Atomic-Scale Insights into the Low-Temperature Oxidation of Methanol over a Single-Atom Pt₁-Co₃O₄ Catalyst

Zeyu Jiang, Xiangbo Feng, Jianlin Deng, Chi He,* Mark Douthwaite, Yanke Yu, Jian Liu,* Zhengping Hao, and Zhen Zhao

Heterogeneous catalysts with single-atom active sites offer a means of expanding the industrial application of noble metal catalysts. Herein, an atomically dispersed Pt₁-Co₃O₄ catalyst is presented, which exhibits an exceptionally high efficiency for the total oxidation of methanol. Experimental and theoretical investigations indicate that this catalyst consists of Pt sites with a large proportion of occupied high electronic states. These sites possess a strong affinity for inactive Co²⁺ sites and anchor over the surface of (111) crystal plane, which increases the metal–support interaction of the Pt₁-Co₃O₄ material and accelerates the rate of oxygen vacancies regeneration. In turn, this is determined to promote the coadsorption of the probe methanol molecule and O₂. Density functional theory calculations confirm that the electron transfer over the oxygen vacancies reduces both the methanol adsorption energy and activation barriers for methanol oxidation, which is proposed to significantly enhance the dissociation of the C–H bond in the methanol decomposition reaction. This investigation serves as a solid foundation for characterizing and understanding single-atom catalysts for heterogeneous oxidation reactions.

1. Introduction

Reducing the emission of volatile organic compounds (VOCs) is one of the biggest challenges facing the chemical industry in the 21st century. Many VOCs are considered to pose significant health hazards (carcinogenic, mutagenic, and teratogenic properties) and linked with various environmental hazards. Methanol, as an oxygenated volatile organic compound (OVOC), is widely used in numerous industrial applications. Catalytic oxidation over supported noble metal catalysts has emerged as a promising method for these compounds removal from industrial waste streams, ascribing to their high specific activity, resistance to deactivation, and ability to be regenerated. In our previous works, we determined that Pt-based catalysts are highly active for the low-temperature decomposition of methyl ethyl ketone. The economic limitations associated with the industrial application of these catalysts have, however, reduced their widespread utilization. As such, an abun-dance of researches are currently ongoing to tackle these challenges. Single-atom catalysts (SACs) offer a means of reducing the costs associated with the preparation and utilization of noble metal catalysts. The recent expansion to research in this area will undoubtedly assist with the development of sustainable catalytic technologies for a host of important industrial applications. Single-atom Pt-based catalysts have already been applied to various heterogeneous oxidation reactions, particularly for CO oxidation. Qiao et al. designed a catalyst consisting of isolated single Pt atoms, which were anchored to the surface of iron oxide nanocrystallites.

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When tested for CO oxidation, this SAC exhibited a turnover frequency H2–3 times higher than a corresponding Pt/Fe2O3 catalyst (with sub-nanometer Pt particles) and remained stable for up to 300 min.[9a] The inherent stability of single-site Pt catalysts for CO oxidation was further demonstrated by Zhang et al.,[9d] who determined that there was no observable deactivation over 60 sequential cycles when operating at temperatures between 100 and 400 °C. The excellent stability of this catalyst was attributed to firmly anchored Pt atoms on the internal surface of the mesoporous Al2O3 support, which were proposed to be stabilized by coordinate unsaturated pentahedral Al3+ centers.[9d]

Regrettably, there are only limited examples for the use of SACs in the oxidation of VOCs.[10] One example was, how-ever, presented by Tang and co-workers,[11] who reported that a single-atom Ag-HMO catalyst (HMO: hollandite manganese oxide) was highly active in the oxidation of formaldehyde. The studies revealed that subtle changes in the support material could lead to electronic perturbations in the single-atom sites, which had a profound effect on the intrinsic reactivity of the cat-alyst. This was a consequence of changes in the silver atoms 4d electronic state, resulting in an increasing of both the metal–support interaction and reducibility of the catalyst.[11]

It is known that the activity, selectivity, and stability of SACs can be tuned by modifying the interactions between the anchored atoms and support surfaces.[12] The geometric location of these metal atoms on the surface of support is considered to impact how strongly the atoms interact with the support.12b The isolated single metal atoms, which are consid-ered to anchor over cationic vacancies/equivalent cationic posi-tions or to other surface defects on the support, are recognized as the active centers in these catalytic systems.12 We therefore proposed that developing a greater understanding of how SACs affect oxidative reaction mechanism is paramount for their application as catalysts for the removal of VOCs.

As such, we began by synthesizing a highly oxygen-deficient single-atom Pt1-Co3O4 catalyst via a facile one step method. A detailed description of the geometric and electronic structure of this catalyst and its role in the oxidation of methanol was established by conducting a series of experimental and theo-retical studies. A link in the structure and the origin of catalytic property was also investigated to demonstrate the excellent per-formance our single-atom Pt1-Co3O4 catalyst in methanol total oxidation reactions.

2. Results and Discussion

2.1. Determination and Modeling of Single-Atom Sites

Herein, we aimed to promote the application of single-atom catal-ysts in VOC destruction. As such, we synthesized the Pt1-Co3O4 catalyst over which the Pt atoms are anchored on the surface defects of Co3O4 due to the existence of inactive Co3+-species. During the coprecipitation process, the Pt atoms are inclined to occupy the inactive Co3+-sites and anchor over the surface of (111) crystal plane through metal–support interaction.[13] The exposed (111) facet of Co3O4 support consisted of queued Co and O atoms in isolated lines (Figure 1a). This was also sub-sequently confirmed to be the most stable Co3O4 conformation by theoretical calculations (Figure S1, Supporting Information). On this surface, two types of threefold-coordinate oxygen atoms (Oa and Ob) are exposed; Oa has three Co3+ ions as nearest neighbors, whereas Ob atoms are bound to one Co2+ ion and two Co3+ ions. Upon impregnation with chloroplatinic acid, the Pt nanoparticles (yellow circles) with around 0.5–3.1 nm particle size were supported on the Co3O4 (111) plane surface, which are highlighted in Figure 1b and Figure S2 (Supporting Informa-tion). No Pt nanoparticles could be observed from high-resolu-tion transmission electron microscopy- (HRTEM) images of the Pt1-Co3O4 catalyst (Figure 1c and Figure S3, Supporting Informa-tion) and the intraplanar spacing (d values) were measured to be H0.45 nm, in good consistence with that of the exposed (111) crystal plane of the Co3O4 sample.[14] The fast Fourier transformation (FFT) of exposed facet is in well agreement with the standard Co3O4 (111) crystal plane.[13] In addition, as shown in Figure S4 (Supporting Information), no changes have occurred of FFT patterns between Co3O4 and single-atom Pt1-Co3O4 materials, which indicates that the single-atom Pt sites are anchored over the surface of Co3O4 supports and the exist-ence of single atoms has a negligible effect on (111) crystal plane. In addition, the content of Pt detected by inductively coupled plasma optical emission spectrometry (ICP-OES) is similar to that determined by X-ray photoelectron spectroscopy (XPS) and X-ray fluorescence (XRF) methods (Table S1, Sup-porting Information), which confirms that the Pt active sites are almost located at the surface of support. Models for the opti-mized structure of the single-atom Pt1-Co3O4 catalyst were sub-sequently proposed through considering all possible synergetic states (Figure 1d–f). Previously, aberration-corrected high-angle annular dark-field (HAADF) microscopy has been used to deter-mine the presence of heavy metals in SACs.[15] Upon utilization of this technique to probe the Pt1-Co3O4 catalyst (Figure 1g–j), the presence of individual uniformly dispersed Pt atoms was observed. Extensive examination of additional catalytic regions in the sample confirmed that only Pt single atoms are present in the sample (Figure S5, Supporting Information).

Infrared (IR) spectr um utilizing CO as the probe mole-cule is a fast and convenient characterization method to dif-ferentiate between metallic single atoms and nanoparticles.[16] When investigating Pt-containing catalysts, it can also be used to acquire information regarding the geometric configuration of the Pt centers.[16] No CO adsorption peak can be detected over the Co3O4 sample. In contrast, two CO adsorption bands pre-sented in the IR spectrum of the Pt1-Co3O4 catalyst (Figure 1k). The adsorption bands centered at 2116 and 2029 cm–1 at room temperature correspond to the adsorption of CO over a single Pt atom and the asymmetric stretching vibrations of CO in the Pt(CO)2 gem-dicarbonyl species on single Pt atoms, respectively.[16] With the Pt/Co3O4 catalyst however, only one peak can be observed at 2089 cm–1, characteristic of linear CO adsorption over metallic Pt nanoparticles.[16a,b] To investigate how the adsorbed CO species on the surface of the Pt1-Co3O4 catalyst was affected by exposure temperature, some additional IR experiments were conducted, and the results of which are displayed in Figure 1l. Interestingly, the intensity of the bands almost remain unchanged as the temperature increased in this experiment, indicating that the quantity of Pt active sites in the sample remained stable in temperature range 25–100 °C.
2.2. Determination of Electronic Structure and Defect Sites

Understanding the electronic structure of elemental components in a SAC is the most important challenge, which is fundamental for leading to the structure–activity correlations. Representative electron energy-loss spectra obtained from the Pt1-Co3O4 and Pt/Co3O4 materials are presented in Figure 2a. Pronounced peaks at 780 and 795 eV are characteristic of L3 and L2-shell ionization edges in Co, respectively. It is evident that approximately a third of the Co present exists as Co2+ in a tetrahedral coordination to oxygen and two-thirds exist as Co3+ in an octahedral oxygen environment in the SAC. In the octahedral coordination, the tetrahedral environment splits the Co 3d states into three t2g and two eg levels. The proportion of tetrahedral/octahedral species present in the Pt1-Co3O4 catalyst is notably higher than that observed in the Pt/Co3O4, indicating that there is a strong interaction between the single-atom Pt sites and Co3O4 supports in the Pt1-Co3O4 catalyst, which in turn, would facilitate the formation of Co3+ species. These results can be supported by XPS experiments presented in Figure 2b. The asymmetrical Co 2p3/2 region can be deconvoluted into two components centered at binding energies of 779.4 and 780.7 eV, characteristic of surface Co2+ and Co3+ species, respectively. The proportion of Co3+/Co2+ in each catalyst was subsequently quantified and the ratios were determined to be 1.71, 1.97, and 2.42 for the Co3O4, Pt/Co3O4, and Pt1-Co3O4 catalysts, respectively. Furthermore, the XPS spectra of samples with different Pt loadings indicated that the ratio of Co3+/Co2+ was enhanced with the increasing of Pt content, attributing to the support–metal strong interaction in these SACs (Figure S6, Supporting Information). The Co3+ species is clearly dominant, which is unsurprising given that this species is known to be pivotal in low-temperature oxidation reactions over Co3O4 catalysts. According to previous reports, the Pt0 species were seen as the preponderant active sites compared with Pt2+ and Pt4+ species in VOC total oxidation. In this work, a higher
Electron paramagnetic resonance (EPR) spectroscopy was subsequently used to probe the materials; see Figure 2c. Each of the catalysts exhibited a broad, intense resonance at $g = 2.17$ mT, which is characteristic of a Co(II) with $S = 3/2$ species. According to a previous study, the quantity of high spin Co(II) species presenting in a given catalyst is proportional to the content of oxygen vacancies.\([22]\) It is clearly evident from Figure 2c, that the intensity of Co(II) peak in the Pt$_1$-Co$_3$O$_4$ catalyst is significantly greater than that observed for the Co$_3$O$_4$ and Pt/Co$_3$O$_4$ catalysts. Furthermore, an additional peak at $g = 2.24$ mT is observed in the EPR spectrum of the Pt$_1$-Co$_3$O$_4$ catalyst; the intensity of this peak in the spectra of the Co$_3$O$_4$ and Pt$_1$-Co$_3$O$_4$ materials is considerably smaller. This peak is considered to be characteristic of electrons trapped in oxygen vacancies.\([19a,23]\) This could indicate that the Pt$_1$-Co$_3$O$_4$ catalyst possesses a higher proportion of oxygen vacancies than the Co$_3$O$_4$ and Pt/Co$_3$O$_4$ catalysts. This aligns with additional DFT calculations (Figure 2g.h), which reveal that the formation of oxygen vacancies over Co$_3$O$_4$ and Pt$_1$-Co$_3$O$_4$ samples (red, pink, and blue spheres represent O, Pt, and Co atoms, respectively).

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energy for an oxygen vacancy (\(E_{\text{vac}}\); energy needed to form one oxygen vacancy on the surface and half an oxygen molecule in the gas phase) on a Pt1-Co3O4 (111) surface (1.52 eV) is much lower than that on a Co3O4 (111) surface (2.41 eV).

The Bader charge analysis confirmed that the formation of an oxygen vacancy leads to an increased electron density on the Pt (0.4 e). After adsorption of methanol and O\(_2\), the electron on the Pt atoms decreased by 0.2 and 0.5 e, respectively. It is likely that the Pt atoms exhibit a stronger adsorption capacity on the defective surface than that on the Co3O4 (111) and Pt1-Co3O4 (111) surfaces. To further investigate this conclusion, the density of states (DOS) of the Pt and Co 3d orbital was compared, as displayed in Figure 2d. The Pt atom with a higher occupied orbital exhibits a stronger affinity to methanol and O\(_2\), which can be ascribed to the decreased filling of adsorbate-metal antibonding states. Thus, methanol and O\(_2\) molecules are more likely to coadsorb onto a defective Pt1-Co3O4 surface and the adsorption capacity is greatly enhanced by the oxygen vacancies. Further evidence of this is provided by O\(_2\)-temperature-programmed desorption (TPD), methanol-TPD, and in situ Fourier transform infrared (FTIR) of methanol adsorption experiments, which are illustrated in Figures S8–S10 (Supporting Information), respectively.

The optimized configurations of methanol and O\(_2\) over the Pt1-Co3O4 (111) surface are presented in Figure S11 and Table S2 (Supporting Information). The calculated adsorption energy of methanol is 0.03 eV. The O atom of the methanol molecule bonds to the Pt atom and the distance between Pt and O atoms is calculated to be 2.78 Å. O\(_2\) molecules can also adsorb onto Pt sites with an exothermic energy of 0.3 eV and exhibit a Pt O bond length of 2.13 Å. In contrast, the adsorption energy of methanol and O\(_2\) onto the Co3O4 (111) surface is 0.91 and 0.49 eV, respectively (Figure S12, Supporting Information). The adsorption of methanol and O\(_2\) on the defective Pt1-Co3O4 (111) surface was subsequently investigated (Figure S13, Supporting Information). With this catalyst, it was determined that methanol adsorbs onto a Pt site with an energy of 1.48 eV and the resultant Pt O bond length was calculated to be 2.08 Å. This notable enhancement is likely to be attributed to the OH group in the methanol participating in hydrogen bonding interactions with the lattice oxygen species. The adsorption energy of O\(_2\) over this catalyst was calculated to be 0.66 eV. As such, it can be concluded that increasing the quantity of oxygen vacancies in the catalyst greatly enhances both the methanol and O\(_2\) adsorption significantly, which is obviously very well important for total oxidation catalysts.\(^\text{(4)}\)

In the Pt1-Co3O4 material, we have confirmed that the Pt atoms instigate the charge imbalance and increase the proportion of oxygen vacancies in the catalyst. In Figure 2e, the Pt1-Co3O4 catalyst, which exhibits AB2O4 structure (where A and B represent Co\(^{2+}\) and Co\(^{3+}\), respectively), consists of five Raman-active modes: A\(_{1g}\), E\(_g\), and three F\(_{2g}\) modes, centered at 675, 467, 513, 189, and 608 cm\(^{-1}\), respectively.\(^\text{(24)}\) The F\(_{2g}\) and E\(_g\) modes suggest that there are different motions in the tetrahedral sites (CoO\(_4\) unit); the F\(_{2g}\) mode at 189 cm\(^{-1}\) is assigned to a complete translation of the CoO\(_4\) unit within the lattice. The A\(_{1g}\) mode can be attributed to the octahedral CoO\(_6\) sites of crystalline Co3O4 phase. Interestingly, when the spectra of the Co3O4 and Pt/Co3O4 modes were investigated, the positions of all recorded Raman peaks of Pt1-Co3O4 shift to higher frequencies and the intensity of A\(_{1g}\) symmetry increases, which is indicative of defects in the crystal structure. Analysis of in situ Raman spectra, the intensity of the peak at about 665 cm\(^{-1}\), characteristic of Co O bond vibration, decreases in the spectrum of the Pt/Co3O4 as temperature increases from 0 to 230 °C; see Figure S14 (Supporting Information). This is likely to be attributed to a change in the crystalline phase as the temperature is increased. Interestingly, in the experiment over the Pt1-Co3O4 catalyst, this peak goes through a max at 110 °C; see Figure 2f. This suggests that at this temperature, the presence of oxygen vacancies in the catalyst is at its highest, which in turn, indicates that the electron transfer from Pt to Co is at its most efficient at this temperature.

### 2.3. Catalytic Performance of Prepared Materials

Methanol was used as a substrate to assess the catalytic potential of the Pt1-Co3O4 SAC. The results from these initial experiments are displayed in Figure 3 and Table 1. It was determined that over the Pt1-Co3O4 catalyst, the complete oxidation of methanol into CO\(_2\) and H\(_2\)O could be achieved at only 110 °C (Figure 3a). The reaction rate and turnover frequency of Pt-based catalysts prepared in this study were subsequently assessed (Figures S15 and S16, Supporting Information). A reaction rate of 0.013 mol g\(^{-1}\)Pt s\(^{-1}\) was determined for the Pt1-Co3O4 catalyst at 70 °C, which was over four times higher than that observed over the Pt/Co3O4 catalyst (0.03 mol g\(^{-1}\)Pt s\(^{-1}\); Figure 3b and Figure S15, Supporting Information). The turnover frequencies (TOFs) exhibited by both catalysts were based on the Pt dispersion (Figure 3b and Figure S16, Supporting Information). The measurements indicate that the activity exhibited by each Pt site in Pt1-Co3O4 was substantially higher at 70 °C, which to the best of our knowledge is higher than other catalysts previously reported for this reaction to date (Table S3, Supporting Information). These data were subsequently used to construct Arrhenius plots and establish activation energies (\(E_a\)) for methanol oxidation over each catalyst (Figure S17, Supporting Information). The activation energy of the single-site Pt1-Co3O4 catalyst was substantially lower than that observed over the Pt/Co3O4 catalyst. Based on the observed activity of Pt1-Co3O4 catalyst, we conduct the detailed correlation analysis between catalytic performance and structure properties including: high specific surface area (Figure S18, Supporting Information, and Table 1), sufficient surface-bound hydroxyl species (Figure S19, Supporting Information), high quantity of lattice oxygen (Figures S20–S22, Supporting Information), superior low-temperature reducibility (Figure S23, Supporting Information), high oxygen/methanol adsorption capacity (Figure S24, Supporting Information), and abundant oxygen vacancies (Figure S25, Supporting Information). As discussed, based on the experimental and theoretical studies conducted in this work, we consider that the significant increase in the quantity of oxygen vacancies over the Pt1-Co3O4 catalyst is likely to be the most crucial factor, the electron transfer over the oxygen vacancies reduces both the methanol adsorption energy and activation barriers for methanol oxidation, which was proposed to significantly enhance the dissociation of the C–H bond in methanol decomposition reaction.
The stability test of prepared catalysts was subsequently performed. The resistance of water was examined by introducing 3 vol% of water vapor to the catalysts between a heating and cooling cycle (Figure 3c). After additional tests, notable losses in methanol conversion of H15.0 and 18.0% were observed over the Pt/Co3O4 and Pt/CoO4 catalysts, respectively. Interestingly, however, the original activity was restored after only 30–45 min of stopping H2O in the feed, suggesting that the reductions in activity were not indicative of permanent changes to the catalysts. In addition, the single-atom Pt/Co3O4 catalyst possesses a better CO2 (5 vol%) resistance (Figure 3d), which can be attributed to the abundant oxygenvacancies around Pt active sites. The adequate oxygen vacancies around single-atom Pt provide sufficient adsorption sites for O2 molecules during methanol oxidation, which restrains the competitive adsorption of CO2 and O2 molecules to some extent. These catalysts are evidently highly stable, which was supported by in situ X-ray diffraction (XRD) experiments (Figures S26 and S27, Supporting Information). In addition, the aberration corrected scanning transmission electron microscopy (ac-STEM) images of Pt1-Co3O4 after oxidation reaction were performed in order to assess the aggregation of Pt active sites. As displayed in Figure S28 (Supporting Information), the Pt sites remain stable after the oxidation reactions at 150 and 200 °C; however, the Pt atoms aggregated to some extent when it involved in 250 °C. As such, the stability of SACs remains a challenge when involving VOC oxidation at high temperature.

We have explored the possibility and pushed to the limit activity of this catalyst in order to explore the application of SACs with higher metal loadings in VOC elimination. As shown in Figure 3e, the catalytic performance of prepared catalysts was promoted within 0.02–0.15 wt% of Pt content for methanol oxidation; however, unobvious activity change can be found when further increasing the Pt content to 0.20 or 0.30 wt% due to the formation of Pt nano-particles, identified by the hydrogen and oxygen titration.

**Table 1. Textural property and catalytic performance of prepared materials.**

<table>
<thead>
<tr>
<th>Sample</th>
<th>Pt content [wt%]</th>
<th>Dispersion [%]</th>
<th>$S_BET$ [m$^2$ g$^{-1}$]</th>
<th>$V_{pore}$ [cm$^3$ g$^{-1}$]</th>
<th>$D_{pore}$ [nm]</th>
<th>$T_{S0}$ [°C]</th>
<th>$T_{S90}$ [°C]</th>
<th>$r_i$ [mol g$^{-1}$ s$^{-1}$]</th>
<th>$TOF_{Pt}$ [x 10$^4$] [s$^{-1}$]</th>
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<tr>
<td>Co3O4</td>
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<td>–</td>
<td>55.5</td>
<td>0.53</td>
<td>18.4</td>
<td>107</td>
<td>128</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Pt/Co3O4</td>
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<td>63.7</td>
<td>58.9</td>
<td>0.57</td>
<td>19.1</td>
<td>87</td>
<td>109</td>
<td>0.03</td>
<td>7.2</td>
</tr>
<tr>
<td>Pt1-Co3O4</td>
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<td>91.5</td>
<td>59.5</td>
<td>0.51</td>
<td>18.4</td>
<td>81</td>
<td>96</td>
<td>0.13</td>
<td>29.3</td>
</tr>
</tbody>
</table>

a) Actual Pt content detected by ICP-OES; b) The dispersion of Pt active sites measured by hydrogen and oxygen titration (HOT); c) Specific surface area obtained at $P/P_0 = 0.05–0.30$; d) Total pore volume estimated at $P/P_0 = 0.99$; e) BH pore diameter calculated from the adsorption branch; f) Temperatures at which 50 and 90% conversion of methanol; g) Reaction rate of methanol molecules transformed per Pt metal per unit time of various catalysts at 70 °C; h) Turnover frequency based on the dispersion of Pt obtained at 70 °C.
We further calculated the turnover frequency of prepared materials at 62 °C (conversion lower than 20%), as shown in Figure 3f. The TOF of Pt sites maintained at around 0.03–0.05 s⁻¹ within 0.02–0.15 wt% of Pt and declined when Pt loading increased (0.0225 s⁻¹ for 0.20 wt% P1-Co3O4; 0.0159 s⁻¹ for 0.30 wt% P1-Co3O4). Above results indicate that the actual active sites of these P1-Co3O4 catalysts are Pt single atoms. The Pt loading of 0.15 wt% is the limit of single-atom P1-Co3O4 catalyst prepared by this coprecipitation method in this work, over which the complete oxidation of 700 ppm methanol could be achieved at only 90 °C and with a 0.0356 s⁻¹ TOF at 62 °C.

2.4. Insight on the Atomic-Level Mechanism

In this work, the low-concentration VOC oxidation over the single-atom materials follows a first-order reaction, which is ascribed to the excess of oxygen in the reaction system and the reaction rate is only proportional to the first power of VOC concentration. A detailed understanding of the oxidative surface mechanisms for VOCs over SACs would be incredibly beneficial from a catalyst design perspective. For this reason, a series of in situ diffuse reflectance infrared Fourier transform spectros-copy (DRIFTS) experiments were conducted over the P1-Co3O4 and Pt/Co3O4 catalysts, in an attempt to derive information regarding the intermediate products formed during the reaction (Figure 4a,b). The spectra of both catalysts consist of bands characteristic of C-H bond vibrations (2892 cm⁻¹) and surface-bound hydroxyl species (3050 and 1431/1422 cm⁻¹).[26] The peaks centered at 1257/1230 cm⁻¹ are attributed to the vibration of methoxy species.[26a] Interestingly, an additional band is observed at 1494 cm⁻¹ in the spectrum of P1-Co3O4, which is typically characteristic of a symmetric carbonyl groups.[27] It is likely that after coadsorption of CH3OH and O2, the OH group of CH3OH molecule can dissociate, leading to the formation

![Figure 4](image-url)
of Pt-methoxyl species. The oxygen vacancies regeneration and electron transfer over Pt atoms accelerate the C–H bond activation over methoxyl species, which go through the formaldehyde as intermediate and then oxidized into H$_2$O and CO$_2$.

The $^{18}$O isotope tracer technique was introduced to investigate the lattice oxygen transformation. Figure 4c displays the products containing isotopic species (C$^{18}$O$_2$, C$^{16}$O $^{18}$O, and C$^{16}$O$_2$) during methanol oxidation. All these species showed the temperature-dependent variation. The C$^{16}$O$_2$ species took a vigorous increasing after the reaction beginning, meanwhile, the signal of C$^{15}$O$^{18}$O and C$^{18}$O$_2$ species grew tardily during the oxidation reaction. This phenomenon provides the evidence that lattice oxygen is the primary active oxygen species during methanol elimination over Pt$_{1}$-Co$_{3}$O$_4$ catalyst. It is widely accepted that the process of oxygen vacancies regeneration provides abundance of hydroxyl groups for sam-ples, which plays a significant role in intermediates decomposition.[28] As such, some additional theoretical experiments were conducted in order to investigate how the regenera-tion of oxygen vacancies on the surface of the Pt$_{1}$-Co$_{3}$O$_4$ was affected by the presence of hydroxyl groups (Figure 4d–g). The Pt$_{1}$-Co$_{3}$O$_4$ surface was first saturated with hydroxyl groups (OH or O bH); evidence for their introduction is displayed in Figure 4d. For a H atom to migrate from O$_b$ to a lattice oxygen atom bonded to Pt (O$_p$H), the energy barrier was determined to be 0.3 eV (Figure 4e). This allows for the subsequent migra-tion of an additional H atom from a lattice oxygen species to the O$_p$H, resulting in the production of a H$_2$O molecule (Figure 4f). This process was determined to be endothermic (0.95 eV). The sequential desorption of the H$_2$O on the surface of the Pt$_{1}$-Co$_{3}$O$_4$ catalyst leads to the liberation of the lattice oxygen sites (Figure 4g).

Based on the DFT calculations, a catalytic cycle for the oxidation of CH$_3$OH and the corresponding transition states over the Pt$_{1}$-Co$_{3}$O$_4$ catalyst is proposed in Figure 5. First of all, a surface phase diagram was provided for Pt$_{1}$-Co$_{3}$O$_4$ as the surface can be with an oxygen defect or adsorption phase. The calculated surface free energies of the three possible terminations are shown in Figure S29 (Supporting Information). The temperature of $T =$ 600 K corresponds to the annealing temperature employed in our experiment. When $\Phi$O becomes extremely low (lower than 1.46 eV), the Pt$_{1}$-Co$_{3}$O$_4$ surface is the most stable structure. Accordingly, Pt$_{1}$-Co$_{3}$O$_4$ surface with an oxygen vacancy is used to investigate the oxidation of methanol in this work. After coad-sorption of the CH$_3$OH and O$_2$, the OH group of CH$_3$OH molecule can dissociate (CH$_3$OH → CH$_3$O + H), leading to the formation of Pt-methoxyl species and H atom required for the formation of the O$_3$H species (Figure 5a). In the absence of Pt, over the Co$_3$O$_4$ (111) surface, a significantly higher energy barrier is required for this reaction pathway (0.2 eV). O$_2$ activation over the Pt$_{1}$-Co$_{3}$O$_4$ (111) surface leads to the formation of a peroxide-type O$_2$ $^{2-}$ species. One oxygen atom of O$_2$ molecule fills the oxygen vacancy and the other oxygen atom binds to an adjacent Pt atom. The Pt O and O bonds lengths were determined to be 2.09 and 1.40 Å, respectively. With the assistance of the active oxygen species O$_2$ $^{2-}$, the methoxyl species partake in a C–H bond cleavage, leaving a surface-bound CH$_2$O species (CH$_3$O $\rightarrow$ CH$_2$O + H), the energy barrier of which was determined to be 1.14 eV.

**Figure 5.** Proposed reaction pathways and the corresponding transition states for methanol oxidation over Pt$_{1}$-Co$_{3}$O$_4$ catalyst.
(1.37 eV on Co₃O₄ (111) surface). The surface-bound hydrogen atoms can also interact with the O₂⁻²⁻ species, leading to the formation of OH groups (Figure 5b). Through interaction with these surface hydroxyls, the CH₂O interme-diate species can undergo a sequential C-H dissociation to produce CHO and H₂O. The calculated activation barrier for this step over the Pt₁-Co₃O₄ (111) surface is 0.44 eV, but is significantly higher in the absence of Pt, 1.78 eV. The CHO group subsequently binds to the Pt atom to form a Pt-OCH species (Figure 5c). Following this, H₂O desorbs and Pt-OCH undergoes a geometric distortion to form Pt-CHO (Figure 5d). This process is endothermic (0.46 eV). The C-H bond of CHO can then undergo further decomposi-tion (CHO + OH → CO + H₂O) with an energy barrier of 0.54 eV and the hydrogen atom adsorbs onto another O₃C₄ group to form water (Figure 5e). Water desorbs with an energy of 0.35 eV, leaving only the O₃C₄ molecule bound to a Pt atom and an oxygen vacancy on the surface (Figure 5f). Finally, an additional O₂ molecule adsorbs to the oxygen vacancy and dis-sociates, providing the additional oxygen atom required for oxidizing CO to CO₂. This step releases the energy of 2.65 eV (Figure 5g). The CO₂ subsequently desorbs, releasing heat (1.67 eV) and the surface of the Pt₁-Co₃O₄ SAC returns to ini-tial state without oxygen (Figure 5h). As such, the surfaces of the oxide particles are covered with (OH) formed by the dis-sociative chemisorption of water. Following a microkinetics simulation has been carried out by MKMCXX software. In the whole catalytic reaction cycle, the energy barrier of the recovery of Pt₁-Co₃O₄ surface is only 0.3 eV, which is easy to achieve. Thus, we ignored the process of surface recovery and assumed that the reaction cycle finished after the desorption of CO₂. The initial concentration of CH₃OH and O₂ is set as 2:3, and the pressure is set as 1 atm. The forward and back-ward rates of each elementary reaction step of methanol oxidation at 375 K are shown in Table 2. It can be seen that the step of CH₃O*HO → CH₂O*HO₂H has the largest degree of rate control value, indicating that the cleavage of C-H bond of (CH₃O*) influences the reaction the most. It is related with the largest energy barrier of this step in the whole reaction cycle. The corresponding surface coverage results show that CH₃O*H and CH₂O*HO₂ are predominate surface species (73.1 and 26.4%), which attributes to the weak adsorption of O₂. The calculated apparent activation energy decreases with the increase of temperature, and the apparent activation energy for CH₃OH oxidation at 375 K is 55.1 kJ mol⁻¹.

3. Conclusion

Through a combination of theoretical and experimental investigation, a detailed description for the geometric and electronic structure of a Pt₁-Co₃O₄ single-atom catalyst was derived. This catalyst was determined to consist of isolated single Pt atoms anchored to Co₃O₄ (111) planes, which occupied positions on the Co³⁺ atoms. The Pt₁-Co₃O₄ exhibited an excellent activity for the oxidation of methanol. One-third of the Co atoms in this catalyst were determined to exist as Co²⁺ in a tetrahedral coor-dination to oxygen; the remaining Co atoms were present as Co³⁺ in an octahedral oxygen environment. The Pt sites, which had high occupied electronic states, exhibited a strong affinity for the 3d orbital of adjacent Co atoms, resulting in significant electron transfer from Pt to Co, ultimately, increasing the pro-portion of oxygen vacancies present on the catalysts surface. Regeneration of these oxygen vacancies was determined to promote the coadsorption of methanol and O₂, leading to an increase in the rate of C-H bond dissociation. DFT calcula-tions revealed that the electron transfers reduced the activation barriers for methanol oxidation. The discovery of this single-atom Pt catalyst highlights that single-site heterogeneous cat-alysts can be exceptionally effective for the oxidation of VOCs and provides a method for reducing the cost of noble-metal catalysts in industrial applications.

4. Experimental Section

All experimental details can be found in the Supporting Information.

Supporting Information

Supporting Information is available from the Wiley Online Library or from the author.
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Conflict of Interest

The authors declare no conflict of interest.

Keywords

developed oxygen vacancies, DFT calculations, mechanism, single-atom catalysts, VOC low-temperature oxidation