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Modification of TiO₂ Nanoparticles with Organodiboron Molecules Inducing Stable Surface Ti³⁺ Complex



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HIGHLIGHTS Organodiborons are used

to reshape the surface electronic state of semiconductor oxides

Diboron adsorption leads to spontaneous charge transfer and reduced surface metal ions

Photodetector based on diboron material affords 10³ fold higher visible light response

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Modification of TiO₂ Nanoparticles with Organodiboron Molecules Inducing Stable Surface Ti³⁺ Complex

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SUMMARY

As one of the most promising semiconductor oxide materials, titanium dioxide (TiO₂) absorbs UV light but not visible light. To address this limitation, the introduction of Ti³⁺ defects represents a common strategy to render TiO₂ visible-light responsive. Unfortunately, current hurdles in Ti³⁺ generation technologies impeded the widespread application of Ti³⁺ modified materials. Herein, we demonstrate a simple and mechanistically distinct approach to generating abundant surface-Ti³⁺ sites without leaving behind oxygen vacancy and sacrificing one-off electron donors. In particular, upon adsorption of organodiboron reagents onto TiO₂ nanoparticles, spontaneous electron injection from the diboron-bound O²⁻ site to adjacent Ti⁴⁺ site leads to an extremely stable blue surface Ti³⁺–O⁻⁻⁻ complex. Notably, this defect generation protocol is also applicable to other semiconductor oxides including ZnO, SnO₂, Nb₂O₅, and In₂O₃. Furthermore, the as-prepared photoelectronic device using this strategy affords 10³-fold higher visible light response and the fabricated perovskite solar cell shows an enhanced performance.

INTRODUCTION

Owing to its abundance, nontoxicity, and stability, semiconductor oxide (TiO₂, ZnO, SnO₂, etc.) nanoparticles and films have been widely used as wide-band-gap semiconductor photocatalysts for a variety of solar-driven clean energy and environmental technologies, such as photovoltaics and photocatalytic fuel generation (Chen and Mao, 2007; Grätzel, 2001). However, pristine wide band semiconductor oxides is not an appropriate candidate for practical applications since it only adsorbs UV light. The optical response of TiO₂ nanocrystal has been tuned to visible region using band gap engineering techniques, including metallic (Dahl et al., 2014; Hoffmann et al., 1995) or nonmetallic (Asahi et al., 2001, 2014; Chen and Burda, 2008; Khan et al., 2002) impurity doping, solid solution formation (Maeda et al., 2006; Wang et al., 2008), and self-structural modification (Liu and Chen, 2014). Among various self-modification techniques, in situ formation of self-doped Ti³⁺ in the bulk phase through the introduction of oxygen vacancy at high temperature is an effective strategy for band gap engineering (Figure 1A) (Zuo et al., 2010). Additionally, hydrogenation of TiO₂ nanocrystals can also result in visible-light-responsible materials (Chen et al., 2011). The latest studies show that these low-energy absorption materials arouse utilities in visible light water splitting and microwave and terahertz absorption (Green et al., 2018a, 2018b, 2019a, 2019b; Guan and Chen, 2018; Tian et al., 2017). Unfortunately, harsh reaction conditions and long-time treatment have limited the practicality of these methods, especially in the context of on-site fabrication and reprocessing of light harvesting devices. Alternatively, UV irradiation of TiO₂ nanoparticles (Schrauben et al., 2012) or visible light irradiation of dye-sensitized systems can also produce visible-light-responsive blue-surface TiO₂ (Figure 1B) (Yan et al., 2017). In both cases, organic sacrificial agents are required, and these agents are irreversibly oxidized by photoinduced holes (e.g., alcohol to aldehyde and I^- to I_2). Moreover, such nascent optically active Ti^{3+} center is highly unstable toward O_2 owing to instantaneous oxidation to Ti⁴⁺. To our knowledge, the formation of air-stable surface Ti³⁺ centers using these techniques has not been previously described. We surmise that this is largely due to the difficulty in stabilizing the key Ti³⁺- O^{2-} molecy formed in these processes, as such molecy is highly reactive toward O_2 to afford Ti⁴⁺-O- O^{-1} . In light of these constraints, the ability to access air-stable surface Ti³⁺ defects within TiO₂ would represent a paradigm shift for the field of TiO_2 -based oxide semiconducting materials. In this report, we show that this goal can be accomplished through the simultaneous modification of surface O^{2-} and Ti^{4+} sites in TiO₂ to afford a persistent, optically active $Ti^{3+}-O^{--}$ species.

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Continued



Organodiboron compounds constitute a class of stable and highly versatile reagents commonly used in organic synthesis (Neeve et al., 2016). Despite their unique reactivity, to date, the use of these diboron species in the modification of inorganic materials remains surprisingly scarce. Herein, we report a new method for the facile generation of Ti^{3+} defects on TiO_2 surface under mild conditions (<80°C) enabled by the use of diboron reagents (Figure 1C). In this process, facilitated by the intimate interaction of the diboron reagent's B center with the surface bridging O_{2c} of TiO_2 , the adsorption of organodiboron reagents onto TiO_2 nanoparticles leads to spontaneous electron transfer, resulting in a stable, blue surface Ti^{3+} -TiO₂. Notably, this process does not require complicated synthetic manipulations such as anaerobic photo irradiation or high temperature calcination, which are commonly employed by previously developed techniques.

Previously developed strategies for TiO₂ surface modification have mainly focused on the modification of surface Ti sites with organic nucleophiles (e.g., phenol, alcohol, and carboxylic acids). In contrast, surface bridging oxygen (O_{2c}) sites have seldom been functionalized with electrophilic organic reagents. We posited that electrophilic organic adsorbates with an appropriate reduction potential might favorably interact with the surface O_{2c} sites. Importantly, this binding event might overcome the barrier of electron transfer from O^{2-} sites to adjacent Ti⁴⁺ sites and further stabilize the primary charge separation state. We were particularly interested in the use of organic diboron reagents because of their unique Lewis acidity and reducing ability (*vide supra*). Previous work in the area of synthetic organic chemistry showed that, upon binding to a Lewis basic oxygen atom, these organic diboron species can function as single electron reducing agents, thus allowing for various important transformations (Liu et al., 2019; Mo et al., 2010, 2018; Pietsch et al., 2015; Wang et al., 2016; Zhang and Jiao, 2017). Based on these reasons, we envisioned that, upon the coordination of such diboron compounds with the surface oxygen atom in metal oxide materials, the formation of surface diboron-oxygen Lewis pair may induce single electron transfer from the *ipso*-O_{2c} site to the adjacent Ti site. Furthermore, the interaction of diboron species with the oxygen atom may stabilize the resulting low-valent Ti–high-valent O pair.

RESULTS AND DISCUSSION

Spectroscopic Characterizations

We use P25 TiO₂ to study the surface modification behavior, for P25 TiO₂ is typical TiO₂ material applied in photochemical applications and shows enhanced performance based on fine nanoparticles and heterointerface between anatase and rutile phases (Xia et al., 2013, 2014). Mechanical mixing of commercial Degussa TiO₂ P25 and bis(pinacolato)diboron (B₂Pin₂, B1) in a nitrogen-filled glovebox at room temperature resulted in a rapid color change from white to blue within 1 min (Figure 1E and Video S1). This blue titania, labeled as B1-TiO₂-N₂, was highly sensitive to O_2 and faded immediately upon exposure to air. Unexpectedly, heating B1-TiO₂-N₂ at 80°C under vacuum for 3 h resulted in a blue sample that is stable under air, which we labeled as B1-TiO2. This sample could be stored outside the glovebox for weeks, and the blue color persisted. Similar transformations were observed with three other diboron compounds as shown in Figures 1 and S1. To fully disperse the diboron compound onto the TiO₂ nanoparticles, this heterogeneous reaction was performed in a diboron-soluble solvent, such as methanol or diethyl ether. Same blue samples were obtained after removal of organic solvents under vacuum. On the basis of previous studies (lorio et al., 2012), the blue color was indicative of the formation of Ti³⁺ defects. To probe whether B1 was oxidized in this process, the newly prepared blue B1-TiO₂ sample was extracted with CDCl₃, and B1 was found to be the only boron-containing species present in the extract (Figure S2). Additionally, the use of this recovered B1 from the extract for TiO₂ modification also gave rise to the same blue titania. Thus, these experiments suggested that diboron compound B1 is not a one-off sacrifice agent. Based on these findings, a new mechanism must be responsible for the formation of this blue TiO₂.

To probe the existence of Ti^{3+} species, electron paramagnetic resonance (EPR) spectra (Zuo et al., 2010) of the **B1**-TiO₂ sample were acquired. Low-temperature (2 and 100 K) EPR data feature transitions with g = 1.97-1.99, whereas the room temperature EPR is silent (Figure 1F). This behavior is characteristic of Ti^{3+} species as documented by previous studies (Li et al., 2008). The observed g values show the presence of a strong anisotropic paramagnetic Ti^{3+} center, indicating that the local symmetry of Ti^{3+} is vastly broken. Furthermore, this Ti^{3+} EPR signal would be split if B replaces the bridging O or is directly bound to Ti (Gopal et al., 2008). However, in all the cw-EPR measurements, we did not observe the hyperfine coupling originating from the ¹⁰B or ¹¹B nucleus. This suggests either no or very weak coupling between the B atom

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Figure 1. Conventional Methods and Present Work for the Preparation of Self-Doped Ti³⁺-TiO₂ Materials

(A) Ti^{3+} generation as a result of oxygen vacancy formation. High-temperature combustion with reductant leads to oxygen vacancy in TiO_2 bulk phase (Zuo et al., 2010).

(B) Ti^{3+} generation by photo irradiation. By removal of photoinduced holes using organic sacrificial agents, electrons are cumulated at the conduction band of TiO_2 , resulting in the reduction of Ti^{4+} to Ti^{3+} (Schrauben et al., 2012; Yan et al., 2017).

(C) Present work. Diboron compound (B₂Pin₂) adsorption on the surface of TiO₂ nanoparticles induces spontaneous electron injection for the formation of optically active $Ti^{3+}-O^{--}$ species. The light blue, blue, red, orange, earth yellow, green, and pink balls stand for the Ti^{4+} , Ti^{3+} , O^{2-} , O^{--} , C, B, and H atom, respectively.

(D) Four commercially available diboron compounds used in this study: B_2Pin_2 (bis(pinacolato)diboron), $B_2(OH)_4$

 $(tetrahydroxydiborane), B_2Cat_2 \ (bis(catecholato)diboron), \ and \ B_2(NMe_2)_4 \ (tetrakis(dimethylamino)diboron).$

(E) Photograph comparing TiO₂ (P25), B_2Pin_2 (B1), B1-TiO₂-N₂ sample under an inert atmosphere (mixture of solid TiO₂ and B_2Pin_2) and B1-TiO₂ under air after heating at 80°C under vacuum for 3 h.

(F) EPR spectra of $\textbf{B1}\text{-}\text{TiO}_2$ at various temperatures.

(G) HRTEM of **B3**-TiO₂. A short-dashed curve is applied to outline a portion of the interface between the crystalline core and the outer layer (marked by white arrows).

(H) The corresponding HADDF-STEM image of B3-TiO₂ with distribution of the C element mapping.

and the Ti³⁺ center. Moreover, we found that **B3**-TiO₂ shows a *g* signal at 2.003 at room temperature (Figure S3), which could be characteristic of a Ti³⁺-O⁻⁻ radical. We also applied the ACTEM method to study the existence and status of diboron molecule on the surface of TiO₂. In the high-resolution transmission electron microscopy (HRTEM) image of **B3**-TiO₂ (Figure 1G), we found that the surface of TiO₂ nanocrystal is wrapped by an outer layer with ~1-nm thickness. In addition, the EDS mapping of C element under the HADDF model (Figure 1H) shows a carbon-rich surface shell representing adsorbed organic diboron molecule on the TiO₂ surface.



Figure 2. Spectroscopic Characterizations

(A) UV-vis spectra of the white TiO_2 P25 and diboron-adsorbed TiO_2 P25. The inset shows the UV-vis spectra of four free diboron compounds.

(B) In situ ATR-FTIR spectrum of B1 onto TiO_2 over 15 min. (C) FTIR spectra of free B1 and $B1\text{-TiO}_2.$

Figure 2. Continued

(D) Solid-state ¹¹B MAS NMR of diboron compounds B and B-TiO₂.

(E) XRD pattern of the diboron compounds modified TiO₂.

(F) Ti 2p, O 1s, and valence band structure X-ray photoelectron spectroscopy spectra of TiO₂ and B1-TiO₂.

(G) Normalized Ti-L edge XANES spectra of TiO $_2$ and B1-TiO $_2$.

(H) Normalized O-K edge XANES spectra of TiO_2 and B1- TiO_2 .

We next investigated the structures and properties of these diboron-TiO₂ nanoparticles with powder X-ray diffraction (PXRD), solid-state UV-vis spectroscopy (UV-vis), in situ attenuated total reflectance-Fourier transform infrared spectroscopy (in situ ATR-FTIR) and solid-state ¹¹B magic angle spinning nuclear magnetic resonance (¹¹B MAS NMR). PXRD analysis (Figure S4) shows that the commercial P25 sample is a mixture of anatase-phase and rutile-phase TiO2. The PXRD pattern of TiO2 remained the same upon adsorption of the diboron reagent, indicating that no phase change occurred in this process. Thus, this result suggests that the blue species is likely related to surface engineering rather than bulk phase modification. The solid-state UV-vis spectra in Figure 2A showed that neither the pristine white TiO₂ nor the diboron compound absorbs visible light (>400 nm). First, absorption bands beginning at 400 nm and steadily growing into the near-infrared (NIR) region were observed for all four B-TiO₂ samples. These adsorption bands were different from that arising from metal to ligand charge transfer of the surface complex between Ti⁴⁺ sites and organic ligands commonly observed in previous work (Lang et al., 2012). The UV-vis absorption spectra associated with Ti³⁺ materials can differ between different preparation methods (Howe and Gratzel, 1985). In our case, the diboron-modified TiO₂ samples were found to absorb visible light with higher intensity in the NIR region. This suggested that the Ti³⁺ species in our case is different from the previously reported Ti³⁺-O²⁻ structures. In the *in situ* ATR-FTIR spectra (Figure 2B), the emergence of new signals corresponding to the diboron compound was observed over the time, clearly demonstrating the adsorption process. In the FTIR spectra (Figure 2C), signals at 1280 and 1170 cm^{-1} are assigned to two different types of B-O vibration. Based on density functional theory (DFT) calculations, the B-B stretching vibration appears at ca. 1000 cm⁻¹ (Figure S5). As shown in Figure 2C, this signal is absent in **B1** owing to the C_2 symmetry of this diboron molecule. In contrast, a new peak at 1010 cm⁻¹ corresponding to B–B stretching emerged in the B1-TiO₂ sample, thereby demonstrating the formation of the oxygen-diboron Lewis pair.

Solid-state ¹¹B MAS NMR spectroscopy was next used to elucidate the binding details of the diboron reagent with the TiO₂ surface. Upon adsorption, the B(sp²)-B(sp³) adducts should exhibit two distinct boron environments in the solid-state NMR spectra. As shown in Figure 2D, in all four cases, new peaks appear and shift upfield, indicating changes in the chemical environment of the two B atoms. Previous studies revealed that the ¹¹B signal of sp³ hybridized tetracoordinate boron undergoes an upfield shift relative to the sp² hybridized tricoordinate one (Nöth and Wrackmeyer, 1978). Thus, we attribute the peaks close to 0 ppm to B atom bound to the surface O atom of TiO₂ nanoparticles. X-ray diffraction (XRD) was applied to examine the structure of the TiO_2 by diboron compound modification (Figure 2E). According to the XRD results, no change of the lattice was characterized. To further our understanding of the surface interaction between the diboron molecule and TiO₂ nanoparticles, we carried out XPS and SXANES studies. The Ti 2p spectra shows that the peaks were shifted for 0.2 eV toward the lowerbinding-energy region; this also happened in O 1s spectra (Figure 2F). The valence band spectra show that the band level shifts to lower energy, a shift from 2.5 to 2.16 eV. The Ti 2p, O 1s spectra change means that the surface modification shows significant influence on the surface electron structure of the TiO₂. As shown in Figure 2G, the t_{2g} (L₃) peak of Ti-L edge shows that the B1-TiO₂ is slightly lower than that of pristine TiO₂ and the peak of e_a (L₂) is shifted to higher energy, which can be attributed to the existence of Ti³⁺ (Kronawitter et al., 2011; Stewart et al., 2006). The O-K edge originates from the hybridization of O_{2p} ligand-hole states with the coordinating atoms s, p, d states (Karvonen et al., 2010). In our case, the O-K edge shows that the peaks of B1-TiO₂ that index to the Ti–O bond around 545 eV and the O 2p around 539 eV are much higher than that of pristine TiO₂ (Figure 2H), demonstrating more electron location in O 2p orbital and Ti–O bond instead of the O $e_{\rm q}$ orbitals (de Groot et al., 1989). Meanwhile, the Ti-K edge line shows no significant change between B1-TiO₂ and pristine TiO₂ in the preedge region and the EXANE region (Figure S6). The reason is that the diboron compound is bonded with surface ipso-O²⁻ atoms. Moreover, the B 1s XPS also provides consistent results indexing of B species with different binding energies (Figure S7). On the basis of these spectroscopic evidences, we propose that the coordination of diboron compound with surface O sites on TiO₂ facilitates the electron transfer from the *ipso*- O^{2-} to the adjacent Ti⁴⁺ site and this coordination further stabilizes the newly formed Ti³⁺-



Figure 3. DFT Calculation Results

(A) Optimized geometric structures and charge difference density mappings for no adsorption.
(B) Optimized geometric structures and charge difference density mappings for adsorption of **B1** on the TiO₂ {101} surface. The numbers in the figure stand for the Bader charges on the various atoms. The isosurfaces of the electron density difference plots are all 0.001 e Å⁻³. The yellow and light blue surfaces represent electron depletion and accumulation. The blue, red, earth yellow, green, and pink spheres stand for the Ti, O, C, B, and H atoms, respectively.
(C) Projected density of state (PDOS) plots for no adsorption.

(D) Projected density of state (PDOS) plots for adsorption of **B1** on the TiO_2 {101} surface (the insets on the right are the magnification of the inter band at near the Fermi energy level). The dashed line shows the Fermi level.

 O^{-} species. We note that the activation of inert lattice O^{2-} and the stabilization of surface Ti³⁺ enabled by the formation of $\equiv B-O^{-}-Ti^{3+}$ moiety is a novel process that has not been reported in TiO₂ modification.

DFT Calculation Results

We next investigated the surface Ti^{3+} -O⁻⁻ structure in the blue TiO_2 nanoparticles by simulating the adsorption of B1 on the TiO $_2$ surface and the accompanying charge transfer between them using DFT calculations. According to the literature, the {101} facet in anatase TiO₂ is the dominant facet, and we set up the adsorption model of anatase TiO₂ {101} facet (Lazzeri et al., 2001). Our calculations showed that the Fermi levels in no adsorption is located in the TiO₂ band gap (Figures 3A and 3C). Importantly, the diboron-adsorbed TiO₂ model shows a clear charge transfer from B1 to the TiO₂ {101} surface (Figure 3B). Furthermore, the Bader charge calculation showed substantial electron transfer from the B_1 atom to the bridging O atom. Furthermore, those electrons (light blue region) are localized between the bridging O atom and the neighbor surface Ti atom according to the calculated charge difference density mapping. The calculated projected density of state (PDOS) plots showed that the Fermi level rises to the bottom of the TiO₂ conduction band, which is mainly derived from Ti 3d states (Figure 3D). Some Ti 3d and O 2p states below the Fermi level appear, which are filled with electrons. The filled Ti 3d states indicate the partial reduction of the surface Ti sites, which explains the existence of the Ti³⁺ species observed in our EPR experiments. Moreover, the adsorption model of B2, B3, and B4 on TiO_2 {101} surface was also calculated (Figures S8A-S8C and S9). These data showed that a new electron-filled band-gap state consisting of Ti 3d appears in all the three models (Figures S8D-S8F and S9). This strongly suggested that the adsorption of organic diboron compounds can lead to the formation of Ti^{3+} species on the TiO_2 surface. The adsorption model based on rutile TiO_2 shows similar results (Figures S10 and S11). The adsorption model of the interface between anatase TiO_2 and rutile TiO_2 was also set up to evaluate the heterojunction in P25 nanoparticles (Figures S12 and S13). To interpret the stability of the surface Ti^{3+} in B-TiO₂ in the air atmosphere, the oxygen adsorption models were set. The calculation result shows that O_2 molecule is



Figure 4. Photodetectors and Perovskite Solar Cells Performance

(A) Schematic of the FTO/TiO $_2/diboron/Spiro-OMeTAD/Au \ photodetector.$

(B) Time-dependent current curves for the detectors under light illumination (λ > 400 nm) with 10 s on-off switching intervals.

(C) Time-dependent current curves for these photodetectors under illumination intensities (λ > 400 nm) of 19.5, 26.5, 35.0, and 44.5 mW·cm⁻².

(D) Nyquist plots from the photodetectors in the dark.

(E) Schematic of the ITO/SnO₂/diboron/Perovskite/Spiro-OMeTAD/Au solar cell.

(F) Current density-voltage curves (J-V) of the diboron-modified perovskite solar cell.

(G) Photovoltaic parameters of the diboron-modified perovskite solar cell. FF, fill factor; PCE, power conversion efficiency.

hesitant to adsorb to the adjacent Ti_{5c} site of the molecule (energy rising process), which represents the protection role of the molecule (Figure S14 and Table S1).

Photodetectors and Perovskite Solar Cells Performance

To further explore the optoelectronic properties of this diboron-adsorbed TiO_2 material under visible-light excitation, thin films were then fabricated into photodetectors (denoted as $FTO/TiO_2/di-boron/Spiro-OMeTAD/Au$, Figure 4A). The time-dependent current curves (*I-t*) for these diboron-type



photodetectors under visible-light illumination (>400 nm) exhibited excellent on-off switching repetitions through five cycles (interval = 10 s) (Figure 4B). The photocurrent for these four diboron-type detectors increased linearly with the light intensity (Figures 4C and S15 and Table S2). In contrast, the observed photocurrent for TiO₂ without diboron modification did not change regardless of the on-off state, which was ascribed to the intrinsic band gap of anatase TiO_2 (3.2 eV). Furthermore, the charge transport process in diboron-modified devices was investigated using electrochemical impedance spectroscopy measurements in the dark at 0 V (Figure 4D). The larger semicircle radius of Nyquist plot in the low-frequency region for the B3 device compared with the B1, B2, and B4 devices indicated less electron recombination (Kim et al., 2012). This observation agreed well with the highest photocurrent for the B3 device (Figure 4B). The IPCE (incident photon-to-electron conversion efficiency) was also evaluated, and the B3-TiO2 device shows the best performance (Figure S16). The B-TiO₂ photodetectors show good response toward longtime on-off tests compared with the TiO₂ photodetector (Figures S17 and S18), and the response of the photodetectors does not attenuate after being stored in a dry air atmosphere for at least 7 months (Figure S19). We also fabricated solar cells based on the diboron molecule interface modulation, and the J-V performance of the device was characterized showing an improved solar cell performance (Figures 4E-4G). The data show that the B3-treated device based on the B3-SnO2 interface has a better performance compared with the reference device of SnO₂-based perovskite solar cell.

We note that previously developed solar cells such as dye-sensitized solar cells and perovskite solar cells inevitably involve the use of expensive dyes and/or toxic and unstable reagents. Thus, our approach to generating visible-light active center via the adsorption of stable and inexpensive diboron compounds holds great advantages over these traditional techniques. Furthermore, a diverse range of diboron compounds can be conveniently synthesized from **B4**, potentially allowing for the fine-tuning of semiconductor-based materials. More importantly, the present strategy is not limited to TiO_2 . In our studies, we have determined that a broad range of other common semiconducting oxides, including ZnO, SnO_2 , Nb_2O_5 , and In_2O_3 , were also successfully modified by diboron reagents, thus clearly demonstrating the generality of this method for semiconducting metal oxide modification (Figure S20).

Limitations of the Study

In the current study, perovskite solar cells assembled from B-TiO₂ display relatively poor performance (Figure S21). Many factors such as device fabrication could influence the performance of such device. Additionally, in our present work, mesoporous TiO₂ was used in B-TiO₂-based devices, whereas planar SnO₂ was used in B-SnO₂-based devices. We are uncertain if this deviation led to the contrasting device performance based on B-TiO₂ and B-SnO₂ materials. Nonetheless, the three orders of magnitude improvement observed in the B-TiO₂-based photodetector clearly demonstrates the utility of this diboron-based modification strategy. Currently, we are actively pursuing other applications based on diboron-modified semiconductor oxide materials featuring unique surface defects.

METHODS

All methods can be found in the accompanying Transparent Methods supplemental file.

SUPPLEMENTAL INFORMATION

Supplemental Information can be found online at https://doi.org/10.1016/j.isci.2019.09.024.

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AUTHOR CONTRIBUTIONS

F.M. conceived the idea. Y.C., P.Z., Y.T., Z.L., and B.-W.D. performed experiments. Y.C., P.Z., and X.Z. conducted the DFT calculation. All authors analyzed data. Y.C., P.Z., Y.T., X.Z., Y.Y., S.-D.J., F.M., and W.M. wrote the manuscript. F.M. directed the whole project.

DECLARATION OF INTERESTS

The authors declare no competing interests.

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Supplemental Information

Modification of TiO₂ Nanoparticles

with Organodiboron Molecules Inducing

Stable Surface Ti³⁺ Complex

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Supplementary Materials

This PDF file includes:

Transparent Methods Figures S1 to S21 Tables S1 to S3

Transparent Methods

EXPERIMENTAL PROCEDURES

Sample preparations

B1-TiO₂-N₂ sample: In a glovebox, 50 mg B₂Pin₂ was dissolved in 2 mL dry CH₃OH. Then 0.5 g TiO₂ (Degussa P25, 20 nm) was added into the solution. The mixture was stirred for overnight at room temperature. After evaporation of the solvent by an oil pump, the titled sample was obtained as a blue powder. This blue titania is sensitive to oxygen and will fade when exposed to air. However, when the sample is again isolated from air, the blue color returns. By the same procedure, the corresponding **B2**-TiO₂-N₂, **B3**-TiO₂-N₂, **B4**-TiO₂-N₂ were prepared from **B2** B₂(OH)₄, **B3** B₂Cat₂, **B4** B₂(NMe₂)₄, respectively.

B1-TiO₂ sample: This sample was obtained by heating the above sample at 80 °C under high vacuum for 3 h. **B2**-TiO₂, **B3**-TiO₂ and **B4**-TiO₂ were obtained by the same procedure.

Device Fabrication

Photodetector fabrication:

The pre-patterned FTO substrates (NSG) were cleaned by UV-ozone treatment for 15 min, followed by cleaning ultrasonically with deionized water, detergent, acetone, and isopropanol in succession for 20 min. A 30 nm TiO₂ compact layer was deposited on the pre-cleaned FTO glass by spray pyrolysis using O₂ as the carrying gas at 450 °C from a precursor solution of 0.6 mL titanium diisopropoxide bis(acetylacetonate) and 0.4 mL acetylacetonate in 9 mL anhydrous ethanol. A 150 nm mesoporous TiO₂ was coated on the substrate by spin-coating with a speed of 5000 rpm for 10 s with a ramp rate of 2000 rpm s⁻¹, from a diluted 30 nm TiO₂ particle paste (Dyesol 30 NR-D) in ethanol with the weight ratio of TiO₂ paste/ethanol = 1:6, and then the substrate was sintered at 500 °C for 30 min. For the adsorption of diboron compounds, the mesoporous TiO₂ films were immersed in 5 mg mL⁻¹ B₂Pin₂, B₂Cat₂, B₂(OH)₄ in anhydrous isopropanol at room temperature for 24 h in the nitrogen glove box, respectively. Then the diboron-sensitized TiO₂ films were heated

at 100 °C for 5 min. For the sample of B₂(NMe₂)₄-sensitized TiO₂ films, B₂(NMe₂)₄ solvent was spin-coated on the TiO₂ film with a speed of 2500 rpm for 20 s, then annealed at 40 °C for 2 min to remove extra solvent. The amount of diboron compounds on each film were examined by ICP method (Table S3). After cooling down to room temperature, 2,2',7,7'tetrakis(N,N-di-pmethoxyphenylamine)-9,9'-spirobifluorene (spiro-OMeTAD) solution was spin-coated onto the active layer at 4000 rpm for 30 s. A spiro-OMeTAD solution was prepared by dissolving 72.3 mg of spiro-OMeTAD in 1 mL of chlorobenzene, to which 28.8 mL of 4-tert-butyl pyridine and 17.5 mL of lithium bis(trifluoromethanesulfonyl)imide (Li-TFSI) solution (520 mg LI-TSFI in 1 mL acetonitrile) were added. All devices were stored in a desiccator (humidity < 15 %) for 12 h. Finally, 80 nm of gold was deposited under vacuum through a shadow mask.

Solar cell fabrication:

SnO₂-based perovskite solar cells fabrication. The ITO substrates were cleaned with ultrapure water, acetone, ethanol and isopropanol subsequently. The cleaned ITO substrates were treated under ozone irradiation for 45 min. Then the subsequent were coated with a SnO₂ nanocrystal solution at 4000 rpm/min for 30 second forming a 50 nm thick layer, then annealed at 150 °C for 30 min in air atmosphere. Then 0.1 mM B3 DMF solution were spin coated onto the layer for 5000 rpm for 30 second in N₂ atmosphere, then annealed at 70 °C for 5 min. The perovskite film was fabricated by a two-step solution process: the 1.3 M PbI₂ in DMF/DMSO (9:1) mixture were spin-coated on the ITO substrate for 2500 rpm/min for 30 second, then annealed for 1 min at 70 °C in N₂ glovebox. A mixed organic cation isopropanol solution (MAI 0.12 M; MABr 0.05 M; MACl 0.07 M; FAI 0.23 M) were spin coated for 30 second at 2300 rpm/min and subsequent annealation at 150 °C. Then a spiro-OMeTAD/chlorobenzene solution including 35 µL Li-TFSI/acetonitrile, 30 µL 4-tertbutylpyridine, was coated onto the perovskite layer with 3500 rpm/min for 30 second. After that, 80 nm Au were deposit onto the substrate by thermo evaporation method. The fabricated solar cell was stored in a desiccator (humidity < 15 %) for 12 h before performance measurement.

<u>TiO₂-based perovskite solar cells fabrication.</u> The pre-patterned FTO substrates (NSG) were cleaned by UV-ozone treatment for 15 min, followed by cleaning ultrasonically with deionized water, detergent, acetone, and isopropanol in succession for 20 min. A 30

nm TiO₂ compact layer was deposited on the pre-cleaned FTO glass by spray pyrolysis using O₂ as the carrying gas at 450 °C from a precursor solution of 0.6 mL titanium diisopropoxide bis(acetylacetonate) and 0.4 mL acetylacetonate in 9 mL anhydrous ethanol. A 150 nm mesoporous TiO₂ was coated on the substrate by spin-coating with a speed of 5000 rpm for 10 s with a ramp rate of 2000 rpm s⁻¹, from a diluted 30 nm TiO₂ particle paste (Dyesol 30 NR-D) in ethanol with the weight ratio of TiO₂ paste/ethanol = 6:1, and then the substrate was sintered at 500 °C for 30 min. After cooling down to room temperature, 5 mg mL⁻¹ B₂Pin₂ in anhydrous isopropanol was spin-coated onto the TiO₂ film with a speed of 2000 rpm for 20 s in N₂ atmosphere, then annealed 100 °C for 5 min. The cesium-containing triple cation perovskite precursor solution was prepared by mixing PbI₂ (1.15 M), FAI (1.09 M), PbBr₂ (0.20 M), MABr (0.14 M) and CsI (0.06 M) in a mixed solvent of DMF/DMSO/NMP (DMF/DMSO, 4/1, v/v) and stirred at 70 °C for 10 min before use. The perovskite solution was deposited on TiO₂ substrate with two-step program: 2000 rpm for 10 s with a ramping up speed of 2000 rpm/s, and 6000 rpm for 30 s with a ramping up speed of 2000 rpm/s, respectively. 100 µL of chlorobenzene was poured on the center of the spinning substrate during the spin-coating step at 15 s before the end of the procedure. The substrate was then immediately transferred on a hot plate and heated at 100 °C for 1 h. After cooling down to room temperature, 20 µL of 60 mM Spiro-OMeTAD solution was spin-coated on the perovskite layer at 4000 rpm for 30 s. A Spiro-OMeTAD solution was prepared by dissolving 72.3 mg of Spiro-OMeTAD in 1 mL of chlorobenzene, to which 28.8 µL of 4-tert-butyl pyridine and 17.5 µL of lithium bis(tri-uoromethanesulfonyl)imide (Li-TFSI) solution (520 mg LI-TSFI in 1 mL acetonitrile) were added. All devices were stored in a desiccator (humidity < 15%) in the dark for 12 h. Finally, 80 nm of gold was deposited under vacuum through a shadow mask to ensure the same active layer area (0.0863 cm^2) .

Materials and Methods

The characterization details:

Electron paramagnetic resonance was collected on a Bruker EMX plus under continuous wave model of X wave at room temperature or low temperature hold by liquid Helium.

FT-IR spectroscopy were collected on a Thermal Nicolet iS 50 spectrometer equipped with a MCT detector with high resolution, which is cooled to 77 K by liquid N₂ during the detecting period. For each sample, the data were collected for 128 scans at 4 cm⁻¹ resolutions. Solid-state ¹¹B NMR spectra were performed on a JEOL JNM-ECZ600R NMR spectrometer (14.1 T) equipped with a 3.2 mm probe at room temperature. The corresponding ¹¹B Larmor frequency is 192.4 MHz. A standard solid single pulse sequence was employed, and the $\tau/2$ pulse widths was 4 µs. The ¹¹B NMR experiments were carried out with an effective sample volume of 50 µL and a sample spinning rate of about 20 kHz. Chemical shifts are referenced to 0.1 M B(OH)₃ in D₂O at 19.6 ppm, solid NaBH₄ at -42.1 ppm. All spectra were acquired using 1024 scans with a recycle delay time of 5s. The collected NMR data were processed using Delta software. X-ray diffraction spectra were collected on a Rigaku RINT 2000 spectrometer. Photo detection measurement were performed on a CHI 670E electrochemical workstation from Shanghai Chen Hua Company. The light source was irradiated from a 300 W solar simulator from Newport Company with a 400 nm cut off. Impedance spectroscopy measurements were carried out by CHI670E electrochemical workstation. Solid state UV-VIS-NIR spectrograph were collected on a UV-3600 plus Shimadzu UV-VIS-NIR spectroscopy. XPS spectra were collected on a Thermo Scientific Escalab 250Xi spectrometer. The XAS measurements of the TiO₂ and diboron modified TiO₂ sample were operated in transmittance mode at the XAS beamline of BL14W of Shanghai Synchrotron Source (3.5 GeV). The energy range of the beamline were 4 to 15 keV. The collect data were fitted using the ATHENA software package. The test sample were prepared by mixing of 50 mg LiF and 7 mg TiO₂ or **B**-TiO₂ and then press to a tablet. A Si (311) monochromator were used to collect the data and the energy were calibrated by using a Ti metal foil. We collect the Ti K line data around 4966 eV. The Ti L line and O K line were collected on the BL08U1 of Shanghai Synchrotron Source. Boron K-edges, oxygen K-edges, Titanium L-edges sXANES spectra were collected in total electron yield (TEY) mode at beamline 08U1A of Shanghai Synchrotron Radiation Center. The spectra were calibrated with a clean gold mesh which

give a simultaneous photocurrent based on the photon flux.

ATR-FTIR measurements:

The ATR- FTIR measurements were performed on a Nicolet 6700 FTIR instrument with a mercury cadmium telluride (MCT) detector. IR spectra ranging from 4000 to 1000 cm⁻¹ were recorded by averaging 32 scans with a resolution of 4 cm⁻¹. The TiO₂ film was coated on a ZnSe crystal. 2 mL of methanol was dripped onto the film surface and then purged with Ar for 5 min at a flow rate of 20 mL min⁻¹. After adsorption equilibrium was achieved, the background spectrum was collected. Then a B₂Pin₂-containing methanol solution was added and the corresponding IR spectrum was recorded.

Computational details:

The electronic properties of the no adsorption and adsorption models of B1 on the TiO₂ {101} surface were investigated using the Vienna Ab-initio Simulation Package (VASP). The Perdew-Burke-Ernzerhof (PBE) of the generalized gradient approximation (GGA) was used as the exchange-correlation function (Perdew et al., 1996; Perdew and Wang, 1992). The interaction between the valence electrons and the ionic core was described by the PAW pseudo-potential. The anatase TiO_2 {101} facet is a typical model used to simulate the TiO₂ crystal surface. The interactions between the two adjacent **B1** molecules on the periodic TiO₂ surface were modeled using one 2×4 {101} supercell with four O-Ti-O layers (Figure 3a and 3b). Two models were optimized with adsorption and without adsorption. Considering the balance between the supercell size and the calculational cost, the geometry was only optimized at the gamma point. The energy cutoff of the plane-wave basis was set to 350 eV and the energy convergence was set to 1.0×10^{-10} ⁴ eV. The interactions between the two adjacent **B1** molecules on the periodic rutile $\{110\}$ TiO₂ surface were modeled using one 2×4 {110} supercell with four O–Ti–O layers. The heterojunction adsorption model by joining anatase {112} and rutile {101} of 8A/3R (phase ratio of anatase/rutile phase: 8/3) were also modeled to simulate the adsorption on the heterojunction region. (Zhao et al., 2015) The adsorption energy (E_{ads}) of the **B1** molecules adsorbed on the TiO₂ $\{101\}$ surface was calculated using Equation (1) where E_{total} is the total energy of the **B1**-adsorbed TiO₂ {101} surface, $E(\text{TiO}_2)$ is the free energy of the TiO₂ {101} surface, and E(B1) is the free energy of one B1 molecule in vacuum. $E_{\text{ads}} = E_{\text{total}} - E(\text{TiO}_2) - E(\mathbf{B1})$ (1)

After the geometry optimization, the Monkhorst-Pack $2 \times 2 \times 1$ k-point mesh was used for the partial density of state (PDOS) calculations for the optimized adsorption models. The charge difference density mapping and the bader charge were also calculated to investigate the charge transfer between the **B1** and the TiO₂ surface.



Figure S1. EPR signal of (A) **B2**-TiO₂-N₂ (B) **B3**-TiO₂-N₂ (C) **B4**-TiO₂-N₂ at room temperature or 100 K (The sample were protected under N₂ atmosphere during measurement). Related to Figure 1.



Figure S2. ¹H NMR spectra indicate that **B1** is unchanged upon extraction from the adsorbed sample. Related to Figure 1.



Figure S3. EPR signal of B3-TiO₂. Related to Figure 1.



Figure S4. XRD pattern of boron compound adsorbed on TiO₂ by heat treatment at 80 °C and pristine TiO₂ (Degussa P25, A: anatase, R: rutile). Related to Figure 2.



Figure S5. Calculated vibrational spectra of **B1** (B₂Pin₂, blue line) based on G09 package and experimental FT-IR transmission spectra (red line) (The frequency were corrected by a factor of 0.9614, at 6-31G* level). Related to Figure 2.



Figure S6. (A) Ti K edge line of **B3**-TiO₂ and pristine TiO₂. (B) Extend XAFS spectra of **B3**-TiO₂ and pristine TiO₂. Related to Figure 2.



Figure S7. B 1s XPS spectra of **B1**-TiO₂ and simulate multiple peak fitting curve. Related to Figure 2.



Figure S8. The optimized geometry structures of (A) **B2** B₂(OH)₄, (B) **B3** B₂Cat₂ and (C) **B4** B₂(NMe₂)₄ adsorbed TiO₂ {101} surface. Projected density of state (PDOS) plots of (D) **B2** B₂(OH)₄, (E) **B3** B₂cat₂ and (F) **B4** B₂(NMe₂)₄-adsorbed TiO₂ {101} surface. The dashed line stands for the Fermi level. Related to Figure 3.



Figure S9. The optimized geometric structures and charge density difference plots of (A) **B1** B₂Pin₂ (B) **B2** B₂(OH)₄, (C) **B3** B₂Cat₂ and (D) **B4** B₂(NMe₂)₄ adsorbed TiO₂ {101} surface. The isosurface levels are set at 0.001 e Å⁻³. The yellow and skyblue surfaces represent electron depletion and accumulation, respectively. Related to Figure 3.



Figure S10. The optimized geometric structures and charge density difference plots of (A) **B1** B₂Pin₂ (B) **B2** B₂(OH)₄, (C) **B3** B₂Cat₂ and (D) **B4** B₂(NMe₂)₄ adsorbed on rutile TiO₂ {110} surface. The isosurface levels are set at 0.001 e Å⁻³. Related to Figure 3.



Figure S11. Projected density of state (PDOS) plots of (A) pristine TiO₂, (B) B1 B₂Pin₂,
(C) B2 B₂(OH)₄, (D) B3 B₂cat₂ and (E) B4 B₂(NMe₂)₄-adsorbed rutile TiO₂ {110}
surface. The dashed line stands for the Fermi level. Related to Figure 3.



Figure S12. The isosurfaces of the electron density difference plots of B3 molecule adsorption on (A) the interface of anatase {112} (right) and rutile {101} (left) TiO₂ (phase ratio of anatase/rutile=8/3), (B) the surface of anatase {101}, (C) the surface of rutile {110}. The isosurfaces of the electron density difference plots are 0.001 e Å⁻³. The yellow and light blue surfaces represent electron depletion and accumulation. The blue, red, yellow, green and pink spheres stand for the Ti, O, C, B and H atoms. Related to Figure 3.



Figure S13. Projected density of states (PDOS) plots of (A) pristine B3 adsorption on the anatase {112}-rutile {101} interface, (B) B3 B2cat2-adsorbed anatase {112}-rutile {101} interface, (C) pristine anatase {101} surface, (D) B3 B2cat2-adsorbed anatase {101} surface, (E) pristine rutile {110} surface, (F) B3 B2cat2-adsorbed rutile {110} surface. The dashed lines stand for the Fermi level. Related to Figure 3.



Figure S14. Optimized geometry structures of O₂ adsorption on the TiO₂ {101} plane of (A) **B1**-TiO₂, (B) **B2**-TiO₂, (C) **B3**-TiO₂, (D) **B4**-TiO₂ (adsorb to the Ti₅ site adjacent to the O₂ site with diboron compound adsorption). Related to Figure 3.



Figure S15. The simulated linear dynamic response SNR curve (signal to noise response). Related to Figure 4.



Figure S16. IPCE spectra of the photodetectors and the integration of the IPCE spectra for the photodetectors. Related to Figure 4.



Figure S17. The transient photocurrent curve of the diboron compound devices and TiO₂ device alone with on off light switches ($\lambda > 400$ nm). Related to Figure 4.



Figure S18. The comparation time dependent current curves for the **B3**-TiO₂ and TiO₂ detectors under light illumination with 10 s on/off switching intervals. (no cut off filter). Related to Figure 4.



Figure S19. The transient photocurrent curve of the diboron compound device alone with on off light switches stored in the air atmosphere for 6 months (light source: 300 W Xe lamp with a 400 nm cut filter). Related to Figure 4.



Figure S20. (A-D) EPR spectra of **B1**-metal oxides at various temperatures. (E-H) UV-Vis spectra of **B**-metal oxides. In these preliminary studies, by adopting a same adsorption operation to other common semiconductor oxides, such as ZnO, In₂O₃, SnO₂ and Nb₂O₅, similar phenomena were observed. EPR spectra demonstrated the appearance of reduced metal species, such as Zn⁺, In²⁺, Sn³⁺ and Nb⁴⁺. Additionally, the UV-Vis of these modified materials were also collected. These diboron-modified metal oxides samples were found to absorb visible light with higher intensity in the NIR region. Related to Figure 4.



Figure S21. (A) Schematic of the FTO/TiO₂/Perovskite/Spiro-OMeTAD/Au solar cell (B) Current density-voltage curves (J-V) of the diboron modified perovskite solar cell. Related to Figure 4.

Table S1. The adsorption energy of oxygen molecule to the diboron molecule preabsorption TiO_2 {101} plane. Related to Figure 3.

	$E(TiO_2-B)$	$E(O_2)$	$E(TiO_2-B-O_2)$	E_{ad}
B1- TiO ₂	-2063.56119598	-9.44011081	-2071.61712553	1.38418126
B2- TiO ₂	-1883.20183442	-9.44011081	-1891.54740527	1.09453996
B3- TiO ₂	-2002.25704951	-9.44011081	-2011.19765769	0.49950263
B4- TiO ₂	-2028.25727155	-9.44011081	-2036.34654405	1.35083831

The adsorption energy of oxygen molecule to the diboron molecule pre-absorption TiO_2 {101} plane was calculated:

 $E_{ad} = E(TiO_2-B-O_2) - E(TiO_2-B) - E(O_2)$

Photodetector	R (μA/W)	D (10^{12} Jones)	Ilight/Idark	SNR
TiO ₂	0.18	27.7	3.06	2.05
B4- TiO ₂	35.4	3.2	5.39	4.39
B2- TiO ₂	151	9.3	214	213
B1- TiO ₂	406	10.0	726	725
B3- TiO ₂	814	8.9	778	777

Table S2. The parameter of the photodetector devices. Related to Figure 4.

The illumination power was calculated as P_{ill} = 44.5 mW/cm² × 0.0863 cm² = 3.84 mW. The detail of the origin of the photocurrent

1. The responsivity R is calculated according to the equation:

 P_{ill} is definite as the illumination power on the photo harvest area of the device.

2. The light-detection ability of the photodetector can be evaluated by D:

$$D=R \times (S/2qI_{dark})^{1/2}$$

3. The signal-to-noise ratio (SNR) can be defined as:

SNR=(Ilight-Idark)/Idark

We get the amount of the diboron compounds by detection of B element concentration using ICP method (ICP-MS, iCAP Q, Thermo, Waltham, USA). We modified the mesoporous TiO₂ layer with diboron molecules as depicted in the experimental part. The mesoporous TiO₂ layer were then removed from the FTO glass and this sample was subject to ICP measurement. The results are given below.

Table S3. Amount of the diboron compounds on FTO glass. Related to Figure 4.

	c (B) μg/L (original data, amount of 4 FTO)	Diboron compound molecule weight	Diboron compound mass per FTO (µg)	Diboron compound molar per FTO (µmol)
B1- TiO ₂	794.8	253.94	22.9354	0.090318
B2- TiO ₂	1027	89.65	10.46256	0.116705
B3- TiO ₂	1037.6	237.81	28.03996	0.117909
B4- TiO ₂	775	197.92	17.43045	0.088068

Reference:

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