Photoconductivity of Solution-Processed MoS$_2$ Films

Graeme Cunningham, Umar Khan, Claudia Backes, Damien Hanlon, David McCloskey, John Donegan and Jonathan N Coleman*

School of Physics, CRANN & AMBER, Trinity College Dublin, Dublin 2, Ireland

*colemaj@tcd.ie

ABSTRACT: Solution-exfoliated MoS$_2$ nano-platelets were formed into thin films by deposition onto a water surface followed by transfer to indium tin oxide coated glass. After drying, a gold electrode was evaporated on top to give a sandwich structure with quasi-Ohmic contacts. Illumination of this device with broadband light of ~1 kW/m$^2$ intensity gave a fourfold increase in conductivity. The photocurrent increased sub-linearly with intensity and exponentially with time indicating the presence of traps. The photo-responsively was $\sim$10$^{-4}$ A/W at 15 V, competitive with other 2-dimensional photoconductors. This work demonstrates the potential for liquid-exfoliated, inorganic nanosheets to be fabricated into low-cost optoelectronic devices.

ToC entry

We have prepared solution-processed thin films of MoS$_2$ nano-platelets which show four-fold conductivity increase under 1-sun illumination.
Introduction

Over the last few years interest in 2-dimensional (2D) nano-materials has developed rapidly, partly due to the unprecedented success\(^1\) of graphene. Of the many types of 2D material,\(^2\) the family of transition metal dichalcogenides (TMDs) is particularly interesting.\(^3, 4\) TMDs consist of 2D monolayers which are constructed from a plane of transition metal atoms, usually Mo, W or Nb, sandwiched between two planes of chalcogen atoms i.e. S, Se or Te. Under normal circumstances, TMD monolayers are stacked together to form 3-dimensional crystals. However, it has recently been shown that such crystals can be exfoliated to give mono- or few-layer flakes on surfaces.\(^5\) Alternatively, mono- or few-layer-thick films can be grown directly by CVD.\(^6\) Monolayers of TMDs such as MoS\(_2\) have generated much interest due to their exceptional electronic and optical properties.\(^3\) For example MoS\(_2\) and WS\(_2\) monolayers are semiconductors and have been used as the basis of a wide range of electronic and optoelectronic devices including transistors, logic circuits, sensors and light emitting diodes.\(^3, 7\) In particular, mono-and few-layer MoS\(_2\) and WS\(_2\) display photoconductivity and so may find applications as sensitive photodetectors.\(^8-14\) This is an application to which these materials appear to be particularly suited. For example, individual MoS\(_2\) monolayers have very high photosensitivity, up to 880 A/W at low illumination power.\(^10, 12\) This, coupled with their extremely low noise equivalent power, makes them ideal for very low level photo-detection.\(^10, 12\) In addition, their spectral range can be tuned by control of the MoS\(_2\) flake thickness,\(^15\) leading to a broad photo-response which is expected to extend beyond 900 nm for thick MoS\(_2\) flakes.\(^9\) In addition, MoS\(_2\) photodetectors are extremely robust, working at high temperatures\(^12\) and in harsh environments with high radiation fluences.\(^16\)

All of the work described above was performed on CVD-grown or mechanically-exfoliated TMDs. Another way to produce 2D materials is to exfoliate layered materials in
liquids. Liquid exfoliation has been demonstrated for graphite, TMDs and a range of other layered crystals and is a cheap and simple way to produce large quantities of 2D materials. One big advantage of this process is that liquid exfoliation gives suspensions of 2D nanosheets which can easily be processed into composites, films or other structures. Such techniques have been used to prepare samples for a range of applications, mainly in electrochemistry. However, relatively few electronic or optoelectronic device applications have been demonstrated for liquid exfoliated nanosheets. This is unfortunate as devices based on such materials could be fabricated easily and cheaply using a range of techniques such as spraying, inkjet printing or gravure coating. Here we show that liquid exfoliation techniques can be used to give partially exfoliated, but solution processable MoS$_2$ suspensions. These suspensions can be formed into thin films which, when electrically connected, show promising photoconductive properties.

**Experimental procedure**

The MoS$_2$ starting powder was purchased from Sigma Aldrich (<2μm particle size, product number 234842) and used as supplied. Nano-platelets of MoS$_2$ were exfoliated and suspended in liquid using well-established solvent exfoliation techniques. MoS$_2$ powder (1 g) was added to HPLC-grade isopropanol (IPA, 100 ml) in a glass beaker. The sample was sonicated with a stand-mounted ultrasonic tip (Heilsher model UP200S, 200W, 24 kHz) for 24 hours at 60% amplitude in pulsed mode with 2s on and 1s off. This results in formation of a dispersion of MoS$_2$ nano-platelets with a wide range of sizes and thicknesses.

When exfoliating layered materials in liquids, a centrifugation step is normally employed post sonication to remove the larger nano-platelets and any partially exfoliated crystallites. However, we found that films prepared from centrifuged MoS$_2$ dispersions always displayed pinholes extending from top to bottom as evidenced by electrical shorting.
when electrodes were applied in a sandwich structure. Here, we avoid centrifugation as we found films of un-centrifuged MoS\textsubscript{2} to be generally pinhole free. Shorting can occur in films consisting of discrete nano-scale entities, even when the films are relatively uniform and free of well-defined holes, due to diffusion of metal through the porous internal structure of the film. That shorting occurs for centrifuged films but not for non-centrifuged samples implies that the film microstructure depends strongly on the details of sample processing. Further work is needed to fully understand such effects. Because our dispersions were not centrifuged, they are expected to contain larger MoS\textsubscript{2} nano-platelets and small crystallites as well as thin nanosheets. However, most importantly, they are still solution processable. To investigate the nature of the dispersed MoS\textsubscript{2}, we performed TEM characterisation (figure 1A) which showed the presence of micron sized nano-platelets. However, these were largely electron opaque suggesting the nano-platelets to be relatively thick. To measure the nano-platelet thickness, we deposited a few drops of dispersion onto Si/SiO\textsubscript{2} (200 nm oxide thickness), just dried them in ambient air before imaging with AFM (figure 1B). Measurement of the thickness of many nano-platelets allowed statistical analysis (figure 1C) showing a broad distribution peaked around 50 nm. This is consistent with the presence of both well- and poorly-exfoliated nano-platelets as expected. In addition, the lateral nano-platelet dimensions were measured by TEM (figure 1D) and showed typical nano-platelets to be approximately 1 \(\mu\)m in length.

These dispersions were used to form thin films for electrical testing. In order to avoid the formation of pinholes, a number of film formation methods were attempted before settling on a technique based in principle on the Langmuir-Blodgett method. A 250 ml conical flask was filled to the brim with Millipore Water and, using a pipette, a few drops of the MoS\textsubscript{2} dispersion were slowly dropped onto the rim of the flask and allowed to flow down into the water. This technique employs MoS\textsubscript{2}’s inherent hydrophobicity: interaction with the water
drives the MoS$_2$ to the surface while the IPA enters the water subphase. The conical flask is ideal for keeping the concentration of IPA low in a larger body of water whilst allowing film formation over a smaller area. At first the MoS$_2$ aggregates form at a number of independent nucleation sites. However, as more dispersion is added these sites coalesce resulting in the entire liquid surface becoming covered with a film of MoS$_2$ nano-platelets.

To transfer this film onto a substrate, an indium tin oxide (ITO)-covered glass slide (1.2 cm×1.2 cm) was inserted below the surface using a set of tweezers and slowly pulled out at an angle of ~75° to the horizontal. This resulted in the ITO being coated with a thin film of MoS$_2$ nano-platelets. This film was allowed to dry naturally in ambient air until appearing dry whereby it was then placed onto a hotplate at 250°C for 3 minutes before being removed and allowed to cool. The MoS$_2$ film thickness was increased by re-dipping a number of times. However, the water had to be changed every 2-3 coatings to avoid excessive IPA content compromising the film formation.

Results and Discussion

The resultant films were dark grey and visually quite uniform (figure 1F). SEM examination (Zeiss Ultra Plus SEM) showed them to be continuous over length scales greater than a few microns (figure 1G). However, when viewed at higher magnification it becomes clear that they are composed of a disordered array of nano-platelets of MoS$_2$ (figure 1H). The film for which we present electrical data below had a thickness of ~6 μm (achieved after 9 dips). However, the thickness varied by ~20% over a line scan of 800 μm (Dektak 6M Stylus Profiler) showing that this film formation method could certainly be improved. Raman spectra were recorded for such films (figure 1I). These were very similar to the equivalent spectra for the starting powder confirming the films to consist of relatively thick flakes (the Raman spectra of exfoliated MoS$_2$ flakes approaches that of bulk MoS$_2$ for flakes thicker than
~6 layers.\textsuperscript{24} We note the peak positions are slightly shifted from those expected for crystalline MoS\textsubscript{2} as is sometimes observed for powdered samples.\textsuperscript{25} We also measured the absorbance spectrum of a thin layer prepared on a glass substrate by a single dipping cycle (figure 1J). This spectrum displays the features expected from MoS\textsubscript{2} although the spectrum is probably superimposed on a scattering background.\textsuperscript{18}

Devices were fabricated by evaporating a gold electrode (~50 nm thick, 1.98 cm\textsuperscript{2} electrode overlap area) on top of the MoS\textsubscript{2} film to give an ITO/MoS\textsubscript{2}/Au sandwich structure arrangement (figure 2A inset). After evaporation, films were annealed in air at 200°C for 1 hour. Typically, such a mild anneal resulted in almost an order of magnitude current increase and a considerable reduction in current noise.

We expect this combination of ITO and gold electrodes to give roughly Ohmic contacts on both sides of the device. Bulk MoS\textsubscript{2} is generally n-type with a band gap of 1.2 eV.\textsuperscript{3} For bulk MoS\textsubscript{2} (appropriate here because of the partial exfoliation) the conduction band edge, Fermi energy and valence band edge are at energies of 4.4, 4.8 and 5.6 eV below the vacuum level.\textsuperscript{26} This means that the workfunction of the ITO (~4.5-4.8 eV)\textsuperscript{27} is expected to lie between the conduction band edge and the Fermi energy of MoS\textsubscript{2}. This is expected to result in an Ohmic contact.\textsuperscript{28} Although the work function of the gold does not lie between MoS\textsubscript{2}’s Fermi level and conduction band edge, a number of papers have shown good electron injection at Au/MoS\textsubscript{2} interfaces, which is consistent with the presence of Ohmic contacts.\textsuperscript{29-32} In fact, detailed measurements have shown that the potential barrier at the MoS\textsubscript{2}/Au interface is extremely small.\textsuperscript{33} This may be partially explained by chemical bonding between gold and sulphur atoms, located at flake edges or defects, leading to a very low injection barrier and so a quasi-Ohmic contact.
Shown in figure 2A is an I-V curve for an ITO/MoS2/Au sandwich structure device taken in the dark. This curve is symmetric around the origin, as expected if both contacts are Ohmic.\textsuperscript{34} The dark conductivity is \( \sim 2 \times 10^{-10} \, \text{S/m} \), three orders of magnitude lower than measured for similar films but with in-plane current flow.\textsuperscript{23} This is consistent with the anisotropy between in-plane and out of plane conductivities in graphite and suggests the current to be limited by inter-flake transport.\textsuperscript{35}

In this paper we are interested in the photoconductivity of films of MoS\textsubscript{2} nano-platelets. As such, we have studied the effect of illuminating the sample through the ITO with a full spectrum solar simulator (Newport 96000), fitted with a Xe arc discharge lamp (150W) and air mass (AM) 1.5 filter. The lamp was calibrated against a standard Si based reference cell. Light intensity at the sample was controlled using a set of neutral density filters (NDFs) purchased from Newport. Shown in figure 2A are I-V curves measured for an MoS\textsubscript{2} film illuminated at different light intensities compared to the dark curve. All curves are symmetric about zero and linear at low voltage although some curvature is observed at higher voltages. Assuming the contacts are indeed Ohmic, such curvature is probably due to space charge effects.\textsuperscript{28} However, such behaviour will not be analysed in this work.

We measured the conductivity (measured in the -2 to 2 V range) as a function of light intensity, \( F \), as shown in figure 2B. The conductivity increases significantly from its dark value of \( 2 \times 10^{-10} \, \text{S/m} \), reaching \( 8 \times 10^{-10} \, \text{S/m} \) for \( F=1100 \, \text{W/m}^2 \). We note that this increase is not linear as is usually observed for semiconductors.\textsuperscript{28} Rather, the conductivity increase (i.e. the photoconductivity, \( \sigma_{\text{ph}} \)) scales roughly as \( \sigma_{\text{ph}} \propto F^\gamma \) with \( \gamma=0.5 \). In the simplest case, such behaviour is usually associated with bimolecular recombination,\textsuperscript{28, 36} which is usually observed in insulators where the density of optically generated carriers greatly exceeds that of free carriers in the dark.\textsuperscript{37} The majority of papers studying photoconductivity of individual
monolayer (or few-layer) MoS$_2$ or WS$_2$ have observed linear (or near-linear) dependence of photocurrent on light intensity.$^{8,9,13,14}$ However, a number of papers on such samples report sub-linear behaviour with $\gamma$ between 0.5 and 0.71.$^{10-12}$ However, the materials studied in these papers are semiconductors with reasonable high dark carrier densities making bimolecular recombination unlikely. In such cases, sub-linear intensity dependence is likely to be associated with the presence of traps.$^{28}$

Simple analyses of bimolecular recombination which predict $F^{1/2}$ behaviour generally assume trap free systems.$^{28}$ However, in many real systems traps are present and manifest themselves through non-linear dependence of photocurrent on intensity and a relatively slow increase and decay of photocurrent with time.$^{28}$ To test for the presence of traps, we measured the current as a function of time under constant bias of 15 V. During the measurement, the light was switched on and later switched off. Examples of the resultant curves are shown in figure 2C. In all cases, on illumination a fast photocurrent increase was observed followed by a slow photocurrent increase. Similarly, on cessation of illumination there is a fast decay component followed by a slow decay. Such a slow rise and decay of photocurrent are strong indicators of the presence of traps.$^{28}$

Fast rise and decay of photocurrent has been observed by a number of authors in experiments on individual monolayer (or few-layer) MoS$_2$ or WS$_2$, and usually in systems with linear intensity dependence that are assumed to be trap free. In addition, slow photocurrent rise and decay have also been observed.$^{10,14}$ This has been attributed to traps associated with the interaction of the surface of the flake with either the substrate or the ambient gas. We note that a combination of fast and slow photocurrent rise and decay has not been reported for MoS$_2$ or WS$_2$. However, such behaviour is known for semiconductors such as n-type silicon.$^{28,38}$ In such behaviour, the fast component is attributed to the initial
equilibration of carrier generation and recombination rates and occurs on the timescale of the carrier lifetime. This can be as fast as \(~1\) ns in MoS\(_2\) and so is hard to measure experimentally.\(^9\) The slow photocurrent rise is associated with the effects of carrier trapping.\(^{28}\) Once the light is switched off, the fast decay is due to the recombination of free carriers while the slow decay represents the thermal detrapping of carriers.

As described above, the presence of traps leads to slow rise (decay) of photocurrent as traps fill (empty), slowing the approach to steady state behaviour. A mono-exponential rise (or decay) implies the presence of a single type of discrete trapping state within the gap where re-trapping of freed carriers is negligible.\(^{28, 39-41}\) The dark current on decay can be slightly higher than before illumination. This persistent photoconductivity (PPC) is a common effect and is associated with very deep traps.\(^{40, 42, 43}\)

For all light intensities these slow photocurrent increases can be fit to an exponential function of the form

\[
\Delta I_R = \Delta I_{f,R} + \Delta I_{s,R}(1-e^{-t/\tau_R})
\]

where \(\Delta I_{f,R}\) is the fast component of the rising photocurrent, \(\Delta I_{s,R}\) the slow component of the rising photocurrent and \(\tau_R\) the time constant. Similarly, the slow photocurrent decay can be fit to

\[
\Delta I_D = \Delta I_{s,D}e^{-t/\tau_D} + \Delta I_{p,D}
\]

where \(\Delta I_{s,D}\) is the slow component of the decaying photocurrent, \(\tau_D\) is the time constant and \(\Delta I_{p,D}\) is the persistent photocurrent.\(^{28}\) Note, the fast component (i.e. \(\Delta I_{f,D}\)) of the decaying photocurrent can be read off the graph.
The fast components of photocurrent rise and decay have been plotted against \( F \) in figure 2D. In both cases they sit on the same line consistent with \( \Delta I_f \propto F^\gamma \) where \( \gamma=0.54 \). While this is very close to the 0.5 expected for bimolecular recombination care must be taken as the relationship between this fast component and the trap distribution is not clear. Indeed power-law behaviour whose exponent is controlled solely by the details of the trap distribution is well known.\(^{28}\)

The slow component of the rise and decay of photocurrent was also plotted versus \( F \) in figure 2E. Both \( \Delta I_{s,R} \) and \( \Delta I_{s,D} \) behave similarly with no obvious correlation to light intensity. This behaviour is not surprising as the slow component is thought to be controlled by the trap density.\(^{28}\) The PPC is not plotted but was close to 0.5 nA for all samples.

We note that after long illumination times, the photocurrent will saturate at 
\[
\Delta I_{R,sat} = \Delta I_{f,R} + \Delta I_{s,R}.
\]
This saturation photocurrent increases weakly with intensity. As a result, the saturation photoresponsivity (usually defined as \( P_R = \Delta I_{R,sat} / F \) where \( F \) is the incident intensity) falls with increasing intensity. However, at low intensity \( P_R \approx 10^{-4} \) A/W (measured at 15 V), similar to both the lower end of the range reported for CVD grown TMDs\(^{11}\) and to early graphene photodetectors.\(^{44}\) However, we note that this does not take the wavelength dependence of the photoconductivity into account. To test this we measured the action spectrum of the photoconductive response using a set of notch filters and the solar simulator. Because the photocurrent scales roughly as \( F^{1/2} \), we calculate the measured photocurrent divided by the square root of the integrated incident intensity. This is plotted versus wavelength in figure 1J (inset). We find the spectral response to increase with decreasing wavelength as expected from the absorbance spectrum. We see an apparent peak around 650 nm consistent with the A exciton position in MoS\(_2\).\(^{45}\) In addition, the spectral response increases sharply below 500 nm as observed previously for MoS\(_2\) monolayers.\(^{10}\)
This suggests that the photoresponsivity calculated using the absorbed power would be considerably higher than the value quoted above.

In figure 2F time constants for the rise and decay of photocurrent are plotted against F and tend to fall off as roughly $F^{-1/2}$. It has been shown that when the $\tau_D$ scales with intensity as $\tau_D \propto F^{-\gamma}$, this means the observed time constant equals the carrier lifetime (including time spent in traps). That this lifetime is so long indicates that trapping is a problem in these films. The main difference between the films used here and the MoS$_2$ or WS$_2$ samples previously studied is that these films consist of a disordered array of nano-platelets. It is entirely possible that states associated with flake edges are acting as traps. Future work will study methods of passivation of the flake edges with the aim of reducing the number of traps and so minimising their effect.

Conclusion

In summary, we have demonstrated a method to prepare solution-processed thin films of MoS$_2$ which can be fabricated into photodetectors. These films show a four-fold increase in conductivity on exposure to ~1 kW/m$^2$ broadband excitation. However, due to the presence of traps the photo-response scales sub-linearly with intensity and increases slowly with time. However, the photo-responsivity is reasonable, matching that of early monolayer graphene reports. The ability to prepare photo-detectors from inorganic nano-materials such as MoS$_2$ by solution processing techniques represents an important step forward. Such techniques allow the preparation of cheap, large area devices quickly and at low cost. We consider this the first step toward processing solution-exfoliated 2D materials into a range of optoelectronic devices.

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Figure 1: A) TEM image of solution processed MoS$_2$ nano-platelets. B) AFM image of an MoS$_2$ nano-platelet deposited on a substrate. C) Thickness distribution for MoS$_2$ nano-platelets deposited on a Si/SiO$_2$ substrate as measured by AFM. D) Length distribution as measured by TEM. E) Conical flask containing water with a film of MoS$_2$ nano-platelets floating on top. F) Photograph of a film prepared using 9 dipping cycles. G) Widefield and H) close-up SEM images of a film of MoS$_2$ nano-platelets. I) Raman spectra of both starting
powder and film prepared from an uncentrifuged MoS$_2$ dispersion. J) Optical absorbance spectrum of an MoS$_2$ film prepared by a single dipping cycle. Inset: Action spectrum for a film prepared from 9 dipping cycles. The photoconductive behaviour is represented by the photocurrent divided by the square root of the integrated incident intensity (because $\sigma_{ph} \propto F^{\gamma}$ with $\gamma=0.5$).

Figure 2: A) IV curves for an MoS$_2$ film in the dark and under various light intensities, F. Inset: Electrode structure. B) Measured conductivity as a function of light intensity. The line illustrates square root behaviour. C) Photocurrent, PC, (dark current removed) plotted versus time for three different light intensities (V=15 V). For the red curve, the red arrows indicate
the times the light was switched on and off while the black double-headed arrows illustrate the fast portions of the rise and decay. D) Fast components of both rise and decay of photocurrent plotted versus light intensity. The dashed line is a power law fit. E) Slow components of both rise and decay of photocurrent plotted versus light intensity. F) Time constants for both rising and decaying photocurrents plotted versus light intensity. The dashed line represents $F^{-1/2}$ behaviour.
References

33. Personal communication from A. Kis, 2013.