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Ferrierite and Its Delaminated and Silica-Intercalated Forms Modified with Copper as Effective Catalysts for NH₃-SCR Process

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Abstract: The main goal of the study was the development of effective catalysts for the low-temperature selective catalytic reduction of NO with ammonia (NH₃-SCR), based on ferrierite (FER) and its delaminated (ITQ-6) and silica-intercalated (ITQ-36) forms modified with copper. The copper exchange zeolitic samples, with the intended framework Si/Al ratio of 30 and 50, were synthetized and characterized with respect to their chemical composition (ICP-OES), structure (XRD), texture (low-temperature N₂ adsorption), form and aggregation of deposited copper species (UV-vis-DRS), surface acidity (NH₃-TPD) and reducibility (H₂-TPR). The samples of the Cu-ITQ-6 and Cu-ITQ-36 series were found to be significantly more active NH₃-SCR catalysts compared to Cu-FER. The activity of these catalysts in low-temperature NH₃-SCR was assigned to the significant contribution of highly dispersed copper species (monomeric cations and small oligomeric species) catalytically active in the oxidation of NO to NO₂, which is necessary for fast-SCR. The zeolitic catalysts, with the higher framework alumina content, were more effective in high-temperature NH₃-SCR due to their limited catalytic activity in the side reaction of ammonia oxidation.

Keywords: NH₃-SCR; ferrierite; ITQ-6; ITQ-36; copper

1. Introduction

Nitrogen oxides (NOₓ) belong to the most dangerous air pollutants responsible for various environmental problems, such as photochemical smog, acid rain and ozone depletion. Moreover, NOₓ emission contributes to the global warming problem. Nitrogen oxides are formed mainly as side-products of fossil fuel combustion in power plants, vehicles and factories. Thus, various technologies, including the optimization of the fuel combustion process and post-combustion technologies, used for the elimination of NOₓ from flue gases, were developed and implemented [1]. Among the post-combustion technologies of NOₓ emission control, the selective catalytic reduction of nitrogen oxides with ammonia (NH₃-SCR) is one of the most important methods. In this case, NOₓ present in flue gases are selectively reduced by ammonia to dinitrogen and water vapor, which are the desired products of this process. This technology was first applied in Japan in the 1970s for the conversion of NOₓ emitted by electric power stations and has since spread to the USA, Europe and several Asian countries [2]. The major industrial catalyst for NH₃-SCR is based on the V₂O₅-WO₃/TiO₂ metal oxide system and effectively operates in a relatively narrow temperature range from 300 to 400 °C [2–4]. At temperatures below 300 °C, the efficiency level of NOₓ conversion is not satisfying.
but while above 400 °C the side reaction of direct ammonia oxidation decreases the efficiency of NOx conversion. Industrial NH3-SCR catalysts are used mainly in the form of monoliths, located upstream of electrostatic precipitator (ESP). Therefore, such units for NOx conversion operate with dusty gas stream (high-dust SCR), which may result in the clogging of channels in monolith by particles of dust. To avoid this problem, flue gases prior to the NH3-SCR unit, should be dedusted in the ESP unit (low-dust SCR). The disadvantage of such a solution is the relatively low temperature of flue gases—about 250 °C or even lower, downstream of EPS [5]. Thus, for the low-dust SCR technology development of the NH3-SCR catalysts, effective operation at a temperature lower than 250 °C is necessary. Copper belongs among the most promising active components of the low-temperature NH3-SCR catalysts [1], especially in the case of copper deposited into zeolites [6–10]. It is not surprising, taking into account the ion-exchange properties of zeolites important for the deposition of metal in highly dispersed forms, surface acidity important for chemisorption and the activation of basic ammonia molecules, high surface area and uniform porous structure important to generate large number of active sites. The catalytic activity of copper strongly depends on its form and aggregation. Monomeric copper cations in the ion-exchange positions of the zeolitic framework are reported to be catalytically active in low-temperature NH3-SCR [11–13]. Shan et al. [11] reported that monomeric Cu2+ cations in Cu-SSZ-13 zeolites are catalytically active in the oxidation of NO to NO2, which is necessary for fast-SCR (2NH3 + NO + NO2 → 2N2 + 3H2O), important for low-temperature NH3-SCR. On the other hand, it was reported that aggregated CuO species are significantly less active in the NO2 formation [11]. Clark et al. [12] reported that stable solvated Cu+ cations are responsible for the limitation of low-temperature NH3-SCR efficiency, while at higher temperature, the reduction of Cu2+ to Cu+ is the rate limiting step in the case of Cu-exchanged small-pore SSZ-13 zeolites. Another problem related to the application of small pore zeolites, such as Cu-SSZ-13, in NH3-SCR is the possible aggregation of Cu-species with the formation of bulky CuO aggregates, which not only decrease catalytic activity, but may also result in the destruction of the zeolite skeleton and lead to a breakdown in the long-range order of its structure [13]. Thus, copper is very promising in low-temperature NH3-SCR, but the controlled deposition of active metal species is necessary. The high-temperature NH3-SCR process is limited by the side reaction of direct ammonia oxidation by oxygen present in flue gases. Thus, the NH3-SCR catalysts operating in the high-temperature range should be active in NOx reduction with ammonia and should be inactive in direct ammonia oxidation. The formulation of such catalyst is a demanding task, because in both these processes ammonia chemisorption and activation is one of the main steps in the majority-postulated reaction mechanisms of NH3-SCR [2] and ammonia oxidation (AMOx) [14].

Recently, layered zeolites with delaminated or intercalated structures aroused great interest as catalysts [15] or catalytic supports [16,17], due to the increased rate of internal diffusion. Delaminated layered zeolites consist of ordered zeolite layers, which are chaotically oriented, forming a so-called “house of cards” structure with the bimodal pore structure. Micropores are located in zeolite layers, while the larger pores are interlayer spaces. Pillared layered zeolites can be produced by the surfactant directed method. In this case amorphous silica aggregates are deposited into the interlayer space of swollen layered zeolite precursors, which, during calcination, are converted into stable silica pillars, which permanently open interlayer space in the zeolitic sample. Such zeolites are characterized by bimodal porous structure—micropores in zeolitic layers and larger pores in interlayer spaces. It should be mentioned that not only pure silica pillars can be intercalated into the interlayer space of zeolites but the successful intercalation of multicomponent pillars into layered zeolites was also reported (e.g., silica–titania pillars [18]). As previously mentioned, zeolitic materials with the hierarchical porous structure are very promising due to the increased rate of internal diffusion, which is important for the overall reaction rate. However, our recent studies [7,19] proved that the deposition of metal species in highly dispersed form was more effective for MFI zeolites with micro-mesoporous structures than for classical microporous MFI zeolites. It is very important to assume that the form and aggregation of the deposited metal play a crucial role in the NH3-SCR process.
The presented study reports the catalytic properties of copper modified ferrierite and its delaminated (ITQ-6) and silica-intercalated (ITQ-36) forms in the NH$_3$-SCR process and associated reactions of NO to NO$_2$ oxidation and ammonia oxidation (AMOx). The influence of porous structure, Si/Al ratio, as well as the form and aggregation of copper species introduced into the zeolitic samples, were analyzed and discussed.

2. Results and Discussion

2.1. Characterization of Catalyst Precursors and Catalysts

The zeolites were obtained from two layered precursors (PREFER) with the intended molar Si/Al ratios of 30 (PREFER_30) and 50 (PREFER_50). The calcination of such precursors resulted in the condensation of layers with the formation three dimensional (3D) microporous ferrierite zeolites (FER_30 and FER_50). The second part of the zeolite precursors was swollen with alkylammonium surfactants to open interlayer space for the intercalation of the silica pillars precursor (tetraethyl orthosilicate, TEOS), which after calcination resulted in ITQ-36_30 and ITQ-36_50 (depending on the intended Si/Al ratio in the zeolite framework). On the other hand, swollen PREFER_30 and PREFER_50, after sonication and calcination, resulted in delaminated ITQ-6_30 and ITQ-6_50 zeolites, respectively.

The process of PREFER swelling, condensation, delamination and pillaring was monitored by the X-ray diffraction method, as shown in Figure 1. In the diffractograms of the layered precursors, the (200) reflection at $2\Theta = 6.8^\circ$, corresponding to the interlayer distance of about 1.3 nm, was identified [20]. After the swelling of PREFER_30 with surfactant, the main (200) reflection was shifted to $2\Theta$ at about 3.0°, indicating the successful intercalation of the surfactants into the interlayer space of zeolites and resulting in the extending of the interlayer distance to about 2.1 nm. This interlayer distance is very close to the length of the hexadecyltrimethylammonium cations (CTMA$^+$), indicating that the surfactant molecules are oriented perpendicularly to the zeolite layers. Apart from the main (200) reflections, the diffraction peaks at $2\Theta$ at about 6.0, 6.8 and 8.9°, characteristic of the parallel ordering of the zeolite layers, are also present in the diffractogram of the swollen PREFER_30 sample. The first diffraction peak indicates the extension of the interlayer distance to about 1.5 nm and shows that part of the zeolite was intercalated with the surfactant molecules oriented non-perpendicularly to the zeolite layers [15]. The diffraction peak at 6.8° corresponds to a small fraction of the PREFER_30 sample, which was not swollen, while the diffraction peak at 8.9° is possibly related to the partially condensed zeolite layers. The swelling process causes a structural disorder in PREFER_30 indicated by broadening peaks and the loss of their intensities. The calcination of PREFER_30 resulted in a shift in the (200) reflection to 9.7°. The high intensity of this basal reflection, as well as the diffraction peaks characteristic of the 3-dimensional (3D) structure of ferrierite, indicates the successful synthesis of this zeolite (FER_30). The sonication of swelled PREFER_30 in acidic conditions, followed by calcination, resulted in a nearly completely delaminated structure (non-parallel ordering of the zeolite layers). The low-intensive reflection at 9.7° indicates that part of the zeolite layers was condensed with the formation of ferrierite. However, the comparison of this reflection intensity in diffractograms of FER_30 and ITQ-6_30 shows that the contribution of the ferrierite in the delaminated ITQ-6_30 sample is relatively small. The intercalation of swollen PREFER_30 with silica pillars resulted in ITQ-36_30. The diffractogram of this sample is characteristic of the delaminated structure, due to the lack of the basal reflections typical of the pillared samples with the parallel ordering of the zeolite layers. The (200) diffraction peak at about 9.3° is only slightly shifted in relation to the (200) reflection in the diffractogram of FER_30 (9.7°), indicating the presence of small micropores, with a diameter of about 0.5 nm located in the interlayer space of the zeolite. However, the very low intensity of this reflection, in comparison to the (200) reflection recorded for FER_30, shows only a small contribution of this fraction of material in the ITQ-36_30 sample. The comparison of diffractograms recorded for FER_30 and FER_50, with respect to the intensity of the reflections, shows that the calcination of
PREFER_50 resulted in the ferrierite zeolite with better crystallinity, as shown in Figure 1. However, the disappearance of the (200) reflection in the diffractogram of the swollen PREFER_50 sample indicates the delamination of its layered structure. The very small diffraction peak, in the position characteristic of the (200) reflection, in the diffractogram of PREFER_50 indicates that only a small fraction of this sample was not swollen. Moreover, the swelling process, conducted in strong basic media, resulted in the significant destruction of the zeolite structure, which corresponded to decreased intensity or the disappearance of the reflections characteristic of the intralayer ordering of the zeolite structure. The broad reflection in the range of 15–30° indicates the presence of amorphous silica [21], which was possibly formed by the extraction of silica from the zeolite precursor in strong basic media used for the swelling of PREFER_50. Possibly part of the dissolved silica was precipitated in a solution and deposited on the external zeolite surface in the form of amorphous silica aggregates. Thus, it seems that the stability of the ferrierite precursors (PREFER) in a basic media strongly depends on the Si/Al ratio. The high-silica ferrierite precursors are significantly less stable in basic media than the low-silica PREFER samples. The delamination and intercalation of the swollen PREFER_50 sample resulted in delaminated materials, consisting of chaotically organized defected zeolite layers with a significant contribution of amorphous silica. The low intensive (200) reflection at 9.7° indicates the presence of a ferrierite phase in these samples.

Figure 1. X-ray diffraction patterns of ferrierite zeolite-based samples (PREFER, swelled PREFER, delaminated ITQ-6, pillared ITQ-36, FER) with Si/Al molar ratios of 30 (a) and 50 (b).

The dinitrogen adsorption–desorption isotherms of the studied samples are presented in Figure 2, while their textural parameters are compared in Table 1. The isotherms of FER_30 and FER_50 are type I, according to the IUPAC classification [22], characteristic of microporous materials. The microporous character of the samples is indicated by a steep adsorption at low relative pressure, assigned to dinitrogen capillary condensation in micropores [23]. The isotherms of the ITQ-6_30 and ITQ-6_50, as well as the ITQ-36_30 and ITQ-36_50 samples, presented in Figure 2, are type IV, according to the IUPAC classification [22], characteristic for mesoporous materials. However, an increase in adsorbed dinitrogen volume, observed at very low relative dinitrogen pressure, also indicates a significant contribution of micropores in these series of the zeolitic samples. Micropores are located in the zeolitic layers, while mesopores are the spaces between intercalated and delaminated zeolite layers in ITQ-36 and ITQ-6, respectively. As it was shown by XRD analysis, the swelling of high-silica PREFER_50 resulted in the partial extraction of silicon from the ferrierite layers with the formation of amorphous silica, identified in ITQ-6_50 and ITQ-36_50, as shown in Figure 1. Thus, an increase in adsorbed dinitrogen volume, observed at higher relative pressure for these samples, indicates not only the presence of interlayer mesopores but also mesopores of different size in amorphous silica [24].
The hysteresis loops present in the adsorption–desorption isotherms of ITQ-6_30 and ITQ-36_30 are type H4, typical for mesoporous zeolites and micro-mesoporous materials, while the hysteresis loops recorded for ITQ-6_50 and ITQ-36_50 are type H3, characteristic of non-rigid aggregates of plate-like particles [22]. The characteristic step down in the desorption isotherms, associated with hysteresis loop observed for ITQ-6_50 and ITQ-36_50, indicates the presence of partially plugged mesopores [25].

Profiles of pore size distribution in the micropore and mesopore ranges for ferrierites and their delaminated and intercalated forms are shown in Figure 3. In the case of FER_30 and FER_50, the maximum micropore size distribution, located at about 0.53–0.55 nm, fits very well to the diameter of 10 MR (member ring) channels in ferrierite [16,26]. In the mesopore range, the only small contribution of pores with the diameter below 15 nm was found for both ferrierite samples. The delamination and intercalation of PREFER_30, resulting in ITQ-6_30 and ITQ-36_30, respectively, reduced the intensity of micropore size distribution profile, as shown in Figure 3. This effect is related to the ordered microporous structure limited only to the zeolite layers in the case of ITQ-6_30 and ITQ-36_30, and the three dimensional microporous structure of FER_30 and FER_50. In the mesopore range, the maxima of pore size distribution are at about 3.6 and 2.6 nm for ITQ-6_30 and ITQ-36_30, respectively. For both these samples, the tails up to about 18 nm indicate the high heterogeneity of pore sizes, which is characteristic of delaminated materials. The microporous character of ITQ-6_50 and ITQ-36_50 is significantly different than the ITQ-6_30 and ITQ-36_30 samples, as shown in Figure 3. The maximum
characteristic of the ferrierite 10 MR channels disappeared, while broad and low-intensive peaks at about 0.6–0.7 nm with long tails from the side of larger pores indicate the high heterogeneity of micropore sizes. These results are consistent with the XRD studies, which showed the partial destruction of the ferrierite layers during PREFER_50 swelling. The mesopore size distribution in ITQ-6_50 and ITQ-36_50 is much broader than for the analogous zeolitic samples with the lower Si/Al ratio. The maxima are located at about 4.5 and 2.6 nm for ITQ-6_50 and ITQ-36_50, respectively. It was shown by the XRD analysis of these samples that part of the silicon was extracted from the zeolite layers during PREFER_50 swelling and was then deposited in the form of amorphous silica aggregates on the zeolite external surface, as shown in Figure 1. Thus, possibly the mesopores of various size, present in amorphous silica, contribute in the mesopore size distribution profiles of ITQ-6_50 and ITQ-36_50. The textural parameters of all the studied samples are presented in Table 1. Micropores dominate in FER_30 and FER_50, while in their delaminated and intercalated forms the contribution of micropores is reduced, while the mesopore volume significantly increased. The introduction of copper into these zeolitic samples decreased their textural parameters.

The elemental compositions of the zeolitic samples, presented in Table 2, show that the real Si/Al ratios, determined by the ICP-OES method, are 22 for FER_30 and 64 for FER_50, so slightly different from the intended values of 30 and 50, respectively. The delamination of PRERER_30 did not change the Si/Al ratio in ITQ-6_30. However, in the case of ITQ-6_50, the Si/Al ratio increased to 214, indicating a decrease in aluminum content in this sample. As previously discussed, the swelling conducted in strong basic conditions resulted in the partial leaching of silicon from the zeolite layers of PREFER_50, which in solution possibly formed amorphous silica aggregates deposited on the zeolite surface. This effect was observed only for high-silica zeolite, which is possibly less stable in basic conditions. In the next step of ITQ-6_50 synthesis, the swollen PREFER_50 was treated in acidic conditions, which possibly resulted in the effective leaching of aluminum from the partially degraded zeolite layers. Additionally, the partial dealumination of PREFER_50 in swelling conditions, as the process assisting silicon leaching, cannot be excluded. The intercalation of the zeolite precursors with silica to obtain ITQ-36_30 and ITQ-36_50 resulted in an increase in the Si/Al ratio. The content of copper introduced to the zeolitic samples is in the range of 0.9–2.7 wt %, as shown in Table 2. Taking into account that one Al\(^{3+}\) cation in the zeolite framework generates one negative charge in such a framework, as well as taking into account that copper is deposited in the form of Cu\(^{2+}\) cations, the number of copper cations introduced by the ion-exchange mechanism should be two-times lower than the Al\(^{3+}\) cations in the zeolite framework. The Cu/Al molar ratio for FER_30 and its delaminated and pillared forms, as well as for FER_50, are in the range of 0.55–0.64, thus it seems that, in these cases, the ion-exchange mechanism dominated in copper deposition. Much larger Cu/Al ratios, determined for ITQ-6_50 and ITQ-36_50 indicate a contribution, aside from ion-exchange, also other mechanisms of copper species deposition.

The surface acidity of the zeolitic samples was studied by the temperature-programmed desorption of ammonia (NH\(_3\)-TPD). Ammonia is a basic molecule with respect to both Brønsted and Lewis theory and therefore may interact with both these types of acid sites. Moreover, the diameter of the ammonia molecule is about 0.3 nm and therefore may easily penetrate all microporous channels in ferrierite [27,28]. The ammonia desorption profiles, presented in Figure 4A, consist of two desorption peaks and are characteristic of ferrierite zeolites [26]. The low-temperature maximum was centered at about 205–215 °C for the samples with the intended Si/Al ratio in the zeolite framework of 30, and at about 175–195 °C for the zeolites with the intended framework Si/Al ratio of 50, which indicates the presence of the relatively weak acid sites. The second, much broader and less intensive maxima, centered at about 405–445 °C for the zeolites with the intended framework Si/Al ratio of 30 and at about 300–405 °C for the samples with the Si/Al ratio in the zeolite framework of 50, are related to the stronger acid sites. The most significant difference in desorption profiles is related to their intensity depending on the Si/Al ratio. It is not surprising, assuming that the surface acidity depends on the aluminum content in the zeolitic samples. The deposition of copper into zeolites resulted in a very significant modification of the ammonia desorption profiles, as shown in Figure 4B. In the case of FER_30,
the intensity of the low-temperature maximum was significantly reduced, while the high-temperature maximum was shifted to lower temperature. The similar effect of the high-temperature maximum shift to lower temperature was observed for FER_50; however, in this case the deposition of cooper resulted in an increase in ammonia adsorption. The ammonia desorption profiles of delaminated and intercalated zeolites consist of one broad and asymmetric maximum, centered at about 245 °C in the case of the samples with the lower Si/Al ratio, while for zeolites with the higher Si/Al ratio at a temperature about 15–25 °C lower. Moreover, the surface concentration of adsorbed ammonia increased after copper deposition to these samples. Thus, the introduction of copper into the zeolitic samples very significantly modified their surface acidic properties. It should not be surprising, taking into account that possibly all ion-exchange positions were substituted by copper species, as shown in Table 2, and in the Cu-ITQ-6_50 and Cu-ITQ-36_50 samples the content of deposited copper is a few times higher than their ion-exchange capacity, as shown in Table 2. Copper species introduced to zeolites play a role in the Lewis acid sites, which can accommodate a free electron pair of ammonia molecules into unoccupied d-orbital. Of course the content of ammonia chemisorbed in this way depends on the surface availability of the copper species and, therefore, their dispersion. The results of NH3-TPD should be discussed not only as an evaluation of the surface acidity of the samples, but also as important steps in the NH3-SCR and AMOx processes, in which chemisorption and the activation of ammonia is one of the main stages [2,14]. This topic will be discussed together with the results of the catalytic tests.
Additionally, the shoulders in the range of 280–400 nm, observed in the spectra of other samples, (e.g., Cu(H2O)62+) are attributed to such small copper oxide aggregates. Moreover, the shoulder above 400 nm could be assigned to the presence of small CuO crystallites [31]. Copper oxide crystallites were not observed in the spectra of Cu-ITQ-6 and Cu-ITQ-36 series, the bands above 400 nm and below 600 nm are assigned to the d–d transition of Cu2+ level of this experimental method. For the samples of the Cu-ITQ-6 and Cu-ITQ-36 series, the bands above 600 nm are assigned to the d–d transition of Cu2+ ions in pseudo-octahedral coordination (e.g., Cu(H2O)62+) [31–33], and therefore cannot be used for the analysis of acid sites located inside such channels.

The coordination and aggregation of copper species introduced into the zeolitic samples was studied by UV-vis-DR spectroscopy, as shown in Figure 5. The intensive band centered at about 220–245 nm, present in the spectra of all copper-containing samples, is related to monomeric Cu2+ ions interacting with the oxygen of the zeolite framework (O2−→Cu2+) [31–33]. In the case of Cu-FER_30, the band centered at 340 nm indicates the presence of small oligomeric copper oxide species [31–33]. Additionally, the shoulders in the range of 280–400 nm, observed in the spectra of other samples, are attributed to such small copper oxide aggregates. Moreover, the shoulder above 400 nm could be assigned to the presence of small CuO crystallites [31]. Copper oxide crystallites were not identified by XRD analysis of the samples, which indicates that their size is below the detection level of this experimental method. For the samples of the Cu-ITQ-6 and Cu-ITQ-36 series, the bands above 600 nm are assigned to the d–d transition of Cu2+ ions in pseudo-octahedral coordination (e.g., Cu(H2O)62+) [31–33], and therefore indicate, similarly to the bands at 220–245 nm, the significant contribution of copper deposited in the form of monomeric cations.

Figure 3. Pore size distributions of FER, ITQ-6 and ITQ-36 in micropore (left—(a,c,e), respectively) and mesopore range (right—(b,d,f), respectively).

Table 2. Silicon, aluminum and copper content in FER, ITQ-6 and ITQ-36 samples.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Si [wt %]</th>
<th>Al [wt %]</th>
<th>Cu [wt %]</th>
<th>Si/Al [mol/mol]</th>
<th>Cu/Al [mol/mol]</th>
</tr>
</thead>
<tbody>
<tr>
<td>FER_30</td>
<td>44.9</td>
<td>2.0</td>
<td>-</td>
<td>22</td>
<td>-</td>
</tr>
<tr>
<td>FER_50</td>
<td>46.1</td>
<td>0.7</td>
<td>-</td>
<td>64</td>
<td>-</td>
</tr>
<tr>
<td>ITQ-6_30</td>
<td>44.9</td>
<td>2.0</td>
<td>-</td>
<td>22</td>
<td>-</td>
</tr>
<tr>
<td>ITQ-6_50</td>
<td>46.7</td>
<td>0.2</td>
<td>-</td>
<td>214</td>
<td>-</td>
</tr>
<tr>
<td>ITQ-36_30</td>
<td>45.0</td>
<td>1.9</td>
<td>-</td>
<td>23</td>
<td>-</td>
</tr>
<tr>
<td>ITQ-36_50</td>
<td>46.5</td>
<td>0.2</td>
<td>-</td>
<td>224</td>
<td>-</td>
</tr>
<tr>
<td>Cu-FER_30</td>
<td>43.4</td>
<td>2.0</td>
<td>2.6</td>
<td>21</td>
<td>0.55</td>
</tr>
<tr>
<td>Cu-FER_50</td>
<td>45.6</td>
<td>0.6</td>
<td>0.9</td>
<td>73</td>
<td>0.64</td>
</tr>
<tr>
<td>Cu-ITQ-6_30</td>
<td>43.4</td>
<td>1.9</td>
<td>2.7</td>
<td>22</td>
<td>0.60</td>
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<tr>
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<td>44.9</td>
<td>0.2</td>
<td>2.7</td>
<td>216</td>
<td>5.74</td>
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<tr>
<td>Cu-ITQ-36_30</td>
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<td>1.8</td>
<td>2.5</td>
<td>24</td>
<td>0.59</td>
</tr>
<tr>
<td>Cu-ITQ-36_50</td>
<td>46.5</td>
<td>0.2</td>
<td>2.7</td>
<td>224</td>
<td>5.20</td>
</tr>
</tbody>
</table>
The analysis of the pyridine pre-adsorbed samples by FTIR spectroscopy, which is a standard method used for determination of Brønsted and Lewis acid site contribution, is very problematic in the studies of the ferrierite based samples. According to Wichterlova et al. [29] and Starzyk et al. [30], the kinetic diameter of pyridine is about 0.57 nm, which is bigger than the diameter of 10 MR channels in ferrierite [16,26], and therefore cannot be used for the analysis of acid sites located inside such channels.

The coordination and aggregation of copper species introduced into the zeolitic samples was studied by UV-vis-DR spectroscopy, as shown in Figure 5. The intensive band centered at about 220–245 nm, present in the spectra of all copper-containing samples, is related to monomeric Cu^{2+} ions interacting with the oxygen of the zeolite framework (O^{2-} → Cu^{2+}) [31–33]. In the case of Cu-FER_30, the band centered at 340 nm indicates the presence of small oligomeric copper oxide species [31–33]. Additionally, the shoulders in the range of 280–400 nm, observed in the spectra of other samples, are attributed to such small copper oxide aggregates. Moreover, the shoulder above...
maxima, centered at about 235–245 °C, as mentioned earlier, could be attributed to the direct reduction of CuO to metallic copper and the reduction of monomeric Cu$^{2+}$ cations to Cu$. However, the intensive high-temperature maxima, centered at about 475–505 °C and related to the reduction of isolated Cu$^+$ to Cu$^0$, may indicate the presence of mainly highly dispersed copper species in these samples. The peak at about 370 °C in the reduction profile of Cu-ITQ-6_30 and the arm at a similar temperature in the profile of Cu-ITQ-6_50 are possibly attributed to the reduction of finely dispersed CuO particles to elemental Cu$^0$ [34,35]. The maxima in the reduction profile of the sample with the higher Si/Al ratio, Cu-ITQ-6_50, are shifted to higher temperatures, compared to Cu-ITQ-6_30. A similar effect was reported in the scientific literature and was explained by the lower electronic density of the samples with lower aluminum content, which makes the reduction of the deposited metal species more difficult [36,37]. In the case of Cu-ITQ-36_30 and Cu-ITQ-36_50, the reduction profiles contained low-temperature peaks, centered at about 230–245 °C, and high-temperature maxima at about 445–470 °C, as shown in Figure 6. The intensities of the high temperature peaks are significantly higher compared to the low-temperature ones, which indicates not only the significant contribution of highly dispersed copper species, but also the presence of Cu$^+$ cations in these samples. It is possible that part of the highly dispersed copper species was thermally reduced from Cu$^{2+}$ to Cu$^+$ during the treatment of the samples in a flow of pure argon at 500 °C, prior to the H$_2$-TPR runs. A similar effect was reported and analyzed for the Cu-ZSM-5 samples by Yang et al. [38]. This effect may indicate a very high lability in Cu$^{2+}$ → Cu$^+$ reduction.
The decrease in the NO conversion, observed at higher temperatures, is related to the side reaction of ammonia oxidation by oxygen present in the reaction mixture. The selectivity to dinitrogen, also possibly attributed to the better reducibility of copper species deposited into zeolites. The catalytic performance of these samples is significantly different at higher temperatures. In the presence of ferrierite-based catalysts, the NO conversion started at about 150–175 °C, while for Cu-ITQ-6_50 a decrease in the NO conversion, due to the side reaction of ammonia oxidation, was observed at higher temperatures. Similar results were obtained for the series of Cu-ITQ-36 samples. It is possible that part of the highly dispersed copper species was thermally reduced from Cu2+ to Cu+ during the treatment of the samples in a flow of pure argon at 500 °C, prior to the H2-TPR determination of the types of copper species introduced into the zeolitic samples. The reduction of Cu2+ to Cu+ at temperatures below 300 °C and, at higher temperatures, the reduction of CuO particles to elemental Cu0 [34,35]. The maxima in the reduction profile of the sample with the higher aluminum content, Cu-ITQ-36_30, presented better catalytic activity at temperatures below 175 °C, possibly due to the better reducibility of copper species deposited into this sample compared to the copper species in the catalysts with the lower aluminum loading. It can be seen that the catalyst with the higher aluminum content, Cu-ITQ-6_30, was more effective in the low-temperature range, and then intensively increased to 275 °C, reaching 99 and 86% for Cu-FER_30 and Cu-FER_50, respectively. The decrease in the NO conversion, observed at higher temperatures, is related to the side reaction of ammonia oxidation by oxygen present in the reaction mixture. The selectivity to dinitrogen, also in the range characteristics of ammonia oxidation, was very high and did not drop below 97%. Intercalated and delaminated ferrierite zeolites were found to be significantly more catalytically active in the NH3-SCR process. First of all, the NO conversion in the presence of the Cu-ITQ-6 catalysts started at about 125 °C—so at a temperature lower by about 25–50 °C than in the case of FER based catalysts—and then intensively increased to 275 °C, reaching about 97 and 95% for Cu-ITQ-6_30 and Cu-ITQ-6_50, respectively. Thus, in the low-temperature range the NO conversion profiles for both catalysts of this series are very similar, indicating the limited role of their surface acidity and copper loading. It can be seen that the catalyst with the higher aluminum content, Cu-ITQ-6_30, was more...
catalytically active at temperatures below 150 °C. As it was shown, the reduction of copper species in the zeolitic catalysts with the higher content of framework aluminum occurs at lower temperatures than for zeolites with the lower aluminum content, as shown in Figure 6. Thus, the increased catalytic activity in the low-temperatures range is possibly attributed to the better reducibility of copper species deposited into zeolites. The catalytic performance of these samples is significantly different at higher temperatures. In the presence of Cu-ITQ-6_30, the NO conversion on the level 97–100% was observed in the range of 275–400 °C, while for Cu-ITQ-6_50 a decrease in the NO conversion, due to the side reaction of ammonia oxidation, occurred at about 325 °C. The selectivity to dinitrogen is higher for Cu-ITQ-6_30. Thus, the sample with the higher surface acidity, Cu-ITQ-6_30, is a more effective catalyst in the high-temperature NH3-SCR process. Similar results were obtained for the series of the Cu-ITQ-36 samples. The catalyst with the higher aluminum content, Cu-ITQ-36_30, presented better catalytic activity at temperatures below 175 °C, possibly due to the better reducibility of copper species deposited into this sample compared to the copper species in the catalysts with the lower aluminum content, Cu-ITQ-36_50, as shown in Figure 6. The NO conversion in the presence of Cu-ITQ-36_30 increased to about 400 °C, reaching 97%, while in the case of Cu-ITQ-6_50, the NO conversion increased to 93% at 300 °C and then decreased due to the side reaction of ammonia oxidation, as shown in Figure 7. Thus, in this case, the acid properties of the catalysts also play a very important role in the high-temperature NH3-SCR process. The comparison of the results, presented in Figure 7, shows that the catalysts based on ITQ-6 are more active and selective than the analogous catalysts of the ITQ-36 series. On the other hand, the comparison of the results obtained for the FER series and their intercalated and delaminated forms shows that the most significant differences are related to the lower NO conversion for Cu-FER_30 and Cu-FER_50 in the low-temperature range, compared to the catalysts of the Cu-ITQ-6 and Cu-ITQ-36 series. The results of UV-vis DRS, as shown in Figure 5, and H2-TPR, as shown in Figure 6, show a significantly higher contribution of highly dispersed copper species, mainly monomeric copper cations, in the catalysts of the Cu-ITQ-6 and Cu-ITQ-36 series, while the presence of more aggregated copper oxide species was postulated for the samples of the Cu-FER series. Such highly dispersed copper species, which can be reduced at relatively low temperatures, are possibly needed for the low-temperature conversion of NO. One of the possible mechanisms of the low-temperature NH3-SCR is based on fast-SCR (2NH3 + NO + NO2 → 2N2 + 3H2O); however, for this reaction NO2 is needed.
2.3. Catalytic Studies of NO to NO$_2$ Oxidation

The results of the catalytic NO to NO$_2$ oxidation in the presence of the studied catalysts are presented in Figure 8. The normalized intensity of the FTIR band at 1593 cm$^{-1}$ was used for the analysis of NO$_2$ formation. It can be seen that the intensity of the band characteristic of NO$_2$ is significantly higher for the Cu-ITQ-6 and Cu-ITQ-36 catalysts than for the Cu-FER samples. Moreover, NO oxidation in the presence of delaminated and intercalated zeolites started at temperatures significantly lower compared to the Cu-FER catalysts. The maximal NO to NO$_2$ conversions, determined from the normalized intensity of the 1912 cm$^{-1}$ band (characteristic of NO), are about 24% for the Cu-FER catalysts, 43–45% for the Cu-ITQ-6 samples and about 33% for both catalysts of the Cu-ITQ-36 series. Thus, the catalytic activity of the studied zeolitic samples in NO to NO$_2$ oxidation very well correlates with the activity order of these samples in the low- and middle-temperature NH$_3$-SCR.
The ammonia oxidation in the presence of the catalysts of the Cu-FER series started at about 300 °C. Catalysts 2020, 10, x FOR PEER REVIEW

The ammonia oxidation to dinitrogen, as shown in Figure 9, which explains the high selectivity towards dinitrogen, as shown in Figure 7.

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The ammonia oxidation to dinitrogen, as shown in Figure 9, which explains the high selectivity towards dinitrogen, as shown in Figure 7.

2.4. Catalytic Studies of Ammonia Oxidation (AMOx)

Analysis of the NH3-SCR results, as shown in Figure 7, shows that the high-temperature efficiency of this process is determined by the side reaction of direct ammonia oxidation (AMOx). Catalysts for high-temperature NH3-SCR should be active in NO reduction with ammonia and simultaneously inactive in ammonia oxidation. The comparison of the NH3-SCR results, obtained for Cu-ITQ-6_30 and Cu-ITQ-36_30, with the results obtained for Cu-ITQ-6_50 and Cu-ITQ-36_50, shows that ammonia oxidation possibly starts at temperature significantly lower for the catalysts with the larger Si/Al ratio. On the other side, the results of NH3-TPD, as shown in Figure 4, show that ammonia is more effectively chemisorbed on the surface of Cu-ITQ-6_30 and Cu-ITQ-36_30 than on Cu-ITQ-6_50 and Cu-ITQ-36_50, which indicates that ammonia coordinated on copper species is effectively activated for the NH3-SCR and not activated for direct oxidation. In order to verify the activity of the studied catalysts in the side reaction of ammonia oxidation, which limits the efficiency of this process is determined by the high-temperature NH3-SCR.

The ammonia oxidation in the presence of the Cu-FER series started at about 300 °C and increased to the level of 80–90%. Cu-FER_30 presented higher activity and selectivity to dinitrogen at higher temperatures, catalytic tests of ammonia oxidation were performed, as shown in Figure 9. The ammonia oxidation in the presence of the Cu-FER series started at about 300 °C and increased to the level of 80–90%. Cu-FER_30 presented higher activity and selectivity to dinitrogen than Cu-FER_50. More significant differences in the results of ammonia oxidation were found for the catalysts of the Cu-ITQ-6 and Cu-ITQ-36 series. The ammonia conversion in the presence of the zeolitic catalysts with the higher Si/Al ratio, Cu-ITQ-6_50 and Cu-ITQ-36_50, started at lower temperatures compared to the samples with the lower Si/Al ratio, Cu-ITQ-6_30 and Cu-ITQ-36_30, which is in full agreement with the results of the NH3-SCR tests, as shown in Figure 7. Thus, it is supposed that stronger acid sites in the delaminated (Cu-ITQ-6) and intercalated (Cu-ITQ-36) zeolitic samples decrease their activity in ammonia oxidation and, therefore, extend the temperature window of the effective NH3-SCR process to higher temperatures. Another important issue is a very high selectivity of ammonia oxidation to dinitrogen, as shown in Figure 9, which explains the high selectivity towards dinitrogen in the NH3-SCR process, and also in the temperature range characteristic of the side reaction of direct ammonia oxidation, as shown in Figure 7.
Figure 9. Results of AMOx tests for FER (a), ITQ-6 (b) and ITQ-36 (c) samples modified with Cu.
For the most promising catalyst, Cu-ITQ-6_30, selected on the basis of the catalytic tests of the NH$_3$-SCR reaction, as shown in Figure 7, the stability test in NH$_3$-SCR was done at 275 °C for 15 h, as shown in Figure 10. The NO conversion changed in the range of 92.5–94%, while selectivity towards dinitrogen was in the range of 96–99%. Thus, the Cu-ITQ-6_30 presented high catalytic stability in the NH$_3$-SCR reaction.

3. Experimental

3.1. Catalysts Preparation

3.1.1. Synthesis of PREFER and FER

Zeolite precursor materials, PREFER, with different intended Si/Al molar ratios, were prepared using fumed silica Aerosil 200 (Evonik Industries AG, Essen, Germany) as a silicon source, hydroxy(oxo)alumane Catapal B (Sasol, Johannesburg, South Africa) as an aluminum source, 4-amino-2,2,6,6-tetramethylpiperidine (R, Fluka, Germany) as a structure directing agent, NH$_4$F (98%, Sigma-Aldrich, St. Louis, MO, USA), HF (49.8%, Sigma-Aldrich, St. Louis, MO, USA) and distilled water in the molar ratios of 1 SiO$_2$: x Al$_2$O$_3$: 1 R: 1.5 NH$_4$F: 0.5 HF: 10 H$_2$O, where x = 0.015 and 0.01 for the Si/Al molar ratios of 30 and 50, respectively. The obtained gels were mixed in an autoclave at 135 °C for 3 days. The resulting products were filtered, washed with distilled water and dried overnight at 60 °C. Three dimensional (3D) ferrierite zeolites (FER) were obtained by the calcination of PREFER at 600 °C for 6 h. The FER zeolites with the intended Si/Al molar ratios of 30 and 50 are denoted as FER_30 and FER_50, respectively.

3.1.2. PREFER Swelling

For the swelling of PREFER, an aqueous solution of hexadecyltrimethylammonium bromide (CTMABr, ≥99%, Acros Organics, Geel, Belgium) and tetrapropylammonium bromide (TPABr, 98% Sigma-Aldrich, St. Louis, MO, USA) were used. The solution was prepared by dissolving
25 g CTMABr/OH (CTMABr with 50% Br\(^-\) ions exchanged for OH\(^-\) anions) and 12 g TPABr/OH (TPABr with 30% Br\(^-\) ions exchanged for OH\(^-\) anions) in 113 g distillated water. The PREFER sample of 5 g was dispersed in this solution and obtained slurry and was stirred under reflux at 95 °C for 16 h.

### 3.1.3. Synthesis of ITQ-6

First part of the slurry, containing swelled PREFER, was sonicated in an ultrasound bath (50 W, 40 kHz) for 1 h. Then, the mixture was acidified to pH = 3.0 with concentrated HCl (high purity grade, Honeywell, Charlotte, NC, USA), separated by centrifugation (12,000 rpm, 20 min) and washed with distillated water. Finally, the solid product was calcined at 600 °C for 6 h, resulting in delaminated ITQ-6 zeolite. The ITQ-6 zeolites with the intended Si/Al molar ratios of 30 and 50 are denoted as ITQ-6_30 and ITQ-6_50, respectively.

### 3.1.4. Synthesis of ITQ-36

The second part of the slurry, containing swelled PREFER, was mixed with tetraethyl orthosilicate (TEOS, ≥99%, Merck KGaA, Darmstadt, Germany), used as a pillaring agent, with a swollen PREFER:TEOS weight ratio of 1:5. The obtained mixture was intensively stirred at 90 °C for 16 h in dinitrogen atmosphere. Then, the solid product was separated by filtration, washed with ethanol and dried overnight at 60 °C. In the next step, solid material was hydrolyzed by its dispersing in distillated water in the weight ratio of 1:10. The obtained slurry was kept at 100 °C for 10 h, then the solid product was separated by filtration, washed with distillated water, dried overnight at 60 °C and finally calcined at 600 °C for 6 h, resulting in ITQ-36 zeolite. The ITQ-36 zeolites with the intended Si/Al molar ratios of the zeolite layers of 30 and 50 are denoted as ITQ-36_30 and ITQ-36_50, respectively.

### 3.1.5. Modification of the Zeolitic Samples with Copper

The obtained samples of the FER, ITQ-6 and ITQ-36 series were modified with copper by ion-exchange method. The zeolitic samples were dispersed in 0.06 M aqueous solution of Cu(CH\(_3\)COO)\(_2\)·H\(_2\)O with a zeolite:liquid ratio of 10 g/1000 g. The obtained slurry was stirred at 80 °C for 6 h, then the solid sample was separated by filtration, washed with distillated water, dried overnight at 60 °C and finally calcined at 550 °C for 8 h. The samples modified with copper are denoted: Cu-FER_30 and Cu-FER_50; Cu-ITQ-6_30 and Cu-ITQ-6_50; Cu-ITQ-36_30 and Cu-ITQ-36_50.

### 3.2. Characterization of the Samples

The textural properties of the samples were determined by N\(_2\) sorption at −196 °C using the 3Flex (Micrometrics, Norcross, GA, USA) automated gas adsorption system. Prior to the analysis, the samples were outgassed under a vacuum at 350 °C for 24 h. The specific surface area was determined using the BET (Braunauer–Emmett–Teller) model. Micropore size distribution was determined using the Horvath–Kawazoe model, while mesopore volume and mesopore area were enumerated using the BJH (Barrett–Joyner–Halenda) model. The micropore volume (at p/p\(_0\) = 0.98) and specific surface area of the micropores were calculated using the t-plot method. The X-ray diffraction patterns of the samples were obtained using a Bruker D2 Phaser diffractometer (Bruker, Billerica, MA, USA). The measurements were performed in the 2θ range of 2–40° with a step of 0.02° and a counting time of 1 s per step. The chemical compositions of the samples were analyzed by inductively coupled plasma—optical emission spectrometry (ICP-OES) using iCAP 7400 instrument (Thermo Scientific, Waltham, MA, USA). The solid samples were dissolved in a mixture of hydrofluoric (high purity grade, Honeywell, Charlotte, NC, USA), hydrochloric (high purity grade, Honeywell, Charlotte, NC, USA) and nitric acid (high purity grade, Honeywell, Charlotte, NC, USA) solution, assisted by microwave radiation using the Ethos Easy system (Milestone, Sorisole, Italy). The chemical analysis was done with respect to silicon, aluminium and copper content in the samples. The coordination and aggregation of the copper species were studied by UV-vis-DR spectroscopy. The measurements were performed using an Evolution 600 spectrophotometer (Thermo Scientific, Waltham, MA, USA) operating in the range of 200–900 nm.
with a resolution of 2 nm. The reducibility of the samples was studied by temperature-programmed reduction using H₂ as a reducing agent (H₂-TPR). The measurements were carried out in a fixed-bed quartz microreactor system in the temperature range of 100–700 °C, with a linear heating rate of 10 °C/min in a flow of gas mixture (5 mL/min) containing 5.0 vol.% of H₂ diluted in argon. The flow of gas mixture was monitored by mass flow controller (Brooks Instrument, Hatfield, PA, USA). Before H₂-TPR runs, each sample (50 mg) was outgassed in a flow of pure argon at 500 °C for 30 min. The hydrogen conversion was analyzed using a thermal conductivity detector—TCD (VICI, Houston, TX, USA).

3.3. Catalytic Tests

The copper modified zeolitic samples were tested as catalysts of the selective catalytic reduction of NO with ammonia (NH₃-SCR) as well as the catalytic oxidation of ammonia (AMOx) and NO to NO₂ oxidation, which are reactions associated with the NH₃-SCR process. Prior to the catalytic tests, each sample (100 mg) of catalyst was placed in a fixed-bed quartz microreactor and outgassed in a flow of pure helium at 550 °C for 30 min. Then, after cooling the reactor to 100 °C, the flow of pure helium was changed for the flow of a gas mixture containing: 0.25 vol.% NO; 0.25 vol.% NH₃; 2.5 vol.% O₂; 97.0 vol.% He (total flow rate of 40 mL/min) in the case of the NH₃-SCR tests— or 0.5 vol.% NH₃; 2.5 vol.% O₂; 97.0 vol.% He (total flow rate of 40 mL/min) in the case of the AMOx test. Both in the NH₃-SCR and AMOx tests, the reactants and product concentrations were continuously measured using a quadrupole mass spectrometer, QMS (PREVAC, Rogów, Poland), connected directly to the reactor outlet. In the case of NO to NO₂ oxidation, the gas mixture containing 0.5 vol.% NO and 2.5 vol.% O₂ diluted in helium (total flow rate of 40 mL/min) was used. For the analysis of the NO and NO₂ concentrations in the reaction mixture, before and after the reactor, a FTIR spectrometer Nicolet iS5 (Thermo Scientific, Waltham, MA, USA) equipped with a gas cell of 10 cm length was used. The spectra were recorded in the wavenumber range of 625–4000 cm⁻¹ with a resolution of 4 cm⁻¹ (averaged seven scans for each spectrum) every 9 s. The bands at 1593 and 1912 cm⁻¹ were used for the detection of NO and NO₂, respectively. All catalytic tests were conducted with a linear temperature increase of 10 °C/min. For the most promising catalyst, an additional isothermal stability test of NH₃-SCR was performed. The test was conducted at 275 °C for 15 h with the same composition and total flow of the reaction mixture as in polythermic catalytic tests.

4. Conclusions

Ferrierite and its delaminated (ITQ-6) and silica intercalated (ITQ-36) forms with different Si/Al ratios were synthesized and, after copper deposition, studied as NH₃-SCR catalysts. It was shown that copper modified ITQ-6 and ITQ-36 zeolites effectively operated in NH₃-SCR in much broader temperature ranges compared to the Cu-FER catalysts. For the most active catalyst, Cu-ITQ-6_30, the NO conversion above 90% was measured in the range of 250–400 °C with the selectivity to dinitrogen on the level of 98–99%. High catalytic activity of the delaminated and silica intercalated zeolites in low-temperature NH₃-SCR was related to the better activity of highly dispersed copper species—mainly monomeric copper cations—which were deposited into these series of zeolites. Such copper species are more active in the NO to NO₂ oxidation compared to aggregated copper oxide species present in microporous ferrierites. NO₂ was necessary for fast-SCR (2NH₃ + NO + NO₂ → 2N₂ + 3H₂O), which is one of the main pathways of low-temperature NH₃-SCR. In the high temperature NH₃-SCR, delaminated and silica intercalated zeolites with higher aluminum content presented better catalytic performance compared to low-alumina Cu-ITQ-6_50 and Cu-ITQ-36_50. This effect was related to the higher activity of low-alumina zeolites in the side process of direct ammonia oxidation, which decreased the efficiency of high-temperature NH₃-SCR. Thus, the catalysts with a higher concentration of acid sites are possibly able to more effectively protect chemisorbed ammonia against oxidation. The stability NH₃-SCR test for Cu-ITQ-6, conducted at 275 °C for 15 h, shows only small changes in the NO conversion and selectivity towards dinitrogen, proving the high
stability of this catalyst under reaction conditions. Taking into account the results of the presented studies, it could be concluded that delaminated Cu-ITQ-6_30 is a very promising catalyst for the low-temperature NH$_3$-SCR reaction.

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