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Oxygen-blown operation of the TwoStage Viking gasifier

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Abstract
The TwoStage Viking gasifier from the Technical University of Denmark is being further developed for biofuel synthesis applications. In order to optimize the gasification process, it is suggested to apply an O2-CO2 gas mixture as gasification medium, instead of air, to limit N2-dilution of the product gas. It is found through a modeling study that the system is expected to achieve operating conditions in the range of air-blown operation, when 21v% O2 in CO2 is applied, and nearly identical parameters as the concentration is increased to 30v%. An experimental campaign with the 80kWth Viking pilot plant using 21v% oxygen confirms this, as operation temperatures are seen to slightly decrease the partial oxidation (POX) temperature by 52-69⁰C and grate temperature by 31-36⁰C. Tests with 25v% oxygen were also carried out with slightly higher temperatures. Detailed gas analysis showed that N2 had effectively been reduced to a few percent and that tar and sulphur levels were similar to the very high standards of the air-blown operation: only a few mg/Nm3 of tar and <3 ppm sulphur were detected. The lone gas cleaning, a bag filter, was found to be virtually inactive for capturing these impurities. Hence, the gasifier had been successfully demonstrated with O2-CO2 mixtures and is expected to be able to maintain its simple design, whilst enabling very high system efficiency.

Keywords: Biomass gasification, Two-stage gasifier, Thermodynamic analysis, Experimental, Gas quality

1 Introduction
It is very cost-effective to use biomass-based energy to reduce the impact on climate change, because it to a large extent can be directly utilized into the current fossil infrastructure. Biomass as a flexible resource can be used for heat and power production, but is especially relevant as a carbon source for transport fuels. The fuels can be produced in a number of ways, but the thermal gasification platform offers maximum fuel and product flexibility along with very high conversion efficiency.

The production of biofuels via biomass gasification is a well-studied area and there are a number of limitations and challenges in this coupling of technologies which are namely associated with gas quality. Synthesis reactors are very sensitive to harmful gas impurities such as tars and inorganics and while no hard conclusions can be made on tolerances some overall considerations can be made. These reactors utilize catalytic material at elevated temperatures of ≈200-700⁰C for synthesis [1][2][3], which are typically in the range of the dew points of tar species in the product gas at ≈200-500⁰C, which can cause fouling of

1 Tolerances for gas impurities are set based on catalyst, operating conditions and economic analysis and will vary
equipment via condensation [4][5][6] and deactivate catalysts via carbon deposition [1][7]. Hence tar concentrations should be kept very low, in range of <100mg/Nm$^3$, depending especially on the dew point at the given conditions when applying catalytic reactors [1][8]. Sulphur and chlorine are both poisons to catalyst reactors and should be removed to <0.1-1.0ppmv and preferably completely removed to ensure years of catalyst lifetime [9][10].

Another critical gas component, nitrogen ($N_2$) is of special interest when coupling gasification and biofuel synthesis. It is especially central when synthesizing gaseous fuels such as synthetic natural gas (SNG), as nitrogen separation is expensive and dilution of the gas product might bring the Wobbe index and relative density out of the limits of the local natural gas grid\(^2\). This typically limits the nitrogen content to a few percent of the final product. As an inert gas, nitrogen is also problematic as it will increase the size of the costly synthesis reactors and is recommended around <2v% [10].

1.1 Modifying the TwoStage Viking gasifier

The TwoStage Viking gasifier has been developed for many years at the Technical University of Denmark [11]. The air-blown downdraft system is designed with separate pyrolysis and gasification, with a partial oxidation (POX) in between and is namely characterized by its ability to process biomass into product gas at a very high cold gas efficiency of 87-90% (dry) and a very low tar content of <0.1mg/Nm$^3$ [12][13]. The plant is presented in detail in Section 3.1. In order to develop the fuel synthesis application platform of the system, oxygen-blown operation is desired to avoid nitrogen dilution of the product gas - which coupled with the high efficiency and low tar content would make the system ideal for biofuel production. This study will therefore investigate an oxygen-blown configuration of the gasifier system.

The oxygen is initially projected to be used with a carrier gas such as $CO_2$ or steam in order to ease the implementation, obtain operating parameters similar to air and protect equipment, as it acts as a thermal buffer during oxidation and reduces potential erosion from the hot oxygen flow in the heat exchanger and piping. $CO_2$ is initially preferred over steam as carrier gas, as it: 1) can be easily implemented from gas bottles; 2) is much less energy intense (no need for vaporization) to implement; 3) and can possibly be recycled from a downstream product upgrader ($CO_2$ separator) and led back to the POX.

Replacing $N_2$ with $CO_2$ is expected to affect the POX zone, as the gases have significantly different properties - see Error! Reference source not found.. While no studies have been found on $O_2$-$CO_2$ partial oxidation of pyrolysis gas, several references [14][15][16][17][18] have dealt with flame studies of $O_2$-$CO_2$ burners in oxy-fuel combustion of coal. The main differences between these burners and typical air burners are related to the gas properties that can cause (21% $O_2$ and similar flow rates as reference):

- Higher flame instability and flame retarding via a lower flame propagation speed [14][15][17]
- Generally lower gas temperatures of $\approx$100-200$^\circ$C [15][16]
- Increased diffusion resistance of $CO_2$ that can cause a more compact flame [16]
- Similar operational parameters for air and $O_2$-$CO_2$ are reached at $O_2$ concentrations of 28-35v% [14][15][16][17]

\(^2\) According to Danish law, the Wobbe Index must be higher than 50.76 MJ/Nm$^3$ (HHV) and the relative density must be higher than 0.555 [34].
Thus it will be important for the POX zone to optimize the mixing/contact between reactants to obtain a stable flame and maximize the tar exposure to high-temperatures in the potentially smaller flame zone, which will be even more important as the temperature is expected to decrease.

<table>
<thead>
<tr>
<th></th>
<th>Density [kg/m³]</th>
<th>Heat capacity [J/mol-K]</th>
<th>Mass diffusion coefficient of O₂ in N₂/CO₂ [m³/s]</th>
</tr>
</thead>
<tbody>
<tr>
<td>N₂</td>
<td>0.244</td>
<td>34.18</td>
<td>1.7 · 10⁴</td>
</tr>
<tr>
<td>CO₂</td>
<td>0.383</td>
<td>57.83</td>
<td>1.3 · 10⁴</td>
</tr>
<tr>
<td>Ratio N₂/CO₂</td>
<td>0.640</td>
<td>0.59</td>
<td>1.31</td>
</tr>
</tbody>
</table>

Table 1 - Properties of N₂ and CO₂ at 850⁰C and 1bar [18]

Gasification with O₂-CO₂ mixtures has been studied on several occasions and compared to air-blown operation. Pohorely et al. [19] studied air and O₂-CO₂ (21% O₂) gasification in a limestone-fed bubbling fluid bed at 850⁰C and found that replacing N₂ with CO₂ generally heightens both carbon conversion and cold gas efficiency compared to air. This could partially be due to an increased inlet energy content, as the CO₂-mix has a higher heat capacity and same inlet temperature as the air. Higher CO and lower H₂ yields were also found, likely due to a shift in the water-gas shift equilibrium. Tar and light hydrocarbon concentrations were found to be significantly higher for O₂-CO₂ operation, which the authors explain by increased CO and CO₂ partial pressures that could shift the reforming reaction equilibrium away from conversion. Similar findings of carbon conversion, efficiency and CO₂ and H₂ yields were found by Hanaoka et al. [20]. The study gasifies aquatic biomass at 900⁰C in a downdraft fixed bed reactor with O₂-CO₂ and O₂-He operation and also found that an increased CO₂ flow caused larger concentrations of sulphur species (H₂S, COS), which could be related to increased carbon conversion, as the sulphur content of the char were found to be relatively high.

This study will present the use of O₂-CO₂ in the TwoStage Viking gasifier and compare it to its typical air-blown operation - both via mathematical modeling and experimentally. An experimental campaign will show the effects on the operating conditions of the partial oxidation and the char bed, and also assess the impact on gas quality including gas composition, tars and sulphur compounds. These aspects are not well-studied and will be investigated further here as a link in the development of the TwoStage gasifier and further development towards fuel synthesis applications.

2 Modeling O₂-CO₂-blown TwoStage gasification

In order to project and plan out experimental work, the TwoStage Viking gasifier is modeled and projected with air and O₂-CO₂ mixtures. The core process is modeled via three main components: pyrolyzer, POX and gasifier. The thermodynamic modeling is carried out in DNA (Dynamic Network Analysis) that is a zero-dimensional modeling tool for simulating energy systems [21][22].

Initially a model using air is constructed and calibrated to match current gasifier data. The main assumptions are given in Error! Reference source not found. and are mainly based on the very detailed experimental campaign report by Bentzen et al. [13]. Note that this report deals with a previously constructed, but very similar, TwoStage gasifier plant. The pyrolysis features a reactor that based on atomic
balances and input calculates a char and volatile fraction at the specified temperature. The POX and gasifier utilizes Gibbs minimization for calculating the gas composition and process parameters.

Fuel

49% moisture, 20°C, composition: C=49% H=6% O=44% N=0.2%
Ash=0.8%(balance) LHV=18.28MJ/kg(dry)

Pyrolyzer
Volatiles are assumed to have a H₂ and CH₄ content of 5v% and 13v% [13], tars are represented by n-hexane (C₆H₁₄), char yield of 25% [23], 1% heat loss of fuel input, LHVᵢₐ=LHVᵩᵪᵦ, outlet temperature of volatiles and char=600°C
Char composition: C=93%.0 H=2.2% N=0.2% HHV=33.6MJ/kg(dry) [23]

Gasification media
Air: O₂=21% N₂=77% H₂O=1% Ar=1%, O₂-CO₂: O₂=21% CO₂=79%, 20°C input, 450°C after heat exchanger

Partial oxidation
Assumes thermal equilibrium at outlet temperature via Gibbs minimization of volatiles-air mixture – method described in [24][25].

Gasifier
3% fuel input heat loss [13], gas outlet thermal equilibrium=800°C, gas composition calculated via chemical equilibrium (water-gas shift), carbon conversion=99% when using air [12], pressure loss is 30mbar, outlet temperature=750°C

Table 2 – Modeling parameters. *Composition from [12] and moisture from [13].

The model calibration included adjusting gasifier equilibrium temperature and allocating the total heat loss of 4% on to the pyrolyzer and gasifier. The air-blown model is pictured in Figure 1 with resulting key data. Main model results including gas composition are compared to operational data for the TwoStage gasifier in Table 3 showing an overall satisfying fit. One of the key differences is the CH₄ content that is negligible in the model, but in practice typically 1-2% methane is formed in and/or slips by the partial oxidation zone, which converts most of it and other hydrocarbons. This difference will namely cause a slightly higher hydrogen content in the model as well as a slight difference in partial oxidation temperature due to endothermic reforming reactions. The POX (maximum) temperature is seen to be within range of the measured temperature, but this value will in practice fluctuate between ≈1100-1300°C.
Wood chips
49% H₂O
25°C

Pyrolysis

Heat

600°C
Volatiles

Partial oxidation

Char

450°C

1191°C

Gasification

Product gas

Air
20°C

750°C

Ash
(1% of input C)

Figure 1 – Model of TwoStage gasifier

<table>
<thead>
<tr>
<th>Unit</th>
<th>TwoStage Viking gasifier</th>
<th>Model</th>
</tr>
</thead>
<tbody>
<tr>
<td>Air/fuel flow [kg/kg]</td>
<td>1.36</td>
<td>1.33</td>
</tr>
<tr>
<td>POX temperature [°C]</td>
<td>1150 ±100</td>
<td>1191</td>
</tr>
<tr>
<td>Cold gas efficiency [%] (LHV)</td>
<td>90</td>
<td>89.1</td>
</tr>
<tr>
<td>H₂ [v%]</td>
<td>34</td>
<td>36</td>
</tr>
<tr>
<td>CO [v%]</td>
<td>17</td>
<td>17</td>
</tr>
<tr>
<td>CO₂ [v%]</td>
<td>17</td>
<td>17</td>
</tr>
<tr>
<td>CH₄ [v%]</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td>N₂ [v%]</td>
<td>31</td>
<td>30</td>
</tr>
<tr>
<td>LHV [MJ/kg]</td>
<td>6.6</td>
<td>6.4</td>
</tr>
</tbody>
</table>

Table 3 – Comparison of TwoStage gasifier [13] and model. All flows and fractions are dry.

The calibrated model is used to project various O₂-CO₂ mixtures, which are compared to the air-blown mode by implementing similar parameters. Two main weaknesses by using this direct comparison are namely that the CH₄ concentration can be affected by a change in operation and that the product gas and air/O₂-CO₂ temperatures are assumed constant across gasification media.

As seen in Table 4 the direct substitution of CO₂ for N₂ with 21v% O₂ will cause a decline in gasifier performance with lower temperatures and subsequent carbon conversion and efficiency. Namely the higher heat capacity is responsible for this decline, as the gasifier exhaust will have a higher content of sensible heat. In order to keep the efficiency and POX temperature at similar levels, a ≈12% larger volume flow is required. Generally it is seen that the efficiency and POX temperature are in the same range as for the air-blown mode.

In line with the literature review, it is seen that an O₂-concentration of 30v% obtain very similar parameters to those of the air-blown mode. As an extreme case, pure oxygen might be added to the process, which is seen to obtain higher performance across parameters, as the otherwise large amounts of N₂/CO₂ does not need to be heated and carried through the system. Increasing the O₂ concentration for higher cold gas efficiency is in line with experimental studies e.g. [20][26]. The use of pure oxygen on the plant might
however be challenging with regards to the present plant design (temperatures, materials, gas flows etc.) and highly dependent on the fuel moisture levels in order to avoid hot spots. Therefore it is seen as reasonable to blend the oxygen with a carrier gas in order to make the system more robust and allow dryer fuels and potential other fuels with a lower volatile fraction that both will increase the POX temperature.

<table>
<thead>
<tr>
<th>Unit</th>
<th>Air</th>
<th>O₂-CO₂</th>
<th>O₂-CO₂</th>
<th>O₂-CO₂</th>
<th>O₂</th>
</tr>
</thead>
<tbody>
<tr>
<td>Oxygen fraction [v%]</td>
<td>21</td>
<td>21</td>
<td>21</td>
<td>30</td>
<td>100</td>
</tr>
<tr>
<td>Gas/fuel flow [m³/kg(dry)]</td>
<td>1.13</td>
<td>1.13</td>
<td>1.26</td>
<td>0.80</td>
<td>0.22</td>
</tr>
<tr>
<td>Gas preheat to 450°C [kWₚ,h]</td>
<td>2.6</td>
<td>3.6</td>
<td>4.0</td>
<td>2.6</td>
<td>0.5</td>
</tr>
<tr>
<td>POX temperature [⁰C]</td>
<td>1191</td>
<td>1085</td>
<td>1144</td>
<td>1197</td>
<td>1307</td>
</tr>
<tr>
<td>Carbon conversion [%]</td>
<td>99.0</td>
<td>90.2</td>
<td>99.0</td>
<td>99.0</td>
<td>99.0</td>
</tr>
<tr>
<td>Cold gas efficiency [%](dry, LHV)</td>
<td>89.1</td>
<td>80.9</td>
<td>87.8</td>
<td>89.2</td>
<td>91.0</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th></th>
<th>H₂ [v%(dry)]</th>
<th>CO [v%(dry)]</th>
<th>CO₂ [v%(dry)]</th>
<th>CH₄ [v%(dry)]</th>
<th>N₂ [v%(dry)]</th>
<th>LHV₅₅ [MJ/kg]</th>
<th>LHV_vol [MJ/Nm³]</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>36</td>
<td>17</td>
<td>17</td>
<td>0</td>
<td>30</td>
<td>6.4</td>
<td>5.8</td>
</tr>
<tr>
<td></td>
<td>30</td>
<td>25</td>
<td>46</td>
<td>0</td>
<td>0</td>
<td>5.1</td>
<td>6.1</td>
</tr>
<tr>
<td></td>
<td>28</td>
<td>26</td>
<td>46</td>
<td>0</td>
<td>0</td>
<td>5.1</td>
<td>6.1</td>
</tr>
<tr>
<td></td>
<td>36</td>
<td>27</td>
<td>37</td>
<td>0</td>
<td>0</td>
<td>6.6</td>
<td>7.0</td>
</tr>
<tr>
<td></td>
<td>52</td>
<td>25</td>
<td>23</td>
<td>0</td>
<td>0</td>
<td>10.8</td>
<td>8.5</td>
</tr>
</tbody>
</table>

Table 4 – Model comparison using air or O₂-CO₂. aAt 20⁰C, 1bar. bBased on 80kWₚ,h fuel input (LHV).

Based on the literature and modeling studies presented, the TwoStage Viking gasifier plant was modified and experimental campaigns were carried out over 3 days. The campaign details are presented in the following sections.

3 Experimental methods and materials

3.1 The gasifier plant

The TwoStage Viking gasifier has been developed for many years at the Biomass Gasification Group at the Technical University of Denmark. The 80kWₚ,h gasifier pilot plant processes wood chips in two stages: moving bed pyrolysis and downdraft fixed bed char gasification with a POX zone in between – see Figure 2. The pyrolyzer employs an externally heated screw conveyer that processes the fuel up to 600⁰C by utilizing engine exhaust for heat supply. The released volatiles and char are led to the second reactor where they are exposed to a POX zone where 99% of the residual tar is converted [27]. The hot gases and char are then led to the char bed where the char is gasified and the remaining tars are reduced to a minimum by a 95-99% [27], resulting in a reported tar content of ≤0.1mg/nm³ [12]. Coupled with a high carbon conversion of 99%, the cold gas efficiency reaches 87% on dry basis and 93% on wet basis (34wt% moisture) [12]. The gasifier applies only a simple bag filter for gas cleaning in order to capture particles and a condenser to dry the product gas.
As part of the development of the system, a steam dryer has been installed on the Viking plant. This will enable the use of fuel with high moisture contents up to ≈60-70% and also enable separation of the high-temperature pyrolysis heat exchanger area as shown in scaled up designs [28]. The steam dryer utilizes a steam loop, where it is moved and heated by a blower and an electrical heater. As seen on Figure 2, the steam is then passed through a screw conveyer where the fuel moisture evaporates. The main fraction of the steam is then recirculated via a blower and reheated, while the produced moisture-steam is carried with the dry fuel to the pyrolyzer. The inlet steam temperature to the steam dryer was 173°C.
The standard gasification medium is atmospheric air that is delivered to the system via a blower. Replacing
the air injection with an $O_2$-CO$_2$ mixture is done via the following setup. The $O_2$ and CO$_2$ are supplied via gas
bottles, reduced to 10 bar via reduction valves and led to a gas mixer (Dansensor MAP Mix Provectus.) The
gas mixer is based on two mass flow controllers, which secure the correct composition within 1%. The
mixer feeds a 100L buffer tank with a reduction valve, that secures a stable outlet pressure at 3 bar. The
system feeds a thermal mass flow controller (Aalborg Model GFC) that uses the original air blower signal
from the PLC (Siemens Step7) to dose the mixture near atmospheric pressure levels. The flow controller has
an accuracy of ±1%. The mixture composition is manually set at the mixer and has been thoroughly tested
beforehand. The equipment is shown in Figure 3.

![Figure 3 – Experimental setup for converting the air-blown gasifier to $O_2$-CO$_2$-blown. Left: Gas bottles. Upper right: gas mixer. Lower right: mass flow controller.](image)

The gasifier can be operated via various strategies, but a constant air/$O_2$-CO$_2$ flow into the POX zone is
chosen as the primary setpoint. The fuel feed is automatically set to maintain a bed height within a
specified interval. The ash is discharged by the grate as the pressure difference builds up. When switching
to the 21v% oxygen-mix, the volume flow is maintained similar to air. The 25v% oxygen-mix volume flow is
set to match the absolute oxygen flow, meaning a smaller total gas flow is applied.

### 3.2 Fuel analysis

Standard Danish spruce wood chips were used as fuel for the gasifier. The main fuel elements (C, H, N, S, O)
was analyzed with the VarioEL III (Elementar Analysensysteme GmbH, Germany) and results are shown in
Table 5. The inorganic elements were measured by inductively coupled plasma optical emission
spectrometry (ICP-OES) analysis and are shown in Table 6. The analysis showed a high amount of Ca
(1063.0 and 1061.9 μg/g), whereas the presence of K, Mg and Na were also significant in the fuel.

<table>
<thead>
<tr>
<th></th>
<th>C [%]</th>
<th>H [%]</th>
<th>N [%]</th>
<th>S [%]</th>
<th>O (rest) [%]</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Table 5 – Composition of dry spruce wood chips used in the experiments based on 4 samples.

<table>
<thead>
<tr>
<th>Element</th>
<th>Al</th>
<th>B</th>
<th>Ca</th>
<th>Cu</th>
<th>Fe</th>
<th>K</th>
<th>Mg</th>
<th>Mn</th>
<th>Na</th>
<th>P</th>
<th>S</th>
<th>Zn</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sample #1</td>
<td>38.6</td>
<td>1.6</td>
<td>1063.0</td>
<td>1.2</td>
<td>38.4</td>
<td>295.0</td>
<td>116.5</td>
<td>8.2</td>
<td>116.5</td>
<td>76.2</td>
<td>130.3</td>
<td>8.7</td>
</tr>
<tr>
<td>Sample #2</td>
<td>36.9</td>
<td>1.3</td>
<td>1061.9</td>
<td>1.5</td>
<td>36.2</td>
<td>252.1</td>
<td>113.5</td>
<td>8.0</td>
<td>109.3</td>
<td>77.2</td>
<td>125.5</td>
<td>8.5</td>
</tr>
</tbody>
</table>

Table 6 – Inorganic elements detected in the wood chips [μg/g] dry

The moisture content was relatively low compared to previous tests and was measured to 16-17wt% on 4 occasions during testing, by weighing samples before and after at least 24 hours residence in a 105°C oven.

3.3 Gas analysis

Several types of analyses were performed on the collected product gas. For tar measurements, solid phase adsorption (SPA) was used. Sulphur was measured via gas samples and gas chromatography. For gas composition measurements, both gas chromatography from pipettes and online measurements were used, because the gas composition were out of the online equipment measurement range when applying O$_2$-CO$_2$.

Solid phase adsorption (SPA)

The tar measurements were taken at two locations: before the bag filter and after the condenser. Sampling was done through a needle connected to the filter and subsequently a 100ml syringe – see Figure 4. The needle was placed in the middle of the hot gas flow as the samples were taken. Sample was done over approximately 30 seconds.

![Figure 4 – SPA sampling: needle with cartridge fastened to syringe via rubber band](image)

The received SPA filters were removed, extracted and resulting samples were analyzed by gas chromatography – mass spectrometry (GC-MS). The cartridges were removed from the SPE tubes and were stored overnight after addition of 10 ml acetone, 1ml phenol D5 and 1ml polycyclic aromatic hydrocarbon (PAH) standard mixture. (The PAH standard mixture included: naphthalene D8, acenaphthene D10, anthracene D10, phenanthrene D10, fluorene D10, pyrene D10). The acetone was then evaporated and the samples were redissolved in 1 ml acetone. The samples were analyzed by GC-MS using a Hewlett Packard HP 6890 gas chromatograph interfaced to a HP5973 Mass Selective Detector (Agilent, Denmark). Samples (1 μl) were injected in split mode (1:20) using an HP 7683 autosampler (Agilent, Denmark). The source and rod temperatures were 230°C and 150°C, respectively. The products were separated using a 0.32-mm i.d.×30 m WCOT-fused silica column coated with VF-23ms at a thickness of 0.25 μm (Analytical Instruments as, Denmark). The carrier gas was He at a flow rate of 1.2 ml/min. Separation of products was achieved using a temperature program from 70 to 250 °C at 10°C/min. The applied ionization energy was 70 eV. Full
mass spectra were recorded every 0.3s (mass range m/z 40–450). Products were identified using NIST search engine version 2.0 f. (Agilent, Denmark). The deuterated stable isotopes were used as internal standards and used for quantification, as they were added to the system in a known amount.

**Sulphur measurements**

Gas sampling for analysing the sulphur components H₂S and COS were performed Day 1 (air blown gasifier operation) and Day 3 (25% oxygen blown gasifier operation). Both days sampling were performed upstream (before particle filter) and downstream (after the condenser) of the gasifier gas cleaning equipment. Three samples were taken at each location comprising a total number of six samples each of the two days. The gas samples were extracted into dedicated 5 litre Supel™ Inert Foil gas sampling bags with screw cap valves from SUPELCO. Upstream gas cleaning the product gas pressure was negative and an EX gas pump was used to extract the gas into the SUPELCO bags. Downstream gas cleaning the product gas pressure was positive and large enough to fill the sampling bags by just opening the sampling port valve. Prior to the sampling the sampling connection tubes were flushed to get rid of air and accumulated condensed water in the sampling ports.

The sampling on Day 1 was performed as described above, but the subsequently analysis revealed sensitivity issues using the Agilent GCMS due the high water content in the gas samples. Therefore it was decided to add an amount of helium (He) to the gas sampling bags before performing the actual measurements at October 12. The result was diluted gas samples with water dew point lower than 100% (unsaturated gas samples) which caused lesser issues during analysis.

Analysis was done with an Agilent 7890A gas chromatograph combined with 5975C mass selective detector. Pre-concentration of the sample was done with a Markes Unity 2 thermal desorber with air sampler. Pre-concentration conditions: Markes T6SUL-2S cold trap at -30°C; gas flow 50 ml/min during 5 minutes. Injection: trap heating at maximum rate to 300°C; total split ratio 29.4. Analysis: separation of H₂S and COS was done at an Agilent DB-5ms column (20 m x 180 mm x 0.14 mm) at 35 °C with He as carrier gas. Other components were removed from the column by heating to 200°C at 25°C/min.

**Gas analysis from gas pipettes**

The gas sampling was done after the product gas condenser, where the pipettes were connected to the sampling port with a rubber tube – see Figure 5. The vessel was flushed thoroughly for roughly 1 minute by opening both ends and afterwards filled. Gas analysis was carried out within 1-2 hours to avoid leakage of H₂.

![Gas pipette sampling with rubber tube connection to sampling port](image-url)
The samples were analyzed by an Agilent Technologies 7890A gas chromatograph (Agilent, Denmark). Samples (volumes 100-1000 µl) were injected in split mode (1:25) using Pressure-Lok® Syringes (Vici, Baton Rouge, USA). For measurements of H₂, CO and CH₄, the gas components were separated using a 0.32-mm i.d. × 25 m PLOT-fused silica column coated with Molsieve 5A (Analytical Instruments as, Denmark). The carrier gas was Ar at a flow rate of 1 ml/min. For CO measurements, He carrier gas was used. Separation of products was achieved using a temperature program from 50 to 235°C at 10°C/min. For measurements of CO₂, the gas components were separated using a 0.32-mm i.d. × 25 m PLOT-fused silica column coated with Poraplot U (Analytical Instruments as, Denmark). The carrier gas was He at a flow rate of 1 ml/min. Separation of products was achieved using a temperature program from 75 to 235°C at 16°C/min. Gas components were detected with a thermal conductivity detector (TCD).

**Online gas measurements**

On dry and tar-free basis, the gas composition were measured by an Advance Optima 2020 Modular continuous process gas analyzer system. The system was equipped with a Caldos 15 cell for analysis of H₂ and an Uras 14 cell for CO, CO₂, and CH₄ (ABB). The only issue with the Uras 14 cell was that the CO₂ measurement range is 30%, which makes the gas composition data unprecise above this limit. An PMA 10 O₂-analyzer were also applied. N₂ was calculated as difference. All of the equipment has a ±1% uncertainty range.

**4 Experimental results**

The gasifier was heated up to stable operating conditions over roughly 24 hours using initially gas burners and afterwards air-blown operation from afternoon till next morning. The reported tests were carried out over 3 following days: Day 1 - air-blown, Day 2 – 21 and 25v% O₂-CO₂-blown, Day 3 - 21v% O₂-CO₂-blown. While tests were run during work hours, the system was kept thermally stable by operating it fully automated and unmanned overnight with air.

**4.1 Operating temperatures and gas composition**

Time dependent temperature data for air and O₂-CO₂ operation is shown in Figure 6 and Figure 7 respectively, while the corresponding online gas measurements are given in Figure 8 and

![Figure 9](https://example.com/figure9.png)

Figure 9. Temperature measurements were taken after the air preheater, at the POX zone, just above the gasifier grate and at the reactor outlet and all parameters showed satisfying process stability. In- and
decreasing trends can be due to changes in fuel moisture and/or thermal stability. The operating data is within range of previous tests with the gasifier [12][13][25].

Figure 6 – Operating temperatures for air-blown operation during Day 1.

Figure 7 – Operating temperatures for 21v% and 25v% O₂-CO₂-blown operation during the time of testing. The two periods before the switch to 25v% O₂-CO₂ is 21v% O₂-CO₂. Day 2
Figure 8 – Gas composition for air-blown operation during Day 1. The spikes of N₂ are caused by filter flushing and these are minimized around 17:30 where the gas analysis is taken after the product gas buffer tank instead of before.

Figure 9 – Gas composition for 21v% and 25v% O₂-CO₂-blown operation during Day 2. Test were interrupted by a power outage around 9:20. Note that the measurements are out the measurement range and therefore not precise.

The operating temperatures and gas compositions are summarized as averages in Table 7 and gas pipette gas compositions are given in Table 8. As discussed in the modeling study (Section 2), it is seen that the general trend is that the temperatures are decreasing in the gasifier as the process is switched from air to 21v% O₂-CO₂: a POX temperature reduction of 52-69°C with grate temperatures decreasing with 31-36°C. The preheating temperature is generally somewhat lower, which is expected as the heat capacity is significantly higher compared to air. At 25v% oxygen, both preheat, POX and grate temperatures are increased and are more similar to air-blown data.

The N₂ content is drastically reduced, but a couple of percentages remain, which is mainly due to fuel feed ash silo N₂ purging. The level is expected to be reduced to a negligible content if e.g. CO₂-purge is applied instead. As expected from the modeling, the H₂ content is slightly lower, which is likely a result of a displacement in the water-gas shift reaction as the CO₂ concentration is much higher. The CO₂ and CO concentrations are in line with the predictions of the model, but slightly lower though. This small difference
can – amongst several minor differences – be partly attributed to a lower steam content (model fuel moisture 49%, while 16-17% experimentally) content that will promote CO₂ conversion in the char bed. As mentioned, the oxygen flow is constant across tests, meaning that the CO₂ will be lower at 25v% O₂, which is the main cause of the lower CO₂ concentration at this condition. The lower CO level at 25v% O₂-CO₂ is likely due to reduced char-CO₂ gasification reactions. Methane contents are low and similar for both tests as the POX temperature remains high.

The samples during the 25v% O₂-CO₂ operation is seen to be somewhat inaccurate, as the sum of gas components did not close in on 100% and it is still unknown why this is the case.

<table>
<thead>
<tr>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Air</td>
<td>12:45-18:45</td>
<td>1</td>
<td>686</td>
<td>1177</td>
<td>792</td>
<td>736</td>
</tr>
<tr>
<td>Air #2</td>
<td>0:00-6:13</td>
<td>2</td>
<td>680</td>
<td>1188</td>
<td>766</td>
<td>719</td>
</tr>
<tr>
<td>21v% O₂-CO₂</td>
<td>6:44-9:20</td>
<td>2</td>
<td>683</td>
<td>1137</td>
<td>730</td>
<td>714</td>
</tr>
<tr>
<td>21v% O₂-CO₂</td>
<td>13:05-15:24</td>
<td>2</td>
<td>604</td>
<td>1136</td>
<td>735</td>
<td>692</td>
</tr>
<tr>
<td>25v% O₂-CO₂</td>
<td>15:24-16:24</td>
<td>2</td>
<td>616</td>
<td>1149</td>
<td>745</td>
<td>689</td>
</tr>
<tr>
<td>Air #3</td>
<td>0:00-6:23</td>
<td>3</td>
<td>640</td>
<td>1183</td>
<td>766</td>
<td>695</td>
</tr>
<tr>
<td>21v% O₂-CO₂</td>
<td>10:42-17:00</td>
<td>3</td>
<td>595</td>
<td>1114</td>
<td>731</td>
<td>708</td>
</tr>
</tbody>
</table>

Table 7 – Day-by-day Averaged temperature and online gas data for the test campaigns. *Test data for overnight operation without gas analysis.

<table>
<thead>
<tr>
<th>Day - time</th>
<th>H₂ [v%]</th>
<th>CO₂ [v%]</th>
<th>CO [v%]</th>
<th>CH₄ [v%]</th>
<th>N₂ [v%]</th>
<th>SUM</th>
</tr>
</thead>
<tbody>
<tr>
<td>Air</td>
<td>1 – 12:45-18:45</td>
<td>27.0</td>
<td>14.7</td>
<td>15.3</td>
<td>0.4</td>
<td>42.1</td>
</tr>
<tr>
<td>21v% O₂-CO₂</td>
<td>2 – 11:15</td>
<td>24.0</td>
<td>42.6</td>
<td>26.9</td>
<td>0.05</td>
<td>4.6</td>
</tr>
<tr>
<td></td>
<td>2 – 13:20</td>
<td>21.2</td>
<td>43.2</td>
<td>24.9</td>
<td>0.16</td>
<td>4.7</td>
</tr>
<tr>
<td></td>
<td>2 – 13:22</td>
<td>20.6</td>
<td>44.3</td>
<td>25.8</td>
<td>0.22</td>
<td>4.6</td>
</tr>
<tr>
<td>25v% O₂-CO₂</td>
<td>2 – 15:50</td>
<td>21.2</td>
<td>39.3</td>
<td>18.6</td>
<td>0.04</td>
<td>3.8</td>
</tr>
<tr>
<td></td>
<td>2 – 16:50</td>
<td>22.8</td>
<td>39.4</td>
<td>20.6</td>
<td>0.09</td>
<td>4.6</td>
</tr>
</tbody>
</table>

Table 8 – Data from online gas analysis (air) and gas chromatography data from gas pipette samples (21v% and 25v% O₂-in-CO₂).

### 4.2 Tars and inorganics

Tar measurements are shown in Table 9. For air-blown operation, the results show expected low results in the low mg/m³-range with only PAH compounds present. The particle filter is seen to not cause any significant reduction in tar concentration, however on day 1 no tars could be measured after the filter. While the relative difference between tar concentrations in the air and O₂-CO₂ samples is high, the absolute
difference is seen to be very small. Hence no significant difference is seen between the two states. It is however interesting to provide a brief overview of the parameters that could cause a difference (char bed height and product gas flows were seen to be similar across the samples and are not expected to impact the tar):

- Reduced grate and POX temperature with \( \text{O}_2\) to \( \text{CO}_2 \) of \( \approx 60^\circ\text{C} \) and \( \approx 40^\circ\text{C} \) respectively will decrease tar conversion, as the conversion by thermal and chemical (char) means will be reduced.
- Applying a \( \text{CO}_2 \)-rich gasification medium will affect the char structure, as it has shown to produce char with a larger microstructure, specifically increase micro- and mesopore area e.g. [29], which will likely increase tar conversion [29][30].
- By increasing the \( \text{CO}_2 \) partial pressure, it is likely that an increase in dry reforming of tar over the char will be present and hence decrease the tar concentration.

<table>
<thead>
<tr>
<th>Time</th>
<th>Location</th>
<th>Gasifier medium</th>
<th>Pyrene</th>
<th>Naphthalene</th>
<th>Sum [mg/Nm³]</th>
<th>Sum [ppm]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Day 1</td>
<td>Pre-liminary</td>
<td>Before filter</td>
<td>4.9 ±0.2</td>
<td>0</td>
<td>4.9</td>
<td>N/A</td>
</tr>
<tr>
<td>Day 1</td>
<td>Pre-liminary</td>
<td>After filter</td>
<td>4.2 ±0.5</td>
<td>0</td>
<td>4.2</td>
<td>N/A</td>
</tr>
<tr>
<td>Day 2</td>
<td>10:29</td>
<td>Before filter</td>
<td>2.8</td>
<td>0</td>
<td>2.8</td>
<td>0.003</td>
</tr>
<tr>
<td>Day 1</td>
<td>11:00</td>
<td>Before filter</td>
<td>3</td>
<td>0</td>
<td>3</td>
<td>0.003</td>
</tr>
<tr>
<td>Day 1</td>
<td>10:05</td>
<td>After filter</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Day 1</td>
<td>10:17</td>
<td>After filter</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Day 2</td>
<td>Preliminary</td>
<td>Before filter</td>
<td>21v% ( \text{O}_2)-( \text{CO}_2 )</td>
<td>5.7 ±0.8</td>
<td>3.5 ±2.5</td>
<td>9.2</td>
</tr>
<tr>
<td>Day 2</td>
<td>Preliminary</td>
<td>After filter</td>
<td>21v% ( \text{O}_2)-( \text{CO}_2 )</td>
<td>3.8 ±0.2</td>
<td>6.5 ±0.4</td>
<td>10.3</td>
</tr>
<tr>
<td>Day 2</td>
<td>13:53</td>
<td>Before filter</td>
<td>21v% ( \text{O}_2)-( \text{CO}_2 )</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Day 2</td>
<td>13:42</td>
<td>After filter</td>
<td>21v% ( \text{O}_2)-( \text{CO}_2 )</td>
<td>1</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>Day 2</td>
<td>13:47</td>
<td>After filter</td>
<td>21v% ( \text{O}_2)-( \text{CO}_2 )</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
</tbody>
</table>

Table 9 – Tar measurements [mg/Nm³] before and after the bag filter of the gasifier. Preliminary samples were taken during the initial tests of the system 2 months prior to the main experimental work that is reported here – operation conditions were very similar.

Gas samples were taken during Day 1 and 3 to assess the sulphur load and results are shown in Table 10. The range of 0.6-2.8 ppm total sulphur is within previous measurements of the gasifier of 3.7 ppm of COS (no \( \text{H}_2\text{S} \)) [25], 0.17-0.28 ppm of COS (no \( \text{H}_2\text{S} \)) [32] and <2ppm \( \text{H}_2\text{S} + \text{COS} \) [33], which is also in line with the sulphur content of the applied wood fuels (Table 6) and is similar to previous analysis of wood fuel for the Viking [12]. It was expected that the bag filter might capture some of the sulphur species, as it will be partially coated with char from the gasifier and hence act as a carbon filter. It is however seen that this is not the case, as the filters’ capture, if any, is negligible. The sampling and analysis were carried out by
Danish Gas Technology Center and the relative uncertainty was estimated based on experiences to 40% for the first three samples in Table 10 and 25% for the remaining samples.

Between the media, the difference in sulphur species is negligible, with an additional 1ppm extra on average for the O\textsubscript{2}-CO\textsubscript{2} blend. This is due to additional COS, that could be slightly promoted with the given gas composition. As mentioned, previous tests have shown higher COS levels when air was applied, and hence the difference might also be due to small variations in operation from Day 1 to 3.

<table>
<thead>
<tr>
<th>Sampling time</th>
<th>Location</th>
<th>Gasification media</th>
<th>H\textsubscript{2}S [ppm]</th>
<th>COS [ppm]</th>
<th>Total S [ppm]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Day 1 12:20</td>
<td>Before filter</td>
<td>Air</td>
<td>0.1</td>
<td>0.6</td>
<td>0.7</td>
</tr>
<tr>
<td>Day 1 12:23</td>
<td>After filter</td>
<td>Air</td>
<td>0.1</td>
<td>0.5</td>
<td>0.6</td>
</tr>
<tr>
<td>Day 1 12:27</td>
<td>Before filter</td>
<td>Air</td>
<td>0.1</td>
<td>1.0</td>
<td>1.1</td>
</tr>
<tr>
<td>Day 1 12:32</td>
<td>Before filter</td>
<td>Air</td>
<td>0.4</td>
<td>1.1</td>
<td>1.5</td>
</tr>
<tr>
<td>Day 1 12:36</td>
<td>After filter</td>
<td>Air</td>
<td>0.4</td>
<td>1.0</td>
<td>1.4</td>
</tr>
<tr>
<td>Day 3 10:52</td>
<td>Before filter</td>
<td>21v% O\textsubscript{2}-CO\textsubscript{2}</td>
<td>0.3</td>
<td>1.8</td>
<td>2.1</td>
</tr>
<tr>
<td>Day 3 10:57</td>
<td>After filter</td>
<td>21v% O\textsubscript{2}-CO\textsubscript{2}</td>
<td>0.4</td>
<td>2.4</td>
<td>2.8</td>
</tr>
<tr>
<td>Day 3 11:36</td>
<td>Before filter</td>
<td>21v% O\textsubscript{2}-CO\textsubscript{2}</td>
<td>0.2</td>
<td>1.6</td>
<td>1.8</td>
</tr>
<tr>
<td>Day 3 11:41</td>
<td>After filter</td>
<td>21v% O\textsubscript{2}-CO\textsubscript{2}</td>
<td>0.2</td>
<td>1.3</td>
<td>1.5</td>
</tr>
<tr>
<td>Day 3 12:07</td>
<td>Before filter</td>
<td>21v% O\textsubscript{2}-CO\textsubscript{2}</td>
<td>0.2</td>
<td>1.8</td>
<td>2.0</td>
</tr>
<tr>
<td>Day 3 12:12</td>
<td>After filter</td>
<td>21v% O\textsubscript{2}-CO\textsubscript{2}</td>
<td>0.2</td>
<td>1.3</td>
<td>1.5</td>
</tr>
</tbody>
</table>

Table 10 – Measurements for sulphur in the product gas.
5 Conclusions
The Viking gasifier has been successfully converted from its original air-blown configuration to using O$_2$-CO$_2$ as gasification medium. Literature, modeling and experimental studies showed that operating conditions were expected to be in the range of air-blown values at 21-30v% O$_2$-in-CO$_2$, with partial oxidation and grate temperatures reduced by 52-69°C and 31-36°C respectively at 21v% O$_2$. Detailed gas analysis for tar and sulphur species showed that the gas qualities during O$_2$-CO$_2$ operation were comparable to the very high standards of the typical air-blown mode at <11mg/Nm$^3$ and <3ppm respectively – without any downstream gas cleaning equipment.

Hence the system can be successfully converted to operate with an O$_2$-CO$_2$ blend without major additions to existing design. Compared to the more typically applied O$_2$-H$_2$O medium in the literature, applying CO$_2$ might be better suited for some applications, as the media can be: 1) conveniently recirculated back to the oxygen source without need for high evaporation heat; 2) be completely converted into biofuels by addition of electrolytic hydrogen downstream of the system.

6 Acknowledgements
The authors would like to thank the ForskVE-programme of Energinet.dk for financial support through the Biomass Gasification Polygeneration project (ForskVE-12205).

7 References


Highlights of the article “Oxygen-blown operation of the TwoStage Viking gasifier”

- Air and 21-25v% O$_2$-in-CO$_2$ as gasification media yielded comparable operation
- Operational temperatures were slightly lower for O$_2$-CO$_2$–blown operation
- Tar and sulphur concentrations measured to <11mg/Nm$^3$ and <3ppm respectively