Impact of the chemical description on a Large Eddy Simulation of a lean partially premixed swirled flame.

Benedetta Franzelli a, Eleonore Riber a and Bénédicte Cuenot a

aCERFACS, CFD Team, 42 Avenue G. Coriolis, 31057 Toulouse Cedex 01, France

Received *****; accepted after revision +++++
Presented by B. Franzelli

Abstract

Six different chemical reduced mechanisms are used in a Large Eddy Simulation of a lean partially premixed swirled methane/air flame in order to investigate their capability to describe the flame structure and the species concentrations comprising the pollutant CO species. The mechanisms range from a two-step fitted mechanism to an analytical scheme composed by 13 species and 73 reactions. Following the classical approach, the performances of the mechanisms have been preliminary analyzed on laminar unstrained free flames. In addition, results for strained premixed counterflow flames have been discussed in order to evaluate their response to turbulence in a very simple way. This work demonstrates that the capability of a mechanism to describe three-dimensional complex turbulent premixed flames could be estimated on results for laminar one-dimensional unstrained and strained flames.

Résumé

Impact du modèle cinétique dans une Simulation aux Grandes Echelles d’une flamme méthane-air swirlée en régime partiellement prémélangé pauvre.

Six cinétiques chimiques réduites sont utilisées pour le calcul d’une flamme swirlée en régime partiellement prémélangé pauvre avec l’approche de Simulation aux Grandes Echelles afin d’évaluer leur capacité à décrire la structure de flamme et la composition chimique, y compris l’espèce polluante CO. Les mécanismes cinétiques testés diffèrent par leur complexité, le plus simple étant un mécanisme global à deux étapes, le plus complexe étant un schéma analytique comprenant 13 espèces et 73 réactions. Pour évaluer leurs performances, les mécanismes sont tout d’abord testés classiquement dans des flammes laminaires prémélangées non étirées se propageant librement. Puis, des calculs de flammes laminaires prémélangées étirées à contre-courant sont analysés pour évaluer simplement la réponse des différents schémas à la turbulence. Ce travail montre que la capacité d’un mécanisme à décrire correctement une flamme turbulente prémélangée dans une configuration complexe peut être anticipée en analysant les réponses du mécanisme dans des flammes prémélangées laminaires non étirées et étirées.

Key words: chemical description; partially premixed swirled flame; response to stretch

Mots-clés: cinétique chimique; flamme swirlée prémélangée; réponse à l’étirement
1. Introduction

A growing need for simulations based on reliable chemistries has been underlined in the last years [1] since restrictions on pollutant emissions motivate request for more accurate results. Detailed kinetic mechanisms, comprising hundreds of species and thousands of reactions, correctly predict multiple aspects of flames over a wide range of cases (i.e. one-dimensional flame structure, gas composition in a stirred reactor, ignition delay, etc...) and are available for most hydrocarbons [2]. Unfortunately, using them to simulate turbulent reacting flows in complex geometries is still prohibitive mainly due to their computational cost. On the one side, the computational time drastically increases with the number of species to be solved; on the other side, complex schemes are usually very stiff and demand specific (implicit) algorithms to avoid unreasonably small time steps.

Two approaches have been proposed to overcome this problem: reduced chemistry [3,4,5] where a detailed mechanism is simplified in order to obtain an accurate chemical behavior using fewer species and reactions, and tabulated chemistry [6,7,8] which is based on the idea that information from academic laminar configurations could be stored into a table and used for turbulent calculations. Using tabulation method is still difficult in complex configurations since determining the prototype flame to create the table can be a complicated task if the combustion regime is unknown. For this reason, only reduced chemistries are considered in this work. Two families could be distinguished: fitted schemes and analytical mechanisms. Fitted schemes are characterized by reaction parameters which have been fitted to capture some global quantities such as flame speed or burnt gas state. Analytical mechanisms are systematically derived from detailed or skeletal schemes to include more details on the flame providing a physical insight into the chemical processes. Schemes belonging to the first family are generally easier to develop and implement into a CFD solver compared to the second family’s mechanisms. These simplified chemical descriptions should be carefully used when simulating three-dimensional turbulent complex flames since some pieces of information are neglected to reduce the computational cost possibly affecting the accuracy [9,10]. Moreover, all these reductions have been developed and evaluated for laminar configurations and their impact on turbulent unsteady flames has not been completely evaluated [11].

The objective of this work is to identify the characteristics of a reduced mechanism mostly impacting the LES of realistic turbulent premixed flames in order to build from one-dimensional tests a reliable reduced scheme for three-dimensional turbulent configurations or select in a priori way the mechanism offering the best compromise between CPU cost and result accuracy. Six chemical mechanisms for methane/air flames are tested in this work with increasing complexity (Section 2.1): the two-step fitted schemes 2S\textsubscript{CH4,BFER} [12] and 2S\textsubscript{CH4,BFER*}, the four-step fitted JONES mechanism [13] and the analytical schemes PETERS [14], SESHADRI [15] and LU [16]. Following the classical approach, their performances on laminar premixed free flames are first discussed in Section 2.2 in terms of flame speed, burnt gas temperature and species profiles. Turbulence mainly interacting with the flame by straining and wrinkling its front, the response of the mechanisms to strain is analyzed in strained premixed counterflow flames to evaluate their response to turbulence in a very simple way (Section 2.3). Once the performances of the mechanisms have been discussed on laminar configurations, their behavior is evaluated on Large Eddy Simulation of the experimental partially premixed swirled burner named PRECCINSTA [17] (Section 3) to verify the impact of the mechanism characteristics identified on the laminar cases.

Email addresses: franzell@cerfacs.fr (Benedetta Franzelli), riber@cerfacs.fr (Eleonore Riber), cuenot@cerfacs.fr (Bénédicte Cuenot).
2. Reduced chemical mechanisms

2.1. Main characteristics of the six mechanisms

The six mechanisms are described in the following in order of increasing complexity.

The simplest scheme is the two-step 2S CH4 BFER mechanism [12] developed following the methodology described in [18]. It accounts for six species (CH4, O2, CO2, CO, H2O and N2) and two reactions, the methane oxidation and the CO – CO2 equilibrium. Both reactions follow an Arrhenius law and unity Lewis numbers are assumed for all species (Table 4). This scheme correctly predicts the flame speed for laminar free flames on a wide range of fresh gas temperature (300 K < Tf < 700 K), pressure (1 atm < P < 10 atm) and equivalence ratio (0.6 ≤ φ ≤ 1.5) as well as the burnt gas temperature for φ < 1.4.

The 2S CH4 BFER scheme does not respond well to strain (Section 2.3). Therefore, a modified version (2S CH4 BFER*) is proposed where the species Lewis numbers are modified (Leφ = 1.65) since the flame response to strain is strongly affected by species Lewis numbers [19]. The pre-exponential factor for the methane oxidation reaction of the original version has been adjusted multiplying it by a factor 0.8 to correctly reproduce the flame speed.

The JONES scheme [13] is slightly more complex, comprising seven species (CH4, O2, CO2, CO, H2O, H2 and N2) and accounting for four reactions. Each reaction follows an Arrhenius law whose parameters have been chosen in order to correctly describe the flame structure of both premixed and diffusion flames at ambient temperature (Tf = 300 K) and atmospheric pressure.

The analytical PETERS mechanism [14] accounts for eight species (CH4, O2, CO2, CO, H2O, N2, H2 and H) and four reactions. It has been systematically derived on a skeletal mechanism for lean methane/air flames composed of 18 reactions and 13 species. This mechanism has been tested on laminar premixed and diffusion flames at ambient temperature and atmospheric pressure.

The SESHADRI scheme is an improved version of the PETERS scheme proposed by Seshadri and Peters [15] by using a computer program for optimization and reduction of detailed mechanisms. It takes into account the same species and reactions as the PETERS mechanism but the global reaction rates are based on a reduction of a more complex skeletal mechanism composed of 25 reactions.

The LU mechanism [16] is the most complex reduced mechanism analyzed in this work. It takes into account 13 resolved species (CH4, O2, CO2, CO, H2O, N2, H2, H, OH, O, HO2, CH3 and CH2O) and 73 elementary reactions. It has been derived from the detailed mechanism GRI1.2 applying the directed relation graph method, the sensitivity analysis and the computational singular perturbation approach [16]. This scheme shows a satisfactory behavior on lean premixed methane/air flames, perfectly stirred reactor for Tf = 300 K and auto-ignition configurations from 1000 K to 1800 K at atmospheric pressure.

Simplified transport properties are used for all reduced mechanisms: the Prandtl number is assumed constant (Pr = 0.7) whereas the Lewis numbers are assumed constant but not equal for each species except for the two two-step schemes (Table 4). The molecular viscosity μ follows a power law in temperature μ = μ0(T/T0)α, where α = 0.6759 is the power law constant, T0 = 300 K is the reference temperature and μ0 = 1.8405 × 10−5 kg/m/s is the reference dynamic viscosity. This set of parameters enables to fit the dependence on temperature over the whole range of temperature considered in this work.

The performances of the six chemical mechanisms are compared for laminar premixed free (Section 2.2) and counterflow flames (Section 2.3) to results obtained using the detailed GRI3.0 mechanism [20] chosen as reference. It is a compilation of 325 elementary chemical reactions and associated rate coefficient expressions, thermochemical parameters and complex transport properties for the 53 species involved in them. Results correspond to the operating point of the PRECCINSTA burner, i.e. initial temperature
$T_f = 320 \, K$ and atmospheric pressure. All calculations presented in this section have been performed with CANTERA [21].

2.2. Results on premixed free flames

Figure 1a. shows the flame speed obtained with the reduced schemes together with results of the detailed GRI3.0 mechanism over the whole range of flammability at initial temperature $T_f = 320 \, K$ and atmospheric pressure. For lean and stoichiometric mixtures, all mechanisms correctly reproduce the flame speed. For rich mixtures, the decrease in flame speed is well predicted by all mechanisms except the JONES and PETERS schemes which greatly overpredict it. However, for the composition of interest ($\phi = 0.83$), the agreement with the detailed GRI3.0 scheme is satisfactory for all mechanisms and the largest error (about 15%) is found for the SESHADRI scheme.

The same comparison is displayed in Fig. 1b. for the burnt gas temperature. The agreement with the GRI3.0 reference scheme is satisfactory for all the mechanisms except the two two-step schemes which overestimate the burnt gas temperature for very rich flames ($\phi > 1.4$). The introduction of additional species in the JONES and the analytical mechanisms enables to improve the description of the burnt gas temperature. Discrepancies are detected for near-stoichiometric mixtures except for the most complex reduced LU scheme which is the only one to correctly predict the burnt gas temperature on the whole range of equivalence ratio. The error on the burnt gas temperature at $\phi = 0.83$ is less than 1% for all the schemes.

Spatial profiles of $CH_4$, $CO_2$ and $CO$ species representing respectively reactants, products and intermediate species for the reduced schemes are compared to the GRI3.0 scheme at $\phi = 0.83$, $T_f = 320 \, K$ and $P = 1 \, atm$ in Fig. 2. It has been chosen to zoom in the reaction zone where the main discrepancies occur. Therefore the equilibrium state can not be evaluated from Fig. 2 but it has been verified that it is correctly described by all schemes. The $CO_2$ spatial profile predicted by the GRI3.0 scheme shows two different zones: a first reaction zone characterized by a high gradient, and a second post-flame zone where recombination takes place and $CO_2$ increases slowly. This structure is not captured by the two-step mechanisms which are not able to reproduce the slower recombination zone and reach equilibrium too quickly. The JONES scheme shows good agreement with the detailed mechanism but better results are obtained with all analytical mechanisms.

A correct description of the $CO$ concentration is necessary to predict pollutants but reproducing the maximum value of $CO$ species is a hard task since it is first produced in the reaction zone and then oxidized.
into $CO_2$ in the recombination region. The two-step schemes predict an unphysical monotonous profile and greatly underestimate its maximum value. The value at equilibrium is however correctly described. The JONES results are in good agreement with the detailed mechanism, with a reasonable maximum level although the equilibrium in the post flame region is reached too quickly. The analytical mechanisms predict almost perfectly the flame structure: the maximum level of CO species is well captured and the equilibrium is reached slowly, as predicted by the GRI3.0 scheme.

2.3. Results on premixed counterflow flames

The response to strain rate is evaluated for all reduced mechanisms on classical strained one-dimensional premixed flames (Fig. 3a). A fresh premixed methane/air mixture is injected at $T_f = 320$ K and $\phi = 0.83$ on the left side and combustion products are injected on the right side. The strain rate $a$ is evaluated in the reaction zone identified by an isoline of the progress variable based on $O_2$ species and equal to $c = 0.65$:

$$a = \frac{\partial v}{\partial y}.$$  \hfill (1)

As the mass flow rate increases, the flame is more strained by the velocity field, making the flame front generally thinner and modifying its structure. The response of the mechanisms to strain rate is evaluated in terms of the consumption speed $S_C$ (Fig. 3b):

$$S_C = \frac{1}{\rho^f Y_{O_2}^f} \int \dot{\omega}_{O_2} dx,$$  \hfill (2)

where $\rho^f$ is the fresh gas density, $Y_{O_2}^f$ is the mass fraction of $O_2$ in the fresh gases and $\dot{\omega}_{O_2}$ is the consumption rate of species $O_2$.

From the results of the detailed GRI3.0 mechanism, the consumption speed is expected to decrease when the strain rate increases. This behavior is not correctly reproduced by all schemes: the JONES mechanism shows the opposite tendency, i.e. the consumption speed increases with the strain rate; the 2S,CH4,BFER scheme is not sufficiently affected by the strain rate and too high values of consumption speed are found even for really high strain rates. On the contrary, the modified two-step 2S,CH4,BFER* mechanism shows a great improvement compared to the 2S,CH4,BFER scheme and the agreement with the GRI3.0 mechanism is now satisfactory. The PETERS mechanism is too much affected by strain rate and for high strain rate values the flame quenches. The response to strain rate in terms of consumption speed
speed is better predicted by the SESHADRI mechanism, at least for relatively small strain rates, and the LU scheme.

![Sketch of a premixed strained counterflow flame.](image)

**Figure 3.** a) Sketch of a premixed strained counterflow flame. b) Consumption speed and c) maximum value of the CO mass fraction for a premixed strained methane/air flame as a function of local strain rate at \( T_f = 320 \) K. Comparison between the reduced mechanisms and the GRI3.0 detailed scheme.

The maximum value of CO mass fraction, corresponding to \( c = 0.65 \), is also studied as function of strain rate (Fig. 3c). It is a good indication of the impact of strain rate on the flame structure since intermediate species and radicals are more affected by strain than reactants and products. The two-step mechanisms being unable to predict the CO concentration in the reaction zone for unstrained flames, the response to strain rate in terms of CO mass fraction is completely lost. All other mechanisms show the same tendency, when the strain rate increases the CO mass fraction in the reaction zone decreases. More specifically, the JONES scheme overestimates the CO mass fraction in the reaction zone and this error increases with strain rate. The maximum value of CO is on the other hand correctly predicted by the PETERS mechanism, at least for low strain rates, and by the SESHADRI and LU schemes for the whole range of strain rates compared to results of the GRI3.0 scheme.

Globally, it has been observed that the more complex the mechanism is the higher is the accuracy of results on laminar flames and not all schemes correctly respond to strain in terms of consumption speed (2S\_CH4\_BFER and JONES) or CO mass fraction (2S\_CH4\_BFER, 2S\_CH4\_BFER\* and JONES). A good description of laminar freely propagating flames is suspected not to be sufficient to correctly describe a turbulent flame. More specifically, the turbulent flame structure is expected to strongly depend on the capability of a mechanism to predict the recombination zone and the intermediate species concentrations, strongly affected by strain, on laminar strained flames. Moreover, the turbulent flame length is supposed to depend on the response to strain of the consumption rate and, as a consequence, the consumption speed \( S_C \) has to be correctly reproduced on laminar strained flames. The impact of the result discrepancies on laminar strained flames is evaluated on LES of the PRECCINSTA burner in the following.

### 3. LES of the PRECCINSTA burner

#### 3.1. Experimental and numerical setup

The target experimental burner has been specifically designed to study the impact of partially premixed mixture in flame stability [17]. As sketched in Fig. 4a, air is injected into the plenum through one large...
intake while methane is injected through twelve small tubes of diameter 1 mm directly into the swirler. Methane and air are then mixed by the high momentum flow of the swirler and a methane/air mixture enters the combustion chamber. The flow in the chamber is characterized by an inner recirculation zone (IRZ) and two outer recirculation zones (ORZ), which enables to stabilize a swirled flame with a classical conical shape at the nozzle exit. The burnt gases finally exit the chamber through the exhaust tube. Two regimes have been experimentally observed, depending on the global equivalence ratio. For $\phi = 0.7$, the flame is pulsating where for $\phi = 0.83$ the flame keeps stable. Laser Raman scattering measurements are available for concentration of the major species ($CH_4$, $CO$, $CO_2$, $H_2O$, $H_2$, $N_2$ and $O_2$) and for temperature in vertical planes ($y, z$) at eight different axial positions downstream of the injector ($h = 6, 10, 15, 20, 30, 40, 60, 80$ mm, where $h = 0$ mm corresponds to the exit plane of the nozzle) for at least five radial positions $r$ (Fig. 4b). The systematic and statistical uncertainties are less than 4% and 2.5% respectively for temperature and less than 5% and 7% respectively for almost all species except for $CO$ and $H_2$ for which statistical uncertainty is between 20 − 50% [17].

Numerous simulations have been performed [22,23,24,25,26] assuming a perfect mixing between methane and air at the nozzle exit, which simplifies the computational work. Only recently, fuel/air mixing has been explicitly computed including fuel jets into the swirler in order to estimate the impact of the perfect premixing assumption on prediction of $NO_x$ formation [27] and thermo-acoustic instabilities [12]. In this work, LES are performed without the perfect mixing assumption on the same unstructured mesh shown in Fig. 4b. Dry air and pure methane are injected separately at ambient temperature ($T_f = 320$ K) with air mass flow $\dot{m}_{air} = 734.2$ g/min and methane mass flow $\dot{m}_{CH_4} = 35.9$ g/min, corresponding to the stable operating point ($\phi = 0.83$). The numerical setup proposed in [12] has been used to perform all computations to guarantee consistent comparisons of the results and to correctly identify the impact of the reduced chemical mechanisms. A Taylor-Galerkin weighted residual central distributions scheme is used for numerical integration [28]. The interaction between chemistry and turbulence is modeled by the Dynamically Thickened Flame (DTFLES) model [29]. The sensor activating the flame thickening is based on the net production rate of $CO$ and $CO_2$ species, which guarantees an equivalent thickening for all chemistries in both the reaction and the post-flame zones. The inlets for methane and air and the outlet are described by Navier-Stokes Characteristic Boundary Conditions (NSCBC) [30] to ensure a physical representation of the acoustic wave propagation and reflection. An adiabatic no-slip condition is applied to the walls.
3.2. Analysis of results

In Section 2, the GRI3.0 detailed mechanism has been used as the reference and the LU analytical scheme has shown very good agreements with it. As using the GRI3.0 mechanism to perform LES in the target burner is computationally too expensive, the LU scheme is used as the reference in the following. When analyzing the results, the LU scheme is first compared to the experiments before comparing the performances of the other five mechanisms to the LU results. Figure 5 compares the mean temperature profiles in the vertical mid-plane cut. In Fig. 5.a, the comparison between experiments and the LU results shows good agreement. The discrepancies are likely to be due to some numerical simplifications, such as combustion model, numerical discretization and adiabaticity assumption. Indeed evaluating the temperature both on the ORZ and in the near wall region is inaccurate when neglecting wall heat losses and radiation effects.

Figures 5.b-5.f compare the LU mechanism with the five other mechanisms. Although the overall agreement is acceptable, the chemical models show non-negligible discrepancies. As already said, the flame length is expected to be related to the consumption rate of the reactants, i.e. the consumption speed $S_C$. The higher the consumption speed, the quicker the reactants are burnt and consequently the shorter the flame is. The flames predicted by the 2S_CH4_BFER and the JONES mechanisms confirm the results of the laminar counterflow flames where both schemes overestimate $S_C$. As expected, the modified 2S_CH4_BFER* scheme predicts a longer flame than the 2S_CH4_BFER mechanism. The analytical schemes reproduce a correct flame length. Second, in terms of post flame region, the 2S_CH4_BFER flame reaches rapidly the equilibrium state in agreement with results for laminar free flames, whereas a recombination zone

![Figure 5. Mean temperature field in the vertical mid-plane. a) Comparison between experiments and LU scheme. b-f) Comparison between reduced schemes and reference LU mechanism. The black iso-line of progress variable $c = 0.65$ represents the mean flame surface position.](image)
touching the wall downstream of the flame ($-40 \text{ mm} < r < -30 \text{ mm}$ and $25 \text{ mm} < h < 50 \text{ mm}$) is detected when using analytical schemes. Globally, the flame structure strongly depends on the chemical description used.

Figures 6 and 7 display the mean and fluctuating profiles at five sections ($h = 6, 10, 20, 30$ and $60 \text{ mm}$) in terms of temperature and CO mass fraction. The experiments [17] and the LU results are compared to the results obtained with the five other mechanisms. For the sake of clarity, the 2S\_CH4\_BFER, 2S\_CH4\_BFER* and JONES results are displayed at the top in black whereas the analytical schemes (PETERS and SESHADRI) are shown at the bottom in grey.

Figure 6. a) Mean and b) fluctuating temperature profiles at five sections in the chamber. The experimental (○) and LU (■) results are compared to numerical data for the other mechanisms: top) 2S\_CH4\_BFER (--), 2S\_CH4\_BFER* (—) and JONES (······); bottom) PETERS (— —) and SESHADRI (— —).

Concerning the mean and fluctuating temperature profiles (Figs. 6a. and 6b. respectively) the overall agreement is satisfactory. However, there are some discrepancies between experimental data and LU results: the underestimation of the IRZ extension at $h = 10 \text{ mm}$ and $h = 20 \text{ mm}$, as well as the overestimation of the temperature in the ORZ that could be due to wall heat losses and radiation effects which are neglected. The temperature fluctuations are slightly underestimated but the global behavior is well captured. On the one side, results obtained with the analytical mechanisms are in good agreement with the LU results showing that these two schemes could be used instead of the LU mechanism to reduce the computational cost. On the other side, larger discrepancies are detected when using the fitted schemes which overestimate the mean temperature in the layer between the fresh gases and the ORZ ($x > 20 \text{ mm}$ at $h = 20 \text{ mm}$). Since a smaller flame is predicted by the 2S\_CH4\_BFER and JONES schemes, the equilibrium temperature is already reached in this region contrary to the experiments. The modified 2S\_CH4\_BFER* scheme behaves better than the original scheme but discrepancies are still visible. The results for mean and fluctuating profiles of major species ($\text{CH}_4$ and $\text{CO}_2$ for example) present the same characteristics and are not shown in this paper.

In Fig. 7, the mean and fluctuating profiles of CO species predicted by the five mechanisms are compared to the measurements (note that the experimental error on CO was estimated at $50\%$) and the LU results. Results are in agreement with the behavior of mechanisms in laminar strained flames (Section 2.3). The
simplest 2S\_CH4\_BFER and 2S\_CH4\_BFER\* schemes greatly underestimate the CO mass fraction in the reaction zone but recover the correct level at equilibrium ($h = 60$ mm). On the contrary, the JONES mechanism greatly overestimates the maximum value of mean and fluctuating CO mass fractions in the reaction zone as expected from the laminar analysis. Only the analytical schemes (PETERS, SESHADRI) correctly predict the CO mass fraction although the mean and fluctuating values are slightly overestimated in the IRZ. Moreover, only one peak of CO is experimentally detected in the layer between fresh gases and IRZ whereas a smaller second peak of CO is predicted by the analytical schemes, also pointed out by the LU mechanism. The adiabacity assumption and the mesh refinement could be at the origin of this discrepancy.

4. Conclusion

In this work the impact of using reduced chemical mechanisms in LES of a three-dimensional partially premixed flame has been analyzed. The origin of the discrepancies between the different simulations has been identified in the performances of these mechanisms on unstrained and strained premixed flames. Even if using reduced mechanisms decreases the computational cost compared to the reference LU calculation (Table 2), it could affect the results quality.

Six reduced chemical schemes for methane-air flames have been first tested in academic configurations of one-dimensional premixed free and strained counterflow flames. Results have been analyzed by comparison with the GRI3.0 detailed scheme in terms of laminar flame speed and burnt gas temperature as well as the response to strain. The 2S\_CH4\_BFER scheme has the lowest computational cost but it shows a non-physical behavior for CO and almost no response to strain. This second point can be modified (2S\_CH4\_BFER\*) changing the Lewis numbers without any additional cost. No major improvements are obtained when using the JONES mechanism for an higher computational cost, except a correct
description of the flame structure only for unstrained flames. The analytical schemes (PETERS and SESHADRI) accurately predict the consumption speed, the concentration of intermediate species and the flame structure for a reduced computational cost of about 20% compared to the LU scheme.

These behaviors have then been confirmed in the LES of a 3D partially premixed flame at ambient temperature and atmospheric pressure. These results on both academic and industrial configurations allow to propose a methodology to evaluate the capability of a mechanism to predict some three-dimensional chemical phenomena basing on one-dimensional unstrained and strained laminar flames configurations:

- an accurate description of the consumption speed for laminar strained flames is necessary to correctly reproduce the length and surface of a turbulent flame;
- to predict the CO concentration in turbulent flames, the CO mass fraction in the reaction zone for unstrained and strained flames must be correctly described;
- the presence of a recombination zone for an unstrained flame guarantees the presence of a small temperature gradient region characterized by product formation in a turbulent configuration.

This methodology allows to a priori choose a chemical scheme depending both on the quality of the results required and the computational cost affordable. It has been however evaluated only on premixed flames and still needs validation for diffusion flames and more specific combustion phenomena such as auto-ignition or flame extinction.

Acknowledgements

This research project has been supported by a Marie Curie Early Stage Research Training Fellowship of the European Community’s Sixth Framework Programme under contract number MEST-CT-2005-020426. This work was granted access to the HPC resources of CINES under the allocation 2010-025031 made by GENCI (Grand Equipement National de Calcul Intensif).

References


### Tables

**Table 1**

<table>
<thead>
<tr>
<th>Scheme</th>
<th>CH$_4$</th>
<th>O$_2$</th>
<th>CO$_2$</th>
<th>CO</th>
<th>H$_2$O</th>
<th>H$_2$</th>
<th>N$_2$</th>
<th>H</th>
<th>O</th>
<th>OH</th>
<th>HO</th>
<th>H$_2$</th>
<th>CH$_3$</th>
<th>CH$_2$O</th>
</tr>
</thead>
<tbody>
<tr>
<td>2S_CH4_BFER</td>
<td>1.00</td>
<td>1.00</td>
<td>1.00</td>
<td>1.00</td>
<td>1.00</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td></td>
</tr>
<tr>
<td>2S_CH4_BFER*</td>
<td>1.65</td>
<td>1.65</td>
<td>1.65</td>
<td>1.65</td>
<td>1.65</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td></td>
</tr>
<tr>
<td>Others</td>
<td>0.97</td>
<td>1.06</td>
<td>1.35</td>
<td>1.07</td>
<td>0.78</td>
<td>0.29</td>
<td>0.17</td>
<td>0.7</td>
<td>0.7</td>
<td>0.7</td>
<td>0.81</td>
<td>0.81</td>
<td>1.25</td>
<td></td>
</tr>
</tbody>
</table>

**Table 2**

<table>
<thead>
<tr>
<th>Scheme</th>
<th>Computational time normalized by the computation time of the LES using the LU scheme.</th>
</tr>
</thead>
<tbody>
<tr>
<td>2S_CH4_BFER</td>
<td>0.72</td>
</tr>
<tr>
<td>2S_CH4_BFER*</td>
<td>0.72</td>
</tr>
<tr>
<td>JONES</td>
<td>0.79</td>
</tr>
<tr>
<td>PETERS</td>
<td>0.81</td>
</tr>
<tr>
<td>SESHADRI</td>
<td>0.81</td>
</tr>
<tr>
<td>LU</td>
<td>1.00</td>
</tr>
</tbody>
</table>