Sensing Mechanism of H₂O, NH₃, and O₂ on the Stability-Improved Cs₂Pb(SCN)₂Br₂ Surface: A Quantum Dynamics Investigation

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ABSTRACT: Although the perovskite sensing materials have shown high sensitivity and ideal selectivity toward neutral, oxidative, or reductive gases, their structural instability hampers the practical application. To exploit perovskite-based gas-sensing materials with improved stability and decent sensitivity, three adsorption complexes of H₂O, NH₃, and O₂ on the Cs₂Pb(SCN)₂Br₂ surface are built by doping Br⁻ and Cs⁺ in the parent (CH₃NH₃)₂Pb(SCN)₂I₂ structure and submitted to quantum dynamics simulations. Changes in the semiconductor material geometric structures during these dynamic processes are analyzed and adsorption ability and charge transfer between Cs₂Pb(SCN)₂Br₂ and the gas molecules are explored so as to further establish a correlation between the geometrical structure variations and the charge transfer. By comparing with the previous CH₃NH₃PbI₃ and (CH₃NH₃)₂Pb(SCN)₂I₂ adsorption systems, we propose the key factors that enhance the stability of perovskite structures in different atmospheres. The current work is expected to provide clues for developing innovative perovskite sensing materials or for constructing reasonable sensing mechanisms.

1. INTRODUCTION

The metal–organic halide perovskite ABX₃ has attracted much interest due to its excellent photovoltaic properties.¹⁻⁵ At present, the power conversion efficiency (PCE) of solar cells based on ABX₃ structures has exceeded 25.5%;⁶ however, the structural instability of perovskite materials hinders their further development and application in solar cells.⁷⁻¹⁰ Zhao¹¹ and Bao¹² et al. used CH₃NH₃PbI₃ to explore the gas-sensing properties of NH₃, which usher in an era of applying perovskite materials in gas sensing. Since then, the detection limit of O₂ concentration has been reduced to as low as 70 ppm based on CH₃NH₃PbI₃ films,¹³ while CH₃NH₃PbI₃−ₓClₓ has been proved to be able to detect an ultralow ozone concentration of few ppb.¹⁴ These typical experiments have proved the extraordinary sensitivity of the perovskite materials to gases with different reductive or oxidative properties.¹¹⁻¹⁸ Differing from the application in solar cells, the range for designing stable perovskite materials for gas sensing can break through the strict limitations on band gaps of photovoltaic materials and hence the interest in the development of such perovskite materials has been triggered.

Despite the extraordinary sensitivity or selectivity of CH₃NH₃PbI₃ to gases with various physical or chemical properties, it has been found that the sensing material could be damaged when putting in NH₃ (reductive), H₂O (neutral), and O₂, NO₂, or O₃ (oxidative) environment. The stability improvement of the gas sensors based on perovskite materials, therefore, remains a major challenge. Our previous quantum dynamics simulations show that Pb²⁺ with an intermediate valence state exhibits both reduction and oxidation ability, thus it offers active sites that are attacked by both oxidizing and reducing gases, and the weak Pb—I framework fails to resist attack by strong oxidative or reductive gases. Further experiments have proved that (CH₃NH₃)₂Pb(SCN)₂I₂, obtained through the partial substitution of the X-site elements in CH₃NH₃PbI₃, can improve the perovskite material moisture resistance.¹⁰ In the simulations of (CH₃NH₃)₂Pb(SCN)₂I₂ adsorbing gases with different properties, i.e., neutral H₂O, reductive NH₃, and oxidative NO₂, O₂, and O₃, it is found that what underpins the mechanism of the SCN⁻ groups effectively enhancing the stability of perovskite skeletons is that both the S atoms and the CN groups of SCN⁻ can stably bond with Pb. Therefore, the structural stability in the dynamics process is maintained in the form of Pb—NCS or Pb—SCN—Pb network structures. However, the weak Pb—I bonds in the structure are still the targets of oxidizing or reducing gases. It has been found that both the neutral H₂O and oxidative gases (NO₂, O₂, and O₃) can form a stable Pb—O connection by directly attacking the Pb—I bonds.¹⁰ Anyway, in the study of the adsorption of reductive NH₃ by CH₃NH₃PbI₃ it is found...
that the A-site CH$_3$NH$_3^+$ groups rapidly adsorb NH$_3$ and undergo H proton exchanges, thus blocking the interactions between the gas molecules and the skeletons; nevertheless, NH$_3$ can eventually break through the CH$_3$NH$_3^+$ barrier and interact with Pb to achieve stable adsorption and charge exchange. In addition, during the adsorption process, it has been confirmed that the perovskite materials normally receive charges from reductive gases but donate charges to oxidative gases, which is believed to be related to the electron structural changes of the semiconducting materials.$^{21}$

In recent years, the substitution of the original A-site organic groups with inorganic elements or groups to improve the perovskite stability has also been utilized in gas sensing. Experiments have proved that Cs$_2$Pb(SCN)$_2$I$_2$ exhibits enhanced ambient stability compared with (CH$_3$NH$_3$)$_2$Pb(SCN)$_2$I$_2$. In addition, CsPbBr$_3$ has been used to detect...
volatile organic compounds (acetone and ethanol) with a detection limit as low as 1 ppm. In addition, it also provides a fast response and rapid recovery in the detection of O2. The excellent stability of CsPbBr3 in air and oxygen has also been verified.

Investigating the gas-sensing mechanisms is the first step leading to the development of ideal sensing materials, while research on adsorption and desorption of gas molecules on the perovskite material surface is the starting point for probing gas-sensing mechanisms. According to Yamazoe et al., the initial move in the sensing process—gas molecules is to calibrate gas molecules and their active sites where the gas operates via the electronic changes introduced by gas—solid interactions. Our previous work shows that the structural instability of the perovskite materials causes different dynamic changes around the adsorption sites. Such a property challenges the traditional static first-principles calculations, therefore, the quantum dynamics methodology is employed to deal with such special properties.

In the current work, to improve the structural stability of CH3NH3PbI3 and to clarify the sensing mechanism origin of the perovskites toward gases with different oxidation—reduction characteristics, the original organic CH3NH3+ groups are replaced by inorganic Cs+ and the weak Pb—I bonds are substituted by Pb—SCN and Pb—Br. The previously simulated typical neutral (H2O), reductive (NH3), and oxidative (O2) gases are reselected to form three complexes of Cs2Pb(SCN)2Br2—H2O, Cs2Pb(SCN)2Br2—NH3, and Cs2Pb(SCN)2Br2—O2, respectively. All of the systems are submitted to dynamics simulations. The structural stability of Cs2Pb(SCN)2Br2 under the attack of strong reductive, oxidative, and neutral gas molecules, is described in detail via monitoring the bond breaking and formation during the dynamics processes. The adsorption energy and charge transfer between the perovskite materials and the gases are both quantified. The adsorption details of H2O, NH3, and O2 with CH3NH3PbI3 and (CH3NH3)2Pb(SCN)I2 are also recalled and compared with systems in the current work. Based on the comparisons, a relationship between the geometrical structure changes of the semiconductor materials and the charge transfer is established. The purpose of this research is to reveal the adsorption—desorption processes and the stability enhancement mechanisms of Cs2Pb(SCN)2Br2 in an environment containing gases bearing different properties, so as to provide new ideas for developing perovskite sensing materials with improved stability and decent sensitivity.

2. RESULTS AND DISCUSSION

The root mean square deviations (RMSDs) of all of the complexes are demonstrated in Figure 1. The RMSD curves of the Cs2Pb(SCN)2Br2—H2O, Cs2Pb(SCN)2Br2—NH3, and Cs2Pb(SCN)2Br2—O2 systems tend to be stable at 125, 95, and 145 ps, respectively, indicating that the adsorption of the gas molecules on the perovskite skeletons reaches an equilibrium. It takes different time spans for the RMSD curves of the three systems to obtain an equilibrium, and the fluctuations of which are distinct as well, which might suggest that gases with different chemical properties exhibit different adsorption behaviors. The RMSD of the atoms on the perovskite material surface, including Pb, Br, S, C, and N, have also been extracted and described in Figure 1. From the figure, it is clear that the fluctuations of S, C, and N on the surface are larger than those of the atoms inside the skeleton. This is in agreement with the structural stability analyses below, where the —SCN groups on the surface stick out to and are disturbed by the gas molecules to form new Pb—SCN—Pb connections.

2.1. Adsorption Properties of Water Molecules on the Cs2Pb(SCN)2Br2 Surface. The simulations show that, compared with those of the CH3NH3PbI3 and (CH3NH3)2Pb(SCN)I2 systems, the stability of Cs2Pb(SCN)2Br2 is further improved in a humid environment. The study of the adsorption of H2O on CH3NH3PbI3 and (CH3NH3)2Pb(SCN)I2 demonstrates that the breakage of the weak Pb—I bonds in the dynamic process is the main reason for the distortion and collapse of the perovskite skeletons under the attack of water molecules. Therefore, in the current work, the changes in the Pb—Br bonds in Cs2Pb(SCN)2Br2 are first investigated. In the current dynamics process, a Pb—I connection is considered broken when the bond length is stretched longer than 3.4 Å and the threshold is 3.25 Å for a Pb—Br connection. At the end of the simulations, there are 21 intact Pb—Br bonds, out of the initial 32 bonds in the system, and the integrity rate is 65.63% (21/32). The integrity rates of Pb—I bonds in the (CH3NH3)2Pb(SCN)I2 and CH3NH3PbI3 systems are 62.50% (20/32) and 58.33% (28/48), respectively. Therefore, the substitution of the original...
Pb–I bonds with stronger Pb–Br connections can effectively decrease the broken Pb–X bond proportion, thus reducing the opportunity for the exposed metal ions to connect with H₂O molecules. Anyway, a small portion of the Pb–Br bonds are still broken under the attack of H₂O molecules, and H₂O is found to attack Pb atoms and form Pb₂−O₁, Pb₇−O₂₁, Pb₇−O₁₆, Pb₆−O₉, Pb₆−O₂₄, Pb₅−O₂₃, and Pb₅−O₅ connections (Figure 2a). The distances between these Pb and O atoms are mostly slightly longer than the Pb−O bond length (2.49 Å), indicating that the interactions between Pb−OH₂ are weaker than that of normal Pb−O bonds (Figure 3a). In the simulations of CH₃NH₃PbI₃−H₂O and (CH₃NH₃)₂Pb(SCN)₂I₂−H₂O, the release of reactants CH₃NH₃I and CH₃NH₃SCN is observed at 5 and 94 ps (Figure 4), respectively; no reactant is, however, observed in the Cs₂Pb(SCN)₂Br₂−H₂O system. This implies that the doping of SCN⁻ and Br⁻ is conducive to improving the perovskite structural stability.

Furthermore, it has been proved that the formation of stable Pb−NCS bonds and Pb−SCN−Pb networks in the (CH₃NH₃)₂Pb(SCN)₂ system is the crucial reason to effectively maintain the stability of perovskite structures. In the Cs₂Pb(SCN)₂Br₂−H₂O complex, it is also noticed that SCN⁻ groups rotate in the dynamic process to form Pb−NCS bonds (Figure 2b). In view of this phenomenon, the structures of Cs₂Pb(SCN)₂Br₂ and Cs₂Pb(NCS)₂Br₂ are both optimized, and the formation energies are −124.475 eV and −124.050 eV, respectively (Figure 5). These energies are very similar, revealing that both connections can exist stably. Furthermore, Pb₁−(SCN)₄, Pb₆−(SCN)₁₁, and Pb₂−(SCN)₃ connections are successively observed (Figure 3b). Moreover, the Pb₄−(SCN)₇−Pb₂ network is also detected at 19 ps (Figure 2c).

The stability improvement mechanisms of the perovskite structure by replacing the polar stick-like CH₃NH₃⁺ groups with the nonpolar spherical CS⁺ atoms have been proved previously. The rotation of the former is shown to damage the weak Pb−I bonds in three-dimensional (3D) skeletons to a certain extent. Li et al. suggested that when CS⁺ is inserted into a two-dimensional (2D) (PbX₄(SCN)₂) framework, smaller Br⁻ that is more strongly bonded with Pb²⁺ can be used, without serious lattice distortion in air, which is also verified by our simulation results. The comparison of the dynamic fluctuations of the three systems shows that the amplitude of fluctuation of Cs⁺ in the Cs₂Pb(SCN)₂Br₂−H₂O system is much smaller than that of CH₃NH₃⁺ in the CH₃NH₃⁺Pb₁−H₂O and (CH₃NH₃)₂Pb(SCN)₁−H₂O complexes (Figure S1). In addition, the increased amplitudes of the RMSD curves of Pb and Br atoms are smaller than those of Pb and I in the CH₃NH₃⁺Pb₁−H₂O and (CH₃NH₃)₂Pb-
(SCN)2I2−H2O systems, indicating that the substitution of Cs+ can stabilize such materials.

2.2. Adsorption Properties of NH3 on the Cs2Pb-(SCN)2Br2 Surface. The dynamics results show that without the blockage of CH3NH3+, NH3 can rapidly and directly attack Pb atoms on the Cs2Pb(SCN)2Br2 surface, which is the biggest difference between the Cs2Pb(SCN)2Br2−NH3 system and the previous CH3NH3PbI3−NH3 system. On the CH3NH3PbI3 surface, NH3 molecules do not stably bond with Pb atoms until 150 ps. Meanwhile, the CH3NH3+ groups, the products of CH3NH3+−NH3 proton exchange, attack Pb atoms to form a Pb−N connection, which survives the rest of the simulations. These Pb−NH2CH3 bonds rotate continuously to break the nearby Pb−I bonds, which directly leads to the damage of the Pb−I skeleton. The reactant CH3NH3I has been observed at 5 ps, as seen in Figure 6. In the current system, the bond length of Pb5−N17 reaches 2.41 Å at 20 ps (Figure 7a), which is smaller than the standard Pb−N bond length of 2.91 Å. After that, stable connections of Pb3−N36, Pb6−N23, Pb8−N35, and Pb6−N30 are successively generated at 21, 29, 51, and 60 ps, respectively (Figure 8). Pb6−N30 is formed at 60 ps and remains stable until the end of the simulation, and no new Pb−

N connections are observed. The Pb−NH2 bonding does not rotate like Pb−NH2CH3 in the CH3NH3PbI3−NH3 complex, while the Pb−Br bonds are only partially broken. Therefore, the octahedrons in the perovskite skeletons remain intact without serious distortion or collapse (Figure 9b). At the end of the simulations, the integrity rate of the Pb−Br bonds in the Cs2Pb(SCN)2Br2−NH3 system is 71.88% (23/32), while that in the CH3NH3PbI3−NH3 complex is 56.25% (27/48).

In addition, both the Pb−NCS and Pb−SCN−Pb networks have been observed in the Cs2Pb(SCN)2Br2−NH3 system (Figure 7b), which also make contributions to improving the stability of the perovskite structures.

2.3. Adsorption Properties of O2 on the Cs2Pb-(SCN)2Br2 Surface. In contrast to H2O and NH3 that can form stable interactions with exposed lead atoms, the O2 molecules on the perovskite surface are found to be frequently adsorbed and desorbed. As shown in Figure 10a, the distance

Figure 6. Release of the reactant CH3NH3I in the CH3NH3PbI3−NH3 complex.

Figure 7. Typical snapshots of the Cs2Pb(SCN)2Br2−NH3 complex in the current work: (a) Pb and NH3 connection and (b) formation of Pb−NCS and Pb−SCN−Pb networks.
of the Pb6-O7 connection is 2.59 Å at 83 ps, which is longer than the Pb−O bond length of 2.49 Å, suggesting a weak interaction between Pb6 and O7 (Figure 11a). Such adsorption properties of O2 are also observed on the surfaces of CH3NH3PbI3 and (CH3NH3)2Pb(SCN)2I2; however, in these two systems, the O2 molecules manage to enter the perovskite lattices by attacking and breaking the Pb−I bonds to temporarily bond with Pb2+, which severely damages the material structures (Figure S2). In this study, O2 still cracks the Pb−Br bonds. At the end of the simulations, the integrity ratio of Pb−Br connections is 62.50% (20/32), which is slightly higher than that of 58.33% (28/48) in the CH3NH3PbI3−O2 system and 43.75% (14/32) in the (CH3NH3)2Pb(SCN)2I2−O2 system. This indicates that the Pb−Br bonds are still not

Figure 9. Starting and final structures of the adsorption systems: (a) Cs2Pb(SCN)2Br2−H2O, (b) Cs2Pb(SCN)2Br2−NH3, and (c) Cs2Pb(SCN)2Br2−O2.

Figure 10. Typical snapshots of the Cs2Pb(SCN)2Br2−O2 complex: (a) weak Pb−O2 connection and (b) Pb−NCS and Pb−SCN−Pb networks.
strong enough under strong oxidizing gas attacks. Despite this, in the Cs₂Pb(SCN)₂Br₂−O₂ system, the O₂ molecules are blocked and reside on the surface of perovskite materials and cannot enter the internal of the lattice throughout the simulations (Figure 9c), which is attributed to the contribution of the Pb−SCN (Figure 10b).

At the end of the dynamics process of the Cs₂Pb(SCN)₂Br₂−O₂ system, 11 out of 16 Pb−SCN transform to Pb−NCS connections, that is, Pb₁−(NCS)₁, Pb₂−(NCS)₂, Pb₂−(NCS)₃, Pb₁−(NCS)₄, Pb₃−(NCS)₅, Pb₄−(NCS)₆, Pb₄−(NCS)₇, Pb₆−(NCS)₁₀, Pb₈−(NCS)₁₅, Pb₇−(NCS)₁₆, and Pb₇−(NCS)₁₃ (Figure 11b). In the Cs₂Pb(SCN)₂Br₂−H₂O and Cs₂Pb(SCN)₂Br₂−NH₃ complexes, the number of Pb−NCS is 1 and 8, respectively. The reason of which is further analyzed in the following charge transfer section. In addition, the Pb₆−(SCN)₁₁−Pb₈, Pb₅−(SCN)₁₂−Pb₁, and Pb₄−(SCN)₃−Pb₂ networks are observed (Figure 11c, the normal bond length of Pb−S is 3.70 Å⁴³). It is these special forms that are deemed to prevent O₂ molecules from entering the perovskite lattice, thus ensuring the structural integrity of the semiconductor materials.

Figure 11. Evolution of the newly formed connections in the Cs₂Pb(SCN)₂Br₂−O₂ complex: (a) interaction of Pb−O, (b) connection of Pb−NCS, and (c) formation of Pb−SCN−Pb networks.

Figure 12. Adsorption energies of (a) Cs₂Pb(SCN)₂Br₂−H₂O, (b) Cs₂Pb(SCN)₂Br₂−NH₃, and (c) Cs₂Pb(SCN)₂Br₂−O₂ complex.
Furthermore, considering that the total number of H$_2$O$^-$ is relatively lower than the above discussed two complexes, which dynamic process, the adsorption energy of the system is the three systems.

The following quantitative data of the adsorption energy and charge transfer are used to depict the adsorption properties of the three systems.

2.4. Adsorption Ability of H$_2$O, NH$_3$, and O$_2$ on the Cs$_2$Pb(SCN)$_2$Br$_2$ Surface. Figure 12 and Table S1 demonstrate the data of the adsorption energies of the Cs$_2$Pb(SCN)$_2$Br$_2$--H$_2$O, Cs$_2$Pb(SCN)$_2$Br$_2$--NH$_3$, and Cs$_2$Pb(SCN)$_2$Br$_2$--O$_2$ complexes. The negative adsorption energy ($E_{ad}$) implies the reactions are exothermic and spontaneous. This indicates that the gas molecules are adsorbed on the Cs$_2$Pb(SCN)$_2$Br$_2$ surface; the larger adsorption energy suggests that more energy is released during the reaction and more stable connections are established. For the Cs$_2$Pb(SCN)$_2$Br$_2$--NH$_3$ system, in the initial phase of the dynamics process, the NH$_3$ molecules attack Pb atoms to form a stable Pb--N connection; however, the presence of Pb--SCN and Pb--NCS connections on the surface prevents more NH$_3$ molecules from attacking Pb atoms and the average adsorption energy of the system in the equilibrium stage is $-57.611$ eV. For the Cs$_2$Pb(SCN)$_2$Br$_2$--H$_2$O system, Pb atoms are still active sites that are first attacked by the H$_2$O molecules. Although the number of the newly formed Pb--OH$_2$ connections is larger than that of the Pb--N bonds mentioned above, the average adsorption energy of the system in the equilibrium stage is slightly lower, $-53.678$ eV. This is mainly due to the relatively weak Pb--OH$_2$ connections as discussed in Section 2.1. Furthermore, considering that the total number of H$_2$O molecules is 25 and that of NH$_3$ is 20, the adsorption ability of a single NH$_3$ is stronger than that of a single H$_2$O molecule. In contrast, the Cs$_2$Pb(SCN)$_2$Br$_2$--O$_2$ system is more special. Due to the frequent adsorption--desorption behaviors of O$_2$ in the dynamic process, the adsorption energy of the system is relatively lower than the above discussed two complexes, which is $-2.781$ eV. This phenomenon seems to fulfill the requirements of an ideal gas-sensing system, where the gas molecule under detection is not only easily adsorbed but also easily released from the sensing material surface.

2.5. Charge Transfer between Cs$_2$Pb(SCN)$_2$Br$_2$ and H$_2$O, NH$_3$, and O$_2$. Charge transfer is one of the most crucial factors influencing the resistivity of sensing materials. The calculated Bader charges are shown in Figure 13 and Table S2. In the initial stage, the quantity of the charge transfers from H$_2$O to Cs$_2$Pb(SCN)$_2$Br$_2$ is 0.616 e. In this case, both Pb$^{2+}$ ($-4.869$ e) and Cs$^+$ ($-9.870$ e) in the Cs$_2$Pb(SCN)$_2$Br$_2$--H$_2$O system lose charges. The charges gained by SCN$^-$ and Br$^-$ are $7.967$ e and $7.352$ e, respectively. At the end of the dynamics process, the charge transferred from H$_2$O to Cs$_2$Pb(SCN)$_2$Br$_2$ is 1.647 e, and those of Pb$^{2+}$, Cs$^+$, SCN$^-$, and Br$^-$ are $-4.954$ e, $-9.913$ e, $9.253$ e, and $7.261$ e, respectively. Therefore, during the dynamics processes, the donors in the Cs$_2$Pb(SCN)$_2$Br$_2$--H$_2$O system are H$_2$O($-1.031$ e), Pb$^{2+}$($-0.085$ e), Cs$^+$($-0.043$ e), and Br$^-$($-0.091$ e), respectively, while the acceptor is SCN$^-$($1.286$ e). Similarly, for the Cs$_2$Pb(SCN)$_2$Br$_2$--NH$_3$ system, the donors are NH$_3$, Pb$^{2+}$, and Cs$^+$, donating $-0.848$ e, $-0.141$ e, and $-0.088$ e, respectively. The acceptors are SCN$^-$ and Br$^-$, respectively, accepting $0.370$ e and $0.714$ e. In the Cs$_2$Pb(SCN)$_2$Br$_2$--O$_2$ complex, the donors are Pb$^{2+}$ and Br$^-$, which separately provide $-0.209$ e and $-0.708$ e, respectively. The acceptors are O$_2$, Cs$^+$, and SCN$^-$, accepting $0.824$ e, $-0.150$ e, and $0.244$ e, respectively.

Pb$^{2+}$ has both oxidizing and reducing properties. In the current work, Pb$^{2+}$ loses charge, showing different degrees of reducibility, which may be related to the strong electron-withdrawing halogen or the pseudo-halogen groups connected with Pb$^{2+}$. In view of the formation of the multiple Pb--NCS connections and the strong electron-withdrawing capacity of the --CN groups, the relationship between the charge loss of Pb atoms and the number of Pb--NCS connections is proved, as shown in Table 1 and Figure 14. The data show that there is an obvious linear correlation between them in all three systems. Based on the above analysis, Pb$^{2+}$ is a crucial active site and the gas molecules--Pb$^{2+}$--SCN$^-$ are the main charge-transfer channels. Combined with the above structural stability analysis, although the stability of Pb--SCN and Pb--NCS is similar, considering the stronger electron-withdrawing properties of the --CN groups, more Pb--NCS connections indicate stronger electron-accepting ability during...
the adsorption of various gas molecules. In the three systems, with the increase in the number of Pb−NCS bonds, Pb$^{2+}$ loses more charge, showing stronger oxidizability. For the neutral H$_2$O and reductive NH$_3$, the systems receive charges. In terms of the strong oxidizing O$_2$, these systems still lose charges.

2.6. Influences on the Band Gap of Cs$_2$Pb(SCN)$_2$Br$_2$ from the Adsorption of H$_2$O, NH$_3$, and O$_2$. The band gap values of Cs$_2$Pb(SCN)$_2$Br$_2$ before and after the adsorption of the gas molecules were calculated and are listed in Table 2. To clearly show the band gap tuning of the semiconductor materials from gas sensing, the structures, in which the stable connections between the gas molecules and the surface of Cs$_2$Pb(SCN)$_2$Br$_2$ were formed (Figure S3), were selected and calculated. The band gap value of 0.794 eV refers to the pure perovskite material. After adsorption, as seen in Figure 13 and Table 2, when reductive NH$_3$ or neutral H$_2$O molecules were involved, Cs$_2$Pb(SCN)$_2$Br$_2$ receives charge from NH$_3$ or H$_2$O, the band gap increases; when oxidative O$_2$ molecules were absorbed, Cs$_2$Pb(SCN)$_2$Br$_2$ losses charge, the band gap decreases. It is well known that organometallic halide perovskites are ambipolar charge transporters due to the comparable effective masses of the electrons and holes. The adsorption of both the electron-donating (NH$_3$ and H$_2$O) and the electron-withdrawing (O$_2$) molecules can increase the electron- and hole-doping level of Cs$_2$Pb(SCN)$_2$Br$_2$, causing the materials to behave as n- or p-type semiconductors.

3. CONCLUSIONS
In the current work, the influences of the adsorption of typical neutral H$_2$O, reductive NH$_3$, and oxidative O$_2$ on the surface of Cs$_2$Pb(SCN)$_2$Br$_2$ on the perovskite geometric structures are investigated. The adsorption energy and charge-transfer processes between the gas molecules and semiconductor materials are calculated as well. The main conclusions are as follows:

1. The Br$^-$ doping, the Pb−NSC connection formation, and the replacement of the polar and stick-like CH$_3$NH$_3^+$ groups with the nonpolar and spherical Cs$^+$ cations together can effectively improve the structural stability of the perovskite materials. The quick adsorption of the gas molecules on the Cs$_2$Pb(SCN)$_2$Br$_2$ surface and the fast charge transfer between them indicate ideal molecular recognition and, therefore, potentially high sensitivity to the three types of gas. When put in a humid or reductive gas environment, sensors based on Cs$_2$Pb(SCN)$_2$Br$_2$ are expected to present enhanced stability; however, under the attack of strong oxidizing gas molecules, the Pb−Br bonds are still not strong enough, which is a problem that warrants attention in the design of perovskite materials sensitive to strong oxidizing gases.

2. Although the simulation time of 200 ps is still too short compared with the actual adsorption−desorption process, H$_2$O and NH$_3$ can be stably adsorbed on the surface of Cs$_2$Pb(SCN)$_2$Br$_2$ in the dynamics simulations, and no desorption is observed. The electrons flow from the gases to the semiconductor materials. O$_2$ is...
frequently adsorbed or desorbed, and the charge flows from the semiconductor materials to O₂. Based on these two different adsorption properties, the desorption properties of O₂ on the Cs₂Pb(SCN)₂Br₂ surface are more favorable to improving the recovery time of gas sensors.

(3) The simulations suggest when put in a humid or reductive gas environment, sensors based on Cs₂Pb(SCN)₂Br₂ are expected to present enhanced stability; however, under the attack of strong oxidizing gas molecules, the sensors can show high sensitivity but less stability than those in the neutral or reductive gases. This is because the Pb−Br bonds are still not strong enough, which is a problem that warrants attention in the future design of perovskite materials sensitive to strong oxidizing gases.

4. THEORETICAL METHODS

The modeling structure of Cs₂Pb(SCN)₂Br₂ was first optimized by Vienna ab initio simulation package (VASP) 46, which is built on the parent perovskite material (MA)₂Pb(SCN)₂H₂O crystal structure. Slabs of 2*2 from the bulk structure of Cs₂Pb(SCN)₂Br₂ were cut for Cs₂Pb(SCN)₂Br₂−H₂O, Cs₂Pb(SCN)₂Br₂−NH₃, and Cs₂Pb(SCN)₂Br₂−O₂ systems. The Cs⁺-terminated (001) 48 perovskite surface was employed as the gas-exposed surface and a 5 Å vacuum space was set along the c-direction, which was filled with 25 H₂O, 20 NH₃, and 20 O₂ molecules, respectively (Figure 15). The cell lattice parameters a = 12.05 Å, b = 12.55 Å, and c = 21.98 Å for the complexes were used. In total, 163, 168, and 128 atoms are included in the Cs₂Pb(SCN)₂Br₂−H₂O, Cs₂Pb(SCN)₂Br₂−NH₃, and Cs₂Pb(SCN)₂Br₂−O₂ systems, respectively.

All of the ab initio dynamics simulations were carried out using the Car−Parrinello molecular dynamics (CPMD) 49 module in the Quantum-Espresso package. 50 Ultrasoft scalar relativistic pseudopotentials 51 and the generalized gradient approximation with the Perdew−Burke−Ezernhof (PBE) function 52 were applied describing the exchange interaction between atoms. Plane-wave basis set cutoffs for the augmented density and the smooth part of the wave functions are 200 Ry and 25 Ry, respectively. A fictitious electronic mass corresponds to 400 au. All of the dynamics calculations were conducted at a Nose−Hoover 53 constant temperature of 300 K. An integration time of 5 au was employed and the total simulation time was 200 ps for each system.

To describe quantitatively the influences of gas on the perovskite skeletons, the adsorption energy and charge transfer of the semiconductor−gas systems were computed every 5 ps. The structure optimization and comparisons of the stability of the Pb−SCN and Pb−NCS bonding were performed by the projected augmented wave (PAW) 54 plane-wave basis in VASP. The PBE calculations including the spin−orbit coupling (SOC) 55,56 were employed for all of the periodic complexes. The cutoff of the kinetic energy was set as 500 eV, and the electronic minimization was carried out with a tolerance of 10⁻⁴ eV. A Monkhorst−Pack k-point grid of 3 × 3 × 1 was applied for the convergence of the energies and forces of the three adsorption systems. To reflect accurately the geometry variations and the electronic states of the adsorbed gas molecules on the perovskite materials, all of the coordinates of the atoms in the snapshots were left unoptimized. The adsorption energy was computed via equation

\[ E_{\text{ads}} = E_{\text{perovskite+gas}} - E_{\text{perovskite}} - E_{\text{gas}} \]

where \( E_{\text{ads}} \), \( E_{\text{perovskite+gas}} \), \( E_{\text{perovskite}} \), and \( E_{\text{gas}} \) are the adsorption energy of the complex, the total structural energy of perovskite and gas, the total structural energy of perovskite, and the structural energy of gas, respectively. The nonlocal van der Waals (VDW) contributions were considered with Grimme’s DFT-D3 correction 57 to probe the interactions between Cs₂Pb(SCN)₂Br₂ and the gas molecules.

The Bader charge population analysis 58,59 was adopted to calculate the charge transfer between the gas molecules and the semiconductor material. The net charge transfer \( \Delta Q \) is defined as

\[ \Delta Q_X = Q_X^{1=200} - Q_X^{i=0} \]

where \( X \) represents Pb²⁺, Br⁻, SCN⁻, Cs⁺, or gas, respectively.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acsomega.1c03952.

The RMSD curves of CH₃NH₃PbI₃−H₂O and (CH₃NH₃)₂Pb(SCN)₁₋H₂O: the final structure of CH₃NH₃PbI₃−O₂ and (CH₃NH₃)₂Pb(SCN)₁₋O₂ adsorption complexes; the snapshots with the stable connections between H₂O (a)−(c), NH₃ (d)−(f), and O₂ (g)−(i) and the surface of Cs₂Pb(SCN)₂Br₂, respectively; the adsorption energies of Cs₂Pb(SCN)₂Br₂−H₂O, Cs₂Pb(SCN)₂Br₂−NH₃, and Cs₂Pb(SCN)₂Br₂−O₂ adsorption complexes (PDF)

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Author Contributions

B.Z. contributed to the conception of the study, wrote the paper together with B.H., X.W., Y.Y., and L.T., and performed the data analyses and figure preparation. L.Z. and Q.L. contributed to the constructive discussions and data analyses.

Notes

The authors declare no competing financial interest.

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