Fate of NO and Ammonia in Chemical Looping Combustion—Investigation in a 300 W Chemical Looping Combustion Reactor System

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1. INTRODUCTION

One approach to reduce CO₂ emissions is to implement carbon capture and storage (CCS), whose purpose is to capture CO₂ emissions and store them. The conventional methods of carbon capture are however highly energy-intensive processes and require a larger fuel input to sustain the increased energy demand. To achieve so-called negative CO₂ emissions, CCS can be conducted in combination with carbon-neutral fuels, i.e., biomass or waste streams of biogenic origin. This is called bioenergy with carbon capture and storage (BECCS).

1.1. Chemical Looping Combustion (CLC).

CLC is a novel combustion technology that can be used for the purpose of CO₂ capture, without the large energy penalty of conventional capturing methods. It consists of two inter-
connected fluidized bed reactors, an air reactor and a fuel reactor, where the oxygen carrier is cycled between the reactors (Figure 1). A typical reactor system with interconnected fluidized beds is shown in Figure 2.

In the air reactor, the oxygen carrier will be oxidized and the oxidation of a generic oxygen carrier with oxygen provided from the air can be represented by

$$2\text{Me}_x\text{O}_{y-1} + \text{O}_2 \rightarrow 2\text{Me}_x\text{O}_y$$ (1)

The gas stream out of the air reactor consists of N\textsubscript{2} and some remaining oxygen. The reaction in the air reactor, eq 1, is always exothermic. In the fuel reactor, the oxygen carrier will be reduced by the fuel.

The following two equations describe the reduction of a generic oxygen carrier with syngas:

$$\text{Me}_x\text{O}_y + \text{CO} \rightarrow \text{Me}_x\text{O}_{y-1} + \text{CO}_2$$ (2)

and

$$\text{Me}_x\text{O}_y + \text{H}_2 \rightarrow \text{Me}_x\text{O}_{y-1} + \text{H}_2\text{O}$$ (3)

The flue gases from the fuel reactor consist of mainly CO\textsubscript{2} and steam, where the steam can easily be condensed to create a CO\textsubscript{2}-rich stream. The reactions in the fuel reactor can be either endothermic or exothermic, depending upon the combination of the oxygen carrier and fuel. The net energy release in the CLC unit is the same as in conventional combustion.

### 1.2. Oxygen Carrier

The oxygen carrier is crucial in CLC and needs to perform its duty of transferring oxygen, without being rapidly mechanically or chemically degraded. The number of materials examined in laboratory are of the order of a thousand, whereas the number of materials examined in pilot operation are in the order of a hundred. Oxygen carrier materials investigated in pilot operation include manufactured monometallic oxide systems, i.e., oxides of Ni, Fe, Mn, Cu, and Co, normally using an inert support as well as several combined oxide systems, primarily combined Mn oxides. Moreover, many low-cost natural ores and waste materials have been examined, including ilmenite, manganese, and iron ores. Total pilot operation includes 12,000 h in 49 CLC pilots of sizes up to 3 MW, presented in more than 200 publications. Normally but dependent upon the oxygen carrier, the gas coming from the fuel reactor is not fully oxidized to CO\textsubscript{2} and H\textsubscript{2}O. Therefore, an oxy-polishing step may be required, where any remaining combustibles are burnt by oxygen added.

### 1.3. Chemical Looping with Oxygen Uncoupling (CLOU)

A variant of CLC is CLOU. In CLOU, the oxygen carrier spontaneously releases molecular oxygen to the gas phase instead of being reduced by the fuel. Thus, the reduction of the oxygen carrier in CLOU can be expressed as

$$2\text{Me}_x\text{O}_y \rightarrow 2\text{Me}_x\text{O}_{y-1} + \text{O}_2$$ (4)

whereas the oxidation of the oxygen carrier is similar to normal CLC. To work, the oxygen carrier needs to have an equilibrium partial pressure that is below the oxygen concentration in the air reactor and above the oxygen concentration in the fuel reactor, which is close to zero as oxygen is consumed by the fuel. Thus, a suitable CLOU material can have an equilibrium partial pressure corresponding to around 1–3% oxygen. Thus, the fuel burns in oxygen being released.

Potential oxygen carriers for CLOU are based on copper and combined manganese oxides. The latter includes manganese oxides combined with iron, nickel, silicon, calcium, and magnesium as well as ternary systems. Copper oxide is the most potent CLOU material, and full gas conversion has been reported from pilot operation. For combined manganese materials, best results thus far were obtained with calcium manganite, CaMnO\textsubscript{3}, a perovskite with oxygen-releasing properties. More than 700 h of operation with calcium manganates in seven different pilots of 0.3–1000 kW, mostly with gaseous fuels, clearly demonstrates that these materials show high performance, long lifetime, and, under good conditions, even full gas conversion.

### 2. NITROGEN OXIDES

Nitrogen oxides (NO\textsubscript{x}) include nitric oxide (NO), nitrogen dioxide (NO\textsubscript{2}), and nitrous oxide (N\textsubscript{2}O). NO is the main NO\textsubscript{x}...
Table 1. Nitrogen Species in the Bottom of a 12 MW Circulating Fluidized Bed Boiler

<table>
<thead>
<tr>
<th>Nitrogen Species</th>
<th>Bottom (ppm)</th>
<th>Flue Gas (ppm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>nitric oxide, NO</td>
<td>520</td>
<td>55</td>
</tr>
<tr>
<td>ammonia, NH₃</td>
<td>1090</td>
<td></td>
</tr>
<tr>
<td>hydrogen cyanide, HCN</td>
<td>240</td>
<td></td>
</tr>
<tr>
<td>isocyanic acid, HNCO</td>
<td>250</td>
<td></td>
</tr>
<tr>
<td>nitrous oxide, N₂O</td>
<td>15</td>
<td>140</td>
</tr>
</tbody>
</table>

Figure 4. Overview of important reactions in nitrogen chemistry starting from compounds such as HCN or NH₃: red, thermal NOₓ; blue, prompt NOₓ; brown, fuel NOₓ; coal; green, fuel NOₓ; biomass; yellow, N₂O formation/reduction; and cyan, reburning.

product formed during combustion. In normal combustion processes, there are three distinct paths of NOₓ formation: thermal NOₓ, prompt NOₓ, and fuel NOₓ, which are further described below.

2.1. Thermal NOₓ. Thermal NOₓ is created when molecular oxygen and molecular nitrogen react at high temperatures, according to the Zeldovich mechanism.

\[
\begin{align*}
N₂ + O & \rightarrow NO + N \quad (5) \\
N + O₂ & \rightarrow NO + O \quad (6) \\
N + OH & \rightarrow NO + H \quad (7)
\end{align*}
\]

The formation of thermal NOₓ is only significant at high temperatures, above 1500°C. In combustion, local high temperatures are associated with flames. In combustion with no fuel-bound nitrogen, thermal NOₓ is responsible for most of the NOₓ emissions produced.

In CLC, flames are absent and the temperature in both the air and fuel reactors is far below that needed for thermal NOₓ formation. In the fuel reactor, the absence of oxygen and the minute presence of N₂ would also prevent thermal NOₓ formation. Thus, thermal NOₓ formation can be excluded in CLC, except possibly for the oxy-polishing step, downstream of the fuel reactor.

2.2. Prompt NOₓ. In fuel-rich environments, hydrocarbon radicals, such as C and CH, are formed that can react with molecular nitrogen to produce carbon-containing nitrogen species:

\[
C + N₂ \rightarrow CN + N \quad (8)
\]

and

\[
\text{CH} + N₂ \rightarrow HCN + N \quad (9)
\]

NOₓ is subsequently formed by oxidation of these nitrogen compounds. This mechanism is referred to as prompt NOₓ. Prompt NOₓ only contributes to a small fraction of the total NOₓ emissions. In the air reactor, the formation of prompt NOₓ is not possible in the absence of volatile organic compounds. In the fuel reactor, N₂ is only present in minute amounts, and therefore, it would be highly unlikely that prompt NOₓ formation is significant in CLC.

2.3. Fuel NOₓ. Nitrogen is found in almost all solid fuels, is bound to the fuel structure, and can be released both from the char conversion as well as in the form of volatile matter released during pyrolysis. The temperature, the heating rate, and the fuel structure affect how nitrogen is divided between the volatile matter and the char. The nitrogen species that are present within the volatiles are mainly hydrogen cyanide (HCN) and ammonia (NH₃). For high-rank fuels, such as coal, HCN is considered to be the main nitrogen species released during pyrolysis, whereas for low-rank fuels, such as biomass, NH₃ stands for the majority of nitrogen species in the volatiles. In addition to the fuel type, the heating rate, and the final temperature, the residence time also affects the distribution of the nitrogen species released during pyrolysis.

The intermediate nitrogen species formed during combustion (HCN, NH₃, etc.) can both react with oxygen species to create NO and with NO produced to form N₂. For NH₃, the general reaction involves oxidation of NH₃ to NH radicals (i = 0, 1, and 2), where the radicals can be further oxidized to produce NO or react with NO to form N₂ (cf. Figure 3).

In situ gas measurements in the bottom of a 12 MW circulating fluidized bed boiler fired with bituminous coal are shown in Table 1. The concentrations measured indicate that the bottom zone is highly reducing, which is because there is a large bypass of the fluidizing air. The high velocity of the bypassing gas means that this flow is underrepresented in the gas measurement, which is made by a suction probe that is most of the time in the dense phase, where the fuel releases volatiles and the char removes oxygen. Thus, the gas concentrations measured do not represent the gas flow. On the basis of measured total carbon of around 21%, the concentrations in Table 1 can be normalized to 15% CO₂ by division by 1.4.

Even with normalization, the data in Table 1 indicate high concentrations of NO as well as other precursors discussed above. The comparison to the flue gas shows that the NO concentration falls dramatically, whereas N₂O rises. It can also be noted that there are several potential NO precursors present in high concentrations.

2.4. Nitrous Oxide, N₂O. N₂O emissions have been found to be of importance in fluidized bed combustion but of minor importance in other combustion systems, which is explained by the low temperature in fluidized bed combustion. Thus, an increase in the temperature from normal operating temperature at 850 to 925°C decreased N₂O by a factor of 3 at 2% O₂. The results presented here for N₂O come from work in a 12 MW circulating fluidized bed boiler. It was also found that N₂O could be decreased by more than 90% by afterburning, i.e., adding fuel that raises the cyclone temperature to 950°C. Five different fuels were added, and the decrease in N₂O versus the cyclone temperature in the range of 850–950°C was identical for these. With so-called reversed air staging, it was possible to reduce N₂O emissions from 60 ppm...
to just a few parts per million at 970°C. The method involved keeping the riser at slightly overstoichiometric conditions and then adding air after the cyclone outlet. Moreover, experiments with injection of N₂O in the bottom of the riser found that 80–97% of N₂O was reduced.

Further, there seems to be essentially no formation of N₂O when burning high-volatile fuels. Thus, at 950°C, 130–140 ppm of N₂O was found in the flue gas for bituminous coal and coke, whereas for wood chips, 0–10 ppm was found. With the high fuel reactor temperatures typical for CLC, e.g., >950°C, and the absence of oxygen, it can be expected that no N₂O will form, i.e., unless it is formed from reactions with the oxygen carrier.

2.5. Reburning. At understoichiometric conditions, NO can be reduced. Introduction of fuel immediately after the main combustion stage to create a fuel-rich zone can be used to reduce NO to HCN. The process is called reburning.

2.6. Summary. The important reactions for the nitrogen chemistry in combustion are summarized in Figure 4. The scheme is simplified and does not consider all potential or existing reaction paths. It includes the main precursors formed when fuels are devolatilized, HCN and NH₃. However, in addition to the gas-phase reactions outlined, various nitrogen compounds are also formed during the char combustion, including N₂, N₂O, NO, HCN, and NH₃.

As indicated above, there is good reason to assume that neither thermal NOₓ nor prompt NOₓ formation takes place in CLC. Moreover, the formation of N₂O is not likely, and in any way, a reduction of N₂O should be efficient at the high temperature and slightly understoichiometric conditions of a fuel reactor. In the case of fuel NOₓ, the main reactions leading to NO are obviously promoted by the presence of oxygen, which is absent in the fuel reactor. Moreover, the formation of oxidized nitrogen compounds during char conversion, i.e., gasification, is less likely in the absence of oxygen. At the slightly understoichiometric conditions of the fuel reactor, reburning of NO could be possible, depending upon the presence of radicals. As seen in Figures 3 and 4, the gas phase reactions involve a number of radicals, i.e., O, OH, H, N, and CH₃. However, there are two reasons to assume that radical concentrations are lower in fluidized beds: first, the temperature is lower than in normal flame combustion, and second, radicals may be quenched by the surface of the bed material. Johnsson investigated NOₓ formation and reduction in fluidized bed combustion (FBC) and concluded that it is unknown how the high solid concentration will affect the radical chemistry. Löffler et al. developed a mechanism describing NOₓ formation in a fluidized bed (FB), in which they proposed that the high solid concentration is causing radical quenching. The mechanism included the radical recombination reactions of O, H, OH, and HO₂ that occur on the surfaces of the solid bed material. Radical quenching is also included in the work of Källpinen et al., where NOₓ formation and reduction are modeled in FBC.

3. EQUILIBRIUM CONCENTRATIONS

In the absence of oxygen or under the more or less reducing conditions in a fuel reactor, any nitrogen species, i.e., except for N₂, is below or well below 1 ppm, as shown in Figure 5. Here, the gas is a 50:50 mixture of steam and CO₂ with 1% N₂, and the temperature is 950°C. Oxygen is stepwise-removed from the system, which gradually increases the gas content of CO and H₂. NO dominates at stoichiometry, whereas NH₃ becomes dominant as soon as the CO/(CO + CO₂) ratio becomes higher than 0.005. Equilibrium calculations were made with HSC Chemistry 5.11 for Windows. The conditions in the experiments presented in this paper all have a CO/(CO + CO₂) ratio within the range of 0–0.1, which means that any measurable amounts of NO or NH₃ represent a deviation from equilibrium.

4. CLC RESEARCH RELATED TO NO

NOₓ formation in CLC applications has been investigated experimentally, in both laboratory-scale fluidized bed reactors and connection with operation of 0.5–100 kW CLC units.

4.1. Batch Reactor Experiments. Ishida and Jin were the first to study NOₓ formation in CLC and found that no NO is found when oxidizing a nickel-based oxygen carrier at 1200°C.

Cheng et al. investigated the possible reactions between NH₃ and ilmenite in a laboratory-scale fluidized bed reactor, in the presence of syngas. They found that around 18% of NH₃ was converted to NO independent of the temperature in the investigated range, 850–950°C. The remainder was converted to N₂ except at 850°C, where small amounts of unconverted NH₃ were seen. When the NH₃ inlet concentration was lowered from 1 to 0.2%, there was a strong increase in the fraction of NO and a corresponding decrease in N₂. Not unexpected, the conversion to NO fell with the increasing syngas concentration in the range of 0–40%. However, even at the highest syngas concentrations, NO was formed. Also, the ability of ilmenite to reduce NO was studied as a function of the degree of oxidation and in the absence of fuel. Fully oxidized ilmenite did not reduce NO, and the reduction of NO was almost linear to the conversion of ilmenite.

Normann et al. studied the oxidation of NH₃ for three different oxygen carriers based on oxides of Fe, Mn, and Ni in a laboratory-scale fluidized bed reactor. The oxygen carrier was alternatingly exposed to syngas and 1% NH₃, with periods of inert gas in between. Thus, the oxidation of NH₃ could be examined as a function of the degree of conversion of the oxygen carriers. For NiO, no NO was formed independent of the oxidation level of the oxygen carrier. For iron oxide, the conversion to NO was initially around 40%, i.e., for fully oxidized iron oxide, and gradually decreased to zero as the oxygen carrier became fully reduced. For manganese oxide, on the other hand, the conversion was well above 60% for most of the conversion and only dropped to zero in the last part of the reduction.
Table 2. Nitrogen Compounds Measured in Operation of CLC Pilots

<table>
<thead>
<tr>
<th>reference</th>
<th>size (kW)</th>
<th>oxygen carrier</th>
<th>temperature (°C)</th>
<th>fuel</th>
<th>inlet N fraction</th>
<th>NOx (ppm, normalized)</th>
<th>N conversion to NOx (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Song et al.57</td>
<td>1</td>
<td>NiO</td>
<td>850—950</td>
<td>bituminous coal</td>
<td>1.03% ad</td>
<td>400—630</td>
<td></td>
</tr>
<tr>
<td>Song et al.57</td>
<td>1</td>
<td>NiO</td>
<td>850—950</td>
<td>anthracite</td>
<td>1.39% ad</td>
<td>1—20</td>
<td></td>
</tr>
<tr>
<td>Song et al.58</td>
<td>1</td>
<td>Fe2O3</td>
<td>970</td>
<td>bituminous coal</td>
<td>1.03% ad</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Song et al.58</td>
<td>1</td>
<td>Fe2O3</td>
<td>970</td>
<td>anthracite</td>
<td>1.39% ad</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Markström et al.</td>
<td>100</td>
<td>ilmenite</td>
<td>960</td>
<td>bituminous coal</td>
<td>1.65% daf</td>
<td>1000—2000</td>
<td>10—20</td>
</tr>
<tr>
<td>Linderholm et al.</td>
<td>100</td>
<td>ilmenite</td>
<td>950—970</td>
<td>wood char</td>
<td>0.4% daf</td>
<td>500 (850)</td>
<td>7—32</td>
</tr>
<tr>
<td>Linderholm et al.</td>
<td>100</td>
<td>ilmenite</td>
<td>960</td>
<td>petcoke</td>
<td>1% daf</td>
<td>3—8/20</td>
<td></td>
</tr>
<tr>
<td>Linderholm et al.</td>
<td>100</td>
<td>ilmenite</td>
<td>970</td>
<td>bituminous coal</td>
<td>1.6% daf</td>
<td>11 of gas/5.6 of total</td>
<td></td>
</tr>
<tr>
<td>Linderholm et al.</td>
<td>100</td>
<td>ilmenite + manganese ore</td>
<td>963—971</td>
<td>bituminous coal</td>
<td>1.6% daf</td>
<td>29 of gas</td>
<td></td>
</tr>
<tr>
<td>Linderholm et al.</td>
<td>100</td>
<td>iron ore</td>
<td>895—950</td>
<td>bituminous coal</td>
<td>1.6% daf</td>
<td>10 of gas</td>
<td></td>
</tr>
<tr>
<td>Linderholm et al.</td>
<td>100</td>
<td>manganese ore</td>
<td>950—979</td>
<td>bituminous coal</td>
<td>1.6% daf</td>
<td>4—11 of gas</td>
<td></td>
</tr>
<tr>
<td>Mendiara et al.65</td>
<td>0.5</td>
<td>ilmenite</td>
<td>875/930</td>
<td>lignite</td>
<td>0.6%</td>
<td>20</td>
<td>1</td>
</tr>
<tr>
<td>Mendiara et al.66</td>
<td>0.5</td>
<td>iron ore</td>
<td>875/930</td>
<td>pine sawdust</td>
<td>0.28%</td>
<td>4—14</td>
<td></td>
</tr>
<tr>
<td>Mendiara et al.66</td>
<td>0.5</td>
<td>iron ore</td>
<td>875/930</td>
<td>olive stone</td>
<td>0.18%</td>
<td>4—14</td>
<td></td>
</tr>
<tr>
<td>Mendiara et al.66</td>
<td>0.5</td>
<td>iron ore</td>
<td>875/930</td>
<td>almond shell</td>
<td>0.16%</td>
<td>4—14</td>
<td></td>
</tr>
<tr>
<td>Gu et al.67</td>
<td>1</td>
<td>Fe2O3</td>
<td>880—980</td>
<td>bituminous coal</td>
<td>1.03% ad</td>
<td>24—68</td>
<td>0.5—0.8</td>
</tr>
<tr>
<td>Bayham et al.68</td>
<td>25</td>
<td>Fe2O3</td>
<td>970</td>
<td>sub-bituminous coal</td>
<td></td>
<td>1148—1669</td>
<td>10—15</td>
</tr>
<tr>
<td>Ohlemüller et al.</td>
<td>1000</td>
<td>ilmenite and iron ore</td>
<td></td>
<td>coal and biomass</td>
<td></td>
<td>1.81 and 0.17%</td>
<td>0, 20, and 8</td>
</tr>
</tbody>
</table>

“daf = dry and ash-free.  ad = air dried.

Normann et al.55 investigated the oxidation of NH3 using an oxygen carrier with CLOU properties, copper oxide. The experiments were conducted in a laboratory-scale fluidized bed reactor with syngas as fuel, and the inlet fractions of NH3 and syngas were varied. They concluded that the nitrogen selectivity is highly dependent upon the oxidation level of the oxygen carrier; a higher oxidation level promotes the nitrogen selectivity toward NO. They also found that the nitrogen selectivity is affected by syngas; a higher concentration of syngas seems to hinder NO formation.

Mayrhuber et al.60 compared a rock ilmenite and a sand ilmenite, using both NH3 and NO with syngas as fuel. NO formation was found to decrease with an increased temperature, and the mass-based conversion of the oxygen carrier needs to be above 0.99 to achieve NO formation. Further, the outgoing concentration of NO was approximately the same when adding NO and NH3. The NO formation was investigated using two different approaches: one with alternating pulses of syngas and NO/NH3, similar to Normann et al.55 and in the other, pulses of syngas together with NO/NH3 were used. With this approach, it was possible to conclude that more NO was found in the presence of syngas, for the same degree of oxygen carrier conversion. This was clearly demonstrated for the addition of both NO and NH3. This indicates that the syngas or H2O and CO2 formed by the syngas oxidation hinder the reduction of NO.

4.2. Nitrogen Compounds in Continuous CLC Operation. Song et al.57 studied the conversion of fuel nitrogen in a 1 kW CLC unit using two coals and a Ni-based oxygen carrier. In the fuel reactor, all fuel nitrogen is converted to N2 for both fuels. This agrees well with the lab reactor results of Normann et al.55 above. However, NO was found in the air reactor as a result of the carryover of char. Song et al.58 also operated the 1 kW CLC unit with an iron ore, using bituminous coal and anthracite. Again, no NO was produced in the fuel reactor, while some NO is formed in the air reactor.

Markström et al.59 conducted solid fuel operation with a 100 kW CLC unit using bituminous coal and ilmenite as the oxygen carrier. They found that typically 10—20% of fuel nitrogen is converted to NO. Also, NO increased when the oxygen demand decreased, e.g., at increased solids circulation. Further, increased addition of the steam fluidizing the fuel reactor led to a rise in NO formation.

Linderholm et al.60 investigated NO formation in a 100 kW CLC unit with low-volatile solid fuels and ilmenite. With petcoke, they found that 3—8% of fuel nitrogen was converted to NO at 148 kW, while a 20% NO conversion was obtained at 72 kW. With wood char, fuel N conversion to NO was 7—32%. For both fuels, NO increased with solids circulation.

Linderholm et al.61 made nitrogen balances for CLC operation in a 100 kW unit with a Columbian bituminous coal and ilmenite as the oxygen carrier and were able to account for 62% of incoming fuel nitrogen. Nitrogen converted into gaseous compounds had the distribution of 1% HCN, 11% NO, and 26% NH3 with the remainder assumed to be N2. In a later operation with a Polish coal62 but with a mixture of ilmenite and manganese ore, the distribution was 0% HCN, 29% NO, and 11% NH3. Higher NO is expected, because the oxygen demand was significantly lowered in the presence of manganese ore. Further, operation with Columbian coal and an iron ore in this unit found 10% NO and 37% NH3.63 Operation with Columbian coal and sintered manganese ore found 4—11% NO and 15—29% NH3.64 NH3 fell and NO rose with rising circulation. Interestingly, NO was lower than in the previous cases, despite the fact that the oxygen demand was at a similarly low level as for ilmenite and manganese ore.
The ability to oxidize NH₃ and reduce NO is strongly linked to instance, highly reactive nickel oxide did not give any NO. The not necessarily linked to the ability to oxidize fuel gases; for entering the fuel reactor.

The NO reactor may form NO in the air reactor. CLC pilot operation shows that high concentrations of NO the degree of conversion of the oxygen carrier. Furthermore, between different oxygen carriers. The ability to form NO is not necessarily linked to the ability to oxidize fuel gases; for instance, highly reactive nickel oxide did not give any NO. The laboratory tests also shows that oxygen carriers can reduce NO. The ability to oxidize NH₃ and reduce NO is strongly linked to the degree of conversion of the oxygen carrier. Furthermore, CLC pilot operation shows that high concentrations of NO can form in the fuel reactor, despite the fact that equilibrium concentrations are very low. In addition, char leakage to the air reactor may form NO in the air reactor.

The purpose of this work is to expand the understanding of the NOx chemistry in CLC, by investigating the fate of NO and NH₃ in a 300 W CLC pilot operated with gaseous fuel. This is accomplished by adding NO/NH₃ to the gas stream entering the fuel reactor.

5. MATERIALS AND METHODS
5.1. Linz–Donawitz (LD) Slag. LD slag is a ground steel converter slag from the Linz–Donawitz process. Millions of tons of LD slag could make it relevant for use in CLC. Moldenhauer et al. investigated LD slag in chemical looping with syngas, methane, and biomass fuel. Full conversion of syngas at 900°C and 60% methane conversion at 950°C were obtained. CO₂ yields of 75–82% were achieved when operating with biomass fuel. LD slag was also used as bed material in the 12 MW circulating fluidized bed (CFB) boiler at Chalmers University of Technology to investigate oxygen-carrier-aided combustion (OCAC) with biomass.

The LD slag was delivered by SSAB Merox, Sweden. The elemental composition of LD slag and ilmenite is given in Table 3. The oxygen carriers were calcined in air at 950°C for 24 h to ensure full oxidation. The calcined particles were sieved to obtain around 500 g of particles smaller than 212 μm, for use in the 300 W reactor.

5.2. Ilmenite. Ilmenite has been subject to investigations in many studies related to CLC using various types of fuel since 2008, and more than 1500 h of CLC pilot operation with ilmenite has been reported. Ilmenite readily oxidizes syngas, with higher conversion of H₂ to CO. However, the conversion of CH₄ is moderate. Ilmenite is a low-cost natural iron titanium ore, which, in its natural state, mainly consists of FeTiO₃. This is also the reduced state. In the air reactor, ilmenite can be oxidized to its full oxidation state, according to

\[ 2\text{FeTiO}_3 + 1/2\text{O}_2 \rightarrow \text{Fe}_2\text{TiO}_4 + \text{TiO}_2 \]  

Just like LD slag, ilmenite has been used in OCAC, including 20 000 h of burning municipal solid waste in a 75 MW CFB boiler.

5.3. Chemical Looping Fluidized Bed Reactor System at 300 W. The experiments were performed in a dual fluidized bed 300 W CLC reactor system, with syngas fuel under continuous circulation of oxygen carrier particles. The schematic illustration of the 300 W chemical looping reactor used in this work is shown in Figure 6.

The reactor is 300 mm high, with a cross section of 25 × 25 mm for the fuel reactor (FR) and 25 × 42 mm for the air reactor (AR). The riser of the AR is 25 × 25 mm and is operated at a gas velocity sufficient to transport the particles upward, thus accomplishing the circulation of bed material. This is due to the difference in density between the two reactors to secure that no gas can leak between them. There are two wind boxes located in the bottom of each reactor to inject the gas via porous quartz plates, which act as gas distributors. There is a top part above both reactors, where the cross section of the gas path widens and gas velocity consequently decreases, allowing elutriated particles to fall back into the reactors. On the AR side, the section is designed to lead part of the particles into a loop seal that connects the AR to FR. On the FR side, particles drop directly back into the FR. From the bottom of the FR, particles pass through the lower loop seal and return to the AR, where the cycle starts over, completing the loop. The two loop seals are fluidized with inert gas, a total of 0.65 L/min of argon, when operating with fuel to ensure the fluidization and circulation of the particles. The exit pipe of the FR is connected to a water seal with a column height of less than 20 mm to maintain a slightly higher pressure, 0.1–0.2 kPa, in the FR compared to AR to minimize the risk of gas leakage from AR to FR. The bed temperatures of the AR and FR are measured by means of K-type thermocouples. The 300 W reactor is located inside an electrical furnace that controls the temperature. The fluidization and circulation of the particles are monitored via pressure difference measurement at different heights of the AR and FR.

5.4. NH₃ Measurements. The setup was devised to make it possible to measure ammonia directly using Dräger tubes. To avoid the dissolution of ammonia in condensed water, the gas stream must be dried while still being above the point of condensation. Further, the drying of the gas must not remove any ammonia. Therefore, the moist hot gas stream was passed over a desiccant, i.e., sodium hydroxide (NaOH), which is a strong base at the same time, allowing water vapor to be absorbed, while ammonia leaves the drying unit unreacted. A stream of gas from the FR was led through the drying step. This stream was well-insulated and heated to stay above 100°C to avoid any water condensation. After the drying, a Dräger tube is used to measure the concentration of ammonia in the dry gas stream. The
setup included a valve that was opened for 5 min for each measurement.

The reactor is run with syngas at a maximum flow of 1 L/min. Assuming that the syngas conversion to steam and CO₂ is 100% and that all of the flue gas is led through the drying unit for 5 min, 9 g of sodium hydroxide in total is needed for the reaction with CO₂ and absorbing water. On the basis of this, adequate amounts of drying agent were added to the moisture trap and exchanged regularly. Although CO₂ reacts with hydroxide, it does not interfere with the procedure because there will be enough hydroxide in the drying unit to keep the pH very low and, thus, prevent reaction with ammonia. Figure 7 shows a schematic plan of the setup. The online gas analyzer

**Figure 7.** Setup for NH₃ measurements.
is equipped with a pump that sucks 1 L/min. With the opening of the valve leading to the moisture trap and closing of the other valve leading to the gas analysers, the gas is forced through the moisture trap and can be sampled by Dräger tubes.

5.5. Circulation. The solids circulation was estimated using the char injection method, whereby 0.5 g of char was added to the fuel reactor and the circulation is derived from the response in the CO2 concentration from the air reactor. For each oxygen carrier, two air flows were used in the air reactor, 8 and 9 L/min, giving two different circulations below called high and low. For LD slag, the low and high circulations were 120 and 144 g/min, and for ilmenite, they were 75 and 84 g/min.

The solids inventory of the fuel reactor was estimated to 100 g for LD slag and 95 g for ilmenite, corresponding to a residence time in the fuel reactor of somewhat below 1 min for LD slag and a little more than 1 min for ilmenite.

5.6. Experimental Procedure. Prior to the experiments, oxygen carriers were activated with syngas, 50% hydrogen in CO, at 900°C for 2–3 h. Oxygen carriers were circulating during the activation, and the air flow rate to the AR was 8 L/min. Reaching a stable concentration of CO2 from the FR was used to confirm the activation.

The study involved the addition of a stream of NO or NH3, diluted in argon, to the fuel reactor, normally at a concentration of 1% but in a few cases at 0.2%. The tests involved the addition of these flows to the fuel reactor during chemical looping operation of the 300 W reactor system, normally together with fuel and in some cases also an additional inert flow. Further, a part of the inert gas from the fluidization of the loop seals also mixes into the gas stream in the fuel reactor. Normally, the air reactor was fluidized with air that was diluted with gas from the loop seals. The test matrix involved two oxygen carriers, two temperatures, two circulation flows, and three or four fuel flows.

Furthermore, experiments were made without fuel addition as well as with an empty reactor system.

5.7. Data Evaluation. Gas conversion of the syngas was based on the CO conversion to CO2, because H2 was assumed to be essentially complete for these tests. Thus, gas conversion was attained as

$$\eta_{\text{gas}} = 1 - \frac{x_{\text{CO}}}{x_{\text{CO}_2}}$$

(11)

where \(x_i\) is the volumetric concentration of species \(i\).

6. GAS CONVERSION IN CHEMICAL LOOPING OPERATION

The work involved almost 20 h of fueled operation with LD slag as well as almost 20 h with ilmenite. Figure 8 shows the gas conversion in operation with LD slag with either NH3 or NO addition. Conversion was complete for the lowest fuel flow. As expected, it was higher for a high temperature and
high circulation. The trends are similar for the tests with NH₃ and NO. Given the small amount of NO or NH₃ added, similar results would be expected for both cases. There is no consistent difference: in two cases, the tests with NO show higher conversion, and in the other two cases, they show lower or slightly lower conversion.

Similarly, the gas conversion for ilmenite is shown in Figure 9. It is clear that the conversion is somewhat lower than for LD slag, but in all cases, the gas conversion is above 92%. The difference is more pronounced at lower fuel flows, e.g., 0.5 Lₐ/min, where LD slag has full conversion in all cases, whereas ilmenite only shows full conversion in one case. A possible explanation could be that LD slag has higher reactivity but lower oxygen transfer capacity. It is also evident that the temperature has a larger impact on conversion for ilmenite. Similar to LD slag, higher circulation is consistently giving higher conversion.

7. NH₃ TESTS WITH AN EMPTY REACTOR

To validate the measurement procedure, measurements were made with an empty reactor, i.e., in the absence of an oxygen carrier. The investigation involved the addition of NH₃ in argon, at both varying the temperature and constant temperatures of 850 and 900°C.

7.1. Empty Reactor with NH₃: Temperature Ramping. The reactor temperature was increased gradually while 1.5 Lₐ/min of 1% NH₃ in argon was added to the fuel reactor. The aim was to observe when ammonia starts to decompose in the fuel reactor. Below 625°C, the measured level of NH₃ was 1%, i.e., 10,000 ppm, and hydrogen was 0%. However, at 625°C, NH₃ decreased to 9500 ppm and 0.1% hydrogen appeared. At 700°C, NH₃ dropped to 15 ppm and the gas chromatograph (GC) showed 0.3% nitrogen, indicating the breakdown of NH₃ to N₂ and H₂. When the temperature reached 850°C, hydrogen was 0.31% and NH₃ was still 15 ppm.

7.2. Empty Reactor with NH₃: Stable Temperature of 900 and 850°C. NH₃ was added while the empty reactor was stable at 900°C to observe the decomposition of ammonia. The added flow was higher than in the temperature ramping, 1.5 Lₐ/min of 1% NH₃ plus 1 Lₐ/min of pure argon, thus giving 6000 ppm. Concentrations were measured every 5 min, and the results are shown in Table 4.

The temperature was then lowered to 850°C, and a stable concentration of 55 ppm was seen, while the GC found 0.36% hydrogen.

Thus, around 99% of ammonia breaks down at both 850 and 900°C. This is consistent with thermodynamic equilibria, showing that both NO and NH₃ have negligible concentrations in the absence of oxygen (see Figure 5). However, the expected concentration of H₂ for breakdown to N₂ and H₂ would be 0.9%, but measured values are 10–20% of that. A possibility could be that H₂ reacts with the reactor walls.

The residence time of the gas in the wind box and the pipe leading to the windbox is small compared to the residence time in the reactor and the top part above the reactor. In addition, the incoming gas is cold, and the wind box is cooled; therefore, it is likely that the major breakdown of ammonia takes place after entering the fuel reactor. Consequently, it is assumed that all or a major part of ammonia reaches the bed of the oxygen carrier in the tests below.

8. EXPERIMENTS WITH LD SLAG

8.1. Tests with NO or NH₃ without Fuel. A total of 1% of NH₃ in argon was added to the fuel reactor, while the oxygen carrier was circulated between the air and fuel reactors at two different temperatures, 850 and 900°C. It was found that 6–10% of NH₃ was oxidized to NO. A high circulation and high temperature gave higher conversion to NO. There was no NH₃ in the gas sampled.

Similarly, 0.2% of NO was added to the fuel reactor. The NO concentration measured corresponded to 87–93% of incoming NO. The numbers were lower for 900°C and also lower for high circulation. It can be concluded that a majority

Table 4. Gas Concentrations Measured

<table>
<thead>
<tr>
<th>sample</th>
<th>T (°C)</th>
<th>NH₃ (ppm)</th>
<th>H₂ (%)</th>
<th>NO (ppm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>900</td>
<td>15</td>
<td>0.1</td>
<td>0</td>
</tr>
<tr>
<td>2</td>
<td>900</td>
<td>15</td>
<td>0.12</td>
<td>0</td>
</tr>
<tr>
<td>3</td>
<td>900</td>
<td>50</td>
<td>0.13</td>
<td>0</td>
</tr>
<tr>
<td>4</td>
<td>900</td>
<td>70</td>
<td>0.19</td>
<td>0</td>
</tr>
</tbody>
</table>

Figure 10. NH₃ addition at 900°C and high circulation.
of NO passes through the fuel reactor when the oxygen carrier is fully oxidized. NO is not expected to be reduced by the fully oxidized oxygen carrier. The deviation from 100% could be a result of error in the mass balance. When bed material is circulated, inert gas is added to upper and lower loop seals and a part of the gas goes to the air reactor, while the rest of the gas goes to the fuel reactor. Thus, the gas to the fuel reactor is diluted with this flow, and the mass balance will be dependent upon the assumption of the gas fraction going to the fuel reactor. This fraction may also be affected by the temperature, solids circulation, and pressure difference.

8.2. Tests with LD Slag, Fuel, and NH₃. Figures 10−13 show the effect of the temperature, fuel flow rate, and circulation rate on the NO emissions with the addition of 10,000 ppm of NH₃ in argon. The argon/NH₃ flow was 1 L/min, and the syngas flow is shown in the diagrams. Further, the inert gas added to the loop seals is assumed to give an additional 0.42 L/min. Three syngas flows were used, 0.5, 0.75, and 1 L/min, which means that NH₃ was diluted by a factor of 1.9–2.4 depending upon syngas flow. Figures 10 and 11 show the results for the higher solids circulation. The highest conversion of NH₃ to NO was obtained for the lowest fuel flow, where around 17% is converted to NO at the higher temperature and 8% is converted to NO at the lower temperature. On the other hand, there was little NO at the highest fuel flow. This is also expected because higher fuel flows mean a more reduced oxygen carrier with reduced ability to oxidize NH₃ to NO. The results for the lower circulation (Figures 12 and 13) are fairly similar to the higher flow. Several of the shifts to higher fuel flow are accompanied by a peak in NO. The cause for this peak is not clear, but the peak subsides as CO gradually increases after the shift. The increase in NO when CO again decreases at 215 min should also be noted (Figure 10). Also, Figures 11−13 show how decreases in NO are associated with increases in CO. For CO concentrations above 0.2–0.3, the NO concentrations are very low.

The results are summarized in Figure 14, including data for no fuel addition. The data for fuel addition were each collected in sequence on the same day, whereas the data for zero fuel flow were collected at a later date. The lower oxidation for NO
in the absence of fuel is somewhat unexpected and may possibly be an artifact explained by some difference in the experimental conditions, e.g., lower bed content because of the loss of particles or loss in reactivity. However, an additional test series was made at 900°C with high circulation, where the fuel flow was varied from zero the same day, with the following sequence of fuel flow: 0, 0.5, 0.75, 1, 0, 0.5, 1, and 0. The sequence is shown in red in Figure 14, and there is no significant difference between no fuel and 0.5 L/min, which should be noted. In contrast to the test in Figure 10, this sequence was operated with a constant gas flow of 2 L/min to the fuel reactor. While the flow with NH₃ was constant at 1 L/min, lower fuel flows were compensated by the addition of argon. Thus, for flows other than 1 L/min, both fuel and NH₃ are more diluted in comparison to the other tests. The general expectation of course is that a higher state of oxidation would promote the oxidation of NH₃ to NO. However, the detailed reaction paths are not known, and the presence of fuel also involves the presence of high concentrations of steam and CO₂, which could potentially promote NO formation or hinder reduction of NO. As previously noted, Mayrhuber et al. found increased conversion of NH₃ to NO in the presence of syngas, for a similar degree of oxidation of the oxygen carrier ilmenite.⁵⁶

However, with the spread in the data, no safe conclusion can be made whether the formation of NO really increases when going from no fuel flow to low fuel flow, i.e., 0.5 L/min.

There is a clear decrease in oxidation to NO as more fuel is added. Conversion to NO is consistently higher at a higher temperature, whereas the effect of circulation is small. The oxidation to NO is strongly correlated to the CO concentration, as shown in Figure 15. In the presence of CO, there is essentially no formation of NO, whereas in the absence of CO, the formation of NO is significant in all cases.

8.3. Tests with LD Slag, Fuel, and NO. Figures 16 and 17 show the effect of the fuel flow rate and circulation rate on the NO emissions with the addition of 10 000 ppm of NO in argon at 900°C. The argon/NO flow was 1 L/min, and the syngas...
Figure 15. NO and versus CO for the addition of 1 L/min of 1% NH₃ in argon.

Figure 16. NO addition at 900°C and high circulation.

Figure 17. NO addition at 900°C and low circulation.
flow is shown in the diagrams. Further, the inert gas added to
the loop seals is assumed to give an additional 0.42 L\textsubscript{n}/min.
Three syngas flows were used, 0.5, 0.75, and 1 L\textsubscript{n}/min, which
means that NO was diluted by a factor of 1.9–2.4 depending
upon syngas flow. It is clear that the reduction of NO increases
when more syngas is added.

Similar results are found at 850°C, although the breakdown
of NO is higher at lower temperatures.

The results are compared in Figure 18. It is clear that lower
circulation and higher fuel flow give increased reduction of
NO. This is expected; more syngas and lower circulation
should both contribute to give a more reduced oxygen carrier
as well as higher concentration of reduced gaseous species. It is
also clear from the figure that a lower temperature is associated
with a higher breakdown of NO. At a lower temperature, the
CO concentration is also higher.

Normally, a lower temperature is associated with slower
reactions, but there are two possible explanations for this.
Either the presence of CO enhances the breakdown, or the
surface of the oxygen carrier is more reduced, e.g., as a
consequence of less rapid transfer of oxygen inside the oxygen
carrier. The latter then also explains higher CO.

Figure 19 shows the correlation between NO and CO for the
tests shown in Figure 18. It is clear that, in the presence of
CO, the breakdown of NO is complete, whereas in the absence
of CO, at least some NO escapes through the bed.

8.4. Addition of NO and NH\textsubscript{3} in the Absence of Fuel
but with Reduced Oxygen Carrier Particles. From the
studies with the addition of fuel, it is clear that oxidation of
NH\textsubscript{3} to NO is lower in the presence of fuel. The experiment
shown in Figure 20 was performed to see whether the reduced
oxygen carrier is able to oxidize NH\textsubscript{3} to NO also in the absence
of fuel. The temperature was 900°C, and the gas flow to the air
reactor was set to give high circulation. Initially, the 300 W
CLC unit was operated with syngas. Then, the air flow to the
air reactor is changed to argon, thus ending the regeneration of
the oxygen carrier coming from the fuel reactor. For 2 min, the
system is run with continued syngas flow and no air flow, thus
reducing the oxygen carrier. After 2 min, the syngas flow is
stopped, and with only inert gases added, the reduced state of
the oxygen carrier is preserved. A few minutes later, NH\textsubscript{3}
addition is started, and it is clear that there is no oxidation of NH\textsubscript{3} to NO. After 12 min, feeding of air to the air reactor commenced, and 3 min later, NO starts to rise and stabilize at around 1000 ppm of NO, which corresponds to a conversion of NH\textsubscript{3} to NO of 15%.

The 2 min without regeneration only gives a partial reduction of the oxygen carrier, which should be noted. Thus, the gas conversion is essentially complete during the 2 min, and only at the end of the 2 min, a minor peak of 0.1% CO\textsubscript{2} was seen. This is not shown in the figure. A syngas flow of 1 L\textsubscript{min}/min during 2 min will remove 1.4 g of oxygen from the oxygen carrier. Assuming approximately 500 g of oxygen carrier, this means a reduction in mass-based conversion of approximately 0.3%. However, the oxygen carrier is not fully oxidized when the 2 min period starts, which means that the mass-based reduction is higher than 0.3%. This can be compared to the theoretical maximum mass-based conversion of LD slag of around 1%\textsuperscript{70} (see Figure 21). The theoretical maximum seems to be below that determined from experiments. It is not clear why, but it may possibly be associated with the formation of various mixed oxide phases involving Fe, Ca, Si, Mn, and Al.

Similar to the test above, where it was examined whether the partly reduced oxygen carrier could oxidize NH\textsubscript{3}, it was examined whether the partly reduced oxygen carrier could reduce NO. The test is shown in Figure 22, and again, a period of 2 min of reduction of the oxygen carrier is made. A few minutes after this, the addition of gas containing NO is started. It is evident that all NO is converted. Several minutes later, air addition to the air reactor is started, and after some delay, the reduction of incoming NO stops and the NO concentration quickly exceeds the maximum concentration range of the NO analyzer, explaining the plateau at 2400 ppm. At this point, the concentration of NO in the inlet gas is lowered to 0.2%, i.e., 2000 ppm. Bearing in mind that the gas is diluted with gas coming from the loop seals, the NO concentration then stabilizes at a level that is reasonably close to that expected with no reduction.
9. EXPERIMENTS WITH ILMENITE

9.1. Tests with Ilmenite and NH₃, with and without Fuel. Similar to the experiments with LD slag, the effect of fuel flow on the oxidation of NH₃ was investigated for ilmenite. The test matrix was expanded with an additional low flow, 0.25 Lₐ/min, which is half of the lowest flow used for LD slag. In contrast to the LD slag results, no significant conversion of NH₃ to NO was found for any of the fuel flows. The results are summarized in Figure 23. The levels of NO in the presence of fuel are all below 0.4% of added NH₃, which is also below 1% of the measurement range of the analyzer. The very low levels measured are believed to be an artifact caused by cross sensitivity with CO, which could also explains the consistent increase in NO with fuel flow. It can be concluded that NO levels during all cases of fuel addition are very low and below the detection limit.

9.2. Tests with NH₃ and Ilmenite with No Fuel but with a Reduced Oxygen Carrier. To examine whether the oxygen carrier in its partly reduced form but in the absence of fuel is able to oxidize NH₃ to NO, the oxygen carrier was reduced before introducing NH₃. The procedure is similar to the test LD slag previously shown in Figure 20 and is shown in Figure 24. First, syngas is added to the fuel reactor, and then air addition to the air reactor is stopped, while fuel addition is allowed to proceed for another 2 min to reduce the oxygen carrier. The CO concentration during fuel addition, not shown, was 0.5−0.6% both before and after turning off the air. After the stop of fuel addition, the reduced state is conserved because only inert gas is added to the air and fuel reactors. After a few minutes, the addition of NH₃ to the fuel reactor is started. The figure clearly indicates that the reduced oxygen carrier cannot oxidize any NH₃ to NO. After 3 min of NH₃ addition, fluidization of the air reactor is shifted from argon to air. Oxidation of NH₃ only starts after several minutes of air addition to the air reactor and then slowly increases. Thus, the figure indicates that oxidation to NO only happens when ilmenite is highly oxidized.

In comparison to the results shown with LD slag (Figure 20), it is clear that the return of the ability to oxidize NH₃ is much slower for ilmenite. An important difference between ilmenite and LD slag is that the maximum mass-based conversion of ilmenite is significantly higher, 5% for pure
ilmenite and 1.7% for fully phase-separated iron and titania oxides.\textsuperscript{82} The maximum mass-based conversion of ilmenite in actual operation is likely in the range of 3–4%. However, in both cases, the oxygen carrier was reduced for 2 min, which should give a similar change in mass-based conversion. However, the mass-based conversion at the starting point of the reduction is not necessarily the same.

9.3. Tests with Ilmenite and NO, with and without Fuel. Similar tests with and without fuel flow and a flow of 1 L\textsubscript{N}/min with 1% NO were made. Figure 25 shows the ratio of NO\textsubscript{out}/NO\textsubscript{in} versus fuel addition. Within the uncertainty of the mass balance, it can be said that, in the absence of fuel, all NO seems to pass through the bed, but in the presence of fuel, all NO is reduced independent of the fuel flow, temperature, or circulation. In all cases, measured NO\textsubscript{out}/NO\textsubscript{in} is 0.003 or lower. Just as with NH\textsubscript{3} addition previously shown in Figure 23, there is a slight increase with fuel flow, which is believed to be caused by cross sensitivity between NO and CO. NO measured is in these cases below 20 ppm, i.e., less than 1% of the measurement range of 0–2700 ppm of the analyzer.

In contrast to the results with LD slag, where some NO always survived the passage through the bed of the fuel reactor when CO was zero (see Figure 18), here, NO is zero in six cases where CO is zero. In particular, it can be noted that, while some NO passed through the bed in most of the cases at fuel additions between 0.5 and 1 L\textsubscript{N}/min for LD slag, incoming NO was fully reduced by ilmenite, even at the low fuel flow of 0.25 L\textsubscript{N}/min.

Thus, it is clear that ilmenite is much more effective in reducing NO than LD slag.

Similar to the test with NH\textsubscript{3} and reduced oxygen carrier, it was investigated whether the reduced oxygen carrier could reduce NO. To examine whether the oxygen carrier in its partly reduced form but in the absence of fuel is able to reduce NO, the oxygen carrier was reduced before introducing NO. The procedure is shown in Figure 26. First, syngas is added to the fuel reactor, and then air addition to the air reactor is stopped, while fuel addition is allowed to proceed for another 2 min to reduce the oxygen carrier. After the stop in fuel addition, the reduced state is conserved because only inert gas is added to the air and fuel reactors. A few minutes after the
Figure 26. Ability of the reduced oxygen carrier to reduce NO in the absence of fuel.

10. DISCUSSION

The most conspicuous result of the study is the fact that NO is completely reduced and that no NH$_3$ was oxidized to NO at any fuel flow when using ilmenite. This is in agreement with the pilot results of, i.e., Ohlemüller et al. but in disagreement with the results of Mendiara et al., Linderholm et al., and Markström et al. The latter found 1000–2000 ppm of NO (see Table 2). Most likely, this is explained by the fluidizing conditions, allowing for a significant fraction of the volatiles to bypass the bed. As pointed out in section 3, all nitrogen compounds, except N$_2$, are well below 1 ppm at equilibrium under the conditions in a CLC fuel reactor.

11. CONCLUSION

The oxidation of NH$_3$ and the breakdown of NO were studied in operation of a 300 W chemical looping reactor system, by adding either NH$_3$ or NO to the fluidization gas. Two different oxygen carriers, LD slag and ilmenite, were compared under varying fuel flows, temperatures, and solids circulation.

Experiments with ammonia addition and LD slag indicated the following: (1) Oxidation to NO fell with fuel flow and was essentially zero at the highest fuel flow. (2) Oxidation to NO rose with increasing temperatures. (3) The effect of solids circulation was less clear. At the higher fuel flows, increased circulation raised conversion to NO, whereas the effect was the opposite at the lowest fuel flow. (4) In absence of CO, NO was always formed, but when conversion was incomplete, i.e., in the presence of CO, conversion to NO was always less than 1%. (5) In the absence of fuel but with the oxygen carrier partially reduced, no NO was formed.

Experiments with NO addition and LD slag indicated the following: (1) Reduction of NO increased with fuel flow. (2) Reduction of NO was higher at the low temperature. (3) Reduction of NO was higher at low circulation. (4) In the absence of CO, NO was not completely reduced, whereas in the presence of CO, NO was fully reduced. (5) In the absence of fuel but with a partly reduced oxygen carrier, NO was completely reduced.

Experiments with ammonia addition and ilmenite indicated the following: (1) In contrast to LD slag, there was no oxidation to NO at any fuel flow, even though the lowest fuel flow was half of the lowest fuel flow used for LD slag. Thus, the effect of the temperature, circulation, and fuel flow could not be assessed. (2) This also means that, in contrast to LD slag, there was no oxidation to NO for the cases when CO was zero. (3) In the absence of fuel but with the oxygen carrier partially reduced, no NO was formed. When air addition was started, it took 5 min until NO started to increase slowly. This contrasts with LD slag, where oxidation to NO started after 3 min with a rapid increase.

Experiments with NO addition and ilmenite indicated the following: (1) In contrast to LD slag, NO was completely reduced at all fuel flows, even though the lowest fuel flow was half of the lowest fuel flow used for LD slag. Thus, the effect of the temperature, circulation, and fuel flow could not be assessed. (2) This also means that, in contrast to LD slag, reduction of NO was complete also when CO was zero, i.e., in six cases. (3) In the absence of fuel but with a partly reduced oxygen carrier, NO was completely reduced. When air addition was started, it took 4 min until NO started to increase, in contrast to LD slag, where a steep increase started after 3 min.
Thus, ilmenite is much more efficient in reducing NO and much less prone to oxidize NH₃ to NO. The latter is somewhat unexpected, considering that LD slag is more efficient in oxidizing CO.

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