Study of the structure and characteristics of mesoporous TiO₂ photocatalyst, and evaluation of its factors on gaseous formaldehyde removal by the analysis of ANOVA and S/N ratio

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This study differs from previous studies of TiO₂/SiO₂ in that 0.5–10 μm microsized TiO₂-rutile based catalysts (TR catalysts) with varying proportions of titanium and silicon were synthesized using a one-step modified hydrothermal method. At Ti/Si = 1/9, a two-dimensional channel-structured catalyst with a morphology resembling that of SBA-15 was obtained. In contrast, at Ti/Si = 3/7 or 5/5, a three-dimensional porous structure was formed, and Ti–O–Si–C bonds appeared. The structure of the TR catalyst transformed due to the decrease in C–Si bond content and the increase in C–C bond content with increasing Ti/Si ratio. The results indicated that the rutile phase was the main crystal phase of the TR catalyst. The small crystal size and large rutile phase content of the mesoporous TR catalyst contributed to the low band gap energy below 3.0 eV. Under 2 × 10 W lamp irradiation with either UVA or visible light, the three TR catalysts showed better formaldehyde (HCHO) removal efficiency than P25. Furthermore, the Taguchi method was employed to evaluate the catalytic factors by analysis of variance (ANOVA) and S/N ratio. The results revealed the contributions of each of the three factors to HCHO removal efficiency over TR catalysts to be as follows: space velocity (62%), Ti ratio (32%), and time on stream (5%). The TR catalyst with Ti/Si = 1/9 showed good HCHO removal efficiency with a high S BET (787.1 m² g⁻¹) and large pore volume (0.95 cm³ g⁻¹) for a residence time of over 2.29 × 10⁻² s under visible light irradiation. Microwave-assisted EG reduction was successfully applied to dope a TR catalyst with nanosized Pt particles in a short synthesis time. After Pt doping, the removal efficiency in the stream improved and stabilized. The Pt particles were Pt⁰ and proved effective for improving the photocatalytic removal of HCHO over the TR catalyst by prolonging the separation time of the electron–hole pairs. Overall, the Pt/TR catalyst is a potential material for pollutant removal and can be easily separated from the pollutant removal system since the catalysts are microsized.

1. Introduction

With improvement in people's social living conditions, the pace of building renovations has increased. The pollution emitted by building-material manufacturing plants as well as furniture, decorations, and building materials used in homes all contain volatile organic compounds that indirectly release organic pollutant gases, deteriorating indoor air quality and damaging personal health. Considering that humans spend 80% of their time indoors,¹ such as in offices, laboratories, or homes, improving the indoor air quality has become one of the most significant problems of this century. Of the various indoor pollutants, formaldehyde [HCHO] has attracted the most attention.²

To efficiently remove hazardous organic gases, their deep oxidation and conversion to harmless CO₂ and H₂O is better than their adsorption since the absorbed organic gases may desorb due to some indoor factors such as temperature, moisture, volatilization of gases, and saturated absorption of the material. Among the various oxidation methods, photocatalytic oxidation has the potential to remove indoor organic gases with low energy consumption. Titanium dioxide (TiO₂) is a classic photocatalyst; however, due to its extremely low activity under visible light, TiO₂ reactions are performed only under UV light, which comprises less than 5% of the sun's total light output. If catalytic materials can be modified to function under visible light, photocatalysis can be extended to indoor applications and related uses.

Common modification methods include (1) addition of noble metals such as Pd, Au, Ag, and Pt³–⁵ (2) combining TiO₂ and other semiconductors such as CdS, CdSe, WO₃, ZnO, SnO₂, C₃N₄⁶–¹⁰ (3)
doping transition metals such as Co, Fe, Cu, and Cr;\textsuperscript{12–13} (4) adding nonmetallic molecules, e.g., C, N, S, and F;\textsuperscript{14–18} and (5) increasing the sensitization of TiO\textsubscript{2} surface.\textsuperscript{19,20} Considering this, the current research directions involving surface modification/doping are as follows:\textsuperscript{17–20} (1) modification of absorption wavelength to shift the absorption range of photocatalyst from UV to visible wavelength; (2) reducing the band gap between the valence band and the conduction band; (3) effective electron–hole pair separation methods; (4) controlling reaction selectivity \( \sigma \) to reduce the production of intermediate products and increase the production of final products.

Besides the above studies, some investigations focused on the development of efficient photocatalysts with high specific surface areas \( S_{BET} \) and large pore volumes, such as TiO\textsubscript{2} hollow spheres,\textsuperscript{21} TiO\textsubscript{2} nanofiber,\textsuperscript{22} Zr–TiO\textsubscript{2} monolith,\textsuperscript{23} 3D TiO\textsubscript{2} foam,\textsuperscript{24} and mesoporous silica supported TiO\textsubscript{2} catalyst.\textsuperscript{25}

An ideal structure of a catalyst contributes to efficient catalysis by improving the four underlying catalytic mechanisms—absorption, migration, oxidation/reduction, and desorption.\textsuperscript{26–28} Santa Barbara Amorphous (SBA15), one of the silica family materials, has been modified with Ti atoms for broad-spectrum photocatalysis in recent years. Various mesoporous Ti-SBA15 catalysts have been synthesized by post-synthesis,\textsuperscript{25} sol-gel method,\textsuperscript{26} hydrothermal process,\textsuperscript{27–28} impregnation,\textsuperscript{29,30} grafting method,\textsuperscript{31} and gas deposition.\textsuperscript{32} These studies concluded that Ti-SBA15 synthesized with a well ratio of Si/Ti was advantageous for high photocatalytic activity such as high Ti/Si precursors, TEOS and TTIP, were pre-mixed in HCl solution, which was stirred at 20 °C. P123 was dissolved in HCl solution by agitation for 30 min. The solution of TEOS and TTIP was then added to the acidic solution of P123, and the mixture was heated at 100 °C for 24 h. The obtained material was filtered and washed with deionized water until \( \text{pH} = 7 \). Finally, the sample was calcined at 600 °C to remove the organic template and improve crystallinity. Three TR catalysts, TR(10), TR(30), and TR(50), were prepared with Ti/Si ratios of 1/9, 3/7, and 5/5, respectively.

2. Experimental

2.1. Synthesis of TR photocatalysts

TiO\textsubscript{2}-rutile based (TR) photocatalysts were synthesized using the modified hydrothermal method reported for SBA15 preparation. TR catalysts were obtained from the reaction of triblock copolymer Pluronic P123 (EO\textsubscript{20}PO\textsubscript{70}EO\textsubscript{20}, Aldrich), tetraethyl orthosilicate (TEOS, 98%, Aldrich), and titanium tetraisopropoxide (TTIP, 97%, Aldrich) under acidic conditions (HCl, 2 M) via the modified hydrothermal method. Ti/Si precursors, TEOS and TTIP, were pre-mixed in HCl solution, which was stirred at 20 °C. P123 was dissolved in HCl solution by agitation for 30 min. The solution of TEOS and TTIP was then added to the acidic solution of P123, and the mixture was heated at 100 °C for 24 h. The obtained material was filtered and washed with deionized water until \( \text{pH} = 7 \). Finally, the sample was calcined at 600 °C to remove the organic template and improve crystallinity. Three TR catalysts, TR(10), TR(30), and TR(50), were prepared with Ti/Si ratios of 1/9, 3/7, and 5/5, respectively.

2.2. Synthesis of Pt/\( \text{TR} \) photocatalysts

Pt/\( \text{TR} \) photocatalysts were prepared by a microwave-assisted EG reduction method. The Pt precursor (\( \text{H}_{2}\text{PtCl}_{6} \cdot 6\text{H}_{2}\text{O} \)) was added to EG and evenly stirred. This mixture was then added into the TR photocatalyst solution, prepared as mentioned above. After sufficient stirring, it was heated in a microwave at 500 W for 60 s. Subsequently, the Pt/\( \text{TR} \) photocatalyst was rinsed in alcohol and dried in an oven at 110 °C. Finally, the product was calcined to obtain the Pt/\( \text{TR} \) catalyst.

2.3. Activity tests

Pre-prepared imitation gas was released into a catalytic reactor to initiate photocatalytic reaction. The catalytic reactor used in this study had an internal diameter of 20 mm, an external diameter of 22 mm, and a length of 310 mm. The reactor was placed upright to prevent bypass flow caused by catalyst sedimentation and to provide actual catalytic environment with minimal external interference from concentration and temperature gradients. A quartz filter plate was inserted into the quartz reaction tube to prepare the catalyst for photocatalytic reaction. Furthermore, the gaps in the filter plate allowed sufficient gas molecules to pass through. Following the reaction, portable gas analyzers (MIC-800) and Agilent gas chromatographs (Agilent GC-7890) were used to detect the change in gas concentration before and after the reaction. The data were entered online immediately using a personal computer. Finally, the reactor was irradiated with two low-power lamps from both the sides of reactor. The UVA light source was a T8 10 W black light lamp (365 nm), and the visible light source was a 10 W fluorescent lamp.
The following activity tests were performed in this study. First, TR(10) catalyst was tested for HCHO removal and compared with P25 under UVA irradiation. A mixture of HCHO/oxygen in He + N₂ was chosen to represent one of the organic pollutants. The HCHO was carried by He from a saturator filled with liquid HCHO. The concentration of HCHO was 40 ± 10 ppm, which was adjusted by the saturator temperature and mixed air flow rate. Second, three TR catalysts were tested under visible light irradiation. One of the catalysts with superior performances chosen for the study of optimum space velocity in a later experiment. Further, two TR catalysts with superior performances were chosen for Pt/TR catalyst preparation and HCHO removal. The HCHO removal efficiency was calculated from the pollutant consumption obtained by comparing the inlet and outlet concentrations of the pollutant.

2.4. Characterization of catalysts

Particle size, morphologies, and elemental composition of TR and Pt/TR catalysts were investigated by field emission scanning electron microscopy (FESEM; Model JSM-6700F, JEOL) operated at 5 kV accelerating voltage and equipped with X-ray energy-dispersive spectrometry (EDS). The dispersion of Pt particles in TR was observed using transmission electron microscopy (TEM, Philips 400T) operated at 120 keV. The samples were crushed in an agate mortar and suspended in ethanol. After ultrasonic dispersion, a droplet was deposited on a copper grid supporting a perforated carbon film.

An X-ray powder diffractometer (XRD) (SIEMENS D5000) was used to identify the crystalline species of TR and Pt/TR catalysts. A Cu tube serving as the X-ray source was employed to estimate the active site phase at a working voltage and current of 40 kV and 30 mA, respectively. The powdered samples were pressed onto suitable holders and scanned in the 2θ range of 1 to 60° and 20° to 80°, respectively, at a scanning speed of 2° min⁻¹. The diffraction patterns were manually analyzed with the Joint Committee of Powder Diffraction Standard (JCPDS) cards.

The specific surface areas of the TR catalysts were calculated by graviometric methods with an ASAP 2010 vacuum microbalance. The Brunauer–Emmett–Teller (BET) surface area was calculated from the adsorption isotherm, and the ratio of pore volume and average pore diameter was obtained from the pore size-volume distribution. UV-Vis diffuse reflectance (DRS) spectrum of the TR catalyst was measured from 190 to 1100 nm, and the photoluminescence (PL) spectrum of Pt/Ti-SBA15(I) catalyst was measured using a fluorescence spectrometer (Hitachi F-7000) with an excitation (EX) wavelength of 250 nm. X-ray photoelectron spectroscopy (XPS) was employed to analyze the chemical states of Ti, O, C, and Pt in the TR catalysts via Kratos Axis Ultra DLD.

2.5. Experimental design of Taguchi method

In order to study the effect of catalytic factors in this experiment, the Taguchi method was employed to efficiently design a representative experiment. Three factors were chosen as experimental parameters: the ratio of Ti, space velocity, and time on stream. According to the orthogonal arrays (OA) of Taguchi method, a L9 OA table with three levels was considered, as described in Table 1. The above three factors were labelled as A, B, and C, respectively, and the three levels were designed with different parameters. For factor A, 1, 2, and 3 were TR(10), TR(30), and TR(50); for factor B, 1, 2, and 3 were 3932 h⁻¹, 15 727 h⁻¹, and 31 454 h⁻¹; and for factor C, 1, 2, and 3 were 20, 40, and 60 min, respectively.

Table 1. Three factors and their three levels for Taguchi method

<table>
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<tr>
<th>Level</th>
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<th>Space velocity (h⁻¹)</th>
<th>Time on stream (min)</th>
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<tr>
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3. Results and discussion

3.1. Characterization of TR catalysts

3.1.1. XRD analysis. In this study, Ti was incorporated into a silica framework by a one-step modified hydrothermal method; thus, the classic hexagonal structure of SBA-15 was expected to change after the incorporation of Ti. The small-angle XRD pattern was examined to analyze the structure of the TR catalysts, as shown in Fig. 1(a). The characteristic diffraction peaks at 2θ = 0.80°, 1.40°, and 1.60° corresponding to the (100), (110), and (200) planes of a hexagonal straight pore (P6mm) structure disappeared in three TR catalysts with increasing Ti/Si ratio. In theory, four steps are involved in classic SBA-15 preparation:

1. At pH < 1, P123–H⁺ formed from a neutral triblock copolymer surfactant, Pluronic 123, and H⁺ (HCl + H₂O → H₃O⁺ + Cl⁻).
2. P123–H⁺ reacted with Cl⁻ and H₂O–TEOS to form P123–H⁺Cl⁻Si⁴⁻.
3. At longer reaction times, a hexagonal columnar structure (P6mm) formed through self-assembly due to condensation from the static electricity effect of 2[P123–H⁺Cl⁻·×H₂O–Si⁴⁻]ₘ at a high temperature at longer reaction times.
4. The obtained sample was calcined to remove the copolymer template.

The three TR catalysts were synthesized with different ratios of TTIP (Ti)/TEOS (Si). The hydrolysis rate and condensation of Ti atoms affected the framework of the TR catalyst. In Fig. 1(a), when the Ti/Si ratio was 1/9, only (110) and (200) peaks were observed in TR(10). When the Ti/Si ratio was increased to 5/5, all characteristic peaks disappeared in TR(50). As the Ti doping weight increased during the synthesis process, the results indicated that the structures of TR(30) and TR(50) changed from hexagonal straight pores (P6mm) to other structures, whereas TR(10) may have retained the main structure of SBA-15 since the (110) and (200) peaks remained.

Fig. 1(b) shows the XRD patterns of TiO₂–P25 and the three TR catalysts measured from 20° to 80°. Comparison of the diffraction patterns shows that the crystalline species of Ti added in various proportions during the hydrothermal
ions (Si$^{4+}$ and Ti$^{4+}$) a crystal phase of TiO$_2$, anatase or brookite, transforms easily and the purpose of calcination was to remove the surfactant template calcination temperatures higher than 600°C. In this study, when Ti/Si was 1/9, the main crystalline species in the TR(10) catalyst was anatase ($2\theta = 25.2^\circ$, 36.9°, 48.0°, and 53.8°). When the Ti/Si ratio was increased to 3/7, both the anatase and rutile crystalline species appeared in TR(30). Compared to that of P25, the crystalline diffraction peak of TR(30) was significantly higher for the rutile phase ($2\theta = 27.5^\circ$, 36.0°, 41.2°, 44.0°, 54.3°, 56.6°, 64.0°, and 69.0°) than for the anatase phase. However, the intensity of the anatase phase increased in TR(50) compared with that in TR(30) when the Ti/Si ratio increased to 5/5.

Acosta-Silva et al. reported that Si improves thermal stability and inhibits the transformation of anatase to rutile since the surrounding Si stabilizes the Ti–O bonds at the interface with SiO$_2$. In addition, Resende et al. reported that a homogeneous silica layer was formed in contact with TiO$_2$, which inhibited the appearance of the rutile phase because of insufficient atomic diffusion. The atomic mixing of Ti and Si retards phase separation and crystallization. Thus, at Ti/Si = 5/5 (TR(50)), the transformation of large amounts of anatase phase to the rutile phase was successfully inhibited.

3.1.2 Morphological studies. FESEM was employed to observe the surface morphologies and particle sizes of TR catalysts with varying Ti–Si proportions. Fig. 2 displays the FESEM images of SBA-15 and TR catalysts with varying Ti/Si ratios at 3000× to 100 000× magnification. Fig. 2(a) shows the classic pillar shape of SBA-15, with a particle size of approximately 1–3 μm, as shown in the inset of Fig. 2(a). SBA-15 has a two-dimensional channel structure with uniform pore channels, as shown in the 50 000× image in Fig. 2(a). The pore channels could be clearly observed on the surface of SBA-15. Moreover, a small region of the samples were analyzed by EDS, as shown in Fig. (3). The EDS patterns of the three samples show that the elements in the particles are the same, including Ti, O, and Si.

In contrast to traditional SBA-15, deformation occurred after the titanium precursor, TTIP, was premixed with the silica precursor, TEOS, during the hydrolysis/condensation process; the material no longer possessed the original structure of SBA-15. The FESEM images confirm the results of small-angle XRD analysis (Fig. 1(a)). As shown in Fig. 2(b), the TR(10) catalyst is irregularly shaped but has numerous holes distributed along its surface, similar to SBA-15. The structure of surface channels can be seen to have changed due to the embedding of Ti into the Si–O framework in TR(10). Moreover, the TR(10), with a diameter of approximately 10 μm, is larger than SBA-15, as shown in the inset of Fig. 2(b). The ionic radii of Ti and Si were 0.064 nm and 0.042 nm, respectively, which suggested that the bonding length of Ti–O–Si was longer than that of Si–O–Si. The Ti incorporated into the silica framework contributed to the large porous structure, as shown in the inset of Fig. 2(b).

With increasing Ti content, the structures of TR(30) and TR(50) transformed into a three-dimensional porous structure. Fig. 2(c) shows that the shape of TR(30) is a sphere formed by many long cylinders, resembling a cauliflower. The diameter of TR(30) is approximately 0.8 μm. Similar to TR(30), TR(50) with Ti/Si = 5/5 is composed of large amounts of long cylinders and is shaped like an individual sphere. Although both TR(30) and TR(50) are spheres comprising innumerable long cylinders, TR(50) is slightly smaller at 0.5 μm, as shown in the inset of Fig. 2(d). Additionally, the cylinders in TR(50) are shorter than those in TR(30). The results suggest that the structure changed from two-dimensional to three-dimensional as the Ti/Si ratio increased to 3/7 or 5/5. The high concentration of Ti formed a different framework from that of the incorporated silica.
Thereafter, the decreased number of Si sites around the P123 surfactant and the shorter hydrolysis time of Ti resulted in a three-dimensional porous structure after the condensation process.\(^{27}\)

3.1.3 Raman spectra. XRD analysis showed that the anatase and rutile phases coexisted in the TR catalysts. To confirm the XRD results, the Raman spectra of the TR catalysts were obtained and are presented in Fig. 4. In Fig. 4(a), the characteristic Raman-active modes of anatase are observed at 143, 196, 395, 513, and 637 cm\(^{-1}\) in TR(10). Compared with the standard spectrum (bands at 144, 197, 399, 519, and 639 cm\(^{-1}\)), the Raman spectra of TR(10) show a blueshift, which might be due to the lattice distortion of the Si–O–Ti bond and the ratio of anatase and rutile phases.\(^{44}\) Choi \textit{et al.}\(^{42}\) indicated that particle size is inversely proportional to phonon momentum distribution. Thus, an increase in the phonon momentum distribution would result in a smaller particle size of the TR catalyst (TR(30) or TR(50)); consequently, the extended scattered phonon momentum would shift the Raman spectra.

In contrast, the peaks at 237, 440, and 609 cm\(^{-1}\) are attributed to the rutile phase in TR(30) and TR(50). Furthermore, in TR(30) and TR(50), the anatase peaks are almost overlapped by the rutile peaks (237, 440, and 609 cm\(^{-1}\)). In addition, the peak at 143 cm\(^{-1}\), which is characteristic of the anatase phase, shows a stronger intensity for TR(50) than for TR(30); this reveals that TR(50) possesses a greater amount of anatase phase than TR(30). The Raman results thus confirm the XRD analysis results.

Since the anatase and rutile phases coexisted in the TR catalysts, the ratio of the anatase/rutile phase was expected to affect the photocatalytic activity.\(^{45,46}\) Thus, the content fractions of anatase and rutile in the TR catalysts were calculated by the Spurr and Myers equation,\(^{47}\) as shown in Table 2. The fraction of rutile phase is in the order of TR(30) (81.5%) > TR(50) (77.9%) > TR(10) (56.9%).

The rutile phase content varied with the Ti/Si ratios of the TR catalysts. According to earlier studies,\(^{45,29}\) Si enhances the

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![Fig. 2 FESEM images of: (a) SBA15; (b) TR(10); (c) TR(30); (d) TR(50).](image1)

![Fig. 3 EDS analysis of: (a) TR(10); (b) TR(30); (c) TR(50).](image2)

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thermal stability and crystallinity of TiO₂ during calcination, as shown in Table 2. By comparing with the intensity of standard XRD patterns of TiO₂, crystallinities of anatase and rutile phase were calculated. The higher degree of crystallinity would reduce the possibility of electron and hole recombination at a defective site and enhance the photocatalytic activity.

As shown in Fig. 2(b), embedded Ti was observed on the surface of the two-dimensional channel structure over TR(10). At a low Ti content, Ti is incorporated into the silica framework, and Si inhibits the transformation of the anatase phase to the rutile phase. After calcination, the anatase phase content of TR(10) was 43.1%.

However, the suppression of the TiO₂ phase transformation in TR(10) was not the same as in TR(30) and TR(50). The transformation inhibition did not decrease with large amounts of added TiO₂. In general, the low inhibition efficiency was mainly attributed to the reduced assistance from the silica framework. The three-dimensional porous structures of TR(30) and TR(50) also contributed to the different inhibition effects during calcination. The long cylinders of TR(30) led to a lower anatase content (18.5%) than the short cylinders of TR(50) (22.1%). The result suggests that the longer Si–O–Ti bonding length in a three-dimensional porous structure is one of the factors affecting the TiO₂ crystal phase during calcination, leading to the lower inhibition of TiO₂ phase transformation. Furthermore, the compositions of the TiO₂ crystalline phase in the three TR catalysts were not the same as those in commercial TiO₂–P25 (Table 2). The fraction of anatase phase in P25 (86.1%) is higher than that in the three TR catalysts. Moreover, the photocatalytic activity of the TR catalyst is different from that of traditional TiO₂.

Fig. 4(b) shows the Raman spectra of TR catalysts measured from 500 to 2000 cm⁻¹. The band at 797 cm⁻¹ corresponds to the symmetric stretching mode of Si–O–Si in the three TR catalysts. With increasing Ti/Si ratio, the intensity of the Si–O–Si peak decreased in TR(30) and TR(50). The band located at 1085 cm⁻¹ was assigned to the symmetric stretching vibration of the Ti–O–Si bond in TR(10). These results are in agreement with the results of FESEM and XRD. At a low Ti/Si ratio, the Ti ions were incorporated into the silica framework to form a two-dimensional channel structure (Fig. 2(b)). Thus, the thermal stability of the anatase phase improved and contributed to the inhibition of the phase transformation and sintering of TR(10).

In Fig. 4(b), the peak at 1602 cm⁻¹ corresponds to the IG mode of carbon, whose sources are the precursors TEOS and TTIP and the surfactant P123. Graphite crystal was clearly observed in TR(30) and TR(50), whereas no defect carbon (ID mode of carbon) appeared at 1350 cm⁻¹. Thus, the shift in the spectra was not affected by carbon. Without defect carbon, the excited electron would not scatter during analysis. This result indicates that C–C bonds existed in the structures of TR(30) and TR(50), while the decreased Si sites around the P123 surfactant and the shorter hydrolysis time of Ti contributed to the structure formed after the condensation process. These factors contributed to the transformation of a two-dimensional channel column to a three-dimensional porous spherical structure.

### Table 2 The crystal characteristics of TR catalysts according to the XRD patterns

<table>
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<th>Catalyst</th>
<th>Crystal size of anatase (nm)</th>
<th>Crystal size of rutile (nm)</th>
<th>Average crystal size (nm)</th>
<th>Component of anatase (%)</th>
<th>Component of rutile (%)</th>
<th>Crystallinity of anatase (%)</th>
<th>Crystallinity of rutile (%)</th>
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3.1.4 XPS analysis. The chemical states of the elements Ti, O, and C were analyzed by XPS, as shown in Fig. 5–7. The Ti2p spectra are shown in Fig. 5, where the two characteristic peaks at 456.5 eV and 462.4 eV in TR(10) correspond to Ti2p_{3/2} and Ti2p_{1/2} of the octahedrally coordinated anatase phase. The energy width of the two peaks was approximately 5.9 eV, which revealed the chemical state of Ti to be Ti^{4+}. Compared with the standard spectrum, the measured spectrum is slightly blue-shifted. The Ti2p_{3/2} or Ti2p_{1/2} peak shifted to a lower binding energy by approximately 2 eV compared to the anatase phase of Ti^{4+}. Gao et al. indicated that the blue-shift might be associated with the Ti–O–Si bonds, resulting in a more positive charge on Ti, a less negative charge on O, and more negative charge on Si. With increasing Ti content, the intensity of the peak increased as the Ti/Si ratio improved. In addition, the Ti2p_{3/2} and Ti2p_{1/2} peaks blueshifted to 456.3 eV and 462.0 eV due to the quantum size effect that resulted from the decrease in TiO_2 particle size. As shown in Fig. 2, TR(30) and TR(50) were smaller than TR(10). The average crystal size was in the order TR(50) (13.4 nm) < TR(30) (14.3 nm) < TR(10) (39.2 nm), as shown in Table 2. Thus, a blueshift was observed in the XPS spectra of Ti2p.

Similar to the Ti2p spectra, the O1s spectra shifted downward to a binding energy of −2.0 eV for the three catalysts, as shown in Fig. 6. The band at 527.9 eV corresponded to the lattice oxygen of the Ti–O–Ti bond. As the Ti/Si ratio increased, the intensity of the Ti–O–Ti bond became stronger for TR(50). The O1s peak at 530.1 eV was deconvoluted into three peaks. The resolved peaks of TR(10) and TR(50) were similar, and the binding energies of the three peaks were 530.0, 530.6, and 531.3 eV, respectively. For TR(30), the O1s peak decomposed into three peaks with binding energies of 530.5, 531.0, and 531.6 eV.

The peak at 530.0 eV was related to the absorbed oxygen on the surface. Mazur et al. reported that the absorbed oxygen originated from H_2O and hydroxyl radicals. No decomposed peak appeared at 530.0 eV in TR(30); however, there was a peak at 531.6 eV related to the Si–OH bond or absorbed H_2O. The oxygen bond content arising from the H_2O or hydroxyl groups was calculated in relation to the total oxygen bond content, and the order was TR(10) (23.9%) > TR(30) (15.6%) > TR(50) (13.0%). This result was in agreement with earlier studies, which suggested that Si facilitated the bonding of OH groups onto the TR catalyst. The peaks at 530.5/530.6 eV and 531.0/531.3 eV...
corresponded to the Ti–O–Si and Si–O–Si bonds, respectively. As mentioned above, the peak for defect oxygen arose from the Ti atoms incorporated into the silica framework of TR(10). In TR(30) and TR(50), the strong interaction of the Ti–O–Si bonds between TiO₂ and SiO₂ resulted in the formation of a three-dimensional porous structure.²¹

The C1s spectra of the three catalysts are shown in Fig. 7. The C1s peak was deconvoluted into three peaks related to the C–Ti, C–Si, and C–C bonds at 282.1, 283.1, and 284.3 eV, respectively.²⁴ The C1s peak of TR(10) was weaker than that of the other two catalysts, possibly because the C–C bond, which appeared in samples with a Ti/Si ratio greater than 3/7, contributed to the stronger C1s peak. For TR(10), the carbon bond content was 71.8% for C–Ti and 28.2% for C–Si. With increasing Ti loading weight, the hydrolysis and condensation of Ti and Si surrounded by the P123 surfactant contributed to the production of C–C bonds after calcination. The C–C bond content was calculated in relation to the total carbon bond content, and the order was TR(50) (19.1%) > TR(30) (17.6%). The proportions of C–Ti and C–Si in TR(30) and TR(50) were lower than those in TR(10). The C–Si bond content was calculated in relation to the total carbon bonds, and the order was TR(10) (28.2%) > TR(30) (20.1%) > TR(50) (19.4%). Although the amount of carbon in the catalysts was very small, the analysis of the C–Si bonds revealed that the framework of the catalysts could be transformed by reducing the C–Si bond content and increasing the C–C bond content. This transformation might be the other factor responsible for the formation of a three-dimensional porous structure. Ti–O–Si–C bonds would appear in the three-dimensional structured catalyst. The results revealed that the observed G-band of the Raman spectra, which appeared only in TR(30) and TR(50), arose from the C–C bond, as shown in Fig. 4(b).

3.1.5 S_{BET} and structure. Further analysis of the specific surface area (as shown in Table 3) showed that the S_{BET} and pore volume of the three TR catalysts with varying Ti/Si ratios synthesized in this study were greater than those of the commercial photocatalyst P25. The S_{BET} followed the order TR(10) (787.1 m² g⁻¹) > TR(30) (463.5 m² g⁻¹) > TR(50) (419.0 m² g⁻¹), and the pore volume varied in the order TR(10) (0.95 cm³ g⁻¹) > TR(30) (0.88 cm³ g⁻¹) > TR(50) (0.54 cm³ g⁻¹). Compared with those of the classic SBA-15, the physical characteristics of SBA-15 included a higher specific surface area (an increase from 630 to 1040 m² g⁻¹) and a greater pore volume (an increase from 0.56 to 1.26 cm³ g⁻¹).²² TR(10) showed similar textural characteristics and morphology (Fig. 2(b)) to SBA-15 due to the Ti incorporated in the silica framework. The particles of TR(10) were larger than those of SBA-15 because the Si–O–Ti bonds were longer than the Si–O–Si bonds. In contrast, the specific surface area and pore volume decreased as the metallic Ti content increased. The morphologies of TR(30) and TR(50) transformed to a three-dimensional porous structure; however, they still maintained good S_{BET} and large pore volumes because of the strength of the Si–O–Ti bonds. TiO₂ was rigidly held in place by its interactions with SiO₂, and sintering during calcination was inhibited by the improved thermal properties owing to Si.²³,²⁸,²⁹

The average pore diameter of the three samples decreased in the order TR(30) (7.61 nm) > TR(50) (5.18 nm) > TR(10) (4.82 nm). The FESEM images (Fig. 2) revealed that the pores of TR(30) and TR(50) mainly arose from the gaps between cylinders, whereas hexagonal channels formed the pores of TR(10). The proportion of mesoporous volume followed the order TR(30) (74.9%) > TR(50) (67.4%) > TR(10) (64.2%). The proportions of mesoporous structures for all TR catalysts were above 64%. This result suggested that the gaseous molecules could react efficiently with active sites over mesoporous TR catalysts during photocatalytic oxidation since the molecular size of HCHO is 0.27 nm.²⁵

3.2. Formaldehyde removal efficiencies of TR catalysts

3.2.1 HCHO removal of TR(10) and P25 under UV light. Formaldehyde was selected as the reaction gas, and the formaldehyde removal efficiency over TR catalysts at room temperature was examined. Fig. 8(a) shows the formaldehyde removal efficiency of commercial TiO₂–P25 and TR(10) under UV irradiation. The reaction conditions were as follows: 40 ppm HCHO and 20% O₂ (balance He + N₂) with a total space velocity of 23 591 h⁻¹ under 2 × 10 W UVA irradiation. The results of the experiment indicated that TR(10) possessed superior formaldehyde photocatalytic removal efficiency to that of P25. Within 20 min, TR(10) had removed 99% of the formaldehyde, whereas P25 had removed only 20% to 30%.

The results demonstrated that the shift in the light wavelength from the UVB to the UVA region contributed to the low photocatalytic activity of P25 with a retention time of 1.52 × 10⁻³ s. In general, the maximum absorption wavelength of light for commercial TiO₂–P25 was 320 nm; thus, little energy was obtained from UVA (365 nm) irradiation. Therefore, few photons were excited for the separation of electron–hole pairs, which contributed to the low photocatalytic oxidation efficiency of HCHO.²⁵ In contrast, the proportion of the rutile phase was above 56.9% in the three TR catalysts, whereas that in P25 was only 13.9%, as shown in Table 1. The results under near-visible light irradiation suggested that more superoxide radical anions would be produced from the photoexcitation process e⁻ + O₂ads.
order of the photocatalytic formaldehyde removal efficiency for the catalysts was as follows: TR(10) > TR(30) > TR(50) > P25. The higher retention time, $9.16 \times 10^{-1}$ s, improved the removal efficiency of P25 from 38% at 10 min to 59% at 10 min under UVA irradiation (Fig. 8(a)). However, P25 still exhibited a specific downward trend to a final value of 0 to 5%, as seen initially and at 60 min for HCHO removal under visible light irradiation. Photoexcitation on the surface of P25 could not be driven effectively under visible light irradiation.

The removal efficiencies of the TR catalysts varied after a period of reaction time in the HCHO stream. After 45 min of time on stream, TR(10) attained a removal efficiency of 87–96%.

TR(30) attained a removal efficiency of approximately 82–97% after 35 min. TR(50) attained a removal efficiency of 56–89% after 20 min. Multiple factors were assumed to contribute to the differences in the removal efficiencies. First, the morphologies and textural structures of the three samples were different, as shown in Fig. 2 and Table 3. The $S_{BET}$, pore volume, and average pore diameter directly affect the photocatalytic efficiency. All samples were mesoporous structures, and the HCHO molecule would be most easily adsorbed on a catalyst with a high $S_{BET}$ and large pore volume, such as TR(10), which is why TR(10) showed the most stable HCHO removal for the longest time, as shown in Fig. 8(b).

The literature recommends adding an appropriate amount of SiO$_2$ to improve the photocatalytic properties of TiO$_2$ because increasing the SiO$_2$ content may increase the surface acidity of TiO$_2$. SiO$_2$-modified TiO$_2$ exhibited the greatest surface acidity improvement, not only increasing the adsorption of organic matter on the TiO$_2$ surface but also forming strong OH groups in the hole capture sites on the surface, thereby reducing the probability of electron–hole pairing. This process generated OH groups with strong oxidation activity and improved the photocatalytic properties. According to the XPS analysis of O1s (Fig. 6), TR(10) showed 23.9% oxygen bonds arising from H$_2$O$_{abs}$ or hydroxyl groups, TR(30) showed 15.6%, and TR(50) showed 13.0%. The amount of absorbed oxygen corresponded to the removal efficiency of HCHO. Finally, comparing Fig. 8(b) and (a) could suggest that the photocatalytic activity of the TR catalyst might be affected by the different absorption of light energy under visible light irradiation since the proportion of the rutile phase was not the same in all TR catalysts, as shown in Table 2. Overall, the TR catalysts showed better photocatalytic efficiency than commercial P25 regardless of whether the light source was UVA or visible light, showing that TR catalysts have potential for use as photocatalytic materials.

### 3.2.3 UV-Vis DRS analysis of TR catalysts.

To study the effects of the different proportions of rutile phase in the three TR catalysts on the photocatalytic reaction, UV-Vis DRS analysis of the samples was carried out. Fig. 9(a) shows the absorption spectra of three samples: the absorption edge was 418 nm, 423 nm, and 432 nm for TR(10), TR(30), and TR(50), respectively. Obviously, a redshift occurred as the Ti/Si ratio increased. XRD analysis (Table 2) showed that the redshift was due to the small crystal size of the rutile phase. In general, the band gap energy of the anatase phase is 3.2 eV. Commercial P25 is
the anatase/rutile phase ratio in the photocatalyst. Thus, the optical absorption edge would vary with the strong optical absorption occurs at the visible light wave-

In contrast, the band gap of the rutile phase is 3.0 eV, and the calculated by the formula of 432 nm, longer than that of the others. This result indicated that TR(50) would separate electron–hole pairs under visible light irradiation more easily than the others.

Moreover, the UV-Vis DRS analysis results can be employed to calculate the band gap energy of the TR catalysts from the plot of [ahv]² vs. the photon energy, as shown in Fig. 9(b). The band gap energy of the three TR catalysts was consistently below 3.2 eV (pure anatase phase) or 3.0 eV (pure rutile phase). The band gap energy of the TR catalyst followed the order TR(50) (2.83 eV) < TR(30) (2.87 eV) < TR(10) (2.89 eV). The results revealed that a higher rutile phase content contributed to a lower band gap. The band edge positions of the TR catalyst can be calculated by the empirical equation $E_{CB} = X - E_v - 0.5E_g$ where $E_{CB}$ is the conduction band edge potential; $X$ is the absolute electronegativity of the semiconductor (X value for TiO₂ is 5.81 eV); $E_v$ is the energy of free electrons on the hydrogen scale (approximately 4.5 eV); and $E_g$ is the band gap of the TR catalyst. The predicted band positions at the point of zero charge of TR(10), TR(30), and TR(50) were $-0.135$ eV–$2.755$ eV, $-0.125$ eV–$2.745$ eV, and $-0.105$ eV–$2.725$ eV, respectively.

However, earlier studies indicated that the recombina-
tion of electron–hole pairs was faster for the rutile phase than for the anatase phase since photoexcitation would occur easily over the rutile phase with low band gap energy. This observation was corroborated by comparing Fig. 8(b) with Fig. 9(b): TR(50) showed the lowest band gap energy, but its HCHO removal efficiency decreased with reaction time due to the rapid recombination rate of the electron–hole pairs. Therefore, the drawback of the short recombination time of electron–hole pairs for the rutile phase needs to be resolved to prolong the reaction time of photocatalytic oxidation.

3.3. Taguchi method for optimizing the catalytic factor over TR catalysts

To tune the operational parameters for HCHO removal over TR catalysts, three catalytic factors were studied in nine tests designed by the orthogonal array L9 of the Taguchi method, as shown in Tables 1 and 4. The experimental results were analyzed by using the S/N ratio and one-way analysis of variance (ANOVA).

![Absorption spectra of TR(10), TR(30), and TR(50) catalysts; (b) band gap calculation of (ahv)² vs. photon energy for TR catalysts.](image)

**Fig. 9**

<table>
<thead>
<tr>
<th>Run</th>
<th>Ratio of Ti</th>
<th>Space velocity</th>
<th>Time on stream</th>
<th>Removal (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Level 1</td>
<td>Level 1</td>
<td>Level 1</td>
<td>99.1</td>
</tr>
<tr>
<td>2</td>
<td>Level 1</td>
<td>Level 2</td>
<td>Level 2</td>
<td>99.0</td>
</tr>
<tr>
<td>3</td>
<td>Level 1</td>
<td>Level 3</td>
<td>Level 3</td>
<td>43.8</td>
</tr>
<tr>
<td>4</td>
<td>Level 2</td>
<td>Level 1</td>
<td>Level 3</td>
<td>82.8</td>
</tr>
<tr>
<td>5</td>
<td>Level 2</td>
<td>Level 2</td>
<td>Level 1</td>
<td>79.0</td>
</tr>
<tr>
<td>6</td>
<td>Level 2</td>
<td>Level 1</td>
<td>Level 2</td>
<td>49.1</td>
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<tr>
<td>8</td>
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<td>Level 2</td>
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</tr>
<tr>
<td>9</td>
<td>Level 3</td>
<td>Level 3</td>
<td>Level 1</td>
<td>25.2</td>
</tr>
</tbody>
</table>
The S/N ratio is the ratio of signal to noise and was calculated according to the degradation efficiencies for the corresponding factors at each level tested. S/N serves as an indicator of the experimental quality, has the additive effect of an experimental factor and reduces interactions between factors. The larger the value of the S/N is, the smaller is the loss of experimental quality or the smaller is the variation in the quality. The corresponding chart of experimental results for the individual effects of various parameters at each level is shown in Fig. 10. For factor A (proportion of Ti), the S/N values followed the order TR(10) > TR(30) > TR(50). For factor B (space velocity), the S/N values were 3932 h⁻¹ > 15 727 h⁻¹ > 31 454 h⁻¹. For factor C (time on stream), the S/N values were 40 min > 20 min > 60 min. The more extreme the difference in the S/N values was, the more significant was the effect. The level effects of the three factors were in the order B (6.5) > A (4.6) > C (1.7). The ranks of the three factors varied in the order B > A > C. Factor B, space velocity, had a significant effect on the photocatalytic activity for HCHO removal, which indicated that mass transfer was the principal factor for TR catalysts, since the mesoporous structures of TR catalysts were variable. Factor A, the ratio of Ti, had a minor effect, which demonstrated that a low Ti loading wt% would result in better catalytic activity than a high Ti loading. Factor C, time on stream, had the least impact on the photocatalytic activity. The lack of an obvious decay effect on the TR catalysts during the reaction was suggested to be because appreciable coke deposition did not occur on the TR catalysts due to the amount of surface acidity on TR catalysts (discussed in Section 3.2.2). In this study, the optimum test parameter for the ratio of Ti was level 1 (TR(10)); that for the space velocity was level 1 (3932 h⁻¹); and that for the time on stream was level 2 (40 min).

Then, ANOVA was used to analyze the influence of various test parameters on the HCHO removal efficiency in nine runs. Table 5 shows the ANOVA results for the photocatalytic removal of HCHO with a 95% confidence interval and two degrees of freedom. The degree of residual value in the ANOVA analysis was 6. The contributions of the three factors to HCHO removal efficiency over TR catalysts were 32%, 62%, and 5%, respectively. Space velocity had a significant influence on HCHO removal. The contributions of factors could be ranked as follows: space velocity > ratio of Ti > time on stream. The time on stream had very little effect on the TR catalyst and could be neglected for HCHO removal. Furthermore, the error contribution to the total variance was 1%, which is lower than 15%, and this result indicated that no significant parameters were ignored in the experimental design. The three factors chosen in this study were reasonable and credible. The effects of the studied factors on the response function were evaluated by the F value of the three factors, which varied in the order B (7.35) > A (2.13) > C (0.24). Only the F value of factor B was found to be larger than the critical value of Fₐ, where Fₐcritical (0.05, 2, 6) was 5.143. This finding rejected the null hypothesis that the standard deviations were equal and demonstrated differences in the variances in theory. The order of the effects of the three factors on HCHO removal over TR catalysts was factor B > factor A > factor C, which was the same as that obtained by investigation of the S/N ratio. Finally, the P value was studied. The F value of 7.35 and the P value of 0.02 for factor B (space velocity) demonstrated the real and high significance of factor B for HCHO removal over TR catalysts. In contrast, a P value > 0.05 meant that HCHO removal efficiency was not significantly affected by factor A and factor C, even though factor A showed a 32% contribution in the catalytic test.

In the following study, the better-performing catalysts TR(10) and TR(30) were chosen for Pt/TR catalyst preparation to evaluate the assisting effect of metal-Pt particles on HCHO removal over TR catalysts. Before testing the activity of Pt/TR catalysts, the optimum space velocity should be considered to assess whether Pt/ST(10) or Pt/ST(30) could be a potential catalyst for HCHO removal.

### 3.4. Optimum space velocity for HCHO removal over TR catalysts

In this study, TR catalysts showed good photocatalytic activity for HCHO removal at a reaction space velocity of 3932 h⁻¹ under visible light irradiation. To confirm the existence of a mass transfer gradient effect on the chemical reactions in the continuous reactor used in this study, we selected the better catalyst—TR(10)—to measure the actual effects of the catalytic reaction weight and gas flow velocity on the photocatalytic reactions. The primary factors controlling the heterogeneous photocatalytic reaction rates were the dispersion and surface reaction. When the size of the

![Fig. 10](image.png) The effect of each factor (ratio of Ti, space velocity, and reaction time) on the values of S/N ratio.

<table>
<thead>
<tr>
<th>Factor</th>
<th>DF</th>
<th>SS</th>
<th>MS</th>
<th>Contribute</th>
<th>F value</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>2</td>
<td>1715.76</td>
<td>857.88</td>
<td>0.32</td>
<td>2.13</td>
<td>0.20</td>
</tr>
<tr>
<td>B</td>
<td>2</td>
<td>3310.38</td>
<td>1655.19</td>
<td>0.62</td>
<td>7.35</td>
<td>0.02</td>
</tr>
<tr>
<td>C</td>
<td>2</td>
<td>266.08</td>
<td>133.04</td>
<td>0.05</td>
<td>0.24</td>
<td>0.80</td>
</tr>
<tr>
<td>Residual</td>
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<td>44.84</td>
<td>7.47</td>
<td>0.01</td>
<td></td>
<td></td>
</tr>
<tr>
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<td>5337.06</td>
<td></td>
<td></td>
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</tr>
</tbody>
</table>
photoreactor and the light intensity were fixed, variations in the gas flow affected the photocatalytic reaction. A varying gas flow affects not only the residence time but also the reaction gas flow. Because low gas flows are slow, the flow in the reactor was similar to a laminar flow, hindering interaction between the reaction gas and the photocatalyst and limiting the ability of the reaction gas to diffuse onto the surface of the photocatalyst; thus, the reaction was deemed to have been caused by reaction gas diffusion. Conversely, as the gas flow was gradually increased, the flow created turbulence equivalent to a mixing process. Therefore, the probability of interaction between the reaction gas and the photocatalyst increased, accelerating diffusion. For low flows, the reaction rates are affected by mass transmission. Increasing the flow increases the reaction rate. For high flows, the reaction rates are not limited by mass transmission but are controlled by surface reactions.

Fig. 11(a) shows the HCHO removal efficiency of TR(10) under visible light irradiation using various spatial velocities. The reaction conditions were as follows: 40 ppm HCHO and 20% O₂ (balance He + N₂) under 2 × 10 W fluorescent irradiation. The results of the experiment indicated that the photocatalytic reaction efficiency decreased with increasing the spatial velocity during the reaction. Spatial velocities of 3932 h⁻¹ and 15727 h⁻¹ provided the best reaction removal efficiency (greater than 99%). The HCHO removal efficiency began to decline when the spatial velocity exceeded 31 454 h⁻¹. The photocatalytic activity was clearly affected by the contact time between the HCHO and TR catalysts when the residence time was less than 1.14 × 10⁻² s. A short residence time contributed to a low mass transfer efficiency and lowered the removal of HCHO. The minimum efficiency was obtained at 78 635 h⁻¹ (residence time of ~0.46 × 10⁻¹ s); however, the minimum efficiency was still 30%. The efficiency at a high spatial velocity still exceeded that of the commercial photocatalyst P25 (Fig. 8(b)). Fig. 11(b) shows the photocatalytic kinetic curves for the TR catalysts at different space velocities. A linear correlation was observed at high space velocities of 31 454 h⁻¹ and 78 635 h⁻¹; the values of the pseudo-first-order reaction rate constant for these two space velocities were 0.0127 m⁻¹ s⁻¹ and 0.0117 m⁻¹ s⁻¹, respectively. In contrast, no linear correlation was observed at low space velocities. To render subsequent tests comparable, this study set the spatial velocity for the following HCHO photocatalytic reaction tests at 31 454 h⁻¹.

3.5. Formaldehyde removal efficiency of Pt/TR catalysts

3.5.1 TEM images of Pt/TR catalysts. FESEM analysis can be used to explain differences in the external appearance of the TR catalysts with varying Ti proportions synthesized using the modified hydrothermal method. However, the size and dispersion of Pt particles in the TR catalysts were difficult to observe by FESEM. Therefore, TEM analysis was conducted to analyze the internal structure and gain a better understanding of the distribution of metallic Pt particles on the TR photocatalysts.

Fig. 12(a) shows TEM images of Pt/TR(10) at 200 000× magnification, while the inset is at 30 000× magnification. A comparison of the inset image of Fig. 12(a) with Fig. 2(b) showed that the morphology of TR(10) did not change after Pt doping. The hexagonal pore structure was retained, and the size of the Pt particles ranged from 4 to 30 nm (as indicated by red arrows). Moreover, X-rays were used to perform EDS, as shown in Fig. 12, which shows a good dispersion of Pt particles (light spots in Fig. 13(a)) and peak signals for Pt, Ti, O, and Si; therefore, the Pt particles on the surface of Pt/TR(10) were observable, and EDS analysis verified that Pt particles were indeed doped on Pt/TR(10). TEM images and EDS analysis showed that some Pt particles were dispersed on the surface of the Ti–O–Si structure, and some Pt particles were embedded in the two-dimensional channels of TR(10). The microwave-assisted ethylene glycol reduction method effectively dispersed Pt particles onto the catalysts.

Fig. 12(b) shows TEM images of Pt/TR(30) at 100 000× magnification, and the inset is at 400 000× magnification. Similar to TR(30) (Fig. 2(c)), the shape of Pt/TR(30) was similar to that of a cauliflower. However, the internal texture was not observed clearly because the porous three-dimensional...
structure led to low electron beam penetration during TEM analysis. Also, EDS analysis shows that Pt, Ti, O, and Si were appeared on TR(30), as shown in Fig. 13(b). At high magnification (inset of Fig. 12(b)), the Pt particles were dispersed into the splits between cylinders, and the sizes of the Pt particles were approximately 5–10 nm. The results showed that when 1 wt% Pt was attached to TR photocatalysts with varying proportions of Ti/Si, the external shape of the photocatalysts remained relatively unchanged. The internal structure of the catalysts also remained unchanged after calcination, which indicated that TR catalysts possess good thermal stability.

3.5.2 XRD and XPS analysis of Pt/TR catalysts. The XRD patterns of Pt/TR catalysts are shown in Fig. 14. Compared to observations made based on Fig. 1(b), no apparent pattern change in the TR catalysts was noted after Pt doping. The anatase and rutile phases still coexisted in the Pt/TR catalysts. Moreover, the presence of the crystal phase of zerovalent Pt was confirmed by diffraction peaks at $2\theta = 39.8^\circ$ (111), 46.3$^\circ$ (200), and 67.5$^\circ$ (220). The Pt diffraction peaks of Pt/TR(10) were stronger than those of Pt/TR(30), and that the Pt particles were suggested to be aggregated into larger particles in Pt/TR(30) since the Pt doping weight was the same in the two samples. The Scherrer equation was applied to calculate the mean Pt particle size from the strongest peak Pt (111) at $2\theta = 39.8^\circ$. The calculated sizes of the Pt particles were 12.6 nm and 73.6 nm in Pt/TR(10) and Pt/TR(30), respectively. As shown in Fig. 12, the Pt size in Pt/TR(10) was calculated to be between 4 and 30 nm according to the TEM images, which implied that nanoscale Pt particles were effectively supported on Pt/TR(10). In contrast, the calculated Pt size in Pt/TR(30) was larger than that observed in Fig. 12(b). The inconsistent size calculated for the Pt particles was attributed to the overlap of the rutile phase (111) and Pt diffraction peaks. Thus, the FWHM of Pt/TR(30) was 0.152, smaller than that of Pt/TR(10) (0.679); the small value of FWHM contributed to the large crystal size of Pt, as calculated by the Scherrer equation.

Fig. 12 TEM images of Pt/TR catalysts: (a) Pt/TR(10); (b) Pt/TR(30).

Fig. 13 EDS analysis of: (a) Pt/TR(10); (b) Pt/TR(30).

The spectra for Pt4f of the Pt/TR catalysts are shown in Fig. 15. Similar to the Ti2p spectra, the Pt4f spectra showed a downward shift to a value of −2.0 eV in the binding energy for Pt/TR(10) and Pt/TR(30). For Pt/TR(10), the bands at 69.0 eV and 72.3 eV were attributed to Pt4f7/2 and Pt4f5/2. The gap between the two bands was 3.3 eV, and the peak area ratio was 1.26, which was assigned to zerovalent Pt (Pt0) formed in TR(10). A blueshift of the two bands of Pt/TR(30) was observed, possibly due to the quantum effect resulting from the smaller size of Pt particles, as seen in Fig. 12(b). This result revealed that the size of Pt particles calculated using the Scherrer equation would be affected by the diffraction of neighboring crystals. The gap and peak area ratio of the two bands were 3.3 eV and 1.30, respectively. Thus, Pt0 was also the main phase in TR(30). This result was confirmed by XRD.

Fig. 14 XRD patterns of Pt/TR(10) and Pt/TR(30) catalysts.
3.5.3 HCHO removal by Pt/TR catalysts under visible light.

The experiment described above showed that the TR(10) and TR(30) catalysts possessed better photocatalytic capacity than TR(50). Therefore, this section examines the HCHO removal efficiency of these two photocatalysts after doping with Pt particles. Fig. 16 depicts the results of HCHO removal over Pt/TR catalysts under visible light irradiation. The reaction conditions were as follows: 40 ppm HCHO and 20% O₂ (balance He + N₂) with a total space velocity of 31 454 h⁻¹/C₀ under 2 × 10 W fluorescent irradiation.

Comparison of Fig. 8(b) and 11 showed that Pt doping on the TR catalysts effectively increased the HCHO photocatalytic efficiency at a residence time of 1.14 × 10⁻¹ s, increasing the removal efficiency of Pt/TR(10) and Pt/TR(30) to 90.4% and 80.0%, respectively, at a reaction time of 60 min. Furthermore, the physical and chemical advantages of TR catalysts led to a synergistic effect on HCHO removal. A possible reason for this increase in the photocatalytic efficiency was that the use of the microwave-assisted EG reduction method to synthesize Pt/TR catalysts can enhance Pt particle dispersion and control the particle size, as shown in Fig. 12. These uniformly dispersed and nanosized Pt particles can effectively capture electrons during electron–hole separation, thereby prolonging the electron–hole separation time and ultimately improving the photocatalytic efficiency of HCHO removal. Sun et al.⁶¹ reported that Pt particles would be photoactivated by visible light irradiation; then, the photoinduced electrons would be easily excited to the impurity band and conduction band. The result was also in agreement with reports made by Zhang et al.⁵ and suggested that nanoscale Pt⁰ particles would serve as electron sinks to reduce the recombination time of electron–hole pairs; then, an assisting effect would be created by the production of additional superoxide radical anions by electron transfer from TiO₂ to the surface of the TR catalysts. To confirm the effect of the Pt particles, the PL spectra of Pt/TR(10) were analyzed, as shown in Fig. 17. The intensity of the Pt/TR(10) spectra was weaker than that of the TR(10) spectra. This result correlated with the longer separation time for electron–hole pairs noted for Pt/TR(10). The argument presented above was thus confirmed.

4. Conclusions

TiO₂ is a common photocatalyst and is widely used in photocatalytic reactions performed under UV light with a wavelength of 325 nm. To expand the range of photocatalysis applications and to reduce their deleterious effects on health, in this study, TiO₂-rutile-based catalysts were synthesized using the one-step modified hydrothermal method and then doped with metallic Pt particles. The resulting photocatalysts provided greater efficiency than that of traditional P25 photocatalysts, even under visible light. The following conclusions could be drawn based on the results obtained in this study:

(1) The shape and structure of TR catalysts with various Ti proportions synthesized using the modified hydrothermal method differed from those of the porous material in traditional SBA-15. When the ratio of Ti/Si was 1/9, TR(10)
maintained a two-dimensional hexagonal channel structure; however, Ti–O–Si bonds enlarged the TR(10) structure. When the ratio of Ti/Si was 3/7 or 5/5, a three-dimensional porous spherical structure was formed instead of a two-dimensional channel structure, with small particle sizes of less than 1 μm.

(2) XRD and Raman analyses showed that the rutile phase was the primary crystalline species in the TR catalysts. Si atoms inhibited the transformation of the anatase phase and improved the thermal stability of the TR catalysts during calcination. Raman and XPS analysis results demonstrated that the Ti–O–Si–C bond contributed to the transformation of the two-dimensional structure to the three-dimensional structure during TR preparation. The mesoporous structure of the TR catalysts improved their photocatalytic activity owing to its large pores volume and high BET. DRS analysis showed that a higher rutile phase content and smaller crystal size of the rutile phase of the TR catalysts contributed to the lower band gap energy.

(3) The activity test results showed that when HCHO was used as the reactant, the TR catalysts were removed HCHO more effectively than the commercial photocatalyst P25 regardless of whether the reaction occurred under UV or visible light. The mesoporous structure and surface-absorbed oxygen of the TR catalyst might increase the adsorption of HCHO, improving its photocatalytic activity. Then, photoexcitation would occur easily because of the low band gap energy of the TR catalyst (≤3.0 eV) under visible light irradiation.

(4) The microwave-assisted EG reduction method was shown to effectively disperse nanoscale Pt particles with low energy consumption and a short synthesis time. After the Pt/TR catalyst underwent calcination, the TR catalyst structure and the crystalline Ti remained unchanged and undamaged, demonstrating the excellent stability of TR, and the chemical state of platinum was Pt⁴⁺. The Pt particles could increase the electron–hole pair separation time, thereby effectively improving the catalytic efficiency of the photocatalysts.

(5) The size of the TR catalysts followed the order TR(10) (10 μm) > TR(30) (8 μm) > TR(50) (0.5 μm). The Taguchi method was employed to study the effect factors for the microscale TR catalysts during the photocatalytic removal of HCHO. The results of ANOVA and the analysis of the S/N ratio were the same and showed that the level effect followed the order space velocity > ratio of Ti > time on stream.

Conflicts of interest
There are no conflicts to declare.

Acknowledgements
This work was performed under the auspices of the Ministry of Science and Technology, ROC, under Contract MOST 98-2221-E-040-001-MY3 and MOST 102-2221-E-040-011-MY3, to which the authors wish to express their thanks.

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