Alternative deNOx catalysts and technologies

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Publication date:
2010

Document Version
Publisher's PDF, also known as Version of record

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Alternative \( \text{deNO}_x \) catalysts and technologies

Johannes Due-Hansen

Ph.D. Thesis
A section in *Zirconia-based catalyst systems* (section 3.2), regarding enhanced acidification of the sulfated zirconia originally presented here, cannot currently be made freely available due to a concurrent patent application. The material has thus been removed from this version.
Preface and acknowledgments

This thesis has been submitted in candidacy for the Ph.D. degree in chemistry at the Technical University of Denmark (DTU). The majority of the work summarized herein was carried out at the Department of Chemistry, from June 2007 to May 2010 under the joint supervision of Assoc. Prof. Anders Riisager and Prof. Rasmus Fehrmann. This project is financed through a Ph.D.-scholarship from DTU, Energinet.dk, DONG, Vattenfall and Haldor Topsøe.

During March and April 2009, I worked with the group of Pedro Ávila at Instituto de Catálisis y Petroquímica, Consejo Superior de Investigaciones Científicas (ICP-CSIC), Campus de Universidad Autónoma de Madrid in Spain.

First and foremost, I would like to thank my two supervisors for many fruitful discussions and suggestions during the project and for a constant source of inspiration.

I would further like to thank Steffen Buus Kristensen and Siva Sankar Reddy Putluru (DTU Chemistry) for a productive lab-collaboration and many interesting catalysis discussions, Olivier Nguyen van Buu (DTU Chemistry) for lending me his vast expertise in organic synthesis, and Kenny Ståhl (DTU Chemistry) for his help and inspiration with X-ray diffraction and Rietveld refinement. Warm thanks to Soghomon Boghosian (FORTH/ICE-HT and University of Patras, Greece), Rolf W. Berg (DTU Chemistry), Bodil Holten (DTU Chemistry), and Berit Wenzell (DTU-CEN) for an always enjoyable collaboration. Special Thanks to Pedro Ávila and Søren Birk Rasmussen (ICP-CSIC) for many inspiring ideas and an exciting collaboration from which I have learned a lot. Furthermore, I am indebted to Julie Skotte for assistance with Adobe Photoshop and Illustrator issues, as well as moral support.

I am grateful to the Travel Foundation of The Danish Chemical Society, the foundation of Fabrikant P.A. Fisker, the Idella foundation and Berg, Nielsens scholarship, for financial support for the external stay.

I have been very fortunate to have some dedicated students, Selcuk Mert, Søren Heintz and Thorey Anna Grétarsdóttir, with whom I have collaborated on various projects throughout the project period.

Finally I would like to thank all the people at the department and at the Centre for Catalysis and Sustainable Chemistry (CSC) for advice, feedback, co-work, and an always enjoyable atmosphere which has made the past three years memorable.

Johannes Due-Hansen
Lyngby, July 26, 2010
Abstract

The present thesis entitled *Alternative deNO₃: Catalysts and technologies* revolves around the topic of removal of nitrogen oxides. Nitrogen oxides, NOₓ, are unwanted byproducts formed during combustion (e.g. in engines or power plants). If emitted to the atmosphere, they are involved in the formation of acid rain and photochemical smog. Some basic concepts and reactions regarding the formation and removal of NOₓ are presented in chapter 1 and 2.

Two approaches are undertaken in the present work to reduce the emission of NOₓ: by means of catalytic removal, and by NO absorption in ionic liquids.

The commercial catalyst used for the selective catalytic reduction (SCR) of nitrogen oxides exhibits high activity and selectivity towards N₂. However, the vanadia-titania-based catalyst used is very sensitive to deactivation by alkali-species (primarily potassium), which are typically present in high amounts in the flue gas when biomass is combusted. By co-firing with large amounts of CO₂-neutral straw or wood (to meet stringent CO₂ emission legislation), the lifetime of the traditional SCR catalyst is thus significantly reduced due to the presence of deactivating species originating from the fuel. To develop a catalyst less susceptible to the poisons present in the flue gas, a number of catalysts have been synthesized and tested in the present work, all based on commercially available supports. A highly acidic support consisting of sulfated zirconia was chosen based on preliminary studies. A number of different active species distributed on the support were investigated, such as iron, copper and vanadium oxides. However, based on the catalysts performance in the SCR reaction and their resistances towards potassium, the most promising candidate of the formulations studied was the vanadia-loaded catalyst, i.e. V₂O₅·SO₄²⁻·ZrO₂. This work, together with an introduction to the catalytic removal of NOₓ, are described in chapter 3.

The remainder of the first part is concerned with the catalytic NOₓ removal (chapter 4) and it addresses the upscaling of the best catalyst candidate. The catalyst was mixed with the natural binding clay (sepiolite) to upscale the selected catalyst to the monolithic level, suitable for installation in gas stream with high flows, e.g. a flue gas duct of a power plant. A series of catalyst pellets with increasing levels of sepiolite were produced to evaluate the optimum mixing ratio. Based on these results, a monolith containing V₂O₅·SO₄²⁻·ZrO₂ in 25 wt% sepiolite was produced, and evaluated with respect to the influence of space velocity, reaction temperature, and NH₃/NO feed ratio on the NO reduction efficiency.

The last part of this thesis deals with the non-catalytic sorption of NOₓ in ionic liquids, collected in chapter 5. Since no previous studies of the absorption of NO in ionic liquids have been reported, a preliminary study was conducted to identify suitable solvents. Two result-
ing imidazolium-based candidates, namely [BMIM]OAc and [BMIM]OTf, were selected due to their impressively high sorption capacities. Both solvents examined here revealed solubilities about twofold higher than those previously reported for e.g. CO₂-capture in ionic liquids. Especially the [BMIM]OAc demonstrated extraordinary absorption capabilities, being able to retain around four NO molecules per molecule ionic liquid. However, [BMIM]OTf exhibited promising behavior due to its reversible absorption/desorption properties. This in principle allows recycling of the ionic liquid as well as harvesting the NO. The accumulated NO could hereby be used in e.g. the synthesis of nitric acid allowing production of value-added chemicals from waste flue gas effluent. Although additional understanding of the mechanisms of the presented system is required, the perspective of a selective NO stripping technology is a very interesting alternative to the catalytic removal of NO from industrial flue gases.
Foreliggende afhandling med titlen *Alternative deNO\textsubscript{x}-katalysatorer og -teknologier* omhandler fjernelse af nitrogen oxider fra røggasser. Nitrogen oxider, NO\textsubscript{x}, er uønskede biprodukter dannet under forbrænding, f.eks. i motorer eller kraftværker. Ved udledning til atmosfæren er de medvirkende til dannelsen af syreregn og fotokemisk smog. Nogle grundlæggende koncepter og reaktioner vedrørende dannelsen og fjernelsen af NO\textsubscript{x} er beskrevet i kapitel 1 og 2.

To forskellige metoder til at reducere emissionen af NO\textsubscript{x} bliver undersøgt i denne afhandling: katalytisk fjernelse samt absorption i ioniske væsker.

Den comercielle katalysator som idag primært bliver brugt i den selektive katalytiske reduktion (SCR) af nitrogen oxider udviser høj katalytisk aktivitet og selektivitet for omdannelse til N\textsubscript{2}. Men denne vanadia-titania baserede katalysator er meget følsom overfor deaktiverende alkali metal specier (primært kalium), som typisk er tilstede i høje koncentrationer i røggassen når biomasse afbrændes. Afbrænding af store mængder CO\textsubscript{2}-neutralt træ eller halm (for at imødekomme CO\textsubscript{2} emissionskrav) vil derfor reducere levetiden af den traditionelle SCR katalysator betydeligt pga. tilstedeværelsen af deaktiverede specier fra brændslet. For at udvikle en katalysator, der er mindre følsom overfor forgiftende stoffer i røggassen, er en række katalysatorer blevet fremstillet og testet, alle baseret på kommercielt tilgængelige bærematerialer. En særlig bærer bestående af sulfatiseret zirkonia blev udvalgt, baseret på indledende studier. En række forskellige aktive metaller fordelt på bæreren blev undersøgt, såsom oxiderne af jern, kobber og vanadium. Baseret på katalysatorernes virkningstid og deres modstandsdygtighed overfor kalium, var den vanadium-baserede katalysator (V\textsubscript{2}O\textsubscript{5}-SO\textsubscript{2}–4-ZrO\textsubscript{2}) den mest lovende kandidat. Dette arbejde, sammen med en introduktion til den katalytiske fjernelse af NO\textsubscript{x}, er beskrevet i kapitel 3.

Resten af første del (kapitel 4), som omhandler den katalytiske fjernelse af NO\textsubscript{x}, håndterer opskalingen af den mest lovende katalysator, V\textsubscript{2}O\textsubscript{5}-SO\textsubscript{2}–4-ZrO\textsubscript{2}. Katalysatoren blev sammenblandet med en naturlig lerbinder (sepiolit) for at opskalere den valgte katalysator til monolit niveau, som er egnet til atindsættelse i gasstrømme med høje hastigheder f.eks. i en røggaskanal på et kraftværk. En serie af katalysator-ekstrudater med stigende indhold af sepiolit blev fremstillet for at evaluere det optimale sammenblandingsforhold. På baggrund af disse resultater, blev en monolit indeholdende V\textsubscript{2}O\textsubscript{5}-SO\textsubscript{2}–4-ZrO\textsubscript{2} i 25 wt% sepiolit fremstillet og indflydelsen af hhv. gas hastighed, reaktions temperatur, NH\textsubscript{3}/NO forholdet i fødegassen, og NO reduktions effektivitet, blev undersøgt.

Sidste del af denne afhandling (kapitel 5) beskæftiger sig med den ikke-katalytiske sorption (absorption/desorption) af NO\textsubscript{x} i ioniske væsker. Da absorption af NO\textsubscript{x} i ioniske væsker ikke
tidligere har været beskrevet i litteraturen, blev der udført et indledende studie for at identifi-
cere egnede solventer. To imidazolium-baserede kandidater ([BMIM]OAc og [BMIM]OTf) blev
herved udvalgt på baggrund af deres imponerende høje sorptions kapaciteter. Begge solventer,
som blev undersøgt, udviste NO-opløseligheder omkring dobbelt så høje som dem rapportet
for f.eks. CO$_2$ indfangning i ioniske væsker. Specielt [BMIM]OAc udviste ekstraordinære ab-
sorptions kapaciteter, og var i stand til at fiksere fire NO molekyler per molekyle ionisk væske.
Omvendt, så udviste [BMIM]OTf lovende opførsel pga. dens reversible egenskaber. Dette vil i
princippet tillade genbrug af den ionisk væske samt opkoncentrering af NO. Den akkumulerede
NO kunne herved blive udnyttet til f.eks. fremstillingen af salpetersyre, hvorved et affaldspro-
dukt fra gassen bliver værdiskabende. Selvom en yderligere forståelse af mekanismerne i det
omtalte system er nødvendig, så er perspektivet for en teknologi der selektivt kan isolere NO
meget lovende som alternativ til den katalytiske fjernelse af NO fra industrielle røggasser.
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Chapter 1

Basic concepts

The work embedded in this thesis is the continuation of a long tradition of continuous optimizing and development of the methods for removal of nitrogen oxides. To ensure a common denominator, some basic terms are examined in the following sections.

1.1 \(\text{NO}_x\) sources

Nitrogen oxides, \(\text{NO}_x\), are among the most troublesome atmospheric pollutants and represents a major environmental problem, being partly responsible for the depletion of ozone (see reaction R-1.1), acid rain and town smog\(^{[1]}\).

\[
\text{NO} + \text{O}_3 \xrightarrow{hv} \text{NO}_2 + \text{O}_2 \quad \Delta H_{298K}^0 = -201 \text{ kJ/mol} \quad (\text{R-1.1})
\]

\(\text{NO}_x\) is not only produced in all combustion processes involving fossil- and bio-fuels, but also in lightning, microbiological denitrification processes in the soil, and by volcanic activity. However, the man-made \(\text{NO}_x\) from combustion processes contribute to almost three-quarters of the total amount of \(\text{NO}_x\) emitted into the atmosphere\(^{[2,3]}\). Fig. 1.1 depicts the \(\text{NO}_x\)-contribution from different sources in the US and Europe.

![Fig. 1.1: Emission of \(\text{NO}_x\) by source category in USA and Europe (based on data from 1993 and 2006). Adapted from\(^{[4]}\)](image)

The term ‘\(\text{NO}_x\)’ covers all the seven known oxides of nitrogen, \(\text{NO}\), \(\text{NO}_2\), \(\text{NO}_3\), \(\text{N}_2\text{O}\), \(\text{N}_2\text{O}_3\), \(\text{N}_2\text{O}_4\) and \(\text{N}_2\text{O}_5\). However, the resulting gas from a combustion is predominantly the oxides \(\text{NO}\) and \(\text{NO}_2\), with nitric oxide representing 90-95% of the total \(\text{NO}_x\)\(^{[3,5,6]}\).
Nowadays, catalysis is a fundamental part of modern chemistry.

A catalyst is defined as a substance that increases the rate of reaction without modifying the overall standard Gibbs energy change of the reaction, i.e. a catalyst does not affect the position of the chemical equilibrium\(^7\). The concept of catalysis was established by Ostwald, but was coined by Berzelius as early as 1836\(^8\). Catalysis is of major significance to our society as we rely on catalytic reactions in almost every part of the chemical industry - more than 90% of the processes in the chemical manufacturing utilize catalysts in some form\(^9\).

Catalysis can be subcategorized as homogeneous, in which reactants and catalyst are in the same phase, and heterogeneous catalysis, in which the reaction occurs at or near an interface between phases. However, in this study only heterogeneous catalysis is examined.
Chapter 2

Nitrogen oxides - a mechanistic approach

Before embarking upon the catalytic removal of the undesired NO\textsubscript{x}, it is worthwhile looking into the fundamentals of the reactions taking place. This chapter is subdivided into sections dealing with the reactions regarding the formation and removal of NO\textsubscript{x}.

The calculations of thermodynamic reaction parameters employed in this chapter, such as $\Delta H^o$ and $\Delta S^o$, are all based on the standard values of the CRC Handbook of Chemistry and Physics\textsuperscript{10}. An overview of the reactions and calculated thermodynamic entities are collated in the appendix tables: Tab. B.1 and B.2 (page 97).

### 2.1 Formation

Already in 1988 Bosch and Janssen\textsuperscript{3} categorized the formed NO\textsubscript{x} into three types, all formed during the combustion process. The first category, \textit{thermal NO\textsubscript{x}}, is responsible for the majority of the nitrogen oxides formed during combustion and is formed by oxidation of the atmospheric nitrogen at very high temperatures via the overall reaction:

$$N_2 + O_2 \rightarrow 2 \text{NO}, \quad \Delta H^o_{298K} = -183 \text{ kJ/mol}$$

\textbf{(R-2.1)}

\begin{figure}[h]
\centering
\includegraphics[width=0.5\textwidth]{fig2_1.png}
\caption{Theoretical equilibrium concentration of NO in air under the assumption that $\Delta H^o_{298K}$ and $\Delta S^o_{298K}$ are temperature-independent}
\end{figure}

Reaction R-2.1 is very unfavored thermodynamically at ambient temperatures, and only allows formation of a few hundreds ppm of NO at temperatures of 1300 K, but the NO evolution at higher temperatures is considerable, as illustrated in Fig. 2.1, since flame combustion temperatures in excess of 1700 K can occur\textsuperscript{11}. Reaction R-2.1 can be catalyzed by materials...
Chapter 2. Nitrogen oxides - a mechanistic approach

(e.g. particles from the fuel) in contact with the flame during the combustion, some endorsing the formation of NO, others inhibiting. This is highly relevant for fluidized bed solid combustors, as reported by Olofsson et al. [12]. Efforts have been made to minimize the amount of thermal NO\textsubscript{x} by combustion control, achieved by e.g. redesign of the combustor itself or low NO\textsubscript{x}-burners, where the combustions peak temperature are smoothed by partial premixing [13].

The second, and more complex, category of NO\textsubscript{x} is called prompt NO\textsubscript{x}, which involves the reaction of molecular nitrogen with organic hydrocarbon fragments to yield intermediate products, such as HCN, which subsequently is oxidized to NO. The reaction occurs at 1500 K or above and is responsible for the formation of NO in the range of tens of ppm [14], and due to the high temperature, the reverse reaction of reaction R-2.1, i.e. the decomposition of NO, does not take place. Furthermore, at lower temperatures NO can react with oxygen to form NO\textsubscript{2} (reaction R-2.2) or N\textsubscript{2}O (reaction R-2.3).

\[
2 \text{NO} + \text{O}_2 \rightarrow 2 \text{NO}_2, \quad \Delta H^0_{298K} = -116 \text{ kJ/mol} \quad (R-2.2)
\]

Reaction R-2.2 is shifted to the left at high temperatures due to its exothermic character, whereby only NO is formed during combustion. The equilibrium temperature for the reaction, where the Gibbs free energy is zero, can be calculated:

\[
\Delta G^0 = \Delta H^0 - T \Delta S^0 \Rightarrow \\
0 = \Delta H^0 - T \Delta S^0 \leftrightarrow \\
T_{\Delta G^0=0} = \frac{\Delta H^0}{\Delta S^0} = -116200 \frac{\text{kJ}}{\text{mol}} -146.6 \frac{\text{kJ}}{\text{mol} \cdot \text{K}} = 793 \text{ K} \quad (2.1)
\]

Consequently, NO\textsubscript{2} is practically completely decomposed to NO and O\textsubscript{2} at temperatures above 800 K. Although the enthalpy and entropy values utilized in the calculation only holds valid at 298 K and 1 bar, the result comply well with other reported findings for this reaction [1,11,14]. When air is present, the equilibrium of the NO oxidation (reaction R-2.2) is shifted to the right. However, the reaction is slow and as a result of this, most of the NO\textsubscript{x} is essentially composed of NO [14].

The prompt NO can also form N\textsubscript{2}O via reaction R-2.3.

\[
2 \text{NO} \rightarrow \text{N}_2\text{O} + \frac{1}{2} \text{O}_2, \quad \Delta H^0_{298K} = -101 \text{ kJ/mol} \quad (R-2.3)
\]

This reaction can be mediated by dimeric species of NO, such as N\textsubscript{2}O\textsubscript{2} [5], and is only favored at low temperatures [11]. Thus, during combustion at high temperatures the reverse reaction can take place, whereby N\textsubscript{2}O can be intermediate in the formation of NO. However, N\textsubscript{2}O can also be (but not necessarily) intermediate in the formation of NO at low temperatures via reaction R-2.4.

\[
2 \text{N}_2\text{O} \rightarrow 2 \text{NO} + \text{N}_2, \quad \Delta H^0_{298K} = 19 \text{ kJ/mol} \quad (R-2.4)
\]

The third category of NO\textsubscript{x} is called fuel NO\textsubscript{x} and is formed by oxidation of nitrogen in the fuel. Substantial amounts of fuel nitrogen is present in coal and heavy oils as well as in biomass, such as wood [14]. In contrast to the temperature controlled thermal NO\textsubscript{x}, fuel NO\textsubscript{x} is relatively
independent of temperatures at normal combustion temperatures\(^{[4]}\). On the other hand it is strongly influenced by reactor design\(^{[14]}\), but can be moderated by use of e.g. staged burning or reburning\(^{[15]}\).

### 2.1.1 Ammonia oxidation

Another source to NO\(_x\), which is highly relevant for the selective catalytic reduction reaction where NH\(_3\) or urea (decomposes to equimolar amount of ammonia and isocyanic acid, HNCO) is used as reductant in the presence of oxygen, is the reaction of ammonia with oxygen, and can take place through the following ways:

\[
\begin{align*}
4 \text{NH}_3 + 3 \text{O}_2 & \rightarrow 2 \text{N}_2 + 6 \text{H}_2\text{O}, & \Delta H_{298}^o = -1267 \text{ kJ/mol} & \text{(R-2.5)} \\
4 \text{NH}_3 + 4 \text{O}_2 & \rightarrow 2 \text{N}_2\text{O} + 6 \text{H}_2\text{O}, & \Delta H_{298}^o = -1104 \text{ kJ/mol} & \text{(R-2.6)} \\
4 \text{NH}_3 + 5 \text{O}_2 & \rightarrow 4 \text{NO} + 6 \text{H}_2\text{O}, & \Delta H_{298}^o = -902 \text{ kJ/mol} & \text{(R-2.7)} \\
4 \text{NH}_3 + 6 \text{O}_2 & \rightarrow 2 \text{N}_2\text{O}_3 + 6 \text{H}_2\text{O}, & \Delta H_{298}^o = -1094 \text{ kJ/mol} & \text{(R-2.8)} \\
4 \text{NH}_3 + 7 \text{O}_2 & \rightarrow 4 \text{NO}_2 + 3 \text{H}_2\text{O}, & \Delta H_{298}^o = -1134 \text{ kJ/mol} & \text{(R-2.9)}
\end{align*}
\]

These unwanted secondary reactions to the SCR reaction can occur as a result of the presence of oxygen, the temperature, the catalyst nature and the ratio of NO/NH\(_3\)\(^{[1,5,16,17]}\).

Reaction R-2.5 to R-2.9 are all displayed with four stoichiometric ammonia as reactants even though e.g. reaction R-2.6 strictly speaking only should contain half the amount of molecules. This is done for easy comparison of the resulting enthalpies. Furthermore, note that increasing the partial pressure of oxygen leads to a corresponding increase in the oxidation number of the resulting nitrogen species.

All the oxidations of ammonia shown above are fully accepted thermodynamically (\(\Delta G_{298}^o < 0\)), although the oxidation to N\(_2\) is most favored in enthalpy. Reaction R-2.5 is the so-called selective catalytic oxidation (SCO) of ammonia and is interesting because it in principle allows abatement of excess ammonia (ammonia slip) from the NH\(_3\)-SCR process. Several active SCR catalysts also perform well in SCO, although at slightly higher temperatures\(^{[5]}\). The oxidation of ammonia to NO in reaction R-2.7 is applied industrially in the synthesis of HNO\(_3\) using Pt as the catalyst, although several transition metals can catalyze the reaction\(^{[5,17,18]}\).

Due to the higher thermodynamic stability of N\(_2\) with respect to NO and N\(_2\)O, reaction R-2.5 is in general favored at lower temperature compared to reaction R-2.6 and R-2.7, whereas the opposite holds true at higher temperatures.

Even though nitrogen trioxide formed in reaction R-2.8 only exist in its pure form below the melting point at -101°C\(^{[10]}\), it disproportionates at higher temperatures according to reaction R-2.10.

\[
\text{N}_2\text{O}_3 \xrightarrow{\text{K}_{298}} \text{NO} + \text{NO}_2, \quad \Delta H_{298}^o = 38 \text{ kJ/mol} \quad \text{(R-2.10)}
\]

At ambient conditions the reaction is shifted to the right (\(\text{K}_{298} = 3 \text{ bar}\)), and equilibrium is quickly settled\(^{[19]}\).
Chapter 2. Nitrogen oxides - a mechanistic approach

Formation of NO\(_2\) in reaction R-2.9 does generally not compose a problem in correlation with the SCR reaction, other than the removal of ammonia (accounting for a non-stoichiometric conversion of NO and NH\(_3\) in the SCR reaction), since the formed nitrogen dioxide can participate in an ‘improved’ SCR process in concert with NO and ammonia, the so-called fast SCR. This aspect is further treated in the following section.

### 2.2 Removal

The most simple way to remove NO is in principle by its decomposition reaction, i.e. the reverse of reaction R-2.1.

\[
\text{NO} \rightarrow \frac{1}{2} \text{N}_2 + \frac{1}{2} \text{O}_2, \quad \Delta H^o_{298K} = -91 \text{ kJ/mol} \quad (R-2.11)
\]

The simple NO decomposition is thermodynamically favored at lower temperatures (cf. Fig. 2.1) but does not occur spontaneously due to slow kinetics. Many efforts have been made to catalyze the reaction, and several materials have been found (e.g. iron or copper oxides supported on ZSM-5), but a suitable catalyst which is active and resistant enough are yet to be found\(^{[1,5,20–23]}\). Consequently, to reduce NO to N\(_2\) selectively at sufficiently low temperatures, a catalyst must be applied. Although the specific type of catalyst may vary, the reduction of NO can efficiently be accomplished by a range of different reductants:

\[
\begin{align*}
\text{NO} + \text{H}_2 & \rightarrow \frac{1}{2} \text{N}_2 + \text{H}_2\text{O}, \quad \Delta H^o_{298K} = -333 \text{ kJ/mol} \quad (R-2.12) \\
\text{NO} + \text{CO} & \rightarrow \frac{1}{2} \text{N}_2 + \text{CO}_2, \quad \Delta H^o_{298K} = -374 \text{ kJ/mol} \quad (R-2.13) \\
\text{NO} + \text{CH}_4 + \frac{3}{2} \text{O}_2 & \rightarrow \frac{1}{2} \text{N}_2 + \text{CO}_2 + 2\text{H}_2\text{O}, \quad \Delta H^o_{298K} = -894 \text{ kJ/mol} \quad (R-2.14) \\
4\text{NO} + 4\text{NH}_3 + \text{O}_2 & \rightarrow 4\text{N}_2 + 6\text{H}_2\text{O}, \quad \Delta H^o_{298K} = -1632 \text{ kJ/mol} \quad (R-2.15)
\end{align*}
\]

Comparing the above reactions with the direct NO decomposition in reaction R-2.11, an overall decrease in enthalpy is generally observed when applying a reductant, i.e. an increase in the exothermic character of the reactions. A corresponding trend is also observed in the Gibbs free energy change of the reduction reactions, which is collated in Table 2.1. With the decrease in Gibbs free energy change using H\(_2\), CO, CH\(_4\), and NH\(_3\) as reductants, respectively, the corresponding equilibrium of the reactions is shifted further to the right, also compared with the Gibbs free energy change of the direct NO decomposition in reaction R-2.11 of \(\Delta G^o_{298K} = -88 \text{ kJ/mol}\).

Use of hydrogen as the reducing agent to remove NO (reaction R-2.12) is only applicable under anaerobic conditions, which is not the case with the SCR process. Reaction R-2.13 is used in reducing conditions. The reaction is exploited in the catalytic converters of cars, the so-called three-way catalyst\(^{[5]}\). The use of hydrocarbons, such as methane (reaction R-2.14), can constitute a suitable alternative to the traditional ammonia, which today is used in large stationary plants working in excess oxygen. However, the so-called HC-SCR is still not available for stationary plants on a commercial basis\(^{[14]}\).

\[
6\text{NO} + 4\text{NH}_3 \rightarrow 5\text{N}_2 + 6\text{H}_2\text{O}, \quad \Delta H^o_{298K} = -1815 \text{ kJ/mol} \quad (R-2.16)
\]
Table 2.1: Thermodynamic entities of the NO removal with different reductants

<table>
<thead>
<tr>
<th>Reaction no.</th>
<th>Reductant</th>
<th>$\Delta H_{298K}^o$ [kJ/mol]</th>
<th>$\Delta S_{298K}^o$ [J/mol K]</th>
<th>$\Delta G_{298K}^o$ [kJ/mol]</th>
<th>Eq. const., $K_{298K}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>(R-2.12)</td>
<td>H₂</td>
<td>-333</td>
<td>-127</td>
<td>-295</td>
<td>$5.3 \times 10^{51}$ bar^{-1/2}</td>
</tr>
<tr>
<td>(R-2.13)</td>
<td>CO</td>
<td>-374</td>
<td>-99</td>
<td>-345</td>
<td>$2.6 \times 10^{60}$ bar^{-1/2}</td>
</tr>
<tr>
<td>(R-2.14)</td>
<td>CH₄</td>
<td>-894</td>
<td>-158</td>
<td>-847</td>
<td>$2.2 \times 10^{148}$ bar⁰</td>
</tr>
<tr>
<td>(R-2.15)</td>
<td>NH₃</td>
<td>-1632</td>
<td>-340</td>
<td>-1531</td>
<td>$1.6 \times 10^{268}$ bar¹</td>
</tr>
</tbody>
</table>

Even though NO can be reduced with NH₃ based on the exothermic reaction R-2.16, the SCR process is usually performed with ammonia in the presence of oxygen, yielding reaction R-2.15, which has a substantial thermodynamic drive to the right (cf. Tab. 2.1). Furthermore, reaction R-2.16 is also performing much slower than the standard SCR reaction on vanadia-based catalysts (R-2.15), and therefore not relevant under lean (oxygen-containing) conditions [24].

If NO₂ is present in the gas, equimolar amounts of NO and NO₂ will react in a manner much faster than the standard SCR reaction (on a traditional vanadia-based catalyst) [25,26], the so-called fast SCR reaction:

$$2 \text{NO} + 2 \text{NO}_2 + 4 \text{NH}_3 \rightarrow 4 \text{N}_2 + 6 \text{H}_2\text{O}, \quad \Delta H_{298K}^o = -1516 \text{kJ/mol} \quad (R-2.17)$$

NO₂ can be produced by introducing e.g. a platinum oxidation catalyst upstream of the SCR catalyst. However, if the NO₂ concentration downstream of the oxidation catalyst surpasses 50%, which can be removed via reaction R-2.17 [24,26], excessive NO₂ can only react with ammonia.

$$6 \text{NO}_2 + 8 \text{NH}_3 \rightarrow 7 \text{N}_2 + 12 \text{H}_2\text{O}, \quad \Delta H_{298K}^o = -2734 \text{kJ/mol} \quad (R-2.18)$$

However, reaction R-2.18 is slower than the standard SCR reaction [26], but may promote formation of other side products, such as N₂O, all ultimately decreasing the N₂-selectivity of the SCR reaction. The formation of the byproduct N₂O depends on the reaction temperature. At intermediate temperatures (~200-350°C) NO₂ will react with ammonia in the absence or presence of oxygen in the following manner:

$$8 \text{NO}_2 + 6 \text{NH}_3 \rightarrow \text{N}_2\text{O} + 9 \text{H}_2\text{O}, \quad \Delta H_{298K}^o = -1595 \text{kJ/mol} \quad (R-2.19)$$

$$4 \text{NO}_2 + 4 \text{NH}_3 + \text{O}_2 \rightarrow 4 \text{N}_2\text{O} + 6 \text{H}_2\text{O}, \quad \Delta H_{298K}^o = -1074 \text{kJ/mol} \quad (R-2.20)$$

At high temperatures (>350°C) the thermodynamic equilibrium in reaction R-2.2 (Page 16) will settle at low space velocity, forming NO [26].
Chapter 3

Catalytic removal of NO\textsubscript{x}

3.1 Introduction

Nitrogen oxides belong to a class of troublesome pollutants and also have substantial damaging
effects on the environment. NO\textsubscript{x} is also involved in the formation of photochemical smog and
acid rain, which is perturbing the ecosystem.

Nitrogen oxides plays an important role in the photochemistry involving ozone. NO\textsubscript{x} con-
tributes to the depletion of stratospheric (located above the troposphere) ozone. The photo-
chemical destruction of ozone is catalyzed by NO via the reactions\textsuperscript{[4]}:

\[
\begin{align*}
\text{NO} + \text{O}_3 & \xrightarrow{h\nu} \text{NO}_2 + \text{O}_2 & \text{(R-3.1)} \\
\text{NO}_2 + \text{O} & \rightarrow \text{NO} + \text{O}_2 & \text{(R-3.2)}
\end{align*}
\]

Great efforts have been applied to reduce the emission of these pollutants through avail-
able technologies or by developing new methods\textsuperscript{[1,5]}. Different catalytic abatement technolo-
gies have in particular been applied to both mobile sources\textsuperscript{[4,27–29]} and stationary sources\textsuperscript{[30]}. However, the focus here will be on catalysis for stationary applications, which will be described
below along with a general introduction to catalysts for the selective catalytic reduction (SCR) in
section 3.1.1. Finally, the results obtained during this work are presented in section 3.2.

3.1.1 SCR catalysts

The direct decomposition of NO into N\textsubscript{2} and O\textsubscript{2} (reaction R-2.11) does not occur selectively at
reasonable temperatures without an appropriate catalyst, as discussed in Chapter 2. Although
the NO\textsubscript{x} can be removed without the use of catalyst, such as in the so-called selective non-
catalytic reduction (SNCR) which takes place at high temperatures between 850 to 1100°C\textsuperscript{[31]},
there are certain challenges with the SNCR process. The SNCR process can achieve the same
efficiency in the NO\textsubscript{x} removal as in SCR, but practical constraints (longer residence time, narrow
temperature window) limits the overall flexibility of the process\textsuperscript{[4,15]}. However, when utilizing a
catalyst, the temperature window can be decreased to around 200-650°C, while still maintaining
a high conversion of NO\textsubscript{x}\textsuperscript{[30,31]}. The SCR process was invented and patented in the US in 1959\textsuperscript{[32]}
and has since the 1970s been used commercially on a worldwide basis (primarily in Japan and
Europe, but the technology is developing in the USA and Asia\textsuperscript{[30]}) to meet the legally proscribed
NO\textsubscript{x} emission limits from stationary sources\textsuperscript{[5,6]}.

Ammonia has a key role in both the SCR and SNCR processes, due to the ability of ammonia
to react with NO\textsubscript{x} to selectively form N\textsubscript{2}, instead of being oxidized further by oxygen to N\textsubscript{2}O as
other reactants (CO or hydrocarbons) would do\[^6,30\].

The fact that roughly 80% of the stationary plants in Europe are removing NO\(_x\) by use of the SCR process illustrates the size and importance of the technology\[^14\]. However, there are still challenges that need to be addressed regarding the existing commercial catalyst, and further development is always possible in terms of improved efficiency, selectivity, and resistance to deactivation.

### 3.1.1.1 Commercial catalysts

In general, three types of catalyst systems have been developed for commercial SCR: noble metals, metal oxides, and zeolites\[^5,30\].

The noble metals, developed in the 1970s, were very active in the selective reduction of NO\(_x\). Unfortunately, they were also active in the oxidation of ammonia, and were soon succeeded by metal oxide catalysts\[^30\].

Of the many catalyst combinations investigated, those based on vanadia supported titania promoted with tungsta (WO\(_3\)) or molybdena (MoO\(_3\)) were found to possess genuine advantages in NO reduction and SO\(_2\) oxidation\[^30\]. Today, after three decades of studies and industrial application, the most widespread commercial SCR catalyst is based on V\(_2\)O\(_5\)/TiO\(_2\) formulations, such as V\(_2\)O\(_5\)−WO\(_3\)/TiO\(_2\) and V\(_2\)O\(_5\)−MoO\(_3\)/TiO\(_2\). The present catalysts are constituted of TiO\(_2\) anatase supports with a monolayer surface coverage of V\(_2\)O\(_5\) and WO\(_3\) (or MoO\(_3\)), deposited by impregnation. The surface area of the catalysts is around 50-100 m\(^2\)/g, with a V\(_2\)O\(_5\) content of 0.5-3 wt% and WO\(_3\) or MoO\(_3\) loading of 5-10 wt%. Vanadia, especially as bulk, is responsible for the undesired oxidation of SO\(_2\), and the vanadia content is thus generally kept low in the presence of high SO\(_2\) concentrations\[^14,30\]. Monolayer catalysts are more efficient than bulk V\(_2\)O\(_5\) and utilizing TiO\(_2\) anatase as support material relies on the fact that TiO\(_2\) is only weakly and reversibly sulfated in presence of SO\(_2\) and oxygen\[^33\]. Furthermore, TiO\(_2\)-supported V\(_2\)O\(_5\) possesses high acid site density and the vanadia is more readily distributed on the surface of TiO\(_2\) anatase than on SnO\(_2\) and ZrO\(_2\) (only monoclinic zirconia tested)\[^30,34,35\].

Commercial SCR catalysts are typically shaped in form of honeycomb monoliths, plates and coated metal monoliths with parallel channels. This gives several advantages over i.e. a packed bed: lower pressure drop, higher attrition resistance and lower tendency to plugging of fly ash\[^1\].

Since Iwamoto et al. and Held et al. independently discovered in the 1990s that selective reduction of NO\(_x\) takes place over a Cu-ZSM-5 catalyst\[^36\] (and references herein), the SCR by hydrocarbons have been studied intensely. An unique feature of zeolite catalysts for SCR is that NO first has to be oxidized to NO\(_2\), which is the rate-determining step of the reaction. The actual reaction taking place over zeolites is therefore the fast SCR reaction (reaction R-2.17). The NO\(_2\) is produced on the metal centers (i.e. iron) of metal-exchanged zeolites, but the SCR reaction takes place in the zeolite framework. Zeolites on the acid form with transition metal centers present rather high SCR activity at high temperatures. However, their durability under hydrothermal conditions is still a challenge. Deactivation at temperatures above 500°C is always observed, where sintering of metal centers and dealumination initiates with resulting loss of surface area.
3.1. Introduction

and Brønsted acidity (cf. Tab. 3.1). Although vanadia-based catalysts have been introduced to
diesel vehicles, zeolites do not have similar problems with SO₂ to SO₃ oxidation, low selectivity
at higher temperature and the volatilization of the toxic vanadium species above 650°C [23,30]
(and references therein). Even though zeolites exhibit high activity in the SCR process, they are
primarily being developed for the automotive industry due to the higher cost of zeolites than
conventional catalyst supports, such as TiO₂.

<table>
<thead>
<tr>
<th>Acid</th>
<th>Base</th>
</tr>
</thead>
<tbody>
<tr>
<td>Brønsted</td>
<td>Proton donor</td>
</tr>
<tr>
<td>Lewis</td>
<td>Proton acceptor</td>
</tr>
<tr>
<td></td>
<td>Electron acceptor</td>
</tr>
<tr>
<td></td>
<td>Electron donor</td>
</tr>
</tbody>
</table>

### Tab. 3.1: General Brønsted/Lewis acid/base properties

#### 3.1.1.2 Vanadium-based catalysts

The properties of the TiO₂-supported V₂O₅ catalyst promoted with WO₃ (or MoO₃) have been
extensively studied by numerous laboratories since its discovery as an active SCR catalyst sev-
eral decades ago. The catalytic properties of the supported vanadium catalysts are evidently
strongly affected by the vanadia loading, preparation method, and type of support and pro-
moter. The type of metal oxide species present on the support surface is determined by the
content of vanadium. At low loadings the isolated and monomeric VOₓ species predominate,
which, with increasing loading, will react to form dimeric and polymeric species until mono-
layer coverage is reached, illustrated in Fig. 3.1. The crystalline phase of V₂O₅ will dominate on
the surface above monolayer coverage [37].

![Fig. 3.1: Schematic figure of dispersed surface vanadates in their dehydrated form. Vanadia loading increasing from left to right](image)

Although there have been significant contributions to the understanding of the SCR process
over vanadium-based catalysts, an exhaustive elucidation of the mechanism has not yet been
achieved.

It is widely accepted that both the titania support and V₂O₅ contain Lewis acidity, and that
Lewis sites on vanadia under adsorption of H₂O can be converted into Brønsted acid sites [38,39],
as depicted in Fig. 3.2. The ammonia species adsorbed on vanadia/titania-based catalysts there-
fore show two forms: coordinated NH₃ on the Lewis acid site and ammonium (NH₄⁺) on the
Brønsted acid site (Fig. 3.3). In accordance, two types of mechanisms are suggested, Lewis-
NH₃-based and Brønsted-NH₄⁺-based mechanisms.

Even though one of the first reaction schemes proposed for the SCR reaction was based on
a Langmuir-Hinshelwood mechanism [40], there is a today general consensus that the SCR re-
Chapter 3. Catalytic removal of NO\textsubscript{x}x

![Image of catalytic removal of NO\textsubscript{x}x](image)

**Fig. 3.2:** Generation of Brønsted acid sites from Lewis acid site (surface vanadyl center). Adapted from Busca et al.\cite{Busca5}. However, the vanadium specie formed is here over coordinated, which should have resulted in cleavage of the weakest bond: the V-O-support bond.

**Fig. 3.3:** Proposed structures for ammonia adsorbed on titania supported V\textsubscript{2}O\textsubscript{5}. Adapted from Busca et al.\cite{Busca5}

action occurs via an Eley-Rideal type mechanism, in which ammonia is first adsorbed on the vanadium-based catalyst, followed by activation with gas-phase NO\cite{Inomata41,Inomata42,Inomata43,Inomata44,Inomata45,Inomata46,Inomata47,Inomata48,Inomata49,Inomata50}. Different proposals have been suggested for the adsorption mode of ammonia, which is still not fully elucidated. Inomata et al.\cite{Inomata41,Inomata42} suggested that ammonia is first adsorbed at a Brønsted V\textsuperscript{5+}—OH site adjacent to a V\textsuperscript{5+}—O site. With isotopic labeling studies, Janssen et al.\cite{Janssen43,Janssen44} suggested a dual site mechanism with V\textsuperscript{5+}—O as the easy reducible active species. According to the mechanism, ammonia is adsorbed on a V=O site as V—ONH\textsubscript{2} by reduction of the neighboring V=O site to V—OH. Ozkan et al.\cite{Ozkan49,Ozkan50} applied isotopic labeling of \textsuperscript{18}O\textsubscript{2}, \textsuperscript{15}NH\textsubscript{3}, \textsuperscript{15}NO, and \textsuperscript{15}N\textsuperscript{18}O to suggest that ammonia adsorbs on Brønsted pairs of V—OH, forming ammonium ion species (cf. Fig. 3.3, right). Topsøe et al.\cite{Topsøe45,Topsøe46,Topsøe47} and Gasior et al.\cite{Gasior48} proposed that predominantly Brønsted acid sites (V—OH) were responsible for the ammonia adsorption and subsequent formation of an activated complex in concert with an adjacent vanadyl group. The mechanism they suggested (reproduced in Fig. 3.4) is a perfected version of that proposed by Inomata et al.\cite{Inomata42}. Density functional theory (DFT) performed on V\textsubscript{2}O\textsubscript{5} show that ammonia interacts with Bronsted acid sites, and together with NO reacts to form the intermediate NH\textsubscript{3}NHO, which undergoes a series of hydrogen-transfer steps on the vanadia cluster to form adsorbed NH\textsubscript{2}NO species that decomposes to N\textsubscript{2} and H\textsubscript{2}O and a reduced vanadium oxide cluster\cite{Inomata53,Inomata54}. In contrast, Ramis et al. suggested that the presence of Bronsted acid sites is not a prerequisite for the SCR reaction to occur\cite{Ramis38,Ramis51,Ramis52}. Based on FTIR studies, the authors propose that ammonia is adsorbed on Lewis acid sites and is activated to from adsorbed amide NH\textsubscript{2} species, which then can react with gaseous NO to give an intermediate nitrosamide (NH\textsubscript{2}NO) that is known to decompose into N\textsubscript{2} and H\textsubscript{2}O. The authors investigated the spectra region where ammonia is coordinated (1300–1100 cm\textsuperscript{-1}), a region which Topsøe et al. did not include in their publications, even though Lewis coordinated NH\textsubscript{3} show
3.1. Introduction

Fig. 3.4: Catalytic cycle of the SCR reaction on vanadia/titania in the presence of O$_2$, proposed by Topsøe et al.\textsuperscript{[45,46]}

![Catalytic cycle of the SCR reaction on vanadia/titania in the presence of O$_2$.](image)

a significant band in this spectral range. Recent DFT calculations performed by Soyer et al.\textsuperscript{[53]} on a V$_2$O$_9$H$_8$ cluster have, however, showed that Lewis acidic adsorption of ammonia is highly unfavorable.

Even though the \textit{Brønsted-NH$_4^+$} mechanism has gained the major support in the literature, it should be noted that it is still a hypothesis and lacks solid experimental evidence under \textit{in situ} conditions. It can thus not be ruled out that the SCR reaction on vanadia/titania (dependent on the reaction conditions) proceeds through several mechanisms involving both types of acid sites.

3.1.1.3 Role of promoter and support material

As already mentioned, the commercial SCR catalyst is based on vanadia/titania formulations, such as V$_2$O$_5$–WO$_3$/TiO$_2$ and V$_2$O$_5$–MoO$_3$/TiO$_2$. According to Chen and Yang\textsuperscript{[55]} (and references herein), the advantages of adding WO$_3$ to the catalyst as promoter are numerous: stabilization of the metastable anatase phase of TiO$_2$; broadening the operating temperature window; increase of the surface acidity; decrease of the ammonia oxidation activity; decrease of the activity for SO$_2$ oxidation; and improved resistance towards alkali poisoning. DFT has been performed on the V$_2$O$_5$–WO$_3$ system, where it was suggested that addition of tungsten oxide lead to enhanced probability of the dissociation of water molecules, thereby forming more Brønsted acid sites\textsuperscript{[56]}. Lietti et al.\textsuperscript{[57]} performed a comparative study of V$_2$O$_5$–WO$_3$/TiO$_2$ and V$_2$O$_5$–MoO$_3$/TiO$_2$ catalysts in the SCR reaction, and found that although MoO$_3$-based catalysts are more active they are also less N$_2$-selective than the corresponding WO$_3$-based catalysts. However, both ‘additives’ acts as structural promoters, by preventing the sintering of the TiO$_2$ matrix.

Sulfate has also been applied as promoter for the SCR catalyst, and is suggested to have a promoting effect on the SCR activity due to significant increase in Bronsted acidity (if water is present. The surface sulfate species on the catalyst can originate from impregnation or treatment with i.e. H$_2$SO$_4$ or (NH$_4$)$_2$SO$_4$ solutions, or from the gaseous SO$_2$ if present in flue gas\textsuperscript{[5]}. The surface acidity of the sulfate-promoted catalysts is even higher than the correspond-
ing WO₃-promoted catalysts. Sulfate (like WO₃) was shown to have an inhibiting effect on the phase transformation from TiO₂ anatase to rutile, and to increase the number and strength of surface acid sites. However, TiO₂ is only weakly and reversibly sulfated at normal SCR conditions due to the low sulfate stability on the titania surface. If WO₃ or MoO₃ is present on the surface, they will compete and displace SO₃ on the basic sites of titania, consequently limiting the sulfate absorbed from gaseous oxidized SO₂. On the other hand, sulfation of ZrO₂ or Al₂O₃ is more favorable with regard to sulfate stability. Titania-based catalysts are therefore only partially surface-sulfated during SCR operation in the presence of SO₂.

Hamada et al. were the first to report of SCR active sulfated metal oxides (TiO₂, Al₂O₃, ZrO₂, Fe₂O₃). The most promising catalysts of this type have been shown to consist of transitions metals supported on ZrO₂ or SO₄²⁻/ZrO₂ for the SCR of NO with NH₃ and with hydrocarbons. Sulfated zirconia was first reported by Hino and Arata to possess unusually high acidic properties, and was suggested to be a so-called solid superacid. A superacid is a medium having an acidity higher than 100 wt% H₂SO₄, such as HF or HSO₃F. Superacidity can also be expressed with the Hammet scale, H₀ ≤ -11.93, although some authors disagree on whether sulfated zirconia is in fact a solid superacid (and references herein). However, a general issue regarding the characterization of sulfate-doped samples is that the acidic and catalytic properties depend on a large number of preparative variables, such as; calcination temperature; sulfate content; concentration of sulfate solution; sulfate precursor; crystal phase; surface area and sample activation. The most abundant use of this type of solid acids has been for alkane isomerization reactions, but a review on the wide range of reactions that sulfated zirconia can catalyze have been collected by Yamaguchi et al. and Jin et al.

![Fig. 3.5: Proposed Lewis acidic site structure of sulfated zirconia. Reproduced from Yamaguchi](image)

![Fig. 3.6: Proposed structure where both Brønsted (a) and Lewis acid sites (b) are present. Reproduced from Arata and Hino](image)

Although there is a generally consensus that the presence of sulfate species with covalent S=O bonds are necessary to achieve superacidity (and references herein), the exact nature of the acidic sites remains unknown. Dunn et al. report that the sulfated species present on TiO₂, ZrO₂, and Al₂O₃, with similar vanadia loading, all have identical molecular structures, i.e. (M–O)₃=S. Morterra et al. reported that the monoclinic phase of ZrO₂ is not active in the isomerization of butane. Thus, the sulfate acid site formation is weaker on phases of zirconia other than the tetragonal. This has been confirmed recently by DFT calculations, where it was found that the affinity for SO₃ absorption on t-ZrO₂ was significantly larger than on the cor-
responding cubic and monoclinic crystal configurations, due to stabilization by abstraction of oxygen atom from the tetragonal phase\textsuperscript{[78]} (knowing this can be an useful indicator of the crystal phase of zirconia in acid-catalyzed studies where the ZrO\textsubscript{2}-phase is not reported, e.g. in\textsuperscript{[79]}). Jin et al.\textsuperscript{[75]} suggested that the acid centers are Lewis sites, and associated with the metal cation. The strength of this Lewis site is enhanced by the strong ability of the sulfur complex to absorb the electrons as outlined in Fig. 3.5. Others suggest the simultaneous presence of Bronsted and Lewis acid sites at hydrated conditions\textsuperscript{[76,80]}, depicted in Fig. 3.6. However, the Bronsted acid sites are readily interconverted to Lewis sited by evacuation at temperatures above 150°C\textsuperscript{[71]}.

Even though sulfated zirconia show stronger acidic sites than tungstated zirconia\textsuperscript{[72,79]}, sulfate species have been shown desorb from the surface during thermal treatment, above ca. 700°C\textsuperscript{[71,72]}. Therefore, in terms of thermally stability, WO\textsubscript{3}/ZrO\textsubscript{2} can be suggested as an alternative solid acid catalyst. Sohn and Park\textsuperscript{[81]} found that a tungstate concentrations of 20 wt% on zirconia, resulted in maximum surface area and acidity. Surface coverage, \(n_s\), can be calculated with expression 3.1, where \(N_A\) is Avogadro’s number, \(S_{BET}\) the surface area, and \(c_W\) the metal loading [g/g].

\[
n_s = \frac{c_W N_A}{M_W S_{BET} \cdot 10^{18} \text{nm}^2/\text{m}^2} = \frac{0.159 \cdot \frac{\text{dWO}_3}{\text{g}}}{183.84 \cdot \frac{\text{g}}{\text{mol}}} \cdot \frac{6.022 \cdot 10^{23} \text{mol}^{-1}}{155 \text{m}^2/\text{g} \cdot 10^{18} \text{nm}^2/\text{m}^2} = 3.35 \text{WO}_3/\text{nm}^2 \tag{3.1}
\]

Assuming surface area of 155 m\textsuperscript{2}/g, the surface coverage of 20 wt% WO\textsubscript{3}/ZrO\textsubscript{2} can be calculated to 3.4 W/\text{nm}\textsuperscript{2}, which according to Wachs measurements of monolayer coverage outlined in Tab. 3.2, corresponds to submonolayer coverage. However, Zhou et al.\textsuperscript{[82]} have reported polytungstate cluster formation at loadings below monolayer coverage, already at 2.9 W/\text{nm}\textsuperscript{2}. Thus, the reported two-dimensional polytungstated species which has been suggested to be important for generation of strong acid sites\textsuperscript{[83]}, could possibly be formed below monolayer coverage.

### Tab. 3.2: Monolayer surface coverages of supported metal oxides ([atoms/\text{nm}\textsuperscript{2}]) obtained by Raman and IR studies. Adapted from Wachs\textsuperscript{[84]} (and references herein)

<table>
<thead>
<tr>
<th></th>
<th>Al\textsubscript{2}O\textsubscript{3}</th>
<th>TiO\textsubscript{2}</th>
<th>ZrO\textsubscript{2}</th>
<th>SiO\textsubscript{2}</th>
</tr>
</thead>
<tbody>
<tr>
<td>V</td>
<td>7.3</td>
<td>7.9</td>
<td>6.8</td>
<td>0.7</td>
</tr>
<tr>
<td>W</td>
<td>4.0</td>
<td>4.2</td>
<td>4.0</td>
<td>0.1</td>
</tr>
<tr>
<td>Mo</td>
<td>4.6</td>
<td>4.6</td>
<td>4.3</td>
<td>0.3</td>
</tr>
</tbody>
</table>

Naturally, the dispersion and surface species strongly depend on the nature of the support, as can be envisaged in Tab. 3.2. For example, distributing vanadia on the surface of SiO\textsubscript{2} results in formation of polycrystalline V\textsubscript{2}O\textsubscript{5} already at very low loadings, whereas on Al\textsubscript{2}O\textsubscript{3}, TiO\textsubscript{2}, and ZrO\textsubscript{2} much more vanadia (almost an order of magnitude) can be added to the support before crystalline V\textsubscript{2}O\textsubscript{5} is formed. The low surface coverage of metal oxides on silica are due to the lower reactivity and density of surface hydroxyls compared to the other metal oxides\textsuperscript{[84]}. A comparative study of the surface acidity of various promoted metal oxides was performed by Niwa et al.\textsuperscript{[79]}. The authors found that for the metal oxides studied (SnO\textsubscript{2}, ZrO\textsubscript{2}, TiO\textsubscript{2}, and Al\textsubscript{2}O\textsubscript{3}) strong acidity is obtained following the sequence, SO\textsubscript{4}\textsuperscript{2–} > WO\textsubscript{3} > MoO\textsubscript{3} > V\textsubscript{2}O\textsubscript{5}. Dunn et al.\textsuperscript{[33]}
performed a comparative study of the SO$_2$ oxidation activity ability of 1 wt% loaded V$_2$O$_5$ cata-
ysts, and found the following trend: V$_2$O$_5$/CeO$_2$ > V$_2$O$_5$/ZrO$_2$ > V$_2$O$_5$/TiO$_2$ > V$_2$O$_5$/Al$_2$O$_3$ > V$_2$O$_5$/SiO$_2$.

Finally, even though tungsta-promoted titania has traditionally been used as a commercial
SCR catalyst, other catalyst combinations options exist that can be tuned depending on the op-
erational requirements, e.g. temperature of operation, SO$_2$ or deactivating species in flue gas.

### 3.1.2 Catalyst deactivation

The mechanism of catalyst deactivation can be subdivided into three classes: poisoning, coking and sintering. Other types of deactivation include masking, volatilization of active elements, erosion and attrition\cite{85}. Throughout the thesis, the primary area of focus will be on the chemi-
cal deactivation by poisoning.

#### 3.1.2.1 Inorganics in coal and biomass

Naturally, the SCR catalyst is exposed to many different gaseous species, depending on the type
of fuel combusted in the power plant. Natural gas comprise a relatively clean combustion gas, 
whereas combustion of e.g. municipal waste or straw often contain species that can lead to pos-
sible deactivating of traditional vanadia-titania-based SCR catalysts. The composition of various
species of both fuel and ash associated with commercial or pilot plants currently combusting or 
co-combusting biomass have recently been reported\cite{86–88}. All studies show a significant in-
crease in the quantity of alkali and alkaline-earth elements (K, Na, Mg, Ca), phosphorous and 
chlorine in the biomass-based fuels compared to coal. The quantity of potassium and chlorine

| Tab. 3.3: Fuel composition. Moisture, volatiles and ash-concentration are given on a wet basis, all other elements on dry basis. Adapted from Zheng et al.\cite{87} |
|-----------------|-----------------|-----------------|-----------------|-----------------|
| Moisture        | 2.00            | 2.00            | 2.00            | 6.70            |
| Volatiles       | 39.53           | 24.07           | 51.85           | 68.76           |
| Ash             | 5.00            | 14.49           | 5.28            | 7.37            |
| C               | 76.14           | 71.51           | 62.70           | 44.80           |
| O               | 11.59           | 6.55            | -               | -               |
| N               | 1.41            | 1.75            | 0.71            | 1.10            |
| S               | 0.81            | 0.63            | 0.40            | 0.20            |
| Cl              | 0.01            | 0.01            | 0.05            | 0.73            |
| P               | 0.004           | 0.11            | 0.02            | 0.14            |
| Ca              | 0.06            | 0.93            | 1.17            | 0.42            |
| Mg              | 0.03            | 0.16            | 0.19            | 0.10            |
| Na              | 0.01            | 0.02            | 0.01            | 0.02            |
| K               | 0.07            | 0.08            | 0.01            | 1.90            |

\[1\] Columbia high volatile bituminous coal  
\[2\] South Africa medium volatile bituminous coal

in the straw can be more than an order of magnitude higher than that of coal (cf. Tab. 3.3). Espe-
ially the high content of potassium in e.g. wood and straw is of concern, which will be further
3.1. Introduction

discussed in section 3.1.2.2. On the other hand, the sulfur content in biomass is typically lower in biomass than in coal. Zheng et al.\cite{87} showed a clear relationship between sulfur in fuel and SO\textsubscript{2} emission. Thus, by firing or co-firing with biomass a reduction of SO\textsubscript{2} in the flue gas is expected.

There are several practical concerns when these constituents enter the boiler in higher concentrations: high-temperature corrosion due to chlorine; accelerated slagging; and fouling\cite{88}. A major problem regarding the application of SCR catalysts has been their deactivation by different components in the flue gas. This issue is addressed in the following.

3.1.2.2 Effect of alkali on vanadia-based catalysts

Alkali metals were identified early as poisons for vanadia-titania-based SCR catalysts by Chen and Yang\cite{89}, where they found that the strength of the poison is related to the increase of basicity. Since then, the effect of alkali species on the vanadia-based catalyst have been reported by several groups\cite{55,72,85,86,88,89–98}. Although the alkali-vanadium system has been intensively studied, the exact role of the dopants on the supported vanadium oxide at the molecular level are still a subject of discussion. Most authors agree that potassium is severely deactivating the traditional V\textsubscript{2}O\textsubscript{5}−WO\textsubscript{3}−TiO\textsubscript{2} catalyst, although tungsta is partially inhibiting the deactivating effect\cite{55,85,86,90–92,94,95,98,99}. Numerous TPD and spectroscopic studies show that the deactivating nature of potassium is clearly connected with a reduction of the V\textsubscript{2}O\textsubscript{3} Brønsted acid sites of the catalyst\cite{90–92,98,99}. However, only a limited number of studies have been performed on WO\textsubscript{3}-containing catalysts\cite{85,91,92,94,95,98}. Loss of Brønsted acid sites, as discussed in section 3.1.1.2, would reduce catalyst activity severely, since they are vital for the SCR reaction to occur.

The deactivating effect of potassium on an industrial reference catalyst with the formulation 3 wt% V\textsubscript{2}O\textsubscript{5}−WO\textsubscript{3}−TiO\textsubscript{2} is illustrated in Fig. 3.7, where an increasing degree of potassium is deposited on the catalyst. Although a high SCR activity (expressed by the first order rate constant) is obtained for the fresh catalyst, the impact of potassium is significant, almost completely deactivating the catalyst at K/V ratio of 0.3 mol/mol. Experimental details regarding catalyst extraction from the commercial monolith is described in section A.2.1 (page 89).

![Fig. 3.7: SCR performance of commercial 1.7VWT_HT catalyst with increasing potassium loading.](image)

Theoretical studies regarding the adsorption of potassium on the pure surface of TiO\textsubscript{2} ru-
tile\(^{100,101}\) were initially performed, but recent theoretical DFT studies have addressed the alkali-vanadia system\(^{72,99,102–104}\). Zheng et al.\(^{105}\) suggested an early mechanism, based on a substitution reaction on the Brønsted site, whereby the reactive site is blocked by potassium:

\[
-V - OH \xrightarrow{KCl} - V - O - K
\]  

(R-3.3)

This mechanism requires an equimolar ratio of vanadium and potassium to completely deactivate the catalyst. Although the model provides a very intuitive understanding of the deactivation, several groups report of almost complete loss of catalyst activity already at K/V \(\approx 0.5\) (mol/mol)\(^{85,89,93,94}\) (see also Fig. 3.7), which suggests that potassium blocks multiple active sites.

Recent DFT studies suggest mechanisms in better agreement with experimental findings. Witko et al.\(^{102}\) and Nicosia et al.\(^{99}\) both studied the addition of one alkali atom to clusters of V\(_{10}\)O\(_31\)H\(_{12}\) and V\(_8\)O\(_{20}\)H\(_{10}\), respectively, which in practice would resemble crystalline V\(_2\)O\(_5\). Based on their findings, they suggested that the deactivating agent preferentially occupy hole sites between four V\(_2\)O\(_5\) centers, thereby inhibiting both Brønsted and V\(^{5+}\)=O sites. Calatayud and Minot\(^{103}\) used smaller V\(_2\)O\(_5\) clusters, but added the substrate titania anatase to the model. With a K/V concentration of 0.5, they suggested that potassium is interacting with the oxygens of two neighboring vanadia units (bridging V-O-Ti and terminal V=O) and the support, as shown in Fig. 3.8.

![Fig. 3.8: Suggested structure for coordination of potassium to several vanadia and titania sites. Yellow: K, blue: V\(_2\)O\(_5\), Red: oxygen, Gray: Ti\(^{103}\)](image)

All the proposed models of the alkali doping of vanadia can explain the observed elongation of the V=O bond, readily observed in Raman spectroscopy, and a reduction in the reducibility of the K-doped catalyst compared to the fresh catalyst\(^{72,91,104,106,107}\). Lewandowska et al.\(^{104}\), Kiwi-Minsker et al.\(^{106}\), and Calatayud et al.\(^{103}\) report that alkali doping would counteract the polymerization of vanadia-units by stabilization of the vanadium cluster. Furthermore, Kiwi-Minsker et al. reported of the disappearance of monomeric vanadium species upon potassium
doping, which fit well with the findings suggested in Fig. 3.9. This suggests that potassium is also involved in the reaction and cleavage of bridging V-O-V bonds.

The apparent disagreements between the reactivity of different vanadium-based clusters reported in the theoretical literature was recently addressed by Calatayud et al.\textsuperscript{[108]}. Based on periodic calculation of V\textsubscript{2}O\textsubscript{5} compounds as gas-phase clusters, single crystals and supported on titania, they report significant differences in reactivity dependent on the choice of model cluster. This issue can be extrapolated to the conflicting reports in the literature of how alkali atoms could bind to the active sites. The nature of potassium coordination to the active sites of the vanadium catalyst would strongly depend on the vanadium oxide cluster model for the DFT calculation. Hence, a study of the alkali coordination on molecule models with various vanadia loadings, promoted with e.g. sulfate, tungsta, supported on substrate is strongly required.

### 3.1.2.3 Deactivation overview

The content of possible poisons in the fuel for e.g. municipal solid waste (MSW) or biomass incineration can vary significantly. Many parameters besides the fuel composition influence the activity of the SCR catalyst e.g. particle size of poison in flue gas; placement of catalyst in high- or low-dust positions; temperature of operation; scrubbing etc. Hence, it is feasible to evaluate the potential deactivation of the catalyst in a comparable way for the different fuel types, which is attempted in table 3.4, where the impact on the traditional SCR catalyst for different fuel types, based on the fuel composition is collected.

| Table 3.4: Estimation of the degree of deactivation ($\chi$) of the traditional vanadia-based SCR catalyst based on fuel composition (dry basis). $\chi=5$: Severe deactivation, $\chi=1$: Negligible effect |
|---|---|---|---|---|
| **MSW** | **Wood** | **Forest Residues** | **Straw** | **Coal** |
| Na | 1–2 | 0.2 | 1 | 0.02–0.05 | 1–2 | 0.02–0.05 | 1–2 | 0.02–0.05 | 1–2 | 0.02–0.05 |
| S | 1–2 | 0.2 | 1 | 0.02–0.05 | 1–2 | 0.02–0.05 | 1–2 | 0.02–0.05 | 1–2 | 0.02–0.05 |
| Cl | 2–3 | 0.62–0.9 | 1 | 0.02–0.05 | 1–2 | 0.02–0.05 | 1–2 | 0.02–0.05 | 1–2 | 0.02–0.05 |
| K | 1–2 | 0.2 | 1 | 0.02–0.05 | 1–2 | 0.02–0.05 | 1–2 | 0.02–0.05 | 1–2 | 0.02–0.05 |
| Zn | 2 | 0.09–0.16 | 1 | 0.02–0.05 | 1–2 | 1.5 ppm$^\dagger$ | 1–2 | <0.01 | 1–2 | <0.01 |
| Pb | 2 | 0.01–0.04 | 1 | 0.02–0.05 | 1–2 | 0.01–0.04 | 1–2 | <0.01 | 1–2 | <0.01 |

$^\dagger$ Measured in flue gas, particle diameter <1μm

Considering the incineration of biomass, it can be concluded from table 3.4 that alkali, especially potassium, has to be considered as potential poison for the traditional SCR catalyst when significant amounts of straw is burned.

### 3.1.2.4 Alkali-resistant SCR catalyst

Although tungsta is added to the V\textsubscript{2}O\textsubscript{5}–TiO\textsubscript{2} formulation to decrease the poisoning effect of alkali, the traditional catalyst is still far from being resistant to deactivating agents typically found in the flue gas of biomass, confirmed by several full-scale experiments with the V\textsubscript{2}O\textsubscript{5}–WO\textsubscript{3}–TiO\textsubscript{2} catalyst\textsuperscript{[83,88,95,110]}. As concluded in previous sections, the basic potassium is blocking the Brønsted V-O-V acidic sites of the working SCR catalyst, as schematically illustrated in Fig. 3.10.

Several approaches can be applied to design a SCR catalyst more resistant to alkali. One
procedure is to increase the acidity of the substrate, which would have a stronger interaction with potassium than the active vanadia sites, which is illustrated in Fig. 3.11. As suggested in section 3.1.1.3, sulfated zirconia possesses very strong acidity, and is thus a suitable candidate, alongside tungstated zirconia. Another approach is based on substituting the active species of the catalyst. A crucial part of the mechanism is the Brønsted acid sites of vanadia, which is much more affected than the Lewis acid sites by potassium doping. Hence, by using metal oxides which mainly possess Lewis acidity, the catalyst could avoid deactivation by alkali. CuO and Fe$_2$O$_3$ are attractive under these conditions, since they are well known Lewis acids\[51\] and have also exhibited NH$_3$-SCR activity\[1,5\]. This approach method has already been applied by Kustov et al.\[93,114\]. However, only sulfated monoclinic ZrO$_2$ was investigated in the study, whereas the sulfated tetragonal zirconia contain much higher acidity (discussed in section 3.1.1.3). Tungstated zirconia has also previously been investigated in our group as a support material for vanadia loaded SCR catalysts with promising results\[115\]. Finally, as a more exotic non-catalytic method of NO removal, is by means of gas capture in ionic liquids as illustrated in Fig. 3.12. The technology has already been used for CO$_2$ (although some challenges are still to be overcome\[116\]) and SO$_2$ absorption, but has never been applied to the removal of NO. All three approaches will be explored in the following thesis.
3.2. Zirconia-based catalyst systems

This section is dedicated to the study of ternary SCR catalysts, consisting of metal oxide as active phase, sulfate as promoter and zirconia as support material. The sulfated zirconia is loaded with the oxides of vanadium, iron and copper.

3.2.1 Vanadium-based catalysts

Early in the study of zirconia-based catalysts, we found that a line of commercial zirconia-based samples from Saint-Gobain Norpro (NP) proved suitable as support material, competitive with the SCR activities of vanadia-loaded synthesized zirconia which was previously published[72]. Since the zirconia already was commercially available on extruded form (pellets), the exploitation of this material as possible SCR catalyst is a step closer to industrial relevance. Hence, the commercial zirconia is forming the frame for all the ternary catalysts synthesized in the following sections.

The initial selection of suitable support materials, was primarily based on the surface area and crystalline form. As earlier discussed (section 3.1.1.3), the monoclinic phase of ZrO$_2$ provides weak acid sites when sulfated, whereas the tetragonal form is responsible for the generation of strong acid sites[70,117]. Zirconia has three stable polymorphs: tetragonal (t–ZrO$_2$), monoclinic (m–ZrO$_2$), and cubic (c–ZrO$_2$), and the occurrence and transformation between the phases depend on doping and/or thermal treatment. The monoclinic phase is stable up to 1050°C, where it transforms into the tetragonal phase, which further changes into the cubic phase at 2400°C. The cubic phase of zirconia is stable up to the melting point at about 2700°C. Thus, tetragonal and cubic ZrO$_2$ are only metastable phases below 1050°C, where monoclinic is the thermodynamically preferred form. In the case of i.e. sulfate or tungstate promoted zirconia catalysts, the crystalline structures of the samples behave differently from that of pure ZrO$_2$. Amorphous zirconia has been reported to undergoes crystallization around 350-450°C[81,118], but when doping the matrix with e.g. a promoter, the phase transition temperature from amorphous to tetragonal zirconia is shifted to higher temperatures[72,76,81,115]. Zirconia is thus a grateful material to work with, because it can easily be stabilized in its tetragonal phase by e.g. a promoter, which hinders the phase transition to monoclinic.

3.2.1.1 ZrO$_2$ crystal phases

With XRD it is easy to identify the monoclinic phase of ZrO$_2$, whereas tetragonal and cubic zirconia can be difficult to distinguish between. However, by refining the XRD data with the Rietveld method (further description in Appendix A), it is possible to discriminate the abundance of the c–ZrO$_2$ from t–ZrO$_2$. In Fig. 3.13 is the diffraction patterns of the commercial supports depicted together with the difference curve from the Rietveld fit. If the Rietveld analysis results in a perfect fit, accounting the presence of all phases present, the difference curve would yield a straight line, located on the baseline.
Chapter 3. Catalytic removal of NO\textsubscript{x}

From the diffraction patterns in Fig 3.13 it is clear that the pure ZrO\textsubscript{2} samples primarily consist of monoclinic ZrO\textsubscript{2}, whereas the samples containing sulfate mainly contain tetragonal and cubic ZrO\textsubscript{2}. The numbers in the name of the pure zirconia samples correspond to their surface area (m\textsuperscript{2}/g), and in the sulfated samples they denote the loading of sulfate measured as SO\textsubscript{3} (wt%) (from here on, sulfate is abbreviated as S, and zirconia as Z). Through Rietveld refinement the phase distributions can be obtained, which is collected in Tab. 3.5. Although synthesis details for these commercial materials not can be obtained, a trend can be extracted regarding the concentration of sulfate versus the crystallinity, being inverse proportional. Assuming that the sulfated materials are calcined under similar conditions, the degree of crystallinity obtained agrees well with results from literature, where increasing the dopant would retard the temperature of crystallization\textsuperscript{[81]}.

<table>
<thead>
<tr>
<th>Sample</th>
<th>BET Area</th>
<th>Crystallinity\textsuperscript{†}</th>
<th>Phase distribution</th>
<th>Surface acidity</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>[m\textsuperscript{2}/g]</td>
<td>[%]</td>
<td>t–ZrO\textsubscript{2}\textsuperscript{‡}</td>
<td>c–ZrO\textsubscript{2}\textsuperscript{‡}</td>
</tr>
<tr>
<td>52Z_NP</td>
<td>52</td>
<td>75</td>
<td>3</td>
<td>0</td>
</tr>
<tr>
<td>97Z_NP</td>
<td>97</td>
<td>100</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>3.45SZ_NP</td>
<td>145</td>
<td>73</td>
<td>10</td>
<td>67</td>
</tr>
<tr>
<td>3.95SZ_NP</td>
<td>130</td>
<td>60</td>
<td>5</td>
<td>36</td>
</tr>
<tr>
<td>4.95SZ_NP</td>
<td>125</td>
<td>53</td>
<td>3</td>
<td>58</td>
</tr>
<tr>
<td>1.7VWT</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

\textsuperscript{†} Calculated in software program Winprep
\textsuperscript{‡} Phase distribution found through Rietveld refinement
\textsuperscript{‡} Total amount of desorbed NH\textsubscript{3} in temperature range 100-1000°C

The phase distribution of the three crystalline ZrO\textsubscript{2} configurations of the commercial samples are outlined in Tab. 3.5, where the two pure ZrO\textsubscript{2} samples consist almost entirely of monoclinic ZrO\textsubscript{2}. The sulfated samples are all polymorph mixtures, comprising mostly of monoclinic
3.2. Zirconia-based catalyst systems

Only 3.4SZ_NP holds significant amounts of the tetragonal phase, namely 10 vol%. However, the high content of c-ZrO$_2$ in the samples is slightly surprising when comparing with literature. The majority of XRD-studies on promoted zirconia report only of the monoclinic and tetragonal ZrO$_2$ phase, only using the strong (111) reflection around 30° 2θ of the latter phase, without mentioning the cubic phase$^{[65,70,117–119]}$. Only a few distinguishes the phases$^{[120,121]}$. On the other hand, differentiation between the two is not straightforward, and only few differences exist, as can be seen in Fig. 3.14, where the diffraction pattern from the three phases have been calculated. Only using the (111) reflection line would therefore not be enough to discriminate t-ZrO$_2$ from c-ZrO$_2$, which is located at almost the same angle. However, the relative intensity of the (111) reflection to the line of ≈35° 2θ or the (004) reflection at 73° 2θ can be used to identify the tetragonal phase.

![Fig. 3.14: Calculated diffraction patterns of the three phases of ZrO$_2$. The plots are for CuK$_{α1}$ X-rays. Unit cells of zirconia: m-ZrO$_2$ (spacegroup P2$_1$/c), a = 5.107 Å, b = 5.174 Å, c = 5.332 Å, $α=γ=90^\circ$, $β=99^\circ$; t-ZrO$_2$ (spacegroup P4$_2$/nmc), a = b = 3.6 Å, c = 5.18 Å, $α=β=γ=90^\circ$; c-ZrO$_2$ (spacegroup Fm3m), a = b = c = 5.118 Å, $α=β=γ=90^\circ$.](image)

3.2.1.2 Determination of surface acidity

The ammonia desorption profiles of the commercial samples are depicted in Fig. 3.15. The total amount of adsorbed ammonia, which is listed in Tab. 3.5, corresponds to molecular adsorbed ammonia or ammonium ions on Lewis or Brønsted acid sites, respectively. The relative strength of the acid sites are reflected by the temperature of maximum ammonia desorption, these are listed in the figure legend.

The broad nature of the low-temperature desorption peaks of sulfate zirconia in Fig. 3.15 can be attributed to a wide range of surface acid sites present with different acid strengths. However, categorizing the exact nature of the acid sites are not possible with NH$_3$-TPD alone, but a commonly used technique is infrared spectroscopy (IR) coupled with pyridine adsorption, where the lone pair of nitrogen in pyridine can coordinate to Lewis acid sites and react with Brønsted sites (forming the pyridinium ion), giving yield to different IR signals which can be used to
differentiate between the two acid sites. Barthos et al.\textsuperscript{[122]} used combined TPD and frequency response (further description of method in\textsuperscript{[123]}) on sulfated zirconia and determined a number of Lewis acid sites desorbing ammonia around 300-430°C and a peak around 470°C attributed as the Brønsted acid site.

The TPD curve for the supports 52Z\_NP and 97Z\_NP showed a bi- and tri-modal desorption profile with peaks at about 190, 300, and 427°C, whereas no desorption peaks are observed at higher temperatures. The overall acidity (listed in Tab. 3.5) of the two pure monoclinic ZrO\textsubscript{2} samples contain an order of magnitude less surface acidity compared to the sulfated ZrO\textsubscript{2}. Only desorption peaks above 430°C are observed for sulfated zirconia, which tentatively can be attributed to the presence of Brønsted acid sites\textsuperscript{[122]}, which does not exist on the pure monoclinic ZrO\textsubscript{2}. Moreover, the absence of any ammonia desorption peak of 1.7V\_WT around 430°C, suggests a low concentration of Bronsted acid sites on this type of catalyst.

At lower temperatures, roughly two types of acid sites can be identified on the sulfated samples, one around 165°C, which is likely to originate from very weakly bound or physisorbed ammonia, and one at around 270-300°C. A number of high-temperature peaks exist for the sulfated samples, the strongest at 660°C, 732°C, and 763°C for 1.7V\_4.9SZ\_NP, 1.7V\_3.9SZ\_NP, and 1.7V\_3.4SZ\_NP, respectively. The temperature of desorption from these strong acid sites are consistent with the findings of Barthos et al.\textsuperscript{[122]}, which detected ammonia desorption on sulfated zirconia at about 630°C, attributed to be of Lewis-acid nature. Besides the very low temperature desorption peaks observed around 165°C in Fig. 3.15, the Bronsted and Lewis acid site designation of Barthos et al. could tentatively be extrapolated to the results observed of the commercial sulfated ZrO\textsubscript{2}. However, the NH\textsubscript{3}-desorption peak of 3.9SZ\_NP and 4.9SZ\_NP from the apparent Brønsted acid site are located at a slightly higher temperature than they reported, namely above 500°C.

Ammonia that is strongly bound to the acid site is not likely to participate in the SCR reaction, which takes place at much lower temperatures, e.g. the vanadia-titania system oper-
ates around 350-400°C. Also, many authors using TPD for characterization of SCR catalysts used lower maximum temperature, e.g. 400°C\textsuperscript{[60]} or 700°C\textsuperscript{[122]}, and Barthos et al.\textsuperscript{[122]} and Katada et al.\textsuperscript{[124]} reported of Brønsted acid sites on sulfated zirconia which releases ammonia below 500°C. Furthermore, the endothermic decomposition of sulfate on \(\text{SO}_2^-/\text{ZrO}_2\) have been found to take place at temperatures above 550°C\textsuperscript{[125,126]}, whereby the surface acidity could change during the TPD experiment. On the other hand, Khodayari et al.\textsuperscript{[60]} suggests that ammonia may promote the decomposition of surface sulfate species during the TPD experiment. Therefore, only peaks with \(\text{NH}_3\)-desorption maxima below 500°C is reasonable to use for the determination of surface acidity relevant to SCR operation. Hence, the surface acidity collated in Tab. 3.5 are subdivided into total surface acidity and surface acidity below 500°C.

Haw et al.\textsuperscript{[127]} postulates that the number of Brønsted acid sites on sulfated zirconia are moderate, consisting only of about 7% of the number of sulfur atoms on the basis of \(^{15}\text{N}\) NMR of adsorbed pyridine-\(^{15}\text{N}\). This was confirmed by Katada et al.\textsuperscript{[124]}, which found a slightly higher number of Brønsted acid sites, namely 12–18%, whereas Wachs and Weckhuysen\textsuperscript{[128]} reported that around 5–10% of the surface vanadia species are surface Brønsted acid sites. Assuming that the peaks observed at 489–526°C of the sulfated zirconia supports are due to Brønsted acid sites. An integration of the peak area versus the total desorption area would then result in the fraction of Brønsted acid sites, where the number of Brønsted acid sites versus total acidity is in the range of 9–15%. Although this number is encumbered with some uncertainty, it is close to the observed in literature.

### 3.2.1.3 Catalytic activity

The commercial samples were impregnated with vanadyl (incipient wetness impregnation, IWI) corresponding to a vanadia loading of 3.0 wt% and calcined at 400°C for 2 hrs in air. Further synthesis description of the catalysts and the 3wt% \(\text{V}_2\text{O}_5\)-10wt% \(\text{WO}_3\)-\(\text{TiO}_2\) (1.7VWT) reference catalyst is given in section A.2 (page 88). Experimental description of the SCR test rig, gas composition and first-order rate constant, \(k\), are given in appendix, section A.3 (page 92).

The SCR activity of the vanadium-impregnated samples have been measured from room temperature to 550°C, presented in Fig. 3.16. The SCR activity of unpromoted 1.7V-97Z_NP reaches maximum conversion of NO around 500°C, only yielding a \(k\)-value around 170 cm \(^3\) g \(^{-1}\) s \(^{-1}\). 1.7V-52Z_NP yields even lower activities, and is therefore not shown here. In contrast, the sulfate-promoted catalysts render somewhat higher SCR activities, where the total activity follow the order 1.7V-4.9SZ_NP < 1.7V-3.9SZ_NP < 1.7V-3.4SZ_NP, with a concurrent decrease in temperature of maximum NO conversion. Thus, 1.7V-3.4SZ_NP present the highest SCR activity of the vanadium-loaded sulfated-zirconia catalysts at the lowest temperature, namely 430°C. In the absence of potassium, the reference catalyst 1.7VWT, based on the traditional vanadia-based catalyst, display a somewhat higher activity with a maximum at 400°C. However, the first order rate constant is here given on a weight basis of the catalyst (cf. section A.3), and the density of titania is somewhat lower than for zirconia (particle density for titania-based catalyst \(\approx 0.2\) \text{g/ml}, zirconia-based catalyst \(\approx 0.5\) \text{g/ml}). Since similar catalyst mass is used for the activity mea-
measurements, the difference in density (catalyst volume) should be taken into consideration in the evaluation of catalyst performance.

The thermal deactivation is an issue for all the catalysts, where lower SCR activity is observed while cooling the catalyst down after treatment at 550°C, except for 1.7V-97Z_NP, which show reversible behavior. The low reversibility of the sulfate-type catalysts could be associated with thermal loss of sulfate species. Based on FTIR measurements at hydrated conditions, Kustov et al.\textsuperscript{[129]} reported a decrease in the intensity of the $\nu_{S=O}$-band at 500°C, and concluded that 25% of the sulfate groups on zirconia were eliminated due to thermal decomposition. On the other hand, the (sulfate free) reference catalyst also suffers from thermal deactivation, which suggests sintering of active centers. The solid-state diffusion of vanadia can be assessed with the so-called Tamman and Hüttig temperatures\textsuperscript{[130]}. Based on the melting point (in Kelvin), it is possible to estimate the temperature where atoms become mobile and sintering can occur, based on the semi-empirical relation:

$$T_{\text{Tamman}} = 0.5T_{\text{mp}}$$  \hspace{1cm} (bulk atoms becomes mobile)$$
$$T_{\text{Hüttig}} = 0.3T_{\text{mp}}$$  \hspace{1cm} (atoms at defects becomes mobile)  \hspace{1cm} (3.2)

For V$_2$O$_5$ ($T_{\text{mp}} = 681°C$\textsuperscript{[10]}) the Tamman temperature, where bulk vanadia exhibit mobility, is obtained at 477°C. Although vanadia does not exist as bulk on the catalyst, but as monolayer with stronger vanadia-support interaction which would increase the actual Tamman temperature, some degree of sintering can not be avoided at 550°C.

![Fig. 3.16: SCR profile of various vanadia-loaded commercial supports and a 1.7VWT reference catalyst. Markers indicate SCR activity while heating, lines the activity while cooling from 550°C](image1)

![Fig. 3.17: SCR activity of various potassium-doped commercial supports and the reference catalyst, K/V = 0.4](image2)

Impregnation of the catalysts with potassium (K/V = 0.4) leads to a considerable decrease of their catalytic activity (Fig. 3.17). This deactivation is pronounced for all catalysts, but in the case of the unpromoted K-1.7V-97Z_NP, an almost complete deactivation observed. In the case of the potassium-doped reference catalyst, severe deactivation is also detected, whereas the sulfate-promoted samples still exhibit SCR activity, especially K-1.7V-3.4SZ_NP. Moreover, the addition
of potassium results in a shift of maximum catalytic activity towards lower temperatures compared with the corresponding catalyst. A possible explanation of such a shift in temperature is that potassium reduces the activity of the main SCR reaction while the rate of the competitive oxidation of ammonia remained constant or even increased.

The SCR activity at 360°C (which is close to the operating temperature of the traditional SCR catalyst) of the fresh and potassium-doped catalysts are listed in Tab. 3.6 for comparison. Based on the measured deactivations, it is clear that the binary vanadia-zirconia catalysts are very sensitive to potassium, which can be explained by the supports low surface acidity combined with the apparent absence of Brønsted acid sites and lack of tetragonal ZrO$_2$. Severe deactivation was also observed for the 1.7VWT catalyst, where loss of more than 80% of the original SCR activity is observed, whereas the catalysts based on sulfated zirconia are less affected by the potassium. Although 1.7V-4.9SZ_NP only deactivated 32%, the overall SCR activity of 1.7V-3.4SZ_NP is higher and still exhibit high resistance towards potassium. The 1.7V-3.4SZ_NP only deactivated 38%, more than a factor of two less deactivated than the vanadia-tungsta-titania catalyst.

**Tab. 3.6:** SCR activities at 360°C of fresh vanadium-loaded zirconia-based catalysts and the corresponding potassium-doped (K/V = 0.4)

<table>
<thead>
<tr>
<th>SCR activity</th>
<th>Deactivation†</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Fresh kg⁻¹s⁻¹</td>
</tr>
<tr>
<td>1.7V-3.4SZ_NP</td>
<td>347</td>
</tr>
<tr>
<td>1.7V-3.9SZ_NP</td>
<td>170</td>
</tr>
<tr>
<td>1.7V-4.9SZ_NP</td>
<td>63</td>
</tr>
<tr>
<td>1.7V-97Z_NP</td>
<td>86</td>
</tr>
<tr>
<td>1.7VWT</td>
<td>794</td>
</tr>
</tbody>
</table>

† Deactivation = \( \frac{k_{360°C \text{ fresh}}}{k_{360°C \text{ K-doped}}} \)

A rough indicator for high resistance toward the poisoning of the basic alkali can be found in the initial surface acidity of the supports. The un-promoted ZrO$_2$-based catalysts possess relatively low surface acidity (less than 650 \( \mu \text{mol/g} \)) and suffer from significant deactivation, as does the 1.7VWT reference catalyst (total surface acidity = 1324 \( \mu \text{mol/g} \)).

If we take into account that the catalytically active Brønsted acid sites consist of less than ca. 20% of all surface acid sites\[^{124,127,128}\], it emphasize the vulnerability of the catalyst. Thus, even a small amount of alkali potentially can deactivate the majority of Brønsted sites, which are essential for the SCR reaction on vanadia catalysts.

### Concise conclusion

Binary V$_2$O$_5$–ZrO$_2$ catalysts were shown to provide low catalytic activity and almost complete deactivation was observed upon alkali-doping due a combination of low surface acidity and/or high abundance of monoclinic ZrO$_2$ in the matrix.

The 3wt% V$_2$O$_5$-10wt% WO$_3$–TiO$_2$ reference catalyst also suffered from significant deacti-
Chapter 3. Catalytic removal of NOx

vation, where less than 20% of the initial SCR activity was present after potassium doping (K/V = 0.4). The low overall surface acidity and apparent absence of Brønsted acid sites are likely contribute to a high sensitivity to the basic potassium.

The sulfated zirconia catalysts of commercial grade presented here contain high surface acidity, and especially 3.4SZ_NP contain the least fraction of monoclinic zirconia among the samples studies and the highest surface area. The corresponding vanadia-loaded catalyst exhibited much higher resistance toward potassium (only deactivating 38%) than the two titania-based reference catalysts. Utilization of sulfated zirconia as support material is thus a promising candidate for increasing the catalysts resistance when confronted with deactivating potassium in the flue gas.

3.2.2 Iron and copper mixed oxide catalysts

Studies similar to those reported previously for vanadium-based catalysts in section 3.1.1.2, have recently also been performed with other oxide catalysts with copper-[51,64,93,131,132] and iron-based catalysts[93,133,134]. Since the metal oxides of copper and iron mainly possess Lewis acidity[51] they belong to an interesting class of SCR catalysts in principle less sensitive to deactivation of the Brønsted acid sites. This was exploited by Kustov et al.[93,114] who studied the effect of potassium on vanadium, copper and iron oxides supported on sulfated zirconia (monoclinic), where a slight increase in alkali resistance was found using the metal oxides of copper and iron compared to vanadium. However, samples with less or no monoclinic phase of ZrO2 were not investigated, and sulfation was performed after calcination of zirconia, which has been found to result in less acidic samples than sulfation of the hydroxide Zr(OH)_2[71]. Thus, in continuation of the study of copper and iron oxides on sulfated zirconia, a series of mixed Cu and Fe oxides supported on the best commercial sulfated zirconia support from previous section were investigated in the following.

The catalysts studied in this section consist of iron-copper mixed oxides in different molar ratio, with a constant total metal loading of 364 µmol/g (similar loading as in[93]) and doped with potassium to K/M = 0.4. The optimal support, 3.4SZ_NP, were used in all catalysts. Further synthesis details are given in appendix A.2.2 (page 89).

3.2.2.1 Temperature programmed desorption

The TPD profiles of the iron-copper mixed oxide on sulfated zirconia are collected in Fig. 3.18. The fresh catalysts primarily consisting of copper oxide (i.e. samples with the iron mole fraction: X_{Fe} = 0.00–0.33) show roughly a tri-modal desorption profile and a shoulder around 260–290°C, whereas for the catalysts comprising mainly iron oxides (i.e. X_{Fe} = 0.50–1.00) only two major desorption peaks are observed and the shoulder is disappeared. The first major desorption peak can be attributed to acid sites generated by the copper and iron oxides, considering that only the small peak at 489°C was present on the TPD profile of 3.4SZ_NP. Moreover, as can be concluded from Tab. 3.7, a gradual shift in temperature of desorption is observed with the relative content of iron oxide in the samples. For the pure CuO-based catalyst (i.e. X_{Fe} = 0.00) the temperature of
maximum desorption below 600°C is located at 345°C, which is increasing slowly with the content of iron up until 402°C for \( X_{Fe} = 1.00 \), suggesting an increase in the strength of acid sites as iron oxide loading displaces copper oxide on the sulfated zirconia. The gradual displacement of CuO with Fe\(_2\)O\(_3\) does almost not affect the overall acidity (cf. inset figure in Fig. 3.18). The disappearance of the minor peak around 490–520°C, combined with the small decrease in overall surface acidity when increasing the fraction of iron oxide, could be attributed to physical coverage of acid sites with Fe\(_2\)O\(_3\) species. At \( X_{Fe} = 0 \), the CuO loading is 3.5 wt% with a corresponding surface density of 2.4 CuO/nm\(^2\) (cf. Tab 3.7, and equation 3.1), whereas at \( X_{Fe} = 1 \), the equimolar concentration of Fe\(_2\)O\(_3\) is 7.7 wt%, corresponding to a surface density of 5 FeO/nm\(^2\). Although monolayer capacity probably is not reached for either of the pure metal oxide catalysts \([84]\), the coverage of iron species is higher, covering or weakening the acid sites of sulfated zirconia.

For all the catalysts, the addition of potassium results in a significant reduction in total acidity, in agreement with previous observations \([72,93]\). Especially the amount of ammonia desorbed at low-temperature peaks around 340–400°C is reduced considerably, whereas the peaks observed at temperatures above 650°C does not seem to have changed upon treatment with alkali. This could suggest that the unaffected desorption peak originates from decomposing sulfate. Furthermore, it should be noticed that the desorption-peak around 340–400°C is shifted towards lower temperatures for the K-treated samples, whereby the acid sites and ammonia adsorption are weakened, presumably due to electron donation from the basic potassium oxide, previously discussed in section 3.1.2.2. An interesting behavior in the surface acidity of the K-

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**Fig. 3.18:** NH\(_3\)-TPD profiles for fresh iron-copper oxides hybrid catalyst and the pure support material as reference (top) and the corresponding potassium-treated samples (bottom), same scale. Inset figure shows the trends in the surface acidity of the Fe-Cu mixed oxide catalysts.
Tab. 3.7: Characteristics of fresh and potassium-treated Fe-Cu mixed oxide catalysts: surface area, surface acidity, and temperature of NH$_3$-desorption during TPD

<table>
<thead>
<tr>
<th>$X_{Fe}$</th>
<th>BET surface area</th>
<th>Total surface acidity</th>
<th>Major TPD peaks &lt;600°C</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Fresh</td>
<td>K-doped</td>
<td>Fresh</td>
</tr>
<tr>
<td></td>
<td>[m$^2$/g]</td>
<td>[µmol/g]</td>
<td>[µmol/g]</td>
</tr>
<tr>
<td>0.00</td>
<td>127</td>
<td>132</td>
<td>1873</td>
</tr>
<tr>
<td>0.20</td>
<td>1999</td>
<td>1706</td>
<td>362</td>
</tr>
<tr>
<td>0.33</td>
<td>1888</td>
<td>1719</td>
<td>367</td>
</tr>
<tr>
<td>0.50</td>
<td>1923</td>
<td>1643</td>
<td>377</td>
</tr>
<tr>
<td>0.67</td>
<td>1969</td>
<td>1628</td>
<td>364</td>
</tr>
<tr>
<td>0.80</td>
<td>1842</td>
<td>1598</td>
<td>365</td>
</tr>
<tr>
<td>1.00</td>
<td>115</td>
<td>128</td>
<td>1829</td>
</tr>
</tbody>
</table>

doped compared to the fresh catalysts is the trend in overall acidity (cf. inset figure in Fig 3.18). Linear fitting of the total surface acidity versus iron mole fraction of the fresh series, reveals an almost constant slope, corresponding to a difference between the pure CuO-based and Fe$_2$O$_3$-based catalysts of on 75 µmol/g, whereas the overall decline of the K-treated catalysts show a difference of around 200 µmol/g. In other words, copper oxide catalysts maintain the surface acidity better upon potassium doping than catalysts consisting mainly of iron oxide.

3.2.2.2 Catalytic activity

The activity of the Fe-Cu oxide catalysts in the catalytic removal of NO were measured in the temperature range of 25–500°C, presented in Fig. 3.19. Here, the catalysts based solely on CuO (on sulfated zirconia) is denoted $X_{Fe} = 0.00$, whereas the iron-based catalyst without any copper oxide is denoted $X_{Fe} = 1.00$. In general, the temperature of maximum activity is shifted from 400°C for the reference catalyst to around 460°C for the hybrid catalysts, along with a decrease in maximum activity. All the fresh catalysts roughly possess similar SCR activity up to around 440°C, at which point the SCR activity is decreasing rapidly for catalysts comprising mainly copper oxide ($X_{Fe} = 0.00–0.67$), whereas the two samples with highest content of iron oxide maintain, or even increase, their high-temperature activity. The shift of the catalytic activity towards higher temperatures for the catalysts with most iron oxide, agrees well with the increase of the overall strength of the acid sites with iron content, as reflected by TPD measurements. The different behavior in the high-temperature region, where sudden loss of activity on the catalysts with most copper oxide, could be a result of thermal deactivation. The Tamman temperature of CuO is 477°C ($T_{CuO}^{mp} = 1227°C$[10]), whereas for Fe$_2$O$_3$ it is 633°C ($T_{Fe2O3}^{mp} = 1539°C$[10]). However, the catalytic activity upon stepwise cooling from 500°C (only shown in figure for the pure iron- or copper-based catalyst) of the fresh and K-doped samples, reveal almost complete reversibility of copper-based catalysts, whereas the iron-based catalysts exhibit some degree of deactivation after treatment at 500°C. Thus, the copper-based catalyst are not thermally deactivated at the highest temperatures, and the reversibility suggests that competing reactions to SCR are dominating at higher temperatures. In contrast, the high coverage of Fe$_2$O$_3$-species (5
3.2. Zirconia-based catalyst systems

Fig. 3.19: SCR activity profile of fresh Fe-Cu mixed oxide catalysts supported on 3.4SZ_NP. The activity when cooling from 500°C for \( X_{Fe} = 0.00 \) and 1.00 is indicated by gray lines, dotted is CuO-based, and solid Fe\(_2\)O\(_3\)-based catalyst.

Fig. 3.20: Temperature dependency of the SCR activity of K-treated Fe-Cu mixed oxide catalysts. K/M = 0.4. Dotted gray line is CuO-based, solid Fe\(_2\)O\(_3\)-based catalyst.

The SCR activity of the Fe-Cu catalysts after impregnation of potassium are shown in Fig. 3.20, where a significant reduction in activity are observed for all catalysts, especially for the hybrid catalysts with highest concentration of iron oxide. However, catalysts comprising primarily copper oxide, namely \( X_{Fe} = 0.00–0.33 \), seem less deactivated.

To obtain a more clear trend in the performance of the Fe-Cu series supported on sulfated zirconia, the activities of the fresh and K-doped catalysts at two selected temperatures are compared and presented in Fig. 3.21. Based on the interpolated activities\(^1\) at 380 and 440°C, the activity of the fresh Fe-Cu mixed oxide catalyst at these temperatures is almost constant, regardless of composition, although catalysts containing most iron oxide exhibit slightly higher activity at 440°C. However, a different trend is observed when the catalysts have been deactivated with potassium, where it is clear that predominantly the copper oxide-containing catalysts (\( X_{Fe} = 0.00–0.50 \)) exhibit higher activity than the corresponding iron-samples.

The degree of deactivation through the series, based on the interpolated values, are depicted in Fig. 3.21 (right). Here, the catalysts with pure CuO as active component, or a high relative content of copper oxides, display somewhat better alkali resistance than catalysts with increasing content of iron oxide. Compared to the \( \text{V}_2\text{O}_5-\text{WO}_3-\text{TiO}_2 \) reference catalyst, which deactivated 83% at 360°C (cf. Tab. 3.6), slightly lower activity loss is observed for the CuO–SO\(_4^2–\)ZrO\(_2\) (\( X_{Fe} = 0.00 \)), deactivating around 75%, whereas the Fe\(_2\)O\(_3\)–SO\(_4^2–\)ZrO\(_2\) is loosing almost all SCR activity, around 90%.

For the potassium-treated catalysts, a similar decreasing trend is observed in surface acidity as well as in SCR activity with increasing iron-oxide content (i.e. inset figure in Fig. 3.18 and Fig. 3.21), suggesting a relationship. Based on TPD observations, the basic potassium is

\(^1\)The temperature obtained during experiment deviate slightly from set-point, and to obtain comparable values the activities are linear interpolated.
3.2.2.3 Concise conclusion

All the catalysts with equimolar copper and iron loadings, a mixture of 3.5 wt% CuO−SO$_2^2$−ZrO$_2$ and 7.7 wt% Fe$_2$O$_3$−SO$_4^2$−ZrO$_2$, display similar surface acidity, although iron oxides displaces the apparent Bronsted sites (490–520°C), originating from the sulfated zirconia support. Increasing the molar fraction of iron oxides in the catalysts, is accompanied by a strengthening of the low-temperature acid site, desorbing ammonia around 345–402°C, which is reflected by the corresponding higher SCR activity of high iron content catalysts (i.e. $X_{Fe} = 0.80$–1.00) at elevated temperatures (<460°C).

SCR activities up until ca. 440°C are almost similar for all fresh catalysts. However, after K-doping it is evident that catalysts comprising mainly copper oxide ($X_{Fe} = 0.00$–0.50) exhibit deactivating a higher number of acidic sites on iron-based catalysts than on copper-based. On the pure Fe$_2$O$_3$-based catalyst, the potassium treatment resulted in complete removal of the TPD peak at 402°C on the fresh catalyst. This indicates that potassium is attacking the lewis sites generated by iron oxides or because of the high surface coverage of iron-oxides, whereby sulfate-generated acid sites are weakened or covered by iron-oxides, unable to effectively host the incident potassium. Furthermore, the presence of the stronger acid site (TPD peak at 489°C for $X_{Fe} = 0.00$), introduced by sulfated zirconia, is likely to appear as a more attractive site for potassium than relatively weaker sites, such as the Lewis sites of the active catalyst.

Finally, only linear behavior were observed in the TPD experiments and SCR activity measurements of the iron-copper mixed oxide series as a function of the molar Cu-Fe content, suggesting that no predominant metal alloys are formed on the catalysts or other symbiotic effects occur.

Fig. 3.21: Left: interpolated SCR activities of Fe-Cu 'alloy' catalysts at 380°C and 440°C versus iron loading for fresh hybrid catalyst and corresponding K-doped samples. Total metal loading 364 µmol/g, equivalent of 3.5 wt% CuO loading. Right: loss of SCR activity versus Fe-Cu oxide composition at 380°C and 440°C.
higher resistance to potassium-poisoning, which were attributed to the presence of strong acid-sites of the sulfated zirconia. These sites were either covered or chemically weakened by the Fe₂O₃-species at higher iron mole fractions.

Although interesting trends were found in the Fe-Cu mixed oxide catalyst series, the ‘best’ catalyst, namely CuO−SO₂⁻−ZrO₂, deactivated around 70%, and were thus only marginally more resistant than the 1.7VWT reference catalyst which deactivated 83%.

### 3.2.3 Concluding remarks

The commercial sulfated zirconia 3.4SZ_NP was selected as support based on its low content of monoclinic ZrO₂, high crystallinity, acidity and surface area. Furthermore, loaded with V₂O₅, the initial activity in the catalytic removal of NOₓ is lower than of the titania-based reference catalyst, 1.7VWT. However, when potassium is deposited (K/V = 0.4 mol/mol), the 1.7V-3.4SZ_NP catalyst exhibit twice better alkali resistance, only deactivating 38% at 360°C, whereas in the case of 1.7VWT, less than 20% of the original activity were left after K-doping. Unpromoted V₂O₅−ZrO₂ catalysts suffer from almost complete deactivation, stressing the significance of surface acidity enhancement, provided by sulfation of the zirconia. For comparison, SCR activities and deactivation due to K-doping of some selected catalysts investigated here, are collated in Tab. 3.8.

**Tab. 3.8:** Comparison of SCR activity at 360°C of the most promising catalysts based on sulfated zirconia (3.4SZ_NP) investigated

<table>
<thead>
<tr>
<th>Catalyst</th>
<th>Fresh SCR activity</th>
<th>Deactivation ‡</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.7VWT</td>
<td>794</td>
<td>83</td>
</tr>
<tr>
<td>1.7V-3.4SZ_NP</td>
<td>347</td>
<td>38</td>
</tr>
<tr>
<td>2.8Cu-3.4SZ_NP†</td>
<td>260</td>
<td>75</td>
</tr>
</tbody>
</table>

† k-value at 380°C
‡ Deactivation after doping, K/M 0.4, IWI

Using the alternative transition metal oxides of iron and copper, which mainly possesses Lewis acidity (in contrast to V₂O₅-based catalysts), show that Cu-based catalysts in general maintain their acid strength better upon K-doping compared to the iron-based catalysts, which is reflected by the corresponding higher resistance toward alkali poisoning in the SCR reaction.

The depositing of potassium on the catalysts leads to a significant decrease of their SCR activity, in concert with an observed shift in the maximum SCR activity to lower temperatures and a reduction in acid strength, probably due to electron donation from the basic potassium oxide. In general, vanadium-based catalysts supported on sulfated zirconia reveal maximum SCR activity at lower temperatures than that of the corresponding copper and iron-based catalysts. However, thermal deactivation is an issue for both iron- and vanadium-based catalysts, revealed by hysteresis in the temperature profile of the SCR activity. On the other hand, the copper-based catalysts investigated here, all exhibit extremely high reversibility, even after treatment at 500°C, making CuO-based catalysts more suitable for high-temperature exposure than e.g. vanadia-
based catalysts.

Fig. 3.22: Comparison of the SCR activity of some selected potassium-poisoned catalysts investigated

The effect of potassium-deposition on the catalytic SCR activity for some selected zirconia-based catalysts are collected in Fig. 3.22 for comparison. From the activity profiles it is clear that vanadium is the most promising active metal in terms of alkali-resistance, and low-temperature activity. It is noteworthy that the maximum in SCR activity of the K-doped V-based catalysts, as well as the titania-based reference catalyst, all are at the same temperature, ca. 380°C, whereas Fe- and Cu-based catalysts typically display maximum activity at higher temperatures. Since the operating catalyst typically performs around 350-400°C, the development of an alternative alkali-resistant catalyst operating within similar temperature range would expedite the implementation. Therefore, for industrial use, vanadium-based SCR catalysts seem to be the best candidate of the catalysts studied here. Following the identification of vanadium supported on sulfated zirconia as a suiting alkali-resistant catalysts, is the issue of upscaling. Until now, all the catalysts have been on particle form and tested at relatively low flows, which is unsuitable for the high flows used in e.g. power plants. Therefore, the next chapter is addressing the issue of catalyst upscaling.
Chapter 4

**ZrO$_2$-sepiolite composite catalysts**

This chapter is dedicated to the upscaling of the best alkali-resistant catalyst found in chapter 3, namely V$_2$O$_5$–SO$_4^{2-}$–ZrO$_2$. For practical reasons, the previous investigations were done on small catalyst particles. However, utilizing such particles under industrial conditions, i.e. high flow and in the presence of fly ash, would quickly result in massive pressure drop and ash-plugging. Hence, the catalysts are shaped into monoliths to avoid these problems. The natural clay material, sepiolite, is used as a binder to shape the catalyst. An introduction of the properties of sepiolite is given below.

### 4.1 Introduction

The incorporation of sepiolite into the catalyst significantly enhances the mechanical properties of both the paste to be extruded and the final monolith, an important factor for its adaptation in industrial applications.

Although sepiolite has only once before been used as binding material for ZrO$_2$-based catalysts$^{[135]}$, the use of sepiolite in TiO$_2$-based catalysts is a well-known technology$^{[136–140]}$. As can be extracted from the references on the subject, a considerable amount of the work done within the use of sepiolite as catalyst binder, are carried out at ICP-CSIC$^1$ in Madrid, Spain. There, I had the pleasure to visit the institute for a few months to learn the technology and produce some ‘beta’-catalysts in collaboration with Pedro Ávila and Søren Birk Rasmussen.

Since almost all the extruded catalysts in this chapter are based on sepiolite, its structural and thermal properties are briefly highlighted here.

#### 4.1.1 Sepiolite structure and thermal behavior

The sepiolite used as binder in this study, Mg$_8$(Si$_{12}$O$_{30}$(OH)$_4$)$\cdot$(H$_2$O)$_{12}$, is a natural hydrous magnesium silicate clay mineral, with fibrous morphology. Sepiolite has a layered structure with phyllosilicate modules composed of two layers of tetrahedral silicate that sandwich the octahedral MgO strip (Fig. 4.1). The ribbons are linked by inversion of the SiO$_4$ tetrahedra Si-O-Si bonds. This structure induces needle-like particle shapes, containing large channels with the dimensions 3.7 x 10.6 Å, running in parallel to the phyllosilicate ribbons$^{[141]}$. In the hydrated form, these tunnels are occupied by zeolitic H$_2$O molecules.

The structure of sepiolite was first reported by Brauner and Preisinger$^{[142]}$, but because of its

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$^1$Instituto de Catálisis y Petroleoquímica, Consejo Superior de Investigaciones Científicas
Fig. 4.1: Dehydration of sepiolite. **Top:** Sepiolite structure at ambient conditions with 8 zeolitic H$_2$O and four structural H$_2$O molecules per unit cell, Mg$_8$(Si$_{12}$O$_{30}$(OH)$_4$)(OH$_2$)$_4$·(H$_2$O)$_8$, viewed along the orthorhombic c-axis. Unit cell $a = 13.405$ Å, $b = 27.016$ Å, $c = 5.2750$ Å, $\alpha = \beta = \gamma = 90^\circ$. **Bottom:** Structure of the 'dehydrated' dihydrate form of sepiolite, Mg$_8$(Si$_{12}$O$_{30}$(OH)$_4$)·(H$_2$O)$_2$ viewed down the monoclinic c axis. Unit cell $a = 23.446$ Å, $b = 11.352$ Å, $c = 5.2782$ Å, $\alpha = \gamma = 90^\circ$, $\beta = 89.06^\circ$. Crystallographic information file (CIF) obtained from[141]

poor crystallinity, resulting in low crystallographic resolution, some details regarding the structure were unanswered. However, recently the structure was resolved by synchrotron powder X-ray, with various degrees of hydration on a purified sepiolite by Post et al.[141]. They found that the hydrated form contain almost 16 H$_2$O molecules per unit cell, where the zeolitic water is distributed over four sites in the channels, and four as structural H$_2$O bound to the terminal Mg atoms in the ribbons. Applying vacuum to the hydrated structure at room temperature only removes about 10% of the zeolitic water, whereas heating to around 200°C results in loss of most zeolitic water, and until about 350°C, the dehydration of sepiolite is reversible, whereby zeolitic water quickly is replaced by humidity.[141–143]. Around 380°C the sepiolite structure collapses under the loss of half of its structural water, causing the phyllosilicate ribbons to rotate around the linking Si-O-Si bond, reducing the cross sectional area of the channels and changing the structural symmetry from orthorhombic to monoclinic (Fig. 4.1, bottom). The last structural water is slowly removed by additional heating to about 600°C, obtaining the ‘anhydrous’ structure (OH...
groups still present)\textsuperscript{[141]}.  

The dehydration temperatures of the purified form of sepiolite suggested by Post et al.\textsuperscript{[141]} do not agree well with those in the literature, mainly because the purity of natural sepiolite is rather low and depends on the origin. Frost et al.\textsuperscript{[144]} compared the gravimetric decomposition profiles of sepiolite from Nevada, Nairobi and Spain, and found that the dehydration and dehydroxylation temperatures varied significantly. However, most authors agree that phase transition occurs below 500°C\textsuperscript{[141,142,144]}, which is the temperature often used for the calcination of sepiolite-titania composite catalysts\textsuperscript{[145]}.

![Fig. 4.2: TGA profile of commercial Sepiolite from Tolsa S.A.](image1)

To evaluate the thermal behavior of the sepiolite applied in this study, the TGA-DSC profile of the as-received clay was performed, the TGA result of this is presented in Fig. 4.2. However, due to its low purity (60%), the dehydration mass losses are significantly damped. To assist in the exact temperature-determination of the weight profile, the curve gradient is calculated and plotted, whereby roughly five steps of mass loss are obtained. The presence of other impurities can be confirmed by the relative molar weight between the hydrated and completely dehydrated sepiolite. The molar mass of Mg\textsubscript{8}(Si\textsubscript{12}O\textsubscript{30}(OH)\textsubscript{4}) is 1080 g/mol and the hydrated Mg\textsubscript{8}(Si\textsubscript{12}O\textsubscript{30}(OH)\textsubscript{4})·12 H\textsubscript{2}O is 1295 g/mol, 17% higher. Therefore, upon complete dehydration of the sepiolite, the mass should decrease by a maximum of 83%, but the relative mass loss at 1000°C is 76%, suggesting simultaneous removal or decomposition of other contaminants, and clouding the assignment of dehydration steps of the sepiolite. However, based on the reported thermal behavior of sepiolite\textsuperscript{[141,144]}, the endothermic mass loss step observed at 416°C in Fig. 4.2 can tentatively be attributed to the phase transition to the monoclinic dihydrate form. It should be mentioned that the choice of the low purity sepiolite is not only for economic concern, the other clay-contaminants play an active role in providing rheological properties to the paste. The before mentioned fibrous character of sepiolite is clearly seen in the SEM micrograph, Fig. 4.3, of a pure extruded sepiolite after calcination.
Chapter 4. ZrO$_2$-sepiolite composite catalysts

### 4.1.2 Pore distribution

Details regarding the porosimetry of the sepiolite can be extracted with mercury intrusion porosimetry (MIP). In Fig. 4.4 the MIP profile is shown for the extruded and calcined sepiolite and reveals a small pore diameter of the pure sepiolite after extrusion, with both meso pores (2–50 nm) and ‘small’ macro pores (50–100 nm). However, it has previously been observed that composite zirconia and sepiolite retain some interparticular porosity even if diminished by the sepiolite component\[^{140}\]. In other words, the pore sizes of the composite catalyst is a blend of pores originating from sulfated zirconia, containing interparticular porosity from the zirconia particles and meso and macro porosity from the sepiolite (cf. Fig. C.2).

In the following section, where sulfated zirconia were mixed into the matrix of sepiolite, the ratio of sulfated zirconia/sepiolite is studied, to find the optimal formulation.

### 4.2 Optimization of sepiolite content

A series of pelletized catalysts with mixed ratios of sepiolite and V$_2$O$_5$–SO$_4^{2–}$–ZrO$_2$ (VSZ) were synthesized for this study. Details regarding the synthesis of the catalysts are described in section A.2.3 (page 90). The catalysts were denoted by the mass fraction of sepiolite, ranging from 0/8 to 8/8, the latter being the pure sepiolite and 0/8 corresponds to V$_2$O$_5$–SO$_4^{2–}$–ZrO$_2$, which is on powder form due to the lack of binder. Some properties of the series are collected in Tab. 4.1, including the elemental analysis of the samples. Based on the content of sulfur, the sulfate concentration on 0/8 is 5.5 wt% (measured as SO$_3$), and therefore slightly higher than the 3.4 wt% on the commercial sulfated zirconia, 3.4SZ_NP. The high content of silicon in the 8/8, suggests the presence of impurities, expected considering the 60% purity of sepiolite.

High BET surface areas are obtained for all samples, which slowly decrease with the sepiolite content.

The SCR activities at 360°C as a function of sepiolite content are presented in Fig. 4.5. The
4.2. Optimization of sepiolite content

Tab. 4.1: Properties of \( V_2O_5-SO_4^{2-}-ZrO_2 \)-sepiolite composite catalysts

<table>
<thead>
<tr>
<th>Sample</th>
<th>Sepiolite content [wt%]</th>
<th>BET area ([\text{m}^2/\text{g}])</th>
<th>IEP*</th>
<th>Elemental analysis</th>
<th>Surface acidity Total &lt;550°C [(\mu\text{mol/g})]</th>
<th>Major TPD peaks†</th>
<th>α</th>
<th>β</th>
<th>γ</th>
<th>δ</th>
</tr>
</thead>
<tbody>
<tr>
<td>0/8 Sep_VSZ</td>
<td>0</td>
<td>165</td>
<td>-</td>
<td>m(_Mg) 5.8 0.0 2.2 0.0 3.9</td>
<td>2131 1141 473 801</td>
<td>-</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1/8 Sep_VSZ</td>
<td>12.5</td>
<td>152</td>
<td>5.4 1.4</td>
<td>m(_S) 4.8 2.7 2.0 6.0 2.7</td>
<td>2495 1169 436 789</td>
<td>-</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2/8 Sep_VSZ</td>
<td>25</td>
<td>154</td>
<td>2.5 5.7</td>
<td>m(_Si) 1.6 1.0 12.6 1.9</td>
<td>2966 715 789</td>
<td>-</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3/8 Sep_VSZ</td>
<td>37.7</td>
<td>153</td>
<td>-</td>
<td>m(_V) 1.6 1.4 14.1 1.5</td>
<td>1985 571 646 790</td>
<td>-</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>4/8 Sep_VSZ</td>
<td>50</td>
<td>153</td>
<td>-</td>
<td>m(_Zr) 2.3 12.0 0.4 25.7 0.0</td>
<td>2255 562 661 797</td>
<td>625 149 7.1 19.1</td>
<td>0.9 18.2</td>
<td>2140 594</td>
<td>642 791</td>
<td>901</td>
</tr>
<tr>
<td>5/8 Sep_VSZ</td>
<td>62.5</td>
<td>149</td>
<td>-</td>
<td>m(_V) 2.2 12.0 0.4 25.7 0.0</td>
<td>2255 562 661 797</td>
<td>625 149 7.1 19.1</td>
<td>0.9 18.2</td>
<td>2140 594</td>
<td>642 791</td>
<td>901</td>
</tr>
<tr>
<td>6/8 Sep_VSZ</td>
<td>75</td>
<td>143</td>
<td>-</td>
<td>m(_Si) 2.0 1.4 14.1 1.5</td>
<td>1985 571 646 790</td>
<td>-</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>7/8 Sep_VSZ</td>
<td>87.5</td>
<td>136</td>
<td>-</td>
<td>m(_V) 2.2 12.0 0.4 25.7 0.0</td>
<td>2255 562 661 797</td>
<td>625 149 7.1 19.1</td>
<td>0.9 18.2</td>
<td>2140 594</td>
<td>642 791</td>
<td>901</td>
</tr>
<tr>
<td>8/8 Sep_VSZ</td>
<td>100</td>
<td>130</td>
<td>-</td>
<td>m(_Zr) 2.3 12.0 0.4 25.7 0.0</td>
<td>2255 562 661 797</td>
<td>625 149 7.1 19.1</td>
<td>0.9 18.2</td>
<td>2140 594</td>
<td>642 791</td>
<td>901</td>
</tr>
</tbody>
</table>

† Sample name correspond to weight fraction of sepiolite mixed with \( V_2O_5-SO_4^{2-}-ZrO_2 \)‡

* Isoelectric point charge

Sample on powder form, i.e. 0/8, were pressed at two tons and fractionated. However, since the sample was prepared slightly different than the extrudates (i.e. not extruded with sepiolite, but pressed and fractionated), the fresh activity is only written in the figure since direct comparison should not be done. As expected, the activity is declining as the fraction of \( V_2O_5-SO_4^{2-}-ZrO_2 \) decreases. However, the drop in activity is higher than can be accounted for by the mere dilution of the active \( VO_x \) species, which would ideally follow a linear dependence, as illustrated in the figure. The apparent discrepancy suggests that sepiolite (or an impurity in sepiolite) either chemically deactivates the active phase or physically blocks the access for the SCR gas molecules. However, the BET areas do not decrease substantially with increasing sepiolite content, ruling out that sepiolite is blocking the pores to the catalyst. The relative surface coverage of sepiolite

![Fig. 4.5: SCR activity at 360°C of fractionated pellets (180-300 µm)](image)

![Fig. 4.6: Apparent surface coverage of sepiolite on the VSZ-sepiolite particle versus sepiolite content, based on Zeta potential measurements](image)
can be elucidated by measuring the zeta potential\(^2\) as a function of pH of the composite catalysts suspended in water. Thereby the isoelectric points (IEP) can be obtained, from which the molar surface coverage of sepiolite can be calculated\(^{[137]}\). The results are shown in Fig. 4.6. The addition of sepiolite entails a dramatic decrease of VSZ at the surface, even by addition of small quantities of sepiolite. At 25 wt% loading, the VSZ particles are almost completely coated by the sepiolite, which might partly be responsible for the rapid decrease in catalytic activity. The coverage of the active particle by sepiolite was also previously observed for TiO\(_2\)-sepiolite mixtures, reported by Knapp et al.\(^{[137]}\). Avila et al.\(^{[146]}\) suggested the kneading of the catalyst and sepiolite, and subsequent calcination causes 'delamination' of the sepiolite fibers in a manner where the catalyst particles were dispersed among the fibers, accounting for the coverage of VSZ particles, even at low loadings of sepiolite.

### 4.2.1 Structural characterization

The samples were further characterized with SEM (Fig 4.7), where the surface morphology of the 0/8 and 1/8 are almost similar, consisting of particles in the 5-10 µm size region. However, with increasing sepiolite content there is a sudden shift in particle appearance and size, reminiscent of particle bundles around 20-50 µm in diameter, thus confirming the sudden shift in surface coverage observed with zeta potential measurements. On the surface on the 2/8–4/8 samples, although it roughly resembles the morphology of the pure sepiolite, it is still possible to identify particles of VSZ linked to the surface, which emphasizes that the zeta potential measurements

\(^2\) difference in potential between dispersed particle and medium
4.2. Optimization of sepiolite content

are based on an average interpolation between the two pure phases. Thus, the particles of zirconia are, more or less, enclosed into the sepiolite cluster at loadings surpassing ca. 12.5 wt%.

The surface acidity of the composite catalysts were measured with NH\textsubscript{3}-TPD and depicted in Fig. 4.8, and reveal a number of different desorption sites, especially at high temperatures. Four distinct acid sites were denoted according to their acid strength: α, β, γ, and δ. The exact desorption temperatures are collected in Tab. 4.1. The samples consisting mostly of VSZ (i.e. 0/8–4/8) display a broad desorption profile around 150–400°C (similar to the profile observed previously for 3.4SZ_NP), whereas only the 0/8–1/8 samples contain an acid site desorbing ammonia around 435–475°C, corresponding to the α-acid site. The elimination of the α-acid site at sepiolite loadings above 25 wt%, suggest that the coverage of the VSZ-particles in sepiolite impede the acid site. The β-acid site is only present on samples containing more than 50% sepiolite, and can thus be attributed to a sepiolite acid site. All samples desorb ammonia around 790–800°C at the γ-site, but only mixtures of sepiolite and VSZ contain sites desorbing ammonia at the strongest δ-site, peaking for the 4/8 sample. The δ-site can thus be attributed to a composite acid site, probably formed at the sepiolite/VSZ interface.

![Fig. 4.8: NH\textsubscript{3}-desorption profile for sepiolite-VSZ composite catalysts. Four desorption sites are observed in the high temperature region](image)

Comparing the overall acidity of the series reveal almost identical total acidity across the series. However, for catalysis applications, the strongest sites of the active catalyst (i.e. γ and δ sites) are assumed to bind ammonia too strongly to participate in the SCR reaction. Therefore, only counting acid sites below ca. 550°C (cf. Tab. 4.1) is much more representative of the catalytic performance observed. The development in acidity, desorbing NH\textsubscript{3} below 550°C, with sepiolite content show a sharp decrease in acidity between the 1/8 and 2/8 sample, which can be explained by the sudden surface coverage of sepiolite, whereas at higher sepiolite loadings, a more moderate decline in acidity is observed as the sepiolite progressively covers the remaining VSZ particles.

The X-ray patterns of the calcined samples are presented in Fig. 4.9. The reflexes from the two phases resembling zirconia, cubic and tetragonal are located at 2θ 30.4°, sandwiched by the
two reflexes from the monoclinic ZrO$_2$ at 28.4$^\circ$ and 31.6$^\circ$. Sepiolite only has few reflexes between 25 and 55$^\circ$, but is acknowledged by the reflex at 39$^\circ$ 2$\theta$. However, several reflexes besides those of sepiolite and zirconia exist. Catalysts with low content of sepiolite, reveal a trace of solid solutions of vanadium in zirconia. At intermediate sepiolite loadings, reflexes from CaSO$_4$ emerges, where Ca$^{2+}$ from sepiolite react with sulfate from the sulfated zirconia. Samples consisting predominantly of sepiolite, reveal reflexes from CaCO$_3$, SiO$_2$, as well as diminutive reflexes, suggesting the presence of iron oxides, and silicates, verifying the high concentration of foreign compounds in the sepiolite. Electron spin resonance (ESR) spectroscopy performed on the samples (presented in Appendix C.1, Fig. C.1) also confirm the presence of Fe$^{3+}$-species in the sepiolite.

To estimate the amount of the aforementioned phases present in the composite catalysts, Rietveld analysis has been performed on the series, the results of which are presented in Fig. 4.10. However, it must be noted that only the crystalline fraction can be observed in XRD, and the results obtained with Rietveld refinement are therefore not necessarily representative of these partially amorphous materials. With increasing sepiolite fraction, the concentration of crystalline m-ZrO$_2$ and c-ZrO$_2$ obtained through Rietveld refinement (Fig. 4.10), are gradually reduced in a linear manner, suggesting that the sepiolite to some degree inhibits formation of crystalline compounds by perturbing the matrix, resulting in amorphous species not detectable by XRD. The increasing fraction of sepiolite, represented by the hydrated and dehydrated conformations, only shows linear behavior for the dehydrated form of sepiolite, whereas the linearity of the hydrated sepiolite breaks off at sepiolite loadings above 50%, suggesting that the orthorhombic structure is stabilized by the zirconia at low sepiolite loadings. The low concentration of dehydrated monoclinic sepiolite, could suggest that the zirconia-based particles perturb the dihydrate-structure of sepiolite, forming amorphous monoclinic sepiolite, not detected with XRD. This is supported by the structure of TiO$_2$-particles embedded within the fibers of sepiolite proposed by Ávila et al.[140,146]. Thereby, the actual concentration of the monoclinic structure of
4.2. Optimization of sepiolite content

The formation of CaSO\(_4\) in composite samples consisting mostly of sepiolite, i.e. 5/8-7/8, constitute a rather large fraction of the total crystalline fraction, responsible for around 10 wt% (Fig. 4.10, right). A considerable amount of sulfate is therefore abstracted from the sulfated zirconia and diminishing its acidity, probably responsible for the enhanced decrease in SCR activity with sepiolite content observed in Fig. 4.5. The extraction of sulfate from zirconia, reducing its acidity, imply that additional sulfate should be added to the mixture during synthesis to compensate for this effect. This was attempted by impregnating the synthesis mixture of the 2/8 catalyst with additional ammonium sulphate. However, the effect on the SCR activity was negligible, although a small maximum was found, the results are shown in appendix C (Fig. C.4). Due to the insignificant impact on activity, optimization by means of additional sulfation were not pursued any further.

Based on the results obtained with MIP of the composite catalysts (shown in appendix, section C.1), the pore sizes of the catalysts can be extracted, and are presented in Fig. 4.11. Increasing the sepiolite content of the V\(_2\)O\(_5\)−SO\(_4^{2-}\)/ZrO\(_2\)/sepiolite composites has a marked effect on the observed pore volumes and macro pore diameter. By introducing 25 wt% of sepiolite the characteristic interparticular macro pore diameter decreased from 911 nm (0/8) to 320 nm, whereas the meso pore diameter remained almost constant throughout the series.

Finally, the physical strength of the pelletized catalysts were evaluated, the results from this are depicted in Fig. 4.12. As expected, the overall robustness of the pelletized catalysts increase with the sepiolite content. However, an abrupt increase in pellet strength were observed for the 2/8 sample, constituting a fair compromise between catalytic activity and physical strength. Furthermore, the coating effect of sepiolite at low loadings may provide some degree of shielding effect, enhancing the durability of the operating catalyst. This is further explored in the following section.
Chapter 4. \(\text{ZrO}_2\)-sepiolite composite catalysts

\[
\text{Fig. 4.11: Development in pore diameter of macro- and meso pores obtained from MIP with increasing sepiolite content}
\]

\[
\text{Fig. 4.12: Physical strength of produced pellets. Robustness measured by applying vertical force on identical sized pellets until breakage, as outlined in figure}
\]

\[
\text{4.2.2 Alkali-resistance of composite catalysts}
\]

The resistance toward potassium-poisoning of the composite catalysts were evaluated. Two methods of alkali-depositing were explored: incipient wetness impregnation (IWI) and hydrothermal treatment of the catalysts pellets submerged in fine particles of KCl as outlined in Fig. 4.13. The latter method, inspired by Zheng et al. [95], were performed to mimic the deactivation thought to occur through surface diffusion of alkali-species hitting the catalyst surface and migrating to the active sites of the SCR catalyst during operation [88]. An actual \textit{in situ} deactivation would require catalyst exposure to aerosol alkali-particles, so this method is a crude simplification, but considerably easier to carry out.

The impact of potassium introduced via IWI and hydrothermal treatment in KCl under humid air at 350°C for 188 hrs (7.8 days) and 384 hrs (16 days) on the catalytic performance, are presented in Fig. 4.14 together with the 1.7VWT reference catalyst. Since a threshold of physical strength is required of the catalysts, only the 2/8, 4/8 and 6/8 composite catalysts were tested here. The catalytic activity of the 1.7VWT reference catalyst yielded considerably higher SCR activity than all of the composite catalysts ‘diluted’ with sepiolite, and the 6/8 catalyst with 75% sepiolite exhibited zero activity. The effect of hydrothermal treatment with KCl only moderately retarding the SCR activity, although potassium is present on the catalyst, as can be evaluated by SEM-EDS images provided in appendix (Fig. C.3), though potassium could be present as lumps of potassium, not effectively bound to the active sites. However, the IWI treatment significantly
4.2. Optimization of sepiolite content

decreased the SCR activity of all catalysts. The relative activities present after alkali-poisoning are depicted in Fig. 4.15, and show that all catalysts exhibited roughly similar loss of activity due to the KCl-treatment. The reference catalyst deactivated 21%, and 2/8 composite catalyst deactivated 16% after 384 hrs submersed in KCl.

The low degree of deactivation suggests that the treatment temperature should be increased, at least to above the Hüttig temperature of KCl (422°C). At 350°C, the relative large KCl particles may never reach the majority of the active sites, e.g. due to pore blockage, where the macro pores of the catalysts (pore diameter of 1.7VWT: 298 nm, 2/8: 320 nm) could hinder the access of the KCl species. In contrast, K-depositing by IWI effectively penetrates both macro and meso pores, deactivating the reference catalyst 80% (i.e. remaining activity 20%), whereas the composite catalysts 2/8 and 4/8 only deactivates around 60%.

The reproducibility of the IWI method is confirmed by the 1.7VWT catalyst, which were earlier K-doped by IWI in a different experiment, where a deactivation of 83% were observed (cf. Tab 3.6). The relatively high deactivation of the sepiolite-containing composite catalysts is somewhat higher than previously observed for the 1.7-3.4SZ_NP catalyst, which only deactivated 38% under similar conditions. This suggests that sepiolite further reduces the alkali resistance of the catalysts, i.e. by deacidification through sulfate abstraction. However, the high vanadia content of the composite catalysts would also be more vulnerable to potassium deactivation than catalysts with lower loading\(^{[90]}\).

### 4.2.3 Concise conclusion

The increasing content of sepiolite-binder in the catalyst certainly provide beneficial effects in terms of physical strength and possibility to manipulate the macro pore diameter of the extruded catalyst. On the other hand, the introduction of sepiolite also entails loss of surface acidity (damping of the $\alpha$-acid site) through abstraction of sulfate from $V_2O_5(SO_4)^2- - ZrO_2$, forming $CaSO_4$, and thereby reducing the SCR activity of the composite catalysts.
By electrophoretic migration experiments it was shown that zirconia-based catalysts containing sepiolite concentrations above 12.5 wt% are, more or less, coated by the clay fibers. Although the suggested shielding effect of the enveloping sepiolite could be affirmed here, the concept is an interesting approach in reducing the impact of potassium particles in biomass fired power plants.

The most promising candidate of the composite catalysts studied here, as a compromise between SCR activity and physical stability, is the 2/8 catalyst, comprising 75% VSZ in sepiolite. To further evaluate the performance of the optimal catalyst, the 2/8 formulation were extruded as monolith and the results are presented in the following section.

### 4.3 Evaluation of monolithic catalyst

The optimal catalyst, in terms of catalyst stability and SCR activity, was in section 4.2 identified as the composite catalyst comprising 75% VSZ in sepiolite, the 2/8 sample. The 2/8 formulation was extruded as cylindrically-shaped monoliths with 24 parallel channels, as presented in Fig. 4.16.

#### 4.3.1 Catalytic performance

To ensure that the monolithic catalyst was comparable to the catalysts synthesized in connection with the series with increasing sepiolite content, the monolith were crushed to the same particle size as the 2/8 pellet catalyst from previous section. The catalytic activity of the two samples are depicted in Fig. 4.17, which reveal almost identical SCR activities. High-temperature annealing at 500°C in both cases result in significant thermal deactivation to the catalysts, as observed previously for V-based catalyst.

![Fig. 4.16: Various shapes of the extruded monolith. Only the cylindrically shaped monolith were used for the results presented in this section](https://example.com/fig416.jpg)

![Fig. 4.17: Comparison of the SCR activity profile of mechanically and manually extruded catalysts, subsequently fractionated to 180–300µm](https://example.com/fig417.jpg)

Interpolation of the k-values to 360°C show only an insignificant difference in activity, af-
firming that the monolithic catalyst produced is indeed representative of the 2/8 catalyst formulation. Considering that the catalyst synthesis has been scaled up about 50 times, it is remarkable that the SCR activities are identical to those of the catalyst manually kneaded and extruded.

The SCR activity of the extruded 2/8 monolith was tested as described in appendix, section A.2.3.3. The activity at different gas hourly space velocities (GHSV, STP) are obtained by varying the gas flows (but keeping the linear velocity above 1.2 m/s), presented in Fig. 4.18. The overall activity is naturally increased with decreasing GHSV (i.e. longer residence time). At GHSV = 10700 h⁻¹ the conversion profile stagnates at temperatures above 300°C, suggesting that the SCR reaction become diffusion limited and/or mass transfer limited on the 2/8 catalyst under these circumstances. In other words, the SCR reaction is proceeding at a rate faster than the reactants can be transported to the reactive sites.

A convenient representation of the conversion is as a function of the gas residence time, $\tau$, which can be expressed as:

$$\tau = \frac{1}{\text{GHSV}} = \frac{V_{\text{cat}}}{q} = \frac{L_{\text{cat bed}}}{v_L}$$  \hspace{1cm} (4.1)

where $V_{\text{cat}}$ is the catalyst volume, $q$ the total gas flow, and $v_L$ the linear velocity. Under the given test circumstances, it is not possible to reach a higher contact time due to the diffusion limitation observed at 10700 h⁻¹ ($\tau = 0.34$ s). However, based on the rate expression for the first order reaction provided in equation A.5 (page 93), it is possible to extrapolate the conversions to longer residence times. Unit analysis reveal that the rate expression of the first order rate constant, $k$, can be rewritten:

$$k = -\frac{F_0}{[\text{NO}]_0 W} \ln (1-X) \Rightarrow -\frac{q}{W} \ln (1-X)$$  \hspace{1cm} (4.2)

The catalyst weight is given by its volume and bulk density ($W = \rho_{\text{bulk}} V_{\text{cat}}$). By applying equa-
tion 4.1 the residence time emerges, which can be rearranged as a function of $\tau$:

$$k = -\frac{q}{\rho_{\text{bulk}} V_{\text{cat}}} \ln (1-X) \Leftrightarrow k = -\frac{1}{\rho_{\text{bulk}}} \frac{\ln (1-X)}{\tau}$$ (4.3)

Thus, by plotting $-\ln (1-X)$ as a function of residence time, $k \rho_{\text{bulk}}$ can be obtained from the slope. The relationship is presented in Fig. 4.19. The first order rate constants of the monolith can be calculated since the bulk density is known, $\rho_{\text{bulk}} = 1.45 \text{ g/cm}^3$. The results are outlined in Table 4.2, where the $k$-values obtained from the measurement of the fractionated monolith ($k_{\text{powder}}$) are also provided. The ratio between the $k$-values of the monolith and fractionated catalyst represents the efficiency of the monolith, under the assumption that no mass transfer limitation is occurring on the fractionated catalyst, as was the case at 350°C for the monolith (cf. Fig. 4.18). The efficiencies observed here correspond well with those found in literature\[147].

<table>
<thead>
<tr>
<th>Catalyst temperature</th>
<th>250°C</th>
<th>300°C</th>
<th>350°C</th>
</tr>
</thead>
<tbody>
<tr>
<td>$k_{\text{mon}}$ [cm$^3$g$^{-1}$s$^{-1}$]</td>
<td>2.4</td>
<td>4.2</td>
<td>5.0</td>
</tr>
<tr>
<td>$k_{\text{powder}}$ [cm$^3$g$^{-1}$s$^{-1}$]</td>
<td>4.7</td>
<td>31.8</td>
<td>131.7</td>
</tr>
<tr>
<td>$\eta$ † [%]</td>
<td>51.3</td>
<td>13.2</td>
<td>3.8</td>
</tr>
</tbody>
</table>

† Catalyst efficiency, $\eta = k_{\text{mon}}/k_{\text{powder}}$

By rearranging equation 4.4, the NO conversion ($X$) can finally be expressed as a function of $\tau$ (outlined in equation 4.5) which can be extrapolated to higher residence times, assuming absence of mass transfer limitations, as presented in Fig. 4.20.

$$X = 1 - \exp(-k \rho_{\text{bulk}} \tau)$$ (4.5)

Based on the parameters found, a projection of the catalyst activity at lower space velocities can be calculated. 98% conversion is achieved at $\tau = 0.6$ s in the 2/8 monolithic catalyst at 350°C, and assuming a linear gas velocity of 2 ($v_L = 1–4 \text{ m/s}$\[148]) over honeycomb $V_2O_5$–$WO_3$–$TiO_2$ catalysts under industrial conditions) would require a catalyst length ($L_{\text{cat bed}}$) of 1.2 m to ensure total NO$_x$ removal. As an example of industrial conditions, a 250 MW coal-fired power plant in Belgium is used. The plant is operating with a Haldor Topsøe SCR catalyst in the temperature regime 320–390°C under a total flow of 820,000 Nm$^3$/h with a catalyst volume 268 m$^3$ in two layers of 2 m in total length, and a NO$_x$-concentration of 1000 ppm\[149], yielding a GHSV (STP) of 3060 h$^{-1}$ ($\tau = 1.2$ s). Thus, based on the projections in Fig. 4.20, the extruded 2/8 monolith would under industrial conditions ($\tau = 1.2$ s, 350°C) remove all the NO present, and more than 98% at 250°C. Considering that the ZrO$_2$-based monolith exhibit somewhat higher tolerance to potassium-poisoning than the traditional $V_2O_5$–$WO_3$–$TiO_2$ catalyst, the lifetime of the zirconia-based catalyst may be considerably longer than the traditional catalyst in biomass-fired
4.3. Evaluation of monolithic catalyst installations. However, long time exposure of the sepiolite-based VSZ catalyst has not yet been carried out, and of course this would need to be addressed.

![Figure 4.20: NO conversion as a function of residence time by varying reactor temperature and space velocity](image1)

![Figure 4.21: Conversion of NO versus the ammonia/NO ratio. NH<sub>3</sub>-slip and evolution of N<sub>2</sub>O concentrations on right axis](image2)

### 4.3.2 Influence of feed ratio

The gas inlet concentration of NH<sub>3</sub> was varied, while fixing the NO concentration at 700 ppm in order to evaluate the experimental impact of the feed ratio \( \alpha (\text{[NH}_3\text{/}[\text{NO}]) \) on the SCR activity of the 2/8 monolith at 350°C. The NO conversion as a function of the the feed ratio is presented in Fig. 4.21, where it can be observed that the NO conversion is stoichiometrically and kinetically limited by the NH<sub>3</sub>-concentration at substoichiometric feed ratios (\( \alpha < 1 \)), in line with results reported by Bahamonde et al. on VWT-sepiolite monoliths\textsuperscript{[136]}. Furthermore, at \( \alpha < 1 \), the rate expression used so far (equation A.1, page 93), \( r = k[\text{NO}]^1[\text{NH}_3]^0 \), fails to describe the reaction rate effectively, because it was derived assuming a NH<sub>3</sub>-surplus. However, under these circumstances, a Rideal-type reaction rate has previously been used to describe the rate of reaction\textsuperscript{[150,151]}:

\[
r = k[\text{NO}]\theta_{\text{NH}_3}
\]

where \( \theta_{\text{NH}_3} \) is the fractional surface coverage of absorbed ammonia on the active sites, and represents the impact of ammonia on the activity.

In excess ammonia (\( \alpha > 1 \)) the NO conversion becomes independent of the feed ratio (cf. Fig. 4.21), and the rate expression in equation A.1 holds true, where the ammonia concentration is described by zero order kinetics. Due to the 1:1 stoichiometry of the SCR reaction, an ammonia slip is observed at \( \alpha > 1 \). It is worth noticing that almost 90% of the NH<sub>3</sub>-surplus at \( \alpha = 1.11 \) is not oxidized to NO\textsubscript{x}, but remains on NH<sub>3</sub> form. However, at industrial conditions the feed ratio is operated at \( \alpha = 0.8 \)\textsuperscript{[136,152]} to avoid ammonia slip. Under these conditions (350°C, GHSV = 10700 h\textsuperscript{-1}), a high catalytic activity is still provided by the 2/8 monolith, removing around 80% of the initial NO, the same behavior as the titania-based catalyst\textsuperscript{[136,147]}.
4.3.3 Summary and conclusion

The catalyst formulation containing $V_2O_5\text{--SO}_4^{2--\text{ZrO}_2}$ in 25% sepiolite was successfully upscaled to a monolith. The monolithic configuration exhibited exactly the same SCR activity (when fractionated) as the manually kneaded and extruded pellet, and were thus representative of the optimal 2/8 formulation.

Catalytic characterization of the cylindric monolith at different space velocities and feed ratios revealed fairly high SCR activities, although mass transfer and/or diffusion limitations in this system commence around 350°C with the linear velocity $1.2\ m/s$ and $\text{GHSV} = 10700\ h^{-1}$. However, extrapolation of the activity to lower space velocities were carried out by incorporating the residence time, $\tau$, into the expression for the first order rate constant. Thus, the activity of the 2/8 monolith could be predicted for conditions similar to those of the industrial $V_2O_5\text{--WO}_3\text{--TiO}_2$ catalyst ($\tau = 1.2\ s$, 350°C). According to these predictions, the ZrO$_2$-based catalyst would perform equally well, removing more than 99.9% of the incident NO$_x$ gases. Furthermore, the advantage of the ZrO$_2$-based catalyst become most evident if alkali species are present in the flue gas, due to the higher potassium resistance when compared to the titania-based catalysts. Thus, in biomass-fired installations the catalyst lifetime could be considerably prolonged. However, long term exposure of the catalyst under in situ conditions are yet to be addressed for the 2/8 catalyst.

4.4 Conclusion

The mixing of $V_2O_5\text{--SO}_4^{2--\text{ZrO}_2}$ with the the binding clay, sepiolite, provided physical stability to the catalyst, rendering the extrusion as monolith or pellets possible. Electrophoretic migration experiments and SEM revealed that already at small concentrations of sepiolite (above 12.5 wt%) the VSZ particles were enveloped by the clay fibers. Beyond the ‘threshold’ clay-concentration, other noticeable structural changes of the composite samples entailed: increased physical stability particle aggregation; decrease of surface acidity; and macro porosity. As a compromise between catalytic activity and physical stability, the formulation containing VSZ in 25% clay was selected as the optimal catalyst.

Although sepiolite was found to abstract sulfate from the VSZ particles in mixtures comprising predominantly of sepiolite (>50%), thus reducing the overall acidity and SCR activity, the presence of sepiolite on the surface of the active particles could provide some protective effect.
due to its meso porosity, prohibiting larger alkali particles access to the active sites, or simply postponing the poisoning. However, only indicative evidence for this effect was found through the slightly higher alkali-resistance of the sepiolite-based catalyst over the VWT-reference catalyst when submersed in fine particles of KCl and hydrothermally treated.

The zirconia-sepiolite formulation could be extruded in monolithic form, and contained promising SCR activity under conditions similar to industrial circumstances. This study thus opens for the possibility of further investigation into this type of SCR catalyst as a candidate, less susceptible to potassium-poisoning than the traditional titania-based catalyst.
Chapter 5

NO absorption in ionic liquids

This chapter is devoted to study NO gas removal by absorption in ionic liquids. It should be noted that absorption of gaseous NO in ionic liquids have not yet been reported in literature. Thus, due to the novelty of the field, the system is not completely understood. The removal of NO by ‘wet’ absorption is however not a new phenomena and some other processes are summarized in the following introduction as well as a preface to ionic liquids.

5.1 Introduction

Although the catalytic removal of NO\textsubscript{x} from the flue gas is a very effective process, the overall high operating expenses of the SCR process and possibility of ammonia slip, have motivated a search for other methods to abate emissions of nitrogen oxides\textsuperscript{[153,154]}. A promising concept is wet scrubbing systems for removal of SO\textsubscript{2} and NO\textsubscript{x}. Some aqueous scrubbing systems have been developed for the simultaneous removal of NO\textsubscript{x} and SO\textsubscript{2}\textsuperscript{[155,156]}. The wet flue gas desulfurization (FGD) typically exhibit high SO\textsubscript{2} removal efficiencies, but the FGD can only remove a small amount of NO\textsubscript{x} because about 90–95% in a typical flue gas is present as insoluble NO and only the remaining 5-10% NO\textsubscript{2} is water soluble. Attempts to oxidize NO to water soluble NO\textsubscript{2} have been made by adding strong oxidizing additives, such as MnO\textsubscript{4}\textsuperscript{-} and H\textsubscript{2}O\textsubscript{2}, but the treatment cost involved herein has been too high for practical utilization\textsuperscript{[155]} (and references herein).

Promising results of the simultaneous NO and SO\textsubscript{2} removal in a Co(NH\textsubscript{3})\textsubscript{2}\textsuperscript{2+} solution have been reported by Long et al.\textsuperscript{[155,157]}, which operates below 80°C. Another approach is complexation of NO with Fe\textsuperscript{2+}-chelates based on ethylenediaminetetraacetic acid (EDTA) or nitrilotriacetate (NTA)\textsuperscript{[158]}, as outlined in reaction R-5.1 for the case of iron-EDTA complex\textsuperscript{[154]}.

\[
\text{Fe}^{II}(\text{EDTA}) + \text{NO} \rightarrow \text{Fe}^{II}(\text{EDTA})(\text{NO}) \quad (\text{R-5.1})
\]

The metal-chelate can be electrochemically regenerated after absorption or reduced by sulfite ions to sulfate and nitrogen\textsuperscript{[154]}. Recently a biotechnological approach were described to regenerate the iron-complex, the so-called BioDeNO\textsubscript{x} process. In this process the NO-saturated iron-chelate solution is brought in close contact with bacteria that regenerate the iron-EDTA complex and converts the bound nitrosyl to N\textsubscript{2}\textsuperscript{[159]}. The Fe\textsuperscript{III}(EDTA) solution need to be somewhat diluted (<200 mM) due to the presence of microorganisms, which naturally limits the absorption capacity.
Chapter 5. NO absorption in ionic liquids

5.1.1 Ionic liquids

All of the above proposed technologies are based on liquids with a vapor pressure, which means that the solvent, to some extent, vaporizes. One promising solution to this problem could be the use of a relatively new class of solvents referred to as ionic liquids (ILs). The expression ‘ionic liquid’ in principle encompasses any liquid entirely composed of ions (e.g. molten salts), but within the context of this work the term will only be used to describe materials which are liquid in their pure state at room temperature. This class of solvents are often considered as ‘green’ solvents because of their immeasurably low vapor pressure[160] (and references herein). This feature gives the ILs an essential advantage over traditional solvents used for absorbing gases. Ionic liquids have already demonstrated promising behavior in a number of reactions where gaseous reactants enter the IL solution (such as hydrogenation, hydroformylation, and oxidations) despite low gas solubilities of the gases in the IL at ambient conditions[160,161].

Another application of the ILs is to utilize them to separate gas mixtures. Recently, a promising solid ionic cation (1,1,3,3-tetramethylguanidinium) have been identified[162] for the absorption of SO$_2$. Anthony et al.[160] reported of the solubilities of a number of gases (such as CO$_2$, CO, O$_2$) in imidazolium-based ILs.

Ionic liquids tend to be more viscous compared to conventional solvents[163], which can result in challenges regarding the mass transfer of gas into the IL. In the case of low-soluble gases, the mass transfer into the IL will likely be a rate limiting step, which can be minimized by increasing the interfacial gas-IL area and/or use high pressure systems[160].

So far, only little information regarding the gas solubilities in ILs is reported. Besides the reports regarding CO$_2$ capture[164] (and references herein) the focus of most works evolve around the reactions taking place in the IL with the gas already absorbed. Only few reports exist on gas solubilities[160,165,166], but especially Brennecke et al. have contributed with a number of seminal studies on absorption of a number of gases in imidazolium-based ILs[160,166–168].

The focus of the present work is to study the absorption and desorption behavior of NO in ILs based on the cation 1-ethyl-3-methylimidazolium (EMIM$^+$) and 1-butyl-3-methylimidazolium (BMIM$^+$).

5.2 Screening of ionic liquids

The absorption of NO in a number of different ILs were investigated in this preliminary study, all based on the cations and anions presented in Fig 5.1. Some physical properties of the pure ILs used (supplier and purity listed in Appendix, section A.4.3) are collected in Tab. 5.1, all revealing viscosities many times higher than that of water. The experimental procedure to measure the NO absorption capacity in the ILs are based on a rudimentary setup, described in section A.4.1. The absorption time was several days due to slow mass transfer from gas to IL, and the screening-experiments were therefore stopped before complete equilibrium were obtained. The resulting solubilities presented in this section are therefore based on pseudo-equilibrium
5.2. Screening of ionic liquids

Fig. 5.1: The cations and anions used in this study. Lower box, from left to right: acetate, triflate, tetrachloroaluminate and thiocyanate

measurements. However, the apparent solubilities obtained here still provide sufficient information to identify suited candidates for further studies.

Tab. 5.1: Properties of the ionic liquids used in the preliminary study. Data supplied by BASF†

<table>
<thead>
<tr>
<th>Compound</th>
<th>M [g/mol]</th>
<th>Viscosity at RT [mPa·s]</th>
<th>Density at RT [g/cm³]</th>
<th>T mp [°C]</th>
</tr>
</thead>
<tbody>
<tr>
<td>[EMIM]AlCl₄</td>
<td>280.0</td>
<td>26</td>
<td>1.30</td>
<td>9</td>
</tr>
<tr>
<td>[BMIM]AlCl₄</td>
<td>308.0</td>
<td>32</td>
<td>1.24</td>
<td>-10</td>
</tr>
<tr>
<td>[EMIM]SCN</td>
<td>169.3</td>
<td>22</td>
<td>1.11</td>
<td>&lt;-20</td>
</tr>
<tr>
<td>[BMIM]SCN</td>
<td>197.3</td>
<td>54</td>
<td>1.07</td>
<td>&lt;-20</td>
</tr>
<tr>
<td>[EMIM]OAc</td>
<td>170.2</td>
<td>93</td>
<td>1.10</td>
<td>&lt;-20</td>
</tr>
<tr>
<td>[BMIM]OAc</td>
<td>198.3</td>
<td>398</td>
<td>1.06</td>
<td>&lt;-20</td>
</tr>
<tr>
<td>[EMIM]OTf</td>
<td>260.2</td>
<td>41</td>
<td>1.34</td>
<td>-11</td>
</tr>
<tr>
<td>[BMIM]OTf</td>
<td>288.3</td>
<td>84</td>
<td>1.30</td>
<td>17</td>
</tr>
<tr>
<td>[OMIM]OTf†</td>
<td>344.4</td>
<td>n.a.</td>
<td>n.a.</td>
<td>n.a.</td>
</tr>
<tr>
<td>H₂O</td>
<td>18</td>
<td>1</td>
<td>1</td>
<td>0</td>
</tr>
</tbody>
</table>

† BASF: www.basionics.com/en/ionic-liquids/products
‡ 1-octyl-3-methylimidazolium triflate

5.2.1 Absorption/desorption study

All the ILs presented in Tab 5.1 were evaluated with respect to their NO sorption performance. However, not all ILs were suitable for the liquid sorption studies. The ILs based on AlCl₄⁻ and SCN⁻ all formed precipitate upon contact with the NO/N₂ gas mixture and the absorption results were thus discarded. Although the NO-induced crystallization of the ILs observed are interesting and definitely worthwhile to study further, the focus were here maintained on absorbents which remained in their liquid state upon NO exposure. Hence, only the sorption results of the ILs with the acetate and triflate anion are presented in Fig. 5.2 as a function of the apparent mole fraction of NO absorbed (X₉₅/NO). Considering the acetate-based ILs, [EMIM]OAc and [BMIM]OAc, it is clear that significant amounts of NO can be absorbed, corresponding to about 2 moles of NO per IL molecule (equal to a NO mole fraction of 0.667). Interestingly, varying the cation of the acetate-based ILs only seem to influence the NO solubility to a minor extent,
in contrast to the triflate-based ILs where the absorption capacity increased with alkyl length. Thus, the [OMIM]OTf exhibit highest solubility of NO of the triflate-based ILs, absorbing about 1 mole of NO per mole of IL (mole fraction 0.5). NO absorption capacities in ionic liquids of this size have not previously been reported, to our knowledge. Only the absorption of SO$_2$ in the tetramethylguanidium-based ILs, [TMGHPO]$_2$N or [TMGHPO$_2$]$_2$N, have been reported to yield similar solubilities, i.e. mole fractions about 0.67$^{[162,169]}$.

The reversibility of the gas absorption were examined by applying vacuum to the NO-absorbed IL at room temperature for at least 24 h. The remaining NO in the IL, expressed by the mole fraction of NO, is also presented in Fig. 5.2. Only a small amount of NO could be removed from the acetate-based ILs by reducing the pressure, suggesting that the NO is strongly bound in the [EMIM]OAc and [BMIM]OAc. On the other hand, the reversibility of the NO sorption in the ILs based on triflate is somewhat higher. Almost all the absorbed NO could be desorbed in the case of [BMIM]OTf, but only about $1/3$ of the NO could be removed from the [OMIM]OTf under the examined conditions.

Preliminary examination of the absorption of N$_2$ in [BMIM]OAc were also undertaken. This did not result in any measurable weight increase, from which it can be concluded that only NO is absorbed when passing NO/N$_2$ through the [BMIM]OAc.

### 5.2.2 Influence of metal-promotion on absorption capacity

It is well known that iron can bind NO in form of the brown iron nitrosyl complex Fe(NO)$_{2+}$$^{[19]}$. The approach of incorporating a complexation or redox agent into the ionic liquid was explored in the following section where enhancing the absorption capacity of the ILs was attempted by metal promotion. Based on the preliminary sorption study (section 5.2.1), the [BMIM]OTf was selected due to its high sorption reversibility. The metal promotion was carried out by adding the triflate-salt of a metal (Fe(OTf)$_3$, Co(OTf)$_2$, and Cu(OTf)$_2$) to the [BMIM]OTf to obtain a metal concentration of 0.5 mol/l. The absorption capacities of these metal-promoted ILs were examined, and the results are depicted in Fig. 5.3 as a function of ‘time-on-stream’. The results
5.2. Screening of ionic liquids

![Fig. 5.3: NO absorption time-profile at 20°C for [BMIM]OTf with metals of iron, copper and cobalt dissolved. Absorption performed with 10% NO/N\textsubscript{2} on a setup similar to the one used to obtain the results presented in Fig. 5.2 (described in section A.4.1), but without a glass frit on the pipette. Thus, the time of absorption are longer due to lower gas-IL interfacial area.]

The solubility of NO in [BMIM]OTf is not further enhanced by promotion with cobalt or copper triflate. As can be observed in Fig. 5.3, the addition of cobalt actually decreases the solubility of NO in [BMIM]OTf significantly, although only 0.5 M cobalt was added. The weight decrease observed for the Co-doped IL, suggests that the [BMIM]OTf reacted with the NO (facilitated by Co\textsuperscript{2+}), resulting in the generation of a liquid with a vapor pressure, or a gradually decomposition of the IL to gaseous components.

An 20% increase in the absorption capacity was observed in the case of absorption in Fe(OTf)_3\textsuperscript{3-}/[BMIM]OTf, which together with an observed darkening of the brown-colored IL, suggests that the iron-nitrosyl complex has actually been formed.

Considering the use of the ionic liquids as NO\textsubscript{x}-removing media, a reversible absorption where the IL could be reused would be essential due to presently high cost of ILs\textsuperscript{170}. The iron nitrosyl is a rather stable complex with an equilibrium constant around 450 M\textsuperscript{-1} (in water)\textsuperscript{171}, which inhibits the desorption. The small increase in absorption achieved when utilizing iron-promoted ILs may thus be counterbalanced by the corresponding additional energy needed to desorb the NO. Therefore, for the remaining of the absorption study, only the pure ILs were examined.
Concluding remarks on preliminary sorption study

Based on the experiences from the preliminary study (section 5.2), only [BMIM]OAc and [BMIM]-OTf were selected for further examination. [BMIM]OAc was chosen based on its capability to retain large amounts of NO, and [BMIM]OTf due to its high reversibility in the absorption/desorption. Although [BMIM]OAc and [EMIM]OAc exhibited similar NO solubilities, the purity of the commercial [EMIM]OAc was only >90%, whereas [BMIM]OAc was >95% pure. Thus, to minimize the effect from any pollutants, the purest was selected for further examination.

NO sorption in [BMIM]OAc and [BMIM]OTf

The absorption of NO into [BMIM]OAc and [BMIM]OTf was measured at various temperatures at 1 bar (description of the experimental test rig and conditions is presented in section A.4.2). The absorption results presented in this section are all based on equilibrium measurements.

The NO solubilities in [BMIM]OAc and [BMIM]OTf as a function of temperature are presented in Fig. 5.4. The ILs were initially saturated at room temperature and subsequently heated stepwise under a continuous flow of 10 mol% NO/N₂ (i.e. NO partial pressure 0.1 bar). Comparing the solubilities of NO obtained here (room temperature and 0.1 bar) with e.g. CO₂ absorbed in [BMIM]BF₄ (X₇₆0 = 0.0012, 0.1 bar) reported by Anthony et al. [168], reveal that NO sorption capacities of [BMIM]OAc and [BMIM]OTf are several magnitudes higher. A more suitable comparison with literature data could not be performed due to the lack of gas sorption data on [BMIM]OAc and [BMIM]OTf. However, gas solubilities can be predicted with a quantum chemistry method known as COSMO-RS (further description in section A.4.4), which can estimate thermophysical data, such as physisorption, in a number of ILs [172–174]. This method was applied to estimate the contribution from physisorbed NO species in [BMIM]OAc and [BMIM]OTf, the results are presented in Fig. 5.5. As can be observed, only a very small fraction of the absorption (<1%) can be ascribed to the physical binding of NO in the ILs. Thus, the NO-sorption in [BMIM]OAc and [BMIM]OTf is more of a chemical nature than physical.

The overall NO absorption capacity of the ILs at room temperature was very high (cf. Fig. 5.4). The [BMIM]OTf could host about 1.5 mol NO per mol IL (X₇₆0 = 0.6), whereas [BMIM]OAc could retain about four molecules of NO per IL molecule (X₇₆0 = 0.8). By heating to 70°C is was possible to remove about 3/4 of the absorbed NO molecules. In the case of [BMIM]OTf, it was possible to completely desorb all NO by heating to about 60°C, thus suggesting a weak interaction between solute and [BMIM]OTf. In contrast, the solute-solvent interaction between NO and [BMIM]OAc is stronger and not all NO could be desorbed in these experiments by increasing the temperature. This is in agreement with results from the preliminary study where NO could easily be removed from the absorbed [BMIM]OTf, as opposed to [BMIM]OAc.

The solubility of NO in a liquid can be described by Henry’s law [160]:

\[
H_{NO}(T,P) \equiv \lim_{X_{NO} \to 0} \frac{f_{NO}}{X_{NO}} \approx \frac{p_{NO}}{X_{NO}} \quad (5.1)
\]
5.3. NO sorption in [BMIM]OAc and [BMIM]OTf

Fig. 5.4: NO absorption profile of [BMIM]OAc and [BMIM]OTf with temperature. Both experiments performed with two sorption reactors

Fig. 5.5: Predicted contribution from NO physiosorption in [BMIM]OAc and [BMIM]OTf

where $H_{NO}$ is the temperature and pressure dependent Henry coefficient, $f_{NO}^L$ is the fugacity of the vapor in the liquid, $p_{NO}$ the partial pressure of the gas, and $X_{NO}$ the mole fraction of NO dissolved in the liquid. Equation 5.1 thus describes a linear relationship between gas concentration and pressure, but only holds valid for solubilities approaching zero. The decrease in solubility observed with temperature (increasing NO partial pressure) in Fig. 5.4 are thus in agreement with the expected decrease with temperature.

The enthalpy ($\Delta H_{sol}$) and entropy ($\Delta S_{sol}$) of gas dissolution can be found by considering the effect of the temperature on the solubility. Information on the strength of interaction between the IL and solute can be found in the enthalpy, and the entropy indicates the level of order present in the NO/IL mixture. The change in enthalpy and entropy upon dissolution can be determined from the solubility data with the following thermodynamic van’t Hoff relations in equation 5.2 and 5.3.

\[
\Delta H_{sol} = R \left( \frac{\partial \ln H_{NO}}{\partial (1/T)} \right)_P = -R \left( \frac{\partial \ln X_{NO}}{\partial (1/T)} \right)_P
\]

\[
\Delta S_{sol} = -R \left( \frac{\partial \ln H_{NO}}{\partial \ln T} \right)_P = R \left( \frac{\partial \ln X_{NO}}{\partial \ln T} \right)_P
\]

The equations hold valid in the Henry's law regime at infinite dilution, where the activity coefficient of the gas is independent of the amount of dissolved gas ($X_{NO}$)$^{[167]}$. The calculated enthalpies and entropies, based on measurements at highest temperatures (i.e. lowest gas solubility), are collected in Tab. 5.2. The values are based on a linear regression of the data, suggested by the van’t Hoff relationships in equation 5.2 and 5.3. However, the values obtained from the linear regression are only based on solubilities at three temperatures, and the uncertainty may thus be somewhat larger than suggested here.

By applying the van’t Hoff equations, a negative enthalpy change upon dissolution of NO in [BMIM]OTf was found together with an increase in the order of the system. The observed decrease in entropy suggests that NO is more ordered within the IL than in the gas. Compared to the average bond energy of the NO bond of 631 kJ/mol$^{[175]}$, the energy released upon sorption
is thus only about 8% of the intramolecular energy.

The magnitude of the enthalpy observed in Tab. 5.2 are in the same range as those typically observed upon absorption of gases in ILs\cite{160,162,168}. However, the solubility data available for NO in [BMIM]OAc (cf. Fig. 5.4) are far from the conditions where the van’t Hoff equations apply (Henry regime), and the resulting enthalpy and entropy can not be calculated without an considerable margin of error. However, based on the sorption results, the NO is more strongly bound in [BMIM]OAc than in [BMIM]OTf, which should result in a more negative enthalpy of absorption (more exothermic).

<table>
<thead>
<tr>
<th></th>
<th>$\Delta H_{\text{sol}}$ [kJ/mol]</th>
<th>$\Delta S_{\text{sol}}$ [J/(mol·K)]</th>
</tr>
</thead>
<tbody>
<tr>
<td>[BMIM]OTf</td>
<td>-53 ± 4</td>
<td>-167 ± 12</td>
</tr>
<tr>
<td>[BMIM]OAc†</td>
<td>n.a.</td>
<td>n.a.</td>
</tr>
</tbody>
</table>

† Values are based on NO solubilities obtained at $X_{\text{NO}} = 0.43–0.78$, and thus far from infinite dilution where $\Delta H_{\text{sol}}$ and $\Delta S_{\text{sol}}$ can be calculated.

Based on the results obtained here and in the screening, the effect of cation and anion on the NO-solubility can be evaluated. Although similar amounts of NO can be absorbed in [BMIM]OAc and [BMIM]OTf, the reversibility is only characteristic of the triflate anion, suggesting that NO is stabilized in the solution by the acetate ion. On the other hand, in the preliminary study of the triflate-based ILs the cation was also shown to have a significant effect on the overall NO sorption and reversibility. Consequently, it can not be determined whether the anion or cation is responsible for the absorption, based on the ILs studies here. Anthony et al. suggested that the IL-anions are accountable for the gas sorption properties\cite{168}, whereas Huang et al. implied that the cations were responsible\cite{162}. As a result, no consensus seem to exist at present about which ion to attribute for the sorption capacity, and the answer may lie within the individuality of ionic liquid and/or gas.

### 5.3.1 Characterization of pure and NO-saturated ionic liquids

The dissolution of NO in the ionic liquids resulted in a number of changes in the physical properties. The viscosity of the fresh and NO-saturated ILs were evaluated and the results presented in Fig. 5.6. Saturating [BMIM]OAc with NO results in a dramatic decrease in viscosity from $517 \pm 5.2$ mPa·s to $29.2 \pm 0.2$ mPa·s, a 94% reduction. Meanwhile, the density increased from $1.053 \text{ g/cm}^3$ to $1.134 \text{ g/cm}^3$. A similar trend was observed upon saturation of the [BMIM]OTf, although the initial viscosity was somewhat lower than [BMIM]OAc. The fresh [BMIM]OTf revealed a viscosity of $84 \pm 2.5 \text{ g/cm}^3$, which decreased about 70% to $25.1 \pm 0.1 \text{ g/cm}^3$, together with a small density increase from $1.298 \text{ g/cm}^3$ to $1.315 \text{ g/cm}^3$. Although the density of the ILs increased upon saturation with NO, i.e. higher compactness, an inflation of the volume (about 20–30% increase) could be observed, as shown in Fig. 5.8, confirming the presence of a large fraction of solute in the IL.

The viscosities of the ILs presented here are several tens to hundred times higher than that of water at room temperature. Bonhote et al.\cite{163} pointed out that the high IL-viscosities are
5.3. NO sorption in [BMIM]OAc and [BMIM]OTf

largely governed by the anion-cation hydrogen bonding and van der Waals forces\(^1\). The significant decrease in viscosity observed upon gas sorption suggests that NO acts as a dispersive agent, decreasing the van der Waals interaction or by suppressing the CH\(_3\)COO\(\cdots\)H bonding, known to exist in [BMIM]OAc\(^{163}\). This would be in agreement with the significant reduction in viscosity observed in Fig. 5.6.

The thermal profile of the fresh and NO-dissolved ILs were examined with TGA. The results are presented in Fig. 5.7 (experimental description given in section A.1.8). Considering the [BMIM]OTf, a decrease in mass of 15 wt% was observed at about 200°C, implying the desorption of NO taking place. This is in agreement with the absorption results (cf. Fig 5.4) where a weight increase of 15 wt% was observed at room temperature for [BMIM]OTF. It can thus be concluded that the NO absorbed in the IL desorbs readily. Furthermore, the high degree of sorption-reversibility observed previously for [BMIM]OTf, was confirmed by the almost identical temperature of decomposition around 445°C. In other words, the molecular configuration of NO-saturated [BMIM]OTf after NO desorption is roughly identical to the fresh sample, thus

\(^1\)The attractive or repulsive interaction between closed-shell molecules, such as permanent and induced dipole forces, dipole-dipole interactions
decomposing at similar temperature.

The TGA-profile of the NO-absorbed [BMIM]OAc in Fig. 5.7 is composed of two apparent NO desorption steps around 125°C and 180°C. Around 250°C the desorption stagnates, at which point the weight had decreased 36 wt%. This agrees well with the absorption results (cf. Fig. 5.4), where the weight of the IL increased 40% upon saturation. Thus, about 4 wt% NO is still trapped in the [BMIM]OAc at 250°C. The residual NO bound in the IL is likely to be responsible for the increase in decomposition temperature from about 240°C of the commercial [BMIM]OAc to ca. 325°C. A small fraction of the NO thus remain bound in the IL, suggesting formation of stronger bonds of covalent character between the solute and solvent, which confirms the irreversibility of the NO absorption in [BMIM]OAc observed.

Finally, some additional results from the characterization of the two ILs with $^1$H-NMR and Raman spectroscopy are collected in section C.4.

## 5.4 Concluding remarks

Although some ‘dots’ still need to be connected regarding the understanding of the exact nature of the NO absorption in [BMIM]OAc and [BMIM]OTf, the system shows promising behavior. Some issues that need to be addressed involves: a) selectivity measurements of gases present in flue gas stream, e.g. N$_2$, O$_2$, CO$_2$, SO$_2$, H$_2$O, N$_2$O, NO$_2$; b) reversibility tests during temperature-swing (absorption/desorption); c) mass-transfer limitation need to be minimized through absorption design or by depositing the IL on an inert supporting material to increase the interfacial gas-IL area, d) effect of contaminants reaching the IL, such as alkali species, chlorine volatile organic compounds (VOCs).

The challenges of the mass-transfer limitation could be overcome by increasing the ‘surface area’ of the IL. The concept of the supported ionic liquid phase (SILP) system was first suggested by Riisager et al. [176] and is schematically outlined in Fig. 5.9. The method have already been applied to the system studied here, where 20 wt% of [BMIM]OAc and [BMIM]OTf have been distributed on high-surface area SiO$_2$ particles (180–355 µm), but the evaluation of the NO-absorption capacities of the SILPs will be further investigated. By applying the SILP concept, the dynamics of the absorption/desorption should be significantly enhanced due to the shorter...
diffusion distances and the increased absorption area generated\textsuperscript{177}. Furthermore, even though a large amount of IL have been distributed onto SiO$_2$, the apparent properties of the SILP particles on a macroscopic scale are hereby similar to those of ‘dry’ particles, which significantly simplifies the handling as well as increasing the surface of the IL.

The absorption capacities of NO in IL reported here reveal a promising system for NO-scrubbing of dust-free tail-end gases, not requiring the addition of reductant such as ammonia or urea. Since no additive is needed to reduce the NO in the gas stream, the complexity of the NO removal system is somewhat reduced compared to the NH$_3$-SCR process. Assuming that the selectivity toward NO sorption is sufficiently high, it opens new possibilities of NO removal. The captured NO could easily be removed by increasing the temperature (temperature swing), as observed previously, where only a small increase in temperature was required to remove the NO from [BMIM]OTf. Transport of the NO-saturated IL (i.e. by exploiting the differences in viscosity) or the SILP particles to another container or location followed by desorption of NO would render possible to concentrate the NO gas. Thereby the NO could subsequently be used in e.g. the Ostwald process for the production of HNO$_3$ in a ‘from waste to value’ concept.

The conditions at the previously discussed Electrabel’s Langerlo Power Station in Belgium\textsuperscript{149} can be used as an example for considering the implementation of an IL scrubber based on the reversible [BMIM]OTf system. Here, the removal of 1000 ppm NO under a total flow of 820,000 Nm$^3$/h would only require an effective IL volume of around 4.2 m$^3$/h to remove all the incident NO. Even though the mass-transfer issue is not considered here (which is expected to be minimized by using the SILP technology), the IL-volume is considerably less than the SCR catalyst volume of 268 m$^3$.

For comparison of solubilities, the absorption capacities of CO$_2$ in ILs are considered. CO$_2$ capture is an area which has attracted a considerable interest over the latest years. The best IL-candidates present solubilities which are in the range of 0.5–1 mol CO$_2$ per mol ionic liquid (although a high partial pressure typically is required). A challenge concerning the ILs typically used is that CO$_2$ is strongly bound in the IL (typically $\Delta H_{\text{sol}} \approx -80$ kJ/mol), requiring a significant amount of energy to release the CO$_2$\textsuperscript{116,178,179}. Thus, the absorption capacities of the two ILs presented here, [BMIM]OAc and [BMIM]OTf, represent considerably higher absorption capacities, being able to retain 1.5 and 4 NO molecules of NO per IL molecule at room temperature, respectively. Particularly the high sorption reversibility of [BMIM]OTf (and to some extent also [BMIM]OAc) represent a novel technology for the stripping of NO. Although the results presented here are slightly fragmented, and the nature of NO absorption still somewhat unknown, the first steps in the potential use of ILs for NO$_x$-sorption have been taken.
Concluding remarks and outlook

The objective of the present thesis has been to contribute to the knowledge about alternative alkali-resistant technologies for the removal of NO, with focus on zirconia-based catalysts. The attention has been on alternative methods and catalysts to eliminate or harvest the nitrogen oxide from the gas stream in e.g. power plants. During this work, a number of different zirconia-based catalysts have been evaluated based on their performance in the SCR reaction before and after deactivation with potassium. In general, the resistance toward potassium found in vanadia-based catalysts supported on high-acidity sulfated zirconia were somewhat higher than that of the traditionally used SCR catalyst, which typically is based on a formulation of $V_2O_5$-WO$_3$-TiO$_2$. The most promising candidate, based on highest potassium resistance (the vanadia-based sulfated zirconia with high crystallinity and low content of monoclinic ZrO$_2$), have successfully been upscaled to a monolith, stabilized with sepiolite clay material. The monolith exhibited sufficiently high activity to perform well under industrially relevant conditions.

A different approach for the removal of gaseous NO was undertaken by employing ionic liquids as NO-absorbing liquids. Although gas-sorption in ionic liquids is not a new phenomena, it has never been reported with NO. In the present work, the two ILs, [BMIM]OAc and [BMIM]OTf were selected, based on a preliminary absorption study. These two liquids exhibited very promising behavior in the absorption of NO, being able to retain considerable amounts of nitric oxide, about 1.5–4 molecules of NO per molecule of IL. The reversibility demonstrated by [BMIM]OTf is especially notable, where almost all absorbed NO can be desorbed by a small increase in temperature (to about 60°C). This can potentially establish a new platform technology of NO sequestration, i.e. for the synthesis of nitric acid.

Although the examples for the removal of NO presented here, will not in any way solve the major challenges we face, they are small contributions to the environmental field, and hopefully this thesis will inspire other engineers and scientists in the continuous development of technologies for the removal of nitric oxide.
Chapter 7

References


Chapter 7. References


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Chapter A

Experimental

A.1 Characterization techniques

A.1.1 X-ray diffraction

Powder diffraction data was collected on a Huber G670 Guinier camera using CuK$_\alpha_1$ radiation, $\lambda = 1.5406$ Å. The sample was placed in a thin layer on scotch tape and rotated during data collection. Data were collected in the range of 3-100° in steps of 0.005° in 2θ. Time of exposure was varied from 20–60 min. In case of data collection for Rietveld refinement, exposure was a minimum of 60 min. Rietveld refinement was done with WINPOW (Software by Kenny Ståhl), where crystallographic information files (CIF) from Fachinformationszentrum Karlsruhe are fitted against experimental XRD data, yielding information about the phase distribution and unit cell parameters.

The CIF files used for Rietveld refinement are listed by their identification code in the Inorganic Crystal Structure Database:
- t−ZrO$_2$: 93030, m−ZrO$_2$: 94886, c−ZrO$_2$: 92094,
- Mg$_8$(Si$_{12}$O$_{30}$(OH)$_4$)·(H$_2$O)$_2$ (Sepiolite 2H$_2$O)$^{[141]}$: 156200,
- Mg$_8$(Si$_{12}$O$_{30}$(OH)$_4$)·(H$_2$O)$_{12}$ (Sepiolite 12H$_2$O)$^{[141]}$: 156199,
- SiO$_2$: 174, CaCO$_3$: 16710, CaSO$_4$: 68592, (V$_{0.12}$Zr$_{0.88}$)O$_2$: 96542.

A.1.2 Scanning electron microscopy

The morphology and particle size of the catalysts were investigated by SEM using a FEI Inspect ‘S’ with tungsten filament. Images were typically obtained with acceleration voltages of 2 to 10 kV. The specimens were prepared by dry dispersing the catalyst powder on carbon tape with subsequently carbon coated to avoid charging.

A.1.3 Temperature programmed desorption

To compare the strength and the number of active sites for samples, the surface acidity of the catalysts was measured with a Micromeritics AutoChem II 2920. Approximately 100 mg of sample was initially pretreated at 200°C in helium for 30 min to remove any physisorbed species and subsequently saturated with 50 ml/min dry 1% NH$_3$/He at 100°C for 60 min. Loosely bound ammonia was hereafter removed by purging with 50 ml/min He for 60 min, before performing temperature programmed desorption (TPD) at a heating rate of 10°C/min in the range 100-1000°C in a He flow of 25 ml/min.
Appendix A. Experimental

### A.1.4 Nitrogen physisorption

Determination of the BET surface area was done by nitrogen physisorption. $N_2$ adsorption and desorption were performed at liquid nitrogen temperature on a Micromeritics ASAP 2020. Prior to measurement, the samples were degassed at 200°C under vacuum for 6 h.

### A.1.5 Mercury intrusion porosimetry

The porosity was determined by mercury intrusion porosimetry (MIP) on a CE Instruments Pascal 140/240 porosimeter on samples dried overnight at 150°C.

### A.1.6 Electrophoretic migration

The zero point charges (ZPC) of the composite materials were determined by measuring the zeta potential as a function of the solution pH. The zeta potentials were calculated by the Helmholtz-Smoluchowski equation: \[ \tilde{V} = \frac{\xi}{4\eta}, \] where $\tilde{V}$ is the electrophoretic mobility, $\varepsilon$ the permittivity, $\xi$ the zeta-potential and $\eta$ the dynamic viscosity. The electrophoretic migration rate was measured on a Zeta-Meter Inc. Instrument model 3.0+, provided with an automatic sample transfer unit to avoid sample sedimentation problems. Experiments were carried out using 30 mg of <2 µm powder samples suspended in 200 mL of mM KCl, adjusting the pH with 0.2 M KOH and HCl solutions. Each curve was recorded at least three times to ensure the reproducibility of the results.

### A.1.7 NMR spectroscopy

The ionic liquids were characterized with $^1$H-NMR, recorded on a Varian Unity Inova-500 MHz spectrometer. To avoid mixing of the samples with the solvent, pure samples were placed in closed glass capillaries. The close ampule containing the sample was placed in NMR tubes together with the solvent CDCl$_3$, which also was used as an internal standard.

### A.1.8 TGA-DSC

TGA-DSC curves were obtained from measurements on a Mettler Toledo TGA/DSC 1 SF instrument equipped with a SDTA (single differential thermal analysis) sensor. The about 25 mg of sample were placed in 70 µl alumina crucibles and heated from room temperature up to 600°C with a heating rate of 10 °C/min under a $N_2$ flow of 20 ml/min.

### A.2 Catalyst synthesis

A number of catalysts were synthesized for this study. The synthesis details are listed in the following.
A.2.2 Catalysts based on commercial zirconia

A.2.2.1 Vanadium-based catalysts
Vanadia-based catalysts were synthesized by incipient wet impregnation (IWI) of vanadyl solution on fractionated zirconia (180-300 μm), to obtain a vanadium concentration of 1.7 wt% (3.0 wt% V₂O₅) unless otherwise stated. The vanadyl solution were obtained from ammonium metavanadate (Sigma-Aldrich, >99%) and oxalic acid (Merck, p.a.), where the ammonium meta vanadate is slowly added to a heated solution (50-70°C) of oxalic acid with a small amount of water. After impregnation, the sample was allowed to settle for an hour before drying at 100°C followed by calcination at 400°C in air for at least 2 hrs.

A.2.2.2 Iron-copper mixed oxide catalysts
Synthesis of iron-copper mixed oxide catalysts were carried out by sequential IWI of the aqueous nitrate precursor [Cu(NO₃)₂, Fe(NO₃)₂] on particles of 3.4SZ_NP (180-300 μm). The total molar concentration of metal species was kept constant at 364 μmol/g (equivalent to a CuO loading of...
3.5 wt%), but the Cu-Fe ratio was varied, producing Fe-Cu metal oxide catalysts with the iron molar fraction, $X_{Fe}$, 0.00, 0.20, 0.33, 0.50, 0.67, 0.80, 1.00. After drying at 100°C, the catalysts were calcined at 450°C, 4 hrs (3.5°C/min), resulting in the catalysts presented in Fig. A.2.

**Fig. A.2:** Fe-Cu mixed oxide catalysts supported on 3.4SZ_NP. From left to right decreasing copper oxide loading, increasing iron oxide loading

### A.2.3 Catalysts mixed with sepiolite

#### A.2.3.1 Catalysts for optimization of VSZ-sepiolite ratio

The sulphated zirconia was prepared by impregnating 101.6 g of the zirconia hydroxide (Melcat) with 8.3 g $(NH_4)_2SO_4$ (Panreac, >99%) by the incipient wetness method. After overnight standing in ambient conditions, the samples were dried at 150°C for 3 hours. Calcination of the samples at 500°C for 4 hours in air was performed by placing the material directly into a pre-heated furnace, in order to facilitate the preferred formation of metastable tetragonal phases. Subsequent chemical analysis (ICP-MS) revealed a sulfate loading of 5.3 wt% (measured at wt% SO$_3$). The catalysts were prepared by wet impregnation of VOSO$_4$ onto the sulphated zirconia, to produce a slurry that was allowed to settle for an hour with stirring. Thereafter sepiolite, $\alpha$-sepiolite (Pansil 100, Tolsa S.A., 60%) and water were added in the desired amounts to obtain a homogeneous paste with an adequate viscosity for extrusion from a 20 ml syringe with a 2 mm orifice. The extrudates were slowly dried, sealed in a wet atmosphere for 48 hours then dried overnight at 150°C in air, resulting in the extrudates presented in Fig. A.3. Calcination was carried out at 500°C for 4 hours in air. Finally the extruded material was broken into 3–5 mm cylindrical pellets.

#### A.2.3.2 Sepiolite-based catalysts for optimization of sulfate content

A series of nine catalysts were synthesized by kneading 0–9 g $(NH_3)_2SO_4$ (Panreac, >99%) in water with the sulfated zirconia from the same batch prepared above, and 0.745 g VOSO$_4$·5H$_2$O. After the settling of the slurry, 1.67 g sepiolite (Pansil 100, Tolsa S.A., 60%) was added and after homogenising the slurry extruded and calcined as described above. The resulting catalysts thereby contained a total sepiolite concentration of 25 wt%, a sulfate loading of 4.1–8.2 wt%, measured at SO$_3$, and a total V$_2$O$_5$ loading of 4 wt%.

#### A.2.3.3 Monolith extrusion and activity measurements

Monolithic catalysts were made of the 2/8 formulation, with composition, drying, and calcination procedure as described above. The extruded monoliths were cylindrically shaped with 24
A.2. Catalyst synthesis

**Fig. A.3:** Sepiolite-VSZ catalyst series with increasing weight fraction of sepiolite from 0/8 to 8/8, the latter being sepiolite without VSZ. Notice that the $V_2O_5$-$SO_4^{2-}$-$ZrO_2$ (0/8) catalyst is on powder form due to the lack of sepiolite binder.

parallel channels, with the geometric dimensions: monolith length 15–30 cm, square cell size 1.6 mm, wall thickness 0.92 mm, geometric surface $9.91 \text{ cm}^2/\text{cm}^3$, cell density $14.57 \text{ cell/cm}^2$ and bulk density 1.45 g/cm$^3$.

The SCR activity of the monolith were evaluated in a reactor system, illustrated in Fig. A.4, consisting of a pyrex tube with three electrically heat zones to ensure isothermal conditions. The catalyst temperature was monitored with two thermocouples inserted above and below the catalyst bed. Furthermore, the temperature at the entrance point of $NH_3$ was constantly monitored, in order to be sure that no precipitation of $(NH_4)_2NO_2$ took place. Free space surrounding the monolith and the reactor wall were filled with inert material to prevent bypass flow. Reactant gases were regulated with mass-flow controllers providing the typical inlet concentrations: NO 700 ppm, $NH_3$ 700 ppm (except when stated otherwise), $O_2$ 3.5 vol%, $H_2O$ 1.8 vol%, balanced in $N_2$. The inlet and outlet NO and NO$_2$ concentrations were determined by chemiluminescence with a Signal NO + $O_2$ analyzer (Series 4000). Analysis of $N_2O$ and $NH_3$ was carried out by IR spectroscopy with a Signal 7000FT GFC Analyzer and with an A.D.C. Double Beam Luft Type Infrared Gas Analyzer, respectively. Stable data was acquired under steady-state conditions after approximately 90 min of stabilization for each point after changing the space velocities. Before
the analyzers, a peltier element is inserted. Only one temperature per day was measured, in order to ensure complete steady state equilibrium of NH₃ on the catalyst surface.

### A.2.4 Catalyst poisoning

Doping with potassium was done by IWI with a solution of KNO₃ (Merck, >99%) on the fractionated catalysts to obtain a potassium loading, relative to the active metal, of 0.4 mol/mol, unless otherwise noted. After settling and drying at 100°C, the sample were calcined at 400°C for 2 hrs.

### A.3 Catalyst testing

The experimental setup used in the examination of catalytic activity is schematically illustrated in Fig. A.5 and the actual test rig is shown in Fig. A.6. SCR activities were measured with a Thermo Environmental chemiluminescence analyzer, model 17C, NH₃-analyzer.

![Fig. A.5: Left: Schematic drawing of the continuous flow experimental setup for catalytic measurements. Right: Quartz reactor with loaded sample diluted in SiO₂, fixed between two layers of quartz wool. Gas direction from top to bottom. Scale 1 cm](image)

About 10 mg of catalyst (if catalyst was synthesized in powder form, it was pressed at 2 tons, Ø = 13 mm before fractionation) fractionated to 180-300 µm, were fixed in a quartz plug flow reactor with a total ambient flow of 300 ml/min containing 1000 ppm NO, 1100 ppm NH₃, 3.5% O₂, 2.3% H₂O in He. The catalysts were diluted with about 125 mg quartz (Merck, 99.95%, BET surface area <0.1 m²/g) to ensure a cylindrical shaped catalyst bed, see Fig. A.5, right.

The catalyst testing was performed with a specially designed feed-back controlled Labview program, monitoring the total flow, NO-, NO₂-, NH₃- and total NOₓ-signals, and controlling mass flow controllers and temperature. Catalyst temperature was measured directly above the catalyst bed (thermocouple not shown in Fig. A.5). A typical program was initiated by activation in the SCR gas at 200°C for minimum 60 min, followed by cooling to room temperature. From ambient conditions the temperature was stepwise increased from 25 to 500°C in steps of 20°C. Steady-state conversion was obtained at each temperature (ensured with PID feed-back controller embedded in Labview) and data collected for 5 min (frequency 10 sec) before sequential temperature increase. After the maximum temperature, the catalyst was cooled stepwise and data collected every 50°C.
A.3. Catalyst testing

Fig. A.6: Setup for testing the catalytic SCR activity at various temperatures. NH$_3$-NO$_x$-analyzer located below table along with gas diluting unit

### A.3.1 Rate expression

A convenient method of representing the catalysts activity is by its rate constant, $r$, which is first order with respect to NO$^{[42,89]}$ (and references therein):

$$r = -\frac{1}{W} \frac{d[\text{NO}]}{dt} = k[\text{NO}]^1[\text{NH}_3]^0$$  \hspace{1cm} (A.1)

where $W$ is the amount of catalyst, and $k$ the first-order rate constant. Expression A.1 can be rearranged and solved:

$$\int_{[\text{NO}]}^{[\text{NO}]} \frac{1}{[\text{NO}]} d[\text{NO}] = -kW \int_{t=0}^{t} dt \Leftrightarrow$$

$$\ln \frac{[\text{NO}]}{[\text{NO}]} = -kWt \Leftrightarrow$$

$$k = -\frac{1}{Wt} \ln \frac{[\text{NO}]}{[\text{NO}]}$$  \hspace{1cm} (A.2)

Introducing the fractional conversion $X$, of NO, which is expressed as:

$$X = \frac{[\text{NO}]}{[\text{NO}]} - \frac{[\text{NO}]}{[\text{NO}]} \Rightarrow \frac{[\text{NO}]}{[\text{NO}]} = 1 - X$$ \hspace{1cm} (A.3)

Upon substitution of $\frac{[\text{NO}]}{[\text{NO}]}$ from equation A.3 into equation A.2, we obtain the following:

$$k = -\frac{1}{Wt} \ln (1-X)$$ \hspace{1cm} (A.4)

By assuming plug flow in the reactor, i.e. time is proportional to a volume flux through the reactor, we can introduce the inlet NO molar concentration, $[\text{NO}]_0$ (mol/l), which is unit-counterbalanced by the molar inflow rate of NO, $F_0$ (mol/s). This is collected in the expression of the first order rate constant, $k$:

$$k = -\frac{F_0}{[\text{NO}]} W \ln (1-X)$$ \hspace{1cm} (A.5)

Thereby we obtain a rate expression wherein the catalyst weight is incorporated, in contrast to the conversion of NO, which is often used in the literature, but difficult to compare at different space velocities.
**Appendix A. Experimental**

**A.4 NO gas absorption in ionic liquids**

**A.4.1 Preliminary screening**

Preliminary gas absorption was carried out in a ca. 5 ml test tube with 1 ml ionic liquid (IL) at room temperature (22–23°C). The 10% NO/N$_2$ were introduced into the IL through a glass pasteur pipette retrofitted with a glass frit to reduce the bubble size and increase the gas-liquid contact area (interfacial area). The flow was maintained at around 5 ml/min to contain the bubbles formed within the test tube. The acetate-based ILs were exposed to the NO gas for about 100 h, whereas the triflate-based only were subjected to the gas for about 43 h. The absorption of NO was monitored in real time by logging the weight increase with a PC interfaced with the microbalance (accuracy 0.1 mg). However, to avoid balance drifting, the absorption results are supplied by measuring the total weight of the test tube with IL and pipette before and after NO absorption. Any increase in weight of the sample after NO exposure were attributed solely to the absorption of NO. The weight increase are in all cases given as the fraction molar increase, defined in equation A.6.

\[
X_{\text{IL}}^{\text{NO}} = \frac{n_{\text{NO}}}{n_{\text{NO}} + n_{\text{IL}}} \quad (A.6)
\]

Subsequent to the absorption procedure, the NO-absorbed ionic liquids were subjected to vacuum in order to estimate the reversibility of the bound NO. The samples were desorbed at pressures below 10 mbar for minimum 24 h at room temperature.

**A.4.2 Equilibrium gas absorption-desorption**

The absorption of 10% NO/N$_2$ were performed at 1 bar at room temperature, by passing the gas stream (ca. 10 ml/min under ambient conditions) via a glass frit (diameter 6 mm) through 1000 µl of ionic liquid. The ionic liquid were loaded in individual cylindric glass tubes (length 105 mm, inner diameter 10 mm) with inlet and outlet valves. The exit of the glass frit were located just above the bottom of the cylinder so the gas bubbles would supply ‘stirring’ of the solution during absorption. The absorption experiments were performed with three glass reactors with the same ionic liquid, for reproducibility. All reactors were placed in a glass tube rack in an oven where temperature could be varied to an certainty of ± 0.5°C, as outlined in Fig. A.7. The ionic liquid were saturated at room temperature before the temperature were increased in a step wise manner, where the solubility of the NO were determined at each step. The gas flow to each reactor were supplied by Bronkhorst flow controllers. The temperature were monitored continuously via Labview with a thermocouple located close to the ionic liquids in the oven. The mass increase of the IL phase were determined by weighing the reactors with IL on a microbalance (accuracy 0.1 mg) on a regular basis. Before removal from the oven for weighing, the gas inlet and outlet valves were closed to avoid gas desorption due to changes in pressure. To ensure that the thermodynamic equilibrium were obtained, the weighing procedure were repeated until the weight were constant within 6 h between consecutive measurements.
A.4. NO gas absorption in ionic liquids

Fig. A.7: Setup for equilibrium absorption experiments. The ionic liquid are placed in glass reactors and placed in oven

A.4.3 Chemicals
The ILs used in the preliminary study were all commercial grade and were used without any further purification. The ILs were: 1-ethyl-3-methylimidazolium tetrachloroaluminat (>95%, Fluka/BASF), 1-ethyl-3-methylimidazolium thiocyanate (>95%, Fluka/BASF), 1-ethyl-3-methylimidazolium acetate (>90%, Fluka/BASF), 1-ethyl-3-methylimidazolium triflate (99%, Io-li-tec), 1-butyl-3-methylimidazolium tetrachloroaluminat (>95%, Aldrich/BASF), 1-butyl-3-methylimidazolium thiocyanate (>95%, Fluka/BASF), 1-butyl-3-methylimidazolium acetate (>95%, Aldrich/BASF), 1-butyl-3-methylimidazolium triflate (>98%, Solchemar), 1-octyl-3-methylimidazolium triflate (99%, Io-li-tec).

A.4.4 NO gas solubility predictions with COSMO-RS
Theoretical calculations of NO gas solubility in ionic liquids were performed using the software COSMOthermX Version C21_0110. The program predicts thermophysical data for mixtures and is based on a method called COSMO-RS (Conductor-like Screening Model for Real Solvents) which combines quantum chemical calculations with statistical thermodynamics[174]. The COSMO files used for the calculations were obtained from the TZVP database in the program. These files contain results of quantum chemical calculations, including the optimized geometries of the molecules, performed with the DFT method using TZVP basis sets.
### Tab. B.1: CRC standard thermodynamic properties of gases at 298.15K and 1 bar. Values from [10]

<table>
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<tr>
<th>Molecular formula</th>
<th>Name</th>
<th>$\Delta_f H_{298K}^o$ [kJ/mol]</th>
<th>$\Delta_f G_{298K}^o$ [kJ/mol]</th>
<th>$\Delta_f S_{298K}$ [J/mol K]</th>
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<tr>
<td>$O_2$</td>
<td>Oxygen</td>
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<td>$O_3$</td>
<td>Ozone</td>
<td>142.7</td>
<td>163.2</td>
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<td>$N_2$</td>
<td>Nitrogen</td>
<td>0</td>
<td>191.6</td>
<td></td>
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<td>$N_2O$</td>
<td>Nitrous oxide</td>
<td>81.6</td>
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<td>NO</td>
<td>Nitric oxide</td>
<td>91.3</td>
<td>87.6</td>
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<td>$N_2O_3$</td>
<td>Nitrogen trioxide</td>
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<td>Nitrogen dioxide</td>
<td>33.2</td>
<td>51.3</td>
<td>240.1</td>
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<td>$N_2O_4$</td>
<td>Nitrogen tetroxide</td>
<td>11.1</td>
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<td>Nitrogen pentoxide</td>
<td>13.3</td>
<td>117.1</td>
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<td>Ammonia</td>
<td>-45.9</td>
<td>-16.4</td>
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<td>CO$_2$</td>
<td>Carbon dioxide</td>
<td>-393.5</td>
<td>-394.4</td>
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<td>Methane</td>
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<td>-50.5</td>
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<td>CO</td>
<td>Carbon monoxide</td>
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<td>H$_2$O</td>
<td>Water</td>
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</table>
### Appendix B. Standard thermodynamic properties

#### Tab. B.2: Thermodynamic calculations based on data reported in Tab. B.1

<table>
<thead>
<tr>
<th>Reaction</th>
<th>( \Delta H_{298K}^0 ) [kJ/mol]</th>
<th>( \Delta S_{298K}^0 ) [kJ/mol K]</th>
<th>( \Delta G_{298K}^0 ) [kJ/mol]</th>
<th>Eq. const., ( K_{298K} ) †</th>
<th>( T_{\Delta G^o=0} ) [K]</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>NO(_x) removal</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>4NO + 4NH(_3) + O(_2)</td>
<td>4 N(_2) + 6H(_2)O</td>
<td>-1632</td>
<td>-340</td>
<td>-1531</td>
<td>1.6E+268 bar(^1)</td>
</tr>
<tr>
<td>6NO + 4NH(_3)</td>
<td>5 N(_2) + 6H(_2)O</td>
<td>-1815</td>
<td>-365</td>
<td>-1706</td>
<td>7.9E+298 bar(^1)</td>
</tr>
<tr>
<td>NO</td>
<td>( \frac{1}{4} N(_2) + \frac{1}{4} O(_2))</td>
<td>-91</td>
<td>-12</td>
<td>-88</td>
<td>2.2E+15 bar(^0)</td>
</tr>
<tr>
<td>2NO + 2NO(_2) + 4NH(_3)</td>
<td>4 N(_2) + 6H(_2)O</td>
<td>-1516</td>
<td>-194</td>
<td>-1458</td>
<td>3.2E+255 bar(^{-2})</td>
</tr>
<tr>
<td><strong>NO removal with different reductants</strong></td>
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<td></td>
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<td></td>
</tr>
<tr>
<td>NO + CO</td>
<td>( \frac{1}{2} N(_2) + CO(_2))</td>
<td>-374</td>
<td>-99</td>
<td>-345</td>
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<tr>
<td>4NO + 4NH(_3) + O(_2)</td>
<td>4 N(_2) + 6H(_2)O</td>
<td>-1632</td>
<td>-340</td>
<td>-1531</td>
<td>1.6E+268 bar(^1)</td>
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<tr>
<td>NO + CH(_4) + ( \frac{1}{2} ) O(_2)</td>
<td>( \frac{1}{2} N(_2) + CO(_2) + 2H(_2)O)</td>
<td>-894</td>
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<td>-847</td>
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<td>NO + H(_2)</td>
<td>( \frac{1}{2} N(_2) + H(_2)O)</td>
<td>-333</td>
<td>-127</td>
<td>-295</td>
<td>5.3E+51 bar(^{-1/2})</td>
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<td><strong>NO(_x) formation</strong></td>
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<td>O(_2) + N(_2)</td>
<td>2 NO</td>
<td>183</td>
<td>25</td>
<td>175</td>
<td>2.0E-31 bar(^0)</td>
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<td>2NO + O(_2)</td>
<td>2 N(_2)</td>
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<td>-99</td>
<td>-71</td>
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<td>3NO</td>
<td>N(_2)O + NO(_2)</td>
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<td>-172</td>
<td>-108</td>
<td>7.5E+18 bar(^{-1})</td>
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<tr>
<td>4NH(_3) + 4NO + 3O(_2)</td>
<td>4 N(_2)O + 6H(_2)O</td>
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<td>-637</td>
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<td>NO + O(_3)</td>
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<td>-4</td>
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<td>2N(_2)O</td>
<td>2 NO + N(_2)</td>
<td>19</td>
<td>365</td>
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<tr>
<td>N(_2)O(_3)</td>
<td>NO + NO(_2)</td>
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<td>-3</td>
<td>3.0E+00 bar(^1)</td>
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<td><strong>NO(_x) formation from air</strong></td>
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<tr>
<td>N(_2) + ( \frac{1}{2} ) O(_2)</td>
<td>N(_2)O</td>
<td>82</td>
<td>-74</td>
<td>104</td>
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<tr>
<td>( \frac{3}{2} N(_2) + \frac{3}{2} O(_2))</td>
<td>NO</td>
<td>91</td>
<td>12</td>
<td>88</td>
<td>4.5E-16 bar(^0)</td>
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<tr>
<td>N(_2) + ( \frac{3}{2} ) O(_2)</td>
<td>N(_2)O(_3)</td>
<td>87</td>
<td>-185</td>
<td>142</td>
<td>1.5E-25 bar(^{-3/2})</td>
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<tr>
<td>( \frac{3}{2} N(_2) + O(_2))</td>
<td>NO(_2)</td>
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<td>-61</td>
<td>51</td>
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<td>N(_2)O(_4)</td>
<td>11</td>
<td>-298</td>
<td>100</td>
<td>3.2E-18 bar(^{-2})</td>
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<td><strong>NH(_3) oxidation</strong></td>
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<td>4NH(_3) + 3O(_2)</td>
<td>2 N(_2) + 6H(_2)O</td>
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<td>-973</td>
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<td>-897</td>
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<td>1.3E+25 bar(^-1)</td>
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<td>5 N(_2)O + 3H(_2)O</td>
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<td>-772</td>
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<tr>
<td>4NH(_3) + 4NO + 3O(_2)</td>
<td>4 N(_2)O + 6H(_2)O</td>
<td>-1306</td>
<td>-637</td>
<td>-1116</td>
<td>3.3E+195 bar(^-1)</td>
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<td><strong>Various NO(_2) reactions</strong></td>
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<td>6NO(_2) + 8NH(_3)</td>
<td>7 N(_2) + 12H(_2)O</td>
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<td>-216</td>
<td>-2669</td>
<td>( \infty ) bar(^5)</td>
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<tr>
<td>2NO(_2) + 4NH(_3) + O(_2)</td>
<td>3 N(_2) + 6H(_2)O</td>
<td>-1334</td>
<td>-169</td>
<td>-1283</td>
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<td>4 N(_2)O + 6H(_2)O</td>
<td>-1074</td>
<td>-344</td>
<td>-971</td>
<td>1.3E+170 bar(^{-1})</td>
</tr>
</tbody>
</table>

1. Calculated with \( K = \exp \left( \frac{\Delta G_{298K}^o}{RT} \right) \)

2. Temperature where spontaneous reaction takes place under the coarse assumption that \( \Delta H^0 \) and \( \Delta S^0 \) are temperature independent. \( \Delta G^0 = 0 \), thus \( \Delta G^0 = \Delta H^0 - T \Delta S^0 \Rightarrow 0 = \Delta H^0 - T \Delta S^0 \Rightarrow T = \frac{\Delta G^0}{\Delta S^0} \)
Chapter C

Supplementary figures

C.1 Optimization of sepiolite-VSZ ratio

Fig. C.1 show the paramagnetic species, such as V\textsuperscript{4+} and Fe\textsuperscript{3+}, that are present in the composite catalyst. With increasing sepiolite content, the strong signal from Fe(III) develops around 1500 Gauss, confirming the presence of impurities in the sepiolite. Simultaneous, the signal from vanadyl species is gradually displaced. At highest sepiolite loadings, the presence of Mn\textsuperscript{2+} species in the sepiolite is also confirmed.

Fig. C.1: Room temperature electron resonance spectroscopy (EPR) profiles of the sepiolite/V\textsubscript{2}O\textsubscript{5}−SO\textsubscript{4}\textsuperscript{2−}−ZrO\textsubscript{2} composite catalysts with increasing weight fraction of sepiolite. **Left:** EPR signal due to the presence of Fe\textsuperscript{3+}-species in sepiolite. **Right:** Signal originating from V\textsuperscript{4+}. Dotted red curve: simulated EPR signal of V\textsuperscript{4+}. At highest sepiolite concentrations, the samples display signal from a Mn\textsuperscript{2+}-impurity.
The porosity of the manually extruded composite catalysts can be determined based on Fig. C.2. The $V_2O_5$–$SO_4^{2-}$–$ZrO_2$ sample (0/8) display large interparticular pores of around 1 µm due to its powder configuration. However, the large macro pores of the mixed catalyst are gradually reduced in diameter with increasing sepiolite content. Thus, the porosity can be designed, based on the content of sepiolite.

**C.2 SEM-EDS of 2/8 catalyst after KCl hydrothermal treatment**

The distribution of KCl on the catalysts were performed subsequent to hydrothermal treatment to evaluate the distribution of potassium. According to the results obtained with EDS, the potassium 'density' seem to be almost homogeneously distributed over the particle surface. However, whether potassium is actually reaching the active vanadia sites or blocked by sepiolite (that is partly covering the surface) can not be determined here.
C.3 Optimization of sulfate content

With increasing sulfate content, the catalytic activity proceeds through a maximum sulfate concentration around 6.3 wt% (as SO$_3$), at where the sepiolite probably is neutralized. However, the overall impact on the SCR activity with increased sulfate loading is moderate. This stresses the importance of sulfating the Zr(OH)$_2$ for obtaining high acidity, in contrast to sulfation of ZrO$_2$ [73], as were done here.

![Graph showing SCR activity at 360°C of the extruded 2/8 catalyst pellets with additional ammonium sulfate added](image)

**Fig. C.4:** SCR activity at 360°C of the extruded 2/8 catalyst pellets with additional ammonium sulfate added

C.4 Characterization of ionic liquids for NO absorption

The pure and NO-saturated [BMIM]OAc and [BMIM]OTf were evaluated with Raman spectroscopy. NO exhibit a stretching frequency around 1870 cm$^{-1}$ [181], which could not be observed in any of the obtained spectra (cf. Fig C.5 and C.6). However, in case of [BMIM]OAc a peak developed after NO saturation in the OH stretching region at 2965 cm$^{-1}$, implying the presence of water in the IL. The water ‘contamination’ could originate from the gas mixture.

![Graph showing Raman spectroscopy of the fresh and NO saturated [BMIM]OAc. NO from the saturated [BMIM]OAc was attempted removed by reducing the pressure at room temperature](image)

**Fig. C.5:** Raman spectroscopy of the fresh and NO saturated [BMIM]OAc. NO from the saturated [BMIM]OAc was attempted removed by reducing the pressure at room temperature

In case of the [BMIM]OTf, only one band suggest that the IL has been altered, which confirms that NO is only weakly bound to the IL. The band at 1312 cm$^{-1}$ disappears subsequent to
saturation. Further investigation is being undertaken to identify the remaining bands.

![Raman spectroscopy](image1.png)

**Fig. C.6:** Raman spectroscopy of the fresh and NO saturated [BMIM]OTf

$^1$H-NMR were performed on the pure and NO-saturated ILs, the chemical shifts of the protons are presented in Fig. C.7. The protons of the [BMIM]OTf are only involved in minor low-field shifts of around 0.2 ppm. Apparently, only the alkyl chains are affected by the absorption of NO, suggesting that the imidazolium-part is not or only weakly involved in the ‘coordination’ of NO. The overall small proton shifts observed here are in agreement with the weak interaction previously between NO and [BMIM]OTf, e.g. in the reversible absorption experiments and Raman spectroscopy.

![H-NMR chemical shifts](image2.png)

**Fig. C.7:** $^1$H-NMR chemical shifts before and after NO-absorption. Experimental description in section A.1.7

In contrast, saturating the [BMIM]OAc with NO results in somewhat larger proton shifts of the both the acetate anion and the cation, implying that both ions are involved in the strong
C.4. Characterization of ionic liquids for NO absorption

... sorption of nitric oxide. Moreno et al.\textsuperscript{[182]} performed a NMR study on the effect of water on the \([\text{BMIM}]\text{BF}_4\). Even at high water loadings (mole fraction 0.36) they report of almost unaltered chemical shifts. Only in the case of the singlet imidazolium proton, a small shift was observed of around 0.2 ppm. Thus, even if small amounts of water is present in the IL (as implied by Raman spectroscopy), it would not induce such a significant shift as observed in the case of NO-saturated \([\text{BMIM}]\text{OAc}\). Hence, the dramatic impact on the protons can only be due to the presence of NO in the IL.

After saturation in NO, a new low-field peak was observed in the proton NMR spectra for both ILs. In the case of the NO-saturated \([\text{BMIM}]\text{OAC}\), the singlet was located at \(\delta = 14.489\) ppm, and for the \([\text{BMIM}]\text{OTf}\) at \(\delta = 11.951\) ppm. The two singlets could be attributed to a weakly bound proton, maybe involving a NO molecule ("HNO").
Chapter D

List of publications

List of included publications

**Paper 1** • *Structural characterization of 1,1,3,3-tetramethylguanidinium dichromate*

**Paper 2** • *Pore design of pelletised VO₄/ZrO₂-SO₄/Sepiolite composite catalysts*

**Paper 3** • *SCR activity of conformed CuOₓ/ZrO₂-SO₄ catalysts*

**Paper 4** • *Multidisciplinary Determination of the Phase Distribution in the VOₓ-ZrO₂-SO₄-Sepiolite Catalytic System (Distribución de las Fases en el Sistema VOₓ-ZrO₂-SO₄-Sepiolita Determinada Por Varios Métodos)*
Søren B. Rasmussen, Johannes Due-Hansen, Mirza Villaroel, F. Javier Gil-Llambías, Rasmus Fehrmann, Pedro Ávila, *XXII CICAT*, **2010** (accepted for oral + paper)

**Paper 5** • *Impact of support and potassium-poisoning on the VₓO₅-WO₃/ZrO₂ catalyst performance in ammonia oxidation*
Johannes Due-Hansen, Arkady L. Kustov, Claus Hviid Christensen, Rasmus Fehrmann, *Catalysis Communications*, 10, **2009**, 803–806
Appendix D. List of publications

Miscellaneous publications

• **Vanadia-based SCR catalysts supported on tungstated and sulfated zirconia: Influence of doping with potassium**
  Johannes Due-Hansen, Soghomon Boghosian, Arkady Kustov, Peter Fristrup, George Tsilomelelis, Kenny Ståhl, Claus Hviid Christensen, Rasmus Fehrmann *Journal of Catalysis*, 251, **2007**, 459–473

• **Fe-BEA Zeolite Catalysts for NH$_3$-SCR of NO$_X$**
  Anne Mette Frey, Selcuk Mert, Johannes Due-Hansen, Rasmus Fehrmann, Claus Hviid Christensen, *Catalysis Letters*, 130, **2009**, 1–8

• **On the redox behaviour of the V$_2$O$_5$-WO$_3$ / TiO$_2$ SCR at moderate temperatures: Influence of Hydrogen-Oxygen Treatment as observed by Operando & in-situ Spectroscopy at 250°C**
  Johannes Due-Hansen, Søren Birk Rasmussen, Ewelina Mikolajska, Miguel Angel Bañares, Pedor Ávila, Rasmus Fehrmann, (manuscript)

• **Biobrændsel - et varmt alternativ**
  Anne Mette Frey, Asbjørn Klerke, Johannes Due-Hansen, Claus Hviid Christensen, Nye Kemiske Horisonter, DTU, Litotryk Svendborg A/S, **2007**

• **Alkali Resistant Catalyst**
  Søren Birk Rasmussen, Arkady Kustov, Rasmus Fehrmann, Johannes Due-Hansen, Patent (WO/2008/037255), September **2007**
Structural characterization of 1,1,3,3-tetramethylguanidinium dichromate

Johannes Due-Hansen, Kenny Ståhl, Soghomon Boghosian, Anders Riisager, Rasmus Fehrmann

_Dalton Transactions_, 2010 (submitted)
Structural characterization and catalytic properties of 1,1,3,3-tetramethylguanidinium dichromate

Johannes Due-Hansen*, a, Kenny Ståhl a, Soghomon Boghosian b, Anders Riisager a, Rasmus Fehrmann*, a

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Abstract
The structure of 1,1,3,3-tetramethylguanidinium dichromate was determined from powder X-ray diffraction data. The compound crystallizes in the monoclinic system (space group P2_1/n) with a = 10.79679 (19) Å, b = 11.7580 (2) Å, c = 8.15076 (14) Å, β = 109.5247 (6)°. The structure consists of dichromate anions (Cr_2O_7^{2-}) stabilized by tetramethylguanidinium cations (NH_2C(N(CH_3)_2)_2^+ or TMGH^+). Phase transitions of [TMGH]_2Cr_2O_7 were determined by differential scanning calorimetry, thermal gravimetric analysis and in situ Raman spectroscopy, where the decomposition of the matrix into CrO_x was found at 171–172°C. Further heat treatment to above 400°C resulted in formation of the thermodynamically stable Cr_2O_3, most likely with the TMGH^+ cation as reductant. The catalytic activity of [TMGH]_2Cr_2O_7 supported on TiO_2 anatase in the selective catalytic reduction (SCR) of nitrogen oxide was also investigated, however only moderate activity was observed in the temperature range 100–400°C compared to the activity of e.g. vanadia supported on titania.

1. Introduction
Selective catalytic reduction (SCR) plays an important role in the removal of nitrogen oxides from flue gases generated by fuel combustion in stationary sources. The traditional, and widely commercialized, catalyst for the SCR reaction is composed of V_2O_5 and WO_3 or MoO_3 supported on TiO_2 which is operated in the temperature window of 300–400°C [1]. Due to the relatively high operating temperatures the catalyst unit is often placed upstream of cleaning units (e.g. desulfurizer, electrostatic particle remover) to avoid costly reheating of the flue gas. This environment, however, promotes deactivation of the catalyst because of high concentrations of dust. Plac-
ing the SCR unit downstream of the desulfurization and dust removal units would thus solve the deactivation, but also lower the temperature of the flue gas and consequently the activity of the catalyst, since the low-temperature activity of the traditional SCR catalysts is negligible. Accordingly, efforts have been made to develop catalysts operating in the range 80-280°C [2–8], which would render placement of the SCR unit possible in the tail end after the desulfurizer and particulate control installations.

Recent studies have shown that chromium and/or manganese oxides, such as MnOx [9], MnOx/TiO2 [10], Fe-Mn-TiO2 [8], MnOx–CeO2 [11], exhibit good low-temperature SCR potential, but their N2-selectivity and sulfur resistance need to be improved. Sreekanth et al. [12] investigated the NO conversion over a range of mono and bimetallic transitions metals, namely Cr, Mn, Cu, Mn-Cu and Mn-Cr, and found that the monometallic catalysts, Cr, Mn and Cu yielded better SCR performance than the mixed oxides, especially at low-temperature (100°C). Schneider et al. [13] reported higher selectivity to N2 (>90%) for a range of titania-supported chromia catalysts (i.e. CrO2, Cr2O3, CrOOH), but the catalytic performance, such as N2O selectivity, was very sensitive to the phase of the chromium oxide.

In this paper we have examined the low-temperature SCR activity of a newly characterized compound of chromium, tetramethylguanidinium dichromate, which was first reported by Kim et al. [14] for oxidation of alcohols. The physiochemical properties of 1,1,3,3-tetramethylguanidine are well documented, e.g. by Anderson and Hammer [15], but the 1,1,3,3-tetramethylguanidinium dichromate salt has not been structurally characterized. The crystal structure of the resembling guanidinium dichromate has been reported by Wajsman et al. [16]. However, addition of four methyl groups to the arrangement changes the structure and there are therefore some differences in the matrix.

2. Experimental section

2.1 Synthesis

[TMGH]2Cr2O7 was synthesized by mixing 1,1,3,3-tetramethylguanidine, TMG (Aldrich, 99%) with absolute ethanol in the molar ratio 1:5 (slightly exothermic). Equimolar amounts of ammonium chromate (Fluka, 99%) were added to the solution under stirring, resulting in evolution of ammonia gas due to the reaction (1)

\[2\text{TMG} + 2(\text{NH}_4)\text{CrO}_4 \rightarrow \text{[TMGH]}_2\text{Cr}_2\text{O}_7 + 4\text{NH}_3(\text{g}) + \text{H}_2\text{O} \quad (1)\]

whereby the solution changed color from yellow to orange within about 30 minutes indicative of transformation of the chromate to dichromate. To ensure complete reaction the slurry was stirred overnight before the ethanol and water was subsequently removed under reduced pressure with a rotary evaporator. Finally the residue was dried in a vacuum oven at 60°C.

The preparation of 20 wt% [TMGH]2Cr2O7 supported on titania (8.2 mol% Cr/TiO2) was performed by initial mixing of TMG, (NH4)2CrO4 and absolute ethanol in a molar ratio of 1:1:35. To the resulting slurry, TiO2 (Aldrich, nanopowder, 99.7% anatase) was then added and the mixture
stirred overnight before solvent and water were removed under reduced pressure in a rotary evaporator.

2.2 Characterization

FTIR spectra were obtained at room temperature in KBr on a Perkin Elmer 1710 FTIR spectrometer. All samples, in the KBr tablet, were dried overnight at 100°C before recording of the spectra. 32 consecutive scans in the range 4000–400 cm$^{-1}$ were performed at 2 cm$^{-1}$ resolution.

Powder diffraction data was collected on a Huber G670 Guinier camera using CuK$_{α1}$ radiation. The sample were placed in a thin layer on scotch tape and rotated during data collection. Data were collected for 4h in the range 3-100$^\circ$ in steps of 0.005$^\circ$ in 2$\theta$. The pattern were indexed by the program ITO [17] and the crystal structure solved by EXPO [18,19]. The structure was refined by WINPOW (a local variation of LHMP) [20]. Hydrogen positions were added in calculated positions and the methyl group hydrogens were given large thermal parameters to mimic rotational disorder. Moreover, as the dichromate part was found disordered the bridging O4 was split in two positions (the average position would give a 180$^\circ$ Cr-O4-Cr angle). The individual disordered positions of O1, O2 and O3 could not be resolved, but showed increased thermal parameters. The methyl groups were found rotated out of the central plane to minimize the H-H interactions in agreement with related tetramethylguanidinium structures [21].

Thermal stability of the samples was determined by thermogravimetry (simultaneous TGA-DSC, Netzsch, model STA 409) with a heating rate of 20°C/min under a nitrogen atmosphere. The decomposition gases were further analyzed with mass spectrometry.

For recording of the Raman spectra of [TMGH]$_2$Cr$_2$O$_7$/TiO$_2$, approximately 100 mg of the sample was pressed into a wafer and mounted on the sample holder of an in situ Raman CSTR-like optical cell. The spectra were recorded at various temperatures in the range 25–400°C under flowing O$_2$ using the 488.0 nm line of a Spectra Physics Stabilite 2017 Ar$^+$ laser operated at a power of 40 mW, collected on the sample by a slightly defocused cylindrical lens in order to reduce sample irradiance. The scattered light was collected at 90$^\circ$, analyzed with a 0.85 Spex 1403 double monochromator and detected by a -20°C cooled RCA PMT equipped with EG&G photon counting electronics [22].

SCR activities were measured with a Thermo Environmental chemiluminescence NO$_x$ and NH$_3$-analyzer, model 17C. About 40 mg of catalyst (pressed at 2 tons, Ø = 13mm, fractionated to 180-300 µm) were fixed in a quarts plug flow reactor with a total ambient flow of 300 ml/min containing 1000 ppm NO, 1100 ppm NH$_3$, 3.5% O$_2$, 2.3% H$_2$O in He. The gas hourly space velocity (GHSV) was 70,000 h$^{-1}$. The catalysts were diluted with 125 mg quartz 180-300 µm particles (Merck, 99.95%) to ensure a cylindrical shaped catalyst bed with plug-flow conditions. The catalytic activity was reported as the first-order rate constant $k$, in the first order removal of NO using the definition reported previously [22].
3. Results and Discussion

3.1 Characterization

The synthesized $\text{[TMGH]}_2\text{Cr}_2\text{O}_7$ was characterized by various techniques, such as XRD, DSC, TGA and FTIR, to support the structure determination.

![FTIR spectra](image1)

Figure 1. FTIR spectra of $\text{[TMGH]}_2\text{Cr}_2\text{O}_7$ and the synthesis-precursors

The infrared spectral data of $\text{[TMGH]}_2\text{Cr}_2\text{O}_7$ in Figure 1 showed that the chromate bands of the precursor $(\text{NH}_4)_2\text{CrO}_4$ at 951, 936, 883 and 769 cm$^{-1}$ were replaced with characteristic bands for dichromate, i.e. 945, 925, 880, 771 cm$^{-1}$ [23] in the product, as expected. The same bands (except the band at 771 cm$^{-1}$) were also determined by Kim et al. [14]. The elimination of the $>\text{N-H}$ stretch vibration around 3330 cm$^{-1}$ of TMG and the broad N-H bands around 3140 cm$^{-1}$ of NH$_4^+$ in ammonium chromate further confirmed full protonation of TMG occurred during synthesis of $\text{[TMGH]}_2\text{Cr}_2\text{O}_7$.

![DSC](image2)

Figure 2. DSC of $\text{[TMGH]}_2\text{Cr}_2\text{O}_7$. Heating rate 5°C/min in Ar

Besides the bands from dichromate, the $\text{[TMGH]}_2\text{Cr}_2\text{O}_7$ spectra also contained an additional imine (C=N-R$_2$) band at 1657 cm$^{-1}$ [24]. This suggests the electrons to be delocalized between all three nitrogen atoms and the sp$^2$ carbon instead of being confined to one nitrogen atom as
in pure TMG.

$[\text{TMGH}]_2 \text{Cr}_2 \text{O}_7$ was investigated by DSC analysis, as presented in Figure 2. When heating from $-60^\circ$C to $172^\circ$C a continuous endothermic heat flow was found, suggesting a slow phase transformation or crystal reorganization. At $172^\circ$C a distinctive energy uptake was observed for the crystal due to decomposition of the salt into organic residues and $\text{CrO}_x$. The irreversible decomposition of the $[\text{TMGH}]_2 \text{Cr}_2 \text{O}_7$ was confirmed by the DSC profile upon cooling from $500^\circ$C to $-60^\circ$C, since no phase transitions occurred here.

![Figure 3. XRD diffractogram of $[\text{TMGH}]_2 \text{Cr}_2 \text{O}_7$ and the difference pattern from the Rietveld refinement. Markers indicate K\(\alpha_1\) Bragg positions](image)

The results of the Rietveld refinement of the powder diffraction data of $[\text{TMGH}]_2 \text{Cr}_2 \text{O}_7$ are displayed in Figure 3 ($2\theta = 10-50^\circ$). Here a good agreement between the experimental and fitted data was found for the monoclinic cell. The cell dimensions found through refinement are collected in Table 1.

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The determined crystal structure of $[\text{TMGH}]_2 \text{Cr}_2 \text{O}_7$ is presented in Figure 4 with both of the two possible O4 positions depicted (more projections of the structure are collected in Supplementary Material). Note that the anion therefore is not depicted correctly from a stoichiometric point of view, but should be viewed as two different configurations, presented simultaneously. Thus, the average position of the bridging oxygen in the Cr1-O4-Cr1 is the $180^\circ$ configuration.
The guanidine part of the tetramethylguanidinium cation group had a planar configuration due to the C1=N3 double bond. However, the terminal C1-N3 bond distance appeared rather long (1.595 (11) Å), which is probably due to the delocalization of the π-electrons over the C-N bonds, in accordance with observations from FTIR spectra. Single-crystal XRD studies (i.e. higher resolution) of [TMGH]$_3$Sb$_2$Br$_9$ done by Bujak and Zaleski [21], report of a double bond length of 1.330 (7) Å.

![Figure 4](image_url)

Figure 4. (a) The atom labels of Cr$_2$O$_2^2$– anion displayed with the bridging O4 atom in two possible positions and the tetramethylguanidinium cation with calculated hydrogen atoms. (b) Crystal structure and hydrogen bonds of [TMGH]$_2$Cr$_2$O$_7$ (without hydrogen atoms) viewed along the b-axis projection.

For the Cr$_2$O$_7^{2–}$ group, the length of the Cr-O bond with the bridging O4 atom was 1.725 (9) Å and thus longer than the terminal Cr-O bonds which were in the range 1.414-1.581 Å. Hence, the dichromate group configuration was comparable with the monoclinic β-K$_2$Cr$_2$O$_7$ system, where the length of the bridging oxygen atom in Cr-O has been reported to be 1.7819 Å and the cornering oxygens to be somewhat shorter in the range of 1.605-1.624 Å [25]. The reason for the shorter bond length from chromium to a cornering oxygen atom could be a result of displacement of charge from the dichromate polyhedra to [TMGH] $^+$ through hydrogen bonds.

The hydrogen bonds H1···O2 and H2···O3 cross-linked the iminium-part of [TMGH] $^+$ with dichromate ions in the (101) plane, thus forming layers connected by ionic and van der Waal’s bonds (Figure 4b). A similar cation-anion linkage was previously reported for [TMGH]$_3$Sb$_2$Br$_9$ [21]. The hydrogen bond lengths were 1.998 Å (N3-H1···O2 angle: 150.5°) and 2.018 Å (N3-H2···O3 angle: 153.8°), respectively. The hydrogen bonds were thus of same strength as those found in liquid water. This agrees well with FTIR spectra of [TMGH]Cr$_2$O$_7$ in Figure 1, where a splitting of the N-H bonds was observed at 3302 and 3229 cm$^{-1}$.

The weight loss curves (TGA) for the [TMGH]$_2$Cr$_2$O$_7$, TiO$_2$-supported [TMGH]$_2$Cr$_2$O$_7$ and reference materials are shown in Figure 5. The corresponding DSC results are presented in Supplementary Material. The thermal decomposition of [TMGH]$_2$Cr$_2$O$_7$ took place in two steps. The first decomposition occurred at 171°C, as also observed in Figure 2, where the dichromate possibly formed intermediate CrO$_3$ [26]. During the second weight reduction from 171-400°C, the remaining organic residues was decomposed and CrO$_3$ reduced to Cr$_2$O$_3$. 

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Figure 5. Relative TGA profiles and the remaining weight at 800 °C

Assuming that the final product after thermal treatment of \([\text{TMG}H]_2\text{Cr}_2\text{O}_7\) is the thermodynamically most stable \(\text{Cr}_2\text{O}_3\), then the mass loss would result in a final weight of 33.9 wt% \((M_{\text{Cr}_2\text{O}_3} \times \frac{m_{\text{TMG}H_2\text{Cr}_2\text{O}_7}}{m_{p}})\) which complies very well with the observed 33.5 wt%. Hence, a redox reaction took place with the reduction of chromium and a resulting oxidation of the organic material. Similar argument for the other samples, and exploiting that the weight loss of TiO\(_2\) was known, namely 2.4 wt%, results in the calculated yields a) CrO\(_3\)/TiO\(_2\): 93.7 wt%, b) \([\text{TMG}H]_2\text{Cr}_2\text{O}_7\)/TiO\(_2\): 84.9 wt%. Thus, all TGA-results were within about 1% of the expected values.

The first decomposition of \([\text{TMG}H]_2\text{Cr}_2\text{O}_7\) at 171°C was accompanied by the mass spectroscopy signals \(m/z = 28, 30, 44, 72\), suggesting liberation of some TMG synthons (C=NH\(_2\), \(m/z = 28\), N(CH\(_3\))\(_2\), \(m/z = 44\), [NH\(_2\)CN(CH\(_3\))\(_2\)], \(m/z = 72\)) as expected. Possibly CO\(_2\) (\(m/z = 44\)) was also formed in connection with the reduction of Cr(VI) to Cr(III).

Figure 6 shows sequential in situ Raman spectra obtained with increasing temperature for a fresh \([\text{TMG}H]_2\text{Cr}_2\text{O}_7\)/TiO\(_2\) catalyst sample under flowing O\(_2\) gas. At temperatures up to 150°C (i.e. below the decomposition temperature of 171°C) only the 638 cm\(^{-1}\) (\(E_g\)) band due to TiO\(_2\)-anatase is seen in the recorded region of the Raman spectra (550–1100 cm\(^{-1}\)). Starting from 200°C the Raman spectra are gradually being transformed and new bands emerge in the Cr-O stretching region (i.e. a broad band at 850 cm\(^{-1}\) and a band at 1000 cm\(^{-1}\)), which are reminiscent of the respective spectral features of the deposited chromia phase of chromia/titania catalysts. Thus, a progressive deconstruction of the organic framework of \([\text{TMG}H]_2\text{Cr}_2\text{O}_7\) and liberation of a deposited \((\text{CrO}_x)_n\) phase take place. The Raman “snapshots” obtained at higher temperatures show that the bands due to the deposited chromia phase gain intensity relative to e.g. the 635 cm\(^{-1}\) \((E_g)\) anatase band that undergoes (as expected) a slight red shift with increasing temperature. At 400°C the organic matrix is almost completely decomposed (see Figure 5), leaving a chromia phase deposited on titania corresponding to a loading of 8.2 mol% Cr/TiO\(_2\) (5.5 wt% Cr/TiO\(_2\)). The final spectrum obtained at 400°C (after in situ calcination in the Raman cell) exhibits a band due to Cr=O stretching at 999 cm\(^{-1}\) (with a shoulder at 1020 cm\(^{-1}\)) and a much broader band at 850 cm\(^{-1}\) due to Cr-O-Cr functionalities \([27,28]\). Although the occur-
Figure 6. Sequential in situ Raman spectra of [TMGH]_2Cr_2O_7/TiO_2 obtained under flowing O_2 at temperatures as indicated. Laser wavelength, λ_0 = 488.0 nm; time constant, τ = 4 s; laser power, w = 40 mW; resolution, 7 cm\(^{-1}\)

The appearance of two bands in the Cr=O stretching region (separated by 20 cm\(^{-1}\)) may indicate a di-oxo Cr(=O)_2 configuration [29], it has been shown that the Raman data combined with ^18O/^16O vibrational isotope effects are consistent with a mono-oxo configuration and an existence of two types of mono-oxo species [30]. Thus, the deposited chromia phase consists of two distinct mono-oxo species, namely a monochromate and associated (polymeric) chromates (bands at 999 cm\(^{-1}\) and at 1020 cm\(^{-1}\) due to Cr=O and at 850 cm\(^{-1}\) due to Cr-O-Cr functionalities of the polychromates). Although, by general consensus the high frequency Cr=O band is assigned to the monochromate [27,30] there have been also cases in which the high frequency Cr=O band has been ascribed to the polychromates [28]. Bands observed at 396 and 511 cm\(^{-1}\) originate from the TiO_2 anatase support, corresponding to the Raman vibrating modes B\(_{1g}\) and A\(_{1g}\), respectively [31].

### 3.2 SCR reactions

The performance of [TMGH]_2Cr_2O_7/TiO_2 (8.2 mol% Cr/TiO_2) in conversion of NO by reduction with ammonia is depicted in Figure 7 along with a CrO_3/TiO_2 (4.3 mol% Cr/TiO_2) reference catalyst. Above 200°C both samples exhibited increased NO conversion until 300°C, whereas the degree of conversion decreased at temperatures from about 325°C. The achieved low-temperature activity was considerably higher than reported by Schneider et al. [13] for various chromium oxides on titania (10 wt% chromium oxide content), where no conversion of NO were detected below 350°C, even with a relatively low GHSV at 28,000 h\(^{-1}\).
At all temperatures $[\text{TMGH}]_2\text{Cr}_2\text{O}_7/\text{TiO}_2$ displayed higher SCR activity than $\text{CrO}_3/\text{TiO}_2$, which partly can be explained by the higher loading of chromium. Furthermore, it is noteworthy that no SCR activity of $[\text{TMGH}]_2\text{Cr}_2\text{O}_7/\text{TiO}_2$ was detected at temperatures below the decomposition temperature at 171°C, thereby making the $[\text{TMGH}]_2\text{Cr}_2\text{O}_7/\text{TiO}_2$ unsuitable for SCR catalysis, although a dispersive effect of the chromium species due to the $[\text{TMGH}]_2\text{Cr}_2\text{O}_7$ can be imagined. However, after decomposition the sample exhibited activity in the reduction of NO with fairly high selectivity, resulting in formation of less than 2% NO$_2$ at temperatures below 325°C. At higher temperatures the chromium-based catalysts exhibited though considerable oxidation of both NO and NH$_3$ ($\Delta$[NH$_3$]/$\Delta$[NO] ratio presented in supplementary material), and seems thus to be more promising candidates for oxidation reactions under normal SCR operation conditions.

4. Conclusion

The crystal structure of 1,1,3,3-tetramethylguanidinium dichromate was found to consist of chromate dimers charge counter-balanced by tetramethylguanidinium cations. The matrix decomposed at 171–172°C into chromium oxides, whereas further thermal treatment at temperatures up to 700°C resulted in formation of Cr$_2$O$_3$.

The activity in the SCR process was investigated by supporting 20 wt% of $[\text{TMGH}]_2\text{Cr}_2\text{O}_7$ on TiO$_2$. Minor conversion of NO was found only at temperatures above $\approx$170°C (after decomposition of the salt) and thus attributed to the activity of formed chromium oxides. Furthermore, above $\approx$325°C the $[\text{TMGH}]_2\text{Cr}_2\text{O}_7/\text{TiO}_2$ functioned as an oxidation catalyst which oxidized the NH$_3$ to nitrogen oxides, thus making it unsuited as SCR catalyst at normal operation conditions.

Acknowledgments

Financial support from Energinet.dk (PSO project 7318) is gratefully acknowledged. The authors are grateful to Mo Hongling Sønnichsen, DTU Chemistry for the TGA data collection.
References

Supplementary material

Structural characterization and catalytic properties of 1,1,3,3-tetramethylguanidinium dichromate

Johannes Due-Hansen¹, Kenny Ståhl², Soghomon Boghosian³, Anders Riisager⁴, Rasmus Fehrmann⁵

April 24, 2010

Abstract

In the following is the supplementary results collected from the study of [TMGH]₂Cr₂O₇: Projections of the crystal structure along the a and c axis, the NO₂ selectivity and NH₃/NO ratio from the activity experiment, and the heat flow from the TGA-DSC experiment.

1. Crystal structure
Figure 2. Crystal structure and hydrogen bonds of [TMGH]$_2$Cr$_2$O$_7$ (without hydrogen atoms) viewed along the c-axis projection

2. NO$_2$ selectivity and NH$_3$/NO ratio

Figure 3. NO$_2$ selectivity of CrO$_3$/TiO$_2$, [TMGH]$_2$Cr$_2$O$_7$/TiO$_2$. As reference a 3wt% V$_2$O$_5$–WO$_3$/TiO$_2$ is used
Figure 4. $\Delta [\text{NH}_3]/\Delta [\text{NO}]$ ratio of CrO$_3$/TiO$_2$, [TMGH]$_2$Cr$_2$O$_7$/TiO$_2$. As reference a 3wt% V$_2$O$_5$–WO$_3$/TiO$_2$ is used. A $\Delta [\text{NH}_3]/\Delta [\text{NO}]$ ratio above unity suggests catalytic oxidation of ammonia.

3. DSC results from TGA-DSC experiment
Pore design of pelletised VOx/ZrO$_2$-SO$_4$/Sepiolite composite catalysts

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*Studies in Surface Science and Catalysis*, 2010 (accepted)
Pore design of pelletised VOX/ZrO2-SO4/Sepiolite composite catalysts

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Abstract

The NH3-SCR activities of a series of extruded and calcined VOX/ZrO2-SO4 - sepiolite catalysts were determined. The pore structures were heavily influenced by the clay content with macropore sizes ranging from 50 to >1000 nm. Mechanical strength and SCR activity measurements suggested that 25% w/w sepiolite is the optimal catalyst composition.

Keywords: Composite, extrusion, biomass, NH3-SCR, ZrO2-SO4, sepiolite.

1. Introduction

The NH3-SCR process is established as a robust and useful technique for the elimination of NOX from off-gases. Commercially used V2O5-WO3/TiO2 catalysts reduce NO selectively to N2 and H2O by the following reaction:

\[
\text{O}_2 + 4\text{NO} + 4\text{NH}_3 \rightarrow 4\text{N}_2 + 6\text{H}_2\text{O}
\]

(1)

The reaction involves two cycles, an acid cycle and a redox cycle [1,2]. Thus, the redox capacity of V(V) and V(IV) is of key importance, but also it is crucial to have sufficient acidic surface sites in order to chemisorb and administer NH3 for the SCR reaction [3]. To comply with the Kyoto protocol, substitution of fossil fuels with biomass constitutes a practical, economical and environmentally viable solution, as biomass (straw, wood chips, saw dust etc.) attains its carbon from air during photosynthesis. Thus, biomass can be regarded as a CO2 neutral fuel, and making attempts to increase the biomass content in energy production is of great interest. Though biomass combustion technology is relatively easy to implement in coal and oil-fired power plants, there are drawbacks. Among others, potassium fly ash particles, originating from the firing of biomass, poison the traditional SCR catalysts [4]. Thus, there is an urgent need for alternative potassium fly ash resistant catalysts. We report herein some special features of composite VOX-ZrO2-SO4/sepiolite catalysts. The catalyst exhibit increased surface acidity as well as a shielding effect of the working catalyst, induced by the sepiolite clay enhancing the durability of the SCR catalyst while working under exposure to fly ash that contains potassium.
2. Experimental

2.1. Catalyst preparation

Melcat $\text{Zr(OH)}_4$ with a particle size $d_{50} = 15\mu m$ was used as the zirconia source and the ammonium sulphate was from Panreac ($> 99\%$). The sulphated zirconia was prepared by impregnating 101.6 g of the zirconia hydroxide with 8.3 g $(\text{NH}_4)_2\text{SO}_4$ by the incipient wetness method. After digestion/drying in ambient conditions for 2 hours the samples were dried at 150°C for 3 hours. Calcination of the samples at 500°C for 4 hours in air was then performed by placing the material directly into a pre-heated furnace, in order to facilitate the preferred formation of metastable tetragonal phases. The catalysts were prepared by wet impregnation of $\text{VOSO}_4$ onto the sulphated zirconia, to produce a slurry that was allowed to settle for an hour under stirring. Thereafter sepiolite, $\alpha$-sepiolite Pansil 100 supplied by Tolsa S.A. and water were added in the desired amounts to obtain a homogeneous paste with an adequate viscosity for extrusion from a 20 ml syringe with a 2 mm orifice. The extrudates were allowed to dry slowly, sealed in a wet atmosphere for 48 hours then dried overnight at 150°C in air. Calcination was carried out at 500°C for 4 hours in air. Finally the extruded material was broken into 3-5mm cylindrical pellets. The synthesis procedure is outlined in Figure 1.

2.2. Catalyst characterisation

The specific surface areas ($S_{\text{BET}}$), were obtained from the corresponding nitrogen adsorption isotherms at $-196^\circ\text{C}$ using a Micromeritics Tristar apparatus, after application of the BET equation in the relative pressure range 0.05–0.35 $p/p^\circ$. Prior to $\text{N}_2$ adsorption, the samples were outgassed overnight at 150°C to a vacuum of $<10^{-4}$ Pa to ensure a dry clean surface, free from any loosely held adsorbed species. The determination of the surface area of the external ($A_{\text{extern}}$), the cumulative pore volumes ($V_p$), and the macropore porediameters ($D_{\text{macro}}$) were complemented by mercury intrusion porosimetry (MIP) using a CE Instruments Pascal 140/240 and applying the Washburn equation for cylindrical pores [5], with the values recommended by the IUPAC of 141° and 484 mNm$^{-1}$, for the contact angle and surface tension of mercury.

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</table>
2.3. NO SCR with ammonia
The SCR activity measurements were carried out on crushed samples sieved to obtain fractions of 0.18–0.300 mm. These measurements were carried out in a fixed-bed reactor, with 10 mg of the catalyst loaded between two layers of inert quartz wool. The reactant gas composition was 1000 ppm NO, 1100 ppm NH₃, 3.5% O₂, 2.3% H₂O and He balance. The total flow rate was 300 mL/min (ambient conditions), with a continuous monitoring of the NO and NH₃ concentration with a Thermo Electron model 17C chemiluminescent NO–NOx gas analyser. The catalytic activity, represented as a first-order rate constant (k), can be calculated from the NO conversion, X, as Eq. 2:

\[
k = -\frac{F_0}{[NO]_0 V_{cat}} \ln(1 - X)
\]

where \(F_0\) is the molar inflow of NO, \([NO]_0\) is the initial molar concentration of NO, and \(V_{cat}\) is the catalyst bed volume.

2.4. KCl poisoning
To simulate poisoning by potassium in the flue gas stream, the sample pellets were submerged in fine particles of KCl (<200 nm) and placed in a furnace for 248 hrs at 350°C in a water saturated air flow. Hereby the temperature is above the Tamman temperature for KCl (m.p. = 771°C, \(T_{Tamman} = 258.5°C\)). Thus, the salt is surface mobile, and will diffuse into the catalyst pores deactivated any accessible active sites.

3. Results & Discussion
3.1. Catalyst pore system
The characteristic data from the pore analyses are collated in Table 1. The \(S_{BET}\) areas were maintained close to 150 m²g⁻¹ for samples with a sepiolite content below 62.5% w/w, whereupon the surface area gradually reduces to a minimum of 130 m²/g - the surface area of pure sepiolite. Since the catalysts were treated at 500°C, little microporosity remained although all exhibited high external surface areas, useful for a catalytic system. The N₂ adsorption-desorption isotherms (\(V_{N₂}\) versus \(P/P^0\)) and MIP cumulative pore volumes (\(V_p\)) are shown in Figure 2 for samples with 0-50% w/w sepiolite content.

![Figure 2: A) Nitrogen adsorption-desorption isotherms of composite catalysts with 0, 25, 50% w/w sepiolite content measured as the volume of N₂ (\(V_{N₂}\)) versus the relative pressure (\(P/P^0\)). B) MIP cumulative pore volumes (\(V_p\)) for 0, 12.5, 25, 37.5 and 50% w/w sepiolite.](image-url)
Varying the sepiolite content of the VO$_x$/ZrO$_2$-SO$_4$/sepiolite composites has a marked effect on the observed pore volumes and macropore diameter. For instance, by introducing 25% w/w of sepiolite the characteristic interparticular macropore diameter decreased from 911 nm (0/8, pure VO$_x$-ZrO$_2$-SO$_4$) to 320 nm.

### 3.2. Catalytic activity and relation to the pore system and crushing strength

Comparing the activity data in Figure 3, before and after KCl exposure, it may be appreciated that almost no deactivation was observed. An extruded reference catalyst (3% w/w) V$_2$O$_5$-WO$_3$/TiO$_2$ under the same conditions suffered an 11% deactivation. It therefore seems that the sepiolite provides a protective effect of the catalytically active centres, or at least delays the deactivation by potassium salts. Furthermore, the effect is present even for the sample with only 12.5% w/w sepiolite, suggesting that the sulfated zirconia itself might have some chemical affinity for the potassium species, which enhance the resistance to deactivation. The crush strengths of the catalysts are also provided in Figure 3. The optimal composition of the composite system, regarding both activity and crushing strength is deduced to be a catalyst consisting of 25% w/w sepiolite.

### References

SCR activity of conformed CuOx/ZrO₂-SO₄ catalysts

Søren B. Rasmussen, Malcolm Yates, Johannes Due-Hansen, Pedro Ávila, Rasmus Fehrmann

Studies in Surface Science and Catalysis, 2010 (accepted)
SCR activity of conformed CuO\textsubscript{X}/ZrO\textsubscript{2}-SO\textsubscript{4} catalysts

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Abstract

CuO\textsubscript{X}/ZrO\textsubscript{2}-SO\textsubscript{4} catalysts have been synthesised as conformed materials with the use of sepiolite as agglomerant and the performance in the NH\textsubscript{3}-SCR reaction with relation to biomass fired boiler units have been studied. The optimal Cu-loading of the catalysts is 3 wt.% CuO, both in terms of activity and selectivity. This catalyst constitutes a possible solution for NO removal in biomass-related applications, since it posses mainly Lewis acid sites, and therefore might be less subjected to deactivation by potassium containing fly ash particle produced during biomass combustion.

Keywords: CuO\textsubscript{X}, sepiolite, biomass, NH\textsubscript{3}-SCR, ZrO\textsubscript{2}-SO\textsubscript{4}.

1. Introduction

The use of biomass in fossil fuel based power plants is of increasing interest, since it is considered as a CO\textsubscript{2} neutral fuel, having zero human impact on the carbon release to the atmosphere. Simultaneously, continuous efficient selective catalytic reduction of NO with ammonia (NH\textsubscript{3}-SCR) remains a very important condition for the implementation of biomass fuel as a sustainable alternative for energy production. However, the NH\textsubscript{3}-SCR catalyst suffers from a number of deactivation phenomena when installed in boiler units based on biomass combustion, due to exposure to potassium containing fly ash\cite{1}. The active V=O and V-OH sites on the commercially used V\textsubscript{2}O\textsubscript{5}-WO\textsubscript{3}/TiO\textsubscript{2} based catalyst reacts with the potassium salts and form inactive alkali vanadates, which are unable to adsorb ammonia.

Therefore new NH\textsubscript{3}-SCR catalysts more resistant to deactivation by potassium salts metals are needed. One of the possible ways to increase catalyst resistance to alkaline poisons is the use of supports revealing high or super-acidic properties, which would interact more strongly with potassium than vanadium species. Potassium oxide affects the Brønsted acid sites of the catalyst to a much larger extent than Lewis sites. Therefore, another possible solution for NO removal in biomass-related applications is the use of other metal oxides as active components, which posses mainly Lewis acidity\cite{2,3}. In this context we have synthesised and studied CuO\textsubscript{X}/ZrO\textsubscript{2}-SO\textsubscript{4} catalysts with respect to the performance of NH\textsubscript{3}-SCR in biomass fired boiler units.
2. Experimental

2.1. Catalyst preparation
The zirconia source was freshly precipitated Zr(OH)₄ from Mel Chemicals. The α-sepiolite employed (Pansil 100) was supplied by Tolsa S.A. Ammonium sulphate and the Cu-precursor, Cu(NO₃)₂ were from Panreac, (> 99%). The sulphated zirconia was prepared by dissolving (NH₄)₂SO₄ and Cu(NO₃)₂ into a part of the water. Thereafter the solution was mixed with the homogenised powder blend of the oxides and pore generating agent (PGA, starch), and extra water was added in order to obtain a paste of adequate rheology. After digestion/drying in ambient atmosphere for 2 hours with occasional kneading, the paste was extruded from 20 ml plastic syringes with 2 mm orifices. The samples were then dried at 150°C for overnight. Calcination of the samples was performed at 600°C for 4 hours in air. Finally the extruded materials were chopped into 3-5mm cylindrical pellets. The synthesis procedure is outlined in Figure 1.

2.2. Catalyst characterisation
The thermal gravimetric analysis of the samples were carried out on a Netzsch 409 EP Simultaneous Thermal Analysis device. The DSC curves were measured using approximately 20-30 mg of powered sample which were heated in an air flow of 75 ml·min⁻¹ at a rate of 5 ºC·min⁻¹ from room temperature to 1000 ºC, using α-alumina as reference. The specific surface areas, S_BET, were obtained from nitrogen adsorption at −196 ºC using a Micromeritics Tri-Star apparatus, after application of the BET equation at relative pressures in the range 0.05–0.35 p/p°.

Table 1. Textural characteristic of conformed catalysts

<table>
<thead>
<tr>
<th>Sample</th>
<th>S_BET (m²g⁻¹)</th>
<th>S_Extren (m²g⁻¹)</th>
<th>V_meso (cm³g⁻¹)</th>
<th>V_macro (cm³g⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1% CuO</td>
<td>110</td>
<td>104</td>
<td>0.20</td>
<td>0.71</td>
</tr>
<tr>
<td>2% CuO</td>
<td>80</td>
<td>68</td>
<td>0.19</td>
<td>0.68</td>
</tr>
<tr>
<td>3% CuO</td>
<td>79</td>
<td>70</td>
<td>0.20</td>
<td>0.73</td>
</tr>
<tr>
<td>5% CuO</td>
<td>84</td>
<td>77</td>
<td>0.19</td>
<td>0.74</td>
</tr>
</tbody>
</table>
SCR activity of conformed CuOₓ/ZrO₂-SO₄ catalysts

Prior to N₂ adsorption the samples were outgassed overnight at 150 °C to a vacuum of <10⁻⁴ Pa to ensure a dry clean surface, free from loosely held adsorbed species.

2.3. NO SCR with ammonia

Activity measurements of the cylindrical pellet shaped catalysts were carried out in a continuous tubular glass reactor that operated at an integral regimen close to an isothermal axial profile. The reactor was 75 cm long and had an internal diameter of 2.54 cm. Ammonia was fed directly into the reactor bed to avoid the formation of ammonia salts [4]. Pellet beds 10 cm long were used. The inlet and outlet NO and NO₂ concentrations were determined by chemiluminescence with a Signal NO + NO₂ analyser (Series 4000). Analysis of N₂O and NH₃ was carried out by IR spectroscopy with a Signal 7000FT GFC Analyser and with an A.D.C. Double Beam Luft Type Infrared Gas Analyser, respectively.

3. Results & Discussion

3.1. Characterisation

The characteristic data from the pore analyses of the CuOₓ/ZrO₂-SO₄ catalysts are collated in Table 1. The BET surface areas are around 84 m²g⁻¹, except for the 1% CuOₓ, which exhibits 110 m²g⁻¹. The samples exhibit high external surface areas, close to the BET surface areas, which suggests that no micro porosity remains. A thermo gravimetric analysis was carried out in order to determine how the sepiolite, copper oxide, PGA and sulphation modify the phase transitions of the zirconia hydroxide gel (Fig. 2). The phase change from amorphous zirconia to the monoclinic phase was observed to occur at around 440 °C for pure zirconia hydroxide gel. By introducing the sulphate as stabilizer for zirconia instead the transformation into the tetragonal phase is achieved at around 580 °C, which appeared to be unaltered by the presence of the binder, and thus treatment at 600 °C was valid to ensure its formation. Furthermore we tested if the PGA and the introduction of Copper salts had influence on the transformations. It was observed that introduction of CuOₓ lead to lower combustion temperatures for the starch, but the transition temperature for formation of the tetragonal zirconia phase remained constant. Based on these data 600 °C was chosen as the temperature for calcinations.

![Figure 2: A) CuOₓ/ZrO₂-SO₄ sepiolite catalyst B) ZrO₂-SO₄ sepiolite carrier material with PGA. C) ZrO₂-SO₄ sepiolite carrier material without PGA. D) Pure ZrO₂.](image-url)
3.2. Catalytic activity
Varying the CuO\textsubscript{X} content on the CuO\textsubscript{X}/ZrO\textsubscript{2}-SO\textsubscript{4}/sepiolite catalysts has some effects on the catalytic activities and selectivities. The NH\textsubscript{3}-SCR activity is steadily increasing with the CuO content until the catalyst contains 3% CuO. Thereafter formation of crystalline CuO sets in according to complementary powder XRD data. Thereafter, any additional copper is deposited as crystalline CuO, which doesn’t contribute to the catalytic activity, since the conversion remains stable. However, this also implies that the crystalline copper formed is merely a spectator species- hence there is no significant decline in activity nor in selectivity. However, the slight decrease in NO/NH\textsubscript{3} consumption ratio observed for the catalysts with 3% compared to 5% CuO could be interpreted as a slight loss in selectivity suggesting that for this catalytic system 3% CuO provides the optimal loading of the active species.

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Multidisciplinary Determination of the Phase Distribution in the VOx-ZrO2-SO4-Sepiolite Catalytic System

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XXII CICAT, 2010 (accepted for paper + oral)
Distribución de las fases en el Catalizador VO$_x$-ZrO$_2$-SO$_4$-Sepiolita determinada por varios métodos.

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Resumen
Se ha preparado y caracterizado una serie de catalizadores V$_2$O$_5$/ZrO$_2$/SO$_4$ – sepiolita, cuya actividad catalítica en la reacción NH$_3$-SCR se ha relacionado con el contenido de la fase activa VO$_x$SO$_4$ en ZrO$_2$. La distribución de la sepiolita y óxido de zirconio en la superficie de la mezcla mecánica fue estudiada por la técnica de migración electroforética. Las muestras se caracterizaron además por adsorción de nitrógeno, difracción de rayos X y por microscopía electrónica de barrido. Los resultados de migración electroforética mostraron que la adición de cantidades crecientes de sepiolita, la cual se amasa en agua junto con la zirconia, tiene un efecto dramático sobre el sistema, porque produce una sensible disminución de la fracción molar de zirconia en la superficie, y se alteran las propiedades electroforéticas de las mezclas. Las microfotografías SEM, los patrones de difracción de rayos X y las medidas de la actividad de las muestras estudiadas concuerdan con los resultados de migración electroforética, verificándose que hasta una composición del 75% en VO$_x$ZrO$_2$SO$_4$, las fibras de sepiolita cubren las partículas de zirconia. Esto tiene una influencia ligeramente negativa sobre la actividad del catalizador global de estos catalizadores; sin embargo, se considera que puede ser una excelente propiedad que permitiría mejorar la vida de los catalizadores De-NOx, expuestos a partículas de cenizas volantes, que contienen sales de potasio malignos.

Palabras clave: NH$_3$-SCR; sepiolita; zirconia sulfatada; potencial zeta

Abstract
A series of V$_2$O$_5$/ZrO$_2$/SO$_4$ - sepiolite mixtures were extruded, calcined and characterized. NH$_3$-SCR activity was found to be related to the content of the active VO$_x$ZrO$_2$SO$_4$ phase. The distribution of sepiolite and zirconia at the surface of the mechanical mixtures was studied by the electrophoretic migration technique. The samples were further characterised by nitrogen adsorption, XRD and scanning electron micrography techniques. The electrophoretic migration results showed that the addition of sepiolite to zirconia, kneaded in water, had a dramatic effect on the quantity of zirconia present at the support’s surface, both strongly decreasing the molar fraction of zirconia at the surface, and altering the electrophoretic properties of the mixtures. The SEM micrographs, XRD patterns and activity measurements of the water-kneaded samples agreed with the electrophoretic migration results. Furthermore, it was verified by electrophoretic migration experiments that up until a composition of 75% VO$_x$ZrO$_2$SO$_4$, the sepiolite fibres coats the zirconia particles. This has a slightly negative influence on overall catalyst activity of fresh catalysts. However, it is believed that this can enhance lifetime of DeNOx catalysts exposed to fly ash particles, which contain malignant potassium salts.

Keywords: NH$_3$-SCR; sepiolite; sulfated zirconia; zeta potential
Distribución de las Fases en el Sistema VO$_x$ZrO$_2$SO$_4$-Sepiolita Determinada Por Varios Metodós

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Resumen

Se ha preparado y caracterizado una serie de catalizadores V$_2$O$_5$/ZrO$_2$/SO$_4$ – sepiolita, cuya actividad catalítica en la reacción NH$_3$-SCR se ha relacionado con el contenido de la fase activa VO$_x$SO$_4$ en ZrO$_2$. La distribución de la sepiolita y óxido de zirconio en la superficie de la mezcla mecánica fue estudiada por la técnica de migración electroforética. Las muestras se caracterizaron además por adsorción de nitrógeno, difracción de rayos X y por microscopía electrónica de barrido. Los resultados de migración electroforética mostraron que la adición de cantidades crecientes de sepiolita, la cual se amasa en agua junto con la zirconia, tiene un efecto dramático sobre el sistema, porque produce una sensible disminución de la fracción molar de zirconia en la superficie, y se alteran las propiedades electroforéticas de las mezclas. Las microfotografías SEM, los patrones de difracción de rayos X y las medidas de la actividad de las muestras estudiadas concuerdan con los resultados de migración electroforética, verificándose que hasta una composición del 75% en VO$_x$ZrO$_2$SO$_4$, las fibras de sepiolita cubren las partículas de zirconia. Esto tiene una influencia ligeramente negativa sobre la actividad del catalizador global de estos catalizadores; sin embargo, se considera que puede ser una excelente propiedad que permitiría mejorar la vida de los catalizadores De-NO$_x$, expuestos a partículas de cenizas volantes, que contienen sales de potasio malignos.

Palabras clave: NH$_3$-SCR; sepiolita; zirconia sulfatada; potencial zeta

Introducción

El catalizador comercial generalmente usado para la Reducción Catalítica Selectiva de NOx con amoníaco, NH$_3$-SCR, está basado en V$_2$O$_5$/TiO$_2$ y reduce selectivamente al NO en presencia de aire, según la siguiente reacción:

$$O_2 + 4NO + 4NH_3 \rightarrow 4N_2 + 6H_2O$$  \hspace{1cm} (1)

Esta reacción consta de dos ciclos, un ciclo de ácido y un ciclo redox [1-9]. Para el ciclo de oxidación-reducción, la capacidad redox de V (V) y V (IV) es de importancia clave, mientras que para el ciclo ácido, el elemento clave es contar con suficientes centros ácidos superficiales para quimisorber y distribuir el NH$_3$ en la superficie del catalizador V$_2$O$_5$/TiO$_2$.

En un intento por cumplir con el Protocolo de Kyoto, la sustitución de combustibles fósiles con biomasa constituye una solución viable, práctica, económica y ambiental, ya que la biomasa (en forma de paja, virutas de madera, polvo, etc.) conserva su carbono del aire durante la fotosíntesis. Así, la biomasa, en este sentido puede ser considerada como un combustible neutro de CO$_2$, y actualmente en Dinamarca DONG Energy A / S está intentando aumentar el contenido de biomasa en la producción de energía hasta un 20%.

Aunque la tecnología de combustión de biomasa es relativamente fácil de implementar en las centrales eléctricas de carbón y petróleo, existen algunos inconvenientes que deben ser resueltos. Por un lado, el potasio de las partículas de cenizas volantes es muy agresivo a altas temperaturas, e induce problemas a los materiales, como la corrosión y el deterioro de las baldosas cerámicas en el quemador. Por otra parte, también es
extremadamente venenoso para los catalizadores de la reacción NH₃-SCR mencionados.

**Experimental**

Preparación de la muestra: Como fuente de zirconio se utilizó hidróxido de zirconio Zr(OH)₄ Melcat, con un tamaño de partícula d₅₀ = 15 µm y el sulfato amónico de Panreac (> 99%). La zirconia sulfatada fue preparada por impregnación de 101,6 g de hidróxido de zirconio, con 8,3 g de (NH₄)₂SO₄ por el método de humedad incipiente. Después de la digestión y secado en condiciones ambientales durante 2 horas, las muestras se secaron a 150°C durante 3 horas. Se llevó a cabo la calcinación de las muestras a 500°C durante 4 horas colocando el material directamente en un horno precalentado, a fin de facilitar la formación preferente de las fases cúbica metaestable y tetragonal.

Los catalizadores fueron preparados por impregnación húmeda de VOSO₄ en la zirconia sulfatada, para producir una pasta que se deja durante una hora en agitación. Posteriormente, se añaden la sepiolita, α-sepiolita Pansil 100 suministrada por Tolsa SA y el agua en los niveles deseados para obtener una pasta homogénea con una viscosidad adecuada para la extrusión de una jeringa de 20 ml con un orificio de 2 mm. Se dejan secar lentamente los extruidos, manteniéndolos en un ambiente húmedo durante 4 horas en atmósfera de aire. La calcinación se realizó a 500°C durante 4 horas en atmósfera de aire. Por último, el material resultante fue dividido en piezas cilíndricas 3-5 mm de largo.

**Actividad catalítica (Reducción Catalítica Selectiva de NO con NH₃):** Las medidas de la actividad SCR se llevaron a cabo en un reactor de lecho fijo, con 10 mg de catalizador, colocado entre dos capas de lana de cuarzo inerte. La composición de los gases de reacción fue de 1000 ppm de NO, 1100 ppm de NH₃, 3,5% de O₂, 2,3% de H₂O y He hasta balance. El caudal total fue de 300 ml/min. (en condiciones normales), con un seguimiento continuo de la concentración de NO por quimioluminiscencia con un analizador de gases Thermo Environmental, modelo 17C. La actividad catalítica, representada como una constante de velocidad de orden 1(k), puede calcularse a partir de la conversión de NO, X, a partir de la ecuación 2:

\[
k = \frac{F_0}{[NO]_0 V_{cat}} \ln(1 - X)
\]

(2)
donde \(F_0\) es el flujo molar de entrada de NO, \([NO]_0\) es la concentración molar de entrada de NO, y \(V_{cat}\) es el volumen de lecho catalítico.

**Textura:** Se determinaron los valores de área específica (BET) mediante isotermas de adsorción/desorción de nitrogeno con un Micromeritics TriStar 3000 y la distribución de volumenes de micro, meso y macroporos combinando los resultados de esta técnica con los obtenidos mediante porosimetría de intrusión de mercurio.

**Migración Electroforética:** El Punto de Carga cero (ZPC) de los materiales composites fue determinado midiendo el potencial zeta en función del pH de la solución. Los potenciales zeta fueron obtenidos usando la ecuación de Helmholtz-Smoluchowski midiendo la velocidad de migración electroforética de las partículas en un Zeta-Meter Inc. Instrument, modelo 3.0+, provisto de una unidad de transferencia de muestra automática (bomba peristáltica y pinzas eléctricas especiales) para evitar los problemas de sedimentación de muestra.

Los experimentos se realizaron usando 30 mg de muestras de polvo (<2 µm) suspendidas en 200 ml de KCl 10⁻³ M, ajustando el pH con disoluciones 0.2 M de KOH y HCl. En cada muestra no se detectó más de un tipo de partícula, excepto para la que contenía 12,5% de sepiolita. Cada curva fue repetida al menos dos veces para confirmar la reproducibilidad de los resultados.

Se ha calculado la fracción molar de cada componente en la superficie de la muestra de acuerdo al método previamente descrito [10], que se basa en el hecho de que el punto de carga cero (ZPC) de una mezcla cae entre los puntos isoeléctricos (IEP) de los materiales individuales (A y B). Según este principio se desarrollaron las ecuaciones (3) y (4).

\[
ZPC = X_A (IEP)_A + X_B (IEP)_B
\]

(3)

\[
X_A + X_B = 1
\]

(4)

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Figura 1: Difractogramas de Rayos-X de los composites VO\textsubscript{X}-ZrO\textsubscript{2}-SO\textsubscript{4}/sepiolita después de la extrusión, secado y calcinación a 500°C.

Esto permite determinar la fracción molar de cada componente expuesta en la superficie, relacionando dicho valor con la composición total de la muestra. No se realizaron cálculos posteriores para obtener el grado de cubrimiento aparente (% en peso en la superficie) como se han realizado en estudios previos, debido a la dificultad de asignar una masa molecular a la sepiolita, cuya fórmula estructural es Si\textsubscript{12}Mg\textsubscript{8}O\textsubscript{30}(OH)\textsubscript{4}(H\textsubscript{2}O)\textsubscript{4}·8H\textsubscript{2}O [11].

Micoscopía Electrónica de Barrido (SEM): La morfología y el tamaño de las partículas de los catalizadores fueron investigados por SEM, utilizando un FEI Inspect ‘S’ con filamento de tungsteno. Las imágenes fueron obtenidas con voltajes de aceleración de 5 a 10 kV. sobre las muestras previamente grafitizadas.

Difracción de rayos-X (XRD): Se obtuvieron los difractogramas de polvo de cada muestra usando un difractómetro Huber G670 Guinier analizando el rango de 2\(\theta\): 3-100\(^\circ\) en etapas de 0,005\(^\circ\), durante un tiempo de 120 min a temperatura ambiente y longitud de onda \(\lambda = 1,54051\)\(\AA\). La muestra se coloca en una capa fina de cinta adhesiva y se gira durante la recolección de datos.

Resultados y Discusión

En la Figura 1 se muestran los difractogramas de rayos X de los composites VO\textsubscript{X}-ZrO\textsubscript{2}-SO\textsubscript{4}/sepiolite después de la extrusión, secado y calcinación a 500°C. En ella se aprecian los picos de difracción de ZrO\textsubscript{2} a 2\(\theta\) = 28-32\(^\circ\). En las muestras se aprecia la formación de la fase monoclínica, pero la fase dominante es la fase cúbica. Esta última fase cristalina se parece mucho a la tetragonal de ZrO\textsubscript{2} (2\(\theta\) = 31,586), que se ha sugerido que es necesario para la generación de la alta acidez en la zirconia sulfatada. Por lo tanto, esta reflexión se asigna a menudo a esta fase. Sin embargo, en un análisis previo de Rietveld, se concluyó que dicha reflexión corresponde de hecho a la fase cúbica, la cual probablemente presenta propiedades similares.

Por otra parte, con el aumento del contenido de sepiolita se observa la aparición progresiva de sepiolita cristalina, CaCO\textsubscript{3}, SiO\textsubscript{2} y CaSO\textsubscript{4} (Ca\textsuperscript{2+} de la fase de sepiolita que reacciona con sulfato de la fase de zirconia). La formación de CaSO\textsubscript{4} podría sugerir que la acidez de la fase de zirconia puede disminuir, y debería añadirse sulfato extra para compensar este efecto. Esta observación puede ser importante para la actividad catalítica del catalizador, pero no modifica el alcance de este trabajo, ya que aquí se concentra la atención en la distribución de fases entre VO\textsubscript{X}-ZrO\textsubscript{2}-SO\textsubscript{4} y sepiolita.
Figura 3: Variación de los puntos de carga cero ZPC y del recubrimiento superficial aparente (ASC en gráfica insertada), en función del contenido en sepiolita.

En la Figura 2 se muestran las curvas del potencial zeta en función del pH del sistema VOₓ-ZrO₂-SO₄₃, la sepiolita y sus mechas amasadas con agua, todas ellas tratadas a 500°C durante 4 h. Los Puntos isoeléctricos (IEPs) de la sepiolita, el ZPC del sistema VOₓ-ZrO₂-SO₄₃ y de sus mezclas, así como los recubrimientos superficiales aparentes resultantes, se muestran en la Figura 3. Estos valores del ZPC se calcularon por interpolación de la curva de potencial Zeta.

El valor de ZPC= 5,8 del sistema VOₓ-ZrO₂-SO₄₃ es función del IEP de la fase pura de zirconía [12], el V₂O₅, y el efecto del sulfato. El IEP de la sepiolita, 2,2, ha sido previamente reportado [13]. El efecto de la adición de sepiolita a la zirconia que se muestra en la Figura 2, indica que la presencia de sepiolita provoca un gran desplazamiento del ZPC a valores bajos, incluso con bajos contenidos del silicato. Este efecto fue menos pronunciado para contenidos altos de sepiolita.

El ZPC alcanza un valor muy cercano al de la sepiolita pura con sólo el 25% (w/w) de este silicato en la composición global. Este efecto se muestra en la Figura 3, donde se representa el ZPC en función del contenido de sepiolita. Las fracciones molares de sepiolita en la superficie, calculadas mediante la aplicación de la ecuación 3 y 4 a los resultados (véase la sección experimental) se presentan en la Figura 3 (gráfica insertada), expresados como porcentajes.

Figura 4: Microfotografías SEM de la serie de VOₓZrO₂-SO₄/sepiolite.

No es de extrañar que el porcentaje de zirconia en la superficie se redujera en gran medida por la adición de pequeñas cantidades de sepiolita.

Este efecto de fuerte enmascaramiento de la sepiolita a un óxido (zirconia) en la superficie se ha demostrado antes para titanio [14], si bien en este caso de la fase VOₓ-ZrO₂-SO₄₃ es mucho más pronunciado que en el caso de la titania. El porcentaje de zirconia necesario para alcanzar fracciones molares de 50% de zirconia en la superficie fue de más del 75% en peso.

Dado que la fase activa de la reacción estudiada es el sistema VOₓ-ZrO₂-SO₄₃, sería importante conseguir una protección parcial de los sitios activos optimizando la composición de este tipo de catalizadores con el fin de aumentar la vida útil del catalizador. De esta forma se ha considerado que la sepiolita podría ejercer esta función protectora siempre que se mantenga una buena accesibilidad del zirconio presente en la superficie.

El comportamiento de la mezcla de los dos sólidos podría explicarse a partir de la termodinámica y las fuerzas motrices que operan en la interacción de los dos componentes y la morfología del sistema durante el proceso de amasado. Esto ha sido investigado por este grupo en un estudio anterior [14] para el caso de un catalizador SCR sobre la base de titania-sepiolita. La sepiolita tratada a 500°C se compone de partículas con tamaños que van desde 100 a 250 µm. Estas partículas están compuestas, a su vez, por haces de fibras, que varían en tamaño desde 0,2 hasta 1µm de longitud y 10-100 nm de ancho. Las partículas de titano (0,05-0,1µm) están formadas por agregados de 0,3-1 µm, en el que las partículas originales mantienen su identidad.
El objetivo de este trabajo es utilizar estos fenómenos para incorporar parcialmente la fase activa de zirconia dentro de los haces de fibras de sepiolita y proteger así a los centros de óxido de vanadio de los compuestos de potasio básicos presentes en los gases de combustión de biomasa.

El recubrimiento físico de la fase de zirconia por la sepiolita puesto de manifiesto por las medidas del potencial zeta ha sido confirmado por análisis SEM de los catalizadores calcinados (Figura 4). A partir de estas micrografías se puede ver que la superficie de la muestra 1/8 (12,5% sepiolita), básicamente, presenta una morfología similar a la de la muestra 0/8 (sistema VOX-ZrO2-SO4). Algunas fibras de la sepiolita se puede ver adjunto a las partículas de óxido de zirconio, lo cual explica plenamente por qué el punto de carga cero de la muestra de 1/8 es un promedio simple entre la composición y los potenciales de carga cero de las fases puras.

Sin embargo, desde la muestra 2/8 (25% en sepiolita), se observa un cambio morfológico espectacular. El material compuesto se observa como cristales de zirconio rodeados de haces de fibras de sepiolita con un aspecto externo muy similar al de la sepiolita pura (muestra 8/8). Esto concuerda totalmente con la variación de los valores de ZPC observada en las Figuras 2 y 3. En ellas se aprecia que todas las muestras con un porcentaje de sepiolita igual o superior al 25% en peso (muestras 2/8 - 8/8) tienen el punto de carga cero muy cerca del valor del PIE de la sepiolita pura.

En la Figura 5, se muestran los valores de actividad en el proceso NH₃-SCR a 360°C para los catalizadores estudiados, en función del contenido de sepiolita. Como es bien sabido que las especies activas en este proceso son los VOₓ distribuidos sobre óxidos de metales de transición (TiO₂, Al₂O₃, o ZrO₂), la tendencia general observada en la Figura 5 es la lógica. En general, al aumentar el contenido de sepiolita se observa una disminución de la actividad. Sin embargo, la tendencia no parece seguir completamente una relación lineal entre el contenido de fase activa (VOₓ/ZrO₂-SO₄) y la actividad. La disminución de la actividad parece más fuerte de lo esperado al aumentar el contenido de sepiolita.

Esta pérdida de linealidad podría deberse a un bloqueo severo de los centros activos por parte de la sepiolita que dificulte el acceso de los gases, lo cual no parece muy probable teniendo en cuenta los espacios interfibrilares observados por SEM. En la Figura 5 también se puede confirmar que no hay cambios drásticos con respecto a los volúmenes de poros o áreas específicas, y parece que la caída de la actividad es debida a una desactivación química de la fase activa de la sepiolita. Esta segunda opción está en concordancia con la formación de las fases CaSO₄ observadas por difracción de rayos X en la Figura 1.

**Conclusiones**

La serie de catalizadores híbridos de V₂O₅-ZrO₂-SO₄/sepiolita obtenidos por extrusión de sus mezclas, secados y calcinados a 500°C, demostraron ser activos para el proceso NH₃-SCR. La velocidad de reacción de cada catalizador está directamente relacionada con la cantidad de fase activa de VOₓ-ZrO₂-SO₄ presente, pero se observó la existencia de cierta desactivación debida a la formación de CaSO₄ confirmada por los resultados de DRX. Se ha verificado, por experimentos de migración electroforética, que las fibras de sepiolita recubren externamente las partículas de ZrO₂-SO₄ hasta una composición de 75% de ZrO₂-SO₄.

La influencia negativa de la adición de sepiolita sobre la actividad del catalizador global probablemente puede ser compensada mediante la adición de sulfato extra durante la síntesis o el uso de sepiolita de alta pureza, con bajo contenido en Ca. Por otra parte, es especialmente interesante la posible mejora de la vida de catalizador, debido al efecto inducido por el recubrimiento de la fase activa por una capa de sepiolita, que sería especialmente efectiva en plantas térmicas de biomasa, en las que los catalizadores se ven sometidos a una fuerte desactivación causada por el potasio que contienen las partículas de cenizas volantes.
Agradecimientos
Los autores agradecen al Ministerio español de Ciencia e Innovación la financiación del Proyecto CTMT2008-06876-CO2-O2/TECNO, a la Comunidad de Madrid (CAM) el Programa de investigación S0505/AMB/0406, JAE (Junta de Ampliación de Estudios), PSO (Proyectos FU5201 y FU7318) y el acuerdo de colaboración CSIC-USACH 2008CL0017.

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Catalysis Communications, 10, 2009, 803-806
Impact of support and potassium-poisoning on the $V_2O_5$–$WO_3$/ZrO$_2$ catalyst performance in ammonia oxidation

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A R T I C L E   I N F O

Article history:
Received 22 October 2008
Received in revised form 28 November 2008
Accepted 2 December 2008
Available online 9 December 2008

Keywords:
SCR catalysts
Tungsten oxide
Vanadia
Potassium
Deactivation
N$_2$O oxidation
Zirconia
Crystallinity

A B S T R A C T

A series of WO$_3$-promoted zirconia supports were synthesized and calcined between 400 and 800 °C. Subsequently vanadium oxide was introduced to obtain 3.5 wt% $V_2O_5$. The influence of the calcination temperature and potassium-poisoning (K/V = 0.2, molar ratio) on the catalytic activity in the selective catalytic reduction (SCR) of NO with NH$_3$ was previously studied. Here, we focus on the influence of these parameters on the catalytic activity for the undesirable oxidation of ammonia, which decrease the $N_2$ selectivity of the SCR process. It is found that potassium doping of the catalysts results in a considerable decrease in the overall ammonia conversion while selectivity to NO increases.

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1. Introduction

The selective catalytic reduction (SCR) of NO$_x$ by NH$_3$ in the presence of O$_2$ remains the state of the art technology for controlling NO$_x$ emissions from stationary sources. Transition metal oxides (most commonly vanadia) promoted with tungsten or molybdenum oxides supported on anatase are among the most active SCR catalysts used in industrial scale at temperatures 300–450 °C. It is generally agreed by most researchers that the SCR reaction proceeds through an Eley–Rideal mechanism with activation of ammonia by adsorption and NO reacting directly from the gas phase or from a weakly adsorbed state. However, the selectivity of the SCR reaction can be compromised by the undesired oxidation of ammonia to $N_2$, $N_2O$, NO, and NO$_2$.

$$4NH_3 + 3O_2 \rightarrow 2N_2 + 6H_2O$$  \hspace{1cm} (1)

$$2NH_3 + 2O_2 \rightarrow N_2 + 3H_2O$$  \hspace{1cm} (2)

$$4NH_3 + 5O_2 \rightarrow 4NO + 6H_2O$$  \hspace{1cm} (3)

$$4NH_3 + 7O_2 \rightarrow 4NO_2 + 6H_2O$$  \hspace{1cm} (4)

A significant work on the catalytic oxidation of ammonia over various metal oxides, including $V_2O_5$, $WO_3$, and TiO$_2$, was undertaken already in the 1970s by Il’ichenko and Golodets [1]. They discussed the correlation between the selectivity of the NH$_3$ oxidation and the binding energy of lattice oxygen in metal oxides, since a higher binding energy of oxygen would cause a poorer oxidation ability of the metal oxide. The study showed that high lattice oxygen binding energy induces a low NH$_3$ oxidation rate, but high N$_2$ selectivity. The formation of nitrous oxides during ammonia oxidation has recently been described over $V_2O_5$–$WO_3$/TiO$_2$ catalysts at high temperatures [2]. The authors observed that the selectivity towards N$_2$O increased with temperature and vanadia loading. Ozkan et al. [3,4] suggests three combinational sites responsible for the ammonia adsorption, by isotopic labeling studies, namely the following neighboring sites: (a) two V=O, responsible for NO formation from ammonia, (b) V=O and V–OH, responsible for oxidative formation of N$_2$ and N$_2$O from ammonia, and (c) two V–OH, accountable for formation of N$_2$ from ammonia. The authors therefore concluded that the V=O sites are responsible for the oxidation of ammonia to NO and N$_2$O. Jung and Grange [5] showed by DRIFT spectroscopy of $V_2O_5$, that above 400 °C almost all vanadium is fully oxidized, in other words, all V–OH sites are oxidized to V=O sites at temperatures higher than 400 °C. This implies that both increased rate of NO formation during the ammonia oxidation and decreased selectivity to N$_2$ in the SCR reaction can be explained by the increased number of V=O sites above 400 °C. Thus,
the oxidation state of vanadium, i.e. the distribution between the dual active V–OH and V@O sites, is an essential factor for controlling the reactivity and ammonia oxidation.

Although the adsorption, activation and oxidation of ammonia over vanadia-based catalysts have been investigated to a certain degree, the literature of the potassium influence on the reaction is only scarcely covered. The effect of alkaline metals on the SCR activity of V₂O₅/TiO₂ catalysts has been reported by some authors [6–9]. Most of them conclude that the poisons are affecting the Brønsted acid sites (V–OH), thus decreasing both their number and activity in NO reduction. Chen and Yang [10] reported on the alkali influence on the NO conversion of V₂O₅/TiO₂ and V₂O₅-WO₃/TiO₂ catalysts and found that doping with 0.3–2.2 wt% K₂O resulted in an overall “negative nitrogen monoxide conversion” in the NH₃ SCR reaction at temperatures above ~450 °C, i.e. additional NO was produced by the competitive reaction of ammonia oxidation than consumed by SCR reaction. The authors thus claimed that K₂O additive was an effective promoter for ammonia oxidation. However, they only measured the actual ammonia oxidation at 500 °C on V₂O₅/TiO₂, WO₃/TiO₂ and V₂O₅-WO₃/TiO₂ samples, and thus not on potassium-doped samples.

Recent studies have shown that the catalysts supported on tungstated zirconia have improved resistance towards alkali poisoning in the reaction of NH₃ SCR with ammonia [11]. Especially the crystallinity of the zirconia-supported catalyst was found to be an important parameter accounting for NOx reduction activity. Nevertheless, the catalyst doping with potassium in all cases lead to a considerable decrease of their catalytic activity. It was suggested that potassium adsorbs to the same sites as ammonia, thus blocking a fraction of the sites necessary for the SCR reaction. Therefore, a support with a high surface acidity is required, to host the basic potassium, leaving the active vanadia sites available for reaction. The traditional support used for SCR catalysts is titania, promoted with tungsten or molybdenum oxides. This system reveals only moderate surface acidity. On the contrary, tungstated zirconia is known to display high surface acidity and its surface properties, e.g. acidity, specific surface area, and crystallinity can be tuned in a wide range by varying preparation procedure, WO₃ content and calcination temperature [12,13].

To our knowledge, there are no studies of the ammonia oxidation over vanadia-loaded zirconia-based catalysts and their corresponding potassium-doped samples. In the present paper, we report on the activity of fresh and potassium-doped vanadia–zirconia-based catalysts for the oxidation of ammonia.

2. Experimental

2.1. Catalyst preparation

Tungstated zirconia (WZ) was synthesized by coprecipitation of ammonium metatungstate and zirconyl nitrate hydrate and calcined at different temperatures ranging from 400°C (WZ_400) to 800°C (WZ_800). The resulting supports, containing 20 wt% WO₃, were pressed at 10 tons (Ø = 13 mm) and crushed into 0.180–0.295 mm particles and consequently impregnated to obtain 2.0 wt% vanadium (VWZ). Doping with potassium was performed using incipient wetness impregnation with a KNO₃ solution to obtain samples with a K/V molar ratio of 0.2, resulting in the poisoned catalyst K₂O-V₂O₅-WO₃/ZrO₂ (KVKWZ). After doping, the samples were dried at 100°C for 2 h. For further synthesis details, see [11].

The commercial reference catalyst shaped as fiber-reinforced monolith system, consists of a 5 wt% V₂O₅ and 7 wt% WO₃ supported on TiO₂, anatase (1.7VWT_ref). The active material on the monolith-surface was separated from the fiber material and pressed into tablets at 5 tons, Ø = 13 mm and crushed into 0.180–0.295 mm particles.

2.2. Catalyst characterization

Room-temperature X-ray powder diffraction (XRD) measurements were recorded on a Huber G670 powder diffractometer using Ni-filtered Cu Kα radiation (λ = 1.5406 Å) with 2θ scan range 10–65° in steps of 0.02°. The XRD-patterns of the crystalline structures were refined with the Rietveld technique using WINPOW software [14]. The crystalline peak profiles were represented by a pseudo-Voigt profile function and a five-coefficient Chebyshev polynomial background fitting.

2.3. Catalytic activity measurements

The ammonia oxidation reaction was carried out in a continuous flow fixed bed quartz reactor, operating at atmospheric pressure.

In the ammonia oxidation experiments, the ammonia concentration were measured with a Jasco V-570 UV/VIS/NIR spectrophotometer, using the characteristic ammonia absorption band around λ = 201 nm. Thermo Electron’s Model 10A Rack-Mounted Chemiluminescent NO–NO₂ Gas Analyzer was used to detect NO concentrations. The operating gas composition for the ammonia oxidation was: 1100 ppm NH₃, 3.5% O₂, 2.3% H₂O and balance N₂. The total flow rate was maintained at 300 ml/min (ambient conditions).

3. Results and discussion

3.1. Surface area and structural characterization

The morphological characteristics of the WO₃/ZrO₂ supports calcined at different temperatures and the results of the BET surface area measurements are outlined in Table 1 and show the effect of calcination temperature on the surface area, crystallinity and phase composition of the support. After synthesis, the samples were annealed in air between 400 and 800°C. Calcination at 400 and 500°C resulted in essentially amorphous samples, whereas calcination at 600–800°C yielded more crystalline samples. With increasing calcination temperature, there is a significant increase in sample crystallinity from 37.1% to 54.7% when going from 500 to 600°C, respectively. Further increase in calcination temperature leads to increased crystallinity, resulting in highest crystallinity (67.5%) for the sample calcined at 800°C (cf. Supplementary material Fig. 1).

The crystalline samples are composed of three ZrO₂ crystal phases with different zirconium/oxygen atom positions and interatomic distances. XYZ coordinates of the space groups are avail-

### Table 1

<table>
<thead>
<tr>
<th>BET area (m²/g)</th>
<th>Crystallinity (%)</th>
<th>Vₖ₂o₃</th>
<th>V₅OH</th>
<th>Vₓ mm</th>
<th>Vₓ tot</th>
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<tbody>
<tr>
<td>WZ_400</td>
<td>226.8</td>
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<td>–</td>
<td>–</td>
</tr>
<tr>
<td>WZ_500</td>
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<td>–</td>
<td>–</td>
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<td>WZ_800</td>
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<tr>
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<td>–</td>
<td>(0.19)</td>
<td>(0.99)</td>
<td>(0.40)</td>
</tr>
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</table>

* Calculated with the diffraction pattern program ‘Winprep’ 2006-01-17 from 2θ 10–65°.
able in Supplementary material, Table 1. The volumetric amount of crystalline phase of monoclinic WO₃ and the three zirconia phases (tetragonal, monoclinic and cubic) are calculated with the Rietveld method for WO₃/ZrO₂ samples calcined at temperatures between 600 and 800 °C and presented in Table 1.

Cortés-Jacome et al. [15] have also performed Rietveld analysis of WO₃/ZrO₂ samples, synthesized according to similar recipe. The authors tried to fit the XRD results with a model, first based on tetragonal and cubic zirconia, which resulted in a poor fit. Better results were obtained when fitting with two combined tetragonal phases, by varying the oxygen position (corresponding to the z-position of t-ZrO₂). In this study, however, the cubic and monoclinic phases of ZrO₂ are included in the model in order to describe the possible influence of different crystal phases on the catalytic behavior in the NH₃ SCR reaction.

For the sample calcined at 600 °C, the majority of the crystalline zirconia is present in the tetragonal form, 91.25 vol%, whereas only 8.69 vol% exists as cubic ZrO₂, with hardly any monoclinic impurities in the matrix. As a function of the calcination temperature, the relative phase composition of the polymorph mixture changes. Increasing the calcination temperature results in a decrease in tetragonal ZrO₂, compensated by increased amount of cubic and monoclinic ZrO₂ along with appearance of monoclinic tungsten oxide phase. The latter can be related to the decrease of the surface area with increasing calcination temperature and therefore higher surface concentration of WO₃ species, far exceeding monolayer coverage. The observed results are in good correlation with previous studies of the temperature effect on the crystalline zirconia [13,16]. The authors found that zirconia was present in mainly tetragonal form at lower temperatures and transformed to monoclinic phase above 600 °C. It was found that sintering of zirconia crystallites at elevated temperatures leads to the stabilization of ZrO₂ in the monoclinic phase.

3.2. NH₃ oxidation measurements

The ammonia oxidation on the fresh and poisoned catalyst samples has been measured between room temperature and 550 °C. Below 400 °C, only negligible NH₃ oxidation was observed for all catalysts tested. Both NH₃ conversion and selectivity towards NO formation are increasing with calcination temperature. The ammonia conversion for different catalysts along with the crystallinity and fractional vanadium monolayer (ML) coverage of the corresponding tungstated zirconia support are presented in Fig. 1 at 450 and 500 °C as function of the support calcination temperature. Vanadium monolayer coverage on ZrO₂ is reported at 6.8 V/nm² [17], which can be recalculated to 15–59% vanadium ML coverage for the 2.0VWZ_400 and 2.0VWZ_800 samples respectively (1.0–4.0 V/nm²). All fresh catalysts show noticeable activity in ammonia oxidation reaction, which increases parallel with the calcination temperature of the support (WZ). The crystallinity of the support and vanadium surface density seems to influence the NH₃ oxidation ability to a much larger extent than surface acidity (see Table 2). Amorphous samples (VWZ_400, VWZ_500), even though they possess high specific surface area, show quite low ammonia conversion.

Poisoning of the V₂O₅–WO₃/ZrO₂ system with K₂O leads to a decline of the overall NH₃ oxidation activity. Previously, a similar decrease of the activity in NH₃ SCR reaction was observed after potassium-poisoning [11]. This indicates that potassium is either deactivating both types of the sites used for the NH₃ SCR reaction and the NH₃ oxidation or both reactions proceed in parallel on the same type of active sites which are sensitive towards poisoning with potassium.

In order to compare selectivities of different catalysts at the same levels of NH₃ conversion the selectivity is plotted versus the conversion (Fig. 2), where the main products of NH₃ SCR reaction are NO and NO. Commercial SCR catalyst consisting of 3 wt% V₂O₅–7 wt% WO₃/TiO₂ (denoted as 1.7VWT_ref) is used as reference. It is clearly seen that all crystalline unpioned catalysts, including the anatase-based reference catalyst, fit into almost identical selectivity vs. conversion curves. This indicates that the conversion of

<table>
<thead>
<tr>
<th>SCR activity (s⁻¹)</th>
<th>Ammonia desorbed (μmol/g)</th>
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</thead>
<tbody>
<tr>
<td>Fresh</td>
<td>K-doped</td>
</tr>
<tr>
<td>Fresh</td>
<td>K-doped</td>
</tr>
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<td>385</td>
</tr>
<tr>
<td>2.0VWZ_900</td>
<td>411</td>
</tr>
</tbody>
</table>

**Fig. 1.** NH₃ conversion, vanadium monolayer surface coverage and crystallinity for fresh V₂O₅–WO₃/ZrO₂ and corresponding K-doped catalysts calcined from 400 to 800 °C at two operating temperatures (450 and 500 °C). Experimental conditions: 50 mg sample, 1100 ppm NH₃, 3.5% O₂, 2.3% H₂O and balance N₂ to 300 ml/min.

**Fig. 2.** NO Selectivity versus the NH₃ conversion for fresh and K-doped catalysts measured in the temperature range 450–550 °C. Open symbols represent fresh catalysts, solid symbols denote the corresponding K-doped samples.
ammonia is proceeding by a similar mechanism over all fresh catalysts and that the selectivity to NO is determined by similar factors. Two amorphous catalysts, VWZ_400 and VWZ_500, display negligible ammonia conversion and are therefore not shown here. Although all the fresh catalysts show low selectivity to NO at temperatures below 525 °C, a significant increase in the selectivity is observed at higher temperatures and consequently higher NH₃ conversions. For example, the VWZ_800 catalyst converts almost 100% of the ammonia already at 525 °C with high selectivity to NO, particularly 47% at 550 °C.

Doping the zirconia-based catalysts with potassium to K/V = 0.2 results in an increased selectivity to NO for all catalysts, while the conversion of ammonia decreases significantly, as indicated by the gray dotted lines in Fig. 2, representing the data recorded at 450 °C. The increase in NO selectivity with reaction temperature (i.e. with increasing NH₃ conversion in Fig. 2) is associated with the expected decrease in N₂ selectivity at elevated reaction temperatures. Madia et al. [18] also reported this trend for V₂O₅–WO₃/TiO₂ catalysts in the ammonia oxidation reaction. The increase in the NO selectivity due to K-doping is highest for the reference V₂O₅–WO₃/TiO₂ and VWZ_800 catalysts, which both have high vanadium surface concentration of all the samples tested, 3.3 and 4.0 V/nm², respectively. This complies well with the results of Nova et al. [2], who reported of decreasing N₂ selectivity for samples with high surface vanadium density. Surprisingly, the overall conversion of NH₃ on the reference catalyst is almost unaffected by the potassium doping in contrast to the more vulnerable SCR reaction, where the catalyst deactivates almost completely. According to Ozkan et al. [3,4], the results suggest that only V=O sites are available on the catalyst surface and responsible for the increase in NO selectivity, meaning that all the V=OH sites have been deactivated by potassium.

Apparent activation energies for the ammonia oxidation to NO have been calculated and are presented in Table 3. The reaction is assumed first-order in NH₃, due to the high concentration of oxygen compared to ammonia. For the fresh zirconia-based catalysts, all calculated activation energies are in the range of 114–160 kJ/mol, and 217 kJ/mol for the reference catalyst. For the comparison, apparent activation energy of the NH₃ SCR reaction on the 3V₂O₅–7WO₃/TiO₂ reference catalyst is calculated to be 64 kJ/mol, which correlate well with those reported in the literature for V₂O₅/TiO₂ catalysts, which are in the range 69–82 kJ/mol for a V₂O₅–MoO₃/TiO₂ catalyst [19]. The large difference between activation energies for SCR reaction and ammonia oxidation explains why the SCR reaction is prevailing at temperatures below 400 °C. Doping the catalyst with potassium (K/V = 0.2) leads to an increase in the corresponding apparent activation energy of about 30% in average, i.e. a decrease in ammonia oxidation rate, as observed in Fig. 2. However, care must be taken in the interpretation of the activation energies since the calculated apparent activation energies in the present work are based on relative high ammonia conversions (up to ~10%) and therefore not entirely in the kinetic region. For example, a slight increase in activation energy is calculated for the reference catalyst, from 217 to 227 kJ/mol when doping with potassium despite slightly higher NH₃ conversion is detected after being poisoned with potassium.

4. Conclusion

The Rietveld refinement of the tungstated zirconia calcined from 400 to 800 °C confirm that the majority of the crystalline phase is present as tetragonal ZrO₂. Phase transformation of the tetragonal zirconia into cubic and monoclinic forms occurs with increasing calcination temperature, as well as formation of monoclinic WO₃ at temperatures above 700 °C.

All samples tested for ammonia oxidation exhibited negligible NH₃ oxidation at operating temperatures below 400 °C. At temperatures above 450 °C a noticeable increase in ammonia oxidation rate is observed.

The activity of the catalysts in NH₃ oxidation correlate well with the vanadium surface density and the crystallinity of the support, which implies that the support is either involved in the NH₃ oxidation mechanism or facilitates the formation of specific active sites responsible for ammonia oxidation. For the same NH₃ conversion levels, selectivity towards NO is similar for all fresh zirconia- and anatase-based catalysts indicating that the conversion of ammonia is proceeding by a similar mechanism over all fresh catalysts and that the selectivity to NO is determined by similar factors.

Doping of the catalysts with potassium (K/V = 0.2) results in reduction in the NH₃ conversion for all samples, which correlate well with the calculated apparent activation energies. Since poisoning with potassium also induces a decrease in the overall SCR activity, this could indicate that potassium is affecting the sites used for both the NH₃ SCR and ammonia oxidation reaction. Although doping with potassium decreases the overall activity in ammonia oxidation, a significant increase in the selectivity towards NO is observed for the potassium-poisoned samples.

Acknowledgements

The authors thank Energinet.dk (PSO Project FU5201 and FU7318) and the Danish National Research Foundation for financial support.

Appendix A. Supplementary data


References


<table>
<thead>
<tr>
<th>Table 3</th>
<th>Apparent activation energies for the apparent first-order NH₃ oxidation to NO.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Eₚ [kJ/mol]</td>
<td>Fresh</td>
</tr>
<tr>
<td>2.0VWZ, 600</td>
<td>114</td>
</tr>
<tr>
<td>2.0VWZ, 700</td>
<td>123</td>
</tr>
<tr>
<td>2.0VWZ, 800</td>
<td>160</td>
</tr>
<tr>
<td>1.7VWT, ref</td>
<td>217</td>
</tr>
</tbody>
</table>

* Calculated for temperature interval 400–450 °C.