The Simple Synthesis of High Metal Loading Single-Atom Catalysts via Chelate Fixation

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Abstract: Supported atomically dispersed metal catalysts are at the frontier of catalysis research, promising combined advantages of well-defined sites and high selectivity, with high stability and easy separation characterizing solid catalysts. We report that addition of an organic chelating agent in the final stage of impregnation, allows the metal precursor to be anchored and isolated without the need for anchoring sites on the support. The resulting isolated atoms are characterized with scanning transmission electron microscopy, X-ray photoelectron spectroscopy, and thermal desorption of carbon monoxide. This “chelate fixation” (CheFi) method can yield relatively high loadings of single atoms (e.g., Pt, Ru, Ir), up to at least 1 atom nm⁻² on a variety of carbon (e.g., black, activated carbon, diamond) and oxide (e.g., TiO₂, SiO₂) supports.

One-Sentence Summary:
A simple, novel synthesis method yields isolated atoms without the need for anchoring sites on the support.
Supported single-atom catalysts (SACs) are at the frontier of catalysis research, promising the combined advantages of homogeneous (e.g., well-defined site, high selectivity) and heterogeneous (e.g., high stability, easy separation) catalysts (1–7). Such well-defined materials have the potential to provide unique catalytic performance due to the intimate electronic interaction between the isolated active site and a given support material, which serves as a ligand. Some examples of such materials include a 2wt% Pt SAC for preferential oxidation of CO in hydrogen-rich fuel (8), a highly active, selective, and coke-resistant Rh single-atom catalyst dispersed in Cu for low-temperature, nonoxidative propane dehydrogenation (9, 10), and a stable heterogeneous single-atom Pd catalyst supported on graphitic carbon nitride for Suzuki coupling (11). A general limitation in these studies has been the inability to produce loadings of accessible single atoms that are relevant to industrial applications (12). In addition, in order to produce and retain isolated single metal atom sites in the resulting materials, novel and/or complicated synthesis methods have been employed (13–22). To fully exploit the potential for commercial applications of SACs, the development of scalable and facile synthesis methods that can produce high loadings of isolated active sites is required.

There are many synthetic methods available to deliver metal precursors to catalytic supports (i.e., oxides, carbon). For example, strong electrostatic adsorption (SEA), ensures an even distribution of metal ions on the support surface through control of solution pH (23–26). This method can reliably produce small metal nanoparticles (27, 28), as well as isolated single atoms if the metal loading is low enough (29, 30). However, at loadings greater than ~0.1 wt%, activation of such catalysts (i.e., through calcination and/or reduction treatments) results in cluster and nanoparticle formation (29, 31). One strategy to achieve high single metal atom loadings involves the use of chelating agents that complex with the metal precursor and are then pyrolyzed (sometimes along with other organic components) to produce a carbon material with single metal atoms dispersed throughout. However, there are some concerns that not all of the single atoms in such a carbon support matrix are accessible (1, 32), and the approach is limited to carbon. Another strategy is to employ supports with anchoring sites and to carefully tune the treatment protocols to maintain isolation, as has recently been demonstrated by Perez-Ramirez and co-workers (33).

Promising results have been reported using non-aqueous organic solvents to deliver metal salt and organometallic precursors to carbon support surfaces. Hutchings and co-workers (34) were able to achieve isolated Au, Pt, Pd and Ru up to a loading of 1 wt%, and demonstrated their activity for acetylene hydrochlorination. More recently, Gates and co-workers achieved isolated Ir sites for 14.8 wt% Ir/rGA which were effective for ethylene hydrogenation (35). Due to the instability of the employed precursors in aqueous solution, “extra dry” organic solvents and air exclusion technique were employed (34, 35). Indeed, solution removal at the last step of the incipient wetness impregnation (i.e., before catalyst activation) has been reported to form crystallites because of weak interactions and redistribution of precursors ions (36, 37). Even for SEA, incomplete drying of the adsorbed precursors on the support during activation (Fig. S1) results in the formation of large nanoparticles.

We hypothesized that formation of reduced metal clusters and particles (as opposed to isolated atoms), even when starting with well-dispersed, adsorbed precursors obtained with SEA, is significantly influenced by the presence of water (Fig. 1A). In addition to any bulk water that might remain in support pores even after drying, there is also a significant amount of water associated with each precursor in the form of a hydration sheath (38, 39). For example, PtCl$_6^{2-}$ is known to retain one hydration sheath, while Pt(NH$_3$)$_4^{2+}$ is known to retain two hydration sheaths, upon drying at room temperature. In order to retard (or eliminate) the negative influence of such
water on the formation of supported single atoms, a chelating agent was introduced immediately following the precursor adsorption step. This chelating agent, 8-hydroxyquinoline (8-HQ), has a high affinity for transition metals (40–43), and has been shown to adsorb on a variety of oxide and carbon surfaces for applications in metal extraction (44–46). We surmised that 8-HQ, which is not soluble in water, will isolate the metal precursor on the support surface without the need for defects or anchoring sites (Fig. 1B). The strong metal-8-HQ interaction coupled and high normal boiling point of 8-HQ (267°C), would then allow the metal ion to be reduced with H₂, while avoiding formation of clusters or nanoparticles. A comparison of SEA with this new chelate fixation (CheFi) method for Pt on VXC-72 and TiO₂ support substrates is provided in Fig. S2. It can be seen that there is significant particle/cluster formation for the SEA samples upon reduction in H₂ at 300°C, albeit with a very tight particle size distributions (Figs. S2B,D). However, the CheFi samples (Figs. S2A,C), primarily consist of isolated single atoms, along with a few agglomerates as dimers, trimers, etc. It was further found that SEA was not necessary for the CheFi method to be successful. A set of experiments conducted for VXC-72 at pH 2.9 (where maximum SEA of the Pt precursor occurs) and 9 (the point of zero charge where there is no SEA) shows identical results (Figs. S3A-B). In addition, contrary to SEA, the CheFi synthesis method proved successful at high ionic strength using both anionic and cationic Pt precursors (Figs. S3C-D). Because the chelating agent is not soluble in water, a polar solvent like acetone with a higher dipole moment than water is needed to dissolve the chelating agent prior to its addition to the impregnating solution.

Fig.1. Schematic of hypothesis and synthesis methods. (A) effect of water during drying and activation of the catalyst, (B) chelate fixation during drying and activation. Chemical formulas and structures (based on van der Waals spheres) prepared using tools available at https://molview.org/.

In this contribution, we demonstrate that this proposed CheFi method produces high densities (up to at least ~1 atom/nm²) of isolated single atoms of Pt, Ir, and Ru on a variety of carbon (e.g., black, activated carbon, diamond) and oxide (e.g., TiO₂, SiO₂) supports with no anchoring sites. The results open up a simple, cheap and scalable method to produce SACs, for both fundamental catalysis studies and practical industrial applications.

Aberration corrected scanning transmission electron microscopy (STEM) has been used to characterize the samples. Fig. 2A-F show the images of CheFi-prepared 0.8%Pt/VXC-72, 3.5%Pt/VXC-72, and 7.7%Pt/VXC-72 corresponding to densities of 0.1, 0.45 and 1.0 atom/nm². With a reduction temperature of 170°C the atoms are seen to be completely isolated (Figs. 2A-
C), while a higher reduction at 300°C maintains isolation (Figs. 2D,E), except for a small amount of agglomeration at the highest metal loading (Fig. 2F). Dispersing 1 atom/nm² on a Norit ROX activated carbon with a surface area of ~1200 m²/g resulted in 30%Pt/C SACs shown in Fig. 2G-H. The higher resolution images show well distributed single atoms. The higher temperature reduction (300°C) again causes a small amount of Pt-Pt coordination (Fig. 2H). The highest magnifications contain overlapping images of well-defined single atoms at the focal plane and blurry atoms off the focal plane. Fig. S4 shows the same area of sample with either the left-hand side (Fig. S4A) or right-hand side (Fig. S4B) in focus. Atoms are clearly resolved when any section of the sample is brought into focus. Such imaging strongly suggests that the metal atoms are not only present on the outer support surface, but within the support pore framework.

Further characterization has been performed with X-ray photoelectron spectroscopy (XPS) and temperature programmed desorption of carbon monoxide (CO-TPD). The XPS measurement of Pt 4f peaks of 2%Pt/VXC-72 prepared by standard dry impregnation after in-situ reduction in hydrogen, is shown in Fig. 3A. After reduction at 300°C only one peak at 71.2 eV is observed, and is associated with the Pt 4f of metallic Pt nanoparticles (47). Indeed, reduction of the sample to 600°C resulted in the same position of the Pt 4f peak, confirming it as a reference for Pt nanoparticles. A 2%Pt/VXC-72 catalyst prepared using CheFi was subjected to in-situ reduction at 170°C, 300°C, and 600°C (Fig. 3B). After reduction at 170°C, a Pt 4f peak showed up at the characteristic position of 72.4 eV for ionic Pt₁, attributed to single atoms (48), which is consistent with the associated STEM images (Fig. S5A). A small peak at 70.9 eV is attributed to formation of single atom Pt carbide atoms (49). The sample reduced at 300°C, demonstrated two Pt 4f peaks at both 72.4 eV and 71.2, indicating the presence of both ionic Pt₁ atoms and neutral Pt₁ atoms. Some of the neutral Pt atoms might be taking part in the formation of small nanoparticles, consistent once again with the associated STEM images (Fig. S5B). The 30% Pt/CNorit catalyst reduced at 100°C (Fig. 3C) resulted only in the characteristic Pt 4f peak for ionic Pt₁ at 72.4 eV, attributed to single atoms. Reduction at 170°C led in the formation of mostly neutral Pt₁, as indicated by the 71.1 eV peak (Fig. 3C). A higher reduction at 300°C again resulted mainly in neutral Pt₁ atoms (Fig. 3C). Although, the Pt₁ neutral state at 71.1 eV can be attributed to nps or Pt clusters, the STEM images (Figs. 2G, 2H) clearly show that after reduction at 170°C -300°C, the main feature on the surface is still Pt₁ single atoms. The CO-TPD spectra (described later) from the same samples (Fig. 3F), also indicate that the majority of Pt is not in the form of nanoparticles.

Computational studies (50) of Pt single atoms and dimers on carbon help to explain how ionic Pt₁, corresponding to single atoms, can become more electron rich and detected as neutral Pt₁ on carbon. The DFT-calculated density of states of Pt single atoms predicts that hydrogen adsorption on defect sites and other oxygen containing surface functionalities, results in the appearance of Pt d-band energy levels closer to the Fermi edge. This is consistent with the conductive nature of single atom Pt after in-situ reduction, as there is evidence from the O 1s (OH peak contributions Fig. 3D) that hydrogen is added on the substrate surface. A strong indication that the above prediction plays a major role in the current measurements, is also the fact that the effect of Pt neutralization upon reduction, is more evident on the defect-rich Norit ROX than on VXC-72. There are also predictions in the literature (51) which show that when Pt single atoms react with the carbon substrate, making stronger and shorter Pt-C bonds, the Pt charge increases. There is some evidence of stronger Pt-C interaction after reduction at 170 C (Figs. 3B, 3C) through the formation of small amounts of carbides.


Fig. 2. Various densities of single atom supported catalysts. STEM images of 0.8%Pt/VXC-72 CheFi, 3.5%Pt/VXC-72 CheFi, 7.7%Pt/VXC-72 (1 Pt atom/nm²) CheFi reduced at (A-C) 170°C, and (D-F) 300°C, 30%Pt/CNorit (1 Pt atom/nm²) CheFi-reduced at (G) 170°C, and (H) 300°C.
However, the effect of neutralizing the ionic Pt\(_1\) atoms is in a larger extent, which indicates that the addition of hydrogen on defects and carbon functionalities is the main reason for this observation. Computational studies (50) of Pt single atoms and dimers on carbon help to explain how ionic Pt\(_1\), corresponding to single atoms, can become more electron rich and detected as neutral Pt\(_1\) on carbon. The DFT-calculated density of states of Pt single atoms predicts that hydrogen adsorption on defect sites and other oxygen containing surface functionalities, results in the appearance of Pt d-band energy levels closer to the Fermi edge. This is consistent with the conductive nature of single atom Pt after in-situ reduction, as there is evidence from the O 1s (OH peak contributions Fig. 3D) that hydrogen is added on the substrate surface. A strong indication that the above prediction plays a major role in the current measurements, is also the fact that the effect of Pt neutralization upon reduction, is more evident on the defect-rich Norit ROX than on VXC-72. There are also predictions in the literature (51) which show that when Pt single atoms react with the carbon substrate, making stronger and shorter Pt-C bonds, the Pt charge increases. There is some evidence of stronger Pt-C interaction after reduction at 170°C (Figs. 3B, 3C) through the formation of small amounts of carbides. However, the effect of neutralizing the ionic Pt\(_1\) atoms is in a larger extent, which indicates that the addition of hydrogen on defects and carbon functionalities is the main reason for this observation.

The CO-TPD can further distinguish between SACs and catalysts containing nanoparticles, as it used here as a probe molecule that provides information about adsorption sites and energies (52). The results in Fig. 3E demonstrate the difference in CO desorption temperature from completely isolated SACs and nanoparticles. The sample from which the top (red) curve is obtained, contains only nanoparticles (Figs. 3A and S6). The bottom (black) curve shows the CO desorption profile from the 2%Pt/VXC-72 catalyst that was first reduced at 170°C. A main CO desorption peak appears at 75°C, and is attributed to desorption from single Pt atoms (47). A small shoulder at 130°C can be attributed to the Pt carbide in the catalyst(47), consistent with the XPS results (49). The middle (blue) curve shows the CO desorption profile for the 2%Pt/VXC-72 catalyst reduced at 300°C. Desorption peaks observed at 75°C and 205°C can be attributed to single atoms and nanoparticles, respectively. These observations are consistent with the STEM images of the catalyst treated at 170°C (Fig. S5A) and 300°C (Fig. S5B) the latter of which is a combination of isolated atoms and nanoparticles. The lower integrated area of the CO desorption peak is consistent with the loss of adsorption sites due to the formation of nanoparticles.

CO-TPD from the reduced 30% Pt/C\text{Norit} sample, shows the wide range of adsorption sites with different binding strengths, indicating that this catalyst will be effective in a wide range of reactions and catalytic processes. Fig. 3F (blue) gives the CO-TPD pattern of that sample reduced at 100°C. This sample is imaged in Fig. S7 and shows only isolated atoms. The CO-TPD pattern of this sample exhibits the peaks at 75°C and 130°C indicative of isolated atoms (atoms and Pt\(_1\) carbide). Further reduction at 170°C resulted in CO desorption at a wide range of temperatures (Fig. 3F dark green), expanding from 75°C to almost 300°C. These spectra indicate stronger interaction of the CO with the Pt atoms, which becomes even more pronounced after reduction at 300°C. The above-mentioned observations exclude the possibility of nanoparticle formation after reduction, because in that case there would be only a distinct CO desorption at ~220°C characteristic of CO adsorbed on nanoparticles. On the contrary the CO-TPD confirms predictions from DFT (50) that the changes in the Pt d-band center due to the addition of hydrogen on defects and surface functionalities, affects the binding strength of the CO adsorption on Pt single atoms and dimers. TPD observations are consistent with the associated STEM images in Fig. 2H which clearly show that there are no nanoparticles on the catalyst after
reduction (a total of 25 separate areas of the 30wt%Pt/C$_{\text{Norit}}$ were imaged, and none contained even a single nanoparticle). Finally, following the N 1s peak intensity in XPS (not shown here), we detected that as the reduction temperature increases, the chelating agent is removed, leaving Pt single atoms on the support with a wide variety of adsorption sites available to act as reaction active centers.

\[ \text{Fig. 3. In Situ XPS studies and CO TPD. In situ XPS spectra of (A) Pt 4f fresh, reduced at 170}^\circ\text{C, 300}^\circ\text{C, and 600}^\circ\text{C 2%Pt/VXC-72 DI, (B) Pt 4f fresh, reduced at 170}^\circ\text{C, 300}^\circ\text{C, and 600}^\circ\text{C 2%Pt/VXC-72 CheFi, (C) Pt 4f fresh, reduced at 100}^\circ\text{C, 170}^\circ\text{C, and 300}^\circ\text{C 30%Pt/C CheFi, (D) O 1s fresh, reduced at 100}^\circ\text{C, 170}^\circ\text{C, and 300}^\circ\text{C 30%Pt/C CheFi, (E) CO-TPD of 2%Pt/VXC-72 DI in situ reduced at 300}^\circ\text{C and CheFi in situ reduced at 170}^\circ\text{C and 300}^\circ\text{C, (F) CO-TPD of 30%Pt/C CheFi in situ reduced at 100}^\circ\text{C, 170}^\circ\text{C and 300}^\circ\text{C.}} \]

The CheFi method appears to be very versatile with respect to both supports and metals. This is perhaps not surprising, given that 8-HQ is well-known to chelate metal ions (41, 45, 53) or metal ion complexes with ligands (54–56). As noted previously, 8-HQ has been shown to adsorb on a variety of typical support surfaces (45, 46). In addition to VXC-72 (i.e., carbon black) and Norit ROX (activated carbon), Figures 4A and B show representative STEM images for Pt on
nanodiamond prepared with nitrate and chloride precursors, respectively. The result shows that single Pt atoms can be produced on a carbon support that has no surface functional groups. Silica (Fig. 4C and Fig. S8) and titania (Fig. 4D) are two other common industrial oxide support substrates for which CheFi also yielded isolated Pt atoms. Finally, Figures 4E and 4F show isolated metal atoms in 3.5%-Ir and 1.8%-Ru on VXC-72 prepared by the CheFi method.

Fig. 4. Versatility of the CheFi method. (A) 2%Pt/Diamond (nitrate precursor), (B) 1% Pt/Diamond (chloride precursor), (C) 0.78% Pt/SiO$_2$, (D) 1.9% Pt/TiO$_2$, 3.5%Ir/VXC-72, and 1.8%Ru/VXC-72.

This facile, inexpensive method which does not rely on anchoring sites should be generalizable to many metal precursors and supports to prepare SACs at high metal loading.

References and Notes


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Supplementary Materials
Materials and Methods
Supplementary Text
Figs. S1 to S9