Modelling of Laminar Diffusion Flames in Vitiated Environments

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ABSTRACT
A laminar flamelet model has been used to calculate chemistry of methane-air flamelets in vitiated and non-vitiated environments. Calculated results for temperature and stable species concentrations agree well with those obtained using other similar models. The present results also agree quite well with recent laboratory measurements on both laminar and turbulent diffusion flames. The largest discrepancies between measured and calculated results are for CO-concentrations.

KEYWORDS: Flamelet model, Combustion models, Diffusion flames, Chemical kinetics

INTRODUCTION
About two thirds of deaths in fires are caused by (CO) poisoning. The mechanism of formation of CO in fire is quite complicated and largely depends on fuel type, oxygen supply in surroundings, and how the fire develops, i.e. turbulence intensity.

To understand the formation of CO and other species in fires, one has to study the micro-structure of flames. Ignition, combustion and extinction occur in a turbulent flame at the same instant in time. These phenomena can have spatial separations of only a few millimeters. Locally the mixture can be diluted by complete or incomplete products of combustion, whose temperatures can vary considerably. This influences the chemistry, and hence the combustion efficiency at that location. To predict combustion in diffusion flames, finite rate chemical kinetics must be modelled to take account of combustion efficiency, which varies considerably inside the same flame. This work [1] is concerned with diffusion flames under the influence of vitiation, i.e. fire gas recirculation, which always occurs in room fires. The modified version of Cranfield SNECKS-code (Solver for Non-Equilibrium Chemical-Kinetic Systems) is used to calculate the chemistry [2]. The model can be used with any flow field calculation model.
LAMINAR FLAMELET MODEL

A laminar diffusion flame exhibits unique relationships for chemical species, temperature, enthalpy, viscosity, soot concentration, etc., in terms of mixture fraction (a conserved scalar). By assuming that a turbulent diffusion flame consists of microscopic elements that have the structure of an undisturbed laminar diffusion flame, these relationships can be averaged by using of an appropriate shape of the probability density function (PDF). It is assumed that the chemistry is fast and the chemical reactions occur mainly in these thin flamelets. These assumptions allow us to decouple the statistical uncertainties of a turbulent flow field from a complex multicomponent chemistry and make it possible to calculate the chemical kinetics and flow field separately. The calculated flamelets with different degrees of vitiation, strain rate (i.e. different turbulent intensity) etc., can be stored in a data library, from where they can be easily accessed during the flow field calculations.

A special numerical model of co-flowing laminar diffusion flame was used in the present study. This model was calculated using SNECKS [2]. The flame model is a modified version of the model used in Pratt's CREK code, which is described in ref [3]. SNECKS differs from CREK in that the stream-wise distance, $x$, in CREK is replaced by time in SNECKS. SNECKS is therefore a 1D unsteady code while CREK is a 2D steady code. Both codes use the non-dimensionalized stream function of Spalding as the cross-stream independent variable.

In such model flames, the thermally hypergolic ignition and combustion of co-flowing streams of methane and air is considered. The schematic of the flame is shown in fig. 1. Initial temperatures of 300 K in fuel oxidizer streams were used in non-vitiated cases. In vitiated cases, the air stream is diluted by combustion products (for simplicity assumed to consist of only CO$_2$ and H$_2$O) at either 300 K (cold vitiation) or 1000 K (hot vitiation), so that the oxidizer stream temperature is dependent on the degree of vitiation. An initial temperature of 2000 K in the burning zone, the initially very thin sheet of stoichiometric proportions of fuel and oxidizer between the two co-flowing streams, was used to initiate combustion. The computationally efficient "expanding grid" was utilized, so that computational field occupies the entire shear layer as it grows wider downstream.

FIGURE 1. Axisymmetric shear flow mixing and combustion of initially unmixed fuel and air.
The governing set of differential equations for this special kind of diffusion flame consists of one-dimensional unsteady, parabolic differential equations of the form:

\[ \frac{\partial \phi}{\partial t} = \frac{\partial}{\partial \psi} \left( \rho \Gamma_\phi \frac{\partial \phi}{\partial \psi} \right) + \frac{S_\phi}{\rho} \]  

where \( \phi \) is a symbol for any atomic element or enthalpy, \( \Gamma_\phi \) is a transport coefficient, \( S_\phi \) is the source term, and \( \psi \) is a stream function.

The present chemical model for combustion of methane consists of 13 elementary reaction steps involving 11 species. The reaction scheme is shown in table 1.

Table 1. A 13-step reaction scheme for combustion of methane involving 11 species used in the present study. The reactions 1-12 are reversible (sign '='), and reaction 13, the fuel destruction reaction, is irreversible (sign '-'). The parameters \( A_r \) and \( A_b \) denote Arrhenius pre-exponential factors (cm^3) for forward and backward reactions, \( \beta_r \) and \( \beta_b \), temperature exponents and \( E_r \) and \( E_b \), activation energies (cal).

<table>
<thead>
<tr>
<th>Forward rate</th>
<th>Backward rate</th>
</tr>
</thead>
<tbody>
<tr>
<td>( A_r )</td>
<td>( A_b )</td>
</tr>
<tr>
<td>( \beta_r )</td>
<td>( \beta_b )</td>
</tr>
<tr>
<td>( E_r )</td>
<td>( E_b )</td>
</tr>
</tbody>
</table>

1. O+H_2O=OH+OH  \( 1.76 \times 10^7 \)  -0.02  106747.3  1.55 \times 10^8 \) 0.0  0.0
2. H+H=H_2  \( 1.00 \times 10^{12} \)  -1.0  0.0  7.15 \times 10^6 \)  -0.82  103328.3
3. O+O=O_2  \( 6.00 \times 10^6 \)  0.0  0.0  5.83 \times 10^{10} \)  -1.0  1198367
4. H+OH=H_2O  \( 2.00 \times 10^7 \)  -2.0  0.0  2.20 \times 10^4 \)  0.0  1052179
5. H+O_2=OH+O  \( 2.00 \times 10^{11} \)  0.0  16665.8  6.5 \times 10^6 \)  0.27  0.0
6. O+H_2=OH+H  \( 6.00 \times 10^8 \)  0.0  10008.6  1.94 \times 10^{10} \)  -0.03  8059.4
7. O_2+H=H_2O  \( 1.50 \times 10^9 \)  0.0  994.5  2.10 \times 10^{12} \)  0.0  46039.3
8. H+HO_2=OH+OH  \( 1.59 \times 10^{11} \)  0.0  1080.0  1.26 \times 10^{10} \)  0.0  59873.4
9. CO+OH=CO_2+H  \( 1.50 \times 10^8 \)  0.0  100.5  1.50 \times 10^{10} \)  0.0  23576.9
10. H_2+OH=H_2O+H  \( 1.50 \times 10^8 \)  0.0  5004.3  5.55 \times 10^{10} \)  -0.01  19802.4
11. CO+O=CO_2  \( 6.00 \times 10^7 \)  0.0  0.0  1.31 \times 10^{10} \)  -0.58  125843.7
12. H+OH=HO_2  \( 3.00 \times 10^8 \)  0.0  0.0  6.96 \times 10^{10} \)  0.21  101379.1
13. 2CH_4+O_2 -2CO+4H_2  \( 1.75 \times 10^7 \)  1.0  24265.

The species transport properties are calculated from Chapman-Enskog expressions by using the Lennard-Jones parameters [2]. The Chapman-Enskog method is based on the three basic assumptions: a) molecular collisions are binary collisions, b) translational energy is treated using classical mechanics and c) spatial gradients of the microscopic or continuum properties of the gas are assumed to be small. These conditions are best satisfied by monoatomic molecules. The flames easily satisfy this requirement except at very high pressures. The method proceeds from a series expansion of the velocity distribution function about the Maxwellian distribution in order to obtain explicit expressions for the transport vector in terms of the gradients of the dependent variables of fluid dynamics from the Boltzmann equation [4]. The result is a set of Navier-Stokes equations, which is applicable for large deviations from the equilibrium.

For calculating heat capacities and enthalpies of species, the polynomial fit for temperature
is used. The polynomial coefficients are stored in a data base. There are 14 coefficients for each species stored in the data base: seven coefficients for the temperature range 300 K to 1000 K and seven for the temperature range 1000 K to 5000 K.

RESULTS AND DISCUSSIONS

In the present work, flamelet chemistry was calculated for methane-air laminar diffusion flames in both non-vitiated and vitiated atmospheres. The results have been compared with both laminar and turbulent diffusion flames in non-vitiated atmospheres. At the time this study was made, nothing could be found in the literature concerning flamelet problems in vitiated atmospheres.

Three degrees of vitiation were used: the oxidizer stream was diluted by 20%, 40% and 60% of combustion products. When using hot combustion products, the temperature of the mixture of combustion products + air was calculated as the arithmetic mean value of hot gas at 1000 K and air at 300 K (equal heat capacities for hot gas and air were assumed), which gives oxidizer stream temperatures 450 K, 570 K and 680 K for 20%, 40% and 60% vitiation, respectively.

Due to vitiation, the stoichiometric mixture fraction is shifted towards the lower values. By introducing a vitiation factor $\psi_{vit}$, the $O_2$ mass fraction $Y_{O_2,2}$ in the oxidizer stream can be expressed as a function of $\psi_{vit}$:

$$Y_{O_2,2} = \frac{\psi_{vit} M_{O_2}}{\psi_{vit} M_{O_2} + M_{N_2} / M_{N_2}} \left(1 - \psi_{vit}\right); \quad 0 \leq \psi_{vit} \leq 1;$$

(2)

where $M$ and $\nu$ are molecular weight and stoichiometric coefficient and indices $O_2$ and $N_2$ denote oxygen and nitrogen, respectively. The stoichiometric mixture fraction for a vitiated oxidizer stream can be calculated from a simple formula

$$\xi_{st, vit} = \left(1 + \frac{4}{Y_{O_2,2}(\psi_{vit})}\right)^{-1}$$

(3)

The stoichiometric mixture fraction $\xi_{vit}$ and the oxygen mass fraction $Y_{O_2,2}$ in the oxidizer stream for $\psi_{vit} = 0.00, 0.20, 0.40$ and 0.60 are shown in Table 2.

TABLE 2. Variation of stoichiometric mixture fraction and oxygen mass fraction with different degrees of vitiation.

<table>
<thead>
<tr>
<th>Vitiation factor</th>
<th>0.00</th>
<th>0.20</th>
<th>0.40</th>
<th>0.60</th>
</tr>
</thead>
<tbody>
<tr>
<td>Oxygen mass fraction</td>
<td>0.233</td>
<td>0.186</td>
<td>0.140</td>
<td>0.093</td>
</tr>
<tr>
<td>Stoichiometric mixture fraction</td>
<td>0.055</td>
<td>0.045</td>
<td>0.034</td>
<td>0.023</td>
</tr>
</tbody>
</table>
Comparisons with other studies

Comparison between theoretical models and Raman measurements

The temperatures and major species concentrations calculated in the present study agree well with results from earlier calculations on laminar opposed jet methane-air flames by Peters and Kee [5]. Peters and Kee used the five scalar, four step, reduced mechanism, which is a systematically condensed form of an 18 elementary step reaction mechanism. The rate coefficients for resulting global reactions are combinations of species concentrations and unmodified elementary reaction rate coefficients. Thus, by using the reduced scheme, the concentrations of the intermediate species such as OH, O, etc. cannot be seen (but their effects are accounted for in calculations) as they can in the "full" scheme.

Figure 2 shows a comparison of the present calculations with those of Peters and Kee for strain rates of 450 s⁻¹, 300 s⁻¹ and 100 s⁻¹. Figure 2 also shows the scatter plot data from laser Raman experiments by Dibble et al. [6,7] on turbulent high-momentum methane jet flames, with a 41 m/s central jet velocity, at the point x/D = 20 and r/D = 1.14 and 1.55, where x and r are the distances from burner top and the axis of symmetry, respectively, and D is the burner exit diameter.

FIGURE 2. Joint pdf scatter plot for a) temperature, b) CO₂, and c) CO versus mixture fraction. Lines denote calculations and dots denote Raman experiments [6].
Near stoichiometry, $\xi = \xi_{st}$, the temperatures predicted in the present study are quite near to those of Peters and Kee. For $\xi > 0.4$ the temperature is about 100 to 150 K lower in the present study.

The concentration of CO$_2$ at $\xi = \xi_{st}$ predicted by the present work is about the same as Peters and Kee's calculation for a strain rate of 100 s$^{-1}$. For higher values of $\xi$, the CO$_2$ concentration in the present study moves towards the 300 s$^{-1}$ curve. This indicates that combustion is more incomplete for rich mixtures in the present model than in Peters and Kee's model. The concentration of H$_2$O follows the same pattern as that of CO$_2$ and comparisons with the present results are therefore omitted.

In the present study, somewhat higher concentrations of CO were found compared with Peters and Kee's calculations. There is a large discrepancy in CO concentrations between the calculated results and those measured with Raman methods. In the blue regions of hydrocarbon flames the laser signals are contaminated by fluorescence, the intensity of which is of the same order as the Raman signals. As the source molecules for this fluorescence are not fully known, this can lead to a large uncertainty in measured species concentrations. Discrepancies in CO concentrations between measured and calculated results can be attributed to this fluorescence.

**Comparison with measurements in laminar and turbulent diffusion flames in non-vitiated atmospheres**

Smith and Cox [8] made detailed measurements of time-averaged concentrations of major species produced in different sizes (18 to 111 kW) of turbulent diffusion flames of natural gas (94% methane) on a 0.3x0.3 m burner. All species measurements were time-averaged over a period of at least 10 min. The curves show generally the same behaviour as the laminar diffusion flames measured by Mitchell et al. [9] and Tsuji and Yamaoka [10], except that the peak values of temperature and of CO and CO$_2$ concentrations are shifted towards the higher mixture fraction values. As expected, the time averaging of the species fluctuations in the turbulent flames reduces peak values of concentrations.

Mitchell et al. [9] used a cylindrical diffusion flame burner consisting of two concentric tubes: an inner tube of radius 0.633 cm (fuel) and an outer tube of radius 2.54 cm (air). Tsuji and Yamaoka [10] used a rectangular combustion chamber with a cross section of 3 x 12 cm and an uncooled porous (sintered bronze) cylinder, length 3 cm and diameter 6 cm. Mitchell et al made radiation and conduction corrections to the thermocouple bead temperature to determine the unperturbed gas temperature at the bead location. The fact that Tsuji and Yamaoka did not make such a correction explains their lower peak values of measured temperatures, as can be seen in figure 3.

At $\xi = \xi_{st}$ the results from laminar flames obtained by Mitchell et al. yielded the same results for temperature and CO concentrations as computed using the present model. For $\xi > \xi_{st}$, calculated values of temperature are about 200-300°C higher and CO concentration about 25-40% higher. The temperatures measured by Tsuji and Yamaoka agree well with calculated results for $\xi > \xi_{st}$. At stoichiometry, Tsuji and Yamaoka measured considerably lower values of temperature and CO and CO$_2$ concentrations than Mitchell et al. For $\xi > 0.2$ the temperatures measured by Tsuji and Yamaoka are higher than those calculated. The radiation

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FIGURE 3. Comparison of current calculations with measured centreline mean temperature (a) and mean mass fractions of CO$_2$ and CO (b and c) as a function of mean mixture fraction. Mitchell et al. and Tsuji and Yamaoka; laminar flames, Smith and Cox; turbulent flames.

energy losses are omitted in the present calculation model. These losses would shift the temperature to somewhat lower values and would increase the concentration of CO slightly.

The predicted mass fraction of CO$_2$ at stoichiometry ($\xi = 0.055$) is about 0.10, and the peak value is 0.11 at $\xi = 0.06$. This is very close to Tsuji and Yamaoka's results on laminar diffusion flames. Mitchell et al., however, measured very high peak values of CO$_2$ concentrations with a mass fraction of nearly 0.160 [9]. This value is even higher than the theoretical value (which should be about 0.150), if it is assumed that all carbon after combustion is in the form of CO$_2$.

Most of the difference between values obtained from theoretical models (the present model and Peters and Kee's) and measured values can be attributed to carbon balance in combustion products. At stoichiometry, the calculated CO concentrations are about 3-4%, having peak values at $\xi = 0.10$ between 5 and 6%, which are much higher than those measured. This large discrepancy may depend on the theoretical model. Maybe a more detailed chemical scheme is needed to calculate CO chemistry. On the other hand, it is difficult to measure CO concentration, due to its relatively low value and difficulties in determining the exact position of the stoichiometric mixture in the flame sheet.
Vitiated flamelets

Fuel and stable species (CO₂, CO, H₂O, O₂) concentrations and temperatures are shown in figure 4. As expected, temperature decreases with increasing $\psi_{vt}$. The peak temperature decreases approximately 100 K for every 10% increase of $\psi_{vt}$ in the case of hot gas vitiation, i.e. from about 2000 K for $\psi_{vt} = 0$, to less than 1400 K for $\psi_{vt} = 0.6$ (see fig. 4 e-f). When cold vitiating gas is used, the temperature decrease is slightly larger for a given increase in vitiation (fig 4 a-d).

In the case of hot vitiation, the peak value of CO mass fraction, $Y_{CO, top}$, decreases from 0.056 to 0.035 with an increase in $\psi_{vt}$ from 0 to 0.6. In the case of cold vitiation, $Y_{CO, top}$ is almost constant. It should be noted that vitiation reduces the relative concentration of O₂ in the oxidizer stream and hence a greater mass of oxidizer gas is needed for combustion of a certain portion of fuel. Since CO concentrations in fig. 4 are presented as a fraction of total mass in the mixture, CO production per unit of combusted fuel increases with increased vitiation. The predicted peak concentrations of CO per combusted amount of fuel increase by about 53% for hot and by about 58% for cold vitiation when $\psi_{vt}$ is increased from 0 to 0.6. The peak values of CO mass fractions and CO/CH₄ for different vitiations are shown in table 3.

<table>
<thead>
<tr>
<th>$\psi_{vt}$</th>
<th>Hot vitiation</th>
<th>Cold vitiation</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$Y_{CO, top}$</td>
<td>$\xi_{top}$</td>
</tr>
<tr>
<td>0.0</td>
<td>0.056</td>
<td>0.100</td>
</tr>
<tr>
<td>0.2</td>
<td>0.050</td>
<td>0.086</td>
</tr>
<tr>
<td>0.4</td>
<td>0.044</td>
<td>0.054</td>
</tr>
<tr>
<td>0.6</td>
<td>0.035</td>
<td>0.041</td>
</tr>
</tbody>
</table>

The CO₂ concentration increases slightly with increased vitiation with hot gas, while remaining nearly constant for rich mixtures in the case of cold vitiation. The H₂O-concentration seems to be unaffected by whether the vitiation gas is cold or hot for the rich side of stoichiometry.

The mass fractions of O₂ decrease from initial values in oxidizer streams at $\xi = 0$ to zero at $\xi_{st, vt}$. The decrease is a linear function of $\xi$ for $\xi > \xi_{st, vt}$ at which the concentration of O₂ is zero indicating that O₂ is consumed totally.

The concentrations of H₂ and the radicals O, H, OH and HO₂ are presented in figure 5. The maximum H₂ mass fractions occur at about the range $\xi = 0.1$ to 0.2. The temperature has a significant effect on H₂ concentration. For hot gas vitiation at mixture fraction $\xi = 0.1$ to 0.2 the mass fraction of H₂ is 0.0035 for $\psi_{vt} = 0.2$. For $\psi_{vt} = 0.6$, the H₂ mass fraction is about 0.0027. For cold vitiation, the $Y_{H₂}$ is about 0.0040 for all $\psi_{vt}$. The peak value of H₂ is also shifted slightly towards the richer mixture fraction values when the vitiating gas is cold.

O radicals exist only within a narrow band at the lean side of stoichiometry as a result of
FIGURE 4. Temperature, CO, CO$_2$, H$_2$O, $O_2$ and fuel concentrations as a function of mixture fraction.
FIGURE 5. H, OH, O, HO₂ and H₂ concentrations as a function of mixture fraction.
dissociation of O$_2$ due to heat from the flame. The peak values of O concentrations occur at $\xi = 0.95 \xi_{st}$. The mass fraction of O at that point is about 0.0015 for $\Psi_{vit} = 0$ and 0.0004 for cold vitiation with $\Psi_{vit} = 0.6$. At $\xi = \xi_{st}$ ($\xi = \xi_{st,vit}$ in vitiated cases) the O concentration has almost vanished. For hot vitiation, O concentrations are somewhat lower than for cold vitiation.

As expected, the maximum concentrations of H radicals are located on the rich side of the stoichiometry, not far from the stoichiometric mixture in the flame sheet, where the temperature is high enough making methane molecules to lose their first H-atoms.

OH radical concentrations in the flamelets follow the same pattern as O radical concentrations, except that the peak occurs at $\xi = \xi_{st}$. The peak concentration levels are about 2 to 3 times those of O-radical levels. OH concentrations vanish at about $\xi = 0.2$.

HO$_2$ radicals exist only on the lean side of stoichiometry. The peak concentrations $\approx 1.5 \times 10^5$ are located at the oxidizer boundary of flamelet $\xi = 0$. At the mixture fraction $\xi = \xi_{wt}$ the HO$_2$ concentration is zero.

CONCLUSIONS

The stable species concentrations and the temperatures calculated using the present thirteen-step reaction model agree well with the earlier calculations of Peters and Kee using a four-step, reduced-reaction mechanism [5]. Near stoichiometry, agreement with recent measurements on laminar diffusion flames [9] is also quite good. The largest discrepancy was found for CO concentrations. For $\xi \geq 0.2$, the calculated CO concentrations are overestimated by 30%.

The agreement of the present model with turbulent diffusion flames is also quite good [8], except that, due to time averaging, measured peak values of temperature and CO and CO$_2$ concentration are considerably lower and occur further to the rich side of stoichiometry than values predicted by the model.

Vitiation affects the temperature and hence the chemistry. In the case of hot vitiation, the temperature is reduced approximately 100 K for every 10% increase in vitiation. In the case of cold vitiation, the temperature effect is larger. For hot vitiation, CO production is increased by 53% per combusted mole of methane when the vitiation factor is increased from 0 to 0.6. For cold vitiation, the increase in CO is about 58%.

REFERENCES


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