Large Eddy Simulation of a 100 kW_{th} Swirling Oxy-coal Furnace

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Abstract

Large Eddy Simulation (LES) has been applied to the swirling 100 kW_{th} OXYCOAL-AC test facility of Aachen University. A set of models to represent devolatilisation, volatile combustion, char combustion and radiation for oxy-coal combustion in an LES framework has been implemented and tested. A qualitative analysis of the flow behaviour and the overall coal combustion processes occurring within the furnace was made. The LES results for the flow field were compared to axial and tangential mean velocity measurements, showing good agreement, particularly in the upstream regions of the flame. The LES results were also compared to oxygen concentrations and gas temperature. Overall good agreement was observed in the upstream central regions of the flame, whilst downstream the LES overestimated the combustion rates. The results show the potential of using LES for more complex oxy-coal combustion burners and opens the way for applications to industrial furnaces.

Keywords: Large Eddy Simulation, Coal Combustion, Oxy-coal, Turbulent Combustion
1. Introduction

Electricity production from coal-fired power plants is expected to keep on playing a dominant role in the future. Carbon Capture and Storage (CCS) can drastically reduce emissions from coal-fired power plants, but leads to lower plant efficiency and increased overall costs. Coal combustion with pure oxygen ($O_2$) and recirculated exhaust gases facilitates the separation of carbon dioxide ($CO_2$) from the exhaust gases. The main disadvantages of oxy-coal combustion are the capital costs and the loss in energy efficiency related to the pure $O_2$ production. Maximising boiler efficiency is thus essential to regain the efficiency lost in the CCS plant, which necessitates an in-depth understanding of the chemical and physical mechanisms involved in oxy-coal combustion.

Experimental measurements inside a Pulverised Coal Combustion (PCC) boiler are difficult and expensive due to poor optical access, making simulations the primary source of detailed information. To date Reynolds-Averaged Navier Stokes (RANS) remains the preferred way for getting in-situ information from within the flame, [1, 2, 3, 4] as it is relatively cheap. Predictions based on Large Eddy Simulations (LES) promise to be more accurate than RANS, and with the rapid advancement of High Performance Computing have recently gained more attention in the PCC research community. Kurose and Makino [5] performed the first LES of a hypothetical solid fuel flame, where the fuel was modelled as pure methane and the simulation results were presented without any comparison to experimental data. Yamamoto et al. [6] performed a LES of a pre-heated pulverised coal flame. Edge et al. [7] and Gherebaghi et al. [8] carried out the first LES of a 1 MWth scale test facility. Both studies showed that LES can provide more detailed information than a RANS simulation. Recently, Franchetti et al. [9] and Stein et al. [10] presented LES of a pulverised coal jet flame with good agreement with experimental data. Pedel et al. [11] also performed an LES of the pulverised coal jet flame. Their work combined LES with the direct quadrature method of moments (DQ MOM) to predict the flame ignition mechanisms. Finally, Rabacal et al. [12] performed a simulation of a large scale laboratory furnace comparing their LES results with experimental measurements of species concentrations and temperature.

The combustion of pulverised coal in an $O_2/CO_2$ atmosphere changes from air combustion due to the different thermo-physical properties of the gas mixture. The density and heat capacity of the gaseous mixture is higher
for oxy-PCC due to the higher molecular weight and heat capacity of \( \text{CO}_2 \) compared to nitrogen (\( \text{N}_2 \)). Moreover, in oxy-PCC, radiative heat transfer will be a priori stronger as \( \text{CO}_2 \) has higher radiative emission power than \( \text{N}_2 \). The higher \( \text{CO}_2 \) concentrations in the gas mixture will also affect the char reaction mechanisms.

The purpose of this study is to apply LES to an oxy-coal swirl burner. The 100 kW\( _{th} \) OXYCOAL-AC test facility at the Aachen University [13, 14] is an ideal test case thanks to the relatively large amount of quantitative experimental data available, and the fact that it is in the heart of the new large-scale research program “Transregio 129 Oxyflame” of Aachen, Bochum and Darmstadt Universities [www.oxyflame.de]. The OXYCOAL-AC burner has been previously modelled in a RANS framework [14, 15]. Chen et al. [16] performed a LES with RANS-like symmetry conditions, where they simulated only a section of the burner and then transposed the results to the other sections using periodic boundary conditions, which made the simulations affordable but suppresses some large-scale turbulent modes, and under-predicts the turbulent stresses on the periodic boundaries.

The present simulations were performed using the PsiPhi code, which has been previously used to simulate a laboratory-scale pulverised coal jet flame [9, 10] and a large scale laboratory furnace [12]. The present simulation extends our proven modelling approach from previous work to a swirled oxyfuel flame that is likely to become the benchmark for oxyfuel combustion, due to the Aachen flames role in the “Oxyflame” [www.oxyflame.de] project. The results of our simulations permit a first assessment of how well the developed modelling framework can be transferred to oxy-combustion, in a system that is well suited for validation due to its detailed velocity measurements. At the same time, the LES provides valuable insights into the instantaneous velocity, scalar and particle fields and their statistics, which have not been available for this flame before - neither from the experiments or from the RANS simulations.

2. Experimental set-up

A schematic of the furnace and inner quarl is shown in Fig. 1, and a more detailed description is given by Toporov et al. [14]. The furnace is cylindrical with an inner diameter of 0.4 m and a vertical height of 2.1 m. The burner consists of four inlets, where the \( \text{O}_2/\text{CO}_2 \) mixture enters. The coal is injected together with the primary gas stream through an annular orifice. The highly
swirled (swirl number 1.2) secondary gas stream is injected through an annulus surrounding the primary stream. For scavenging purposes a tertiary gas stream is injected at very low flow rates. Finally a heated staging gas stream is injected at the outer diameter of the furnace, right at the wall. The purpose of the staging stream is to provide the necessary heat to compensate the higher heat capacity of the gas mixture and to reduce the local stoichiometry of the burner quarl. The flow rates, temperatures and gas compositions of the streams are summarised in Table 1. The furnace is fired with pre-dried Rhenish lignite; its proximate analysis (PA) and ultimate analysis (UA) are shown in Table 2.

The Aachen group [13, 14] measured the particle velocity using a Laser Doppler Anemometer (LDA), measuring the velocity of the pulverised coal particles, so that no velocity data is available for the staging stream. Species concentration were measured with a water-cooled suction probe. The gas temperature was obtained using a traversable suction pyrometer. The experimental paper [14] provided data for the particle size distribution, which is composed of 28 classes ranging from 0.9 - 123 microns. The same particle size distribution provided by the experimental paper [14] was retained in the simulations.

<table>
<thead>
<tr>
<th>Table 1: Experimental Conditions [14].</th>
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<tbody>
<tr>
<td>$\dot{m}$</td>
</tr>
<tr>
<td>[kg/h]</td>
</tr>
<tr>
<td>Coal</td>
</tr>
<tr>
<td>Primary</td>
</tr>
<tr>
<td>Secondary</td>
</tr>
<tr>
<td>Tertiary</td>
</tr>
<tr>
<td>Staging</td>
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<table>
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<tr>
<th>Table 2: Rhenish lignite PA and UA [14].</th>
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<tbody>
<tr>
<td>PA</td>
</tr>
<tr>
<td>Moisture</td>
</tr>
<tr>
<td>Ash</td>
</tr>
<tr>
<td>Volatile Matter</td>
</tr>
<tr>
<td>Fixed Carbon</td>
</tr>
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</table>
3. Numerical Modelling

3.1. Gas Phase

The simulations were performed using the PsiPhi code, which has been validated extensively for gas combustion [17, 18], coal combustion [9, 12] and spray combustion [19]. The generalised Favre filtered governing equations were solved for mass, momentum, species mass fraction and enthalpy. In the momentum equation, the viscous stresses were modelled with Smagorinsky’s classical turbulent eddy viscosity model [20]. The transported species include: oxygen ($O_2$), carbon dioxide ($CO_2$), carbon monoxide ($CO$), water vapour ($H_2O$) and the volatile gases ($VG$), modelled as a single postulated substance ($C\alpha H\beta O\gamma$).

The convective fluxes are approximated using a Central Differencing Scheme for momentum and a Total Variation Diminishing scheme for the scalars. Time integration is performed using a third order explicit low storage Runge-Kutta scheme.
3.2. Solid Phase

A Lagrangian approach is used to treat the dispersed phase, which involves tracking the coal particles in time and space. The spray-pdf approach by Jones et al. [21, 22] is used, which consists of solving a set of differential equations for the trajectory of the particles and evolution of the particle properties. The position and acceleration of a stochastic particle is obtained from:

\[ dx_p = v_p dt \]
\[ dv_p = \frac{\tilde{u}_f - v_p}{\tau_p} dt + \sqrt{C_o \frac{k_{sgs}}{\tau_t}} dW \]

where \( v_p \) and \( \tilde{u}_f \) are the particle and filtered gas velocity respectively, and \( \tau_p^{-1} = 3/4 (\tilde{\rho}_g C_D)/(\rho_p d_p) \) is the particle relaxation time. The drag coefficient \( C_D \) is evaluated using the Yuen-Chen empirical relationship [23]. The first term on the right hand side (RHS) of Eq. (2) represents the influence of the resolved scales of the carrier gas on the particles. The second term represents the influence of the sub-grid scales on the particle motion. The dispersion constant \( C_o \) is set to unity [22]. The timescale \( \tau_t = \tau_p (\tau_p k_{sgs}/\Delta)^{0.6} \) represents the rate of interaction between the particle and gas-phase turbulence and \( k_{sgs} = 2\Delta^2 C_s^{2/3} \tilde{S}_{ij} \tilde{S}_{ij} \) is the unresolved kinetic energy of the gas phase, which can be obtained by an equilibrium assumption for the sub grid turbulence [21, 22, 24], where \( \tilde{S}_{ij} \) is the strain rate tensor. Finally, \( dW \) represents the incremental Wiener process.

The rate of change of temperature of a particle is determined from:

\[ \frac{dT_p}{dt} = \frac{Nu}{3Pr} \frac{c_{p,f}}{c_{p,p}} \left( T_g - T_p \right) \]  
\[ - \frac{m_{ev} h_{dev}}{m_p c_{p,p}} + \frac{\dot{Q}_{char}}{m_p c_{p,p}} + \frac{\dot{Q}_{rad}}{m_p c_{p,p}} \]  

The first term on the RHS of Eq. (3) represents the heat exchange with the gas phase due to convection and conduction, where \( c_{p,f} \) is the specific heat of the gas mixture and \( c_{p,p} = 1100 \text{ J/kgK} \) is the specific heat of the coal particle, which was assumed constant for this work. The Prandtl number \( Pr \) was set to 0.6, \( Nu \) denotes the Nusselt number calculated by the Ranz-Marshall correlation [25], and \( \tau_m = \rho_p d_p^2/(18 \mu_g) \) is the diffusion relaxation time. The second term represents the heat loss due to the volatiles being released from the particle, where \( h_{dev} \) is the latent heat of evaporation of the
volatiles. The first two terms on the RHS of Eq. (3) are based on the heat transfer equation of a liquid spray [24]. Finally, $\dot{Q}_{\text{char}}$ represents the heat exchange between the gas and particle due to char combustion, and $\dot{Q}_{\text{rad}}$ represents the heat exchange of the particle due to radiation, as explained in section 3.6.

3.3. Two way coupling terms

A source term $\bar{S} = 1/\Delta^3 \sum_{i=1}^{P} S^p$ accounts for the contribution of the dispersed phase on the gas phase in each computational cell, where $P$ is the number of particles present in a cell’s volume and $S^p$ is the source term arising from the $p^{th}$ particle. Each coal particle is assumed to be composed of volatile matter, char and ash. Ash acts as an inert substance, and the change in mass of the $p^{th}$ particle is related to the yield of the volatile gases ($dm_{vg}/dt$) and char burning rate ($dm_{char}/dt$), such that:

$$\frac{dm_p}{dt} = \frac{dm_{vg}}{dt} + \frac{dm_{char}}{dt}$$

(4)

The effects of particle momentum exchange with the continuous phase have been studied [26] and were found to have a negligible influence due to the low particle concentration; consequently they have been ignored in this study. Finally, the enthalpy change of the gas phase due to the coal particles burning is given by:

$$\dot{S}_{p,h} = -\dot{Q}_{\text{con}} - \dot{Q}_{\text{rad}} + \frac{dm_p}{dt} h(T_p)_{vap}$$

$$- \frac{dm_p}{dt} [h_s(T_g)_{vap} + h_s(T_p)_{vap}]$$

(5)

The first term on the RHS of Eq. (5) accounts for the convective effect of the coal particle on the surrounding gas, which can be obtained from the first term on the RHS of Eq. (3), and the second term accounts for the radiative effects. The third term is responsible for the enthalpy increase due to the added mass in the gas phase from the coal particle at the particle temperature. The fourth term represents the energy required to bring the sensible enthalpy of the released gases ($h_{s,vap}$) from the particle temperature to the gas temperature.
3.4. Coal combustion model

Previous studies [27, 28] found that the exchange of $N_2$ for $CO_2$ in oxy-fuel combustion does not have much influence on devolatilisation. It was thus deemed sufficient for the purpose of this study to represent the devolatilisation of the volatile gases from the coal particle using the single-step reaction rate model [29], which assumes a single reaction for the total weight loss of the volatiles:

$$\frac{dm_{vg}}{dt} = k_v(VM - VG)$$  \hspace{1cm} (6)

$$k_v = A_v T_p^\beta \exp \left( - \frac{E_v}{RT_p} \right)$$  \hspace{1cm} (7)

The constants $A_v$, $E_v$ and $\beta$ follow a modified Arrhenius expression, $VG$ is the total mass of volatile gases that have left a coal particle and $VM$ is the initial mass of volatile matter inside the coal particle adjusted by a factor $Q = VM_{CPD}/VM_{PA}$ to account for the higher yield of volatile gases at higher temperatures [29]. The values used were $Q = 1.15$ and the single rate parameters $A_v = 4.727 \times 10^9 \text{ (1/s)}$, $E_v = 10.256 \times 10^3 \text{ J/(kmolK)}$ and $\beta = -0.9503$ were obtained from a fit of the Chemical Percolation Devolatilisation (CPD) Model [30]. The input values for the CPD model were obtained from the proximate and ultimate analysis through the correlation of Genetti et al. [31].

Char combustion in a $CO_2$ enriched environment can differ considerably compared to combustion in air. The char burning temperatures and residence times are lower in a $O_2/CO_2$ environment compared to a $O_2/N_2$ environment [28]. This difference can be attributed to the char-$CO_2$ and char-$H_2O$ gasification reactions. Three heterogeneous reactions were considered in the present work:

$$C_{char} + 0.5O_2 \rightarrow CO$$  \hspace{1cm} (8)

$$C_{char} + CO_2 \rightarrow 2CO$$  \hspace{1cm} (9)

$$C_{char} + H_2O \rightarrow CO + H_2$$  \hspace{1cm} (10)

The reaction rates of Eqs. (8-10) are obtained from the Baum and Street model [32]:

$$\frac{dm_{char}}{dt} = A_p p_{ox} \left( \frac{1}{R_c} + \frac{1}{D_o} \right)$$  \hspace{1cm} (11)

Here, $D_o = C_{diff}/d_p[(T_p+T_g)/2]^{3/4}$ is the diffusion rate term. The chemical reaction rate term is obtained via the Arrhenius expression $R_c = A_c \exp(-E_c/RT_p)$. 

8
The diffusivity constants and Arrhenius coefficients for the reactions (8-10) were provided by Toporov et al. [14] and are consistent with their RANS simulations, and are reproduced in Table 3.

Table 3: Rate parameters and diffusion coefficients for the oxy-char surface reactions [14].

<table>
<thead>
<tr>
<th>Reaction</th>
<th>n</th>
<th>$A_c$</th>
<th>$E_c$</th>
<th>Temp (°C)</th>
<th>$C_{diff}$</th>
<th>Ref</th>
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<tbody>
<tr>
<td>8</td>
<td>1</td>
<td>0.005</td>
<td>174,000</td>
<td>&gt;677</td>
<td>$4.41 \times 10^{-12}$</td>
<td>[14, 33]</td>
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<tr>
<td>9</td>
<td>1</td>
<td>$0.135 \times 10^{-3}$</td>
<td>135,500</td>
<td>850-950</td>
<td>$2.47 \times 10^{-12}$</td>
<td>[14, 34]</td>
</tr>
<tr>
<td></td>
<td>1</td>
<td>$6.35 \times 10^{-3}$</td>
<td>162,000</td>
<td>&gt;950</td>
<td></td>
<td></td>
</tr>
<tr>
<td>10</td>
<td>1</td>
<td>0.319</td>
<td>208,000</td>
<td>860-960</td>
<td>$2.47 \times 10^{-12}$</td>
<td>[14, 34]</td>
</tr>
<tr>
<td></td>
<td>1</td>
<td>$1.92 \times 10^{-3}$</td>
<td>147,000</td>
<td>&gt;960</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

3.5. Gas phase turbulent combustion

The volatile gases were modelled as a single postulated substance $C_aH_bO_cN_d$ ($a = 2.58$, $b = 3.98$, $c = 0.87$, $d = 0$). The enthalpy of formation of the volatile gases was determined to be $h_{f,vol} = -587.86$ MJ/kmol. The homogeneous chemistry was described by a two-step mechanism:

$$C_{2.58}H_{3.98}O_{0.87} + 1.85O_2 \rightarrow 2.58CO + 1.99H_2O \quad (12)$$

$$CO + \frac{1}{2}O_2 \rightarrow CO_2 \quad (13)$$

The reaction rate $\bar{w}_{vg}$ was modelled using the Eddy Break Up (EBU) [35] model adjusted for LES [36].

$$\bar{w}_{vg} = c_{EBU} \bar{\varphi} |\bar{S}| \min \left( \frac{\bar{Y}_{fu}}{s}, \frac{\bar{Y}_{ox}}{s}, c_2 \frac{\bar{Y}_{pr}}{1 + s} \right) \quad (14)$$

The model constants are $c_{EBU} = 4.0$ and $c_2 = 0.5$, while $s$ is the stoichiometric coefficient. Ideally the volatile gases should be modelled as an ensemble of multiple species, however this would require very large reaction mechanism, resulting in a considerable increase in computational cost, which would be unfeasible for the already very expensive simulation (see Table 4). Moreover, using a large mechanism for this flame would require dedicated sub-grid modeling of turbulence-chemistry interaction which is done by the EBU model but not with a direct application of the mechanism.
3.6. Radiative heat transfer

The Radiative Transfer Equation (RTE) is solved using the Discrete Ordinates Method (DOM) [37, 38, 39]. In the DOM, the RTE is discretized and solved for \( n \) different direction \( \hat{s} \), and summed over all directions in the integral \((4\pi)\). In this work the RTE was discretized and solved for 24 directions using the S\(_4\) approximation. The total incident radiation \( G \) is calculated from the sum of all the intensities impinging on a given control volume:

\[
G = \int_{4\pi} I(\hat{s})d\Omega \approx \sum_{i=1}^{n} w_i I_i
\]

where \( w_i \) is the quadrature weight associated for each direction.

The spectral properties of the participating media were treated with a grey gas model. Gosman and Lockwood [40] found that in the context of coal combustion \( CO_2, H_2O \) and the volatile gases have a dominant impact over other gaseous species. In this work the volatile gases are treated as a single postulated substance \( C_\alpha H_\beta O_\gamma \) and the gas absorption co-efficient was determined using an empirical relationship [40]:

\[
\kappa_g = 0.2X_{vg} + 0.1(X_{CO_2} + X_{H_2O})
\]

The heat exchange of a particle with the gas phase due to radiation \( \dot{Q}_{rad} \) (Eq. 5) can be obtained by:

\[
\dot{Q}_{rad} = \frac{1}{4\varepsilon_p \pi d_p^2} (4\pi I_{b,p} - G)
\]

The particle’s absorption coefficients \( \kappa_p \) and scattering coefficient \( \sigma_p \) in a given cell are determined according Chui et al. [41]:

\[
\kappa_p = \varepsilon_p \sum_i N_i \frac{\pi d_{p,i}^2}{4}
\]

\[
\sigma_p = (1 - \varepsilon_p) \sum_i N_i \frac{\pi d_{p,i}^2}{4}
\]

where \( N_i \) is the particle number density relevant to the size class \( d_i \), and \( \varepsilon_p \) is the particle emissivity which depends on the char-burnout, and the proportions of volatile content and ash found in the particle. Finally, the particle blackbody intensity \( I_{b,p} \) is obtained via:

\[
\kappa_p I_{b,p} = \varepsilon_p \sum_i N_i \frac{\pi d_{p,i}^2}{4} \frac{\sigma T_{p,i}^2}{\pi}
\]

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3.7. Simulation

The furnace geometry was described by a Cartesian mesh $0.4 \times 0.4 \times 0.8$ m in size, which represents a reduced domain length compared to the whole furnace length (2.1 m). Coarse and fine grid simulations were performed with equally sized cubic cells ($\Delta = 2$ mm and $\Delta = 1$ mm), leading to 16M and 128M cells respectively. The coarse grid simulations were all performed on 96 processors and took approximately 15 days ($\approx 35,000$ CPU hours) for 100,000 steps. Without radiation, the equivalent simulation took approximately half of the time. The fine grid simulation was performed on 384 processors and was only performed without radiation, as the coarse grid simulation showed the effect of radiation to be small as explained in more detail in section 5.1. The fine grid simulations took approximately 27 days ($\approx 250,000$ CPU hours) to perform 170,000 iterations, which were necessary to achieve a converged solution. A summary of the computational cost of the fine and coarse grid simulations is shown in Table 4.

At the inlets, top hat velocity profiles were imposed, corresponding to the nominal volume fluxes. Pseudo-turbulent inflow conditions were generated [42] with an inlet fluctuation $u' = 0.5$ m/s and a length scale $L_{turb} = 3$ mm. Laminar flow conditions were assumed for the staging stream. Immersed boundaries were set at the burner walls. For radiation, the boundary conditions were taken from Toporov et al. [14], who provided temperature and emissivity values for the furnace and burner of 1000 °C ($\epsilon = 0.7$) and 300 °C ($\epsilon = 0.2$) respectively. For the open boundaries at the inflow and outflow of the domain, blackbody conditions were assumed ($\epsilon = 1.0$). Approximately 11

<table>
<thead>
<tr>
<th></th>
<th>Coarse grid</th>
<th>Fine grid</th>
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<tbody>
<tr>
<td>$N_o$ of cells</td>
<td>$\Delta = 2$ mm</td>
<td>$\Delta = 1$ mm</td>
</tr>
<tr>
<td>$N_o$ of CPUs</td>
<td>16 M</td>
<td>128 M</td>
</tr>
<tr>
<td>Time steps</td>
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<td>384</td>
</tr>
<tr>
<td>Accumulation time</td>
<td>100,000</td>
<td>170,000</td>
</tr>
<tr>
<td>Days (without Rad.)</td>
<td>3.25 sec</td>
<td>3.00 sec</td>
</tr>
<tr>
<td>CPU hours (without Rad.)</td>
<td>18,000</td>
<td>250,000</td>
</tr>
<tr>
<td>Days (with Rad.)</td>
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<td>27</td>
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<tr>
<td>CPU hours (with Rad.)</td>
<td>15</td>
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</tbody>
</table>
million computational particles per second were fed into the domain, where each computational particle represents an ensemble of ten real particles. Approximately 1.8 million particles were present in the domain at a given time during the simulation. The particle size distribution given by Toporov et al. [14] was used in the simulations.

4. Results

4.1. Flow field

Figure 2 shows the volume rendered particle positions across the whole domain. The highest particle density is found in the burner quarl, where the particles are fed in from the inlet. The particle cloud expands axially and radially almost uniformly, and at \( z \approx 0.35 \) m the particles are almost evenly spread throughout the domain cross section, see Figure 2(b). Upstream (\( z < 0.2 \) m) very few particles are found close to the walls as the incoming staging stream pushes the particles towards the center of the domain.

Figure 3 shows the contour plots of the instantaneous mean axial and tangential gas velocities. The coal carrying primary stream and the swirling secondary stream merge together almost immediately, and form a single stream that expands radially outwards and starts to decay at approximately \( z = 0.1 \) m downstream from the burner outlet. The swirling secondary stream and the quarl geometry are responsible for the very strong internal recirculation zone (IRZ) which is clearly visible inside the quarl.

Figure 4 shows the mean axial (a) and tangential (b) velocity profiles predicted by the LES, and compared to experimental measurements at four axial locations from the burner outlet. Figure 4(a) shows that the LES correctly predicts the magnitude and size of the axial velocity maximum and tangential velocity minimum at \( z = 0.025 \) m and \( z = 0.05 \) m. Another axial velocity peak is observed close to the furnace walls, caused by the staging stream.

A strong internal recirculation zone (IRZ) is clearly visible inside the quarl in Fig. 1. The experimentalists measure a strong backward axial velocity \( \approx -3 \) m/s at the two upstream planes, \( z = 0.025 \) m and \( z = 0.05 \) m, which is correctly captured by the LES. At \( z = 0.2 \) m, the experiments show a slower (\( \approx -1 \) m/s) but much larger external recirculation zone (ERZ), which extends from \( r = 0.1 \) m to the furnace wall. The ERZ predicted by the LES is smaller in size and magnitude. Moreover the LES over-predicts the velocity profiles at \( z = 0.3 \) m. The discrepancies between the LES results and the experimental measurements in the downstream planes can be attributed to...
uncertainties in the staging stream. The behaviour of the staging stream is of crucial importance for this burner given that it transports more than 50% of the fluid mass into the domain. However, the experiments used the coal particles in the flow for the LDA measurements, which (see Fig. 2) will be mostly present upstream, in the central region of the furnace. Consequently the added mass (and velocity) provided by the staging stream will not be captured by the LDA measurements, turning the staging air into an “invisible” mass flow. To analyse the effect of the staging stream, a coarse adjustment to the LES results was made. The volume flow of the staging stream was subtracted throughout the domain width at the $z = 0.3\,\text{m}$ plane and throughout half of the domain width ($r > 0.1\,\text{m}$) for the $z = 0.2\,\text{m}$ plane. In Fig. 4(a) the LES results are closer to the experimental measurements when the staging air flow is subtracted from the experimental data. A real compensation of the experimental uncertainty introduced by the “invisible” staging air cannot be reproduced by LES-statistics of particle velocity, since it is not clear

Figure 2: Fine grid LES of (a) volume rendering of individual particle positions, white regions show high particle density. (b) Individual particle position, in cross section at $z = 0.35\,\text{m}$, showing equal distribution of particles along the cross section of the burner. (Dimensions in mm.)
how large the particle (local, instantaneous) concentration needs to be for the experiment to get a valid signal (one should note that the axial velocity and the local concentration of mass from the staging stream are likely to be correlated). The velocity fluctuations at $z = 0.05\, \text{m}$ and $z = 0.3\, \text{m}$ have been included for completeness in Figure 5.

Figure 6 shows the ratio between turbulent viscosity and laminar viscosity ($\mu_t/\mu_l$) for the fine grid simulation. Regions of intense turbulence can be observed in the corner between the quarl and the inlet plane. It should be noted that in this small region, the viscosity ratio reaches higher values ($\approx 40$) than what is normally aimed for ($\approx 20$) [43] in the high-quality LES of laboratory scale gas burners, indicating that achieving a good grid resolution of swirled coal burners is still not an easy feat.
Figure 4: Comparison between LES fine (Δ = 1 mm) (solid line) and coarse (Δ = 2 mm) (dotted line) (a) mean axial and (b) tangential gas velocity profiles with experimental measurements. (c) Mean radial gas velocity profiles, radial profiles were not available experimentally.

4.2. Devolatilisation and volatile combustion

Figure 7 shows the volatile yield and volatile gas concentration in a cutting plane through the central axis of the burner. The volatile yield represents the rate at which volatiles are being released by the pulverised coal particles based on the single step devolatilisation model eq. 6. The volatile gas concentrations, represent the mass concentrations of volatile gases present on average at a given point in the burner. The volatile gases are released rapidly, close to the burner inlet in the quarl and are mostly consumed in the central up-stream region of the flame z < 0.1 m.

Figure 8 shows the contour plots of the mean (a) and instantaneous (b) O₂ concentrations in the furnace in a cutting plane through the central axis of
the burner. The oxygen enters the quarl through the primary and secondary stream and is transported radially outwards, carrying the coal particles along. Most of the oxygen from the primary and secondary stream, together with the coal particles, are carried into the highly swirling flow surrounding the IRZ, and the particles start heating up very rapidly, thereby releasing rapidly the volatile gases (see Figure 7 (a)). Once the volatile gases are released in the vortex surrounding the IRZ, they burn fast, limited only by the rate of mixing according to the EBU model. In fact, the volatile combustion process only occurs in the vortex surrounding the IRZ but hardly within the IRZ itself. In the vortex surrounding the IRZ most of the combustion occurring will relate to the burning of the volatile gases to form CO via reaction (12). This is observable in Figure 9, which shows a high CO concentration in the IRZ region. Within the IRZ itself, there is no oxygen left (see Figure 8) to burn the remaining CO and volatile gases, which causes parts of the unburned volatile gases and CO to be transported back into the quarl.

Figure 8 (d) shows the $H_2O$, volatile gases ($VG$) and $O_2$ mole fraction profiles obtained from the LES and the experimental measurements of $O_2$ concentrations at four axial distances from the burner outlet. Both the experiments and the LES show very low $O_2$ concentrations at $z < 0.2$ m in the central region of the furnace. In the same region, the LES results show diminishing $VG$ and increasing $H_2O$ concentrations, which confirms that intense volatile combustion must be occurring there. For all of the upstream planes the $O_2$
levels predicted by the LES follow the experimental measurements reasonably well between 0 < r < 0.05 m, indicating that the volatile combustion process is being captured satisfactorily by the LES. Nonetheless, in the region between 0.05 < r < 0.15 m the LES over-estimates the O\textsubscript{2} concentration. This can be attributed to the devolatilisation and EBU models, which might over-estimate the volatile release rate and their combustion rate. If the volatile gases are present outside the IRZ they can burn with the surrounding O\textsubscript{2}, lowering its concentration between 0.05 < r < 0.15 m. A second reason for the deviations might be related to the devolutilisation model and will be discussed in section 5.
Figure 8: Left: Fine grid LES contour plots of $O_2$ concentration in a cutting plane through the central axis (a) mean, (b) instantaneous and (c) instantaneous cross section at $z = 0.35$ m. (Dimensions are in mm.) Right (d): Fine $O_2$, $VG$ and $H_2O$ and coarse $O_2$ concentrations predicted by LES and $O_2$ experimental measurements.

4.3. Char Combustion

More than half of the $O_2$ entering the furnace comes from the staging stream, which drives most of the char combustion process, especially downstream. The $O_2$ in the staging stream flows inwards towards the centre of the furnace filling both the upstream outer region of the furnace and downstream throughout the furnace width. After all the volatile gases have been released, the char particles that escape the IRZ surrounding vortex are transported axially and radially outwards and start reacting with the $O_2$ from the staging stream. Figure 10 shows that the bulk of the char combustion occurs outside the IRZ throughout most of the domain. This is also evident from Fig. 9, which shows several small regions of high instantaneous $CO$ concentrations surrounding the IRZ. These are due to local region of intense char combus-
tion, where CO is primarily produced by char combustion according to Eqs. (8-9). Figure 11 compares the mean values of the char-O$_2$ reaction and the char-CO$_2$ reaction along the central plane of the furnace. The char-O$_2$ reaction rate contributes about an order of magnitude more than the char-CO$_2$ gasification to the total char combustion rate. The exact contributions of the char-O$_2$ and char-CO$_2$ reactions to char consumption are 92.91% and 7.00% respectively, which is very close to the values observed by Chen et al. [16], who predicted 91.90% and 7.46% and only 0.68% for the char-H$_2$O reaction. As the contribution of the char-H$_2$ reaction was found by Chen et al. [16] to be < 1% it was ignored in the present calculations to avoid the need for transporting H$_2$. For the present test case the char oxidation reaction is particularly dominant due to the relatively low temperatures throughout the domain (see Figure 12), which does not necessarily imply that such is the case for every oxy-coal combustion process. The char-CO$_2$ gasification process may become important in regions of high temperatures and low O$_2$ concentrations. As shown in Figure 11, within the central region of the flame, some char-CO$_2$ gasification is observed even in the absence of O$_2$, which may contribute considerably to the overall combustion process.

Most of the volatile gases leave the coal particles in the vortex surrounding the IRZ, and the devolatilisation process terminates close to the burner inlet. Figures 10 and 11 show that most of the char combustion occurs in the region of $z < 0.3$ m and that little char is burnt downstream of $z > 0.3$ m.
This provides additional justification for simulating the boiler only to 0.8 m downstream from the burner, excluding the regions even further downstream.

4.4. Temperature

Figure 12 shows the contour plots of the mean (a) and instantaneous (b) gas temperature in the furnace in a cross-sectional plane along the central axis and (c) an instantaneous cross-sectional image of the gas temperature at $z = 0.35$ m. The combustion of the volatiles results in the rapid temperature increase in the location of the primary/secondary stream jet in Fig. 12. The temperature is highest at this point within the quarl, reaching values of approximately $\approx 1400$ K. At the centre of the quarl the temperature is lower $\approx 1000$ K, which confirms that limited combustion is occurring there and the temperature rise is mainly due to the recirculating hot products (see section 4.2). The combustion of the volatiles extends to $z \approx 0.2$ m, contributing to the rapid temperature rise occurring in the centre of the furnace between $z = 0.05$ m and $z = 0.2$ m. In the downstream region of the furnace ($z > 0.3$ m), overall high temperatures are observed everywhere along the central axis of the flame, Fig. 12(a-b), and throughout the furnace cross-section, Fig. 12(c). The temperature rise is a result of the CO produced by the char combustion (see Fig. 10, and section 4.3) burning with the oxygen from the staging stream.

In Figure 12(d) the gas temperature profiles predicted by the LES are compared to experimental measurements at four axial locations from the burner outlet. At the $z = 0.05$, 0.1 and 0.2 m planes, a good agreement is ob-
served between the LES and experiments. At the $z = 0.3$ m plane, the LES overestimates the experimental measurements by up to 200 K. The high temperatures predicted by the LES cause an over-expansion of the gases, which could explain the discrepancy in axial velocity at the $z = 0.3$ m plane in Fig. 4. An attempt to understand the discrepancies in the downstream temperatures between the LES and experiments is made in the next section.

5. Parametric study

A parametric study has been performed, to understand what affects the results and which models must be improved to enhance the predictions. The effect of grid refinement on the simulation results was assessed by performing two grid simulations with cell size $\Delta = 1$ mm and $\Delta = 2$ mm. Figures 4, 8 and 12 compare the fine and coarse grid simulation results for the velocity, temperature and oxygen concentration. The oxygen concentration predicted by the fine grid simulation follows more closely the experimental data. For the fine grid simulations, the better description of mixing meant that the EBU model can provide a better description of the reaction rates.
Figure 12: Left: Fine grid LES (a) mean and (b) instantaneous contour plots of gas temperature in a cross-sectional plane along the central axis and (c) cross section of instantaneous gas temperature at \( z = 0.35 \) m. (Dimensions are in mm.) Right (d): Comparison between LES fine (solid line) and coarse with radiation (dotted red line) and without radiation (dotted black line) gas temperature with experimental measurements.

However, for the downstream planes the temperature results of the coarse and fine grid simulations are almost identical. This suggests that the downstream temperature discrepancies are not related to the gas combustion rates but to processes independent of mixing such as char combustion or radiation. The results between the two simulations did not differ considerably, and therefore in all further simulations for the parametric study, the coarse grid was retained.

5.1. Effect of radiation

Figure 12(d) shows the gas temperature profiles for the coarse grid LES with and without radiation (dotted lines). The temperature at the furnace
walls were set to 1000 °C as suggested by Toporov et al. [13, 14], which is slightly colder than the upstream temperature measurements close to the walls. The upstream gases close to the furnace walls thus lose heat, which results in the slight drop in temperature observed at the \( z = 0.05 \) m plane and close to the walls at the \( z = 0.1 \) m plane. Throughout the central region of the flame, radiation has a negligible impact. In this region the walls are too far (and not cold or hot enough) to have a considerable impact. Moreover, the transporting medium temperature is not very hot, which limits the amount of radiative heat dispersed to either the surroundings or transferred to the dispersed phase.

The current results are unable to assess alone whether radiative effects are really negligible within the central regions of the flame. Treating the radiative properties of the participating medium as grey is somewhat simplistic. A more advanced treatment of the radiative properties would be to use the Weighted Sum of Grey Gas (WSGG) method, where the radiative properties of the medium are represented by a number (usually four or five) grey gases, and where the RTE is solved for each grey gas, having its own absorption coefficient and weight, rather than a single value to represent the whole mixture. Cavallo Marincola [44] compared both the grey gas and WSGG approaches for the IFRF No.1 burner [45] and found that the grey medium assumption overestimated the temperatures downstream, whilst the WSGG approach gave overall better results. The WSGG method however, leads to a considerable increase in computational times which would be currently prohibitive for this test case.

Moreover, Toporov et al. [13, 14] provided the radiation boundary conditions that were used in this work. A more thorough analysis of the radiative properties at the burner walls could show that more appropriate radiative boundary conditions are required for the burner, possibly resulting in a bigger impact on the temperature predictions.

5.2. Char Combustion

Figure 13 compares the temperature and oxygen concentration results for two LES, where in one the char combustion rate was artificially diminished by a factor of ten and in another the char combustion was ignored. In the upstream \((z < 0.2 \text{ m})\) central region of the flame the results with the different char combustion rates show negligible differences. This follows the previous findings, which concluded that in the IRZ and in its surrounding vortex the driving combustion processes are devolatilisation and the burning
of the volatile gases.
The results in section 4 showed that the driving combustion process in the downstream region of the furnace \((z > 0.2 \text{ m})\) is char combustion. The results observed in Figure 13 confirm this, indicating a considerable drop in temperature throughout the domain width at the downstream planes when
the char combustion rate is diminished. The downstream temperature for the slower char combustion process matches the experimental data at the \( z = 0.3 \) m plane, which suggests that the current char combustion model over predicts the char combustion rate. A more detailed analysis could involve implementing more advanced char combustion models that account for multiple competing char oxidation reactions (Shaddix et al. [28, 46]), but it should be stressed that the baseline case presented here was necessary first, before further model improvements can be justified and tested reliably.

5.3. Devolatilisation

To test the effect of the uncertainties in the devolatilisation model, the rate was adjusted. Figure 13 shows the LES results with the devolatilisation rate constant \( A_v \) of Eq. 6 reduced by a factor of ten. In the upstream \((z < 0.2 \) m) central region of the flame the results with the slower devolatilisation rate show a large drop in temperature. The previous findings concluded that most of the volatile gases are released and burn within the vortex surrounding the IRZ, leading to the high temperatures observed in the upstream central region of the flame. When the devolatilisation rate is diminished, the particles release most of the volatiles outside the IRZ and burn upstream between \( 0.05 \) m < \( r \) < \( 0.15 \) m, producing the high temperatures observed in the same region. The oxygen from the primary and secondary stream is not fully consumed and flows downstream resulting in the higher \( O_2 \) concentrations and lower temperatures observed throughout the central axis of the furnace. Figure 13 shows that the LES results with the slower devolatilisation rate deviate further from the experimental measurements. However, especially in the plane \( z = 0.1 \) m the temperatures between \( 0.05 \) m < \( r \) < \( 0.15 \) m are in closer agreement with the experimental data. This might be attributed to the current devolatilisation model, which does not distinguish between the release of lighter and heavier volatiles. In the experiments the lighter volatiles may be released within the vortex surrounding the IRZ, whilst the heavier tars are released outside. This would have the effect of slightly dropping the high temperatures in the central upstream region of the flame and slightly increasing the temperatures at the sides of the flame between \( 0.05 \) m < \( r \) < \( 0.15 \) m, thereby potentially matching more closely the temperature measurements. Ideally, the volatile gases would be represented as light hydrocarbons, tar, carbon dioxide and water vapour, thus removing the need to model them as a single postulated substance. This would allow the use of more advanced devolatilisation models that account for different species and release rates.
However, currently transporting multiple species and solving their reactions using multiple reaction chemistry mechanisms, is prohibitively expensive, and could introduce additional uncertainties into the simulation.

6. Conclusions

Large Eddy Simulations were performed of a swirling oxy-coal flame. A working set of LES models for oxy-coal combustion with radiative heat transfer and Lagrangian particle transport have been implemented, tested and verified in comparison to experimental data. A qualitative analysis was made of the flow behaviour in the furnace and the overall PCC processes occurring. The volatiles are mostly released in the quarl and are burnt rapidly upstream by the $O_2$ from the primary and secondary streams. Most of the downstream combustion is driven by the char burning with the $O_2$ from the staging stream. The LES results were compared to experimental data at several locations, showing an overall good agreement in the flame region. A grid resolution study confirmed that almost grid independent results have been obtained. A parametric study was conducted to investigate the effect of the devolatilisation and char combustion rates on the LES results. The current LES was able to predict the flow and combustion, but also provided a wealth of additional time resolved data that can be accessed for further analysis. Such data is particularly relevant for improving boiler performance and for reducing pollutant emissions from flame regions that are not accessible to experimental investigation.

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