RESEARCH ARTICLE

TiO₂ Band Restructuring by B and P Dopants

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Abstract

An examination of the effect of B- and P-doping and codoping on the electronic structure of anatase TiO₂ by performing density functional theory calculations revealed the following: (i) B- or P-doping effects are similar to atomic undercoordination effects on local bond relaxation and core electron entrapment; (ii) the locally entrapped charge adds impurity levels within the band gap that could enhance the utilization of TiO₂ to absorb visible light and prolong the carrier lifetime; (iii) the core electron entrapment polarizes nonbonding electrons in the upper edges of the valence and conduction bands, which reduces not only the work function but also the band gap; and (iv) work function reduction enhances the reactivity of the carriers and band gap reduction promotes visible-light absorption. These observations may shed light on effective catalyst design and synthesis.

Introduction

Because of its high chemical stability, high photocatalytic activity, and non-toxicity, titanium dioxide has been considered one of the more promising photocatalysts for photoelectrochemical water splitting, dye-sensitized solar absorption, degradation of pollutants, and fuel conversion. However, the undesirably fast rate of carrier recombination and the large band gap (3.0 eV for the rutile phase, 3.2 eV for the anatase phase) for photon absorption limits its practical applications. In order to improve the catalytic efficiency of TiO₂, one needs to consider the following three parameters: (i) how to narrow the band gap from 3.2 eV to match the energy of high-intensity visible light, (ii) how to increase the electroaffinity of the specimen to prolong the lifetime of the carriers, and (iii) how to decrease the work function to improve the reactivity of the carriers. The electroaffinity is the separation between the bottom of the conduction band and the vacuum level, which describes the ability of the specimen to hold electrons captured during reaction or to entrap the conduction electrons that localize and polarize nonbonding electrons [1]. In the process of photocatalytic, when the reduction and oxidation do not proceed simultaneously, there is an electron accumulation in the conduction band, thereby causing a fast recombination of e-h pairs. Therefore, improve the utilization rate of sunlight by modulating band gap and enhance electrons lifetime by reducing the work function and increasing the
electroaffinity via locally pinning the polarized electrons are essential to promote photocatalytic reaction.

We have recently shown [2] that atomic undercoordination, such as a defect or a terrace edge formation, and nanocrystallization could improve the three quantities in the previous paragraph due to the bond order-length-strength correlation and nonbonding electron polarization (BOLS-NEP) mechanism [1]. The BOLS-NEP notation indicates that a loss of bond order shortens and stiffens the bonds between undercoordinated atoms [2, 3]. This event leads to the local densification and quantum entrapment of the bonding electrons to deepen the bottom edge of the energy bands, which polarizes the nonbonding electrons to shift up the upper edge of the bands. The conduction band edge depression increases the electroaffinity, which prolongs the carrier life. The polarization decreases the work function. On the other hand, atomic heterocoordination, such as what occurs with foreign-ion doping or codoping [4–10], surface sensitization by metal complexes or organic dyes [11, 12], noble metal loading [13, 14], and semiconductor coupling [15, 16], also exhibit the BOLS-NEP effect but to different extents.

Opinions about the B- and P-doping effects on the photoactivity of TiO₂ remain controversial though studies have been frequent. Chen et al. [17] observed that B-doping widens the band gap and attributed this observation to the quantum size effect. Zhao et al. [18] explored a redshift in the absorption spectrum of B-doped TiO₂ and explained the redshift as arising from crystal geometry modification of the electronic structures. In fact, B dopants can substitute for either O or Ti atoms, or can even sit in the interstitial sites. Density functional theory (DFT) calculations [19–21] suggest that B substitution for Ti atoms is energetically least favorable, while interstitial substitution and substitution for O are energetically comparable, so they can both occur in real situations. Patel et al. [21] suggested that at low concentration B preferentially occupies interstitial positions, but with increasing concentration B occupies lattice sites by replacing O.

Yang et al. [22] reported that P substitution for O narrows the band gap slightly by introducing impurity P 3p levels in the bandgap which could induce absorption edge redshift. Impurity level creation not only decreases the band gap but also provides centers promoting carrier recombination. Band gap reduction enhances visible light absorption but shortens carrier life by recombining carriers. The competition of these two effects determines the catalytic performance of TiO₂. However, P replacement of Ti produces no impurity energy levels though it narrows the band gap of the anatase TiO₂ slightly which is ineffective for photocatalytic improvement.

Recent progress indicates that codoping with B and P or with other nonmetals [8, 23–27] enhances UV-Vis light absorption when compared with monodoped TiO₂ due to “p-n pairs” [8] or the “synergistic effect” [25]. However, the effects of doping and codoping with B and P on the band structure of TiO₂ remain unclear. Here we systematically examine the effect of B- and P-doping and codoping on the electronic structures and optical properties of the anatase phase of TiO₂ using DFT calculations.

**Materials and Methods**

The present work was done using the spin-polarized DFT model of the CASTEP package [28]. The electron-ionic core interaction was represented using the ultra-soft pseudo-potential and the electron-electron interactions were described by the generalized gradient approximation (GGA) via the PW91 functional [29]. The structural optimizations were calculated using the plane-wave basis cutoff energy of 400 eV.

After structure optimization, we calculated the electronic structures and the optical properties of B- and P-doped and B/P-codoped TiO₂. However, traditional DFT usually cannot
accurately describe the band gap of semiconductors. We considered that an on-site correction can get a band gap match more in line with experimental results as well as provide a more economical calculation than a hybrid functional correction, so the GGA+U method was an efficient way to study the electronic and optical properties [30–32]. It was shown that when $U = 4.25$ eV is applied to the Ti 3d electrons, the band gap of pure anatase TiO$_2$ is 3.20 eV, which corresponds well with experimental results. This $U$ value has proven to be reasonable for dealing with impurity states in the bandgap [31] and in redox catalysis by TiO$_2$ [33].

### Results and Discussion

#### B- or P-Doping

We replaced one O atom with a B or P atom in the 48-atom $2 \times 2 \times 1$, 24-atom $2 \times 1 \times 1$, and 12-atom $1 \times 1 \times 1$ supercells of anatase TiO$_2$ shown in Fig 1 and denoted as (a) and (b) to simulate doping concentrations of 8.3% and 4.2%, respectively. Pure TiO$_2$ was used as a reference. The Monkhorst-Pack mesh was set as $5 \times 5 \times 4$, $10 \times 10 \times 4$, and $10 \times 10 \times 4$ for these supercells. Optimized structural parameters of $a$, $c$, and $d$ ($d_{eq}$ and $d_{ap}$ are the equatorial and apical Ti–O bond lengths, respectively) for the primitive unit cell of anatase TiO$_2$ (space group I4$_1$/amd) are shown in Table 1 which fits well with experimental observations [34] and HSE06 hybrid functional calculation results [35].

![Fig 1: B(P) replacement of an oxygen atom in the (a) 1 × 1 × 1 and (b) 2 × 1 × 1 anatase TiO$_2$ supercells, the corresponding doping level is 8.3% and 4.2%, respectively. The $d_{eq}$ and $d_{ap}$ denote the apical and the equatorial bond lengths, respectively.](image)

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| Exp. [34] | HSE06 [35] | This work |
|-----------|------------|-----------|
| $a$/Å     | 3.78       | 3.80      | 3.78      |
| $c$/Å     | 9.50       | 9.47      | 9.49      |
| $d_{eq}$/Å| 1.93       | 1.94      | 1.95      |
| $d_{ap}$/Å| 1.98       | 1.98      | 2.00      |
| $c/a$     | 2.51       | 2.49      | 2.51      |

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For comparison, the defect formation energy ($E_{\text{form}}$) can be calculated using the following formula

$$E_{\text{form}} = E_{\text{tot}(\text{doped})} - E_{\text{tot}(\text{pure})} + \mu_x - \mu_O$$  \hspace{1cm} (1)

where $\mu_x$ represents the chemical potential of $x$ ($x = \text{B, P, O, and Ti}$) and $E_{\text{tot}(\text{pure})}$ and

| Concentration (%) | Formation energy (eV) | Bond length (Å) | Mulliken population | Band gap (eV) |
|-------------------|-----------------------|-----------------|---------------------|--------------|
|                   | Ti-rich | O-rich | Ti–O | B–Ti | B–O | Ti | O | B |      |
| 0                 | -       | -      | 1.979/1.932(2) | - | - | 1.48/1.34[22] | -0.74/-0.67[22] | - | 3.2 |
| 2.1               | 7.09    | 10.69  | -     | 1.371(2) | 1.36(2) | -0.72(2) | 0.27 | 2.84 |
| 4.2               | 10.89   | 14.49  | 2.352/2.074(2) | 1.34(2) | 1.41 | -0.26 | 2.64 |
| 8.3               | 11.31   | 14.91  | 2.170/1.964(2) | 1.21(2) | 1.38 | -0.37 | 0 |

B-Doping

For comparison, the defect formation energy ($E_{\text{form}}$) can be calculated using the following formula

$$E_{\text{form}} = E_{\text{tot}(\text{doped})} - E_{\text{tot}(\text{pure})} + \mu_x - \mu_O$$  \hspace{1cm} (1)

where $\mu_x$ represents the chemical potential of $x$ ($x = \text{B, P, O, and Ti}$) and $E_{\text{tot}(\text{pure})}$ and

Fig 2. Partial geometries of the B-doped TiO$_2$ at (a) 2.1%, (b) 4.2%, (c) 8.3% doping concentrations.

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$E_{\text{tot}}(\text{doped})$ are the total energies of pure and doped TiO$_2$, respectively. It is noteworthy that $E_{\text{form}}$ is not fixed but that it changes with different growing conditions, such as Ti-rich, O-rich, or mixed conditions. The B chemical potential was given by the equation $\mu_B = \frac{1}{2}E(B_2O_3) - \frac{3}{4}E(O_2)$. Under Ti-rich conditions, the Ti chemical potential is assessed from the energy of bulk Ti while the O chemical potential was obtained from the following equation:

$$\mu_{\text{Ti}} + 2\mu_{\text{O}} = \mu(\text{TiO}_2)$$  \hspace{1cm} (2)

Under O-rich conditions, the $\mu_{\text{O}}$ value is taken from the chemical potential of the free O$_2$ molecule, while the chemical potential of Ti is fixed by the conditions set by Eq 2. It is well known that the smaller the $E_{\text{form}}$ value, the greater the stability of the doped TiO$_2$ supercell. The formation energies for various concentrations of B-doped TiO$_2$ are reported in Table 2. It is noted that for all concentrations of B doped into TiO$_2$, substitution of the lattice O is more
reasonable under the Ti-rich growth conditions and B-doping is increasingly difficult as the concentration increases.

The partial structures derived from the optimized B-doped TiO$_2$ with different doping concentrations are given in Fig 2. It is shown that high concentration (8.3% and 4.2%) doping does not distort the optimized crystal structure with elongation of the Ti–B bond. In the $d_{eq}$ direction, the distances between B and Ti atoms stretch from the initial 1.932 Å to 1.964 and 2.074 Å, meanwhile, in the $d_{ap}$ direction, the distances elongate from the initial 1.979 Å to 2.170 and 2.352 Å for 8.3% and 4.2% B-doping, respectively.

When the doping concentration is low (2.1%), the B atom deviates from its original position to instead bind with two neighboring O atoms to form a Ti–O–B bond in accordance with XPS measurements [21, 36]. Both B–O bonds are 1.371 Å. Furthermore, the Mulliken population study shows that the charges on the B atoms are about −0.37 and −0.26 for the 8.3% and 4.2% doping, respectively (see Table 2). The charge density is mainly distributed on the B atom rather than on the neighboring Ti and O atoms. At the 2.1% doping level, B atoms have a positive charge of 0.27 and form a Ti–O–B structure after the transfer of electrons from B to the adjacent O atoms.

The band structures of pure and B-doped anatase TiO$_2$ are shown in Fig 3. The conduction band minimum (CBM) and the valance band maximum (VBM) of pure TiO$_2$ appear on the gamma (G) and M points, respectively. The DFT+U calculations confirm an indirect band gap.

### Table 3. Formation energy, Mulliken populations on the P atom and the Ti atoms bonded to the P atom, and bond lengths of the P–Ti bonds for different doping concentrations.

| Concentration (%) | Formation energy (eV) | Bond length (Å) | Mulliken population | Band gap (eV) |
|-------------------|-----------------------|-----------------|---------------------|--------------|
|                   | Ti-rich | O-rich | Ti–O | P–Ti | Ti | O | P |             |
| 0                 | - | - | 2.003/1.945(2) | - | 1.48(1.34[22]) | −0.74/(−0.67[22]) | - | 3.2 |
| 2.1               | 11.47 | 15.07 | 2.264/2.362/2.412 | 1.22/1.26/1.32 | -0.17 | 3.12 |
| 4.2               | 12.86 | 16.46 | 2.235(2)/2.367 | 1.24(2)/1.36 | −0.17 | 2.46 |
| 8.3               | 13.85 | 17.45 | 2.222(2)/2.318 | 1.09(2)/1.30 | −0.26 | 0.75 |

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Fig 3B–3D shows that the band gap decreases gradually with an increase in the doping concentration. The band gap approaches zero at the 8.3% doping level. Fig 4 shows the projected density of states (PDOS) near the dopants and the total density of states (DOS) for B-doped TiO₂. The PDOS plots indicate that the O 2p states form most of the valence band while the conduction band is mainly composed of the Ti 3d states in B-doped TiO₂, which is consistent with pure TiO₂. B doped into TiO₂ does not produce a simple superposition of valence electrons but the process instead exchanges and polarizes electrons. First, some impurity energy levels emerge in the bandgap and they are mainly composed of B 2p electrons and neighboring Ti 3d and O 2p electrons. Second, the strong delocalization and bonding characteristics are formed by B 2s and O 2p electrons. Third, the valence and conduction bands gradually broaden with increased concentration of B-doping.

P-Doping
Substituting O sites with P in the TiO₂ was also examined for the same set of anatase TiO₂ supercells. The formation energies for P-doped anatase TiO₂ were also calculated according to Eqs (1) and (2), in which the chemical potential of phosphorus was obtained from the equation \( \mu_P = 1/2E(P_2O_5) - 5/4E(O_2) \). Table 3 lists the optimal structural parameters. For all concentrations of P
doped into TiO₂, substitution of the lattice O is reasonable under the Ti-rich growth conditions and P-doping becomes increasingly difficult as the concentration increases.

**Fig 5** shows the local geometries taken from different P-doping concentrations in the anatase TiO₂ supercells. This case is similar to B-doping because the distances between the P atom and adjacent Ti atoms shorten with increasing concentration, while an asymmetric structure deformation occurred for 2.1% P-doping (d<sub>eq</sub> = 2.222 Å, d<sub>ap</sub> = 2.318 Å for 8.3% doping level; d<sub>eq</sub> = 2.235 Å, d<sub>ap</sub> = 2.367 Å for 4.2% doping level; d<sub>eq</sub> = 2.264 Å and 2.362 Å, d<sub>ap</sub> = 2.412 Å for 2.1% doping level). These differences for both the B- and P-doped structures are larger than the original Ti–O bond length due to the larger B and P atomic radii. Moreover, considering the lower electronegativity of P than B and O, the charge on P in P-doped anatase TiO₂ is more mobile than on B and O due to weaker ionic interactions between the P anion and adjacent Ti cation, and thus fewer electrons are transferred from Ti to P (see Table 3).

**Fig 6** shows that P-doping reduces the band gap to 3.12 eV for the 2.1% doping level, 2.46 eV for the 4.2% doping level, and 0.75 eV for the 8.3% doping concentrations. **Fig 7** shows the PDOS and DOS due to different concentrations of dopant. Characteristics of the B-doped TiO₂ DOS
also similarly appear in the P-doped TiO$_2$. P doped into TiO$_2$ also exchanges and polarizes valence electrons of pure TiO$_2$. First, some impurity energy levels emerge in the bandgap that are mainly composed of the P 3p electrons and neighboring Ti 3d and O 2p electrons. Second, the strong delocalization and bonding characteristics are formed by P 3p and O 2p electrons. Third, the valence and conduction bands gradually broaden with increased concentration of P-doping.

In Fig 8, zone-selective photoelectron spectroscopy (ZPS) technology [37] is used to determine the changes in the electronic structure for different concentrations of B and P doped into TiO$_2$. The results clarified that additional DOS features are formed in all B- and P-doped TiO$_2$. Entrapped states within the band gap as impurity states enhance the utilization of visible light.

**Fig 7.** PDOS and normalized DOS for P-doped anatase TiO$_2$. The energy is measured from the top of the valence band of pure anatase TiO$_2$. The gray dotted line represents the actual Fermi level. PDOS is the average density of states of each atom near the defects.

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**Fig 8.** The residual DOS for B- (a) and P- (b) doped TiO$_2$ at 2.1%, 4.2%, and 8.3% doping concentrations.

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Fig 9. (a) Optimized geometries for the B- and P-codoped TiO$_2$, and (b) corresponding difference electron density maps for the plane with B and P atoms.

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Fig 10. Calculated imaginary part of the dielectric function ($\varepsilon_2$) for pure, monodoped, and codoped TiO$_2$. The region between the horizontal dashed lines marks the visible light region.

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and the antibonding states polarize the Ti 3d electrons by entrapped states, thus decreasing the work function contributing to catalytic ability, which have been proven in previous reports [38].

B- and P-Codoping

To investigate the stabilities of B- and P-codoped systems, we studied the $E_{\text{form}}$ for two O atoms: one replaced by a B atom and one replaced by a P atom in the 48-atom (2 × 2 × 1) supercell. Note that the increasing distance between the B and P atoms in both Ti-rich and O-rich growing conditions leads to an increase in $E_{\text{form}}$. B and P atoms substituting the nearest neighboring O atoms are energetically more favorable. Fig 9 shows the optimized structure and the electron density distribution. The B and P atoms form a new B–P bond of length $d = 1.775$ Å. This bond is shorter than the sum of the atomic radii: 0.95 (B) + 1.30 (P) = 2.25 Å, which means a stronger B–P single bond is formed as compared to the original O–O separation in pure TiO$_2$ of 2.474 Å. The difference charge map also shows that B gains electron from P.

To further validate the influence of doping on light absorption, we compared the imaginary part of the dielectric function ($\varepsilon_2$), which is an important parameter for photoresponse testing (Fig 10). The calculated $\varepsilon_2$ for pure TiO$_2$ is in agreement with a recent experiment [39]. It is obvious that all B-, P-, and B/P-codoped TiO$_2$ can achieve absorption of visible light and the absorption edges decrease relatively from pure TiO$_2$ by 1.89, 1.32, and 1.15 eV, respectively.

The band structures of B-doped, P-doped, and B/P-codoped are compared in Fig 11. Because of the synergistic effect of B and P, unoccupied energy levels at the bottom of the CBM are quenched which act as a recombination center in B-doped and P-doped TiO$_2$. Moreover, it is worth noting that the CBM and the VBM determine the reducing and oxidizing ability. As the VBM gets lower compared with the hydrogen production level, the oxidizing ability increases. Similarly, as the CBM gets higher, the reducing ability increases. Since the CBM of pure TiO$_2$ is slightly higher than the hydrogen production level, it is best to either keep the position of the CBM unchanged or raise it in the process of band structure modulation. Because B and P passivated codoping can decrease the band gap, as well as avoid changes to the CBM compared with the B-doped and P-doped TiO$_2$, codoping produces a suitable visible-light absorption region and doesn’t handicap the reducing power of TiO$_2$. 

![Fig 11. Band structures of (a) B-, (b) P-, and (c) B/P-codoped. The energy is measured from the top of the valence band of pure anatase TiO$_2$. The gray dotted line represents the actual Fermi level.](doi:10.1371/journal.pone.0152726.g011)
Conclusions
In summary, we investigated the effects of B-doping, P-doping, and B/P-codoping on the electronic structures and optical properties of anatase TiO₂ by employing DFT calculations. Non-metal doping, such as that with B and P, does not produce a simple superposition of valence electrons, but instead the process exchanges and polarizes electrons compared to pure TiO₂. Localized entrapped states in the band gap can absorb visible light and the polarization states induced by entrapped states decrease the work function to improve the reactivity of the carriers. Moreover, B and P passivated codoping quenches the unoccupied energy levels to prolong the carrier lifetime and cause less perturbation to the CBM, thus resulting in a suitable visible-light absorption region that does not handicap the reducing power of TiO₂.

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Author Contributions
Conceived and designed the experiments: LL FLM HWT WTZ CQS. Performed the experiments: LL LQ XYH. Analyzed the data: LL HWT LQ XYH WTZ CQS. Wrote the paper: LL CQS.

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