Ti$_3$BN monolayer: the MXene-like material predicted by first-principles calculations†

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The discovery of graphene and other two-dimensional (2D) materials has set the foundation for exploring and designing novel single layered sheets. The family of 2D materials encompasses a wide selection of compositions including almost all the elements of the periodic table and they have the potential to play a fundamental role in the future of electronics, composite materials and energy technology. Therefore, searching for new 2D materials is a big challenge in materials science. In this work, we theoretically designed a monolayer of Ti$_3$BN following the strategy of “atomic transmutation”. The Ti$_3$BN monolayer can be considered as three Ti-atomic layers being interleaved with one N-atomic layer and one B-atomic layer, in the sequence of Ti$_1$–N–Ti$_2$–B–Ti$_3$. The moderate cohesive energy, positive phonon frequencies and high melting point are the best guarantees for good stability of Ti$_3$BN. Based on a global minimum structures search using the particle-swarm optimization (PSO) method, Ti$_3$BN is the lowest energy structure in 2D space, which holds great promise for the realization of layered Ti$_3$BN in experiment. Based on density functional theory (DFT) calculations, Ti$_3$BN is intrinsically metallic and its electronic properties can be modulated by varying the surface groups, such as OH or F-terminations. If realized in experiment, it may find applications in many aspects.

1. Introduction

Since graphene was realized experimentally in 2004,1 2D materials have attracted significant interest and a large variety of freestanding monolayer solids have been successfully fabricated, such as hexagonal boron nitride (h-BN),2 metal chalcogenides (MoS$_2$, WS$_2$),3,4 silicene5 and so on. Due to their unique electronic, chemical, and mechanical properties, those 2D materials have found wide applications in supercapacitors, solar cells,6 lithium ion batteries7 and water splitting applications8 and other areas of energy and environment.

Recently, a new family of graphene-like 2D materials termed as MXenes were successfully synthesized by selectively extracting the “A” element from the layered MAX phases (A is an A-group element, mostly Al or Si) in the aqueous HF.9 MAX phases are a large (>60 members) family of layered ternary early transition-metal carbides, nitrides, and carbonitrides with P6$_3$/mmc symmetry.10 To date, several MXenes have been synthesized successfully, including Ti$_3$C$_2$,11 Ti$_2$C$_3$,12 Ta$_4$C$_3$,13 V$_2$C$_5$,14 TiNbC$_6$,15 Nb$_2$C$_7$ and Mo$_2$C.16

With the increasing interest in MXenes, a mass of experimental and theoretical efforts related to their synthesis, structures, properties and potential applications have been made experimentally and theoretically.17,18 Among the as-synthesized MXene phases, the most studied MXene is Ti$_3$C$_2$, prepared by immersing Ti$_3$AlC$_2$ in HF solutions at room temperature.11 Ti$_3$C$_2$ was predicted theoretically to be good electrical conductors and its electrical conductivities can be tuned by different surface terminations.21 What’s more, Ti$_3$C$_2$ have been proved to be very promising as anode materials for Li-ion batteries and as hydrogen storage media.22,23

Motivated by the Ti$_3$C$_2$ monolayer, which is composed of three Ti-atomic layers being interleaved with two C-atomic layers, herein we designed a new 2D material of Ti$_3$BN by performing density functional theory (DFT) calculations following the strategy of “atomic transmutation”, which means substituting certain types of elements with their neighboring elements in the periodic table but the total number of valence electrons is kept unchanged.24,25 A major breakthrough has been made in finding and designing novel materials. For example, if one were to substitute the oxide ions in ZnO with N and F, one would ultimately obtain Zn$_3$NF whose conduction and valence band edges are more favorable for water splitting.26 When two C atoms in graphene were transmutated with one B atom and one N atom, h-BN with wide band gap and new functionalities will be obtained.27 Therefore Ti$_3$BN monolayer...
can be thought of as obtained by substituting the two C-atomic layers of Ti1C2 monolayer with one nitrogen-atomic layer and one boron-atomic layer, respectively, in consideration of that nitrogen and boron are two nearest-neighbors of carbon in the periodic table and Ti1BN is isoelectronic to Ti1C2.

2. Computational method

DFT calculations were performed within the Perdew–Burke–Ernzerhof (PBE) generalized gradient approximation\textsuperscript{9} and the projected augmented wave (PAW) method\textsuperscript{29,30} as implemented in the Vienna \textit{ab initio} simulation package (VASP). To reach convergence criteria for both energy and forces, a special \(k\)-point sampling with a \(k\)-point separation of 0.04 Å\(^{-1}\) is applied for the Brillouin-zone integration and the cutoff energy for the plane wave basis set is 408 eV. The ground state geometries of the 2D Ti1BN are obtained with all the atomic positions relaxed until their residual forces are less than 0.01 eV Å\(^{-1}\). In building the monolayer models, a vacuum thickness of 15 Å is adopted to avoid the interactions between adjacent layers. The phonon dispersion curves of Ti1BN monolayer was calculated using the density functional perturbation theory (DFPT)\textsuperscript{31,32} as implemented in the PHONOPY program interfaced with VASP. To obtain the accurate electronic properties of Ti1BN monolayer, hybrid functional calculations using Hedin–Scuseria–Ernzerhof (HSE06) functional\textsuperscript{33} have been employed.

The thermal stability of Ti1BN monolayer was assessed by first-principles molecular dynamics (MD) calculations using the PAW pseudo-potential and PBE functional as implemented in VASP.\textsuperscript{14} For each temperature, a preheating for 1 ps was applied for the initial geometry structure. And the MD calculations were in NVT ensemble, lasting for 10 ps with a time step of 1.0 fs. To control the temperature, Nose–Hoover method was applied.\textsuperscript{35}

The global minimum structure for Ti1BN monolayer was searched by particle-swarm optimization (PSO) method within the evolution algorithm which was implemented in CALYPSO code.\textsuperscript{36,38} The population size was set as 30, and the number of generation was maintained at 25. Unit cells containing 5, 10, and 15 atoms were considered. The structure relaxations during the PSO searching were carried by using PBE functional as implemented in VASP.

3. Results and discussion

3.1 Structural properties and stability of Ti1BN monolayer

The optimized structure of our designed Ti1BN monolayer is shown in Fig. 1(a). Similar with Ti1C2 monolayer, Ti1BN monolayer is crystallized in the space group \(P3m1\) (no. 156) and each unit cell contains five atoms, all in different atomic planes with the sequence of Ti1–N–Ti2–B–Ti1. To avoid the interactions between the Ti1BN monolayer and its periodic images among the normal direction, a vacuum layer of 15 Å is used and the calculated lattice constants for the unit cell are \(a = b = 3.095\) Å. In Ti1BN monolayer, the length of Ti1–N and N–Ti2 bonds (2.054 and 2.195 Å) are shorter than those of corresponding Ti–C bonds (2.064 and 2.212 Å)\textsuperscript{39,40} in Ti1C2 monolayer, while the Ti2–B and B–Ti3 bond lengths are 2.249 and 2.117 Å, respectively, longer than those of corresponding Ti–C bonds in Ti1C2 monolayer. The thickness of Ti1BN monolayer is 4.78 Å with the atomic layer distances of 1.01, 1.27, 1.36 and 1.13 Å, respectively.

To elucidate the chemical bonding and stabilization mechanism of Ti1BN monolayer, the deformation electronic density\textsuperscript{42} has been calculated. As shown in Fig. 1(b), there is remarkable electron transfer from Ti atoms to N atoms and from Ti atoms to B atoms, indicating the electronically stabilization in the Ti1BN monolayer. Bader charge analysis shows that the net charges on C, N and B atom are \(-1.65\) e and \(-1.57\) e and those on Ti1, Ti2, and Ti3 atom are \(-0.97\) e, +1.34 e and +0.91 e, respectively. The electron localization function\textsuperscript{44} of Ti1BN monolayer is also calculated to highlight the electron distribution. As seen from the isosurfaces of electron localization functions presented in Fig. S1,\textsuperscript{1} the electrons are complete delocalized around the Ti atoms, and widely distributed in the N and B frameworks, which also suggests the electron transfer from Ti atoms to N and B atoms.

Although Ti1BN monolayer possesses similar structure properties with Ti1C2 monolayer, the question whether Ti1BN monolayer is as stable as Ti1C2 or not need to be answered. The cohesive energy of Ti1BN monolayer is a useful argument for evaluating its stability, defined as \(E_{\text{coh}} = (xE_{\text{Ti1}} + yE_{\text{N}} + zE_{\text{B}} - E_{\text{Ti1BN}})/(x + y + z)\), where \(E_{\text{Ti1}}, E_{\text{N}}, E_{\text{B}}\) and \(E_{\text{Ti1BN}}\) are the total energies of a single Ti1 atom, a single N atom, a single B atom and Ti1BN monolayer, \(x, y\) and \(z\) are the number of Ti1, N, and B atoms in the supercell, respectively. Based on our DFT calculations, the cohesive energy of Ti1BN monolayer is 7.46 eV per atom, which is a little smaller than that of graphene (7.95 eV per atom)\textsuperscript{45} and higher than that of Ti1C2 (about 7.00 eV).\textsuperscript{13} The relatively large cohesive energy of Ti1BN monolayer indicates that Ti1BN monolayer is a stable phase with strong chemical bonds. What’s more, the small cohesive energy difference between Ti1BN monolayer and Ti1BN bulk (7.46 eV per atom vs. 7.82 eV per atom) means that it’s favorable to obtain Ti1BN monolayer from its bulk phase. The elastic constants of Ti1BN monolayer were
then calculated to be $C_{11} = C_{22} = 202.11 \text{ N m}^{-1}$, $C_{12} = C_{21} = 59.24 \text{ N m}^{-1}$, which satisfy the mechanical stability criteria, indicating that the Ti$_3$BN monolayer is also mechanically stable.

The kinetic stability of Ti$_3$BN monolayer has been further confirmed by its phonon dispersion along the high-symmetry directions in the first Brillouin zone. As shown in Fig. 2(a), there is no appreciable imaginary frequency in the phonon dispersion curves, implying the good kinetic stability of Ti$_3$BN monolayer. The highest frequency of Ti$_3$BN monolayer is 653.60 cm$^{-1}$, higher than that of the widely studied MoS$_2$ monolayer (473 cm$^{-1}$) and silicene (580 cm$^{-1}$). The high frequency suggests that the related bonds in Ti$_3$BN monolayer are strong.

Finally, the thermal stability of Ti$_3$BN monolayer was investigated by first-principles molecular dynamics (MD) calculations. A $5 \times 5$ supercell containing 125 atoms was used here and three individual MD calculations for Ti$_3$BN monolayer at temperatures of 500 K, 800 K, and 1000 K were performed. Fig. 3 presents the snapshots of Ti$_3$BN monolayer at the end of 10 ps MD calculations. These snapshots show that Ti$_3$BN monolayer can maintain its structural integrity throughout a 10 ps dynamical calculation up to 800 K, however will be disrupted at the temperature of 1000 K. Those results reveal that the Ti$_3$BN monolayer has good thermal stability and the melting point of Ti$_3$BN monolayer is between 800 K and 1000 K.

### 3.2 Global minimum search for Ti$_3$BN monolayer in 2D space

Although Ti$_3$BN monolayer has good stability based on the above results, the doubt about that the Ti$_3$BN monolayer is a local minimum or a global minimum needs to be solved. It’s well known that the global minimum structure is more likely to be realized experimentally. Therefore, we carried out a global search for the lowest energy structure of Ti$_3$BN monolayer in the 2D space by adopting the first-principles based particle-swarm optimization method. After 25 generations, three low-energy structures for 2D Ti$_3$BN were obtained, labelled as Ti$_3$BN-I, Ti$_3$BN-II, and Ti$_3$BN-III in the order of increasing energy.

As shown in Fig. 4, in which the relative energy per atom is presented, the global minimum structure is Ti$_3$BN-I, which is just the above discussed Ti$_3$BN monolayer. Interestingly, Ti$_3$BN-II is also crystallized in the space group $P31m$ (no. 156). The geometric construction, thickness, atomic layer distances of Ti$_3$BN-II is similar with that of Ti$_3$BN-I, while the biggest difference between them is the N-atomic layer just located above the B-atomic layer and the Ti$_3$-atomic layer above the Ti$_3$-atomic layer in Ti$_3$BN-II monolayer. The length of Ti$_3$-N, N–Ti$_2$, Ti$_2$–B, and B–Ti$_3$ bond in Ti$_3$BN-II monolayer is 2.043 Å, 2.202 Å, 2.254 Å, and 2.116 Å, respectively. Considering structure Ti$_3$BN-III, it is 0.057 eV per atom higher in energy than Ti$_3$BN-I. This high energy might be due to the more nonbonding electrons of Ti$_3$ atoms.

### 3.3 Electronic properties of Ti$_3$BN monolayer

Since Ti$_3$BN-I monolayer holds great potential to be realized in experiment, does it have intriguing properties and promising applications? To figure out this issue, we have studied the electrical properties of the Ti$_3$BN-I monolayer using the hybrid density functional calculations. Fig. 5 illustrates the band structure and partial density of state (PDOS) of Ti$_3$BN-I monolayer. Obviously, a conduction band and a valence band cross through the Fermi level, indicating that Ti$_3$BN monolayer has metallic properties. The PDOS analysis shows that mainly the Ti-3d states, especially the 3d states of Ti atoms from surface atomic layer, contribute to the high density of electron states around the Fermi level. Note that the high density of states near the Fermi level means available carriers which are beneficial to the high electric conductivity of the Ti$_3$BN monolayer.

![Fig. 2](image1.png)  
**Fig. 2** Phonon dispersion curves of (a) bare Ti$_3$BN monolayer; (b) Ti$_3$BN(OH)$_2$-IV monolayer; (c) Ti$_3$BNF$_2$-I monolayer.

![Fig. 3](image2.png)  
**Fig. 3** Top (upper) and side (lower) views of snapshots of the Ti$_3$BN monolayer equilibrium structures at the end of 10 ps MD simulations: (a) 500 K, (b) 800 K, and (c) 1000 K. Gray, pink and red balls represent Ti, N and B atoms, respectively.

![Fig. 4](image3.png)  
**Fig. 4** Side (upper) and top (lower) views of low-energy 2D structures of Ti$_3$BN obtained from the PSO calculations. For Comparison, the relative energy per atom is given. The blue dash lines denote a unit cell where $a$, $b$ represents the lattice constants. Gray, pink and red balls represent Ti, N and B atoms, respectively.
3.4 Surface modification of Ti3BN monolayer by OH and F termination

Moreover, the high DOS at the Fermi level mainly originated from Ti1 and Ti3 atoms indicates the high activity of surface Ti atoms, suggesting the feasibility in the surface modification and even in composite materials for Ti3BN monolayer. For example, our theoretical studies have shown that the electronic properties of Ti3BN can be tuned by surface functional groups (–OH, –F). Herein, four possible geometry structures of Ti3BN monolayer with hydroxylated, and fluorinated surfaces are considered, as presented in Fig. 6. The top views of Ti3BN(OH)2-I and Ti3BNF2-II configurations are depicted in Fig. S2.† In configuration I, all the –OH or –F groups are located above the hollow sites between the three neighboring C/B atoms or point directly toward the Ti2 atoms on both sides of Ti3BN monolayer. In configuration II, all the –OH or –F groups are oriented above the topmost sites of C and B atoms on the two sides of Ti3BN monolayer, respectively. Afterwards, we can view the symmetric arrangement of configuration III and IV as the combination of configuration I and II.

The structural stability of different Ti3BN(OH)2 and Ti3BNF2 configurations can be estimated by comparing their relative total energies. For Ti3BN(OH)2, configuration IV is energetically most favorable. Ti3BN(OH)2-IV is energetically lower than Ti3BN(OH)2-I, Ti3BN(OH)2-II and Ti3BN(OH)2-III by 0.047, 0.320, and 0.382 eV per unit cell, respectively. While for Ti3BNF2, configuration I is energetically most favorable, with its energy lower than that of Ti3BNF2-II, Ti3BNF2-III and Ti3BNF2-IV by 0.584, 0.507, and 0.054 eV per unit cell, respectively. Phonon dispersions of Ti3BN(OH)2-IV and Ti3BNF2-I has been further calculated to investigate their kinetic stability, as shown in Fig. 2(b) and (c). As expected, there are no imaginary frequencies in the phonon dispersion curves. The phonons at about 3654.56 cm⁻¹ and 3620.86 cm⁻¹ for Ti3BN(OH)2-IV should be dominated by the OH groups, and the phonons at about 744.73 cm⁻¹ for Ti3BNF2-I should be due to the F groups. These high-frequency phonons indicate the strong bond nature of the related bonds (Ti–O and Ti–F).

In comparison with bare Ti3BN, the OH or F-terminated Ti3BN monolayer have smaller lattice constants. With terminal groups, the bond lengths of N–Ti2 and Ti2–B shrink, while the bonds between the Ti1–N and B–Ti3 are elongated except the Ti1–N bonds of Ti3BNF2-I and Ti3BNF2-III. Those results imply that the surface groups strongly interact with the original Ti3BN block, in accordance with the corresponding phonon dispersions. For clearance, the calculated lattice constants and bond lengths are presented in Table S1.†

Though the Ti3BN monolayer is metallic, its hydroxylated or fluorinated derivatives may be narrow-gap semiconductors or metals, depending on the geometrical arrangements of surface F and OH groups. Seen from Fig. 7(a), for Ti3BN with surface OH and F termination...

Fig. 5 Band structure (left) and partial density of states (PDOS) of Ti3BN-I monolayer obtained from hybrid functional calculations. For reference, the Fermi level is set at 0 eV. Ti1 and Ti3 represent the Ti atoms from the surface Ti-atomic layer and Ti2 denotes the Ti atoms from inner Ti-atomic layer.

Fig. 6 Side views of optimized geometries of the hydroxylated and fluorinated Ti3BN monolayer: (a) Ti3BN(OH)2-I, Ti3BN(OH)2-II, Ti3BN(OH)2-III, Ti3BN(OH)2-IV. (b) Ti3BNF2-I, Ti3BNF2-II, Ti3BNF2-III, Ti3BNF2-IV. Gray, pink and red balls represent Ti, N and B atoms, respectively. The H, O, and F atoms are indicated by white, purple, and cyan balls.

Fig. 7 Band structures of the hydroxylated and fluorinated Ti3BN monolayer obtained from hybrid functional calculations: (a) Ti3BN(OH)2-I, Ti3BN(OH)2-II, Ti3BN(OH)2-III, Ti3BN(OH)2-IV. (b) Ti3BNF2-I, Ti3BNF2-II, Ti3BNF2-III, Ti3BNF2-IV.
groups, the most favorable Ti$_3$BN(OH)$_2$-IV has a semi-conducting character, with a direct band gap of 0.09 eV. Ti$_3$BN(OH)$_2$-II also shows the electronic properties of direct semiconductor, and the band gap is 0.16 eV. However, Ti$_3$BN(OH)$_2$-I and Ti$_3$BN(OH)$_2$-III is metallic. For Ti$_3$BN with F-termination (Fig. 7(b)), the most stable configuration of Ti$_3$BNF$_2$-I and the metastable configuration of Ti$_3$BNF$_2$-III are metals. On the contrary, the band structures of Ti$_3$BNF$_2$-II and Ti$_3$BNF$_2$-IV demonstrate their indirect semiconducting characters, with the band gap of 0.07 eV and 0.37 eV, respectively. Those theoretical results prove that the electronic structure of Ti$_3$BN monolayer can be modulated by varying the surface functional groups.

4. Conclusions

In summary, we designed a new inorganic 2D material of Ti$_3$BN monolayer following the strategy of “atomic transmutation” by performing DFT calculations. In Ti$_3$BN monolayer, the Ti-B and Ti-N bonds are strong and the geometry structure is similar with the reported MXenes. The high cohesive energy, elastic constants, and absence of imaginary phonon frequencies prove that the Ti$_3$BN monolayer possess dynamic and mechanical stability. Particularly, MD simulation results verify that Ti$_3$BN monolayer can maintain its structural stability up to at least 800 K. The PSO method revealed that Ti$_3$BN monolayer is the global minimum structure in 2D space, suggesting the possibility to realize Ti$_3$BN monolayer in experiment. What's more, the electronic properties of Ti$_3$BN monolayer can be modified by varying the surface functional groups and their geometrical configurations. Therefore, if Ti$_3$BN is realized, it may find wide applications, such as in electrics, composite materials, Li-ion batteries and so on.

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