LARGE EDDY SIMULATIONS OF COAL JET FLAME IGNITION
USING THE DIRECT QUADRATURE METHOD OF MOMENTS

by

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ABSTRACT

The Direct Quadrature Method of Moments (DQMOM) was implemented in the Large Eddy Simulation (LES) tool ARCHES to model coal particles. LES coupled with DQMOM was first applied to nonreacting particle-laden turbulent jets. Simulation results were compared to experimental data and accurately modeled a wide range of particle behaviors, such as particle jet waviness, spreading, break up, particle clustering and segregation, in different configurations. Simulations also accurately predicted the mean axial velocity along the centerline for both the gas phase and the solid phase, thus demonstrating the validity of the approach to model particles in turbulent flows.

LES was then applied to the prediction of pulverized coal flame ignition. The stability of an oxy-coal flame as a function of changing primary gas composition ($CO_2$ and $O_2$) was first investigated. Flame stability was measured using optical measurements of the flame stand-off distance in a 40 kW pilot facility. Large Eddy Simulations (LES) of the facility provided valuable insight into the experimentally observed data and the importance of factors such as heterogeneous reactions, radiation or wall temperature. The effects of three parameters on the flame stand-off distance were studied and simulation predictions were compared to experimental data using the data collaboration method.

An additional validation study of the ARCHES LES tool was then performed on an air-fired pulverized coal jet flame ignited by a preheated gas flow. The simulation results were compared qualitatively and quantitatively to experimental observations for different inlet stoichiometric ratios. LES simulations were able to capture the various combustion regimes observed during flame ignition and to accurately model the flame stand-off distance sensitivity to the stoichiometric ratio. Gas temperature and coal burnout predictions were also examined and showed good agreement with experimental data.
Overall, this research shows that high-fidelity LES simulations combined with DQMOM can yield a deeper understanding of complex coal flames and their ignition mechanisms, indicate where experimental uncertainties lie and in the end, be a valuable tool for the design, retrofit and scale-up of oxy-coal boilers.
Je souhaite dédier la réalisation de ce mémoire à mes parents qui ont toujours été là pour moi et qui m’ont infailliblement soutenu. Je tiens par ces quelques mots à leur exprimer mon amour, ainsi qu’à ma très chère soeur et à ma femme.
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NOMENCLATURE

\( \nu \)   gas kinematic viscosity \((m^2/s)\)
\( d_p \)   particle diameter \((m)\)
\( d \)   primary nozzle diameter \((m)\)

DQMOM Direct Quadrature Method of Moments

\( \epsilon \)   turbulence energy dissipation rate \((m^2/s^3)\)
\( \rho_g \)   gas density \((kg/m^3)\)
\( \mathbf{u}_g \)   gas velocity \((m/s)\)
\( \tau_p \)   particle relaxation time \((s)\)
\( h_p \)   particle enthalpy \((J)\)
\( \xi \)   internal coordinate vector
\( \kappa \)   wave number \((m^{-1})\)
\( \eta \)   Kolmogorov length scale \((m)\)

LES Large Eddy Simulation

\( n \)   number density function \((\text{number of particles/m}^3/\text{internal coordinate})\)
\( N_\xi \)   number of internal coordinates
\( N \)   number of quadrature nodes
\( \Delta \)   Nyquist cut-off length \((m)\)
\( \mathbf{u}_p \)   particle velocity in real space \((m/s)\)
\( \mathbf{v}_p \)   particle velocity in internal coordinate space \((m/s)\)
\( \alpha_c \)   particle mass of raw coal \((kg)\)
\( \alpha_h \)   particle mass of char \((kg)\)

\( P_{O_2, pri} \)   partial pressure of oxygen in the primary stream
\( Q_{\text{conv}} \)   net convective and conductive flux to particle \((J/s)\)
\( Q_{\text{rad}} \)   Net radiative flux to particle \((J/s)\)
\( T_g \)   gas temperature \((K)\)
\( T_p \) particle temperature (K)
\( T_{pri} \) primary stream temperature (K)
\( T_{wall} \) wall temperature (K)
\( \tau_f \) fluid characteristic time (s)
\( U_0 \) central jet maximum gas velocity (m/s)
\( U_a \) annular jet maximum gas velocity (m/s)
\( VR \) inlet velocity ratio
\( w_\alpha \) weight for quadrature node \( \alpha \) (number of particles/m\(^3\))
\( \langle \xi \rangle_\alpha \) abscissa for internal coordinate \( \xi \) and quadrature node \( \alpha \)
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CHAPTER 1

INTRODUCTION

Global warming, one of the largest environmental challenges of our time, is due to increased carbon dioxide levels in the atmosphere [66]. The world is currently depending on the use of fossil fuels, especially coal, for its energy supply, and will continue to do so for a long time. According to the Kyoto Protocol, by 2008-2012, Annex I countries have to reduce their greenhouse gas emissions by a collective average of 5% below their 1990 levels. Therefore, oxy-fuel combustion, as one of the promising technologies to reduce CO$_2$ emissions, has attracted worldwide attention [1,20,48,66]. Oxy-coal combustion, in which an O$_2$/CO$_2$ mixture replaces air, is one of the few possible capture technologies to enable CO$_2$ sequestration for existing coal-fired boilers. Burning coal with relatively pure oxygen, together with recycled flue gases, can produce a highly concentrated (up to 95% CO$_2$) flue gas stream, which makes carbon sequestration more economical. One issue of interest in rapidly implementing a strategy to retrofit existing air-fired burners or designing new ones is to understand how replacing air by a O$_2$/CO$_2$ mixture affects the kinetics, aerodynamics, pollutants formation, heat fluxes and flame ignition in the near burner region. Large Eddy Simulation has the potential to predict oxy-coal flame characteristics and to be an important tool in the retrofitting and designing processes. In a turbulent coal flame, accurate prediction of the particle behavior is critical for predicting the flame characteristics. Stokes number analyses suggest that LES has the capabilities to resolve all the turbulent length scales affecting coal particles in a coal flame, and could therefore lead to more accurate simulations.

The main purpose of this research is to perform reliable Large Eddy Simulations (LES) of oxy-coal flames. ARCHES, the LES tool developed at the University of Utah, initially only included gaseous phase simulations, so implementation of particle-modeling capabilities was
required. The Direct Quadrature Method of Moments (DQMOM) was chosen to represent particles in an Eulerian-Eulerian framework. This method allows the transport of multidimension probability density functions in a computationally cheap manner. The first step of this research project was thus the implementation of DQMOM in Arches in a stable and efficient way. The next one was to implement all the physical models related to coal particles combustion within the DQMOM approach and couple them with the gaseous phase. A non-exhaustive list of those models includes mass, momentum and heat transfer between the gas and solid phases, devolatization and char oxidation.

As a general rule, simulations results should be compared to experimental data to assess the validity and accuracy of the models. This process, called Validation and Uncertainty Quantification, can be achieved be selecting relevant experimental data and comparing them with the predictions made by the simulations. This research has two main objectives.

The first one is to show that DQMOM can be used in combination with LES to accurately model particles in complex turbulent flows. It was proposed to use Budilarto’s data [14] to validate this approach. Budilarto investigated the effect of fluid aerodynamics on particle motion in the near field region of coaxial particle-laden jets. The fluid aerodynamics was modified by varying the inlet velocity ratio of the annular to central jet velocity at 0, 1.0 and 1.5. The dispersion of particles was enhanced with decreasing particle size and with increasing the velocity ratio greater than 1.0. The particle-phase radial velocity fluctuation was determined as an important parameter to characterize the particle-turbulence interaction and the spatial distribution of particles in coaxial laden jets. The investigation of particle size distribution effects showed that the radial spreading of the coarse particles was reduced by the presence of the fine particles. LES simulation of Budilarto’s cases are presented in the first chapter and show that this LES combined with DQMOM is able to predict particle dispersion, particle clustering and gas-solid interactions.

The second objective aims to show that LES coupled with DQMOM can be used to perform reliable simulations of oxy-coal flames. One issue of interest in developing oxy-coal technology is to understand and predict the effects of near burner zone environment, now
consisting of $O_2$ and $CO_2$ instead of $N_2$, on flame stability and ignition. Flame stability is an important issue in designing burners because it affects the lifespan of the burner as well as pollutant formation and heat fluxes in the furnace. Experimental data on oxy-coal flame ignition that would meet the criteria for LES simulations are very limited. Zhang [112,113] studied how composition of the coal transport medium (primary $CO_2$ and $O_2$) affects the observed ignition stand-off distance in coaxial turbulent diffusion, oxy-coal flames. Results show that flame stability is affected by primary $P_{O_2}$, wall temperature, secondary preheat temperature, secondary $P_{O_2}$ and transport medium. The burner and the furnace geometry were chosen to be simple enough for LES simulations. It was therefore proposed to perform simulations of Zhang’s cases and to compare predictions on the flame stand-off distance with the experimental data. To further assess the capabilities of LES coupled with DQMOM to predict coal flame ignition, another validation study was also performed for an air-fired pulverized coal flame. LES was applied to a coal jet flame ignited by a preheated gas flow. The simulation results were compared qualitatively and quantitatively to experimental observations for different inlet stoichiometric ratios.

The details for each simulation performed in this thesis can be found in the input files available online at: https://software.crsim.utah.edu/svn/LES_Coal/Pedel.
2.1 Abstract

Modeling coal particles in turbulent coal flames is a challenging but nonetheless critical step for predicting flame characteristics. Traditional approaches such as Reynolds-Averaged Navier-Stokes (RANS) coupled with Lagrangian solvers are unable to correctly predict particle concentrations and velocities in complex systems. This study shows how Large Eddy Simulations (LES) coupled with an Eulerian solver can address those issues. Previous Direct Numerical Simulations and Stokes number analysis suggest that LES has the capabilities to resolve all the turbulent length scales affecting coal particles in a coal flame and could therefore lead to more accurate simulation. The effects of fluid aerodynamics on the particle
motion in the near field region of nonreacting coaxial particle-laden jets were simulated using LES. The Direct Quadrature Method of Moments (DQMOM) was used to track the particle phase in an Eulerian framework. Simulation results were compared to experimental data and accurately modeled a wide range of particle behaviors, such as particle jet waviness, spreading, break up, particle clustering and segregation, in different configurations. Simulations also predicted the mean axial velocity along the centerline for both the gas phase and the solid phase with a maximum error of 12% relative to experimental data. This study therefore provides a solid validation of the LES with DQMOM approach to model particles in turbulent flows and justifies its use for coal flame simulations.

2.2 Introduction

Coal is an important and indispensable energy resource for electricity production as its reserves are far more abundant than those of other fossil fuels. As environmental regulations get stricter, coal thermal power plants need to control pollutants emissions such as $NO_x$, $SO_x$ and ash particles. Coal also plays an important role in global $CO_2$ emissions, which will need to be reduced to limit the effects of climate change [66]. In order to meet those requirements, understanding the pulverized-coal combustion mechanisms and developing advanced combustion technology is necessary. However, the combustion of pulverized-coal is a complex phenomenon compared with that of gaseous or liquid fuels, since dispersion of coal particles, devolutilization and oxidation reactions take place simultaneously [45,92]. Moreover, experimental measurements of pulverized-coal flame characteristics are extremely difficult [38,78,80,88,97,100,107]. Accordingly, the development of new combustion furnaces and burners is still empirical and requires a significant amount of time and money.

In coal-combustion systems, finely pulverized coal particles are conveyed by a medium, air in general, that enters the combustor as a jet. The distribution of particles and its dispersion affect the local environment surrounding each particle and therefore its combustion kinetics [84]. Although the reaction kinetics involved in coal combustion have been studied extensively, the fundamental mechanics of coal trajectories and of particle-laden jets in
general are as yet poorly understood and have not been included in optimizing the design and operating flow condition for coal combustors. This issue can be addressed by using Computational Fluid Dynamics (CFD), which was shown to be a very useful and accurate modeling framework for the description of particulate systems, and for the description of turbulent flows in general [24,29].

RANS (Reynolds-Averaged Navier-Stokes), LES (Large-Eddy Simulation) and DNS (Direct Numerical Simulation) are the three common CFD methods for turbulence modeling. DNS requires a very fine grid spacing and even though it has the highest numerical accuracy, its application to practical combustion systems is made impracticable by the high computer load required. RANS on the opposite end of the spectrum is widely used for practical applications. There is however evidence that RANS can be inaccurate in predicting particle-laden flows [5,86]. As a result, interest for LES is growing, as it directly solves the transport equations for the large eddies and models only the smallest eddies. Unsteady turbulent motions are evaluated and the number of model parameters is greatly reduced. LES has been convincingly shown to be superior to RANS in accurately predicting turbulent mixing and combustion dynamics [56,75]. Although LES requires a high computer load compared to RANS, its superiority in terms of prediction will likely make it the best choice for practical combustion applications as computational cost decreases.

In order to describe the evolution of the dispersed particulate phase, CFD must be coupled with a population-balance equation (PBE). Several numerical approaches can be used to solve this equation, such as classes methods (CM) or Monte-Carlo methods (MCM) [77]. Classes methods, in which the internal coordinate (e.g., particle length or volume) is discretized into a finite series of bins, are the most popular [40,53,54], whereas MCM [11] are well known for their ease of implementation. In order to get reasonable results, the CM method requires a large number of classes (e.g., 20–30), so it is a computationally expensive approach for CFD calculations. Although the Monte-Carlo method is theoretically applicable, especially for Lagrangian-Eulerian simulations, in order to reduce the statistical error, a very large number of particles must be used. Due to limitations on the computational
resources, the number of particles which can possibly be tracked is often too low to get accurate particle statistics for applications such as coal boilers.

The method of moments (MOM) offers an attractive alternative where the PBE is tracked through its moments by integrating the internal coordinate [43]. The main advantage of MOM is that the number of scalars required is very small (i.e., usually 4–6), which makes the implementation in CFD codes feasible. However, due to the difficulties related with expressing transport equations in terms of the moments themselves, the method has been scarcely applied until recently. The so-called closure problem [62,94,98,103] has been reviewed [19] and a new method known as quadrature method of moments (QMOM) has been proposed [64], validated [8,61,63] and applied for studying a wide range of practical problems [62,94,98,101,102]. The main limitation of the classical QMOM is that it can treat only PBE tracking one property of the population of particles, such as particle mass, size, volume or area (that is, monovariate PBE). QMOM has then been extended to bivariate problems [103,108,109]. However, in a number of practical cases, it is interesting to describe the population of particles with multivariate PBE, where two or more properties of the population are simultaneously tracked. In order to address these issues, the direct quadrature method of moments (DQMOM) has been formulated and validated [26,59].

DQMOM is based on the direct solution of the transport equations for weights and abscissas of the quadrature approximation and presents the advantage of being directly applicable to multivariate PBE. It has been shown to be a powerful approach for describing polydisperse solids undergoing segregation, growth, aggregation and breakage processes in the context of CFD simulations [26,32,60]. Pulverized coal flames are considered dilute systems as the volume of particles is less than 1% of the volume of the fluid. Aggregation and breakage phenomena are rare events and are therefore usually neglected. However, even in this context where drag is the main process affecting particles, the DQMOM equations are equivalent to the widely used Lagrangian particle method [23], but it has the advantage of precisely controlling the statistical noise in the lower order moments (e.g., particle number density, mass density, Sauter radius). For a given desired accuracy, this greatly reduces
the computational cost since it does not require a large number of particles [17,18]. In previous work, the advantages of using DQMOM for treating particle populations with low Stokes numbers (i.e., the dispersed-phase velocity follows closely the velocity of the continuous phase) have been clearly demonstrated [61,63,98,115]. However, coal particles, which typically range from 1 micrometer to 200 micrometers in pulverized coal flames, have a finite Stokes number and their velocity depends on the particle size. In quadrature methods, this dependency is accounted for by solving an Eulerian model where each quadrature node has its own velocity field. Fox et al. [32] have applied DQMOM to droplets spray with a finite Stokes number and compared the results with the multifluid method and a classical Lagrangian solver. Results show that DQMOM can provide a comparable accuracy with a significantly lower computational cost, making it a very good candidate for more complex two-phase combustion applications.

The goal of this study is to show that DQMOM can be used in combination with LES to predict particle behavior in coal flames. Since few data on particles in coal flames are available, experimental data of nonreacting turbulent flows with particles were chosen. The study presents LES simulations of coaxial particle-laden jets and compares the results to experimental data [14]. Special interest is given to the following important characteristics of particle-laden jets: gas and particle velocity, particle size segregation and particle clustering. These phenomena affect operational considerations within a coal-fired boiler such as flame stability, pollutants formation or heat flux characteristics [84]. Simulation of coaxial particle-laden jets are performed with three different velocity ratios between the primary and secondary stream and two particle size distributions (25 microns and 70 microns). Simulation results show predictions of both the gaseous and solid phase averaged velocity and are compared to experimental data.

2.3 Numerical modeling

Simulations were performed using ARCHES, a LES tool resulting from a ten-year partnership with the Department of Energy and the University of Utah. It is a massively parallel
code that solves conserved quantities (mass, momentum, energy, scalar) spatially and temporally in a turbulent flow field, allowing for detailed and accurate simulations of fires and flames [87]. The code is integrated into a C++ framework called Uintah which provides large-scale parallelization tools for physics components [9,16,73]. The code is maintained in a repository and distribution is freely available [41]. The governing equations and numerical methods for the gas phase and the dispersed phase are described in the following sections. Interactions between the two phases are taken into account.

2.3.1 Governing equations of the gas phase

Since the particles considered are nonreacting, the gas-phase governing equations are solved using filtered continuity and momentum equations for incompressible flows in a finite volume formulation. The mass balance can be written as:

\[ \frac{\partial \bar{u}_{g,i}}{\partial x_i} = 0 \]  

(2.1)

where \( u_{g,i} \) is the \( i^{th} \) gas velocity component, and the momentum equations as:

\[ \frac{\partial \bar{u}_{g,i}}{\partial t} + \frac{\partial}{\partial x_j}(\bar{u}_{g,i} \bar{u}_{g,j}) = -\frac{1}{\rho_g} \frac{\partial \bar{p}}{\partial x_i} + 2\nu \frac{\partial}{\partial x_j} \bar{S}_{ij} - \frac{\partial \tau_{ij}}{\partial x_i} + \bar{S}_{u_i} \]  

(2.2)

where \( \rho_g \) is the gas density, \( \nu \) is the gas kinematic viscosity and \( S_{ij} \) is the rate-of-strain tensor. \( \tau_{ij} \) is the residual stress tensor and is modeled using the dynamic subgrid scale model of Germano et al. [36]. The source term \( S_u \) is a momentum source term that accounts for momentum transfer from the dispersed phase.

2.3.2 Governing equations of the dispersed phase

The DQMOM method was implemented into ARCHES to solve the dispersed phase. A detailed description of the particles representation and of the DQMOM equations is given in the following paragraphs.
2.3.2.1 Number density function (NDF)

The particle Number Density Function (NDF) describes the number of particles per volume as a function of the spatial location and of other independent variables. These independent variables are intrinsic characteristics of the particles (e.g., particle diameter, particle velocities, etc.) and are called internal coordinates. The vector of internal-coordinate random values is denoted by $\zeta$, and the internal coordinate sample space is denoted by $\xi$.

When the particle velocities are considered as internal coordinates, the random values are denoted by $u_p$, and the particle velocity sample space is denoted by $v_p$. The full NDF is denoted $n(v, \xi; x, t)$. At a given location in space and time $(x_0, t_0)$, the total number of particles per volume is:

$$n_p(x_0, t_0) = \int_{v_p} \int_{\xi} n(v_p, \xi; x_0, t_0) d\xi dv_p$$

(2.3)

NDFs can be univariate or multivariate. Univariate NDFs are only functions of one internal coordinate, so the internal coordinate sample space $\xi$ has a single dimension. Multivariate NDFs, however, are functions of multiple internal coordinates, so the internal coordinate sample space has $N_\xi$ dimensions:

$$n(\xi; x, t) = n(\xi_1, \ldots, \xi_{N_\xi}; x, t)$$

(2.4)

or, including the particle velocity as an internal coordinate,

$$n(v_p, \xi; x, t) = n(v_{p,1}, v_{p,2}, v_{p,3}, \xi_1, \ldots, \xi_{N_\xi-3}; x, t)$$

(2.5)

The particle Probability Density Function (PDF) $p_{u\zeta}$ denotes the probability of the velocity-scalar vector taking on a particular value. At a fixed point in space and time $(x_0, t_0)$, the PDF is related to the NDF by:
\[ n(v_p, \xi; x, t_0) = n_p(x_0, t_0)p_{u\zeta}(v_p, \xi; x_0, t_0) \quad (2.6) \]

### 2.3.2.2 NDF transport equation

The PDF transport equation is given by [76]:

\[
\frac{\partial}{\partial t} (p_{u\zeta}(v_p, \xi; x, t)) + \frac{\partial}{\partial x_i} (\langle v_{p,i}|v_p, \xi \rangle p_{u\zeta}(v_p, \xi; x, t)) = -\frac{\partial}{\partial v_i} \left( \langle A_i|v_p, \xi \rangle p_{u\zeta}(v_p, \xi; x, t) \right) \\
- \frac{\partial}{\partial \xi_k} \left( \langle G_k|v_p, \xi \rangle p_{u\zeta}(v_p, \xi; x, t) \right) 
\]  

(2.7)

In this equation, the variable \( \langle v_{p,i}|v_p, \xi \rangle \) represents the entire velocity sample space and is thus a full distribution. The quantities \( \langle A_i|v_p, \xi \rangle \) and \( \langle G_k|v_p, \xi \rangle \) are conditional quantities that describe the “velocity” of the PDF in the phase space \( (v_p, \xi) \). \( A_i \) is defined by:

\[ A_i = \frac{dv_{p,i}}{dt} \quad (2.8) \]

As the right side depends on \( v_p \) and \( \xi \), this quantity is a distribution. The particular value of \( A_i \) depending on \( v \) and \( \xi \) can be expressed as \( \langle A_i|v_p, \xi \rangle \). Likewise, \( G_k \) is defined by:

\[ G_k = \frac{d\xi_k}{dt} \quad (2.9) \]

and is also a distribution, with the particular value of \( G_k \) expressed as a conditional quantity \( \langle G_k|v_p, \xi \rangle \). Eq. (2.7) can be multiplied by the function \( n_p(x, t) \) and combined with the number balance equation to yield the NDF transport equation [77]:

\[
\frac{\partial}{\partial t} (n(v_p, \xi; x, t)) + \frac{\partial}{\partial x_i} (\langle v_{p,i}|v_p, \xi \rangle n(v_p, \xi; x, t)) = -\frac{\partial}{\partial v_i} \left( \langle A_i|v_p, \xi \rangle n(v_p, \xi; x, t) \right) \\
- \frac{\partial}{\partial \xi_k} \left( \langle G_k|v_p, \xi \rangle n(v_p, \xi; x, t) \right) \\
+ h(v_p, \xi; x, t) 
\]  

(2.10)
where $h$ is a source term representing the birth and death of particles in the domain. This term is set to zero as particle breakage and aggregation effects are neglected.

### 2.3.2.3 Direct Quadrature Method of Moments

The Direct Quadrature Method of Moments (DQMOM) approximates the NDF by a summation of multidimensional Dirac delta functions. Figure 2.1 gives an example for bivariate Gaussian distribution (i.e., two internal coordinates) of the NDF and its quadrature approximation with three quadrature nodes. The quadrature approximation can be regarded as an extension of the mono-kinetic assumption for the multifluid model. In the multifluid model, the closure of the system is obtained through the following assumptions [57]:

![Fig. 2.1: DQMOM approximation of the NDF with 3 quadrature nodes](image-url)
- For a given particle size, at a given point \((x,t)\), there is only one characteristic averaged velocity,
- The velocity dispersion around the averaged velocity is zero in each direction at every point.

This is equivalent to presuming that the NDF is described by:

\[
n(v_p, \xi; x, t) = n_p(\xi; x, t) \delta (v_p - \langle v_p \rangle) \tag{2.11}
\]

The quadrature approximation extends the mono-kinetic assumption to all the internal coordinates:

- For a given quadrature node \(\alpha\), at a given point \((x,t)\), there is only one characteristic averaged internal coordinate value \(\langle \xi \rangle_\alpha\), also called abscissa,
- The internal coordinate dispersion around the averaged value \(\langle \xi \rangle_\alpha\) is zero.

With those assumptions, the NDF is written as :

\[
n(v_p, \xi; x, t) = \sum_{\alpha=1}^{N} \left( w_\alpha \delta (\xi - \langle \xi \rangle_\alpha) \delta (v_p - \langle v_p \rangle_\alpha) \right) \tag{2.12}
\]

\[
n(v_p, \xi; x, t) = \sum_{\alpha=1}^{N} \left( w_\alpha \prod_{i=1}^{N_\xi-3} \delta (\xi_i - \langle \xi_i \rangle_\alpha) \prod_{i=1}^{3} \delta (v_{p,i} - \langle v_{p,i} \rangle_\alpha) \right) \tag{2.13}
\]

where \(w_\alpha\) is the number of particles per volume associated with the quadrature node \(\alpha\).
\(w_\alpha\), \(\langle \xi \rangle_\alpha\) and \(\langle v_p \rangle_\alpha\) depend on space and time, but the dependence is omitted for clarity of notation.

The validity of the mono-kinetic assumption has been discussed for the multifluid method [17]; the hypothesis is valid for particles with low Stokes number and becomes inappropriate when Stokes number increases \((St > 0.4)\) and particle trajectory crossing occurs. It is however important to note that this limitation can be partially overcome with DQMOM. In the multifluid model, only the particle size space is discretized, whereas DQ-
MOM allows us to discretize a multidimensional space. It is therefore possible to discretize the particle velocity space if the particle velocity is chosen as an internal coordinate and then to have several quadrature nodes with different velocities and otherwise identical internal coordinates values. As quadrature nodes can cross each other, the error associated with finite Stokes number particles crossing each others is reduced. This strategy was applied for large particle-size distribution and its validity is discussed in the second part of this work.

2.3.2.4 Quadrature-approximated NDF transport equation

The quadrature-approximated NDF transport equation can be derived for a multivariate NDF by combining the NDF transport equation (Eq. (2.10)) with the quadrature approximation (Eq. (2.13)). The quadrature node averaged phase space velocities $\langle A_i \rangle_\alpha$ and $\langle G_i \rangle_\alpha$ are introduced:

$$
\langle A_i \rangle_\alpha = \langle A_i | v_p = \langle v_p \rangle_\alpha, \xi = \langle \xi \rangle_\alpha \rangle
$$ (2.14)

$$
\langle G_i \rangle_\alpha = \langle G_i | v_p = \langle v_p \rangle_\alpha, \xi = \langle \xi \rangle_\alpha \rangle
$$ (2.15)

The multivariate quadrature-approximated NDF transport equation is then given by a set of equations:

$$
\frac{\partial}{\partial t} (w_\alpha) + \frac{\partial}{\partial x_i} (\langle v_{p,i} \rangle_\alpha w_\alpha) = a_\alpha
$$ (2.16)

$$
\frac{\partial}{\partial t} (\varsigma_{n\alpha}) + \frac{\partial}{\partial x_i} (\langle v_{p,i} \rangle_\alpha \varsigma_{n\alpha}) = b_{n\alpha}
$$ (2.17)

where $\varsigma_{n\alpha} = w_\alpha \langle \xi_n \rangle_\alpha$ is the weighted abscissa for the $n^{th}$ internal coordinate. The terms on the right-hand side are determined by the following equations:
where

\[ S_\xi = -\sum_{\alpha=1}^{N_\xi} \frac{\partial}{\partial v_i} \left[ \langle A_i \rangle_{\alpha} w_{\alpha} \left( \prod_{j} \delta \left( v_{p,j} - \langle v_{p,j} \rangle_{\alpha} \right) \right) \left( \prod_{j} \delta \left( x_j - \langle x_j \rangle_{\alpha} \right) \right) \right] \\
- \sum_{\alpha=1}^{N_\xi} \frac{\partial}{\partial x_i} \left[ \langle G_i \rangle_{\alpha} w_{\alpha} \left( \prod_{j} \delta \left( v_{p,j} - \langle v_{p,j} \rangle_{\alpha} \right) \right) \left( \prod_{j} \delta \left( x_j - \langle x_j \rangle_{\alpha} \right) \right) \right] \]  \tag{2.19}

### 2.3.2.5 Moment-transformed NDF transport equation

The quadrature-approximated NDF transport equation (Eq. (2.18)) is a single equation but multiple unknowns need to be determined (weights and abscissas). In order to obtain a number of independent equations equal to the number of weights and abscissas, a set of independent moments are chosen and the moment transform of the quadrature-approximated NDF transport equation yields a set of independent equations, equal in number to the independent moments. A number of moments equal to \((N_\xi + 1)N\) is required, where \(N_\xi\) is the number of internal coordinates and \(N\) the number of environments. This procedure results in the moment-transformed quadrature-approximated NDF transport equation [60]:

\[
\sum_{\alpha=1}^{N_\xi} \left[ \prod_{j=1}^{N_\xi} \langle x_j \rangle_{\alpha} \right] - \sum_{m=1}^{N_\xi} \left[ \prod_{j=1,j\neq m}^{N_\xi} \langle x_j \rangle_{\alpha} \right] a_{\alpha} \\
+ \sum_{\alpha=1}^{N_\xi} \sum_{n=1}^{N_\xi} \left[ k_n \left( \prod_{j=1,j\neq n}^{N_\xi} \langle x_j \rangle_{\alpha} \right) \right] b_{n\alpha} = S_k \]  \tag{2.20}

where:
\[
S_k = \sum_{\alpha=1}^{N} \sum_{n=1}^{N} \left( \sum_{j=1, j \neq n}^{N} \left( k_n \langle \xi_j^{k_n} \rangle \right) \right) w_\alpha (G_n)_\alpha 
\]

(2.21)

This set of equations forms a linear system that can be solved to get the source terms of the weights and weighted abscissas transport equations (Eqs. (2.16) and (2.17)) \(a_\alpha\) and \(b_{n\alpha}\).

### 2.3.2.6 Particle velocity model

In this work, the particles considered are nonreacting glass beads. The aggregation/breakage phenomena are neglected and the only physical process modeled is the drag force, which can be described by the Stokes drag law. For a mesoscale size particle, other forces such as lift can be neglected and the momentum equation for the particle can be expressed by an ordinary differential equation.

\[
\frac{du_{p,i}}{dt} = \frac{f_{\text{drag}}}{\tau_p} (u_{g,i} - u_{p,i}) + \frac{g_i (\rho_p - \rho_g)}{\rho_p}
\]

(2.22)

where \(g\) is the gravity force acting on the particle and \(f_{\text{drag}}\) is the coefficient of the drag force, which has a close relationship with the particle Reynolds number [85]:

\[
f_{\text{drag}} = \begin{cases} 
1 & \text{Re}_p < 1 \\
1 + 0.15Re_p^{0.687} & 1 < \text{Re}_p < 1000 \\
0.0183Re_p & \text{Re}_p > 1000 
\end{cases}
\]

(2.23)

\[
\text{Re}_p = \frac{\rho_p d_p |u_p - u_g|}{\mu_g}
\]

(2.24)

where \(\mu_g\) is the gas dynamic viscosity.

In Eq. (2.22), \(\tau_p\) is the particle relaxation time:
\[
\tau_p = \frac{\rho_p d_p^2}{18 \mu_g}
\]  

(2.25)

The particle density \(\rho_p\) and particle diameter \(d_p\) are assumed to be constant. This model can easily be implemented into the DQMOM formulation through the term:

\[
\langle A_i \rangle_\alpha = \frac{f_{\text{drag}}}{\tau_p} (\bar{u}_{i,g} - \langle v_{p,i} \rangle_\alpha) + \frac{g_i (\rho_p - \rho_g)}{\rho_p}
\]  

(2.26)

The gas phase and the particle phase are then coupled through the source term \(\bar{S}_{u_i}\) in the gas momentum equation (Eq. (2.2)):

\[
\bar{S}_{u_i} = -\frac{1}{\rho_g} \sum_{\alpha=1}^{N} w_\alpha \rho_p \frac{f_{\text{drag}}}{\tau_p} (\bar{u}_{i,g} - \langle v_{p,i} \rangle_\alpha)
\]  

(2.27)

### 2.3.3 Stokes number analysis

The major advantage of LES over RANS is that a large part of the turbulent kinetic energy spectrum (usually about 80%) is resolved as opposed to RANS where it is modeled. However, a subgrid scale model (SGS) is still needed to model the effects of the smallest eddies on the flowfield. The unclosed subgrid scale terms appear when the spatial filter is applied to the Navier-Stokes equations (Eq. (2.2)). A similar issue arises when filtering the NDF transport equation (Eq. (2.10)):

\[
\frac{\partial}{\partial t} (\bar{n}(v_p, \xi; x, t)) + \frac{\partial}{\partial x_i} (\bar{v}_{p,i} \bar{n}(v_p, \xi; x, t)) = \frac{\partial}{\partial v_i} \left( \langle A_i | v_p, \xi \rangle \bar{n}(v_p, \xi; x, t) \right) - \frac{\partial}{\partial \xi_k} \left( \langle G_k | v_p, \xi \rangle \bar{n}(v_p, \xi; x, t) \right) + \frac{\partial \tau_{\text{sgs}, i}}{\partial x_i} + \frac{\partial \tau_{\text{sgs}, u_i}}{\partial u_i} + \frac{\partial \tau_{\text{sgs}, \xi_k}}{\partial \xi_k}
\]  

(2.28)

The subgrid scalar flux \(\tau_{\text{sgs}, i}\) represents the flux of the number density as a result of unresolved velocity fluctuations. Likewise, the subgrid scalar fluxes \(\tau_{\text{sgs}, u_i}\) and \(\tau_{\text{sgs}, \xi_k}\) represent the subgrid flux of the number density in phase space \((v_p, \xi)\).
Few studies have focused on the effects of subgrid scale velocity fluctuations on particle motion in LES so far. Yei and Lei [106] used LES to investigate the motion of particles in isotropic and homogeneous shear flows. They generated particle statistics by neglecting the subgrid scale (SGS) effects on particles and showed that LES can successfully predict second-order statistics of the particle motion, such as root-mean-square velocity fluctuations, the mean square displacement and the dispersion coefficient. They also showed that particle dispersion was dominated by the large eddies. Later, Yand and Lei [105] studied the phenomenon of particle accumulation in low vorticity regions and found that it is controlled mainly by the small scales of turbulence with wave number $k_\omega$ corresponding to the maximum of the dissipation (vorticity) spectrum. The small eddies with wave number greater than $2.5k_\omega$ had no effect on the particle accumulation and the average settling velocity. Since the length scale $l_\omega = 1/k_\omega$ is typically one order of magnitude greater than the Kolmogorov length scale $\eta$, they concluded that LES was adequate for particle-laden flows and that the subgrid-scale effects can be neglected, provided that the cutoff wave number is greater than $2.5/l_\omega$. As a result, many LES studies of particle-laden flows neglected the SGS effects for various flow configurations [5, 10, 82, 99]. More recently, Shotorban and Mashayek [81] used a stochastic model for particle motion in LES and found that this assumption may break down when particles have small time constant and/or the filtered energy is significant. The error was however important for $\Delta = 8\eta$ but very limited for $\Delta = 4\eta$, which tends to agree with Yang and Lei’s results.

The previous studies only considered a limited range of turbulence intensity or particle size. A general theoretical justification for direct use of the drag law in LES can be made by a Stokes number analysis. The particle Stokes number combines the effects of particle size, particle density and turbulence intensity, and simultaneously accounts for the variety of turbulent timescales. The Stokes number is defined as:

$$St = \frac{\tau_p}{\tau_f} \quad (2.29)$$
where $\tau_f$ is a characteristic time of the fluid. The difficulty of Stokes number analysis often lies in determining $\tau_f$. The turbulent kinetic energy (TKE) spectrum, denoted as $E_U$, is convenient for this analysis since it breaks down the turbulent fluctuations into contributions of different sizes – with each size exhibiting a corresponding eddy turnover time. When the eddy turnover time is used as the fluid characteristic time in the Stokes number, and when each wave number in the turbulent-energy spectrum has an associated eddy turnover time, then a single particle of fixed radius and density will exhibit a unique Stokes number value for each different wave number. To illustrate this concept, the model energy spectrum for isotropic turbulence is shown in Figure 2.2a. This figure shows a dotted line corresponding to the wave number for which the eddy turnover time is equal to a particle’s relaxation time – Stokes number equals one for that particle. For wave numbers much larger than the dashed line, the eddy turnover time is much smaller than the particle relaxation time and the particle behaves ballistically (with respect to the fluctuations at those wave numbers). On the other hand, for wave numbers much smaller than the dashed line, the eddy turnover time is much larger than the particle relaxation time and the particle behaves as a tracer element. It is only for a relatively narrow bandwidth that the turbulence interacts nontrivially with the particle.

The implications of this concept on LES modeling are illustrated in Figure 2.2b. In these three plots, the model energy spectrum is shown with a hypothetical Nyquist cut-off. Three particle sizes are considered, one for each of the three plots, with the overlap bandwidth of nontrivial Stokes number illustrated in relation to the Nyquist cut off. The small particles, top plot, behave as tracers for all of the resolved scales. Therefore, the drag of particles in this unresolved regime should be modeled in the subgrid-scale flux. In the extreme of zero Stokes number, the subgrid-scale flux is often modeled using gradient diffusion. The large particles, bottom plot, are in a regime such that they are ballistic with regard to all the unresolved scales – their trajectories are essentially unaffected by the presence or absence of the unresolved scales. For this reason, particles in the resolved regime should be modeled using the drag directly – as done in direct numerical simulations. The
Fig. 2.2: Illustration of the different particle kinematic regimes as a function wave number for a fixed particle relaxation time (a) and the implications on LES modeling (b)
middle plot shows particles of intermediate size for which some of the nontrivial drag effects are resolved and some are not. Particles in this partial-resolution-of-drag regime present a challenging modeling problem. It will be shown in the results section, 2.5.2, that particles in the current study are in the resolved regime. Therefore, the drag law is used directly and the subgrid scale effects of turbulence on particles were neglected.

2.4 Experimental and simulation conditions

In this study, LES coupled with DQMOM is applied to simulate particle-laden coaxial jets and results are compared to experimental data obtained by Budilarto [14].

2.4.1 Experimental setup

In the experiments, particles were conveyed with air in the primary nozzle whereas the secondary nozzle was fed only with air (Figure 2.3). Air and particles were injected at room temperature in an 18x18 inch chamber. The experiments focused on the effects of the inlet velocity and the particle sizes on the flow-field and the particle motion in the near field region of the nozzle. The velocity ratio, defined as the ratio of the annular to central inlet velocity \( VR = U_a/U_0 \), was modified by varying the volumetric flow rate of the annular jet. Three different velocity ratios were studied: 0, 1.0 and 1.5. The Reynolds number based on the average central inlet velocity was 8,400. Table 2.1 lists the flow conditions for the experiments. Two particle-size distributions were used for this purpose, one with a mean diameter of 25 microns and the other with a mean diameter of 70 microns. Particles used were glass beads with a density of 2500 kg/m³. The particle mass loading was maintained at 0.5 for all velocity ratios. Laser Doppler Velocimetry and Phase Doppler Particle Analyzers were used to measure time-averaged velocities.

Those experimental data were chosen for several reasons. First, measuring coal particles in coal flames is challenging and very few studies have been reported. Even though our final goal is to simulate coal flames, nonreacting systems offer a more reliable and accurate source of data for particle velocities, segregation and clustering effects.
Table 2.1: Experimental setup for the different velocity ratios

<table>
<thead>
<tr>
<th>Case</th>
<th>d (inch)</th>
<th>D (inch)</th>
<th>Central jet maximum inlet velocity $U_0$ (m/s)</th>
<th>Annular jet maximum inlet velocity $U_a$ (m/s)</th>
<th>Velocity ratio (VR)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0.56</td>
<td>1.254</td>
<td>11.7</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>2</td>
<td>0.56</td>
<td>1.254</td>
<td>11.7</td>
<td>11.6</td>
<td>1.0</td>
</tr>
<tr>
<td>3</td>
<td>0.56</td>
<td>1.254</td>
<td>11.7</td>
<td>18.0</td>
<td>1.5</td>
</tr>
</tbody>
</table>

Budilarto’s data include extensive results in different operating conditions (3 velocity ratios) with two different particle sizes distributions, which allows us to test the model sensitivity and accuracy with respect to turbulence intensity and particle size. Then, the experiments offer a simple geometry with well-defined boundary conditions and are therefore suited for numerical simulations. Finally, the operating conditions are very close to those of pulverized coal flames in terms of velocities, turbulence intensity, particle mass loading,
particle sizes and volume fraction. In addition, coaxial burners are also used in coal flame experiments on a laboratory scale [72,112,113], so this study provides a first step towards the simulation of coaxial coal flames.

2.4.2 Simulation details

The computational domain is a 0.28 m wide cube which is divided with a cartesian mesh into 8 millions cells, resulting in a 1.42 mm cellsize. A total of 9 simulations were performed; for each velocity ratio, simulations without particles, with 25-micron particles and with 70-micron particles were run on 512 CPUs for about 24 hours. Each simulation was performed for at least 15,000 timesteps, each timestep being about $25 \mu s$ and at least 5,000 steps were used for time averaging. Measurements have shown that the common power law velocity profile of fully developed turbulent flow can be applied at the central inlet:

$$\frac{U}{U_0} = \left(1 - \frac{2r}{d}\right)^{1/7} \tag{2.30}$$

The dispersed phase was modeled using the DQMOM method with four internal co-ordinates: the particle diameter $d_p$ and the three particle velocity components $v_x, v_y, v_z$. Particles were assumed to be perfect spheres of constant size and density.

Special care must be taken when choosing the quadrature nodes to represent the full NDF. As discussed earlier, for small particles, the mono-kinetic assumption is applicable and all the particles of a same size can be assumed to have a unique velocity at a given location. On the other hand, larger particles have more inertia and their trajectory history may result in multiple velocities for same size particles at a given location. In this case, the mono-kinetic assumption would not be valid anymore. It is however possible to overcome this limitation with the DQMOM approach by discretizing the velocity phase space into several quadrature nodes. Taking those considerations into account, a different modeling approach is used to represent the 25-micron particle size distribution and the 70-micron distribution.
For the 25-micron distribution, the particle diameters range from a few microns up to 45 microns. The mass of the smallest particles is several orders of magnitude less than the mass of the largest particles. Since a 10-micron particle would behave significantly differently from a 40-micron particle, the size distribution was represented by three quadrature nodes in the DQMOM approach, each node corresponding to a different particle size. Table 2.2 shows the particle diameter associated with each quadrature node as well as their weight at the inlet. Even though the mean diameter is 25 microns, it can be noticed that there are more 14-micron particles than 25-micron particles.

For the 70-micron distribution, the particle diameters range from 50 to 90 microns. The mass of all the particles is of the same order of magnitude, so the size distribution can be represented by a single particle size of 70 microns. However, these particles have more inertia than the small ones and, as they collide and bounce against the walls inside the nozzle, their initial velocity at the exit of the nozzle follows a distribution that should be taken into account. For the 25-micron particles, measured velocity fluctuations at the inlet were indeed small, so assuming that all the particles have the same inlet velocity is a good approximation. On the contrary, for the 70-micron particles, velocity fluctuations at the inlet were important and it is necessary to represent their velocity distribution. Based on the measured root mean square of velocity fluctuations at the inlet, a 2-D normal distribution in particle velocity components $v_{p,y}$ and $v_{p,z}$ was assumed. Fluctuations in $v_{p,x}$ were neglected since they are small compared to the mean velocity in this direction. The normal distribution was approximated by 5 quadrature nodes in the DQMOM approach, as shown in Table 2.3.

Table 2.2: Quadrature approximation for 25-micron particles (inlet values)

<table>
<thead>
<tr>
<th>Quadrature node</th>
<th>Particle diameter ($\mu$m)</th>
<th>Weight (# particles/m$^3$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>14</td>
<td>4.11E10</td>
</tr>
<tr>
<td>1</td>
<td>25</td>
<td>1.44E10</td>
</tr>
<tr>
<td>2</td>
<td>34</td>
<td>2.90E9</td>
</tr>
</tbody>
</table>
Table 2.3: Quadrature approximation for 70-micron particles (inlet values)

<table>
<thead>
<tr>
<th>Quadrature node</th>
<th>Particle diameter ($\mu$m)</th>
<th>$v_{p,x}$ (m/s)</th>
<th>$v_{p,y}$ (m/s)</th>
<th>$v_{p,z}$ (m/s)</th>
<th>Weight ( # particles/m$^3$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>70</td>
<td>9.6</td>
<td>0</td>
<td>0</td>
<td>3.8E8</td>
</tr>
<tr>
<td>1</td>
<td>70</td>
<td>9.6</td>
<td>-0.55</td>
<td>0</td>
<td>2.3E8</td>
</tr>
<tr>
<td>2</td>
<td>70</td>
<td>9.6</td>
<td>0.55</td>
<td>0</td>
<td>2.3E8</td>
</tr>
<tr>
<td>3</td>
<td>70</td>
<td>9.6</td>
<td>0</td>
<td>-0.55</td>
<td>2.3E8</td>
</tr>
<tr>
<td>4</td>
<td>70</td>
<td>9.6</td>
<td>0</td>
<td>0.55</td>
<td>2.3E8</td>
</tr>
</tbody>
</table>

2.5 Results and discussion

Simulations were performed for single-phase coaxial jets and for particle-laden coaxial jets to assess the uncertainties associated with turbulence modeling and particle modeling.

2.5.1 Single phase coaxial jet

Simulations were performed for the three velocity-ratios cases without particles. The two main objectives are first to validate the modeling approach by ensuring that the gas mean velocities are predicted correctly and secondly to have a reference case to then understand the effect of particles on the gas dynamics. Figures 2.4, 2.5 and 2.6 show instantaneous profiles of the x-component of the gas velocity. For VR=0, the flow field is identical to a round jet, whereas for VR=1.0 and VR=1.5, the effect of the secondary stream on the flowfield can be observed. Simulations results were evaluated against Budilarto’s experimental data by comparing the mean gas velocity along the centerline (Figure 2.7). Velocity profiles are predicted with a maximum error less than 10%. This error can be attributed to several factors, among which grid resolution, turbulence modeling or experimental measurements uncertainty. Results however predict the velocity profile and jet spreading accurately enough to validate the modeling approach.
Fig. 2.4: Gas x-velocity profile (m/s) for VR=0, no particles
Fig. 2.5: Gas x-velocity profile (m/s) for VR=1.0, no particles
Fig. 2.6: Gas x-velocity profile (m/s) for VR=1.5, no particles
Fig. 2.7: Comparison between LES and experimental data of gas averaged x-velocity along the centerline for different velocity ratios / No particles

### 2.5.2 Stoke number analysis

Results from the single phase coaxial jet can be used to perform a Stokes number analysis *a posteriori*. For this analysis, we assume local isotropy of turbulence. The turbulent kinetic energy spectrum can be modeled by a simple model spectrum [76]:

\[
E(\kappa) = C\varepsilon^{2/3}\kappa^{-5/3}f_L(\kappa\lambda)f_\eta(\kappa\eta)
\]  

(2.31)
where $\varepsilon$ is the dissipation rate, $\kappa$ is the wave number, $L$ is the characteristic length scale of the large eddies and $\eta$ is the Kolmogorov length scale. $f_L$ and $f_\eta$ are specified nondimensional functions:

$$f_L(\kappa L) = \left( \frac{\kappa L}{(\kappa L)^2 + c_L} \right)^{5/3+p_0}$$  \hspace{1cm} (2.32)

$$f_\eta(\kappa \eta) = \exp \left( -\beta \left( \left(\frac{\kappa \eta}{c_{\eta}}\right)^4 + c_{\eta}^{1/4} - c_\eta \right) \right)$$  \hspace{1cm} (2.33)

The constant values of $C$, $p_0$ and $\beta$ are taken to be $C = 1.5$, $p_0 = 2$, $\beta = 5.2$. For high Reynolds number, the values of the constants $c_L$ and $c_\eta$ can be approximated by $c_L \approx 6.78$ and $c_\eta \approx 0.40$ [76]. For specified values of $\varepsilon$, $L$ and $\eta$, the model spectrum is determined by Eqs. (2.31), (2.32) and (2.33).

The dissipation rate can be estimated from the LES simulations by the simplified formula:

$$\varepsilon = 15\nu \left( \frac{\partial u_{g1}}{\partial x_1} \right)^2$$  \hspace{1cm} (2.34)

For the LES simulation of the single phase coaxial jet with the highest turbulence intensity, i.e., $VR=1.5$, the maximum of the dissipation rate along the centerline is calculated to be $\varepsilon \approx 40 m^2/s^3$. The Kolmogorov length scale can be estimated from this value:

$$\eta \approx \left( \nu^3 / \varepsilon \right)^{1/4} \approx 0.1 mm$$  \hspace{1cm} (2.35)

The characteristic length scale of the large eddies is obtained by:

$$L \approx \frac{\eta}{Re^{-3/4}} \approx 0.09 m$$  \hspace{1cm} (2.36)
The resulting normalized model energy spectrum \(E_U = E/\left(\eta u_\eta^2\right)\) is plotted on Figure 2.8a, where \(u_\eta \approx (\varepsilon \nu)^{1/4}\) is the Kolmogorov velocity. The relationship between the dissipation energy and the energy spectrum (Eq. (2.37)) can be used to verify that the estimated values for \(\varepsilon, L\) and \(\eta\) are consistent.

\[
\varepsilon = \int_0^\infty 2\nu \kappa^2 E(\kappa) d\kappa
\]  
(2.37)

The value of the integral in Eq. (2.37) returns \(\varepsilon = 39.4\, J/s\), which is close to the LES simulation value \(\varepsilon \approx 40\, J/s\).

The wave number \(\kappa_\Delta = 1/\Delta\) corresponding to the Nyquist cut-off limit is indicated by a green line in Figure 2.8, where \(\Delta\) is the LES filter length (\(\Delta = 1.4\, mm\)). The model spectrum indicates that more than 90% of the energy spectrum is resolved for the LES simulations performed in this study and can further be used to determine the eddy characteristic time \(\tau_f\) as a function of the wave number by assuming:

\[
\tau_f(\kappa) \approx \frac{1}{\kappa \left[2k(\kappa)\right]^{0.5}}
\]  
(2.38)

where \(k(\kappa) \approx \kappa E(\kappa)\) is the turbulent kinetic energy associated with eddy of wave number \(\kappa\). Eq. (2.38) combined with the Stokes number definition (Eq. (2.29)) allows us to plot the Stokes number as a function of the wave number for 25-micron and 70-micron particles, as shown on Figures 2.8b and 2.9b, respectively, \((\rho_p = 2500\, kg/m^3)\). The black dotted lines represent the wave number for which the particle Stokes number is equal to one. The top images of Figures 2.8 and 2.9 show the respective locations of the Nyquist limit and the unity Stokes number wavelength for the 25-micron particles (Figure 2.8a) and the 70-micron particles (Figure 2.9a). In both cases, the Nyquist cut-off wave number is higher than the unity Stokes wave number by a safe margin. It can be concluded that both particle sizes are in the resolved regime and that the specified mesh resolution for the LES simulations in this study allows us to resolve all the eddies affecting the particles motion.
Fig. 2.8: Location of the Nyquist cut-off and the unity Stokes wave number on the model energy spectrum for 25-micron particles (a) and Stokes number as a function of wave number for 25-micron particles (b)
Fig. 2.9: Location of the Nyquist cut-off and the unity Stokes wave number on the model energy spectrum for 70-micron particles (a) and Stokes number as a function of wave number for 70-micron particles (b)
2.5.3 Two-phase coaxial jet

Simulation results of the coaxial jet for the three velocity ratios and the two particle size distributions are presented in this section. Results for the six simulations are analyzed both qualitatively and quantitatively. In general, the qualitative analysis of the particle behavior predicted by the simulations shows very good agreement with the experimental observations.

Figures 2.10 and 2.11 show instantaneous snapshots of particle concentration for a given quadrature node in the XZ plane for the simple jet case (VR=0). Images for the 25-micron particles case are presented in Figure 2.10, whereas images for the 70-micron particles case are shown in Figure 2.11. For the 25-micron particle size distribution, particle segregation is observed: the smaller particles (quadrature node 0) disperse faster than the larger ones (quadrature node 2). The 14-micron particles follow closely the gas flowfield and form a wider jet than the 34-micron particles which remain mostly along the centerline. The 70-micron particles have more inertia and tend to keep their initial velocity. Figures 2.11a, 2.11b and 2.11c, respectively, show particle concentration for quadrature nodes 0, 3 and 4. Quadrature nodes 1 and 2 are not shown but would be similar to quadrature nodes 3 and 4 in the XY plane. For each quadrature node, little dispersion is seen around the main jet as particles are less affected by turbulence. The effect of their initial velocity is more prominent and is felt on their trajectory until they reach a semi-equilibrium with the gas phase.

Figure 2.12 presents time-averaged particle concentrations at different locations for VR = 0. \( w_{\text{sum}} \) is the sum of the weights of each quadrature node and represents the total number of particles and is defined as:

\[
w_{\text{sum}} = \sum_{\alpha=1}^{N} w_{\alpha}
\]  

(2.39)

Figure 2.12a shows particle concentration in the radial direction at \( x = 15d \) for 25-micron particles. This plot confirms that particle dispersion decreases with particle size and that the dispersion of the three particle sizes is significantly different.
Fig. 2.10: Instantaneous profiles of weights (\# particle/m$^3$) for 25-micron particles, VR=0
Fig. 2.11: Instantaneous profiles of weights ($\# \text{ particle/m}^3$) for 70-micron particles, VR=0
Fig. 2.12: Time-averaged particle concentration at X = 15d [(a), (b)] and along the centerline [(c), (d)] for VR = 0.
Figure 2.12b presents a similar plot for the 70-micron particles case. The particles with no initial radial velocity \((w_0)\) show very little dispersion, whereas the particles with an initial y-velocity \((w_1, w_2)\) show a significant dispersion along the y-axis. As a result, the overall profile of \(w_{sum}\) is wider than the profile of \(w_0\) and it can be concluded that particle dispersion for 70-micron particles occurs because of their initial velocity whereas for the 25-micron particles, it is due to gas phase interactions. Figures 2.12c and 2.12d show averaged particle concentration profiles along the centerline. For the 25-micron particles, a peak is observed before the particle concentration decreases. This peak is located further down the jet as the particle size increases. As a matter of fact, the peak for \(w_0\) in the 70-micron particles case is not shown on the figure but would be located at about \(x = 20d\).

To validate the LES with DQMOM approach to model particles-laden jets, we first compare qualitatively the particle behavior predicted by the simulations in the different configurations to that observed experimentally. Figure 2.13 presents instantaneous screen-shots of the total particle concentration \(w_{sum}\) from LES simulations and photographs of particles taken during the Budilarto’s experiment for the three velocity ratios cases with the 25-micron particle size distribution. For the velocity ratio of 0, Budilarto noted that the region with the highest particle number density was located along the centerline and its shape resemble a cone (Figure 2.13b). The 25-micron particles were dispersed symmetrically and gradually in the radial direction as the axial distance increased. He also observed that the conical region did not have a smooth interface, but its shape followed a “pine-tree” like structure where wedges of particles with backward slope were present. This effect resulted from the partial interaction of the particles with the large eddies: particles were thrown outwards at the leading edge of the large eddies, but the large eddies did not have enough energy to bring the particles into the high vorticity region, i.e., the center of the large eddies. As can be seen on Figure 2.13a, the LES simulation also predicts the conical and symmetrical shape and the “pine-tree” structures are present as well. For \(VR=1\), Budilarto reported that for \(x<8d\), there was no change in the cross-sectional area of the structures of the 25-micron particles and that the particle number density was approximately constant.
in this region (Figure 2.13d). The presence of the annular jet inhibited the spreading of the particles in the radial direction. At x>8d, the jet of particles started to become wavy as the turbulent structures from the outer shear layer interact with the particles. The LES simulation predicts the same behavior (Figure 2.13c); the jet of particles remains compact with a constant particle concentration until x=8d and then it becomes wavy. For VR=1.5, Budilarto pointed out that the particles jet started becoming wavy at the end of the potential core, which was located at x=4d (Figure 2.13f). This waviness was attributed to interactions between particles and the large eddies of the inner shear layer. The radial dis-
persion of particles was significantly enhanced at \( x > 6d \), as particles interacted with eddies of the length scale on the order of the jet diameter. Due to the presence of more energetic turbulent structures, the radial spreading of the particles in \( VR = 1.5 \) was higher than that in \( VR = 1.0 \). The more energetic eddies were able to drag the particles to follow their motion and to create large displacement in the radial direction, which caused the particle jet to break and become asymmetrical. For \( VR = 0 \), the large eddies only had enough energy to displace particles near the jet edges whereas for \( VR = 1.5 \), the large eddies are able to displace particles with no preference if they are at the center of the jet or near the edges. All those observations are in agreement with the LES simulation: the particles jet becomes wavy at \( x = 4d \) and the waves grow at \( x > 6d \) (Figure 2.13e). The jet eventually breaks up into clusters of particles as observed experimentally.

Figure 2.14 presents the same results for the three cases with the 70-micron particle size distribution. For \( VR = 0 \), Budilarto reported that the particles flowed downstream with a straight-line trajectory and were not affected by the presence of the turbulent structures in the flow (Figure 2.14b). The 70-micron particles spread in the radial direction and the spreading was attributed to the presence of radial velocity fluctuations at the nozzle inlet. The LES simulation results (Figure 2.14a) confirm those statements; the particle jet is little affected by turbulence and the spreading results mainly from the initial particle velocity, as shown previously in Figure 2.10. For \( VR = 1.0 \), particles flowed with straight-line trajectories until \( x = 9d \). At \( x > 9d \), the turbulent structures formed in the outer shear layer began to affect the particles motion and small waves in the particles flow appeared (Figure 2.14d). The waviness was less pronounced than for the 25-micron particles case because of the particle inertia and did not increase the radial spreading. As can be seen on Figure 2.14c, LES results agree with the observations since small waves appear after \( x = 9d \) and the jet spreading is limited. Finally, for \( VR = 1.5 \), the particles structure became wavy at \( x > 7d-8d \) because of the higher turbulence intensity (Figure 2.14f). A significant increase in the radial spreading was observed, exceeding other velocity ratios, as the particles were dragged and dispersed by the large eddies. The enhancement in the radial spreading for
Fig. 2.14: Qualitative comparison of particle behavior for the 70-micron particles cases, LES screenshots of particle concentration on the left ($W_{sum}$), photographs from experiments on the right.
the 70-micron particles was less important than that for the 25-micron particles. The LES simulation (Figure 2.14e) also predicts that the particles jet becomes wavy at $x=7d$ and that the radial dispersion is greater compared to other velocity ratios but less important compared to the 25-micron particles case.

Overall, simulations were able to capture very different particle behaviors, such as jet spreading, waviness, break up or particle clustering, in the six configurations studied.

To further validate the LES coupled with DQMOM approach to model turbulent particle-laden flows, we now compare the time-averaged velocities along the centerline predicted by the simulations to the experimental measurements. Figure 2.15 presents plots of the measured and predicted mean gas and particles velocities. For the simulation results, the mean particle velocity was calculated by averaging the first moment of the particle velocity $v_i^1$ over time, where:

$$v_i^1 = \frac{\sum_{\alpha=1}^{N} w_{\alpha} \langle v_i \rangle_{\alpha}}{\sum_{\alpha=1}^{N} w_{\alpha}}$$

(2.40)

Since the inertia of the 25-micron particles is smaller than that of the 70-micron particles, they respond faster to the variations in the gas motion. This is clearly shown by the similarity in the centerline development of the axial mean velocity between the gas and the particles for the smaller particles (Figures 2.15a, 2.15c and 2.15e). In general, the development of the particles velocity along the centerline includes a first region where the gas-phase velocity is faster than the solid-phase velocity and, as the gas phase velocity decreases and the particles retain more momentum, the two curves cross each other and the particle velocity becomes higher than the gas velocity. The length of this region for the 25-micron particles is short because of their small slip velocity at the nozzle inlet. Since the 70-micron particles have a velocity at the nozzle exit that is about 20% lower than the gas velocity, the length of this region is greater, about $7d$ for $VR=0$ and $1.0$ and is increased to $13d$ for $VR=1.5$ (Figures 2.15b, 2.15d and 2.15f). A comparison with the single phase results (Figure 2.7) also shows that the particles have a strong influence on the gas phase velocity.
Fig. 2.15: Comparison between simulations and experimental data of time-averaged gas and particles velocities along the centerline for 25 µm particles [(a) (c) (e)] and 70 µm particles [(b) (d) (f)]
Particles indeed transfer some of their momentum to the gas phase and thus the gas velocity along the centerline decreases slower. For instance, for VR=0, the gas velocity magnitude at x=15d is less than 40% of its initial magnitude at the nozzle exit whereas it is more than 50% for both cases with particles. The LES simulations results are in good agreement with the experimental data; the maximum error over all the datapoints between the predicted velocities and the measured velocities for both phases is less than 12%. The uncertainty of the measured velocities can be assumed to be less than 5%, so the simulation predictions are slightly in disagreement with the experimental data. However, a comparison with Figure 2.7 shows that this disagreement can be attributed to the gas phase turbulence modeling since the maximum error for the single phase simulations was about 10%.

The radial spreading of particles is an important effect in coaxial-laden jets but also in coal flames as smaller particles will disperse faster and ignite first. Figure 2.16 presents the time-averaged concentration of particles $w_{sum}$ predicted by LES simulations for the different velocity ratios along the radial direction at x = 15d. The predicted particle concentrations are compared to the particle data rate obtained from the LDV measurements by Budilarto. Values are normalized by the centerline value where the particle concentration is the highest. For every velocity ratio, the fine particles have a wider distribution whereas the coarse particle jet remains more compact. Overall, simulations give good predictions and show that the modeling approach is able to capture particle segregation, even though the spreading rate is over-predicted for VR = 0 and under-predicted for VR = 1.0 and VR = 1.5. The comparison with the experimental data is however difficult to interpret for several reasons. First, the LDV data rate is not an accurate measurement of the number of particles as the laser resolution is not high enough to distinguish two small particles from a large one. Then, the error due to the gas phase turbulence modeling may be responsible for the error in the prediction of the particle dispersion. Finally, the predicted distribution is highly sensitive to the initial choice of environments in the quadrature approximation, as shown in Figure 2.12, and a larger number of environments to represent the particle distribution in size and velocity would likely lead to more accurate results. For instance, for the 70-micron particles,
Fig. 2.16: Radial profiles of particles number density taken at x = 15d
the concentration of particles at r/d>1 tend to be under-predicted. This error could be reduced if the small fraction of particles with initial radial velocity greater than 0.55 m/s were taken into account.

2.6 Conclusion

In this work, the Direct Quadrature Method of Moments was implemented in a massively parallel LES code to model finite Stokes number particles in turbulent flows. The derivation of the DQMOM equations was shown for a multivariate NDF using the particle velocity components as internal coordinates. Equations were simplified by extending the mono-kinetic assumption used in the multifluid approach to all the internal coordinates and for each environment and by neglecting the subgrid-scale terms in the NDF filtered transport equations. This approach was tested by comparing simulation predictions to experimental results in nonreacting coaxial particle-laden jets. Simulations were performed for three different velocity ratios and two particle size distributions. Results showed very good agreement with the experimental observations, both qualitatively and quantitatively. The LES coupled with DQMOM approach was able to accurately model a variety of different particle behaviors, such as particle jet waviness, spreading, break up and particle clustering, for both fine and coarse particle size distributions. The development of the mean axial velocity along the centerline for the gas phase and the solid phase was also analyzed. The experimental values were predicted by the simulations with a maximum error below 12% and single phase simulations suggest that a significant part of this error can be attributed to the gas phase modeling. Segregation effects were also captured as small particles dispersed faster than the large ones. Moreover, simulations showed the importance of correctly discretizing the particle size phase-space for small particles and the particle velocity phase-space for large particles in the quadrature approximation. Overall, this study provides a solid validation of the LES with DQMOM approach to model particles in turbulent flows and justifies its further application to coal flame simulations.
2.7 Acknowledgements

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CHAPTER 3

IGNITION OF COAXIAL TURBULENT
DIFFUSION OXY-COAL JET FLAMES:
EXPERIMENTS AND SIMULATIONS
COLLABORATION

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3.1 Abstract

The primary purpose of the study is to obtain physical insight into the stability of an oxy-coal flame as a function of changing primary gas composition (CO₂ and O₂). Flame stability was measured by Zhang et al. using optical measurements of the flame stand-off distance in a 40 kW pilot facility. Large Eddy Simulations (LES) of the facility were performed using a multiscale simulation tool and provide additional insight into the experimentally observed data. The importance of factors such as heterogeneous reactions, radiation or wall temperature can be better understood thanks to simulations. The effects of three parameters on the flame stand-off distance were studied. Simulation predictions were compared to experimental data using the data collaboration method. Overall, this study shows that high-fidelity LES simulations combined with experimental data can yield a deeper under-
standing of very complex coal flames, indicate where experimental uncertainties lie and be a valuable tool for design, retrofit and scale-up of oxy-coal burners.

3.2 Introduction

Coal is an important energy resource for electricity production as its reserves are far more abundant than those of other fossil fuels. As environmental regulations get stricter, coal thermal power plants need to control pollutants emissions such as $NO_x$, $SO_x$ and ash particles. Coal also plays an important role in global $CO_2$ emissions, which will need to be reduced to limit the effects of climate change [66]. Oxy-coal combustion, in which an $O_2/CO_2$ mixture replaces air, is one of the few possible capture technologies to enable $CO_2$ sequestration for existing coal-fired boilers. Despite the numerous research efforts that have been conducted on oxy-fuel combustion, several gaps of knowledge have been identified and need to be addressed in more detail to obtain a fundamental understanding of the changes between oxy-fuel combustion and conventional air-fired combustion [79,95]. One area of interest is understanding how the replacement of air by a $O_2/CO_2$ mixture affects the flame ignition and stability for retrofit and new design of oxy-coal units [15]. Flame stabilization is a key technology to control $NO_x$ emissions because it increases the size of the fuel-rich flame region where $NO_x$ is reduced to nitrogen [6]. Furthermore, flame stability of oxy-fuel combustion strongly depends on oxygen concentration of the gas mixture [21] and can also affect the heat transfer performance of the plant. Thus, flame stabilization plays a very important role in controlling both $NO_x$ and $CO_2$ emissions.

Ignition of turbulent coal flames is a complex phenomenon involving multiphysics aspects. The aerodynamics, kinetics and thermodynamics of both the gas and solid phase play an important role in understanding the flame characteristics. In the past, there have been numerous studies of single coal particle ignition mechanisms [7,22,25,37,49,50,67,69,89,96,110,111,114]. Both homogenous and heterogeneous mechanisms have been proposed to understand the ignition process. In the homogenous mechanism, the initial step is pyrolysis and subsequent ignition of the volatiles, followed by ignition of the char. The heterogeneous
reaction involves the direct attack of oxygen on carbon at the surface of the particle. The heterogeneous mechanism neglects the influence of volatile matter and depends on heat generation by combustion on the coal surface, whereas the homogeneous mechanism involves the combustion of evolved volatile matter in the vicinity of the particle [37]. The dominant ignition mechanism is subject to controversy as Smoot and Smith [85] believed that reaction of oxygen on the coal surface was not a major factor in the ignition process while others identified surface oxidation as very important [22,25]. In addition, numerous factors, such as coal properties, particularly particle size and volatile content [7,25,49,89], heating rate [25,89], oxygen concentration [22,67,68,89], pressure [69,89,110] and nature of the gas flow surrounding the particle [50,67,68] may all impact coal ignition.

Oxy-coal combustion introduces an additional variable to those present in air-fired configurations, namely the concentration of $O_2$ in the inlet oxidant mixture [15]. In practical turbulent diffusion flames, the overall inlet oxidant stream is split into at least two streams: a primary stream that transports the pulverized coal to and through the coal injector within the burner and a secondary stream that mixes with the primary coal jet downstream of the burner. In contrast to air combustion, the amount of oxygen within the primary jet in oxy-coal combustion is a variable, which can be set anywhere from zero up to any value that is considered safe. Previous work showed that flame stability is influenced by the $P_{O_2}$ in primary $O_2/N_2$ streams [15,39,51,68,96]. However, with the exception of the recent work of Molina and Shaddix [67,68], most studies have not considered the effect of elevated levels of $CO_2$, as present in oxy-coal combustion systems, on the ignition of coal particles. The different properties of $CO_2$ in comparison to $N_2$ have been shown to cause differences in flame and furnace operation parameters such as ignition time and gas temperature profiles [58]. Molina and Shaddix [68] have suggested that differences between air- and oxy-fired coal particle ignition times and char burnout times are due to differences in molecular diffusion rates of $O_2$ in $N_2$ and $CO_2$. Hu et al. [42] and Zeng et al. [110] have shown that the longer ignition delay of coal/char particles can be attributed to reduced $O_2$ diffusion to the particle surface.
Although the reaction kinetics involved in coal particle ignition have been investigated extensively, most of the studies were done on single coal particles and on laminar flows. The single coal particle ignition mechanism cannot, however, sufficiently elaborate on the real turbulent flames due to certain limitations. First, the particles injected always contain a size distribution which should be taken into account to obtain correct kinetics [7]. Second, interactions between particles play a crucial role in the ignition and combustion behavior and pollutant formation and destruction. Annamalai et al. [3,4] studied isolated coal/char particles and found that the particles with relatively low volatile matter, which ignite heterogeneously when isolated, ignite homogeneously under cloud conditions. Finally, in real field boilers, with the mixing effect introduced by the large eddies in turbulence, single coal particle ignition mechanism must be coupled with turbulent mixing to understand or predict the ignition and the flame stability. Turbulence can easily wipe off the gradients of concentrations, temperatures or velocities, which dramatically changes the ignition mechanism. The fundamental mechanics of coal particles in turbulent flows in general are as yet poorly understood [14,24,52] and have not been included in optimizing the design and operating flow condition for coal combustors. Smith et al. [84] showed that particle behaviors such as particle dispersion and particle clustering affected operational considerations within a coal-fired boiler such as flame stability, pollutants formation or heat fluxes.

The gap between the fundamental research and the real industrial applications is thus still large and to understand how turbulent coal flames are stabilized, one must consider numerous complex physical processes and their interactions. This issue can be addressed by using Computational Fluid Dynamics (CFD), which was shown to be a very useful and accurate modeling framework for the description of particulate systems, and for the description of turbulent flows in general [24,29]. Turbulence models based on the Reynolds-Averaged Navier–Stokes (RANS) equation are commonly used with gas and coal combustion models for the simulation. There is however evidence that RANS can be inaccurate in predicting particle-laden flows [5,86] and a turbulence model based on large eddy simulation (LES) has
been convincingly shown to be superior to RANS in accurately predicting turbulent mixing and combustion dynamics [56, 75].

This work is directed towards understanding the ignition mechanisms of coaxial turbulent diffusion, oxy-coal flames and how operating parameters such as the composition of the coal transport medium (primary $CO_2$ and $O_2$) or the furnace temperature affect the flame standoff distance. The focus is on interaction mechanisms between turbulent mixing in coaxial jets and coal particle ignition, rather than on each physical or chemical process taken individually. Flame stability was measured by Zhang et al. [113] using optical measurements of the flame stand-off distance in a 40 kW pilot facility. Experimental results suggest that flame stability is affected by primary $P_{O_2}$, secondary preheat temperature, secondary $P_{O_2}$ and the transport medium. Simulations of the facility are performed using a massively parallel LES simulation tool and provide additional insight into the experimental data. The importance of factors such as heterogeneous reactions, radiation, primary stream temperature or wall temperature is evaluated through the simulation results. A formal Validation and Uncertainty Quantification study is conducted with the Data Collaboration method [27, 33, 35]. The goal of this study is to show that LES can be a valuable tool for predicting and understanding oxy-coal flames characteristics and that combining experimental data with high-fidelity modeling can guide experimentalists in their measurements by indicating where uncertainties lie.

3.3 Experiments

This section briefly summarizes the experimental results published by Zhang et al. [112, 113] and the experimental conditions. A special attention is given to the uncertainties associated with the measurements.

The experiments were carried out in the University of Utah 40 kW (100 kW maximum rating) down-fired oxy-coal combustor. The furnace consists of an oxy-fuel combustion chamber, followed by downstream-controlled temperature cooling to simulate practical furnace conditions [15]. It allows the systematic control of burner momentum and velocity
variables as well as wall temperatures. Quartz/sapphire windows allow nonintrusive optical
diagnostics to measure the flame stability, quantified by the stand-off distance. The burner
applied was a coaxial Type 0 (zero swirl) burner with a transport stream and pulverized
coal in the center and a secondary stream in the annular sleeve.

The coal employed was Utah bituminous pulverized coal. The mass average size of the
particles is approximately 60 $\mu$m (the coal particle size distribution is given in Table 3.1).
The coal analysis is as follows:

1. Ultimate (wt%, daf): C 77.75%, H 5.03%, N 1.44%, S 0.45%, O 15.33%.
2. Proximate: moisture 3.03%, volatile matter 38.81%, fixed carbon 46.44%, ash
11.72%.
3. Higher (gross) heating value (HHV) = 27,286 kJ/kg.

The flame stability was quantified by the observed stand-off distance, the distance
between the burner tip and the visible ignition of the flame. An EPIX CMOS camera was
set up at an exposure time of 8.3 ms and a frame rate of 30 fps to take sequences of flame
images. Flame envelope/structure was defined at the above camera settings, which are
consistent with the averaging process of the human eye. An image processing methodology
was developed to quantify the flame stand-off distance from image data.

The following operating parameters were fixed: coal feeding rate 4.84 kg/h, stoichio-
metric ratio 1.15, wall temperature 1283 K, primary stream velocity 5.4 m/s and primary
stream temperature 305 K. Both the primary and secondary streams were $O_2/CO_2$ mixtures
and the overall oxygen concentration was maintained to 40 vol%. In this study, we are inter-

<table>
<thead>
<tr>
<th>Particle size ($\mu$m)</th>
<th>Volume %</th>
</tr>
</thead>
<tbody>
<tr>
<td>178</td>
<td>1.2</td>
</tr>
<tr>
<td>152</td>
<td>2.5</td>
</tr>
<tr>
<td>106</td>
<td>13.8</td>
</tr>
<tr>
<td>75</td>
<td>12.7</td>
</tr>
<tr>
<td>63</td>
<td>58.6</td>
</tr>
<tr>
<td>44</td>
<td>5.3</td>
</tr>
<tr>
<td>37</td>
<td>5.9</td>
</tr>
</tbody>
</table>

Table 3.1: Utah bituminous coal particle size distribution
ested in a specific set of experiments where the secondary stream preheat temperature was set to 544 K and the primary $P_{O_2}$ varied from 0% to 20.9%. Variations in primary $P_{O_2}$ were obtained by transferring oxygen flows from the primary to the secondary stream, with only small effects on jet momentum. The averaged flame stand-off distance was measured with the CMOS camera for different values of primary $P_{O_2}$. Each experiment was replicated five times.

Results are presented in Table 3.2 and show that the flame stand-off distance decreases with increasing primary $P_{O_2}$. Figure 3.1 shows the averaged values over the five replicates of the stand-off distance with associated error bars, where the upper and lower bounds correspond respectively to the maximum and minimum average value over the five replicates. This set of data was chosen as the main focus of this study because it is unclear how the primary $P_{O_2}$ affects the flame stand-off distance. High-fidelity computer simulations may provide insights on the ignition mechanisms and help design further experiments. This set of data will then later be used for the validation and uncertainty quantification analysis with the ARCHES LES tool provided in Section 3.6.2.

Sources of uncertainty in the operating conditions were identified during those experiments. First, the wall temperature was measured with a thermocouple located behind the wall heater. The reported wall temperature value of 1283 K is therefore the value measured behind a 2 inches thick layer and not the wall temperature inside the furnace. The later one could indeed be much higher. An infrared camera was used to determine the wall temperature inside the University of Utah furnace in other experiments with similar operating conditions and measured wall temperatures at about 1800 K. In addition, thermocouple

Table 3.2: Average standoff distances (cm) for overall $P_{O_2}$=40%, secondary preheat 544 K

<table>
<thead>
<tr>
<th>$P_{O_2, pri}$ (vol%)</th>
<th>0</th>
<th>5.4</th>
<th>9.9</th>
<th>15.4</th>
<th>20.9</th>
</tr>
</thead>
<tbody>
<tr>
<td>replicate 1</td>
<td>29.7</td>
<td>29.0</td>
<td>28.8</td>
<td>28.5</td>
<td>28.2</td>
</tr>
<tr>
<td>replicate 2</td>
<td>30.3</td>
<td>29.9</td>
<td>21.2</td>
<td>7.6</td>
<td>7.3</td>
</tr>
<tr>
<td>replicate 3</td>
<td>32.0</td>
<td>32.2</td>
<td>30.1</td>
<td>9.6</td>
<td>7.7</td>
</tr>
<tr>
<td>replicate 4</td>
<td>28.0</td>
<td>27.6</td>
<td>8.7</td>
<td>7.2</td>
<td>7.6</td>
</tr>
<tr>
<td>replicate 5</td>
<td>28.9</td>
<td>16.1</td>
<td>8.9</td>
<td>8.6</td>
<td>8.5</td>
</tr>
<tr>
<td>average</td>
<td>29.8</td>
<td>26.8</td>
<td>19.5</td>
<td>12.3</td>
<td>11.9</td>
</tr>
</tbody>
</table>
measurements can be inaccurate and have up to 50K uncertainty. The primary stream was injected at room temperature and its temperature was assumed to be 305 K but was not measured directly. Operators noticed that the tip of the burner was glowing red after the experiments. The burner penetrates inside the furnace by about 10 cm and its tip is therefore exposed to a high radiative flux. As a result, it is possible that the burner transfers heat to the primary stream and the coal particles, which would result in a higher primary stream temperature.

3.4 Numerical modeling

Simulations were performed using ARCHES, a LES tool resulting from a ten-year partnership with the Department of Energy and the University of Utah. It is a massively parallel code that solves conserved quantities (mass, momentum, energy, scalar) spatially and temporally in a turbulent flow field, allowing for detailed and accurate simulations of fires and flames [87]. ARCHES includes particle phase representation through the use of a moment
method, the direct quadrature method of moments (DQMOM). DQMOM combined with LES allows ARCHES to perform highly accurate simulations of coal combustion and gasification applications. The code is integrated into a C++ framework called Uintah which provides large-scale parallelization tools for physics components [9,16,73]. ARCHES is maintained in a repository and distribution is freely available [41]. The governing equations and numerical methods for the gas phase and the dispersed phase are described in the following sections.

3.4.1 Governing equations of the gas phase

The gas phase is solved in a Eulerian manner by a finite volume method. Favre-filtered governing equations for continuity, momentum, enthalpy and scalars transport are solved. A second-order central difference scheme is used to calculate the convection and diffusion terms of the transport equations.

For the continuity equation, the mass source term due to coal combustion is taken into account. Coal combustion models are described in Section 3.4.2.

For the momentum equation, the Navier–Stokes equations are solved by taking the momentum source due to coal combustion into account. The subgrid scale (SGS) stress tensor is calculated by the dynamic subgrid scale model [36].

For the enthalpy equation, radiation, heat exchange between gas and coal particles, and heat transfer due to coal combustion are taken into account. The radiation source term is calculated by the discrete ordinate radiation method [2,46,47].

The gas-phase combustion is modeled through a mixture fraction approach and chemical equilibrium for the gas phase properties is assumed [93]. Spinti et al. [87] gave a more detailed description of the gas phase modeling in ARCHES.

3.4.2 Governing equations of the dispersed phase

In order to describe the evolution of the disperse particulate phase, LES equations must be coupled with a population balance equation (PBE). The method of moments (MOM)
offers an attractive alternative to the traditional Lagrangian method where the PBE is tracked through its moments by integrating the internal coordinate [43]. The Direct Quadrature Method of Moments allows the description of population of particles with multivariate PBE, where two or more properties of the population are simultaneously tracked [26,59]. DQMOM has been shown to be a powerful approach for describing polydisperse solids undergoing segregation, growth, aggregation and breakage processes in the context of CFD simulations [26,32,60] and to greatly reduce the computational cost compared to the Lagrangian approach [17,18].

In previous work, the advantages of using DQMOM for treating particle populations with low Stokes numbers (i.e., the dispersed-phase velocity follows closely the velocity of the continuous phase) have been clearly demonstrated [61,63,98,115]. However, coal particles, which typically range from 1 micrometer to 200 micrometers in pulverized coal flames, have a finite Stokes number and their velocity depends on the particle size. In quadrature methods, this dependency is accounted for by solving an Eulerian model where each quadrature node has its own velocity field. Fox et al. [32] have applied DQMOM to droplets spray with finite Stokes number and compared the results with the multifluid method and a classical Lagrangian solver. Results show that DQMOM can provide a comparable accuracy with a significantly lower computational cost, making it a very good candidate for more complex two-phase combustion applications.

The DQMOM method was implemented into ARCHES to solve the dispersed phase in an Eulerian manner. ARCHES coupled with DQMOM has been proven to be a reliable and accurate method to simulate particles in turbulent flows [74]. A brief description of the particles representation and of the DQMOM equations is given in the following paragraphs. A more detailed description of DQMOM implementation in ARCHES was previously given by Pedel et al. [74].
3.4.2.1 Number density transport equation

The particle Number Density Function (NDF) $n$ describes the number of particles per volume as a function of the spatial location and of other independent variables. These independent variables are intrinsic characteristics of the particles (e.g., particle diameter, particle velocities, etc.) and are called internal coordinates. The internal coordinates are contained in the internal coordinate vector $\xi$, and are dependent on space and time. At a given location in space and time $(x_0, t_0)$, the total number of particles in a volume $V$ is given by:

$$n_p(x_0, t_0) = \int \int n(\xi; x_0, t_0) d\xi dV \quad (3.1)$$

The velocity of the NDF in both real and internal coordinate space can be described using a set of internal coordinates $\xi = [\xi_1, \xi_2, ...]$, and external coordinates $x = [x_1, x_2, x_3]$. For a single particle, the time derivatives of these variables may be written as

$$\frac{dx_i}{dt} = v_{p,i} \quad (3.2)$$

$$\frac{d\xi_j}{dt} = G_j \quad (3.3)$$

where $v_{p,i}$ is the $i^{th}$ component of the particle velocity vector $v_p$.

The NDF transport equation is given by:

$$\frac{\partial}{\partial t} (n(\xi; x, t)) + \frac{\partial}{\partial x_i} (\langle v_{p,i} | \xi \rangle n(\xi; x, t)) = -\frac{\partial}{\partial \xi_j} (\langle G_j | \xi \rangle n(\xi; x, t)) + h(\xi; x, t) \quad (3.4)$$

In this equation, the variable $\langle v_{p,i} | \xi \rangle$ represents the velocity in real space of the NDF. This quantity is an average over particles having the same coordinates $(\xi; x, t)$ and is a distribution conditional to the value of the vector $\xi$. Similarly, the quantity $\langle G_j | \xi \rangle$ describes
the velocity of the NDF in the phase space. The term $h$ is a source term representing the birth and death of particles in the domain. This term is set to zero as particle breakage and aggregation effects are neglected.

### 3.4.2.2 Direct Quadrature Method of Moments

The Direct Quadrature Method of Moments (DQMOM) approximates the NDF by a summation of multidimensional Dirac delta functions. Figure 3.2 gives an example for bivariate Gaussian distribution (i.e., two internal coordinates) of the NDF and its quadrature approximation with three quadrature nodes. In this approximation, the following assumptions are made:

- For a given quadrature node $\alpha$, at a given point $(x,t)$, there is only one characteristic averaged internal coordinate value $\langle \xi \rangle_\alpha$, also called abscissa,

- The internal coordinate dispersion around the averaged value $\langle \xi \rangle_\alpha$ is zero.

With those assumptions, the NDF is written as:

$$
n(\xi; x, t) = \sum_{\alpha=1}^{N} \left( w_\alpha \prod_{i=1}^{N_\xi} \delta \left( \xi_i - \langle \xi_i \rangle_\alpha \right) \right)
$$

where both the weight $w_\alpha$, which defines the number of particles per volume associated with the quadrature node $\alpha$, and the average internal coordinate value $\langle \xi \rangle_\alpha$ depend on space and time, but the dependence is omitted for clarity of notation. $N$ is the number of quadrature nodes and $N_\xi$ the number of internal coordinates.

The validity of those assumptions has been discussed for particle velocity in the multifluid method [17]; the hypothesis is valid for particles with low Stokes number and becomes inappropriate when Stokes number increases ($St > 0.4$) and particle trajectory crossing occurs. It is however important to note that this limitation can be partially overcome with DQMOM. In the multifluid model, only the particle size space is discretized, whereas DQMOM allows us to discretize a multidimensional space. It is therefore possible to discretize the particle velocity space if the particle velocity is chosen as an internal coordinate and then
Fig. 3.2: DQMOM approximation of the NDF with 3 quadrature nodes

to have several quadrature nodes with different velocities and otherwise identical internal coordinates values. As quadrature nodes can cross each other, the error associated with finite Stokes number particles crossing each others is reduced.

DQMOM solves transport equations for the weights and weighted abscissas of the quadrature approximation. These transport equations are simply scalar transport equations, and are easily implemented into a CFD code:

\[
\frac{\partial}{\partial t} (w_\alpha) + \frac{\partial}{\partial x_i} (\langle v_{p,i} \rangle_\alpha w_\alpha) = a_\alpha 
\]  (3.6)

\[
\frac{\partial}{\partial t} (s_{n\alpha}) + \frac{\partial}{\partial x_i} (\langle v_{p,i} \rangle_\alpha s_{n\alpha}) = b_{n\alpha} 
\]  (3.7)
where \( s_{n\alpha} = w_{\alpha} \langle \xi_n \rangle_\alpha \) is the weighted abscissa for the \( n^{th} \) internal coordinate and \( \langle v_{p,i} \rangle_\alpha \) is the average internal coordinate velocity. The right-hand side terms \( a_\alpha \) and \( b_{n,\alpha} \) are obtained by solving a linear system of equations.

### 3.4.2.3 Single coal particle description

Following Smoot and Smith [85], a single coal particle can be characterized using several particle independent variables. These are denoted:

- Raw coal \( \alpha_{c_j} \)
- Char \( \alpha_{h_j} \)
- Moisture \( \alpha_{w_j} \)
- Ash \( \alpha_{a_j} \)
- Particle diameter \( d_{p_j} \)
- Particle enthalpy \( h_{p_j} \)
- Particle velocity vector \( v_{p_j} \)

The above quantities use the subscript \( j \) to denote the \( j^{th} \) particle. The variable \( r \) can be used to denote reaction rates, so that \( r_{h_j} \) would be the net char reaction rate for the \( j^{th} \) particle. Using this nomenclature, physical processes important to coal particles are depicted in Figure 3.3. The raw coal can react to form gaseous volatile matter in devolatilization reactions and solid char; the solid char can be oxidized to form more gaseous products. The ash mass is fixed, and ash is treated as inert.

In this study, a total of seven internal coordinates were chosen to represent particle characteristics in the DQMOM formulation:

- mass of raw coal \( \alpha_{c} \)
- mass of char \( \alpha_{h} \)
- particle diameter \( d_{p} \)
- particle enthalpy \( h_{p} \)
- particle velocity components \( v_x, v_y, v_z \)
The particle diameter $d_p$ was assumed to be constant. To reduce the number of internal coordinates and the computational cost, the moisture in the particles was assumed to be evaporated before entering the domain and the water content was added to the inlet stream.

### 3.4.2.4 Particle velocity model

In this work, the aggregation/breakage phenomena are neglected and the only physical process modeled is the drag force, which can be described by the Stokes drag law. For a mesoscale size particle, other forces such as lift can be neglected and the momentum equation for the particle can be expressed by an ordinary differential equation.

\[ \frac{dv_{p,i}}{dt} = \frac{f_{\text{drag}}}{\tau_p} (u_{g,i} - v_{p,i}) + \frac{g_i (\rho_p - \rho_g)}{\rho_p} \]  \tag{3.8}

where \( g \) is the gravity force acting on the particle, \( \rho_g \) and \( \rho_p \) are respectively the gas and particle densities, \( u_g \) is the gas velocity, and \( f_{\text{drag}} \) is the coefficient of the drag force, which has a close relationship with the particle Reynolds number [85]:

Fig. 3.3: Illustrative schematic of coal particle components and reactions.
\[
\begin{align*}
\tau_p &= \frac{\rho_p d_p^2}{18 \mu_g} \\
Re_p &= \frac{\rho_p d_p |u_p - u_g|}{\mu_g} \\
\end{align*}
\]  

where \(\mu_g\) is the gas dynamic viscosity.

In equation (3.8), \(\tau_p\) is the particle relaxation time:

\[
\begin{align*}
\tau_p &= \frac{\rho_p d_p^2}{18 \mu_g} \\
Re_p &= \frac{\rho_p d_p |u_p - u_g|}{\mu_g} \\
\end{align*}
\]  

3.4.2.5 Devolatilization model

In this study, an empirical devolatilization model proposed by Yamamoto et al. is used [104]. The approach was proven accurate and inexpensive by comparison to more sophisticated models such as the distributed activation energy model, FLASHCHAIN [71] or chemical percolation devolatilization (CPD) [31]. The devolatilization process is described as:

\[
\begin{align*}
r_v \\
(\text{raw coal}) &\rightarrow Y (\text{volatiles}) + (1-Y) (\text{char})
\end{align*}
\]  

where \(Y\) is a stoichiometric coefficient. The devolatilization rate of the model \(r_c\) is based on a single reaction:

\[
r_v = \frac{d\alpha_c}{dt} = -Fk_0 exp \left(-\frac{E_v}{RT_p}\right) \alpha_c
\]  

where \( k_0 \) is the frequency factor, \( E_v \) is the activation energy and \( R \) is the gas constant. \( F \) is the modification factor and is expressed as a function of fraction devolatilized \( G_v = \frac{(\alpha_{c,0} - \alpha_c)}{\alpha_{c,0}} \). Here \( \alpha_{c,0} \) is the initial mass of raw coal.

The model parameters \( Y \), \( k_0 \), \( E_v \) and \( c_i \) are determined by minimizing the difference between CPD and the present model at the heating rate of 100,000 K/s. Figure 3.4 shows results calculated by the Yamamoto model and CPD. The coal used here is the same as the coal used for the experiments. The results obtained by the present model show good agreement with CPD results. The maximum error of the fraction devolatilized is below 1.5% for both heating rates of 100,000 and 20,000 K/s. This model thus offers the advantage to predict the fraction devolatilized with the same accuracy as the CPD model but with the low computational cost of a single-rate reaction model.

### 3.4.2.6 Char oxidation model

After the raw coal in a particle has devolatilized, it forms char. Historically, most kinetic data on burning coal char particles have been interpreted using an \( n^{th} \) order Arrhenius model of char combustion. In this representation, the global surface reaction rate follows the expression

\[
q = k_s (T_p) P_{O_2,s}^n
\]  

(3.15)

where \( n \) is the reaction order, \( T_p \) is the particle temperature, and \( k_s \) is a temperature-dependent rate coefficient, assumed to follow an Arrhenius form:

\[
k_s (T_p) = A_s exp\left(-\frac{E_s}{RT_p}\right)
\]  

(3.16)
Fig. 3.4: Verification results of the Yamamoto devolatilization model with comparisons to results by the chemical percolation devolatilization model (CPD)

The parameter values were chosen from the experimental study of Murphy and Shaddix [70] for Eastern bituminous coal: $A_s = 344 \text{ mol/s/m}^2/\text{atm}^n$, $E_s = 45.5 \text{ kJ/mol}$ and $n = 0.18$. To be consistent with the experiments, the same equations used to determine these parameter values [70] are solved to calculate the surface reaction rate $q$. The partial pressure of oxygen at the char surface $P_{O_2,s}$ is obtained by solving the gas-phase diffusion,

$$\frac{P_{O_2,s}}{P} = \gamma + \left(\frac{P_{O_2,\infty}}{P} - \gamma\right) \exp\left(-\frac{qd\eta}{2CD_{O_2,\text{mix}}}\right)$$

(3.17)

In this equation, $C = P/RT_f$ is the average molar concentration of the gases in the boundary layer and $\gamma = \frac{\psi - 1}{2}$ where $\psi = CO_2/CO$ is the fraction of carbon that becomes $CO_2$. The $CO_2/CO$ production ratio at the particle surface is determined from the correlation $CO_2/CO = A_0 P_{O_2,s}^{\eta_0} \exp(-B_0/T_p)$ with the coefficients suggested by Tognotti et al. [91]. Given a trial value of $q$, Equation 3.17 must be solved iteratively because $\gamma$ depends on the $CO_2/CO$ ratio, which in turn depends on $P_{O_2,s}$. 
The char oxidation rate is then given by

\[ r_h = \frac{d\alpha_h}{dt} = -\pi d^2 W_C q \]  

(3.18)

where \( W_C \) is the molecular weight of carbon.

To account for the eventuality of heterogeneous ignition, the oxidation reaction was also applied to the raw coal before ignition. As shown in Sections 3.6.1 and 3.6.2, the oxidation rate was negligible before ignition, even though the activation energy value \( E_s \) is significantly lower than oxidation activation energy values reported in previous studies at low temperature. There is indeed a relative agreement about the magnitude of the global intrinsic activation energy for carbon oxidation at a low temperature regime where direct measurement is possible; it falls between 105 and 180 kJ/mol with many values in the range 130 to 150 kJ/mol [30, 44, 83].

3.4.2.7 Particle heat transfer

The particle is heated by convection, radiation and reaction enthalpy changes:

\[ \frac{dh_p}{dt} = Q_{rad} + Q_{conv} + r_j h_j, \]  

(3.19)

where \( Q_{rad} \) represents the net radiation flux to a particle, \( Q_{conv} \) represents energy transfer due to convection and conduction between the gas and the particle, and \( r_j h_j \) represents both the amount of energy lost by the particles due to lost mass and the enthalpy released during devolatilization and oxidation reactions.

Convection

The convection term is expressed as:

\[ Q_{conv} = N u \pi k_g (T_g - T_p) d_p \frac{\kappa/2}{e^{\kappa/2} - 1} \]  

(3.20)
where \( T_g \) is the gas temperature, \( k_g \) is the gas thermal conductivity, and \( \kappa = -W_C q d_p C_{pg} / k_g \) is a modified version of the Peclet number [70]. \( C_{pg} \) is the gas heat capacity. The Nusselt number \( Nu \) is calculated with the following correlation [55]:

\[
Nu = 2.0 + 0.65 Re_p^{1/2} Pr^{1/3}.
\] (3.21)

The particle temperature \( T_p \) is obtained from the particle enthalpy by assuming Merrick’s correlations for the particle heat capacity [65].

Radiation

The radiative flux is given by:

\[
Q_{rad} = Q_{incident} - Q_{emitted}
\] (3.22)

where \( Q_{incident} \) is the incident radiative flux to the particle and \( Q_{emitted} \) is the radiative heat flux emitted by the particle. The absorption coefficient of the coal particles is assumed constant: \( Q_{abs} = 0.8 \). The incident flux is obtained through the Discrete Ordinates radiation model whereas the emitted flux is calculated using the blackbody emissive power:

\[
Q_{emitted} = Q_{abs} \pi d_p^2 \sigma T_p^4.
\] (3.23)

Following [85], the coupling with the gas phase is done by modifying the gas absorption coefficient:

\[
k_{abs,g} = k_{abs,g} + k_{abs,p}
\] (3.24)

with

\[
k_{abs,p} = \sum_{j=1}^{n_p} \frac{\pi}{4} Q_{abs} d_{pj}^2
\] (3.25)
3.5 Validation and uncertainty quantification

The common approach to model validation consists in comparing the predictions made by the model to available experimental data. Typically, such comparisons result in mixed outcomes: some show a good agreement and some do not. In the latter case, the apparent inconsistency between the model and the experiment is argued to imply either that the model is inadequate or that the experiment is incorrect. Before a decision can be made as to whether or not to use a model, it is essential for the decision maker to understand the uncertainty in the predictions made by the model. Simulation error or uncertainty arises from four types of uncertainties:

- Model uncertainty, or error in the mathematical description of the physical process being modeled. This error includes uncertainty in the parameters used in these models.
- Uncertainty in boundary conditions. This uncertainty derives from the uncertainty in the inputs and boundary conditions used for the simulation.
- Numerical error (verification error). This error arises from the numerical approximations used in producing the simulation (i.e., grid resolution, discretization error, convergence error, etc.)
- Experimental error in validation data. The quantification of simulation error is achieved by comparing the simulation output to experimental observations; since the experimental measurements themselves are inexact, the simulation accuracy can not be known to any better degree of accuracy than the measurements themselves.

Because of these inaccuracies, it is essential to simultaneously estimate the sensitivity of the assessed variability in the model output to changes in the assumptions concerning the model uncertainties, as well as to changes in the experimental measurements.

This approach is illustrated in Figure 3.5. It tightly couples simulation and experimental data with formal verification and validation / uncertainty-quantification (V&V/UQ) methodologies. More specifically, given a bounded parameter space, this methodology seeks to find the minimum and maximum error by propagating the error/uncertainty in the input parameter space (priors) to the simulation output or posterior subject to the constraints.
of the experimental validation and verification uncertainty. This methodology provides the lower and upper bound on the predicted efficiency by requiring consistency between the simulation output and the measured experimental data.

This work presents a formal validation and uncertainty quantification analysis of the ARCHES LES tool with the experimental data on oxy-coal flame ignition presented in Section 3.3. The Data Collaboration approach [27, 34, 35] is used to study the uncertainty associated with a couple of scenario parameters which were identified as potential sources of uncertainty. The following section first provides a brief description of the Data Collaboration method and then explains the methodology chosen to perform the analysis. Finally, a detailed description of the simulations is provided.

### 3.5.1 Data Collaboration method

Recently, the Data Collaboration approach has been developed to provide quantitative and reliable uncertainty bounds on predictions from multiphysics, multiscale simulations [27, 34, 35]. Data Collaboration is a method that unites process models and associated admissible parameter values with experimental data and accompanying uncertainties. The
approach focuses on transferring the uncertainties of the experimental data into the model directly. Data obtained from various sources are integrated through models that depend on common parameters. This ensemble is called a dataset. Optimization-driven techniques can then be applied to dataset analysis, including performing dataset-based predictions [34] and determining dataset consistency [28]. Doing so allows one to harvest substantially more of the information content of the data [35] and determine more realistic bounds on model predictions [34].

### 3.5.1.1 Dataset

In the Data Collaboration approach [28], each experiment $e$ is associated with a dataset unit $U_e = (d_e, u_e, l_e, M_e)$ where $d_e$ is the measured value, $u_e$ and $l_e$ are respectively the upper bound and lower bound of the reported uncertainty and $M_e$ is a mathematical model of the experiment. The model depends only on a parameter vector to generate the prediction $M_e$ for $y_e$. The true value of the experimental observable, $y_e$, satisfies $l_e \leq y_e - d_e \leq u_e$. A dataset is a collection of dataset units. The dataset is denoted as $D$ and the set of indices $e$ as $E$, i.e., $D = \{U_e\}_{e \in E}$.

Complex models usually have a large number of parameters. In the Data Collaboration method, a subset of model parameters which have an influence on the outcome is chosen. The parameters contained within this subset are called active variables. An individual active variable is designated as $X_j$ and $x_j \in \mathbb{R}$ refers to a specific value of $X_j$. Individual dataset units may have different sets of active variables. Thus, $X_j$ might be an active variable for one experiment but not another. We denote the list of active variables for experiment $e$ as $X_e$. The total number of active variables is denoted as $n$ and the vector $x \in \mathbb{R}^n$ represents the active variable values. Associated with a vector $x$, $x_e$ are the values extracted from $x$ that correspond to the active variables for experiment $e$. 
3.5.1.2 Initial hypercube and feasible region

The Data Collaboration approach assumes that prior information on the possible values of the dataset active variables is available. This prior information can be expressed as the confinement of possible values of the active variables to an n-dimensional “hypercube,” $H = \{ x \in \mathbb{R}^n : \alpha_j \leq x_j \leq \beta_j \}$, where $\alpha_j$ and $\beta_j$ are the lower and upper bounds on $x_j$ for $j = 1, 2, ..., n$. Each edge of the hypercube $H$ represents the presumed interval of physically allowed values of the corresponding active variable.

Some parameter values drawn from $H$ may result in model predictions that lie outside the experimentally determined ranges. In other words, not every $x \in H$ predicts all experimental observations of the dataset within their specified uncertainties. The collection of parameter values that are both contained in the hypercube and satisfy $l_e \leq M_e(x_e) - d_e \leq u_e$ for every $e \in E$ form the feasible region, $F_D$. A point $x$ that is not contained in $F_D$ can thus be eliminated from consideration as a possible value for the dataset active variables. An experiment may therefore eliminate portions of the hypercube $H$ from consideration, thereby decreasing the uncertainty in the values of the parameters.

3.5.1.3 Dataset consistency

The Data Collaboration approach also allows us to determine whether the data contained in the dataset are mutually consistent. A dataset $D$ is said to be inconsistent if there is no single point $x$ in the hypercube $H$ that satisfies $l_e \leq M_e(x_e) - d_e \leq u_e$ for all $e \in E$. Otherwise, the dataset is consistent. In other words, the dataset is inconsistent if the feasible region is empty. The consistency of a dataset depends on the size of the hypercube. A consistent dataset may become inconsistent with a decrease in the intervals of active variables and/or in the level of experimental error.

3.5.2 Methodology

In this study, we are interested in understanding the ignition phenomena in oxy-flames and the influence of operating conditions on the flame stability. The experimental data with
their associated uncertainties presented in Section 3.3 were selected for the validation study. Experiments suggest that the primary $P_{O_2,pri}$ has an influence on the flame stand-off distance and was therefore chosen as a parameter of interest. Two other parameters were chosen: the wall temperature $T_{wall}$ and the temperature of the primary stream $T_{pri}$. Those parameters were selected because experiments show a high sensitivity to the wall temperature and the stream temperature [112,113], but also because of their potential uncertainty. As described in Section 3.3, the wall temperature was measured at 1283 K but could be as high as 1800 K. The uncertainty range for $T_{wall}$ was thus assumed to be 1200 K - 1800 K. Similarly, the primary stream temperature could be higher than reported in the experiments since the burner tip is heated by radiation. We assumed that $T_{pri}$ could be as high as the secondary stream temperature (544K). The range associated with each parameter is reported in Table 3.3.

### 3.5.3 Surrogate model

The Data Collaboration method uses optimization procedures to determine the effects of input variables and associated uncertainties of system responses. This procedure requires a large number of function evaluations. Ideally, the function evaluations would be performed with ARCHES LES tool. However, LES simulations are computationally expensive and running hundreds of LES simulations is not feasible.

In this situation, it is useful to have a simpler model, also called a “surrogate model,” that is much cheaper to evaluate and much simpler in form, such as a polynomial. These can be constructed based on limited information about the larger scale computer simulation model. Box and Wilson first proposed the response surface methodology (RSM) [13]. RSM uses polynomial surfaces to represent the response as a function of input variables. Polyno-

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Range</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wall temperature ($T_{wall}$)</td>
<td>1200-1800 K</td>
</tr>
<tr>
<td>Primary stream temperature ($T_{pri}$)</td>
<td>305-544 K</td>
</tr>
<tr>
<td>Primary oxygen partial pressure ($P_{O_2,pri}$)</td>
<td>0-20.9 %</td>
</tr>
</tbody>
</table>
mials are very general and work well for many responses, and as a result, RSM has thrived as a widely used modeling technique for complex systems.

In this work, a Box-Behnken design [12] was chosen to obtain a surrogate model with three parameters: $P_{O_2, pri}$, $T_{wall}$ and $T_{pri}$. The design provides information required to build a second-order (quadratic) model. The Box-Behnken design places sample points on a parameter hypercube. One sample point is placed at the center of the hypercube and each other point is placed on the middle of each edge (Figure 3.6). In our case, for a design with three variables, the Box-Behnken design requires 13 points.

Following the Box-Behnken design, 13 LES simulations of the oxy-coal reactor are performed with the ARCHES model in order to construct a response surface surrogate model for use in the Data Collaboration validation method. The values of the three parameters $P_{O_2, pri}$, $T_{wall}$ and $T_{pri}$ for each simulation are described in Table 3.4.

### 3.5.4 Simulation details

The computational domain is a cylinder of diameter 0.6 m and length 0.9 m. The domain is decomposed into a cartesian mesh with a cellsize $\triangle = 1.7 \text{ mm}$. The primary and secondary inlets are located at the center of the $X=0$ face and outlet boundary conditions are applied at the $X=0.9$ m face. All other boundaries are set as walls with a fixed and uniform temperature $T_{wall}$.

The operating conditions are summed up in Table 3.5. According to the simulation matrix (Table 3.4), the oxygen concentration in the primary stream and secondary stream, the wall temperature and the primary stream temperature change with each simulation while the overall partial pressure of oxygen is maintained to 40%. Each of the 13 simulations was run for 24 hours on 1,500 CPUs with timesteps of about $9 \mu s$, which resulted in a total simulation time of about 1.25 s. Time-averaging was done on the last second of data.
Fig. 3.6: A Box-Behnken design for three parameters

Table 3.4: Parameter values for each simulation in the Box-Behnken design

<table>
<thead>
<tr>
<th>Simulation</th>
<th>$T_{\text{wall}}$(K)</th>
<th>$T_{\text{pri}}$(K)</th>
<th>$P_{O_2,pri}$ (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>1500</td>
<td>424.5</td>
<td>10.45</td>
</tr>
<tr>
<td>2</td>
<td>1500</td>
<td>305</td>
<td>0</td>
</tr>
<tr>
<td>3</td>
<td>1500</td>
<td>305</td>
<td>20.9</td>
</tr>
<tr>
<td>4</td>
<td>1500</td>
<td>544</td>
<td>0</td>
</tr>
<tr>
<td>5</td>
<td>1500</td>
<td>544</td>
<td>20.9</td>
</tr>
<tr>
<td>6</td>
<td>1200</td>
<td>424.5</td>
<td>0</td>
</tr>
<tr>
<td>7</td>
<td>1200</td>
<td>424.5</td>
<td>20.9</td>
</tr>
<tr>
<td>8</td>
<td>1800</td>
<td>424.5</td>
<td>0</td>
</tr>
<tr>
<td>9</td>
<td>1800</td>
<td>424.5</td>
<td>20.9</td>
</tr>
<tr>
<td>10</td>
<td>1200</td>
<td>305</td>
<td>10.45</td>
</tr>
<tr>
<td>11</td>
<td>1200</td>
<td>544</td>
<td>10.45</td>
</tr>
<tr>
<td>12</td>
<td>1800</td>
<td>305</td>
<td>10.45</td>
</tr>
<tr>
<td>13</td>
<td>1800</td>
<td>544</td>
<td>10.45</td>
</tr>
</tbody>
</table>
Table 3.5: Operating conditions

<table>
<thead>
<tr>
<th></th>
<th>Primary stream</th>
<th>Secondary stream</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mass flow rate (kg/s)</td>
<td>1.83E-3</td>
<td>7.60E-3</td>
</tr>
<tr>
<td>Temperature (K)</td>
<td>305-544</td>
<td>544</td>
</tr>
<tr>
<td>Mass fraction</td>
<td></td>
<td></td>
</tr>
<tr>
<td>(O_2)</td>
<td>0-0.16</td>
<td>0.37-0.41</td>
</tr>
<tr>
<td>(CO_2)</td>
<td>0.82-0.98</td>
<td>0.59-0.63</td>
</tr>
<tr>
<td>(H_2O)</td>
<td>0.02</td>
<td>0</td>
</tr>
</tbody>
</table>

The coal particles are injected with the primary stream at a mass flow rate of 4.84 kg/h and are assumed to have the same temperature and velocity as the gas phase at the inlet. The initial coal particle density is 1300 kg/m³. Particles are modeled using the DQMOM method with seven internal coordinates, as described in Section 3.4.2, and three quadrature nodes, each quadrature node corresponding to a different particle size. Table 3.6 shows the particle diameter associated with each quadrature node as well as their weight at the inlet. The weights for each particle size were determined from the particle size distribution given in Table 3.1.

3.6 Results and discussion

This section provides an analysis of the flame ignition mechanisms observed in the simulations as well as a validation and uncertainty quantification analysis of the LES model with respect to the chosen parameters.

Table 3.6: Quadrature approximation for 25-micron particles (inlet values)

<table>
<thead>
<tr>
<th>Quadrature node</th>
<th>Particle diameter ((\mu m))</th>
<th>Weight (# particles/m³)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>40</td>
<td>2.51E9</td>
</tr>
<tr>
<td>1</td>
<td>63</td>
<td>3.36E9</td>
</tr>
<tr>
<td>2</td>
<td>100</td>
<td>4.33E8</td>
</tr>
</tbody>
</table>
3.6.1 Ignition mechanisms

Gas temperature is assumed to be a good indicator for flame ignition. This assumption is discussed in Section 3.6.2. Results show that the wall temperature has a strong influence on the flame stand-off distance, whereas the primary oxygen concentration has no effect and the primary stream temperature has a moderate effect.

LES coupled with DQMOM is able to capture complex particle behaviors such as particle dispersion or particle clustering. Figure 3.7 shows the normalized particle concentration (weight) profile of each quadrature node at a selected timestep of simulation 1. The small particles (Figure 3.7a) are strongly influenced by the gas-phase turbulence and their jet spreads faster than the medium and large particles (Figures 3.7b and 3.7c). The large particle jet is only affected by the largest eddies and remains concentrated along the centerline. However, particle clustering effects are observed for each particle size and play an important role in predicting flame characteristics as shown further.

Figures 3.8 and 3.9 show instantaneous snapshots of the gas temperature profile on the central X-Z plane for different cases. The operating conditions of simulations 6 and 7 only differ by the value of $P_{O_2, pri}$. In both cases, the continuous flame region starts at about 30 cm from the burner inlet (Figures 3.8b and 3.8c). Little effect from $P_{O_2, pri}$ can be observed. Similarly, the gas temperature profiles are almost identical for simulations 8 and 9 (Figures 3.9a and 3.9b). The flame stand-off distance is about 10 cm in both cases and no effect of $P_{O_2, pri}$ is detected. On the other hand, the wall temperature appears to be an important parameter to predict flame ignition. When $T_{wall} = 1800K$ (Cases 8 and 9), the flame stand-off distance is short - about 10 cm - and the flame can be considered attached. When $T_{wall} = 1200K$ (Cases 6 and 7), the stand-off distance is about 30 cm and the flame is detached. Simulation 1 (Figure 3.8a) shows a stand-off distance of about 18 cm with an intermediate value $T_{wall} = 1500K$, thus confirming the trend of increasing stand-off distances with decreasing wall temperatures.

Depending on whether the flame is attached or detached, the ignition process is slightly different. When the flame is detached ($T_{wall} = 1200K$), three combustion regions can be
(a) Quadrature node 0 (40-micron particles)

(b) Quadrature node 1 (63-micron particles)

(c) Quadrature node 2 (100-micron particles)

Fig. 3.7: Normalized weight (number of particles/m$^3$) of each quadrature node at a given timestep of simulation 1
(a) Simulation 1 ($T_{wall} = 1500\,K$, $T_{pri} = 424.5\,K$, $P_{O_2, pri} = 10.45\%$)

(b) Simulation 6 ($T_{wall} = 1200\,K$, $T_{pri} = 424.5\,K$, $P_{O_2, pri} = 0\%$)

(c) Simulation 7 ($T_{wall} = 1200\,K$, $T_{pri} = 424.5\,K$, $P_{O_2, pri} = 20.9\%$)

Fig. 3.8: Gas temperature profiles (K) for simulations 1, 6 and 7
Fig. 3.9: Gas temperature profiles (K) for simulations 8 and 9 distinguished. The first one is a preheating region where no flame is observed. Pulverized coal particles are heated by turbulent mixing with the surrounding gas and by radiation from the walls but do not ignite yet. The second one is a growing flame region, where a small cluster of particles first ignites and then grows into an ignited particle cloud. The third one is a continuous flame region where the center of the jet is stably ignited.

Figures 3.10 and 3.11 present the profiles of different variables of interest for a selected timestep of simulation 6. The three combustion zones can be observed on Figure 3.10a. The preheating region ($x < 20 \text{ cm}$) ends with this ignition of a small cluster of particles at $x = 22 \text{ cm}$ which will grow into an ignited cloud, such as the one on Figure 3.8b at $x \approx 30 \text{ cm}$, and then merge with the continuous flame region ($x > 32 \text{ cm}$).
(a) Gas temperature (K)

(b) Particle temperature (K) for quadrature node 0 (40-micron particles)

(c) Particle temperature (K) for quadrature node 1 (63-micron particles)

(d) Particle temperature (K) for quadrature node 2 (100-micron particles)

(e) Devolatilization reactions rate (kg/s)

Fig. 3.10: Profiles of temperatures and devolatilization rate at a given timestep of simulation.
Fig. 3.11: Profiles of oxygen, raw coal mass fraction and char oxidation rate at a given timestep of simulation 6.
Figure 3.11a shows the oxygen mass fraction profile. The primary and secondary streams start to mix at \( x \approx 10\, \text{cm} \) and are fully mixed at \( x > 15\, \text{cm} \). The devolatilization of volatiles creates a rich region with low oxygen concentrations in the core of the continuous flame. Figures 3.10b, 3.10c and 3.10d present the particle temperature profile for each quadrature node. The small particles (quadrature node 0) heat up faster and are the first ones to ignite. The ignition of the small particle cluster at \( x = 22\, \text{cm} \) was favored by a large eddy which broke the main jet and brought hot gases into the center of the jet, as seen on Figure 3.10a. The small particles have a low thermal inertia and respond to this short event (Figure 3.10b), whereas the temperature of the medium and large particles is mostly unchanged by this event (Figures 3.10c and Figure 3.10d). This is confirmed by the raw coal mass profile for each quadrature node (Figures 3.11b, 3.11c and 3.11d). Only the 40-micron particles lose a significant fraction of raw coal during the cluster ignition.

The small particles are also the first ones to devolatilize and ignite in the continuous flame region (Figures 3.10b and 3.11b). The medium particles ignite shortly thereafter (Figures 3.11c) but the large particles are slower to heat up and only ignite at \( x > 40\, \text{cm} \) (Figures 3.10d and 3.11d). Figures 3.10e and 3.11e show respectively the total reaction rate for the devolatilization reactions and the char oxidation reactions. Generally, the devolatilization reactions rate is 2 to 3 orders of magnitude greater than the oxidation reactions rate and we can conclude that ignition occurs homogeneously solely because of devolatilization reactions.

When the flame is attached (\( T_{\text{wall}} = 1800\, \text{K} \)), the preheating region is shorter and no growing flame region is observed. Figure 3.12 presents the gas temperature profile and the particle raw coal mass profiles for each quadrature node at a selected timestep of simulation 8. The preheating region is about 10 cm (Figure 3.12a). The strong radiative flux from the walls ignite the small particles and their ignition induces the ignition of the medium and large particles shortly thereafter, as can be seen on the particle raw coal mass profile (Figures 3.12b, 3.12c and 3.12d). The flame has a cylindrical shape with an inner region that is rich and remains relatively cold until turbulence breaks the jet. Particles in this region devolatilize slower.
Fig. 3.12: Profiles of gas temperature (a) and particle raw coal mass ((b), (c) and (d)) at a given timestep of simulation 8.
The simulations with $T_{wall} = 1500K$ present an intermediate behavior where the ignition of individual particle clusters may occur but less frequent than with the lower wall temperature $T_{wall} = 1200K$.

All the previous images show that the gas-phase turbulence creates local and temporary pocket regions with specific temperature, gas species and particle concentration in the flame. Such effects are important to capture in order to predict the flame characteristics as they affect the stoichiometry and the kinetics of the local mixture and would not be captured in a RANS approach.

Figures 3.13 and 3.14 present time-averaged values of several quantities along the centerline for simulation 1. The gas and particle temperatures are plotted on Figure 3.13a. The peak in the gas temperature profile at $x \approx 0.2m$ corresponds to the flame ignition. After ignition, the gas temperature drops because of a fuel-rich region, as indicated by the oxygen mass fraction on Figure 3.14a. It then starts rising again as the fuel-rich region mixes with the surrounding lean mixture. Before ignition, the particle temperatures are close to the gas temperature. It can be noticed that the small particles are slightly hotter than the gas, whereas the large ones are colder. The small particles are indeed absorbing a significant amount of energy by radiation, energy that they partially transfer to the gas phase by convection, as shown on Figure 3.13b.

In the preheating region, radiation is the main heating process for particles since the net radiative heat flux for each particle size is greater than the net convective flux. The difference of temperature between the gas phase and the particles is the greatest during ignition. The particle thermal response time increases with the particle size. The peak in oxygen concentration observed on Figure 3.14a reveals that the primary and secondary streams mix rapidly and are fully mixed within a distance of 15 cm. Figure 3.14b shows the raw coal mass fraction profile for each particle size. As expected, the small particles devolatilized faster than the large ones.
Fig. 3.13: Time-averaged values of temperatures and heat fluxes along the centerline for simulation 1
Fig. 3.14: Time-averaged values of oxygen and raw coal mass fractions along the centerline for simulation 1
3.6.2 Validation and uncertainty quantification

analysis

This section aims at comparing the experimental results presented in Section 3.3 to the LES simulations. A formal validation and uncertainty analysis is performed with the Data Collaboration method.

3.6.2.1 Stand-off distance

In the experiments, flame ignition was determined by optical measurements. Comparing optical measurements with CFD results is a challenging task since CFD does not predict emission in visible light. Nevertheless, the gas temperature was judged the best criteria to determine if the flame was ignited. The coal jet is assumed to be ignited when the gas temperature reaches 1850K, which is an increase of 50K from the highest wall temperature. Hereafter, this temperature is called ignition judgment temperature. The stand-off distance $L_{SOD}$ is then defined as the first point in the x-direction where the gas temperature is greater than the ignition judgment temperature:

$$L_{SOD}(t) = \min \{x; T_g(x, y, z, t) > 1850 \text{ K}\} \quad (3.26)$$

where $D$ is the computational domain.

The time-averaged stand-off distance was calculated for each simulation. Results are presented in Table 3.7.

3.6.2.2 Surrogate model

From the results in Table 3.7, a quadratic surrogate model can be established to predict the average flame stand-off distance as a function of the parameters $T_{wall}$, $T_{pri}$ and $P_{O_2,pri}$:
Table 3.7: Time-averaged stand-off distance for each simulation

<table>
<thead>
<tr>
<th>Simulation</th>
<th>$L_{SOD}$ (cm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>17.5</td>
</tr>
<tr>
<td>2</td>
<td>19.0</td>
</tr>
<tr>
<td>3</td>
<td>18.9</td>
</tr>
<tr>
<td>4</td>
<td>15.2</td>
</tr>
<tr>
<td>5</td>
<td>16.0</td>
</tr>
<tr>
<td>6</td>
<td>26.9</td>
</tr>
<tr>
<td>7</td>
<td>26.3</td>
</tr>
<tr>
<td>8</td>
<td>9.7</td>
</tr>
<tr>
<td>9</td>
<td>9.7</td>
</tr>
<tr>
<td>10</td>
<td>26.6</td>
</tr>
<tr>
<td>11</td>
<td>25.6</td>
</tr>
<tr>
<td>12</td>
<td>11.3</td>
</tr>
<tr>
<td>13</td>
<td>8.0</td>
</tr>
</tbody>
</table>

$L_{SOD} = a_{00} + a_{11}X_1^2 + a_{10}X_1 + a_{22}X_2^2 + a_{20}X_2 + a_{33}X_3^2$

$+a_{30}X_3 + a_{12}X_1X_2 + a_{13}X_1X_3 + a_{23}X_2X_3$  \hspace{1cm} (3.27)

The parameters were normalized for a more comprehensive comparison:

$X_1 = (T_{wall} - 1200)/(1800 - 1200)$

$X_2 = (T_{pri} - 305)/(544 - 305)$

$X_3 = P_{O_2,pri}/20.9$

The polynomial coefficients are determined by a linear least square regression (Table 3.8). The infinite norm of the residual is 0.4 cm. The surrogate model can thus be judged as a good approximation of the simulation predictions.
Table 3.8: Surrogate model coefficients

<table>
<thead>
<tr>
<th>Coefficient</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>$a_{00}$</td>
<td>0.2739</td>
</tr>
<tr>
<td>$a_{11}$</td>
<td>0.0252</td>
</tr>
<tr>
<td>$a_{10}$</td>
<td>-0.1839</td>
</tr>
<tr>
<td>$a_{22}$</td>
<td>-0.0107</td>
</tr>
<tr>
<td>$a_{20}$</td>
<td>-0.0099</td>
</tr>
<tr>
<td>$a_{33}$</td>
<td>0.0014</td>
</tr>
<tr>
<td>$a_{30}$</td>
<td>-0.0083</td>
</tr>
<tr>
<td>$a_{12}$</td>
<td>-0.0224</td>
</tr>
<tr>
<td>$a_{13}$</td>
<td>0.0059</td>
</tr>
<tr>
<td>$a_{23}$</td>
<td>0.0088</td>
</tr>
</tbody>
</table>

Figure 3.15 shows the response surface of the model for a fixed value of $P_{O_{2, pri}}$ (Figure 3.15a) and for a fixed value of $T_{wall}$ (Figure 3.15b). The sensitivity to $P_{O_{2, pri}}$ is less than 0.5 cm, which is within the error of the surrogate model. The sensitivity to $P_{O_{2, pri}}$ is therefore negligible. The stand-off distance varies almost linearly with the wall temperature, whereas the relationship with the primary temperature is quadratic. The wall temperature can change the stand-off distance by more than 20 cm, whereas the primary stream temperature cannot modify it by more than 3 cm. The surrogate model thus shows a large sensitivity of the predicted stand-off distance to the wall temperature, a moderate sensitivity to the primary stream temperature and no sensitivity to the primary $P_{O_{2}}$.

3.6.2.3 Data collaboration analysis

A consistency analysis is performed using the Data Collaboration method described in Section 3.5.1. The predictions made by the surrogate model are compared to the experimental data presented in Figure 3.1.
Fig. 3.15: Surrogate model response surfaces for $P_{O_2, pri} = 10.45\%$ (a) and $T_{wall} = 1500K$ (b)
The dataset is made of 5 dataset units, each one corresponding to a value of $P_{O_2, pri}$ for which an experimental measurement was made. The experimental uncertainties were determined by replicate experiments, as explained in Section 3.3. The numerical uncertainty due to the time-averaging of the LES results and the surrogate model approximation is estimated to be $\pm 0.01\ m$ and is added to the experimental uncertainty for the consistency analysis. $P_{O_2, pri}$ is needed to predict the stand-off distance with the surrogate model but is not considered as a source of uncertainty in this study. Only $T_{wall}$ and $T_{pri}$ are considered as active variables and form the initial 2-dimensional hypercube $H$. The limits of the hypercube are determined by the uncertainty range of each active variable (see Table 3.4).

The consistency analysis reveals that the dataset is consistent. Therefore, there exists parameter values of $H$ for which the surrogate model predicts all the experimental observations of the dataset within their specified experimental uncertainties. The collection of those values constitutes the feasible region and is shown in Figure 3.16a. The feasible region is a small subset of the hypercube with wall temperatures near 1200 K and primary stream temperatures below 400 K. Figure 3.16b compares the experimental uncertainty ranges to the model predictions. The uncertainty bars in red correspond to the image of the whole hypercube under the surrogate model, whereas the uncertainty bars in green correspond to the image of the feasible set. The red bars cover a large part of the experimental uncertainty range, which suggests that the uncertainty ranges chosen for the active variables were appropriate. On the other hand, the model prediction within the feasible set of parameter values (green bars) has a small uncertainty. It predicts a stand-off distance value of about 27 cm with an uncertainty of about 1 cm. The predicted values are similar for all $P_{O_2, pri}$ values. The experimental results (Table 3.2) show that for almost all $P_{O_2, pri}$ values, the flame stand-off distance can be short ($L_{SOD} < 16\ cm$) or long ($L_{SOD} > 28\ cm$). According to the LES simulations and the data collaboration analysis, short stand-off distances can only be explained by high wall temperatures, whereas long stand-off distances require low wall temperatures.
Fig. 3.16: Feasible set of parameter values (a) and model predictions compared to experimental values (b)

Data collaboration of experiment and simulation indicates that the ignition stand-off distance in this oxy-coal jet is not a strong function of the oxygen content of the primary stream as the experimental data alone would indicate, but is predominately a function of the wall temperature in this small laboratory apparatus. The simulation and experimental data are consistent within the uncertainties of the experimentally measured stand-off distances and the uncertainties in the wall temperatures.

3.7 Conclusion

Large Eddy Simulation of oxy-coal flames were performed to study the ignition mechanisms using the Direct Quadrature Method of Moments. Simulation results demonstrate that LES coupled with DQMOM can capture particle dispersion and particle clustering effects. The flame ignition mechanisms were analyzed under different operating conditions. The local and transitional effects of turbulent eddies on the flame structure and the flame ignition were pointed out. The turbulence greatly modifies the local properties of the fluid
such as temperature or species concentration. LES can resolve those effects directly and should therefore lead to more accurate simulations than traditional RANS methods. LES simulations predict that the flame ignition is controlled by homogeneous reactions and that the heterogeneous reactions play a negligible role. The effects of three parameters on the flame stand-off distance were studied. Simulation predictions were compared to experimental data thanks to a surrogate model. The Data Collaboration method was used to perform a consistency analysis. The simulations were consistent with the experimental data but failed to predict any sensitivity to the primary $P_{O_2}$. The analysis suggests that the flame stand-off distance is mainly controlled by the wall temperature and that the role of primary $P_{O_2}$ is minor. Overall, this study shows that LES simulations can provided valuable insights on coal flame characteristics, guide experimentalists in their measurements by indicating where uncertainties lie and in the end, provide avenues for design, retrofit and scale-up of real boilers.

3.8 Acknowledgements

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4.1 Abstract

The goal of this study is to show that Large Eddy Simulation (LES) combined with the Direct Quadrature Method of Moments (DQMOM) can accurately predict coal flames ignition mechanisms and more generally be a valuable tool for predicting and understanding coal flames characteristics. A validation study of the ARCHES LES tool is performed. LES is applied to a pulverized coal jet flame ignited by a preheated gas flow. A Direct Quadrature Method of Moments has been developed to track the particle phase in an Eulerian framework. The simulation results are compared qualitatively and quantitatively to experimental observations for different inlet stoichiometric ratios. LES simulations are able to capture the various combustion regimes observed during flame ignition and to accurately model the flame stand-off distance sensitivity to the stoichiometric ratio. Gas temperature
and coal burnout predictions are also examined and show good agreement with experimental data. Overall, this study shows that high-fidelity LES simulations combined with DQMOM can yield a deeper understanding of complex coal flames and be a valuable tool for design, retrofit and scale-up of oxy-coal boilers.

4.2 Introduction

Coal is an important energy resource for electricity production as its reserves are far more abundant than those of other fossil fuels. As environmental regulations get stricter, coal thermal power plants need to control pollutants emissions such as $NO_x$, $SO_x$ and ash particles. Coal also plays an important role in global $CO_2$ emissions, which will need to be reduced to limit the effects of climate change [66]. Oxy-coal combustion, in which an $O_2/CO_2$ mixture replaces air, is one of the few possible capture technologies to enable $CO_2$ sequestration for existing coal-fired boilers. Despite the numerous research efforts that have been conducted, several gaps of knowledge have been identified and need to be addressed in more detail to obtain a fundamental understanding of the changes between oxy-fuel combustion and conventional air-fired combustion [79, 95]. One area of interest in rapidly implementing a strategy to retrofit existing air-fired burners or designing new ones is to understand how replacing air by an $O_2/CO_2$ mixture affects the flame ignition and the flame stability [15]. Flame stabilization is a key technology to control $NO_x$ emissions because it increases the size of the fuel-rich flame region where $NO_x$ is reduced to nitrogen [6]. Furthermore, flame stability of oxy-fuel combustion strongly depends on oxygen concentration of the gas mixture [21] and can also affect the heat transfer performance of the plant. Thus, flame stabilization plays a very important role in controlling both $NO_x$ and $CO_2$ emissions.

Although the reaction kinetics involved in coal particle ignition have been investigated extensively, most of the studies were done on single coal particles and on laminar flows. The single coal particle ignition mechanism cannot, however, sufficiently elaborate on the real turbulent flames due to certain limitations. First, the particles injected always contain a
size distribution which should be taken into account to obtain correct kinetics [7]. Second, interactions between particles play a crucial role in the ignition and combustion behavior and pollutant formation and destruction. Annamalai et al. [3, 4] studied isolated coal/char particles and found that the particles with relatively low volatile matter, which ignite heterogeneously when isolated, ignite homogeneously under cloud conditions. Finally, in real field boilers, with the mixing effect introduced by the large eddies in turbulence, single coal particle ignition mechanisms must be coupled with turbulent mixing to understand the flame stability. Turbulence can easily wipe off the gradients of concentrations, temperatures or velocities, which dramatically changes the ignition mechanisms. The fundamental mechanics of coal particles in turbulent flows in general are as yet poorly understood [14, 24, 52] and have not been included in optimizing the design and operating flow condition for coal combustors. Smith et al. [84] showed that particle behaviors such as particle dispersion and particle clustering affected operational considerations within a coal-fired boiler such as flame stability, pollutants formation or heat fluxes.

The gap between the fundamental research and the real industrial applications is thus still large and to understand how turbulent coal flames are stabilized, one must consider numerous complex physical processes and their interactions. This issue can be addressed by using Computational Fluid Dynamics (CFD), which was shown to be a very useful and accurate modeling framework for the description of particulate systems, and for the description of turbulent flows in general [24, 29]. Turbulence models based on the Reynolds-Averaged Navier–Stokes (RANS) equation are commonly used with gas and coal combustion models for the simulation. There is however evidence that RANS can be inaccurate in predicting particle-laden flows [5, 86] and a turbulence model based on large eddy simulation (LES) has been convincingly shown to be superior to RANS in accurately predicting turbulent mixing and combustion dynamics [56, 75].

This work presents a validation study of the LES tool ARCHES developed at the University of Utah to predict coal flame ignition. LES is applied to a pulverized coal jet flame ignited by a preheated gas flow. The simulation results are compared to experimental
data obtained for different inlet stoichiometric ratios [90]. LES simulations of this system were also previously performed by Yamamoto et al. [104] so results from both codes can be compared. ARCHES uses the Direct Quadrature Method of Moments (DQMOM) to track the particle phase in a Eulerian framework, whereas Yamamoto et al. used a Lagrangian framework. The goal of this study is to show that LES coupled with DQMOM can be a valuable tool for predicting and understanding coal flames characteristics and in the end, provide avenues for design, retrofit and scale-up of oxy-coal boilers.

### 4.3 Numerical modeling

Simulations were performed using ARCHES, a LES tool resulting from a ten-year partnership with the Department of Energy and the University of Utah. It is a massively parallel code that solves conserved quantities (mass, momentum, energy, scalar) spatially and temporally in a turbulent flow field, allowing for detailed and accurate simulations of fires and flames [87]. ARCHES includes particle phase representation through the use of a moment method, the direct quadrature method of moments (DQMOM). DQMOM combined with LES allows ARCHES to perform highly accurate simulations of coal combustion and gasification applications. The code is integrated into a C++ framework called Uintah which provides large-scale parallelization tools for physics components [9, 16, 73]. ARCHES is maintained in a repository and distribution is freely available [41]. The governing equations and numerical methods for the gas phase and the dispersed phase are summarized in the following sections.

#### 4.3.1 Governing equations of the gas phase

The gas phase is solved in a Eulerian manner by a finite volume method. Favre-filtered governing equations for continuity, momentum, enthalpy and scalars transport are solved. A second-order central difference scheme is used to calculate the convection and diffusion terms of the transport equations.
For the continuity equation, the mass source term due to coal combustion is taken into account. Coal combustion models are described in Section 4.3.2.

For the momentum equation, the Navier–Stokes equations are solved by taking the momentum source due to coal combustion into account. The subgrid scale (SGS) stress tensor is calculated by the dynamic subgrid scale model [36].

For the enthalpy equation, radiation, heat exchange between gas and coal particles, and heat transfer due to coal combustion are taken into account. The radiation source term is calculated by the discrete ordinate radiation method [2,46,47].

The gas-phase combustion is modeled through a mixture fraction approach and chemical equilibrium for the gas phase properties is assumed [93]. Spinti et al. [87] gave a more detailed description of the gas phase modeling in ARCHES.

4.3.2 Governing equations of the dispersed phase

In order to describe the evolution of the disperse particulate phase, LES equations must be coupled with a population balance equation (PBE). The method of moments (MOM) offers an attractive alternative to the traditional Lagrangian method where the PBE is tracked through its moments by integrating the internal coordinate [43]. The Direct Quadrature Method of Moments allows the description of a population of particles with a multivariate PBE, where two or more properties of the population are simultaneously tracked [26,59]. DQMOM has been shown to be a powerful approach for describing polydisperse solids undergoing segregation, growth, aggregation and breakage processes in the context of CFD simulations [26,32,60] and to greatly reduce the computational cost compared to the Lagrangian approach [17,18].

The DQMOM method was implemented into ARCHES to solve the dispersed phase in an Eulerian manner. ARCHES coupled with DQMOM has been proven to be a reliable and accurate method to simulate particles in turbulent flows [74]. A brief description of the particles representation and of the DQMOM equations is given in the following paragraphs.
A more detailed description of DQMOM implementation in ARCHES was previously given by Pedel et al. [74].

4.3.2.1 Number density transport equation

The particle Number Density Function (NDF) \( n \) describes the number of particles per volume as a function of the spatial location and of other independent variables. These independent variables are intrinsic characteristics of the particles (e.g., particle diameter, particle velocities, etc.) and are called internal coordinates. The internal coordinates are contained in the internal coordinate vector \( \xi \), and are dependent on space and time. At a given location in space and time \((x_0, t_0)\), the total number of particles in a volume \( V \) is given by:

\[
    n_p(x_0, t_0) = \int_V \int n(\xi; x_0, t_0) d\xi dV \tag{4.1}
\]

The velocity of the NDF in both real and internal coordinate space can be described using a set of internal coordinates \( \xi = [\xi_1, \xi_2, ...] \), and external coordinates \( x = [x_1, x_2, x_3] \). For a single particle, the time derivatives of these variables may be written as

\[
    \frac{dx_i}{dt} = v_{p,i} \tag{4.2}
\]

\[
    \frac{d\xi_j}{dt} = G_j \tag{4.3}
\]

where \( v_{p,i} \) is the \( i^{th} \) component of the particle velocity vector \( \mathbf{v}_p \).

The NDF transport equation is given by:

\[
    \frac{\partial}{\partial t} (n(\xi; x, t)) + \frac{\partial}{\partial x_i} ((v_{p,i}|\xi) n(\xi; x, t)) = - \frac{\partial}{\partial \xi_j} ((G_j|\xi) n(\xi; x, t)) + h(\xi; x, t) \tag{4.4}
\]
In this equation, the variable \( \langle v_{p,i} | \xi \rangle \) represents the velocity in real space of the NDF. This quantity is an average over particles having the same coordinates \((\xi; x, t)\) and is a distribution conditional to the value of the vector \( \xi \). Similarly, the quantity \( \langle G_i | \xi \rangle \) describes the velocity of the NDF in the phase space. The term \( h \) is a source term representing the birth and death of particles in the domain. This term is set to zero as particle breakage and aggregation effects are neglected.

### 4.3.2.2 Direct Quadrature Method of Moments

The Direct Quadrature Method of Moments (DQMOM) approximates the NDF by a summation of multidimensional Dirac delta functions. Figure 4.1 gives an example for bivariate Gaussian distribution (i.e., two internal coordinates) of the NDF and its quadrature approximation with three quadrature nodes. In this approximation, the NDF is written as:

\[
n(\xi; x, t) = \sum_{\alpha=1}^{N} \left( w_{\alpha} \prod_{i=1}^{N_{\xi}} \delta (\xi_i - \langle \xi_i \rangle_{\alpha}) \right)
\]  

where both the weight \( w_{\alpha} \), which defines the number of particles per volume associated with the quadrature node \( \alpha \), and the average internal coordinate value \( \langle \xi \rangle_{\alpha} \) depend on space and time, but the dependence is omitted for clarity of notation. \( N \) is the number of quadrature nodes and \( N_{\xi} \) the number of internal coordinates.

DQMOM solves transport equations for the weights and weighted abscissas of the quadrature approximation. These transport equations are simply scalar transport equations, and are easily implemented into a CFD code:

\[
\frac{\partial}{\partial t} (w_{\alpha}) + \frac{\partial}{\partial x_i} (\langle v_{p,i} \rangle_{\alpha} w_{\alpha}) = a_\alpha
\]

\[
\frac{\partial}{\partial t} (s_{n\alpha}) + \frac{\partial}{\partial x_i} (\langle v_{p,i} \rangle_{\alpha} s_{n\alpha}) = b_{n\alpha}
\]
where $\xi_n = w_\alpha \langle \xi_n \rangle_\alpha$ is the weighted abscissa for the $n^{th}$ internal coordinate and $\langle v_{p,i} \rangle_\alpha$ is the average internal coordinate velocity. The right-hand side terms $a_\alpha$ and $b_{n,\alpha}$ are obtained by solving a linear system of equations.

### 4.3.2.3 Single coal particle description

Following Smoot and Smith [85], a single coal particle can be characterized using several particle independent variables. These are denoted:

- Raw coal $\alpha_{c_j}$
- Char $\alpha_{h_j}$
- Moisture $\alpha_{w_j}$
- Ash $\alpha_{a_j}$
- Particle diameter $d_{pj}$
- Particle enthalpy $h_{pj}$
- Particle velocity vector $v_{pj}$

The above quantities use the subscript $j$ to denote the $j^{th}$ particle. The variable $r$ can be used to denote reaction rates, so that $r_{hj}$ would be the net char reaction rate for the $j^{th}$ particle. Using this nomenclature, physical processes important to coal particles are depicted in Figure 4.2. The raw coal can react to form gaseous volatile matter in devolatilization reactions and solid char; the solid char can be oxidized to form more gaseous products. The ash mass is fixed, and ash is treated as inert.

In this study, a total of seven internal coordinates were chosen to represent particle characteristics in the DQMOM formulation:

- mass of raw coal $\alpha_c$
- mass of char $\alpha_h$
- particle diameter $d_p$
- particle enthalpy $h_p$
- particle velocity components $v_x, v_y, v_z$

The particle diameter $d_p$ was assumed to be constant. To reduce the number of internal coordinates and the computational cost, the moisture in the particles was assumed to be evaporated before entering the domain and the water content was added to the inlet stream.

4.3.2.4 Particle velocity model

The aggregation/breakage phenomena are neglected and the only physical process modeled is the drag force, which can be described by the Stokes drag law. For a mesoscale size particle, other forces such as lift can be neglected and the momentum equation for the particle can be expressed by an ordinary differential equation.

$$\frac{dv_{p,i}}{dt} = \frac{f_{drag}}{\tau_p} (u_{g,i} - v_{p,i}) + \frac{g_i (\rho_p - \rho_g)}{\rho_p} \tag{4.8}$$
where $\mathbf{g}$ is the gravity force acting on the particle, $\rho_g$ and $\rho_p$ are respectively the gas and particle densities, $\mathbf{u}_g$ is the gas velocity, and $f_{\text{drag}}$ is the coefficient of the drag force, which has a close relationship with the particle Reynolds number [85]:

$$f_{\text{drag}} = \begin{cases} 
1 & Re_p < 1 \\
1 + 0.15 Re_p^{0.687} & 1 < Re_p < 1000 \\
0.0183 Re_p & Re_p > 1000 
\end{cases}$$ (4.9)

$$Re_p = \frac{\rho_p d_p |u_p - u_g|}{\mu_g}$$ (4.10)

where $\mu_g$ is the gas dynamic viscosity.

In equation (4.8), $\tau_p$ is the particle relaxation time:

$$\tau_p = \frac{\rho_p d_p^2}{18 \mu_g}$$ (4.11)
4.3.2.5 Devolatilization model

In this study, an empirical devolatilization model proposed by Yamamoto et al. is used [104]. The approach was proven accurate and inexpensive by comparison to more sophisticated models such as the distributed activation energy model (DAEM), FLASHCHAIN [71] or chemical percolation devolatilization (CPD) [31]. The devolatilization process is described as:

\[
\frac{r_v}{(\text{raw coal})} \rightarrow Y(\text{volatiles}) + (1-Y)(\text{char})
\]  

(4.12)

where \(Y\) is a stoichiometric coefficient. The devolatilization rate of the model \(r_c\) is based on a single reaction:

\[
r_v = \frac{d\alpha_c}{dt} = -Fk_0 e^{\left(-E_v/RT_p\right)}\alpha_c
\]  

(4.13)

\[
F = e^{\left(\sum_{i=0}^{5} c_i G_v^i\right)}
\]  

(4.14)

where \(k_0\) is the frequency factor, \(E_v\) is the activation energy and \(R\) is the gas constant. \(F\) is the modification factor and is expressed as a function of fraction devolatilized \(G_v = (\alpha_{c,0} - \alpha_c)/\alpha_{c,0}\). Here \(\alpha_{c,0}\) is the initial mass of raw coal.

The model parameters \(k_0\), \(E_v\), and \(c_i\) are determined by minimizing the difference between predictions of the DAEM model and the present model at the heating rate of 100,000 K/s. Figure 4.3 shows results calculated by the Yamamoto model and DAEM. The coal used here is the same as the coal used for the experiments. The results obtained by the present model show good agreement with CPD results. The average error of the fraction devolatilized is about 1.5% for both heating rates of 100,000 and 20,000 K/s. This model
thus offers the advantage to predict the fraction devolatilized with the same accuracy as the DAEM model but with the low computational cost of a single-rate reaction model.

4.3.2.6 Char oxidation model

After the raw coal in a particle has devolatilized, it forms char. Historically, most kinetic data on burning coal char particles have been interpreted using an $n^{th}$ order Arrhenius model of char combustion. In this representation, the global surface reaction rate follows the expression

$$q = k_s (T_p) P_{O_2,s}^n$$

where $n$ is the reaction order, $T_p$ is the particle temperature, and $k_s$ is a temperature-dependent rate coefficient, assumed to follow an Arrhenius form:
\[ k_s (T_p) = A_s e^{E_s / R T_p} \] (4.16)

The parameter values were chosen from the experimental study of Murphy and Shaddix [70] for the Eastern bituminous coal: \( A_s = 344 \text{ mol/s/m}^2/\text{atm}^n \), \( E_s = 45.5 \text{ kJ/mol} \) and \( n = 0.18 \). The partial pressure of oxygen at the char surface \( P_{O_2,s} \) is obtained by solving a gas-phase diffusion equation.

4.3.2.7 Particle heat transfer

The particle is heated by convection, radiation and reaction enthalpy changes:

\[ \frac{d h_p}{d t} = Q_{rad} + Q_{conv} + r_j h_j, \] (4.17)

where \( Q_{rad} \) represents the net radiation flux to a particle, \( Q_{conv} \) represents energy transfer due to convection and conduction between the gas and the particle, and \( r_j h_j \) represents both the amount of energy lost by the particles due to lost mass, and the enthalpy released during devolatilization and oxidation reactions.

Convection

The convection term is expressed as:

\[ Q_{conv} = Nu \pi k_g (T_g - T_p) d_p \frac{\kappa / 2}{e^{\kappa / 2} - 1} \] (4.18)

where \( T_g \) is the gas temperature, \( k_g \) is the gas thermal conductivity, and \( \kappa = -W C q d_p C_{pg} / k_g \) is a modified version of the Peclet number [70]. \( C_{pg} \) is the gas heat capacity. The Nusselt number \( Nu \) is calculated with the following correlation [55]:

\[ Nu = 2.0 + 0.65 Re_p^{1/2} Pr^{1/3}. \] (4.19)
The particle temperature $T_p$ is obtained from the particle enthalpy by assuming Merrick’s correlations for the particle heat capacity [65].

Radiation

The radiative flux is given by:

$$Q_{rad} = Q_{\text{incident}} - Q_{\text{emitted}}$$

(4.20)

where $Q_{\text{incident}}$ is the incident radiative flux to the particle and $Q_{\text{emitted}}$ is the radiative heat flux emitted by the particle. The absorption coefficient of the coal particles is assumed constant: $Q_{abs} = 0.8$. The incident flux is obtained through the Discrete Ordinates radiation model, whereas the emitted flux is calculated using the blackbody emissive power:

$$Q_{\text{emitted}} = Q_{abs} \pi d_p^2 \sigma T_p^4.$$  

(4.21)

Following [85], the coupling with the gas phase is done by modifying the gas absorption coefficient:

$$k_{\text{abs}, g} = k_{\text{abs}, g} + k_{\text{abs}, p}$$

(4.22)

with

$$k_{\text{abs}, p} = \sum_{j=1}^{n_p} \frac{\pi}{4} Q_{abs} d_{pj}^2$$

(4.23)

4.3.3 Simulation details

The computational domain is a 0.6x0.3x0.3m box. The domain is decomposed into a uniform cartesian mesh of 54 million cells, which translates into a cellsize $\Delta = 1.0 \text{mm}$. The mixture of coal and air is injected through the primary nozzle (diameter: 7 mm) and ignited by the surrounding preheated gas. The primary and secondary inlets are located
at the center of the X=0 face and outlet boundary conditions are applied at the X=0.6 m face. The side boundaries are set as open pressure boundaries. The operating conditions are summed up in Table 4.1. The Reynolds number of the primary jet is about 4500. Coal analysis data are given in Table 4.2. The initial coal particle density is 1400 kg/m$^3$. The coal particles are injected with the primary stream and are assumed to have the same temperature and velocity as the gas phase at the inlet.

Table 4.1: Operating conditions

<table>
<thead>
<tr>
<th></th>
<th>Primary stream</th>
<th>Preheated secondary stream</th>
</tr>
</thead>
<tbody>
<tr>
<td>Average velocity (m/s)</td>
<td>10</td>
<td>4.8</td>
</tr>
<tr>
<td>Temperature (K)</td>
<td>300</td>
<td>1510</td>
</tr>
<tr>
<td>Mass fraction</td>
<td></td>
<td></td>
</tr>
<tr>
<td>$N_2$</td>
<td>0.768</td>
<td>0.760</td>
</tr>
<tr>
<td>$O_2$</td>
<td>0.232</td>
<td>0.101</td>
</tr>
<tr>
<td>$CO_2$</td>
<td>0.0</td>
<td>0.093</td>
</tr>
<tr>
<td>$H_2O$</td>
<td>0.0</td>
<td>0.045</td>
</tr>
</tbody>
</table>

Table 4.2: Coal analysis

<table>
<thead>
<tr>
<th>Ultimate analysis (daf, wt%)</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>C</td>
<td>83.1</td>
</tr>
<tr>
<td>H</td>
<td>4.6</td>
</tr>
<tr>
<td>O</td>
<td>9.9</td>
</tr>
<tr>
<td>N</td>
<td>1.9</td>
</tr>
<tr>
<td>S</td>
<td>0.5</td>
</tr>
<tr>
<td>Proximate analysis (dry basis, wt%)</td>
<td></td>
</tr>
<tr>
<td>Volatile matter</td>
<td>31.1</td>
</tr>
<tr>
<td>Fixed Carbon</td>
<td>54.0</td>
</tr>
<tr>
<td>Ash</td>
<td>14.9</td>
</tr>
</tbody>
</table>
The inlet stoichiometric ratio (SR) is modified by varying the coal mass flow rate. Three inlet stoichiometric ratios are considered (0.14, 0.22 and 0.36) and a LES simulation is performed for each value of SR. Each simulation was run for 5 days on 1,500 CPUs with timesteps of about 15 $\mu$s, which resulted in a total simulation time of about 1 s. Time-averaging was done on the last 0.5 second of data. Particles are modeled using the DQMOM method with seven internal coordinates, as described in Section 4.3.2, and three quadrature nodes, each quadrature node corresponding to a single particle size. Table 4.3 shows the particle diameter associated with each quadrature node as well as the volume percentage associated with each particle size. The volume percentages were determined from the experimental particle size distribution [90].

### 4.4 Results and discussion

The flame stand-off distance has been previously measured from images taken by a high-speed camera [90]. In addition, temperature and coal burnout on the centerline were measured. First, LES results are compared qualitatively with the flame structure observed in the experimental images and with the simulation results obtained by Yamamoto et al. [104]. Then simulation results of temperature, coal burnout, and stand-off distance are compared quantitatively with experimental data.

#### 4.4.1 Qualitative analysis

Figure 4.4 shows the distribution of instantaneous gas temperature obtained by LES on the central X-Z plane for each stoichiometric ratio. Assuming gas temperature is a good indicator of flame ignition, it can be noticed that the flame stand-off distance increases with

<table>
<thead>
<tr>
<th>Quadrature node</th>
<th>Particle diameter ($\mu m$)</th>
<th>vol%</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>20</td>
<td>20.0</td>
</tr>
<tr>
<td>1</td>
<td>50</td>
<td>60.0</td>
</tr>
<tr>
<td>2</td>
<td>100</td>
<td>20.0</td>
</tr>
</tbody>
</table>
Fig. 4.4: Gas temperature profiles (K) obtained by LES for various inlet stoichiometric ratios (SRs)
increasing stoichiometric ratios. The stand-off distance is about 12 cm for SR=0.14 and about 22 cm for SR=0.36. As less volatiles devolatilize when the stoichiometric ratio increases, the flame length also becomes shorter. The flame ends at $x \approx 55\,cm$ for SR=0.14 and at $x \approx 38\,cm$ for SR=0.36. Three combustion regions were observed experimentally. The first one is a preheating region where no flame is observed. Pulverized coal particles are heated by turbulent mixing with the surrounding gas but do not ignite yet. The second one is a growing flame region, where a small cluster of particles first ignites and then grows into an ignited particle cloud. The third one is a continuous flame region where the center of the jet is stably ignited. LES predictions agree with those observations, as shown on Figure 4.4.

LES coupled with DQMOM is able to capture complex particle behaviors such as particle dispersion or particle clustering. Figure 4.5 shows the particle concentration (weight) profile of each quadrature node at a given timestep of the LES simulation for SR=0.36. The small particles (Figure 4.5a) are strongly influenced by the gas-phase turbulence and their jet spreads faster than the medium and large particles (Figures 4.5b and 4.5c). The large particles are mostly unaffected by the gas phase turbulence and remain concentrated along the centerline. Particle clustering effects can be observed for each particle size. Small particles almost fill the whole width of the domain for $x \geq 50\,cm$, even if their concentration is small. The common Lagrangian approach to track particles often will not predict the presence of particles in small concentration regions as the number of tracked particles is too low [104]. The DQMOM method predicts a full particle size distribution at any point in the domain.

4.4.2 Quantitative analysis

In the experiments, flame ignition was determined by optical measurements [90]. Comparing optical measurements with CFD results is a challenging task since CFD does not predict emission in visible light. Nevertheless, the gas temperature was judged the best criteria to determine if the flame was ignited. Following Yamamoto et al. [104], the coal jet is judged to be ignited when the temperature reaches 1560 K, which is an increase of 50 K
Fig. 4.5: Instantaneous profiles of each quadrature node weight (number of particles x $10^9/m^3$) for SR=0.36
from the preheated gas temperature (1510 K). This temperature is called the ignition judgment temperature. The stand-off distance $L_{SOD}$ is then defined as the first point in the $x$-direction where the gas temperature is greater than the ignition judgment temperature:

$$L_{SOD}(t) = \min\{x; T_g(x, y, z, t) > 1560 K\}$$

(4.24)

where $D$ is the computational domain.

The time-averaged stand-off distance was calculated for each simulation. Figure 4.6 compares the observed stand-off distance to the averaged stand-off distance predicted by LES simulations. The experimental stand-off distance reported in Figure 4.6 corresponds to the beginning of the growing flame region. Simulations results are in good agreement with experimental data.

Figures 4.7, 4.8 and 4.9 present time-averaged values along the centerline of gas temperature and coal burnout. Figure 4.7 compares the gas temperature predictions to the experimental values for SR=0.14 and SR=0.36. For SR=0.14, gas temperatures predicted by LES are in relative good agreement with the experimental values. A peak is observed at $x \approx 0.18 m$. This peak was not predicted by Yamamoto et al. [104] and may be the result of the fast and simultaneous devolatilization of the 20-micron particles (quadrature node 0), as shown on Figure 4.9 for SR=0.22. Taking into account smaller particles would probably smooth out the peak and increase the predicted temperature at $x = 0.1 m$. For SR=0.36, the devolatilization process is slower (Figure 4.8) and the simulated gas temperature is an good agreement with the measurements for $x \leq 0.2 m$. However, LES simulations predict high temperatures above 1300 K for $x > 0.2 m$, which was not observed experimentally. Figure 4.8 shows coal burnout comparisons for SR=0.36. Measurements were made for three particle size ranges which can be compared to the LES single particle size of each quadrature node. For the medium (37-74 microns) and the coarse (74-150 microns) particles, LES predictions are consistent with the experimental values as they fall within the uncertainty
Fig. 4.6: Validation results for the flame stand-off distance
Fig. 4.7: Comparison between experiments and LES results of gas temperature
Fig. 4.8: Comparison between experiments and LES results of gas coal burnout (SR=0.36)
ranges. For the fine particles (7-37 microns), no uncertainty range were provided in the experimental data. Simulation results seem to under-predict the mass loss for $x \leq 0.2 \text{ m}$. This early mass loss can be attributed to the smallest particles in this range (less than 20 microns) which are not taken into account in the LES simulations. Figure 4.9 compares the total coal burnout predicted by the simulations to the experimental measurements for SR=0.22. The total coal burnout for the simulations is obtained by averaging each quadrature node burnout with their respective weight. As the small particles disperse faster, the relative weight of the medium and large particles become more important as $x$ increases. LES total coal burnout prediction is in good agreement with the experiments.

Overall, the quantitative analysis of the LES results reveals a general consistency with the experimental data. LES simulations were able to accurately predict the flame stand-off.
distance and the coal burnout for different stoichiometric ratios. Further work would be required to understand the disagreement in gas temperatures along the centerline.

4.5 Conclusion

Large Eddy Simulations of a pulverized coal jet flame ignited by a preheated gas flow were performed for three inlet stoichiometric ratios using the Direct Quadrature Method of Moments (DQMOM). Simulation results were compared qualitatively and quantitatively to experimental observations. LES simulations were able to capture the three combustion regimes observed during flame ignition and to accurately model the flame stand-off distance sensitivity to the stoichiometric ratio. Gas temperature and coal burnout data were also compared and showed good agreement with the experimental results. Overall, this validation study shows that LES coupled with DQMOM can accurately predict coal flames ignition mechanisms and more generally be a valuable tool for predicting and understanding coal flames characteristics.

4.6 Acknowledgements

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CHAPTER 5

CONCLUSION

The implementation of DQMOM in the LES code ARCHES provides the ability to perform accurate and fast simulations of particle-laden turbulent flows. Simulation results show that LES combined with DQMOM accurately models complex particle behaviors in turbulent flows such as particle dispersion or particle clustering. LES simulations of air-fired and oxy-fired coal flame were performed to predict flame stability and showed that the LES with DQMOM approach is able to predict coal-flame ignition mechanisms and, more generally, to yield a better understanding of the multiphysics phenomena involved in oxy-coal combustion.

The original contributions of this research can be listed as the following:

- Efficient, accurate and stable implementation of DQMOM. So far, DQMOM has been used with only a few internal coordinates and a limited number of quadrature nodes, due to computational time and numerical stability issues. This research formulates a fast, accurate and stable implementation of DQMOM that can handle a large number of internal coordinates and quadrature nodes.

- Implementation of DQMOM in a LES code. Few LES codes can handle multiphase reacting flows and the ones who can usually use an Eulerian-Lagrangian approach. To the author’s knowledge, Arches is the first LES code able to simulate coal flames with DQMOM.

- LES simulations of coaxial particle-laden jets with DQMOM and comparison with experimental data.

- LES simulations of oxy-coal flames to investigate flame stability. Few LES simulations of oxy-coal flames have ever been performed. This research provides the first LES simulations on oxy-coal flame stability and comparison with experimental data.
Overall, this research shows that high-fidelity LES simulations combined with DQMOM can provide a deeper understanding of complex coal flames and their ignition mechanisms, indicate where experimental uncertainties lie and in the end, be a valuable tool for the design, retrofit and scale-up of oxy-coal boilers. This research focused on the validation of the approach and simulations were limited to the near burner region. Further work will consist of the simulation of laboratory and commercial scale oxy-fired boilers. LES plus DQMOM capabilities also lead the way to accurate simulations of a wide range of other applications since turbulent multiphase flows are encountered in a variety of engineering processes (fluidized beds, aerosols, soot formation, etc.).
REFERENCES


