Modeling of chemical and physical processes in reactive porous bed

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Abstract

This paper contains the description of pyrolysis and flow process through porous media taking into account kinetic rate coefficient and equilibrium state of pyrolysis. Mathematical equation of pyrolysis mechanism including kinetic rate coefficient and equilibrium rate of pyrolysis, and equations like mass, momentum and energy and flow through porous zone are presented. Considered model is based on a simple geometry based on downdraft gasifier and analysed basing on CFD calculation of pyrolysis and flow of volatiles. Numerical analysis of a gas flow in an open system with mass source is carried out by means of Fluent. Calculations of the chemical and physical processes in porous bed are performed for different porosity.

1. Introduction

Biomass, which contains mainly an organic substance, is a common renewable fuel in nature. The most popular kind of biomass is wood, which is available almost everywhere in the world. Biomass is utilised in combustion, gasification and pyrolysis processes. Gas produced in gasification process can be used for power generation in gas engine or directly combusted in boiler, which could lead to minimize the pollutants emission and CO₂ reduction. These advantages generated considerable interest in the gasification of biomass. In a fixed bed gasifier biomass constitutes packed bed. The flow of volatiles and gases in a gasifier can be treated like a flow through porous media. Many investigations [1,2,3,4] turn to CFD modeling thermochemical conversion of biomass in porous media because of the possibility of indepth look in high temperature and highly polluted region and comparing it with experimental data. CFD results could give informations on chemical and physical processes in packed bed and could help to understand experimental results. Finally, numerical solution could be used to optimize biomass gasification process.

2. Biomass pyrolysis

Pyrolysis referred to biomass gasification is the fundamental chemical process which occurs in gasifier but it is still not completely comprehensive.

![Fig.1. Sketch of one-step global mechanism of biomass pyrolysis.](image)

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Taking into account that mass of biomass sample consists of mass of solid $m_s$ and gas phase $m_g$:

$$m = m_s + m_g,$$

(2.1)

where mass of solid $m_s$ consists of mass of volatiles $m_v$ and residual mass $m_r$.

The change of solid phase during pyrolysis process can be defined as function of kinetic rate coefficient $k$, period of time $\Delta t$ and difference between solid mass of sample $m_s$ and residual mass $m_r$:

$$\Delta m_s = -k(m_s - m_r)\Delta t,$$

(2.2)

where the reaction kinetic rate coefficient $k$ is usually expressed by Arrhenius law [1,4]:

$$k = k_0 e^{\left(\frac{E}{RT}\right)},$$

(2.3)

where: $E$ - energy activation [kJ/mol],

$k_0$ - constant,

$R$ - ideal gas constant [J/mol K],

$T$ - temperature [K].

According to literature kinetic rate coefficient can be also calculated by means of thermogravimetry measurements and given by [7,8]:

$$k = -\frac{\Delta m_s}{(m_s - m_r)\Delta t}.$$

(2.4)

where $m_r$ in this case is a residual mass of solid phase in certain temperature for very slow heating rate. The rate of pyrolysis can be defined as [9,10]:

$$Z = \frac{m_s}{m_{s,0}},$$

(2.5)

where: $m_s$ - mass of solid phase [kg],

$m_{s,0}$ - initial mass of solid phase [kg].

For the very low heating:

$$\frac{\partial T}{\partial t} \to 0,$$

(2.6)

pyrolysis rate $Z$ can be function of temperature only and named as equilibrium rate of pyrolysis $Z_e(T)$ [9],[10]. If the pyrolysis process is led with finite rate of heating the pyrolysis rate and equilibrium rate are different:

$$Z(T) \neq Z_e(T).$$

(2.7)

After simple calculations the change of pyrolysis rate can be expressed as a function of kinetic rate coefficient and deviation of pyrolysis rate $Z$ from equilibrium state:

$$\frac{dZ}{dt} = -k(Z - Z_e).$$

(2.8)

The relationship between pyrolysis rate $Z$ and equilibrium rate of pyrolysis $Z_e$ is presented in Fig2.
3. Gasification process

Biomass gasification consists of oxidation and reduction processes [5,6]. In the oxidation zone volatiles are burned with air to CO\textsubscript{2}, CH\textsubscript{4}, CO, H\textsubscript{2}O and H\textsubscript{2} in series of exothermic reactions. The heat obtained in exothermic reactions is used to release volatiles from biomass in pyrolysis process. In the reduction zone reactions between char and CO\textsubscript{2} and H\textsubscript{2}O take place. All chemical reactions occurring in gasification process could be of volumetric or surface type. The main homogenous and heterogeneous reactions which can be found in literature [1,6,12] are:

\begin{align}
CH\textsubscript{4} + H\textsubscript{2}O &= CO + 3H\textsubscript{2}, \\
CO + H\textsubscript{2}O &= CO\textsubscript{2} + H\textsubscript{2}, \\
H\textsubscript{2} + \frac{1}{2}O\textsubscript{2} &= H\textsubscript{2}O, \\
CH\textsubscript{4} + 2O\textsubscript{2} &= CO\textsubscript{2} + 2H\textsubscript{2}O, \\
2CO + 2H\textsubscript{2} &= CH\textsubscript{4} + CO\textsubscript{2}, \\
C + CO\textsubscript{2} &= 2CO, \\
C + O\textsubscript{2} &= CO\textsubscript{2}.
\end{align}

4. Governing equations

Physical and chemical processes in a reactive porous bed were described by the conservation laws of mass, momentum and energy for solid and gas phase and flow through porous media [1,11,14]. For the solid phase governing equations are given by:

\begin{align}
\frac{\partial \varepsilon_r \rho_s}{\partial t} &= \nabla (\varepsilon_r \rho_s \mathbf{v}_s) - S_m, \\
\frac{\partial \varepsilon_r \rho_s \mathbf{U}_s}{\partial t} + \nabla (\varepsilon_r \rho_s \mathbf{v}_s \mathbf{U}_s) &= -\alpha (T_s - T_s^e) \\
&+ \nabla (\lambda \nabla T_s) - S_g.
\end{align}

And respectively for gas phase:

\begin{align}
\frac{\partial \varepsilon_g \rho_g}{\partial t} + \nabla (\varepsilon_g \rho_g \mathbf{v}_g) &= S_m, \\
\frac{\partial \varepsilon_g \rho_g \mathbf{v}_g}{\partial t} + \nabla (\varepsilon_g \rho_g \mathbf{v}_g \mathbf{v}_g) &= -\nabla p + \nabla (\varepsilon_g \mu_g \nabla \mathbf{v}_g) + S_p, \\
\frac{\partial \varepsilon_g \rho_g \mathbf{U}_g}{\partial t} + \nabla (\varepsilon_g \rho_g \mathbf{v}_g \mathbf{U}_g) &= -p_g \nabla \mathbf{v}_g + \alpha (T_s - T_s^e) \\
&+ \nabla (\lambda \nabla T_s) + S_g.
\end{align}
where: $\varepsilon$ - porosity,

$v$ - fluid velocity [m/s],

$\rho$ - density [kg/m$^3$],

$S_m$ - mass source term [kg/m$^3$s],

$p$ - pressure [Pa],

$\mu$ - viscosity [Pa s],

$S_p$ - source term, resistance of solids to fluid flow in a porous medium,

$T$ - temperature [K],

$\lambda$ - thermal conductivity [W/mK],

$U$ - internal energy [J],

$S_U$ - heat source term [J] (due to heat release during chemical reactions),

$s$ - (lower subscript) - solid phase,

$g$ - (lower subscript) - gas phase.

To better understand processes which occur in reactive porous bed presented model was simplified to one dimensional balance equations. Assuming that the solid phase is steady $v_s=0$, so eq.(4.1),(4.2) do not contain convection term. The porosity of gas phase is defined as ratio of gases volume occupied by volatiles $V_g$ and the total volume $V$:

$$\varepsilon_g = \frac{V_g}{V}. \tag{4.6}$$

Taking into account definition of volume fraction of solid, the relationship between porosity of gas and solid phase fulfill formula

$$\varepsilon_g + \varepsilon_s = 1. \tag{4.7}$$

Considering that the rate of pyrolysis (eq.(2.5)) in following form:

$$Z = \frac{m_s}{m_{s,0}} = \frac{\varepsilon_s \rho_s}{\varepsilon_{s,0} \rho_{s,0}}, \tag{4.8}$$

where: $\rho_s$ - density of solid phase [kg/m$^3$],

$\rho_{s,0}$ - initial density of solid phase [kg/m$^3$],

the mass source in eq.(4.1) and (4.3) is given by equation:

$$\varepsilon_{s,0} \rho_{s,0} \frac{dZ}{dt} = -k \varepsilon_{s,0} \rho_{s,0} (Z - Z_s) = S_w. \tag{4.9}$$

Assuming that the pressure loss in the momentum equation (eq.(4.4)) is estimated by Darcy’s law [3], which is based on the gas averaged velocity, the source term $S_p$ can be defined:

$$S_p = \frac{\mu_g \varepsilon_g}{K} v_g. \tag{4.10}$$

where: $v_g$ - gas velocity [m/s],

$K$ - permeability [m$^2$],

$\mu_g$ - gas viscosity [Pa s].

The permeability can be also described [11] with regard to porosity $\varepsilon$ and particle diameter $D$:
transfer was described by P-1 model, which is the simplest formulation of the radiation model.

Described problems were treated as a non-stationary and resolved with usage of Ansys Fluent. For this case the mass, momentum and energy equations for gaseous phase, and mass and energy equations for solid phase were calculated. Mass source in mass balance equations (4.1) and (4.3) is dependent on temperature. Resistance of solids to fluid flow in a porous medium in momentum equation (4.4) is dependent on gas velocity and viscosity and zones permeability.

Referred to eq.(4.9) the change of pyrolysis rate depends on kinetic rate coefficient \( k \), eviation of pyrolysis rate \( Z \) and equilibrium rate \( Z_e \). The equilibrium rate of pyrolysis \( Z_e \) was set by thermogravimetric experimental results [7,8] and approximated by temperature function \( Z_e(T) \). Kinetic rate coefficient \( k(T) \) was determined basic on thermogravimetric experimental results [7], which included residual mass of solid phase \( m_s \) in certain temperature. Kinetic parameters were set at: \( k_0=0.4 \) s\(^{-3}\), \( E/R=2850 \) K. When the pyrolysis rate reached equilibrium state \( Z=Z_e \) then pyrolysis process was stopped. The initial mass fraction of volatiles (ratio of volatiles mass \( m_v \) and the total mass of solid phase \( m_s \)) was set at \( Y_e=0.5 \). The pyrolysis process was implemented by user defined function.

6. Results and discussions

Numerical analysis of pyrolysis process was calculated for two cases. In the first case (c1) porosity was set at \( \varepsilon=0.8 \) for fixed bed of biomass (porous zone 1) and \( \varepsilon=0.4 \) for fixed bed of char (porous zone 2). The diameter of biomass particle was set at \( d=8 \) cm. In the second case (c2) porosity was set at \( \varepsilon=0.5 \) for
fixed bed of biomass and $\varepsilon=0.3$ for fixed bed of char and the diameter of biomass particle was set at $d=2$ cm.

very slow due to porosity. On the left part of chamber filled by gases the temperature increased faster than in porous zones which is caused by gas heating by radiation from heating wall. A distribution of temperature for first case is illustrated in Fig.5. for the three representative points: $x1$-porous zone 1, $x2$-midpoint,$x3$ - porous zone 2 (Fig.3).

Figure 6 illustrates the rate of pyrolysis in porous zones for case1. The distribution of temperature reached a maximum value at point $x3$ and the pyrolysis rate decreased to minimum value $Z=0.05$, which represented equilibrium rate $Z_e$ for maximum temperature, in $t=950s$. In this point the process of devolatilization can be treated as finished. For points $x1$ and $x2$ temperature did not reach maximum value so the rate of pyrolysis changed insignificantly. The deviation of pyrolysis rate $Z$ from equilibrium rate $Z_e$ in point $x3$ is presented in Fig.7.

Figure 4 illustrates the temperature front of pyrolysis. The fixed bed become heated

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**Fig.4.** Contours of static temperature [K] for case1 for the particular time steps.

**Fig.5.** The temperature distribution of pyrolysis in porous zones.

**Fig.6.** The temperature distribution of pyrolysis in porous zones.
Comparison of two cases (c1, c2) of zones temperature for points x1, x2 and x3 are presented in Figs. 8-10. Process of heat propagation was slightly different in each case. The temperature distribution for case 1 was faster in each point which was caused by higher porosity than in case 2.

The differences in distribution of temperature for various porosity provide to the different rate of pyrolysis which is illustrated for point x2 in Fig.11. The pyrolysis was faster for case c1 due to lower...
heat capacity of porous bed in this case and better transport of radiation.

Fig.11. The rate of pyrolysis at point x2 for two cases of porous zones in fixed bed.

The analysis of flow through porous media showed different influence of porosity on magnitude of gas x-velocity which is shown for points x2 and x3 in Fig.12 and Fig.13. For case 1 the x-velocity was higher than for case 2 in points x2 and x3 due to higher zone porosity and (with regard to eq.(4.10) and (4.11)) higher permeability.

For each case increase of gas velocity during flow through porous zones was observed (Fig.14 and 15). It was caused by increase of gas temperature and production of additional gases in pyrolysis process.

Fig.12. Comparison of x-velocity at point x2 for two cases against the time.

Fig.13. Comparison of x-velocity at point x3 for two cases against the time.

Fig.14. Comparison of x-velocity at points x1,x2 and x3 for case1.

Fig.15. Comparison of x-velocity at points x1,x2 and x3 for case2.
7. Summary

A devolatilization model in porous medium has been developed. The pyrolysis mechanism including kinetic rate coefficient \( k \) and equilibrium state of pyrolysis \( Z_e \) were presented. Presented work showed that user defined function makes the modeling of pyrolysis process in CFD calculations possible. The change rate of pyrolysis can be described by function dependent on kinetic rate coefficient \( k \) and deviation of pyrolysis rate from equilibrium state \( Z-Z_e \).

In consequence distribution of temperature in porous zones, the temperature front of pyrolysis was generated and the rate of pyrolysis was variable in fixed bed. In this case different gas velocity was observed. Increase of gas velocity was caused by heating of porous bed and released volatiles - source terms in mass and energy equations.

In summary the numerical modeling of chemical and physical processes in reactive porous bed can be use to predict and analyse the biomass pyrolysis.

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