An Assessment of Gasification of Lipid-Extracted Algae by Thermodynamic Simulation

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Abstract
Thermodynamic simulation of gasification of Nannochloropsis gaditana (after lipid extraction and before lipid extraction) over a wide range of temperatures (800 – 1400 °C) was performed in this study. The reactor temperature variation with respect to the O₂ to fuel ratio and the syngas composition with temperature were evaluated for gasification performance. The results showed the H₂/CO ratio for lipid-extracted algae (LEA) was very similar to the gasification behavior of raw algae (RA) before lipid extraction. Only a slight variation was observed for the lower heating value (9.2 MJ/Nm³ for RA and 8.9 MJ/Nm³ for LEA at 1000 °C) of the syngas from the two feedstocks. The H₂/CO ratio for both RA and LEA remained almost the same over the range of temperatures (800-1400 °C) under consideration. The cold gas efficiency was found between 82 % to 75 % for RA and 75 % to 69 % for LEA at 1000 °C and 1400 °C respectively. As the overall gasification performance did not degrade significantly after extraction of lipid from algae, the LEA holds remarkable potential as a gasification feedstock in a biorefinery set up.

Keywords: Gasification, Lipid-extracted algae, Thermodynamic simulation

1. Introduction
In the past decade, the earth has experienced a drastic rise in the surface temperature due to the greenhouse gas release, reaching more than 400 ppm of CO₂ concentration in the atmosphere [1]. Other than the rising sea level, this might result in serious disasters like unbearable humidity leading to human death in near future [2,3]. To mitigate the CO₂ emission from source, it is of absolute urgency to adopt a sustainable strategy to utilize the current technology infrastructure for the gradual transition from fossil fuel to renewables. The use of the available infrastructure and knowledge will result in minimal requirement for modification and cost allocation allowing the existing energy producers to switch from fossil to renewable energy sources. One such replacement for fossil fuel could be biomass which can take the advantage of utilizing state of the art technology.

Among several types of biomass, the type categorizes as third generation is considered to be the most suitable for energy production as it does not compete with food and not limited by the production rate [4]. The most promising third generation biomass is considered to be microalgae for its high production rate and lipid content making it suitable for biodiesel production [4,5]. However, the process of lipid extraction from algae and biodiesel production can incur significant cost resulting in a biodiesel price 10 times higher than the traditional fossil oil [6]. Only increasing the efficiency in extracting lipid from algae would not suffice to make it economically feasible. On the other hand, it might be possible to produce an array of products from microalgae combining several conversion methods forming a biorefinery to make the whole process economically feasible. A potential next step could be the utilization of the lipid extracted residue from algae through thermochemical pathway [7]. It was reported in several studies [7,8] that more than 90% of the carbon may remain intact in the lipid extracted residue. In some cases, the percentage of carbon even increased due to the removal of oxygen during lipid extraction [8]. This remaining carbon can very well be the source of fuel and fine chemicals if an appropriate conversion method is used. One such conversion process producing energy and chemicals efficiently leaving no residual carbon is gasification.

Being a relatively mature technology for coal utilization, the possibilities with microalgae gasification had been explored by researchers for practical applications [9–11]. While gasification alone might have some prospect, it is rather cost-effective to utilize the residue from a process which can already produce slightly more valuable product. However, only a few studies had tested the lipid extracted algae in laboratory scale [7,8,12]. This indicates the lack of data available for practical scale application to gain an in-depth understanding of large scale utilization of algae (after lipid extraction) under variable process conditions. To address this issue, this study assesses the full scale gasification plant simulation considering lipid extracted algae as feedstock comparing with pure algae gasification. The process parameters, i.e., temperature and oxygen to fuel ratio were varied to
identify the effects on the syngas composition for downstream requirements for further processing.

2. Simulation Basis

Compositions of raw and lipid-extracted *Nannochloropsis oculata* were used for thermodynamic modelling of algae gasification. The elemental compositions and higher heating values of raw algae (RA) and lipid-extracted algae (LEA), taken from the work of Barreiro et al [8], are shown in Table 1. The lipid content of the RA was 13.4%, from which 11.1% mass was during extraction by n-hexane[8].

Table 1: Elemental composition, ash content and mineral elements in wt% (dry basis), and heating value in MJ/kg (dry basis) [8]

<table>
<thead>
<tr>
<th>Element</th>
<th>Raw algae</th>
<th>Lipid-extracted algae</th>
</tr>
</thead>
<tbody>
<tr>
<td>C</td>
<td>47.6</td>
<td>43.9</td>
</tr>
<tr>
<td>H</td>
<td>7.5</td>
<td>7.1</td>
</tr>
<tr>
<td>N</td>
<td>6.9</td>
<td>7.4</td>
</tr>
<tr>
<td>S</td>
<td>0.5</td>
<td>1.1</td>
</tr>
<tr>
<td>O</td>
<td>25.1</td>
<td>26.0</td>
</tr>
<tr>
<td>Ash</td>
<td>12.4</td>
<td>14.5</td>
</tr>
<tr>
<td>Ca</td>
<td>0.50</td>
<td>0.62</td>
</tr>
<tr>
<td>Fe</td>
<td>0.62</td>
<td>0.05</td>
</tr>
<tr>
<td>K</td>
<td>1.30</td>
<td>1.56</td>
</tr>
<tr>
<td>Mg</td>
<td>0.27</td>
<td>0.32</td>
</tr>
<tr>
<td>Na</td>
<td>3.02</td>
<td>3.64</td>
</tr>
<tr>
<td>P</td>
<td>1.43</td>
<td>1.72</td>
</tr>
<tr>
<td>Si</td>
<td>&lt;0.003</td>
<td>&lt;0.003</td>
</tr>
<tr>
<td>Zn</td>
<td>0.01</td>
<td>0.01</td>
</tr>
<tr>
<td>HHV</td>
<td>23.1</td>
<td>21.3</td>
</tr>
</tbody>
</table>

Aspen Plus (Aspen Technology, Inc., Bedford, MA, USA) simulator was used for thermodynamic equilibrium calculations. RA and LEA with 10% moisture content were used as the feedstock. Typical composition of oxygen stream (95% oxygen, 3.4% argon and 1.6% nitrogen [13]) from an air separation unit (ASU) was used as the oxidant for the gasification. Redlich–Kwong–Soave equation of state with Boston–Mathias γ function (RKS-BM) was used for property calculations for the gasification [14]. For solids (algae and ash) HCALIGT and DCOALIGT models were used for enthalpy and density calculations [15].

A two-stage gasifier model, consisting of an R-YIELD and an R-GIBBS reactor, was used for the calculation. The gasifier pressure was assumed to be 10 bar. Gasifier temperature was varied from 800 °C to 1400 °C, in order to cover the whole spectrum from fluidised-beds gasifier to entrained-flow gasifier. Heat loss from the gasifier was assumed to be 4% of the lower heating value (LHV) of the feedstock [16].

3. Results and Discussion

Gasification of RA and LEA were simulated using thermodynamic simulation. For thermodynamic calculations, the temperature of the gasifier is a function of fuel composition, fuel-to-oxidant ratio in the gasifier feed, and the loss of heat to the surroundings from the gasifier. The fuel-oxidant ratios were calculated to achieve gasifier temperatures between 800 °C and 1400 °C with a 50 °C interval. Fuel-oxidant equivalence ratios (φ) for each temperature were calculated using the Eq.(1).

\[ \varphi = \frac{\text{FOR}}{(\text{FOR})_{\text{stoch}}} \tag{1} \]

where, \( \varphi \) is the fuel-oxidant equivalence ratio, \( \text{FOR} \) is the fuel-oxidant ratio on mass basis and \( (\text{FOR})_{\text{stoch}} \) is the fuel-oxidant ratio on mass basis for stoichiometric mixture.

![Fig.1: Gasification temperature versus equivalence ratio](image)

The variation in the gasifier temperature as a function of the fuel-oxidant equivalence ratio is shown in Fig. 1. All \( \varphi \) values are above 1, indicating fuel-rich conditions inside the gasifier. For 800 °C and 850 °C, the simulation showed incomplete conversion of algal biomass to gas and ash. However, at temperatures above 900 °C, complete conversion could be achieved. Therefore, a small decrease in \( \varphi \) (hence a small increase in oxidant) resulted noticeable increase in the gasifier temperature. Fig. 1 also shows that more oxidant was required for LEA to achieve the same temperature as RA gasification. This is due to the lower carbon and heat content of LEA than that of RA, caused by the extraction of lipid. As complete conversion of algae was not achieved at temperatures below 900 °C, only temperatures between 900 and 1400 °C were considered in the following sections.

Gasification is a complex process of incomplete combustion of solid feedstock. Initially, algae undergo pyrolysis to produce volatiles and char. Volatiles reacts further with the oxygen and produce combustion products, i.e., \( \text{CO}_2 \), \( \text{CO} \), \( \text{H}_2 \text{O} \), \( \text{H}_2 \), \( \text{H}_2 \text{S} \). Char also reacts with oxygen producing \( \text{CO} \) and \( \text{CO}_2 \). In addition, char participates in steam and \( \text{CO}_2 \) gasification (Boudouard reaction) and methanation reactions. There are also some gas-phase reactions.
of significant importance: water gas shift and steam-methane reforming reactions.

The equilibrium compositions of syngas generated from RA at various temperatures are shown in Fig 2(a). The mole fraction of CO and H₂O in the syngas increase as the gasifier temperature rises. Both the hydrogen and CO₂ concentrations showed a decreasing trend with increasing temperature. Higher reaction temperatures favour both reverse water-gas shift and reverse Boudouard reactions, and therefore result an increase in the CO and H₂O concentrations at the expense of hydrogen and CO₂. Lower gasification temperatures showed presence of methane in the syngas. Methane mole fractions were 0.016 and 0.002 at 900 and 1000 °C, respectively. Above 1000 °C, presence of methane in the syngas was insignificant. Lower temperatures and higher pressures favour the formation of methane. However, as the temperature increases, methane concentration diminishes due to consumption in combustion and steam-reforming reactions. The presence of methane in the syngas is desirable if the aim is to produce high-Btu syngas. On the other hand, if the syngas is produced as an intermediate feedstock for chemical production, methane will act as an inert. In such cases, methane formation should be minimized. Sulphur present in the algae found to be converted to H₂S, the dominant sulphur containing species, the mole fraction of which remained almost constant for all temperatures. Trace amount of SO₂ and NH₃ formation was also found.

The compositions of syngas produced from LEA are shown in Fig. 2(b). The trends observed in the syngas composition with temperature variation are somehow similar. However, the syngas from LEA found to contain slightly lower CO mole fraction, due to the variation in elemental composition resulting from lipid extraction. Due to removal of some carbon and hydrogen, LEA had slightly lower LHV than RA (see Table 1). Also the ash content increased after lipid extraction. This required more energy to reach the same temperature as RA to heat up the excess ash as ballast in the gasifier at the expense of producing CO₂ from CO. In both Fig. 2(a) and 2(b), YI indicates the mole fraction of the inerts in the syngas, which is a combination of argon and nitrogen. These components were introduced to the system via the oxidant stream from ASU. The fuel nitrogen also contributed to some extent to the inert build-up in the syngas.

H₂-to-CO ratio is an important parameter used for the evaluation of gasifier performance. The H₂-to-CO molar ratio as a function of temperature is shown in Fig. 3. H₂-to-CO ratio for the RA syngas varied from 1.01 to 0.81 for 900 °C to 1400 °C. For LEA, H₂-to-CO ratio was slightly higher at 900 °C, 1.04, than that of RA. LEA containing less carbon than the parent algae was the reason behind this variation. At 1400 °C, the ratio for LEA was similar to that of RA. It must be noted that the syngas is rich in CO, and hence water-gas shift is essential to increase the H₂-to-CO ratio, if it is used for production of chemicals, such as methanol or dimethyl ether. If the syngas is used in a gas turbine, H₂-to-CO ratio is of lesser importance. Both H₂ and CO have similar HHV on volume basis. However, the H₂-to-CO ratio of the syngas fed to the gas turbine can have an impact on the fugitive emissions from the gas turbine [17].
However, the elemental composition of different species is not likely to vary much. With extraction of lipid the carbon and hydrogen contents are most likely to go down. Therefore, the observations made here are likely to hold for most algae species.

Gasification of RA would yield syngas only. On the other hand, RA to LEA to syngas pathway would have two products: lipid from extraction stage followed by syngas from the gasification of LEA. Carbon and hydrogen balance on the RA and LEA provides an approximate composition of the extracted lipid from *Nannochloropsis gaditana*. The heating value then can be calculated from correlations such as Boie’s formula[8]. By performing carbon and hydrogen balance using the elemental composition of RA and LEA, and the amount of lipid extracted, it can be assumed that the lipid has an apparent formula of C$_{56}$H$_{10}$O$_{25.75}$, with a calculated HHV of 38.3 MJ/kg. To make a simple comparison between two pathways (RA to syngas, and RA to LEA and lipid followed by LEA to syngas), the heat content of the syngas from LEA times syngas yield were added to the heat content of the lipid times the lipid yield. This value was then compared with the heat content of syngas from RA multiplied by the syngas yield. It must be noted that none of the products are final products and require further processing to get a final product, such as biodiesel or methanol. However, this comparison will still provide a comparative idea on these two pathways. Considering same quantity of RA as the feedstock for both pathways, it was found that an energy penalty of 18% is realized if the LEA is gasified compared to the gasification of RA. However, LEA is a waste of the lipid-extraction process of biodiesel production from algal oil. Hence, gasification of LEA holds a prospect as a feedstock for gasification.

Fig. 3: Hydrogen-to-CO molar ratio

$\eta_{\text{cold}}$ for RA and LEA syngas are shown in Fig. 4. There is a noticeable difference in the $\eta_{\text{cold}}$ for RA and LEA syngas. This is due to the lower carbon and hydrogen content in the LEA feedstock. $\eta_{\text{cold}}$ for RA and LEA syngas was 84% and 77%, respectively, at a gasification temperature of 900 °C. As the gasification temperature was increased to 1400 °C, these values reduced to 75% and 69%, respectively.

A comparison of the gasifier parameters are therefore presented in Table 2, to compare these two most like scenarios: fluidised-bed gasifier operating at 1000 °C and entrained-flow gasifier operating at 1400 °C. As shown in Table 2, the gasifier running at 1000 °C for both RA and LEA has higher efficiency than the gasifier running at 1400 °C. The H$_2$:CO ratio at lower temperature gasifier is slightly favourable for chemical production from syngas.

Table 2: Performance of gasifiers at 1000 °C and 1400 °C

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Gasification at 1000 °C</th>
<th>Gasification at 1400 °C</th>
</tr>
</thead>
<tbody>
<tr>
<td>H$_2$:CO molar ratio</td>
<td>0.99 (RA); 1.01 (LEA)</td>
<td>0.81 (RA) and LEA</td>
</tr>
<tr>
<td>LHV (MJ/Nm$^3$)</td>
<td>9.2 (RA); 8.9 (LEA)</td>
<td>8.40 (RA); 8.12 (LEA)</td>
</tr>
<tr>
<td>Cold gas efficiency</td>
<td>0.82 (RA); 0.75 (RA)</td>
<td>0.75 (LEA); 0.69 (LEA)</td>
</tr>
</tbody>
</table>

However, downstream water-gas shift reactors are required to shift the ratio to the desired level. The thermodynamic calculations, therefore, suggest that the gasification of algae, whether raw or lipid-extracted, may be operated at lower temperatures in fluidised-bed or, in a catalytic entrained flow reactor. It must be kept in mind that the thermodynamic calculations calculate the equilibrium compositions only and might not be able to predict the formation and retention of tar products at fluidised-bed conditions. The algae under consideration contains high amount of Na and K with low amount of Ca in ash (see Table 1). This quantity becomes even higher after lipid extraction. It is to be noted that the Si content is very low for both before RA and LEA.
Si inhibits the catalytic activity if its quantity is higher than the combined amount of Na and K [11,18,19] as they might take part in forming silicates. For the current species under study, the Si content was significantly lower than Na and K, making it highly suitable for catalytic gasification. Recent studies on alkali catalyzed gasification showed that as the total alkali (Na+K) to carbon ratio in char increases, the reactivity increases drastically during low temperature gasification (< 850 °C) [20]. On the other hand, at higher temperatures, part of the alkali is evaporated to exert other activities such as tar destruction and soot reduction [21,22]. In all scenarios, the lipid extracted algae species (Nannochloropsis gaditana) stands out as a suitable candidate for catalytic gasification. However, as the assessment of the catalytic activity is beyond the scope of this study, this is a topic for future research.

4. Conclusion

Gasification of Nannochloropsis gaditana, before lipid extraction (RA) and after lipid extraction (LEA), between 800 and 1400 °C was simulated using thermodynamic model in Aspen Plus. Temperatures above 900 °C were able to convert algae completely to syngas and residual ash. The RA syngas found to have slightly more CO than that of LEA syngas. The H2-to-CO ratios for both syngases had almost identical values. However, the RA syngas found to have higher heat content and cold gas efficiency than LEA syngas. Simple energy balance on the RA to syngas and RA to LEA to syngas pathways indicate that there is a 18% energy penalty for the RA to LEA to syngas pathway. However, LEA is a waste of the lipid extraction stage with moderate carbon content. A suitable method of extraction of waste of the lipid extraction stage with moderate carbon content. A suitable method of extraction of energy conversion.

5. References