Experiment and Numerical Modeling of Stratified Downdraft Gasification Using Rice Husk and Wood Pellet

Kittipass Wasinarom and Jarruwat Charoensuk *

Stratified downdraft gasification using rice husks and wood pellets was carried out under different air mass flow rates using both experimental and numerical methods. The flame propagation rate was calculated from the temperature profile at different time steps and was used as the prerequisite to calculate the equivalent ratio in modeling the combustion zone. Chemical equilibrium modeling was employed to predict the temperature and composition of the sample in the combustion zone. Finite kinetic modeling was used to simulate the reduction zone. The initial temperature and composition of the reduction zone simulation were obtained from the chemical equilibrium results taken from the combustion zone. The flame propagation speed of the rice husk was found to be around five times greater than wood pellet at the same air flow rate. The peak temperature of both fuels had similar values. For all air mass flow rates, the equilibrium modeling over-estimated the peaks in comparison with the experimental tests. The kinetic model was sensitive to the input temperature at the zone inlet. The predicted temperature in the reduction zone demonstrated high kinetic activity at the top of the zone due to a high gas temperature. The predicted temperature was in agreement with the experimental test results.

Keywords: Stratified downdraft; Chemical equilibrium; Reduction zone; Gasification; Modeling

Contact information: Department of Mechanical Engineering, Faculty of Engineering, King Mongkut’s Institute of Technology Ladkrabang, Bangkok 10520, Thailand;
*Corresponding author: jarruwat.ch@kmitl.ac.th

INTRODUCTION

Thailand is an agricultural country with a distribution of residual biomass over areas of cultivation. This biomass can be used as an energy resource given that there are technologies available for economical and environmentally friendly utilization. Gasification is the thermo-chemical process of converting a biomass fuel to a combustible gas mixture. The process comprises varying degrees of oxidation, pyrolysis, and char heterogeneous reactions. It is governed by heat and the mass transfer of the specific gasification type. The reactors are categorized into fixed bed, moving bed, fluidized bed, and entrained flow. The fixed bed has a simple design, and are more appropriate for small-scale applications (Guangul et al. 2012). The combustion front propagation within packed fuel bed is the key indicator for successful operation. There are various types of the fixed bed gasification reactors, which are updraft, downdraft, and cross draft. In each case, the name indicates the flow direction of the gasification agent in relation to the fuel bed.

An updraft reactor produces a higher amount of tar in comparison to downdraft reactor because the pyrolysis gas passing through the low temperature region does not
allow tar cracking to occur. Therefore, it is widely used with low-volatility fuels. More complicated systems are found in larger-scale updraft gasification. Cerone et al. (2017) conducted the oxy-steam gasification of hydrolytic residues from biorefinery. They reported H2/CO ratios as high as 2.08. Moreover, they found that using steam as the co-gasification agent would benefit in reducing temperatures above the fuel grate, therefore alleviating slag clogging from ash melting.

Downdraft gasification provides lower tar content due to higher temperatures along the pyrolysis gas flow path, which enable the tar cracking process and also allow the water-gas shift reaction, which leads to increasing combustible syn-gas. Downdraft models feature a combustion zone for volatile combustion, and the reduction zone exists underneath it.

According to Reed and Das (1998) there are two different configurations of downdraft gasification, which are the Imbert downdraft and stratified downdraft versions. There has been research on flame propagation under downdraft conditions. Dasappa and Pual (2001) found that the critical air mass flux that provides the peak propagation rate is 0.1 kg/m² for the char bed and that the smoldering velocity will decrease with the increase of air flow rates beyond the peak value. Further increasing the air mass flux will result in flame quenching due to convection loss. Mahapatra et al. (2014) concluded that front velocity was directly correlated with bulk density of the fuel bed. However, it is difficult to specify a single parameter that influences the propagation velocity.

Combustion modeling of solid fuels, such as a biomass particle, is comprised of four sub-processes that simultaneously occur in varying degrees of kinetic rates, which are drying, devolatilization, volatiles homogeneous combustion, and char glowing or heterogeneous combustion. By employing the Arrhenius law, the estimated kinetic rates of each process can be calculated (Blasi 2000). Determining the local temperature and concentrations of related species is a pre-requisite prior to the rate calculation. In the gasification reactor, temperature and concentration distribution are governed by heat and mass transfer. The calculation procedure incorporates many inter-links of fluid dynamics, heat transfer, species transport, and the kinetic rates into the model. The solution is obtained by the iterative method.

By treating the domain as a continuum porous matrix with defined void fraction of the packed fuel bed, the heat transport equation of the packed solid fuel and the gas flow in pores space can be modelled. The calorific equations of state is comprised of Arrhenius rate of reactions, species generation from the combination of reactions, pyrolysis, and heat generation from oxidation. The domain was discretized into small finite volumes. All transport equations were repeatedly solved and updated along with the calorific equation of state of each control volume until the converged solution was obtained. This is known as conventional computational fluid dynamics (CFD) modeling. Many researchers have used CFD modeling to predict the process that occurs in the gasification reactor (Wang et al. 2014; Monteiro et al. 2017; Murugan and Sekhar 2017; Yan et al. 2018). CFD enables investigators to explore the most elaborate details of the process in different locations in the reactor.

However, the development of a robust and reliable mathematical model is required. It must be validated and calibrated with a wide range of operation. Moreover, the gasification process is a complex system with many non-linear source terms. Therefore, the CFD procedure consumes larger computational resources in comparison with other prediction methods.
The thermodynamics equilibrium model is another prediction method. The concept stems from the second law of thermodynamics. It has been widely used to evaluate the performance of gasification systems and usually developed with an assumption that the process is a lumped system. Therefore, it is widely adopted to simulate sub-zone volatile combustion in the reactor. Many researchers have used equilibrium modeling in gasification analysis (Zainal et al. 2001; Jarunghammachote and Dutta 2007; Huang and Ramaswamy 2011; Shabbar et al. 2013; Liu et al. 2016; Atnaw et al. 2017; Ferreira et al. 2019).

The reduction kinetic model was extensively used in downdraft gasification, particularly to simulate char reduction zone. The pyrolysis and combustion were assumed to be completed within the combustion zone. Because there was no oxygen in this zone, the packed fuel particle was completely transformed into char. The reduction kinetic model has been used in various research for reduction zone prediction. The predicted temperature profile and the exit gas composition were in good agreement with the experimental data (Giltrap et al. 2003). The improvement was made by using variable char reactivity factor (exponentially increasing) along the length of reduction zone (Babu and Sheth 2006). Sharma (2001) compared equilibrium modeling and finite kinetic modeling in the char reduction zone and pointed out the similar trend of the calorific values, conversion efficiency and exhaust gas temperature. Kinetic modeling was applied to finite sub-layers in the reduction zone. A number of researchers have used multi-zone modeling in their works (Gao et al. 2008; Masmoudi et al. 2014; Salem and Paul 2018). More elaborate study on the kinetic of steam gasification of various biomass chars has been performed. It was found that the char conversion rate depended on inorganic elements of the feedstock. Potassium acts as the catalyst which increases the reactivity during char conversion. On the contrary, silicon and phosphorous behave as inhibitors, which reduce the conversion rate. Dupont et al. (2016) revealed that the reactivity among different biomass chars were correlated well with the inorganic elements of potassium/(silicon+ phosphorus) ratio. They also compared the experimental result and the model prediction of biomass char conversion by using inorganic element ratio as an indicator. Prestipino et al. (2018) constructed the model prediction for different biomass char. He postulated that the Avrami-Erofeev model could capture sigmoidal behavior of some types of biomass char conversion.

There has been recent research study on the pellet used in energy plants. For instance, Bartocci et al. (2018b) made the series of experimental studies on the pyrolysis of commercial biomass pellet, followed by steam gasification of obtained charcoal pellet. The investigation focused on heat and mass balance of the integrated plant of hydrogen-rich gas production. They demonstrated that all the heat required by all different reactors can be supplied by heat released from volatile oxidation. Bartocci et al. (2018a) studied the pyrolysis of the pellet made of raw glycerol and biomass with different mixing ratio. They concluded that glycerol pellet was a good fuel to be used in Combine Heat and Power (CHP) plants.

In this paper, the interaction between flame propagation, combustion zone temperature, and reduction zone was highlighted. The flame propagation was analyzed under stratified downdraft gasification in different operating conditions with two different feedstocks, rice husk and wood pellet. Equilibrium modeling was employed to predict temperature and gas composition in the combustion zone, and kinetic reduction modeling was used to simulate the reduction zone. The analysis focused on heat and mass transfer at the propagation front and temperature distribution in the reduction zone. The predicted
temperature was considered relative to the experimental data.

EXPERIMENTAL

Methods

*Fixed bed reactor*

The experimental test rigs are shown in Fig. 1. The reactor was made of cylindrical stainless steel tubes. It had an inner diameter of 21 cm and a height of 100 cm and was covered with 5 cm of ceramic fiber to prevent major heat loss. An air distributor manifold was located on top of the reactor. It was piped with the air compressor tank via an air flow meter. Ten type-K thermocouples (T1 through T10) were placed along the axial location. There were 10 cm gaps between the thermocouples. A thermocouple (T0) was placed at the syngas exit port. The temperature was recorded by a YOKOGAWA FX100 data recording system (Yokogawa Electric Corporation, Tokyo, Japan). A metering rod used for monitoring bed height was placed on the top of the reactor.

![Schematic diagram for downdraft gasification and the reactor used for downdraft gasification](image)

*Fig. 1. Schematic diagram for downdraft gasification (a) and the reactor used for downdraft gasification (b)*

*Experimental procedure*

The feedstocks were poured into the reactor from the top. In the case of rice husk, 3 kg was poured, while 20 kg was used in the case of wood pellet. The top surface of the packed fuel was meant to reach a height of 100 cm if it was properly packed into the reactor. Air was fed into the reactor, and the burner was used for firing into the ignite port. T1 was monitored until it reached a temperature of 450 K. The volatile combustion took place and propagated at this temperature. After that, the burner was pulled out and the ignite port was closed. The air input was adjusted to satisfy specific cases of the experiment. A data logger was used to collect temperature information at intervals of 5 min. The peak temperature moved in the upward direction following the propagation
front during the test period, as shown in Fig. 3. Finally, the propagation front reached the top surface of the packed fuel bed. This was indicated by movement of the peak temperature in the downward direction to the grate below. This behavior is further explained in results and discussion section. At this moment, the air supply and data logger were stopped. The total bed movement was measured by the bed height indicator rod. The residual was collected and weighted to perform the mass loss calculation. An experimental test case was completed for a specified condition. In this work, 6 cases of the experimental were performed. The details of all conditions are provided in Table 1.

### Table 1. Experimental Details

<table>
<thead>
<tr>
<th>Feedstock</th>
<th>Initial mass (kg)</th>
<th>Air flow rate (L/min)</th>
<th>Period of experiment (min)</th>
<th>Calculated equivalent ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rice husk</td>
<td>3</td>
<td>20</td>
<td>165</td>
<td>3.17</td>
</tr>
<tr>
<td></td>
<td></td>
<td>30</td>
<td>155</td>
<td>2.41</td>
</tr>
<tr>
<td></td>
<td></td>
<td>40</td>
<td>130</td>
<td>2.24</td>
</tr>
<tr>
<td></td>
<td></td>
<td>50</td>
<td>120</td>
<td>1.98</td>
</tr>
<tr>
<td>Wood pellet</td>
<td>20</td>
<td>40</td>
<td>510</td>
<td>3.60</td>
</tr>
<tr>
<td></td>
<td></td>
<td>50</td>
<td>430</td>
<td>3.50</td>
</tr>
</tbody>
</table>

### Materials

Wood pellet (Jumneansawas 2017) and rice husk (Madhiyanon et al. 2010) were used as fuel in this work. Their properties and heating values are shown in Table 2.

### Table 2. Proximate and Ultimate Analyses of Wood Pellet and Rice Husk

<table>
<thead>
<tr>
<th>Properties</th>
<th>Feedstock</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Rice Husk</td>
</tr>
<tr>
<td>Ultimate analysis (%)</td>
<td></td>
</tr>
<tr>
<td>C</td>
<td>38.00</td>
</tr>
<tr>
<td>H</td>
<td>4.55</td>
</tr>
<tr>
<td>O</td>
<td>32.40</td>
</tr>
<tr>
<td>N</td>
<td>0.69</td>
</tr>
<tr>
<td>S</td>
<td>0.06</td>
</tr>
<tr>
<td>LHV*</td>
<td>20.43</td>
</tr>
<tr>
<td>Empirical formula</td>
<td>CH_{1.40}O_{0.63}N_{0.006}</td>
</tr>
<tr>
<td>Bulk density (kg/m³)</td>
<td>115</td>
</tr>
<tr>
<td>Proximate analysis (%)</td>
<td></td>
</tr>
<tr>
<td>Fixed carbon</td>
<td>20.10</td>
</tr>
<tr>
<td>Volatile</td>
<td>55.60</td>
</tr>
<tr>
<td>Moisture</td>
<td>10.30</td>
</tr>
<tr>
<td>Ash</td>
<td>14.00</td>
</tr>
</tbody>
</table>

*Calculated according to the empirical formula used by Jarungthammachote and Dutta (2007)*

The flame front propagation rate (FPR) was calculated by tracing the location of the peak temperature along the recorded period of time between two adjacent thermocouples (Mahapatra et al. 2014), as indicated below.
\[
FPR(cm/\text{min}) = \frac{\text{Distance thermocouples(cm)}}{\text{Time required to reach the temperature(min)}}
\]

(1)

Bed movement was calculated using Eq. 2.

\[
\text{Bed movement(cm/\text{min})} = \frac{\text{Fuel subsidence in reactor(cm)}}{\text{Time of experimental(min)}}
\]

(2)

**Modeling**

*Equilibrium modeling*

The global species balance for gasification of dry feedstock is expressed by Eq. 3,

\[
CH_xO_yN_z + m(O_2 + 3.76N_2) \rightarrow x_1H_2 + x_2CO + x_3CO_2 + x_4H_2O + x_5CH_4 + (z / 2 + 3.76m)N_2
\]

(3)

where \( x, y, \) and \( z \) are the atomic numbers of hydrogen, oxygen, and nitrogen in the feedstock, respectively, and \( x_1 \) to \( x_5 \) represent the moles of the gasification products per 1 mole of feedstock. The variable \( m \) represents \((4.76 \text{ m})\) the moles of air that were introduced in gasification per 1 mol of feedstock. The atomic balance of carbon, hydrogen, and oxygen atoms could be written in Eqs. 4, 5, and 6, respectively, as shown below.

Carbon balance

\[
x_2 + x_3 + x_5 - 1 = 0
\]

(4)

Hydrogen balance

\[
2x_1 + 2x_4 + 4x_5 - x = 0
\]

(5)

Oxygen balance

\[
x_2 + 2x_3 + x_4 - y - 2m = 0
\]

(6)

There are three major kinetics that contribute to gasification process which are the Boudouard reaction \((C + CO_2 \rightarrow 2CO)\), CO water shift reaction \((CO + H_2O \leftrightarrow CO_2 + H_2)\), and methanation \((C + 2H_2 \leftrightarrow CH_4)\). The equilibrium constants of each kinetic rate are expressed by Eqs. 7, 8, and 9.

Equilibrium constant for CO water shift reaction:

\[
K_1 = \frac{x_2x_4}{x_2x_4}
\]

(7)

Equilibrium constant for methanation combustion,

\[
K_2 = \frac{x_5x_{\text{total}}}{x_3^2}
\]

(8)

where \( x_{\text{total}} \) is the total molar of the gaseous product from Eq. 3, which can be calculated by Eq. 9.
\[ x_{total} = x_1 + x_2 + x_3 + x_4 + x_5 + \left( \frac{Z}{2} + 3.76 \right) \quad (9) \]

The equilibrium constants of any kinetic rates can be calculated from Eq. 10. The parameter \( \Delta G_f^0 \) is the change in Gibbs function of formation obtained from all species involved in the reaction, which are functions of temperature. The calculation of the Gibbs of formation are referred to by Jarungthammachote and Dutta (2007).

\[ K_i = e^{\frac{\Delta G_f^0}{RT}}, i = 1, 2 \quad (10) \]

The energy equation could be written in the form of Eq. 11. The temperature of gas compositions can be calculated by energy conservation principle. \( h_f^0 \) is the enthalpy of formation of the reactant; (kJ/kmol). \( h_f^0 \) is the enthalpy of formation of the products (kJ/kmol). \( \Delta h_p \) is the sensible enthalpy of the products (kJ/kmol). \( C_{p,i} \) is the specific heat of the i\(^{th}\) species in the product, which is a function of temperature (kJ/kmol K). It was calculated in accordance with temperature by using the correlation referred to by Jarungthammachote and Dutta (2007). \( \Delta T \) is the change in temperature with respect to the reference temperature of 298 K.

\[
\Delta h_{f,\text{feedstock}} + m h_{f,\text{H}_2O} + 3.76mh_{f,N_2} = x_1 \left( h_{f,H_2}^o + C_{p,H_2} \Delta T \right) + x_2 \left( h_{f,CO}^o + C_{p,CO} \Delta T \right) + x_3 \left( h_{f,CO_2}^o + C_{p,CO_2} \Delta T \right) + x_4 \left( h_{f,H_2O(\text{super})}^o + C_{p,H_2O(\text{super})} \Delta T \right) + x_5 \left( h_{f,CH_4}^o + C_{p,CH_4} \Delta T \right) + \left( \frac{Z}{2} + 3.76 \right) \left( h_{f,N_2}^o + C_{p,N_2} \Delta T \right) \quad (11)
\]

**Reduction kinetic modeling**

The reduction kinetic rate model was described in previous studies (Sharma 2001; Giltrap et al. 2003). The heat of reaction and the reaction rate of all considering reactions can be written as follows,

\[
C + CO_2 \rightleftharpoons 2CO \quad (172,600 \text{ J/mol}) \quad (12)
\]

\[
C + H_2O \rightleftharpoons CO + H_2 \quad (131,400 \text{ J/mol}) \quad (13)
\]

\[
C + 2H_2 \rightleftharpoons CH_4 \quad (-75,000 \text{ J/mol}) \quad (14)
\]

\[
CH_4 + H_2O \rightleftharpoons CO + 3H_2 \quad (206,400 \text{ J/mol}) \quad (15)
\]

\[
r_1 = CRF \cdot A_1 \exp \left( \frac{-E_1}{RT} \right) \cdot \left( \frac{y_{CO_2}}{K_{R1}} \right) \quad (16)
\]

\[
r_2 = CRF \cdot A_2 \exp \left( \frac{-E_2}{RT} \right) \cdot \left( \frac{y_{H_2O}}{y_{CO}y_{H_2}} \right) \quad (17)
\]
$$r_i = CRF \cdot A_i \exp \left( \frac{-E_i}{RT} \right) \left( y_{H_2}^2 - \frac{y_{CH_4}^2}{K_{R3}} \right)$$ \hspace{1cm} (18)

$$r_i = A_i \exp \left( \frac{-E_i}{RT} \right) \left( y_{H_2O} y_{CH_4} - \frac{y_{H_2}^3 y_{CO}}{K_{R4}} \right)$$ \hspace{1cm} (19)

where \( r_i \) is the rate of reaction for the \( i^{th} \) reaction; (kmol/m\(^3\)/s), \( A_i \) is the frequency factor for the \( i^{th} \) reaction; (kmol/m\(^3\)/s), \( E_i \) is the activation energy of the \( i^{th} \) reaction; (kj/kmol). \( R \) is the universal gas constant; (kj/kmol), \( T \) is the temperature, \( y_x \) is the mole fraction of \( x \) species, \( k_i \) is the equilibrium constant for the \( i^{th} \) reaction, and \( CRF \) is the char reactivity factor.

**Model solution procedure**

The flow chart of the numerical procedure is illustrated in Fig. 2. The chemical equilibrium modeling was employed to predict the temperature and the composition of the combustion zone. Then the obtained temperature and composition was used as the initial value in reduction kinetic modeling in the reduction zone. In obtaining the solution for the chemical equilibrium modeling, there was a need to couple between an energy equation and the system of chemical equilibrium equations. This could be achieved through an iterative technique until the relative error of the predicted temperature was at an acceptable tolerance. For the solution of reduction kinetic modeling, the finite layer height was defined in the reduction zone and the residence time in each layer was estimated. The temperature and the composition were then used as the input for the adjacent layer below. More detail of the zone definition from the experimental information will be discussed in the next section.

Fig. 2. Calculation procedure

**RESULTS AND DISCUSSION**

**Flame Propagation and Kinetic Mechanism in Stratified Downdraft Gasification**

The flame propagation and characteristics of each regime of the stratified downdraft gasification in different conditions were similar. It comprised of a combustion zone at the flame front following by a reduction zone, as depicted in Fig. 3a. The steep
temperature gradient where the temperature rose to its peak was observed in the combustion zone. This was due to major heat release from volatile combustion within the pores of the packed fuel bed. Oxygen was completely consumed within this zone. The fresh fuel was devolatilized, thus leaving char behind as the flame propagation moved counter to the supplied oxidizer stream as shown in Fig. 3b.

The reduction zone was located below the combustion zone. In this zone, the kinetic mechanism took place in the absence of the oxidizer. Flue gas from the combustion zone then reacted with solid char at a high temperature to produce carbon monoxide and hydrogen. While carbon dioxide and steam were consumed in the reduction process, the process was endothermic overall, so a decrease in temperature along the direction of the flue gas was observed in this zone.

**Fig. 3.** Zone and temperature at specific times (a) and zone and temperature movement at different time periods (b)

**Flame Propagation in Stratified Downdraft Gasification**

In stratified downdraft gasification conditions, flame propagation moved counter to the supply oxidizer stream, as calculated by Eq. 1. The speed of the propagation depended on various factors, for example, fuel particle type, fuel size, moisture content in fuel, and air mass flow rate. The flame propagation mechanism was controlled by diffusion heat and mass transports in the direction counter to the supplied oxidizer stream. This was observed at the reaction front where a steep temperature gradient was presented. There were two distinctive regimes revealed in the propagation characteristic regardless the type of feedstock.

In regime I, at a low range of air flow rate, the propagation speed increased with increasing air flow rates. As the air flow rate increased, at a certain point, the peak propagation speed was reached. In regime II, at a higher range of air flow rate, propagation speed decreased with increasing air flow rate (Sadhan et al. 2014). In this scenario, increasing the air flow rate more than that resulted in the peak propagation speed. The propagation speed has a direct variation with the peak temperature under different air flow rates and the same feedstock conditions (Onthong et al. 2016).
Fig. 4. Temperature distribution at different times in the reactor at various air flow rates for rice husk.

After the ignition period, the flame was propagated upwards, as can be seen in the temperature profile evolution in Figs. 4 and 5. The propagation speed of each case was demonstrated in Fig. 6. The values of all cases were calculated by Eq. 1. All cases fell within regime I as stated in previous paragraph. The propagation rate of the wood pellet was around five times greater than rice husk at the same air flow rate. This could be because of the higher bulk density of the wood pellet, which was around six times higher in the randomly packed bed. The higher bulk density of the feedstock required more
thermal energy input to heat it up to de-volatilization temperature level at the reaction front. Heat was transferred from the combustion front in diffusion mode to heat up the fresh biomass until volatile combustion occurred within the pores space at the adjacent layer above the current location of the combustion front.

![Graph showing temperature distribution at different times in the reactor at various air flow rates for wood pellet](image)

**Fig. 5.** Temperature distribution at different times in the reactor at various air flow rates for wood pellet

![Graph showing propagation rate and bed movement at different air flow rates](image)

**Fig. 6.** Propagation rate and bed movement at different air flow rates

The combustion temperature, indicated by the peak temperature as shown in Fig. 3, was found to increase with the increasing air flow rate. A higher temperature would
have allowed a greater rate of heat transfer in the diffusion mode at the reaction front, which had resulted in a higher propagation rate of the flame front. Increasing the air flow rate would enrich the oxidizer concentration, which would have resulted in a higher heat release rate at the flame front. However, Sadhan et al. (2014) found that increasing the air flow rate until reaching a certain value will provide peak propagation rate. Further increasing air flow rate would then result in decreases in the propagation rate and the combustion temperature because the convection loss at the flame front was more dominant. This was considered as regime II, which was explained in previous paragraph.

A spike in temperature was observed for every case, as highlighted in Figs. 4 and 5. This happened when the propagation front reached the top surface of the packed fuel bed. This was the indication for termination in every experimental case. The air flow rate and data logger were stopped, and the data was collected for analysis at this point. The spike in temperature appeared because the flame front had reached the top fuel layer. This prevented the flame from propagating. The flame front stayed there until the volatiles had been completely released and the combustion regime had changed to char burning mode. There was a considerably higher combustion temperature in the char burning mode due to less dilution constituent in comparison with the volatile combustion mode. If the air flow was continued, the flame propagation front of the char burning mode moved downward until it reached the fuel grate, leaving incombustible ash behind (Ryu et al. 2006). Considerable bed movement due to fuel particle shrinkage was observed, as seen in Fig. 6.

**Combustion Zone Prediction (Equilibrium Modeling)**

The combustion zone temperature was defined as the peak temperature for any experimental cases as given in Fig. 7. It was also considered as the beginning of reduction zone, since the combustion zone in the stratified downdraft condition is typically very thin (Blasi 2000).

![Fig. 7. Comparison of the equilibrium analysis and experimental test on temperature](image)

The reduction zone occupied the area underneath the combustion zone downward to the combustor grate. It was necessary to define the $m$ value (4.76 moles of air per a mole of fuel) for equilibrium modeling in Eq. 3. The global kinetic equation can be written from the known fuel composition. The stoichiometric air to fuel ratio was
calculated based on previous studies (Madhiyanon et al. 2010; Jumneansawas 2017). The experimental condition for the air to fuel mass ratio can be estimated by the total amount of air required for a complete batch divided by complete batch of feedstock usage, as seen in Table 1.

All the predicted temperatures were higher than the experimental results as seen in Fig. 7. It was assumed that the model did not account for any heat loss. In fact, there was heat loss at the combustion zone. Thermal energy at the combustion zone had diffused upstream allowing the propagation mechanism to proceed as stated in the previous paragraph. It was also due to the convection downstream that promoted char reduction endothermic kinetics. Increasing the air flow rate had resulted in a leaner air-fuel mixture. The product species tended to have a greater composition of combustion product (CO₂ and H₂O), as can be seen in Fig. 8.

![Fig. 8. Equilibrium analysis on the gas composition of the product](image)

The combustion temperature of the wood pellet was a little lower than that of the rice husk. For wood pellet, volatile combustion was quenched when the supply air was fed at 20 L/min and 30 L/min. This was due to the greater size and higher bulk density of the wood pellet than those of rice husk because of the higher thermal inertia of the fuel particle. However, the limited concentration of the oxidizer had restricted the rate of heat generation during homogeneous volatile combustion within the pore space. If heat generation was lower than the heat loss, the flame would decrease in temperature and finally be quenched.

**Reduction Zone Prediction (Reduction Kinetic Modeling)**

High temperature flue gas from the combustion zone reacted with solid char to produce carbon monoxide and hydrogen species in the reduction zone. The kinetics involved were boudouard, water gas, methanation, and water gas shift reactions, as shown in Eqs. 12, 13, 14, and 15, respectively. The calculation procedure began by defining the reduction zone length and dividing it into a finite number of layers. Property transport between vertically adjacent layers was considered only due to convection where diffusion effect was ignored. This resembled the plug flow reactor model (Turns 1988).

The initial properties of the first layer were included. Gas compositions and temperatures were obtained from the combustion zone calculation. The initial superficial velocity was calculated from the total air supply flow rate, porosity of the packed bed, initial gas mixture, initial temperature, and the averaged mass loss of the case.
The temperature profiles for most cases gave similar trends. The temperature decreased at a higher rate at the first 10 cm from its peak value within the reduction zone, as they exhibited steeper gradients in comparison with other layers, as can be seen in Figs. 9 and 11. This was because the high temperature of flue gas from the combustion zone allowed a greater reduction in the kinetic rate. The reduction kinetics were endothermic so the kinetic rates at certain locations were reflected by the temperature gradient. A higher temperature gradient at the beginning of the reduction zone was observed in the case of higher air flow rates. This was due to the higher temperature in the combustion zone when compared with the cases with lower air flow rates.

**Fig. 9.** Comparison of the temperature along the length of reactor for rice husk at certain time steps
Fig. 10. Effect of air flow rate on the gas composition for rice husk

(a) 20 (L/min)
(b) 30 (L/min)
(c) 40 (L/min)
(d) 50 (L/min)

Fig. 11. Comparison of temperature along the length of the reactor for wood pellet at certain time steps

(a) 40 (L/min)
(b) 50 (L/min)
The reason for the discrepancy between the experimental data and the simulation model solution for the temperature profile in the case of rice husk with 20 liter/min air supply was not clear. It was possible that the reduction kinetic had considerably slower rates due to the low combustion temperature at the beginning of the reduction zone, as indicated by the relatively mild decrease in temperature seen in Fig. 9a. Consequently, the adjacent lower layer with a high gas temperature had a greater amount of heat loss to the lower solid char particle in comparison with the endothermic reduction at the beginning of the reduction zone. Therefore, the temperature decreased with the higher downstream gradient. Kinetic modeling did not account for the heat transfer due to different temperatures between the gas and char.

Kinetic modeling with a CRF value of 10 provided similar trends of temperature profiles as in the experimental results. However, in the case of rice husk at air flow rates of 40 L/min and 50 L/min, a decrease in the temperature at the beginning of the reduction zone was over-predicted, as shown in Figs. 9c and 9d. This was due to the over-prediction in the temperature of the combustion zone by equilibrium modeling, which was used as the initial temperature in reduction modeling. The Arrhenius rate model had an exponential relation with temperature, which meant it had a high sensitivity to the accuracy of the temperature, which was used as initial value. The CO₂ and H₂O mole fraction decreased while CO and H₂ increased along the reduction zone. H₂ slightly decreased when the air flow rate was increased. The exit compositions were not significantly different for all of the cases, as can be seen in Figs. 10 and 12. The compositions and temperature profiles of the modeling results were in agreement with previous research (Giltrap et al. 2003; Babu and Sheth 2006).

**CONCLUSIONS**

Flame propagation under downdraft gasification at different air flow rates was investigated. The effects of different feedstocks on the propagation, namely rice husk and wood pellet, were considered. Flame propagation rate, mass loss, and bed movement was discussed. Equilibrium modeling was employed to predict the temperature and gas composition at the combustion zone. The equilibrium model results were used as the initial conditions for the reduction zone calculation. The temperature and gas composition along the reduction zone were predicted using finite kinetic modeling.
1. Flame propagation increased with increasing air mass flow rates for all the cases with rice husk and wood pellet between 20 L/min to 50 L/min. For wood pellet, the flame propagation (volatile combustion) was quenched in when the supply air flow rate was between 20 L/min and 30 L/min. This was because the bulk density of the wood pellet was considerably higher than rice husk, which resulted in massive heat loss to the fuel particle during the volatile combustion within the cavity space of the packed fuel bed.

2. The propagation rate of wood pellet was around five times greater than rice husk at the same air flow rate. This was because of the high bulk density of the wood pellet, which was around five times higher than the rick husk in the randomly packed bed. The higher bulk density of the feedstock required more thermal energy input to heat up to a de-volatilization temperature level at the reaction front, allowing flame front propagation.

3. Equilibrium modeling tended to overestimate the combustion temperature. This was possibly because it did not account for any heat loss, while there was a certain degree of heat loss at the flame front from upstream diffusion heat transfer to the feed propagation mechanism and downstream convection heat transfer.

4. Reduction kinetic modeling had a high sensitivity to the initial temperature input. The over-prediction of temperature in the combustion zone by equilibrium modeling led to over-prediction of the temperature gradient at the beginning of the reduction layer.

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