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Study of ZrS_3 -based field-effect transistors toward the understanding of the mechanisms of light-enhanced gas sensing by transition metal trichalcogenides

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ABSTRACT

Extending knowledge of the properties of low-dimensional van der Waals materials, including their reactivity to the ambiance, is important for developing innovative electronic and optoelectronic devices. Transition metal trichalcogenides with tunable optical band gaps and anisotropic conductivity are an emerging class among low-dimensional structures with the possibility of gate tunability and photoreactivity. These properties can be combined into light-enhanced field-effect transistor gas sensors. We demonstrated prototype zirconium trisulfide (ZrS₃) sensors for nitrogen dioxide, ethanol, and acetone. Photoconductivity and photogating play a critical role in photoinduced gas sensing, with the dominance of the first for blue (470 nm) and green (515 nm) and the second one prevailing for red (700 nm) irradiations. Our results suggest that surface trap states lead both to trapping and scattering of the charge carriers in the channel. The gas detection is guided by charge transfer and modulation of the carrier mobility, resulting in distinct *I-V* characteristics for selected irradiation conditions.

1. Introduction

Since the discovery of graphene and its potential for applications in electronic devices, the research interest in exploring various twodimensional (2D) materials has grown enormously [1,2]. The exceptional physical and chemical properties of graphene led to the development of carbon-based components in electronic devices, including anodes in batteries and cells, supercapacitors, flexible electronics, and different types of sensors and biosensors [3–5]. The following isolation of 2D transition metal dichalcogenides (TMDCs) opened a new route of using semiconducting monolayers with the possibility of the enhanced light-matter interaction owing to the tunable optical band gaps of TMDCs – an advantage compared to zero-band gap graphene [6–8]. Low-dimensional materials, including graphene and its derivatives, carbon nanotubes, polymer nanostructures, MXenes, TMDCs, nanostructured metal oxides (MOx), and their hybrids, are widely investigated for gas sensing applications. They are reviewed in many works from the last three years, including different fabrication routes and sensing methodologies [9–15]. A relatively new class among low-dimensional structures are transition metal trichalcogenides (TMTCs) with general structural formula MX₃, where M is a transition metal from group IVB (Ti, Zr, Hf) or group VB (Nb, Ta), and X is a chalcogen atom from group VIA (S, Se, Te) [16–19]. Similar to graphene or TMDCs, TMTC monolayers can be fabricated *via* chemical or mechanical exfoliation (top-down techniques) or through chemical vapor deposition (CVD) or chemical vapor transport (CVT) (bottom-up approaches). A recent report suggests an even more straightforward route of fabrication by printing from ink solutions, which was successfully realized for TiS₃ dispersed in the mixture of ethanol and ethylene glycol [20].

The layered structure of TMTCs exhibits in-plane solid anisotropy (*e. g.*, in conducting properties) and quasi-one-dimensional (quasi-1D) electronic properties [20,21]. Moreover, quasi-1D structures are expected to exhibit interesting properties due to quantum confinement,

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such as electron correlation, spin-orbit coupling, charge density wave phenomenon, and superconductivity [22,23]. Thus, the exceptional properties of TMTCs highlight their potential in a wide range of application fields, including field-effect transistors (FETs), photodetectors, photocatalysts, Li- & Na-ion batteries, thermoelectric materials, and sensors [24]. Studies of MX₃ (M = Ti, Zr; X = S, Se) band gaps by first-principle methods revealed an indirect band gap in the range of 0.21–1.83 eV and a direct band gap in the range of 0.73–2.13 eV in the bulk, meaning these materials should effectively absorb visible to infrared radiation [25]. For instance, quasi-1D TiS₃ with \sim 1 eV band gap may constitute an alternative to silicon in miniaturized electronics (the band gap of Si is \sim 1.17 eV) [24,26].

Among other less-studied TMTCs, zirconium trisulfide (ZrS₃) is a ptype semiconductor with a resistivity of 15 Ω ·cm at room temperature in the bulk. However, nanostructured ZrS3 exhibited n-type semiconducting behavior [14,27]. Theoretical investigations revealed that ZrS3 possesses a wide direct band gap of 2.13 eV and an indirect band gap of 1.83 eV as a monolayer, offering the potential for effective visible light absorption [28]. ZrS₃ single crystals were reported to be efficient photodetectors with anisotropic photoresponse proportional to white light optical power at a temperature range of 100–300 K [25]. The same group demonstrated a wavelength-dependent photoresponse of ZrS₃ crystals with the highest responsivity to blue light of 470 nm, compared to green (540 nm) and red (670 nm) irradiation [29]. The anisotropy in the electronic properties has been experimentally investigated for ZrS₃ and TiS₃ [30]. Moreover, it was reported that an exfoliated solid solution of TiS₃ and ZrS₃ has the potential to be used as FETs sensitive to visible light with a tunable optical band gap dependent on the Ti/Zr ratio [31].

Even though TMTCs incorporated in FETs and photodetectors were studied separately, no reports on combining these two applications into light-enhanced gas sensing devices based on quasi-1D ZrS3 were reported to the best knowledge of the authors. The ambiance effect was studied for ZrS3 nanobelts, comparing the current-voltage characteristics in the dark and under selected irradiation for ambient air and vacuum conditions. A few times higher current was observed for the $\mbox{Zr}\mbox{S}_3$ thin films under vacuum, and this effect was magnified under irradiation (405 nm) [32]. Interestingly, the opposite effect was observed for ZrS_3 nanobelts fabricated via the reaction of zirconium and sulfur powders to form *p*-type semiconductors. The authors claim that the fabrication procedure influences the type of conductivity of ZrS₃ films, which may result in different effects of ambient gases [33]. For *p*-type ZrS₃, oxygen molecules in the ambient air are adsorbed onto the nanobelt surface, capturing free electrons from the material to form O₂ ions. This increases the hole concentration and amplifies the current when ZrS_3 is a *p*-type semiconductor. These observations suggest that TMTCs may be responsive toward other gases, with the detection mechanisms presumably guided by charge transfer.

Although there is a lack of reports on ZrS₃-based gas sensors, the gas responsivity of TaS₃ and TiS₃ was demonstrated. TaS₃ sensors showed selectivity toward NO compared to H₂O, N₂, CO₂, and C₆H₆ [34]. Another work reported graphene/TiS₃ heterojunctions investigated toward ethanol sensing [35]. Concentrations of ethanol between 2 and 20 ppm were efficiently detected at room temperature with 55% of relative humidity (RH). The authors observed that the sensor selectively detects polar vapors owing to the molecules' presence of –OH groups. Such a conclusion was suggested after watching much higher sensor responses toward ethanol, methanol, and acetone than small molecules such as H₂, CH₄, and CO. Additionally, the authors suggested cleaning the sensor surface at 170 °C in an argon environment after several cycles of detection to enable full recovery and higher sensing performance, showing the possibility of reversible sensing operation.

Here, we demonstrate a quasi-1D ZrS_3 -based sensor offering quantitative detection of nitrogen dioxide (NO₂), ethanol (C₂H₆O), and acetone (C₃H₆O) enhanced by visible light. We propose a FET configuration of the sensing device, where ZrS_3 film acts as the channel. The possibility of gate voltage tuning enables channel conductivity

modulation to obtain higher sensing responses than in purely DC measurements. Blue (470 nm), green (515 nm), and red (700 nm) LEDs were used for light-enhanced detection to show the influence of wavelength and optical power density on ZrS_3 sensor performance. Our work demonstrates ZrS_3 sensor performance toward selected gases *via* transfer curves and time response measurements. We discuss possible gas sensing mechanisms supporting our considerations with channel current and conductivity studies of light-enhanced sensing. Our work discusses the mechanisms underlying the sensing process of nanostructured ZrS_3 by comparing it to sensing properties exhibited by other 2D metal sulfides. We present a comprehensive study on possible modulation for ZrS_3 sensor performance enhanced by FET configuration and optical tuning by selecting LED enhancement of specific wavelength and irradiance power.

2. Materials and methods

2.1. ZrS₃ sensors fabrication

The ZrS₃ layers were mechanically exfoliated from the crystals produced by the chemical vapor transport (CVT) method using metallic zirconium and sulfur vapor in a vacuum-sealed quartz tube (2D Semiconductors). Details of the standard CVT methods are well described in the literature for TMDC and TMTC materials [30,36]. A thin film consisting of ZrS₃ nanowires (quasi 1D nanoribbons) served as a channel in field effect transistors (FETs). Golden electrodes (source and drain) were deposited to obtain ZrS₃ channels of width and length equal to 2 μ m and 1.5 μ m, respectively. Si substrate served as a back-gate contact in ZrS₃ FET.

2.2. DC characteristics measurements

DC response measurements were performed in a probe station using titanium needles to connect the ZrS3 sensor's electrodes with the measuring and biasing units. A Keithley-4200A-SCS parameter analyzer with two medium power source-measure units (type 4201-SMU) was used for the transfer and output characteristics measurements and timedomain studies. All drain-source current vs. gate voltage curves (I_{DS}-V_G) were collected in the V_G range from -30 to +30 V with a 0.1 V step and 2 s hold time. The drain-source voltage ($V_{\rm DS}$) was kept equal to 6 V. Output characteristics ($I_{\rm DS}$ - $V_{\rm DS}$) were collected in the $V_{\rm DS}$ range from -5 to +10 V. $V_{\rm G}$ was limited to 30 V. Time-domain studies were performed at $V_{\rm DS}$ = 6 V and $V_{\rm G} = 30$ V since the sensing responses were the most pronounced for the high positive gate voltage. During all measurements, the sensor was kept inside a metal box to limit interference by ambient light, power supply lines, and laboratory airflow during gas-sensing experiments. Light-assisted sensing was realized using three LEDs in the visible light range: blue ($\lambda = 470$ nm), green ($\lambda = 515$ nm), and red ($\lambda = 700$ nm). The LEDs were positioned approximately 0.5 cm from the sensor surface to obtain relatively high optical power densities, estimated as 1.44 mW/cm², 4.05 mW/cm², and 0.91 mW/cm² for blue, green, and red LEDs, respectively. Three blue LEDs (470 nm) were positioned at a fixed distance (\sim 0.5 cm) from the sensing surface to investigate the impact of different optical powers at a fixed wavelength. The diode current was changed between 5 mA and 30 mA, which increased the optical power density from 0.93 mW/cm² to 4.30 mW/cm². For more details on the types of LEDs used in our experiments and their optical properties, see Table A.1 in Supplementary Material.

2.3. Gas-sensing experiments

A metal pipe connected to the gas distribution system was placed within 0.5 cm from the sensor surface to transport selected target gases. Nitrogen dioxide (NO₂), ethanol (C_2H_6O), and acetone (C_3H_6O) gases were used for sensing experiments (all diluted in N₂). Dry synthetic air (S.A.) was used as a reference gas and to obtain selected concentrations of target gases by mixing them with S.A. at specific proportions. During all gas sensing measurements, we kept a constant overall gas flow of 50 mL/min regulated by mass flow controllers (Analyt-MTC, model GFC17). Three gases were selected for sensing experiments using a quasi-1D ZrS₃ sensor. We chose nitrogen dioxide (NO₂), one of the toxic gases harmful to both humans and the environment, even at low concentrations, as a representative inorganic gas [37]. In addition, we selected two organic gases: ethanol (C₂H₆O) and acetone (C₃H₆O) vapors, to test the sensor's selectivity by utilizing DC measurements. Both ethanol and acetone are substances that, if present in an exhaled breath at specific concentrations, can provide crucial information for health monitoring and medical diagnosis [38-40]. The DC characteristics were collected after stabilizing the sensor response in the selected environment/irradiation conditions for ~10 min. All experiments were conducted at room temperature (RT \sim 23 °C) and ambient pressure (\sim 1 bar).

3. Results and discussion

We investigated the ZrS₃-based sensor fabricated in a FET configuration according to the scheme provided in Fig. 1a. For all gas sensing experiments, the length and width of the FET channel amounted to 1.5 μ m and 2 μ m, respectively, yielding 3 μ m² of a sensor's active area (Fig. 1b). Fig. 1c and Fig. 1d present the transfer and output characteristics of the ZrS₃-based FET sensor in the reference atmosphere of S.A. in the dark and under selected LEDs from the visible light spectral range. Both characteristics indicate that the ZrS₃ layer exhibits *n*-type semiconducting behavior. Fig. 1c shows a wide hysteresis, which is more pronounced for blue and green irradiation than for red light and dark conditions. Applied irradiation induces photocurrent in the layer, increasing the overall current flowing through the ZrS₃ channel. As nanostructured ZrS₃ was reported to have an optical bandgap of ~1.8–1.9 eV, the structure should efficiently absorb wavelengths below ~690 nm. The longer wavelengths may be absorbed if any discrete defect levels are present below the conduction band, which can be observed for the defective or doped semiconductors. Thus, one can see that the red light (0.91 mW/cm² of optical power density) has the lowest impact on the *I*_{DS} compared to the blue and green irradiation, presumably owing to the mid-gap energy levels in ZrS₃. Interestingly, the green light increases the current the most. It confirms that ZrS₃ response to irradiation depends on both wavelength and optical power of the light source, as the optical power density for the green LED exceeded the value for the blue LED almost three times (1.44 mW/cm² and 4.05 mW/cm² for blue and green LED, respectively).

Fig. 2 depicts IDS-VG characteristics of the ZrS3 sensor toward different concentrations of NO₂ (4-16 ppm) in four cases of light enhancement: in the dark, under red (700 nm), green (515 nm), and blue (470 nm) irradiation. In all cases, the curves shift downwards - the current decrease means the sensing layer increases its resistance, which confirms the oxidizing ability of NO₂ molecules for n-type ZrS₃. As seen from Figs. 2c and 2d, blue and green irradiation increases I_{DS} noticeably, magnifying the hysteresis and stabilizing the sensor response, which can be more vulnerable to current fluctuations at the level of a few nA (dark and red-light case). Thus, blue or green light enhancement is suggested for providing conditions that enable a more stable and easily measurable response (at tens or hundreds of nA) and facilitate room-temperature sensor operation. Analogous sensing experiments were conducted for 30-90 ppm of ethanol and 20-35 ppm of acetone (Figs. A.1 and A.2). In contrast with NO₂, both organic gases increased I_{DS} (equally decreased sensor resistance), suggesting their reducing character toward the ZrS₃ layer. Such direction of changes induced in the sensor was reported before for ethanol and acetone detection by TiS3 flakes deposited as a

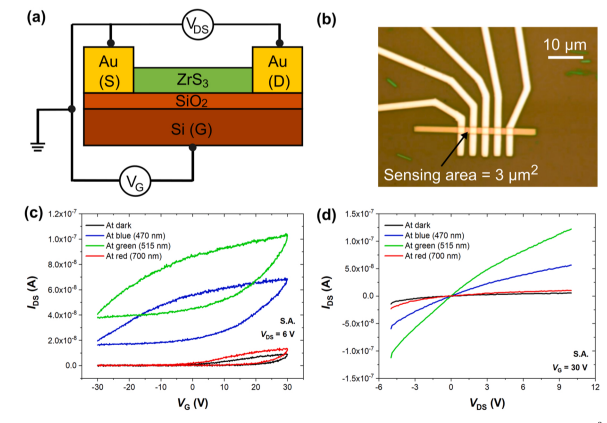


Fig. 1. (a) Schematic representation of the ZrS₃-based sensor, and **(b)** microscopic photograph of the investigated sensing structure with a $3 \mu m^2$ sensing area between each pair of electrodes. Reference DC measurements of the ZrS₃ FET sensor in synthetic air (S.A.): **(c)** transfer curve (drain-source current I_{DS} as a function of gate voltage V_G) with drain-source voltage V_{DS} biased to 6 V, and **(d)** output curve (I_{DS} vs. V_{DS}) with V_G biased to 30 V in the dark and under blue (470 nm, 1.44 mW/ cm²), green (515 nm, 4.05 mW/cm²), and red (700 nm, 0.91 mW/cm²) irradiation.

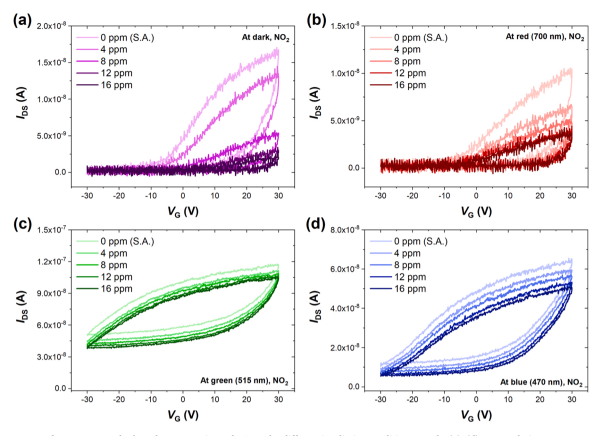


Fig. 2. ZrS_3 sensor performance toward selected concentrations of NO₂ under different irradiation conditions: graphs (a)–(d) present drain-source current I_{DS} vs. gate voltage V_G for NO₂ detection in the dark and under red (700 nm, 0.91 mW/cm²) green (515 nm, 4.05 mW/cm²), and blue (470 nm, 1.44 mW/cm²) light, respectively.

channel between Au electrodes [35]. Moreover, similarly to the NO₂ case, blue and green irradiation stabilize sensor responses toward ethanol and acetone. To address the selectivity of the sensor, we compared the sensor response to the same concentration of ethanol and acetone (30 ppm). The relative change of $I_{\rm DS}$ (the sensor response) under green irradiation is ~4.7% for 30 ppm of ethanol and ~9.2% for 30 ppm of acetone. The difference in values for the blue light case is even more apparent, estimated as ~2.8% and ~14% for 30 ppm of ethanol and acetone, respectively. Such observations suggest a higher affinity between acetone molecules and ZrS₃ structure than for ethanol molecules.

Fig. 3 summarizes I_{DS} changes for selected nitrogen dioxide, ethanol, and acetone concentrations. IDS values were derived from corresponding $I_{\rm DS}$ - $V_{\rm G}$ characteristics for $V_{\rm G} = 30$ V. Dashed lines designate the linear fit functions, with R² values indicating the quality of the linear fitting. The R^2 coefficient represents how well the linear model predicts the experimental data, and R² values close to 1 show the high linearity of the gassensing responses. In the case of NO2, high linearity in sensor response was observed for blue and green irradiation. For ethanol and acetone, all types of irradiation and dark conditions resulted in similarly high linearity of the sensor response. The opposite direction of I_{DS} changes for inorganic NO2 and organic ethanol and acetone can be explained by each molecule's acceptor or donor properties. Generally, the gas detection mechanism for two-dimensional metal sulfides relies primarily on charge transfer between the sensing material and analyte, as reported for nanostructured MoS₂ – the leading representative of the 2D materials with tunable band gaps [14]. The amount of charge donated or extracted from the material by the adsorbed molecules affects the electron carrier density of the semiconducting layer, altering its resistance. Additionally, the FET configuration of the sensor modulates the electric field across the dielectric layer, influencing the channel conductivity.

Apart from that, two main effects were suggested to explain the

changes in I-V characteristics for light-activated FETs consisting of lowdimensional structures. The first, photoconductive effect, results in the induced photocurrent in the channel due to generated charge carriers during irradiation; the second, photogating (photovoltaic) effect, leads to a change in FET threshold voltage, shifting the *I-V* sideways [41,42]. For visible light-enhanced MoS2-based sensors, the dominant photogating effect was demonstrated, which resulted in a negative shift of FET threshold voltage $V_{\rm th}$ for increased irradiation [43]. However, according to the standard charge transfer model of gas detection for metal sulfides, electrons extracted from the sensing material during NO₂ adsorption should shift $V_{\rm th}$ toward positive values without changing the shape (I-V hysteresis slope) of the I_{DS} - V_G curve. Since the channel conductivity depends on both carrier concentration and mobility, the changes introduced in both factors may alter the current flowing through the sensor. The abovementioned observations obtained for NO₂ sensing by MoS₂ were ascribed to the carrier mobility change induced by adsorbed target gas molecules acting as scattering centers for channel electrons. The scattering of electrons reduces their mobility resulting in the drain-source current decrease. At the same time, we cannot exclude the fact that adsorbed molecules may capture electrons during photoexcitation, limiting their concentration, which diminishes the channel current. In the case of our gas sensing results, the change of hysteresis shape is visible primarily for dark and red-light conditions, whereas for green and blue irradiation, the IDS level increases one or two orders of magnitude, but the hysteresis shape is almost unaltered. This suggests that photoinduced charge carriers significantly increase channel current resulting in a positive photoconductivity, so the photoconductive effect play an essential role in the gas detection under blue and green light. At the same time, one can notice the simultaneous sideways shift and the change of hysteresis shape, especially for dark and red-light conditions. Thus, we conducted subsequent studies to develop more detailed

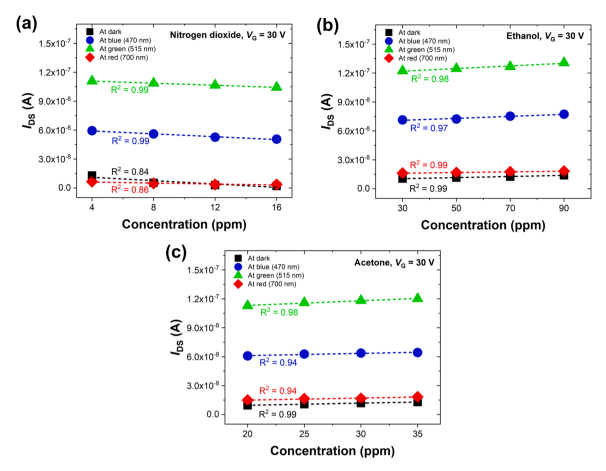


Fig. 3. Comparison of ZrS_3 sensor performance toward different gases: drain-source current I_{DS} response for selected concentrations of **(a)** nitrogen dioxide (NO₂), **(b)** ethanol, and **(c)** acetone under different irradiation conditions. I_{DS} values were taken for the $V_G = 30$ V. V_{DS} was biased to 6 V in all experiments. Dashed lines depict linear functions fitted to the experimental points.

conclusions on ZrS_3 -based gas sensing including the effect of electron concentration and mobility change and taking photoconductive and photogating effects into consideration.

As blue and green light enhanced the ZrS_3 sensor response noticeably, we chose these two LEDs for further gas sensing experiments. Fig. 4a demonstrates time response studies of the ZrS_3 sensor toward repeatable cycles of introducing 16 ppm of NO₂ under green and blue irradiation. The transient curves were collected for V_{DS} and V_G biased to 6 V and 30 V, respectively. Again, we confirm that green light increases $I_{\rm DS}$ more significantly despite a longer wavelength, which can be ascribed to its higher optical power. To check whether optimizing the sensor response is possible by adjusting LED optical power, we used three blue LEDs (470 nm) connected in series to a voltage supplier for sensor irradiation to obtain a wider range of irradiance adjusted by the diodes' polarization current. Fig. 4b shows cycles of 16 ppm of NO₂ introduction during irradiation with blue LEDs of selected optical power densities (marked on the graph with dashed boxes indicating periods for selected irradiation conditions). Increasing optical power densities in

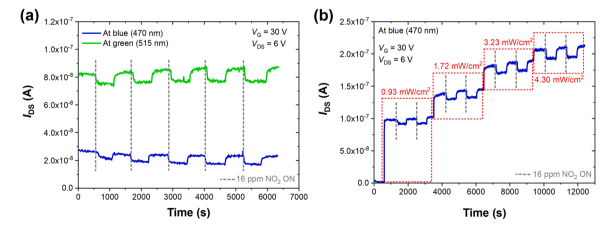


Fig. 4. Time response studies of the ZrS_3 sensor for consecutive cycles of introducing 16 ppm of NO₂ (a) under continuous irradiation using green (515 nm, 4.05 mW/cm²) or blue (470 nm, 1.44 mW/cm²) LED and (b) under continuous blue light (three LEDs) with adjusted optical power density from 0.93 mW/cm² to 4.30 mW/cm² (dashed boxes indicate cycles of NO₂ detection for specific optical power densities with values marked in the figure).

the range of $0.93-4.30 \text{ mW/cm}^2$ led to rapid baseline current growth, and the overall increase of I_{DS} change during exposition to target gas. By examining different optical parameters of LEDs used in our studies, we can confirm that ZrS3 sensor response toward NO2 depends on the wavelength and optical power density, which provide different methods of ZrS₃ performance modulation for optimization of sensor operation in ambient conditions (room temperature and atmospheric pressure). Moreover, the average sensor response (the relative change of I_{DS} before and after exposition to NO₂) did not vary much for increasing irradiance yielding 8.10%, 6.21%, 6.21%, and 7.12% for their optical radiation powers of 0.93 mW/cm², 1.72 mW/cm², 3.23 mW/cm², and 4.30 mW/ cm², respectively. This observation suggests that different optical parameters of LEDs mainly affect the baseline current for measurement modulation but do not significantly influence the detection process. The constant or weak dependence between sensor response and irradiation power may imply carrier mobility modulation during light-enhanced sensing, according to [43], since in a purely charge-transfer mechanism, the sensor response should vary more significantly with increasing optical power density. However, comparing relative sensor responses between repeatable detection cycles for green and blue light, a significant difference between average responses can be observed, amounting to 26.12% for blue and 9.96% for green light. Here, an almost three times higher response for blue light-enhanced sensing stems from a significant difference in the current baseline and indicates that the photoconductive effect shifting DC characteristics upwards becomes a relevant factor when comparing LEDs of varying wavelengths and optical powers, and the sensor response should be normalized to compare the results reliably. Additionally, based on the time response studies, one could see the high stability of the sensor response both to green or blue irradiation and to 16 ppm of NO₂. Despite the very low measured currents (maximum tens or hundreds of nA), the sensor baseline remains stable, indicating the short-time stability of the device with reduced short-time drift. We want to note that the sensor remained responsive to selected gases under lighting conditions for around two months of ongoing measurements, suggesting long-time stability of the sensing structure.

Undoubtedly, the photocurrent induced during sensor irradiation increases the overall IDS noticeably while comparing dark and green conditions, as seen in Fig. 2 and Fig. 4a. Additional irradiation emphasizes the photoconductive effect due to a high number of generated excess carriers. However, as already mentioned, photoinduced carriers may be captured or scattered during gas adsorption. The trap centers may be present due to defects, adsorbents, or dopants and guide the photoconductive and photogating effect [44]. According to [42], the photogating effect arises from charge trapping occurring at the surface binding sites for monolayered MoS₂. Thus, in different ambiances, V_{th} is likely to shift due to capturing or donating electrons by surface states to the target molecules adsorbed on the active material. In our studies, all three gases shift the I_{DS}-V_G curve sideways noticeably for dark conditions and under weak red irradiation enhancement. In general, if the trap centers capture the electrons, the I_{DS} - V_G curve shifts to the right, whereas if the trap centers capture the holes, the opposite effect with the shift to the left is observed. For the ZrS₃ sensor, the shift is positive for NO₂ but negative for ethanol and acetone (Figs. A.1, A.2), which agrees with the acceptor properties of NO2 and donor properties of both organic gases. Moreover, the higher the gas concentration, the greater the $V_{\rm th}$ shift, making quantitative detection possible for the ZrS₃ sensor. The direction of the shift conforms with the charge transfer mechanism. However, the additional change of hysteresis shape implies that carrier mobility modulation also occurs during molecular detection, presumably due to the scattering of carriers. On the other hand, for gas sensing under blue and green irradiation, the shape of the transfer curves remains almost unaltered for different concentrations of all three gases, whereas the visible change in I_{DS} level occurs. Such results may indicate the dominant photoconductive effect responsible for detection processes in these irradiation conditions. For a specific wavelength range and

under sufficient irradiance, the capture and release of carriers by trap states guide the adsorption/desorption processes and dominate over scattering on surface defects and adsorbed molecules. On the other hand, without sufficient light enhancement (dark conditions or longer wavelengths), the lower number of charge carriers flowing through the channel may be more susceptible to scattering, which reduces their mobility making the photogating effect with a change of hysteresis shape dominant.

Furthermore, the dominance of the photoconductive effect for blue or green light-enhanced gas sensing (irradiation of shorter wavelengths) does not entirely exclude the impact of the photogating effect on NO2 detection by quasi-1D ZrS3 since the simultaneous occurrence of both phenomena is the most probable case. Therefore, we conducted additional measurements for various concentrations of NO₂ (2–16 ppm) under blue irradiation of the highest optical power density (4.30 mW/ cm²). We transformed obtained results into sensor conductance response (S_G) derived from I_{DS} values at $V_G = 30$ V and V_{DS} biased to 6 V. We define S_G as a relative change of sensor conductance G in relation to sensor conductance G_0 measured in the reference atmosphere of S.A. – $G - G_0 | / G_0$. It was previously reported for MoS₂ and reduced graphene oxide (rGO) that sensor response depends on gas concentration according to the Langmuir equation, revealing the saturation of the response for a specific concentration threshold [43,45]. In those cases, the sensor response relied on the surface coverage of target molecules on the adsorption sites, which eventually saturated. A similar effect is visible in Fig. 5a for NO₂ concentrations exceeding 8 ppm. The linear relationship between the inverse conductance response and concentration may imply that visible response changes are solely due to channel conductivity change, and any other effects are negligible. According to these considerations, the deviation from the linear dependence between $1/S_{\rm G}$ and $1/{\rm Concentration}$ in the case of the ZrS₃ sensor (Fig. 5b) suggests the coexistence of more than one process responsible for detection, which agrees with our previous assumptions.

4. Conclusions

We investigated a FET-configured gas sensor based on quasi-1D ZrS₃ for qualitative and quantitative detection of nitrogen dioxide, ethanol, and acetone under ambient conditions. Our results show that light-activated ZrS₃ sensor performance can be easily optimized by adjusting the optical parameters of LEDs to ensure high sensitivity and stability of the sensor at room temperature while keeping low-power consumption of the device. This way, we demonstrate another material from the family of 2D metal-chalcogen compounds for light-enhanced gas sensing, proposing the possibility of sensing performance tunability with the FET configuration of the device and optimization of the photoactivation process.

Our gas sensing studies reveal the possibility of up to two orders of magnitude enhancement (from ~1 nA to hundreds of nA) of sensor channel current under green and blue irradiation with irradiances reaching ~4 mW/cm². Such photoexcitation provides the current baseline less susceptible to random fluctuations and a more stabilized sensor response toward ambient gases. Even though the sensor does not exhibit an excellent selectivity to any of the investigated gases, the direction of $I_{\rm DS}$ changes enables the discrimination between NO₂ and organic gases due to their redox properties. Moreover, comparing the sensor response toward the same concentration of ethanol and acetone (30 ppm) supports distinguishing between these two organic gases (Figs. A.1 and A.2). Time-response studies confirmed the sensor stability and repeatability for 16 ppm of NO₂ detected under green and blue irradiation without short-time drift of channel current.

A few aspects should be considered in the discourse on gas sensing mechanisms by quasi-1D ZrS_3 presented in our work. First, the coexistence of photoconductive and photogating effect presumably guide the surface processes during the light-enhanced detection of selected gases. Depending on the external irradiation, mainly the optical parameters of

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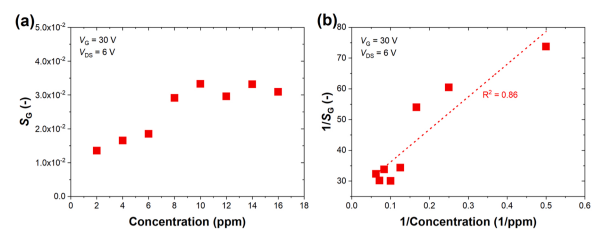


Fig. 5. (a) ZrS₃ sensor response derived from the relative changes of conductance S_G vs. NO₂ concentration (2–16 ppm), and (b) the inverse conductance response vs. the inverse NO₂ concentration. Conductivity values were derived from the transfer curves for $V_G = 30$ V and V_{DS} biased to 6 V under blue irradiation of the highest optical power density (4.30 mW/cm²).

the used LEDs, one of these effects becomes dominant. Although there is no photoexcitation in the dark, the change in the threshold voltage still occurs due to the interaction of channel electrons with gas molecules acting as trapping and scattering centers after adsorption. For the red irradiation, the wavelength of the LED (700 nm) may be insufficient to excite electrons directly from the valence band to the conduction band, but in case of any discrete levels in the band gap (defect levels, midgaps), some excess carriers may participate in molecular detection with a high possibility of being scattered on the adsorbed molecules. For blue and green irradiation, the shorter wavelengths allow photoactivation of the material providing high photoconductive gain and the subtle shift of the threshold voltage of FET. The dominance of the photoconductive effect does not eliminate the photogating in this case; furthermore, the impact of the channel electrons scattered on the adsorbed molecules can be tentatively confirmed by (i) the change of hysteresis shape (I-V hysteresis slope), (ii) low dependence between optical power and sensor response, and (iii) non-linear relationship between the inverse sensor conductance response and gas concentration.

Our findings shed light on light-driven gas sensing mechanisms by quasi-1D ZrS₃, broadening the possibility of using transition metal trichalcogenides in electronic and optoelectronic real-environment applications. Our work also broadens the understanding of surface science among 2D materials in general, extending the range of 2D (quasi-1D) structures as potential candidates for optimized gas sensing.

CRediT authorship contribution statement

J.S. and S.R. conceived the study. K.D. established and ran the experimental electrical and optical studies, evaluated analysis, and proposed interpretation. A.R. and M.W. participated in the fabrication and characterization of the samples. L.B. supervised sample preparation. A.B. contributed to data analysis. K.D., J.S., S.R., and G.C. wrote the manuscript. All authors had an opportunity to edit the manuscript.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

Data availability

Data will be made available on request.

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Appendix A. Supporting information

Supplementary data associated with this article can be found in the online version at doi:10.1016/j.mtcomm.2023.105379.

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