimental data obtained from Perry’s chemical engineers’ handbook [21]. The range of validity for this correlation is T \(\in\) [75,1000] K and p \(\in\) [1,500] bar. The uncertainty of using this data is unknown.

To calculate the compressibility factor of hydrogen, Lemmon et al [22] developed a simple correlation based on experimental results provided by the software REFPROP. Their method is valid in the range of T \(\in\) [200,1000] K and p \(\in\) [1,1200] bar with a maximum error around ±0.01%.

1.3 Thermal conductivity

Air thermal conductivity was also correlated by Lemmon et al. [19] with a formulation similar to the one used for dynamic viscosity with an additional term that includes the enhancement of thermal conductivity near critical conditions. This correlation is valid in the range of T \(\in\) [100,1000] K and p \(\in\) [1,1200] bar with an uncertainty lower than ±5% compared with experimental results.

The correlation developed by Assael et al. [32] to estimate the thermal conductivity of H_2 was used. It has an uncertainty lower than ±7% in the range of T \(\in\) [100,1000] K and p \(\in\) [1,1000] bar compared to experimental results.
1.4 Isobaric Specific Heat

Isobaric specific heat was considered to be independent of pressure. In order to get its evolution with temperature, the correlations found in the book An Introduction to Combustion Concepts and Applications [24] are used. H\textsubscript{2} specific heat is directly obtained from these correlations while for air, N\textsubscript{2} and O\textsubscript{2} specific heats are first obtained and then a weighted average value is calculated using the mass fractions.

1.5 Molecular Diffusivity

Molecular diffusivity describes how two or more fluids mix without the presence of a forced convection field according to the Fick’s law. Since H\textsubscript{2} is the lowest concentration species that diffuses into air before combustion (lean equivalence ratio), the molecular diffusivity coefficient (\(D\)) is taken as the binary diffusion coefficient of H\textsubscript{2} into air.

A pressure-and-temperature-dependent binary diffusion model to estimate \(D_{\text{H}_2/\text{air}}\) is extracted from the book Gas Transport in Solid Oxide Fuel Cells [25]. This correlation is based on the kinetic theory of gases, which could induce some errors depending on the condition. The constant 0.00197 is different from that found in [25] (0.00186) since it has been adjusted to match the experimental data found in [26].

1.6 Mixture models

In order to know the averaged value of \(\mu, k, C_p\) and \(\rho\) for any mixture at a given condition knowing these properties for each species individually, a mixture model is needed. They are based on experimental results and can be as simple as using a weighted average depending on the mole fraction or far more complex depending on the fluid property.

The process that was followed to calculate the properties of the mixture is to firstly obtain the temperature of the mixture by applying energy conservation, then the properties of air and H\textsubscript{2} separately at the corresponding temperature and a chosen pressure, finally to apply the mixture models.

The isobaric specific heat of the mixture is obtained when calculating the temperature through energy conservation as a weighted average of the specific heats of the different species using the mass fractions.

For both dynamic viscosity and thermal conductivity the Wilke’s law of mixing, derived from the kinetic theory of gases, is extensively used and shows good agreement with experimental measurements [27].

Density is obtained from the partial pressures of each species and the ideal gas law. This is equivalent as a weighted average using the mole fractions.

1.7 Laminar Sc, Le and Pr numbers

After correlating all the fluid properties and applying the corresponding mixture models, the non-dimensional numbers were calculated. First, they were calculated for \(T \in [200,1000^\circ\text{C}], p \in (1,5,10,20) \text{ bar}\) and \(\phi = 0.5\) to see how they change with pressure and temperature for a given mixture (figure 1). Finally, they were calculated for \(T \in [200,1000^\circ\text{C}], \phi \in (0.3,0.5,1,2)\) and \(p = 10 \text{ bar}\) to get their variation with the equivalence ratio (figure 2).

![FIGURE 1: Evolution of Prandtl number with \(p\) and \(T\) at \(\phi = 0.5\)]

According to the first calculation (constant \(\phi\)), the fluid non-dimensional numbers are mostly insensitive to pressure and temperature in the studied range. The biggest variation, although not very significant, is observed in the Prandtl number (figure 1). This is because of the small variation in the dynamic viscosity and the thermal conductivity when only increasing pressure. In the case of the Schmidt number, the divergent isobaric lines at low temperature are due to the effect of the compressibility factor (since \(\mu\) depends on \(\rho\) and \(D_{\text{H}_2/\text{air}}\) on \(p\), therefore \(z\) is not neglected). The insensitiveness of these parameters to \(p\) and \(T\) suggests that for a given equivalence ratio, assuming any of these parameters to be constant does not deviate too much from reality. However, the typical assumption of \(Le = 1\) is indeed far from the actual physics, especially for H\textsubscript{2}/air lean mixtures. For the leanest equivalence ratio (0.3) where NO\textsubscript{x} are the lowest, the Schmidt, Lewis and Prandtl numbers are 0.22, 0.3 and 0.75 respectively.

![FIGURE 2: Evolution of Schmidt, Lewis and Prandtl number with \(\phi\) and \(T\) at \(p = 10 \text{ bar}\)]

If different mixtures of H\textsubscript{2} and air are considered, it is possible to find differences in the non-dimensional numbers. However, they do not change more than 0.15 when comparing lean and rich mixtures. \(Le\) takes as much a value of 0.4, still far from 1. The values of \(\phi \in (0.3,0.5,1,2)\) have been chosen to be those that could represent a diffusion flame behaviour since
\( \phi > 2 \) are not expected due to the fast chemistry of hydrogen combustion and the enhanced mixing of micromix combustion system. From the values and the variation of these parameters it is possible to conclude that:

- Since \( Le \sim 0.3 \), the expected flame reactants diffuses around 3 times faster than its energy. This, added to the high laminar flame speed of \( H_2 \), creates a short flame with a thin flame front. However, flame front thickness also depends on the absolute value of the thermal diffusivity, so at low pressure it may increase and produce a short flame but slightly longer than at high pressure because of a thicker flame front. In addition, the high molecular diffusivity, compared to the thermal diffusivity, is likely to generate more unstable flames because the flame is supplied with fuel faster than its capability to evacuate the energy in the flame front.
- \( Sc \sim 0.25 \) in laminar flow means that \( H_2 \) to air mixing by diffusion is 4 times more significant than the capability of the injected \( H_2 \) to get the momentum that the air jet is transmitting to it. This has an effect on the penetration of \( H_2 \) in the air jet given a momentum-flux-ratio. However, the laminar value of \( Sc \) alone plays a minor role in cross-flow since mixing and momentum transport are enhanced by turbulence. It is necessary to estimate the turbulent Schmidt number to model the effective cross-flow mixing characteristics.

### 2 Flame regime characterization

Flame characteristics can be described by means of the Damköhler number (\( Da \)) and the Karlovitz number (\( Ka \)). The former relates the characteristic time of the turbulence integral scales (mixing) with the reaction time (2). High \( Da (>1) \) means high reaction speed flames where mixing controls flame propagation. Although similar, the definition of the latter is related to turbulence-to-chemistry interaction. It is the ratio of the reaction time to the characteristics time of kolmogorov turbulent scale (length scale where the turbulence dissipates (3)). It can also be expressed as the flame thickness to kolmogorov scale. Low \( Ka \) (<1) means the turbulence dissipates before being reduced to a size smaller than the flame thickness, therefore turbulence can not progress inside the flame front and it can be considered as a flamelet, i.e., a turbulent flame than can be considered as a differential set of laminar flames.

\[
Da = \frac{\tau_c}{\tau_{ch}} = \frac{\tau_c}{u' \delta_t} \tag{3}
\]

\[
Ka = \frac{\tau_{ch}}{\tau_\eta} = Da^{-1} Re_t^{1/2} \tag{4}
\]

\[
Re_t = \frac{u' \ell_t}{v} \tag{5}
\]

Hydrogen-fuelled flames are expected to be characterized by high \( Da \) and low \( Ka \) since their laminar flame speed is higher than that of any conventional fuel (2 – 3 m/s). However, chemical, integral and kolmogorov time scales must be estimated to know what type of flame hydrogen-fuelled micromix flames are. The outcome of this analysis served to know what flame regime it is expected and what are the most suitable CFD combustion models that should approach the physical behaviour of these flames.

To estimate \( Da \) and \( Ka \), the laminar flame thickness (\( \delta_l \)), the laminar flame speed (\( S_l \)), integral turbulence scale (\( l_t \)), turbulent characteristic speed (\( u' \)) and kinematic viscosity (\( v \)) are needed. Among these parameters, viscosity is known from the correlations in previous section, integral turbulent scale is taken as the hydraulic diameter of the duct or air gate introducing the air into the combustion region and the turbulent characteristic speed is estimated as the 40\% of the flow mean velocity (30 m/s) from preliminary LES simulations. The laminar flame front is estimated using eq. (6) and the laminar flame speed obtained from Ravi et al. [5]. This correlation follows the form of eq. (7) and has an uncertainty of \( \pm 13 \) cm/s in the range of \( T \in [270,620] \) K, \( p \in [1,30] \) atm and \( \phi \in [0.5,5] \).

\[
\delta_l = 2 \frac{a}{S_l} \left( \frac{T_{ref}}{T_l} \right)^{0.7} \tag{6}
\]

\[
S_l = [a_1 + a_2 \phi + a_3 \phi^2 + a_4 \phi^3 + \ldots + a_5 \phi^4] \left[ \frac{T_{u/T_{ref}}} b_1 + b_2 \phi + b_3 \phi^2 + b_4 \phi^3 \right] T_{ref} = 320 K \tag{7}
\]

**FIGURE 3:** Non-premixed flame diagram. The plotted data corresponds to all the operating conditions where the hydrogen micromix combustor was simulated: \( p \in (1,5,10,20) \) bar and \( \phi \in [0.5,2] \) (diffusion flame) for \( T_{air} = 600 K \) and \( T_{Hz} = 300 K \).

In order to characterize flame regime for hydrogen micromix flames, the Borghi diagram is not suitable because it assumes \( Le = Pr = Sc = 1 \). In hydrogen flames, as it is explained in the previous section, the non-dimensional numbers differ significantly from 1. Furthermore, the line describing \( Ka = 1 \) in the Borghi diagram depends on pressure and equivalence ratio because how it is defined (\( Ka = (\delta_l/\eta^2) \)). Therefore, for different pressures and mixtures there would be several lines of \( Ka = 1 \) in a conventional Borghi diagram. In
order to classify the flame regime of hydrogen-fueled diffusion flames the non-premixed flame diagram is used (figure 3). In this diagram the effects of pressure and equivalence ratio are included in the Da and Re numbers. The definition of Ka in the non-premixed flame diagram is that in eq. (4) obtained from [28] and derived without any assumption on the Le, Sc or Pr numbers. Hence, the line of Ka = 1 is identical for any pressure and mixture.

This diagram consists of 3 different flame regimes depending on Ka and only considers fast chemistry compared to mixing (Da > 1), differently from Borghi diagram. In the first region, where Ka < 1, turbulent flames can be considered as a differential set of laminar flames (pure flamelet) because turbulence dissipates before entering the flame front. Here, the flames can be corrugated (u'/SI > 1) or wrinkled (u'/SI < 1). However, this diagram is not suitable to plot u'/SI = 1 line to differentiate these two types of flamelets since on that line Da = \( l_t/\delta_t = f(p, \phi) \). Therefore, this line would change with \( \phi \) and p and it is impossible to ensure a point in the diagram where u'/SI has an specific value.

In the second region, where 1 < Ka < 100, the flamelet hypothesis still holds to solve the flames. It is a transition region from the pure-flamelet region and the broken flames region [29]. As Ka approaches the limit value of 100, the behaviour of the flames is more different from that of a flamelet and more error is produced by the flamelet hypothesis. This regime can also be called thin reaction flames.

The third region, where Ka > 100, turbulence can be much smaller than the flame front thickness. This allows eddies to progress inside the reaction zone and produce turbulent transport of radicals inside it, producing local quenching of the reactions and breaking the flame front. This regime is characterized by a strong turbulence-to-chemistry interaction and is called broken flames.

In a diffusion flame, it is expected to have a non-homogeneous mixture of fuel and air. That is why in figure 3 the flame non-dimensional numbers for different equivalence ratios are plotted to see the range of regimes that a diffusion flame could be defined by. Hydrogen-fueled mix micromix flames could be defined as transitional flamelets in the range of 1 < Ka < 100. Particularly, relatively-high-\( \phi \) points approach to Ka = 1 line, where the flamelet hypothesis does not deviate significantly from the actual flame regime. Since the flame found in micromix combustors is a diffusion flame, points with \( \phi \) closer to the lower flammability limit are rare, therefore most of the flame is at Ka \( \sim \) 1, i.e., it is a transitional flamelet.

\( Da \) of hydrogen-fueled flames in micromix combustion system is always bigger than 1 (figure 3). This means that chemistry is faster than mixing even though micromix system maximizes the mixing intensity through a cross-flow pattern. However, at points where \( \phi \) is lean \( Da \) is close to 1 despite the fast chemistry of hydrogen flames, proving that mixing is maximized in micromix injectors.

From these data it is possible to conclude that the CFD flamelet model could be suitable for defining the flame behaviour while offering low computational cost as an extra feature of this model. The FGM Kinetic rate model implemented in STAR-CCM+ software also considers turbulence-chemistry interaction through the probability density functions (PDFs). With this, the possibility of local flame quenching due to turbulent species transport inside the flame front is accounted for. This potentially reduces the error of using the flamelet model in the transition regime since these phenomena that describe the transitional flamelet are included in the code. Following this line, using thickened flame model to predict turbulence-chemistry interaction together with flamelet model to generate combustion data could also be suitable. However, since Thickened flame model can only be used in LES simulations it can not be used to perform simulations in a short time.

In conclusion, the CFD combustion model that potentially best reproduces the expected flame physics with the lowest computational cost is the FGM Kinetic rate model in RANS simulations.

### 3 Calibration of \( Sc_t \)

Turbulence Schmidt number is used in RANS simulations to model the turbulent diffusivity of the flow field. Mixing is modelled by means of the sum of molecular and turbulent diffusivities, i.e., the effective diffusivity. In this case, even though \( H_2 \) molecular diffusivity is very high, the turbulent diffusivity of the flow field is even higher, thus controlling the mixing. Since \( Sc \) is fixed by the fluid properties, \( Sc_t \) must be calibrated according to the cross-flow field properties.

Before starting the calibration, it is important to note the following:

- The hypothesis of constant \( Sc_t \) in cross-flow is not true since it changes with the turbulence scale and the downstream distance to the injector and the wall distance [30]. A sensitivity analysis on this should be done.
- For non-reacting RANS simulations it has been observed that for jet-in-crossflow, a constant value of \( Sc_t \) should be around 0.2 [31]. This is for relatively small momentum flux ratio. However, the effect it has in downstream combustion has not been assessed.
- The \( Sc_t \) that was calibrated in this section is not the actual \( Sc_t \) but a numerical value to model the overall mixing behaviour. This is because RANS models tend to underestimate the turbulent viscosity, therefore \( Sc_t \) must be lower than in reality to reproduce a realistic turbulent diffusivity [31].
- Mixing in LES simulations is not controlled by the value of \( Sc_t \) but by the turbulent scales that are solved.

In order to calibrate the value of \( Sc_t \), three RANS simulations were calculated with \( Sc_t \in (0.1, 0.2, 0.7) \). RANS solution was averaged with the iterations since it was unstable. Then, a LES simulation was calculated and its results averaged in time for 8 flow-throughs across the flame. The results of
RANS and LES were compared based on the flame shape, temperature and emissions to finally get a constant value of $Sc_f$. A further study, once experimental results are generated, could include the variation of $Sc_f$ along the flame.

### 3.1 Numerical setup

#### 3.1.1 Geometry

A single injector/flame case was generated to perform all the preliminary simulations as well as the calibration of $Sc_f$ and the evaluation of combustion models. This study was only performed in a single-injector combustor to capture the physics of a single flame.

![Figure 4: Single injector geometry.](image)

#### 3.1.2 Mesh

Three regions can be differentiated with different mesh size: the injector/flame region with base size of 0.125 mm, the intermediate pipe region with base size of 0.6 mm and the downstream pipe region with base size of 1 mm. The base size of the flame region was chosen through a mesh independence study. A base size of 1 mm provides enough resolution to capture the flow field in the pipe. Flame region dimensions were obtained from preliminary simulations. It was estimated a flame length of 20 mm for $p_0 = 20$ bar, $T_0 = 600$ K and $T_0 = 300$ K, therefore the finest mesh region was set to 24 mm long from the injector wall.

The mesh size of the intermediate region is an intermediate value chosen arbitrary to ensure a smooth development of the flow field. Polyhedral mesh was used for the whole fluid domain. There is only wall refinement in the injector region (10 prism layers with a prism layer stretching of 1.1 and a prism layer thickness of 0.125 mm).

#### 3.1.3 Physics models and boundary conditions

The physics models used in STAR-CCM+ version 13.06.014-R8 are:

- For RANS Steady: Steady, Three Dimensional, Turbulent, Reynolds-Averaged Navier-Stokes, K-Omega Turbulence, SST (Menter) K-Omega, Transition Boundary Distance, All y+ Wall Treatment. Most of these models are submodels of the K-Omega Turbulence model.
- For LES: Implicit Unsteady, Three Dimensional, Turbulent, Large Eddy Simulation, Exact Wall Distance, WALE Subgrid Scale and All y+ Wall Treatment.
- Combustion: Reacting, Flamelet, Flamelet Generated Manifold (FGM), FGM Reaction and FGM Kinetic Rate
- NOx emissions: NOx Emission, NOx Thermal and NOx Zeldovich.

The boundary conditions are shown in table 1.

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<th>Boundary Condition</th>
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<tr>
<td></td>
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<tr>
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<td>$m_{H_2}$ [kg/s]</td>
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</tr>
<tr>
<td></td>
<td>$T_0$ [K]</td>
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<td>Turbulence scale [m]</td>
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<tr>
<td>Outlet (Pressure outlet)</td>
<td>Target mass flow rate $\dot{m}<em>{air} + \dot{m}</em>{H_2}$ [kg/s]</td>
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<tr>
<td></td>
<td>Gauge pressure [bar]</td>
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<tr>
<td>Overall</td>
<td>Absolute $p_0$ [bar]</td>
<td>20</td>
</tr>
</tbody>
</table>

TABLE 1: Boundary conditions for the single injector geometry.

The fluid was modelled as an ideal gas. To generate the data for the flamelet model, 0D reactor was used and the chemistry mechanism used to define the reaction was that proposed by C. V. Naik and consists in 9 species and 25 reactions [32]. Kinetic rate was used to calculate reaction rate.

The unnormalized progress variable used to indicate the progress of the combustion is the mass fraction of water. Therefore, the (normalized) progress variable is 1 when the mass fraction of $H_2O$ is maximum, i.e., when the combustion is completed.

The time step for LES simulations was 2 µs to ensure a Courant number close to 1 (from preliminary simulations it was estimated that residence time in the flame region ranges from 1.6 µs and 5 µs). Implicit unsteady solver was set to 2nd order. Each time step consisted on 20 iterations to ensure that all the residuals fall around 2 orders of magnitude within any time step.

### 3.2 Results and discussion

Figure 5 shows the evolution of the mass fraction of OH with the downstream distance in mm. The data displayed in this
According to figure 5, RANS simulation with $S_C = 0.1$ predicts the shortest flame followed by LES, RANS with $S_C = 0.2$ and RANS with $S_C = 0.7$. This is because a lower $S_C$ means a higher turbulent diffusivity, i.e., species are transported much faster than momentum. Faster species transport means they mix in a shorter distance, thus displacing the reaction zone upstream. In addition, due to the high diffusion in the low $S_C$ simulation, some H$_2$ diffuses fast enough into the jet of air that it mixes out of its lower flammability limit. This mainly happens in the reacting region at the center of the jet of air (see figure 6). There, the mass fraction of OH is lower if $S_C = 0.1$ because some of the H$_2$ is not entering that reaction region. Due to this phenomenon the maximum mass fraction of OH when $S_C = 0.1$ is the lowest in figure 5. In contrast, the evolution $Y_{OH}$ when $S_C = 0.7$ is produced by poorer mixing and a longer reaction region compared to $S_C = 0.1$.

The evolution of the mass fraction of OH predicted by the LES simulation is in between the evolution predicted by $S_C = 0.1$ and $S_C = 0.2$. This is indicative that a turbulent Schmidt number in the range $S_C \in [0.1, 0.2]$ should be able to model the mixing characteristics of cross-flow in such a way that it approaches LES results even more than 0.1 or 0.2.

The above discussion is supported by the contours of $Y_{OH}$ in figure 6. The red line shows the predicted flame length by the LES solution, corresponding arbitrarily to $Y_{OH} = 0.25$ where the total temperature starts to fall significantly (this choice is performed since $Y_{OH}$ gradients cannot be easily extracted from CFD simulations). In these contours it is possible to see how $S_C = 0.1$ produces a slightly shorter flame and $S_C = 0.2$ a slightly longer flame than LES. In addition, both $S_C = 0.1$ and $S_C = 0.2$ simulations predict the downward flame propagation showed in LES contours while $S_C = 0.7$ simulation does not.

Interestingly, all the calculated flames are attached to the corner of the wall as can be deduced from figure 5 at $x \in [-2, 0]$ mm, where $Y_{OH}$ increases to its first peak. This implies that combustion in this zone is not due to how turbulence is solved but to the combustion model. The combustion in the region between the H$_2$ injector and the corner is produced because of the low local velocities induced by the cross-flow pattern. This creates a zone where the equivalence ratio is high and the velocity is low. In this case, depending on how the combustion model predicts flame propagation, this region may or may not be burning. Therefore, it is possible that FGM kinetic rate combustion model is overpredicting flame propagation and the wall does not actually burn.

Maximum temperature also depends on mixing (figure 6). In this case, $S_C = 0.1$ produces the closest-to-LES maximum total temperature. Maximum total temperature is produced in the wall where the flame is attached (figure 5 at $x \in [-2, 0]$ mm). It is expected that greater turbulent diffusion ($S_C = 0.1$) produces lower H$_2$ concentrations in the near-to-H$_2$-jet region. This implies that the turbulent diffusivity predicted with $S_C = 0.1$ is closer to that in LES simulations in this region. As explained at the beginning of this section $S_C$ is not actually constant.

Mass of NO$_x$ showed in figure 6 is the averaged mass of NO$_x$ produced in the numerical region that contains the flame. This parameter has no technological application but it allows to qualitatively describe the effect of residence time and flame temperature on NO$_x$ production in the flame. Since emissions trends were calculated in the final simulations, it was necessary to include NO$_x$ production in the calibration process.

The flame predicted through RANS simulation with $S_C = 0.2$ produces slightly lower NO$_x$ than LES simulations while RANS simulations with $S_C = 0.1$ and with $S_C = 0.7$ produce respectively much lower and much higher NO$_x$. NO$_x$ production depends mainly on residence time (flame length) and flame temperatures. In the steady RANS simulations with different $S_C$ the temperature along the flame is very similar, so the controlling parameter for NO$_x$ production is the residence time. This explains why $S_C = 0.1$ simulation shows the lowest mass of NO$_x$ and $S_C = 0.7$ the highest. However, this simple analysis can not be applied to LES time-averaged results (figure 6) since the instantaneous flame temperatures are higher than the average, thus producing higher NO$_x$ (figure 7).
4 Combustion models comparison

In this section, three different simulations were compared. Two of them have the same combustion model (FGM kinetic rate) but different turbulence model (LES and k-ω RANS) while the third one was simulated with Thickened flame model that can only be LES. The aim of this section was to analyse and compare the flame structure and NOx emissions that the most suitable combustion models (according to the predicted flame regime) can predict. This will be helpful when the ENABLEH2 team at Cranfield University carries out experiments in a hydrogen micromix combustor, to be aware of different modelling possibilities of STAR-CCM+ and, if possible, calibrate them to be validated.

4.1 Numerical setup

The geometry, mesh and physics models are mostly the same as in the previous section. There are two main differences in the CFD set up. First, the combustion model was changed from FGM kinetic rate to Thickened flame model. Second, since the flame propagation rate in the Thickened flame model is given by the laminar flame speed, the correlation in [5] was extended with the experimental data at 10 bar and \( \phi = 0.3 \) in [33] and expressed as a function of the mixture fraction (7). Due to the low variation with pressure of laminar flame speed at \( \phi = 0.3 \), it was assumed to be similar at 10 bar and 20 bar.

\[
S_l = a_0 + a_1Z + a_2Z^2 + a_3Z^3 + a_4Z^4 \quad (8)
\]

In LES, the higher instantaneous temperatures together with the thicker flame make that NOx prediction is even higher than that of steady RANS with \( S_c \) = 0.2 where averaged temperatures are higher and the flame longer (compared to LES averaged). Therefore, it is possible to conclude that to approach NOx prediction of RANS simulations to that of LES the turbulent Schmidt number should be slightly higher than 0.2.

From this analysis, it is possible to conclude that \( S_c \) = 0.2 is a good compromise between flame length and emissions production. However, it is important not to forget that in actual flames the turbulent Schmidt number is higher and not constant. In the future, a model describing the variation of \( S_c \) along the flame would be useful to match simulation to experimental data.

4.2 Results and discussion

Contours of \( Y_{OH} \) and \( T_0 \) (figure 8) show that all the combustion models predict flame attachment to the corner of the wall. However, Thickened flame model shows lower temperatures in the wall region than FGM Kinetic rate. This is because each model has a different propagation mechanism. In the case of Thickened flame, the reaction propagates according to the laminar flame speed (based on experimental data) and the turbulence-chemistry interaction while in FGM Kinetic rate it is computed from the reaction kinetics and the turbulence-chemistry interaction. Since flame attachment to the wall is predicted by these two models, it means it does not depend on the combustion model and should be further studied in the future. This analysis is important because the high local temperature near the wall may lead to a decrease in the life of the combustor injector plate. For future work to increase the life of hydrogen micromix combustor, a thermal analysis must be performed, the geometry could be redesigned or the inclusion of a heat shield protecting the wall studied.

The flame structure calculated with the thickened flame model is entirely in the region where the mixed jet is interacting with the vortexes produced between the jet and the walls. Here, the velocity of the fluid is the lowest and the residence time is increased, thus allowing the reaction to start. Higher residence...
time in the flame front increases OH and NO\textsubscript{x} production. This explains that Thickened flame model predicts higher $Y_{OH}$ locally and that, despite the lower total temperatures in the flame, the NO\textsubscript{x} production is similar compared to FGM Kinetic rate results.

In contrast, the flame predicted by FGM Kinetic rate is in the middle of the jet, where the velocity of the fluid is maximum. This implies that there is a noticeable difference in the flame structure when different flame propagation models are used. In the case of FGM Kinetic rate, flame propagation is fast enough to anchor the flame in a high-speed region while in Thickened flame it is slower and can only anchor the flame to the low-speed zone. Consequently, and due to the lack of experimental data, it is uncertain where the flame actually starts.

The maximum total temperature and the mass of NO\textsubscript{x} in the flame region are lower with Thickened flame model than with FGM Kinetic rate in RANS and LES. The maximum total temperature is lower because the flame propagation is slower in the thickened flame model. This causes that the flame is not attached to the wall between the corner and the H\textsubscript{2} injector, where the maximum temperatures are produced, and thickens the flame. A thinner flame means that the thermal diffusivity is high enough to spread the reaction region and evacuate some energy out of it. Since the reaction is produced progressively in a wider space the temperature in the flame front is lower. Finally, Thickened flame model virtually thickens the flame front to solve the turbulence-chemistry interaction inside it. According to the flame regime of hydrogen micromix flames (figure 3) there may be species transport inside the flame front, thus creating local zones poor in H\textsubscript{2} or H\textsuperscript{+} radicals where the reaction is quenched. Therefore, the more detailed resolution of the flame front inner structure could also be responsible for the lower temperatures in the thickened flame model.

Regarding NO\textsubscript{x} production, it is important to note again that in LES it is a time-averaged value that depends on the instantaneous local temperatures along the flame. Since flame propagation in Thickened flame model is slower, overall residence time is increased. However, the fact of thickened flame model producing lower NO\textsubscript{x} means that instantaneous local temperatures are much lower than in RANS and LES with FGM Kinetic rate, where the residence time is lower. These two flame characteristics compensate each other resulting in a very similar NO\textsubscript{x} production.

Finally, among these two combustion models, FGM kinetic rate seems a more promising option to run hydrogen micromix CFD simulations because it can use RANS solvers, thus minimizing the computational cost, and because its flame propagation rate is not based on laminar flame speed, whose behavior at high pressure is not experimentally documented at $p > 10$ bar.

**CONCLUSION**

Along this paper a methodology to estimate the physical properties of H\textsubscript{2}, air and H\textsubscript{2}-air mixtures was given for a range of temperature, pressure and equivalence ratio. For the leanest equivalence ratio (0.3), where minimum NO\textsubscript{x} was expected, the Schmidt, Lewis and Prandtl numbers were estimated as 0.22, 0.35 and 0.75 respectively.

It was concluded that the conventional Borghi diagram is not suitable to characterize the flame regime of hydrogen micromix diffusion flames because it assumed $Sc = Le = Pr = 1$ and is only for premixed flames. The non-premixed flame diagram was defined and used to determine that hydrogen micromix flames are transition flamelets. From this analysis, FGM kinetic rate and Thickened flame combustion models were chosen as appropriate to reproduce the physics of the transitional flamelet flame regime.

Turbulent Schmidt number for cross-flow combustion was set to 0.2 as a compromise between accuracy on NO\textsubscript{x} and flame length prediction. RANS and LES simulations of a single micromix injector were compared to calibrate this parameter. If $Sc_t$ is higher than 0.2, NO\textsubscript{x} production in RANS would be more similar to that in LES. In contrast, if it is lower than 0.2, the flame length in RANS would be more similar to that in LES. If this parameter is calibrated for a single flame, it should also be calibrated for multiple flames because flame-interaction between different injectors depends on mixing, which is equally dependent on $Sc_t$ for both simulations. A sensitivity analysis could be done for multiple-injector configuration.

Finally, FGM kinetic rate and Thickened flame combustion models were compared in a single hydrogen micromix flame. It was concluded that flame structure and temperature distribution along the flame change significantly depending on the combustion and turbulence models. Flame attachment to the wall only depends on the flame propagation predicted by each combustion model. This phenomenon and its implication should be further studied by means of a heat transfer analysis. The addition of a heat shield to protect the wall against high temperature should be considered. Thickened flame model predicts lower maximum total temperatures and higher residence time than FGM Kinetic rate because the flame is developed in a low-speed region. These two phenomena compensate each other resulting in a similar production of NO\textsubscript{x} in the flame region.

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