# RSC Advances



# PAPER

Cite this: RSC Adv., 2020, 10, 2757

# Mechanistic study of N–H- and H–N-codoping of a TiO<sub>2</sub> photocatalyst for efficient degradation of benzene under visible light†

Minghui Li[,](http://orcid.org/0000-0002-6057-5135) **D**<sup>ab</sup> Wulin Song, D<sup>\*ab</sup> L[e](http://orcid.org/0000-0001-5123-3195)i Zeng,<sup>a</sup> Dawen Zeng, D<sup>a</sup> Changsheng Xie D<sup>a</sup> and Qun Yang<sup>c</sup>

Non-metal codoping including nitrogen (N) and hydrogen (H) codoping has been emerging as an effective way to improve the performance of anatase  $TiO<sub>2</sub>$  in solar cell, fuel conversion and pollutant degradation. However, the mechanism of the synergistic effect of N doping and H doping on  $TiO<sub>2</sub>$  is still far from thorough. In this paper, N and H codoped  $TiO<sub>2</sub>$  nanoparticles are obtained by N doping in ammonia and then H doping in hydrogen gas, which achieves substantially boosted efficiency and reaction rate in the photocatalytic degradation of benzene under visible light excitation. The superiority of the  $N-H-TiO<sub>2</sub>$ photocatalyst was fully elaborated by comparing with H–N–TiO<sub>2</sub>, which was obtained by thermal treating in  $H_2$  and then NH<sub>3</sub>. The reaction rate of N-H-TiO<sub>2</sub> in the photocatalytic degradation of benzene was nearly 2 times that of H–N–TiO<sub>2</sub>,  $\sim$ 7 times higher than that of pristine TiO<sub>2</sub>. Furthermore, the cycling test revealed the high repeatability and stability of the N-H-TiO<sub>2</sub> photocatalyst. The excellent performance  $N-H-TiO<sub>2</sub>$  was attributed to an adequate concentration of  $N_iH_i$  defects occupying interstitial sites of the  $TiO<sub>2</sub>$  structure and a disordered surface layer introduced by annealing in  $NH_3$  and H<sub>2</sub> successively. The synergistic effect of  $N-H$ -codoping also increased the separation and migration of electron–hole pairs triggering a photoinduced redox reaction on the surface of  $TiO<sub>2</sub>$ . **PAPER**<br> **EXERCTS AND A SUBLIC STATE CONDUCT THE CONDUCT CONDUCT THE CONDUCT CONDUCT** 

Received 19th November 2019 Accepted 27th December 2019

DOI: 10.1039/c9ra09661b

rsc.li/rsc-advances

# 1. Introduction

The breakthrough of water splitting on a  $TiO<sub>2</sub>$  photoelectrode has triggered great enthusiasm to develop  $TiO<sub>2</sub>$ -based photocatalysts to wrestle with growing environment contamination and the energy crisis.<sup>1</sup> TiO<sub>2</sub> possesses many significant advantages in photocatalytic applications (such as, suitable band energy, non-toxicity, stable to corrosion, and low cost). Unfortunately, the  $TiO<sub>2</sub>$  photocatalyst is extremely limited by its wide bandgap (anatase  $\sim$ 3.2 eV, rutile  $\sim$ 3.0 eV) as it can only absorb the UV fragment of sunlight and thus more than 95% of sunlight energy is not utilised.

To overcome these obstacles, tremendous efforts have been devoted to developing visible light-responsive  $TiO<sub>2</sub>$  by extrinsic element doping, intrinsic self-doping, semiconductor coupling, and noble metal modifications.<sup>2-5</sup> Among these approaches, N doping and H doping have shown the great potential to improve

the photocatalytic properties of  $TiO<sub>2</sub>$  by enhancing visible light absorption. For example, Asahi et al. developed a visible-light responsive TiO<sub>2</sub> photocatalyst by N doping,<sup>6</sup> it ushered a new era of modulation of optical absorption and band structure of  $TiO<sub>2</sub>$  by inter-bandgap element doping. This work revealed that the N doped at the substitutional sites of  $TiO<sub>2</sub>$  could narrow the bandgap, which enhanced the visible light absorption of  $TiO_{2-x}N_x$  and thus improved the photocatalytic activity. Since then, N doping becomes one of the most desirable methods to achieve the better performance of  $TiO<sub>2</sub>$  photocatalysts by forming the inter-bandgap states and enhancing the visible light absorption.<sup>7-10</sup> It was reported that N doping facilitated the formation of oxygen vacancies in  $TiO<sub>2</sub>$ , which could introduce mid-gap states below conduction band.<sup>11</sup> Previous researches also showed that the presence of N dopants benefited for the formation of oxygen vacancies, and meanwhile, the existence of oxygen vacancies might stabilize N impurities, which proved the correlation of N doping and defects formation.12,13 Besides N doping, H doping became more noteworthy as it was found the hydrogenated disorder-engineered  $TiO<sub>2</sub>$  nanocrystals showed excellent optical absorption capability to visible and even infrared lights.<sup>14</sup> The authors highlighted that the disordered surface layer was responsible for the enhanced light absorption and photocatalytic activity of TiO<sub>2</sub>. Since then, hydrogenation becomes a powerful approach to improve the performances of TiO2 nanomaterials.

a State Key Laboratory of Materials Processing and Die & Mould Technology, Huazhong University of Science and Technology, Wuhan, 430074, P. R. China. E-mail: wulins@ 126.com; Fax: +86 27 87792025; Tel: +86 27 87792025

<sup>&</sup>lt;sup>b</sup>Analytical and Testing Center, Huazhong University of Science and Technology, P. R. China

c Wuhan Maritime Communication Research Institute, P. R. China

<sup>†</sup> Electronic supplementary information (ESI) available. See DOI: 10.1039/c9ra09661b

However, N doping and H doping individually are not enough to obtain an efficient  $TiO<sub>2</sub>$  photocatalyst. One issue is the localized states formed by N doping increase the recombination centers of photogenerated electrons and holes, and the other is that visible light activity after H doping is far from practical applications. Consequently, various investigations have been carried out to the development of visible light responsive  $TiO<sub>2</sub>$ photocatalysts by N and H codoping, and then explain their advantages in enhancing light absorption, narrowing bandgap, improving the separation of electron–hole pairs, reducing the recombination centers and so on. For example, Hoang et al. suggested that the enhanced visible light activity of rutile (H, N)–  $TiO<sub>2</sub>$  nanowire arrays was due to the interaction of substitutional N dopants and  $Ti^{3+15}$  Meanwhile, Zhang et al. proposed that the photocatalytic activity of N–H-codoped  $TiO<sub>2</sub>$ was improved by the synergistic effect of oxygen vacancies and N dopants.<sup>16</sup> Moreover, Wang et al. suggested that hydrogenation could induced the defect energy states in  $TiO<sub>2</sub>$  by forming  $Ti<sup>3+</sup>$ and N dopants and combining with disordered structures, which contributed to the enhanced efficiency of hydrogenated N–F doped TiO2. <sup>17</sup> Beyond doubt, previous investigations in N and H codoping TiO<sub>2</sub> provide new insights into utilizing solar light and enlarging the scopes of applications. However, due to lack of comparison under a uniform standard, the mechanism of N and H codoping is still not really clarified. Moreover, though many efforts have been done to investigate the N and H codoping on photocatalytic applications, such as, water splitting and water purification, unsufficient attention has been paid to the applications of volatile organic compounds (VOCs).<sup>18</sup> Benzene, especially, as a typical indoor contaminant and strong cancerogen, is a degradation-resistant VOC gas and may cause serious health concerns. Thus, more efficient photocatalyts to remove it to purify the environment is urgent to develop.<sup>19</sup>–<sup>21</sup> RSC Advances<br>
Despert Article Typeoconysis one issue is the specific in the angular solution was transferred in a 20 nm. Teleform the common article is licensed under the state of the specific in the state of the specific

Given these challenges, we developed a highly efficient N–H– TiO2 photocatalyst for benzene removal utilizing visible light. In this work, the synergetic effect of N and H codoping on  $TiO<sub>2</sub>$ photocatalyst was deeply explored on the basis of the characterizations of structural and properties. The distinct performances of N–H–TiO<sub>2</sub> and H–N–TiO<sub>2</sub> in decomposing benzene under visible light were aroused and the mechanism was fully understood. The facile preparation and the inspiring performance of  $N-H-TiO<sub>2</sub>$  photocatalyst in benzene degradation make us confident in its extensive applications.

## 2. Experimental

### 2.1 Chemicals

All chemicals applied in this experiment including tetra-n-butyl titanate (TTIP, 98%; Sinopharm Chemical Reagent Co., Ltd.), ethanol, ammonia gas (NH<sub>3</sub>, 99.9999%) and H<sub>2</sub>/Ar (5% H<sub>2</sub>/95% Ar) gas mixture were purchased and used without any further purification.

#### 2.2 Synthesis of pure  $TiO<sub>2</sub>$

Typically, 2 mL TTIP was added dropwise to a beaker with 75 mL distilled water under vigorous magnetic stirring and then the final suspension solution was transferred into a 100 mL Teflonlined autoclave to react at 160  $^{\circ}$ C for 12 h. Afterwards, the resulting product centrifuged and rinsed with absolute ethanol and distilled water for several times, and then dried in an oven at 80 $\degree$ C overnight to acquire TiO<sub>2</sub> nanoparticles.

#### 2.3 Preparation of N and H codoping  $TiO<sub>2</sub>$  samples

The as-prepared pure  $TiO<sub>2</sub>$  was placed in a quartz glass tube and followed by thermal treatment in a tube furnace. The tube was filled with the desired gas for 30 min to purge the residual air before thermal treatment. Two kinds of gases, hydrogen and ammonia were used in this experiment. The flow rate of ammonia and hydrogen was 200 mL  $min^{-1}$  controlled by rotormeter. After thermal treatment, the samples were naturally cooled to room temperature. All the samples were treated at 600 °C with a ramp rate 5 °C min<sup>-1</sup>. The details were as follows: firstly, the pure TiO<sub>2</sub> was treated in  $H_2/Ar$  in the former 2 h and sequentially switched to  $NH<sub>3</sub>$  for the latter 2 h; the product was labeled as  $H-N-TiO<sub>2</sub>$ . In the same way, we exchanged the gas feed order of  $H_2/Ar$  and  $NH_3$  in the process of thermal treatment and N-H-TiO<sub>2</sub> was prepared. Secondly, to further investigate the importance of treating order and the effect of hydrogenation, N-H-TiO<sub>2</sub>-E was obtained by annealing in NH<sub>3</sub> for 2 h and then held in  $H_2$  for 4 h. Similarly, H–N–TiO<sub>2</sub>-E was obtained by treating in  $H_2$  for 4 h and NH<sub>3</sub> for 2 h.

#### 2.4 Characterizations

The structures and morphologies of the  $TiO<sub>2</sub>$  before and after the thermal treatment were investigated by X-ray diffraction (XRD, X'Pert powder) with Cu radiation, high-resolution transmission electron microscopy (HRTEM, FEI TecnaiG2 F30). Raman spectroscopy (Raman, LabRAM HR800) was investigated from 100–1000  $\text{cm}^{-1}$  by using a LabRAM HR800 spectrometer and operating at 532 nm. Fourier transform infrared (FT-IR, Bruker VERTEX 70) spectroscopy was investigated in the range of 400-4000  $cm^{-1}$ . Ultraviolet-visible diffuse reflectance spectra (UV-vis DRS, Shimadzu U-3010) was conducted using BaSO<sub>4</sub> as reference. Photoluminescence (PL) emission spectra were acquired under an excitation at 325 nm by using an Edinburgh Instruments PLSP920 spectrometer. The X-ray photoemission spectroscopy (XPS, Shimadzu AXIS-ULTRA DLD-600W) studies were examined to obtain information on the chemical bonding of the samples. The binding energy of XPS spectrum was corrected by the C 1s level at 284.6 eV.

#### 2.5 Photocatalytic activity test

The performance evaluation of the samples in our experiment was conducted in photocatalysis testing system, which was comprised of a home–made reaction chamber with an injection hole to input benzene and a sample loading plate, a gas chromatograph (GC9560) with hydrogen and nitrogen gas supply systems. The light source was a 300W Xe-arc lamp (CEL-HXF300) with a UV cut-off filter ( $\lambda > 400$  nm). The irradiance of the lamp in the visible region was 1200 mW  $cm^{-2}$ . The test was carried out by monitoring the degradation of benzene and the yield of  $CO<sub>2</sub>$  under visible light irradiation within 4 h to

estimate the photocatalytic activity. Generally, 100 mg  $TiO<sub>2</sub>$ catalyst was dispersed on the loading plate  $(50 \times 50 \text{ mm}^2)$  and then injected benzene into the reaction chamber (2 L) when the GC attained the running state according to the operation directions. Finally, we turned on the light source and then examined the photocatalytic activity of  $TiO<sub>2</sub>$  samples by monitoring the change of benzene concentration and the yield of  $CO<sub>2</sub>$ every 15 min intervals in 4 hours. The degradation rate of benzene was calculated by  $C/C_0$ , where C was the concentration of benzene in each time period, and  $C_0$  was the initial concentration of benzene after adsorption equilibrium but before irradiation. Meanwhile, the  $CO<sub>2</sub>$  production was calculated by  $C-C_0$  (unit: ppm), where C was the concentration of  $CO_2$  in each time period, and  $C_0$  was the concentration of  $CO_2$  after adsorption equilibrium but before irradiation. A blank control test without photocatalyst was performed for reference.

## 3. Results and discussion

### 3.1 Basic structural information

The influence of  $N$  doping and  $H$  doping on the structure of  $N-$ H-TiO<sub>2</sub> and H-N-TiO<sub>2</sub> were manifested by XRD and FTEM. High temperature annealing in  $NH<sub>3</sub>$  and  $H<sub>2</sub>$  improves the crystallinity of N–H–TiO<sub>2</sub> and H–N–TiO<sub>2</sub>, as the XRD results shown in Fig. 1. It is obvious that all the TiO<sub>2</sub> samples consist of anatase phase (JCPDS file no. 21-1272) and brookite phase (JCPDS file no. 03-0380). However,  $H-N-TiO<sub>2</sub>$  obtained by H doping contains rutile phase (JCPDS file no. 21-1276). In addition, there is surface disordered layers outside the core of  $H_2$ treated  $TiO<sub>2</sub>$  nanoparticles, as shown by the dotted line of FTEM images in Fig. 2. It confirms the special structure induced by hydrogenation effect. The increase in crystallinity and the unchanged phase structure of  $N-H-TiO<sub>2</sub>$  indicate the distinct effect of N–H- and H–N-codoping on the phase transition of TiO<sub>2</sub>. XRD results confirm that N doping is beneficial to suppress the phase transformation from anatase to rutile, and the following H doping can induce a disordered surface layer.<sup>22</sup>



Fig. 1  $XRD$  patterns of TiO<sub>2</sub> samples.



Fig. 2 FTEM and HRTEM images of (a)  $TiO<sub>2</sub>$ , (b)  $H-TiO<sub>2</sub>$ , (c)  $N-TiO<sub>2</sub>$ , (d)  $N-H-TiO<sub>2</sub>$ , (e)  $H-N-TiO<sub>2</sub>$ , (f)  $N-H-TiO<sub>2</sub>-E.$ 

Instead, in H–N–TiO<sub>2</sub>, H doping first facilitates the phase transformation, and thus increases the numbers of crystal boundaries and bulk defects, which act as the recombination centers of photogenerated electron–hole pairs and thus deteriorate the photocatalytic performance of  $H-N-TiO<sub>2</sub>.<sup>23-25</sup>$ Therefore, N doping followed by H doping is a favorable stratagem to reduce bulk defects and then improves the crystallinity and the stability of anatase  $TiO<sub>2</sub>$ , and it also infers that anatase TiO<sub>2</sub> behaves better photocatalytic activity than rutile TiO<sub>2</sub>.<sup>26,27</sup>

Additionally, the average particle sizes of  $TiO<sub>2</sub>$ , H–N–TiO<sub>2</sub> and N-H-TiO<sub>2</sub> are estimated to be 22.80, 44.21 and 37.70 nm respectively by the Scherrer formula, consistent with the FTEM images presented in Fig. 2. It is evident that the pristine  $TiO<sub>2</sub>$ nanoparticles have the smallest size and the worst crystallinity. The average grain size and the crystallinity of  $H-N-TiO<sub>2</sub>$  and N-H-TiO<sub>2</sub> increase moderately, and the grain growth of N-H-TiO<sub>2</sub> can be restrained by the ordering treatment in  $NH<sub>3</sub>$  and then  $H<sub>2</sub>$ . Moreover, the H-N-TiO<sub>2</sub> nanoparticles agglomerate severely which leads to its inferior performance. This may be ascribed to the thicker disordered surface layer and the worse bulk crystallinity of H-N-TiO<sub>2</sub> than N-H-TiO<sub>2</sub>, as shown in Fig. 2d and e. According to the different microstructures of N-H-TiO<sub>2</sub> and H- $N-TiO<sub>2</sub>$ , it can be deduced that the formation process of disordered layer is different from each other. This special surface layer may be generated in the high temperature annealing process and finally formed in the cooling step. Thus, the different annealing orders in  $H_2$  result in the different bulk structures of N-H-TiO<sub>2</sub> and H-N-TiO<sub>2</sub>. It could be because that H doping would introduce bulk defects, and after N doping, the formed disordered layer may restrain the migration of bulk defects and then lead to the inferior crystallinity of H-N-TiO2. However, N doping increases the crystallinity and decreases the

bulk defects significantly, then after H doping, the H atoms may occupy the interstitial position  $(H<sub>i</sub>)$  bonding with N atoms, which may maintain the advantages of both N doping and H doping.

#### 3.2 Raman spectroscopy

Fig. 3a shows the Raman spectra of  $TiO<sub>2</sub>$  samples. Six typical Raman active modes of anatase TiO<sub>2</sub> occurs at 144 cm<sup>-1</sup> (E<sub>g</sub>), 197 cm<sup>-1</sup> (E<sub>g</sub>), 397 cm<sup>-1</sup> (B<sub>1g</sub>), 518 cm<sup>-1</sup> (A<sub>1g</sub> + B<sub>1g</sub>), and 640 cm<sup>-1</sup> (E<sub>g</sub>).<sup>28</sup> The enlarged Raman spectra in Fig. 3b shows the peak at 440 cm<sup>-1</sup> in H-TiO<sub>2</sub> and 443 cm<sup>-1</sup> in H-N-TiO<sub>2</sub> and  $N-H-TiO<sub>2</sub>$ , which can be ascribed to the Raman modes of rutile.<sup>29</sup> More importantly, these peaks only appear in H doping samples. It proves the successful doping of H atoms after annealing in  $H_2$  atmosphere. In addition, the weak two-phonon scattering bands at about 324  $cm^{-1}$  assigned to anatase Raman feature and the band near 248  $cm^{-1}$  ascribed to A1g mode of rutile can be observed.<sup>30</sup> Generally, the Raman band below 400  $\text{cm}^{-1}$  is ascribed to the Ti-O-Ti bending type and those above belongs to the Ti-O stretching type vibrations.<sup>28</sup> However, the new peak at 480  $cm^{-1}$  has never been found in other researches. Because it has been confirmed that N doping play little influence on the Ti-O stretching by comparing the Raman spectra of H–TiO<sub>2</sub> and H–N–TiO<sub>2</sub>, these emerging changes at 480 cm<sup>-1</sup> in H-TiO<sub>2</sub>, N-H-TiO<sub>2</sub> and H-N-TiO<sub>2</sub> may be induced by H-doping effect. Thus, the sharp peak of  $N-H-TiO<sub>2</sub>$  at  $480 \text{ cm}^{-1}$  provides an evidence for the better synergetic effect of N–H-codoping than H–N-codoping because its higher intensity than that of the N–H bond in N–H–TiO<sub>2</sub> photocatalyst. **PSC Advances**<br>
Unlk defects aignificantly, then after H doping the H atoms are y the mix-yinter respectively. Presidents respected to the same that the common common in the same of the term in the common common common com

### 3.3 FTIR spectroscopy

To investigate the chemical information on the surface functional groups of  $TiO<sub>2</sub>$  samples, the FT-IR transmittance spectra were measured, as shown in Fig. 4. Besides, Fig. 5a and b are the enlarged spectra in the range of 1200-1800  $cm^{-1}$  and 400-700  $\text{cm}^{-1}$  respectively. Previous research has shown that the signals in the range of 400–1250  $cm^{-1}$  are characteristic of O– Ti-O lattice and the peak near  $1628 \text{ cm}^{-1}$  is caused by Ti-O structure.<sup>29,31</sup> In addition, the broad absorption presented at 2900–3600  $\text{cm}^{-1}$  corresponds to O–H stretch.<sup>32</sup> Generally, the typical infrared bands of NO<sub>2</sub> are detected at 1618 cm $^{-1}.$ <sup>33</sup> The  $NO<sub>2</sub>$  band cannot be detected in N doping TiO<sub>2</sub> but instead in  $N-H-TiO<sub>2</sub>$ . Furthermore, as shown in Fig. 5a, the absorption peak at 1402 cm<sup>-1</sup> in N-TiO<sub>2</sub> shifts to 1385 cm<sup>-1</sup> in N and H codoping  $TiO<sub>2</sub>$ , which implies the effect of H doping. Considering the N doping effect, the peaks at about 1385  $cm^{-1}$ in N–H–TiO<sub>2</sub> and H–N–TiO<sub>2</sub> infer the formation of impurity defects like N–H bond in the codoping  $TiO<sub>2</sub>$  lattice. Moreover, Fig. 5b presents the consistent shift direction of  $N-TiO<sub>2</sub>$  and



Fig. 4 FT-IR patterns of  $TiO<sub>2</sub>$  samples.



Fig. 3  $\,$  (a) Raman spectra of TiO $_2$  samples; (b) enlarged Raman spectra from 180 to 600 cm $^{-1}$ .



Fig. 5 Enlarged FT-IR patterns of TiO<sub>2</sub> samples at position (a) and (b) of Fig. 4.

N–H–TiO<sub>2</sub> compared to H–TiO<sub>2</sub> and H–N–TiO<sub>2</sub>. It indicates the distinct effect of N–H- and H–N-codoping on structure and defect states.

#### 3.4 Chemical states

XPS analysis is carried out to determine the chemical states of  $TiO<sub>2</sub>$  samples. N 1s peaks of N or (N, H) doping  $TiO<sub>2</sub>$  are



Fig. 6 (a–d) N 1s XPS spectra of N–TiO<sub>2</sub>, N–H–TiO<sub>2</sub> and H–N–TiO<sub>2</sub>.

Table 1 Atomic concentration of N atoms (at%) of  $N-TiO<sub>2</sub>$ ,  $N-H TiO<sub>2</sub>$  and  $H-N-TiO<sub>2</sub>$ 

Sample	$N_s$ (a) 396 eV	$N_i$ (a) $400$ eV	$N_i$ (a) 403 eV	Total
$N-TiO2$	0.08	0.37	0.11	0.56
$N-H-TiO2$		0.61	0.11	0.72
$H-N-TiO2$	0.09	0.40	0.11	0.6

presented in Fig. 6. The concentrations of N dopants calculated by N 1s peak area are listed in Table 1. It is generally believed that the doped N species results in the narrowed bandgap and the enhanced photocatalytic activity of  $TiO<sub>2</sub>$ , in which the N 1s peak at a binding energy around 396 eV is the characteristic of  $N<sub>s</sub>$  occupying substitutional sites in the form of TiN but the peak at a binding energies above 399 eV suggests the  $N_i$  dominating interstitial position attributed to NO or  $NO<sub>2</sub>$ ,  $10,34,35$ Obviously, only two peaks located at 399.7 eV and 402.5 eV in N 1s XPS spectral region in N-H-TiO<sub>2</sub>, indicating the interstitial nitrogen dopants. More importantly, the content of  $N_i$  in N-H-TiO<sub>2</sub> is higher than N–TiO<sub>2</sub> and H–N–TiO<sub>2</sub>. It is highly likely that the interaction between N and O is strong for the interstitial N doping in a type of  $NO<sub>x</sub>$  species.<sup>36</sup>

However, one more peak located at 396 eV appeared in H–N– TiO<sub>2</sub> and N-TiO<sub>2</sub>, which is usually attributed to N<sub>s</sub> species.<sup>6,37</sup> Previous theoretical calculations showed that the different treatment conditions could tend to result in the different doping positions of N atoms in  $TiO<sub>2</sub>$ , such as excess oxygen and nitrogen, and the doped N would contribute to the formation of Ni, and otherwise the strong reduction atmosphere would benefit the introduction of  $N_s$  and oxygen vacancies.<sup>38</sup> The theoretical inference is well consist with our experimental results. In our work, the  $N_i$  doping is definitely preferred after the annealing treatment in ammonia, while the  $N_s$  species coupled with oxygen vacancies is favoured under the highly reducing conditions such as the annealing treatment in  $H<sub>2</sub>$ . That's why more interstitial nitrogen species in N-H-TiO<sub>2</sub> when the N doping is firstly applied, while  $H-N-TiO<sub>2</sub>$  exists more substitutional nitrogen species when the H doping is firstly applied. Besides, the lower formation energy of  $N_i$  than  $N_s$ implies that  $N_i$  is more favorable in N-TiO<sub>2</sub>; but when phase transformation from anatase to rutile, it becomes more and more difficult to form  $N_i$  while the  $N_s$  doping becomes easier.<sup>39</sup> Thus, there are more  $N_s$  atoms in H–N–TiO<sub>2</sub> than N–H–TiO<sub>2</sub> because of phase transformation, which is consistent with the results of XRD and Raman spectroscopy. In other words, the H doping benefits the formation of  $N_s$ . Although  $N_s$  dopants can lead to narrowed bandgap, the resultant localized states may decrease the mobility of charge carriers and then lead to more recombination centres. From the perspective of energy, the N species in TiO<sub>2</sub> occupy the interstitial sites after the N doping.<sup>34</sup> However, when interstitial position reaches saturation, the excess N atoms may enter into substitutional sites to replace O atoms. Thus, N-TiO<sub>2</sub> contains both  $N_i$  and  $N_s$  dopants. After continuous hydrogenation, the surface of the  $N-H-TiO<sub>2</sub>$ 

nanoparticles becomes amorphous and the O atoms escape from the original lattice to form the surface oxygen vacancies, which makes the  $N_s$  atoms escape from the primary position and then enter the interstitial position of  $TiO<sub>2</sub>$  lattice again.

#### 3.5 PL emission

The room temperature PL spectra of  $TiO<sub>2</sub>$  samples are showed in Fig. 7. The PL spectra have been widely used to investigate the dynamic of electron–hole pairs in semiconductors because the PL emission is caused by the recombination of free carriers.<sup>40</sup> The results clearly indicate that a substantial increase of PL emission occurs in  $H$ -TiO<sub>2</sub>, which may result from its tremendous recombination centers including bulk oxygen vacancies and thicker disordered layer after the  $H_2$  annealing. However, as  $N-TiO<sub>2</sub>$  has a good crystallinity after N doping, its bulk defect density should be dramatically reduced compared with the pristine  $TiO<sub>2</sub>$ . In addition, higher PL emission intensity of H–N–  $TiO<sub>2</sub>$  suggests it owns more recombination centres than N-H-TiO2 because the recombination of charge carriers usually occurs at the grain boundary of two phases or in the bulk of nanomaterials.<sup>41</sup> Specially, H-N-TiO<sub>2</sub> exhibits a higher emission than N–TiO<sub>2</sub>, and N–H–TiO<sub>2</sub> exhibits a lower emission than H-TiO<sub>2</sub>. It further demonstrates that the H doping may lead to create more recombination centres due to the formation of bulk defects. Besides, it is generally accepted that N doping facilitates the formation of oxygen vacancies in  $\text{TiO}_2$ ,<sup>11</sup> and N doping first may reduce the formation energy of oxygen vacancies and in turn the formed O vacancies can stabilize the N impurities.<sup>38</sup> RSC Advances Common access Article of New Article is licensed in the original interior in the simulation of New Article is licensed in the original interior of New Articles Common Sources Are also associated to the simula

Thus, as to  $N-H-TiO<sub>2</sub>$ , the N doping could improve the stability of the H doping-introduced oxygen vacancies, and then facilitate the separation and the transfer of photogenerated charge carriers to participate in the surface photocatalytic reactions. However, when it comes to  $H-N-TiO<sub>2</sub>$ , much more defects are imported by the first H doping, but these defects cannot exist stably and then become recombination centers. Therefore, the higher PL intensity of H-N-TiO<sub>2</sub> can be assigned to its inferior bulk crystallinity and the phase transition from



Fig. 7 PL of  $TiO<sub>2</sub>$  samples.

anatase to rutile. Moreover, it does not necessarily that the more defects, the better photocatalytic activity because some surface defects can serve as charge carrier traps and absorption sites to prevent electron–hole pairs from recombination while some bulk defects tend to act as recombination sites to annihilate photogenerated electron–hole pairs.<sup>42</sup> It has been calculated that H doping may replace oxygen to form  $H_0$  dopants.<sup>43</sup> Though subsequent N doping can narrow the bandgap and then increase the visible light absorption, the deep energy states formed by  $N_iH_o$  in H–N–TiO<sub>2</sub> may promote the recombination of electron–hole pairs. On the contrary, more NiHi defects in N–  $H$ –TiO<sub>2</sub> would generate the shallow states above the top of valence band, which is beneficial to the separation of charge carriers.

#### 3.6 Optical property

Fig. 8 displays the light absorption ability and the estimated band gaps of TiO<sub>2</sub> samples. Compared to the pure TiO<sub>2</sub>, N-TiO<sub>2</sub> exhibits typical characteristic of N doping with the absorption from 400 nm to 500 nm and even extending to 550 nm.<sup>13</sup> In addition, the absorption edge of  $H$ –TiO<sub>2</sub> extending to infrared wavelength confirms the hydrogenation effect on the enhanced light absorption. $44$  After the N and H codoping, the samples behave the visible light absorption and the light harvesting of N–H–TiO<sub>2</sub> in UV region is better than H–N–TiO<sub>2</sub>, while their abilities of light harvesting are opposite in visible light region. The reason is perhaps due to that the H doping promotes the interstitial  $H_i$  bonding with the interstitial  $N_i$  and then form the  $N_iH_i$  defects in  $N-H-TiO_2$ ,<sup>39,43</sup> and thus, the visible light absorption is partially eliminated compared with the sample doped only with N species.

However, compared with  $N-H-TiO<sub>2</sub>$ , the concentration of N atoms in H-N-TiO<sub>2</sub> tends to become lower due to the surface disordered structure. Therefore, there are fewer  $N_iH_i$  bonds are formed in H-N-TiO<sub>2</sub>, however, compared with and finally it leads to the stronger visible light harvesting than  $N-H-TiO<sub>2</sub>$ . The results are also confirmed by XPS and PL. The bandgaps of N-H-TiO<sub>2</sub> and H-N-TiO<sub>2</sub> are estimated by Kubelka-Munk equation according to the UV-vis DRS spectrum. As shown in Fig. 8b, both their bandgaps are narrowed. It is because the new

states formed by an interaction of defect states introduced by N doping and H doping that changes the band structure of N–H–  $TiO<sub>2</sub>$ , which not only ensures the visible light absorption but also increases the photogenerated electron–hole separation and facilitates the electron migration to surface.<sup>43</sup> However, although with the relative higher light absorption, the deep energy level and the thicker surface disordered layer caused by H–N-codoping may result in more recombination of photoformed charge carriers in  $H-N-TiO<sub>2</sub>$  and lead to poorer activity than  $N-H-TiO<sub>2</sub>$ .

#### 3.7 Evaluation of photocatalytic activity

Fig. 9 shows the photocatalytic activity of  $TiO<sub>2</sub>$  samples when degrading benzene within 4 hours under visible light. In detail, Fig. 9a and b showed the degradation of benzene and the production of  $CO<sub>2</sub>$ . It reveals that N-H-TiO<sub>2</sub> exhibit the best photocatalytic activity. The excellent reaction rate constant and the stability of  $N-H-TiO<sub>2</sub>$  are presented in Fig. 9c and d, respectively. Generally, the photocatalytic degradation of pollution obeys the first-order kinetics, and the reaction rate constant could be calculated by the following equation:  $-\ln(C/$  $C_0$ ) =  $k_{\text{app}}t$ , where  $C/C_0$  is the normalized benzene concentration,  $t$  is the reaction time, and  $k$  is the reaction rate constant  $(10^{-3} \text{ min}^{-1})$ .<sup>45</sup> As shown in Fig. 9c, the rate constant of N-H-TiO<sub>2</sub> towards benzene is calculated to be 0.0193  $\mathrm{min}^{-1},$  which is more than twice the rate constant of  $H-N-TiO<sub>2</sub>$ , and especially, about 7 times higher than that of pristine  $TiO<sub>2</sub>$ . For the practical application of photocatalysts, the stability of photocatalyst is one of the key issues, such as the decrease of active sites on the surface of photocatalyst or the unsufficient photocatalytic activity after long time storage. Paper<br>
Macchess Articles. Nonetheight that the state article is detectively the comment of the state article is detectively the state interest are proposed to the state interest are expected by the state interest are expec

To evaluate the stability of N-H-TiO<sub>2</sub>, recycle experiments under same conditions were conducted. According results shown in Fig. 9d, N-H-TiO<sub>2</sub> photocatalyst exhibits excellent repeatability and stability. The slightly fluctuated data may be on account of the variation of the initial concentration of benzene. The different activity of N-H-TiO<sub>2</sub> and H-N-TiO<sub>2</sub> indicates that  $TiO<sub>2</sub>$  annealing in different atmospheres plays obviously different roles in the final products. The characterizations of microstructures and chemical states demonstrate



Fig. 8 (a) UV-vis DRS and (b) estimated band gap of  $TiO<sub>2</sub>$ 



Fig. 9 (a) Photocatlytic degradation of benzene (C/C<sub>0</sub>) by TiO<sub>2</sub> samples under visible light; (b) CO<sub>2</sub> production (C–C<sub>0</sub>) corresponding to Fig. 1a; (c) the reaction rate constants of all samples; (d) the cycling test of  $N-H-TiO<sub>2</sub>$  under visible light.

that phase transformation, surface disordered structure, and concentration of N species have influence on the performance of N–H–TiO<sub>2</sub> and H–N–TiO<sub>2</sub>, especially defect states introduced by N and H codoping. Moreover, stronger visible light absorption of N doping than N–H- or H–N-codoping reveals that property of N and H codoping is not the simple superposition of N doping and H doping. The excellent performance N–H–  $TiO<sub>2</sub>$  was attributed to adequate concentration of  $N<sub>i</sub>H<sub>i</sub>$  defects occupying interstitial sites of  $TiO<sub>2</sub>$  structure and disordered surface layer introduced by annealing in  $NH<sub>3</sub>$  and  $H<sub>2</sub>$  successively. The synergistic effect of N–H-codoping also increased the separation and migration of electron–hole pairs to trigger photoinduced redox reaction on surface of  $TiO<sub>2</sub>$ , confirmed by the PL results.

Furthermore, the contrast tests of N–H–TiO<sub>2</sub>-E and H–N–  $TiO<sub>2</sub>$ -E in Fig. S1† confirm the special influence of annealing atmosphere orders on the photocatalytic activity of  $TiO<sub>2</sub>$ . N–H–  $TiO<sub>2</sub>$ -E shows a better performance compared with H–N–TiO<sub>2</sub>-E in accordance with the results of N–H–TiO<sub>2</sub> and H–N–TiO<sub>2</sub>. The positive effect of H doping is confirmed by the contrast of H-N-TiO<sub>2</sub> and H–N–TiO<sub>2</sub>-E, as well as the difference of N–H–TiO<sub>2</sub> and N-H-TiO<sub>2</sub>-E, as shown in Fig. S2.<sup>†</sup> These results demonstrate

that the visible light activity of  $TiO<sub>2</sub>$  photocatalyst and the corresponding mineralization rate of benzene increase with the extension of annealing time in hydrogen gas. It proves the positive effect of H doping on the photocatalytic properties of  $TiO<sub>2</sub>$  nanoparticles.

#### 3.8 Mechanism discussion

A schematic diagram has been established to explain the photocatalytic reaction paths of the N-H-TiO<sub>2</sub> nanoparticles, as presented in Fig. 10. It clearly shows the related processes that finally decide the reduction and oxidation reactions on N-H- $TiO<sub>2</sub>$  photocatalyst under the visible light excitation. The crystallinity of  $N-H-TiO<sub>2</sub>$  nanoparticles is greatly enhanced and the internal defects are dramatically reduced after N doping, meanwhile the surface oxygen vacancies are introduced and stably exist after the H doping.

More importantly, the surface defective structure introduced by the H doping (defect states) stabilizes the N-doped structure and leads to the increase in the trapping sites and the surface reactive sites, which is beneficial to the photocatalytic reactions by reducing the recombination of charge carriers and



Fig. 10 Schematic diagram of energy states and photocatalytic reaction paths on  $N-H-TiO<sub>2</sub>$  nanoparticles.

increasing the absorption of small molecules like  $H_2O$  and  $O_2$ , so as to produce hydroxyl radicals (OH') or active oxygen  $\mathrm{(O_2}^{\text{-}-})$ to degrade organic pollutants like benzene, as shown in Fig. 10.

The bandgap of N–H–TiO<sub>2</sub> is mainly engineered by the new emerged NiHi states (doping states) above the valence band edge according to the N-H-codoping. The existence of  $N_i$  is beneficial to the incorporation of  $H_i$  and will facilitate the formation of  $N_iH_i$  defects in N–H–TiO<sub>2</sub>, which have been confirmed by Raman and FT-IR spectra. The  $N_iH_i$  defects in N- $H$ –TiO<sub>2</sub> give rise to the shallow states above VBM.<sup>40</sup> However, as to H-N-TiO<sub>2</sub>, the structure of disordered layer on the surface is formed by the H doping, which is similar to a fence on the surface and suppresses the subsequent  $N_i$  doping. Consequently, the concentration of interstitial  $N_i$  in H–N–TiO<sub>2</sub> is less than that in  $N-H-TiO<sub>2</sub>$ , which is confirmed by XPS results. In addition, the surface amorphous layer has a localized effect on the internal defects of nanoparticles, which blocks the charge carriers to transfer to the surface to participate in the following reactions. Therefore, the performance of  $H-N-TiO<sub>2</sub>$  is not fully improved after the H–N-codoping. In fact, the pristine  $TiO<sub>2</sub>$  has many defects because of its poor crystallinity; H may replace the lattice oxygen to form the  $H_0$  defects and then leave the bulk oxygen vacancies after the H doping. The N doping after H doping may cause the reduction of bulk defects due to the increased crystallinity, while the newly generated  $N_iH_0$  defects may result in the formation of deep states above valence band maximum (VBM), which not only increase the recombination centers of photogenerated charge carriers but also eliminate the positive effect on the H doping. Thus, the good cyrstallinity, higher concentration of  $N_iH_i$  dopants, lower recombination of photogenerated charge carriers, and the beneficial surface disordered layer synergistically contribute to the better photocatalytic activity of N-H-TiO<sub>2</sub> in degrading benzene under visible light. Paper Maccess Articles. Published on 15 Articles. Published on 14 Articles. Published on 14 Articles. The common Society per common and the set of the common and the set of the

# 4. Conclusion

In summary, a highly efficient photocatalyst N-H-TiO<sub>2</sub> for benzene degradation driven by visible light has been successfully synthesized by annealing in  $NH<sub>3</sub>$  and  $H<sub>2</sub>$  atmospheres. We have illustrated that the reverse annealing orders of  $NH<sub>3</sub>$  and  $H<sub>2</sub>$ 

induce the different structures of N–H–TiO<sub>2</sub> and H–N–TiO<sub>2</sub>, and thus results in their different photocatalytic performances. The excellent photocatalytic performance of N-H-TiO<sub>2</sub> is a final embodiment of the synergistic effect of good crystallinity, adequate concentration of  $N_iH_i$  defects and disordered surface layer, which lead to the lowered recombination of photogenerated charge carriers, increased amount of surface active sites and more surface absorption sites in  $N-H-TiO<sub>2</sub>$ . Through the investigation on the characteristics of N–H- and H–Ncodoping, we provide a fundamentally new perspective to elucidate the role of codoping effect on  $TiO<sub>2</sub>$  photocatalyst. The as-prepared N-H-TiO<sub>2</sub> photocatalyst by a facile method with low energy consumption is meaningful for expanding the application prospect in environmental improvements, especially in the elimination of volatile organic compounds.

# Conflicts of interest

There are no conflicts to declare.

# Acknowledgements

This work was financially supported by the National Basic Research Program of China (Grant No. 2009CB939705). The authors are also grateful to Analytical and Testing Center of Huazhong University of Science and Technology.

## References

- 1 A. Fujishima and K. Honda, Nature, 1972, 238, 37–38.
- 2 F. Zuo, L. Wang, T. Wu, Z. Zhang, D. Borchardt and P. Feng, J. Am. Chem. Soc., 2010, 132, 11856–11857.
- 3 Y. Wu, J. Zhang, L. Xiao and F. Chen, Appl. Catal., B, 2009, 88, 525–532.
- 4 J. Pan, Z. Dong, B. Wang, Z. Jiang, C. Zhao, J. Wang, C. Song, Y. Zheng and C. Li, Appl. Catal., B, 2019, 242, 92–99.
- 5 Y. Cao, Z. Xing, Y. Shen, Z. Li, X. Wu, X. Yan, J. Zou, S. Yang and W. Zhou, Chem. Eng. J., 2017, 325, 199–207.
- 6 R. Asahi, Science, 2001, 293, 269–271.
- 7 G. Barolo, S. Livraghi, M. Chiesa, M. C. Paganini and E. Giamello, J. Phys. Chem. C, 2012, 116, 20887–20894.
- 8 Y. Chen, X. Cao, B. Gao and B. Lin, Mater. Lett., 2013, 94, 154–157.
- 9 M. Khan and W. Cao, J. Mol. Catal. A: Chem., 2013, 376, 71– 77.
- 10 T. Boningari, S. N. R. Inturi, M. Suidan and P. G. Smirniotis, Chem. Eng. J., 2018, 350, 324–334.
- 11 A. Fujishima, X. Zhang and D. Tryk, Surf. Sci. Rep., 2008, 63, 515–582.
- 12 S. Livraghi, M. C. Paganini, E. Giamello, A. Selloni, C. Di Valentin and G. Pacchioni, J. Am. Chem. Soc., 2006, 128, 15666–15671.
- 13 Y. Chen, X. Cao, B. Lin and B. Gao, Appl. Surf. Sci., 2013, 264, 845–852.
- 14 X. Chen, L. Liu, P. Y. Yu and S. S. Mao, Science, 2011, 331, 746–750.
- 15 S. Hoang, S. P. Berglund, N. T. Hahn, A. J. Bard and C. B. Mullins, J. Am. Chem. Soc., 2012, 134, 3659–3662.
- 16 Z. Zhang, J. Long, X. Xie, H. Zhuang, Y. Zhou, H. Lin, R. Yuan, W. Dai, Z. Ding, X. Wang and X. Fu, Appl. Catal., A, 2012, 425–426, 117–124.
- 17 W. Wang, C. Lu, Y. Ni, M. Su and Z. Xu, Appl. Catal., B, 2012, 127, 28–35.
- 18 W. Lin, X. Xie, X. Wang, Y. Wang, D. Segets and J. Sun, Chem. Eng. J., 2018, 349, 708–718.
- 19 J. Wang, H. Ruan, W. Li, D. Li, Y. Hu, J. Chen, Y. Shao and Y. Zheng, J. Phys. Chem. C, 2012, 116, 13935–13943.
- 20 H. Huang, D. Li, Q. Lin, W. Zhang, Y. Shao, Y. Chen, M. Sun and X. Fu, Environ. Sci. Technol., 2009, 43, 4164–4168.
- 21 P. Devaraji, N. K. Sathu and C. S. Gopinath, ACS Catal., 2014, 4, 2844–2853.
- 22 M. Li, W. Song, L. Zeng, D. Zeng and C. Xie, Mater. Lett., 2014, 136, 258–261.
- 23 D. A. H. Hanaor and C. C. Sorrell, J. Mater. Sci., 2011, 46, 855– 874.
- 24 S.-D. Mo and W. Y. Ching, Phys. Rev. B: Condens. Matter Mater. Phys., 1995, 51, 13023–13032.
- 25 T. Luttrell, S. Halpegamage, J. Tao, A. Kramer, E. Sutter and M. Batzill, Sci. Rep., 2014, 4, 4043.
- 26 K. Nagaveni, M. S. Hegde, N. Ravishankar, G. N. Subbanna and G. Madras, Langmuir, 2004, 20, 2900–2907.
- 27 B. Sun, P. G. Smirniotis and P. Boolchand, Langmuir, 2005, 21, 11397–11403.
- 28 T. Ohsaka, F. Izumi and Y. Fujiki, J. Raman Spectrosc., 1978, 7, 321–324.
- 29 X. Chen, Y.-B. Lou, A. C. S. Samia, C. Burda and J. L. Gole, Adv. Funct. Mater., 2005, 15, 41–49.
- 30 G. A. Tompsett, G. A. Bowmaker, R. P. Cooney, J. B. Metson, K. A. Rodgers and J. M. Seakins, J. Raman Spectrosc., 1995, 26, 57–62.
- 31 H. Li, J. Li and Y. Huo, J. Phys. Chem. B, 2006, 110, 1559– 1565.
- 32 M. Zheng, M. Gu, Y. Jin and G. Jin, Mater. Sci. Eng., B, 2000, 77, 55–59.
- 33 T. Shimanouchi, J. Phys. Chem. Ref. Data, 1977, 6, 993–1102.
- 34 C. Di Valentin, E. Finazzi, G. Pacchioni, A. Selloni, S. Livraghi, M. C. Paganini and E. Giamello, Chem. Phys., 2007, 339, 44–56.
- 35 J. Ananpattarachai, P. Kajitvichyanukul and S. Seraphin, J. Hazard. Mater., 2009, 168, 253–261.
- 36 J. A. Rodriguez, T. Jirsak, J. Dvorak, S. Sambasivan and D. Fischer, J. Phys. Chem. B, 2000, 104, 319–328.
- 37 H. Irie, Y. Watanabe and K. Hashimoto, J. Phys. Chem. B, 2003, 107, 5483–5486.
- 38 C. Di Valentin, G. Pacchioni, A. Selloni, S. Livraghi and E. Giamello, J. Phys. Chem. B, 2005, 109, 11414–11419.
- 39 H. Pan, Y.-W. Zhang, V. B. Shenoy and H. Gao, J. Phys. Chem. C, 2011, 115, 12224–12231.
- 40 H. Tang, K. Prasad, R. Sanjinès, P. E. Schmid and F. Lévy, J. Appl. Phys., 1994, 75, 2042–2047.
- 41 L. Zeng, Z. Lu, J. Yang, M. Li, W. Song and C. Xie, Appl. Catal., B, 2015, 166–167, 1–8.
- 42 M. Kong, Y. Li, X. Chen, T. Tian, P. Fang, F. Zheng and X. Zhao, J. Am. Chem. Soc., 2011, 133, 16414–16417.
- 43 M. Vasilopoulou, N. Kelaidis, E. Polydorou, A. Soultati, D. Davazoglou, P. Argitis, G. Papadimitropoulos, D. Tsikritzis, S. Kennou, F. Auras, D. G. Georgiadou, S.-R. G. Christopoulos and A. Chroneos, Sci. Rep., 2017, 7, 17839. PSC Advances Commons Articles. Articles. Articles. Published on 15 January 2020. Downloaded on 16 January 2020. The state is likensed under a creative Commons Articles. Are also the state is likensed under a creative Commo
	- 44 O. Elbanna, M. Fujitsuka, S. Kim and T. Majima, J. Phys. Chem. C, 2018, 122, 15163–15170.
	- 45 Z. Lu, L. Zeng, W. Song, Z. Qin, D. Zeng and C. Xie, Appl. Catal., B, 2017, 202, 489–499.