WOOD-CHAR GASIFICATION: EXPERIMENTS AND ANALYSIS ON SINGLE PARTICLES AND PACKED BEDS

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Introduction

Gasification has been studied with the aim of designing reactors, gasifiers, and other combustion systems. In a cocurrent gasifier [1], air and solid fuel move in the same direction, and the flame front moves in the opposite direction. Air first reacts with the solid fuel either in the heterogeneous mode (e.g., in the case of a charcoal gasifier) or with the volatiles generated from the solid fuel in the gas phase, releasing heat and helping in the propagation of a flame front into the unreacted solid aided by axial heat transfer by conduction and radiation. The hot combustion products (CO 2 and H 2O) are further reduced by the char. These endothermic reactions generate carbon monoxide and hydrogen, and the exit gas can be utilized as a gaseous fuel. Similar processes also occur during fire spread in permeable materials. A number of workers [2–4] have examined the propagation rate of a flame front against airstream through a packed bed of solids such as wood, foam, or biomass. The primary emphasis in these studies has been in predicting the flame spread through the media. Only the first process described earlier, namely, the oxidation, is of importance in predicting the flame spread rate. However, for design and operation of a gasifier, both oxidation and reduction processes are of equal importance. Hence, the present paper is aimed at studying these processes in an isolated single particle and extending the results to a bed of particles to predict the various features of an operating gasifier, namely, the flame front movement, the profiles of different species concentrations and temperature, and the exit gas composition from the gasifier. The present work is limited to charcoal gasification only. The model developed would be of use for understanding and designing biomass gasifiers.

Several designs of wood gasifiers exist [1,5-7], with modeling aspects addressed by a few [5,8,9] using overall kinetics in a packed bed. Predictions are compared with experiment results by tuning several kinetic and bed-related parameters.

The wood char reactions in CO 2–N 2 mixtures and O 2–N 2 mixtures have been studied in detail in our earlier studies [10,11]. The steam-carbon reaction has been studied by several researchers in the late 1940's and early 1950's [12–14] for extracting suitable rate expression. Kinetic expressions of varying complexity have been derived by these researchers [14–17] for steam-carbon reaction at temperatures of 1200 to 1500 K and in an environment of mixtures including CO, CO 2, and O 2. Satyanarayana and Keairns [15] have conducted experiments on char gasification using CO 2 and H 2O. They show from the results that the rate constants of the C-H 2O reaction are about 2.5-5 times faster than of the C-CO 2 reaction.

The Experiments

Single-Particle Char Experiments

The experimental setup consists of a 40-mm-diameter quartz reactor placed in a temperature-controlled furnace [11] through which measured flow of gases consisting of H 2O, CO 2, O 2, and N 2 in the desired proportions are passed, as shown in
Fig. 1. Preparation of the char samples (of 4-15 mm diameter from Ficus wood), their characterization for porosity, and the other experimental aspects including the qualitative behavior of conversion are the same as in the earlier work on C–CO_2 conversion [11]. The present experiments were conducted at 1250 and 1390 K to extract the temperature effect on the conversion rate.

Experiments with Packed Bed

For measuring the propagation rates, experiments similar to those performed by Reed and Markson [8] were conducted. A 65-mm-diameter and 300-mm high quartz reactor insulated with an observation slit 10 mm wide throughout the length was used as shown in Fig. 1. This corresponds to 1/40 scale version of the state-of-the-art wood gasifier of 275 kW (thermal) [18]. The other elements used are similar to the commercial large gasifier—a grate to hold the charge and tubing to draw the gas to the cooler through a blower into a flare. Charcoal pieces of approximately cubical, 8 mm size were used in the experiments. After initial light-up through the ignition port above the grate, the system was run at fixed flow rates, and the rate of progress of the flame front upward was measured. It was observed that the glowing zone was approximately 25 to 35 mm (3-4 particle depth) and the peak bed temperature measured was in the range of 1000-1230 K depending upon the mass flux.

The Model

The modeling is done in two parts: for the single particle and for the packed bed of particles. The model for single particle is similar to the earlier work [10] with some minor modifications for representing particles at different heights in a packed bed. The additional equations to be solved are for axial transport of heat and mass through the bed.

Single-Particle Modeling

The processes taking place during the combustion or gasification of porous carbon spheres are diffusion and convection of the species and energy in the porous medium and heterogeneous reaction between the gaseous species and the char. These are modeled using unsteady, spherically symmetric one-dimensional conservation equations for species and energy [11,10]. The representative equation for a species is

\[
\frac{\partial (\rho c \rho_i Y_i)}{\partial t} = \frac{1}{r^2} \frac{\partial}{\partial r} \left( r^2 \rho c \rho_i Y_i \right) + \omega'''
\]

(1)

where \( \rho = \rho_c (1 - \varepsilon) + \rho \varepsilon \) is the average density of porous char, \( \rho_c \) is nonporous char density, \( \varepsilon \) is the porosity of the char, and \( \omega''' \) is the volumetric reaction rate of specie \( i \) due to the heterogeneous reactions with the internal surface of the porous char and the gas-phase reactions in the pores. The reaction rates of CO_2 and O_2 with char
have been discussed and validated earlier [11, 10].

The surface reaction rate of carbon with steam is given by Blackwood and McGrory [14],

\[
\omega_{\text{C}_2\text{H}_2O} = -k_1 p_{\text{H}_2\text{O}} + k_2 p_{\text{H}_2} + k_3 p_{\text{C}_2\text{H}_2O}
\]

The water gas shift reaction,

\[
\text{CO} + \text{H}_2\text{O} \rightleftharpoons \text{CO}_2 + \text{H}_2
\]

is assumed to be in equilibrium within the pores [5,9,1]. Blackwood and McGrory [14] showed that the presence of even a small fraction of ash helps in the generation of CO\(_2\) during steam gasification by catalyzing the water-gas reaction.

Boundary conditions for the foregoing conservation equations are obtained by considering the heat and mass transfer in the gas film surrounding the char sphere [11]; these take into account free and forced convection and net mass flow out of the sphere surface. Heat radiation from the surface of the sphere is also accounted for.

The equations are solved using finite-difference procedure, and the results of profiles of species concentration, temperature, and porosity as functions of time are obtained. The results are compared with the experimentally obtained mass loss rate and temperatures at fixed locations in the sphere.

**Modeling of Particles in Packed Bed**

The single-particle model described earlier can be extended to modeling a packed bed of particles discussed in the section Experiments with Packed Bed. The bed is divided into a number of computational cells, and conservation equations for a typical particle representing each cell are solved. Two additional features in the boundary conditions need to be considered for a particle in a bed compared to the single particle, namely, the heat transfer between the particle and the surrounding particles, and the other is the properties of the bulk fluid surrounding the particles that vary continuously. These are determined by solving a set of conservation equations for the bulk gases assuming variations only with the height of the bed. With coordinates fixed to the particles, the conservation equations can be written as

\[
\frac{\partial \rho T}{\partial t} + \frac{\partial (\rho v_i T)}{\partial x_i} = \frac{\partial}{\partial x_i} \left( k \frac{\partial T}{\partial x_i} \right) + W_{\text{gas}}^{\prime\prime} + H_R
\]

where \( n \) is the number of particles per unit volume, \( \varepsilon_b \) is the bed porosity, \( m_p \) is the gasification rate of one particle, \( m' \) is the superficial mass flux of the gases of the bed, \( k \) and \( h \) are the mass and heat transfer coefficients, respectively, through the gas film surrounding the particle [11], and \( W_{\text{gas}}^{\prime\prime} \) and \( H_R \) are the gas-phase reaction rate and the heat generation rate due to gas-phase reactions, respectively.

The subscript \( s \) denotes the properties at the surface of the particle. For calculating \( \varepsilon_b \), the particles are considered to be nonporous. The internal porosity of the particles is considered separately when solving for individual particles. In the continuity equation 3, the time derivative term has been neglected.

For handling particle-to-particle heat transfer, it is assumed that radiation is the major mode of heat exchange among particles [19, 8, 9]. Conduction is likely to play only a minor role because the area of contact between particles is small in randomly packed bed, and emissivity of char particles is large (close to unity). A particle views the surrounding particles at various heights with different temperatures. To account for this, the surface of the particle is divided into strips of latitudes of width \( \delta \theta \). It is assumed that view factor \( f_j \) of the sphere with all the particles whose centers reside in the
latitudinal width $dh$ is equal to the ratio of the area of the strip to the total surface area of the sphere. All such particles are assumed to have a uniform surface temperature representing the average height at which these particles reside within the bed. Assuming further that emissivities of all the surfaces are equal, the total radiative flux falling on the sphere and the net radiation absorbed can be obtained as

$$Q = \sum_j f_j \sigma T_j^4$$  \hspace{1cm} (6)

$$H_R = A_s \alpha (Q - \sigma T_j^4)$$  \hspace{1cm} (7)

where $A_s$ is the surface area of the sphere and $\alpha$ the absorptivity (or emissivity) of the surface. To determine $T_j$, it is assumed that the particle views, on average, other particles at a center-to-center distance of a constant multiple of the particle diameter. This effective distance can be different from the average center-to-center distance of the particles in the bed because of the $T^4$ relationship for radiation and due to the nonuniformity of the surface temperature of a sphere. From the data on the effective thermal conductivity of packed beds due to radiation [19], it has been estimated as 0.65$d$.

The solution of equations for packed beds is performed using a time-split technique. The initial condition is ambient temperature for all the particles in the bed except for some cells near the bottom for which higher temperatures are assigned for ignition. In the first fractional time step, conservation equations for the porous sphere are solved (equation 1). Local conditions of the gas form the boundary conditions for the reacting porous spheres. One representative sphere is solved for each computational cell along the height of the bed. Solution of the equations for the particle gives the conditions at the surface of the sphere and the net mass flow from the sphere surface, which are used in the next fractional time step when equations 3-5 are solved.

**Choice of Parameters**

The choice of physical, thermodynamic, and transport properties is based on the mean char properties presented as follows:

$$Q_c = 41900 \text{ kg/m}^3$$

$$R_p = (t 4 0) 450 \text{ lm}$$

$$c_p = 1.25 \text{ kJ/kg K}$$

$$H_c = 32.60 \text{ MJ/kg}$$

$$K_c = 1.85 \text{ W/m K}$$

Rate parameters for $\text{C +H}_2\text{O}$ reaction are

$$k_1 = 3.7 \times 10^7 \exp(130,000/T) \text{ mol/cm}^3\text{atm}$$

$$k_2 = 35 \text{ atm}^{-1}$$

$$k_3 = 2.1 \times 10^{-3} \exp(-10.055/T) \text{ atm}^{-1}$$

$$k_4 = 91.8 \exp(-15.083/T) \text{ atm}^{-1}$$

**Fig. 2.** Variation of conversion with time—experiments (points) and predictions (line) for $d_0 = 8$ mm and $T_{\text{amb}} = 1250 \text{ K}$ and 1388 K.

**Fig. 3.** Normalized conversion time versus $d_0$—experiments and predictions for $T_{\text{amb}} = 1250$ and 1388 K.
ks = 2.5 * 10^{-8} \text{ atm}^{-1}

The porous char conductivity is 0.4–0.5 W/m K [20,21] and accounts for conduction and radiation inside the char. The thermal conductivity of the gas phase, ks, is calculated locally taking into account the presence of hydrogen. Conductivity of the mixture increases by a factor of 1.2–1.5 with the addition of 10% H_2 in the mixture [22]. The initial porosity of the wood char considered is in the range of 0.75–0.85, consistent with the present measurements as well as those of Groeneveld [5]. The initial radius of the pore is obtained from Groeneveld [5] where wood char was used for measurements. The parameters in the kinetic expression used presently are obtained from Blackwood and McGrorey [14]. The rate constant of the backward reaction, k_2, is obtained from the appropriate equilibrium constants. The emissivity in the expression for radiant heat loss is taken at 0.95. Following earlier work for the packedbed [9], bed porosity is chosen at 0.5, consistent with the randomly packed system [19]. The heat loss coefficient from the reactor to the ambient is estimated from the model reactor experiments as 6 W/m^2K.

Results and Discussion

Figure 2 shows the results of char conversion (X_c) with time for an 8-mm-diameter char particle at two different ambient temperatures. Char conversion is the ratio of the difference between the initial weight and weight at any time t to the initial weight. The parameters that influence the conversion time curve are the activation energy and the char conductivity. Activation energy affects the initial slope, whereas the conductivity affects the point at which the curve departs from the linearity. Because gas-phase conductivity is calculated depending on the local gas composition and temperature, activation energy is the only parameter that is uncertain. The suggested value of activation energy from Blackwood and McGrorey is 121 kJ/mole obtained from experiments on coconut char. Using the experimental results at two different temperatures 1250 and 1388 K, the activation energy was evaluated. The scatter in the experimental data is due to inherent structural differences in wood. Because of this feature, the predictions can be taken to be in reasonable agreement with the experimental results at an activation energy of 212 kJ/mole, matching closely with Groeneveld's [5] experimental results for wood char (217 kJ/mole). This comparison supports the choice of kinetic and transport parameters chosen for the model.

Figure 3 shows the experimental data and the prediction of burn time normalized with respect to density of char with initial char diameter for combustion.
at two different temperatures on a log-log plot. The experimental results were restricted to a particle diameter range of 4–15 mm. The results show a conversion time dependence on the diameter as $t_c \propto d^{10.2}$; at 1250 K and at 1388 K. These results indicate the departure from the diffusion-controlled $d^2$ law. The model prediction on the diameter dependence at the two temperatures compares well with the experimental data. These results further confirm the choice of kinetic and transport parameters. The conversion time–diameter correlations are summarized by

$$t_c / \rho = d_{f}^{10.00} \exp \left[ -3.26 + 15470 \left( \frac{1}{T} - \frac{1}{1273} \right) \right]$$

with $t_c$ in s, $\rho$ in kg/m$^3$, $d_0$ in mm, and $T$ in K. The indication of higher reactivity and a shift toward a diffusion-limited condition.

Figure 4 shows comparative data on the conversion time versus diameter for various reactants at 1273 K. The diameter dependence is nearly $d^2$ in case of pure oxygen and air environment, indicating the diffusion-dominated exothermic char conversion. The conversion time for CO$_2$ ambient is about 3.5 times that of H$_2$O ambient, comparing well with the results of Satyanarayana and Keairns [15]. The consumption rate of char in H$_2$O ambient is comparable or higher than that of char in air beyond a particle diameter of 8 mm. The results for CO$_2$ and H$_2$O show lower slopes than those for air, and therefore, the process is controlled by both diffusion and chemical reaction. Further, kinetic dominance increases with the reduction in particle diameter below 4 mm.

Figure 5, showing the temperature and reactant profile inside the particle, is used to examine the thermochemical behavior with different reactants. Temperature is highest for O$_2$, followed by air and CO$_2$, and lowest for H$_2$O. The reaction between oxygen and char is exothermic, and hence, char temperature is higher than ambient for oxygen and air. Consistent with the relative reactivities, pure oxygen creates higher char temperature than air. H$_2$O and CO$_2$ reactions with char are endothermic, and consequently, the temperatures within the char are below ambient. Further, the variation in temperature through the particle is highest for O$_2$ and lowest for CO$_2$. This difference is due to the diffusion-controlled process for O$_2$ and significantly reaction-controlled process in the case of CO$_2$. In the case of O$_2$ and air, the mass fraction of oxygen becomes close to zero at the surface, and in the case of H$_2$O and CO$_2$, the mass fraction at the core is a significant fraction of that at the surface. The relatively lower slope for H$_2$O compared to CO$_2$ is due to higher diffusivity of the gas reacting inside the sphere for H$_2$O. The ratio of reaction rates between surface and core demonstrates the combined effects of temperature and reactant distribution described earlier. The ratio exceeds by a factor of 10 for O$_2$ and air and is of the order of unity (2, in fact) for both H$_2$O and CO$_2$.

Calculations were made for the dependence of the reactant mass fraction (with inert being the other component) on char conversion time. These dependencies can be described by $t_c / \rho \propto X_{O_2}^{-1}$, $X_{CO_2}^{-0.65}$, and $X_{H_2O}^{-0.7}$, respectively. For high-temperature environment, the relative conversion times are described by the following relations:

$$\left( \frac{t_c}{\rho} \right)_{CO_2} = d_{f}^{10.5} \exp \left[ -1.715 + 35,300 \left( \frac{1}{T} - \frac{1}{1273} \right) \right]$$

Fig. 7. Experimental and model predictions on the propagation rate versus mass flux in a packed-bed char reactor along with peak bed temperature and CO concentration. Also shown are the measured [5] and predicted values of CO.
(filled triangle) and peak temperature (filled circle) for arrested flame propagation. Points indicate experimental data, and lines are predictions.

\[
\left( \frac{t_c}{\rho} \right)_{H_2O} = d_0^{0.1050} \\
\cdot \exp \left[ -3.26 + 15.47 \left( \frac{1}{T} - \frac{1}{1273} \right) \right] X_i^{0.5} \Theta_0^{0.25} \\
\left( \frac{t_c}{\rho} \right)_{O_2} = 0.0022\,d_0^{3}X_i^{1.0} 
\]

In the foregoing equations, \( t_c \) is in s; \( \rho \), the density of the particle, is in kg/m\(^3\); \( d_0 \), the diameter of the particle, is in mm; \( T \), the ambient temperature, is in K; and \( X_i \) is the mole fraction of species \( i \). The previous correlation is accurate to 5\% and is valid for the following range of parameters: \( d_0 > 4 \) mm, \( T = 1000-1400 \) K, \( X_i < 0.3 \).

**Propagation rate in packed bed**

The profiles of temperature and mole fractions of CO and CO\(_2\) at three different mass fluxes are shown in Fig. 6. The profiles are chosen at a time when the rate of propagation of the reaction front through the bed is constant. It can be seen that the peak temperature increases as the air mass flux increases. It is also evident that the thickness of the propagation front increases with air flux, which is consistent with the qualitative observations during the present experiments and earlier references [5]. At very low air fluxes, CO is not generated at any significant levels. At larger fluxes, the level of CO concentration in the exit gas increases.

Groeneveld [5] conducted experiments in a reactor with 0.3 m diameter with wood char bed at an air flux of 0.015 kg/m\(^2\) s and reported exit CO mole fraction of about 10\% and the peak bed temperature of 1200 K. Such high temperature is not obtained in a propagating flame front because of the reasons described in the previous paragraph. However, when air is provided through a distributor in the middle of the bed, the front does not propagate beyond this point. Since the front is arrested at this point, the front temperature increases to much higher value compared to the propagating front at the same air flux, because the amount of unburned char being heated becomes limited. Figure 7 also contains data for such a situation along with the experimental data of Groeneveld [5]. It can be seen that peak temperature and CO mole fraction are increased significantly in an arrested front compared
to the propagating front. The lone experimental data point of Groeneveld [5] shows good match with the predictions.

Conclusions

Char gasification, which forms an important part of biomass gasification, has been modeled. The reactions of char with O₂, CO₂, and H₂O have been studied independently on single particles, and a model for these has been developed and validated for different ambient temperatures, compositions, and external convection. The model has been extended to packed bed of particles, which has also been validated with the present and previous experiments. The model can be used for understanding and designing biomass gasifiers.

Nomenclature

\( A \) surface area of the particle (m²)
\( d_0 \) initial diameter (m)
\( E_{1,2} \) activation energy (J/mol)
\( H_c \) heat of combustion of carbon (J/kg)
\( k, k_c, \bar{k} \) thermal conductivity of gas, carbon, and porous char (W/m K)
\( k_1-k_4 \) rate constants
\( M_i \) molecular mass of species \( i \)
\( \dot{m} \) superficial mass flux in the bed (kg/m² s)
\( n \) number of particles per unit volume of the bed
\( p \) pressure (Pa)
\( r \) radial coordinate (m)
\( R \) universal gas constant (J/kg mol K)
\( t \) time (s)
\( T \) temperature (K)

\( u_p \) velocity of movement of the char bed (m/s)
\( x_c \) fraction of char consumed
\( x \) distance (m)
\( Y_i \) mass fraction of species \( i \)
\( q \) density of gas (kg/m³)
\( e_b \) bed porosity
\( a \) absorptivity of the surface
\( x \dot{i} \) volumetric reaction rate, equation 2 (kg/m³ s)

REFERENCES


