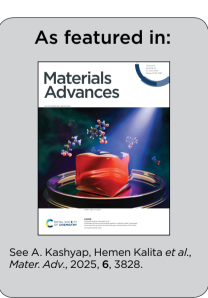


Showcasing research from Professor Hemen Kalita's laboratory, Department of Physics, Gauhati University, Assam, India.

Highly selective ammonia sensing at room temperature using DC plasma-modified MoS_2 nanoflowers

This cover art features the research work of Anurag Kashyap and Hemen Kalita from the Nanomaterials and Nanoelectronics Laboratory, Department of Physics, Gauhati University, Assam, India. It visually represents the enhancement in ammonia sensing by DC plasma-modified MoS $_2$ nanoflowers at room temperature. It highlights the effect of plasma treatment on the hydrothermally synthesized MoS $_2$ nanoflowers in enhancing sensitivity and selectivity towards very low concentrations of NH $_3$. This advancement holds potential for the development of efficient, low-cost gas sensors for environmental monitoring applications.

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Highly selective ammonia sensing at room temperature using DC plasma-modified MoS₂ nanoflowers†

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This study explored the enhancement of ammonia (NH₃) sensing properties of MoS₂ nanoflowers through a direct current (DC) plasma treatment with nitrogen incorporation. Plasma treatment induced sulfur vacancies and introduced nitrogen atoms into the MoS₂ surfaces, enhancing the number of active sites and improving the charge carrier mobility. The structural and chemical alterations were confirmed by characterisations using FE-SEM, XPS, and Raman spectroscopy. The resulting plasma-treated MoS₂ sensor demonstrated highly selective detection of NH₃ at room temperature with a rapid response time of 22 s and a recovery time of 23 s. The experimental limit of detection was achieved at 5 ppm (theoretically ≈ 80 ppb), which was lower than the safety threshold set by the National Institute for Occupational Safety and Health (NIOSH). Results of DFT studies also agreed with the experimental results. Thus, this work highlights DC plasma treatment as an efficient, cost-effective approach to enhance gas sensing performance, with implications for developing highly sensitive and selective sensors for environmental monitoring and safety applications.

1. Introduction

In recent years, the detection of toxic gases has gained increasing importance in various fields, ranging from air quality control and workplace safety to medical diagnostics. 1-3 In particular, detection of ammonia (NH3) has received significant interest owing to its adverse health effects, including severe respiratory conditions like asthma, emphysema, and chronic bronchitis, even when exposed to low levels. Different types of sensing technologies, such as electrochemical cells and metal oxide chemiresistors, have been reported for the commercial detection of toxic gases.⁵ However, these sensors often face challenges, such as the need for high operating temperatures and poor selectivity for differentiating between various gases.

To address these limitations, researchers have focused on alternative materials, particularly low-dimensional materials. Owing to their unique physiochemical properties, such as large

specific surface areas, distinct electrical properties, and ease of functionalisation, 2D materials have gained significant importance in gas sensing in recent years. 4,6,7 Among the 2D materials, transition metal dichalcogenides (TMDs) are promising candidates for fabricating highly sensitive gas and volatile organic compound (VOC) sensors.8 These materials provide weak gas-molecule binding and efficient charge transfer, enabling gas detection at room temperature. 9,10 Among the TMDs, molybdenum disulfide (MoS2) has sparked a lot of research interest as a prominent sensing material owing to its high surface-to-volume ratio, high surface activity and sensitivity, quick response time, and outstanding stability. 11-16 In fact, MoS₂ is more appealing than graphene-based gas sensors owing to its semiconducting nature and the appropriate, adjustable band gap energies.¹⁷ Despite its potential, MoS₂ still suffers from selectivity issues owing to similar VOC adsorption energies, making it difficult to detect one gas without interference from the other. 18,19 To overcome this, advanced fabrication techniques have been implemented, including doping, intercalation, and the creation of heterostructures, enhancing the material's ability to target specific gases more effectively.20-24

Plasma treatment is a prevalent method to alter the inherent physical and chemical properties of TMDs for application in gas sensing, ion storage, and oxygen reduction and hydrogen evolution reactions.^{5,25–28} RF plasma treatment is commonly

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employed to alter surfaces because of its ability to provide homogeneous treatments applicable to 2D, complicated, and 3D materials.²⁹ The direct current plasma (DC plasma) treatment is another method by which material sensing properties can be modified by incorporating surface defects.^{30,31} The procedure involves high-energy ions and reactive plasma species that alter the surface morphology and electronic characteristics. The creation of defects such as vacancies, surface states, and dangling bonds results in such changes, which increase the surface area. These modifications increase the sensitivity of the ability to adsorb the analyte, facilitate charge transfer, and thus increase the sensitivity and selectivity of the sensor. Moreover, plasma treatment with DC offers an avenue to modify sensor surfaces because it is extremely adaptable, less expensive than RF plasma, and suitable for a variety of materials.31 Plasma processing has recently demonstrated itself as a rapid, convenient, straightforward, and effective approach for modifying the surface chemical characteristics of materials, particularly for the treatment of 2D materials without solvent contamination.32,33 The chemical properties of the plasma process gas are crucial for determining the reaction mechanisms involved in plasma surface modification. Various gases, including oxygen, argon, helium, nitrogen, ammonia, carbon dioxide, and water, are used as process gases in surface modification techniques using plasma.³⁴ This approach may produce additional edge sites and heteroatoms, enhancing their charge carrier mobility and catalytic and optoelectronic capabilities.35,36 Among the many existing methodologies, nitrogen plasma treatment has recently been recognized as an excellent approach for achieving controlled nitrogen atom incorporation into TMDs. 5,28,34,37 Zhao et al. developed an Ndoped MoS2 nanosheet-based sensor with improved sensitivity to NO₂ at room temperature.²⁸ Initially, MoS₂ nanosheets were

hydrothermally fabricated and N-doping was performed via nitrogen plasma to synthesize N-MoS2 nanosheets. Nitrogen atoms were introduced using a plasma-assisted technique. The atoms were adsorbed on the surface and doped into the lattice defects of MoS2. This modified the electronic properties, such as the reduction of the Fermi level and the narrowing of the band gap.36 The structure increased the conductivity and sensitization with NO2, thereby increasing the response and reducing the detection limit.

This study reported the effects of DC plasma treatment on the NH₃ sensitivity of the MoS₂ nanoflower structure. Plasma-treated MoS₂ shows selective detection of NH₃ at RT (25 °C), greatly improving its sensing performance. N-MoS₂ sensor has a faster response time of 22 s and takes 23 s to recover. Moreover, plasma treatment significantly stabilized the sensing response of the sensor. The experimental and theoretical limits of detection were 5 ppm and 80 ppb, respectively. An effective way of improving room temperature NH₃ detection is described in this study.

Experimental section

2.1. Materials

Sodium molybdate (Na₂MoO₄·2H₂O), thioacetamide (CH₃CSNH₂), and oxalic acid (H2C2O4) were obtained from Sisco Research Laboratories (SRL) and used as received. The plasma treatment was performed using a glow discharge plasma reactor to produce N2 and Ar plasma. Fig. 1 shows a schematic of the experimental assembly. Deionised water (DIW) was used throughout the experiment.

2.2. Synthesis of MoS₂

A wet chemical route is followed to synthesize MoS2 nanoflowers. In a typical reaction, 1 g of sodium molybdate and 1.2 g

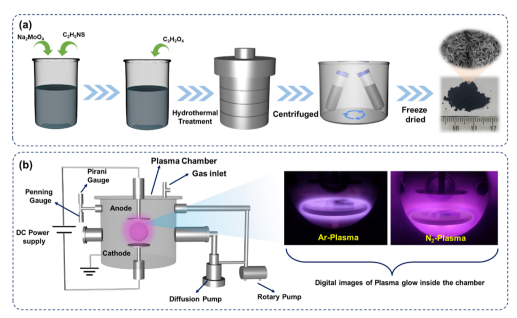


Fig. 1 Schematic of (a) synthesis of MoS₂ nanoflowers following the wet chemical route and (b) plasma treatment assembly with digital images of Ar and N₂ plasma during treatment

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of thioacetamide are stirred in 50 mL of DI water for 30 min at ambient temperature. Then, 0.6 g of oxalic acid was added to the mixture, which was further stirred for another 30 min. The solution was then hydrothermally treated for 24 h at 200 °C by being put in a 100 mL Teflon beaker and sealed inside a stainless steel (SS) container. After cooling to room temperature, the sample is centrifuged with ethanol and DI water and freeze-dried for 24 h.

2.3. Sensor electrode fabrication

Interdigitated electrode structures (IDE) have been fabricated following a standard copper (Cu) etching procedure using a ferric chloride (FeCl₃) solution. The desired IDE structure was transferred to the copper printed circuit board (PCB), which was dipped in FeCl₃ solution following agitation for the etching of the unwanted copper. Finally, the IDE structure remains on the board and is rinsed with ethanol and DI water and dried by purging nitrogen before deposition of the sensing material.

2.4. Synthesis of N-MoS₂

As nitrogen plasma is treated on the sensor, the sensor is termed as N-MoS2 in the entire report. Sensors were treated with an argon/nitrogen plasma produced from a custom-built glow discharge reactor. The sensor was fabricated using the drop-casting method. The synthesized MoS2 sample was dispersed and drop-cast on sensors composed of copper IDEs and dried in a vacuum oven at 60 °C for 3 h before plasma treatment. Fig. 1(b) shows a schematic of the experimental setup. The plasma chamber, a cylindrical stainless steel (SS 304L) vessel with a volume of approximately 3.5×10^3 cm³ (diameter: 30.0 cm, height: 50.0 cm), was evacuated using a diffusion pump (1000 L min⁻¹) backed by a rotary pump (540 L s⁻¹). A base pressure of $\sim 10^{-6}$ mbar was achieved.

Two circular stainless steel (SS 304L) electrodes initiated the discharge and were powered by a DC power supply (1000 V, 2.5 A). Ultra-pure argon/nitrogen gas (99.99%) was introduced into the chamber through a digital flow controller to maintain a working pressure of 1.0×10^{-1} mbar. The sensors were exposed to Ar and N2 plasma for 2 and 10 minutes, respectively.

A Langmuir probe was used for plasma diagnostics. The measured plasma density was 4.35 \times 10¹⁵ m⁻³ and 4.43 \times 10¹⁵ m⁻³ in the presence and absence of sensors, respectively. The electron temperature remained relatively constant at 1.38 eV. The slight decrease in plasma density observed in the presence of sensors is attributed to plasma-sample interactions.34,38

2.5. Computational details

The computations conducted utilised the density functional theory (DFT) - based Cambridge Serial Total Energy Package (CASTEP) module, which is integrated into the Materials Studio software suite. 4,7,39 The Kohn-Sham equations were solved using the plane-wave pseudopotential method developed within the DFT framework.40 For the exchange-correlation function, the generalized gradient approximation (GGA), as outlined by Perdew-Burke-Ernzerhof (PBE), was used for

geometry optimization.41 Throughout the computations, the DFT-D dispersion correction suggested by Grimme and used in CASTEP was used to account for the van der Waals (vdW) intermolecular forces. 42 A 6 \times 6 \times 1 k-point mesh was used to sample the Brillouin zone with an energy cutoff of 517 eV. Geometry optimization, which is essential for achieving a configuration's equilibrium state with minimum energy, was performed using the Broyden-Fletcher-Goldfarb-Shanno (BFGS) algorithm, which is highly effective for energy minimization in crystalline materials.⁴³

A medium convergence tolerance was adopted for geometry optimization, and electronic minimization parameters utilized the Gaussian smearing scheme with a smearing width of 0.1 eV. Additionally, a 20 Å vacuum was introduced perpendicular to the interface surface to eliminate interactions between the interfaces and their periodic images.

The difference in charge density between the MoS₂ system and the adsorbed gas is computed as:

$$\Delta \rho = \rho_{\text{(MoS}_2 + \text{gas_molecule)}} - \rho_{\text{gas_molecule}} \tag{1}$$

where $\rho_{(MoS_2+gas_molecule)}$ and $\rho_{gas_molecule}$ are the charge densities of the MoS₂ + gas molecule system and the gas molecule, respectively.

The binding energy of the adsorbed gas molecules is calculated using the equation:

$$E_{\rm b} = E_{\rm (MoS,+gas_molecule)} - E_{\rm (gas_{molecule})}$$
 (2)

where $E_{(MoS_2+gas_molecule)}$ and $E_{(gas_{molecule})}$ are the total energies of the MoS₂ + gas molecule system and the gas molecule, respectively.

2.6. Characterization

The morphology of the synthesized samples were observed using field emission-scanning electron microscope (FE-SEM, ΣIGMA 300 VP, Carl Zeiss). The structural characteristics of the fabricated samples were characterized using an X-ray diffractometer (XPERT PRO, Philips/PANalytical) with Cu Kα radiation (1.54 Å) and with a laser micro-Raman system (Make: Horiba JobinYvon, Model: LabRam HR). XPS analyses were performed using an X-ray Photo Spectrometer (Model-K Alpha, Make: Thermo Fisher Scientific).

3. Results and discussions

3.1. Structural and morphological characterizations of MoS₂ and N-MoS₂

Fig. 2 shows the FESEM images and EDX analysis of MoS₂ and N-MoS₂. As shown in Fig. 2(a), the MoS₂ nanosheets are arranged in a flower-like structure. EDX analysis (Fig. 2(c)) provides elemental information about the synthesized materials. In the case of N-MoS₂ (Fig. 2(b)), the flower-like structure is intact with a bit of deformation, which may be due to the bombardment of energetic ions during argon plasma treatment, followed by nitrogen plasma. The EDX spectrum (Fig. 2(d)) shows the presence of nitrogen, which is absent in

MoS. N-MoS₂ 17.44 12.67 35,50 22.18 45.71 4.12 0.94 2.18 2.56 26.53 3.76 0.63 56.74 16.39 37.03 (e) (f) (g) (h)

Fig. 2 FESEM images of (a) MoS₂ nanosheet and (b) N-MoS₂ nanosheet; EDX spectra of (c) MoS₂ and (d) N-MoS₂; and (e) –(h) colour mapped images of Mo, S, N and O elements of N-MoS₂ sample.

the EDX spectrum of MoS₂ (Fig. 2(c)). The colour-mapping images of different elements present in the nitrogenincorporated MoS₂ sample are shown in Fig. 2(e-h).

Fig. 3(a) shows the Raman spectra of MoS₂ and N-MoS₂, in which two distinct peaks E_{2g}^{1} and A_{g}^{1} represent the vibration modes of hexagonal MoS_2 . The A_g^1 peak that appears at $\sim 402 \text{ cm}^{-1}$ is attributed to the out-of-plane vibration of Mo and S atoms, whereas E_{2g}^1 appears at $\sim 375~\text{cm}^{-1}$ and is attributed to the in-plane vibration of Mo and S atoms.44 The peak spacing between the E_{2g}^1 and A_g^1 peaks for both MoS₂ and N-MoS₂ was the same (26.4 cm⁻¹), confirming their similar layer numbers.44 The Raman spectra of MoS2 after various nitrogen plasma treatment times are presented in the ESI† (Fig. S1). The X-ray diffraction patterns of the synthesized MoS₂ nanoflowers and bulk MoS₂ are presented in the ESI† (Fig. S2). Fig. 3(b) shows the XPS survey spectra of MoS₂ and N-MoS₂. From this figure, the N/Mo ratios are 0.5 and 3.3 for MoS₂ and N-MoS₂, respectively. Fig. 3(c)-(h) shows the deconvoluted XPS spectra of the MoS₂ and N-MoS₂ samples. In Fig. 3(c), the Mo 3d spectrum of MoS₂ exhibits four peaks, among which Mo⁴⁺ 3d_{5/2} and Mo4+ 3d3/2 correspond to 228.9 eV and 232.0 eV, respectively. 45 Along with this, a small peak around 235 eV appears due to the oxidation of Mo on the MoS₂ surface and is assigned to $\mathrm{Mo^{6+}}\ 3d_{3/2}$ and $\mathrm{Mo^{6+}}\ 3d_{5/2}$, respectively. 46 The peak at 226.1 eV was assigned to the S 2s of MoS₂. 45 For N-MoS₂, the deconvoluted spectrum of Mo 3d in Fig. 3(f) shows the peak present at 233.5 eV along with the peaks present in Fig. 3(c),

which arise due to bonds between Mo and N. 28,47,48 Moreover, the intensity of the Mo⁶⁺ 3d peaks increased, which may have been due to the oxidation of Mo after treatment.

Fig. 3(d) and (g) show the S 2p spectra of MoS₂ and N-MoS₂, respectively. Two peaks appear at 163.1 and 161.7 eV, corresponding to S^{2-} $2p_{1/2}$ and S^{2-} $2p_{3/2}$, respectively.²⁸ Fig. 3(e) and (h) show the Mo 3p spectra of MoS₂ and N-MoS₂, respectively. The peaks at 394.7 eV arise due to Mo 3p_{3/2} in both spectra. Along with that, two distinct and intense peaks in Fig. 3(h) at 398.5 eV and 402.0 eV appear due to characteristic Mo-N bond and N-O, respectively, which are less intense in the case of the Mo 3p deconvoluted spectrum of MoS₂ (Fig. 3(e)). 47,49

3.2. Gas sensing measurements of the MoS₂ and N-MoS₂ sensors:

The gas sensing measurements of the MoS₂ and N-MoS₂ sensors were performed in a homemade sensing assembly (Fig. 4). The sensing material was dropcasted onto the interdigitated electrode fabricated on the PCB, as described in Section 2.3. The sensor was then dried in a vacuum oven at 60 °C for 3 h. The sensor was then plasma-treated with argon and nitrogen plasma in a plasma chamber. The sensor was then exposed to VOCs in the gas sensing chamber (Fig. 4). The required amount of ammonia, calculated using the standard formula (ESI,† eqn (S1)), was inserted inside the sensing chamber.4,7

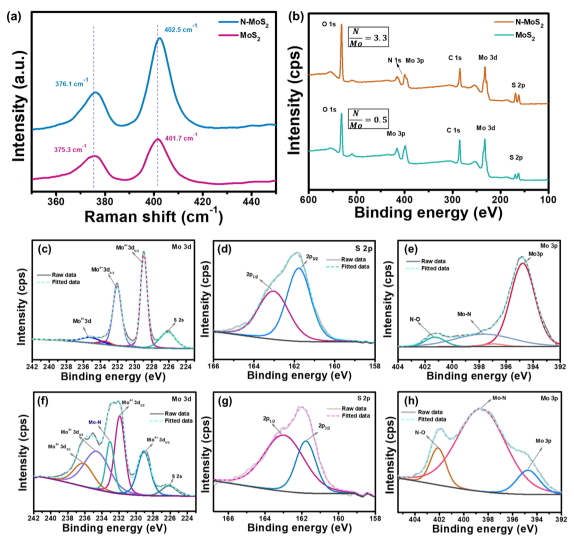


Fig. 3 (a) Raman spectra of MoS_2 and $N-MoS_2$ XPS; (b) XPS survey spectra of MoS_2 and $N-MoS_2$; XPS deconvoluted spectra of (c) Mo 3d, (d) S 2p, and (e) Mo 3p of the MoS_2 sample and (f) Mo 3d, (g) S 2p, and (h) Mo 3p of the $N-MoS_2$ sample.

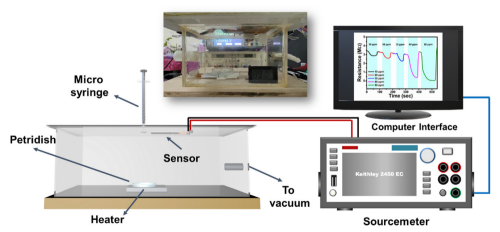


Fig. 4 Schematic of the gas sensing setup assembly; inset: digital image of the sensing assembly.

Fig. 5 shows the dynamic responses of the MoS₂ and N-MoS₂ sensors at room temperature towards ammonia (NH₃),

formaldehyde (HCHO), and triethylamine (TEA). Fig. 5(a) shows the responses of the MoS₂ and N-MoS₂ sensors to ammonia. N-

(a) 2.8 (b) Response $(R_a/R_g)_{\tau}$ Response (R 0.8 0.8 1200 200 600 800 1000 200 600 400 800 1000 Time (sec) Time (sec) 2.8 NH₂ 2.6 Response (R_a/R 2.4 Response

Fig. 5 Dynamic sensing responses of the MoS₂ and N-MoS₂ sensors towards (a) NH₃, (b) HCHO and (c) TEA at room temperature and (d) comparison of the sensing responses of MoS₂ and N-MoS₂ sensors towards various analyte VOCs.

1000

MoS₂ sensor exhibits an enhanced sensing response to NH₃ compared to the MoS₂ sensor. In contrast, in the case of HCHO and TEA (Fig. 5(b) and (c)), there is little change in the response. Fig. 5(d) shows a comparison histogram of the sensing responses of MoS2 and N-MoS2 towards NH3, HCHO and TEA, respectively. To check the selectivity of N-MoS₂ sensor towards common interfering VOCs, acetone and ethanol were also exposed to measure the sensing response.

200

400

600

Time (sec)

800

The effect of plasma exposure time on the sensing performance of the MoS₂ nanoflower structure was analysed by exposing the nitrogen plasma at three different times (5 min, 10 min and 15 min). The sensing responses of the plasmatreated sensors and the non-treated MoS₂ sensor (MS sensor) are shown in Fig. 6. The different plasma-treated sensors were termed NMS 05 (5 min plasma exposure), NMS 10 (10 min plasma exposure), and NMS_15 (15 min plasma exposure), depending on the plasma exposure time. The results show that NMS_05 exhibits no significant change in response compared with the MS sensor. However, NMS_10 and NMS_15 sensors have similar responses to 10 ppm NH₃ vapour at room temperature.

Fig. 7(a) shows the selectivity of the synthesized N-MoS₂ sensor towards various VOCs. As shown in the figure, the sensor is highly selective towards NH3 compared to other common VOCs in the ambient environment. Fig. 7(b) shows the response and recovery times of N-MoS2 towards 10 ppm NH₃ vapour. The response time of a sensor is the time taken for the resistance to reach 90% of its total change when the sensor

is in the proximity of the target gas, while the recovery time is the duration needed for the resistance to return to 90% of its original baseline after the gas is removed. 4,7 The response time of the sensor was 22 s, whereas the recovery time was 23 s.

нсно

MoS, N-MoS,

MoS₂ N-MoS₂

TEA

The dynamic changes in the response of the sensor to 10 ppm NH₃ vapour are shown in Fig. 7(c). Up to five cycles, the sensor showed an almost constant response with a little bit of deviation from the initial response. Fig. 7(d) shows the variation in sensor resistance towards 10 ppm NH₃ vapour. The changes in the resistance values were almost constant from the baseline value, with a deviation at the initial cycle.

Fig. 8(a) shows the dynamic response of N-MoS₂ sensor to varying NH₃ concentrations during continuous measurement. The sensor has a very high response with increasing concentrations of the analyte vapour, which may be due to humidity. Fig. 8(b) shows the linear response with the fitting curve of N-MoS₂ sensor to different VOC concentrations. From the calibration curve with increasing concentration of the NH₃ vapour, it can be observed that the curve is statistically linear with a correlation coefficient value $R^2 = 0.97$. The theoretical estimation of the limit of detection (LOD) of the sensor can be estimated using the following eqn (3):⁵⁰

$$LOD = 3 \frac{RMS_{\text{noise}}}{Slope}$$
 (3)

where RMS_{noise} is estimated using the root mean square deviation from the variation of the relative sensor response in the baseline before exposing the sensor to the NH₃ vapour and the

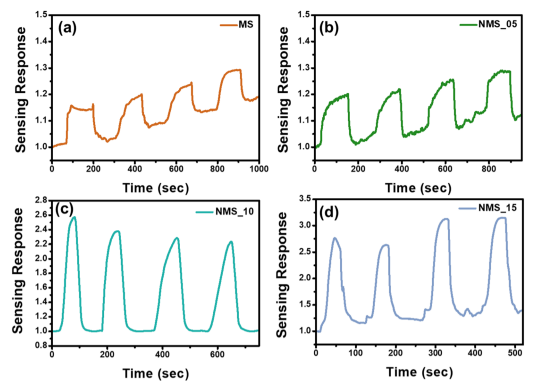


Fig. 6 Dynamic sensing response of (a) MoS₂ (MS sensor) and N-MoS₂ sensors with different nitrogen plasma exposure times of (b) 5 min (NMS_05), (c) 10 min (NMS_10), and (d) 15 min (NMS_15) towards 10 ppm NH₃ vapour.

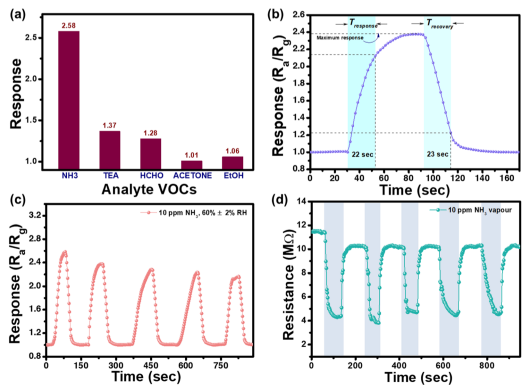


Fig. 7 (a) Selectivity of N-MoS₂ sensor towards different analyte VOCs; (b) response and recovery time measurements of N-MoS₂ sensor; (c) dynamic sensing response of N-MoS₂ sensor towards 10 ppm NH₃; and (d) resistance change during the exposure of 10 ppm NH₃ vapour to N-MoS₂ sensor.

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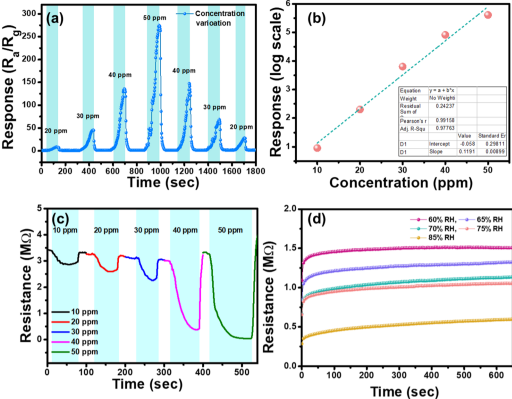


Fig. 8 (a) Response of N-MoS₂ sensor to varying concentrations of NH₃ in a continuous measurement; (b) linear response with fitting curve of N-MoS₂ sensor to different VOC concentrations; (c) measurement of resistance changes from its baseline resistance with increasing concentration of analyte VOC at room temperature; and (d) variation in the baseline resistance of the sensor with varying relative humidity

slope refers to the slope of the linear fitting curve in Fig. 8(b). The slope was calculated as 0.1191. The RMS_{noise} is calculated using eqn (4):

$$RMS_{\text{noise}} = \sqrt{\frac{S^2}{N}}$$
 (4)

where S is the standard deviation (S.D.) of the data before exposing the sensor to formaldehyde vapour and N is the corresponding data point. From Fig. 8(a), the standard deviation for N = 80 (the data before the sensor is exposed to NH_3 vapour) is \approx 0.0284. Therefore, RMS $_{\rm noise}$ is 0.00319, and using eqn (3), the LOD is 0.080 ppm \approx 80 ppb.

Fig. 8(c) shows the dynamic resistance changes of N-MoS₂ sensor with varying concentrations of NH₃ vapour. The sensor is sensitive to humidity, and with an increase in the analyte VOC's volume, the humidity inside the chamber also increases, which enhances the sensor's response to high concentrations of NH₃. Fig. 8(d) shows the variation in baseline resistance at different relative humidity values inside the sensing chamber at room temperature.

3.3. Computational analysis

For the computational analysis, a $4 \times 4 \times 1$ supercell of MoS₂ containing 48 atoms was used. In the equilibrium state, the Mo and S atoms of the MoS2 monolayer form a hexagonal lattice.

The lattice constants of the MoS_2 monolayer are a = 3.21 Å and c = 12.14 Å. The values are in good agreement with earlier experimental and theoretical results. 51 To depict the incorporation of nitrogen and sulphur vacancy in the MoS2 monolayer, one sulphur atom was replaced by a nitrogen atom, and one sulphur atom was removed.

The optimised geometrical configurations of nitrogenincorporated MoS2 monolayers with sulfur vacancy and with different gases are presented in Fig. 8. All possible binding sites of the MoS₂ monolayer were considered, and the configuration with the highest binding energy was considered in further calculations to reflect the most probable experimental scenario.

The binding energies of all gases in the nitrogenincorporated MoS₂ monolayer, as calculated using eqn (4), are shown in Fig. 9. The binding energy of the NH3 molecule with the nitrogen-incorporated MoS2 monolayer was the highest (-0.81 eV). The binding energy of the TEA molecule was the lowest (-0.18 eV). This finding is in close agreement with the sensing response of the nitrogen-incorporated MoS2 monolayer. To delve deeper into the interaction between the various gases and the nitrogen-incorporated MoS2 monolayer, a substantial charge transfer occurs between them, as shown in Fig. 9. In addition to the N-MoS₂ system with S vacancy, the effect of NH3 adsorption on the N-MoS2 system without S

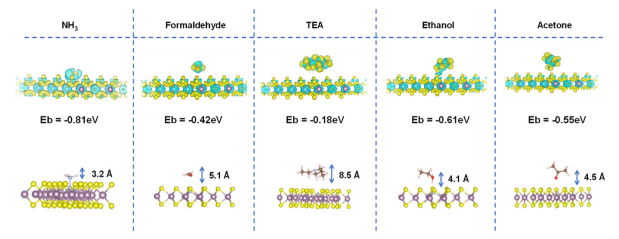


Fig. 9 Optimized configuration structures with electron charge density differences (EDDs) ($\Delta\rho$) of N-MoS₂ sample with S vacancy, along with intermolecular distances of N-MoS₂ with various VOCs. The yellow colour corresponds to charge accumulations, and the cyan colour corresponds to charge depletion.

vacancy was investigated. The charge density difference plots of all gases and the MoS₂ and nitrogen-incorporated MoS₂ monolayer without sulfur vacancies are shown in the ESI† (Fig. S3).

Similarly, from the intermolecular distances between gas molecules and nitrogen-incorporated MoS_2 monolayers with S vacancy, it was found that the NH₃ molecule exhibited the lowest distance (3.2 Å), whereas the TEA molecule exhibited the highest distance of 8.5 Å. This result can be explained by the increased electrostatic interaction between the NH₃ molecule and the nitrogen-incorporated MoS_2 monolayer.

For comparison, the binding energy values of all three systems (*i.e.*, only MoS₂, N-MoS₂ without S vacancy, and N-MoS₂ with S vacancy) are summarized in Tables 1, 2 and 3, respectively.

To examine the changes in the electronic properties of the nitrogen-incorporated MoS_2 monolayer upon NH_3 molecule adsorption, the total density of states (TDOS) was plotted (Fig. 10). The adsorption of the NH_3 molecule on the nitrogen-incorporated MoS_2 monolayer leads to a significant shift in the Fermi level towards higher energy. This also introduces new peaks at the conduction band at 0.52 eV and 1.97 eV. This shows that the introduction of the NH_3 molecule results in changes in the electronic properties of the nitrogen-incorporated MoS_2 monolayer.

To delve deeper into the mechanism behind the change in the density of the state of N-MoS₂ after NH₃ adsorption, we calculated the PDOS of N-MoS₂ with a sulphur vacancy before

 $\mbox{\bf Table 1} \ \mbox{\bf Adsorption energy and intermolecular distances of pure } \mbox{\bf MoS}_2 \mbox{\bf monolayers}$

System	Adsorption energy (eV)	Intermolecular distance (Å)
MoS ₂ + ammonia	-0.38	5.2
MoS ₂ + formaldehyde	-0.31	5.1
MoS ₂ + TEA	-0.18	9.4
MoS ₂ + ethanol	-0.27	5.8
MoS ₂ + acetone	-0.35	5.5

Table 2 Adsorption energy and intermolecular distances for nitrogen-incorporated MoS₂ monolayers (without sulphur vacancy)

System	Adsorption energy (eV)	Intermolecular distance (Å)
MoS ₂ + ammonia	-0.76	3.9
MoS ₂ + formaldehyde	-0.45	5.7
$MoS_2 + TEA$	-0.12	9.1
MoS ₂ + ethanol	-0.53	4.7
MoS ₂ + acetone	-0.42	5.2

Table 3 Adsorption energy and intermolecular distances of nitrogenincorporated MoS₂ monolayers (with sulphur vacancy)

System	Adsorption energy (eV)	Intermolecular distance (Å)
MoS ₂ + ammonia	-0.81	3.2
MoS ₂ + formaldehyde	-0.42	5.1
$MoS_2 + TEA$	-0.18	8.5
MoS_2 + ethanol	-0.61	4.1
MoS_2 + acetone	-0.55	4.5

 NH_3 adsorption and $N-MoS_2$ with a sulphur vacancy after NH_3 adsorption. From Fig. 11(a), it can be seen that the major contribution in the valence band near the Fermi level is from the d_z^2 orbital of molybdenum atoms, followed by a contribution from the p_z orbital of sulphur atoms. In contrast, in the case of $N-MoS_2$ with S vacancy after NH_3 adsorption, the major contribution in the conduction band near the Fermi level is from the $d_{x^2-y^2}$ orbital of Molybdenum atoms, followed by a contribution from the p_y orbital of sulphur atoms (Fig. 11(b)).

3.4. Gas sensing mechanism:

The sensing mechanism of $N-MoS_2$ sensor is based on the change in the electrical resistance during the interaction of analyte vapour and the sensing material. Argon plasma treatment before nitrogen plasma may create sulfur vacancy due to

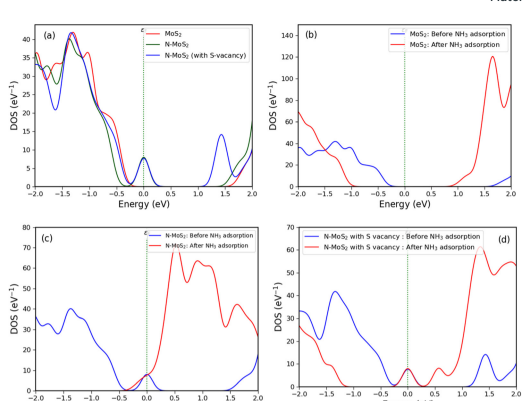


Fig. 10 Total density of state (TDOS) of the (a) MoS₂ system with N attachment and N attachment with S vacancy before NH₃ adsorption; (b) MoS₂ upon NH_3 molecule adsorption; (c) $N-MoS_2$ without sulfur vacancy upon NH_3 molecule adsorption; and (d) $N-MoS_2$ with S vacancy upon S molecule adsorption; and (d) S with S vacancy upon S molecule adsorption; and (e) S with S vacancy upon S molecule adsorption; and (e) S with S vacancy upon S molecule adsorption; and (e) S molecule adsorption S with S vacancy upon S molecule adsorption; and (e) S molecule adsorption S molecule adsorption S molecule adsorption S molecule adsorption S molecule S molecu adsorption.

Energy (eV)

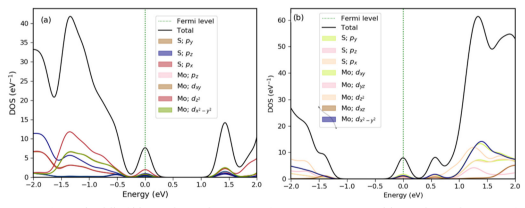


Fig. 11 Projected density of states (PDOS) of (a) N-MoS₂ with S vacancy: before NH₃ adsorption and (b) N-MoS₂ with S vacancy: after NH₃ adsorption.

the bombardment of energetic ions, contributing to the n-type characteristics of N-MoS₂. 52-54 Additionally, oxygen may get adsorbed on the sulfur vacancies, creating oxygen adsorbates, which may also contribute to the n-type behaviour of the sensor.55,56 Along with this, nitrogen plasma treatment may attach nitrogen to the MoS2, which enhances electron carrier concentration in MoS2 by introducing donor levels adjacent the conduction band, hence augmenting n-type conductivity. 57-59 Moreover, the incorporation of nitrogen can make the sensor exhibit a more stable response than a nontreated MoS₂ sensor.⁶⁰ Fig. 12 shows the plausible sensing mechanism of N-MoS2 sensor towards NH3 vapours.

Energy (eV)

The I-V characteristic plot (ESI,† Fig. S4) also shows an increase in current after plasma treatment, which is also an ntype behaviour of N-MoS₂ sensor. When the NH₃ molecules interact with the sensing material, electron transfer occurs from NH₃ to the sensing material, which changes the resistance of the sensing material.4 Hence, plasma treatment can enhance the sensing capabilities of MoS₂ sensors. The theoretical investigations are also consistent with the experimental results.

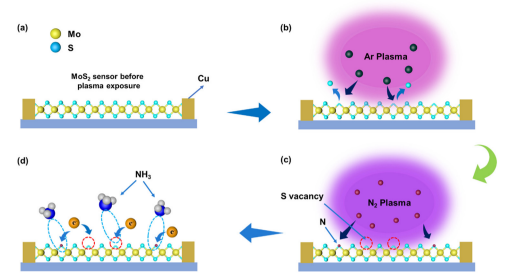


Fig. 12 (a) MoS_2 sensor before plasma treatment; (b) Ar plasma treatment on the MoS_2 sensor creating S vacancies; (c) N_2 plasma treatment on the sensor with nitrogen attachments taking place at some vacancies; and (d) interaction of NH_3 molecules with S vacancies and nitrogen attachment sites.

Table 4 Comparison of various MoS₂ sensors doped using plasma treatment and other chemical routes with this work

Sl no.	Title description	Targeted Gas/VOCs	Operating temperature	Limit of detection (LOD)	Response/recovery time	Ref.
1	Electronic modulation of MoS ₂ nanosheets via N-doping	NO_2	Room temperature	62.5 ppb	53 s/323 s (10 ppm)	28
2	Synergetic phase modulation and N-doping of MoS ₂	NO_2	Room temperature	0.13 ppm	43.1 s/301.2 s (10 ppm)	61
3	Metal-doped MoS ₂ nanoflower	SO_2	Room temperature	250 ppb	_	62
4	Zinc-doped MoS ₂ /RGO composites	NH_3	200 °C	6 ppm	21.3 s/44.9 s (50 ppm)	63
5	N dopant-triggered MoS ₂ nanosheets	NO_2	Room temperature	125 ppb	20 s/113 s (10 ppm)	37
6	Dual functionalized flower-like MoS_2 nanospheres with Pd and $g\text{-}C_3N_4$	TEA	225 °C	17 ppb	38 s/26 s (30 ppm)	64
7	N-doped MoS ₂ hierarchical structures	NO_2	25 °C	10 ppb	146 s/52 s (10 ppb)	60
8	Cu-functionalized MoS ₂ nanoworm thin films	NO_2	100 °C	2 ppm	54 s/82 s	65
9	Co-incorporated MoS ₂ nanosheets	NO_2	Room temperature	7 ppm	10 s/600 s	18
10	DC Plasma-modified MoS ₂ nanoflower	NH_3	Room temperature	80 ppb	22 s/23 s	This Work

Table 4 presents a comparison among various MoS₂-based sensors modified by plasma treatment, as well as doping by chemical methods, with the work presented here.

4. Conclusion

In summary, we fabricated a nitrogen-incorporated MoS₂ sensor following DC plasma treatment with a highly selective sensing response towards NH₃ at room temperature. MoS₂ was synthesized following a wet chemical approach following argon and nitrogen plasma treatment, which created sulfur vacancies and nitrogen incorporation into the MoS₂ matrix. The XPS spectra of the plasma-treated sensor showed an increase in nitrogen content compared with the non-treated sample. Moreover, the FESEM images also show some deformation after plasma treatment. EDX spectra also show nitrogen content. Plasma treatment enhances the number of active sites on the MoS₂ surface, which enhances the transfer of electrons from NH₃ to the active sites and increases the sensing response of

the sensor compared to the pristine MoS₂ sensor. The sensor shows a fast response (22 s) and recovery time (23 s). The sensor also shows very selective detection of NH₃ compared to other common toxic VOCs, with a limit of detection of 80 ppb. This DC plasma treatment method can be applied for the fabrication of very highly sensitive VOC sensors in a controllable manner.

Author contributions

A. K. – conceptualization, methodology, investigation, formal analysis, visualization, data curation, and writing of the original draft. B. C. – methodology (theoretical), investigation, and writing – original draft. T. H. – methodology, investigation, data curation, and writing – original draft. S. C. – investigation, data curation. B. K. – resources, investigation, H. K. – resources, conceptualization, visualization, supervision, funding acquisition, and writing – review and editing*.

Data availability

The data supporting this article have been included as part of the ESI.†

Conflicts of interest

The authors declare no competing financial interest.

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References

- 1 S. Freddi, A. V. Emelianov, I. I. Bobrinetskiy, G. Drera, S. Pagliara, D. S. Kopylova, M. Chiesa, G. Santini, N. Mores, U. Moscato, A. G. Nasibulin, P. Montuschi and L. Sangaletti, Adv. Healthcare Mater., 2020, 9, 2000377.
- 2 L. Zhang, K. Khan, J. Zou, H. Zhang and Y. Li, Adv. Mater. Interfaces, 2019, 6, 1901329.
- 3 V. Van Tran, D. Park and Y. C. Lee, Int. J. Environ. Res. Public Health, 2020, 17, 2927.
- 4 A. Kashyap, H. Sarma, B. Chakraborty and H. Kalita, ACS Appl. Electron. Mater., 2024, 6, 6916-6931.
- 5 K. Wang, L. Lee, S. L. Loo, T. Y. Yang, C. T. Chen, T. W. Kuo, J. L. Chen, H. C. Kuo and Y. L. Chueh, ACS Appl. Nano Mater., 2023, 6, 5336-5344.
- 6 W. C. Tan and K. W. Ang, Adv. Electron. Mater., 2021, 7, 1–21.
- 7 A. Kashyap, B. Chakraborty, M. S. Siddiqui, H. Tyagi and H. Kalita, ACS Appl. Nano Mater., 2023, 6, 7948-7959.
- 8 S. Acosta and M. Quintana, *Sensors*, 2024, 24, 1817.
- 9 H. Schmidt, F. Giustiniano and G. Eda, Chem. Soc. Rev., 2015, 44, 7715-7736.
- 10 W. Zheng, X. Liu, J. Xie, G. Lu and J. Zhang, Coord. Chem. Rev., 2021, 447, 214151.
- 11 A. Shokri and N. Salami, Sens. Actuators, B, 2016, 236, 378-385.
- 12 J. Chen and W. Xu, eScience, 2023, 3, 100178.
- 13 P. Li, G. Zhang, Z. Kang, X. Zheng, Y. Xie, C. Liang, Y. Zhang, X. Fang, R. Sun, Z. Liu, Y. Bu, Y. Lu and Y. Zhang, InfoMat, 2023, 5, e12457.
- 14 R. Duan, W. Qi, K. Tang and W. Liu, InfoMat, 2025, 7, e12610.

- 15 J. Zhao, Z. Chen, S. Liu, P. Li, S. Yu, D. Ling and F. Li, BMEMat, 2024, 2, e12066.
- 16 L. Luo, J. Gao, L. Zheng, L. Li, W. Li, M. Xu, H. Jiang, Y. Li, H. Wu, H. Ji, X. Dong, R. Zhao, Z. Liu, X. Wang and W. Huang, InfoMat, 2024, 6, e12605.
- 17 L. Chacko, E. Massera and P. M. Aneesh, J. Electrochem. Soc., 2020, 167, 106506.
- 18 P. Bharathi, S. Harish, M. Shimomura, S. Ponnusamy, M. Krishna Mohan, J. Archana and M. Navaneethan, Sens. Actuators, B, 2022, 360, 131600.
- 19 P. Bharathi, S. Harish, G. Mathankumar, M. Krishna Mohan, J. Archana, S. Kamalakannan, M. Prakash, M. Shimomura and M. Navaneethan, Appl. Surf. Sci., 2022, 600, 154086.
- 20 D. Zhang, Z. Yang, P. Li, M. Pang and Q. Xue, Nano Energy, 2019, 65, 103974.
- 21 M. Ikram, L. Liu, H. Lv, Y. Liu, A. Ur Rehman, K. Kan, W. J. Zhang, L. He, Y. Wang, R. Wang and K. Shi, J. Hazard. Mater., 2019, 363, 335-345.
- 22 K. Rathi and K. Pal, Adv. Mater. Interfaces, 2020, 7, 2000140.
- 23 Y. Liang, L. Zhang, K. Wang, J. Ren, L. Yu and M. Yin, Sens. Actuators, B, 2023, 378, 133137.
- 24 J. Le Fan, X. F. Hu, W. W. Qin, M. Zhou, Y. S. Liu, S. Cheng, S. J. Gao, L. P. Tan, G. Q. Wang and W. Zhang, J. Mater. Chem. C, 2023, 11, 2364-2374.
- 25 Z. Sheng, P. Qi, Y. Lu, G. Liu, M. Chen, X. Gan, Y. Qin, K. Hao and Y. Tang, ACS Appl. Mater. Interfaces, 2021, 13, 34495-34506.
- 26 P. Tao, J. He, T. Shen, Y. Hao, J. Yan, Z. Huang, X. Xu, M. Li and Y. Chen, Adv. Mater. Interfaces, 2019, 6, 1900460.
- 27 S. Deng, Y. Zhong, Y. Zeng, Y. Wang, Z. Yao, F. Yang, S. Lin, X. Wang, X. Lu, X. Xia, J. Tu, S. J. Deng, Y. Zhong, Z. J. Yao, X. L. Wang, X. H. Xia, J. P. Tu, Y. X. Zeng, X. H. Lu, Y. D. Wang, F. Yang and S. W. Lin, Adv. Mater., 2017, 29, 1700748.
- 28 K. Zhao, X. Chang, J. Zhang, F. Yuan and X. Liu, ACS Sens., 2024, 9, 388-397.
- 29 A. E. Abusrafa, S. Habib, I. Krupa, M. Ouederni and A. Popelka, Coatings, 2019, 9, 145.
- 30 C. Ma, A. Nikiforov, D. Hegemann, N. De Geyter, R. Morent and K. Ostrikov, Int. Mater. Rev., 2023, 68, 82-119.
- 31 R. V. Dabhade, D. S. Bodas and S. A. Gangal, Sens. Actuators, B, 2004, 98, 37-40.
- 32 S. Dou, L. Tao, R. Wang, S. El Hankari, R. Chen and S. Wang, Adv. Mater., 2018, 30, 1705850.
- 33 X. Ji, X. Yuan, J. Wu, L. Yu, H. Guo, H. Wang, H. Zhang, D. Yu and Y. Zhao, ACS Appl. Mater. Interfaces, 2017, 9, 24616-24624.
- 34 T. Hazarika, B. Kakati, D. Pal, R. Saikia, A. Rawal, M. K. Mahanta and S. Biswas, Sci. Rep., 2024, 14, 1-23.
- 35 L. Tao, X. Duan, C. Wang, X. Duan and S. Wang, Chem. Commun., 2015, 51, 7470-7473.
- 36 A. Azcatl, X. Qin, A. Prakash, C. Zhang, L. Cheng, Q. Wang, N. Lu, M. J. Kim, J. Kim, K. Cho, R. Addou, C. L. Hinkle, J. Appenzeller and R. M. Wallace, Nano Lett., 2016, 16, 5437-5443.
- 37 R. Wu, J. Hao, S. Zheng, Q. Sun, T. Wang, D. Zhang, H. Zhang, Y. Wang and X. Zhou, Appl. Surf. Sci., 2022, **571**, 151162.

- 38 T. Hazarika, P. Das, J. Jose, B. Kakati, D. Pal, R. Saikia and M. K. Mahanta, Polym. Adv. Technol., 2023, 34, 2862-2878.
- 39 S. J. Clark, M. D. Segall, C. J. Pickard, P. J. Hasnip, M. I. J. Probert, K. Refson and M. C. Payne, Z. Kristallogr., 2005, 220, 567-570.
- 40 C. Yang, Z. Y. Zhao, H. T. Wei, X. Y. Deng and Q. J. Liu, RSC Adv., 2021, 11, 4276-4285.
- 41 J. P. Perdew, K. Burke and M. Ernzerhof, Phys. Rev. Lett., 1996, 77, 3865.
- 42 S. Grimme, J. Comput. Chem., 2006, 27, 1787-1799.
- 43 T. H. Fischer and J. Almlöf, J. Phys. Chem., 1992, 96, 9768-9774.
- 44 S. Y. Cho, H. J. Koh, H. W. Yoo, J. S. Kim and H. T. Jung, *ACS* Sens., 2017, 2, 183-189.
- 45 Z. Chen, D. Yin and M. Zhang, Small, 2018, 14, 1703818.
- 46 Y. Zhang, Y. Jiang, Z. Duan, Y. Wu, Q. Zhao, B. Liu, Q. Huang, Z. Yuan, X. Li and H. Tai, J. Hazard. Mater., 2022, 434, 128836.
- 47 L. Feng, L. Zhang, S. Zhang, X. Chen, P. Li, Y. Gao, S. Xie, A. Zhang and H. Wang, ACS Appl. Mater. Interfaces, 2020, 12, 17547-17556.
- 48 F. Y. Meng, H. Wu, M. Qiao, X. F. Zeng, D. Wang and J. X. Wang, Langmuir, 2022, 38, 1567-1577.
- 49 L. Lin, Z. K. Yang, Y. F. Jiang and A. W. Xu, ACS Catal., 2016, 6, 4449-4454.
- 50 D. Liu, J. Pan, J. Tang, W. Liu, S. Bai and R. Luo, J. Phys. Chem. Solids, 2019, 124, 36-43.
- 51 E. S. Kadantsev and P. Hawrylak, Solid State Commun., 2012, **152**, 909-913.
- 52 J. He, Y. Wen, D. Han, P. Zeng, P. Zheng, L. Zheng, W. Su, Z. Wu and Y. Zhang, Mater. Sci. Semicond. Process., 2023, **158**, 107347.

- 53 K. Cho, M. Min, T. Y. Kim, H. Jeong, J. Pak, J. K. Kim, J. Jang, S. J. Yun, Y. H. Lee, W. K. Hong and T. Lee, ACS Nano, 2015, 9, 8044-8053.
- 54 H. Qiu, T. Xu, Z. Wang, W. Ren, H. Nan, Z. Ni, Q. Chen, S. Yuan, F. Miao, F. Song, G. Long, Y. Shi, L. Sun, J. Wang and X. Wang, Nat. Commun., 2013, 4, 1-6.
- 55 A. Wu, Q. Song and H. Liu, Comput. Theor. Chem., 2020, 1187, 112906.
- 56 G. Sun, F. Li, T. Wu, L. Cong, L. Sun, G. Yang, H. Xie, A. Mauger, C. M. Julien and J. Liu, Inorg. Chem., 2019, 58, 2169-2176.
- 57 A. A. Kotsun, V. A. Alekseev, S. G. Stolyarova, A. A. Makarova, M. A. Grebenkina, A. P. Zubareva, A. V. Okotrub and L. G. Bulusheva, J. Alloys Compd., 2023, 947, 169689.
- 58 Q. Yang, Z. Wang, L. Dong, W. Zhao, Y. Jin, L. Fang, B. Hu and M. Dong, J. Phys. Chem. C, 2019, 123, 10917-10925.
- 59 M. W. Jung, W. Song, D. S. Jung, S. S. Lee, C. Y. Park and K. S. An, J. Nanosci. Nanotechnol., 2016, 16, 2756-2759.
- 60 M. Yin, K. Wang, C. Gao, R. Yang, Y. Huang and L. Yu, Mater. Res. Bull., 2024, 179, 112943.
- 61 J. Kim, M. Li, C. H. Lin, L. Hu, T. Wan, A. Saeed, P. Guan, Z. Feng, T. Kumeria, J. Tang, D. Su, T. Wu and D. Chu, Adv. Sci., 2025, 12, 2410825.
- 62 D. Zhang, J. Wu, P. Li and Y. Cao, J. Mater. Chem. A, 2017, 5, 20666-20677.
- 63 S. P. Linto Sibi, M. Rajkumar, K. Govindharaj, J. Mobika, V. Nithya Priya and R. T. Rajendra Kumar, Anal. Chim. Acta, 2023, 1248, 340932.
- 64 W. Guo, K. Chen, S. Wang, H. Zhang and D. Wu, Sens. Actuators, B, 2025, 433, 137490.
- 65 S. Tyagi, A. Kumar, A. Kumar, Y. K. Gautam, V. Kumar, Y. Kumar and B. P. Singh, Mater. Res. Bull., 2022, 150, 111784.