The Potential Application of BAs for a Gas Sensor for Detecting SO$_2$ Gas Molecule: a DFT Study

Jian Ren*1, Weijia Kong2 and Jiaming Ni3

Abstract

Different atmospheric gas molecules (e.g., N$_2$, O$_2$, CO$_2$, H$_2$O, CO, NO, NO$_2$, NH$_3$, and SO$_2$) are absorbed on the pristine hexagonal boron arsenide (BAs) through density functional theory calculations. For each gas molecules, various adsorption positions were considered. The most stable adsorption depended on position, adsorption energy, charge transfer, and work function. SO$_2$ gas molecules had the best adsorption energy, the shortest distance for BAs surface in the atmospheric gas molecule, and a certain amount of charge transfer. The calculation of work function was important for exploring the possibilities of adjusting the electronic and optical properties. Our results presented BAs materials can be the potential gas sensor of SO$_2$ with high sensitivity and selectivity.

Keywords: Electronic structure, Density functional theory, Adsorption energy, Gas molecule, BAs

Introduction

BAs (hexagonal boron arsenide) is composed of groups III and V elements. The groups of III–V elements have excellent properties, such as excellent photoelectric properties, mechanical properties, and large band gap [1]. The promising potential applications of 2D materials [2–5] were well documented in recent studies [6–20]; these materials had been used to recognize various biomolecules [21, 22], pollutants [23, 24], and gas molecules [25, 26] to develop suitable sensing devices. We had found more and more the groups of III–V element materials, for example, BN, AlN, GaN, GaAs, and BP, and it has more and more studies for the gas molecules by theoretical calculation. Strak et al. [27] discovered AlN(0001) was a powerful catalyst for high-pressure-high-temperature synthesis of ammonia, and the work also confirmed the possibility of the efficient synthesis of ammonia at the AlN(0001) surface. Diao et al. [28] presented adsorption of H$_2$O, CO$_2$, CO, H$_2$, and N$_2$ on (10–10) surfaces of pristine and Zn-doped GaAs nanowires; the effect of the adsorption of CO$_2$ and N$_2$ on absorption coefficients was the largest. Cheng et al. [29] showed the adsorption of most gas molecules on pure BP and doped BP by first principle study and concluded that N-BP was more suitable as a gas sensor for SO$_2$, NO, and NO$_2$ due to the existence of the desorption process. Kamaraj and Venkatesan [30] studied the structure and electronic properties of the BAs by the DFT and LDA; although considerable progress had been made in the experimental synthesis and theoretical study of BAs, the results of BAs nanosheets endowed the system with promising applications in nanoelectronics and photovoltaics.

In this work, we firstly investigated the gas sensing properties to fully exploit the possibilities of BAs as gas sensors by density functional theory (DFT) calculations. We predicted the adsorption properties of atmospheric gases (e.g., CO$_2$, O$_2$, N$_2$, H$_2$O, NO, NO$_2$, NH$_3$, CO, and SO$_2$) on BAs based on first principle calculations. Our work demonstrated the apparent adsorption behavior, moderate charge transfers, and unique transmission characteristics of SO$_2$ adsorption on BAs. The results suggested that monolayer BAs possessed great potential for SO$_2$ sensing application.

* Correspondence: 916181396@qq.com
1School of Computer Science and Technology, Huaiyin Normal University, Chang Jiang West Road 111, Huaiian 223300, Jiangsu, China
Full list of author information is available at the end of the article

© The Author(s). 2019 Open Access This article is distributed under the terms of the Creative Commons Attribution 4.0 International License (http://creativecommons.org/licenses/by/4.0/), which permits unrestricted use, distribution, and reproduction in any medium, provided you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made.
Theory and Method of Simulations

The system was modeled as a 4 × 4 supercell of BAs and atmospheric gas molecules adsorbed onto it. In DMol$^3$ [31] calculation process, exchange-correlation factions were calculated within a general gradient approximate (GGA) with the Perdew-Burke-Ernzerhof (PBE) [32]. The Brillouin zone was sampled using a 5 × 5 × 1 Monkhorst-Pack k-point grid and Methfessel-Paxton smearing of 0.01 Ry. All the atomic structures were relaxed until the total energy and the Hellmann-Feynman force converged to 1.0 × 10$^{-5}$ eV and 0.06 eV/Å [33].

To evaluate the interaction between gas molecules and adsorption sheet surface, we calculated the adsorption energy ($E_{\text{ad}}$) of adsorbed systems, which was defined as:

$$E_{\text{ad}} = E_{\text{BAs+gas molecule}} - (E_{\text{BAs}} + E_{\text{gas molecule}})$$

where $E_{\text{BAs+gas molecule}}$ is the total energy of BAs-adsorbed system, $E_{\text{BAs}}$ is the energy of BAs, and $E_{\text{gas molecule}}$ is the energy of a gas molecule. All energies were calculated for optimized atomic structures. The charge transfer was investigated by Mulliken’s population analysis.

Result and Discussion

Three adsorption sites were considered for BAs in this work, namely top of a boron atom (B), the top of an arsenic atom (As), and the center of a hexagonal B-As (center), as indicated in Fig. 1a. We studied the presence of the atmosphere and found the best gas sensor.

First of all, the geometric structure of pristine BAs monolayer had been optimized, and as shown in Fig. 1b, BAs bond length was 1.967 Å. There was an indirect band gap of 1.381 eV to exhibit in the band structure of BAs sheet, which was better than that of the bulk structure. These values were in good agreement with the previously reported values (Fig. 2) [34, 35].

Meanwhile, we had analyzed the adsorption energy, the charge transfer, and the distance between the molecules and BAs surface. The final result was as shown in Table 1.

**N$_2$ adsorption:** Adsorption of N$_2$ gas molecule on BAs was studied for three configurations of N$_2$/BAs, viz. top side of the B atom, top side of As atom, and the center of a hexagonal ring above the BAs surface, and the nearest distance was 3.764 Å, 3.549 Å, and 3.65 Å and corresponding adsorption energy was −0.24 eV, −0.27 eV, and −0.24 eV, respectively. The center had the best adsorption energy and the most stable structure. The adsorption energy of N$_2$BAs was −0.24 eV, the charge transfer from BAs to N$_2$ gas molecule was 0.014e, and the distance of the N$_2$-BAs was 3.65 Å. Fig. 3a showed that there were many lines under the Fermi energy level, and the corresponding density of states had several peaks under the Fermi energy level. As shown in the figure, the N$_2$ gas molecule had four peaks, which had a certain influence on BAs, mainly from −5 to 0 eV, and had great contributions to the DOS. Overall, the effect of N$_2$ gas molecule adsorption on BAs was poor.

**O$_2$ adsorption:** O$_2$ gas molecule tended to adsorb on the central point. The adsorption energy of O$_2$/BAs was −0.35 eV, and the distance of the O$_2$-BAs was 2.90 Å. The total band structure and DOS for O$_2$/BAs were plotted in Fig. 3. It was obvious that an extra line crossed the zero point and reduced the band gap; O$_2$ gas molecule had a peak at −1 to 0 eV and had an effect on the density of states above the Fermi level. The population analysis for Mulliken charge transfer showed that −0.172e was transferred from BAs surface to the O$_2$ gas molecule, suggesting that O$_2$ gas molecule acted as an acceptor. In general, the O$_2$ gas molecule adsorption on BAs was better than N$_2$.

**CO$_2$ adsorption:** CO$_2$ gas molecule tended to adsorb on the top of As atom. The adsorption energy of CO$_2$/BAs was −0.28 eV, the charge transfer from BAs to CO$_2$...
gas molecule was $-0.018\text{e}$, and the distance of the CO$_2$-BAs was 3.55 Å. As shown in Fig. 3, compared to pristine BAs, the structure had no apparent change, and there were some obvious wave crests of the energy of $-9\text{ eV}$ in DOS, which had great contributions to the DOS. This point also highlighted the adsorption of CO$_2$ gas molecule by BAs. The results showed that the adsorption effect and sensitivity of BAs to CO$_2$ gas molecule were general.

**H$_2$O adsorption:** H$_2$O gas molecule tended to adsorb on the top of As atom. The adsorption energy of H$_2$O/BAs was $-0.38\text{ eV}$, the charge transfer from BAs to H$_2$O gas molecule was $-0.03\text{e}$, and the distance of the H$_2$O-BAs was 3.63 Å. As shown in Fig. 3, there were no great changes in the structure compared to pristine BAs. The Fermi level of Al-G increased obviously and moved to the valence band. In general, the H$_2$O gas molecule adsorption on BAs was ignored.

**CO adsorption:** CO gas molecule tended to adsorb on the top of As atom. The adsorption energy of CO/BAs was $-0.27\text{ eV}$, the charge transfer from BAs to CO gas molecule was $-0.024\text{e}$, and the distance of the CO-BAs was 3.50 Å. The total density of states (DOS) and band structure for BAs-CO were plotted in Fig. 3. CO gas
molecule and As atom played a huge role in the effect of a peak of 3 to 4 eV on the DOS. However, there was no deviation in DOS in −7 to 4 eV range, which suggested that CO was weakly adsorbed on BAs. There was some obvious wave crest of the energy of −3 to 1 eV and 3 eV, which had great contributions to the DOS. The population analysis for Mulliken charge transfer showed that −0.024e charge was transferred from BAs surface to the CO gas molecule, and it suggested that CO gas molecule acted as an acceptor. Overall, the effect of CO gas molecule adsorption on BAs was not special.

**NO adsorption:** NO gas molecule tended to adsorb on the top of B atom. The adsorption energy of NO/BAs was −0.18 eV, the charge transfer was −0.01e from NO gas molecule to BAs, and the distance of the NO-BAs was 2.86 Å. There were a lot of lines upon the Fermi energy level. It found that the energy gap in the middle band reduced the band gap value. From the density diagram of states, there was an extra wave peak above the Fermi energy level, but there was little change under the Fermi energy level, relatively stable in Fig. 3. The mixing

| System   | $E_{ad}$ (eV) | $Q$ (e) | $D$ (Å) | Style   |
|----------|--------------|--------|--------|---------|
| N\textsubscript{2}  | −0.24        | −0.014 | 3.65   | Acceptor |
| O\textsubscript{2}  | −0.35        | −0.0172| 2.90   | Acceptor |
| CO\textsubscript{2} | −0.28        | −0.018 | 3.55   | Acceptor |
| H\textsubscript{2}O  | −0.38        | −0.030 | 3.63   | Acceptor |
| CO        | −0.27        | −0.024 | 3.50   | Acceptor |
| NO        | −0.18        | 0.010  | 2.86   | Acceptor |
| NO\textsubscript{2} | −0.43        | −0.231 | 2.47   | Acceptor |
| NH\textsubscript{3} | −0.34        | 0.007  | 3.27   | Donor   |
| SO\textsubscript{2} | −0.92        | −0.179 | 2.46   | Acceptor |

**Fig. 3** Density of states (DOS) of N\textsubscript{2}/BAs (a), O\textsubscript{2}/BAs (b), CO\textsubscript{2}/BAs (c), H\textsubscript{2}O/BAs (d), CO/BAs (e), NO/BAs (f), NO\textsubscript{2}/BAs (g), NH\textsubscript{3}/BAs (h) and SO\textsubscript{2}/BAs (i).
of orbitals caused small charge transfer and redistribution over the interacting region. The population analysis for Mulliken charge transfer showed that 0.01e charge was transferred from BAs surface to the NO molecule, suggesting that NO acted as a donor. There was no deviation in DOS in −7 to 4 eV range, which suggested that NO was weakly adsorbed on BAs.

**NO2 adsorption:** NO2 gas molecule tended to adsorb on the top of As atom. The adsorption energy of NO2/BAs was −0.43 eV, and the distance of the NO2-BAs was 2.47 Å. The interesting was that the zero point in the band crossed a straight line directly after the adsorption of NO2 gas molecule, which meant that the BAs, which is a semiconductor, was transformed into the gold attribute; band gap was 0 eV. There was no great change in the whole, and a peak was generated at about −3 eV due to NO2 gas molecular adsorption. There was some obvious wave crest of the energy of −7 eV and 2 eV, which had great contributions to the DOS. In general, the adsorption of NO2 by BAs was better than that of several molecules above.

**NH3 adsorption:** NH3 gas molecule tended to adsorb on the top of As atom. The adsorption energy of NH3/BAs was −0.34 eV, the charge transfer from NH3 gas molecule to BA was 0.007e, and the distance of the NH3-BA was 3.27 Å. There was no clear change in the energy band and the density of states, except that there was an obvious peak of adsorption of NH3 gas molecule below the Fermi level. The NH3 gas molecule had a little impact on BAs at −8 to −4 eV, forming a 15 eV peak. The adsorption effect and sensitivity of its BAs to NH3 gas molecule were general.

**SO2 adsorption:** SO2 gas molecule tended to adsorb on the central point, the adsorption energy of SO2/BAs was −0.92 eV, and the population analysis for Mulliken charge transfer showed that −0.179e charge was transferred from BAs surface to the SO2 gas molecule, suggesting that SO2 gas molecule acts as an acceptor. The distance of the SO2/BAs was 2.46 Å. Compared to other gas molecules, the SO2/BAs had the biggest adsorption energy, the second largest electron transfer, and the shortest distance of the SO2-BAs. As shown in Fig. 3, the valence band of BAs had an obvious up and band gap decreased, and due to the adsorbed SO2 gas molecule, it could be seen from the density of states that there was one more wave peak at −7.5 eV and certain transfer at the Fermi level. The adsorption of SO2 by BAs had the excellent effect.

Fig. 4i showed the electron density diagram of SO2/BAs and the electron local overlap between BAs and SO2 gas molecule. On this basis, we drew the conclusion that the adsorption of SO2 by BAs was physical adsorption. The calculation of WF shown in Fig. 5 was of great significance in exploring the possibility of regulating the electronic and optical properties (such as absorption spectra and energy loss functions) by adsorbing small molecules. The work function was defined in solid physics.
as the minimum energy required to move an electron from the interior of a solid to the surface of the object. The work function of pristine BAs was 4.84 eV. NO and NH₃ gas molecules were donors in charge transfer, and their work function decreased; the work function was 4.80 eV and 4.68 eV, respectively. The work function of N₂/BAs, CO₂/BAs, and CO/BAs was the same as that of BAs. The work function of O₂/BAs, NO₂/BAs, and SO₂/BAs was higher than BAs. Combined with the above adsorption energy, distance of gas molecules and BAs surface, charge transfer, and work function, we found that SO₂ gas molecule was most suitable for BAs materials.

**Conclusion**

We have presented the structural and electronic properties of BAs with adsorbents N₂, O₂, CO₂, H₂O, CO, NO, NO₂, NH₃, and SO₂ gas molecule, using density functional theory method. In the adsorption energy, SO₂ > NO₂ > H₂O > O₂ > NH₃ > CO₂ > CO > N₂ > NO and SO₂ < NO₂ < NO<CO₂ < NH₃ < CO < CO₂ < H₂O < N₂ in the adsorption distance. NO₂ has the largest Q and work function, maybe it could be detected by the proposed material because of good electrical response. SO₂ gas molecule had the best adsorption energy, the shortest distance for gas molecule and BAs surface, and a certain amount of charge transfer. Combined with the above adsorption energy, distance of gas molecule and BAs surface, charge transfer, and work function, the current and the adsorption-induced current change of BAs exhibit strong anisotropic characteristics. Such sensitivity and selectivity to SO₂ gas molecule adsorption make BAs a desirable candidate as a superior gas sensor.

**Abbreviations**

BAs: Hexagonal boron arsenide; DOS: Density of states; WF: Work function

**Acknowledgements**

We thank the College of Materials Science and Engineering, Anhui University of Science and Technology, for its assistance with the MD simulations.

**Funding**

The financial support for this work was from the Natural Science Fund for Colleges and Universities in Jiangsu Province (17KJB510007).

**Availability of Data and Materials**

All data are fully available without restriction.

**Authors’ Contributions**

JR and WK designed and carried out the experiments and drafted the manuscript. JMN participated in the work to analyze the data and gave the materials and supporting software. All authors read and approved the final manuscript.

**Authors’ Information**

Not applicable

**Competing Interests**

The authors declare that they have no competing interests.

**Publisher’s Note**

Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

**Author details**

¹School of Computer Science and Technology, Huaiyin Normal University, Chang Jiang West Road 111, Huaian 223300, Jiangsu, China. ²Department of Chemistry, Beijing Normal University, No.19, Waidajie, Xinjiekou, Haidian District, Beijing 100875, China. ³School of Mechanical and Electrical Engineering, Guilin University of Electronic Technology, Jinji Road No.1, 54100 Gui, China.
