Fuel reactor modelling in Chemical-Looping Combustion of coal:

1. Model formulation

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Abstract
A fundamental part of the reliability of the Chemical-Looping Combustion system when a solid fuel, such as coal, is fed to the reactor is based on the behaviour of the fuel reactor, which determines the conversion of the solid fuel. The objective of this work is to develop a model describing the fuel reactor in the Chemical–Looping Combustion with coal (CLCC) process. The model is used to simulate the performance of the 1 MWth CLCC rig built in the Technology University of Darmstadt. The fuel reactor is a fluidized bed working at high velocity regime, using ilmenite as oxygen carrier. The developed model is based on semi-empirical correlations, and considers the reactor fluid dynamics, the coal conversion and the reaction of the oxygen carrier with evolved gases from coal. The efficiency of a carbon separation system is also considered in order to analyze this parameter on the fuel reactor performance.
The main outputs of the model are presented in this work, i.e. (1) the fluid dynamics structure of the reactor; (2) the axial profiles of gas composition and flows (volatiles, CO, H₂, CO₂ and H₂O); (3) the conversion of the oxygen carrier and char in the reactor; (4) the char concentration in the reactor; (5) the gas composition and solids flow in the upper reactor exit; and (6) the char flow to the air reactor. From these outputs the oxygen demand of the flue gases and the CO₂ capture efficiency are calculated. Simulations on the effect of the efficiency of the carbon separation system are presented. A highly efficient carbon separation system should be used to reach a high carbon capture value. Also incomplete combustion of gases is predicted in the fuel reactor, mainly from unconverted volatile matter. The model can be later used to obtain basic design parameters of the fuel reactor and optimize its operation.

Keywords: Combustion, Fluidization, Mathematical modelling, Simulation, Chemical-Looping Combustion, Coal.
1. Introduction

Chemical-Looping Combustion (CLC) is one of the most promising technologies to carry out the CO₂ capture at a low cost. CLC is based on the transfer of the oxygen from air to the fuel by means of a solid oxygen carrier that circulates between two interconnected fluidized beds: the fuel and the air reactors. In the fuel reactor the oxygen carrier is reduced through oxidation of the fuel. The oxygen carrier is regenerated in the air reactor as the inlet air flow reacts with the solids. The stream of combustion gases from the fuel reactor contains primarily CO₂ and H₂O. Water can be easily separated by condensation and a highly concentrated stream of CO₂ ready for sequestration is achieved. The CO₂ capture is inherent to this process, as the air does not get mixed with the fuel, and no additional costs or energy penalties for gas separation are required.

In the last years increasing interest is found about the application of CLC using coal as fuel, regarding the intensive use of this fuel (Adánez et al., 2012). Important progress has been made in continuous operation of CLC with coal (Adánez, 2012). Among them, different iron based oxygen carriers were used in several CLC systems, being the fuel reactor designed as a bubbling fluidized bed (Berguerand and Lyngfelt, 2008a, 2008b and 2009; Cuadrat et al., 2011, 2012a and 2012b), spouted bed (Gu et al., 2011) or high velocity fluidized bed (Markström et al., 2012). In all these cases, gasification of coal is an intermediate step happening in the fuel reactor. In these studies, complete combustion of gases was not reached in the fuel reactor, making necessary to take additional actions, as to add a final gas polishing step with pure oxygen. Often the “oxygen demand” parameter, $\Omega_T$, is used to describe the fraction of oxygen required in the oxygen polishing step. Besides, high carbon capture can be accomplished by using high temperatures and/or implementing
a carbon separation system between the fuel and air reactors. The carbon separation system
is used to separate char particles from oxygen carrier and to return them to the fuel reactor,
diminishing the bypass of char particles into the air reactor.

A fundamental part of the reliability of a Chemical–Looping Combustion of Coal (CLCC)
system is based on the behaviour of the fuel reactor. This will determine the unburnt gas
losses obtained at the exit of the reactor and the remaining char in the stream of solids
going out from the fuel reactor. The modelling of the reactor would be helpful for the
design, optimization, and scale–up of the process, in order to obtain high coal conversion in
the fuel reactor of a CLCC system.

Some works have been presented in the literature for the modelling of the process involved
in the fuel reactor of a CLCC system. These models can be included in three general
groups: (1) simple models that does not considers the complex fluid dynamic existing in a
fluidized bed (Cuadrat et al., 2012c; Kramp et al., 2012; Ströhle et al., 2009 and 2010); (2)
macroscopic models based on empirical correlations for the fluid dynamic of a fluidized
bed (Brown et al., 2010; Schöny et al., 2011); and (3) multiphase computational fluid
dynamic (CFD) models (Mahalatkar et al., 2011).

Simple models consider a highly simplified description of the fuel reactor fluid dynamic,
but they do not include the complex distribution of gas and solids in a fluidized bed. Thus,
Cuadrat et al. (2012c) predicted the carbon capture and oxygen demand reached in a
bubbling fluidized bed with continuous feeding of coal and oxygen carrier. The difficulty
for gas in bubbles to reach the particles in the emulsion phase was considered with a
parameter for the contact efficiency between volatile matter and oxygen carrier particles.
Kramp et al. (2012) presented the effect on the carbon capture by using a carbon stripper as
carbon separation system. In this work, the fuel reactor was considered in perfect mixing
for solids and gas, and complete combustion of gases evolved in the fuel reactor was assumed. A high-velocity fluidized bed was modelled as fuel reactor by Ströhle et al. (2009 and 2010). Both dense phase and lean phase were modelled as perfectly stirred reactors, the second one after the first one.

In other cases, more elaborated models have been used. A two phase model was used by Brown et al. (2010) to predict the carbon conversion in a batch fluidized bed with continuous feeding of coal. The work done by Mahalatkar et al. (2011) presented simulation and validation of the CFD model on a batch fluidized bed reactor. Schöny et al. (2011) built an empirical model which considers the complex solids and gas flow in a high-velocity fluidized bed. Schöny et al. (2011) focused their results in the combustion of gases evolved in the fuel reactor, and point out the relevance of the coal particle size on the gas conversion.

Nevertheless, model considering the complex fluid dynamics of a fluidized bed being able to predict the behaviour regarding the carbon capture and combustion efficiency of a continuously operated fuel reactor with solid fuels is not found in the literature.

In this work, a mathematical model for the fuel reactor of the 1 MWth CLCC pilot rig erected at TU Darmstadt is developed (Galloy et al., 2011). This reactor is a fluidized bed working at high velocity regime. The process carried out in the reactor is the coal combustion using ilmenite as oxygen carrier. The model considers the complex fluid dynamics of a circulating fluidized bed, the coal conversion, i.e. coal pyrolysis and char gasification, and the rate of the oxygen carrier reduction by gaseous compounds. The model is able to predict the carbon capture and combustion efficiency of the CLCC system.
2. Fuel reactor model

The model developed is focused on the fuel reactor behaviour, being a fluidized–bed reactor working at the high–velocity regime. The reactor is based on the design of a 1 MWth CLCC unit at TU Darmstadt (Abdulally et al., 2012). A diagram of this reactor is presented in Fig. 1. The coal stream is physically mixed with the oxygen carrier in the fuel reactor and the oxygen carrier reacts with the gas products from coal pyrolysis and gasification. The stream of combustion gases from the fuel reactor contains primarily CO2 and H2O, together with some unburnt compounds (CO, H2, hydrocarbons –CₓHᵧ–). At the high–velocity regime, a fraction of the particles having reached the exit duct height follows the gas stream towards the two step cyclone system, where solid particles are separated. The cyclone system is composed by: (1) a low-efficiency cyclone where mostly oxygen carrier particles are separated; and (2) a high-efficiency cyclone which separates char particles not recovered by the low-efficiency cyclone, which are recirculated to the fuel reactor. The solids stream separated by the low-efficiency cyclone consists of oxygen carrier mixed with a small fraction of char. It is desirable that the solids flow recovered by the cyclone was higher than the solids circulation flow rate to the air reactor. Thus, a fraction of the solids exiting the cyclone are externally recirculated to the bottom part of the reactor. The other fraction of solids is introduced to a carbon stripper unit in order to improve the carbon separation before oxygen carrier entering to the air reactor. Carbon separated in the carbon stripper is sent back to the fuel reactor. Thus, the cyclone system and the carbon stripper act together as a carbon separation system. The flow of char in the solids stream to the air reactor depends on the efficiency of separation of the carbon separation system. The processes happening in the components of the carbon separation system, i.e. cyclones + carbon stripper, are not modelled in this work,
but the whole efficiency of char removal is considered in the model to simulate the
performance of the fuel reactor.

In this work a macroscopic model is developed, which is based on empirical and semi-
empirical expressions. The model considers the fluid dynamics of the fuel reactor in the
high-velocity fluidized bed regime, as well as the mixing of gas and solid particles in the
reactor. The mixing of gas and solid particles influences the performance of a fluidized–bed
reactor; a high mixing rate contributes to an effective distribution of reactants, whereas an
insufficient mixing can lead to uncompleted reaction. Therefore, an adequate understanding
of the mixing behaviour is important to predict the performance of the reactor.

The modelling of the high–velocity fluidized–bed reactor is divided in two fields: fluid
dynamics and mass balances. At this point, it is remarkable that the gas velocity, $u_g$, is not
constant with the reactor height. The gas velocity increases by the gas entering together
with the char from the carbon stripper, the gases from the pyrolysis of coal, the gas
generated during char gasification, the gas expansion during the conversion of
hydrocarbons in the volatile matter and the gas expansion due to the pressure drop in the
reactor. This affects the fluid dynamics of the fluidized bed. As the increase in gas velocity
depends on the reaction rate of gases with the oxygen carrier and char, fluid dynamics and
mass balances in the reactor must be solved simultaneously. In mass balances the kinetics
of char gasification and oxygen carrier reaction with gaseous products, i.e. $\text{H}_2$, $\text{CO}$ and
$\text{CH}_4$, are included.

2.1. Fluid dynamics of the fuel reactor

The fuel reactor is considered to be in the high–velocity fluidisation regime. The model is
based on the fluid dynamics modelling presented by Pallarès and Johnsson (2006), which
considers the gas and solids flows inside the reactor and the gas–solids mixing patterns in
the different regions in which it can be divided. This model was based on studies especially
devoted to large scale circulating fluidized beds, and gives a simple and complete model of
the circulating loop fluid dynamics. This model was successfully used to predict the
behaviour of a 12 MWth CFB unit (Adánez et al., 2003).

In the model, the reactor is divided into two vertical zones with respect to axial
concentration and backmixing of solids, see Fig. 2: 1) a bottom bed with a high and roughly
constant solids concentration; and 2) a freeboard above the bottom bed, where there is a
pronounced decay in solids concentration with height. The model can be considered as 1.5
dimensional, with the main dimension the axial direction, but takes into account lateral
exchange of solids in the freeboard between the core and an annulus close to the reactor
wall. Moreover, gas distribution and mixing between the emulsion and bubble in the
bottom bed is considered. Thus, the gas flow in the bottom bed is shared between the
emulsion and bubble phases, with gas mixing between them. Solids are in the emulsion
phase, where gas flow maintains the minimum fluidizing conditions; whereas the rest of gas
goes through bubbles, where there are not solids.

The freeboard is composed by the splash or cluster phase and a transport or dispersed
phase. Both the splash and transport phases are superimposed but with different mixing
behaviour. The splash phase has a strong solids backmixing with solids in the bottom bed.
The transport phase is characterized by a core/annulus flow structure. A net flow up of
solids goes through the core, \( F_c \), and solids backmixing occurs at the reactor walls, \( F_w \).

The presence of three fluid dynamically different zones implies that they should be
investigated separately with respect to gas mixing.
The hypotheses considered for the fluid dynamic model are: (1) steady state; (2) isothermal bed at macroscopic level; (3) perfect mixing of the solids in the bottom bed and splash phase; (4) plug flow of gas in all zones of the reactor but lateral exchange of gas between bubbles and emulsion in the bottom bed; (5) gas stagnant in the annulus is considered, that is, the gas flows only through the core; (6) plug flow up of solids through the core in the transport phase with lateral flow of solids from the core to the annulus; and (7) no existence of particle fragmentation or attrition.

2.1.1. Fluid dynamics in the bottom bed

The bottom bed is defined as the zone located at the bottom of the bed characterized by a roughly constant solids concentration. The gas flow in the bottom bed is considered by a modified two phase flow as proposed by Johnsson et al. (1991). The total gas flow is expressed in terms of gas velocity considering the whole reactor section as: (1) the flow in the particulate or emulsion phase at the minimum fluidization velocity, \((1-\delta_b)u_{mf}\); (2) the visible bubble flow, \(u_{vis}\), related to the gas in the bubbles at the rising velocity of the bubbles; and (3) the gas throughflow, \(u_{tf}\), corresponding to the excess of inlet gas over the gas in the emulsion and bubbles. The gas throughflow can be understood as a stream of gas passing through bubbles. Thus, in terms of superficial gas velocities the total gas flow, \(u_g\), is divided following the equation

\[
u_g = (1 - \delta_b)u_{mf} + u_{vis} + u_{tf}
\]

\(\delta_b\) being the volumetric fraction of bubbles in the bottom bed.

A gas exchange between bubbles \((u_{vis}\) and \(u_{tf})\) and emulsion \((u_{mf})\) is considered allowing the exchange of products and reactants between these phases. The model assumes that the
emulsion phase remains under minimum fluidization condition. Thus, the flow in excess
over the minimum gas velocity, $u_{mf}$, is shared between the visible gas in the rising bubbles,
$u_{vis}$, and the throughflow, $u_{tf}$. The minimum fluidization velocity, $u_{mf}$, is calculated using
the correlation proposed by Wen and Yu (1966)

$$\text{Re}_{mf} = \frac{u_{mf}}{\mu_g} \frac{\rho_g d_p}{\rho} = \sqrt{C_1^2 + C_2 Ar} - C_1$$  \hspace{1cm} (2)

The values for $C_1 = 27.2$ and $C_2 = 0.0408$ suggested by Grace (1986) are used. The porosity
at the minimum fluidization conditions, $\varepsilon_{mf}$, is calculated with the expression proposed by
Broadhurst and Becker (1975):

$$\varepsilon_{mf} = 0.586\phi^{-0.72} Ar^{-0.029} \left( \frac{\rho_g}{\rho_s} \right)^{0.021}$$  \hspace{1cm} (3)

The presence of bubbles gives an additional expansion of the bed compared to the
minimum fluidization conditions. For a high–velocity fluidized bed, two fluidization
regimes were considered depending on the gas velocity. At low velocities, a modified
model developed for a stationary fluidized bed can be used (Johnsson et al., 1991). The
visible bubble flow and the throughflow are written:

$$u_{vis} = \Psi \left( u_g - u_{mf} \left(1 - \delta_b\right) \right)$$  \hspace{1cm} (4)

$$u_g = (1 - \Psi) \left( u_g - u_{mf} \left(1 - \delta_b\right) \right)$$  \hspace{1cm} (5)

$\Psi$ being the ratio of the visible bubble flow, $u_{vis}$, to the total flow through the bubbles, $u_{vis} +
$$u_{tf}$, and it is calculated as

$$\Psi = f_b \left( z + 4\sqrt{A_0} \right)^{0.4}$$  \hspace{1cm} (6)

The parameter $A_0$ is the gas-distributor area per nozzle. The function $f_b$ is calculated from
the following equation:
The bubble fraction in the bottom bed is:

\[
\delta_b = \frac{u_{vis}}{u_{vis} + u_{b\infty}}
\]  

(8)

With the assumption of the minimum fluidization porosity in the emulsion phase, the single bubble velocity, \(u_{b\infty}\), is

\[
u_{b\infty} = 0.71\sqrt{gd_b}
\]  

(9)

the bubble size being calculated with the correlation by Darton et al. (1977):

\[
d_b = 0.54 \left(u_g - u_{mf}\right)^{0.4} \left(z + 4\sqrt{A_b}\right)^{0.8} g^{-0.2}
\]  

(10)

Once obtained the fraction of bubbles in the bed, \(\delta_b\), the average bed porosity at a fixed position in the bottom bed can be obtained as:

\[
\varepsilon_b = (1-\delta_b)\varepsilon_{mf} + \delta_b
\]  

(11)

At high velocities, the average bed voidage levels out. In this regime, bubbles are of an exploding type, providing a large and almost free passage of gas through the bottom bed during the time of bubble eruption. Thus, an increase in the gas velocity in the bottom bed does not produce a further increase in the voidage because the excess of gas is used to increase the throughflow. The porosity at the bottom bed is constant and equal to the saturation porosity, calculated as:

\[
\varepsilon_{b,\text{sat}} = 0.5452 + \frac{495.5}{\Delta P_0 + \frac{4.9 \times 10^{-6}}{d_p}}
\]  

(12)
At this regime, the porosity obtained using Eq. (11) is higher than $\epsilon_{b,\text{sat}}$. In this case, the porosity of the bed is taken to be equal to the saturation porosity, $\epsilon_b = \epsilon_{b,\text{sat}}$. The bubble fraction, $\delta_b$, becomes the value obtained at saturation condition, $\delta_{b,\text{sat}}$

$$\delta_b = \delta_{b,\text{sat}} = \frac{\epsilon_{b,\text{sat}} - \epsilon_{mf}}{1 - \epsilon_{mf}} \tag{13}$$

The visible bubble flow and the throughflow are obtained as:

$$u_{vis} = u_{vis,\text{sat}} = \frac{\delta_{b,\text{sat}}}{1 - \delta_{b,\text{sat}}} u_{ke} \tag{14}$$

$$u_g = u_g - u_{vis} = u_{mf} \left(1 - \delta_b\right) \tag{15}$$

and the share between $u_{vis}$ and $u_{tf}$, given by the dimensionless visible bubble flow, $\Psi_{\text{sat}}$, is:

$$\Psi = \Psi_{\text{sat}} = \frac{u_{vis}}{u_g - u_{mf} \left(1 - \delta_{b,\text{sat}}\right)} \tag{16}$$

2.1.2. Fluid dynamics in the freeboard

The freeboard starts from the upper limit of the bottom bed, and it is characterized by a decrease in the solids concentration with the reactor height. The solids suspension in the freeboard is divided into two phases, a splash phase and a transport phase, each one with different fluid dynamical behaviour. The backmixing behaviour is different in each of these zones. The splash phase is characterised by a strong backmixing of solids with the bottom bed and it is composed by clusters of solids projected from the bottom bed due to the eruption of bubbles. Instead, the transport phase is characterised by a more disperse zone. The transport phase has a core-annulus structure, where there is a net flux of solids upwards in the core, and downwards in the annulus. Thus, the backmixing in the transport phase occurs mainly at the reactor walls.
The gas flows up through the core, whereas it is assumed to be stagnant in the annulus, near the walls of the reactor. The thickness of the annulus zone increases downwards along the reactor from the exit zone, where it is considered to be zero. The thickness of the annulus zone becomes:

$$\delta_w = 0.01076 (H_r - z)$$  \hspace{1cm} (17)

The thickness of the annulus grows downwards until it reaches a saturation value, which is obtained from the following equation:

$$\delta_{w,sat} = 0.06456 d_{react}$$  \hspace{1cm} (18)

The height from the bottom of the reactor at which the saturation value is reached is:

$$z_{sat} = H_r - 6d_{react}$$  \hspace{1cm} (19)

Therefore, the cross section through which the gas is flowing, $A_c$, increases above $z_{sat}$, affecting the gas velocity in the freeboard, $u_g$.

In the model developed, the splash and transport phases start from the upper limit of the bottom bed. Both phases present a decrease in the solids concentration with the reactor height, but this decrease is stronger for the splash phase. The splash phase is dominating the solids suspension in the lower part of the freeboard and the transport phase prevails in the upper part if flow is completely developed.

The decay in solids concentration of the splash phase is given by the decay factor $a$:

$$\frac{dC_{spl}}{dz} = -a C_{spl}$$  \hspace{1cm} (20)

The decay factor, $a$, depends on the superficial gas velocity and the terminal velocity of the particles, and it is calculated as

$$a = 4 \frac{H_r}{u_g}$$  \hspace{1cm} (21)
Concerning the transport phase, the solids concentration in the core is assumed to follow

the expression:

$$\frac{dC_{tr,i}}{dz} = -K_i C_{tr,i} \quad (22)$$

the decay factor $K_i$ being calculated as:

$$K_i = \frac{0.23}{u_g - u_{i,j}} \quad (23)$$

Both decay factors $a$ and $K$ are not constant along the freeboard, because the cross section
of the core, $A_c$, and the gas velocity, $u_g$, change with the height in the reactor.

Note that the concentration of oxygen carrier and char in the transport phase was calculated
separately by Eqs. (22-23), being $C_{tr,i}$ the concentration of the solids $i$ considered, i.e.
oxygen carrier or char particles. The solids concentration in the transport phase is taken
from the flow rate of entrained solids from the bottom region, $F_{0,i}$:

$$C_{tr,i,H_b} = \frac{F_{0,i}}{A_{c,H_b} (u_g,H_b - u_{i,j})} \quad (24)$$

where the flow rate of solids $F_{0,i}$ is obtained using the equation given by de Diego et al.
(1995):

$$F_{0,i} = 131.1 \left[ \frac{A_c u_g}{\varepsilon_b \rho_g} \left( \frac{Re_{sd}}{Ar_i} \right)^{0.31} \right] \quad (25)$$

A simple correlation proposed by Haider and Levenspiel (1989) was used to calculate the
terminal velocity:

$$u_{i,j} = u_{i,j}^* \left( \frac{\rho_g^2}{\mu_g (\rho_i - \rho_g)} \right)^{1/3}$$

being $u_{i,j}^* = \left( \frac{18}{Ar_i^{2.3} + \frac{2.335 - 1.744\phi}{Ar_i^{0.6}}} \right)^{-1}$ \quad (26)
The initial condition to solve these equations is taken following a continuity expression for
the solids concentration at the upper bottom bed height, $H_b$:

$$C_{\text{spl},H_b} = C_{b,H_b} - C_{\text{tr},H_b}$$

(27)

being the total concentration in the transport phase the sum of the concentration of char and
oxygen carrier particles

$$C_{\text{tr},H_b} = C_{\text{tr,OC},H_b} + C_{\text{tr,C},H_b}$$

(28)

The total solids concentration in the freeboard was the sum of the solids in the splash and
the transport phases:

$$C_{\text{fb}} = C_{\text{spl}} + C_{\text{tr}}$$

(29)

Particles in the splash phase are being in perfect mixing with the bottom bed. Instead, the
transport or dispersed phase presents a net upward flow. Particles entrained up through the
transport region can be separated to the annulus near the reactor wall, where particles are
internally recirculated to the bottom bed, or externally sent to the cyclone after having
reached the exit duct height. From the solids concentration in the core, the upwards flow of
solids $i$, $F_{c,i}$, can be obtained as:

$$F_{c,i} = C_{w,i}A_c\left(u_g - u_{c,i}\right)$$

(30)

The transport phase is assumed to be the only solid phase contributing to the externally
recirculated flow of solids $i$, $F_{s,i}$, which can be obtained from the solid flow at the exit zone,
$$\left(F_{c,i}\right)_{H_r}$$, and the backflow ratio, $k_{b,i}$, for every solid $i$, i.e. oxygen carrier or char particles:

$$k_{b,i} = \frac{\left(F_{w,i}\right)_{H_r}}{F_{s,i}} = \frac{\left(F_{c,i}\right)_{H_r} - F_{s,i}}{F_{s,i}}$$

(31)
The total flow of solids from the fuel reactor is the sum of the flow of oxygen carrier and char.

\[ F_s = F_{s,OC} + F_{s,\text{char}} \]  

(32)

During the discretisation carried out to solve the solids profiles, the lateral diffusion of particles \( i \), \( F_{t,i} \), corresponding with the solids net flow from the core to the annulus in the node \( j \) is obtained from the difference on the solids flow upwards by the core in the node \( j \) and \( j+1 \):

\[ F_{t,i} \bigg|_j = F_{t,i} \bigg|_j - F_{t,i} \bigg|_{j+1} \]  

(33)

Thus, profiles of the solids down flow by the annulus can be estimated as:

\[ F_{w,i} \bigg|_j = F_{w,i} \bigg|_{j+1} + F_{t,i} \bigg|_j \]  

(34)

with initial value at the reactor exit, \( \left( F_{w,i} \right)_{H_r} \), obtained from Eq. (31). Thus, the solids concentration in the annulus can be obtained at every position \( z \):

\[ C_{w,i}(z) = \frac{F_{w,i}(z)}{A_y(z)u_{r,j}} \]  

(35)

Despite its importance, little information is available on the magnitude of the backflow ratio, \( k_b \) (Harris et al., 2003; Johnsson et al., 1999). Factors as the presence of internals near the exit zone, the inlet area of the cyclone, the slip velocity or the value of the solids net flow can affect this parameter. Pallarès and Johnsson (2006) reviewed the effect of the particle slip velocity on the particle entrainment probability, which in turn affects to the backflow ratio. They showed that at high particle slip velocity, the particle entrainment probability varies between 0.6 and 1. Taken an average value for the particle entrainment probability of 0.8, a value of \( k_b \) about 0.2 was calculated, which was taken as a reference value in this work.
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Finally, the mixing gas behaviour in the freeboard has been considered. This phenomenon
is taken into account by the use of a contact efficiency parameter, $\xi_{g-s}$, between gas and
solids in the freeboard, calculated by the equation proposed by Furusaki et al. (1976)

$$
\xi_{g-s}(z) = 1 - 0.75 \left( \frac{C_{frb}(z)}{C_{b}(z = H_{b})} \right)^{0.4}
$$

(36)

$C_{frb}$ being the solids concentration in the freeboard given by Eq. (29) and $C_{b}$ the solids
concentration calculated in the upper limit of the bottom bed. The contact efficiency
parameter is applied on the reaction rate calculated at each axial position into the freeboard.

2.2. Mass balances into the reactor

Mass balances for the different reacting compounds and products are developed for the
phases in the bottom bed and the freeboard. The pathway for coal conversion with the
oxygen carrier is considered to happen in two steps. First drying, pyrolysis and gasification
to produce gaseous compounds:

14. Coal (s) $\rightarrow$ H$_2$O (g) + Volatile matter (g) + Char (s)  
15. Char (s) + H$_2$O (g) $\rightarrow$ H$_2$ (g) + CO (g) + ash (s)  
16. Char (s) + CO$_2$ (g) $\rightarrow$ 2 CO (g) + ash (s)  
17. The initial moisture of coal is evaporated, and steam is generated. CO, H$_2$ and CH$_4$ are
uniquely considered as reducing gases in the mass balances (Cuadrat et al., 2012c). In the
second step, the gaseous compounds react with the oxygen carrier particles towards CO$_2$
and H$_2$O following the pathway showed by Eqs. (40)–(42).

21. CH$_4$ + 3 Me$_x$O$_{y}$ $\rightarrow$ CO + 2 H$_2$O + 3 Me$_x$O$_{y-1}$  
22. CO + Me$_x$O$_{y}$ $\rightarrow$ CO$_2$ + Me$_x$O$_{y-1}$
\[
\text{H}_2 + \text{Me}_x\text{O}_y \rightarrow \text{H}_2\text{O} + \text{Me}_x\text{O}_{y-1} \quad (42)
\]

The pathway for reaction of methane with the oxygen carrier considers that H\textsubscript{2}O is a primary product during the reaction with the metal oxide (Dewaele and Froment, 1999). Thus, the CH\textsubscript{4} conversion was considered to happen in two steps: first towards CO and H\textsubscript{2}O and later CO reacts towards CO\textsubscript{2}.

Moreover, the model can assume, or not, that the reaction products (CO, H\textsubscript{2}, CO\textsubscript{2} and H\textsubscript{2}O) reach instantaneously the water–gas shift (WGS) equilibrium –Eq. (43)– in all phases of the reactor.

\[
\text{H}_2 + \text{CO}_2 \leftrightarrow \text{H}_2\text{O} + \text{CO} \quad (43)
\]

2.2.1. Mass balances in the bottom bed

In the bottom bed, a gas exchange between bubbles (\(u_{vis}\) and \(u_{dl}\)) and emulsion (\(u_{mf}\)) is considered allowing the exchange of products and reactants between these phases by diffusive and/or bulk flow mechanism. Indeed, as the gas suffers a volumetric expansion during the char gasification and conversion of hydrocarbons, some of the gas in the emulsion must move to the bubble phase to maintain the minimum fluidization condition in the emulsion phase. Considering all the above assumptions, the mass balances were given by the following differential equations for each gas \(i\) (CO, H\textsubscript{2}, CO\textsubscript{2}, H\textsubscript{2}O and CH\textsubscript{4}) in the emulsion and bubble phases, respectively:

\[
\frac{dF_{e,i}}{dV} = \frac{d\left[(1-\delta_b)u_{mf}C_{e,i}\right]}{dz} = -(1-\delta_b)\sum (-\bar{r}_{g,i})_{OC} + \sum (-\bar{r}_{g,i})_{char} - \delta_b \bar{K}_{be} \left(C_{e,i} - C_{b,j}\right) - y_{e,j} \frac{dF_{exc}}{dV} - \frac{dF_{WGS,j}}{dV} \quad (44)
\]

\[
\frac{dF_{b,j}}{dV} = \frac{d\left[(u_{vis} + u_{gy})C_{b,j}\right]}{dz} = \delta_b \bar{K}_{be} \left(C_{e,i} - C_{b,j}\right) + y_{e,j} \frac{dF_{exc}}{dV} - \frac{dF_{WGS,j}}{dV} \quad (45)
\]
The first term in the right side on Eq. (44) represents the amount of gas $i$ consumed or generated in the differential volume element $dV$. $\sum (-\tau_{g,i})_{OC}$ is the net consumption or generation rate of the gas $i$ by reaction with the oxygen carrier in the emulsion phase following the scheme given in Eqs. (40)–(42). $\sum (-\tau_{g,i})_{char}$ is the net consumption or generation of the gas $i$ by the char gasification, given by Eqs. (38) and (39). The second term is the net gas amount of gas $i$ flowing from the emulsion to the bubble by gas diffusion. The third term refers to the excess of gas in the emulsion that moves to the bubbles because of the gas expansion produced in the differential volume. Finally, the fourth term is the amount of gas $i$ reacted to fulfil the WGS equilibrium. Similar descriptions can be done for the mass balance in the bubble phase, see Eq. (45), excepting that in this phase there is not gas–solid reaction. These equations allow determining the concentration of gas $i$ in both phases: emulsion and bubbles. The bubble–emulsion gas exchange coefficient, $k_{be}$, is obtained using the correlation given by Foka et al. (1996) for turbulent fluidized beds:

$$k_{be} = 1.631 \ u_g \ V^0.37$$  \hspace{1cm} (46)

2.2.2. Mass balances in the freeboard

In the freeboard, the mass balance for each gas in a differential element of bed yields

$$\frac{dF_{\text{di},i}}{dV} = \frac{d\left[ u_g C_{\text{di},i} \right]}{dz} = - \left[ \xi_{qg} \sum (-\tau_{g,i})_{OC} + \sum (-\tau_{g,i})_{char} \right]_{qg} - \left[ \xi_{gq} \sum (-\tau_{g,i})_{OC} + \sum (-\tau_{g,i})_{char} \right]_{qg} - \frac{dF_{\text{WGS},i}}{dV}$$  \hspace{1cm} (47)

The first and second terms in the right side on Eq. (47) represent the amount of gas $i$ consumed or generated in the differential volume element, $dV$, by reaction of gases with the
oxygen carrier and char particles in the splash phase and in the transport phase, respectively. $\xi_{g-s}$ is the contact efficiency between gas and solids in the freeboard given by Eq. (36). The last term is the amount of gas $i$ reacted to fulfil the WGS equilibrium.

Different behaviour for the solids is assumed for the splash and the transport phases. Solids into the splash phase are in perfect mixing with the bottom bed, whereas solids in the transport phase have a net flow upwards through the core. The average conversion for the splash phase corresponds to the same as solids conversion existing in the bottom bed. The solids conversion in the transport phase is a function of the axial location, and it increases as the solids flow up through the core. The variation of the solids conversion, as much as for oxygen carrier as for char particles, is calculated using the following equation:

$$\frac{dX_i}{dz} = \frac{dX_i}{dt} \frac{dz}{dt} = \frac{dX_i}{dt} \frac{1}{u_{s,i}}$$ (48)

$u_{s,c}$ being the velocity of solids in the core

$$u_{s,i} = u_g - u_{i,i}$$ (49)

2.2.3. Boundary and initial conditions

The boundary conditions to solve the above mass balances are the following:

a) At the bottom of the bed, the gas flow of each compound $i$ is the flow at the reactor inlet.

$$F_i|_{z=0} = F_{i,in}$$ (50)

b) At the feeding point of coal, $H_{coal}$, the flow of the gas $i$ in the volatile matter, $F_{i,vm}$, is instantaneously evolved to the gas phase. If the feeding point is inside the bottom bed, the volatiles flow is joined to the bubbles flow.

$$F_i|_{z=z_{coal}} = F_i + F_{i,vm}$$ (51)
c) The gas flow coming together with the recirculated char from the carbon stripper, $F_{i,CS}$, is added to the gas flow in the reactor at the height of the char feeding point, $H_{CS}$. If $H_{CS}$ is inside the bottom bed, the gas flow is joined to the bubbles flow. 

$$ F_{i}|_{z=z_{CS}} = F_i + F_{i,CS} \quad (52) $$

Moreover, the following initial conditions should be fulfilled:

a) The amount of solids in the fuel reactor should be that required to give the pressure drop initially assumed. Thus, the height of the bottom bed is calculated to fulfil the amount of solids required.

$$ \Delta P = \int_0^{H_{CS}} C_0 \rho_g dz + \int_{H_{CS}}^{H_i} C_{dia} \rho_c dz \quad (53) $$

b) The oxygen supplied by the oxygen carrier must be equal to the oxygen reacted with the reacting gases from coal pyrolysis and gasification:

$$ F_{OC} R_{OC} \Delta X_{OC} = M_O \left\{ (F_{H_2O} + 2F_{CO_2} + F_{CO})_{out} - (F_{H_2O} + 2F_{CO_2} + F_{CO})_{in} \right\} \quad (54) $$

$F_{OC}$ being the circulation flow rate of the oxygen carrier between air and fuel reactors, and $\Delta X_{OC}$ the variation of the mean conversion of solids in the reactor.

$$ \Delta X_{OC} = X_{OC,out} - X_{OC,in} \quad (55) $$

c) The carbon in the char flowing to the air reactor from the carbon stripper, $F_{C,AR}$, must be equal to the ungasified carbon from the coal in the fuel reactor.

$$ F_{C,AR} = \frac{[C]_{coal}}{M_C} \frac{F_{coal}}{F_{C,FR}} \left[ (F_{CO_2} + F_{CO} + F_{CH_4})_{out} - (F_{CO_2})_{in} \right] \quad (56) $$

The carbon flowing to the air reactor from the carbon stripper can be also calculated as a function of the separation efficiency of the carbon separation system, $\eta_{CSS}$:

$$ F_{C,AR} = (1 - \eta_{CSS}) F_{C,FR} \quad (57) $$
In this calculation, it is considered that there is a fraction of char intrinsically returned to the fuel reactor, corresponding to the excess of solids flow exiting the fuel reactor regarding the recirculated solids flow, \( F_{OC} \). Thus, \( F_{C,FR} \) depends on the char flow going to cyclone, \( F_{s,\text{char}} \), the ratio between the recirculated flow and solids flow going to the cyclone system, and the content of carbon in the reacted char, which depends on the initial carbon content in char and the char conversion.

\[
F_{C,FR} = \frac{F_{s,\text{char}} F_{OC}}{M_{C}} \frac{(1 - X_{\text{char}}) [C]_{\text{fix}}}{(1 - X_{\text{char}}) [C]_{\text{fix}} + [\text{ash}]} \tag{58}
\]

The char conversion, \( X_{\text{char}} \), can be calculated as,

\[
X_{\text{char}} = \frac{M_{C} F_{C,\text{gasif}}}{[C]_{\text{fix}} F_{\text{coal}}} = \frac{[C]_{\text{fix}} F_{\text{coal}} - M_{C} F_{\text{C,AR}}}{[C]_{\text{fix}} F_{\text{coal}}} \tag{59}
\]

\( F_{C,\text{gasif}} \) being the carbon gasified in the fuel reactor, calculated as:

\[
F_{C,\text{gasif}} = \int_{0}^{H_{c}} (1 - \delta_{b}) \left[ \left( -r_{g,\text{H}_2O} \right)_{\text{char}} + \left( -r_{g,\text{CO}_2} \right)_{\text{char}} \right]_{s} S_{\text{react}} dz - \int_{H_{c}}^{H_{r}} \left[ \left( -r_{g,\text{H}_2O} \right)_{\text{char}} + \left( -r_{g,\text{CO}_2} \right)_{\text{char}} \right]_{\text{spl}} S_{\text{react}} dz - \int_{H_{r}}^{H_{T}} \left[ \left( -r_{g,\text{H}_2O} \right)_{\text{char}} + \left( -r_{g,\text{CO}_2} \right)_{\text{char}} \right]_{s} S_{\text{react}} dz \tag{60}
\]

2.3. Inputs to the model

The present study focuses on the fuel reactor of the 1 MW\(_{th}\) CLCC test plant at TU Darmstadt (Ströhle et al., 2010; Abdulally et al., 2012). The main inputs to the model are shown in Tables 1-6. The dimensions and main process parameters of this reactor are summarized in Tables 1 and 2. The main physical and chemical properties of solids are summarized in Tables 3-6, including the kinetic parameters for reduction of ilmenite and char gasification.
The solids circulation rate is selected to operate at an oxygen carrier to fuel ratio of $\phi = 1.2$. This value was selected according to the results showed by Cuadrat et al. (2012a), where the carbon capture increased as the $\phi$ ratio decreased. The oxygen carrier to fuel ratio, $\phi$, was defined as the availability of oxygen in the flow of oxygen carrier divided by the oxygen required to fully convert the fuel to CO$_2$ and H$_2$O:

$$\phi = \frac{R_{\text{O,blank}} F_{\text{OC}}}{\Omega_{\text{coal}}}$$  \hfill (61)

So, $\phi = 1$ corresponds to the stoichiometric flow of oxygen carrier needed for a full conversion of the fuel to CO$_2$ and H$_2$O. The oxygen demanded by coal is defined as:

$$\Omega_{\text{coal}} = M_O \left( 2[C]_{\text{coal}} / M_C + 0.5[H]_{\text{coal}} / M_H - [O]_{\text{coal}} / M_O \right) m_{\text{coal}}$$  \hfill (62)

$[C]_{\text{coal}}$, $[H]_{\text{coal}}$ and $[O]_{\text{coal}}$ being the carbon, hydrogen and oxygen fractions in the fuel, whose values are obtained from the proximate analysis.

2.3.1. Oxygen carrier

Ilmenite has been widely studied as oxygen carrier in the CLCC process. Tests in units from 0.5 to 100 kW$_{\text{th}}$ showed the good performance of this material in the process (Berguerand and Lyngfelt, 2008a, 2008b and 2009; Cuadrat et al., 2011, 2012a and 2012b; Markström et al., 2012). Ilmenite showed an increase in its reactivity through redox cycles during the so-called activation process (Adánez et al., 2010). A fast activation was observed in continuous operation in a 0.5 kW$_{\text{th}}$ unit, whereas the oxygen transport capacity was maintained roughly constant (Cuadrat et al., 2011). In this work, activated ilmenite was considered as the oxygen carrier material for coal combustion. The physical properties of the activated ilmenite particles are shown in Table 3.
The kinetic parameters for the reduction of activated ilmenite with CH₄, H₂ and CO are in Table 4. These kinetic parameters were obtained in a previous work (Abad et al., 2011). The grain model with uniform reaction in the particle with changing grain size model in the grains was used to determine the reaction rate of the oxygen carrier particles. Every grain react following the shrinking core model (SCM) controlled by chemical reaction. The equations that describe the reaction rate are the following:

\[
\frac{dX_{OC,i}}{dt} = \left[ \frac{3}{\tau_i} \left(1 - X_{OC,i} \right) \right]^{2/3} \left[ \frac{R_{OC,0}}{R_{OC}} \right]
\]

\[\tau_i = \frac{\rho_{OC,i} \tau_{EC}}{b_i k_{OC,i} C_{p,i}^{n_i}} \]

(63)

\[R_{OC,0} \text{ and } R_{OC} \text{ being the oxygen transport capacity of ilmenite particles used for kinetic determination and of ilmenite particles considered in this work. Thus, } R_{OC,0} = 3.3 \text{ wt.} \% \text{ and } R_{OC} = 4.0 \text{ wt.} \% \text{ (Abad et al., 2011). The kinetic constant, } k_{OC,i}, \text{ was calculated as a function of temperature:}

\[k_{OC,i} = k_{OC,0,i} e^{-\frac{E_{OC,i}}{R_T T}} \]

(64)

The average reaction rate of every reducing gas at each height in the reactor, \((-\tau_{EC})_{OC}\), was calculated from the average reaction rate of solids following the scheme given by Eqs. (40)–(42):

\[\left(-\tau_{EC}\right)_{OC} = \frac{\left(-\tau_{EC}\right)_{i}}{2d_i} \]

(65)

\[d_i \text{ being the stoichiometric coefficient for the combustion of the reacting gas } i \text{ (CO, H₂ or CH₄) with molecular oxygen. The oxidation of CH₄ to CO and H₂O was considered in the mass balances, so } d_{CH₄} = 3/2 \text{ was used for methane. In this sense, the stoichiometric factor } b_{CH₄} \text{ used in the kinetic model (see Eq. (63)) also corresponds to the oxidation of CH₄ to}


CO and H$_2$O, i.e. $b_{CH_4} = 4.34$ (see Table 4), similarly to the modification done by Abad et al. (2010).

To obtain the average reaction rate of the oxygen carrier, $(-\bar{r}_{OC,i})$, the residence time distribution of the solids, $E(t)$, is taken into account. So, the average reaction rates of the oxygen carrier in a determined position in the reactor can be obtained as

$$(-\bar{r}_{OC,i}) = \frac{\rho_{OC} R_{OC}}{M_O} \left(1 - \varepsilon_z \right) \int_0^{t_{r,i}} \left[ \frac{d\left(X_{OC}(t) - \bar{X}_{OC,in}\right)}{dt} \right] E(t) dt$$  \hspace{1cm} (66)

Assuming a perfect mixing of the solids in the bottom bed and the splash regions, the residence time distribution curve of the solids, $E(t)$, is given by the following equation (Levenspiel, 1981):

$$E(t) = \frac{1}{t_{mr}} e^{-t/t_{mr}}$$  \hspace{1cm} (67)

where $t_{mr}$ is the mean residence time of particles in the whole reacting zone of the fluidized–bed reactor.

Eq. (66) has been expressed to consider that the oxygen carrier can be introduced into the reactor with a mean conversion of the oxygen carrier for the reduction reaction, $\bar{X}_{OC,in}$, higher than 0, i.e. particles could not have been fully oxidized in the air reactor. It is assumed that the oxidized compound in the grains is in the outer part, being the inner core the material not oxidized. Therefore, the reduction proceeds from the outer surface of the grain towards the interior of the grain following the SCM (Abad et al., 2007; García-Labiano et al., 2004). Thus, the values of $t_{r,i}$ are defined as the reacting time of an oxygen carrier particle from zero conversion until the maximum variation possible in conversion of the carrier, i.e. $(1 - \bar{X}_{OC,in})$:
To integrate Eq. (66) it is necessary to know \( X_{OC}(t) \), which can be obtained taking into account the mean conversion of the carrier at the reactor inlet and the variation of conversion after a time \( t \):

\[
X_{OC}(t) = \bar{X}_{OC,in} + \left[ 1 - \left(1 - \frac{t}{\tau_m}\right)^3 \right] \tag{69}
\]

The mean conversion of the particles, \( \bar{X}_{OC,out} \), can be obtained as

\[
(1 - \bar{X}_{OC,out}) = \int_0^{\tau_m} (1 - X_{OC}(t)) E(t) dt \tag{70}
\]

The mean reacting time, \( \tau_m \), can be determined by an iterative process to obtain from Eqs. (69) and (70) the corresponding mean conversion at the reactor outlet, \( \bar{X}_{OC,out} \), that fit the mean conversion obtained from the mass balance to the whole reactor given by Eq. (54).

Thus, a distribution of conversions is obtained.

The reaction rate of a particle with a conversion \( X_{OC} \) was calculated by using the gas concentration inside the particle, which was assumed to be constant over the entire particle, and equal to the gas concentration at the particle surface, \( C_p = C_{ps} \). The gas concentration in the particle surface, \( C_{ps} \), can be obtained by a mass balance to the whole particle taking into account the external diffusion through the gas film around the particle:

\[
\left(-\frac{r_{g,j}}{J_{OC}}\right) = \frac{D_{L,OC}}{b_i M_g} \left(\frac{4}{3} \pi r_p^3\right) \left[ \frac{d}{dt} \left( X_{OC}(t) - \bar{X}_{OC,in} \right) \right] = k_{g,j} \left(4 \pi r_p^2\right) \left(C_{z,j} - C_{ps,j}\right) \tag{71}
\]

The mass transfer coefficient for each gas, \( k_{g,i} \), is obtained with equations proposed by Palchonok (1998) and Chakraborty and Howard (1981) for the bottom bed and the freeboard, respectively.
2.3.2. Coal

In a previous work it was shown the relevance of the coal rank on the performance of the CLCC process (Cuadrat et al., 2012b). The carbon capture efficiency increased with increasing the char gasification rate or the volatile content in the coal. Besides, the oxygen demand decreased when the volatile content decreased. For simulations in this work, the bituminous “El Cerrejon” coal (Colombia), which is frequently used in power stations as fuel, is chosen. Its reactivity is between those of lignite and anthracite. The average particle size is assumed to be 125 μm, and the apparent density of char particles is 1100 kg/m³.

Table 5 shows the proximate and ultimate analysis of this coal.

When coal is fed to the fuel reactor different physical and chemical processes happen. First comes the drying and pyrolysis, where the moisture content and the volatile matter are evolved to the gaseous stream, as showed Eq. (37). Second, the remaining solid fraction, i.e. char which is mainly composed by carbon and mineral matter (ashes), is gasified. The drying and pyrolysis processes are assumed to happen instantaneously in the feeding point of the coal. Pillai (1981) showed that the devolatilization times for several coals were lower than 2 s for particle size in the order of 0.5 mm at 1010 °C. It is expected that the devolatilization time for smaller particles was lower, and it is believed that this time was lower than the mixing time of solids in the fuel reactor.
The product distribution during pyrolysis is calculated following the model described by Matthesius et al. (1987). This model assumes the formation of carbon, CH₄, C₂H₆, tar, CO, CO₂, H₂, H₂O, NH₃ and H₂S, which are calculated from the proximate and ultimate analysis of the coal. At existing conditions in CLCC, it was found that no higher hydrocarbons than CH₄ and very small amounts of tar were present in the gases (Cuadrat et al., 2011). Thus, it is assumed that all C₂H₆ and tar predicted in the volatile matter were reformed by the fluidizing gas, i.e. H₂O or CO₂, as it was previously done by Cuadrat et al. (2012c). With these considerations, Table 7 shows the products distribution after drying and devolatilization when H₂O was used as fluidization gas in the fuel reactor. The negative values for H₂O were because some H₂O was taken from the fluidization gas flow for the reforming of C₂H₆ and tar. H₂, CO and CH₄ present in the gaseous compounds are oxidized by reaction with the oxygen carrier –see Eqs. (40-42)–. NH₃ was assumed to be instantaneously converted into N₂, as this was the uniquely N-compound from the fuel reactor (Song et al., 2012). Besides, SO₂ and COS was observed to be in much higher concentration than H₂S in the fuel reactor, being SO₂ the major component (Shen et al., 2010), indicating that sulphur compounds are not at thermodynamic equilibrium. Without knowing the kinetics of oxidation of sulphur compounds by the oxygen carrier, it was assumed that SO₂ was the only sulphur compound as an initial approach.

The carbon in the char is gasified with H₂O and CO₂, Eqs. (38-39). Assuming that gasification proceeds according to the homogeneous model, the rates for the gasification with H₂O or CO₂ are calculated as:

\[
\frac{1}{m_c} \left[ \frac{d m_c}{dr} \right]_{H_2O} = \frac{1}{1 - X_{C}} \left[ \frac{d X_{C}}{dr} \right]_{H_2O} = \frac{k_{H_2O} P_{H_2O}}{1 + K_{H_2O} P_{H_2O} + K_{H_2} P_{H_2}}
\]  

(74)
The gasification proceeds by the adsorption of the reacting gas in an active site in the char surface, with an inhibitory effect of the products of reaction, i.e. H₂ for steam gasification and CO for gasification with CO₂. Thus, the gasification rate decreases with increasing the partial pressure of the product gas (H₂ or CO) because of an inhibitory effect. Kinetic parameters were obtained in a previous work (Cuadrat et al., 2012c), and they are showed in Table 6. The dependence on the temperature of the kinetic and adsorption constants, $k_i$ and $K_i$, was assumed to be Arrhenius type, similarly to Eq. (64) for kinetic constant for the oxygen carrier reduction.

The net consumption or generation of the gas $i$ by the char gasification, $(-\tau_{g,i})_{\text{char}}$ to be used in the mass balances -Eqs. (44) and (47)-, is given by

$$(-\tau_{g,i})_{\text{char}} = f_C \rho_s (1-\epsilon_z) \left[ \frac{1}{m_c} \frac{dm_c}{dt} \right]$$

where $f_C$ being the mass fraction of carbon in the bed at a determined height. It was assumed that the bed is composed by oxygen carrier and char particles. It is considered that ash particles are not re-circulated from the air reactor, and only those in unconverted char are present. This behaviour for ash particles was found by Cuadrat et al. (2011) in a 500 W<sub>th</sub> CLCC unit.

The carbon concentration in the fuel reactor, $f_C$, can be calculated in every phase $j$ (bottom bed, splash region or transport region) at every position $z$ in the reactor from the concentration of char and the char conversion as

$$f_C = \left[ \frac{C_{j,\text{char}}}{C_{j,\text{char}} + C_{j,\text{OC}}} \right] \left[ \frac{(1-X_{\text{char}})[C]_{\text{fix}}}{(1-X_{\text{char}})[C]_{\text{fix}} + [\text{ash}]} \right]$$

The gasification rate is described by the following equation:

$$\frac{1}{m_c} \left[ \frac{dm_c}{dt} \right]_{\text{CO}_2} = \frac{1}{1-X_c} \left[ \frac{dX_c}{dt} \right]_{\text{CO}_2} = \frac{k_{\text{CO}_2} P_{\text{CO}_2}}{1 + K_{\text{CO}_2} P_{\text{CO}_2} + K_{\text{CO}} P_{\text{CO}}}
$$

(75)
2.4. Outputs from the model

The main outputs of the model are (1) the fluid dynamics structure of the reactor, e.g. height of the bottom bed and profiles of concentration and flow of solids in the freeboard; (2) the axial profiles of gas composition and flows (CO, H₂, CH₄, CO₂ and H₂O); (3) the axial profile of char concentration in the reactor; (4) the axial profiles of average conversions for the oxygen carrier and char; (5) the gas composition and solids flow in the upper reactor exit to cyclone; and (6) the char flow to the air reactor. From these outputs the performance of the CLCC system was assessed by analyzing the following parameters:

a) Oxygen demand of the flue gases. It represents the extent that coal is burned to CO₂ and H₂O in the CLCC system. The total oxygen demand, Ωₜ, is the fraction of oxygen required to fully oxidize the unconverted gases exiting the fuel reactor to CO₂ and H₂O with respect the total oxygen demand of the fuel, i.e. the stoichiometric amount of oxygen required in an oxy-fuel process.

\[
Ωₜ = \frac{(F_{H_2} + F_{CO} + 4F_{CH_4})_{out}}{Ω_{coal}}
\]

To calculate the oxygen demand it is assumed that all char passing to the air reactor is fully burned by air. Thus, unconverted products only come from the fuel reactor and the oxygen demand depends on the combustion efficiency of gases generated in the fuel reactor, i.e. both volatiles and gasification products. To better evaluate the oxygen demand in the CLCC system, the combustion efficiency in the fuel reactor is defined as the stoichiometric ratio of the oxygen transferred to the fuel in the fuel reactor versus the oxygen demand of the coal converted in the fuel reactor:
The oxygen transferred to the fuel, numerator in Eq. (79), is the oxygen supplied by the oxygen carrier, and it is calculated from an oxygen balance to gaseous compounds entering to and exiting from the reactor. The oxygen demanded by gaseous compounds evolved in the fuel reactor, denominator in Eq. (79), is calculated as the oxygen demanded by coal to be fully oxidized, $\Omega_{\text{coal}}$, minus the oxygen demanded by carbon not emitted in the fuel reactor, i.e. $F_{\text{CO}_2,\text{AR}}$.

b) CO$_2$ capture efficiency. This parameter considers the physical removal of CO$_2$ that would otherwise be emitted into the atmosphere. It is defined as the fraction of the carbon introduced with coal that is converted to gas in the fuel reactor.

$$\eta_{\text{CC}} = M_c \left( F_{\text{CO}_2} + F_{\text{CO}} + F_{\text{CH}_4} \right)_{\text{out}} - \left( F_{\text{CO}_2} \right)_{\text{in}}$$  \hspace{1cm} \text{(80)}

Note that in this technology, the char that has not been gasified in the fuel reactor passes to the air reactor, it is burnt there to CO$_2$, and therefore it is not captured. The carbon capture efficiency depends on the char conversion, which was calculated in Eq. (59).

2.5. Calculation procedure

To solve the mathematical model developed with the above equations, convergence of the mass balance for solids and gases must occur simultaneously. For its solution, the model is solved using a Visual Fortran© code. The flow diagram for the overall solution of the model is shown in Fig. 3. To obtain the variation of the gases concentration with the height, the entire reactor is divided into compartments with a height of $\Delta z$. 

The algorithm for the reactor model has the following calculation flow structure:

- **Input data:** The input data are the operating conditions and the characteristics of the reactor, oxygen carrier and coal, as it is shown in Tables 1-7.

- Firstly, it is necessary to assign an initial value for the following parameters: (1) dept of the bottom bed, $H_{b,0} = H_b/2$ is assumed as first estimation; (2) average variation of the solids conversion in the reactor, which is estimated assuming full combustion and carbon capture in the CLCC system, i.e.

$$\Delta X_{OC,0} = M_0 \cdot \frac{\Omega_{coal} F_{coal}}{F_{OC} R_{OC}};$$

and (3) the carbon concentration in the bottom bed is calculated as

$$f_{C,0} = \frac{0.5[C]_{fix}}{F_{OC}} \cdot F_{coal}.$$ 

The char concentration in the bottom bed is calculated by using Eq. (77) assuming initially $X_{char} = 0.5$.

- The distribution of conversion for the oxygen carrier in the reactor is calculated from Eqs. (69) and (70) to obtain

$$\Delta X_{OC} = \overline{X}_{OC, out} - \overline{X}_{OC, in}.$$

- **Fuel conversion:** The mass balance for each reacting gas (CH₄, CO, H₂, CO₂ and H₂O) is calculated in every compartment for the emulsion and bubble phases in the bottom bed and the freeboard, see Eqs. (44), (45) and (47). The fluid dynamical characteristics of the reactor (solids concentration and gas flow) are calculated simultaneously to the mass balance. The system of differential equations for the mass balance in every compartment is solved using a finite element method over the entire reactor and starting from the distributor plate. A Runge–Kutta method is used to solve the differential equations in every element of the reactor.

- **Output data:** When the mass balance for all the species is solved, the concentration profiles of the gas species are obtained along the entire reactor, as well as the...
variation with the height of fluid dynamical properties, e.g. gas flow distribution
and solids concentration. Eventually, the gas concentration of gases and solids
conversion at the reactor exit are determined. From these results, the variation of the
solids conversion, $\Delta X_{OC,1}$, is obtained from Eq. (54). Also the carbon gasified in the
whole reactor is calculated from Eq. (60).

- Convergence of pressure drop: from the profile of solids concentration, the pressure
drop in the fuel reactor is calculated, $\Delta P_1$. If $\Delta P_1 \neq \Delta P_0$, a new value of $H_b$ is
considered.

- Convergence of oxygen balance: the variation of the solids conversion, $\Delta X_{OC,1}$,
obtained is compared to the assumed value. If $\Delta X_{OC,1} \neq \Delta X_{OC,0}$, the mass balance is
repeated using a new $\Delta X_{OC,0}$ value until reaching convergence.

- Convergence of carbon balance: from the carbon gasified in the whole reactor, the
carbon transferred to the air reactor is obtained from Eq. (56), $F_{C,AR,1}$. This value is
compared to the assumed $F_{C,AR,0}$ calculated from the assumed carbon concentration,
$f_{C,0}$, see Eqs. (57) and (58). If both values are not equal, a new value for $f_{C,0}$ is
assumed.

- Once the pressure balance and the mass balances for oxygen and carbon have
converged, the final results are obtained.

3. Model results

The developed model has been used to simulate the behaviour of the fuel reactor in the 1
MWth CLCC plant erected in TU Darmstadt (Ströhle et al., 2010). Preliminary predictions
from the model has been obtained by using a value for the efficiency of the carbon
separation system of 0.9. Besides, the water-gas shift (WGS) reaction, see Eq. (43), was not considered in the calculations as a first approximation because in previous works it was found that this reaction had low relevance in the coal conversion using ilmenite as oxygen carrier (Cuadrat et al., 2012d and 2012e). Later, simulations were done to analyze the effect of the carbon separation system efficiency, from 0 to 1, on the CLCC process performance.

3.1. Analysis of the fuel reactor performance

The reference case has been simulated, i.e. the design conditions showed in Table 1 and reference conditions in Table 2 were used. Note that the efficiency of the carbon separation system was fixed to be $\eta_{CSS} = 0.9$. The performance of the CLCC is made by analyzing two parameters: the carbon capture and the oxygen demand. Two additional parameters, i.e. the char conversion and the combustion efficiency in the fuel reactor are also used here to better understand the performance of the fuel reactor. From now on, these four parameters will be appointed as the “evaluating parameters”. In the reference case analyzed in this section, the carbon capture efficiency predicted was $\eta_{CC} = 59\%$. This low value is due to the low char conversion reached in the fuel reactor, $X_{\text{char}} = 42\%$. The oxygen demand was $\Omega_T = 10.5\%$, coming from unconverted volatile matter and also from product of char gasification in the freeboard. The predicted combustion efficiency in the fuel reactor was $\eta_{c,FR} = 83.7\%$.

To evaluate these global results of the CLCC performance, a deeper analysis is done where the gas and solids profiles, and the distribution of reaction rates are presented. Figs. 4(a) and (b) shows the axial profiles of gases and solids flow, and Figs. 4(c) and (d) shows the
axial profiles of gas and solids concentration in the fuel reactor. Also the gas velocity is plotted in Fig. 4(b).

The gas flow increases through the reactor because gases generated during coal gasification. Also, abrupt changes in gas flow are found at the positions where the coal is fed ($z = 0.1$ m) and the char separated in the carbon stripper is recirculated ($z = 0.2$ m), see Fig. 4(a). This was because the evolution of volatile matter in the first case, and the gas coming from the carbon stripper in the second case. Note that the flow from the carbon stripper is almost three times the gas flow introduced by the distributor plate. Although the gas flow is increasing throughout the entire reactor, a decrease in the gas velocity is observed in the upper part of the reactor ($z > 9$ m). This fact was because the core section only increases above the height which the saturation value in the annulus is reached, see Eqs. (17-19). As gas was assumed to flow by the core, the gas velocity decreases as the core section increases above $z_{\text{sat}} = 9$ m.

From the profiles of solids concentration, the separation of the bottom bed and the freeboard can be easily observed. The bottom bed is stretched out to a height of $H_b = 0.58$ m, being characterized by a roughly constant porosity. Above the bottom bed, the solids concentration decreases with the reactor height. Until a height of 3 m the solids concentration is dominated by the splash phase. The solids concentration in the splash phase falls out to values close to 0 at higher heights. In the upper half-reactor, the transport phase prevails and the solids concentration is low. In addition, the solids in the transport phase are enriched in char as they go up through the riser (see Fig. 4), because of the lower terminal velocity of char particles. Regarding the solids flow, which is represented in Fig. 4(b), there is a net solids flow upward through the core region in the transport phase. The flow of solids decreases with the axial position as a part of solids are being transferred from
the core to the annulus region. Besides, a change in the solids flow tendency is appreciated at a height of $z = 9$ m. This fact is linked to the decrease in the gas velocity above this position. In particular, the decay constant for the solids concentration in transport phase, $K_i$, increases as the gas velocity decreases, see Eq. (23). As consequence, both concentration and flow of solids in the transport phase decrease quicker when the gas velocity decreases with the reactor height.

To better appreciate the behaviour of the fuel reactor in the bottom bed, the profiles of gases in the emulsion, the bubbles and the average in the bottom bed are showed in detail in Figs. 5(a)-(c). At the bottom of the bed, the gas is mainly composed by H$_2$O, which is the fluidization gas. The carbon gasification and reaction of gases with the oxygen carrier proceed in the emulsion phase. So, downstream the gas is gaining on CO$_2$, which is concentrated in the emulsion phase. This means that the carbon mixed with the oxygen carrier is being gasified. The product gases (CO and H$_2$) are evolved to the emulsion phase and they are quickly oxidized to CO$_2$ and H$_2$O by the oxygen carrier. Moreover, the rate of CO$_2$ generation is faster than the rate of gas diffusion from the emulsion to the bubbles, indicated by the build up of CO$_2$ in the emulsion phase.

It can be seen a strong change on gas concentration and gas velocity at $z = 0.1$ and $z = 0.2$ m, that is at the coal and recirculated char feeding points, respectively. At the feeding point of coal, the volatile matter is evolved to the gas phase, increasing the concentration of CH$_4$, CO, H$_2$, and CO$_2$, i.e. the main components in the volatile matter (see Table 7). Volatile matter reacts slower than the products from carbon gasification, because the volatile matter is evolved to the bubble phase, and it must diffuse to the emulsion phase to react with the oxygen carrier. At the height where the recirculated char from the carbon stripper is introduced, it is observed an increase of the H$_2$O content, and consequently the other gases
are diluted. This H₂O proceeds from the gas used to fluidize the carbon stripper (steam).

After the feeding point of the recirculated char, the gases in emulsion are being enriched in steam by diffusion from the bubbles, which has a higher concentration and flow of steam. This proceeds until steam concentration is similar both in emulsion and bubble phase. Both gasification products generated in the emulsion and volatile matter diffusing from the bubbles react in the emulsion with the oxygen carrier particles. Nevertheless, CO₂ mainly proceeds from conversion of CO from carbon gasification rather than reaction of volatile matter evolved to the bubble phase, because the slow diffusion of gases among the bubble and the emulsion phases. Thus, CO₂ concentration slightly increases in the emulsion, whereas volatile matter slowly decreased its concentration in bubbles. A counter-current diffusion of CO₂ and H₂O from emulsion to bubbles also happens in this section.

Eventually, in the emulsion phase the concentration of gases level out, because it is reached a pseudo–equilibrium where the rates of generation of CO₂ and H₂O in the emulsion phase are balanced by the amount of these gases disappearing by char gasification and diffusing to the bubble phase.

When the top of the bottom bed is reached, the gas in the emulsion and bubble is mixed, which is observed when Fig. 4(c) and Figs. 5(a-c) are compared. Because the flow through the bubbles is much higher than the flow through the emulsion, the gas composition in the freeboard is dominated by the gaseous components in the bubbles. Thus, the concentration of H₂, CO, and CH₄ seen by the oxygen carrier is increased, increasing the reaction rate of the oxygen carrier. As a consequence, the H₂, CO and CH₄ concentrations decrease rapidly while the splash phase is prevailing in the freeboard. Volatile matter, which was hardly converted in the bottom bed, was mainly oxidized in the splash phase. Where the transport
phase prevails over the splash phase \((z > 3\, \text{m})\) the gas flow and composition barely changes because of the low amount of reacting solids.

Additional information about the conversion of gases in the bottom bed and freeboard can be drawn from the analysis of the oxygen demand of gases as a function of the reactor height, see Fig. 6(a). The oxygen demand slowly decreases in the bottom bed due to the slow conversion of volatile matter, which mainly flows through the bubbles. Likewise, the oxygen demand decreases in the freeboard until a height about 3 m, mainly due to the conversion of volatile matter. Later, the oxygen demand is roughly constant because of the low solids concentration and a pseudo–equilibrium is reached between CO and \(\text{H}_2\) generation by gasification and consumption by reaction with the oxygen carrier.

An analysis of the volumetric rate of gasification, \((-r_c^i)\), and oxygen transference \((-r_o^i)\) can explain the gas concentration profiles in the bottom bed and freeboard, besides the oxygen demand evolution. These reaction rates were calculated as:

In the bottom bed:

\[
(-r_c^i) = (1 - \delta_b) \left[ \left( -\overline{r}_{g,\text{H}_2\text{O}} \right)_{\text{char}} + \left( -\overline{r}_{g,\text{CO}_2} \right)_{\text{char}} \right]_e \tag{81}
\]

\[
(-r_o^i) = (1 - \delta_b) \left[ \left( -\overline{r}_{g,\text{H}_2} \right)_{\text{OC}} + \left( -\overline{r}_{g,\text{CO}} \right)_{\text{OC}} + 4 \left( -\overline{r}_{g,\text{CH}_4} \right)_{\text{OC}} \right]_e \tag{82}
\]

In the freeboard:

\[
(-r_c^i) = \left[ \left( -\overline{r}_{g,\text{H}_2\text{O}} \right)_{\text{char}} + \left( -\overline{r}_{g,\text{CO}_2} \right)_{\text{char}} \right]_{\text{ipl}} - \left[ \left( -\overline{r}_{g,\text{H}_2\text{O}} \right)_{\text{char}} + \left( -\overline{r}_{g,\text{CO}_2} \right)_{\text{char}} \right]_{\text{tr}} \tag{83}
\]

\[
(-r_o^i) = \xi_{g-s} \left[ \left( -\overline{r}_{g,\text{H}_2} \right)_{\text{OC}} + \left( -\overline{r}_{g,\text{CO}} \right)_{\text{OC}} + 4 \left( -\overline{r}_{g,\text{CH}_4} \right)_{\text{OC}} \right]_{\text{ipl}}
+ \xi_{g-s} \left[ \left( -\overline{r}_{g,\text{H}_2} \right)_{\text{OC}} + \left( -\overline{r}_{g,\text{CO}} \right)_{\text{OC}} + 4 \left( -\overline{r}_{g,\text{CH}_4} \right)_{\text{OC}} \right]_{\text{tr}} \tag{84}
\]
Fig. 6(b) shows the axial profiles of both reaction rates, as well as the ratio \( \frac{-r_o}{-r_c} \). The oxygen transference rate and the gasification rate are roughly constant in the bottom bed because solids were assumed to be homogeneously distributed. At \( z < 0.1 \text{ m} \) the ratio \( \frac{-r_o}{-r_c} \) was about 2, indicating that the entire carbon gasified is oxidized to CO\(_2\) (one atom of carbon needs two atoms of oxygen). Above coal feeding point the ratio \( \frac{-r_o}{-r_c} \) was about 3, which indicates that some oxygen transferred is used to oxidize volatile matter diffusing from bubbles; nevertheless, most of oxygen is used to oxidize gasification products generated in the emulsion.

When the gas passes from the bottom bed to the freeboard, the rate of oxygen transference increases and the rate of gasification decreases. Although the contact efficiency, \( \xi_{gs} \), in the region just above the bottom bed is low, the higher concentration of reducing gases in the freeboard regarding the bottom bed the higher oxygen transference rate. Nevertheless, the oxygen transference rate rapidly decreases as the solids concentration in the freeboard falls. The gasification rate decreases when gases enters into the freeboard, because char particles are seeing a higher concentration of H\(_2\) and CO, which inhibits the carbon gasification. The ratio \( \frac{-r_o}{-r_c} \) reaches high values at the beginning of the freeboard, and it decreases as the solids and reducing gas (H\(_2\), CO and CH\(_4\)) concentration decreases. This fact indicates the relevance in the splash phase of volatile matter oxidation. Above the splash phase, a new pseudo–equilibrium is reached where the ratio \( \frac{-r_o}{-r_c} \) approaches to a value of 2, indicating that the CO and H\(_2\) generated by gasification is balanced by its oxidation to CO\(_2\) and H\(_2\)O. Nevertheless, as the upper part of the freeboard the solids
concentration is low and the gas flow rate is high, the gas concentration barely is affected by the gas–solid reactions.

It is relevant to clarify the differences between the use of a gaseous fuel and a solid fuel.

Abad et al. (2010) modelled the combustion of CH₄ with a Cu-based oxygen carrier in a 10 kWth CLC unit. They showed that the conversion rate of methane was particularly reduced in the two-third upper part of the bottom bed. In this zone, most of unconverted gas is going through the bubble phase, and the gas conversion was mainly limited by the gas transfer between bubble and emulsion. Instead, when a solid fuel is used, an important amount of oxygen transferred is used to convert the products from carbon gasification in the bottom bed. Moreover, in the bottom bed the carbon gasification is enhanced by the reaction of H₂ and CO with the oxygen carrier to produce the gasification agents (H₂O and CO₂). In fact, the gasification rate was highest in the bottom bed. Similar to the case for gaseous fuels, the conversion of volatile matter is low in the bottom bed.

In the freeboard, the concentration of reducing gases is low when solid fuels are used, because dilution with a high flow of fluidization gas, i.e. H₂O in this case. However, the fuel conversion is faster for gaseous fuels, because the oxygen carrier reacts with a gas highly concentrated in the fuel gas. As consequence, the relative increase in the rate of oxygen transference when gas passes to freeboard with solid fuels is lower than with gaseous fuels. In addition, CO and H₂ are being continuously generated by carbon gasification, which do not react with the oxygen carrier as quick as in the bottom bed. This fact causes the accumulation of CO and H₂ in the gas, preventing the complete conversion of gases to CO₂ and H₂O. Of the above, the conversion of the gases towards CO₂ and H₂O would be lower with solid fuels than with gaseous fuels.
3.2. Efficiency of carbon separation in the carbon separation system

An external carbon separation system, e.g. a carbon stripper, has been proposed as the better solution to increase the carbon capture in the CLCC process (Kramp et al., 2012). In the reference case analyzed in this work with a carbon separation efficiency $\eta_{\text{CCS}} = 90\%$ it was obtained a carbon capture of $\eta_{\text{CC}} = 59\%$. It seems that a higher value of the carbon separation efficiency should be necessary to increase the carbon capture of the CLCC system. Indeed, the efficiency of the carbon separation system has been identified a key factor in order to reach high carbon capture in a CLCC system (Cuadrat et al., 2012c; Kramp et al., 2012; Ströhle et al., 2009). Thus, an analysis of the carbon separation system efficiency on the CLCC system behaviour is done in this section.

Figs. 7(a) shows the evaluating parameters for the design and operating conditions showed in Tables 1 and 2. The carbon separation efficiency was varied from 0 to 1. It can be seen that all the evaluating parameters increase as the carbon separation efficiency increases. As the carbon separation efficiency increases, the loss of carbon towards the air reactor decreases. Consequently, the carbon evolved to the gas phase in the fuel reactor increases, i.e. the carbon capture increases. The real effect of the carbon separation system is to modify the residence time of char particles into the fuel reactor by varying the char concentration mixed together ilmenite particles, see Fig. 7(b). Thus, an increase of the efficiency of carbon separation causes an increase of the char concentration in the bed, and therefore, an increase of the char flow to the cyclones and the carbon stripper, which should be taken into account for the performance of the carbon separation system. Also, the flow of solids that must be managed by cyclones and/or carbon stripper increases as the efficiency of the carbon separation system increases because a higher flow of solids
internally circulating between the fuel reactor and the cyclone-carbon stripper system. In addition, the gas generated in the fuel reactor increases, and a higher flow of solids is entrained to the cyclone.

On the contrary, the oxygen demand from the flue gases increases because the flow of reacting gases increases, whereas the ilmenite hold–up is maintained constant. This fact is due to the high generation of CO in a freeboard highly concentrated in char particles. Thus, the combustion efficiency in the fuel reactor was decreased as the char concentration in the bed was increased, i.e. the carbon separation system was more efficient. Indeed, when the efficiency of carbon separation was 1, the combustion efficiency would be about 75%, whereas the oxygen demand was 25%. Note that in this case as $F_{\text{CO}_2,\text{AR}} = 0$, it is fulfilled that $\Omega_T = 1 - \eta_{\text{c,FR}}$, see Eqs. (78) and (79). At this condition, the char concentration in the bottom bed was 3 wt.%, reaching 10 wt.% at the top of the fuel reactor, and the CO concentration at the exit increases up to 12.5 vol.%.

4. Discussion

From the results showed in this work, the performance of the CLCC system at the base case produced low carbon capture values ($\eta_{\text{CC}} = 59\%$) and relatively high oxygen demand ($\Omega_T = 10.5\%$). An increase of the carbon separation system efficiency increases the carbon capture, but also increases the oxygen demand. The solids hold-up in the 1 MWth high-velocity fluidized-bed reactor was 260 kg, and the mean residence time of oxygen carrier particles was 100 s. The values of carbon capture were similar to that predicted by a model of a bubbling fluidized bed, but the oxygen demand was lower (Cuadrat et al., 2012c). This fact shows the better performance of a high-velocity fluidized bed burning the gases
generated in the reactor compared to that of a bubbling fluidized bed. The residence time was much lower than the assumed by Kramp et al. (2010), and lower than the required to reach high carbon capture (Cuadrat et al., 2012c; Ströhle et al., 2009). The formulated macroscopic model consisted in semi-empirical equations which have intrinsically some uncertainties. Therefore, it would be very interesting to validate the model against experimental results available in the future, both for the fluid dynamics and the coal conversion process.

In addition, the CLCC system should be optimized in order to improve the carbon capture and to decrease the oxygen demand, e.g. increasing the temperature, the solids hold-up, or using a more reactive oxygen carrier. These issues will be addressed in the next work by using the developed model.

5. Conclusions
A model to describe the behaviour of the fuel reactor of a Chemical–Looping Coal Combustion (CLCC) process has been developed. The model considers the processes affecting the reaction of fuel with the oxygen carrier, such as reactor fluid dynamics, reactivity of the oxygen carrier and the gasification reactivity. The fuel reactor simulation is based on the erected 1 MWth CLCC unit at TU Darmstadt, which is considered to be a fluidized bed at the high velocity regime. Also the effect of a carbon separation system on the fuel reactor performance is evaluated.

Predictions of the model showed that conversion of char particles is enhanced in the bottom bed. In this region, the concentration of gasification products, i.e. H₂ and CO, are maintained at low values because they are continuously removed by reaction with the oxygen carrier, reaching a pseudo-equilibrium state. In the freeboard, there is an
accumulation of the gasification products, and the splash phase was primarily the
responsible of volatile matter conversion. Solids in the upper part of the fuel reactor are
being concentrated in char particles, and extension of chemical reactions is low because of
the low concentration of solids.

The conversion of coal is limited by gasification in the bottom bed. At the reference case
analyzed in this work, the carbon capture efficiency is 59%, being rather a low value
because the low char conversion in the reactor. The oxygen demand is 10.5%, mainly due
to unburnt volatile matter.

The CLCC system should be optimized in order to improve the carbon capture and to
decrease the oxygen demand. A key issue to reach high carbon capture is to increase the
efficiency of carbon separation system. The carbon capture efficiency is 95% if the carbon
separation efficiency was increased to a value as high as 99%.

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Nomenclature

1 \( a \) = decay factor for the solids concentration in the splash phase

2 \( A_0 \) = area of the gas-distributor per nozzle, m\(^2\) per nozzle

3 \( A_c \) = cross section area of core, m\(^2\)

4 \( A_w \) = cross section area of the annulus, m\(^2\)

5 \( Ar \) = Archimedes number

6 \( b_i \) = stoichiometric coefficient for metal oxide in combustion of gas \( i \)

7 \( C_1 \) = constant in Eq. (2)

8 \( C_2 \) = constant in Eq. (2)

9 \( C_b \) = concentration of solids in the bottom bed, kg m\(^{-3}\)

10 \( C_{b,i} \) = concentration of gas \( i \) in the bubble phase, mol m\(^{-3}\)

11 \( C_C \) = char concentration, kg of char per kg of solids

12 \( C_{dil} \) = concentration of solids in the freeboard, kg m\(^{-3}\)

13 \( C_{dil,i} \) = concentration of gas \( i \) in the freeboard, mol m\(^{-3}\)

14 \( C_{e,i} \) = concentration of gas \( i \) in the emulsion phase, mol m\(^{-3}\)

15 \( C_{p,i} \) = concentration of gas \( i \) in the particle, mol m\(^{-3}\)

16 \( C_{ps,i} \) = concentration of gas \( i \) at the particle’s surface, mol m\(^{-3}\)

17 \( C_{spl} \) = concentration of solids in the splash phase, kg m\(^{-3}\)

18 \( C_w \) = concentration of solids in the transport phase, kg m\(^{-3}\)

19 \( C_w \) = solids concentration in the annulus, kg/m\(^3\)

20 \( C_z \) = gas concentration in the bulk gas at the height \( z \), mol/m\(^3\)

21 \( d_b \) = diameter of bubble, m

22 \( d_i \) = stoichiometric coefficient for O\(_2\) in the combustion of gas \( i \)
1. \(d_p\) = average diameter of particles, m

2. \(d_{\text{react}}\) = inside diameter of the reactor, m

3. \(D_g\) = gas diffusivity, \(\text{m}^2/\text{s}\)

4. \(E(t)\) = residence time distribution curve

5. \(E_{a,i}\) = activation energy, kJ/mol

6. \(f_b\) = empirical function given by Eq. (7)

7. \(f_C\) = mass fraction of carbon in the bed (-)

8. \(f_{C,fr}\) = fraction of fixed carbon in coal

9. \(F_0\) = flow rate of solids entrained from the bottom bed, kg/s

10. \(F_{b,i}\) = flow of gas \(i\) in the bubble phase, mol/s

11. \(F_c\) = solids flow by the core, kg/s

12. \(F_{\text{coal}}\) = rate of coal feeding, kg/s

13. \(F_{C,AR}\) = flow of carbon to the air–reactor, mol/s

14. \(F_{C,FR}\) = flow of carbon exiting from the fuel–reactor, mol/s

15. \(F_{C,in}\) = flow of carbon in coal fed, mol/s

16. \(F_{\text{dil},i}\) = flow of gas \(i\) in the dilute phase, mol/s

17. \(F_{e,i}\) = flow of gas \(i\) in the emulsion phase, mol/s

18. \(F_{\text{exc}}\) = excess of flow in the emulsion over the minimum fluidization condition, mol/s

19. \(F_{g,CS}\) = gas flow from the carbon stripper, Nm\(^3\)/s

20. \(F_{g,\text{in}}\) = inlet gas flow, Nm\(^3\)/s

21. \(F_i\) = molar flow of gas \(i\), mol/s

22. \(F_{OC}\) = solids circulation rate, kg/s

23. \(F_s\) = solid flow going to cyclone, kg/s
1. $F_t = \text{solid flow from the core to the annulus, kg/s}$
2. $F_w = \text{solids flow by the wall--layer, kg/s}$
3. $F_{WGS,i} = \text{flow of gas } i \text{ due to the WGS reaction, mol/s}$
4. $g = \text{acceleration due to gravity, m}^2\text{s}^{-1}$
5. $H_b = \text{upper bottom bed height, m}$
6. $H_{\text{Coal}} = \text{height of the coal feeding, m}$
7. $H_{\text{CS}} = \text{height of the char feeding, m}$
8. $H_r = \text{height of the reactor, m}$
9. $k_{0,i} = \text{preexponential factor of the kinetic constant, mol}^{1-n}\text{m}^{3n-2}\text{s}^{-1}$
10. $k_b = \text{backflow ratio}$
11. $k_{be} = \text{mass transfer coefficient between bubble and emulsion, s}^{-1}$
12. $k_g = \text{external gas mass transfer coefficient, s}^{-1}$
13. $k_i = \text{kinetic constant, mol}^{1-n}\text{m}^{3n-2}\text{s}^{-1}$
14. $K = \text{decay factor for the solids concentration in the transport phase}$
15. $n = \text{reaction order}$
16. $m_C = \text{mass of carbon in char particles, kg}$
17. $M_C = \text{atomic mass of carbon, kg/mol}$
18. $M_O = \text{atomic mass of oxygen, kg/mol}$
19. $N_{nz} = \text{number of nozzles in the distributor plate}$
20. $P = \text{pressure at the reactor outlet, Pa}$
21. $P_i = \text{partial pressure of gas } i, \text{atm}$
22. $r_g = \text{grain radius, m}$
23. $(\bar{r}_{g,i})_{OC} = \text{average reaction rate of gas } i \text{ with the oxygen--carrier, mol m}^{-3}\text{s}^{-1}$
\[
\left(-\bar{T}_{g,i}\right)_{\text{char}} = \text{average reaction rate of gas } i \text{ with char, mol m}^{-3} \text{ s}^{-1}
\]
\[
\left(-\bar{T}_{OC,i}\right) = \text{average reaction of oxygen in the oxygen–carrier, mol m}^{-3} \text{ s}^{-1}
\]

\[r_p = \text{particle radius, m}\]

\[R_g = \text{constant of ideal gases, J mol}^{-1} \text{ K}^{-1}\]

\[R_{OC} = \text{oxygen transport capacity of the oxygen–carrier}\]

\[\text{Re} = \text{Reynolds number}\]

\[S_{\text{react}} = \text{cross section area of the reactor, m}^2\]

\[\text{Sc} = \text{Schmidt number}\]

\[\text{Sh} = \text{Sherwood number}\]

\[t = \text{time, s}\]

\[t_{\text{mr}} = \text{mean residence time of solids in the reactor, s}\]

\[t_r = \text{reacting time of solid from zero conversion until the maximum variation in solid conversion, s}\]

\[T = \text{temperature of the fuel–reactor, K}\]

\[u_{b_{\infty}} = \text{velocity of a single bubble, m s}^{-1}\]

\[u_{s,c} = \text{velocity of solids in the core, m s}^{-1}\]

\[u_g = \text{gas velocity in the reactor, m/s}\]

\[u_{\text{mf}} = \text{gas velocity at minimum fluidization condition, m/s}\]

\[u_t = \text{terminal velocity of particles, m/s}\]

\[u_{t_f} = \text{throughflow of gas in the bottom bed, m/s}\]

\[u_{\text{vis}} = \text{visible gas flow in the bubbles, m/s}\]

\[u_{\text{vis, sat}} = \text{visible gas flow in the bubbles at saturation condition, m/s}\]

\[V = \text{volume, m}^3\]
\( x \) = stoichiometric factor for carbon in the hydrocarbon \( \text{C}_x\text{H}_y \)

\( X_c \) = conversion of carbon in char

\( X_{OC} \) = conversion of the oxygen–carrier

\( \bar{X}_{OC,\text{in}} \) = average conversion of the oxygen–carrier at the fuel–reactor inlet

\( \bar{X}_{OC,\text{out}} \) = average conversion of the oxygen–carrier at the fuel–reactor outlet

\( X_s \) = solids conversion

\( y \) = stoichiometric factor for hydrogen in the hydrocarbon \( \text{C}_x\text{H}_y \)

\( y_{e,i} \) = molar fraction of gas \( i \) in the emulsion

\( z \) = height or vertical position in the reactor, m

\( z_{\text{coal}} \) = height of the feeding point of coal, m

\( z_{CS} \) = height of the feeding point of solids from the carbon stripper, m

\( z_{\text{sat}} \) = height at which the saturation value is reached, m

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Greek symbols:

\( \delta_b \) = volumetric fraction of bubbles in the bottom bed

\( \delta_{b,\text{sat}} \) = fraction of bubbles in the bottom bed at saturation condition

\( \delta_w \) = thickness of the annulus, m

\( \delta_{w,\text{sat}} \) = thickness of the annulus at saturation conditions, m

\( \Delta P_0 \) = pressure drop in the reactor, Pa

\( \Delta X_{OC} \) = variation of the conversion of oxygen–carrier in the reactor

\( \varepsilon_b \) = bed porosity

\( \varepsilon_{b,\text{sat}} \) = bed porosity at saturation condition

\( \varepsilon_z \) = porosity in the freeboard
$\varepsilon_{mf}$ = porosity at minimum fluidization condition

$\phi_i$ = sphericity of particles $i$

$\phi_{oc}$ = ratio of oxygen-carrier to fuel

$\phi_{H_2O}$ = ratio of steam to coal, kg of steam per kg of coal

$\mu_g$ = viscosity of gas, kg m$^{-1}$ s$^{-1}$

$\eta_c$ = combustion efficiency

$\eta_{C_{gas}}$ = gasification efficiency

$\eta_{CSS}$ = efficiency of the carbon separation system

$\rho_g$ = gas density, kg/m$^3$

$\rho_i$ = apparent density of solids $i$, kg/m$^3$

$\rho_m$ = molar density, mol/m$^3$

$\rho_s$ = average density of solids, kg/m$^3$

$\sum F_{O,in}$ = molar flow of oxygen atoms in the compounds entering to the fuel-reactor, mol/s

$\sum F_{O,out}$ = molar flow of oxygen atoms in the compounds exiting from the fuel-reactor, mol/s

$\xi_{g-s}$ = efficiency of contact between gas and solids in the freeboard

$\tau_i$ = time for complete solid conversion, s

$\tau_m$ = time for complete conversion of solid at the average gas concentration in the reactor, s

$\Omega_{coal}$ = oxygen demand of coal, mol of oxygen per kg of coal

$\Omega_O$ = oxygen demand of gases

$\Psi$ = ratio of the visible bubble flow to the total flow through the bubbles
$\Psi_{\text{sat}} = \text{ratio of the visible bubble flow to the total flow through the bubbles at saturation condition}$
References


