the surface acidity seems to play an important role for the almost lost by the addition of decreased with the increase of added amount of NaOH and was examined. As shown in Figure 9, the activity was decreased with the increase of added amount of NaOH and was finally lost. Since the curve of the activity decrease in the isomerization (Figure 10) is similar to that in NO reduction (Figure 9), the acid sites which are active for the reduction seem to be mainly Brønsted type. Takagi et al. (1979) reported from an infrared study that NH$_4^+$ formed during the reaction of NO with NH$_3$ on the surface of VO$_2$-Al$_2$O$_3$ is a reactive species. In the present case, the Brønsted acid sites evidenced by infrared study of adsorbed pyridine and the poisoning experiment with alkali for the isomerization of cyclopropane may enhance the activity by facilitating the adsorption of NH$_3$ and the formation of NH$_4^+$. It is concluded that the increase of acidity on the addition of SO$_2^-$ and on mixing with MoO$_3$ is vitally important for the activity enhancement.

**Literature Cited**


Received for review August 29, 1980

Accepted January 29, 1981

This work was supported by a Grant-in-Aid (No. 203001) for Scientific Research from the Ministry of Education, Japan.

**Vanadium Oxide Monolayer Catalysts. 4. The Catalytic Oxidation of Carbon Monoxide**

Fred Roozeboom,*^1^ A. J. van Dillen,^2^ John W. Geus,^2^ and Paul J. Gellings^1^  

Twente University of Technology, Department of Chemical Technology, P.O. Box 217, 7500 AE Enschede, The Netherlands, and State University of Utrecht, Inorganic Chemistry Department, Croesestraat 77A, 3522 AD Utrecht, The Netherlands

The oxidation of CO over vanadium oxide, unsupported and supported as a monolayer (or monomolecular dispersion) on γ-Al$_2$O$_3$, CeO$_2$, TiO$_2$, and ZrO$_2$ was studied between 600 and 800 K in a continuous flow reactor. From catalytic and structural studies it was concluded that the activity of unsupported V$_2$O$_5$ is determined by its crystallinity, while that of supported vanadium oxide is affected by interactions with the support. An additional thermogravimetric investigation supports the conclusion that on all catalysts CO reacts to CO$_2$ in a reduction-oxidation mechanism in which the reduction of the catalyst is the rate-determining step.

**Introduction**

The interaction between two adhering oxides can in general be expected to be stronger than that between a metal and an oxide. Consequently, the catalytic properties of an active oxide can be expected to be affected appreciably by the support. Because vanadium oxides are used extensively in catalytic oxidation (Dadyburjor et al., 1979) investigation of the effect of the support on its catalytic properties is worthwhile.

Recently, Goldwasser and Trimm (1979) studied the activity of vanadia supported on γ-Al$_2$O$_3$, ZrO$_2$, SiO$_2$ for
the oxidation of CO and observed a decrease in activity in this order. It is interesting to extend the range of supports and to compare the activity of supported with that of unsupported vanadia catalysts.

In the oxidation of organic molecules it has been observed (Andreikov et al., 1973; Kera and Kuwata, 1979; Shaporskaya et al. 1970; Stasevich et al., 1969; Simard et al., 1955) that active vanadia catalysts contain vanadium ions with a valency lower than 5. The partial reduction of active vanadia catalysts has been attributed to a Mars–van Krevelen or reduction–oxidation mechanism (Hughes and Hill, 1955; Hirota et al., 1968; Kera and Hirota, 1969) in which the catalyst is alternately reduced and reoxidized according to the reaction scheme

\[
M + O_{\text{act}} \rightarrow "\text{MO}" + V_0 \\
V_0 + 1/2O_2 \rightarrow O_{\text{act}}
\]

where M is the molecule oxidized to products symbolized by "MO", O_{\text{act}} is the active oxygen species of the catalyst, and V_0 is a surface oxygen vacancy.

Other evidence for this type of mechanism is that, in the absence of gaseous oxygen, the oxidation proceeds with the same selectivity as in its presence, though only for a short time. The evidence concerning the oxidation of H_2 and CO is less clear cut (Van Dillen, 1977). Some authors (Marshneva et al., 1972) have put forward evidence pointing to an associative mechanism, where lattice oxygen does not participate, while others (Hirotta et al. 1968; Kera and Hirota, 1969; Hughes and Hill, 1955) prefer a reduction–oxidation mechanism.

In this work the effects of γ-Al_2O_3, ZrO_2, TiO_2, and CeO_2 as supports on the activity of vanadia in the oxidation of CO were studied. Because the interaction between vanadia and the support may be influenced by the valence state of the vanadium, these activities have also been studied after an intermediate reduction (by H_2) and reoxidation treatment. Before this reduction–reoxidation treatment the supports were covered by approximately a monolayer of vanadia (Roozeboom et al., part 1, 1979; part 2, 1978; part 3, 1980, part 5, 1981).

As evidence has been published (Tilley and Hyde, 1970) that the crystal size and defect structure of vanadium pentoxide affect its reducibility, two types of unsupported V_2O_5 were also studied.

Comparison of the catalytic properties of the different unsupported and supported vanadia catalysts made it possible to discriminate between effects of particle size and defect structure on the one hand, and those of a chemical interaction with the supports on the other.

To investigate whether a connection exists between the catalytic activity in CO oxidation and the reducibility of the vanadia, the catalysts have also been studied by temperature programmed reduction.

**Experimental Section**

**Materials.** Two different unsupported V_2O_5 catalysts were used: a Baker product (analytical grade), which was annealed at elevated temperatures, and a catalyst prepared by decomposition of NH₄VO₃ (Merck, analytical grade). This was performed at a relatively low temperature by heating at 600 K for 2 h in a stream of air.

The oxidic carriers, viz., γ-Al_2O_3, CeO_2, TiO_2, and ZrO_2 have previously been described (Roozeboom et al., 1978, 1979). All gases (Loosco, Amsterdam) were dried by BDH molecular sieve Type 4A. Nitrogen and argon were deoxygenated by BTS-catalyst R-3-11 (BASF). Hydrogen was deoxygenated by a Baker "deoxygen" catalyst (Pd/Al_2O_3).

**Catalyst Preparation.** The supported vanadia catalysts were prepared by passing a freshly prepared aqueous solution of 1 wt % ammonium vanadate (acidified with HNO_3 to pH 4) through a bed of particles of the support (0.3–0.6 mm diameter). For further details see part 1 of this series (Roozeboom et al., 1979). The catalysts were dried at 383 K (17 h) and calcined at 723 K (2 h).

**X-ray Fluorescence.** The vanadium content of the catalyst samples was determined with a Philips 1410 X-ray spectrometer by X-ray fluorescence (Van Willigen et al., 1971).

**X-ray Diffraction.** The two types of unsupported V_2O_5 were analyzed with a Philips PW 1050-25 vertical diffractometer.

**Temperature-Programmed Reduction.** TPR profiles of the two unsupported V_2O_5 catalysts were recorded by the procedure described in part 3 (Roozeboom et al., 1980).

**Thermogravimetry (TG).** This was carried out by means of a Mettler Thermoanalyzer with which weight changes (accuracy: 0.02 mg) were continuously recorded. In order to eliminate gas flow and buoyancy effects, blank runs with an empty crucible were performed prior to each measurement. In most cases approximately 70 mg of catalyst was brought into an inert alumina sample holder (Type ME-93523). An argon flow of 50 mL NTP min⁻¹ was passed through the housing of the balance as a protective measure. Oxygen and hydrogen were fed into the furnace surrounding the specimen.

Each sample successively underwent three treatments during which its weight was recorded while the temperature was increased from room temperature to 830 K at a rate of 5 K min⁻¹. The treatments were: (i) dehydration in an oxygen flow (10 mL NTP min⁻¹); (ii) after flushing with argon to remove the oxygen: reduction in a hydrogen flow (10 mL NTP min⁻¹); (iii) after flushing with argon: reoxidation in an oxygen flow (10 mL NTP min⁻¹). If required, the temperature in step (ii) was kept at the maximum value of 830 K in order to complete reduction.

**Catalytic Measurements.** The oxidation of CO was studied in a conventional flow reactor as described elsewhere (Van Dillen, 1977). The reactor was kept at a pre-set temperature within ±1 K. The gas feed usually contained 1 vol % CO, 1 vol % O_2, and 98 vol % N_2 at a total gas flow of 50 mL NTP min⁻¹. The determination of the CO₂ formed was based on the change in electric conductivity of an aqueous NaOH solution, using complete oxidation of CO over a BTS catalyst as a reference for 100% conversion.

**Experimental Results**

**Thermogravimetry.** One of the questions to be answered in establishing a mechanism for catalytic oxidation reactions concerns the relative ease of reduction and reoxidation of the catalyst. Therefore, a thermogravimetric study was performed on all supported vanadia catalysts used in this investigation. As an example Figure 1 gives the TG curves obtained during reduction (after previous dehydration) and subsequent reoxidation of vanadia on ZrO₂. These are typical for the curves obtained with all catalysts. All TG-data are collected in Table I and it is clear that the reduction of all catalysts sets in at considerably higher temperatures than their reoxidation. This is in agreement with the results of Van Dillen (1977) on unsupported V_2O_5 (a Baker product) for which reduction started at 733 K and reoxidation already at 393 K.

The weight changes show that the vanadium is reduced to the trivalent state and that the reoxidation restores the original pentavalent state. Only on γ-Al_2O_3 the weight loss would imply reduction to the quadrivalent state, which, however, is not in agreement with TPR results reported.
Table I. Results of Thermogravimetric Experiments

<table>
<thead>
<tr>
<th></th>
<th>(\gamma\text{-Al}_2\text{O}_3)</th>
<th>(\text{V}/\text{CeO}_2)</th>
<th>(\text{V}/\text{TiO}_2)</th>
<th>(\text{V}/\text{ZrO}_2)</th>
</tr>
</thead>
<tbody>
<tr>
<td>wt % V (XRF analysis)</td>
<td>6.8</td>
<td>5.3</td>
<td>1.7</td>
<td>4.3</td>
</tr>
<tr>
<td>starting weight, mg</td>
<td>79.85</td>
<td>76.85</td>
<td>71.58</td>
<td>74.86</td>
</tr>
<tr>
<td><strong>Dehydration</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>weight loss, mg</td>
<td>4.75</td>
<td>2.56</td>
<td>1.58</td>
<td>5.26</td>
</tr>
<tr>
<td><strong>Reduction</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>temperature of onset of reduction, K</td>
<td>633</td>
<td>633</td>
<td>593</td>
<td>453</td>
</tr>
<tr>
<td>weight loss, mg</td>
<td>0.77</td>
<td>1.43</td>
<td>0.54</td>
<td>1.06</td>
</tr>
<tr>
<td>wt % (from TG-analysis)</td>
<td>3.1</td>
<td>5.9</td>
<td>2.4</td>
<td>4.5</td>
</tr>
<tr>
<td><strong>Reoxidation</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>temperature of onset of reoxidation, K</td>
<td>&lt;303</td>
<td>&lt;303</td>
<td>&lt;303</td>
<td>&lt;303</td>
</tr>
<tr>
<td>temperature of complete reoxidation, K</td>
<td>613</td>
<td>513</td>
<td>673</td>
<td>583</td>
</tr>
<tr>
<td>weight change, mg</td>
<td>0.73</td>
<td>1.45</td>
<td>0.60</td>
<td>1.04</td>
</tr>
</tbody>
</table>

\(\text{a} \) Based on the assumption that V\(^{2+}\) is reduced to V\(^{3+}\) (Roozeboom et al., part 3, 1980).

Figure 1. TG curves for vanadium oxide on \(\text{ZrO}_2\): a. weight decrease during reduction; b. weight increase during reoxidation. (Note: the starting weight of the sample is equal for both cases only for arbitrary computer programming reasons).

Temperature-Programmed Reduction. Because crystallite size has been shown to play an important role in the reduction of vanadia (Tilley and Hyde, 1970) and to allow a comparison with earlier TPR measurements on supported catalysts (Roozeboom et al., part 3, 1980), TPR measurements were performed on pure \(\text{V}_2\text{O}_5\). These were done with well-annealed \(\text{V}_2\text{O}_5\) and with \(\text{V}_2\text{O}_5\) freshly prepared by thermal decomposition of \(\text{NH}_4\text{VO}_3\). The resulting TPR curves in Figure 2 show that the reduction of the well-annealed \(\text{V}_2\text{O}_5\) starts at 725 K, whereas that of the freshly prepared \(\text{V}_2\text{O}_5\) starts at 675 K. This is only slightly higher than that of a monolayer of \(\text{V}_2\text{O}_5\) on \(\gamma\text{-Al}_2\text{O}_3\) (Roozeboom et al., part 3, 1980).

The X-ray diffraction patterns shown in Figure 3 show that in the freshly prepared \(\text{V}_2\text{O}_5\) a very small amount of suboxide is present (both \(\text{V}_2\text{O}_4\) and \(\text{V}_6\text{O}_{13}\) have their strongest XRD line near 2\(\theta\) = 28°). This indicates that the freshly prepared \(\text{V}_2\text{O}_5\) contains a large number of defects and these may well cause the greater ease of reoxidation.

Catalytic Studies. Table II summarizes the results of the catalytic experiments. In all cases the order in CO was found to be 1 or nearly so. The order in oxygen was zero for all catalysts. From the measured conversions the rate constants were calculated by integration. In Figure 4 some typical Arrhenius plots are shown from which the data in Table II were derived.

Repeated experiments on \(\text{V}_2\text{O}_5\) freshly prepared from \(\text{NH}_4\text{VO}_3\) show that after this has been heated to sufficiently high temperatures the activity drops to that of the well-annealed \(\text{V}_2\text{O}_5\) (Baker) with an increase in activation energy and preexponential factor.
Table II. Results on CO Oxidation, Gas Feed: 1 vol % CO, 1 vol % O₂, 98 vol % N₂; Flow 50 mL NTP min⁻¹

<table>
<thead>
<tr>
<th>Catalyst</th>
<th>Vₜ wt %</th>
<th>E_A(±0.5) kcal mol⁻¹</th>
<th>kₚ⁺ s⁻¹ m⁻³ cat.</th>
<th>S_BET m⁻² g⁻¹</th>
<th>order</th>
<th>order</th>
<th>T_start K</th>
</tr>
</thead>
<tbody>
<tr>
<td>V₂O₅, stabilized (Baker)</td>
<td>56.0</td>
<td>25.7</td>
<td>7 × 10⁴</td>
<td>8.2</td>
<td>1.0</td>
<td>0</td>
<td>770</td>
</tr>
<tr>
<td>V₂O₅, freshly prepared (decomp. of NH₄VO₃)</td>
<td>56.0</td>
<td>19.0</td>
<td>3 × 10⁷</td>
<td>9.5</td>
<td>1.0</td>
<td>0</td>
<td>750</td>
</tr>
<tr>
<td>V/γ-Al₂O₃ before and after several reduction/reoxidation cycles ²⁹</td>
<td>6.8</td>
<td>18.6</td>
<td>1.5 × 10⁹</td>
<td>80</td>
<td>0.9</td>
<td>0</td>
<td>650</td>
</tr>
<tr>
<td>V/TiO₂ before red./reox. ⁷</td>
<td>1.7</td>
<td>18.6</td>
<td>2 × 10⁴</td>
<td>45</td>
<td>1.0</td>
<td>0</td>
<td>600</td>
</tr>
<tr>
<td>after red./reox. ⁷</td>
<td>12.9</td>
<td>4 × 10⁴</td>
<td>45</td>
<td>550</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>pure TiO₂</td>
<td>12.7</td>
<td>7 × 10⁴</td>
<td>45</td>
<td>620</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>V/ZrO₂ before and after red./reox. ⁷</td>
<td>4.3</td>
<td>14.0</td>
<td>1 × 10⁵</td>
<td>107</td>
<td>1.0</td>
<td>0</td>
<td>520</td>
</tr>
<tr>
<td>pure ZrO₂</td>
<td>14.0</td>
<td>6 × 10⁴</td>
<td>156</td>
<td>570</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>V/CeO₂ before red./reox. ⁷</td>
<td>5.3</td>
<td>15.0</td>
<td>5 × 10²</td>
<td>48</td>
<td>1.0</td>
<td>0</td>
<td>615</td>
</tr>
<tr>
<td>after red./reox. ⁷</td>
<td>15.0</td>
<td>1 × 10⁵</td>
<td>575</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>pure CeO₂</td>
<td>15.0</td>
<td>6 × 10⁴</td>
<td>475</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

² 1 vol % O₂ fixed, 0.25–1.5 vol % CO. ᵇ 1 vol % CO fixed, 0.5–2 vol % O₂. ᶜ Temperature range 600–800 K. ᵈ Temperature at which 2% conversion of CO was observed. ᵉ Calculated from measured conversions (x) according to \(-\ln(1-x) = k_p x^{0.5} E_A/RT\). ƒ This consists of a treatment consisting of reduction by H₂ at 800 K for 1 h, followed by reoxidation by O₂ at 800 K for 1 h. Each reduction and reoxidation was recorded by a hygrometer in order to verify complete reduction.

Figure 4. Arrhenius plots of CO oxidation on: a, well-annealed (△) and freshly prepared (●) V₂O₅; b, V/γ-Al₂O₃, freshly prepared (○), and after reduction by H₂ and reoxidation by O₂ (●); c, V/TiO₂, freshly prepared (○) after reduction by H₂ and reoxidation by O₂ (●) and pure TiO₂ (△).

Irreversible changes in the V₂O₅ monolayers may be caused by reduction, either due to agglomeration of the vanadia or due to dissolution of vanadium ions in the support. Therefore all supported catalysts were submitted to complete reduction by H₂. After reoxidation with oxygen the catalytic activity was measured again. As shown in Table II the catalytic activity of V₂O₅/γ-Al₂O₃ and of V₂O₅/ZrO₂ is not changed by this treatment. V₂O₅/TiO₂, which before reduction/reoxidation behaves similarly to V₂O₅/γ-Al₂O₃, shows a large change in behavior. The activation energy becomes equal to that of the oxidation of CO by pure TiO₂ but the preexponential factor is significantly greater. For V₂O₅ on CeO₂ only the preexponential factor is observed to increase after reduction/reoxidation.

Pure γ-Al₂O₃ shows no catalytic activity in CO oxidation. The other supports show an activity which increases in the order ZrO₂ < TiO₂ < CeO₂.

Discussion

The observed orders of 1 for CO and 0 for O₂ on all catalysts are in agreement with a reduction–oxidation mechanism in which the reduction of the catalyst by CO (reaction 1) is rate determining. This is supported completely by the thermogravimetric results which show that reduction of the catalysts is slow and only starts at a relatively high temperature, whereas reoxidation is fast and sets in already at low temperatures.

Furthermore, it is important to note that the temperatures at which reduction starts in the TG experiments are for all catalysts very close to the temperatures at which CO oxidation sets in.

From the catalytic experiments it emerges that three types of active sites can be distinguished in these catalysts: (1) sites with low activity (high activation energy) on well-annealed V₂O₅; (2) sites with high activity (intermediate activation energy) on freshly prepared V₂O₅ and on...
the V$_2$O$_5$ monolayers on γ-Al$_2$O$_3$ and, before reduction/reoxidation, on TiO$_2$ (3 sites with very high activity (low activation energy) on some supports, with or without dissolved vanadium ions.

1. The results obtained with a well-annealed V$_2$O$_5$ lead to the conclusion that this contains a large number of active sites (high preexponential factor) with a low activity due to the high energy of activation of 25.7 kcal mol$^{-1}$.

2. The freshly prepared V$_2$O$_5$ has a higher overall activity than the well-annealed V$_2$O$_5$ due to a much lower activation energy of 19 kcal mol$^{-1}$ even though the number of active sites is smaller ($k_0 = 3 \times 10^5$ s$^{-1}$ m$^{-2}$).

As is clear from the X-ray diffraction patterns of Figure 3 the well-annealed V$_2$O$_5$ consists of larger crystallites containing less lattice defects as shown by the sharper and higher diffraction maxima. Thus, in agreement with previous work (Tilley and Hyde, 1970) the rate of reduction of V$_2$O$_5$ increases with decreasing crystal size, as shown by the TPR profiles of Figure 2. At the same time this corresponds to a greater activity in CO oxidation.

Evidently the larger number of lattice defects in the smaller crystallites of freshly prepared V$_2$O$_5$ causes an acceleration of the nucleation of a reduced phase. Keeping this catalyst at a sufficiently high temperature leads to recrystallization and hence to annihilation of the defects. The catalytic activity consequently becomes more like that of well-annealed V$_2$O$_5$.

As can be seen from Table II, a monolayer of vanadium oxide on γ-Al$_2$O$_3$ and TiO$_2$ exhibits initially about the same activation energy (18.6 kcal mol$^{-1}$) as poorly crystallized V$_2$O$_5$ (19.0 kcal mol$^{-1}$). With alumina as a support the $k_0$ value is of the same order of magnitude ($10^5$ s$^{-1}$ m$^{-2}$) whereas with titania it is appreciably higher ($2 \times 10^5$ s$^{-1}$ m$^{-2}$). The agreement in $E_A$ indicates that these catalysts also contain very finely divided vanadia species, containing a large number of lattice defects with a high activity. These species are hardly influenced by a specific interaction with the support, as long as they are present as a monolayer. Evidently this monolayer is very stable on γ-alumina as shown by the absence of any effect of (hydrogen-) reduction and reoxidation on the catalytic activity of vanadia on γ-Al$_2$O$_3$.

3. Upon reduction–reoxidation the activity of vanadium oxide on titania becomes characteristic of that of pure titania, though with a higher $k_0$ value. It is probable that, on reduction treatment, vanadium, probably as V$^{4+}$ ions, dissolves in the titania, which brings about the disappearance of V oxide from the surface and explains the increased number of active sites on the titania surface (higher $k_0$ value).

For vanadia on zirconia and ceria the activation energies are equal to those of the supports irrespective of any reduction–reoxidation treatments.

Like titania, ceria is more active than pure V$_2$O$_5$. The activities of the pure supports run parallel to the order of their reducibilities: CeO$_2$ > ZrO$_2$ ≈ TiO$_2$ ≈ γ-Al$_2$O$_3$ as determined from TPR experiments (Roozeboom et al., part 3, 1980).

Deposition of vanadium oxide poisons the activity of ceria. For a reduction–oxidation mechanism this correlates well with the better reducibility of CeO$_2$ as compared with that of V/CeO$_2$ as shown by means of TPR (Roozeboom et al., part 3, 1980).

On reduction–reoxidation some vanadium is probably incorporated in the ceria, thus increasing the uncovered ceria surface, which is more active than V oxide. Since some of the vanadium remains at the surface the $k_0$ value of the pure ceria is not obtained. In the case of vanadia on zirconia a strong interaction may readily cause the formation of a ZrV$_2$O$_7$-like surface compound already during calcination, the stability of which is reflected by the fact that $E_A$ and $k_0$ are not changed by the reduction–reoxidation treatment. Incorporation of vanadium ions into the zirconia lattice leads to an increase in the number of active sites, as observed above with titania.

The above interpretation is in agreement with the structural changes occurring on heat treatment of the monolayer type catalysts reported previously (Roozeboom et al., part 3, 1980): (i) on γ-Al$_2$O$_3$ the vanadia monolayer structure does not change markedly; (ii) on TiO$_2$ a solid solution is formed; (iii) on ZrO$_2$ there is formation of Zr$_2$V$_2$O$_7$ and of a solid solution; (iv) on CeO$_2$ small amounts of V surface compounds are present. This correlation between the spectroscopic observations of the structural behavior on the one and the catalytic behavior on the other hand is summarized schematically in Figure 5.

In a recent paper, Goldwasser and Trimm (1979) studied the influence of the support for the CO oxidation over vanadium oxide supported on γ-Al$_2$O$_3$, SiO$_2$, and ZrO$_2$. They also concluded that the nature of the support affects the adsorption of gases and the catalytic activity. They reported the activities to decrease in the order V/γ-Al$_2$O$_3$ > V/ZrO$_2$ > V/TiO$_2$ > V/SiO$_2$, whereas in this study the order V/ZrO$_2$ > V/TiO$_2$ > V/γ-Al$_2$O$_3$ > V/SiO$_2$ is observed. Goldwasser and Trimm (1979), however, had some difficulties in the study of V/ZrO$_2$ and report that their method for measuring specific surface areas (based on the oxygen sorption of prereduced catalyst samples at 78 K) may not be appropriate in this case. Moreover, their use of XRD in establishing the presence of a monolayer or solid solution is questionable: for XRD crystallite sizes of at least 50 Å (i.e., 10 monolayers) are required. This would contradict their oxygen sorption measurements which point to a monomolecular dispersion.

In the preceding paper (Roozeboom et al., part 3, 1980) it was shown that at low coverages it is impossible to study the dispersed vanadia by XRD, but that Raman spectroscopy and TPR can be used.

Furthermore, the above authors used V coverages far below the theoretical monolayer capacities. On the basis of a surface area of 10.3 Å$^2$ per VO$_2$ species (Hanke et al., 1975) and with their catalyst surface areas it can be calculated that they applied only 5.0, 1.3, and 12.5% of the
theoretical vanadium oxide monolayer capacity onto alumina, silica, and zirconia, respectively. Thus, especially on zirconia O$_2^-$ (inactive for CO oxidation) may be stabilized by spill-over from vanadia to the support (Fenin et al., 1971) and decrease the activity of V/ZrO$_2$.

**Concluding Remarks**

The above consistent evidence has shown that there exists a close correspondence between the structure and support, and two types of active sites can be distinguished in these catalysts. Vanadium ions dissolving in supports, which are active themselves, may often act as promoters constituting a third type of active site.

On all catalysts CO reacts to CO$_2$ via a redox mechanism in which the rate-determining step is the reduction of the catalyst.

**Acknowledgment**

Thanks are due to A. Broersma (State University of Utrecht) for performing the TG experiments and to M. C. Mittelmeijer-Hazeleger (University of Amsterdam) for recording the TPR profiles. This study was supported by the Netherlands Foundation for Chemical Research (SON) with financial aid from the Netherlands Organization for the Advancement of Pure Research (ZWO).

**Literature Cited**


Received for review July 15, 1980
Accepted January 7, 1981

---

**Butynediol Synthesis. A Kinetic Study**

S. S. Kale, R. V. Chaudhari, and P. A. Ramachandran*

National Chemical Laboratory, Pune 411 008, India

A kinetic study of ethynylation of formaldehyde over copper acetylide catalyst prepared from malachite green was carried out in a stirred slurry reactor. The reaction was found to be zero order with respect to acetylene and 0.4th order with respect to formaldehyde. The rate of reaction was found to be retarded in the presence of product butynediol. A Langmuir–Hinshelwood type rate model has been proposed which explains the fractional order with respect to formaldehyde and also the product inhibition effect. The reaction rate constant $k_1$ and adsorption equilibrium constants of formaldehyde and butynediol $K_f$ and $K_b$ obtained at 70°C were $2.921 \times 10^{-3}$ cm$^3$/g s, 951 cm$^3$/mol, and 1500 cm$^3$/mol, respectively.

**Introduction**

The major route for the commercial manufacture of 2-butylene-1,4-diol is by the reaction of acetylene and aqueous formaldehyde in the presence of copper complex catalysts. This process is also known as the Reppe process in which copper acetylide supported on mineral carriers such as silica gel is used as a catalyst. The stoichiometric reaction is

$$
C_2H_2 + HCHO \rightarrow HC\equiv C-\text{CH}_2\text{OH} \quad (i) \quad \text{propargyl alcohol}
$$

$$
HC\equiv C-\text{CH}_2\text{OH} + HCHO \rightarrow \text{HOCH}_2\text{C} \equiv C-\text{CH}_2\text{OH} \quad (ii) \quad \text{butynediol}
$$

$$
C_2H_2 + 2\text{HCHO} \rightarrow \text{butynediol} \quad (iii)
$$

1 NCL Communication No. 2515.

After the discovery of Reppe et al. (1943), many different catalyst systems have been suggested for this reaction and the major product is butynediol. The various catalysts used are summarized in Table I. Most of the investigations summarized in Table I are concerned with the process for butynediol and only a few attempts to study the kinetics of the reaction systematically have been made. Gelperina et al. (1967) studied the kinetics of this reaction using a silica gel supported copper acetylide catalyst and have proposed a kinetic model. Kirchner (1970) has found that the catalyst prepared from malachite green, a naturally occurring copper ore, is much more active than the conventional catalysts. In the present work we report the kinetics of this reaction using this active catalyst (malachite green). The effect of different variables such as stirring speed, catalyst loading, partial pressure of acetylene, and concentration of formaldehyde has been studied and the absence of diffusional resistance is verified. A rate equa-