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Direct Numerical Simulation of combustion near a carbonaceous surface in a quiescent flow

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Abstract

In this work, unsteady gasification and oxidation of carbonaceous materials with gas-phase combustion under oxy-combustion conditions is studied. An original analytical solution is derived to validate the numerical species boundary condition for the heterogeneous surface reactions with frozen gas-phase. Direct Numerical Simulations accounting for multi-species diffusion and detailed chemistry with both species and energy boundary conditions taking into account the Stefan flux were subsequently performed to study combustion near a carbonaceous wall. Transition from gasification to solid carbon oxidation and gas-phase ignition is evidenced. The competition between homogeneous and heterogeneous reactions is discussed based on the analysis of the time-dependent surface and gas-phase species consumption and production rates. Transient histories of surface species as well as spatial profiles of gas-phase species, heat release and temperature are presented for two identified structures of gas-phase CO/O\(_2\) reaction zone. In order to quantify the error induced by the Hirschfelder & Curtiss approximation for the multi-species diffusion and to evaluate the importance of the Dufour and Soret terms, a library dedicated to the computation of complex diffusion phenomena has been used to rebuild a priori, species and Soret fluxes for mass diffusion as well as Fourier, species and Dufour diffusion heat fluxes. Eventually, characteristic times of gas and surface reactions and those corresponding to the switch between the two structures of the gas-phase reaction zone are obtained over a wide range of initial O\(_2\)/CO\(_2\) concentrations.

Keywords:
Gasification, Oxidation, Ignition, DNS, Reactive surface

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1. Introduction

Oxy-fuel combustion is a cost-effective technology for $CO_2$ capture from fuel fired furnaces. To control operating temperatures, flue gas is typically recycled into the combustion chamber, forming $O_2/CO_2$ atmosphere. Solid carbon ignition in such an atmosphere represents an issue not only for combustion technology but also for safety [1]. Despite the lower oxygen diffusivity in $CO_2$ and the endothermicity of $CO_2$ gasification (carbon conversion into carbon monoxide), the latter can actually lead to the enhancement of the overall carbon burning rate in oxy-fuel combustion [2, 3]. Under steady-state conditions the effect of $CO_2$ gasification on carbon conversion rate has been found to depend on $O_2$ concentration [4]. The interaction between $CO_2$ gasification and oxidation have been extensively studied [5, 6] especially at high pressure and high temperature [7] but remains poorly understood. In $O_2$-enriched conditions, gasification can contribute significantly to carbon consumption [8]. Although the sequential occurrence [9] or respective importance [10] of homogeneous and heterogeneous ignition mechanisms have been evidenced experimentally, no detailed modeling work focused specifically, to our knowledge, on the interplay between $CO$ production and consumption pathways under vitiated oxyfuel combustion. The contribution of heterogeneous and homogeneous reactions to carbon ignition is known to depend on the gas-phase composition [7] as well as on carbon particle size [11] and surface temperature [12, 13]. Makino et al. [14] studied the temporal evolution of oxidation and gasification contributions using global kinetics and transport and formulated a criterion for the existence of a $CO$ flame. Zou et al [15] addressed the homogeneous and heterogeneous ignition processes of pulverized coal in oxy-fuel combustion using an Eulerian-Lagrangian approach with devolatilization kinetic sub-models. The impact of volatiles combustion processes, ambient gas temperatures, coal particle size and $O_2$ concentrations on the ignition type were assessed. It was concluded that heterogeneous ignition is more likely to occur for small coal particles and high $O_2$ concentrations whereas homogeneous ignition launched by initial volatiles ignition is more likely to occur for large particles.

Many research groups still use simplified film models based on averaged diffusivity and global kinetics and neglect $CO$ gas-phase conversion in the boundary layer. However, the predictivity of simplified film models need to be further improved [16, 17]. Lewtak et al. [18] showed that the use of an equimolar counterdiffusion model leads to the overprediction of mass transfer towards the particle. A recent review [7] stressed the need to account for multi-component diffusion effect in carbon oxy-fuel combustion. The sensitivity of carbon consumption to Stefan flow was shown to depend on the relative contributions of gasification and oxidation reactions. Stefan velocity can reach significant values when the surface coverages
undergo drastic change during transient processes [19] which may give rise to gas-phase ignition [20]. Thus it is important to take into account this additional velocity induced by reactive walls.

Theoretical studies have been carried out using global chemistry, highlighting the differences between transient and steady-state behaviour [21], the impact of the relative importance of surface reactions on the shape of burning rate curve and the respective importance of homogeneous/heterogeneous chemistry and transport phenomena on particle ignition [22–24].

The aforementioned works reveal that transient homogeneous/heterogeneous carbon combustion data accounting for multi-species diffusion, detailed surface and gas-phase schemes with proper species and energy boundary conditions taking into account the Stefan flux remain scarce. Few studies emphasize the transient history of the heterogeneous/homogeneous carbon combustion. Hence, the objectives of this paper are twofold: An original analytical solution is first derived to develop and validate the numerical species boundary condition for the heterogenous surface reactions. Second, appropriate boundary conditions consistent with the conservation of gaseous/surface species and energy are applied to the direct numerical simulation of a configuration with both surface and gas-phase chemistry being active.

This study is organized as follows. First of all, an analytical solution is derived for a flow bounded by two reactive surfaces. Analytical solutions are then compared to numerical predictions for validation purposes (Section 3.1). Unsteady gasification and oxidation of carbonaceous materials coupled with gas-phase combustion under oxy-combustion conditions is analyzed through the DNS of a CO/O$_2$ combustion near a reactive carbonaceous surface in a quiescent flow (Section 3.2). Detailed and validated kinetics for surface reactions were used (Section 3.2.2). Transition from endothermic gasification to exothermic oxidation of the carbonaceous surface and CO gas-phase ignition is evidenced. The competition between homogeneous and heterogeneous reactions is discussed based on the analysis of the time-dependant surface and gas-phase species consumption and production rates. Transient histories of surface species concentrations as well as spatial profiles of gas-phase species, heat release and temperature are presented for two identified modes of gas-phase CO reaction zone. The impact of the Hirschfelder & Curtiss approximation on the results is assessed in Section 3.2.5 by post-processing the DNS data thanks to a complex transport library. At last, the impact of initial concentrations of O$_2$ and CO$_2$ on both competition between gas and surface reactions and characteristic times corresponding to the switch between the two structures of the gas-phase reaction zone is invistigated in Section 3.2.6.
## Nomenclature

### Dimensionless numbers

- **$Da$**: Damköhler number
- **$Fo$**: Fourier number
- **$Le$**: Lewis number
- **$Pr$**: Prandtl number
- **$Sc_k$**: Schmidt number of the $k^{th}$ species

### Greek symbols

- $\alpha_j$: Pre-exponential factor of the $j^{th}$ reaction (mol cm s)
- $\delta$: Kronecker symbol
- $\epsilon_{k,j}$: Surface coverage parameter for activation energy of the $k^{th}$ species involved in the $j^{th}$ surface reaction
- $\eta_{k,j}$: Surface coverage parameter for the pre-exponential factor of the $k^{th}$ species
- $\gamma_j$: Sticking coefficient for adsorption reactions
- $\Gamma$: Surface site density (mol m$^{-2}$)
- $\lambda$: Heat conduction coefficient (W m$^{-1}$ K$^{-1}$)
- $\mu$: Dynamic viscosity (Pa s)
- $\nu_{k,j}^r$, $\nu_{k,j}^i$: Molar stoichiometric coefficient of the $k^{th}$ species present as a product/reactant in the $j^{th}$ reaction
- $\Omega_G$: Index set of gas-phase species involved in gas-phase and surface reactions
- $\Omega_s$: Index set of surface species involved in surface-phase reactions
- $\dot{\omega}_k$: Rate of production by gas-phase reactions of the $k^{th}$ species (kg m$^{-3}$ s$^{-1}$)
- $\dot{\omega}_T$: Gas-phase heat release in (J m$^{-3}$ s$^{-1}$)
- $\rho$: Gas-phase density (kg m$^{-3}$)
- $\sigma_k$: Number of sites that each surface species occupies
- $\bar{\tau}$: Viscous stress tensor (kg m$^{-1}$ s$^{-2}$)
- $\theta_k$: Surface species site fraction
- $\tau$: Dimensionless time
- $\chi_k$: Thermal diffusion ratios
<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>$C_k$</td>
<td>Molar concentration of the $k^{th}$ surface species ($mol.m^{-2}$)</td>
</tr>
<tr>
<td>$C_{p,k}, C_{v,k}$</td>
<td>Heat capacity at constant pressure and constant volume ($J.\text{kg}^{-1}.K^{-1}$)</td>
</tr>
<tr>
<td>$D$</td>
<td>Index set of space dimensions</td>
</tr>
<tr>
<td>$D_k$</td>
<td>Mass diffusion coefficient of the $k^{th}$ gas-phase species ($m^2.s^{-1}$)</td>
</tr>
<tr>
<td>$E$</td>
<td>Total non-chemical energy ($J.\text{kg}^{-1}$)</td>
</tr>
<tr>
<td>$E_j$</td>
<td>Activation energy of the $j^{th}$ reaction ($J.mol^{-1}$)</td>
</tr>
<tr>
<td>$h_{s,k}, h_k$</td>
<td>Sensible and total enthalpy of the $k^{th}$ species ($J.\text{kg}^{-1}$)</td>
</tr>
<tr>
<td>$\Delta h_{f,k}$</td>
<td>Standard enthalpy of formation of the $k^{th}$ gas-phase species ($J.\text{kg}^{-1}$)</td>
</tr>
<tr>
<td>$J_k$</td>
<td>Diffusive flux of the $k^{th}$ gas-phase species ($kg.m^{-2}.s^{-1}$)</td>
</tr>
<tr>
<td>$k_{f,j}, k_{f,j}$</td>
<td>Forward rate constants of the $j^{th}$ gas-phase and surface reaction ($mol, cm, s$)</td>
</tr>
<tr>
<td>$K_{eq}$</td>
<td>Equilibrium constant</td>
</tr>
<tr>
<td>$l_{diff}$</td>
<td>Characteristic diffusion length within a chemical time $\tau_{chem}$</td>
</tr>
<tr>
<td>$m_c$</td>
<td>Surface carbon mass (kg)</td>
</tr>
<tr>
<td>$m$</td>
<td>Convected mass rate at the wall ($kg.m^{-2}.s^{-1}$)</td>
</tr>
<tr>
<td>$n$</td>
<td>Reaction order</td>
</tr>
<tr>
<td>$p$</td>
<td>Thermodynamic pressure ($Pa$)</td>
</tr>
<tr>
<td>$\dot{q}$</td>
<td>Heat flux ($J.m^{-2}.s^{-1}$)</td>
</tr>
<tr>
<td>$Q_j$</td>
<td>Reaction rate of the $j^{th}$ gas-phase reaction ($mol.m^{-3}$)</td>
</tr>
<tr>
<td>$Q_{s,j}$</td>
<td>Reaction rate of the $j^{th}$ surface reaction ($mol.m^{-2}$)</td>
</tr>
<tr>
<td>$\dot{s}_k$</td>
<td>Rate of production by surface reactions of the $k^{th}$ species ($kg.m^{-2}.s^{-1}$)</td>
</tr>
<tr>
<td>$R$</td>
<td>Universal gas constant ($J.mol^{-1}.K^{-1}$)</td>
</tr>
<tr>
<td>$T$</td>
<td>Temperature ($K$)</td>
</tr>
<tr>
<td>$\vec{u}$</td>
<td>Convection velocity field ($m.s^{-1}$)</td>
</tr>
<tr>
<td>$U_{Stefan}$</td>
<td>Stefan velocity ($m.s^{-1}$)</td>
</tr>
<tr>
<td>$\vec{V}_k$</td>
<td>Diffusion velocity field of the $k^{th}$ gas-phase species ($m.s^{-1}$)</td>
</tr>
<tr>
<td>$W_k, W$</td>
<td>Molar weight of the $k^{th}$ species and gas-phase mixture ($kg.mol^{-1}$)</td>
</tr>
<tr>
<td>$w$</td>
<td>Index denoting for values of scalars or vector components at the wall</td>
</tr>
<tr>
<td>$X_k, Y_k$</td>
<td>Mole and mass fraction of the $k^{th}$ gas-phase species</td>
</tr>
</tbody>
</table>
2. Modelling approach

2.1. Surface chemistry formalism

2.1.1. Surface reactions kinetics

Only reactions occurring on the top layer of the solid are considered and modeled through an open site formalism; bulk reactions are not considered. For each species, the surface reaction rate of production \( \dot{s}_k \) is the sum of rates of surface reactions:

\[
\dot{s}_k = W_k \sum_{j \in M_k} (v_{j}^s - v_{j}^g)Q_{s,j} \forall k \in \{ \Omega_g \cup \Omega_s \}. \tag{1}
\]

where \( Q_{s,j} \) represents the rate of progress of the \( j \)th surface reaction and is written according to the mean field approximation [25]:

\[
Q_{s,j} = k_{j}^s \prod_{\nu \in \Omega_k} \left( \frac{\rho_{j}^k Y_{k,\nu}}{W_k} \right)^{\nu^s} \prod_{\nu \in \Omega_k} \left( \Gamma^s \theta^s \right)^{\nu^g} \tag{2}
\]

In Eq. (2), \( \theta_k \) are surface-species site fractions and \( k_{j}^s \) is the forward reaction constant of the \( j \)th surface reaction. Only mono-directional steps are considered in this study. The reaction constants are evaluated according to the surface reaction type. The \( j \)th surface reaction constant is given by:

\[
k_{j}^s = A_{s,j}^{cov} \eta_{j}^{s} \exp \left( \frac{-E_{s,j}^{cov}}{RT_w} \right) \tag{3}
\]

where \( A_{s,j}^{cov} \) and \( E_{s,j}^{cov} \) are respectively the coverage dependent pre-exponential factor and activation energy that can be written as follows:

\[
\ln(A_{s,j}^{cov}) = \ln(A_{s,j}) + \sum_{k \in \Omega_k} \eta_k \theta_k \tag{4}
\]

and

\[
E_{s,j}^{cov} = E_{s,j} + \sum_{k \in \Omega_k} \varepsilon_{k,j} \theta_k \tag{5}
\]

Assuming the adsorption to be activated, the parameters \( \beta_{s,j}, E_{s,j}^{cov}, A_{s,j}^{cov} \) become [26–28]:

\[
\beta_{s,j} = \frac{1}{2}, \quad E_{s,j}^{cov} = 0, \quad A_{s,j}^{cov} = \gamma_j \left( \prod_{k \in \Omega_k} \sigma_k \right) \left( \frac{k}{T^0} \right)^{1/4} \sqrt{\frac{R}{2\pi W_{g,ads}}} \tag{6}
\]

where \( \sigma_k \) is the number of sites that each adsorbed species occupies, \( m \) corresponds to the number of surface species involved in the surface reaction and \( W_{g,ads} \) is the gas-phase species molar weight. \( \theta_k \) is related to the surface mole concentration of the \( k \)th surface species, \( C_k \), as follows:

\[
\theta_k = \frac{C_k \sigma_k}{\Gamma} \tag{7}
\]

The equation describing the variation of surface coverage [19] is based on the assumption that site density \( \Gamma \) remains constant, diffusion inside the solid is neglected as surface local diffusion is considered to be very fast with respect to reactions and every \( k \)th surface species occupies only one site, i.e, \( \sigma_k = 1 \):

\[
\frac{\partial \theta_k}{\partial t} = \frac{\dot{s}_k}{\Gamma W_k} \forall k \in \Omega_s \tag{8}
\]

The surface site fractions correspond to a single mono-layer which implies that their sum equals unity:

\[
\sum_{k \in \Omega_k} \theta_k + \theta_{void} = 1 \tag{9}
\]

where \( \theta_{void} \) corresponds to the fraction of void sites.
2.1.2. Accounting for surface chemistry through boundary conditions

The flow solver used for this study is a conservative finite-volume based and known as AVBP [29]. It has been extensively validated over a wide range of applications [30–33]. In Appendix A, the set of the unfiltered equations used in Direct Numerical Simulations (DNS) governing the evolution of a compressible flow with gas-phase chemical reactions is described. The modeling of transport coefficients, diffusion and gas-phase kinetics are also available.

AVBP is a gas-phase solver whereas surface reactions correspond to chemical phenomena occurring at the wall. Therefore, the interaction of the latter with the gas-phase scalars was accounted for through boundary conditions. Heterogeneous reactions occurring at the reactive wall consume or produce gas-phase species. This consumption or production leads to an additional velocity at the interface while it releases or adsorbs heat because of solid-fluid reactions that are not necessarily athermal. Hence, surface chemistry impacts gas-phase species mass fractions, energy and momentum. Details of the approach giving the mass and energy balance at a reactive interface with the associated boundary conditions are provided in Appendix B. 

\( \vec{u}_w = U_{\text{Stefan}} \vec{n} \) (10)

\( \vec{u}_w \) is the velocity vector at the reactive wall. The Stefan velocity is obtained from the species mass conservation balance at the reactive interface.

\[ (\rho_w V_{k,n,w} Y_{k,w}) + (\rho_w U_{\text{Stefan}} Y_{k,w}) = \dot{s}_k \] (11)

\( V_{k,n,w} \) refers to the normal diffusion velocity at the reactive wall. Summing Eq. (11) over all the gas-phase species and considering the mass conservation, an expression for the wall-normal Stefan velocity is obtained:

\[ U_{\text{Stefan}} = \frac{1}{\rho_w} \left( \sum_{k \in \Omega} \dot{s}_k \right) = \frac{\dot{m}}{\rho_w} \] (12)

where \( \dot{m} \) is the convected mass-rate at the wall.

Surface chemistry impacts the gas-phase species mass fraction at the reactive wall. The prescribed normal gradient noted \( \vec{\nabla} Y_{k,w} \cdot \vec{n} \) is obtained by expressing in the mass balance given in Eq. (11) the diffusion velocity in terms of species mass fractions gradients by using the Hirschfelder & Curtiss approximation [34, 35] whose impact is evaluated in Section 3.2.5. Hence, it becomes possible to relate the species mass fraction at the wall \( Y_{k,w} \) to its normal gradient at the boundary surface:

\[ \vec{\nabla} Y_{k,w} \cdot \vec{n} = \frac{Y_{k,w}}{D_k \rho_w} \left( \sum_{l \in \Omega_{\text{gas}}} \dot{s}_l + \frac{Y_{k,w}}{D_k} \vec{V}_{\text{correc}} \cdot \vec{n} \right) + Y_{k,w} W_w \left( \sum_{l \in \Omega_{\text{w}}} \frac{1}{W_l} \vec{\nabla} Y_{l,w} \cdot \vec{n} \right) - \frac{\dot{s}_k}{\rho_w D_k} \] (13)
Eq. (13) corresponds to the boundary condition for species mass fractions and constitutes a non-linear system of equations which is solved thanks to an iterative method [36, 37]. In the following, the energy boundary condition accounting for surface chemistry is developed. Surface chemistry in non-athermal cases impacts the wall temperature since surface reactions may be endothermic or exothermic while the reactive interface undergoes heat exchange including gas-phase thermal diffusion and convection, species diffusion, viscous heat and surface reactions heat release involving both surface and gas-phase species. The heat balance at the reactive wall reads:

\[
\lambda \vec{\nabla} T_w^s \cdot \vec{n} = \sum_{k \in \Omega_w} h_k \dot{s}_k - (\vec{\varpi}_w) \cdot \vec{n} + \frac{m}{2} \sum_{j \in D} u_{j,w}^2 + \lambda \vec{\nabla} T_w^s \cdot \vec{n} \tag{14}
\]

Eq. (14) constitutes the heat flux balance that accounts for surface chemistry impact on energy. The balance enables to write the heat flux to be prescribed to the boundary condition \( q_{BC}^{w} \) as shown in Eq. (17). The rate of production/consumption of species due to gas-phase reactions is set to zero at the wall so that no gas-phase reactions occur at the reactive surface:

\[
\dot{\omega}_{k,w} = 0 \tag{15}
\]

The resulting heat release due to gas-phase reaction presents zero values at the wall:

\[
\dot{\omega}_{T,w} = 0 \tag{16}
\]

The conditions of Eqs. (15, 16) are not related to equilibrium assumptions. \( \dot{\omega}_{k} \) and \( \dot{\omega}_{T} \) are set to zero so that the species and energy boundary conditions are consistent with Eq. (B.5). Therefore, the near-wall grid resolution chosen for this study (sec 3.2.1) is judged fair enough to ensure these conditions.

\[
q_{w}^{BC} \cdot \vec{n} = - \sum_{k \in \Omega_w} h_k \dot{s}_k - \sum_{k \in \Omega_w} \Delta H_f^k \dot{s}_k - \rho_w \sum_{k \in \Omega_w} U_{Stefan} h_{x,k,w} Y_{k,w} - \frac{m}{2} \sum_{j \in D} u_{j,w}^2 + (\vec{\varpi}_w) \cdot \vec{n} - \lambda \vec{\nabla} T_w^s \cdot \vec{n} \tag{17}
\]

**2.1.3. Numerical implementation of the boundary conditions**

A Dirichlet condition is applied to the velocity vector at the wall in order to impose the Stefan velocity as shown in Eq. (18).

\[
\vec{u}_{w}^{cor} = U_{Stefan} \vec{n} \tag{18}
\]

Where \( \vec{u}_{w}^{cor} \) is the corrected velocity at the wall. The normal gas-phase species mass fraction gradients and the heat flux are imposed numerically at the reactive surface as shown in Eqs. (19, 20). First, the numerical scheme predicts the gradient of the \( k^{th} \) species mass fraction \( \vec{\nabla} Y_{k,w}^p \) and the heat flux \( \vec{q}_{w}^p \) at the reactive wall. These gradients are then modified by imposing the
prescribed normal gradients and the normal heat flux \( \hat{\nabla} Y_{k,w}^{BC} \cdot \vec{n} \) and \( \hat{q}_w^{BC} \cdot \vec{n} \) obtained respectively from the mass and energy balance at the reactive interface given in Eqs. (11, 14).

\[
\hat{\nabla} Y_{k,w}^{cor} = \hat{\nabla} Y_{k,w}^{p} + \left( \hat{\nabla} Y_{k,w}^{BC} \cdot \vec{n} - \hat{\nabla} Y_{k,w}^{p} \cdot \vec{n} \right) \cdot \vec{n} 
\]

(19)

\[
\hat{q}_w^{cor} = \hat{q}_w^{p} + \left( \hat{q}_w^{BC} \cdot \vec{n} - \hat{q}_w^{p} \cdot \vec{n} \right) \cdot \vec{n} 
\]

(20)

Where \( \hat{\nabla} Y_{k,w}^{cor} \) and \( \hat{q}_w^{cor} \) are the corrected species mass fraction gradient and heat flux, \( \hat{\nabla} Y_{k,w}^{p} \cdot \vec{n} \) and \( \hat{q}_w^{p} \cdot \vec{n} \) correspond to the normal component of the \( k \)th species mass fraction and the heat flux predicted by the numerical scheme.

3. Results

3.1. Analytical solution for wall-bounded flow with carbonaceous wall gasification

In this section the purpose is to derive an original analytical solution to validate the numerical species boundary condition for the heterogeneous surface reactions with frozen gas-phase. This solution can be used to understand the physics and easily analyze the influence of some physical parameters onto the studied phenomena. Moreover, analytical solutions are of wide interest for experimentalists or computational fluid dynamics users to validate their experimental/numerical procedure. Only the global approach is described and the final solution is given. The complete development is provided as a supplementary material in Appendix C. It is sometimes possible to find the analytical solution of the system of partial differential equations that governs the physics of simple two-dimensional laminar flows. An example is the Poiseuille flow configuration which allows to give the analytical velocity/temperature profiles under the assumption of streamwise-developed steady flow [38]. Introducing the transpiring surface effects, Koh and Hartnett [39] have shown that the analytical solution can also be recovered for flow over permeable wedges. Hamza [40] also presented the solution of channel flow with transpiring and moving walls for biomechanics-fluid interest. Concerning reacting wall effects, Rosner [41] introduced a set of solutions for surface catalyzed problems. Duan and Martín [42] also made use of an analytical solution for a diffusion-tube sidearm reactor in order to validate their numerical simulation of airflow over purely catalytic wall. To our knowledge, surface ablation has received little attention in the literature. Libby & Blake [8] developed an analytical solution for ablation but their case of study was purely diffusive. A special attention is given to the configuration set-up which allows to seek a steady-state solution. The idea consists in considering a channel problem of which the lower surface undergoes gaseous species production and solid carbon recession whereas the upper one encounters carbon deposition. Within this framework, the velocity and molar fraction profiles are derived and discussed. Global surface reactions and kinetic models are used, the purpose in this section being to seek numerical validation and not
kinetic accuracy. The surface consumption has an effect on the flow since the products of the heterogeneous reactions include atoms that were originally embedded in the wall lattice. As a consequence, the surface geometry is changed according to the reaction rates at the surface. This feature makes the derivation of the solution more challenging, especially when a steady-state solution is sought for. As presented hereafter, the configuration set-up has to be properly thought to handle this undesirable effect. To simplify the derivation, one seeks for a solution that satisfies the following assumptions:

- two-dimensional laminar flow,
- steady state,
- channel flow developed in streamwise direction (periodic boundary conditions),
- non-reacting binary mixture (frozen state),
- constant Schmidt numbers,
- pure gasification (no pyrolysis or sublimation of the wall material),
- isothermal gasified surface with finite reaction rates,
- constant pressure and temperature.

In such a configuration, a special attention must be given to the chemical scheme retained for the heterogeneous surface reactions. Indeed, if the same scheme is applied to the two surfaces, this leads to a flow that similarly behaves at the wall surfaces. This means that the two convective wall normal velocity vectors induced by surface gasification are pointing towards opposite directions. This translates into an overall increase (or decrease) of mass that is not compatible with the steady state assumption. For this reason, one sets up a configuration in which one of the surface is consumed by a reaction while the reverse reaction takes place at the facing surface so mass increase or decrease in the domain is avoided. The set-up retained is summarized in Fig. (1). The flow is supposed to evolve in a streamwise periodic channel between two plates separated by the distance \( h \). The wall normal direction is referenced by the letter \( y \) whereas \( x \) is used for the streamwise direction.

Figure 1: Sketch of the configuration.

Making use of the assumption of steady developed non-reacting flow, one can write the multicomponent Navier-Stokes mass/species/momentum conservation equations as:

\[
\frac{\partial (\rho v)}{\partial y} = 0 \quad (21)
\]
\[ \frac{\partial}{\partial y} \left( \rho \left( v + V_{k,y} \right) Y_k \right) = 0, \quad k \in \{1, 2\} \quad (22) \]

\[ \frac{\partial (\rho u)}{\partial y} = \frac{\partial \tau_{xy}}{\partial y} + S \quad (23) \]

Where \( u \) the streamwise velocity component, \( v \) the wall normal velocity component, \( Y_k \) the mass fraction of species \( k \), \( V_{k,y} \) the wall normal diffusion velocity of species \( k \), \( \tau_{xy} = \mu u_d/dy \) the shear stress, and \( S \) is the source term driving the flow. Note that this term must be added because the periodicity condition along \( x \) imposes \( \partial p/\partial x = 0 \). Finally, the perfect gas equation of state is used to close the problem, which supposes that no real gas effects are considered. Moreover, the general formulation of species conservation balance at the reacting surfaces reads:

\[ \left( \rho_j \hat{n}_j Y_{k,j} + \rho_j \hat{V}_{k,j} Y_{k,j} \right) \cdot \hat{n}_j = \dot{s}_{k,j} \quad (24) \]

Index \( j \) refers to the reactive surface number, \( j = 1 \) being the lower surface whereas \( j = 2 \) corresponds to the upper one. \( \hat{n}_j \) is the unitary wall normal vector at surface \( j \) directed towards the gaseous domain as shown in Fig. (1). \( \dot{s}_{k,j} \) is the mass production rate of species \( k \) defined as \( \dot{s}_{k,j} = \overline{v}_{k,j} W_k Q_j \) with \( \overline{v}_{k,j} \) the algebraic stoichiometric coefficient of the \( k \) species at the \( j \) surface (positive for produced species, negative for consumed species). Under these conditions described in Fig. (1), one seeks for an analytical solution for the independent variables \( u(y) \), \( v(y) \), \( Y_k(y) \) (or equivalently \( X_k(y) \)) via the relation \( X_k = Y_k W/W_k \). The integration of Eq. (24) leads to two solutions given in Eq. (25) depending on the values taken by the stochiometric numbers \( v_O \) and \( v_P \) referring respectively to the oxidizer (O) and the product(P).

\[ X_O(y) = \left( X_{O,1} - \frac{v_O}{v_O - v_P} \right) \exp \left[ -\frac{S c_O m}{\mu} y \right] + \frac{v_O}{v_O - v_P} \]

for \( v_O \neq v_P \)

\[ X_O(y) = -\frac{S c_O m}{\mu} \frac{W_P}{W_O - W_P} y + X_{O,1} \]

for \( v_O = v_P \)

(25)

The mole fraction of \( P \) is then given by the relation \( X_P(y) = 1 - X_O(y) \). Here \( X_{O,1} \) is the molar fraction of species \( O \) at surface 1, \( m \) correspond to the total mass convected mass-rate at the reactive surfaces, \( S c_O \) is the Schmidt number of the oxidizer species and \( \eta = \frac{W_P (v_O - v_P)}{v_O - v_P w_P} \). Concerning the determination of the streamwise velocity profile, one introduces the newtonian fluid relation \( \tau_{xy} = \mu u_d/dy \) into Eq. (23) which gives \( \rho \nu \; du/dy = \mu \; d^2 u/dy^2 = S \). Since \( \mu \) is constant under the approximation of constant temperature, and because \( \rho(y) v(y) = m \) is constant according to the continuity equation given in Eq. (21), the latter differential equation is merely of second order with constant coefficients. Its integration with the boundary conditions \( u(0) = u(h) = 0 \) leads to the following solution:

\[ u(y) = \frac{S}{m} \left[ \frac{h}{e^{h/\nu} - 1} \left( 1 - e^{\nu y} \right) + y \right] \quad (26) \]

It is interesting to notice that if \( m \) tends to zero, the use of second-order Taylor series in Eq. (26) allows to demonstrate that the streamwise velocity profile
tends to the classical Poiseuille one, namely \( u(y) = \frac{S}{(2\mu)(yh - y^2)} \). Finally, the maximum streamwise velocity is expected to be observed at the wall normal distance which differs from a classical Poiseuille profile for which the maximum velocity is observed at the channel half-height. This deviation is only due to the wall normal mass flow rate and non-sensitive to the driving pressure gradient. Note also that making use of Taylor series (for \( m \to 0 \)) it is easy to show that \( y_{u_{\text{max}}} \to h/2 \).

\[
y_{u_{\text{max}}} = \frac{\mu}{m} \ln \left[ \frac{\mu}{h m} \left( e^{zh} - 1 \right) \right]
\] (27)

The analytical solution is now discussed in the case of carbon \( \text{CO}_2 \) gasification. Three cases presented in Table 1, for which \( h = 1 \text{mm} \) and \( S = 1 \text{kPa/m} \). The transport coefficients are determined thanks to the EGlib library [43]. All cases represent a realistic solution for which the surface reaction has a finite-rate, in other words, the concentration of the oxidizing species is different from zero at surface 1 and different from unity at surface 2. In order to ensure a balance between both forward and backward reactions occurring respectively at surface 1 and 2, constants of reactions are set equal, i.e, \( K_1 = K_2 \).

The comparisons of analytical and numerical solutions for the wall normal velocity component, the mass fraction of the oxidizing species and the streamwise velocity are presented in Figs. (2-4) for \( \text{CO}_2 \) gasification. It is noticed that the wall normal distance for which the maximum velocity is observed \( y_{u_{\text{max}}} \) is identical to the channel half-height \( h/2 \). Moreover, the fluid encounters the same wall shear stress at both surfaces, i.e, \( \tau_{w,2}/\tau_{w,1} \approx 1 \). These cases correspond to values of mass rates that are too low to modify the streamwise velocity profiles significantly (Table 2). As a result, these profiles are very close to the impermeable wall solution. The analytical solutions are identical to the numerical ones, showing that the boundary conditions taking into account the Stefan velocity are able to predict correctly the influence of surface reactions on gas-phase species and momentum given respectively in Figs. (3, 4).

3.2. Unsteady combustion near a carbonaceous surface

An original analytical solution was derived in the previous section to develop and validate the numerical species boundary condition for the heterogeneous surface reactions. In this section, appropriate boundary conditions consistent with the conservation of gaseous/surface species and energy are applied to the direct numerical simulation of a configuration with both surface and gas-phase chemistry being active.

3.2.1. Numerical setup

The considered problem is purely 1D, variations occurring exclusively between the upper and lower surface and all fields being homogeneous in the \( y \) and \( z \) directions because of periodicities. The configuration studied is a parallelepipedic channel containing initially a quiescent and homogeneous
Table 1: Parameters of the test cases.

<table>
<thead>
<tr>
<th>case</th>
<th>Oxidizer (O)</th>
<th>Product (P)</th>
<th>$v_O$</th>
<th>$v_P$</th>
<th>$S_{cO}$</th>
<th>$S_{cP}$</th>
<th>$K_1 = K_2$ (kg·m$^{-1}$·mol$^{-1}$·s$^{-1}$) [8]</th>
<th>P(atm)</th>
<th>T(K)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>CO$_2$</td>
<td>CO</td>
<td>1</td>
<td>2</td>
<td>0.92</td>
<td>0.59</td>
<td>$6.92 \times 10^{-3}$</td>
<td>1</td>
<td>1400</td>
</tr>
<tr>
<td>2</td>
<td>CO$_2$</td>
<td>CO</td>
<td>1</td>
<td>2</td>
<td>0.92</td>
<td>0.59</td>
<td>$5.2 \times 10^{-2}$</td>
<td>1</td>
<td>1600</td>
</tr>
<tr>
<td>3</td>
<td>CO$_2$</td>
<td>CO</td>
<td>1</td>
<td>2</td>
<td>0.92</td>
<td>0.59</td>
<td>0.49</td>
<td>1</td>
<td>1900</td>
</tr>
</tbody>
</table>

Table 2: Characteristic values of the test cases.

<table>
<thead>
<tr>
<th>case</th>
<th>$\dot{m}$</th>
<th>$\frac{y_{\text{limax}}}{h}$</th>
<th>$\tau_{w,2}/\tau_{w,1}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>$3.87 \times 10^{-4}$</td>
<td>0.5</td>
<td>1</td>
</tr>
<tr>
<td>2</td>
<td>$2.16 \times 10^{-3}$</td>
<td>0.5</td>
<td>1.01</td>
</tr>
<tr>
<td>3</td>
<td>$8.88 \times 10^{-3}$</td>
<td>0.5</td>
<td>1.052</td>
</tr>
</tbody>
</table>

$O_2$/CO$_2$ mixture. Solid carbon consumption and CO/CO$_2$ production through surface gasification and oxidation takes place at the lower wall, whereas the upper surface corresponds to an outlet ensuring a constant pressure within the whole channel. The channel height $h$ is higher than the distance $l_{\text{diff}} = \sqrt{\tau_{\text{chem}}(v/Sc)|_{t=0}}$ characterizing diffusion within a chemical time $\tau_{\text{chem}}$. The latter is evaluated in a homogeneous adiabatic reactor provided with identical initial thermodynamic and thermo-chemical conditions as the present studied case. The corresponding value of $\tau_{\text{chem}}$ coincides with complete depletion of surface carbon, i.e., $\theta_{C(S)} = 0$. $(v/Sc)|_{t=0}$ refers to the ratio of kinematic viscosity and the average Schmidt number of the mixture. According to this approach, the initial Damköhler number ($Da$) is much larger than unity which makes the combustion process overall diffusion-limited.

$$Da = \left( \frac{h}{l_{\text{diff}}} \right)^2 \gg 1 \quad (28)$$

The surface over volume ratio $A_{\text{wall}}/V$ ensures that the gas and surface reactions contributions to CO$_2$ and $O_2$ consumption and production are of the same order of magnitude which implies that the surface species consumption proceeds at a rate as fast as gas-phase chemistry. $A_{\text{wall}}$ being the chemically active surface area of the lower wall (Fig. (5)), the ratio $A_{\text{wall}}/V$ reads:

$$\frac{A_{\text{wall}}}{V} = \frac{a^2}{a^2 h} = \frac{1}{h} \quad (29)$$

Variables initialization is spatially homogeneous, Table 3 summarizes the initial concentrations and thermodynamic parameters for the reference case. The transport coefficients are determined using the EGLIB library [43]. An adiabatic condition is applied to the lower reactive surface which means that the
temperature of the reactive wall presents temporal variations based on the heat flux balance given in Eq. (14). The condition is called adiabatic in the sense that no thermal conduction is accounted for within the solid, which means that $\lambda \nabla T_s \cdot \vec{n} = 0$. The latter expression implies that the prescribed normal heat flux at the wall becomes:

$$q_{BC}^w - \dot{m}U_{Stefan}h_{s,k,w}Y_{k,w} - \frac{m^2}{2} \sum_{j \in D} u_{j \omega}^2 + (\vec{\tau} \cdot \vec{n}) \cdot \vec{n} = 0$$

The prescribed mass fraction gradient at the wall
is given by Eq. (13). The upper face is an outlet surface that relaxes the pressure toward the targeted atmospheric pressure in a partially non-reflecting way using the NSCBC method [44, 45]. This allows to control waves reflections and avoids the propagation of non-physical fluctuations. No conditions are applied to the species mass fractions, heat flux and velocity at the outlet:

\[ Y_k|_{\text{outlet}} = Y_k^p, \quad \vec{q}|_{\text{outlet}} = \vec{q}^p, \quad \text{and} \quad \vec{u}|_{\text{outlet}} = \vec{u}^p \quad (31) \]

Where \( Y_k^p \), \( \vec{q}^p \) and \( \vec{u}^p \) are respectively the values of species mass fractions, heat flux and velocity predicted by the numerical scheme. Periodic boundary patches are imposed to the lateral faces. Regarding the mesh, starting from the wall to \( x/l_{\text{diff}} = 0.865 \), mesh cells have a size of 10\( \mu \text{m} \) which means that 1000 cells populate the near-wall region. For \( 0.865 < x/l_{\text{diff}} < \sqrt{D_a} \), a mesh coarsening has been carried out to reach a maximum cell size of 75\( \mu \text{m} \) at the outlet. As will be discussed in section 3.2.7, a grid convergence study is carried out to determine the proper grid resolution in the reactive near-wall region. Regarding the numerical scheme, a second-order cell-vertex implementation of the Finite Volume Lax-Wendroff convection scheme is used for time and space integration [46–49]. It is associated in this study with a Finite Element diffusion scheme with a 2\( \Delta \) operator that comes from a strict application of the Finite Element Galerkin method [50]. Diffusive terms are resolved in an explicit way.

### 3.2.2. Surface and gas-phase kinetic mechanisms

The surface-phase mechanism (Table 4) was extracted from a previously developed and validated semi-detailed kinetic mechanism [10, 51, 52]. The impact of backward reactions is expected to be negligible under typical oxy-combustion conditions and the validity of this hypothesis over a wider range of conditions is outside the scope of the present study. Two additional reactions representing the adsorption of \( CO_2 \) (S4) and oxygen radicals \( O \) (S5) upon carbon sites have been accounted for. The kinetic parameters of surface reaction (S1 to S3) related to char oxidation are similar to those recommended by Hurt et al [53], whereas those of \( CO_2 \) adsorption (S4) are in line with values proposed by Hecht et al [4]. The resulting detailed kinetic surface mechanism involves a lumped intermediate species, \( C(O) \), standing for oxygenated surface species formed during oxidation and a single type of carbon site, \( C(S) \), was chosen to describe solid carbon sites. The mechanism has been validated against continuously stirred tank reactor (CSTR) experiments by comparing predicted carbon conversion rates to the experimental data for char gasification and oxidation [54]. The simulations were carried out using Surface Chemkin kinetic solver [55]. Comparisons of the carbon conversion are shown in Fig. (6) corresponding to carbon \( O_2 \) oxidation and \( CO_2 \) gasification. The validation covers various concentrations of diluted \( O_2 \) and \( CO_2 \) using chemically inert molecular nitrogen \( N_2 \) as a balance dilution gas. Regarding the gas-phase
mechanism, the three reversible elementary reactions of dry CO oxidation given in Table 5 were extracted from the GRI 2.11 mechanism. This reaction set was then validated against shock tube experiments of Brabbs et al. for CO₂ dissociation [56].

3.2.3. History

The reference case (Table 3) is characterized by two gas-phase reaction zone configurations. The first one corresponds to a wall-attached CO-reaction zone maintained by the competition between the CO/O₂ combustion and surface reactions, whereas the second one is characterized by the extinction of surface reactions and a gas-phase reaction zone evolving at a certain distance of the reactive wall. Therefore, two main combustion modes characterize the reference case and the switch between these modes occurs at \( t_{\text{switch}} = 0.22 \) s under the conditions used in this study. This time coincides with the depletion of surface carbon \( (\theta_c = 0) \) and is the same order of \( \tau_{\text{chem}} \). As a consequence, time is normalized by the time corresponding to the combustion mode switch \( t_{\text{switch}} \). The resulting normalized time being noted \( \tau = t/t_{\text{switch}} \). A sketch of the two combustion modes is given in Fig. 7.

![Figure 6](image-url)

**Figure 6:** Carbon conversion in a \( O_2/N_2 \) and \( CO_2/N_2 \) environment. The volumetric flow rate is 100 SCCM (Standard Cubic Centimeter per Minute) and the heating rate is 10K/min.

![Figure 7](image-url)

**Figure 7:** Gas-phase CO/O₂ reaction zone configurations

Fig. (8a) shows the time evolution of space averaged mass fraction of the \( k^{th} \) species at the reactive
Table 4: Surface reaction mechanism for the configuration of Fig. 5

<table>
<thead>
<tr>
<th>Step</th>
<th>Reaction</th>
<th>$\alpha_j$ ( \text{cm}^{2(n-1)} \cdot \text{mol}^{1-n} \cdot \text{s}^{-1} )</th>
<th>$\beta_j$</th>
<th>$E_j$ (kJ mol$^{-1}$)</th>
<th>ref.</th>
</tr>
</thead>
<tbody>
<tr>
<td>S1</td>
<td>$C(S) + \frac{1}{2} O_2 \rightarrow C(O)$</td>
<td>$5.0 \times 10^6$</td>
<td>0.0</td>
<td>45</td>
<td>[10]</td>
</tr>
<tr>
<td>S2</td>
<td>$C(O) \rightarrow CO$</td>
<td>$2.7 \times 10^6$</td>
<td>0.0</td>
<td>132</td>
<td>[10]</td>
</tr>
<tr>
<td>S3</td>
<td>$C(O) + \frac{1}{2} O_2 \rightarrow CO_2$</td>
<td>$1.33 \times 10^5$</td>
<td>0.0</td>
<td>109</td>
<td>[10]</td>
</tr>
<tr>
<td>S4</td>
<td>$CO_2 + C(S) \rightarrow C(O) + CO$</td>
<td>$1.0 \times 10^{13}$</td>
<td>0.0</td>
<td>225</td>
<td>[54]</td>
</tr>
<tr>
<td>S5</td>
<td>$O + C(S) \rightarrow C(O)$</td>
<td>$\left( \frac{Y}{\Gamma_m} \right) \sqrt{\frac{R}{2\pi W_O}}$</td>
<td>0.5</td>
<td>0.0</td>
<td>[57]</td>
</tr>
</tbody>
</table>

Table 5: Gas-phase reaction mechanism for the configuration of Fig. 5

<table>
<thead>
<tr>
<th>Step</th>
<th>Reaction</th>
<th>$\alpha_j$ ( \text{cm}^{3(n-1)} \cdot \text{mol}^{1-n} \cdot \text{s}^{-1} )</th>
<th>$\beta_j$</th>
<th>$E_j$ (kJ mol$^{-1}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>R1</td>
<td>$2O + M = O_2 + M$</td>
<td>$1.2 \times 10^{17}$</td>
<td>-1.0</td>
<td>0.0</td>
</tr>
<tr>
<td>R2</td>
<td>$O + CO + M = CO_2 + M$</td>
<td>$6.02 \times 10^{14}$</td>
<td>0.0</td>
<td>12.55</td>
</tr>
<tr>
<td>R3</td>
<td>$O_2 + CO = O + CO_2$</td>
<td>$2.5 \times 10^{12}$</td>
<td>0.0</td>
<td>200.1</td>
</tr>
</tbody>
</table>

For $\tau \leq 1$, $CO_2$ and $O_2$ species present in the gas-phase diffuse to the carbonaceous wall, are adsorbed and converted on surface as shown by their respective decreasing mass fractions at the wall shown in Fig. (8a). $CO_2$ and $O_2$ adsorption upon the reactive surface coincides with consumption of surface carbon and $CO$ production through gasification. $CO$ production by the reactive surface is described by the increasing profile of $CO$ mass fraction (Fig. (8a)) and the decreasing profile of carbon site fraction $\theta_c$ (Fig. (8b)). $CO$ diffuses to the gas-phase and reacts with $O_2$ thus producing $CO_2$. Carbon monoxide consumption and carbon dioxide production through gas-phase combustion are illustrated by the respective rise and fall of $CO_2$ and $CO$ concentrations for $0.97 \leq \tau \leq 1.0$. At $\tau = 1$, surface carbon consumption reaches completion which puts an end to surface reactions. Gas-phase $CO_2/O_2$ combustion is the only active chemical process which gradually fades.
3.2.4. Combustion modes

Wall-attached reaction zone:

In order to quantify the competition between gas-phase and surface reactions, an index $I_k$ for each $k^{th}$ species is defined in Eq. (33):

$$I_k = \frac{|\langle \dot{s}_k \rangle|}{|\langle \dot{s}_k \rangle| + |\langle \dot{\omega}_k \rangle|}$$  \hspace{1cm} (33)

where $\langle \dot{\omega}_k \rangle$ and $\langle \dot{s}_k \rangle$ are respectively the gas-phase and surface space averaged production/consumption rates that read:

$$\langle \dot{\omega}_k \rangle = \int_V \dot{\omega}_k dV, \quad \langle \dot{s}_k \rangle = \int_S \dot{s}_k dS \quad (34)$$

$I_k$ characterizes the contributions of surface and gas-phase reaction regardless of its sign (production/consumption). Fig. (9) presents the evolution of $I_{O_2}$, $I_{CO_2}$ and $I_{CO}$ versus the normalized time $\tau$. As can be noticed, the index $I_k$ varies between 0 and 1 thus showing that the existence of a wall-attached reaction zone coincides with the competition between surface and gas-phase chemistry. As a matter of fact, $I_{O_2}$, $I_{CO_2}$ and $I_{CO}$ are higher than 0.5 for $\tau \leq 0.24, 0.7$ and 0.75 respectively, which means that surface-phase chemistry is as much involved into species consumption or production as the gas-phase one. $I_{CO}$ and $I_{CO_2}$ show a stiff decrease when $\tau$ tends to 1 because of the quick drop of surface carbon mass (Fig. (8)) putting an end to surface chemistry. This is due to the thermal runaway caused by the exothermic surface carbon oxidation (S1 to S3) and gas-phase reactions. A slight increase can be noticed before $I_{CO_2}$ decreases rapidly which is due to the production of $CO_2$ by reaction S3 (Table 4).

$CO$ provided by the surface feeds a $CO/O_2$-combustion characterized by a reaction front at proximity of the reactive wall. Fig. (10) presents the reaction zone identified by the gas-phase temperature (a) and the peaks of heat release (b). These peaks intensify and move slightly upward toward rich zones that are less $O_2$ diffusion limited but do not go beyond $x/l_{diff} = 0.215$. Note that the reaction zone thickness (non-zero
values of $\hat{\omega}_T$) corresponds to regions where surface produced CO has diffused enough to react with $O_2$ thus covering a length that is the same order of the a priori-estimated characteristic diffusive length $l_{diff}$ as $\tau$ tends to unity, i.e., as physical time $t$ tends to one chemical characteristic time $\tau_{chem}$. This $CO/O_2$-reaction zone configuration is analogous to the “attached CO-flame” mode described by Makino [58]. The reaction zone is continuously fed by CO provided by the surface ($\langle k_{CO} \rangle > 0$), it remains attached to the wall despite the increase of $O_2$ gradient.

Fig. (11) shows at two different times during the wall-attached reaction zone phase the evolution of normalized species spatial profiles. Normalized scalars noted $\hat{\Phi}_k$ are defined as follows:

$$
\hat{\Phi}_k = \frac{\Phi_k - \Phi_{k,min}}{\Phi_{k,max} - \Phi_{k,min}} \tag{35}
$$

where $\Phi_{k,min}$ and $\Phi_{k,max}$ are respectively the minimum and maximum values of the scalar $\Phi_k$ at a given time $\tau$. $\hat{\Phi}_k$ equals zero when $\hat{\Phi}_k = \Phi_{k,min}$ and 1 if $\hat{\Phi}_k = \Phi_{k,max}$.
Gas-phase combustion occurrence coincides with a decrease of $O_2$ in the reaction zone and the formation of a peak on $CO_2$ profile. At $\tau = 0.442$, the minimum of $CO_2$ concentration is at the wall because of its consumption by surface carbon. As $CO/O_2$ reaction becomes significantly active in the gas-phase, $CO_2$ production contributes to generate a peak in the corresponding concentration spatial profile at $x/l_{diff} = 0.425$ at $\tau = 0.954$ identifying the gas-phase reaction front. The wall-attached reaction phase ($\tau \leq 0.06$) begins with an endothermic process during which $CO_2$ oxidizes carbon. $CO_2$ adsorption through surface reaction causes a drop in both wall (Fig. (12a)) and gas-phase temperatures (Fig. (12b)). Indeed, the temperature decrease is due to the endothermicity of $CO_2$ gasification of carbon. Fig. (12a) shows the decrease of the surface temperature during the endothermic phase ($\tau \leq 0.06$) causing an increase of the positive gas-phase temperature gradient at the reactive wall as described in Fig. (12b). Meanwhile, $CO$ mass fraction gradually increases at the wall, sharpening its negative gradient as can be seen in Fig. (13).
Gas-phase reactions after surface carbon depletion:

An overview of the gas-phase reaction zone displacement is shown in Figs. (14a, 14b). As carbon reaches depletion, surface reaction rates decrease which induces a change in the reaction structure. At $\tau = 1.0$, surface chemistry is not active anymore ($\langle \dot{s}_k \rangle = 0$) as shown by Fig. (9). Temperature in proximity of the wall reaches values higher than $3000K$ (Fig. (14) favoring $CO_2$ gas-phase dissociation thus causing a detachment of the reaction front. Fig. (14a) shows negative heat release rates near the wall region related to $O_2$ and $CO_2$ dissociation by $R1$ and $R2$ (Table 5). $CO$ is not generated anymore by the reactive surface so that the reaction zone loses its attachment to the surface. The resulting reaction zone turns into a reaction front moving along the channel height. Fig. (14b) shows spatial profiles of the gas-phase temperature for different times during the phase where the reaction zone is detached from the wall. Unlike gas-phase temperature profile in Fig. (10a) that presents a peak, temperature profiles have a maximum at the wall resulting from the high temperature history of the phase during which the reaction zone was attached to the reactive wall. The heat-release is not high enough to induce a peak in the temperature profile.

As shown in Fig. (14a), the heat release peak moves upward and decreases in amplitude with time. Indeed,
at $\tau > 1.0$, CO diffuses in the whole domain and mixes with $O_2$ thus creating a partially premixed reactant field where the reaction zone propagates toward $O_2$-rich regions. The magnitude and the displacement velocity of the heat release profiles peak drop because the further the flame gets from the reactive wall, the less conducive are the conditions to sustain the gas-phase combustion. As described in Fig. (15), CO concentration decreases as the distance to the wall increases, while rich $O_2$ regions do not compensate for CO diffusion limitation. The zones where $\dot{\omega}_T(x) \leq 0$ for a distance between 0 and $x/l_{diff} \approx 2.5$ (Fig. (14a)) result from $O_2$ dissociation into $O$ radicals and $CO_2$ dissociation into $CO$ and $O$ described respectively by $R1$ and $R2$ given in Table 5.

According to the DNS results, heterogeneous carbon ignition through $CO_2$ and $O_2$ adsorption can be considered as instantaneous due to the high temperature and $O_2$ concentration. Under these extreme conditions, the resulting CO production and diffusion contribute to trigger the gas-phase combustion with $O_2$. During the competition between the heterogeneous and homogeneous reaction, the reaction zone is adjacent to the reactive wall. Carbon depletion leads to a propagation of the reaction front with a decreasing intensity toward areas that undergo less reactant depletion.

3.2.5. A priori mass and energy fluxes evaluation

In the case of multi-component flows, the diffusive velocity flux $\vec{J}_k$ can be decomposed into several terms that consist in mole fraction, pressure and temperature ($Soret$) gradients, whereas the heat flux $\vec{q}$ includes temperature gradient, enthalpy transport through species diffusion and the $Dufour$ term. In our code, only mole fraction gradients are considered for the diffusive velocity flux through the $Hirschfelder & Curtiss$ while the $Dufour$ term is neglected in the heat flux. Therefore, the impact of the $Hirschfelder & Curtiss$ approximation for the multicomponent diffusion [34, 35] and the importance of the $Dufour$ and $Soret$ terms are assessed by post-processing the DNS data thanks to a complex transport library (EGlib library [43]):

$$\Psi_{\text{EGlib}}^{k} = \sum_{j \in \Omega_{e}} D_{k,j}^{\text{EGlib}} \left( \vec{\nabla} X_j + (X_j - Y_j) \frac{\vec{\nabla} p}{p} \right)$$

$$+ \sum_{j \in \Omega_{e}} D_{k,j}^{\text{EGlib}} \left( \chi_j \frac{\vec{\nabla} T}{T} \right)$$

Soret term

$$\Psi_{\text{EGlib}}^{k}$$

Diffusion velocity due to species and pressure gradients
\[
q^{EGlib} = -\lambda^{EGlib} \nabla T + \rho \sum_{k \in \Omega_c} h_{x,k} Y_k \vec{V}_k^{EGlib}
\]

sensible enthalpy diffusion

\[
+ \rho \sum_{k \in \Omega_c} \lambda_k^{EGlib} \vec{V}_k^{EGlib}
\]

Dufour term

\[
D_{k,j}^{EGlib}
\]

represents the binary diffusion matrix, \( \lambda_{k,j}^{EGlib} \) correspond to the thermal diffusion ratios and \( \lambda^{EGlib} \) refers to the \textit{a priori} assessed thermal conductivity coefficient. Table 6 and Table 7 summarize the differences between the modeling of our present simulation and the \textit{a priori} assessed terms for heat and mass diffusion. Since the studied case is 1D, only components in the \( x \) direction (Fig. (5)) are plotted. Fig. (16) shows the comparisons of the spatial profiles of the diffusion mass flux (\textit{Soret} effect not included) and the \textit{Soret} diffusion term for each gas-phase species during both reaction phases, i.e, wall-attached reaction zone phase and gas-phase reactions phase respectively at \( \tau = 0.65 \) and \( \tau = 1.65 \). Negative mass fluxes tend to compensate for low concentrations by diffusing toward the wall, whereas positive fluxes are due to species diffusing toward the outlet. Note that for all species the present mass diffusion model based on \textit{Hirschfelder & Curtiss} approximation is always underestimated compared to the \textit{a priori} EGLIB based mass diffusion. The pressure gradient being negligible in our case, this overestimation can be attributed to the hypothesis of constant \textit{Schmidt} numbers used by our code to estimate non-binary mass diffusion coefficients. The relative error is around 5 percent which is fairly acceptable. Concerning the \textit{Soret} effect which is not taken into account by our model, \textit{a priori} calculations show that for \( CO_2 \) and \( O_2 \) species, the corresponding term is very small compared to mass diffusion in our case as shown by Figs. (16a, 16b, 16c, 16d) while it is negligible for \( CO \) species as given in Figs. (16e, 16f). As a result, species concentration gradients play in our case a much more important role for species diffusion than the temperature gradient.

Heat fluxes linked to \textit{Fourier}, sensible enthalpy diffusion and \textit{Dufour} diffusion are also calculated \textit{a priori} thanks to the EGLIB library fed by the DNS data obtained under the \textit{Hirschfelder & Curtiss} approximation. The estimation of the heat flux given by the EGLIB library enables to estimate the error related to the \textit{Hirschfelder & Curtiss} approximation for species diffusion, moreover, the assumption of constant \textit{Prandtl} numbers is made in our code to calculate the thermal conduction coefficient \( \lambda \) appearing in the \textit{Fourier} diffusion term while the \textit{Dufour} effect is not taken into account during the calculations. Hence, this section aims at verifying its importance.

Fig. (17) compare the spatial profiles of sensible enthalpy diffusion, \textit{Fourier} and \textit{Dufour} diffusion for both reaction phases, i.e, wall-attached reaction phase and gas-phase reaction phase at \( \tau = 0.65 \) and \( \tau = 1.65 \). Regarding the sensible enthalpy transport through species diffusion, our DNS logically overestimates it since it overpredicts already the mass diffusion flux. The overestimation of the \textit{Fourier} diffusion heat flux can be
attributed to the hypothesis of constant Prandtl number used by our code to estimate thermal conductivity coefficient. Though, the relative error is around 6.5 percent which is satisfying. This analysis points out also that the Fourier and sensible enthalpy diffusion are of the same order of magnitude. Concerning the Dufour effect, which is not taken into account in our code, a posteriori calculations show in our case that it is negligible as given in Fig. (17). The system energy is thus much more sensitive to temperature gradients and species enthalpy diffusion than species diffusion.

3.2.6. Parametric study

A parametric study is carried out by varying initial O$_2$ and CO$_2$ concentrations. In order to characterize the effect of the initial ratio (X$_{O_2}$/X$_{CO_2}$) $|_{t=0}$ on the combustion mode switch. Moreover, the influence on surface and gas-phase chemistry competition, i.e, $I_k(\tau) = 0.5$ is studied. Fig. (18) shows the evolution of $\tau_{switch}$ and $\tau_{1/2,k}$ versus the molar ratio (X$_{O_2}$/X$_{CO_2}$) $|_{t=0}$ for each species.

The parameters $\tau_{switch}$ and $\tau_{1/2,k}$ stand respectively for the physical time corresponding to the combustion mode switch and the physical time below which $I_k \leq 0.5$ both normalized by their values for the reference case corresponding initially to X$_{O_2}$ = 0.7, i.e. (X$_{O_2}$/X$_{CO_2}$) $|_{t=0}$= 2.33. As O$_2$ initial concentration increases, exothermic surface oxidation is enhanced which increases carbon combustion rate thus resulting in a decrease of $\tau_{switch}$ as the (X$_{O_2}$/X$_{CO_2}$) $|_{t=0}$ ratio increases. $\tau_{1/2,O_2}$ then rises because richer O$_2$ mixtures favour surface oxidation reactions (Table 4, reactions S1 to S3) which in turn enhances the surface rate $\dot{s}_{O_2}$. As a consequence, the competition between O$_2$ adsorption by surface carbon and its consumption by the gas-phase combustion is extended to a wider time range.

Table 6: Diffusion mass flux modeling : Present simulation versus a priori assessment

<table>
<thead>
<tr>
<th>Model</th>
<th>$\vec{\nabla}X_k$, $\vec{\nabla}p$</th>
<th>Soret</th>
</tr>
</thead>
<tbody>
<tr>
<td>Present simulation</td>
<td>$Hirschfelder &amp; Curtiss$ approximation</td>
<td>Neglected</td>
</tr>
<tr>
<td>$a priori$ assessment</td>
<td>$\sum_{j \in \Omega_c} D^{EGlib}_{k,j} \left( \vec{\nabla}X_j + (X_j - Y_j) \frac{\vec{\nabla}p}{p} \right)$</td>
<td>$\sum_{j \in \Omega_c} D^{EGlib}_{k,j} \left( X_j \frac{\vec{\nabla}T}{T} \right)$</td>
</tr>
</tbody>
</table>

Table 7: Heat flux modeling : Present simulation versus a priori assessment

<table>
<thead>
<tr>
<th>Model</th>
<th>Fourier</th>
<th>Sensible enthalpy diffusion</th>
<th>Dufour</th>
</tr>
</thead>
<tbody>
<tr>
<td>Present simulation</td>
<td>$-\lambda \vec{\nabla}T$</td>
<td>$\rho \sum_{k \in \Omega_c} h_{s,k} Y_k \vec{V}_k$</td>
<td>Neglected</td>
</tr>
<tr>
<td>$a priori$ assessment</td>
<td>$-\lambda^{EGlib} \vec{\nabla}T$</td>
<td>$\rho \sum_{k \in \Omega_c} h_{s,k} Y_k \vec{V}_{k^{EGlib}}$</td>
<td>$\rho \sum_{k \in \Omega_c} \lambda^{EGlib} \vec{V}_{k^{EGlib}}$</td>
</tr>
</tbody>
</table>
Figure 16: Comparison of *a priori* assessment versus the present simulation: Soret term versus diffusion induced by molar fraction and pressure gradients at \( \tau = 0.65 \) (left) and \( \tau = 1.65 \) (right).
3.2.7. Grid convergence

A global criterion was written in order to estimate the proper grid resolution in the near-wall region. The criterion gave a cell height of \( h_{\text{min}} = 15\mu m \) and was established according to the following equality:

\[
dt_{\text{schem, min}} \approx dt_{Fo, min}\text{. With } dt_{Fo, min} = Fo\left(h_{\text{min}}^2/\nu\right)\text{.}
\]

The resulting minimum cell height is \( h_{\text{min}} = \sqrt{(\nu/Fo)dt_{\text{schem, min}}} \) corresponds to the minimum surface chemistry time step estimated thanks to homogeneous calculations and \( dt_{Fo, min} \) is an a priori estimation of the diffusive minimum time step which is the limiting time step in our case. A constant grid resolution was applied for the diffusive length with a cell height corresponding to \( h_{\text{min}} \) and a mesh coarsening has been carried out between \( x/l_{\text{diff}} = 0.865 \) and the outlet. Nevertheless, a grid convergence was necessary to determine properly the mesh size. The grid convergence was performed for 4 different meshes by solely varying the diffusive length mesh size between \( h_{\text{min}} = 15\mu m \) and \( h_{\text{min}} = 5\mu m \). Fig. (19) shows near-wall gradients of \( CO_2 \) species mass fractions and gas-phase temperature at a time where gradients are strongest, i.e, at \( \tau = 0.955 \). It shows that the grid convergence was reached at \( 7\mu m \). The relative differences between the \( 7\mu m \) grid resolution and the present mesh of \( 10\mu m \) are

Figure 17: Comparison of a priori assessment (thin lines) versus the present simulation (thick lines).

Figure 18: Characteristic times versus initial molar ratio \((X_{O_2}/X_{CO_2}) \mid_{t=0}\). The case \((X_{O_2}/X_{CO_2}) \mid_{t=0}= 2.33\) is taken as a reference.

The resulting minimum cell height is \( h_{\text{min}} = \sqrt{(\nu/Fo)dt_{\text{schem, min}}} \) corresponds to the minimum surface chemistry time step estimated thanks to homogeneous calculations and \( dt_{Fo, min} \) is an a priori estimation of the diffusive minimum time step which is the limiting time step in our case. A constant grid resolution was applied for the diffusive length with a cell height corresponding to \( h_{\text{min}} \) and a mesh coarsening has been carried out between \( x/l_{\text{diff}} = 0.865 \) and the outlet. Nevertheless, a grid convergence was necessary to determine properly the mesh size. The grid convergence was performed for 4 different meshes by solely varying the diffusive length mesh size between \( h_{\text{min}} = 15\mu m \) and \( h_{\text{min}} = 5\mu m \). Fig. (19) shows near-wall gradients of \( CO_2 \) species mass fractions and gas-phase temperature at a time where gradients are strongest, i.e, at \( \tau = 0.955 \). It shows that the grid convergence was reached at \( 7\mu m \). The relative differences between the \( 7\mu m \) grid resolution and the present mesh of \( 10\mu m \) are
4. Conclusions

A boundary condition for the impact of surface reactions on the heat flux at the reactive wall has been developed and applied to Direct Numerical Simulations. An original analytical solution has thus been derived for a flow bounded by two reactive surfaces. A special attention has been given to the numerical setup so that compatibility with a steady-state is achieved. This approach enables to compare analytical solutions with stationary numerical predictions thus validating numerical species boundary condition for the heterogeneous surface reactions.

DNS accounting for multi-species diffusion and micro-kinetics taking into account the Stefan flux were subsequently performed to study carbonaceous wall combustion under oxy-fuel conditions. Transition from endothermic gasification to surface carbon oxidation and gas-phase ignition were evidenced thanks to the analysis of the time-dependent surface and gas-phase species production rates and transient histories of surface species concentrations as well as spatial profiles of gas-phase species, heat release and temperature. The reference case that consists in a quiscient mixture containing 70 percent of O₂ and 30 percent of CO₂ in volume is characterized by two gas-phase reaction zone configurations. The first one corresponds to a wall-adjacent CO-reaction zone maintained by the competition between the CO/O₂ combustion and surface reactions. CO surface desorption contributes to the continous feeding of the reaction zone which in turn produces CO₂ that diffuses to the surface and adsorbs on surface carbon thus providing an additional CO formation pathway. Once carbon has been totally consumed the reaction zone moves toward the outlet.

Figure 19: Grid convergence for four different near wall resolutions at \( \tau = 0.955 \)

around 0.1% only. Therefore, the 10\( \mu m \) resolution was chosen because it is judged fine enough to resolve the stiffest gradients at the reactive wall and the gas-phase reaction zone thickness with reasonable numerical limiting time steps.
while its intensity is progressively dumped. Therefore, there is a strong interaction between the surface and gas-phase kinetics depending on the chemical conditions. Note that this reaction zone occurring at a certain distance of the reactive wall is observed despite the absence of volatiles in the gas-phase.

The impact of the Hirschfelder & Curtiss approximation for the multicomponent diffusion and the importance of the Dufour and Soret terms were assessed by post-processing the DNS data thanks to a complex transport library (EGlib library). A priori post-processing of the DNS showed that species concentration gradients have a much more important contribution to the diffusion velocity than the temperature gradient (Soret effect) whereas the system energy is much more sensitive to temperature gradients (Fourier) and sensible enthalpy diffusion that are of the same order of magnitude. The Dufour effect being also negligible, the Hirschfelder & Curtiss approximation is satisfying since the relative error between the fluxes predicted by our DNS and the ones estimated by the EGLIB library is around 6 percent. The discrepancies can be attributed to the hypothesis of constant Schmidt and Prandtl numbers used to estimate non-binary mass diffusion coefficients and thermal conduction coefficient respectively.

Eventually, the effect of initial concentrations of \( O_2 \) and \( CO_2 \) on characteristic times of gas and surface reactions and times corresponding to the switch between the two gas-phase reactions structures has been assessed. As the \( (X_{O_2}/X_{CO_2}) \) \( |_{t=0} \) ratio increases, the switch between the two modes of gas-phase reactions occurs earlier and the competition between \( CO_2 \) consumption by the reactive surface and its production through gas-phase combustion becomes shorter. In contrast, the surface consumption of \( O_2 \) was seen to compete gas-phase combustion for a wider time range.

Acknowledgement

The authors thank Dr. Guillaume Moula for his contributions to the analytical solution derivation for the species boundary condition validation.
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Appendix A.

Gas-phase governing equations

Multi-species reactive flow equations:

A brief summary of the governing equations of the LES/DNS flow solver AVBP is provided in this section. Please, refer to the AVBP handbook [29] and http://www.cerfacs.fr/avbp/AVBP_V6.X/HANDBOOK/AVBP/handbook.pdf for further details. The AVBP solver uses a conservative finite-volume approach. The following description focuses on a conservative formulation of the unfiltered compressible Navier-Stokes equations used in Direct Numerical simulation (DNS). The set of equations describing the evolution of a compressible flow with gas-phase chemical reactions is written in the following vectorial way:

Momentum:

\[ \forall \ i \in D \ : \ \frac{\partial \rho u_i}{\partial t} + \nabla \cdot (\rho u_i u) = \nabla \cdot \left( \tau_i - \rho \delta_i \right) \quad (A.1) \]

Energy:

\[ \frac{\partial \rho E}{\partial t} + \nabla \cdot (\rho E u) = \dot{\omega}_T - \nabla \cdot \left( \tilde{\tau} - \rho \tilde{\delta} \right) \quad (A.2) \]

Species:

\[ \forall \ k \in \Omega_G : \ \frac{\partial \rho Y_k}{\partial t} + \nabla \cdot (\rho (\tilde{u} + \tilde{V}) Y_k) = \dot{\omega}_k \quad (A.3) \]

\( E \) is the total non-chemical energy defined as follows:

\[ E = \sum_{k \in \Omega_G} h_{s,k} Y_k - (p/\rho) + 1/2 \sum_{j \in D} u_j^2 \quad (A.4) \]

While \( \rho \) is given by:

\[ \rho = \sum_{k \in \Omega_G} \rho_k = \sum_{k \in \Omega_G} \rho Y_k \quad (A.5) \]

\( \rho E \) is transported and the temperature values are tabulated thanks to sensible energy look-up table. The pressure \( p \) is given by the equation of state for ideal gas mixture.

Modeling diffusion of species, heat and momentum:

The diffusion velocities \( V_{k,i} \) are obtained thanks to the Hirschfelder & Curtiss approximation [35]. A correction velocity is introduced so that the compatibility between species and mass conservation is ensured. The heat flux vector \( \tilde{q} \) stands for heat conduction (Fourier term) and heat transport by species diffusion:

\[ \tilde{q} = -\tilde{\tau} + \rho \sum_{k \in \Omega_G} h_{s,k} Y_k \tilde{V}_k = -\tilde{\tau} + \sum_{k \in \Omega_G} h_{s,k} \tilde{J}_k \quad (A.6) \]

With:

\[ J_{k,i} = \rho V_{k,i} Y_k = -\rho \left( D_k W_k \frac{\partial X_k}{\partial x_i} - Y_k V_i^{\text{correc}} \right) \quad (A.7) \]

Transport coefficients:

Temperature dependence is assumed for viscosity through a power law function. Non-binary Gas-phase species mass diffusion coefficients and thermal conductivity noted respectively \( D_k \) and \( \lambda^g \) are determined using constant Schmidt and Prandtl numbers which are considered to be constant in time and space as follows:
Species diffusion through temperature gradients known as Soret effect and the Dufour effect which is the heat flux induced by species mass fraction gradients are not accounted for in the present code [59, 60]. These assumptions that consist in neglecting the Soret and Dufour effects are verified thanks to a priori calculations using the EGlib library in Section 3.2.5.

Gas-phase kinetics:

The rate of production/consumption of the \( k \)th species noted \( \dot{\omega}_k \) corresponds to the sum of rates \( \dot{\omega}_{kj} \) produced by all gas-phase reactions:

\[
\dot{\omega}_k = \sum_{j \in M} \dot{\omega}_{kj} = W_k \sum_{j \in M} (\nu_{kj}' - \nu_{kj}'') Q_j \quad (A.10)
\]

The progress rate \( Q_j \) of reaction \( j \) reads:

\[
Q_j = k_{fj} \prod_{k \in \Omega_G} (\rho Y_k/W_k)^{\nu_{ki}} - k_{rj} \prod_{k \in \Omega_G} (\rho Y_k/W_k)^{\nu_{ki}} \quad (A.11)
\]

\( k_{fj} \) and \( k_{rj} \) are forward and reverse rate constants. The former is modeled using Arrhenius law whereas the latter is computed from the forward rates and the equilibrium constants.

The gas phase heat release rate reads:

\[
\dot{\omega}_T = - \sum_{k \in \Omega_G} \Delta h_{f,k}^\circ \dot{\omega}_k \quad (A.12)
\]

Appendix B.

In this appendix, the approach enabling to obtain mass and energy balance is detailed. The mass balance allows to obtain the expression of the Stefan velocity and the gas-phase species normal gradients accounting for surface reactions whereas the energy balance at the interface gives the normal heat flux at the reactive boundary. Flux balances at the reactive interface are obtained by integrating the transport equations over the control volume sketched in Fig. (B.20):

Figure B.20: sketch of the control volume at the reactive interface

\( g \) is the index standing for gas-phase scalars whereas \( s \) stands for solid-phase scalars. \( w \) is the index standing for variables at the wall, \( \vec{n}^g \) is the vector normal to the interface and directed toward the gas-phase, \( \vec{n}^s \) is vector normal to the interface and directed toward the solid, \( S \) is the elementary reactive surface which is also the fluid/structure interface, \( V \) refers to the control volume and \( \Sigma \) stands for the surface envelope of the control volume.
Appendix B.1. Mass balance at a reactive interface

The balance is obtained as follows: The species conservation Eq. (A.3) is integrated over the control volume \( V \) [26, 61] and the Green-Ostrogradski theorem is applied:

\[
\int_{V} \left( \frac{\partial \rho Y_k}{\partial t} \right) dV + \int_{V} -\dot{\omega}_k dV = \int_{\Sigma} (- (\rho \vec{V}_k Y_k) \cdot \vec{n}) d\Sigma
\]

\[
+ \int_{\Sigma} - (\rho Y_k \vec{u} \cdot \vec{n}) d\Sigma
\]

(B.1)

Eq. (B.1) represents the integral form of the transport equation given in Eq. (A.3). If the volume tends to zero (\( V \to 0 \)), the control volume surface \( \Sigma \) tends to the reactive interface \( S \) (\( \Sigma \to S \)) [61]. Since no mass accumulation occurs in a volume that tends to zero, the first term of the L.H.S of Eq. (B.1) becomes:

\[
\lim_{V \to 0} \int_{V} \left( \frac{\partial \rho Y_k}{\partial t} \right) dV = 0 \quad (B.2)
\]

When surface chemistry occurs at the wall, one can decompose the gas-phase production rates into two terms, the first one would be non-zero in the gas-phase and null at the wall whereas the second one would equal zero in the gas-phase and non-zero at the solid/fluid interface as follows:

\[
\dot{\omega}_k = \dot{\omega}_{k,w} + \dot{s}_k \delta(y) \quad (B.3)
\]

where \( \dot{\omega}_{k,w} \) and \( \dot{s}_k \) represent respectively, for each \( k^{th} \) gas-phase species, the gas-phase rate of production at the wall and the rate of production due to surface reactions per unit area. The dirac function integration gives:

\[
\int_{-\infty}^{+\infty} \delta(y) dy = 1 \quad (B.4)
\]

When the volume tends to zero, one can write:

\[
\lim_{V \to 0} \int_{V} \dot{\omega}_k dV = \int_{S} \dot{s}_k dS \quad (B.5)
\]

According to the Dirac function definition above:

\[
\lim_{V \to 0} \int_{V} \dot{\omega}_k dV = 0 \quad (B.6)
\]

Eventually, the volume integral of the L.H.S of Eq. (B.1) becomes:

\[
\lim_{V \to 0} \int_{V} \left( \frac{\partial \rho Y_k}{\partial t} \right) dV + \int_{V} -\dot{\omega}_k dV = - \int_{S} \dot{s}_k dS \quad (B.7)
\]

Since there are no convection and diffusion in the solid and the boundary of the control volume is static \( (\vec{J}^s_k = \vec{u}^s = 0) \), one obtains for the R.H.S of Eq. (B.1):

\[
\lim_{\Sigma \to S} \left\{ \int_{\Sigma} (- (\rho \vec{V}_k Y_k) \cdot \vec{n}) d\Sigma + \int_{\Sigma} (\rho Y_k (- \vec{u} \cdot \vec{n})) d\Sigma \right\} = \int_{S} (- (\rho Y_k \vec{V}_{k,w} Y_{k,w}) \cdot \vec{n}) dS + \int_{S} (\rho Y_k \vec{u} \cdot \vec{n}) dS \quad (B.8)
\]

As a result, Eqs. (B.7, B.8) give:

34
\[- \int_{S} \dot{s}_k dS = - \int_{S} \left( \rho_w V^r_{k,w} Y_{k,w} \right) \cdot \vec{n} \cdot dS \]
\[\text{rate of production by surface reaction} \]
\[+ \int_{S} \rho_w Y_{k,w} (\vec{n} \cdot \vec{v}^r_{k,w}) dS \]
\[\text{Mass flux diffusion} \]
\[\int_{S} \rho_w Y_{k,w} (\vec{n} \cdot \vec{v}^\ast_{k,w}) dS \]
\[\text{Mass flux convection} \]
\[\text{(B.9)} \]

Where:
\[\begin{cases} 
\vec{V}^r_{k,w} \cdot \vec{n}^\ast = V_{k,n,w} \\
\vec{u} \cdot \vec{n}^\ast = U_{Stefan} 
\end{cases} \]

The mass balance at the reactive interface (Eq. (B.9)) reads then:
\[(\rho_w V_{k,n,w} Y_{k,w}) + (\rho_w U_{Stefan} Y_{k,w}) = \dot{s}_k \quad (B.10)\]

Eq. (B.10) is consistent with the mass balance near a reactive wall in [20, 26, 37, 62]. Summing Eq. (B.10) over all the gas-phase species and considering the mass conservation, an expression for the wall-normal Stefan velocity is obtained:
\[U_{Stefan} = \frac{1}{\rho_w} \left( \sum_{\Omega^k \in \Omega} \dot{s}_k \right) \quad (B.11)\]

Appendix B.1.1. Gas-phase species mass fraction gradient correction

This section shows how the prescribed gradient is written. First, let us consider the mass balance at the interface given by Eq. (B.10) demonstrated in the previous Section Appendix B.1:
\[(\rho_w V_{k,n,w} Y_{k,w}) + (\rho_w U_{Stefan} Y_{k,w}) = \dot{s}_k \quad (B.12)\]

By isolating the species diffusion flux, one obtains:
\[(\rho_w V_{k,n,w} Y_{k,w}) = \dot{s}_k - (\rho_w U_{Stefan} Y_{k,w}) \quad (B.13)\]

The species mass flux is written as in Eq. (A.7) according to the Hirschfelder & Curtiss approximation:
\[J_{k,i} = \rho V_{k,i} Y_{k,w} = - \rho \left( \frac{Dk}{W} \frac{\partial X_k}{\partial x_i} - Y_{k,w} \hat{V}_{w} \right) \quad (B.14)\]

By multiplying Eq. (B.14) by the interface normal \(\vec{n}^\ast\), one obtains:
\[\hat{J}_{k,w} \cdot \vec{n}^\ast = (\rho_w \vec{V}_{k,w} \cdot \vec{n}^\ast Y_{k,w}) = \]
\[- \rho \left( \frac{Dk}{W} \frac{\hat{V}_{w} Y_{k,w}}{W} \cdot \vec{n}^\ast - Y_{k,w} \hat{V}_{w} \right) \quad \text{(B.15)}\]

with \(\hat{V}_{w} \cdot \vec{n}^\ast\) and \(\hat{V}_{w} Y_{k,w} \cdot \vec{n}^\ast\) corresponding respectively to the wall-orthogonal correction velocity and species molar fractions:
\[\hat{V}_{w} \cdot \vec{n}^\ast = \sum_{l \in \Omega^k} D_k \frac{W_l}{W_w} \hat{V}_{w} Y_{l,w} \cdot \vec{n}^\ast \quad (B.16)\]
\[\hat{V}_{w} Y_{k,w} \cdot \vec{n}^\ast = \frac{W_w}{W_k} \hat{V}_{w} Y_{k,w} \cdot \vec{n}^\ast - \frac{W^2_k}{W_k} Y_{k,w} \sum_{l \in \Omega^k} \frac{1}{W_l} \hat{V}_{w} Y_{l,w} \cdot \vec{n}^\ast \quad (B.17)\]

Inserting Eqs. (B.11, B.16, B.15) into the mass balance at the interface given by Eq. (B.13), it becomes possible to relate the species mass fraction at the wall \(Y_{k,w}\) to its
normal gradient at the boundary surface by isolating the latter
\[
\Phi^{BC}_{k,w}, \bar{r}^w = \frac{Y_{k,w}}{D_k \rho_w} \sum_{\Omega_{k,w}} \hat{s}_k + \frac{Y_{k,w}}{D_k} \phi^{\text{correct}}_w, \bar{r}^w
\]
\[
+ Y_{k,w} \rho_w \left( \sum_{\Omega_{k,w}} \frac{1}{W_l} \bar{r} Y^{k,w} \cdot \bar{r}^w \right) - \frac{\hat{s}_k}{\rho_w D_k}
\]
(B.18)

Appendix B.2. Energy balance at the reactive interface and heat flux correction

An energy balance at the reactive interface has been developed in order to prescribe the proper temperature gradient at the wall. The same approach used for the mass balance is applied to the total non-chemical energy given in Eq. (A.2). By integrating over the control volume, applying the Green-Ostogradski theorem to Eq. (A.2) and tending the volume to zero, one obtains:

\[
\lim_{V \to 0} \left[ \int_V \frac{\partial \rho E}{\partial t} dV + \int_V -\omega^T dV \right] = \lim_{V \to 0} \left[ \int_{\Sigma} \left( (\bar{r} \vec{u} - \hat{q}) \cdot \bar{n} \right) d\Sigma + \int_{\Sigma} \left( -p \vec{E} \vec{u} \cdot \bar{n} \right) d\Sigma \right]
\]
(B.19)

As shown in Eqs. (B.2-B.5) the first term of the L.H.S equals zero and the second term corresponds to the heat-release due surface reactions. Regarding the R.H.S, the same approach as in Eq. (B.8) is used thus giving the heat balance at the reactive interface:

\[
-\dot{S}_T = (\bar{r} \vec{u}^w)_{n,w} - q_{n,w}^w + \rho_w E_s U_{\text{Stefan}} + \bar{q}^w_{n,w}
\]
(B.20)

With the following notations:

\[
\begin{align*}
\int_{\Sigma} \dot{S}_T dS &= \lim_{V \to 0} \left( \int_V \omega_T dV \right) \\
(\bar{r} \vec{u})_{n,w} - q_{n,w}^w &= (\bar{r} \vec{u}^w_{n,w} - \bar{q}_n^w) \cdot \bar{r}^w \\
U_{\text{Stefan}} &= \bar{u}_w \cdot \bar{r}^w \\
q_{n,w}^w &= (-q_{n,w}^w) \cdot \bar{r}^w
\end{align*}
\]

If one considers that there is no species diffusion in the solid phase so that only solid thermal diffusion remains and that the surface heat release is due to both solid-phase and gas-phase species:

\[
\begin{align*}
q_{n,w}^w &= -\lambda^s \bar{r} T^w_s \cdot \bar{r}^w \\
\dot{S}_T &= - \left( \sum_{k \in \Omega_k} h_k \hat{s}_k + \sum_{k \in \Omega_k} \Delta h^s_{f,k} \hat{s}_k \right)
\end{align*}
\]

By introducing the terms above into Eq. (B.20), and replacing \( E \) by its expression one obtains:

\[
\begin{align*}
\dot{S}_T &= (\lambda^s \bar{r} T^w_s - \lambda^s \bar{r} T^w_s) \cdot \bar{r}^w \\
&+ \sum_{k \in \Omega_k} (\rho_w (h_{s,k,w} V_{k,n,w} Y_{k,w}) + \rho_w U_{\text{Stefan}} (h_{s,k,w} Y_{k,w})) \\
&+ \left( \sum_{k \in \Omega_k} h_{k,w} \hat{s}_k + \sum_{k \in \Omega_k} \Delta h^s_{f,k} \hat{s}_k \right) \\
&- \left( (\bar{r} \vec{u}^w_{n,w} + p U_{\text{Stefan}}) \right) \\
&+ \frac{\bar{m}}{2} \sum_{j \in D} \bar{u}_{j,w}^2
\end{align*}
\]
(B.21)

The first term in the R.H.S of Equation Eq. (B.21) can be simplified by considering the mass balance at the
interface given in Eq. (B.10). Indeed, if one multiplies Eq. (B.10) by the species sensible enthalpy $h_{s,k}$ and sums all over the gas-phase species, the resulting balance becomes:

$$\sum_{k \in \Omega_{\text{g}}} \left( \rho_w (h_{s,k,w} V_{k,n,w} Y_{k,w}) + \rho_w U_{\text{Stefan}} (h_{s,k,w} Y_{k,w}) \right) = \sum_{k \in \Omega_{\text{g}}} h_{s,k,w} \dot{s}_k$$

(B.22)

As a result, the first and second terms in the R.H.S of equation Eq. (B.21) are simplified as follows:

$$\sum_{k \in \Omega_{\text{g}}} \left( \rho_w (h_{s,k,w} V_{k,n,w} Y_{k,w}) + \rho_w U_{\text{Stefan}} (h_{s,k,w} Y_{k,w}) \right) + \left( \sum_{k \in \Omega_{\text{g}}} h_{s,k,w} \dot{s}_k + \sum_{k \in \Omega_{\text{g}}} \Delta h_{s,k} \dot{s}_k \right) = \sum_{k \in \Omega_{\text{g}}} h_{s,k,w} \dot{s}_k$$

(B.23)

and the third term becomes:

$$\left( (\tilde{t}^e \tilde{t}^e)_{n,w} + p U_{\text{Stefan}} \right) = \left( (\tilde{t}^e \tilde{t}^e)_{n,w} + p \dot{\vec{u}} \cdot \vec{n} \right) = (\tilde{\vec{t}}^e \tilde{t}^e)_{n,w}$$

(B.24)

Eventually, injecting Eqs. (B.23, B.24) into Eq. (B.21), and isolating the gas-phase wall-normal thermal diffusion flux, yields:

$$\lambda^e \nabla T_w^e \cdot \vec{n} = \sum_{k \in \Omega_{\text{g}}} h_{s,k,w} \dot{s}_k - (\tilde{t}^e \tilde{t}^e)_{n,w}$$

$$+ \frac{\dot{m}}{2} \sum_{j \in D} u_{j,w}^2 + \lambda^e \nabla T_w^e \cdot \vec{n}$$

(B.25)

Eq. (B.25) constitutes the heat flux balance that accounts for the effect of surface chemistry.

### Appendix B.2.1. Heat flux correction

The heat flux balance of Eq. (B.25) enables to write the heat flux to be prescribed at the boundary condition $q_w^{BC} \cdot \vec{n}$ by isolating in Eq. (B.25) the heat flux $q_j$ given in Eq. (A.6). Under the Hirschfelder & Curtiss approximation $q_w^{BC} \cdot \vec{n}$ reads:

$$q_w^{BC} \cdot \vec{n} = - \sum_{k \in \Omega_{\text{g}}} h_k \dot{s}_k$$

Surface heat release due to surface species

$$- \sum_{k \in \Omega_{\text{g}}} \Delta h_{s,k} \dot{s}_k$$

Surface heat release due to gas-phase species

$$- \rho_w \sum_{k \in \Omega_{\text{g}}} U_{\text{Stefan}} h_{s,k,w} Y_{k,w}$$

Sensible enthalpy transport by Stefan flux

$$- \frac{\dot{m}}{2} \sum_{j \in D} u_{j,w}^2 + (\tilde{\vec{t}}^e \tilde{t}^e)_{n,w} - \lambda^e \nabla T_w^e \cdot \vec{n}$$

Kinetic energy Viscous heat Solid heat diffusion

(B.26)

### Appendix C.

Making use of the assumption of steady developed non-reacting flow, one can write mass/species/momentum conservation equations as:

$$\frac{\partial (\rho v)}{\partial y} = 0$$

(C.1)

$$\frac{\partial}{\partial y} \left( \rho (V_{k,y}) Y_k \right) = 0 \quad k \in \{1, 2\}$$

(C.2)

$$\frac{\partial (\rho uv)}{\partial y} = \frac{\partial \tau_{xy}}{\partial y} + S$$

(C.3)

$$\frac{p}{\rho} = \frac{R}{W} T$$

(C.4)
The momentum source term $S$ is used for the following numerical and practical reasons:

- Standard Inlet/Outlet boundary conditions imply the estimation of a pressure loss in the streamwise direction to include in the momentum conservation equation which is not an accurate approach for analytical solution derivation and not strictly conservative.

- Periodicity enables momentum conservation by imposing a momentum source thus being strictly conservative.

In the following, the letter $C$, $O$ and $P$ denote the solid wall species (which could be for instance solid carbon), the gaseous oxidizing species and the gaseous species produced by gasification, respectively. The associated stoichiometric coefficients are noted $\nu_C$, $\nu_O$ and $\nu_P$. According to these notations, the following chemical schemes are applied to the surface 1 (gasification):

\[ \nu_C C(s) + \nu_O O \rightarrow \nu_P P \quad \text{(C.5)} \]

and to the surface 2 (lieu of deposition):

\[ \nu_P P \rightarrow \nu_C C(s) + \nu_O O \quad \text{(C.6)} \]

In what follows, variables subscripted by letters $C$, $O$ and $P$ refer to the associated species and subscript numbers 1 and 2 refer to space location, $y = 0$ and $y = h$, respectively. The molar progress rate of reaction 1, $Q_1$, can then be modeled as follows:

\[ Q_1 = \left( \frac{Y_{O,1}}{W_O} \rho_1 \right)^{\nu_O} K_1 = \left( \frac{X_{O,1} p}{RT} \right)^{\nu_O} K_1 \quad \text{(C.7)} \]

where $K_1$ represents the rate of reaction 1 which can be modeled by an Arrhenius law for instance. However, in the present framework it is not necessary to go deeper in the modeling of this coefficient because one assumes a constant temperature implying that $K_1$ can be merely seen as a parameter of the problem. Similarly, one can write the molar progress rate of reaction 2 as:

\[ Q_2 = \left( \frac{Y_{P,2}}{W_P} \rho_2 \right)^{\nu_P} K_2 = \left( \frac{X_{P,2} p}{RT} \right)^{\nu_P} K_2 \quad \text{(C.8)} \]

Moreover, the general formulation of species conservation balance at the reacting surfaces reads:

\[ \left( \rho_j \vec{u} Y_k + \rho \vec{V}_k Y_k \right) \cdot \vec{n} = \dot{s}_k \quad \text{(C.9)} \]

The species conservation balance at surface 1 reads:

\[ \rho_1 v_1 Y_{k,1} + \rho_1 V_{k,1} Y_{k,1} = \dot{s}_{k,1} \quad \text{(C.10)} \]

and at surface 2:

\[ \rho_2 v_2 Y_{k,2} + \rho_2 V_{k,2} Y_{k,2} = -\dot{s}_{k,2} \quad \text{(C.11)} \]

where $\vec{n}$ is the vector normal to the wall and pointing toward the gas-phase. Summing over all the species and making use of the mass conservation constraints $\sum_k Y_k = 1$ and $\sum_k Y_k V_k = 0$, one obtains the two relations $\rho_1 v_1 = \sum_k \dot{s}_{k,1}$ and $\rho_2 v_2 = -\sum_k \dot{s}_{k,2}$. Furthermore, the continuity equation Eq. (C.1) imposes that $\rho(y)v(y)$ is constant (in the forthcoming the notation $\dot{m} = \rho(y)v(y)$ will be used). As a consequence, one finds that $\dot{m} = \sum_k \dot{s}_{k,1} = -\sum_k \dot{s}_{k,2}$ which leads to the relation $Q_1 = Q_2$ (the notation $Q = Q_j$ is thus retained.
for simplification). The two progress rates of the surface reactions are thus identical. The relation between the wall normal mass flow rate and the progress rate of reaction is thus expressed as:

\[ \dot{m} = Q (\nu_P W_P - \nu_O W_O) \]  \hspace{1cm} (C.12)

When the species diffusion due to pressure gradients, temperature gradients (Soret effect) and volume forces are neglected, the binary diffusion between the two species \( O \) and \( P \) can be modeled by the classical relation \[35\]

\[ V_k X_k = -D_{\text{bin}}^{\text{OP}} \frac{\partial Y_k}{\partial y}, \text{ where } D_{\text{bin}}^{\text{OP}} = D_{\text{PO}}^{\text{bin}} \]  \hspace{1cm} (C.13)

is the binary diffusion coefficient. Another formulation is obtained by considering equivalent diffusion coefficients, \( D_k \), that represents the diffusion of species \( k \) into the mixture and expressed as \( D_k = (1 - Y_k)/(\sum_{l=e} X_l/D_{kl}^{\text{bin}}) \). This leads to an equivalent diffusion model which now involves the species molar fractions instead of their mass fractions, \( V_k X_k = -D_k \frac{dX_k}{dy} \). This presents a strong advantage for solving the system of equations. Note also that contrary to the binary coefficient, \( D_k \) is depending on the local concentration of each species which means that \( D_k = D_k(y) \). According to the previous definitions, the following relation stands for the present mixture:

\[ D_O W_O = D_P W_P \]  \hspace{1cm} (C.14)

Moreover, the dynamic viscosity for gases is usually assumed to be solely temperature dependent which means that in the present isothermal framework \( \mu \) is a constant. Since the Schmidt numbers, \( Sc_k = \mu/(\rho D_k) \), are assumed to be constant, we find that \( \rho(y)D_k(y) \) is also constant. This statement allows to simplify the integration of the forthcoming partial differential equations. In order to determine the species concentration profiles, one may concentrate on the variable \( X_O \) and then deduce the concentration of species \( P \) by the relation \( X_P = 1 - X_O \). At this step, it is important to consider molar fractions instead of mass fractions because it leads to a first order partial differential equation with constant coefficients. Indeed, making use of Eq. (C.2), integrating once and identifying with Eq. (C.10), rearranging in terms of molar fractions, and making use of \( \rho(y)D_O(y) = \mu/Sc_O \) yields:

\[ \eta X_O - \frac{\mu}{Sc_O} m \frac{dX_O}{dy} = \frac{\nu_O W_P}{\nu_O W_O - \nu_P W_P} \]  \hspace{1cm} (C.15)

with

\[ \eta = \frac{W_P (\nu_O - \nu_P)}{\nu_O W_O - \nu_P W_P} \]  \hspace{1cm} (C.16)

Since \( \eta = 0 \) for \( \nu_O = \nu_P \), the integration of this differential equation leads to two solutions depending on the values taken by \( \nu_O \) and \( \nu_P \):

\[ X_O(y) = \left( X_{O,1} - \frac{\nu_O}{\nu_O - \nu_P} \right) \exp \left[ \eta \frac{Sc_O \dot{m}}{\mu} y \right] + \frac{\nu_O}{\nu_O - \nu_P} \]  \hspace{1cm} (C.17)

for \( \nu_O \neq \nu_P \)

\[ X_O(y) = -\frac{Sc_O \dot{m}}{\mu} \frac{W_P}{W_O - W_P} y + X_{O,1} \]  \hspace{1cm} (C.18)

for \( \nu_O = \nu_P \)
where $X_{O,1}$ is the molar fraction of species $O$ at surface 1. The value of this constant is determined thanks to the relation between $X_{O,1}$ and $X_{O,2}$ that comes from the identity $Q_1 = Q_2$ which according to Eqs. (C.7, C.8) gives:

$$X_{O,1}^{\nu_{O}} \left( \frac{p}{RT} \right)^{\nu_{O}} \frac{K_1}{K_2} = (1 - X_{O,2})^{\nu_{O}}$$ (C.17)

Furthermore, injecting Eq. (C.7) into the wall normal mass flow rate Eq. (C.12) yields the dependency relation between $\dot{m}$ and $X_{O,1}:

$$\dot{m} = X_{O,1}^{\nu_{O}} \left( \frac{p}{RT} \right)^{\nu_{O}} \frac{K_1}{K_2} (\nu_{P} W_{P} - \nu_{O} W_{O})$$ (C.18)

Hence, Eq. (C.16) expressed for $y = h$ can be injected into Eq. (C.17), and with the use of Eq. (C.18) one obtains an equation in which $X_{O,1}$ is the only unknown. This manipulation allows determining the $X_{O,1}$ constant and Eqs. (C.15-C.18) form the analytical solution for the molar fraction of species $O$. The molar fraction of $P$ is then given by the relation $X_{P}(y) = 1 - X_{O}(y)$. Concerning the determination of the streamwise velocity profile, one introduces the newtonian fluid relation $\tau_{xy} = \mu du/dy$ into Eq. (C.3) which gives

$$\rho v \frac{du}{dy} - \mu \frac{d^2 u}{dy^2} = S.$$ Since $\mu$ is a constant under the approximation of constant temperature, and because $\rho(y)v(y) = \dot{m}$ is a constant according to the continuity Eq. (C.1), the latter differential equation is merely of second order with constant coefficients. Its integration with the boundary conditions $u(0) = u(h) = 0$ leads to the following solution:

$$u(y) = \frac{S}{\dot{m}} \left[ \frac{h}{e^\frac{2h}{\mu} - 1} \left( 1 - e^{\frac{2h}{\mu}} \right) + y \right]$$ (C.19)