Energy-Efficient Production of Cassava-Based Bio-Ethanol

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Abstract

Fuel ethanol is an important renewable and sustainable fuel, produced in China by fermentation of mostly corn, wheat and cassava feedstock. Fermentation produces an ethanol-lean broth (10 to 12 vol%). Ethanol is recovered by distillation, followed by a molecular sieve drying beyond the azeotropic point. The distillation and molecular sieve operations consume most of the total energy used, with the steam consumption currently being ~1.8 kg/kg ethanol, including 0.5 kg/kg ethanol in the final molecular sieve stage during regeneration. The objectives of the paper are fourfold: 1) firstly to study the distillation process of a large-scale cassava-based fuel ethanol production (200,000 tons per year), by field measurements and by using an Aspen Plus V8.2 simulation, with and without energy integration of condensers and reboilers, resulting in a distillation steam consumption of ~1.3 kg/kg ethanol; 2) secondly, to examine the effects of using Very High Gravity (VHG) fermentation of cassava meal mash. By using VHG fermentation, the ethanol concentration in the fermenter broth is significantly increased, to about 19 vol% (15.4 wt%). The steam consumption is then reduced to ~0.94 kg/kg, representing a considerable saving in comparison with the current operation. Applying VHG fermentation needs minor additional investment, rapidly recovered through the energy savings and the smaller size of equipment; 3) thirdly, to assess the application of a hybrid operation, where pervaporation will be used to selectively and continuously remove ethanol from the fermenter broth, thus slightly increasing the fermentation yield by reducing the risk of ethanol inhibition, whilst producing an ethanol-rich permeate (about 30 wt%); and finally 4) to demonstrate that the production cost of cassava-based ethanol can substantially be reduced by applying the proposed improvements.

Keywords

Bio-Ethanol, Hybrid Process, VHG, Membranes, Energy Optimization, Simulation

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1. Introduction

Bio-ethanol, as a clean and renewable fuel, is gaining increasing attention. Biochemically produced by fermentation of an aqueous broth of different raw materials (corn, wheat, cassava, sugar cane...), it is purified by distillation till the water/ethanol azeotropic mix of ~95 wt% ethanol. A further removal of water to obtain >99.5 wt% ethanol is commonly performed using molecular sieves.

Distillation is an energy-intensive separation process, contributing significantly to the production cost of bio-ethanol. Over the past years, a lot of research studied modified production pathways to reduce the cost of bio-ethanol, with hybrid processes and Very High Gravity (VHG) fermentation as the most important recommendations. To quantify the impact of possible improvements, and compare production cost of different alternatives, the present paper examined the 200,000 ton/year cassava-based production facility of Cofco (China) [1]-[3]. Different improvements will be assessed using literature data and own experimental results. All data will be imported into a simulation of the distillation train by using Aspen Plus V 8.2 software. The simulation and field results of the current operation will be compared. Simulations will also be performed when using VHG or pervaporation. All findings will finally be used into a tentative economic assessment.

2. The Significant Potential of Bio-Ethanol

With the globalization of the increasing demand for energy, its shortage is a growing worldwide problem. Bio-ethanol is considered as the alternative renewable fuel with the largest potential to replace fossil fuels. Its world production increased from 50,000 million liters in 2007 to over 100,000 million liters in 2012 [4]. The main reasons to develop bio-ethanol are diverse and linked to its possible uses 1) as a favorable carbon-neutral energy source, hence reducing CO₂ emissions [5]; 2) as octane enhancer in unleaded petrol; and 3) as an oxygenated mix for a cleaner combustion of petrol, reducing tailpipe emissions and improving the ambient air quality.

In addition to the well-described fermentations of food-competing corn, wheat and sugarcane, new resources are being investigated and extensively used such as sweet sorghum and cassava, not competing with the food chain and suitable for growing in dryland conditions.

Together with bio-ethanol from carbohydrate-based feedstock such as e.g. corn, wheat, cassava (first generation), cellulose biomass (second generation) and algal biomass (third generation) are also considered. Recent references about the different generations of bio-ethanol were examined and reveal that the first generation bio-ethanol production will remain the most important production process in the immediate future [6]-[8], whilst second and third generation processes still need further development on laboratory and pilot scale to remove major drawbacks, and increase the bio-ethanol yield per input biomass weight [9]-[13].

3. The Bio-Ethanol Production Processes

3.1. Schematics of Current and Possible Hybrid Operations

The key-parts of the common bio-ethanol production, from pretreatment to dehydration and residue treatments, are illustrated in Figure 1(a), simultaneously indicating the potential use of 2 membrane techniques, being microfiltration and pervaporation (Figure 1(b)). Other membrane techniques can also be introduced (nanofiltration and reverse osmosis), but currently investigated mainly for feedstock pre-treatment and for treatment of process residues [13].

To separate the ethanol from the water phase of the broth, distillation is used until the point of the azeotropic mixture, where after residual water is removed from ethanol by molecular sieves. Distillation and molecular sieve regeneration are energy intensive processes, contributing significantly to the production cost of bio-ethanol, representing nearly 50% of the entire energy consumption [13]-[15]. The substitution of molecular sieves by membrane permeation of H₂O was experimentally investigated and confirmed as a solution with high potential in a separate paper [13]. Whereas molecular sieves require ~0.5 kg steam/kg bio-ethanol during regeneration, steam is not required when using membranes, thus contributing to process energy efficiency.

3.2. Illustration of the Cofco Cassava-Based Distillation Process

The flow sheet of the current distillation process of a large scale cassava-based fuel ethanol production (Cofco,
Figure 1. Schematics of the processes for sugar-based bio-ethanol production (a) as currently applied; (b) with potential membrane applications (DDGS: distillers dried grains with solubles; 1 + 2 + 3: cellulosic materials; 2 + 3: starch or carbohydrate materials).

200,000 tpa) is shown in Figure 2. It consists of a sequence of distillation columns, heat exchangers, pumps and a flash tank. The feed stream enters the crude column C1501. The distillate from C1501 is fed to column C1502. The side stream 4 is fed to the stripping section C1503-1 of the second column. C1503-2 is the rectifying section of the second column. C1503-1 has no condenser and C1503-2 has no reboiler. The bottom stream 10 enters C1502. Stream 15 is the ethanol-rich phase. The distillate from C1502 is separated via the flash tank V1504, whose gas and liquid streams are both fed to C1503. The bottom streams (5, 9, 16) are directed to the wastewater treatment unit and are ethanol-lean (≤0.05 wt% ethanol). Finally the distillate from C1503-2 is further dewatered using molecular sieve absorption to the finished product of 99.7 wt% ethanol. Energy-integration is indicated as streams EI1 and EI2. As can be seen from Table 1, the pressure is progressively increased along the distillation train. It is hence possible to recover condenser heat to feed the reboilers of previous columns. The heat duty difference between the top of C1502 and the bottom of its neighbor C1503 is sufficient to modify the system into a combined reboiler/condenser distillation: the top high pressure condensate of condenser C1502 is used as the heat source for the reboiler of C1501. In this energy recycling scheme and accounting for a required thermal driving force in the reboilers, the remaining steam consumption would be mainly due to the bottom reboiler of C1502.
Figure 2. The distillation process of Cofco.
The mass balance of the Cofco cassava-based process, based upon a single fermenter, is shown in Figure 3. The Cofco process uses 6 fermenters in parallel, each fermenter having a volume of 3025 m$^3$, i.e. ~3025 ton. 865 ton of cassava flour is used per batch of 65 hrs. The production is based upon a 70% cassava starch-content and 90% starch conversion. The amount of starch converted is 549 ton. The fermentation reaction can be represented as:

$$C_6H_{12}O_6 \rightarrow 2C_2H_5OH + 2CO_2$$ (1)

All fermenter broths are collected and dealt with in a single distillation train, working in a continuous mode. The recycle to the fermenter is applied to keep the enzyme and yeast balance at the required level.

The composition of the cassava fermentation broth contains mostly water (81.4 wt%), ethanol (11.6 wt%), other aliphatic alcohols (~0.13 wt%), esters and acids (0.5 wt%), glycerol (0.4 wt%) and about 6 wt% of high boiling point components. The continuous feed rate to the distillation plant is 220,000 kg/hr. Referring to Figure 2 and the different key-parts of the distillation process, the number of theoretical trays and operating pressure of each column are taken from field data, and shown in Table 1.

### 3.3. Comparison of Cofco Operational Data with Chemical Engineering Simulations

The software Aspen Plus V8.2 was used. Due to the presence of the highly polar components, ethanol and glycerol,
both the non-random two liquid (NRTL) and universal quasi-chemical (UNIQUAC) thermodynamic/activity models were used to predict the activity coefficients of the components in a liquid phase [16]. Some unavailable interaction parameter coefficients were estimated using the UNIQUAC liquid-liquid equilibrium module. Simulation results towards operating temperatures in the different columns without internal energy recovery were compared with the real values of Table 1. The comparison of Table 2 shows very fair agreement.

In the non-ideal operation mode, i.e. without internal energy re-use, the total reboiler duty amounts to:

$$Q_{\text{ Reb}} \text{ (kW)} = 9000 + 18222 + 11604 = 38826 \text{ kW} \quad (2)$$

A conservative average enthalpy of steam is taken at 2200 kJ/kg steam. For a total production of 25,375 kg/hr of pure ethanol, the steam requirements are:

$$\frac{38826 \times 3600}{25375 \times 2200} = 2.50 \text{ kg steam/kg bio-ethanol} \quad (3)$$

In the integrated heat-duty operation mode, the top high pressure condensate of condenser C1502 is used as the heat source for the reboiler of C1503, and top high pressure condensate of C1503 is used as the heat source for the reboiler of C1501. In this energy recycling scheme and accounting for a required thermal driving force in the reboilers, the remaining steam consumption would be due to the bottom reboiler of C1502.

The total reboiler heat duty of the heat-integrated distillation train will be

$$Q_{\text{ Reb}} \text{ (kW)} = 18222 \text{ kW} \quad (4)$$

At an average available enthalpy of 2300 kJ/kg of steam, the energy consumption of the integrated reboiler-condenser solution is

$$\frac{18222 \times 3600}{25375 \times 2200} = 1.18 \text{ kg steam/kg bio-ethanol} \quad (5)$$

In the current Cofco operation, this reboiler-condenser integration is applied, with steam duties cited as ~1.3 kg steam/kg bio-ethanol.

Aspen-simulations and field data are in very fair agreement. The steam duty of ~1.3 kg steam/kg bio-ethanol can hence be considered as the energy requirement of the heat duty integrated ethanol-H2O distillation.

**4. Potential Improvements of the Basic Production Scheme**

**4.1. Very High Gravity Fermentation (VHG)**

In traditional starch fermentation, *Saccharomyces cerevisiae* is used as yeast, however with the drawback of inhibition at ethanol concentrations in excess of ~12 vol%. The use of specific bacteria such as *Zymomonas mobilis*, with higher inhibition levels, opened the way to high concentration fermentation, referred to as very high gravity (VHG) fermentation.

The total fermentation yield can be raised by application of VHG. This technique has drawn a lot of attention

<table>
<thead>
<tr>
<th></th>
<th>C1501</th>
<th>C1503</th>
<th>C1502</th>
</tr>
</thead>
<tbody>
<tr>
<td>Top temperature/°C</td>
<td>56.10</td>
<td>53.99</td>
<td>95.60</td>
</tr>
<tr>
<td>Bottom temperature/°C</td>
<td>82.70</td>
<td>82.16</td>
<td>123.30</td>
</tr>
<tr>
<td>Ethanol concentration of side stream (wt%)</td>
<td>16</td>
<td>16.9</td>
<td>16.9</td>
</tr>
<tr>
<td>Final Ethanol concentration (wt%)</td>
<td>93-94</td>
<td>93.2</td>
<td></td>
</tr>
<tr>
<td>Heat duty (kW)</td>
<td>Reboiler 9000</td>
<td>11,604</td>
<td>18,222</td>
</tr>
<tr>
<td></td>
<td>Condenser 5694</td>
<td>11,575</td>
<td>16,944</td>
</tr>
</tbody>
</table>
over the past few years [17]-[23]. In VHG, very high concentrations of sugars (>250 g/l) are loaded into the fermenter compared to classical fermentation. Much higher concentrations of ethanol can be maintained in the fermenter by application of this technique, and ethanol fractions of 23.8 vol% (19 wt%) have been reached [22]. Besides this advantage, a lower risk of contamination and a saving of fermenter space can be realized by using VHG fermentation. At these high ethanol concentrations, the yeasts suffer from extremely harsh conditions in the fermenter, the most important two extreme conditions being the high osmotic pressure and toxic effects of ethanol. A lot of research has been done the past few years to understand these extreme conditions and to anticipate on them. Research has been focusing on the micro-organisms themselves, added nutrients and osmoprotectants. As a result, it became clear that the fermentation rate can be raised [24] by addition of ergosterol, oleic acid or polysaccharides, whilst, higher ethanol concentrations can also be achieved when ureum, yeast-extracts or ammonia salts are added [18]. The major drawback of applying VHG fermentation is the high viscosity in the fermenter as a result of the high sugar concentrations. This high viscosity hinders the solid/liquid separation and the heat exchange during the distillation. To solve this problem Wang et al. suggest using pectine-degrading enzymes or other enzymes like xylase, cellulase and xylanase to lower the viscosity [19].

The previous simulations of the basic distillation train (Section 3.2) account for 11.6 wt% ethanol. If the ethanol content of the broth can be increased, the required heat duty of the distillation can be reduced due to the reduction of H2O content of the broth.

Although VHG results up to 23.8 vol% have been reported [22], we conservatively examined situations between 15 and 19 vol% (12.1 and 15.2 wt%, respectively), at a constant production of 25,375 kg/hr of ethanol.

With VHG at 19 vol%, the minimum reboiler duty is obtained. If again the condenser heat of C1502 recycled to the reboilers of the lower-pressure column, the required heat duty is reduced to

\[
Q_{REB} (\text{kW}) = 15162 \text{ kW}
\]

At an average heat supply of 2300 kJ/kg of steam, the steam requirement is reduced to

\[
\frac{15162 \times 3600}{25375 \times 2300} = 0.94 \text{ kg steam/kg bio-ethanol}
\]

The comparison with the energy optimization by energy-pinch alone demonstrates that a further reduction in steam requirement of ~0.3 kg steam/kg bio-ethanol is achieved.

### 4.2. Energy Optimization by Integrating Partial Pervaporation

#### 4.2.1. Pervaporation Experiments

A hybrid fermentation-distillation mode by integrating pervaporation, could reduce the risk of fermenter ethanol inhibition, whilst also slightly reducing the heat duty of the distillation by-passing the first distillation column with this ethanol-H2O stream at higher ethanol concentration, thus reducing the heat duty of the first column.

To complete a literature review as illustrated in Table 3, a confirmatory set of experiments was carried out by the authors using a pervaporation (hydrophobic) membrane of Pervatech PDMS, in a Sulzer Chemtec (Lab Test Cell Unit). Experiments were carried out with a ~12 vol% ethanol-H2O feed, at temperatures of 37°C, 45°C, 55°C and 61°C. The ethanol concentration in the permeate was measured by calibrated Abbe-Model 2A refractometer. The pervaporate vacuum was maintained constant at 1 mbar using a RVS vacuum pump (Edwards High Vacuum International). To limit mass balance errors, the permeate was fully condensed using liquid N2. The layout of the experimental rig is illustrated in Figure 4. Each experiment was repeated at least three times and results were averaged: based on the relative error of each measurement (permeate flow, membrane surface area, and ethanol concentrations), the experimental error of reported performance factors was below 5%. Each run (at a given temperature) was operated for 3 to 6 hrs. Average results are used, and detailed results are reported elsewhere [25].

Permeate flux and ethanol wt% are presented in Figure 5(a) and Figure 5(b). It is clear that the mobility of ethanol increases at a higher temperature, thus also increasing the obtained flux without negative influence on the membrane selectivity.

A summary of design values is given in Table 4.

From the presented data it is clear that the permeate flux significantly increases as the temperature increases. From the observations of ethanol concentration in the permeate, a higher feed temperature does not result in a
Table 3. Literature review of applying pervaporation in bio-ethanol production.

<table>
<thead>
<tr>
<th>Reference</th>
<th>Objectives</th>
<th>Main Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>[26]</td>
<td>Membrane technology for bio-ethanol production</td>
<td>- A highly selective and energy-saving separation process, with ~15% lower costs in comparison to the batch process.</td>
</tr>
<tr>
<td>[27]</td>
<td>Significant improvement of bio-ethanol recovery using a pervaporation (silicone rubber-coated membrane)</td>
<td>- The recovered ethanol concentration in the permeate was 67 wt%, and the amount of recovered ethanol from the broth was more than 10 times higher than when using a noncoated membrane.</td>
</tr>
<tr>
<td>[28]</td>
<td>Pervaporative separation of bio-ethanol using a polydimethylsiloxane/polyetherimide composite hollow-fiber membrane</td>
<td>- Possible but dependent on the feed concentration and operating temperature for ethanol concentrations of 1% - 10%; - Relatively good pervaporation performance with a total flux of 231 - 252 g/m²h.</td>
</tr>
<tr>
<td>[29]</td>
<td>Enhanced bio-ethanol production by pervaporation using a PDMS membrane bioreactor</td>
<td>- Increase of cell density, decreasing ethanol inhibition, improved productivity and yield, and resumption of clean and concentrated ethanol; - Ethanol productivity increased by 26.83% over conventional batch fermentation; - Ethanol concentration in permeated side was 6 to 7 times higher than that of the broth.</td>
</tr>
<tr>
<td>[30]</td>
<td>Bio-ethanol recovery using the pervaporation separation technique</td>
<td>- Zeolite 13× filled CA membrane has the better pervaporation performance; - Application of CA membranes in industrial scale pervaporation units may be feasible for separation of ethanol/water mixtures; - Bio-ethanol can be produced economically (using distillation and pervaporation by hybrid systems).</td>
</tr>
<tr>
<td>[31]</td>
<td>Membrane process opportunities and challenges in the bio-ethanol industry</td>
<td>- As a highly selective and energy-saving unit operation, membranes have a great potential in the bio-ethanol industry, both for starch-based and 2nd generation technology of bio-ethanol.</td>
</tr>
<tr>
<td>[32]</td>
<td>Membrane technologies for biorefining and bioenergy production</td>
<td>- Pervaporation membrane bioreactors (PVMBRs) and membrane distillation bioreactors (MDBRs) lead to higher ethanol productivity and minimize the inhibition; - Membrane hybrid processes further improve biorefining and bioenergy production by decreasing energy consumption, reducing the number of processing steps, and producing high quality final products.</td>
</tr>
<tr>
<td>[33]</td>
<td>Lignocellulosic ethanol through polydimethylsiloxane (PDMS) membrane</td>
<td>- Yeasts, solid particles, and salts can increase ethanol flux and selectivity; - Integrated process can effectively eliminate product inhibition, improve ethanol productivity, and enhance the glucose conversion rate.</td>
</tr>
<tr>
<td>[34]</td>
<td>Energy demand of biofuel production applying distillation and/or pervaporation</td>
<td>- Multi-stage pervaporation is needed to reach the fuel-grade quality of the bio-ethanol.</td>
</tr>
</tbody>
</table>

Figure 4. The experimental pervaporation rig.
Figure 5. Permeate characteristics as function of operating time (■: 37°C; ●: 45°C; ▲: 50°C; ▼: 61°C).

Table 4. Permeate flux and ethanol concentration at different temperature.

<table>
<thead>
<tr>
<th>Temperature (°C)</th>
<th>Ethanol in Permeate (wt%)</th>
<th>Permeate Flux (g/m²s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>37</td>
<td>28.7</td>
<td>0.32</td>
</tr>
<tr>
<td>45</td>
<td>33</td>
<td>0.91</td>
</tr>
<tr>
<td>55</td>
<td>33</td>
<td>1.24</td>
</tr>
<tr>
<td>61</td>
<td>34</td>
<td>1.61</td>
</tr>
</tbody>
</table>
higher ethanol membrane selectivity. Since ethanol molecules have a larger size than water, the assumption can be made that diffusion effects are dominant over sorption effects, and are thus expected to be the determining parameter in the transport mechanism description through the PDMS-based membrane. It is expected that the minor increase of the selectivity at higher temperature is explained by a dependent decrease of the hydrogen bonding between water and alcohol molecules. When this is the case less water is stimulated to permeate through the membrane. Hence, not only swelling seems to play an important role during ethanol/water pervaporation but also the coupling effects between molecules, which tend to be lower at higher temperatures.

4.2.2. Extrapolation of the Experimental Results to the Target Unit (200,000 t/yr Ethanol)

To reduce the ethanol concentration in the fermenter below the inhibition threshold limit, it is proposed to continuously remove 25\% of the total fermenter flow rates through the membrane. Since the fermenter discharges 220,000 kg/hr of ethanol-H2O mixture at 11.6 wt\% ethanol, the feed rate to the pervaporation unit will be 55,000 kg/hr. Operating parameters for the large-scale application are given in Table 5.

Although it is expected that the continuous removal of ethanol will slightly increase the yield of the fermenter due to suppressing ethanol inhibition, this was not accounted for in the calculations.

With the given experimental results and operation data of Table 5, the following steady-state mass balance prevails.

\[
F_{\text{FEED}} = F_{\text{PERMEATE}} + F_{\text{RETENTATE}} + F_{\text{COLUMN}}
\]

\[
F_{\text{FEED}} \times X_{\text{FEED}} = F_{\text{PERMEATE}} \times X_{\text{PERMEATE}} + F_{\text{RETENTATE}} \times X_{\text{RETENTATE}} + F_{\text{COLUMN}} \times X_{\text{COLUMN}}
\]

\[
F_{\text{PERMEATE}} = 7920 \text{ kg/hr}
\]

\[
F_{\text{RETENTATE}} = 47080 \text{ kg/hr}
\]

\[
F_{\text{COLUMN}} = 165000 \text{ kg/hr}
\]

Both \(F_{\text{RETENTATE}}\) and \(F_{\text{COLUMN}}\) are fed to C1501. With a 33 wt\% ethanol content in the permeate, 2614 kg/hr of ethanol is continuously removed from the fermenter, reducing the effective concentration to <10.8 wt\%.

It is proposed to feed the pervaporation permeate directly to C1503-1 in view of its concentration of ethanol (~33 wt\%, i.e. ~41 vol\%).

The required membrane surface area is 7920/5.8 = 1366 m\(^2\). Although this surface area is considerable, it can be reduced by:

1) Operating the pervaporation at as close a temperature as possible to the azeotropic point (~78\(^\circ\)C), hence e.g. 75\(^\circ\)C. Extrapolating the T-dependent flux to 75\(^\circ\)C, yields values of ~2.4 g/m\(^2\)/s or 8.7 kg/m\(^2\)/h, leading to a reduction in membrane surface by 50\%.

2) The selection of a different membrane material, as suggested in the literature, where fluxes up to 10 kg/m\(^2\)/h at comparable or higher selectivities were indicated.

3) It is not considered to operate the pervaporation membranes at higher ethanol concentrations, hence e.g. in association with VHG, since it was demonstrated by Chovau [35] that the selectivity of all examined membranes is strongly dependent on the alcohol content in the feed: a higher concentration of selective compound (ethanol) in the feed led to a decrease in membrane selectivity. Similar observations were found by Chen et al. [36], who investigated the performance of unsupported PDMS-membranes in the separation of ethanol/water mixtures.
ranging between 4 and 91 wt% of ethanol in the feed, and explained the decrease by an extensive swelling of the PDMS rubber at higher feed ethanol concentrations. According to Jia et al. [37], incorporation of silicalite fillers in the polymeric network results in a reduction of membrane swelling and moreover improves the mechanical strength of the active layer.

4.2.3. Effect on the Overall Steam Requirements of the Distillation

Whilst using the same distillation column outlays (Table 1), the mass and heat balances around the complete unit, as illustrated in Table 2 before for a total 220,000 kg/hr of broth fed at 11.6 wt%, are now slightly modified with \( F_{\text{COLUMN}} + F_{\text{RETENTATE}} \) being fed to C1501 at 10.8 wt% ethanol.

With this operating mode, and internal energy reuse, the Aspen simulations revealed a required steam duty of the complete distillation train as given in Table 6.

Since condenser heat of C1502 is reused in reboiler C1501, and C1503-1, and condenser heat of C1503-2 is also used in C1501, the remaining net heat duty to be supplied from the steam network is reduced to ~18,222 kW, or:

\[
\frac{18222 \times 3600}{25375 \times 2300} = 1.13 \text{ kg steam/kg bio-ethanol}
\]

4.3. Effect on the Overall Steam Requirements of the Distillation

Combining the previous simulation results, the following overall evolution of steam consumption in the distillation emerges (Table 7).

As a conclusion, the steam use per kg of bio-ethanol is reduced from 1.18 to 1.17, 0.94 and 1.13 kg steam/kg bio-ethanol for options VHG (15%), VHG (19%) and pervaporation, respectively.

5. Tentative Economic Assessment

The full production cost of bio-ethanol at Cofco in 2013 is reported at 820 €/ton bio-ethanol. Based upon the predictions of the previous sections, a tentative cost calculation per ton bio-ethanol can be made, for a plant depreciation of 25 years, and other costs calculated on the basis of common ratios or literature data [38]. The efficiency of the steam boiler is estimated at 85%. The following Table 8 summarizes the calculations for the basis Cofco-like flowsheet.

Since the estimated production cost for cassava-based bio-ethanol (815 €/ton) is comparable with the Cofco-cited 820 €/ton, the proposed calculation procedure appears fair and acceptable.

Through a similar approach, and based upon estimated investment costs of the membrane module, the impact of hybrid pervaporation and/or VHG can be calculated. The life span of the membrane modules is estimated at 10 years, with a replacement of the membranes every 4 years. The total steam duty of the distillation section is then reduced to ~0.9 kg/kg bio-ethanol. The steam duty of the molecular sieve operation still needs to be added.

The impact of the VHG operation of the fermenter is given in Table 9.

### Table 6. Simulation results of the hybrid operation mode.

<table>
<thead>
<tr>
<th></th>
<th>C1501</th>
<th>C1502</th>
<th>C1503-1</th>
<th>C1503-2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Reboiler duty (kW)</td>
<td>8676</td>
<td>18,222</td>
<td>15,390</td>
<td></td>
</tr>
<tr>
<td>Condensor duty (kW)</td>
<td>5410</td>
<td>16,944</td>
<td></td>
<td>15,081</td>
</tr>
</tbody>
</table>

### Table 7. Summary results of the simulation of the Cofco distillation process.

<table>
<thead>
<tr>
<th>Simulation</th>
<th>Steam consumption (kg steam/kg ethanol)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Basic (no heat integration)</td>
<td>2.50</td>
</tr>
<tr>
<td>Integrating reboilers and condensers</td>
<td>1.18</td>
</tr>
<tr>
<td>VHG (15%)</td>
<td>1.17</td>
</tr>
<tr>
<td>VHG (19%)</td>
<td>0.94</td>
</tr>
<tr>
<td>Pervaporation</td>
<td>1.13</td>
</tr>
</tbody>
</table>
Table 8. Cost calculations for the current Cofco operation.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Unit</th>
</tr>
</thead>
<tbody>
<tr>
<td>Annual bio-ethanol production</td>
<td>200,000</td>
<td>Ton bio-ethanol/year</td>
</tr>
<tr>
<td>Depreciation</td>
<td>7</td>
<td>€/ton bio-ethanol</td>
</tr>
<tr>
<td>Plant overheads</td>
<td>80</td>
<td>€/ton bio-ethanol</td>
</tr>
<tr>
<td>Insurance</td>
<td>1.8</td>
<td>€/ton bio-ethanol</td>
</tr>
<tr>
<td>Spare parts and maintenance</td>
<td>22.4</td>
<td>€/ton bio-ethanol</td>
</tr>
<tr>
<td>Labour</td>
<td>11</td>
<td>€/ton bio-ethanol</td>
</tr>
<tr>
<td>Electricity</td>
<td>105</td>
<td>kWh/ton bio-ethanol</td>
</tr>
<tr>
<td>Cost of electricity</td>
<td>6.1</td>
<td>€/ton bio-ethanol</td>
</tr>
<tr>
<td>Steam duty (incl. molecular sieves)*</td>
<td>1.8</td>
<td>kg/kg bio-ethanol</td>
</tr>
<tr>
<td>Cost of heavy fuel oil</td>
<td>102</td>
<td>€/ton bio-ethanol</td>
</tr>
<tr>
<td>Demi-water for boiler</td>
<td>23</td>
<td>€/ton bio-ethanol</td>
</tr>
<tr>
<td>Cassava</td>
<td>6.6</td>
<td>Ton/tobio-ethanol [39]</td>
</tr>
<tr>
<td>Purchase of cassava</td>
<td>38.6</td>
<td>€/ton cassava [40]</td>
</tr>
<tr>
<td>Cost of cassava</td>
<td>254.8</td>
<td>€/ton bio-ethanol</td>
</tr>
<tr>
<td>Pre-treatment of cassava</td>
<td>132</td>
<td>€/ton bio-ethanol</td>
</tr>
<tr>
<td>Treatment of residues (solid/wastewater)</td>
<td>144</td>
<td>€/ton bio-ethanol</td>
</tr>
<tr>
<td><strong>TOTAL</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Cassava</strong></td>
<td><strong>815</strong></td>
<td>€/ton bio-ethanol</td>
</tr>
</tbody>
</table>

*The steam requirement for regenerating the molecular sieves amounts to 0.5 kg/kg bio-ethanol.

Table 9. Economic impact of integrating pervaporation (hybrid process) or VHG.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
<th>Unit</th>
</tr>
</thead>
<tbody>
<tr>
<td>Annual bio-ethanol production</td>
<td>200,000</td>
<td>Ton bio-ethanol/year</td>
</tr>
<tr>
<td>Additional depreciation</td>
<td>0.8</td>
<td>€/ton bio-ethanol</td>
</tr>
<tr>
<td>New steam duty (incl. molecular sieve)</td>
<td>1.4</td>
<td>kg/kg bio-ethanol</td>
</tr>
<tr>
<td>Cost of heavy fuel oil</td>
<td>79</td>
<td>€/ton bio-ethanol</td>
</tr>
<tr>
<td>Reduced Demi-water for boiler</td>
<td>11</td>
<td>€/ton bio-ethanol</td>
</tr>
<tr>
<td>Reduced residue treatment (~15%)</td>
<td>123</td>
<td>€/ton bio-ethanol</td>
</tr>
<tr>
<td><strong>Comparison</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Difference with equivalent items of Table 8</td>
<td><strong>−56</strong></td>
<td>€/ton bio-ethanol</td>
</tr>
<tr>
<td>Corrected cost for cassava-based production</td>
<td><strong>−759</strong></td>
<td>€/ton bio-ethanol</td>
</tr>
</tbody>
</table>

The production cost by applying a hybrid operation and/or VHG, can reduce the total production cost by ~56 €/ton bio-ethanol, i.e. about 7%. For an annual production of 200,000 ton of bio-ethanol, savings of ~11,000,000 € are achieved.

Further research on both principles, preferably on pilot scale, appears economically and technically justified.

It should moreover be considered that molecular sieves can also be replaced by a hydrophilic membrane, hence operating without steam requirement. The steam duty is then further reduced by 0.5 kg steam/kg bio-
ethanol, representing a further saving on heavy fuel oil of minimum 28 €/ton bio-ethanol as [13].

6. Conclusions
To reduce the distillation-related energy-impact on the production costs of bio-ethanol, the paper investigated the effect of implementing different possible improvements into the traditional fermentation-distillation sequence. The comparison was made by Aspen Plus (V 8.2) simulation, and the current operation of the 200,000 ton/year bio-ethanol plant of Cofco (China) was used as basis. Literature and experimental data were gathered to define the results of possible VHG and pervaporation improvements.

A first improvement, implementing the energy-pinch approach enables to integrate recycle condenser and re-boiler heat duties. This reduced the steam duty from 2.50 (basic operation, without energy recycle) to 1.18 kg steam/kg bio-ethanol.

Further improvements are still possible, with VHG (at 19%) reducing the steam consumption to <1 kg steam/kg bio-ethanol, whilst operation in a hybrid mode (pervaporation of 25% of the fermenter broth) achieves ~20% of savings, but significantly reduces the risk of inhibition in the fermenter.

Finally, a preliminary economic analysis reveals that the factory-gate price of cassava-based bio-ethanol could be reduced from ~820 €/ton to ~760 €/ton.

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