A Discrete-Continuous Method for Predicting Thermochemical Phenomena in a Cement Kiln and Supporting Indirect Monitoring

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Abstract. Thermochemical phenomena involved in cement kilns are still not well understood because of their complexity, besides technical difficulties in achieving direct measurements of critical process variables. This article addresses the problem of their comprehensive numerical prediction. The presented numerical model exploits Computational Fluid Dynamics and Finite Difference Method approaches for solving the gas domain and the rotating wall, respectively. The description of the thermochemical conversion and movement of the powder particles is addressed with a Lagrangian approach. Coupling between gas, particles and the rotating wall includes momentum, heat and mass transfer. Three-dimensional numerical predictions for a full-size cement kiln are presented and they show agreement with experimental data and benchmark literature. The quality and detail of the results are believed to provide a new insight into the functioning of a cement kiln. Attention is paid to the computational burden of the model and a methodology is presented for reducing the time-to-solution and paving the way for its exploitation in quasi-real-time, indirect monitoring.

Keywords: Cement kiln, computational fluid dynamics, discrete element method.
1. Introduction

Cement represents the fundamental material in many building and civil engineering applications. Among several types, the most common is Portland, which is made of a mixture of gypsum and clinker. The clinker production process includes a pyro-processing treatment taking place inside dry kilns equipped with five or six cyclone preheaters and a precalciner. The operation is dramatically energy-intensive, with a heat use of 2900-3300 MJ per ton of clinker [1]. Here the raw material comes from the grinding operation in the state of powder particles; a first heating up to about 870 °C is performed inside the preheaters then the precalciner, subsequently the material, almost completely calcined, enters the dry kiln, where its temperature is raised up to 1400-1500 °C; follows a rapid cooling, which prevents the decomposition of metastable phases.

The dry kiln is a large reactor consisting of a rotating tube with an outer steel shell and inner refractory lining. It is typically inclined 1-3 degrees (longitudinal axis with respect to horizontal line) and rotates at 1-5 rpm [1]. The typical length-to-diameter ratio is 16:1-13:1 and recently even shorter [1]. The particles of the material bed flow inside the kiln and exchange heat with the rotating wall and a hot countercurrent air flow i.e. the freeboard gas, whose initial enthalpy is provided by heat recovery from the cooling stage and fuel burning. As a result, the temperature of the material is raised while it moves towards the exit and several chemical conversions take place until clinker is obtained. The thermochemical phenomena occurring inside the material bed are heavily affected by its powder state [2]. Indeed, the in-plane recirculation of the particles, which is due to the dragging of the rotating wall, produces a redistribution of heat within the material bed and helps reducing the inhomogeneity of properties in the final product.

A highly aggressive environment and physical inaccessibility prevent the direct monitoring of many process variables inside a cement kiln [3]. The consequent lack of real data impacts negatively on the understanding and optimization of the system, which accounts for a major fraction of the plant heat losses [4]. Nevertheless, process optimization is a key target for the global cement industry, which is facing constantly increasing coal prices and always more demanding environmental regulations [5, 6]. Numerical models could represent effective tools for filling the knowledge gap: indeed, their comprehensive predictions could constitute the theoretical basis of unresolved CFD-DEM methods like those reported in [37–39]. CFD (Computational Fluid Dynamics) studies of the cement kilns. A 1D model based on heat and mass balances is presented in [24]. The same model has been presented, for predicting the flow of granular materials inside rotating drums: the governing equations are similar to those derived for conventional fluids, but conservation of the kinetic energy is also accounted in the formulation along with momentum and mass transport. In [15], an analytical prediction of the particles flow has been obtained for the specific case of rolling-mode operated drums. Discrete Element Method (DEM) has been increasingly used for predicting particles movement in rotating drums [16–22], following the breakthrough of higher computing power. Its undoubted advantage respect to other methods like stochastic or empirical is the possibility of obtaining a deterministic estimation of the trajectory of each particle. However, DEM presents high computation requirements which grow as the number of particles increases, therefore preventing the application to many large-scale scenarios in industry unless periodic boundaries and/or agglomeration [23] are used for limiting both the size of the domain and the total number of particles.

Different authors have discussed numerical studies of the thermochemical phenomena occurring inside cement kilns. A 1D model based on heat and mass balances is presented in [24]. The same model has been resumed with minor changes by other authors [25, 26]. CFD (Computational Fluid Dynamics) studies of the freeboard gas and mostly focused on the burner design are reported in [27–32]. In [33] it is presented the first complete model of a cement kiln, which includes a CFD study of the freeboard gas along with numerical predictions of the material bed and the rotating wall, as well. Analogous models have been lately presented in [34, 35]. However, in these studies the accurate representation of the freeboard gas contrasts with strong assumptions over the material bed, whose flow is described as the rigid translation of a continuum along the axial direction of the kiln, regardless its real powder state and the consequent in-plane recirculation.

In [36] it is presented a retrospective review of mathematical models for the representation of a porous medium and the related heat and mass transfer coefficients by means of an equivalent continuum. These constitute the theoretical basis of unresolved CFD-DEM methods like those reported in [37–39] and others.

The present contribution discusses the application of an eXtended Discrete Element Method (XDEM) approach for predicting the thermochemical phenomena occurring in a dry kiln for clinker production.
XDEM is a numerical framework based on a coupled CFD-DEM method. The model includes: a Lagrangian prediction of the particles flow and the heat exchange between particles, along with heat and mass transport inside particles and their chemical conversions and morphology changes; a CFD prediction of the gas domain including fuel transport and its combustion; an FDM (Finite Difference Method) approach for solving the heat equation on the rotating wall. Particles-gas coupling includes mass, momentum and convective heat transfer. Particles-wall conductive and gas-wall convective heat transfer are included in the formulation, as well. A simplified resistance model is included for predicting the gas-wall-particles radiative heat transfers. XDEM has already been tested in the prediction of the phenomena involved in several industrial processes including pyrolysis [40], metal manufacturing [41, 42], drying [43, 44], CaCO$_3$ decarbonation [45, 46] and others [47–49].

The model presented exploits several assumptions for limiting the computation burden. These will be detailed in the following discussion. An effective numerical methodology is presented, for reducing the time-to-solution to the length scale of hours. The computation efficiency of the numerical tool is expected to allow its integration into the plant SCADA architecture as an indirect monitoring tool, useful for backup and/or integration of the hardware distributed sensors.

The predictions discussed in this paper are consistent with experimental and literature data. To the best knowledge of the authors, the present study is the first application of a combined Eulerian-Lagrangian method to the prediction of the phenomena involved in the clinker production.

2. The Real Kiln

![Fig. 1. Kiln geometry, detail of air-intake and positions of DEM domain at different times.](image)

A real dry kiln for clinker production (length: 54 m; internal radius: 2.1 m; external radius: 2.3 m) has been considered. Its geometry is depicted in Fig. 1. The kiln is operated at a filling level of 15 % and processes a total mass flow of 140 t/h. The material enters the kiln at a temperature of 1220 K. The chemical composition of a fresh sample, i.e. at room temperature, is reported in Table 1. Loss on Ignition (LOI) is defined as the mass loss of a fresh sample ignited at 1173 to 1273 K. This is consequent to the decarbonation process of limestone into calcium oxide (CaCO$_3$→CaO+CO$_2$). As typical of dry kilns, the one considered here receives the input material almost completely calcined with a decarbonation percentage at 98 %, therefore LOI occurring inside the kiln is negligible. The kiln rotates at 4 rpm and is tilted 2 degrees. The retention time (i.e. the mean time spent by the material inside the kiln) is 20 min. The hot gas moves countercurrent to the material bed and the total flow includes: the secondary air from heat recovery, which enters at a rate and temperature of 80000 m$^3$/h and 950 K, respectively; the primary air with the coal particles in suspension, which is injected through the burner at room temperature and at a rate of 5700 m$^3$/h. The kiln is operated at a depression of -200 Pa respect to the environment, in order to force the extraction of the exhaust gases. The set of measurements include the flame temperature (in the range of 2073-2173 K), the mean temperature...
of the material in the sintering zone (1733 K), the mean temperature of the material at the exit (1673 K) and
the speed of the gas at the exhaust (15-17 m/s).

Table 1. Chemical composition of a fresh sample.

| Specie   | Mass fraction |
|----------|---------------|
| SiO$_2$  | 0.14          |
| Al$_2$O$_3$ | 0.03         |
| Fe$_2$O$_3$ | 0.02         |
| CaO      | 0.43          |
| MgO      | 0.03          |
| LOI      | 0.35          |

Cement kilns are giant reactors that represent some of the largest-size, moving machineries ever built. Consequently, the CFD-DEM representation must deal with a dramatically broad range of scales, being the characteristic size of clinker lumps in the order of centimeters [50]. A numerical simulation that brutally addresses the kiln at the whole size is clearly impossible due to the excessive computation burden. Therefore, simplifications have been made and they are discussed in the following subsections.

3. The Model

3.1. DEM Approach for the Powder Domain

Two periodic boundaries orthogonal to the kiln axis have been imposed, spaced of 0.4 m each other. The thin slice constituting the DEM domain is depicted in Fig. 2, together with the frame of reference that will be used from here on. The volume has been filled with 4000 spherical particles with a diameter of 0.06 m, providing the filling level required. The diameter of the particles and the spacing between the periodic boundaries have been chosen by upscaling the analogous setup reported in [22].

![DEM domain](image)
3.1.1. Particles flow

The linear and angular accelerations of each particle are calculated by applying Newton’s laws of motion, then position and orientation are updated by integration. Each particle is subject to: the gravitational force, which is calculated according to the mass densities of the constituting species; contact forces due to the mechanical interactions with neighbors and/or boundaries, which include viscous and elastic components in both the normal and shear direction, more details are provided in [51]; a drag force $F_{\text{drag}}$ from the surrounding fluid, whose analytical expression is detailed in Section 3.4.

3.1.2. Conductive and radiative heat exchange between particles

Each $i$-particle exchanges heat with every other $j$-particle that is comprised within the cutoff distances for conduction and radiation. Conductive and radiative heat exchanges are calculated according to Eq. (1) and Eq. (2), respectively. Here $V_{(i\rightarrow j)}$ and $A_c$ are the reciprocal view factor and the contact area, their expressions are retrieved from [52] and reported in Eq. (3) and Eq. (4), respectively.

$$\dot{q}_{i,\text{cond}} = \sum_{j=1}^{N} A_{c,i,j} \frac{1}{1 + k_i/k_j} \frac{T_{p,i} - T_{p,j}}{\Delta x_{ij}}$$  \hspace{1cm} (1)

$$\dot{q}_{i,\text{rad}} = \sum_{j=1}^{N} A_i V_{(i\rightarrow j)} \sigma (T_{p,i}^4 - T_{p,j}^4)$$

$$V_{(i\rightarrow j)} = \frac{A_i}{\sum_{j=1}^{N} A_j}$$  \hspace{1cm} (3)

$$A_{c,i,j} = \frac{1}{2} \left[ (R_i \tan(5^\circ))^2 + (R_i \tan(5^\circ))^2 \right]$$  \hspace{1cm} (4)

In Eq. (2) the emissivity is set to 1, therefore no reflections occur. The assumption greatly simplifies the computation and stands for all the species involved in the present study.

3.1.3. Heat and mass transport inside particles

Each particle may be composed by solid, liquid and gaseous phase. Thermal equilibrium is assumed inside the particle meaning $T_p(r)$, being $r$ the radial coordinate, is the same for all the different phases [53]. The assumption is justified by the negligible heat capacity of the gas phase compared to the liquid and solid. Hence, the heat transported through bulk movement of the gaseous species within the pore space can be neglected. Heat and mass transport within a particle are described by a set of one-dimensional/transient differential equations, as well as the state equation for the gaseous phase.

Eq. (5) expresses the mass conservation of a gaseous specie $i$. For a solid or liquid specie, the convective and diffusive terms vanish and Eq. (5) reduces to an ordinary differential equation, so no boundary conditions apply. $\omega_k$ is the production or consumption rate consequent to the chemical conversion $k$ and $M$ is the molar mass. The conservation of momentum of the gaseous phase is reported in Eq. (6) and it is obtained by applying the Darcy’s law, for which a hypothesis of laminar flow within the particle is assumed considering the small size scale: here $K_p$ identifies the so-called permeability coefficient, which characterizes the morphology of the porous space. The energy equation is based on the homogeneous model for a porous medium as described in [54] and reported in Eq. (7), where $H_k$ denotes the enthalpy of the reaction $k$. $k_{\text{eff}}$ is the effective thermal conductivity whose expression is reported in Eq. (8): here the summation is over all the solid species and $k_{\text{rad}}$ accounts for the radiation of the porous space, more details are provided in [53]. Symmetry boundary conditions are imposed to Eq. (5) and Eq. (7) at $r = 0$. The respective boundary conditions at $r = R$ are stated in Eq. (9) and Eq. (10). Here $\alpha$ and $\beta_i$ are the heat and mass transfer coefficients, respectively, with the subscript $i$ referring to the correspondent gaseous specie of Eq. (5). The analytical expressions of the transfer coefficients are detailed in Section 3.4.

$$\frac{\partial (\rho_i)}{\partial t} + \frac{1}{r^2} \frac{\partial}{\partial r} \left( r^2 \rho_i u_g \right) = \frac{1}{\varepsilon_p r^2} \frac{\partial}{\partial r} \left( r^2 D_i \frac{\partial \rho_i}{\partial r} \right) + \sum_k M_i \omega_k$$  \hspace{1cm} (5)

$$- \frac{\partial}{\partial r} \left( \varepsilon_p p_g \right) = \frac{\mu_{sp}}{k_p} u_g$$  \hspace{1cm} (6)
\[
\frac{\partial T_p}{\partial t} = \frac{1}{r^2 \rho_p c_p} \frac{\partial}{\partial r} \left( r^2 k_{eff} \frac{\partial T_p}{\partial r} \right) + \frac{1}{\rho_p c_p} \sum_k \omega_k H_k \tag{7}
\]

\[
k_{eff} = \varepsilon_k \rho_B + \sum_i \xi_i k_i + k_{rad} \tag{8}
\]

\[-D_i \frac{\partial \rho_i}{\partial r} \bigg|_{r=R_p} = \beta_{p,i} A_p (\rho_i - \rho_{f,i}) \tag{9}\]

\[-A k_{eff} \frac{\partial T_p}{\partial r} \bigg|_{r=R_p} = q_{p,cond} + q_{p,rad} + \alpha_p A_p (T_p - T_f) \tag{10}\]

3.1.4. Chemical conversion of particles

According to [55], kinetics of the chemical reactions involved in clinker forming are controlled by solid-state diffusions. In particular, the following mechanisms are identified: surface diffusion; joint diffusion; lattice diffusion. The latter is recognized as the slowest, so it is the one that effectively controls the rates of the reactions, with a temperature dependency of the diffusion coefficient of the Arrhenius-type. Further, all the reactions are assumed irreversible, coherently with [24].

Under these hypotheses, the rate of a generic \( k \)-reaction is a function of the molar concentrations of the reactants (solid, liquid or gaseous) as reported in Eq. (11) and the rate constant \( K \), whose expression is shown in Eq. (12). In Eq. (11), \( n \) denotes the order of the reaction respect to the correspondent reactant and the product sequence is over all the reactants. In Eq. (12), \( a \) is the pre-exponential factor, \( E \) is the activation energy and \( R_{gas} \) is the gas constant.

\[
\omega_k = K_k \prod_i C_i^{n_i} \tag{11}
\]

\[
K_k = a \exp \left( -\frac{E}{R_{gas} T_p} \right) \tag{12}
\]

Table 2 summarizes the reactions considered in the present study with their activation energies, pre-exponential factors and enthalpies, data retrieved from [24].

| Reaction                | Enthalpy (J/kg) | Pre-exp (1/s) | Act. En. (J/mol) |
|-------------------------|-----------------|---------------|------------------|
| CaCO\(_3\) \rightarrow CaO + CO\(_2\) | -1.66e6         | 4.55e31       | 7.81e5           |
| 2CaO + SiO\(_2\) \rightarrow C2S       | 6.03e5          | 4.11e5        | 1.93e5           |
| 3CaO + Al\(_2\)O\(_3\) \rightarrow C3A | 3.7e4           | 8.33e6        | 1.94e5           |
| 4CaO + Al\(_2\)O\(_3\) + Fe\(_2\)O\(_3\) \rightarrow C4AF | 1.09e5        | 8.33e8        | 1.85e5           |
| C2S + CaO \rightarrow C3S          | 4.48e5          | 1.33e5        | 2.56e5           |

3.2. CFD Approach for the Freeboard Region

The CFD domain covers the whole freeboard region. The air-intake geometry is shown in Fig. 1. Here the primary and secondary air enter through the sections \( A_1 \) and \( A_2 \), respectively. Coal particles are transported through the primary flow. The area of \( A_2 \) has been adjusted, according to the intake flows and the measured speed at the exhaust. The simplified geometry respect to the real design of the burner introduces an approximation on the local flow conditions, which could affect the predictions even at large scale, since coal ignition and its rate of oxidation are affected by the rate of mixing of the primary and secondary flow in the region close to the burner. However, the area of \( A_1 \) and consequently the intake speed of primary air has been adjusted in order to match the real position of the flame. Consequently, the numerical representation is expected to keep an adequate level of accuracy for the most part of the CFD domain. A no-slip condition is set at the rotating wall and the internal pressure is imposed at the exhaust. Particles interactions are included locally, according to their positions respect to the CFD grid.

A RANS method is exploited for solving the CFD field, which is turbulent in the region close to the burner. The CFD grid includes a refinement there and is consistent with a grid-independence study, more details are provided in Section 4. Grids of similar magnitude have also been found adequate in [27–30, 33].
The fuel is constituted by coal particles and their transport is determined according to the CFD velocity, using a diffusive model similar to the one discussed in [56]. The rate of oxidation of a single particle $\omega_{fuel}$ is calculated according to the combustion model from [24] and reported in Eq. (13) to (15). The heat released is included in the energy equation through a heat source in the right-hand side.

$$\omega_{fuel,p} = K_{fuel} \cdot d_o \cdot C_{O2}$$  \hspace{1cm} (13)

$$K_{fuel} = a_{fuel} \cdot \exp \left( -\frac{E_{fuel}}{R \cdot T_{gas}} \right)$$  \hspace{1cm} (14)

$$d_o = \left( \frac{3 \cdot d_{ox}}{n^2_{fuel}} \right) / \left( \frac{3 \cdot d_{ox}}{n^2_{fuel}} + 1 \right)$$  \hspace{1cm} (15)

3.3. FDM Approach for the Rotating Wall

The heat equation is numerically solved on the rotating wall with an FDM approach. Neumann boundary conditions are imposed at both sides, in particular: the heat exchange coefficient due to the forced convection with the environment is set to 30 W/m²K, value taken from [33] and consistent with experimental studies [57]; the emissivity of the external shell is set to 0.8, value taken from [33]; the convective heat exchange coefficient with the CFD domain is calculated from the Prandtl number using the correlations for flows through pipes provided in [58]; the heat exchange with particles is included through Eq. (1) and Eq. (2), for which the wall is assumed as a particle with infinite radius.

3.4. Particles-gas mass, momentum and convective heat transfer

Particles interactions are accounted where they are located, using an unresolved approach [59]. Consequently, the CFD grid has a spatial resolution bigger respect to the characteristic size of the individual channels between the particles. The CFD porosity $\epsilon_f$ is calculated for each computation cell as the ratio between the void and total volume. The coupling terms $\dot{m}_{f,i}$, $f_{drag}$ and $\dot{q}_f$ are included in the right-hand sides of the Navier-Stokes equations. These sources/sinks account for the species, momentum and convective heat transfer with the particles, respectively.

The expression of $f_{drag}$ is reported in Eq. (16). Here $K_{bed}$ and $C_{bed}$ are calculated according to the characteristics of the packed bed, their expressions are retrieved from [54] and reported in Eq. (17) and Eq. (18). The drag force exerted on a $i$-particle is calculated according to Eq. (19), being $N$ the total number of particles inside the computation cell and $V_{cell}$ the volume of the cell.

$$f_{drag} = \frac{2 \cdot \mu_f}{K_{bed} \cdot \epsilon_f} u_f - C_{bed} \cdot \epsilon_f^2 \rho_f |u_f||u_f|$$  \hspace{1cm} (16)

$$K_{bed} = \frac{2 \cdot \rho_f \cdot \epsilon_f^2}{150(1-\epsilon_f)^2}$$  \hspace{1cm} (17)

$$C_{bed} = \frac{1.75(1-\epsilon_f)}{2 \cdot \rho_f \cdot \epsilon_f}$$  \hspace{1cm} (18)

$$F_{drag,i} = \frac{f_{drag} \cdot V_{cell}}{N}$$  \hspace{1cm} (19)

The sources/sinks $\dot{q}_f$ and $\dot{m}_{f,i}$ are calculated according to Eq. (20) and Eq. (21), respectively, where the summation is over the total number $N$ of particles inside the cell. The transfer coefficients are obtained from the Nusselt, Reynolds and Sherwood numbers of the fluid domain, as indicated in Eq. (22) and Eq. (23) and with the subscript $i$ referring to a generic gaseous specie. The expression of the Nusselt number $Nu$ is reported in Eq. (24) and has been resumed from [60]. The particle Reynolds number $Re_p$ and the hydraulic Reynolds number $Re_h$ are expressed by Eq. (25) and Eq. (26), respectively: the first uses the particle diameter as the characteristic length scale, whilst the second accounts for the packing fraction of the particles bed, also. Substitution of Nusselt by Sherwood number, Prandtl by Schmidt number and the thermal conductivity by the gas diffusion coefficient allows determining the Sherwood number accordingly.

$$\dot{q}_f = \sum_{i=1}^{N} \left( \frac{A_i}{V_{cell}} \alpha(T_f - T_i) \right)$$  \hspace{1cm} (20)

$$\dot{m}_{f,i} = \sum_{i=1}^{N} \left( \frac{A_i}{V_{cell}} \beta_i (\rho_{f,i} - \rho_{f,i}) \right)$$  \hspace{1cm} (21)
\[ \alpha = \frac{Nu}{z_{2p}} \] \hspace{1cm} (22)
\[ \beta_i = \frac{Sh_i}{z_{2p}} \] \hspace{1cm} (23)

\[ Nu = \left[ \left(1.18 \cdot Re_p^{0.58}\right)^4 + \left(0.23 \cdot Re_h^{0.75}\right)^4 \right]^{1/4} \] \hspace{1cm} (24)

\[ Re_p = \frac{\rho_f \cdot u_f \cdot \mu_p}{\rho_p} \] \hspace{1cm} (25)
\[ Re_h = \frac{Re_p}{1-\epsilon_f} \] \hspace{1cm} (26)

Fluid-to-particles heat and mass transfers are accounted through the boundary conditions stated in Eq. (9) and Eq. (10). Particles sharing their volume with different computation cells are handled according to the algorithm described in [61].

3.5. Particles-gas-wall radiative heat transfer

Radiative heat transfers are included through the resistance network shown in Fig. 3. Here the radiating temperatures for gas and wall are taken as the mean values on the \( yz \)-plane. The radiating temperature of the material bed is the mean temperature of the particles in the active layer, which are identified using their velocities. The emissivity values are set to 0.2 for the gas and 1 for the wall and particles, values taken from [24]. The emitting surfaces are \( A_3 + A_4, A_3 \) and \( A_4 \) for the gas, the wall and the material bed, respectively. Gas-wall and gas-material transfer occur through the areas \( A_3 \) and \( A_4 \), respectively, and the correspondent view factors are calculated accordingly. Analogously, the exposed wall and particles exchange radiative heat through the gas (whose absorbance is equal to the emissivity) with a view factor of 1. The net gas- and wall-particles transfer are split between all the particles that are located in the active layer and included into the right-hand sides of Eq. (9) and Eq. (10).

Fig. 3. Resistance network for radiative heat transfers.

Initial and boundary conditions of the model are summarized in Table 3, for sake of clarity. The full list of assumptions is reported in Table 4.
Table 3. Initial and boundary conditions.

| Domain | Variable | IC               |
|--------|----------|------------------|
| DEM    | Temperature | Uniform 1220 K   |
| DEM    | x-velocity  | Uniform 0.045 m/s |
| DEM    | y,z-velocity| Uniform 0 m/s    |
| CFD    | Temperature | Uniform 950 K    |
| CFD    | x-velocity  | Uniform -12 m/s  |
| CFD    | y,z-velocity| Uniform 0 m/s    |

| Domain | Variable | Boundary          | BC                      |
|--------|----------|-------------------|-------------------------|
| DEM    | Temperature | Periodic boundaries | Adiabatic                |
| DEM/Wall | Temperature  | Powder-Immersed wall | Cond. heat transfer     |
| DEM/Wall | Temperature  | Powder-Exposed wall  | Rad. heat transfer      |
| DEM/CFD | Temperature  | Gas-Exposed wall   | Conv./Rad. heat transfer|
| Wall   | Temperature  | External wall      | Conv. heat transfer     |
| CFD    | Temperature  | Intake $A_1$       | 273 K                   |
| CFD    | Temperature  | Intake $A_2$       | 950 K                   |
| CFD    | Pressure    | Exhaust            | 1.01e5-200 Pa           |
| CFD    | Velocity    | Intake $A_1$       | 25 m/s                  |
| CFD    | Velocity    | Intake $A_2$       | 12 m/s                  |
| CFD    | Velocity    | Wall               | No slip                 |

Table 4. Assumptions of the model.

| Domain | Assumption                                    |
|--------|-----------------------------------------------|
| DEM    | Emissivity set to 1                          |
| DEM    | Thermal equilibrium inside particles          |
| DEM    | Arrhenius-type reaction constants             |
| DEM    | Irreversible reactions                        |
| DEM    | Adiabatic BCs at periodic boundaries          |
| CFD    | Simplified geometry of intake                 |
| Powder/Wall/CFD | Simplified resistance network for rad. heat transfers |

4. Numerical Strategy

The DEM domain ideally and uniformly moves along the x-direction at a constant speed of 0.045 m/s, which provides the expected residence time. The quasi-2D DEM domain is converted into 3D by shifting the thin slice along the x-direction at consecutive times. The coupling between the three domains is obtained with the following iteration scheme, which has been taken from [33]: 1) initialization, i.e. DEM run with no heat, mass and momentum transfer; 2) step 1, i.e. CFD-wall computation until steady state, by using last DEM solution; step 2, i.e. DEM computation by using the last CFD-wall solution; 3) iteration of steps 1 and 2 until convergence, which is verified by using the mean temperature of the wall $T_{w,\text{mean}}$ at the internal nodes as done in [33]. More in detail, a default of ten iterations are performed initially; subsequently, at each $n$-iteration an error is computed as follows:

$$\text{error}(n) = 100 \cdot \frac{\text{mean}[T_{w,\text{mean}}(n-10:n)] - \text{mean}[T_{w,\text{mean}}(n-10:n)]}{\text{mean}[T_{w,\text{mean}}(n-10:n)]}$$  \hspace{1cm} (27)$$

being $\text{mean}[T_{w,\text{mean}}(n-10:n)]$ mean of $T_{w,\text{mean}}$ over the last ten iterations; convergence is assumed once error goes below a predefined threshold. DEM computation is the main bottleneck for computing speedup, due to the time step imposed for the explicit contact detection. Having reduced the whole powder region to a quasi-2D domain is effective in decreasing the time-to-solution to the length scale of hours.
Further improvements can be attained by assuming that particles flow is not affected by their thermochemical conversion: this is likely to be true for the present case, considering that the negligible LOI provides conservation of mass and moment of inertia for each particle. Under this assumption, one flow simulation could be exploited for running multiple thermochemical analyses, each one requiring less than one hour on a multi-core workstation. The time requirement is taken as acceptable for quasi-real-time monitoring, considering the large-scale dynamics of the process.

5. Error Analysis

According to [62], errors and uncertainties are introduced in the various steps of a computational simulation i.e.: 1) conceptual modeling of the physical system; 2) mathematical modeling of the conceptual model; 3) discretization and algorithm selection for the mathematical model; 4) computer programming of the discrete model; 5) numerical solution of the computer program model; 6) representation of the numerical solution. These errors sum up and affect the final results, which therefore must be considered in view of the present considerations.

To the first two steps it can be referred the usage of a lumped resistance network for representing the radiative heat transfers. The simplification entails an error which is mostly due to a lower emitting temperature of the gas respect to the actual value with a maximum delta of about 80 degrees and a consequent estimated error of 10% on the assessment of the heat fluxes. This error overcomes all others therefore can be assumed as the limit of accuracy of the results.

DEM and CFD integration schemes belong to the third step. The first is explicit and based on the forward Euler method therefore only conditionally stable. The error is proportional to the integration step, which is chosen small enough to properly resolve the mechanical contacts between particles as detailed in Section 5.1. Considering the small time-scale of the collisions respect to the thermochemical conversions, the numerical stability for integration of equations 5, 7 and 11 is automatically satisfied e the correspondent errors are negligible. Stability and error for particle dynamics are discussed in Section 5.1.

CFD integration is implicit and based on the Crank-Nicholson method. The discretization error is quantified through grid independence study with the local Courant number near the burner progressively reduced from 0.5 to 0.1 and a consequent variation of the maximum predicted temperature of 2%.

The iteration scheme belongs to steps 4 and 5. Threshold for the convergence criteria is set at 2%, based on a compromise between accuracy and the time requirement for computation.

6. Results

6.1. Particles Flow

![Particles velocities](image-url)
A prediction of the velocity distribution is shown in Fig. 4. Here the rolling mode of the material bed is evident. Indeed, as typical to this kind of operation, two zones can be identified: the passive region, where the particles move almost as a rigid body following the rotation of the wall, with the speed that linearly decreases towards the center; the active layer on the top, where particles slip over the free surface of the material bed. As a consequence of such movement, a continuous recirculation of the external particles takes place, whereas the internal ones are trapped in the bulk zone. The thickness of the active layer can be computed from the velocity profile along the radial direction of the kiln section and its trend as a function of the $\xi$-coordinate is reported in Fig. 5. The model is consistent with others from literature like the one discussed in [15] (input data reported in Table 5) whose prediction for the active layer is shown in Fig. 5, as well: as it can be seen, the latter provides a maximum thickness of the active layer of 0.16 m and an average speed of the particles inside the active layer of 0.9 m/s and the values are matched quite accurately by the present simulation (0.19 m and 0.8 m/s, respectively). The slight under-estimation of speed from XDEM can be explained with a higher energy dissipation due to having reduced the Young’s modulus, which makes the viscous component of the contact force more relevant respect to the elastic one. The overestimation of the thickness is the consequence of the lower speed and conservation of mass. Indeed, DEM methods show a potential instability with hard materials because the usage of an explicit contact model could result in particles explosions if the contact is not properly resolved in time. In [63], the maximum time step for the computation stability is found to be $\Delta t_{\text{min}} = 0.14 \cdot \sqrt{m_p / K_H}$, being $K_H$ the Hertz stiffness. Assuming clinker particles, the formula entails a time step of 1e-4 s, which is largely smaller respect to the time scale of clinker manufacturing, i.e. the retention time. Therefore, the Young’s modulus of clinker lumps has been reduced of a factor of 10 and the time step consequently increased to 1e-3 s.

![Graph showing comparison between DEM prediction and analytical model][1]

**Table 5. Inputs for analytical model [15].**

| Parameter               | Value  |
|-------------------------|--------|
| Froude number           | 0.02   |
| Drum diameter           | 2.1 m  |
| Particles diameter      | 6 cm   |
| Filling angle           | 57 deg |
| Dynamic contact angle   | 29 deg |
6.2. Particles Heating and Conversion

While passing inside the kiln, the material undergoes a continuous heating. Using larger particles respect to the real ones (diameter of clinker lumps is around 0.01 m) is not expected to introduce relevant errors. Indeed, a total porosity \( \varepsilon_t = \varepsilon_f + \varepsilon_p \left( 1 - \varepsilon_f \right) \) as been used in Eq. (8). Particles in direct contact with the wall are heated by conduction, whilst those located on the top surface get heat by convection and radiation from the CFD domain and by radiation from the exposed wall. Adiabatic conditions are imposed at the periodic boundaries, the assumption is justified by the low thermal conductivity of the material bed.

Three different times are considered, which correspond to the positions of the DEM domain along the kiln shown in Fig. 1: 1 min after the material enters the kiln; 10 min, which is the time instant when the material is at the middle length of the kiln; 20 minutes, i.e. when the material exits the kiln.

Figure 6 shows the temperature distributions of the particles, on the \( yz \)-plane. At 1 min the temperature is still close to the initial value (1220 K). At 10 min the material reaches a mean temperature of 1423 K. At 20 min the mean temperature is equal to 1687 K. The velocity distribution affects the temperature of the particles: those on the top layer of the material bed, which are directly exposed to the gas and the upper wall, show higher temperatures; the bulk particles are at a lower temperature, because they get heat only by conduction from the outer particles. The standard deviation continuously increases from 0.96 K at 1 min to 5.1 K at 20 min and its trend along the kiln can be well appreciated in Fig. 7, together with a comparison against measurements.

![1 min temperature distribution](image1)

![10 min temperature distribution](image2)

![20 min temperature distribution](image3)

Fig. 6. Temperature of particles at different times and histograms of the respective distributions.
Temperature of the particles drives their chemical conversion, until final clinker is obtained. Figure 8 shows the mass fractions of CaO on the $yz$-plane. CaO decomposition is accelerated in the outer particles, which are at a higher temperature respect to the others. Whilst at 1 min the chemical composition is almost uniform, at 10 min a slight gradient inside the material bed is evident. Similarly, at 20 min some unreacted CaO is still present inside the bulk particles. The mass fractions of C2S and C3S show specular distributions, with higher rates of conversion in the outer particles. Mean values at 20 min (0.15 for C2S and 0.60 for C3S) are in agreement with the composition reported from the production plant.

6.3. CFD Domain and Wall

Figure 9 shows the numerical predictions of the CFD domain. The $x$-temperature trends i.e. the values averaged on the $yz$-plane can be appreciated in Fig. 7. Some considerations follow:

1. The simplified geometry of the burner helps reducing the computational burden but could affect the numerical prediction of the flame. However, the predicted temperature (2200 K) is consistent with the real data, as well its position, which starts from about 10 m downstream respect to the intake. The predicted time for complete coal burnout is around 1.5 s and compatible with the real
experience. Therefore, the error is eventually limited to the local region near the burner and the rest of the CFD representation is expected to be accurate enough.

2. The hot gas exchanges heat with the countercurrent material and the rotating wall. As a result, its temperature decreases towards the exhaust and uniforms through the \(yz\)-plane at a mean value of 1500 K. The same happens for the \(x\)-velocity, with a mean value at the exhaust of about 15 m/s.

3. It is worth to mention the inversion of the heat fluxes at the exit of the kiln, with the temperature of the material exceeding that of the wall and in turn of the gas. The same effect is found in [33].

\begin{align*}
\text{\(x\)-velocity [m/s]} & \\
0.0 & 23.0
\end{align*}

\begin{align*}
\text{Fuel concentration [-]} & \\
0.0 & 0.1
\end{align*}

\begin{align*}
\text{Temperature [K]} & \\
313 & 2280
\end{align*}

Fig. 9. Predictions of the CFD domain.

Figure 10 shows the porosity and the local convective heat transfer coefficient between the CFD domain and the particles. The first is uniformly equal to 1 except where the particles are located. An analogous distribution can be observed for the second, with values different from zero only in the CFD cells that contain at least one particle.
7. Conclusions

In the present contribution, the XDEM method was applied to predict the thermochemical phenomena occurring in a rotary kiln for clinker production. The integration of the DEM approach for the particles with a CFD for the fluid domain allowed an accurate representation of the conductive and radiative heat exchanges between the particles and their chemical conversion, as well as the heat transfers between the fluid domain, the particles and the rotating wall. The motion of the particles was predicted, finding that the velocity distribution in a rolling mode-operated kiln affects the propagation of heat inside the material bed, producing a distribution of properties in the final product.

The thermochemical results were validated by using real data from the production plant. A comparison with literature data demonstrated the effectiveness of the model in providing an accurate representation of the advective mass transport phenomena, therefore of the local heat redistribution within the material bed. In general, the consistency of the predictions respect to the benchmark data proved the effectiveness of an integrated CFD-DEM method in simulating thermal processes involving gases and powder materials.

The time-effective computation methodology could allow the exploitation of XDEM as the theoretical basis in quasi-real-time applications like indirect monitoring and model-based control.

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The results presented in this paper were obtained using the High Performance Computing (HPC) facilities of the University of Luxembourg [64] (see http://hpc.uni.lu).
**Nomenclature**

Unless stated otherwise.

| Letters/Greek letters | Letters/Greek letters |
|-----------------------|-----------------------|
| A | Surface area |
| a | Pre-exp. factor |
| C | Mass concentration |
| c | Specific heat |
| D | Diffusion coefficient |
| H | Enthalpy of reaction |
| h | Enthalpy state variable |
| k | Thermal conductivity |
| p | Pressure |
| R | Radius |
| T | Temperature |
| u | Velocity |
| V | Volume |
| α | Dynamic viscosity |
| μ | Mass density |
| ε | Porosity |
| ε | Emissivity |
| ω | Rate of reaction |

**Subscripts**

| f | Fluid CFD |
| f | Fuel |
| g | Gas particle |
| ox | Oxygen |
| p | Particles |

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