PERFORMANCE PREDICTION AND VALIDATION OF EQUILIBRIUM MODELING FOR GASIFICATION OF CASHEW NUT SHELL CHAR

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Abstract - Cashew nut shell, a waste product obtained during deshelling of cashew kernels, had in the past been deemed unfit as a fuel for gasification owing to its high occluded oil content. The oil, a source of natural phenol, oozes upon gasification, thereby clogging the gasifier throat, downstream equipment and associated utilities with oil, resulting in ineffective gasification and premature failure of utilities due to its corrosive characteristics. To overcome this drawback, the cashew shells were de-oiled by charring in closed chambers and were subsequently gasified in an autothermal downdraft gasifier. Equilibrium modeling was carried out to predict the producer gas composition under varying performance influencing parameters, viz., equivalence ratio (ER), reaction temperature (RT) and moisture content (MC). The results were compared with the experimental output and are presented in this paper. The model is quite satisfactory with the experimental outcome at the ER applicable to gasification systems, i.e., 0.15 to 0.30. The results show that the mole fraction of (i) H₂, CO and CH₄ decreases while (N₂ + H₂O) and CO₂ increases with ER, (ii) H₂ and CO increases while CH₄, (N₂ + H₂O) and CO₂ decreases with reaction temperature, (iii) H₂, CH₄, CO₂ and (N₂ + H₂O) increases while CO decreases with moisture content. However at an equivalence ratio less than 0.15, the model predicts an unrealistic composition and is observed to be non valid below this ER.

Keywords: Equilibrium modeling; Gasification; Cashew shell char; Producer gas,

INTRODUCTION

Cashew (Kingdom - Plantae, Division - Magnoliophyta, Class - Magnoliopsida, Order - Sapindales, Family - Anacardiaceae, Genus - Anacardium, Species - A occidentale, Binomial name - Anacardium occidentale L) (http://en.wikipedia.org/wiki/Cashew), native to eastern Brazil, was introduced into India, just as other commercial crops like rubber, coffee, tea etc. by the Portuguese nearly five centuries ago. The cashew was introduced into India in Goa, from where it spread to other parts of the country. In the beginning it was mainly considered as a crop for afforestation and soil binding to check erosion. The nuts, apple and other by-products of this crop are of commercial importance. Because of its adaptability to a wide range of agroclimatic conditions, it has become a crop of high economic value and has attained the status of an export-oriented commodity, contributing considerably to the foreign exchange earnings of the country (http://dacnet.nic.in/cashewcocoa/cashew.htm).

Presently, India is the largest producer, processor and exporter of cashews (http://www.cashewindia.org). Cultivation of cashews in India is confined mainly to the peninsular areas. It is grown in Kerala, Karnataka, Goa and Maharashtra along the west coast and Tamil Nadu, Andhra Pradesh, Orissa and West Bengal along the east coast (Agricultural
The cashew tree, generally a spreading, evergreen tree, has simple, alternate, glabrous, penninerved leaves. The true fruit of the cashew tree is a kidney or boxing-glove shaped drupe that grows at the end of the pseudo-fruit. The cashew fruit (Figures 1 and 2) is unusual in comparison with other tree nuts since the nut is outside the fruit. The cashew apple is an edible pseudo fruit, attached to the externally born nut by a stem. In its raw state, the shell of the nut is leathery, not brittle. It contains thick vesicant oil, cashew nut shell liquid (CNSL), within a sponge-like interior. A thin skin, called the testa surrounds the kernel and separates it from the inside of the shell. The primary products of cashew nuts are the kernels which are valuable as confectionery nuts, CNSL - an important industrial raw material for resin manufacture and the shells – burnt (combusted) inefficiently for meeting thermal energy requirements.

CASHEW PROCESSING

Traditionally, extraction of the kernel from the shell of the cashew nut has been a manual operation. The nut is roasted, which makes the shell brittle and loosens the kernel from the inside of the shell. By soaking the nuts in water, the moisture content of the kernel is raised, reducing the risk of it being scorched during roasting and making it more flexible so it is less likely to crack. The CNSL is released when the nuts are roasted. The shell can be cracked either manually, using a hammer or mechanically. Manually operated blade openers are relatively inexpensive. Once the kernel is removed from the shell, it is dried, the testa is peeled off and the kernel is graded. Figure 3 gives an overview of cashew nut processing and the various choices of method.

The cashew nut shells (CNS), generated as a waste by-product, are being sold at a throw away price. Due to their low cost and abundant availability, cashew shells are utilized in an inefficient manner. These shells are burnt in a way that is generally regarded as a disposal option rather than a source of energy. The CNSL is reported to be 15-20% by weight of the unshelled nut in Africa, 25-30% by weight in India and 25% overall (Piyali Das and Anuradda Ganesh, 2003).

GASIFICATION OF CASHEW SHELL

Owing to the rise in the cost of conventional fuel and its difficult sourcing problems, attention is now being focused towards an efficient utilization of cashew shells. Gasification is one of the effective and efficient bioenergy conversion technologies - known for its modularity, turn down ratio and fuel versatility - available for harnessing this renewable bioenergy resource. Attempts have been made to convert these shells into a burnable low btu gas. However the inherent CNSL content of these shells make the gasification technology infeasible in the long run (IES, Anna University, 2006). In order to overcome the drawback posed by CNSL, the shells were charred in a chamber, thereby de-oiling them for consequent gasification. The characteristics of raw and charred cashew nut shell (charred CNS) are presented in Tables 1 and 2. The bulk density of charred CNS was measured to be 190 kg/m³, indicating its amenability for gasification.
Figure 3: Cashew Processing – An Outline

Table 1: Proximate analysis of raw and charred cashew shells

<table>
<thead>
<tr>
<th></th>
<th>CNS (wt% on an as-received basis)</th>
<th>Charred CNS (wt% on an as-received basis)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Moisture</td>
<td>moisture</td>
<td>7</td>
</tr>
<tr>
<td>Volatile Matter</td>
<td>10.43</td>
<td>28</td>
</tr>
<tr>
<td>Fixed Carbon</td>
<td>69.31</td>
<td>59</td>
</tr>
<tr>
<td>Ash</td>
<td>19.26</td>
<td>6</td>
</tr>
</tbody>
</table>

Table 2: Ultimate analysis of raw and charred cashew shells

<table>
<thead>
<tr>
<th></th>
<th>CNS (wt% on an as-received basis)</th>
<th>Charred CNS (wt% on an as-received basis)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Carbon</td>
<td>48.7</td>
<td>63</td>
</tr>
<tr>
<td>Hydrogen</td>
<td>6.96</td>
<td>3.6</td>
</tr>
<tr>
<td>Nitrogen</td>
<td>0.36</td>
<td>6.4</td>
</tr>
<tr>
<td>Oxygen</td>
<td>43.98</td>
<td>27</td>
</tr>
</tbody>
</table>
EXPERIMENTAL SETUP

The main objective of this study was (i) to investigate the feasibility of gasification of cashew nut shells in a down-draught gasifier in the long run and (ii) to ascertain the effect of equivalence ratio (ER), moisture content (MC) and reaction temperature (RT) on gasification of cashew nut shells. Figures 4 (a), (b) and (c) depict a schematic diagram and a photograph of the experimental setup.

An induced air, closed-top, 20 kW<sub>e</sub>, down-draught gasifier was chosen for the study. Provision for air entry into the gasification system was made through two inclined tuyeres, unclosed at the throat. A grate made of SS was used for holding the feedstock. The region between throat and grate functioned as a reduction zone. The hot gas generated was passed along the annuli of the reduction zone for maintenance of high reaction temperature and minimization of heat loss. A cyclone separator was placed at the gasifier outlet ahead of the ID blower to remove particulates. An aerated burner was used for flaring the producer gas generated. The whole gasifier assembly was mounted on a toughened helical spring, enabling it to vibrate to ensure a smooth fuel flow. A poking rod was fixed at the bottom and linked to the grate. Pushing and pulling the poking rod created an impact on the grate, thereby disintegrating and discharging any clogged residual particles into the ash box. Provisions were made for measuring the temperature, pressure and gas composition at appropriate location in the gasifier. A stand with a ladder arrangement was used for feeding the raw material in at the fuel feed port.

Figure 4(a): Schematic diagram of Gasifier
PROCESS INSTRUMENTATION AND CONTROL

The base fuel characteristics were established with a proximate analyzer (muffle furnace + micro weighing balance with associated auxiliaries). The parameters studied included moisture content (ASTM D 3173-73), volatile matter (ASTM D 3175-73) and ash content (ASTM D 3174-73). The remainder was fixed carbon (FC). The calorific value of cashew shell was established using a standardized (benzoic acid-based) bomb calorimeter and the result was observed to match well the correlations framed for estimation of calorific value for biofuels (Parikh et al. 2005). A Junkers gas calorimeter was used for determining the calorific value of producer gas. Producer gas composition was analyzed using a Siemens Oxymat 61 online gas analyzer (which estimates O$_2$ using the paramagnetic principle), Ultramat 23 (which estimates CO, CO$_2$ and CH$_4$ using nondispersive infrared multilayer technology) and Calomat 61 (which estimates H$_2$ using the thermal conductivity principle). Details regarding the producer gas composition were recorded on the PC every second using Siprom-GA software. The gas sampling system consisted of a wash bottle, a condensation pot, a coalescence filter, a suction pump, a fine filter, a flame arrestor and a diaphragm pump. Chromel-Alumel (K-type) thermocouples were used for measuring the temperature in different zones (T1 to T6). Thermocouples were fixed permanently and measured continuously in all zones except at the throat (T4). For temperature measurement at the throat, a flexible K-type thermocouple was inserted along the air port at regular intervals. Temperatures from different zones were logged simultaneously using an Agilent (34907 A) data acquisition system. The surface temperature of the gasifier was measured with a Kane infrared thermometer (UEI-INF 200). An Airflow thermoanemometer (TA 35) was used for measuring the air flow to the gasifier. A calibrated S-type Pitot Tube & Comark digital manometer was employed for establishing the producer gas flow. The air entry into the system, and thus the ER, is controlled by a globe valve placed at the discharge end of the blower. Water-filled U-tube manometers were deployed for measuring the pressure buildup across the gasifier bed.

EXPERIMENTAL PROCEDURE

Preweighed batches of CNSC, each weighing approx 25 kgs, were placed near the system for hassle- free operation during fuel loading. Gasification of CNSC was initiated by keeping the gas valve open, followed by operation of the ID blower and holding a flame near the air tuyere. The flame was sucked into the system due to the draught created by the blower. Within three minutes, flue gas was observed at the flare port. Normally after 5 to 10 minutes, the onset of gasification commenced and producer gas emanated from the flare port. Experimental analysis was started once the system become stabilized. Generally it took 60 to 90 min to achieve stabilization, which was ensured by inferring a constant temperature in the raw gas and the
reduction zone. The fuel consumption rate was measured by recharging the gasifier on an hourly basis and filling the gasifier volume to a predetermined level at the top of the gasifier hopper. The ash door was operated at regular intervals to remove the ash that had accumulated on the grate.

The major influential parameters in the performance of gasification system are equivalence ratio, the bed temperature and moisture content of the feed stock. Equivalence ratio was varied by adjusting the air supplied to the gasifier bed. Since, the gasifier used was autothermal, the option of maintaining constant bed temperature for varying equivalence ratio/moisture content was ruled out. The experiments were conducted in various permutations and combinations and the outcomes are compared with the modeled results.

EQUILIBRIUM MODELING

The typical chemical formula of CNS char, based on a single atom of carbon, is observed to be CH$_{0.686}$O$_{0.32}$. Equilibrium modeling was carried out, based on this formula, for predicting its gasification characteristics.

The following assumptions are made in the modeling approach:

- The ideal gas laws are valid.
- All reactions are at thermodynamic equilibrium.
- Gases are in equilibrium during flow through the char bed.
- The pressure in the char bed is atmospheric and constant.
- Reactions proceed adiabatically.
- Nitrogen present in both fuel and air is inert.
- Ash is inert and is not involved in any of the reactions, either as a chemical species or as a catalyst.
- No radial temperature gradients/concentrations exist.
- No gas is accumulated in the char bed.
- There is no resistance to conduction of heat and diffusion of mass inside the char particles.
- There is no tar in the gasification zone.
- Carbon conversion efficiency is 100%.
- Producer gas comprises only CO$_2$, CO, H$_2$, CH$_4$, N$_2$ and H$_2$O.

Based on the above assumptions, the global reaction of CNS char with air could be written as

\[
\text{CH}_{0.686}\text{O}_{0.32} + \text{wH}_2\text{O} + \text{mO}_2 + 3.76\text{mN}_2 = \text{x}_1\text{H}_2 + \text{x}_2\text{CO} + \text{x}_3\text{CO}_2 + \text{x}_4\text{H}_2\text{O} + \text{x}_5\text{CH}_4 + 3.76\text{mN}_2
\]  

The above reaction represents an overall reaction, but a number of competing intermediate reactions take place during the process, of which the following are predominant:

Oxidation
\[
\text{C + O}_2 = \text{CO}_2 \quad (-393.8 \text{ kJ/mol})
\]  

Steam gasification
\[
\text{C + H}_2\text{O} = \text{CO} + \text{H}_2 \quad (+131.4 \text{ kJ/mol})
\]  

Boudouard reaction
\[
\text{C + CO}_2 = 2\text{CO} \quad (+172.6 \text{ kJ/mol})
\]  

Methanation reaction
\[
\text{C + 2H}_2 = \text{CH}_4 \quad (-74.9 \text{ kJ/mol})
\]  

Water gas shift reaction
\[
\text{CO + H}_2\text{O} = \text{CO}_2 + \text{H}_2 \quad (-41.2 \text{ kJ/mol})
\]  

Of these, only four reactions are independent, the oxidation, steam gasification, Boudouard and methanation reactions. The water gas shift reaction can be considered as the subtraction of the steam gasification and Boudouard reactions. According to Von Fredersdorff and Elliot (1963) the oxidation reaction is typically assumed to be very fast and is completed quickly, while the other three reactions, namely boudouard, steam gasification and methanation, are in equilibrium. As observed, the water gas shift reaction is a combination of the boudouard and steam gasification reactions. Hence, the water gas shift and methanation reaction could be considered to be in equilibrium.

The equilibrium constants for the methanation and water gas shift reactions [Eq. (5)] are

\[
K_1 = \frac{[\text{P}_{\text{CH}_4}]}{[\text{P}_{\text{H}_2}]}^2
\]  

\[
K_2 = \frac{[\text{P}_{\text{CO}_2}]\cdot[\text{P}_{\text{H}_2}]}{[\text{P}_{\text{CO}}]\cdot[\text{P}_{\text{H}_2}\text{O}]}
\]  

In the global reaction [Eq. (1)], there are six unknowns, \(x_1, x_2, x_3, x_4, x_5\) and \(m\), representing the
molar composition of five unknown species in the producer gas and the oxygen content for the reaction. Hence, to predict the constituents of producer gas, a set of six equations, formulated by balancing the different constituents involved in the global reaction, is required.

Carbon balancing
\[ 1 = X_2 + X_3 + X_5 \]  

(9)

Hydrogen balancing
\[ 2 w + 0.6847 = 2 X_1 + 2 X_4 + 4 X_5 \]

(10)

Oxygen balancing
\[ w + 0.3217 + 2 m = X_2 + 2 X_3 + X_4 \]

(11)

Equilibrium constant from methanation [Eq. (5)]
\[ K_1 = \frac{X_5}{(X_1)^2} \]

(12)

Equilibrium constant from the shift reaction [Eq. (8)]
\[ K_2 = \frac{(X_3)(X_4)}{(X_2)(X_4)} \]

(13)

The heating value of the fuel \( (H^°_f)^{CNSC} \) was determined experimentally with a bomb calorimeter. The heat of formation of the various gases could be sourced from the JANAF thermochemical tables (Stull and Prophet, 1971) and the \( \Delta H^° \) for the gases constituting the present study are presented in Table 3.

The dependence of specific heat on temperature is given by various empirical equations and the most simplified version (Robert and Don, 1984) is

\[ C_p = R \left[ A + BT_{am} + \frac{C_3}{3} (4T^2_{am} - T_1^2 T_2) + \frac{D}{T_1^2 T_2} \right] \]

The values of the heat capacity constants, applicable to our equations, are shown in Table 4.

The equilibrium constant \( K \) is a function of temperature and could be equated as (Zainal et al. 2001)

\[ -RT \ln K = \Delta G^° \]

(18)

\[ \frac{d \ln K}{dT} = \frac{\Delta H^°}{RT^2} \]

(19)

The values of standard Gibbs function of formation for various gas compositions involved are shown in Table 5 (Stull and Prophet, 1971)

Heat of formation is basically a function of temperature, and hence Eq. (19) can be integrated as follows (Zainal et al. 2001):

\[ \ln K = \frac{\Delta H^°}{RT^2}dT + I \]

(20)

\( \Delta H^° \) could also be equated as (Robert and Don, 1984)

\[ \frac{\Delta H^°}{R} = \frac{J}{R} + (\Delta A) T + \frac{\Delta B}{2} T^2 + \frac{\Delta C}{3} T^3 - \frac{\Delta D}{T} \]

(21)

Substituting Eq. (21) into Eq. (20)

\[ \ln K = \frac{J}{RT} + \Delta A \ln T + \frac{\Delta B}{2} T^2 + \frac{\Delta C}{6} T^2 - \frac{\Delta D}{2T^2} + I \]

(22)
The dependence of $\Delta G^\circ$ on temperature can be analyzed as

$$\Delta G = J - RT (\Delta A \ln T + \frac{\Delta B}{2} T + \frac{\Delta C}{6} T^2 + \frac{\Delta D}{2T^2} + I)$$

(23)

Both $J$ and $I$ are respectively calculated from Eq. 21 and Eq. 23 at temperature 298.15 K.

Two equilibrium equations are required to determine the equilibrium constants $K_1$ and $K_2$. $K_1$ is the equilibrium constant for the reaction in Eqn. (5) and is solved as follows:

$$C + 2H_2 = CH_4$$

$\Delta = CH_4 - C - 2H_2$

$\Delta A$, $\Delta B$, $\Delta C$ and $\Delta D$ can be obtained from the data on heat capacity.

The equations to determine the values of $\Delta A$, $\Delta B$, $\Delta C$ and $\Delta D$ can be expressed as

$$\Delta A = A_{CH_4} - A_C - 2A_{H_2}$$

$$\Delta B = B_{CH_4} - B_C - 2B_{H_2}$$

$$\Delta C = C_{CH_4} - C_C - 2C_{H_2}$$

$$\Delta D = D_{CH_4} - D_C - 2D_{H_2}$$

Application of these equations reveals the equilibrium constants for $K_1$.

Similarly $K_2$ is the equilibrium constant for the water gas shift reaction (Eq. 6) and could be solved by adopting the same procedure. Figure 5 depicts the calculation procedure for the equilibrium model in a concise fashion.

The molar concentration of individual species could be predicted by using $K_1$ and $K_2$ and solving equations 9 to 16.

---

**Table 3: Heat of formation at 298.15 K (Stull and Prophet, 1971)**

<table>
<thead>
<tr>
<th>Chemical Species</th>
<th>Phase</th>
<th>$\Delta H^\circ_{298}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Water</td>
<td>g</td>
<td>-241818</td>
</tr>
<tr>
<td>Water</td>
<td>l</td>
<td>-285830</td>
</tr>
<tr>
<td>Carbon dioxide</td>
<td>g</td>
<td>-393509</td>
</tr>
<tr>
<td>Carbon monoxide</td>
<td>g</td>
<td>-110525</td>
</tr>
<tr>
<td>Methane</td>
<td>g</td>
<td>-74520</td>
</tr>
<tr>
<td>Hydrogen</td>
<td>g</td>
<td>0</td>
</tr>
<tr>
<td>Oxygen</td>
<td>g</td>
<td>0</td>
</tr>
<tr>
<td>Nitrogen</td>
<td>g</td>
<td>0</td>
</tr>
</tbody>
</table>

**Table 4: Heat Capacities for constants A, B, C and D (Robert and Don, 1984)**

<table>
<thead>
<tr>
<th>Chemical Species</th>
<th>$T_{max}$</th>
<th>$A$</th>
<th>$10^3 B$</th>
<th>$10^3 C$</th>
<th>$10^4 D$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Methane</td>
<td>1500</td>
<td>1.702</td>
<td>9.081</td>
<td>-2.164</td>
<td>-</td>
</tr>
<tr>
<td>Hydrogen</td>
<td>3000</td>
<td>3.249</td>
<td>0.422</td>
<td>-</td>
<td>0.083</td>
</tr>
<tr>
<td>Carbon monoxide</td>
<td>2500</td>
<td>3.376</td>
<td>0.557</td>
<td>-</td>
<td>-0.031</td>
</tr>
<tr>
<td>Carbon dioxide</td>
<td>2000</td>
<td>5.457</td>
<td>1.047</td>
<td>-1.157</td>
<td>-</td>
</tr>
<tr>
<td>Nitrogen</td>
<td>2000</td>
<td>3.280</td>
<td>0.593</td>
<td>-</td>
<td>0.040</td>
</tr>
<tr>
<td>Water</td>
<td>2000</td>
<td>3.470</td>
<td>1.450</td>
<td>-</td>
<td>0.121</td>
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<tr>
<td>Carbon</td>
<td>2000</td>
<td>1.771</td>
<td>0.771</td>
<td>-</td>
<td>-0.867</td>
</tr>
</tbody>
</table>

**Table 5: Gibbs functions of formation at 298.15 K (Stull and Prophet, 1971)**

<table>
<thead>
<tr>
<th>Chemical Species</th>
<th>Phase</th>
<th>$\Delta G^\circ_{298}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Water</td>
<td>g</td>
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<tr>
<td>Water</td>
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<tr>
<td>Carbon dioxide</td>
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<tr>
<td>Carbon monoxide</td>
<td>g</td>
<td>-137169</td>
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<tr>
<td>Methane</td>
<td>g</td>
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<tr>
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<tr>
<td>Oxygen</td>
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<td>0</td>
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<tr>
<td>Nitrogen</td>
<td>g</td>
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RESULTS AND DISCUSSIONS

Effect of Equivalence Ratio on Molar Concentration

Equivalence ratio (ER) is the ratio of the actual A/F ratio to the stoichiometric A/F ratio. Discarding the fuel component, it could also be defined as the ratio of actual air supplied to the stoichiometric air requirement. For an ideal combustion, the value of ER is 1, while in practical cases it is always greater than 1, being governed predominantly by the type of fuel being combusted. For gasifiers, the value of ER would range from 0.15 to 0.4. In the process of autothermal gasification, part of fuel is burnt to release energy to sustain the endothermic gasification reactions. The lower limit of ER in an autothermal gasifier is fixed by considering a variety of factors like the minimum quantity of air required to burn part of the fuel to release energy for supporting endothermic reactions, required carbon conversion efficiency, the fixed loss of heat that needs to be accounted for in maintaining the reactor temperature etc., Similarly, the upper limit of ER is fixed by factors like tar quantity, gas quality, reactor temperature and ash fusion point.

The effect of ER on gasification of CNSC at a fixed moisture of 7 % and at a reaction temperature of 1373 K is depicted in Figs 6 to 10.
Figure 6: Molar % of H₂ vs ER

Figure 7: Molar % of CO vs ER

Figure 8: Molar % of CO₂ vs ER

Figure 9: Molar % of CH₄ vs ER

Figure 10: Molar % of N₂ + H₂O vs ER

Figure 11: Molar % of CO₂ vs ER Fuel: Casuarina Wood
a) Effect of ER on Hydrogen and CO

The yield of hydrogen from the model is observed to follow a decreasing trend with increasing ER. A similar trend is reported by other researchers like Turn et al. (1998). The trend is quite misleading at first sight. It is a common reality that as the ER increases, the temperature of the any oxidation reaction is bound to increase. This trend is applicable only to allothermal systems, where the temperature of the gasifier is controlled externally (McKendry, 2002). In equilibrium modeling, it is assumed that temperature is maintained constant even at different ERs. However for autothermal systems, the increase in ER would have two paradoxical phases, one following the other in a sequential manner. When the ER is increased from low values, the temperature of the system increases, resulting in a marked increase in generation of both gas and its H₂ concentration - phase 1. However beyond a governing limit, the oxidation reaction predominates due to availability of excess oxygen and the yield of H₂ drops for conversion of H₂ to H₂O - phase 2. The typical ER reported by Lv et al. (2004) is 0.19 to 0.23 for phase 1 and 0.23 to 0.27 for phase 2. The gasifier adopted for experimenting gasification of CNSC is autothermal. Hence, the bed reaction temperatures varied with different ERs. The model predicts unreasonably high H₂ at lower ERs. Ruggiero and Manfrida (1999) described the irrelevance of equilibrium models, which assume perfect gas behavior for very low ERs as these models cannot describe pyrolysis processes due to the presence of liquid hydrocarbons as pyrolysis products. Desrosiers (1981) predicted thermodynamic ER for gasification of dry wood as 0.28. Considering the range of H₂ yield, as predicted by the model with experimental values in the ER range of 0.25 to 0.35, the model values compare quite reasonable with the experimented values. At an ER of 1, the H₂ is null, indicating the complete combustion and conversion of all H₂ to water vapor. The prevailing conditions and criteria quoted for H₂ are equally applicable for CO, albeit at a different magnitude.

b) Effect of ER on CO₂ and CH₄

The molar percentage of CO₂ is observed to increase with an increase in ER, similar to the trend established by Altafini et al. (2003). In reality, the trend of CO₂ could be correlated with the trend opposite to that for CO. A decrease in concentration of CO₂ indicates better gasification efficiency (Zainal et al. 2002). The modeled results for charred CNS indicated negative values of CO₂ at lower ERs, which is quite unrealistic. Ruggiero and Manfrida’s (1999) statement could be applied to this condition also. However, after a critical analysis of the modeling, it was decided to check the trend of CO₂ for other established fuels, for which modeling results had already been reported. Accordingly, the devised model was applied to casuarina wood (CH₁.₄₄O₀.₆₅), whose proximate and ultimate analyses are listed in Tables 6 and 7. The trend of CO₂ for casuarina wood is observed to be increasing and positive for all ERs (Fig. 11). Other constituents of the global reaction [Eq.1] remain the same, the C/H and C/O ratios of charred CNS are observed to be 2.18 and 2.08 times greater than the C/H and C/O ratios of casuarina. Hence it was inferred that the C/H ratio to some extent affects the model output at lower ERs. This was confirmed by increasing the H/C ratio on the reactant side in the global reaction [Eq.1] by increasing the fuel moisture content and analyzing the molar percentage of CO₂ at different ERs. At a moisture level of 40%, the CO₂ yields are all observed to be positive (Fig. 20), indicating the effect of C/H ratio on modeling at lower ERs. Though the values depicted by the model at lower ERs per se would not be relevant to gasification, the cause for it had been reported. It was found that the model predicts well the CO₂ concentration at higher ERs and that pertaining to gasification.

| Table 6: Proximate analysis of casuarina wood (wt% on an as-received basis) |
|-----------------|-----------------|
| Moisture        | 12.5            |
| Volatile Matter | 67.5            |
| Fixed Carbon    | 18              |
| Ash             | 2               |

| Table 7: Ultimate analysis of casuarina wood (wt% on an as-received basis) |
|-----------------|-----------------|
| Carbon          | 48.5            |
| Hydrogen        | 6.06            |
| Oxygen          | 43.3            |
| Nitrogen        | 2.14            |
Mansaray et al. (1999) inferred that increasing the ER results in a decrease in concentrations of methane and other light hydrocarbons, which have relatively high heating values. The model results validate the claim that CH₄ concentration decreases with increasing ER. Conversely, the prediction on CH₄ made by the model is lower than the actual ones by a large margin. Pellegrini and Oliveira Jr. (2007) experienced similar differences in CH₄ predicted by the model and experiment and referred to the cause as a result of the sudden cessation of gasification reactions at the bottom of the reactor. This cessation is a consequence of the temperatures at the bottom, which are too low to start up the reactions. In an equilibrium model, it is assumed that all reactions achieve a steady-state condition; thus no kinetic effects (such as sudden cessation) are considered. To overcome the differences in mole fractions, a fixed CH₄ molar correction needs to be adopted, as reported by Fock and Thomsen (2000).

c) Effect of ER on N₂ and H₂O

Gas analysis was not carried out on a dry basis for want of facilities. Hence the remainder of the measured components, viz., CO, CO₂, H₂, O₂ and CH₄ is assumed to be mix of N₂ and H₂O. The devised model predicted the yield of N₂ and H₂O individually and the molar concentrations of these constituents were combined as a mix to compare with the experimental values. Generally nitrogen formation could be attributed to the N₂ from fuel and N₂ from air. With the nitrogen in fuel remaining constant, the increase in ER increases the N₂ supplied to the system, and hence the higher yield of N₂ at higher ERs. While the majority of researchers had observed a similar pattern, Zainal et al. (2002) reported a decreasing trend of N₂ with an increase in ER, which is not quite possible. Water in the product gas also increases with an increase in ER for the ascendancy of the oxidation reaction of H₂ at higher ERs. The H₂O and N₂ mix predicted by the model correlates well with the experimental values.

Effect of Temperature on Molar Concentration

Jayah et al. (2003) observed that a lower temperature reduces the reaction rate and thereby the conversion efficiency. Altafini et al. (2003) stated that equilibrium models are especially good at high temperatures where the reaction temperatures are above 1500 K. The temperatures generally assumed for equilibrium modeling by different researchers were not coherent and were predominantly modeled between 800 – 1000 K. Moreover the zone of the gasifier having this temperature was also not reported. Prins et al. (2006) remarked that for fluidized bed gasifiers, the average bed temperature can be used as the process temperature, whereas for downdraft gasifiers, the outlet temperature at the throat exit should be used. The temperature measured in the throat zone [T₄] of the gasifier, using charred CNS, was about 1100°C. Dogru et al. (2002) reported a throat temperature of 1015°C using hazelnut shells.

However keeping in mind the veracity of other work in the literature, the model was framed for a wide range of temperatures from 800 K to 1500 K and compared with the experimental results obtained at 1373K (Figs 12 to 16). It should be noted that the deviation in gas composition within the temperature range selected was not very significant. Both exothermic and endothermic reactions occurred in the gasification system. Based on Le Chatelier’s principle, it is understood that higher reaction temperatures favor the reactants in exothermic reactions while they favor the products in endothermic reactions. Methane formed in the gasifier at high temperatures underwent endothermic reactions with the already formed water vapor and was converted into CO, CO₂ and H₂. Hence the yield of CH₄ decreased at higher temperatures.

\[
\text{CH}_4 + \text{H}_2\text{O} = \text{CO} + 3\text{H}_2
\]

\[
\text{CH}_4 + 2\text{H}_2\text{O} = \text{CO}_2 + 4\text{H}_2
\]

The yield of H₂ in the above reactions is three to four times that of formation of CO or CO₂. Hence the increase in molar concentration of H₂ is significantly higher than the increase observed in CO or CO₂. Hence for gasification systems aimed towards generation of H₂, maintaining of higher reaction temperature would be a remarkable advantage.

The endothermic reactions used the water vapor generated in the preliminary gasification process, as a result of which the yield of the N₂ + H₂O mix was lower at higher temperatures. Since N₂ forms the majority of the mix, the dip caused by reduction in moisture was almost insignificant. Precisely, it could be inferred that the cause for the drop in methane at higher temperatures and the increase in CO and H₂ was due to the utilization of methane in endothermic reactions (rather a reforming process indeed). The trend obtained matches well with the style established by other researchers for other fuels (Türk et al., 1998; Luiz & Silvio, 2007; Schuster et al., 2001; Philippe and Raphael, 2002; Yang, 2005; Sergrapagna & Ajmal, 1997; Natarajan et al., 1988; Hos and Groeneveld, 1987).
Figure 12: Yield of H₂ at different bed T vs ER

Figure 13: Yield of CO at different bed T vs ER

Figure 14: Yield of CO₂ at different bed T vs ER

Figure 15: Yield of CH₄ at different bed T vs ER

Figure 16: Yield of N₂+ H₂O at different bed T vs ER

Legends indicating temperatures

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Effect of Moisture on Molar Concentration

Moisture content is one of the most significant properties of any biomass that are known to affect the gasification process. Hos and Groeneveld (1987) detailed the effect of moisture content of the feed material on the composition of the product gas. Bridgewater et al. (1986) cited that moisture constraints for any gasifier fuel are dependent on type of gasifier used. Higher values are possible in updraft systems, but the upper limit acceptable for a downdraft reactor is generally considered to be around 40% on a dry basis. Reed and Das (1988) specified that moisture content of feedstock should be below 33% (d.b.) for generating a burnable, good quality gas, while moisture contents higher than 67% (d.b.) make the product gas too lean for ignition. McKendry (2002) inferred that moisture content above about 30% makes ignition difficult and reduces the CV of the product gas due to the need to evaporate the additional moisture before combustion/gasification can occur. With this background, the present model was analyzed for the impact of moisture content on gas composition and the results were compared with the experimental values. Moisture content in a biomass could be either intrinsic (inherent) or extrinsic (influenced by weather/handling). The moisture in the producer gas is an amalgamation of the moisture sourced from substoichiometric air (relative humidity), water vapor formed due to oxidation of hydrogen in fuel and the intrinsic and extrinsic moisture associated with fuel. Among these contributing factors, the first two absorb only the sensible form of heat for superheating the moisture, while the last component absorbs sensible, latent and superheat for converting the moisture into a superheated vapor form. The higher the fuel moisture content, the higher is the heat absorbed by the moisture, paving the way for a reduced reaction temperature and associated incomplete cracking of the hydrocarbons released from the pyrolysis zone.

Increased levels of moisture and the presence of CO at lower ERs produce more H₂ and CO₂ by the water gas shift reaction [Eq. 6]. The increased H₂ content of the gas produces more CH₄ by direct hydrogenation. Pellegrini and Oliveira Jr. (2007) inferred that though more H₂ is formed with increased moisture, in order to maintain the process, more energy must be supplied, so exothermic reactions are favored, which promotes CO₂ formation. The gain in H₂ and CH₄ of the product gas does not however compensate for the loss of energy due to the reduced CO content of the gas and therefore gives a product gas with a lower CV (McKendry, 2002). Similar results on the effect of moisture content have also been reported for conventional gasifiers (Bhattacharya et al., 1999; Parikh, 1985; Walawender et al., 1987). The model revealed similar trends for CNSC and the experimental values were observed to follow suit predicted by the model (Figs 17-21).

Consolidation

A comparison of the equilibrium modeling results vs the experimental values obtained upon gasification of CNSC is presented in Fig 22. The deviation observed is very minimal on the ER applicable to gasification, i.e., 0.15 to 0.4 for all constituents except CO₂. However the experimental values of CO₂ can be observed to match the modeled outputs well at ER > 0.3.
Figure 19: Molar % of CH₄ with varied moisture vs ER

Figure 20: Molar % of CO₂ with varied moisture vs ER

Figure 21: Molar % of N₂ + H₂O with varied moisture vs ER

Effective moisture on gas composition
Legends indicate % Moisture Content
CONCLUSIONS

- The concept of equilibrium modeling applies well for prediction of the effect of ER, MC and RT in gasification of CNSC.
- ER is observed to have a large impact on the composition of the producer gas. The model predicts well the trend for H2, CO, N2 and H20. For CO2, the model depicts negative values, possibly, owing to the effect of the carbon-hydrogen ratio, which had been proven by application of the same model for gasification of casuarina wood.
- The contribution of reaction temperature is significant for hydrogen yield. The composition of H2 at an ER of 0.3 maintained at a RT of 800 K is 10.5 %, while at 900 K it is 14 %, an increase of 33 %. However, at the same ER, the yield is not so pronounced between 1100 and 1500 K. Higher temperatures favor the formation of CO and minimize the yield of CH4. For temperatures higher than 1200 K, CH4 generation is practically nil.
- The model does predict the effect of MC in a relevant manner. The CH4 content increases with moisture content. At an ER of 0.3 and with a fixed RT of 1373 K, the yield of CH4 is 0.14 % at 10 % moisture, while it is 0.25 % at 50 % moisture, an 80 % increase.
- The model predicts an unrealistic gas composition at ERs less than 0.15, due to pyrolysis, for all constituents of producer gas. For ER pertaining to gasification and beyond that, the model’s prediction is coherent with experimental values.

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NOMENCLATURE

\[ \Delta G^\circ \] Standard Gibbs function of formation
\[ \Delta A, \Delta B, \Delta C \] Coefficients for determining specific heat
\[ \Delta H^\circ \] Heat of formation
\[ A, B, C \text{ and Constants for the properties of the gases} \]
\[ D \] of the gases
\[ CNS \] Cashew nut shell
\[ CNSC \] Cashew nut shell char
\[ CNSL \] Cashew nut shell liquid
\[ C_p \] Specific heat
\[ C_{p(am)} \] Specific heat at arithmetic mean temperature
\[ ER \] Equivalence ratio
\[ HHV \] Higher heating value \( \text{kJ/kg} \)
\[ I \] Constant of integration
\[ J \] Constant
\[ K_1 \text{ and } K_2 \] Equilibrium constants
\[ m \] Amount of oxygen per kmol of wood
\[ MC \] Moisture content per mol of wood
\[ R \] Universal gas constant \( \text{J/mol k} \)
\[ RT \] Reaction temperature
\[ T_1 \] Reference temperature
\[ T_2 \] Reaction temperature
\[ T_{am} \] Arithmetical mean temperature
\[ w \] Amount of water per kmol of wood
\[ x_1, x_2, x_3, x_4 \] Coefficients of constituents and \( x_5 \) of the producer gas

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