

## Numerical Simulation of a Lifted Methane Jet Flame in a Vitiated Coflow: Lagrangian Approach with Detail Chemistry

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**Abstract:** Numerical calculation of turbulent lifted flame is studied. In this study, mass flow rate, temperature and exact chemical composition of hot products mixed with air sent toward the turbulent flame base are fully determined. The effects of both non-infinitely fast chemistry and partially premixed combustion are taken into account by using the MIL model (Modèle Intermittent Lagrangien) originally introduced by Borghi and Gonzalez with complex chemistry. Here, the model based on a presumed joint PDF (Probability Density Function) shape for both a mixture fraction and a mixing time for production of a reaction. A fully detailed chemistry is constructed about the tabulation of ignition delays and introduced in calculation of the mean chemical rate; this allows us to express the higher probability that fluid particles have to burn for large range around stoichiometric composition. Numerical simulation of the turbulent diluted jet flame of methane studied by R. Cabra and his co-workers at Berclays University is satisfactory; the lift of height prediction are successfully compared with experiments and was around 30-40 D.

**Key words:** Numerical simulation, fine rate chemistry, lifted flame, detailed chemistry, partially premixed turbulent combustion, Cameroon

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### INTRODUCTION

In many practical combustion systems the fuel jets emerge into a hot environment of oxidants and combustion products. These hot products carry thermal energy that may help flame stabilization; they are also useful to dilute the reaction zones which homogenizes the mixture and avoids the high temperature levels responsible for NO<sub>x</sub> emissions. Prediction of turbulent jet flames with complex recirculating flows and autoignition mechanisms pose a great challenge to the current combustion modeling. The vitiated coflow flame is a kind of turbulent reacting flow within a hot environment and low oxygen concentration.

Some studies related to the vitiated coflow from the experimental point of view, research of Cabra *et al.* (2002, 2005) with like fuel a mixture of H<sub>2</sub>/N<sub>2</sub> then CH<sub>4</sub>/air, respectively constitute the reference now-a-days; moreover, Gordon *et al.* (2008) brought temperature measurements of OH and of CH<sub>2</sub>O in methane combustion. Calculations were also carried out on this type of configuration by the model of PDF calculated, Gordon *et al.* (2007, 2008) simulated the combustion of two fuels mentioned earlier and brought information on transport by convection-diffusion of the molecules;

Cao *et al.* (2005) applied its calculation to the only case of H<sub>2</sub>/N<sub>2</sub>, similar to Masri *et al.* (2003) detailed chemistry was taken into account by use of ISAT method (*in situ* Adaptive Tabulation). A calculation with CMC model (Conditional Moment Closure) was carried out by Patwardhan *et al.* (2009) where an analysis of sensitivity of the zone of pre-flame was questioned; another of Michel *et al.* (2009) by use of the flamelet model ADF-PCM where an accent was laid on the study of the diffusion zone thanks to the correction made on the probability distribution of the scalar dissipation rate at stoichiometry  $P(\chi_{st})$  like a log-normal function. Domingo *et al.* (2008) carried out a calculation using LES (Large Eddy Simulation) adding an analysis on the turbulent flame and a new closing of the scalar dissipation rate of the reactive species, then Wang *et al.* (2007) a calculation using the DNS (Direct Numerical Simulation) where an examination of the dynamics of the swirls, structure of the micro-scales and the distribution of the turbulence intensity in the flame are made.

The Lagrangian models in fact the MIL (Modèle Intermittent Lagrangien) model was introduced into the modeling of turbulent combustion for the comprehension of the turbulence-chemistry interaction met in non-premixed flames (Borghi and Gonzalez, 1986; Gonzalez

and Borghi, 1991). A well-known projection was observed when Obounou *et al.* (1994) there introduced detailed chemistry, thus predicting zones of partial extinction then of re-ignition. Gonzalez *et al.* (1997) applied the MIL model within the framework of stabilization to the exit of a burner. This model consists of a formulation of presumed PDF, associates ignition times and makes it possible to follow the history of a particle.

The objective of this study is to apply the MIL model, version developed at the Department of Physics of the University of Yaounde 1 with taking into account of a complex chemistry, then to couple it in a computer code for the simulation of a lifted flame. The study relates to measurements of Cabra *et al.* (2005) which provides several interesting data with initial and boundary conditions quite exact.

**MATERIALS AND METHODS**

**Experimental configuration:** The exact chemical composition and mass flow rate of recirculating burnt gases are often difficult to calibrate and measure accurately in real combustion chambers. To gain greater understanding of flame stabilization in environments where burnt gases dominate, Cabra *et al.* (2005) designed a laboratory vitiated burner shown in Fig. 1, in which the injection conditions of reactants and hot products are fully determined. It consists of an CH<sub>4</sub>/air fuel jet issuing into a hot coflow which is the combustion product of a lean premixed H<sub>2</sub>/air flame. The diameter of the central jet is  $d = 4.57$  mm; the coflow diameter about 210 mm is much larger than the central jet which can isolate the central jet from ambient air for a sufficiently long distance.

The corresponding inlet conditions in fuel stream and coflow are shown in Table 1 in terms of mole fractions, temperature, velocity and Reynolds numbers. The Raman-Rayleigh technique was used for measuring instantaneous mass fractions and temperature; all the mean and instantaneous values have been downloaded from Cabra *et al.* (2002).

**Description of the MIL model:** The combustion being the seat of two phenomena which are chemistry and turbulence, the interaction of the latter requires an adequate modeling. The MIL model takes part by its formulation with the comprehension of this type of combustion integrating this competition at the base, this model was introduced for the modeling of the average reaction rate for non-premixed flames (Borghi and Gonzalez, 1986; Gonzalez and Borghi, 1991; Obounou *et al.*, 1994; Gonzalez *et al.*, 1997).

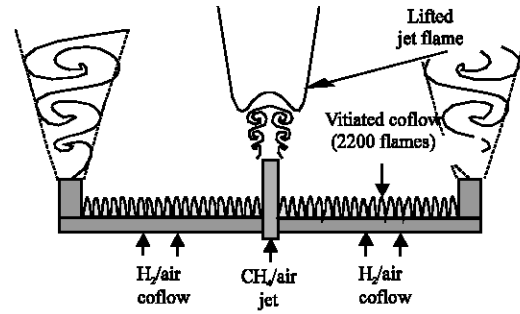


Fig. 1: Experimental arrangement of the Cabra *et al.* (2005) experiment

Table 1: Boundary conditions of the Cabra *et al.* (2005) experiment

Items	R <sub>c</sub>	D (mm)	U (m sec <sup>-1</sup> )	T (K)	X <sub>O<sub>2</sub></sub>	X <sub>N<sub>2</sub></sub>	X <sub>H<sub>2</sub>O</sub>	X <sub>CH<sub>4</sub></sub>
Fuel jet	28,000	4.57	100	320	0.15	0.52	0.0029	0.3300
Coflow	23,300	210	5.4	1350	0.12	0.73	0.1500	0.0003

R<sub>c</sub>: Reynolds number, D: Diameter, T: Temperature, X: mole fraction

**Lagrangian paths in the composition space:** Therefore of the aerothermochemistry, the equation of the mass fraction of the species is:

$$\frac{\partial \rho Y_i}{\partial t} + \frac{\partial \rho u_i Y_i}{\partial x_k} = \frac{\partial}{\partial x_k} \left( \rho D \frac{\partial Y_i}{\partial x_k} - \rho (u_i Y_i) \right) + \rho \omega_Y \quad (1)$$

This equation is in the non-stationary form; (I) convection; (II) diffusion; (III) source; (IV) the term of diffusion being subdivided in molecular diffusion and turbulent diffusion. While returning the term of turbulent diffusion in the member of left which will be paired at the term of convection, all the term of LHS (Left Hand Side) results in a Lagrangian derivative since moreover, the idea is to follow the history of a particle. The laminar term of diffusion (molecular) is modelled by IEM (Interaction par Echange avec la Moyenne) of Villermaux and Devillon (1972); the researchers associate it an equation of the mixing fraction  $\phi$ :

$$\frac{dY}{dt} = \frac{\tilde{Y} - Y}{\tau} + \omega_Y \quad \text{and} \quad \frac{d\phi}{dt} = \frac{\tilde{\phi} - \phi}{\tau} \quad (2)$$

For a time of exchange shown  $\tau$ , the resolution of the Eq. 2 lead to the trajectories of the MIL model in the space of the phases; the assumption of sudden chemistry allows the particles burnt after a certain time, thus leaving branch of mixture by a jump to arrive on the equilibrium branch as shown in Fig. 2. On lines IEM, the reaction rate can be neglected, the previous system shows:

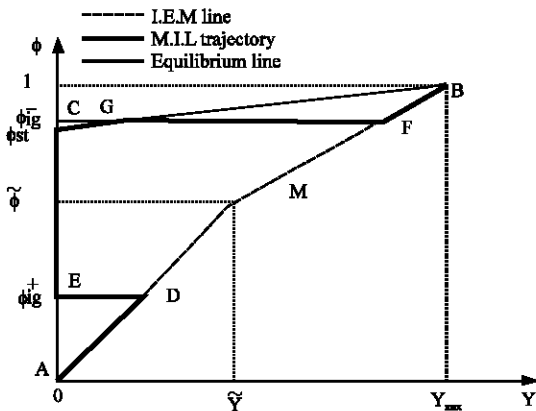


Fig. 2: Conventional representation of the MIL model (thick lines) in the phase plane ( $\phi$ ,  $Y$ ). Equilibrium (plain line), IEM mixing lines (dashed lines) are also depicted

$$\frac{dY}{d\phi} = \frac{\tilde{Y} - Y}{\tilde{\phi} - \phi} \quad (3)$$

Integration gives the branches AM and BM of Fig. 2. The particles need a finite time before ignition; they follow trajectories IEM (AD and BF); then after the ignition occurs, they reach the equilibrium branches instantaneously (DE and FG). Through the simple form of the MIL model, the trajectories in the composition space are regarded as straight segments;  $\phi_{ig}^+$  and  $\phi_{ig}^-$  represent the values of corresponding to the ignition of the particles coming from pure oxidant and pure fuel not A and B, respectively from Fig. 2.

On the equilibrium branch, there are  $Y = 0$  from pure oxidant then from pure fuel following expression  $Y = Y_{max}/(1 - \phi_{st}) (\phi - \phi_{st})$  what corresponds respectively to lines BC and AC. On the equilibrium branch:

$$Y(\phi) = \begin{cases} \frac{\tilde{Y}}{\tilde{\phi}} \phi & \text{If } \phi < \tilde{\phi} \\ \frac{\phi - 1}{\tilde{\phi} - 1} \tilde{Y} + \frac{\tilde{\phi} - \phi}{\tilde{\phi} - 1} Y_{MAX} & \text{elsewhere} \end{cases} \quad (4)$$

**Mean production rates:** The instantaneous reaction rate is reduced to a function of two variables:

$$\omega_{Y_i} = \omega_{Y_i}(\tau, \phi) \quad (5)$$

The average of the reaction rate for each species is shown as follows:

$$\bar{\omega} = \int_0^1 \int_{\tau}^{\infty} \tilde{P}(\tau, \phi) \omega_{Y_i}(\tau, \phi) d\tau d\phi \quad (6)$$

The variables being independent,  $\tilde{P}(\tau, \phi) = \tilde{P}(\tau) \tilde{P}(\phi)$  are written what leads to:

$$\overline{\rho \omega_{Y_i}} = \bar{\rho} \int_0^1 \tilde{P}(\phi) \int_{\tau}^{\infty} \omega_{Y_i} \tilde{P}(\tau) d\tau d\phi \quad (7)$$

By considering a given particle since the idea of the model is to follow the history of it, the Eq. 2 lead to:

$$\omega_Y = \frac{1}{\tau} \left[ \frac{dY}{d\phi} (\tilde{\phi} - \phi) - (\tilde{Y} - Y) \right] \quad (8)$$

The line portions corresponding to this rate of reaction are the mixing (segments AD and BF), jump (segments DE and FG) then equilibrium (segments EC and GC). Mixing branch on the segments AD and BF, particles do not react; the reaction rate zero:

$$\omega_Y^{MIX}(\tau, \phi) = 0 \quad (9)$$

Jump branch on the segments DE and FG, the reaction rate is considered as follows:

$$\omega_Y^{\phi_{ig}^+}(\tau, \phi) = \tilde{P}(\phi) \left( \frac{\tilde{\phi} - \phi_{ig}^+}{\tau} \right) \delta Y_{\phi_{ig}^+} \quad (10)$$

With,

$$\delta Y_{\phi_{ig}^+} = \frac{\tilde{Y}}{\tilde{\phi}} \phi_{ig}^+$$

$$\omega_Y^{\phi_{ig}^-}(\tau, \phi) = \tilde{P}(\phi) \left( \frac{\tilde{\phi} - \phi_{ig}^-}{\tau} \right) \delta Y_{\phi_{ig}^-} \quad (11)$$

With,

$$\delta Y_{\phi_{ig}^-} = \left( \frac{Y_{MAX} - \tilde{Y}}{1 - \tilde{\phi}} - \frac{Y_{MAX}}{1 - \phi_{st}} \right) \phi_{ig}^- + \left( 1 - \frac{\phi_{st}}{1 - \phi_{st}} \right) Y_{MAX} - \frac{1}{1 - \tilde{\phi}} (Y_{MAX} - \tilde{\phi})$$

Equilibrium branch on the segment GC, we have:

$$\omega_Y^{EQU}(\tau, \phi) = - \frac{(\tilde{Y}(1 - \tilde{\phi}) - Y_{MAX}(\tilde{\phi} - \phi_{st}))}{\tau(1 - \phi_{st})} \quad (12)$$

on segment EC, since  $Y = 0$  and  $dy/dZ = 0$ , we have:

$$\omega_Y^{EQU}(\tau, \phi) = -\frac{\tilde{Y}}{\tau} \quad (13)$$

The instantaneous reaction rate is the sum of the three contributions as follows:

$$\omega_Y(\tau, \phi) = \underbrace{\omega_Y^{MIX}(\tau, \phi)}_{\text{mixing}} + \underbrace{\omega_Y^{\phi_{sc}^+}(\tau, \phi) + \omega_Y^{\phi_{sc}^-}(\tau, \phi)}_{\text{jump at } \phi_{sc}^+ \text{ and } \phi_{sc}^-} + \underbrace{\omega_Y^{EQU}(\tau, \phi)}_{\text{equilibrium}} \quad (14)$$

The mean reaction rate is given by the following integral:

$$\tilde{\omega}_Y = \frac{\rho \omega \tilde{Y}}{\bar{\rho}} = \int_0^1 \tilde{P}(\phi) \int_{\tau}^{\infty} \omega_Y \tilde{P}_{\tau}(\tau) d\tau d\phi = \tilde{\omega}_Y^{MIX} + \tilde{\omega}_Y^{\phi_{sc}^+} + \tilde{\omega}_Y^{\phi_{sc}^-} + \tilde{\omega}_Y^{EQU} \quad (15)$$

**Expression of probability density function:** The coupling between turbulence and chemistry results in the pdf appearing in the expression of the mean reaction rate, shown by the MIL model.  $P(\phi)$  who is the probability so that the mixing takes place is taken in this version of the model as being a function rectangle and peak give by Borghi and Moreau (1977); we know that this produced of the satisfactory results in its four configurations as had obtained Obounou *et al.* (1994). Let us specify that a more general formulation is in the use of the beta function Press *et al.* (1986). After studies undertaken by Fox (1995) then by Sabel'nikov and Gorokhovski (2001) on the introduction of a model with multiple scale considering the whole spectrum of the scales of time being located between the micro-scales and the integral scale it arises that  $P_{\tau}(\tau) = \delta(\tau - \tau^*)$  with  $\tau$  which is the scale of a mean mixing time so that the chemical reaction occurs and  $\delta$  fine-grained pdf or probability density of a realization.

This use is justified by the fact that it avoids an integration moreover according to  $\tau$ . Being in a situation of lifted flame, in order to better represent the premixing which is done before effective combustion, we consider like a Log-normal function like Bray *et al.* (1984):

$$P_{\tau} = \frac{1}{\tau_T - \tau_K} \exp\left(\frac{\tau_K - \tau}{\tau_T - \tau_K}\right) \quad (16)$$

With

$$\tau_T = \frac{k}{\varepsilon} \text{ and } \tau_K = \left(\frac{\Theta}{\varepsilon}\right)^{\frac{1}{2}}$$

**Chemical kinetics:** The determination of the ignition delays translating time necessary so that the ignition occurs, passes by the use of a kinetic mechanism. Knowing well that a global kinetic diagram with one equation shows only satisfactory results near to

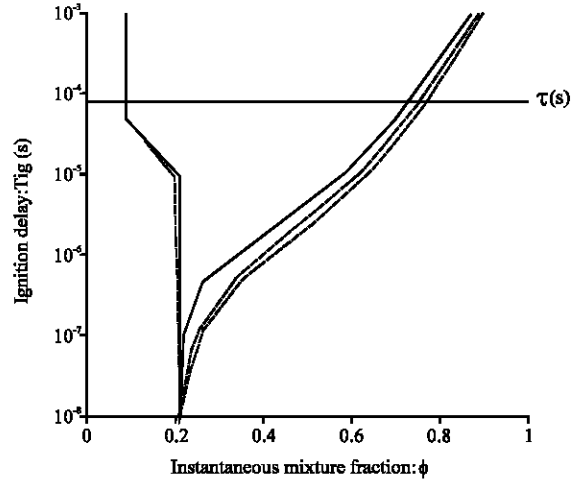


Fig. 3: Ignition delay versus instantaneous mixture fraction;  $\bar{\phi} = 0.13, Y_{O_2} = 0.002812$  (solid line),  $\bar{Y}_{O_2} = 0.01095$  (dashed line),  $\bar{Y}_{O_2} = 0.01637$  (dashdot line)

stoichiometry; Obounou *et al.* (1994) showed that more chemistry is detailed (complexes chemistry), more the precision of the chemical time is good in a rather broad range. Since here we are interested in the auto-ignition, the calculation retained for the tabulation of chemical times is of type PSR (Perfect Stirred Reactor), supporting a homogeneous autoignition; more the number of points taken for ranging between 0 and 1 is large, more the precision is good. The determination then the tabulation of these chemical delays are made by use of package Fortran CHEMKIN2 of the Sandia laboratory; the kinetic mechanism is described by the GRI 3.0 of Bowman *et al.* (1997) corresponding to the combustion of methane in the air which involves 5 elements, 53 species and 325 reactions. The delays are presented in the form of curves in U; Fig. 3 arises the profiles for pure oxidant then for pure fuel; ignition occur by comparison of this chemical time to the mixture time. Thus, it can predict partial extinctions when chemistry is slow (is located in top of the curves in U), we also captures finite chemistry (is located on the curves in U) then infinitely fast chemistry. On the basis of the variable of Schvab-Zeldovich, the mixture fraction for the configuration is shown by:

$$\phi = \frac{2 \frac{M_{O_2}}{M_{CH_4}} Y_{CH_4} - Y_{O_2} - 2 \frac{M_{O_2}}{M_{CH_4}} Y_{CH_4}^{flow} + Y_{O_2}^{flow}}{2 \frac{M_{O_2}}{M_{CH_4}} Y_{CH_4}^{fuel} - Y_{O_2}^{fuel} - 2 \frac{M_{O_2}}{M_{CH_4}} Y_{CH_4}^{flow} + Y_{O_2}^{flow}} \quad (17)$$

With stoichiometry,  $Y_{CH_4} = Y_{O_2} = 0$ ; what leads to  $\phi_{st} = 0.1769$  similar that the value shown by the literature; the curves of the ignitions times are thus shifted towards this position.

RESULTS AND DISCUSSION

**Diagram of the computational domain:** The field of calculation drawn then meshed by GAMBIT 2.4.6 is taken 2D axissymmetric like Fig. 4; this field is then sub-divided in under fields in order to refining the grid in the interesting places such as in the layer of shearing. The researchers implemented the MIL model in the FLUENT 6.3.26 code (2006) through the UDF (User Define Functions) (2006) in order to carry out the numerical simulations.

**Grid independence:** Three grids were tested to determine the solution which does not depend on it any more; the details of the grid are shown in Table 2. Mesh 1 is lightest being used to gauge calculation; meshes 2 and 3 give close results and either may be used to produce a grid-independent solution. However, the finer mesh (Mesh 3) is selected here (although, mesh 2 is optimum) and is used in all subsequent calculations.

**Results of numerical simulations:** The turbulence model retained in FLUENT is the realizable K-ε. Researcher describes the results of simulation per use of the model

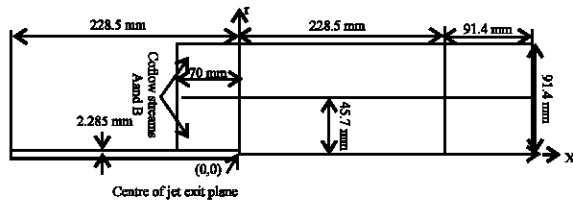


Fig. 4. Diagram of the computational domain

which researcher developed and implemented in FLUENT; a comparison is then made with experimental measurements. The assumption of 2D axissymmetric simulation contrary to a case 3D, product of the tiny differences since the parameters are realised. It is known that the RANS (Reynolds Average Navier-Stokes) simulation led to satisfactory results as in a recent calculation carried out by Mouangue and Obounou (2009). Figure 5 shows the evolution of the radial profile of mean velocity in  $x/D = 15$  and  $30$ . It arises that the MIL model lowers well the values which approach the experimental data; the bringing together is more effective as of the area of pre-flame with  $x/D = 30$ . The MIL model by its aspects Intermittent and Lagrangian, makes it possible to better follow the comparison turbulence-chemistry which translates the position of the flame better. This led on Fig. 6 and 7, respectively arising the

Table 2: Details of the domain mesines

Parameters	Fuel jet	Coflow	Total cells
<b>x</b>			
From (mm)	-228.500	-70.000	-
To (mm)	319.900	319.900	-
<b>r</b>			
From (mm)	0.000	2.285	-
To (mm)	2.285	91.400	-
<b>Mesh 1</b>			
X	55.000	45.000	-
Y	13.000	20.000	1615
<b>Mesh 2</b>			
X	110.000	90.000	-
Y	26.000	40.000	6460
<b>Mesh 3</b>			
X	165.000	135.000	-
Y	39.000	60.000	14,535

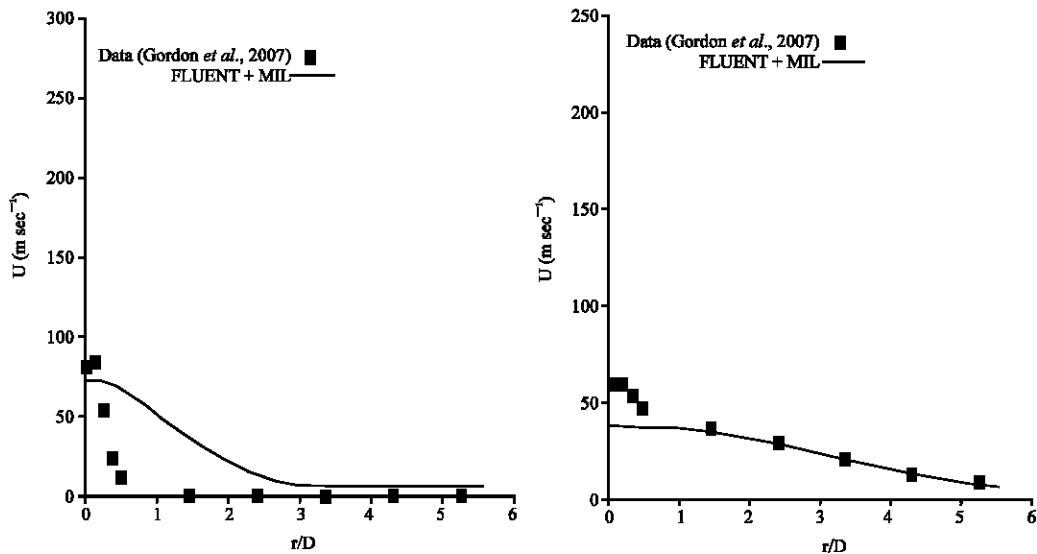


Fig. 5: Radial profiles of the mean velocity at 2 values of  $x/D = 15$  and  $30$  (from left to right); FLUENT+MIL (solid line), measurements (symbols) eddy dissipation (dashed line)

position of the flame and the profiles of temperature. It is observed that the calculation is satisfactory; the radial profiles for temperature in Fig. 7 indicate no reaction

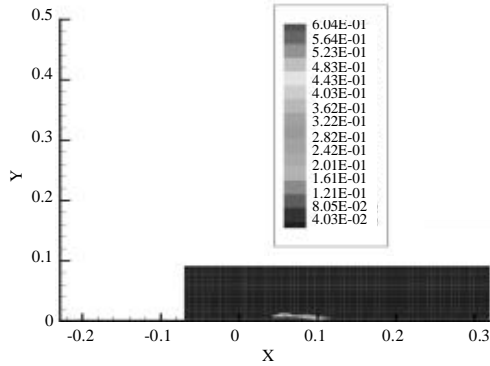


Fig. 6: Mean reaction rate field  $\bar{\omega}_y$  ( $s^{-1}$ )

in the first 30 diameters from the exit nozzle. Initially, mixing without reaction occurs between the vitiated coflow and fuel jet. It is not until  $x/d = 40$  that we see a distinct increase in the temperature fluctuations and a rise in the mean temperature above the coflow condition.

The values remains close, one ignites when it is necessary and moreover, the liftoff height is well agree with experiments between  $x/D = 30$  and  $x/D = 40$ .

In  $x/D = 50$ , we observe a light shift resulting in the difference in positioning of bump materializing ignition; this comes owing to the fact that the model does not arise very well the premixing which takes place before effective combustion. The flame being lifted, the time of diffusion is more significant in the simulation compared to the experiment; the length of mixing is slightly larger in the calculation.

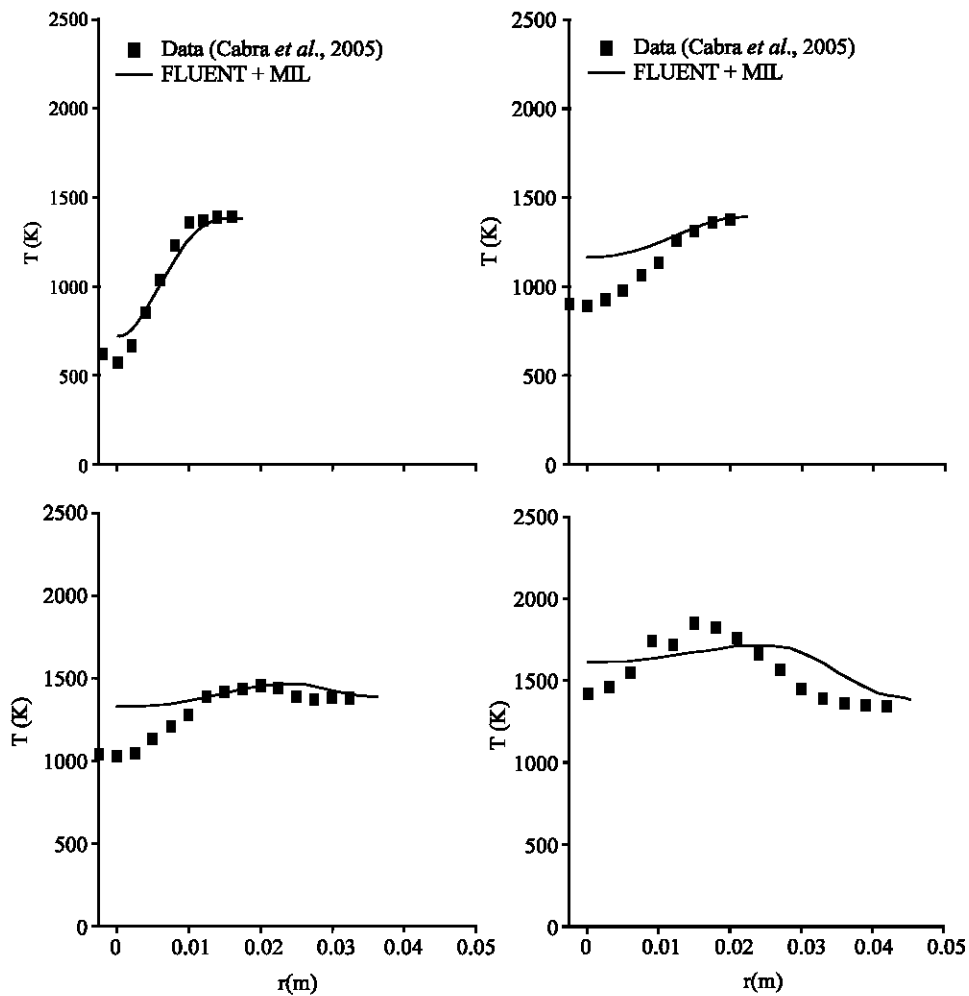


Fig. 7: Radial profiles of the mean temperature at 4 values of  $x/D = 15, 30, 40$  and  $50$  (from left to right then top to bottom); FLUENT+MIL (solid line), measurements (symbols), eddy dissipation (dashed line)

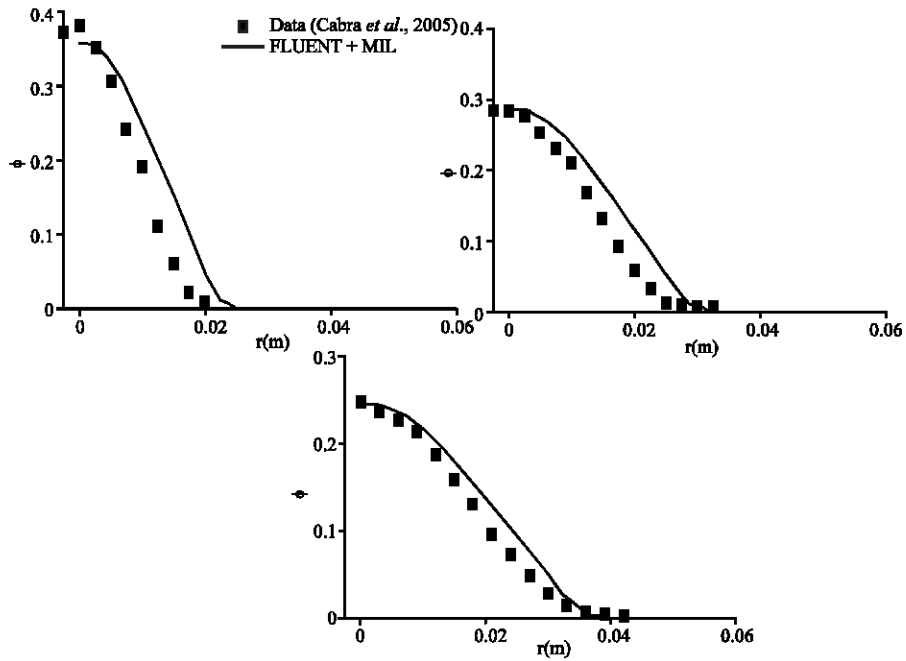


Fig. 8: Radial profile of the mean mixture fraction at 3 values of  $x/D = 30, 40$  and  $50$  (from left to right then top to bottom); FLUENT + MIL (solid line), measurements (symbols), eddy dissipation (dashed line)

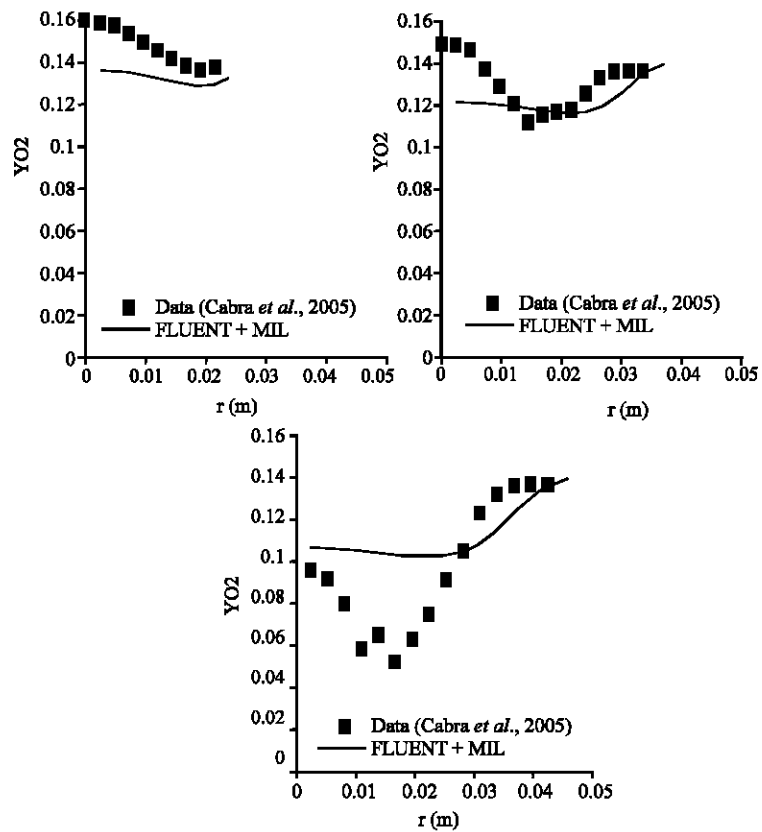


Fig. 9: Radial profile of the  $O_2$  mean mass fraction at 3 values of  $x/D = 30, 40$  and  $50$  (from left to right then top to bottom); FLUENT + MIL (solid line), measurements (symbols), eddy dissipation (dashed line)

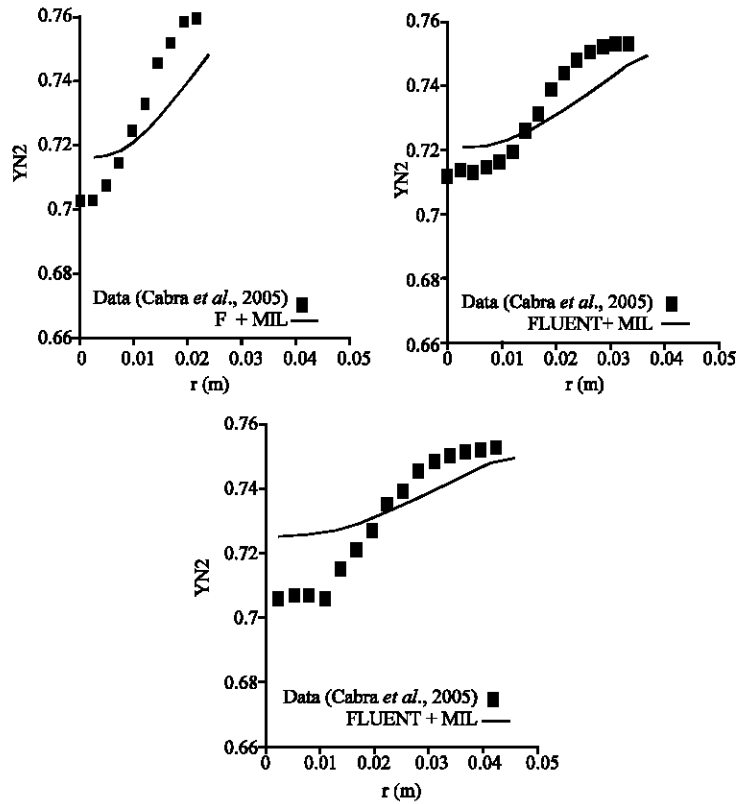


Fig. 10: Radial profiles of the  $N_2$  mean mass fraction at 3 values of  $x/D = 30, 40$  and  $50$  (from left to right and top to bottom); FLUENT + MIL (solid line), measurements (symbols), eddy dissipation (dashed line)

Above the place of stabilization of the flame appears; this for a given kinetics. The flame is puffed up and we observe being separation towards a position enters  $x/D = 30$  and  $x/D = 40$  (Cabra *et al.*, 2005) at the beginning of the calculation, the ignition of the flame occurs downstream of this steady state stabilization point. Afterward, during the convergence of the calculation, the flame front propagates toward this point. The reasons of this shift are confirmed on the curves of Fig. 8 shows the mixing fraction. The dynamics of the vortices strongly improves oxidant and fuel mixing; what is not taken into account in the calculation. We obtain nevertheless, the same evolutions. This influence is observed naturally on the profiles of the species, since it is they which mixing. Fig. 9 shows the radial profile of the average mass fraction of dioxygene. Because the model over-estimates the mixing layer, it predicts low values of the mass fraction with  $x/D = 30$  and  $40$ , then raised values with  $x/D = 50$ . We have the same tendency on Fig. 10 which shows the evolution of the inert. In agreement with Cao *et al.* (2005), then Cabra *et al.* (2005), the turbulence intensity presents a bimodal form Fig. 11 shows this distribution. The bimodal distribution makes it possible the flame to be spread from the outside into the core area.

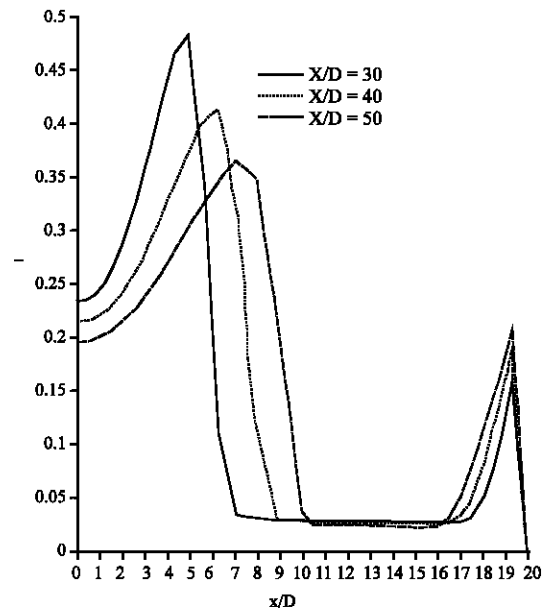


Fig. 11: Distribution of the turbulence intensity (FLUENT + MIL);  $x/D = 30$  (solid line),  $x/D = 40$  (dashed line,  $x/D = 50$  (dash-dot line)



## CONCLUSION

The Lagrangian Intermittent Model was proposed with for objective describing auto-ignition then non-premixed combustion and the partially premixed turbulent flames. Complex chemistry was tabulated based on a calculation PSR with mechanism GRI 3.0 for methane Bowman *et al.* (1997). Their implementation was made in the commercial code FLUENT with a formulation RANS; the simulation of the experiment of Cabra *et al.* (2005) of lifted flame to methane/air is produced. The comparison between calculation and experimental measurements in term of radial profile of mean velocity, temperature, from mixing fraction and average mass fraction of the species is arisen with an agreement; the liftoff is well captured. Concerning the perspective for improvement of this research in the short run, we will propose a calculation on this same flame using the version modified of the MIL model; this version, in addition to its capacity to take into account engines PSR or not already proved reliable within the framework of the lifted flames as in study of Mura and Demoulin (2007). The phenomenon of auto-ignition will be captured better by a calculation SENKIN which moreover, a study of sensitivity of the kinetic diagrams allows. It would be interesting to take a formulation of as a beta function which are more general than the rectangle and peak which were used. Moreover, the chemical we could be improved by taking tables of delays more discretized much using the more recent kinetic diagrams.

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