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Discriminating sensing of explosive molecules using graphene–boron nitride–graphene heteronanosheets†

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Since the synthesis of graphene–boron nitride heterostructures, their interesting electronic properties have attracted huge attention for real-world nanodevice applications. In this work, we combined density functional theory (DFT) with a Green's function approach to examine the potential of graphene–boron nitride–graphene heteronanosheets (h-NSHs) for discriminating single molecule sensing. Our result demonstrates that the graphene–boron nitride–graphene (h-NSHs) can be used for discriminate sensing of the 2,4-dinitrotoluene (DNT), octahydro-1,3,5,7-tetranitro-1,3,5,7-tetrazocine (HMX), pentaerythritol tetranitrate (PENT), and 2,4,6-trinitrotoluene (TNT) molecules. We demonstrate that as the length of the BN region increases, the sensitivity of the heteronanosheets to the presence of these explosive substances increases.

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1. Introduction

The ability to detect explosive materials and their compounds is a major challenge, which has recently become an important priority for security applications.^{1–5} A variety of laboratory-based approaches have been employed for discriminated sensing of explosives,^{4–7} for instance, non-portable devices for detecting such as gas chromatography,^{6,8,9} ion-mobility spectrometer, electronic noses,^{3,10–12} mass spectrometer,^{3,4,6,13} high pressure liquid chromatography-diode array detection,^{14,15} radiation,^{2–4,6,8} and Raman spectroscopy.^{16,17}

To meet the challenge of designing effective, portable and less expensive discriminating sensing nanodevices, there is a need to develop novel nanostructure materials and concepts, with devising new strategies for managing and developing on sensor chips. Nanomaterials might be promising nominees to design chemical sensors with high sensitivity.^{18–23} To avoid the high costs and preclude the need for chemical amendment or separation of the analytes, the label free techniques for

discriminating sensing of molecules are coveted targets in technology.²⁴ Graphene–boron nitride heterostructures have attracted the attention of scientists and research groups due to their unique electrical, mechanical, morphological, thermal properties and optoelectronics.^{25–29} In-plane junctions between single layer boron nitride and graphene are already synthesised,^{30–33} but the heterostructure materials are challenging to recognise experimentally. Doping graphene derivatives with heteroatoms are promising materials for high-performance sensing,³⁴ for example gas and toxic gas sensors.^{35–39} The detection of target analytes using graphene materials is mostly depend on the changes of the conductance after the adsorption of analytes to be sensed.³⁴ Graphene–boron nitride heterostructures have been utilised as gas sensor.^{40–43} For example, it has been reported that graphene–boron nitride heterostructure is stable structure, and also the Fermi level, work function, and conductivity are remarkably changed upon adsorption of the analytes on them compared to the pristine graphene–boron nitride heterostructures. Other studies^{44–46} demonstrated the potential of graphene–boron nitride heterostructures for biosensors. For example, it was shown that the nanopore formed from graphene–boron nitride heterosheet can be used for discriminating sensing of the four nucleotides of DNA.

The 2,4-dinitrotoluene (DNT), octahydro-1,3,5,7-tetranitro-1,3,5,7-tetrazocine (HMX), pentaerythritol tetranitrate (PENT), and 2,4,6-trinitrotoluene (TNT) molecules (shown in Fig. 1, middle sub-figure) are common explosives. The development of suitable detector to sense this materials has received attention from research groups.^{5,7,47–54} In this paper, we demonstrate that graphene–boron nitride–graphene heteronanosheet (h-NSHs)

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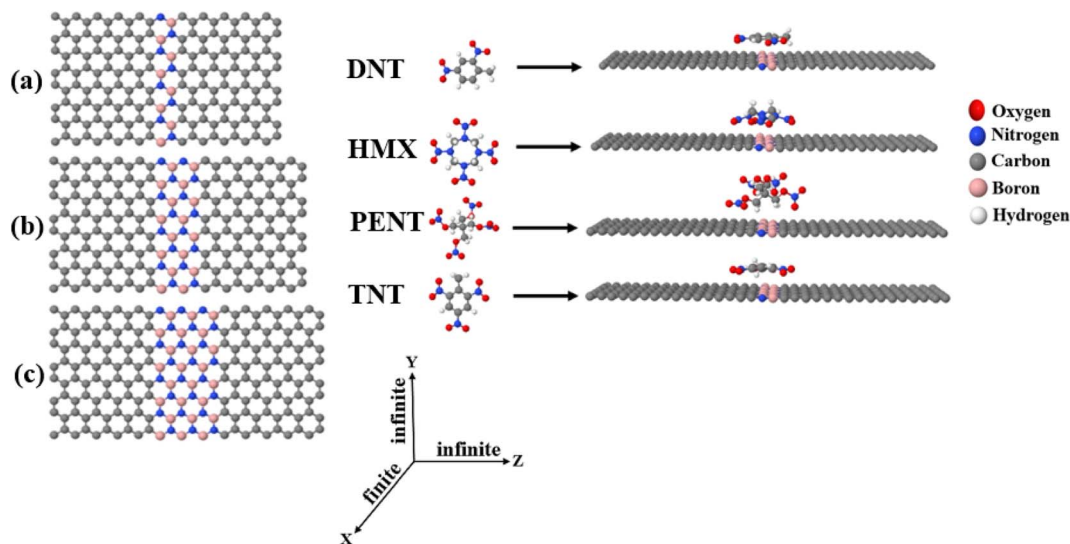


Fig. 1 Shows (a–c) the h-NSH junctions constructed from graphene joined by a perfect insulator BN, ranging from 1BN strip to 3BN strips. (Right) An example of h-NSH with 1BN junction with four explosive molecules (DNT, HMX, PENT and TNT) on 1BN scatterer.

can be a promising candidate for discriminating sensing of these explosive materials. h-NSHs are formed from two graphene nanosheets as left/right leads connecting *via* equivalent boron nitride nanosheet as scattering region as shown in Fig. 1a–c. Electrical current flows from left lead (G, graphene) to the right lead (G, graphene) through the scattering region (BN, boron nitride). To understand the change in conductance of the h-NSHs when single explosive molecule binds to the h-NSHs, we perform first principal computations. In what follows, by analysis the results of the response of the h-NSHs junctions to

different BN-region with varied lengths, a notable fingerprint of each analyte is obtained which can be used for sensing.

2. Characterising of the junctions

Fig. 1 shows the three h-NSHs (labelled a–c) which are composed of graphene (G) nanosheet as left/right leads and connected by boron nitride (BN) nanosheet scatterer of length varying from one strip to three strips (1BN to 3BN, respectively).

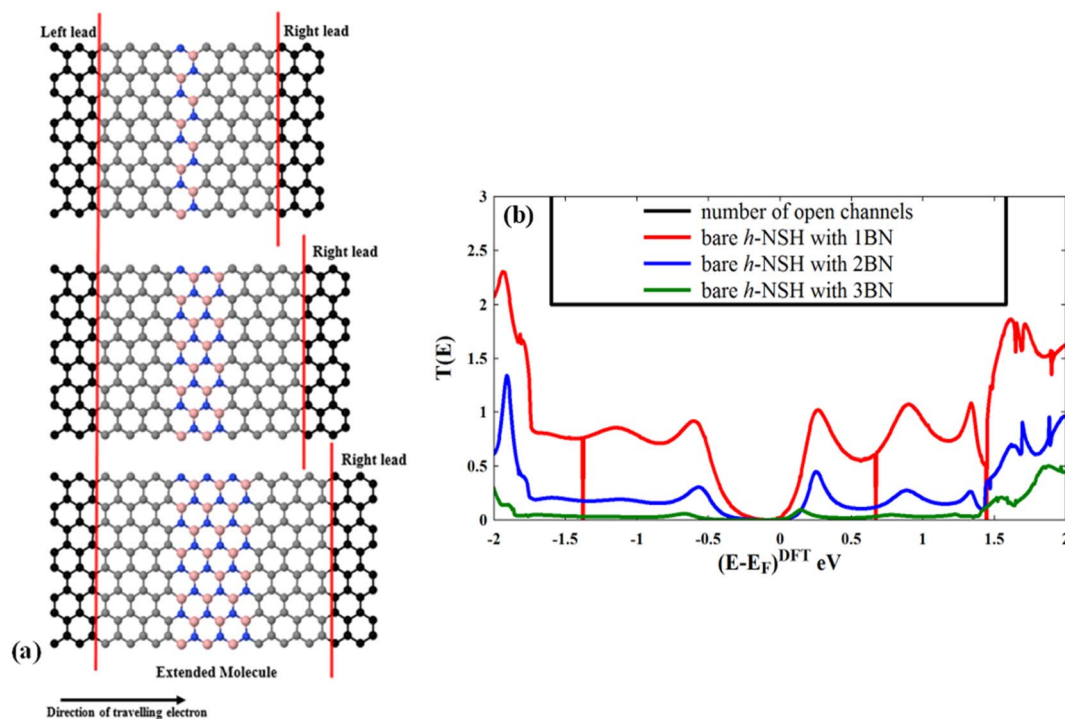


Fig. 2 Shows (a) the characterisation of the bare h-NSH junctions and (b) their electronic transmission coefficients $T(E)$.



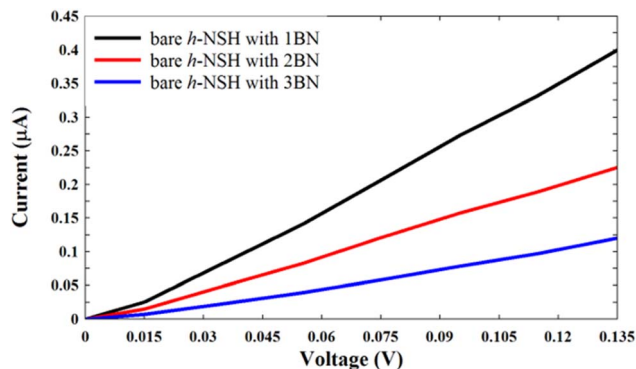


Fig. 3 Shows the I - V curves for the junctions in Fig. 2a.

Table 1 The binding energy (E_{Bin}) of the DNT, HMX, PENT and TNT molecules place on BN region of the h-NSH with 1BN/2BN/3BN junctions

Junction	E_{Bin} (eV)			
	+DNT	+HMX	+PENT	+TNT
h-NSH with 1BN	-0.301	-0.387	-0.410	-0.312
h-NSH with 2BN	-0.411	-0.460	-0.540	-0.480
h-NSH with 3BN	-0.570	-0.490	-0.460	-0.510

To find the ground state geometry of the h-NSHs, we employed the SIESTA⁵⁵ implementation of density functional theory (DFT). For all structures, we used the local density approximation (LDA)⁵⁶ with Ceperley–Alder (CA) parameterisation, the double zeta polarized (DZP) basis sets of pseudoatomic orbitals with norm-conserving pseudopotentials. The

initial h-NSHs were relaxed until the forces below $0.01 \text{ eV } \text{Å}^{-1}$. For lead calculations, a k -point grid of $1 \times 30 \times 30$ is sampled in the Brillouin zone. All the h-NSHs are infinite in Y and Z directions, whereas a vacuum space of 50 Å has been chosen along the X direction to avoid the interaction between neighbouring h-NSHs.

After obtaining the relaxed geometry of each analyte, the analytes were placed on 1BN, 2BN and 3BN scatterers, as shown in Fig. S1,[†] which shows the relaxed h-NSHs with 1BN, 2BN and 3BN scatterer after placing the explosive molecules on BNs regions. To investigate electrical conductance and current through the junctions, we employ Landauer approach,⁵⁷ and calculate transmission coefficients $T(E)$ for electrons of energy (E) passing from left lead to the right lead through the boron nitride (BN) scatterer using the Green's function based quantum transport code GOLLUM⁵⁸ combined with the mean field Hamiltonian (MFH) that obtained from SIESTA.

3. Discriminating single-molecule sensing of the h-NSH junctions

In this section, we investigate the electron transmission coefficients $T(E)$ of the bare h-NSH junctions with varied BN length as shown in Fig. 2a.

Fig. 2b shows the corresponding $T(E)$. We note that the energy gap progressively increases as the BN region length increases, and also the $T(E)$ drops exponentially with the length of the BN region when the energy (E) lies within the energy gap of the BN scatterer.^{24,59}

We also calculate the maximum current (I) at room temperature for small finite voltage of the three bare h-NSH junctions, which carried by the system from the left lead to the right lead using the following equation:⁵⁷

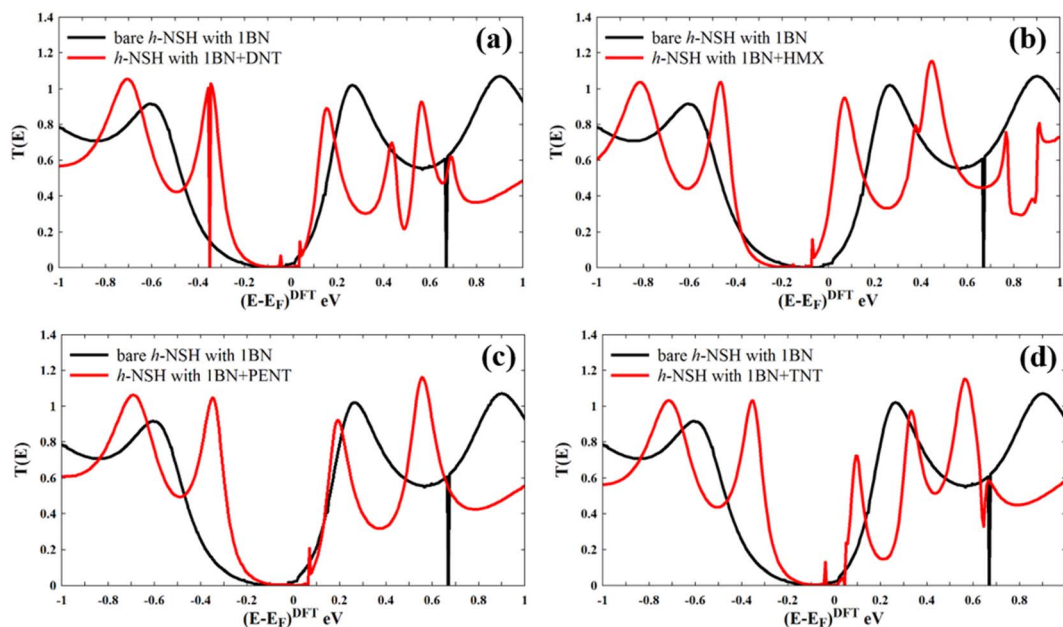


Fig. 4 Shows the $T(E)$ of the relaxed junctions (a) h-NSH with 1BN + DNT, (b) h-NSH with 1BN + HMX, (c) h-NSH with 1BN + PENT, (d) h-NSH with 1BN + TNT. In all figures, the Fermi energy (E_F) is shifted at zero.



$$I = \frac{e}{h} \int dE T(E) (f(E - \mu_{LL}) - f(E - \mu_{RL})) \quad (1)$$

where $e = |e|$ is the electron's charge, h is Planck's constant, $T(E)$ is the electronic transmission coefficient, f is Fermi-Dirac distribution function $f(E - \mu) = \frac{1}{(1 + e^{(E-\mu)/k_B T})}$, which associated with the electrochemical potential μ , k_B is Boltzmann's constant, and T is temperature. The obtained current shown in Fig. 3.

From Fig. 3, we can see that the current decreases as the BN length increases.

Next, we examine the h-NSH with 1BN junction for discriminating sensing. For this we first find the ground state geometry of the DNT, HMX, PENT and TNT molecules by calculating the binding energies (E_{Bin}) and performing geometry optimisation of molecules on 1BN, 2BN and 3BN as shown in Fig. S1.† To calculate the E_{Bin} , we use eqn (2) to minimise the basis set superposition error (BSSE).^{60,61}

$$E_{\text{Bin}} = E^{\text{h-NSH+mol}} - (E^{\text{h-NSH}} + E^{\text{mol}}) \quad (2)$$

where the total energy of the system (h-NSH with molecule) is $E^{\text{h-NSH+mol}}$, $E^{\text{h-NSH}}$ (h-NSH) and E^{mol} (molecule) are the total energies of the isolated subsystems. Table 1 shows the obtained binding energies of the h-NSH with 1BN/2BN/3BN after placing the DNT, HMX, PENT and TNT molecules on the 1BN/2BN/3BN regions.

From Table 1, it is clear that the obtained binding energies for the h-NSH with 1BN are from -0.410 eV to -0.301 eV, for the

h-NSH with 2BN are ranging from -0.540 eV to -0.411 eV, and for the h-NSH with 3BN are from -0.570 eV to -0.460 eV. PENT has the highest binding energy to 1BN and 2BN regions, while DNT molecule shows highest binding energy to 3BN.

Now we calculate transmission coefficient $T(E)$ of electrons with energy E traversing from one electrode to the other through h-NSH with 1BN junction in the absence and presence of analyt.

From Fig. 4a-d, we can see that a notable fingerprint for each molecule around the E_F comparing to the bare h-NSH with 1BN. Similarly, we calculate the $T(E)$ of the h-NSHs with 1BN/2BN in the presence of analytes (DNT, HMX, PENT and TNT) on 2BN and 2BN scatterers as shown Fig. S2 and S3 in the ESI.† Once

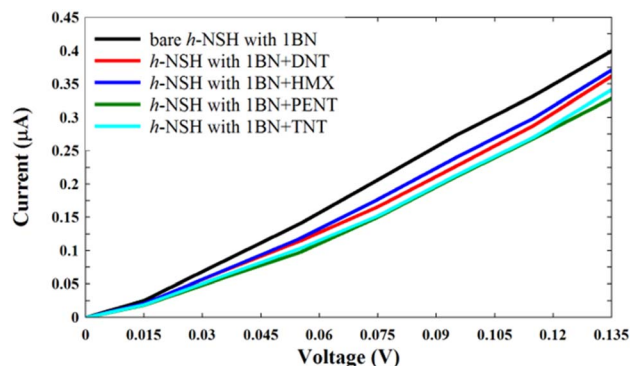


Fig. 6 Shows the current of the h-NSH with 1BN junction after place the DNT, HMX, PENT and TNT on the 1BN scatterer.

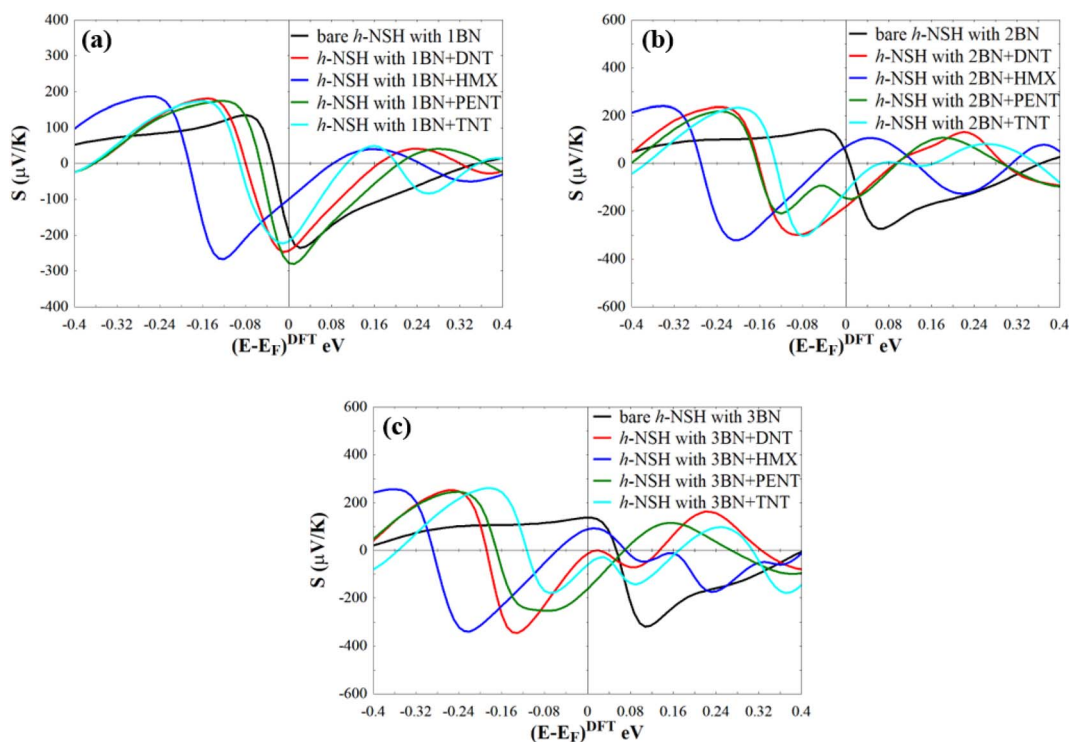


Fig. 5 Shows the room-temperature thermopower (S) of the junctions with/without molecules, (a) h-NSH with 1BN, (b) h-NSH with 2BN, and (c) h-NSH with 3BN. In all figures, bare junctions are in black line and the Fermi energy (E_F) is shifted at zero.



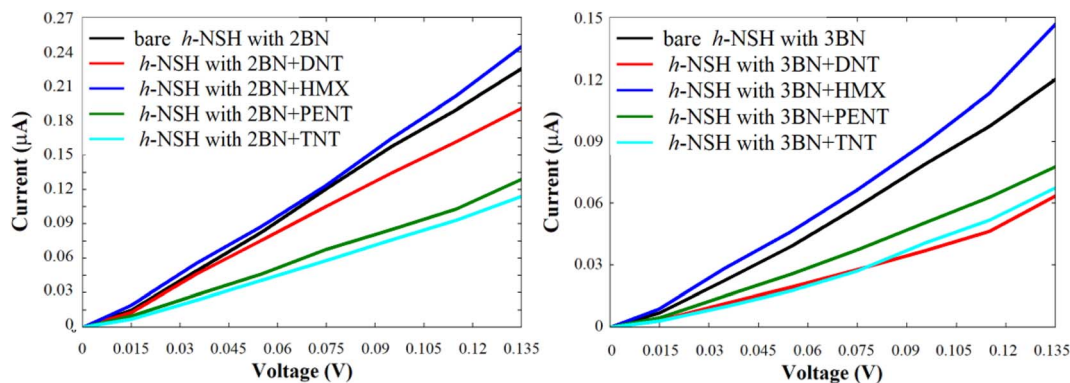


Fig. 7 Shows the current of the h-NSH junctions (a) with 2BN and (b) with 3BN junctions after place the DNT, HMX, PENT and TNT on the BN scatterer.

again, we obtain remarkable fingerprints for each molecule around E_F . These new peaks around the E_F due to the present of explosive molecules can lead to enhancement the Seebeck coefficient (S) as shown in Fig. 5, which can be used for sensing molecules using Seebeck effect (Seebeck sensing). For more clarity, see Table S1,[†] which shows the values of the Seebeck coefficient of the h-NSHs with 1BN/2BN/3BN junctions introducing the analytes.

To confirm the ability of the h-NSH with 1BN junction for discriminating sensing of the analytes (shown in Fig. S1a[†]), we used eqn (1) to calculate the current (I) as shown in Fig. 6.

From Fig. 6, we can see that each molecule has a specific value of current comparing with the current through the bare h-NSH with 1BN junction (black line). These results reinforce that the h-NSH with 1BN system can be used for discriminating sensing of the explosive molecules.

We repeated the same strategy with the h-NSHs with 1BN/2BN junctions shown in Fig. S1b and c,[†] the resulting I shown in Fig. 7.

Again, from Fig. 7, we can see obviously that each molecule has a particular value of the current comparing with the bare h-NSH with 2BN/3BN. The result show that the h-NSH with 1BN/2BN/3BN junctions are potential materials for discriminating sensing nanodevice of the explosive molecules, and the most preferable junctions for selective sensing are the h-NSH with 2BN/3BN junctions comparing with the h-NSH with 1BN junction.

When analyte physisorbed on the membrane, it acts like a locale gate and modifies the potential energy of the membrane locally. Due to the charge transfer between the analyte and the membrane, the charge density on the membrane can also be affected. These can lead to the changes on the electrical conductance. Our calculations show that unlike graphene, these effects significantly influence the electrical conductance of membranes formed by graphene-hBN heterostructures.

4. Conclusions

We demonstrate that the h-NSH with BN (ranging from 1BN strip to 3BN strips) junctions are potential material for

discriminating sensing of four explosive molecules (DNT, HMX, PENT, and TNT). We also show that an energy gap is formed by introducing BNs in graphene as scatterers, and can be turned by changing the length of the BN region. Our result demonstrates that the $T(E)$ decay exponentially with increasing the length of the BN scatterer. Furthermore, our result show that the new peaks appear around the E_F in the $T(E)$ due to the presence of the explosive molecules lead to enhancement the Seebeck coefficient, which can be used for discriminating Seebeck sensing of molecules. Since both electrical current, and the sign and amplitude of Seebeck coefficient are changing upon translocation of analytes on the heterostructures, a combination of these factors can be used to reduce the error in the detection process. This provides a powerful tool for selective sensing of explosive materials with similar electronic fingerprint.

Author contributions

All authors were involved in interpreting the results, wrote and commented on the manuscript.

Conflicts of interest

The authors declare no conflict of interest.

Data availability

All required information to reproduced the simulation data is given in the manuscript and the ESI.[†]

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