Surface-specific interaction by structure-match confined pure high-energy facet of unstable TiO$_2$(B) polymorph

Guolei Xiang, Yang-Gang Wang, Jun Li, Jing Zhuang & Xun Wang

Department of Chemistry, Tsinghua University, Beijing 100084, China.

Surface structures and surface interactions are key factors that influence the reactivity and stability of nanomaterials. Combining experimental and theoretical investigations, we illustrate the roles of surface interactions in the formation and phase stability of an unusual TiO$_2$(B) polymorph that preferentially exposes the plane of the highest surface energy. We find that the favorable bidentate adsorption of ethylene glycol on the TiO$_2$(B)(010) plane enables the formation and confines the phase stability of TiO$_2$(B) ultrathin nanosheets. The essence of such selective generation of the unusual nanostructure with ultrahigh purity both in phase and morphology lies in the specific adsorption driven by the matched interface structures. The general roles of structural match for the activity and stability in physical interactions are elucidated.

Control over phase and morphology of nanomaterials is a key method in improving their performance$^1$. During the preparation and processing of nanocrystals, both growth and phase stability are inherently dominated by thermodynamics, especially on a nanometer scale$^{2,3}$. With decreasing particle size, surface energy becomes the most important parameter that dictates the phase stability of nanocrystals$^{4,5}$. It has been revealed that the phase stability and transformation of common TiO$_2$ polymorphs (rutile, anatase and brookite) are size-dependent; the most stable phase above 35 nm is rutile, but the reversed surface energy makes anatase the most stable phase below 11 nm$^3$. It is well recognized that stable equilibrium shapes of nanocrystals are determined by intrinsic lattice symmetries and the relative order of surface energies of all crystalline facets$^5,6$. Though surface energy largely controls the growth process of crystals, it is still possible to realize shape-controlled synthesis of many nanocrystals in solution by introducing various additives, e.g., ions, surfactants and polymers, to change the growth rates along different orientations$^5,6$. In these processes, certain growth planes are favored due to the selective adsorption of additive molecules$^1$. This provides kinetic approaches to control the growth of nanocrystals by introducing selective or non-selective interactions on the surfaces. Many materials can now be synthesized with well-controlled sizes and shapes$^{1,8,10,11}$.

As an important semiconducting oxide, TiO$_2$ has been widely studied in the fields of heterogeneous catalysis$^{12}$, photocatalysis$^{13,14}$, photovoltaics$^{15}$, etc., involving three major phases$^{16}$. We previously developed a facile method to prepare ultrathin nanosheets of TiO$_2$(B), an unusual phase of TiO$_2$, via a one-step solvothermal reaction$^{17}$. Through the procedure, 1-nm thick sheets with a high exposure percentage of (010) planes can be readily obtained. However, the nature of the surface interactions in the formation of this metastable phase and in preferential exposure of the (010) plane have been elusive. Herein we investigate the surface interactions of TiO$_2$(B) with different solvents to reveal the underlying mechanism of its formation and phase stability. Through a combined experimental and theoretical study, we have found that the stability of this unusual phase with 2D shape is confined by the structure-matched specific interactions on a special crystalline facet. Structural match at different scales is an important mechanism to control specific interactions or processes with definite results, e.g., molecule recognition$^{18}$, and sometimes they are also described in terms of specificity or selectivity. Here we emphasize that it is the structure-matched specific interaction between ethylene glycol (EG) molecules and TiO$_2$(B) over the (010) plane that drives the formation and stability of the special phase with an ultrathin 2D morphology.
Results

Synthesis of TiO$_2$(B) Nanosheets. TiO$_2$(B) was prepared by directly hydrolyzing TiCl$_3$ or TiCl$_4$ in ethylene glycol at 150°C. All the products are uniform 2D nanosheets as shown in Figures 1a–b. The phase character was compared with TiO$_2$(B) nanowires prepared by hydrothermal method and characterized by means of powder X-ray diffraction (XRD) and Raman spectroscopy, which confirm the pure phase of TiO$_2$(B) (Figures 1c–d and S1). Index of the exposed facet was analyzed from a high resolution TEM image (HRTEM) and the corresponding Fourier transformation pattern (Figure 1e). The spots were assigned to 200 and 001, and the angle between them was measured to be 73°, identical to the angle between the reciprocal space axes of $a^*$ and $c^*$ (Figure S2). We further quantified the amount of EG in the product using differential thermal analysis (DTA) and thermogravimetric analysis (TGA) ranging from 25°C to 800°C in air (Figure 1f). EG were completely eliminated at 465°C with a residual weight of 73%. From the TGA and DTA results the total amount of EG on the surface of TiO$_2$(B) nanosheets is estimated to be around 22% by weight (see Figure S3 for more details).

The above results show that TiO$_2$(B) nanosheets can be effectively prepared in EG with high selectivity of phase and morphology. Through contrast experiments we found that TiO$_2$(B) phase could only be directly prepared in EG, while only anatase or the mixture of anatase and rutile were obtained in other hydroxylic solvents, e.g. H$_2$O, CH$_3$OH, C$_2$H$_5$OH. Thus EG is an indispensable component for the formation of TiO$_2$(B). We infer that a strong interaction must exist between EG molecules and TiO$_2$(B)(010) plane that dominates the formation of this unusual phase.

Elucidating the Growth Mechanism by Computational Modeling. To gain insight into the underlying formation mechanism of the TiO$_2$(B) nanosheets, we studied the interactions of EG molecules with TiO$_2$ polymorphs through periodic density functional theory (DFT) calculations using the Quickstep code of CP2K package$^{19,20}$. We systematically studied possible interaction patterns between EG and TiO$_2$(B) by exploring all possible adsorption configurations on the (010) plane. As EG is a bis-hydroxyl compound, both the monodentate and bidentate adsorption configurations on
TiO₂(B)(010) facet were investigated (see Figure 2 for the configuration models).

The geometry optimizations show that for the monodentate adsorption the EG molecule remains in the “chair” configuration as in the gas phase, while for bidentate adsorption the configuration of the EG molecule transforms from “chair” to “boat” configuration to bind with two Ti sites. The calculated adsorption energies for the dissociative configurations (Figure 2c–d) are significantly more negative than those for molecular configurations (Figure 2a–b), indicating that the EG molecules are more favorable to adsorb onto TiO₂(B) (010) facet through dissociative configurations. The dissociative adsorption energies are $-1.92 \text{ eV}$ (or $-0.96 \text{ eV}$ per Ti site) for the bidentate configuration and $-0.95 \text{ eV}$ for the monodentate configuration. These results indicate that the configuration change does not decrease the binding strength of each OH group with the surface Ti sites. It is therefore feasible for EG to interact with the TiO₂(B)(010) plane using both hydroxyl groups to form the most stable configuration. This strong bidentate interaction accounts for the fact that TiO₂(B) can only be synthesized in EG with exposure of a specific (010) plane.

To corroborate this conclusion, we further considered the interactions of EG with TiO₂(B) (001) and (110) planes, which are intrinsically more stable than TiO₂(B) (010) plane in vacuum and in a humid environment. In view of the high reaction temperature, EG molecules are assumed to dissociate after adsorption on these TiO₂(B) facets and the saturated adsorptions are considered in EG solvent. The calculated average adsorption energies per EG molecule on (010), (001), (110) planes (Table 1) are $-1.41, 0.19, -1.22 \text{ eV}$, respectively, confirming the EG molecule binds more favorably to (010) plane compared to the others providing a thermodynamic template to form the (010) plane. We also investigated the surface stability in terms of the change of surface energies ($\gamma$) before and after EG adsorption. For the bare planes without EG adsorption, (001) is the most stable plane with the lowest surface energy, and the order of the surface stability follows (001) > (110) > (010), consistent with previous calculations. The least stable plane (010), however, can be greatly stabilized in EG solvents and becomes more stable than (001) and (110) planes, which agrees well with our experimental results that (010) is the solely exposed plane. We further considered the interactions of EG with other polymorphs of TiO₂, anatase and rutile, although neither are formed during the synthesis of TiO₂ in EG. As the common planes for the two phases are (101), (100), (001) for anatase and (110), (100), (111) for rutile, they were chosen to compare with TiO₂(B) (010) plane. All the adsorption energies for the saturated bidentate adsorptions of EG on the planes of anatase and rutile are found to be less stable than on TiO₂(B)(010) plane (Table 1). This result suggests that the bidentate binding structures of the EG molecules on either anatase or rutile surfaces are less stable than that with TiO₂(B) (010), consistent with the rare phase preference for the later. Clearly the strong bidentate interaction makes EG a specific solvent to generate TiO₂(B) polymorph exposing a plane with the highest surface energy. The structure-match interaction likely dominates the selective formation of ultrathin TiO₂(B) nanosheets with highly pure phase.

### Solvent effects on phase stability of TiO₂(B) nanosheets

Compared with anatase and rutile, TiO₂(B) is a less stable phase of TiO₂. To determine the stability of this metastable polymorph in the forms of ultrathin nanosheets, we investigated its phase transformation behaviors in the solid state and in different solvents. The results were monitored via XRD and Raman spectroscopy (Figure 3). Through the temperature-dependent tests by heat-treating the samples for 5 h in solid state, the nanosheets preserved the metastable phase up to 250°C, but it gradually transformed into anatase phase above 300°C. Pure anatase was obtained at 500°C, and only a little amount of rutile appears at 800°C (Figure 3a). To prepare TiO₂(B) nanowires, hydrogen titanate needs to be treated at 400°C, and they start to transform into anatase at 600°C (Figure S4). Thus TiO₂(B) nanosheets are less stable than nanowires, because the phase transition temperature is reduced by almost 300°C.

On the other hand, the phase stability of the TiO₂ nanosheets depends on the chemical environment of the solvent (Figure 3b–c). The TiO₂ material was prepared in EG by adding H₂O to hydrolyze Ti precursor. The products formed by hydrolysis were separated by centrifugation and washed with the respective solvents, EG and ethanol (EtOH), to minimize the influence of any residual water on the results. Control experiments showed the nanosheets after postprocess in either EG or EtOH preserve the initial phase at temperatures up to 200°C, but are transformed into anatase when 0.5 mL water was introduced into ethanol. The results show that the metastable structure is quite sensitive to water. Through systematic tests we checked the phase stability in pure water and found anatase phase appeared above 55°C. Typical results showing good degrees of crystallization after heating at 60 and 70°C for 15 h are shown in the XRD and Raman spectra (Figures 3b–c). There is also a diffraction peak of the intermediate transitional phase in the XRD pattern.

The key factor dominating different roles of the three solvents in the phase transformation results of TiO₂(B) nanosheets appears to be their different interaction strengths with TiO₂(B) (010) plane, that is H₂O > EG > EtOH. To confirm this, the interactions of ethanol, EG, and water molecules with TiO₂(B) (010) surfaces were investigated computationally. Inasmuch as the phase transformation experiments were conducted in bulk solvents, the coverage density of these solvent molecules on the surface is very high, as revealed from the TGA results (Figure 1d). Therefore, the saturation adsorption configurations with all Ti sites occupied by the solvent molecules are used to model the experimental conditions (Figure 4). The calculated

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**Table 1** Adsorption energies (AEs) for EG saturated adsorption on different planes of TiO₂(B), anatase, and rutile, and the surface energies for TiO₂(B)

| TiO₂(B) | AEs (eV) | $\gamma$ without EG (Jm$^{-2}$) | $\gamma$ with EG (Jm$^{-2}$) |
|---------|---------|-------------------------------|-------------------------------|
| 010     | -1.41   | 0.93                          | 0.32                          |
| 001     | 0.19    | 0.62                          | 0.76                          |
| 110     | -1.22   | 0.83                          | 0.35                          |

| Anatase | AEs (eV) | Rutile | AEs (eV) |
|---------|---------|--------|---------|
| 101     | -1.19   | 110    | -0.93   |
| 100     | 0.20    | 101    | -1.12   |
| 001     | -0.78   | 111    | -1.08   |

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**Figure 2** Adsorption configurations and energies of EG on TiO₂(B) (010) plane from DFT calculations: (a) molecular monodentate adsorption (b) molecular bidentate adsorption (c) dissociative monodentate adsorption and (d) dissociative bidentate adsorption.
average adsorption energies of each Ti site for ethanol, EG, and water molecules (−0.56 eV, −0.70 eV and −0.81 eV, respectively, Figure 4d) indicate that it is thermodynamically plausible for water to replace EG molecules, but it is difficult for ethanol molecules to replace EG. The difference in the binding strength of different solvents can be partly attributed to the steric hindrance effects, which deter efficient adsorption of solvent molecules onto the surface of TiO$_2$(B) nanosheets. As a highly polar molecule, water has the strongest adsorption energy due to its smaller size and the lower steric interaction to the adjacent counterparts. The EG molecules can occupy two Ti sites by bidentate adsorption to decrease the steric hindrance, thus having lower adsorption energy than the ethanol molecules.

**Discussion**

For the formation, phase stability and evolution of inorganic nanocrystals, many examples have been reported from the aspects of surface structure and particle size. Using the models of TiO$_2$ and ZnS, the importance of surface interactions with solvents, ions and impurities is emphasized for the phase stability and transformation processes. The essence lies in altering the energy states of nanocrystals by modifying the surface structures through certain interactions that can affect the thermal stability of nanocrystals under specific circumstances. Since it is a surface-initiating process, phase transformation can be adjusted by changing the surface states. Similarly, it is possible to realize phase transition of nanocrystals by modifying their surfaces via the adsorption or desorption of solvent molecules or impurity ions.

In the current work, we illustrate the roles of surface-specific interaction in the formation and phase stability of an unusual TiO$_2$(B) nanosheet. The bidentate adsorption of EG on the (010) plane is shown to enable the formation and confine the phase stability of TiO$_2$(B). Such a relatively stable state of this metastable polymorph is generated by the matched size and configuration on the surface. Thus through the surface-specific interaction, the TiO$_2$(B) (010) plane can be effectively stabilized under the particular chemical environment. Based on the phase transformation and DFT results, we conclude that EG is a phase-stabilizer for TiO$_2$(B) nanosheets due to bidentate adsorption configurations over the (010) facet. The high ratio of surface adsorbed EG, up to 22% by weight, also indicates a strong binding interaction. On the contrary, water is a potent destabilizer of the TiO$_2$(B) phase and favors the formation of anatase. As ethanol binds more weakly than EG on the surface of TiO$_2$(B), it fails to change the energy state of TiO$_2$(B) or to induce phase evolution.

In general, the specific interaction with structural match as illustrated herein is an important mechanism to analyze the processes with high specificity and selectivity. The underlying driving force is the matched factors at a particular scale. In the physical interactions involving the existence and state evolution of matter, the configuration match of size and shape is an important and potentially dominating factor that induces specific pathways with high
The adsorption energy of EG is calculated according to: $E_{\text{ads}} = (E_{\text{slab}+\text{EG}} - E_{\text{slab}})/n$, where $E_{\text{ads}}$ denotes the energy per surface unit cell of the relaxed slab model with $n$ bulk unit cells, $E_{\text{slab}+\text{EG}}$ the total energy of the adsorption complex including the relaxed surface and the adsorbate EG molecule, and $E_{\text{slab}}$ the energy of the isolated EG molecule. The surface energy without EG adsorption is calculated using $\gamma = (E_{\text{slab}} - nE_{\text{bulk}})/2A$, where $E_{\text{slab}}$ is the energy per unit cell of TiO$_2$(B), $A$ the area of the surface, and $n$ the number of the TiO$_2$ unit cell contained in the slab surface. The surface energy with saturated adsorption of EG is calculated as $\gamma = (E_{\text{relaxed}} - nE_{\text{slab}} - mE_{\text{EG}})/2A$, where $m$ is the number of EG molecules binding to the slab surface and $E_{\text{relaxed}} + mE_{\text{EG}}$ is the total energy of slab surface with EG molecule saturatedly adsorbing on the two sides of the slab.

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Author contributions
G.L.X. and Y.G.W. contributed equally to this work. G.L.X. conducted the experimental parts and Y.G.W. performed the DFT calculations. J.Z. contributed to the characterization processes. J. L. was responsible for the theoretical work. X.W. conceived and designed the work and was responsible for the work. All authors discussed the results, wrote and commented on the manuscript.

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