Scale-Up of Cluster Beam Deposition to the Gram Scale with the Matrix Assembly Cluster Source for Heterogeneous Catalysis (Catalytic Ozonation of Nitrophenol in Aqueous Solution)

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1. INTRODUCTION

Cluster beam deposition (CBD) is a vacuum-based, solvent-free method to synthesize metal clusters on surface.1 The clusters produced have no chemical ligands, and contamination levels should be low, so the intrinsic properties of the clusters can be controlled, investigated, and exploited. This may allow the rapid prototyping of novel, cluster-based catalysts for various reactions, since the capacity to vary cluster composition and size compares favorably with the traditional chemical methods. The clusters are simply formed by condensing individual atoms of the material vaporized,2 so composited clusters (binary, ternary, and beyond) should be relatively easy to synthesize. Recent studies have shown precise control of cluster morphology,3 structure,4 and size5 by regulating the condensation conditions and combining with a time-of-flight mass filter.6 However, the CBD method has been largely limited to surface science studies by the low cluster deposition rate (ng to μg/h). Thus, scale-up of the cluster production rate is a crucial step to translate this technique into practical applications.7 For this reason, a new type of source, the matrix assembly cluster source (MACS), has been developed in our group with potential for considerable scale-up.8,9 The formation of clusters in the MACS has been studied both experimentally8,10 and theoretically.11,12 It is based on the assembly of atoms, initiated by ion beam impacts, in a metal-loaded, cryogenically cooled rare gas (Ar) matrix. The latest study13 shows that the cluster deposition rate from the MACS can reach ∼10 mg of clusters per hour, which is enough to prepare cluster-based catalysts at the gram-scale for R&D studies at the test tube level. While the first investigations of clusters from the MACS have been reported, heterogeneous catalysis in solution has yet to be demonstrated.

Here, the oxidation of nitrophenol by ozone in aqueous solution using clusters deposited on commercial powder supports is demonstrated. Nitrophenol, a nitroaromatic
chemical, is a model compound for the water treatment industry, being toxic, mutagenic, and biorefractory in the aquatic environment. It is widely detected as an organic pollutant in industrial and agricultural wastewaters. The maximum allowable contaminant level (for p-nitrophenol) in water is only around 10 ng/L, since it can irritate eyes and skin, damage kidneys and the liver, cause methemoglobin, etc. The ozonation of nitrophenol is an important reaction commonly used to remove this toxic substance from environmental media. Although the use of catalysts (e.g., metals supported on oxide powders) in chemical reactions (e.g., nitrophenol reduction) has been found effective to accelerate reaction rate, catalytic ozonation with nano-clusters has not yet been reported. One of the issues in heterogeneous catalytic ozonation is catalyst agglomeration due to pH changes in water and catalyst retrieval and reuse, and we address the latter problem here.

In this work, we demonstrate the catalytic activity of Ag and Au clusters physically deposited onto commercial titanium dioxide (TiO₂) powders with the MACS in the catalytic ozonation of nitrophenol. Compared to traditional catalysts reported in the literature, the catalysts produced by the MACS exhibit at least comparable activities and show excellent reusability without evident degradation after five cycles.

2. EXPERIMENTAL METHODS

2.1. Cluster Catalyst Preparation. The Au and Ag cluster catalysts used in this work were prepared with a new MACS system at Swansea University (MACS 3). Figure 1 shows a photograph and schematic diagram of the system. The new MACS 3 system consists of two vacuum chambers: the cluster deposition chamber and a sample load-lock chamber. In the deposition chamber, the matrix is generated, a deposition cup (stainless steel, 10 cm in diameter and 2 cm in height) containing powder supports (1 g, TiO₂, P25 from Sigma-Aldrich, U.K.) is transferred to a position beneath the matrix support to harvest the clusters. The matrix is then sputtered with an Ar-ion beam (~2 mA, Gencoa ion source) to generate a sequence of collision cascades in which the metal atoms are condensed together and ripened in a dynamic irradiated environment. Eventually, the ripened clusters are sputtered out of the matrix. During the deposition, the powder supports were mechanically stirred by a metal comb stirrer to maximize the powder particle surface area exposure to the cluster beam.

Compared to the previous MACS system, the new MACS 3 system shows several improvements: (i) a larger matrix support with a working planar-projected surface area of >100 cm², (ii) a higher cooling capacity cryocooler with 80 W of cooling power at 30 K, (iii) a high-capacity powder system allowing up to 100 g of powders to be mechanically stirred—the stirring system replaces the previous vibration cup, and (iv) a higher-current ion source, which can deliver a sputtering current of >100 mA. The cluster yield in the MACS has been shown to be roughly linearly proportional to the sputtering current (about 1 cluster is emitted per 100 Ar⁺ ion impacts), so the upgraded system should allow the production of up to ~1 g of clusters (Au₁₀₅) per hour, assuming all of the clusters produced are collected. However, in this first experiment with MACS 3, a conservative sputtering current of ~2 mA was used, comparable to the normal level of the previous MACS 2 system.

2.2. Characterization. The morphology imaging of both the Au and Ag cluster catalysts was performed by a Thermo Fisher Talos scanning/transmission electron microscope (STEM) equipped with a high-angle annular dark-field (HAADF) detector operating with an inner angle of 55 mrad at 200 kV. The incident electron beam convergence angle was 21 mrad. Cluster size distributions were directly measured from the diameter cross section of individual clusters in the HAADF-STEM images. The STEM samples were prepared by dispersing the catalyst powders decorated with Au or Ag clusters in deionized water, sonicating for several minutes and drop-casting onto a copper grid coated with an amorphous carbon film.

The metal loading of the cluster catalysts was characterized by inductively coupled plasma-optical emission spectrometry (ICP-OES) following digestion in a mixed solution of hydrofluoric acid and nitric acid. The Brunauer–Emmett–Teller (BET) surface area and pore radius were measured at 77 K using N₂ adsorption/desorption isotherms (Nova 200e, Quantachrome Instruments). The samples were degassed at 125 °C for 12 h before nitrogen adsorption measurements. Quantachrome NovaWin package was used for data acquisition and analysis.

The surface chemistry of the samples was studied by X-ray photoelectron spectroscopy (XPS) after air exposure. The XPS experiments were performed at room temperature and at a base pressure of 1.2 × 10⁻⁹ mbar using a Kratos AXIS Supra equipped with a monochromatized Al Kα X-ray source (1486.6 eV) operated at 15 mA. The photoelectron core-level spectra were acquired using a hemispherical analyzer at a pass energy of 160 eV for the wide scan, while the high-resolution spectra were obtained at a passing energy of 40 eV. All of the spectra were aligned to the C–C carbon peak at 284.8 eV. The deconvolution of the spectra was done by CasaXPS software and used to calculate the surface composition.

2.3. Nitrophenol Ozonation Measurements. Ozonation was carried out at room temperature and neutral pH in a semibatch glass reactor containing 200 mL of 4-nitrophenol solution (0.1 mM) and 50 mg of catalyst under continuous stirring. O₃ was generated from an ozone generator (BMT 803, BMT Messtechnik, Germany) fed by dry air, and the gas concentration was measured by an ozone analyzer.
The gaseous feed was bubbled in the reactor at a flow rate of 500 mL/min. The ozone concentration was adjusted from 3 to 10 g/m^3 NTP (NTP: normal gas temperature and pressure; 0°C and 1 atm) by varying the power of the ozone generator. The catalytic reaction was initiated by introducing ozone into the reactor using a sintered glass diffuser. The optical absorbance of the reaction solution was recorded by an ultraviolet−visible (UV−vis) spectrophotometer (Lambda 35, PerkinElmer) to monitor the kinetics of the reaction. Sample solutions of approximately 3 mL were periodically withdrawn from the reactor, followed by air bubbling to quench the ozone reaction and filtration before transferring into a quartz cuvette. The pH of the sample was adjusted prior to UV−vis spectrum recording. The formation of maleic acid was tracked by liquid chromatography−tandem mass spectrometry (LC/MS/MS) (m/z 115). The LC/MS/MS analysis was performed on an Agilent 1200 series liquid chromatograph system coupled to a 6410 series triple quadrupole mass spectrometer. The chromatographic conditions were: mobile phase, 50% v water:50% v methanol; flow rate, 0.4 mL/min; injection volume, 10 μL.

3. RESULTS AND DISCUSSION

Two different cluster catalysts were prepared by deposition of Ag and Au onto TiO2 powder supports. Figure 2 shows typical HAADF-STEM images (with three magnifications) and the corresponding cluster diameter distributions and X-ray photoelectron spectra of these two samples. In both cases, dense metal clusters are found well dispersed on the supports without severe agglomeration. As the intensity contrast in HAADF images reflects the Z number of the atom (where Z is the elemental atomic number), the observed Au clusters are much brighter than the Ag clusters. For the cluster diameter distribution, Ag and Au cluster catalysts show similar peak sizes, but it is slightly broader in the case of Ag. The peak size for the Ag cluster is around 5.9 and 5.3 nm for the Au cluster. Compared to the previously reported agitation method, the improved mixing method (mechanical stirring by a metal comb) exhibits a much better performance without forming large metal aggregates. Most clusters produced by the MACS are neutral, so they are expected to soft-land on the support. The BET results in Table S1 show that after the cluster deposition, the sample surface area and pore size are decreased as expected, compared to the blank TiO2 supports. This further confirms that metal clusters are well dispersed on the supports. ICP-OES and XPS were used to measure the cluster loading for these two samples, which calculated the average metal loading for the Ag catalyst to be 0.15 and 0.55 wt %, respectively, for Au catalyst, as seen in Table S1.

Figure 3 shows the Ag 3d and Au 4f high-resolution spectra for the Ag and Au cluster catalysts after air bubbling in the reactor. The absorbance spectra during the reaction (~1 min intervals), in the presence of Ag cluster catalyst on TiO2 support. (b) Comparison of nitrophenol conversion as a function of time over the two cluster catalysts as well as blank TiO2 support and without any catalyst. The obtained data were fitted by exponential equations. The inset values are the reaction rate constants. (c) Signal of maleic acid (m/z = 115) in the mass spectra obtained by LC/MS/MS as a function of time for the nitrophenol oxidation by O3 in the presence of Ag cluster catalyst and without catalyst.
exposure. The Ag 3d region for the Ag sample consists of two main doublet peaks in Figure 2d. The peak at 368.2 eV (green) corresponds to the metallic silver and is in good agreement with the literature.25 The peak with a higher intensity (red) located at ~1 eV lower binding energy coincides with the known peak position for silver oxide (AgO).26 Thus, a proportion of the Ag clusters in the Ag/TiO2 catalyst is, as expected, oxidized after air exposure. The clusters may be present as metallic silver and silver oxide. The Ag 3d region for the Ag sample consists of two known peak positions for silver oxide (AgO).26 Thus, a proportion of the Ag clusters in the Ag/TiO2 catalyst is, as expected, oxidized after air exposure. The clusters may be oxidized directly or at the interface with TiO2, as reported previously for Ru and Pd clusters on oxide supports.27 For Au in Figure 2h, fitting the Au 4f spectrum using asymmetric line shape shows the clusters are mainly in the metallic state. The Au 4f region shows well-separated spin–orbit components ($\Delta = 3.7$ eV). The main peak is located at 83.2 eV compared to 84 eV often reported in the literature.28 This negative shift of 0.8 eV in the binding energy can be explained by a reduction of the coordination number of Au atoms29 or by electron transfer from the TiO2 to the Au cluster. The other doublet peaks in the Au 4f spectra at 84.6 and 86.7 eV are commonly observed in Au clusters and attributed to Au1+ and Au3+ states, respectively.

The synthesized cluster catalysts were investigated in the oxidation of nitrophenol by O3. Catalysts (50 mg) were added to the solution followed by introducing O3 gas to the solution. The optical absorbance was recorded to monitor the progress of the reaction. Figure 3a shows typical UV–vis absorbance spectra for nitrophenol oxidation in the presence of Ag cluster catalyst. It can be seen that the peak intensity at 400 nm, which corresponds to the nitrophenol concentration,30 decreases rapidly over the reaction time, and eventually full conversion is achieved. Since nitrophenol can also be oxidized directly by O3 itself,31 the nitrophenol conversion rates over the two cluster catalysts, blank TiO2 support, and in the absence of any catalyst were all obtained and compared in Figure 3b. While 100% nitrophenol conversion was achieved in all of the cases, the complete conversion time was ~13 min without catalysts. It took 9.7 min for the blank TiO2 powder support and 7.6 (7.3) min for Ag (Au) cluster catalysts. This confirms that the cluster catalysts accelerate the reaction.

The mechanism of nitrophenol ozonation is complex, involving mass transfer, direct molecular ozone, and indirect (i.e., hydroxyl radical) reactions, which lead to the formation of several intermediates (e.g., dihydroxybenzenes and unsaturated carboxylic acids); they further react with ozone and hydroxyl radicals. As the reaction progresses over time, interference with the main reaction becomes significant. As a result, the kinetics of nitrophenol oxidation are considered only within the first few minutes of reaction. This was supported by following the kinetics of maleic acid formation, m/z 115, using liquid chromatography–tandem mass spectrometry (LC/MS/MS); maleic acid is a product of the main reaction of ozone with nitrophenol. As shown in Figure 3c, the formation of maleic acid peaks at about 4 min, followed by a decline, so competing reactions manifestly take place after about 4 min from the start of the experiment. A pseudo-first-order reaction with an apparent reaction rate constant, $k_{app}$, was used to describe the experimental data below 4 min in Figure 3c. The values of $k_{app}$ provide a basis to compare the rates of nitrophenol oxidation by the various systems used in the study. According to the experimental results, $k_{app}$ is 0.34 ± 0.01 min$^{-1}$ for ozone alone, and with the addition of Au (Ag) clusters, the rate constant increases to 0.59 ± 0.02 min$^{-1}$ (0.50 ± 0.01 min$^{-1}$). This confirms the addition of cluster catalyst accelerates the reaction. Since the loading of Ag is slightly lower than that for Au cluster catalyst, the normalized reaction rate constant, $k_{norm}$, shows that the Ag catalyst is more active than Au in Table S2. This behavior can be explained by the enhanced decomposition of ozone to strong oxidative radicals, such as *OH and O2*, on the Au3+ and Ag3+ catalyst surfaces and further promoted by the Lewis acid sites25 of AgO, identified in the characterization section.

The rate constants obtained here with the CBD catalysts prepared with the MACS (0.59 ± 0.02 min$^{-1}$ for Au and 0.50 ± 0.01 min$^{-1}$ for Ag) are higher than those reported with catalysts prepared by traditional chemical methods. For example, a calcined bauxite ore catalyst gave a pseudo-first-order rate constant of 0.194 min$^{-1}$,35 while the mesoporous α-MnO2 catalyst gave a rate constant of 0.052 min$^{-1}$.36 This is similar to the value of 0.055 min$^{-1}$ for a TiO2 catalyst37 and 0.0043 min$^{-1}$ for a titanium dioxide–magnetite composite.38 It can hence be concluded that the Ag and Au clusters on TiO2 catalysts are promising candidates for the heterogeneous catalytic ozonation reaction with application in water treatment and environmental protection.

The nitrophenol oxidation reaction was also performed at three different values of pH in Figure 4a. It can be seen that the degradation of nitrophenol is clearly enhanced at alkaline pH. The reaction reaches a conversion level of >95% in 5 min when the pH is ~9. When the pH is decreased to 5, the reaction takes almost double the time, ~9 min, to reach 95%, or ~11 min at pH 3. The strongly pH-dependent rate result is consistent with other reports35,36 that the decomposition of ozone into reactive radicals is catalyzed by OH−. The reusability of catalyst is one of the main advantages of heterogeneous catalysts compared to homogeneous catalysts. To investigate our catalyst reusability, the deposited Ag cluster catalyst was recovered and reused five times in succession, in Figure 4b, which shows the 4-nitrophenol conversion level...
after 2 min of reaction for each cycle. The conversion reached 60% each cycle, which indicates that the cluster catalyst has excellent stability. Error bars shown in the figure (~5%) were obtained from repeated experiments. Thus, no deactivation of the catalyst was apparent after reuse. Together, the results suggest that deposition of metal clusters on oxides is a promising new route to enhance the robust catalytic activity of heterogeneous catalysts for water ozonation.

4. CONCLUSIONS

In conclusion, the catalytic activities of physically deposited Ag and Au clusters on TiO2 were demonstrated in the oxidation of nitrophenol by ozone in aqueous solution. The cluster catalysts were prepared by a new MACS cluster beam deposition system and deposited onto commercial TiO2 powder at the gram scale. Characterization techniques, including STEM, BET, and XPS, revealed that clusters were densely deposited onto the support without severe surface aggregation. The two cluster catalysts show similar size distributions, with peak sizes of around 5.9 nm for the Ag catalyst and 5.3 nm for the Au. Catalytic results demonstrated that both the deposited Ag and Au cluster-based catalysts enhanced the catalytic activity compared to pure ozonation of nitrophenol and blank TiO2 powders. With respect to the catalysts produced by traditional chemical methods reported in the literature, the cluster catalysts made with the MACS exhibited at least comparably high activities. This study demonstrates a new route to prepare physically deposited cluster catalysts with high activities for heterogeneous catalysis in solution and specially the catalytic ozonation of a model pollutant for water treatment.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acsami.0c05955. Summary of the metal loading and BET results (Table S1); comparison of the intrinsic reaction rate constant normalized by the total number of moles of metal for the nitrophenol reduction catalyzed by Au and Ag cluster catalysts (Table S2) (PDF)

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Notes

The authors declare no competing financial interest.

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Theoretical Insight into the Mechanism of p-nitrophenol Degradation by OH Radicals in Presence of Magnetically Recoverable Fe3O4 Nanoparticles.

Calibration of Electron Spectrometers 5-Re-evaluation of the Original calibration of the ESCALAB 220i-XL.

On the Structure of Au/TiO2 Nanoparticles and their Catalytic Application to 4-nitrophenol Reduction.

Effective Degradation of Acid Orange 10 by Catalytic Ozonation in the Presence of Au-Bi2O3 Nanoparticles.

Interaction of Silver Nanoparticles with Ozone in the Presence of Au-Bi2O3 Nanoparticles.

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