

RESEARCH: Short Communication

A strain tunable single-layer MoS₂ photodetector

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Strain engineering, which aims to tune the bandgap of a semiconductor by the application of strain, has emerged as an interesting way to control the electrical and optical properties of two-dimensional (2D) materials. Apart from the changes in the intrinsic properties of 2D materials, the application of strain can also be used to modify the characteristics of devices based on them. In this work, we study flexible and transparent photodetectors based on single-layer MoS_2 under the application of biaxial strain. We find that by controlling the level of strain, we can tune the photoresponsivity (by 2–3 orders of magnitude), the response time (from <80 ms to 1.5 s), and the spectral bandwidth (with a gauge factor of 135 meV/% or 58 nm/%) of the device.

Introduction

Tuning the characteristics of circuit components with an external knob is at the deep core of modern electronics. Good examples of that are the field-effect transistors whose conductance can be adjusted by means of an applied voltage to the gate electrode [1–5]. The development of new tuning knobs have undoubtedly opened up possibilities to design new device concepts [6-11]. Strain engineering provides a powerful route to modify the electrical and optical properties of electronic materials and thus it has the potential to become an excellent external tuning knob [12–15]. Conventional strain engineering approaches, however, typically yield fixed strain levels. This has radically changed with the isolation of 2D materials, which provide an excellent platform for strain engineering as they can be easily stretched and bended to a large extent in a reversible way [16–20]. Moreover, optical spectroscopy techniques have demonstrated that tensile uniaxial or biaxial strain effectively decreases the bandgap in atomically thin transition metal dichalcogenides [21-27].

Previous works on 2D-based flexible photodetectors subjected to mechanical deformations don't provide any direct measurement that ensures that the deformation of the substrate is effectively translated into strain in the 2D material and they don't characterize the effect of the mechanical deformation of the photodetector on their photocurrent spectra [28,29]. Drawing conclusions about the strain-dependent performance of 2Dbased photodetectors from those results is thus challenging.

Here, we exploit the strain tunable bandgap of single-layer molybdenum disulfide (MoS_2) to fabricate a photodetector device whose responsivity, response time, and spectral bandwidth can be adjusted by an externally applied biaxial tensile or compressive strain. We find a remarkably large strain sensitivity of the cut-off wavelength of 58 nm/% of strain (~135 meV/%), making it possible to extend the detection spectral bandwidth with respect to pristine unstrained devices. The method presented here is general and can be applied to photodetectors based on other 2D materials. The case of black phosphorus, for example, could be particularly interesting as a larger strain-dependent band gap tuning and an opposite strain dependence are expected [30,31].

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Results and discussion

We fabricate single-layer MoS₂ photodetectors with a simple photocell (or photoresistor) geometry (Fig. 1a). Single-layer MoS₂ is prepared by mechanical exfoliation of bulk molybdenite (see Materials and Methods for details) and the exfoliated flakes are then transferred to a Gel-Film (from Gel-Pak®) substrate. The single-layer regions are identified by micro-reflectance spectroscopy [32,33] and quantitative analysis of transmission mode optical microscopy images. The selected single-layer flakes are then transferred onto a polycarbonate (PC) substrate with prepatterned drain-source electrodes by a dry-transfer deterministic placement method [34,35] and are annealed at 100 °C to improve the electrical contact between the flake and the electrodes. Fig. 1b shows an optical microscopy image of one of the assembled single-layer MoS₂ photodetector devices fabricated onto PC. Note that it has been previously demonstrated that polymeric substrates, such as PC, polyimide, and polydimethylsiloxane (PDMS), can be used to integrate photonic devices [36-38]. Polycarbonate has been used as substrate because of the trade-off between high Young's modulus and large thermal expansion that allows one to biaxially stretch (or compress) the MoS₂ by moderately warming up (or cooling down) the substrate while ensuring an excellent and homogeneous strain transfer

[39] (see Supplementary Material Fig. S1 to observe the spatial homogeneity of the transferred strain). Substrates with larger thermal expansion but lower Young's modulus cannot effectively transduce their thermal expansion into biaxial strain, as predicted by finite elements analysis (see Supplementary Material Fig. S2). In the case of polycarbonate (E = 2.5 GPa), the calculated efficiency is larger than 80%. The mechanical model, which lacks atomistic details, gives only a coarse estimation of the strain transfer efficiency, and for this reason, we assume in the rest of the paper a perfect transduction of thermal expansion to biaxial strain. Note that by assuming perfect transduction, the gauge factors obtained in this work can be considered as lower bound limits. Fig. 1c shows differential reflectance spectra acquired on a single-layer MoS₂ device at different tensile strain levels from 0% (substrate at room temperature, T = 25 °C) to 0.48% (substrate heated at T = 100 °C). The A and B exciton peaks in the reflectance spectra redshift upon increasing the substrate temperature (and thus the biaxial tensile strain), indicating a narrowing of the MoS₂ bandgap. Fig. 1d shows the relationship between the substrate temperature increase and its biaxial expansion. We address the reader to Ref. [39] and the Supplementary Material (Fig. S3) for details about the measurement of thermal expansion and the calibration of the applied biaxial strain.



FIGURE 1

(a) Schematic picture of the setup used to perform all the measurements. The MoS_2 photodetectors are placed on a thermal stage and illuminated from the top. (b) Optical transmission photograph of a single-layer MoS_2 photodetectors fabricated on polycarbonate (PC). (c) Differential reflectance spectra measured at different temperatures (vertically shifted by 0.08 to facilitate their comparison). Inset: wavelength of the exciton A as a function of temperature. (d) Calibration of the polycarbonate expansion/compression as a function of the temperature in the range from -200 °C (corresponding to a compression of -1.48%) to +100 °C (corresponding to a tension of +0.48%). The line in (d) indicates the best-fit of the experimental data to a linear trend.

In order to test the spectral response of the MoS₂ photocell detectors, we illuminate the devices with a tunable light source (Bentham TLS120Xe) to select the wavelength (with a full-width at half maximum of ~10 nm), while the current across the device (biased at 10 V) is measured. The light is focused into a spot of 400 μ m of diameter with a power density of 8 mW/cm². At each step during the wavelength sweep, we measure both the dark and illuminated current to rule out drifts during the measurement. The responsivity *R* of the device can be extracted from the photocurrent values as

$$R = \frac{I_{\rm ph}}{P_{\rm dens} \cdot A_{\rm dev}}$$

where P_{dens} is the incident light power density and A_{dev} is the area of the MoS₂ channel. The biaxial strain applied to the device is controlled through a thermal stage (see Materials and Methods).

The responsivity spectrum measured for the unstrained device is comparable with the data reported in literature for MoS_2 phototransistors fabricated by electron beam lithography on SiO_2/Si substrates at similar illumination power density and biasing conditions (larger responsivities are reported for devices at very low illumination levels and upon much larger drain-source electric

field biasing conditions) [40-42]. The spectra clearly show two peak features, which are in good agreement with the B and A excitonic resonances also observed in the reflection spectra of single-layer MoS_2 [43–45]. At longer wavelengths (after the A exciton peak) the spectra presents an abrupt drop of the device responsivity. When the polycarbonate substrate is biaxially strained, the MoS₂ photocell responsivity spectrum changes significantly. In Fig. 2a, the overall responsivity values increase by a factor of ~ 100 when the strain is increased from -0.8% to 0.48%(a factor of ~ 1000 for device #2 where the strain ranges from -1.44% to 0.48%). We attribute this effect to the applied strain as similar photocells fabricated on SiO₂ (which have negligible thermal expansion) do not show this strong enhancement of the photoresponse (see a comparison between devices fabricated on SiO₂/Si and on PC in the Supplementary Material, Figs. S4 and **S5**).

To get an insight about the tensile strain-induced increase in the overall responsivity, we studied the response time of the devices. For MoS_2 photodetectors, there are two dominant mechanisms for the photocurrent generation (photoconductance and photogating) that can be easily identified through the response time of the devices [40,46–48]. In the photoconductance, the



FIGURE 2

(a) Responsivity spectra of the single-layer MoS_2 photodetector #1 obtained by measuring under 5 different strains applied (from -0.80% to 0.48%). Each dot corresponds to the value measured under light power of 8 mW/cm² and applying a bias voltage of 10 V. Note that the responsivity values for +0.16%, -0.16%, -0.48% and -0.80% have been multiplied by 1.7, 6, 10, and 10, respectively, to facilitate the comparison between the spectra. Inset: Response time for different strains applied. (b) Cut-off wavelengths extracted from the responsivity spectra of three single-layer MoS_2 photodetectors (#1, #2, and #3) at different strain ranges. Inset: Exciton A wavelengths extracted from the same spectra. (c) Response of the photodetector device to strain changes, cycling the device between two different strain levels 0.16% and 0.48\%, the cycles are measured with an applied voltage of 5 V, and the 740-nm light source density power is 5 mW/cm². (d) Zoom in the third strain cycle of (c) in order to appreciate the rise and fall time of the device, estimated with the 10–90% criterion.

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photoexcited electrons and holes are separated through the bias voltage, leading to an increase in the current flowing through the semiconductor channel. The typical response time of photoconductive devices is <10 ms. In the photogating mechanism, the photoexcited electrons are drifted by the bias voltage to the drain electrode (with a typical timescale of \sim 10–100 ns) and holes are trapped (with a typical timescale of 10 ms-10 s). Due to charge conservation in the channel, one new electron has to jump from the source to the channel once the drifting electron reaches the drain electrode, leading to a photoconductive gain that is proportional to the ratio between the electron drifting time and the hole trapping time (which can be extracted from the response time of the device to pulsed illumination). Therefore, photoconductive-dominated devices have a fast response but low responsivity values, while photogating-dominated ones are typically slow but present ultrahigh values of responsivity.

The response time to pulsed illumination for devices under compressive strain is very sharp (the response is faster than that of our experimental setup, 80 ms, compatible to what one would expect for a photoconductive-dominated device). Tensilestrained devices, on the other hand, show much slower response times ~ 1.5 s (Fig. 2a inset), indicating the presence of long-lived charge traps that could explain the large increase in responsivity upon tension through the gain associated to the photogating mechanism. We rule out the effect of the temperature as photodetectors fabricated on SiO₂, with negligible thermal expansion, do not show a sizeable variation of the response time in the temperature range of 25-100 °C (see Fig. S6 in the Supplementary Material). Therefore, biaxial strain seems to be responsible of a change of the photocurrent generation mechanism from photoconductive (for compressive strain) to photogating (for tensile strain), although the microscopic mechanism is still unknown, and it will be subject of further study. Note that we also studied the effect of strain on the Schottky barrier height through scanning photocurrent (see Figs. S7 and S8 in the Supplementary Material), finding that the Schottky barrier height in our devices is very small (~14 meV for pristine unstrained devices) and they show a moderate decrease in the barrier height upon tensile straining. This mechanism, although small, also contributes to the increase in photoresponse observed for tensile-stressed devices.

The excitonic features, visible in all the photoresponse spectra of strained MoS_2 , are redshifted (blueshifted) when increasing

the tensile (compressive) strain value with a strain gauge factor of 31 nm/% (~94 meV/%) for exciton A (inset Fig. 2b). The shift of the exciton peaks with strain in the responsivity spectra is in good agreement with predictions based on density functional theory calculations [49], with the shift observed in differential reflectance (Fig. 1c and Ref. [39]) and photoluminescence measurements on pressurized blisters of MoS₂ [27]. The observed shift of the A exciton upon straining (for 3 different devices) with a constant gauge factor over the whole strain range studied here indicates that strain is being transferred to the single-layer MoS₂ flake without slippage. Moreover, one might wonder about the presence of buckling upon compressive strain, but the critical (or minimum) strain that is necessary for the buckling to occur is $\varepsilon_c = 0.25 \cdot (3 \cdot E_{substrate}/E_{flake})^{2/3}$ [50,51]. In our case, a compressive strain of ~-2.7% is needed for buckling to occur.

The cut-off wavelength is shifted as well upon straining, as can be seen in Fig. 2b. The device shows a strain gauge factor of the cut-off wavelength in the range of ~58 nm/% (~135 meV/%) of biaxial strain. Therefore, we demonstrated that applying tensile biaxial strain to the MOS_2 device can be an effective strategy to increase both the responsivity and the wavelength bandwidth of the photodetector (at the expense of a slower response time), while compressive strain can be exploited to yield faster photodetectors (although with a lower photoresponse and with a narrower wavelength bandwidth). This adaptable optoelectronic performance of this device can be very useful to adjust the photodetector operation to different lighting conditions, similarly to human eye adaptability (scotopic vision during the night vs. photopic vision during the daylight) [52].

A direct consequence of the strain-induced redshift for tensile biaxial strain is that one can achieve a sizeable response for wavelengths even beyond the cut-off of pristine (unstrained) MoS₂. Fig. 2c shows an example where the photocurrent of the device (with light of 740 nm with a power density of 5 mW/cm², applying a bias voltage of 5 V) is measured while changing the strain level in time. One can appreciate how the photocurrent at 740 nm increases substantially upon increasing the strain level of the device (from 0.16% to 0.48% of tensile biaxial strain) in a reproducible way, which can be attributed to the strain-induced redshift of the device cut-off wavelength. Interestingly, the device can be strain tuned rather quickly (in ~20 s time scale, see Fig. 2d) and this tuning time most likely is only currently limited by the thermalization time of our temperature stage, and it



FIGURE 3

Photodetector #4 responsivities for 740-nm LED light with light power of 12 mW/cm² and applying a bias voltage of 10 V measured for several strain cycles.

could be further improved by employing micro-heaters located underneath the photocell device.

In order to study the reproducibility of the redshift in the responsivity spectra of single-layer MoS_2 photodetectors, several cycles of application/release of tensile strain were performed in another MoS_2 photodetector. Fig. 3 shows the device responsivity for 740-nm illumination wavelength (power density of 12 mW/cm^2 and bias voltage of 10 V) recorded during some of the straining cycles. We observe how the responsivity evolves from negligible values (within the experimental setup noise level) for the unstrained device toward increasingly high values for the tensile-strained devices, similarly to what was displayed for few cycles in Fig. 2c. The device shows a good reproducibility during the whole range of cycles (up to 40 cycles were applied).

Conclusions

In summary, we have harnessed the strain tunability of the band structure of single-layer MoS₂ to fabricate photodetectors with strain-actuated bandwidth. The strain in these devices can be reversibly applied through the thermal expansion (shrinkage) of their substrate material, which induces tensile (compressive) biaxial stress. We find that upon tensile straining the photoresponse increases and that the excitonic features present in the spectra redshift in agreement with previous spectroscopic works. Interestingly, the spectra also develop a slowly decaying tail for long wavelengths, which further increases the bandwidth. We extract a strain gauge factor for the wavelength cut-off shift of up to ~58 nm/% (~135 meV/%). This remarkably large value demonstrates that 2D semiconductors hold a great promise for future straintronic devices where strain is employed as a variable tuning knob. Indeed, the possibility of strain tuning the optoelectronic performance of photodetector devices in a fast timescale opens up the possibility to fabricate artificial photonic devices that mimic the adaptability of the human eye. The photodetectors discussed in this work, which can be tuned from a fast, low responsivity, and narrowband state to a slow, very sensitive, and wideband one, can open the possibility of developing adaptable artificial photonic elements. Indeed, the method presented here can be easily translated to fabricate strain tunable photodetectors based on other 2D materials.

Materials and methods

Materials

The MoS₂ flakes have been obtained by mechanical exfoliation with Nitto tape (SPV 224) from a bulk natural crystal (Moly Hill mine, Quebec, QC, Canada) onto a viscoelastic polydimethyl-siloxane (PDMS) stamp from Gel-Pak[®] (Gel-Film WF ×4 6.0mil).

Device fabrication

The MoS₂ flakes located on the PDMS were characterized by optical transmission microscopy (Motic BA310Met-T metallurgical microscope, equipped with a 18 megapixel digital camera AMScope MU1803 and a fiber-coupled compact spectrometer Thorlabs CCS200/M) to determine the number of layers [32]. The single-layer MoS₂ flakes were then transferred between prepatterned Au/Ti electrodes (fabricated by electron-beam evaporation through a metal shadow mask from Ossila[®], part number E324) on polycarbonate (PC) substrate using an all dry deterministic transfer method [34].

Strain application

The biaxial strain on the photodetector devices is achieved through the thermal expansion of the PC substrate, controlled by a thermal stage (Linkam HFS600-P for the measurements shown in Fig. 2 and a Peltier element with an approximated consumption of 5 W to reach the highest temperature value for the rest of the measurements).

Photocurrent spectroscopy measurements

Photodetectors #1, #2, and #3 were measured in a pure N₂ atmosphere; the rest of the devices were measured under ambient conditions (average humidity 20%). The current vs. voltage characteristics of the devices were measured with a Keithlev 2450 source-meter unit, while the devices were illuminated focusing the light coming from a fiber-coupled light source into a 400-µm spot, covering the whole area of the device and providing a homogeneous power density. The cut-off wavelength is extracted from the wavelength value at which the photocurrent drops below the setup noise level (1pA). A Bentham TLS120Xe tunable light source was used for the measurements shown in Fig. 2, a halogen lamp equipped with a VariSpec[™] Liquid Crystal Tunable Filter was used for the measurement shown in Fig. S6 in the Supporting Information. The rest of the photocurrent measurements were carried out with high power fiber-coupled LED sources from Thorlabs. Fig. S9 in the Supplementary Material shows an example of current vs. voltage characteristics and of response time measurements acquired at different strain levels.

Supplementary material

Scanning micro-reflectance maps; finite element analysis to determine the strain transfer; strain calibration; disentangling temperature change from biaxial strain; scanning photocurrent measurements; examples of strain tuning of the current vs. voltage characteristics and response time measurements; characteristics of other devices (Figs. S10 and S11); and strain tuning of the power-dependent photocurrent generation (Fig. S12).

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Data availability

The raw/processed data required to reproduce these findings cannot be shared at this time due to technical or time limitations.

Data will be made available on request.

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Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.mattod.2019.04.019.

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