

# NUMERICAL SIMULATION OF HIGH PRESSURE BURNER WITH PARTIALLY PREMIXED FLAME

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## ABSTRACT

In this paper we present the results of a 2D axisymmetric parametric study which simulates an atmospheric premixed burner with flame at high pressure, in which methane is burned. A total of nine simulations are performed with different regulators openings of primary and secondary air. Also, it provides a 3D simulation in which entry conditions are the profiles obtained in a 2D axisymmetric simulation, with the intention to note that differences are obtained between 2D and 3D simulation. The simulations are performed using the standard  $k$ - $\epsilon$  model for turbulence, P1 model for radiation and the Finite Rate/Eddy Dissipation model with a simplified 2-step reaction mechanism for combustion. We conclude that when the secondary air regulator is closed, combustion is incomplete. Also, the results of 2D axisymmetric are a good approximation in regards to 3D results.

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**Keywords:** Atmospheric burner, FLUENT, Internal recirculation, Semi-spherical flame, Mathematical models, Premixed flame, Regulators, High pressure, Numerical simulation.

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## RESUMEN

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**E**n este trabajo se presentan los resultados obtenidos de un estudio paramétrico en 2D axisimétrico donde se simula un quemador atmosférico con llama de premezcla a alta presión, en el cual se quema metano. Son realizadas en total nueve simulaciones con diferentes aperturas de los reguladores de aire primario y secundario. También es realizada una simulación 3D que tiene como condiciones de entrada los perfiles obtenidos en una de las simulaciones 2D axisimétrica, con el fin de observar que diferencias se obtienen entre la simulación 2D y 3D. Las simulaciones son realizadas utilizando el modelo  $k-\varepsilon$  estándar para la turbulencia, el modelo P1 para la radiación y para la combustión se utilizó el modelo Finite Rate/Eddy dissipation con un mecanismo reaccional simplificado de 2 pasos. Se concluye que cuando el regulador de aire secundario se encuentra cerrado la combustión es incompleta, y además los resultados obtenidos en 2D-axisimétrico son una buena aproximación con respecto a los resultados en 3D.

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**Palabras claves:** Quemador atmosférico, FLUENT, Recirculación interna, Llama semiesférica, Modelos matemáticos, Llama premezclada, Reguladores, Alta presión, Simulación numérica.

## RESUMO

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**N**este trabalho são apresentados os resultados obtidos de um estudo paramétrico em 2D axissimétrico onde é simulado um queimador atmosférico com chama de pré-mistura a alta pressão, no qual é queimado metano. São realizadas em total 9 simulações com diferentes aberturas dos reguladores de ar primário e secundário. Também é realizada uma simulação 3D que tem como condições de entrada os perfis obtidos em uma das simulações 2D axissimétrica, com o fim de observar que diferenças são obtidas entre a simulação 2D e 3D. As simulações são realizadas utilizando o modelo  $k-\varepsilon$  padrão para a turbulência, o modelo P1 para a radiação e para a combustão foi utilizado o modelo Finite Rate/Eddy dissipation com um mecanismo reaccional simplificado de 2 passos. É possível concluir que quando o regulador de ar secundário está fechado a combustão é incompleta, e, além disso, os resultados obtidos em 2D-axissimétrico são uma boa aproximação com relação aos resultados em 3D.

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**Palavras-chaves:** Queimador atmosférico, FLUENT, Recirculação interna, Chama semi-esférica, Modelos matemáticos. Llama pré-misturada, Reguladores, Alta pressão, Simulação numérica.

## 1. INTRODUCTION

This work proposes the simulation of a furnace used in the refining industry.

The simulation aims to obtain important furnace operating parameters to allow enhanced furnace production. In order to simplify the furnace simulation, the problem is divided in two stages: the first covers the simulation of a furnace burner which results are taken into the second stage to be used as input conditions to simulate the furnace itself. The work of taking the burner results and developing the furnace simulation was made by Arrieta, Cadavid and Arrieta (2011). The currently work involves the first stage, i.e. only the burner simulation.

In this work, the most important burner features are shown: first of all, it is a partially premixed atmospheric burner where the combustion air is entrained from the environment by a high-speed jet. Figure 1 shows a scheme of this burner with its main parts, where a high pressure methane jet is discharged from the nozzle into the venturi or mixer, causing the effect of decreasing the pressure within itself, therefore dragging ambient air that is at atmospheric pressure. The amount of air entering the mixer is regulated by the primary air obturator with an axial displacement. Then the mixture of methane and air moves toward the burner head, being discharged into the furnace where the flame takes place. The air required to complete combustion is entrained from the external environment by the flame and regulated with the secondary obturator. As shown in Figure 1, the discharged of the fuel-air mixture in the burner head is made tangentially on a dome carved into the furnace wall, which gives stability to the flame and causes it to be shorten and with tendency towards a spherical shape.

Although current simulations involving turbulence tend to be modeled with Large Eddy Simulation (LES), models involving Reynolds Average Navier-Stokes (RANS) equations remain as the preferred tool for modeling industrial applications due to lower computational demand that is required by the latter.

Among recent works developed to simulate turbulent flames involving RANS models, the one elaborated

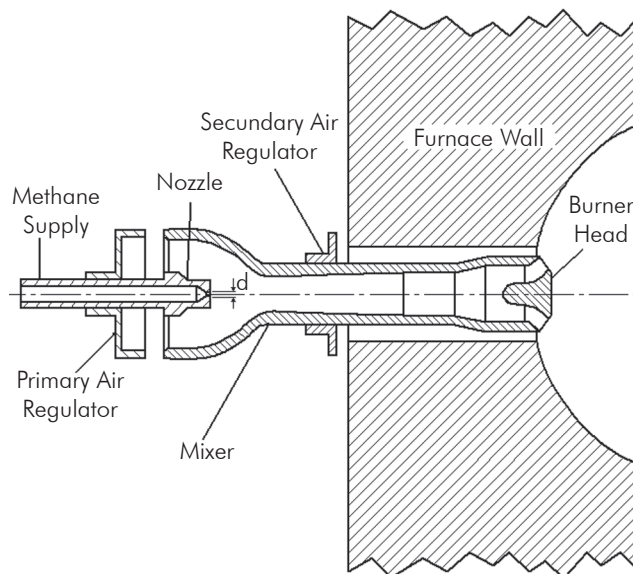


Figure 1. Burner scheme used in the simulations, where major parts are highlighted.

by Herrmann (2006) simulated three methane Bunsen burners which differ only in the Reynolds number. In his paper he uses the turbulence model of two equations standard  $k-\epsilon$ , and finds the concentration of chemical species through a probability density function (PDF). The results obtained are acceptable in comparison to experimental data. Jiang, Liang, Huang and Li (2006) conducted a simulation of a diffusion flame of  $CH_4/air$ , for which three different turbulence models are used ( $k-\epsilon$  realizable,  $k-\omega$ , and the Reynolds Stress Model) finding that the best results are obtained with the  $k-\omega$  model. Bidi, Hosseini and Nobari (2008) simulated a premixed flame of  $CH_4/air$  in a cylindrical combustion chamber using the  $k-\epsilon$  standard model. The reaction rates are found with the Finite Rate/Eddy Dissipation model with a 5 step reactional mechanism, taking into account the effect of radiation by using the Discrete Ordinate Method (DOM). The absorption coefficient is also calculated by using a weighted sum of gray gases (WSGGM), obtaining results that are in excellent agreement with experimental outcomes.

The standard  $k-\epsilon$  model is generally good, but when the flow is restricted by a wall, results lose accuracy, whereas the  $k-\omega$  model is better when the wall is present (Roy & Blottner, 2006). Despite this, the standard  $k-\epsilon$  model is widely used due to its simplicity.

In this work we use the commercial software FLU-ENT 6.2.16 to study the behavior of the burner shown in

Figure 1, using methane as fuel. First, the study develops a parametric 2D axisymmetric model in which different primary and secondary regulators openings are simulated. Also, the simulation of a 3D model is performed, using as input conditions the results obtained from a 2D axisymmetric simulation.

## 2. 2D-AXISYMMETRIC MODEL

It can be considered that both the burner and the flow have rotational symmetry around its axis, but the burner head has divisions which doesn't make it axisymmetric. However, since the divisions are small in size, its effect is considered not significant and, therefore, a 2D axisymmetric burner model is developed. To observe the burner behavior, several simulations in two dimensions were performed by modifying the opening of primary and secondary air regulators, as shown in Table 1.

**Table 1.** Combination of the openings of regulators in different simulations. To observe the effect that the divisions located in the burner head have over the flow, we also made a 3D simulation of the burner using as input conditions the results of 2D simulations.

Simulation	Primary regulator opening (mm)	Secondary regulator opening (mm)
S1	5	0
S2	15	0
S3	30	0
S4	50	0
S5	5	20
S6	15	20
S7	30	20
S8	50	20
S9	15	10

### Operating Conditions

In Table 2, the burner operating conditions are shown, which are useful to determine boundary conditions at the simulations.

All 2D-axisymmetric simulations have a maximum Mach number around 1,19, which is obtained when the fuel is discharged from the nozzle into the mixer, imply-

ing that the flow in this zone is supersonic. To suppose that a gas flow is compressible, we use a criterion where the Mach number should be higher than 0,1, because below this value the density variation in function of pressure can be ignored. In these simulations, the flow must be considered compressible.

**Table 2.** Burner nominal operating conditions, which are used to set up the boundary conditions in simulations.

Fuel	Methane
Oxidier	Air
Thermal power	180 kW
Operation pressure	144,8 kPag
Atmospheric pressure	1 atm
Average air temperature	27° C

### Governing Equations

Equations for a compressible, turbulent and reactive flow in 2D axisymmetric are shown below. The flow is considered compressible because the methane is discharged in the nozzle at supersonic speed, generating a large pressure gradient in this area. The equations governing the problem are the complete Navier Stokes equations, i.e. augmented by the laws of chemical kinetics, measured with the *Favre* average, which takes into account the density fluctuations. For a  $\theta$  property, *Favre* average is defined as shown below.

$$\tilde{\theta} = \frac{\overline{\rho\theta}}{\bar{\theta}} \tag{1}$$

In this equation, the horizontal bar on the variables refers to the time average of these variables, which is shown in *Equation 2*.

$$\bar{\theta} \equiv \frac{1}{\Delta t} \int_{t_1}^{t_2} \theta(t) dt \tag{2}$$

### Mass Conservation

$$\frac{\partial(\bar{\rho}\tilde{u}_x)}{\partial x} + \frac{1}{r} \frac{\partial(r\bar{\rho}\tilde{u}_r)}{\partial r} = 0 \tag{3}$$

## Energy Conservation

$$\frac{\partial(\bar{\rho}\tilde{u}_x\tilde{h})}{\partial x} + \frac{1}{r}\frac{\partial(r\bar{\rho}\tilde{u}_r\tilde{h})}{\partial r} = \frac{\partial}{\partial x}\left(\frac{\mu_e}{S_{c,t}}\frac{\partial\tilde{h}}{\partial x}\right) + \frac{1}{r}\frac{\partial}{\partial r}\left(r\frac{\mu_e}{S_{c,t}}\frac{\partial\tilde{h}}{\partial r}\right) + S_h \quad (4)$$

$$\mu_e = \mu + \mu_t$$

The turbulent viscosity is related to the turbulent kinetic energy ( $k$ ) and the turbulent kinetic energy dissipation rate ( $\varepsilon$ ) as follows:

$$\mu_t = \rho C_\mu \frac{k^2}{\varepsilon} \quad (5)$$

$$C_\mu = 0,09$$

 Momentum Conservation in the Direction  $r$ 

$$\frac{\partial(\bar{\rho}\tilde{u}_x\tilde{u}_r)}{\partial x} + \frac{1}{r}\frac{\partial(r\bar{\rho}\tilde{u}_r\tilde{u}_r)}{\partial r} \quad (6)$$

$$= \frac{\partial}{\partial x}\left(\mu_e\frac{\partial\tilde{u}_r}{\partial x}\right) + \frac{1}{r}\frac{\partial}{\partial r}\left(r\mu_e\frac{\partial\tilde{u}_r}{\partial r}\right)$$

$$- \frac{\partial p}{\partial r} - 2\mu_e\frac{\tilde{u}_r}{r^2} + \frac{\partial}{\partial x}\left(\mu_e\frac{\partial\tilde{u}_x}{\partial r}\right)$$

$$+ \frac{1}{r}\frac{\partial}{\partial r}\left(r\mu_e\frac{\partial\tilde{u}_r}{\partial r}\right) - \frac{2}{3}\frac{\partial}{\partial r}$$

$$\left[\mu_e\left(\frac{1}{r}\frac{\partial(r\tilde{u}_r)}{\partial r} + \frac{\partial\tilde{u}_x}{\partial x}\right) + \bar{\rho}k\right]$$

 Momentum Conservation in the Direction  $x$ 

$$\frac{\partial(\bar{\rho}\tilde{u}_x\tilde{u}_x)}{\partial x} + \frac{1}{r}\frac{\partial(r\bar{\rho}\tilde{u}_r\tilde{u}_x)}{\partial r} \quad (7)$$

$$= \frac{\partial}{\partial x}\left(\mu_e\frac{\partial\tilde{u}_x}{\partial x}\right) + \frac{1}{r}\frac{\partial}{\partial r}\left(r\mu_e\frac{\partial\tilde{u}_x}{\partial r}\right)$$

$$- \frac{\partial p}{\partial x} + \frac{\partial}{\partial x}\left(\mu_e\frac{\partial\tilde{u}_x}{\partial x}\right) + \frac{1}{r}\frac{\partial}{\partial r}\left(r\mu_e\frac{\partial\tilde{u}_r}{\partial x}\right)$$

$$- \frac{2}{3}\frac{\partial}{\partial x}\left[\mu_e\left(\frac{1}{r}\frac{\partial(r\tilde{u}_r)}{\partial r} + \frac{\partial\tilde{u}_x}{\partial x}\right) + \bar{\rho}k\right]$$

## Chemical Species Conservation

$$\frac{\partial(\bar{\rho}\tilde{u}_x\tilde{Y}_i)}{\partial x} + \frac{1}{r}\frac{\partial(r\bar{\rho}\tilde{u}_r\tilde{Y}_i)}{\partial r} = -\frac{\partial}{\partial x}\left(\frac{\mu_e}{S_{c,t}}\frac{\partial\tilde{Y}_i}{\partial x}\right) + \frac{1}{r}\frac{\partial}{\partial r}\left(r\frac{\mu_e}{S_{c,t}}\frac{\partial\tilde{Y}_i}{\partial r}\right) - R_i \quad (8)$$

$R_i$  is the production rate of specie  $i$ , which is calculated using the Finite rate/Eddy dissipation model which computes the Arrhenius reaction rate and a reaction rate based on the work of Magnussen and Hjertager (1976), taking as net reaction rate the smaller of both.

The production rate of specie  $i$ , is given by:

$$R_i = M_{w,i} \sum_{r=1}^M v_{ir} Q_r \quad (9)$$

$$Q_r = k_{f,r} \prod_{i=1}^N [X_i]^{v'_{ir}} - k_{b,r} \prod_{i=1}^N [X_i]^{v''_{ir}} \quad (10)$$

$k_{f,r}$  is the forward rate constant for reaction  $r$  that is calculated using the Arrhenius expression:

$$k_{f,r} = A_r T^{\beta_r} e^{-E_r/RT} \quad (11)$$

The rate of production of the chemical specie  $i$  calculated with the eddy dissipation model is:

$$R_i = \min(R_{i,r}^R, R_{i,r}^P)$$

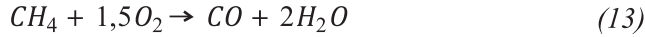
$$R_{i,r}^R = v'_{i,r} M_{w,i} A \rho \left(\frac{\varepsilon}{k}\right) \min_R \left(\frac{Y_R}{v'_{R,r} M_{w,R}}\right) \quad (12)$$

$$R_{i,r}^P = v'_{i,r} M_{w,i} A B \rho \left(\frac{\varepsilon}{k}\right) \left(\frac{\sum_P Y_P}{\sum_j^N v'_{j,r} M_{w,j}}\right)$$

Although multi-step reaction mechanisms may be used with the finite rate/eddy dissipation model, these likely produce incorrect solutions. This is due to multi-step chemical mechanisms based on Arrhenius rates, which differ for each reaction. In the eddy-dissipation model, every reaction has the same turbulent rate; therefore the model should be used only for one-step, or two-step global reactions (Fluent user's guide, 2006).

**Reaction mechanism**

It is used as a methane global reaction mechanism (Westbrook & Dryer, 1981) with two irreversible steps which are shown below.



**Radiation Model**

To calculate the source term in the energy equation ( $S_{ii}$ ) we use the *PI* model (Siegel & Howell, 2002) because of its simplicity and good results obtained by:

$$\begin{aligned} -\nabla \tilde{q} &= aG - 4a \sigma \tilde{T}^4 \\ \tilde{q} &= -\Gamma \nabla G \\ \Gamma &= \frac{1}{3a} \end{aligned} \quad (15)$$

The absorption coefficient  $a$  is considered variable and it is calculated using a weighted sum of gray gases model (WSGGM).

***k-ε Standard Turbulence Model***

Since the turbulence viscosity (*Equation 5*) is in terms of two unknown variables, which are the turbulent kinetic energy and the turbulent kinetic energy dissipation rate, it requires two additional transport equations to find these two variables. The standard *k-ε* model developed by Launder and Spalding (1972) is used. This is a semi-empirical model based on model transport equations for the turbulence kinetic energy ( $k$ ) and its dissipation rate ( $\epsilon$ ). The model transport equation for  $k$  is derived from the exact equation, while the model transport equation for  $\epsilon$  was obtained using physical reasoning and shows some resemblance to its mathematically exact counterpart.

In the derivation of the standard *k-ε* model, the assumption is that the flow is fully turbulent, and the effects of molecular viscosity are negligible. Therefore this model is valid only for fully turbulent flows (Fluent user's guide, 2006).

Turbulent kinetic energy transport equation ( $k$ )

$$\begin{aligned} \frac{\partial(\bar{\rho}\tilde{u}_x k)}{\partial x} + \frac{1}{r} \frac{\partial(r\bar{\rho}\tilde{u}_r k)}{\partial r} & \quad (16) \\ &= \frac{\partial}{\partial x} \left( \frac{\mu_e}{\sigma_k} \frac{\partial k}{\partial x} \right) + \frac{1}{r} \frac{\partial}{\partial r} \left( r \frac{\mu_e}{\sigma_k} \frac{\partial k}{\partial r} \right) \\ &+ \mu_t \left( \frac{\partial \tilde{u}_x}{\partial r} + \frac{\partial \tilde{u}_r}{\partial x} \right)^2 \\ &+ 2\mu_t \left[ \left( \frac{\partial \tilde{u}_x}{\partial x} \right)^2 + \left( \frac{\partial \tilde{u}_r}{\partial r} \right)^2 + \left( \frac{\tilde{u}_r}{r} \right)^2 \right] \\ &- \frac{2}{3} \left[ \mu_e \left( \frac{1}{r} \frac{\partial(r\tilde{u}_r)}{\partial r} + \frac{\partial \tilde{u}_x}{\partial x} \right)^2 \right. \\ &\left. + \bar{\rho} k \left( \frac{1}{r} \frac{\partial(r\tilde{u}_r)}{\partial r} + \frac{\partial \tilde{u}_x}{\partial x} \right) \right] - \bar{\rho} \epsilon - \sqrt{\frac{k}{c^2}} \end{aligned}$$

Turbulent kinetic energy dissipation rate transport equation ( $\epsilon$ ):

$$\begin{aligned} \frac{\partial(\bar{\rho}\tilde{u}_x \epsilon)}{\partial x} + \frac{1}{r} \frac{\partial(r\bar{\rho}\tilde{u}_r \epsilon)}{\partial r} & \quad (17) \\ &= \frac{\partial}{\partial x} \left( \frac{\mu_e}{\sigma_\epsilon} \frac{\partial \epsilon}{\partial x} \right) + \frac{1}{r} \frac{\partial}{\partial r} \left( r \frac{\mu_e}{\sigma_\epsilon} \frac{\partial \epsilon}{\partial r} \right) \\ &+ \frac{\epsilon}{k} \left\{ C_1 \left[ \mu_t \left( \frac{\partial \tilde{u}_x}{\partial r} + \frac{\partial \tilde{u}_r}{\partial x} \right)^2 \right. \right. \\ &+ 2\mu_t \left( \left( \frac{\partial \tilde{u}_x}{\partial x} \right)^2 + \left( \frac{\partial \tilde{u}_r}{\partial r} \right)^2 \right. \\ &\left. \left. + \left( \frac{\tilde{u}_r}{r} \right)^2 \right) \right] - C_2 \rho \epsilon \left. \right\} \end{aligned}$$

The symbols  $C_1, C_2, \sigma_k, \sigma_\epsilon$  are model constants and in this case the Fluent default values are used, which are recommended in the work of Launder and Sharma (1974).

$$C_1 = 1,44, C_2 = 1,92, \sigma_k = 1,0, \sigma_\epsilon = 1,3$$

These values have been determined from experiments with air and water for fundamental turbulent shear flows including homogeneous shear flows and decaying isotropic grid turbulence. They have been found to work fairly well for a wide range of wall bounded and free shear flows.

### Numerical Method

To discretize all the variables, upwind second order schemes have been used, except for the pressure where the PRESTO scheme (Patankar, 1980) was used and momentum where upwind first order scheme was used. To consider the gas compressibility, the ideal gas law was used.

All 2D simulations have the same shape of the mesh, differing only in the openings of the regulators of primary and secondary air as shown in Table 1. The meshes have approximately the same amount of cells (46 000 after two mesh adaptations). The areas needing increased mesh density are the mixer and the furnace dome. This happens because they have the highest gradients of speed, due to gas discharge, and of temperature, due to flame. Regarding the quality of the mesh, the higher aspect ratio is 7,6 and the equi angle skewness does not exceed 0,6. Figure 2 shows a typical mesh used in 2D axisymmetric simulations.

To determine if a simulation has converged, three different criterions have been used: first, the verification that residuals of the iterations are asymptotic. Then, a computed energy balance between inputs and outputs, which should be less than 5% of burner power. Finally, an average temperature and velocity in a line located in reaction zone, unchangeable despite iterations.

Three mesh adaptations have been done to each single simulation in order to outline the quantity of meshes necessary to obtain a mesh-independent solution. The adaptations were performed based on temperature and velocity gradients, determining that after around 45 000 cells, the solution is mesh independent in all simulations. This was concluded by comparing temperature profiles in a line located in the reaction zone with different cell quantities, and observing that there were no differences in these profiles, between successive cell quantities. Also, due to the operating conditions, the burner has a very turbulent flow. Therefore, in order to obtain good results with a standard  $k-\varepsilon$  model, the cells near the wall have been adapted to obtain a Y-plus ( $Y^+$ ) around 50 or less in all simulations.

Since the burner is a premixed burner, to ensure that the flame is generated in the burner head discharge and not inside the mixer, it is necessary to apply a high temperature (3 000K) region in the dome, before the iterations begin, which works like a spark plug to start chemical reactions in a desired region, consequently forming the flame.

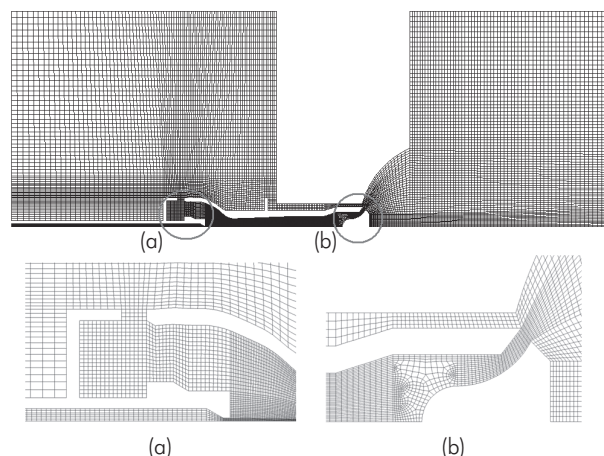


Figure 2. Typical 2D Domain mesh, where all cells are cuadrilaterals and finer mesh zones are inside the mixer.

### Boundary Conditions

Figure 3 shows the boundary conditions used in all 2D-axisymmetric simulations. All pressure inlet conditions have a gauge pressure equal to zero, where air is used as fluid entering on these boundary conditions. As a result, the burner is simulated as discharging into the free atmosphere and not into a furnace. The walls are supposed to be adiabatic, so there is no heat transfer through these. Finally, the boundary conditions called interior conditions are used to take the profiles of velocity, temperature, turbulent kinetic energy, turbulent kinetic energy dissipation rate and methane, oxygen and nitrogen mass fraction. These are used as input conditions in the 3D simulation, which is done through a defined function by the Fluent user.

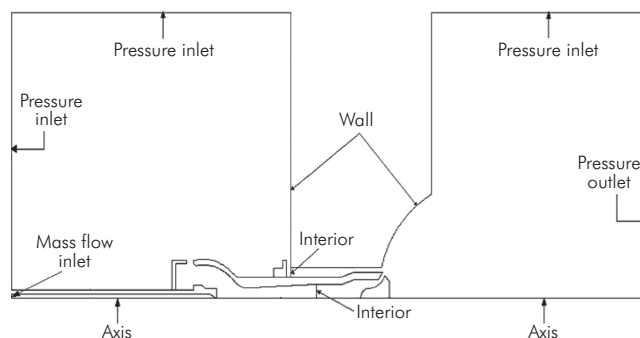


Figure 3. Boundary conditions used in all 2D axisymmetric simulations.

## 3. 2D-AXISYMMETRIC RESULTS

The results presented here correspond to those obtained in 2D simulations with different primary and secondary air regulator openings.

**Burner Flow**

Figure 4 shows the flow in the S8 burner simulation, observing the way it enters the primary and secondary air by the respective regulators.

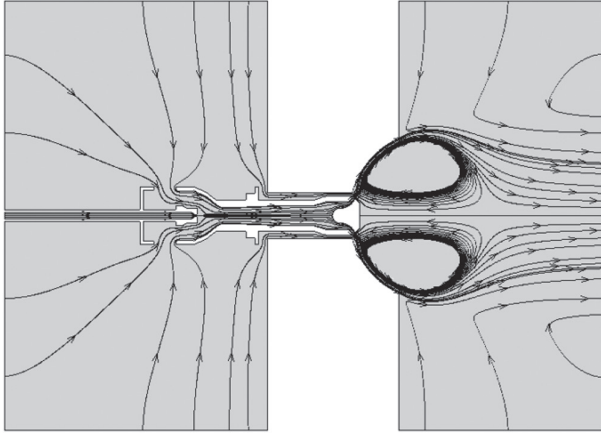


Figure 4. Streamlines of simulation S8, where is shown the strong recirculation zone generated by the furnace dome.

It also highlights the strong recirculation zone that is generated due to the dome in the furnace wall, which helps to provide stability of the flame, shortening it and preheating the air fuel premix.

**Temperature Contours**

Chemical reactions are produced between the exit of the burner head and the dome carved into the furnace wall, whose spherical shape contributes to stability of the flame and recirculation of the combustion products. In this way, the combustion efficiency is improved by preheating the premix discharged.

Figure 5 shows the temperature contours obtained when the primary air regulator is only opened. Figure 6 shows the temperature contours when the secondary air regulator is opened in 20 mm. In general, when the secondary air regulator is closed, the temperatures are higher and the flame tends to be more extended because there is no enough air to completely burn the fuel.

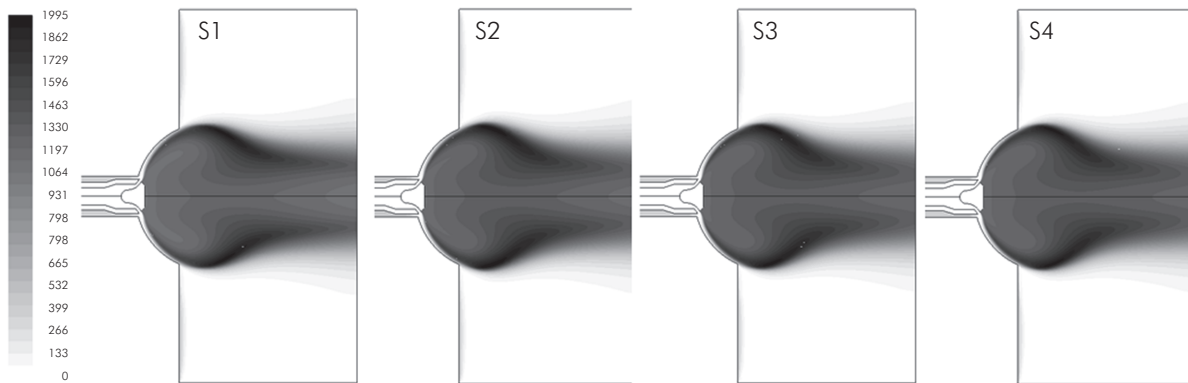


Figure 5. Contours of temperature (°C) with secondary air regulator closed, which correspond to simulations S1, S2, S3 and S4. And also is shown the equivalence factor obtained from each simulation.

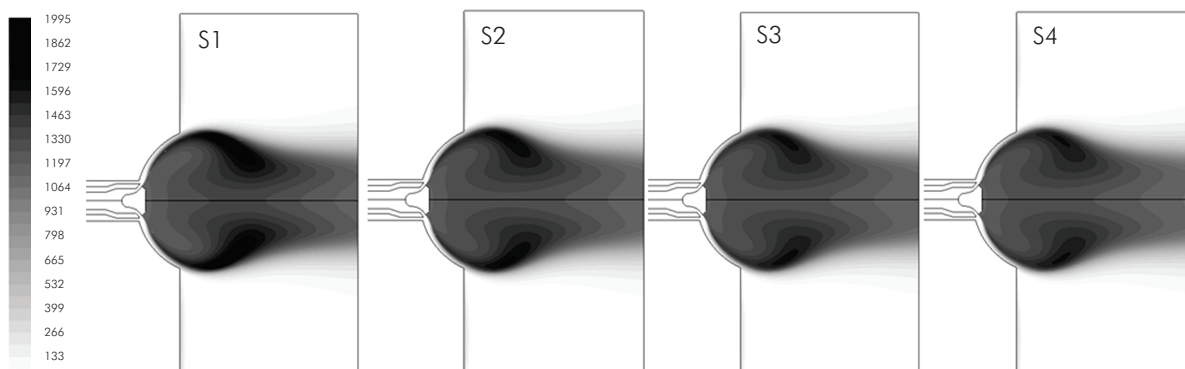


Figure 6. Contours of temperature (°C) with secondary air regulator open at 20 mm, which correspond to simulations S5, S6, S7 and S8. And also is shown the equivalence factor obtained from each simulation.



Figure 7 shows the maximum temperatures obtained in the simulations S1 to S9, observing that when the secondary air regulator is closed, the maximum temperature remains almost constant and close to the adiabatic flame value for methane. This occurs because the equivalence ratio always remains above 1 for all these simulations. Whereas when the secondary air regulator is opened in 20 mm, the maximum temperature tends to decrease as the opening of the primary air regulator increases. This happens because the amount of air is greater than the stoichiometric (except simulation S5), which has the effect of decreasing the temperature. This happens because part of the fuel energy is used to heat not reacting air.

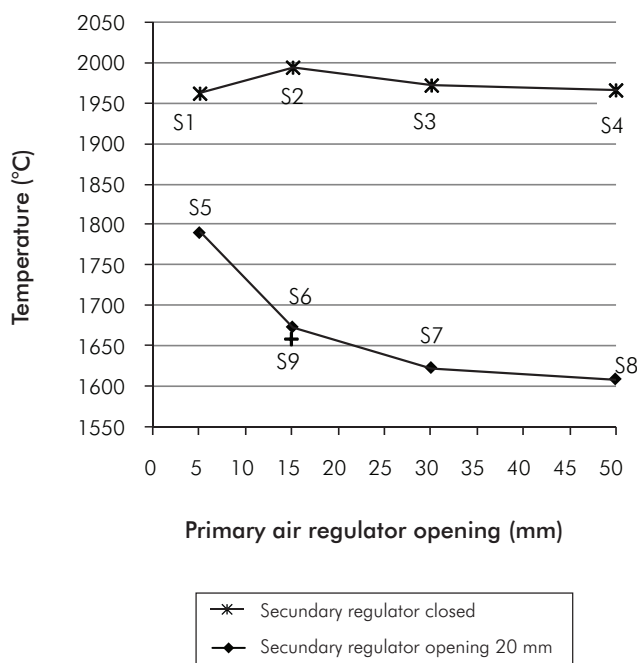


Figure 7. Maximum temperature obtained for all simulations, where there are 2 series corresponding to secondary regulator closed and secondary regulator open at 20 mm.

Another important result that is derived from temperature contours is related with flame length, which in this work is defined as the maximum x-distance where some temperature contour is equal to 1500°C. The results are presented in Table 3 for different simulations in a percent value, which is the relation respect to the distance of ethane pipes inside the furnace. This is useful because it allows seeing if the flame reaches the pipes, which in turn will reduce their lifespan. If this happens, the possibility of changing the burner design should be considered.

Table 3. Flame length (in percentage of distance to ethane pipes) obtained for simulations S1 to S8.

Simulation	Flame length (%)	Simulation	Flame length (%)
S1	82,9	S5	60,7
S2	81,0	S6	48,3
S3	79,0	S7	46,4
S4	79,0	S8	45,1

### CO Contours

In Figure 8 and 9, the CO contours for simulations are presented. These have the secondary air regulator closed and opened in 20 mm respectively, where the amount of CO produced is quite large compared to when the secondary air regulator is opened. This happens because regardless of the openings of the primary air regulator, the air factor does not exceed the stoichiometric value. Therefore, there is not enough air to complete methane combustion.(Figure 10).

### Equivalence Ratio

The equivalence ratio ( $\phi$ ) is defined as the mass ratio of fuel-air ratio present during combustion divided by stoichiometric fuel-air ratio.

$$\phi = \frac{(F/A)}{(F/A)_{stoicsh}} \tag{18}$$

Where  $F$  is the mass of fuel,  $A$  is the mass of air and the suffix stoich stands for stoichiometric conditions.

Figure 10 shows the results obtained for the equivalence ratio with the secondary air regulator closed and opened in 20 m. It is observed that when the secondary air regulator is closed, the equivalence ratio is never lower than the unit, and consequently there is air absence. If air is not added later, combustion remains incomplete. On the other hand, when the secondary air regulator is opened in 20 mm, and the primary air regulator has an opening larger than 15 mm, the equivalence ratio is lower than unity, reaching values of around 9% in air excess. Therefore, the amount of air amount is sufficient to complete combustion.

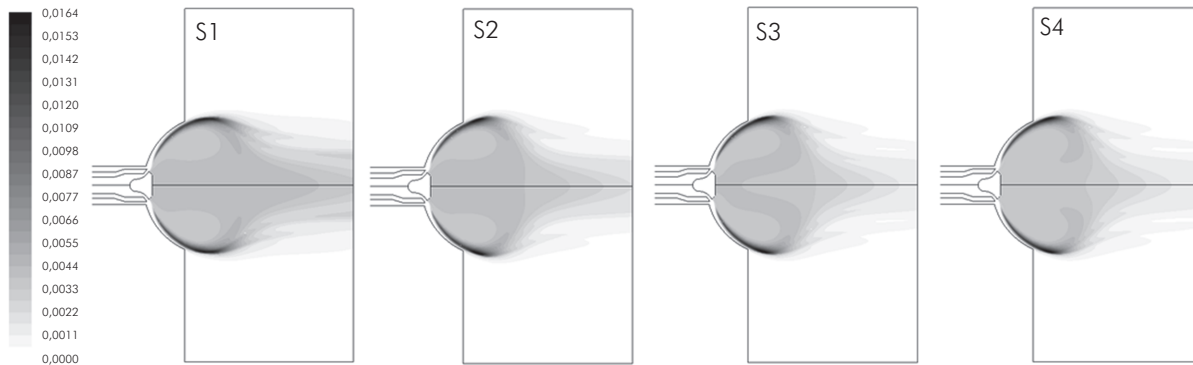


Figure 8. Contours of CO mass fraction with the secondary air regulator closed, which correspond to simulations S1, S2, S3 and S4.

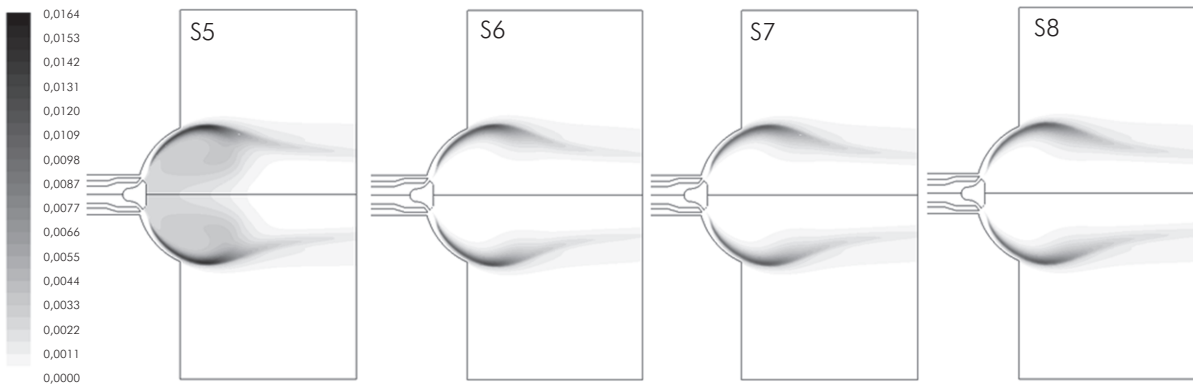


Figure 9. Contours of CO mass fraction with the secondary air regulator open at 20 mm, which correspond to simulations S5, S6, S7 and S8.

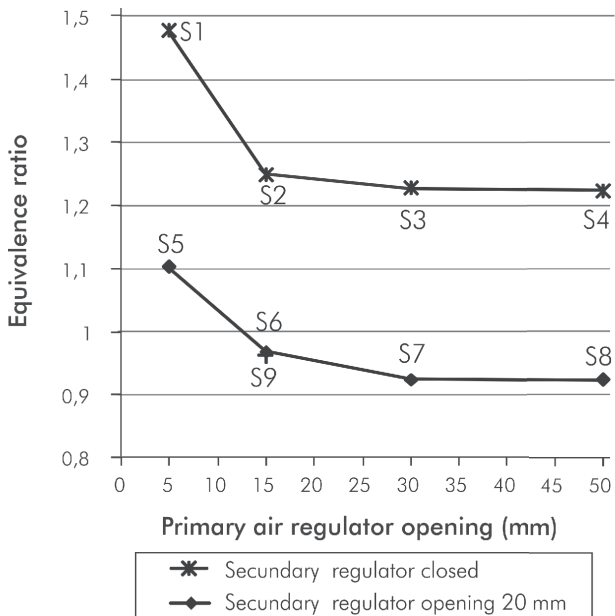


Figure 10. Equivalence ratio obtained for all simulations, where there are 2 series corresponding to secondary regulator closed and secondary regulator open at 20 mm. And also is shown the result obtained for simulation S9, which has an opening of 15 mm in primary regulator and 10 mm in secondary regulator.

#### 4. 3D SIMULATION

Given that in the 2D simulations the effect that divisions of the burner head have on the shape and characteristics of the flame has been neglected, we decided to take the profiles of some flow properties of a representative 2D simulation to serve as entry conditions into a 3D simulation. In this scenario, the flow from the burner head with the formation of the flame is simulated, in order to observe the differences with the corresponding 2D simulation.

To perform the 3D simulation, the following is used as input conditions: velocity profiles, turbulent kinetic energy ( $k$ ), turbulent kinetic energy dissipation rate ( $\epsilon$ ), and density and concentration species ( $O_2$ ,  $CH_4$ ) extracted from the S9 simulation. To use the profiles obtained from the 2D axisymmetric simulation as input conditions into the 3D simulation, a small user defined function (UDF) has been programmed in Fluent.

### 3D Mesh

The creation of the 3D mesh starts from the burner head. This mesh corresponds only to one quarter of the real burner due to symmetry, and includes the burner head, the furnace dome where the flame is formed and a part of discharge zone that is made of cylindrical shape (Figure11).

The original mesh has approximately 39 000 hexahedral cells, with an aspect ratio of 7,72 and an equi angle skewness maximum of 0,52. When performing the adaptation process in FLUENT, the mesh reaches -in the final simulation- a total of about 13 0000 cells.

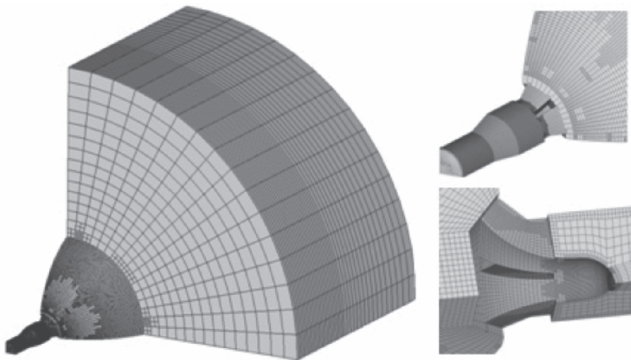


Figure 11. 3D burner mesh, where all cells are hexagonal and structured.

### 3D Results

Figure 12 is a comparison between the flame shape in a 2D simulation and a plane in a 3D simulation passing through the middle of a gas discharge channel. It can be observed that the basic shape of the flame and temperature profiles are preserved, but the flame in the 3D simulation is more robust than the corresponding 2D flame, being the latter more flattened and elongated. This behavior was expected because the output section of the burner head was worked out in detail in the 3D simulation, obtaining more compact jets with greater penetrating power of the reactive mixture air-methane in the internal environment of the furnace.

To observe more clearly the flame shape, Figure 13 shows two iso surface temperature of 1 000°C and 1 300°C, on which the profiles of CO mass fraction are plotted. One of the main effects that the divisions located in the burner head on the flame shape have, is to deform the profiles, making them not completely axisymmetric, differing mainly from the results obtained in 2D simulation.

In conclusion, the velocity, temperature and species profiles obtained with 2D and 3D simulations have differences in shape. Nevertheless, for practical aspects, the 2D simulation results show us a pretty good approximation of the velocity, temperature and chemical species concentration ( $O_2$ ,  $CH_4$ ,  $CO$ ,  $CO_2$ ) taking about 5 times less in computation time.

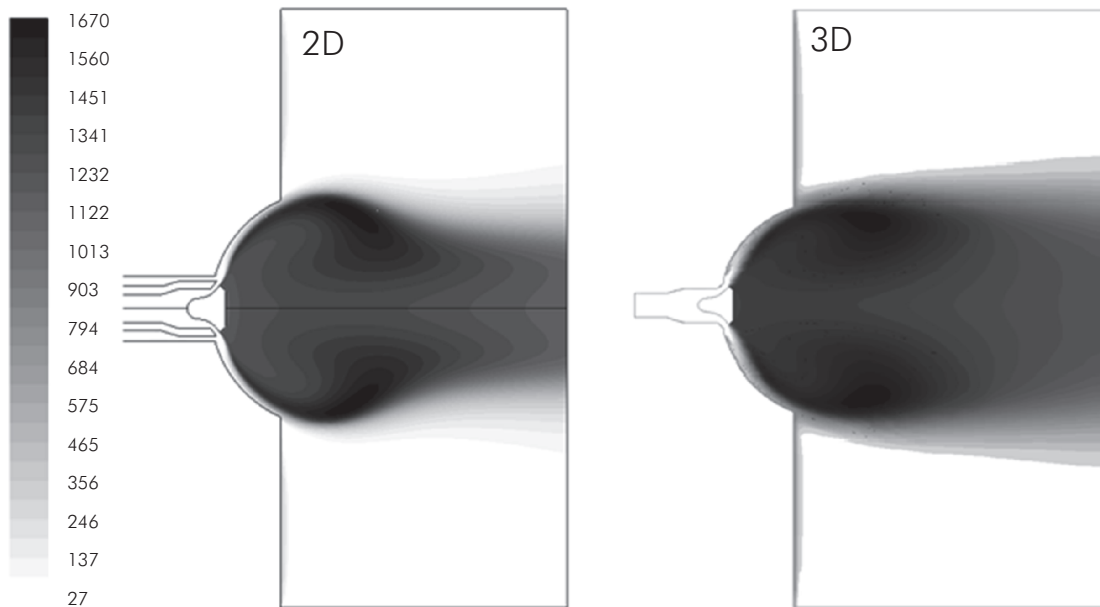
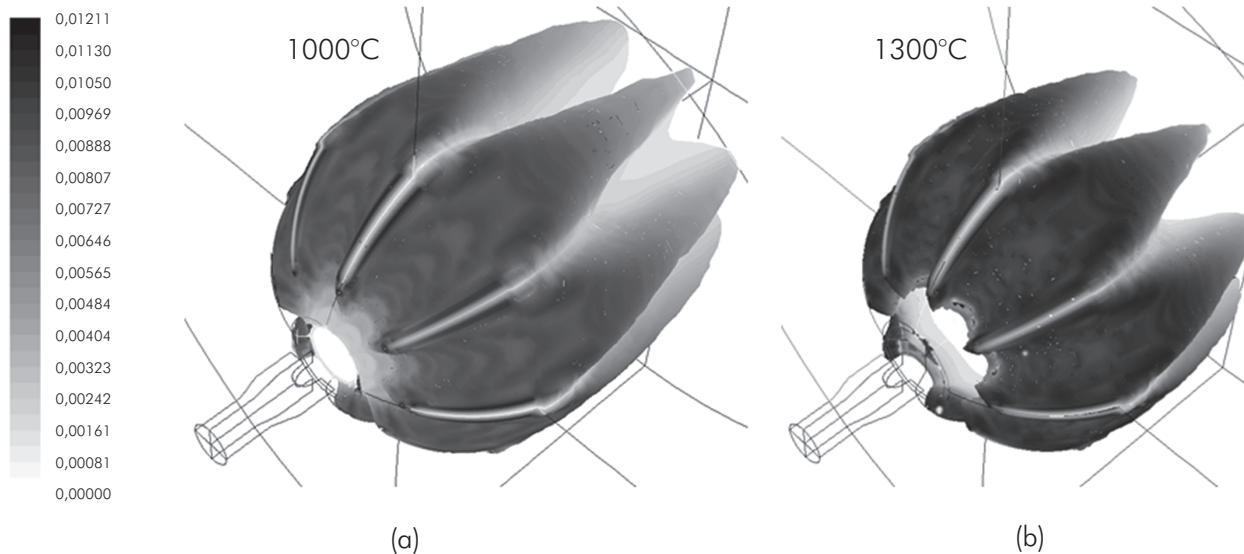


Figure 12. Comparison of temperature contours (°C) between a plane in the 3D simulation and its analogous 2D simulation (S9).



**Figure 13.** CO contours on a iso-surface of 1 000°C (a) and 1 300°C (b), which show in a better way the shape of the flame.

## 5. CONCLUSIONS

In this simulation work, the following aspects were concluded:

- Regarding the opening of the primary air regulator, it has no impact on equivalence ratio from around 30 mm, no matter the position of the secondary air regulator.
- When there is no secondary air entry, regardless of the primary air regulator opening, combustion air will always be lacking, indicating that the secondary air regulator plays an important role by providing remaining air to complete combustion.
- It is possible to obtain equivalence ratio lower than 1 (air excess), having the primary air regulator opened in values higher than 15 mm and maintaining the secondary air regulator open.
- The closure of the secondary air regulator has the following consequences: increased production of  $CO$ , being always greater than the amount produced with the secondary regulator open, causing the combustion

to be more inefficient; increasing flame temperature, since there is a lack of air and higher temperatures for methane are achieved with the absence of air near the stoichiometric (equivalence ratio greater than 1). Due to the absence of air in the combustion, the flame tends to be a bit more elongated than with the secondary air regulator opened, becoming more a diffusion flame.

- Although the shape of the flame differs from the 2D simulation respect to the 3D simulation, the latter takes into account the interaction of flow with the divisions in the burner head. To establish operating parameters, the 2D study is sufficient, since the contours of temperature, speed and other properties in the 3D simulation are similar with the results in 2D, being rather useful 3D simulation to observe more clearly the shape of the flame.

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**NOTATION**

$A$	Empirical constant, equal to 0,4	-
$A_r$	Pre-exponential factor (consistent units)	-
$a$	Absorption coefficient	-
$B$	Empirical constant, equal to 0,5	-
$c$	Sound speed	$\text{m}\cdot\text{s}^{-1}$
$G$	Incident radiation	$\text{W}\cdot\text{m}^{-2}$
$h$	Enthalpy	$\text{J}\cdot\text{kg}^{-1}$
$k$	Turbulent kinetic energy	$\text{m}^2\cdot\text{s}^{-2}$
$k_{f,r}$	Forward rate constant for reaction $r$	$\text{mol}\cdot\text{s}^{-1}$
$k_{b,r}$	Backward rate constant for reaction $r$	$\text{mol}\cdot\text{s}^{-1}$
$M_{w,i}$	Molecular weight of species $i$	$\text{kg}\cdot\text{kmol}^{-1}$
$M_{w,R}$	Molecular weight of a particular specie $R$	$\text{kg}\cdot\text{kmol}^{-1}$
$M_{w,j}$	Molecular weight of reactant species $j$ in reaction $r$	$\text{kg}\cdot\text{kmol}^{-1}$
$N$	Number of chemical species in the system	-
$p$	Pressure	Pa
$Q_r$	Progress rate of $r$ reaction	$\text{mol}\cdot\text{s}^{-1}$
$q$	Heat flux	$\text{J}\cdot\text{m}^{-2}\cdot\text{s}^{-1}$
$R_i$	Net source of chemical species $i$	$\text{kg}\cdot\text{s}^{-1}$
$R_{i,r}$	Arrhenius molar rate of creation/destruction of species $i$	$\text{kg}\cdot\text{s}^{-1}$
$r$	Radial direction	m
$S_{c,t}$	Turbulent Schmidt number	-
$S_h$	Source term in energy equation	-
$T$	Temperature	K
$t$	Time	s
$u_x$	Axial velocity	$\text{ms}^{-1}$
$u_r$	Radial velocity	$\text{ms}^{-1}$
$u_\tau$	Friction velocity	$\text{m}^2\cdot\text{s}^{-1}$
$X_i$	Molar fraction of species $i$	-
$x$	Axial direction	$m$

## NOTATION

$Y_i$	Mass fraction of species $i$	-
$Y^+$	$Y$ -plus, $Y^+ = \rho u_r y / \mu$	-
$\varepsilon$	Turbulent kinetic energy dissipation rate	$\text{m}^2 \cdot \text{s}^{-3}$
$\theta$	Any flow property	-
$\nu_{i,r}$	Stoichiometric coefficient of species $i$ in reaction $r$	-
$\nu'_{i,r}$	Stoichiometric coefficient for reactant $i$ in reaction $r$	-
$\nu''_{i,r}$	Stoichiometric coefficient for product $i$ in reaction $r$	$\text{kg} \cdot \text{m}^{-1} \cdot \text{s}^{-1}$
$\mu$	Molecular viscosity	$\text{kg} \cdot \text{m}^{-1} \cdot \text{s}^{-1}$
$\mu_e$	Effective viscosity	$\text{kg} \cdot \text{m}^{-1} \cdot \text{s}^{-1}$
$\mu_t$	Turbulent viscosity	$\text{kg} \cdot \text{m}^{-3}$
$\rho$	Density	$\text{W} \cdot \text{m}^{-2} \cdot \text{K}^{-4}$
$\sigma$	Stefan-Boltzmann constant ( $5,67 \times 10^{-8}$ )	-
$\phi$	Equivalence factor	-
$C_\mu$	Standard $k$ - $\varepsilon$ turbulence model constant	-
$\beta_r$	Temperature exponent in Arrhenius equation for reaction $r$	-
$E_r$	Activation energy	$\text{j/mol}$
$Y_R$	Mass fraction of a particular reactant $R$	-
$Y_P$	Mass fraction of any product species $P$	-

