New insight into effects of 4d transition metals doping on the photocatalytic activities of anatase TiO2(101) surface

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Abstract

Aiming at improving the visible-light photocatalytic activities of TiO2(101) surface (TiOS) we make an in-dept study on the TiOS doped with 4d transition metal (TM) atoms. It is shown that the 4d TM dopings can not only produce new impurity energy bands in the bandgap but also result in the semiconductor-metal phase transition. Consequently, the visible-light absorption is strongly strengthened due to the dopings of Y, Zr, Nb, Mo, and Ag, while it is only weakly improved for Tc, Ru, Rh, Pd, and Cd dopings. The improvement in visible-light absorption can be attributed to the intraband or interband transition of electrons. Moreover, the photocatalytic activities are explored, and we find Y and Ag dopings can effectively enhance the photocatalytic activity of TiOS. Thus the mechanism of improving photocatalytic activity of TiOS has been clearly addressed, which is beneficial to further experimental and theoretical researches on TiO2 photocatalysts.

1. Introduction

TiO2, a promising semiconductor photocatalytic material, has attracted extensive attention due to its advantages, such as the biological and chemical inertness, stability to corrosion, nontoxicity, relatively low cost, and high photoactivity [1–3]. In nature, there exist three phases of TiO2: rutile, anatase, and brookite TiO2. Among them anatase TiO2 is of the best photocatalytic activity and is widely applied in solar energy conversion, environmental purification, and particularly the degradation of harmful organic pollutants in water [4–6]. However, the popular application of anatase TiO2 is severely restricted due to the following two reasons [7]: (i) The band gap $E_g$ of 3.23 eV is so large that it leads to the low utilization efficiency of visible light; (ii) The quantum yield is very low because of the low electron transfer rate and the high recombination rate of photoexcited electron-hole pairs. Therefore, it is still urgent to improve the photocatalytic activity of TiO2.

In the past decades, much effort has been devoted to extending the optical response range of TiO2 to the visible-light region, by means of impurity doping [8–14], noble metal loading [15], semiconductor compounding [16–18], and organic sensitizing [19,20]. Among all these methods, impurity doping is considered to be one of the most convenient and efficient methods. In particular, transition metal (TM) atoms have been proven to be the ideal dopants to promote the photocatalytic performance of TiO2 catalysts because of their d electronic configuration and unique characteristics. Quantum-sized TiO2 doped with 0.5 at.% Fe³⁺, Mo⁵⁺, Ru³⁺, Os³⁺, Re⁵⁺, V⁴⁺, and Rh³⁺ are found to induce the red shift of absorption edge and can improve photoactivity for both CHCl₃ oxidation and CCl₄ reduction [21]. Mesoporous TiO2 impregnated by Ag⁺, Co²⁺, Cu²⁺, Fe³⁺, and Ni²⁺ can accelerate the degradation of acetophenone and nitrobenzene in...
aqueous solution [22]. TiO$_2$ powders doped by a small amount of Fe$^{3+}$ can clearly enhance the intensity of absorption in the UV-visible light region and the photocatalytic oxidation of acetone in air [23]. Moreover, Ni 8 wt% doped TiO$_2$ has a high photocatalytic activity for the decomposition of 4-chlorophenol in aqueous solution in the presence of both UV and visible light [24]. All the above researches have demonstrated that the TM dopants can extend the optical response range of TiO$_2$ into the visible-light region. However, we want to stress that the above dopings are mainly performed in bulk TiO$_2$.

In fact, surface structure will play an important role in the photocatalytic activity because usually photocatalytic reactions mainly occur on the catalyst surface [25]. It is especially necessary to study the effect of the surface manipulation on the catalytic activity. Hebenstreit et al. [26] found that anatase TiO$_2$(101) surface (TiOS) with few point defects is stable and the fourfold coordinated Ti atoms at step edges are the preferred adsorption sites. By investigating some low-index TiO$_2$ surfaces Labat et al. [27] demonstrated that the TiOS is the most stable surface. Also, Ma et al. [28] found that the TiOS structure with its outermost and second layers terminated by twofold coordinated O atoms and fivefold coordinated Ti atoms, respectively, is much more stable. Although many researches on the TiOS have been carried out, a systematic investigation on the photocatalytic activities of TiOS doped by all 4d TM atoms is still absent. A comparative study of all the 4d TM dopings on TiOS is required to clarify the intrinsic relation between the macroscopic photocatalytic activities and the microscopic TiO$_2$ details.

In this work, we will systematically investigate the surface modification of TiOS doped with 4d TM atoms to uncover how the doping will improve the photocatalytic performance of TiOS. We will first study the geometric structure, the doping manners, and the optical properties of the doped TiOS. Then, the relation of the macroscopic catalytic activity with the microscopic structure is discussed according to the density of states, energy bands, and the effective masses of carriers.

2. Computational methods

The geometric structure, the electronic structure, and the optical properties of the TiOS doped with ten 4d TM atoms Y, Zr, Nb, Mo, Tc, Ru, Rh, Pd, Ag, and Cd are calculated using the plane wave method based on the density functional theory (DFT). First, all the TiOS models doped in different doping manners are optimized, and the optimal doping manners of all TM atoms can be determined according to the impurity formation energy. Second, the band structures, the density of states, and the optical absorption coefficients of all the best doping models are calculated. All the calculations are performed with the Cambridge Serial Total Energy Package (CASTEP) [29]. The exchange-correlation potential is described by the Perdew-Burke-Ernzerhof (PBE) functional of the generalized gradient approximation (GGA) [30]. The cutoff energy for the plane wave basis set is set to be 340 eV. The Monkhorst-Pack scheme k-point grid sampling for the reduced Brillouin zone is set as $2 \times 4 \times 1$, and the convergence criteria for the self-consistent field (SCF) is set to $1.0 \times 10^{-6}$ eV/atom. All the atom coordinates are fully optimized until the forces on every atom are smaller than 0.03 eV/Å. The interaction between the valence electrons and the ionic core is described by the ultrasoft pseudopotential [31].

The formation energy is defined as

$$E(M) = E(M-TiO_2) - E(TiO_2),$$

(1)

where $E(M)$ is the energy of an isolated TM atom, and $E(M-TiO_2)$ is the total energy of the pure [TM-atom-doped] TiOS. For convenience, all of the optimal doping models can be denoted as Ti$_{24}$MO$_{48}$ with M representing the TM atoms Y, Zr, Nb, Mo, Tc, Ru, Rh, Pd, Ag, and Cd.

As widely recognized, the transfer rate of charge carriers can play a vital role in determining the photoactivity of a semiconductor material [32,33], and can be defined as [32,34]

$$v = \frac{\hbar k}{m^*},$$

(2)

where $\hbar$ is the reduced Planck constant, $k$ is the wave vector, and $m^*$ is the effective mass of charge carrier. The effective masses of electrons ($m_e^*$) and holes ($m_h^*$) can be obtained by fitting the energy band around...
the valence band maximum (VBM) and the conduction band minimum (CBM) along a specific direction in the reciprocal space, respectively. It should be noticed that the region used for parabolic fitting is kept in a very small interval to ensure the validity of the parabolic approximation. In addition, to clarify the difference between the effective masses of electrons and holes, an effective mass ratio \( r_{he} \) is introduced as 

\[ r_{he} = \frac{m_h^*}{m_e^*}. \]  

(3)

In general, a larger \( r_{he} \) value indicates \( m_h^* > m_e^* \), resulting in a larger difference of the carrier velocities and a slower recombination rate of the photogenerated electron-hole pairs [32].

3. Results

3.1. Optimal doping manners

Fig. 1 shows the anatase TiO\(_2\) unit cell in which there exist four titanium atoms and eight oxygen atoms. The lattice parameters \( a, c, d_{eq}, d_{ap}, c/a, u (=d_{ap}/c) \), \( 2\theta \), and the unit cell volume \( V_0 \) (see Fig. 1a) are listed in Table 1 so as to compare with the experimental values [35] and other theoretical results [36]. Clearly, the lattice parameters obtained in this work are in better agreement with the experimental data than other theoretical values, demonstrating the reliability of the calculation method and the models adopted here.

Fig. 1 (Color online) (a) The optimized unit cell of anatase TiO\(_2\). (b) Different doping manners on TiOS.

Table 1 Optimized lattice parameters of the anatase TiO\(_2\) unit cell in comparison with experimental and theoretical results.

<table>
<thead>
<tr>
<th>Lattice parameters</th>
<th>Exp.[35]</th>
<th>This work</th>
<th>Theo.[36]</th>
</tr>
</thead>
<tbody>
<tr>
<td>( a ) (Å)</td>
<td>3.782</td>
<td>3.743</td>
<td>3.692</td>
</tr>
<tr>
<td>( c ) (Å)</td>
<td>9.502</td>
<td>9.483</td>
<td>9.471</td>
</tr>
<tr>
<td>( d_{eq} ) (Å)</td>
<td>1.932</td>
<td>1.914</td>
<td>1.893</td>
</tr>
<tr>
<td>( d_{ap} ) (Å)</td>
<td>1.979</td>
<td>1.970</td>
<td>1.948</td>
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<tr>
<td>( c/a )</td>
<td>2.512</td>
<td>2.534</td>
<td>2.566</td>
</tr>
<tr>
<td>( u )</td>
<td>0.208</td>
<td>0.208</td>
<td>0.206</td>
</tr>
<tr>
<td>( 2\theta ) (°)</td>
<td>156.3</td>
<td>155.8</td>
<td>154.4</td>
</tr>
<tr>
<td>( V_0 ) (Å(^3)/TiO(_2))</td>
<td>33.98</td>
<td>33.21</td>
<td>32.27</td>
</tr>
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</table>

Starting from the optimized anatase TiO\(_2\) unit cell, we construct the TiOS model with the outermost and second layers terminated by twofold coordinated O atoms and fivefold coordinated Ti atoms, respectively. The doping concentration of TiOS doped with TM atoms is 1.37 at.% and six typical doping manners in Fig. 1b are considered. A TM atom can be vertically adsorbed on a twofold coordinated O atom (O1-Ad), threefold coordinated O atoms (O2-Ad, O3-Ad), a fivefold coordinated Ti atom (Ti1-Ad), and a sixfold coordinated Ti atom (Ti2-Ad). In addition, TM atoms are only allowed to enter the largest and outermost caves (Cav) because of their large atomic radii. Table 2 shows the impurity formation energies \( E_f \) for ten different 4d TM atom doped models of the TiOS in six different doping manners. We can see only Ag-Cav and Cd-Cav dopings have positive \( E_f \), indicating these two doping processes correspond to endothermic reactions, and are difficult to occur. This should be reasonable since the structures with Ag or Cd atoms doped into the caves are unstable owing to the large atom radii of Ag and Cd. However, all the other doped cases studied here have negative \( E_f \), which indicates the corresponding doping processes are exothermic reactions that can occur more easily. Furthermore, the optimal doping manners for all TM atoms can be readily found according to the lowest \( E_f \), and listed in the following: O1-Ad doping for Ru, O2-Ad doping for Pd and Ag, O3-Ad doping for Y and Rh, Ti1-Ad doping for Tc and Cd, and Cav doping for Zr, Nb, and Mo. Obviously, different TM atoms have different optimal doping manners. In fact, even in the same doping
manner, the impurity atoms may have various positions after being optimized. In particular, Tc, Pd, Ag, and Cd atoms, having stable full or half-full filling outermost orbitals, prefer the O2-Ad or Ti1-Ad doping, while Y, Zr, Nb, Mo, and Rh atoms with unstable outermost orbitals and spin multiplicity, prefer the O3-Ad or Cav doping. The different optimal doping manners are closely dependent on the 4d electron configuration, the atomic radius, and the electronegativity of the TM atom. Thus, the optimal doping manners of all 4d TM atoms have been determined.

Table 2 The formation energies (eV) of TiOS doped with 4d TM atoms.

<table>
<thead>
<tr>
<th></th>
<th>O1-Ad</th>
<th>O2-Ad</th>
<th>O3-Ad</th>
<th>Ti1-Ad</th>
<th>Ti2-Ad</th>
<th>Cav</th>
</tr>
</thead>
<tbody>
<tr>
<td>Zr(4d25s2)</td>
<td>-6.238</td>
<td>-6.544</td>
<td>-6.187</td>
<td>-6.245</td>
<td>-5.684</td>
<td>-7.045</td>
</tr>
<tr>
<td>Nb(4d45s1)</td>
<td>-4.727</td>
<td>-4.725</td>
<td>-4.715</td>
<td>-4.726</td>
<td>-3.377</td>
<td>-2.242</td>
</tr>
<tr>
<td>Rh(4d85s1)</td>
<td>-0.8964</td>
<td>-1.128</td>
<td>-1.364</td>
<td>-0.8950</td>
<td>-1.130</td>
<td>-0.2150</td>
</tr>
<tr>
<td>Pd(4d10)</td>
<td>-1.381</td>
<td>-1.484</td>
<td>-1.434</td>
<td>-1.379</td>
<td>-1.483</td>
<td>-1.014</td>
</tr>
<tr>
<td>Ag(4d105s1)</td>
<td>-0.5412</td>
<td>-0.9686</td>
<td>-0.9660</td>
<td>-0.5548</td>
<td>-0.5511</td>
<td>0.4253</td>
</tr>
<tr>
<td>Cd(4d105s2)</td>
<td>-0.0847</td>
<td>-0.0953</td>
<td>-0.1657</td>
<td>-0.1661</td>
<td>-0.1035</td>
<td>2.376</td>
</tr>
</tbody>
</table>

3.2. Optical absorption

Now we study the optical absorption properties of TiOS doped with 4d TM atoms, as shown in Fig. 2. For comparison, the optical absorption properties of the pure TiOS are also plotted by the dash-dotted curves. The scissors operator, namely, the difference 1.03 eV between the experimental band gap (3.23 eV) and the calculated band gap (2.20 eV) is applied to correct the optical properties [32]. For the pure TiOS, the absorption edge is located at around 392 nm as shown in Fig. 2, in agreement with the experimental observation [37] and the theoretical result [38]. This indicates that the optical response is mainly localized in the UV-light region, implying that the utilization ratio of visible light (approximately 380–780 nm) will be quite low. Therefore, it is necessary to modify the physicochemical properties of TiOS to achieve a better visible-light response. By comparing the solid curve with the dash-dotted one in each panel in Fig. 2, we can find all 4d TM atoms are able to improve the visible-light response of TiOS to different degrees. Totally speaking, their optical absorption coefficients can be classified into two types: (1) Strong improvement case. In Figs. 2a-2e, we can find that high and wide absorption peaks can be formed in the visible-light region, clearly indicating that the visible-light absorption coefficients are significantly increased for Ti24YO48, Ti24ZrO48, Ti24NbO48, Ti24MoO48, and Ti24AgO48. Notably, the Nb doped TiOS shows the best visible-light response. These results imply that the 4d TM dopants with a half-full or less than half-full outer electron configuration can obviously improve the visible-light response of TiOS. (2) Weak improvement case. In Figs. 2f-2j, the visible-light absorption coefficients are only slightly increased for Ti24TcO48, Ti24RuO48, Ti24RhO48, Ti24PdO48, and Ti24CdO48, implying there exists a weak improvement in the visible-light response. Moreover, we can find that the visible-light response intensity will decrease as the atomic number of the dopants increases from Y, Zr, Nb, Mo to Tc, or from Ru, Rh, Pd to Cd. Yet, very surprisingly, Ag with a large atomic number will show unexpected strong visible-light response, different from its neighboring elements. Furthermore, we can find that the 4d TM atom doping induced absorption improvement can appear in a wide area with the wavelength much longer than the visible-light wavelength. This phenomenon is especially outstanding for the strong improvement case in relative to the weak improvement one. Therefore, we can conclude that, among the investigated 4d TM dopings, the TiOS doped with Y, Zr, Nb, Mo, and Ag can remarkably improve the utilization efficiency of visible light.

Fig. 2 (Color online) The optical absorption coefficients of pure (dash-dotted curve) and doped (solid curve) TiOS. The region between two vertical dotted lines represent the visible-light region.
3.3. Density of states

In order to clearly understand the optical absorption properties, Fig. 3 shows the total density of states (TDOS) and partial density of states (PDOS) near the Fermi level of pure and 4d TM atom doped TiOS with spin-up (spin-down) states plotted on the positive (negative) y-axes. In view of the presence of unpaired electrons, it is necessary to take spin into account for the dopings of Y, Zr, Nb, Mo, Tc, Ru, and Rh atoms, while it is not for the dopings of Pd, Ag, and Cd atoms. For the pure TiOS, as shown in Fig. 3a, both the valence band (VB) and the conduction band (CB) are formed by the mixing of O 2p and Ti 3d states. However, the VB is mainly contributed by O 2p states, while the CB is mainly contributed by Ti 3d states. This composition of the VB and CB will not be qualitatively changed by the dopings of 4d TM atoms. However, both VB and CB of all the doped TiOS are seriously shifted to the low energy region in comparison with those of pure TiOS, which should stem from the upward shift of the Fermi energy level. For the dopings of Y, Zr, Nb, Mo, and Ag atoms, the Fermi energy levels move into the CB as shown in Figs. 3b-3f, and they can be viewed as n-type conductors. So we think the dopings can induce the semiconductor-metal phase transition. Therefore, lower energy photons can be absorbed by electron to induce the intra-band transition. Thus, the light absorption response in the entire visible light region or even longer wavelength range may be strengthened, which clearly explains the increase of the optical absorption in Figs. 2a-2e. However, the Fermi energy levels for the dopings of Tc, Ru, Rh, Pd, and Cd atoms seem to stay in the original band gaps in Figs. 3g-3k, implying that only the inter-band transition can occur, sharply different from those of the dopings of Y, Zr, Nb, Mo, and Ag atoms in Figs. 3b-3f. By a close look at Figs. 3g-3k, we can see that the new bands are formed by the coupling of the 4d states of the TM atoms and the O 2p or Ti 3d states. Obviously, this kind of inter-band transition will contribute to the visible-light absorption. Intuitively and reasonably, the relatively small amount of dopants will induce relatively weak light absorption. This is the reason why only the weak improvement can appear in Figs. 2g-2k. Although the doping-induced new bands can also be observed for Y, Zr, Nb, and Mo in Figs. 3b-3e, the Fermi level is deeply immersed into the original CB. In this case it is the intra-band transition in the original CB that dominates the light absorption, which induces the appearing of the strong improvement of the visible-light absorption.

Fig. 3 (Color online) The TDOS and PDOS of pure and 4d TM doped TiOS. For clarity, the Fermi energy level is shown by the vertical dotted lines at zero energy.

3.4. Band structure

In order to explain more intuitively the optical absorption, we further show the energy bands of the 4d TMs doped TiOS in Fig. 4. For pure TiOS in Fig. 4a, the VBM and CBM are located at G point, indicating the nature of the direct band gap. For 4d TM doped TiOS, the VBM and CBM are still located at G point. However, most of dopings except Ag will introduce the impurity energy levels (IELs) in the band gap of pure TiOS. As shown in Figs. 4b-4e, the IELs of Ti_{24}YO_{48}, Ti_{24}ZrO_{48}, Ti_{24}NbO_{48}, and Ti_{24}MoO_{48} are located slightly below the CBM, which can reduce the energy required for the electron transition to the CB and be beneficial for extending the light absorption to visible region and eventually enhancing the visible-light absorption efficiency. Besides, the Fermi energy levels of Ti_{24}YO_{48}, Ti_{24}ZrO_{48}, Ti_{24}NbO_{48}, Ti_{24}MoO_{48}, and Ti_{24}AgO_{48} enter the CB in Figs. 4b-4f, resulting in the intra-band transition, which effectively induces stronger light absorption. For the weak improvement case, the IELs of Ti_{24}TcO_{48} and Ti_{24}RuO_{48} in Figs. 4g and 4h are located near CBM, while the IELs of Ti_{24}RhO_{48} and Ti_{24}PdO_{48} in Figs. 4i and 4j are located near VBM. In the same way, the existence of IELs can reduce the energy required for the electron transition, and hence enhance the optical absorption in the visible-light region.

Fig. 4 (Color online) The band structure of pure and 4d TM doped TiOS. In (b, c, d, e, g, h, i) the solid (dotted) curve represents the spin-up (spin-down) state. The horizontal dashed lines at the energy zero represent the Fermi energy levels.

Next, in Fig. 5 we present the Mulliken charge of the 4d TM doped TiOS. From Fig. 5a we can see that all 4d TM atoms will carry positive charges and there exist equivalent negative charges appearing in TiOS, indicating the electron transferring from 4d TM atoms to TiOS. Clearly Y, Zr, Nb and Mo can carry
much more positive charges than Ru, Rh, Pd and Cd. Large positive charges on doping atoms indicate more electrons on the doping atoms will move from doping atoms to TiOS, further leading to the positive charge centers and making the original CB are filled by electrons. This clearly expounds why the Fermi energy levels are shifted into the CB. Small positive charges on the doping atoms may indicate the electric polarization of the doping atoms and their surroundings. In this case no obvious electron transfer will occur. Also, we present in Fig. 5b-5k the charge density differences of all doped models which vividly depict the redistribution of the electrons in the 4d TM doped TiOS.
Fig. 5 (Color online) (a) Mulliken charge distribution of 4d TM atoms in doped models. (b-k) Charge density difference in 4d TMs doped TiOS with an isovalue of 0.06 e/Å³. Blue (dark) and yellow (grey) areas represent electron accumulation and depletion, respectively.

3.6. Photocatalytic activities

In order to analyze the effect of the 4d TM atom doping on the photocatalytic activity of TiOS, we resort to the effective masses and the transfer rates of the photogenerated electrons and holes. For the pure TiOS, the effective masses are found to be $m_{h^*}(p) = 17.348m_0$ and $m_{e^*}(p) = 2.387m_0$ with $m_0$ being the electron mass, which implies a large $r_{he}(p) = 7.268$. This large difference between $m_{h^*}(p)$ and $m_{e^*}(p)$ will further result in the large difference between the transfer rates of electrons and holes, reducing the recombination of electrons and holes. Therefore, TiOS can be used as a promising photocatalytic material. When TiOS is doped by 4d TM atoms, the effective masses of electrons and holes are denoted by $m_{h^*}(d)$ and $m_{e^*}(d)$ along one certain direction, respectively. We can see in Fig. 6 that the relative variation ratios $r_{h} = m_{h^*}(d)/m_{h^*}(p)$ are all greater than 1, indicating the doping can seriously increase the effective mass of the photogenerated holes. However, the relative variation ratios $r_{e} = m_{e^*}(d)/m_{e^*}(p)$ are all close to 1, indicating only tiny variations of $m_{e^*}$ are observed. Therefore the doping will make stronger influences on the effective mass of the holes than the electrons. For clarity, we also plot the ratio $r'^{he} = r_{he}(d)/r_{he}(p)$ to clearly reflect the overall doping induced variations of the hole and electron effective masses with $r_{he}(d)$ and $r_{he}(p)$ being the effective mass ratios corresponding to the doped and pure TiOS. We can see all $r'^{he}$ are greater than 1, indicating all the dopings can enlarge the effective mass difference between holes and electrons. This phenomenon is more evident for Y, Zr, Nb, Mo, and Ag than Tc, Ru, Rh, Pd, and Cd. Furthermore, this demonstrates that the 4d TM atom doping will further suppress the recombination of the photogenerated electrons and holes, which can improve the photocatalytic activity of TiOS in different degrees. Moreover, the dopings of Y and
Ag can lead to larger $r'_{he}$, indicating both Y and Ag dopings can be viewed as the better schemes used to enhance photocatalytic activity of TiOS.

**Fig. 6** (Color online) The relative variation ratios $r_{bh}$, $r_e$, and $r'_{he}$ for different doping models.

Furthermore, as discussed before, based on the energy bands of the 4d TM atom doped TiOS in Fig. 4 we can see that the couplings of the doping atoms and the TiOS can produce new impurity energy bands in the original band gap and fill the original CB with electrons. The appearing of the impurity energy bands will provide new channels through which the electrons in impurity bands can be pumped to the CB by absorbing the visible-light photons. Therefore, this procedure can strengthen the visible-light induced photocatalytic activities of TiOS. Moreover, the visible-light absorption will also pump the electron below Fermi energy level in the CB into the empty energy levels above Fermi energy level, inducing more electron to appear at high energy levels. This also increases the photocatalytic activities of the TiOS. Thus we can draw that the 4d TM atom dopings are beneficial for increasing the visible-light utilization efficiency and improving the photocatalytic activities of TiOS.

**4. Conclusion**

In summary, we have systematically studied the TiOS doped by all 4d TM atoms. New impurity energy bands can be generated in the bandgap and the semiconductor-metal phase transition can be induced by dopings. The visible-light absorption can be seriously increased by doping Y, Zr, Nb, Mo, and Ag, and only weakly increased by doping other 4d TM atoms. The former should be caused by both interband and intraband transitions of electrons while the latter should be only the interband transition. Furthermore, we discussed the doping-induced improvement of the photocatalytic activities of TiOS. It is verified that among all 4d TM atoms both Y and Ag dopings can be used to effectively improve the visible-light photocatalytic activities of the TiOS.

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