Removal of Volatile Organic Compounds Driven by Platinum Supported on Amorphous Phosphated Titanium Oxide

HUANG Xie-Yi\textsuperscript{1,2,4}, WANG Peng\textsuperscript{2,3,4}, YIN Guo-Heng\textsuperscript{1}, ZHANG Shao-Ning\textsuperscript{1}, ZHAO Wei\textsuperscript{1}, WANG Dong\textsuperscript{1}, BI Qing-Yuan\textsuperscript{1}, HUANG Fu-Qiang\textsuperscript{1,4}

(1. State Key Laboratory of High Performance Ceramics and Superfine Microstructure, Shanghai Institute of Ceramics, Chinese Academy of Sciences, Shanghai 200050, China; 2. University of Chinese Academy of Sciences, Beijing 100049, China; 3. School of Physical Science and Technology, ShanghaiTech University, Shanghai 200031, China; 4. State Key Laboratory of Rare Earth Materials Chemistry and Applications, College of Chemistry and Molecular Engineering, Peking University, Beijing 100871, China)

Abstract: Development of high efficiency catalyst is the key factor to catalytic combustion of volatile organic compounds (VOCs). Herein, amorphous mesoporous phosphated TiO\textsubscript{2} (ATO-P) with high specific surface area supported platinum catalyst was successfully fabricated. P-dopant can increase the surface area (up to 278.9 m\textsuperscript{2}g\textsuperscript{-1}) of ATO-P, which is 21 times higher than that of pristine TiO\textsubscript{2}, and make the amorphous titanium oxide structure. The supported Pt catalyst with amorphous mesoporous feature shows impressive performance and excellent thermostability for VOCs oxidation. The Pt/ATO-P catalyst exhibits outstanding catalytic efficiency, the $T_{50}$ and $T_{90}$ (temperatures required for achieving conversions of 50\% and 90\%) are respectively 130°C and 140°C, for toluene oxidation under high gas hourly space velocity (GHSV) of 36 000 mL\cdot h\textsuperscript{-1}g\textsuperscript{-1} and toluene concentration of 10 000 mL\cdot m\textsuperscript{-3}. The performance is superior to the reference Pt/TiO\textsubscript{2} and comparable with the state-of-the-art catalysts. These findings can make a significant contribution on the new applications of amorphous mesoporous phosphated materials in VOCs removal.

Key words: amorphous mesoporous structure; phosphated TiO\textsubscript{2}; Pt nanoparticles; toluene oxidation; VOCs removal

Volatile organic compounds (VOCs), like toluene, benzene, esters, and hydrocarbons, are emitted from various industrial sources which can cause serious environmental pollution and health problems\textsuperscript{[1-2]}. Toluene, one kind of toxic and strong carcinogenic chemicals, is frequently used in making paints, adhesives, rubbers, and leather tanning processes because of its excellent ability to dissolve organic substances\textsuperscript{[3-4]}. However, toluene is difficult to degrade due to its stable structure\textsuperscript{[5]}. Several techniques, such as physical and chemical adsorption, photocatalytic and catalytic oxidation methods, are widely used for the combustion of VOCs\textsuperscript{[6-7]}. Among them, catalytic oxidation is regarded as a promising approach owing to its high efficiency and convenient operating conditions\textsuperscript{[8]}. Researches on catalysts for toluene oxidation have been conducted, including noble metal and metal oxides catalysts\textsuperscript{[9-10]}. Due to the significant reduction on activation energy during the catalytic oxidation process, noble metal based catalysts, such as Pt, Pd, Au, Rh, and Ir have shown impressive performance in toluene removal\textsuperscript{[11-13]}. It was found that supported Pt catalysts showed the best catalytic performance com-
pared with other noble metals\cite{14-15}. It should be pointed out that the supports play an important role in the catalytic reaction processes\cite{16-18}. Many works have been focused on the metal-support interaction by studying the catalytic properties of TiO$_2$, Al$_2$O$_3$, ZrO$_2$, and ZnO supported Au nanoparticles\cite{19}, and the shape effect of Pt/CeO$_2$ catalysts\cite{10}. Nevertheless, most supports suffer from low specific surface area and few active sites, which are crucial for the overall catalytic activity.

Due to high specific surface area and variable valence, amorphous materials have been attracted increasing interests in VOCs oxidation. And the numerous defects in amorphous structures can offer large quantities of oxygen vacancies, which are beneficial for the adsorption of oxygen and organic molecules. Lee et al.\cite{20} reported that carbon black supported amorphous MnO$_x$ is highly efficient for oxygen involved reaction. Wang et al.\cite{21} found that amorphous MnO$_x$ modified Co$_3$O$_4$ can enhance the catalytic activity for the VOCs oxidation. It was demonstrated that the amorphous structure of bimetallic Pd-Pt/Al$_2$O$_3$-TiO$_2$ could provide more vacancies and active sites for catalytic combustion\cite{22}. Therefore, the amorphous catalysts could show a tremendous potential in practical catalytic reactions. However, it is still a challenge to develop highly active and robust catalysts based on the amorphous materials for the oxidation of VOCs.

Herein, we demonstrate an efficient Pt/ATO-P catalyst for the catalytic removal of VOCs under high gas hourly space velocity (GHSV) and high substrate concentration. It should be pointed out that incorporating phosphorus into the framework of TiO$_2$ is a widely applied strategy for obtaining amorphous mesoporous feature\cite{23-24}. And the P element can stabilize the TiO$_2$ framework and significantly increase the specific surface area\cite{24}.

1 Experimental

1.1 Preparation of Sample

1.1.1 Preparation of support

All regents were of analytical grade and were used without any purification. 3 mL of tetrabutyl titanate was dissolved in 30 mL of ethanol at room temperature, which was marked as solution A. Then 0.125 mL of phosphoric acid (H$_3$PO$_4$) was subsequently dropwisely added into solution A with stirring to form a homogenous mixture, and kept stirring for 24 h. The obtained white solid products were separated by centrifuge and washed by deionized water and ethanol several times, followed by freeze drying overnight. The as-prepared products were calcined in air at 400 °C for 4 h at a heating rate of 5 °C·min$^{-1}$.

1.1.2 Preparation of catalyst

The ATO-P supported platinum (Pt/ATO-P) sample was prepared via impregnation method. A desired amount of ATO-P was transferred into aqueous solution containing appropriate amount of chloroplatinic acid (H$_3$PtCl$_6$). Subsequently, the samples were impregnated at room temperature for 12 h. After drying out the H$_2$O at 80 °C, the samples were treated at 350 °C for 2 h with a H$_2$/Ar mixture (5/95, v/v).

1.2 Characterization

XRD characterization of the samples was carried out on a German Bruker D8 Advance X-ray diffractometer (XRD) using the Ni-filtered Cu Kα radiation at 40 kV and 40 mA. Nitrogen adsorption-desorption isotherms were measured at −196 °C on a Micromeritics ASAP 2460 analyzer. Samples were degassed at 120 °C for 24 h prior to the measurement. The specific surface area of the samples was calculated using the Brunauer–Emmett–Teller (BET) method with the adsorption data at the relative pressure (P/P$_0$) range of 0.05–0.2. The total pore volumes were estimated at P/P$_0$=0.99. The pore size distribution (PSD) curves were calculated from the adsorption branch using Barrett–Joyner–Halenda (BJH) model. The prepared materials were pressed into tablets with KBr powder and then detected by FTIR (Perkin Elmer, USA) in the scanning range from 400 to 4000 cm$^{-1}$. SEM images were obtained by Hitachi-S4800. A JEOL 2011 microscope operating at
200 kV equipped with an EDX unit (Si(Li) detector) was used for the transmission electron microscope (TEM) and high resolution transmission electron microscope (HRTEM) investigations. The samples for TEM testing were prepared by dispersing the powder in ethanol and applying a drop of very dilute suspension on carbon-coated grids. XPS data were recorded with a Perkin Elmer PHI 5000 C system equipped with a hemispherical electron energy analyzer. The spectrometer was operated at 15 kV and 20 mA, and a magnesium anode (Mg Kα, hv = 1253.6 eV) was used. The C1s line (284.6 eV) was used as the reference to calibrate the binding energies (BE). TG measurements were conducted on a Netzsch STA 449C TG-DSC thermoanalyzer. The flow rate of the carrier gas (air) was 30 mL·min⁻¹. The temperature was raised from room temperature to 800 °C at a ramp rate of 10 °C·min⁻¹. Prior to H₂-TPR test, the sample (100 mg) was pretreated at 200 °C for 2 h and cooled to 50 °C in the flowing He. TPR experiment was carried out in 5 vol% H₂/He flowing at 30 mL·min⁻¹, with a ramping rate of 5 °C·min⁻¹ to a final temperature of ca. 800 °C. The signal was monitored using a TCD detector.

1.3 Catalytic activity test

The catalytic activity of samples was evaluated in a continued-flow fixed-bed quartz reactor with 50 mg catalyst. Toluene was introduced into the reactor with bubbling toluene solution in ice bath with pure air. The concentration of toluene was about 10 000 mL·m⁻³, and the flow rate was kept at 30 mL·min⁻¹ by a mass controller, equivalent to a gas hour space velocity (GHSV) of 36 000 mL·h⁻¹·g⁻¹. After steady operation for 100 min, the activity of the catalyst was tested. Toluene concentration was detected by a gas chromatograph equipped with a flame ionization detector. The toluene conversion (X_toluene) was calculated according to the equation:

\[ X_{\text{toluene}} = \frac{C_{\text{in}} - C_{\text{out}}}{C_{\text{in}}} \times 100\% \quad (1) \]

where \( C_{\text{in}} \) and \( C_{\text{out}} \) are the inlet and outlet toluene concentrations, respectively.

2 Results and discussion

2.1 Physicochemical properties of ATO-P support

Figure 1 displays the schematic diagram of amorphous ATO-P prepared via facile co-precipitation. XRD patterns of ATO-P and TiO₂ are shown in Fig. 2. All diffraction peaks of basic TiO₂ sample are indexed to anatase phase (JCPDS 21-1276). Interestingly, there is no TiO₂ crystal phase observed for ATO-P sample (Fig. 2), suggesting that ATO-P sample is typically amorphous and phosphorus dopant can markedly restrain the crystallization of anatase²⁵⁻²⁶.
temperature range of 20–900 °C. The first DSC peak at 30–80 °C is due to the release of physical adsorbed water. When all the water is released, Ti–OH and HPO$_4^{2-}$ groups start to condense[27]. These processes occur simultaneously in the temperatures of 100–220 °C (1.927% of weight loss) and 220–516 °C (0.7% of weight loss), resulting in an overlap of the TG data. There is no further weight loss up to 516 °C. The DSC curve shows two exothermic peaks at 704 °C and 781 °C, corresponding to a two-step exothermic transformation of ATO-P into a crystalline phase.

![TG and DSC curves for ATO-P](image)

Fig. 3 TG (solid line) and DSC (dashed line) curves for ATO-P

Figure 4a and 4b show the SEM images of ATO-P. The ATO-P nanoparticles are homogeneously dispersed with the particle size of ~20 nm, and the sizes are similar to that of TiO$_2$ (Fig.S1a). HRTEM was employed to characterize the nanostructure of samples. No porous structure is observed in the HRTEM image of TiO$_2$ (Fig.S1b), while various porous structure is shown in ATO-P (Fig.4c). Moreover, the pores of ATO-P are uniform and the average diameter is around 10 nm. EDS elemental mappings indicate that the P element is homogeneously distributed in ATO-P.

![SEM and HRTEM images of ATO-P](image)

Fig.4 SEM (a, b) and HRTEM (c) images, and EDS elemental mapping (d) of ATO-P

(Fig.4d). It is found that H$_3$PO$_4$ owns unique effects for synthesizing amorphous mesoporous phosphated TiO$_2$[28-29].

As shown in Fig. 5, the obtained ATO-P sample shows a characteristic type-IV isotherm with clear hysteresis loop located at the $P/P_0$ range of 0.45–1.0, showing the existence of a large amount of mesopore. Notably, the specific surface area of 278.9 m$^2$·g$^{-1}$ for ATO-P is 21 times higher than that of pristine TiO$_2$. The pore diameters of ATO-P center around 10 nm (Fig.5 and Table 1), which is consistent with HRTEM result (Fig.4c).
appears in series ATO-P, which may result from the incorporating effect of phosphorus dopant. There is no distinct peak over the range of 700–800 cm\(^{-1}\) (Fig. 6), indicating the absence of P–O–P groups in the amorphous mesoporous phosphated TiO\(_2\). Therefore, the P element is incorporated into the frameworks of ATO-P by forming Ti–O–P bonds\(^{[24]}\).

![Fig. 6 FTIR spectra of TiO\(_2\) and ATO-P](image)

As shown in Fig. 7a, the full XPS spectra indicate the existence of P in ATO-P. High-resolution XPS spectra of P 2p, Ti 2p and O 1s are depicted in Fig. 7b–d. The peak of P 2p of ATO-P is at 134.0 eV, suggesting that phosphorus in ATO-P gives a pentavalent oxidation state of P\(^{5+}\). No peak observed at 128.6 eV, which is the characteristic binding energy of P2p in TiP, indicating the absence of Ti–P bonds in ATO-P samples. As depicted in Fig. 7c, the peaks of Ti 2p\(_{3/2}\) and Ti 2p\(_{1/2}\) in ATO-P show remarkable blue-shift owing to the incorporation effect of phosphorus element. Figure 7d shows the XPS spectra of O 1s signals of TiO\(_2\) and ATO-P. The single peak at 529.5 eV is corresponded to the oxygen in Ti–O bond of TiO\(_2\). However, the O 1s spectrum of ATO-P contains two peaks at 531.4 and 532.9 eV, which are contributed to Ti–O–P bond and O–H bond, respectively\(^{[32,33]}\).
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2.2 Physicochemical properties of Pt/ATO-P catalysts

Figure 8c shows the XRD patterns of Pt/ATO-P and Pt/TiO$_2$ catalysts. The amorphous structure is still remained for Pt/ATO-P sample. However, no diffraction pattern of Pt nanoparticles is observed, indicating that the Pt nanoparticles are quite small and/or the Pt species are highly dispersed on the ATO-P surface. These results are well consistent with the HRTEM data above mentioned in Figs. 8a and b.

The results of XPS analysis of Pt/ATO-P and Pt/TiO$_2$ samples are depicted in Fig. 8d. It is known that the positions of Pt 4f$_{7/2}$ bind energy at 71.1, 72.4, and 74.2 eV are attributed to Pt$^0$, Pt$^{2+}$, and Pt$^{4+}$ species, respectively[34]. Similiar XPS profiles are rendered as the indication of a mixture of various valence states for Pt species over the small Pt nanoparticles. The existing of Pt$^{6+}$ species reflects the strong metal-support interaction (Pt–ATO-P), especially the prominent electronic interaction between active Pt and underlying phosphated TiO$_2$ support[35]. This probably due to the changes of the metal-support interaction by doping phosphorus atoms which can make an obvious effect on Ti–O–P frameworks.

The H$_2$-TPR profiles depicted in Fig.S3 show that there are two H$_2$-consumption peaks at low and high temperature attributed to weak and strong interaction of Pt and supports, respectively[36]. Notably, two reduction peaks of Pt/ATO-P catalyst at 78 and 601 °C show stronger intensity than that of Pt/TiO$_2$ at 72 and 433 °C, indicative of strong Pt-support interaction for Pt/ATO-P. These results are consistent with the XPS data.

2.3 Removal of VOCs by Pt/ATO-P catalysts

The catalytic efficiencies are depicted in Fig.9. It is clearly observed that reaction temperature can enhance the performance of Pt/ATO-P catalyst. The $T_{50}$ and $T_{90}$ are widely used to evaluate the catalytic performance[37]. As shown in Fig. 9a, Pt/ATO-P shows the excellent catalytic activity. $T_{50}$ and $T_{90}$ values for toluene combustion are 130 °C and 140 °C, which are much lower than those of Pt/TiO$_2$ with $T_{50}$ and $T_{90}$ of 160 °C and 190 °C, respectively. Combined with the above XPS data (Fig.8d), it can be concluded that the existence of phosphorus component plays an
important role in electronic structure of the active Pt species underlying amorphous mesoporous ATO-P support and thus the catalytic oxidation removal of toluene over Pt/ATO-P catalyst.

It is well known that noble metal loading significantly affects the catalytic behavior for many reactions. Pt/ATO-P catalysts with different Pt loadings were examined, and the results are depicted in Fig.10. Compared with 0.5 wt% and 2 wt%, the Pt loading of 1 wt% shows better performance (lower $T_{50}$ and $T_{90}$) for toluene oxidation. The low catalytic activity of 0.5 wt% Pt/ATO-P results from low density of active platinum nanoparticles anchoring on the surface of ATO-P support. For the Pt/ATO-P catalyst with Pt loading up to 2 wt%, larger size of Pt nanoparticles (~5 nm) can be obtained (Fig.54). Larger Pt particles can not only decrease the dispersion of Pt species but also lead to a weaker metal-support (Pt/ATO-P) interactions, thus result in the poor activity.

Stability is critical for the catalysts on the practical application. 1 wt% Pt/ATO-P exhibits excellent thermal stability for toluene oxidation over a 50-h period on stream at 180 °C without visible loss of activity, as shown in Fig. 9b. The toluene conversion remains a high level of 95.4% at the end of reaction process and maintains near full selectivity to final products of CO$_2$ and H$_2$O. The excellent stability of Pt/ATO-P catalyst is attributed to the unique geometric structure of crystalline Pt nanoparticles and amorphous mesoporous phosphated TiO$_2$ with prominent electronic interaction. For the used 1 wt% Pt/ATO-P, TEM measurement and XPS analysis (Figs. S5 and S6) demonstrate no significant change on the morphology, average size of Pt nanoparticles, and the chemical oxidation state of active Pt species. These results suggest the robustness of Pt/ATO-P catalyst for toluene oxidation removal under a relatively mild thermal process.

Given the superb thermocatalytic performance for 1 wt% Pt/ATO-P catalyst toward toluene oxidation, we were curious to examine whether the engineered material would also catalyze the removal of a class of VOCs, especially the complete oxidation of benzene, n-hexane, ethyl acetate, and mesitylene. As depicted in Fig.11, the $T_{50}$ values for the catalytic oxidation of benzene, ethyl acetate, n-hexane, and mesitylene are 216 °C, 331 °C, 271 °C, and 200 °C, respectively. Notably, high temperature is required for ethyl acetate conversion at 90% due to its strong structural stability [39-40]. These results show a broad scope toward
catalytic combustion involving troublesome organic compounds over Pt/ATO-P and indicate that the Pt/ATO-P catalysts can provide a new insight for the oxidation of VOCs.

![Fig.11 Catalytic activity of Pt/ATO-P for the conversion of benzene (a), ethyl acetate (b), n-hexane (c), and mesitylene (d) with respect to reaction temperature](image)

3 Conclusions

In summary, we successfully fabricated the amorphous mesoporous phosphated TiO₂ supported platinum catalysts for efficient removal of volatile organic compounds. The electronic modifications of supported Pt nanoparticles for the underlying amorphous ATO-P material and Pt loading for the whole catalyst were systematically investigated. The phosphorus dopant played an important role for stabilizing the inflated Ti–O–P frameworks as well the electronic structure of Pt species. Compared with pristine TiO₂, ATO-P with high specific surface area showed significant enhancement for Pt/ATO-P samples for catalytic overall oxidation of toluene under practical conditions. The performance of the engineered Pt/ATO-P for toluene combustion was superior to the reference Pt/TiO₂ and comparable with the state-of-the-art catalysts. Additionally, Pt/ATO-P catalyst exhibited excellent stability for toluene oxidation removal under a relatively mild thermal process and could be potentially applied in a broad scope of VOCs. The present work will make a significant contribution on the new applications of amorphous mesoporous phosphated materials in VOCs removal.

Supporting Materials

Supporting Materials related to this article can be found at https://doi.org/10.15541/jim20190154

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掺磷非晶氧化钛负载铂用于高效催化氧化挥发性有机化合物（VOCs）

黄谢意 1,2，王鹏 3,4，尹国恒 1，张绍宁 1，赵伟 1，王东 1，毕庆员 1，黄富强 1,3,4（1. 中国科学院上海硅酸盐研究所，高性能陶瓷和超微结构国家重点实验室，上海 200050；2. 中国科学院大学，北京 100049；3. 上海科技大学物理科学与技术学院，上海 200050；4. 北京大学化学与分子工程学院，稀土材料化学及应用国家重点实验室，北京 100871）

摘 要：高活性催化剂是挥发性有机化合物（VOCs）催化氧化消除的关键因素。本研究通过简单的共沉淀法制备了具有高比表面积的非晶介孔掺杂氧化钛负载铂催化剂（Pt/ATO-P）。通过P掺杂，既可获得非晶介孔结构，又可获得高ATO-P比表面积（可达 278.9 m²·g⁻¹）。非晶介孔Pt/ATO-P催化剂显示出优异的VOCs催化氧化性能和良好的热稳定性。Pt/ATO-P样品在空速为36 000 mL·h⁻¹·g⁻¹、甲苯浓度为10 000 mL·m⁻³的反应条件下，对甲苯催化氧化的T₅₀和T₉₀（实现50%和90%转化率所需的温度）分别为130 °C和140 °C，明显优于无磷催化剂Pt/TiO₂。这些发现为拓展非晶介孔磷化材料在环境净化和能源转化等领域的应用提供有效参考。

关 键 词：非晶介孔材料；磷掺杂非晶氧化钛；铂纳米颗粒；甲苯催化氧化；VOCs 消除

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Supporting Materials:

Fig. S1 SEM (a) and HRTEM (b) images of TiO₂

Fig. S2 HRTEM image of Pt/ATO-P

Fig. S3 H₂-TPR profiles of Pt/TiO₂ and Pt/ATO-P samples

Fig. S4 TEM images of 0.5 wt% Pt/ATO-P (a) and 2 wt% Pt/ATO-P (b), respectively

For the catalyst with low loading of 0.5 wt%, there are few Pt nanoparticles on the ATO-P supports. For the Pt/ATO-P catalyst with Pt loading up to 2 wt%, the Pt particles are larger than that with loading of 1 wt%.
Fig. S5 TEM (a) and HRTEM (b) images of the used Pt/ATO-P catalyst

Fig. S6 XPS Pt 4f spectra of the fresh and used Pt/ATO-P catalysts