Simultaneous $\text{SO}_2/\text{NO}_x$ Abatement Using Zeolite-Supported Copper

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ABSTRACT

The bulk of the results from this project, besides being found in the research reports, are in the form of two theses presented for advanced degrees by two different students at two different institutions. Francis E. Porbeni is a Master's Degree student in Chemistry at Clark Atlanta University and has prepared his Master's Thesis entitled *Simultaneous SO$_2$ Oxidation and NO Decomposition over Copper Oxide on γ-Alumina Catalysts: An Infrared Diffuse Reflectance Study*. Sumit Rao, a Ph.D. student in Chemical Engineering at the Georgia Institute of Technology has defended his thesis entitled *Role of Copper Ensemble Size in Silica and Zeolite Supported Catalysts for Nitric Oxide Decomposition*. These two documents, which will be forwarded when final copies are available, provide the details of the bulk of the research accomplished over the duration of this project. These two documents are summarized, and other results from the project are given, below.

SUMMARY OF RESEARCH ACCOMPLISHMENTS

*Research Presented in the Thesis of Francis E. Porbeni*

Preparation of Novel CuO/γ-Alumina Catalysts using Non-Aqueous Impregnation.

Non-aqueous impregnation using acetonitrile as the solvent and copper acetylacetonate as the copper precursor yield intact complexes on the support, apparently well-dispersed over the surface. The adsorption isotherm for these complexes is linear over the measured concentration range, and elemental analysis and infrared spectroscopy both indicate that the complex is probably intact on the surface before calcination. After calcination, the copper atoms act chemically as copper aluminate, as observed by both SO$_2$ and NO chemisorption.

SO$_2$ Adsorption and Reaction on CuO/γ-Alumina.

SO$_2$ adsorbs on the CuO/γ-alumina sorbent at both copper and aluminum sites. Adsorption at the aluminum sites leads to the formation of both aluminum sulfite and sulfate species, even in the absence of added oxygen. The formation of aluminum sulfate may be due to oxygen spillover effects. The adsorption efficiency increases with temperature up to at least 300 °C. This is in contrast to the situation on pure aluminum oxide or sodium-doped alumina for which 150 °C yielded the highest adsorption efficiency.

Upon oxidation, the adsorbed SO$_2$ is converted to copper sulfate, regardless of its initial oxidation state or associated cation. The exception to this is found at the lowest copper loadings, where adsorbed SO$_2$ may not be able to “reach” surface copper species.

NO Adsorption on CuO/γ-Alumina and on Sulfated CuO/γ-Alumina.

NO adsorbs on the CuO/γ-alumina surface to give rise to three different species. One species corresponds to the complex formed between the surface Cu species, as Cu$^{2+}$, and neutral NO. This species gives rise to an absorption band virtually identical to that observed by others for NO adsorbed on copper aluminate. Additionally, two other bands are formed. A very weak band at approximately 1700 cm$^{-1}$ has been assigned as being due to the formation of Cu$^+$/NO$^-$ species.
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A medium feature at 1640 - 1627 cm\(^{-1}\) is assigned as corresponding to the formation of Cu\(^{2+}\)-ONO\(^{\cdot}\). This latter feature is by far the most intense feature in the spectrum, and is indicative of the participation of the surface Cu species in a redox reaction, leading to the formation of NO\(_2\) from the adsorbed NO. When the surface is sulfated, i.e. by exposure to SO\(_2\) and O\(_2\) at high temperature, the adsorption chemistry of NO is changed dramatically. NO again adsors at the Cu\(^{2+}\) surface sites, but the formation of the Cu\(^{+}\) species is almost eliminated. Only a small contribution from the Cu\(^{2+}\)-ONO\(^{\cdot}\) species is observed in the infrared spectrum, and the band due to the Cu\(^{+}\)/NO\(^{-}\) species is not evident at all. This indicates that sulfation of the surface, locks the Cu\(^{2+}\) species into the +2 oxidation state, and precludes participation of copper species in surface redox chemistry, which is critical if the NO reduction reaction is to occur.

**Research Presented in the Thesis of Sumit N. Rao**

**The Role of the Formation of Cu\(^{2+}\)-OH\(^{-}\) Complexes.**

The state of the art catalyst for NO reduction involves the batch impregnation of ZSM-5 zeolite with copper acetate in aqueous solution. Much discussion has been carried out in the literature regarding the active form of copper in the zeolite, whether the active species is Cu\(^{2+}\) or Cu\(^{+}\), or if the coordination number of the copper is the important parameter, or whether the number of copper atoms in close proximity is important for NO reduction activity. The copper is ion-exchanged into the zeolite, and as such, if it is to reside as Cu\(^{2+}\) in the zeolite, it must find a site where two aluminum ions are close together, so that the local charge in that region of the zeolite framework can compensate its +2 charge.

Some speculation has involved the formation of a Cu\(^{2+}\)-OH\(^{-}\) species, with overall charge +1, which would not need to find two aluminum sites close together, but which could ion-exchange at any single aluminum site to form a neutral complex. Two different experiments were carried out to evaluate the effects of reducing the OH\(^{-}\) solution concentration during impregnation. The first experiment used added acetic acid to shift the water dissociation equilibrium back towards H\(_2\)O and away from the formation of OH\(^{-}\) in solution. Catalysts prepared using an impregnation solution containing added acetic acid showed less activity overall and less activity per copper atom than the state of the art catalyst. The second experiment involved the use of acetonitrile as the solvent for the impregnation of ZSM-5 with copper acetate. The catalyst produced using this solution was almost completely inactive.

**The Role of Cu Atoms Next to Two Aluminum Atoms.**

As mentioned above, speculation regarding the nature of the active copper species for NO reduction has involved whether the copper is sited next to one or two aluminum atoms in the zeolite framework. Experiments were carried out using non-aqueous impregnation with copper ethylenediamine as a precursor for Cu\(^{2+}\). This complex in acetonitrile solution does not dissociate and certainly there are no Cu\(^{2+}\)-OH\(^{-}\) species formed, so that the copper atom, when it does adsorb onto the surface, must find two aluminum atoms in the zeolite framework to compensate its +2 charge. Copper ethylenediamine is a stable source for Cu\(^{2+}\) and there is virtually no possibility that the copper would adsorb as Cu\(^{+}\). Thus, catalysts prepared in this way are a known source of copper in the neighborhood of two framework aluminum atoms. These catalysts yield activity per