Tiled Monolayer Films of 2D Molybdenum Disulfide Nanoflakes Assembled at Liquid/Liquid Interfaces

Joseph Neilson, Michael P. Avery, and Brian Derby*

ABSTRACT: Thin films of MoS2 bilayer nanoflakes, which are predominantly a single flake thick and with flakes in edge-to-edge contact, have been produced via self-assembled tiling at the planar interface between two immiscible liquids. Films of several square centimeters extent can be produced with a total covered area approaching 90% and over 70% of the film covered by single flakes without overlap. Films produced through liquid/liquid assembly are shown to produce a lower uncovered area fraction and more uniform thickness when compared with films of similar areal coverage produced by the “top-down” techniques of spin coating and spray coating. Statistical analysis of flake coverage data, measured by atomic force microscopy (AFM), shows that liquid/liquid assembly produces a distinctly different variation in film thickness than conventional top-down deposition. This supports the hypothesis that the two-dimensional (2D) confinement of liquid/liquid assembly produces more uniform films. Demonstrator field-effect transistors (FETs) manufactured from the films exhibit mobility and on/off current ratios of 0.73 cm² V⁻¹ s⁻¹ and 10⁵, respectively, comparable to FETs of similar layout and chemical vapor deposition (CVD)-grown or mechanically cleaved single-crystal MoS2 channel material. This work demonstrates the use of liquid/liquid interfaces as a useful tool for the self-assembly of high-performance thin-film devices made from dispersions of 2D materials.

KEYWORDS: thin films, 2D materials, Poisson distribution, coverage, self-assembly, device fabrication

INTRODUCTION

Semiconducting transition metal dichalcogenides (TMDC), such as molybdenum disulfide (MoS2), have a layered crystal structure analogous to graphene, with the layers bonded by relatively weak van der Waals bonds, and can be exfoliated to form thin flakes of two-dimensional (2D) materials.1 These materials have unique physical properties that are tunable, depending on the number of atomic layers in the flakes, and have been widely explored for use in the fabrication of electronic and optoelectronic devices such as field-effect transistors (FETs) and optical sensors.2−4

Large-area crystalline MoS2 films can be grown using “bottom-up” methods such as chemical vapor deposition (CVD), leading to high-quality electronic and optoelectronic devices.5,6 However, the high-temperature requirement of the CVD process does not allow the direct deposition of large-area devices onto flexible polymeric substrates. At present, the leading technology for low-cost, large-area flexible electronics manufacture is organic semiconductor technology, which is compatible with a range of relatively low-cost printing methods and roll-to-roll manufacturing. However, current generation organic FETs suffer from relatively low carrier mobility compared to inorganic semiconductor devices, which limits their applications. In contrast, devices produced from liquid dispersions (inks) of TMDCs have potentially higher carrier mobility and could therefore represent an attractive, printable alternative to organic semiconductors with superior performance, while allowing new flexible device types to be produced. To make printed large-area thin films from 2D TMDCs for high-throughput manufacture of electronic devices, solution-based processing provides a low-cost and low-temperature option. The exfoliation and handling of 2D materials as colloids is already widely explored7 and has enabled the formulation of printable inks for use in flexible electronic devices that have the potential to be mass produced, e.g., by roll-to-roll manufacturing.8

The fabrication of large-area films from a colloidal dispersion of nanosheets is a key challenge for the manufacture of electronic devices from solution-processed 2D materials. A number of solution-based methods have been developed to produce thin films of 2D materials; these can be grouped into: (1) “top-down” methods, e.g., spray coating and spin coating,10 where flakes are deposited from the surrounding
medium and fall randomly onto the final substrate, remaining attached at their initial contact position; and (2) “interface assembly” methods at liquid/air (Langmuir–Blodgett) or liquid/liquid interfaces, where flakes are confined to the interface between two phases and assemble through translation within the interface, before transfer of the complete resulting film onto a substrate. Although the Langmuir–Blodgett method for assembly at the liquid/air interface is well established, material can be lost to mixing with the subphase and aggregation can occur during assembly. In addition, the large subphase area is sometimes unsuitable for research scale film assembly. These issues can be ameliorated by assembly at liquid/liquid interfaces, which have been shown to improve confinement of material to the interface without the need for specialized apparatus and can be applied to a range of 2D, 1D, and 0D materials.

Carrier mobility is higher in-plane than out-of-plane for 2D materials, due to the tunnel barrier through the van der Waals gap; thus edge-to-edge contacts between nanoflakes and dense packing within the plane are beneficial to ensure high carrier mobility throughout a 2D material. Improved edge-to-edge contact and reduced overlap can be afforded by interfacial assembly methods, which confine material at the molecularly flat interface, therefore offering a more promising route to high-quality thin films than top-down assembly methods.

The assembly of 2D materials at planar liquid/liquid interfaces can be achieved in a number of ways. Most literature examples report methods that begin by dispersing the 2D material in one liquid, before adding a second immiscible liquid layer to create the interface. The 2D material in the dispersed phase can be induced to transport to the interface using mechanical means, such as sonication or manual shaking, or by the addition of a chemical inducing agent. There have been few reports of assembly by direct injection of the material to the liquid/liquid interface, as described in this work.

Here, we demonstrate the assembly of predominantly bilayer flakes of MoS$_2$ at the interface between two immiscible solvents (hexane and water). Water and hexane were selected to form the interface for film assembly due to their high interfacial tension of around 50.5 mN m$^{-1}$, which can be effectively lowered by the insertion of a monolayer of 2D material, promoting liquid/liquid interfacial assembly. Isopropanol alcohol (IPA) was selected as the solvent for dispersing MoS$_2$, due to its well-matched solubility parameters, low toxicity, and miscibility with both water and hexane. The morphology of the resulting films is compared with those produced by spray coating and spin coating. The films fabricated at liquid/liquid interfaces are shown to have greater film uniformity and single-layer coverage than found with the top-down deposited films.

**RESULTS AND DISCUSSION**

**MoS$_2$ Nanoflake Dispersion.** A dispersion of MoS$_2$ nanoflakes in isopropanol alcohol (IPA) was obtained as described in the Materials and Methods section. The dispersion contained flakes of average area 0.95 $\mu$m$^2$ and thickness 2 nm as determined by scanning electron microscopy (SEM) and atomic force microscopy (AFM), respectively (Figure 1a,b). (histogram of flake lateral area data) is in the Supporting Information, Figure S1). The semiconducting 2H phase of MoS$_2$ is retained as evidenced by the $a$, $b$, and $c$ excitonic peaks at around 673, 610, and 445 nm, respectively (Figure 1c). The $a$ and $b$ peaks correspond to the $\sigma$–d interband electronic transitions associated with the d$_2$ metal in the hexagonal coordination geometry. The $c$ excitonic peak is also typical of 2H phase MoS$_2$ absorption spectra and has been linked to a Van Hove singularity. The Tauc plot in Figure 1d suggests a band gap energy of around 1.75 eV, indicative of one- to two-layer MoS$_2$.

**Stability of Nanoflakes at the Liquid/Liquid Interface.** The adsorption of a solid particle at a fluid–fluid interface is driven by the reduction of the interfacial energy of the system. The energy of detachment of the particle into either liquid 1 or liquid 2, $E_{d1}$ and $E_{d2}$, is given by

$$E_{d1} = \gamma_{12} (A_c + A_{p2} \cos \theta)$$

(1a)

$$E_{d2} = \gamma_{12} (A_c - A_{p1} \cos \theta)$$

(1b)

Here, $\gamma_{12}$ is the interfacial tension between the two liquids, $A_c$ is the change in the liquid 1/liquid 2 interfacial area resulting from the detachment of the particle, $\theta$ is the three-phase contact angle, and $A_{p1}$, $A_{p2}$ are the initial areas of the particle/liquid interface prior to the detachment of the particle for liquids 1 and 2, respectively. The lowest energy of detachment $E_d$ is given by

$$E_d = \begin{cases} E_{d2} & \text{for } 0^\circ \leq \theta \leq 90^\circ \\ E_{d1} & \text{for } 90^\circ \leq \theta \leq 180^\circ \end{cases}$$

(2)

For all values of $\theta$, the detachment energy is positive, with a maximum value at $\theta = 90^\circ$. It is therefore thermodynamically favorable for particles to gather at the interface, and they do so spontaneously. The energy required to detach particles from the interface can be several orders of magnitude greater than $k_BT$, under the correct conditions, making these very stable systems. The detachment energy increases with increasing particle size since the $A_c$, $A_{p1}$, and $A_{p2}$ terms all increase, provided that particles remain small enough (typically...
a few microns in diameter, \(d\), for spherical particles) that the effects of gravity are negligible.\(^{38,39}\) This is confirmed by the Bond number \(<1\) (Supporting Information, eq S1) for particles of diameter 1 \(\mu\)m.

When discussing the behavior of particles at liquid/liquid interfaces, the typical model system, applicable to most colloidal particle behavior, is that of a spherical particle (radius, \(r\)) at a flat horizontal interface. For spherical particles, eq 2 becomes

\[
E_d = \pi r^2 \gamma (1 \pm \cos \theta)^2
\]

(3)

Here, the sign in the parentheses is positive when \(\theta > 90^\circ\) and negative when \(\theta < 90^\circ\).\(^{36,38}\) Note that we have replaced the subscripts 1 and 2 with \(h\) and \(w\) to represent hexane and water, respectively, which are the solvents used in this study.

Spherical particles are, however, not an appropriate description for 2D flake materials. Equation 2 can also be used to describe systems of nonspherical particles, such as rods or disks. Binks provides an analysis for a rounded disklike particle based on the major (\(a\)) and minor (\(b\)) semiaxes of the particle such that \(a/b\) is the particle aspect ratio.\(^{36}\)

\[
E_{dw} = \pi b^2 \gamma (1 - \cos \theta)^2 \left[ 1 + \frac{(\frac{a}{b} - 1)^2}{1 - \cos \theta} \right] + \frac{2(\frac{a}{b} - 1) \sin \theta - \theta \cos \theta}{(1 - \cos \theta)^2}
\]

(4a)

\[
E_{dh} = E_{dw} + 2\pi b \gamma \cos \theta \left[ \frac{a}{b} - 1 \right]^2 + \pi \left( \frac{a}{b} - 1 \right) + 2
\]

(4b)

Although flakes of 2D materials at liquid/liquid interfaces have been described by this rounded disk model previously,\(^{10}\) we suggest a simpler approximation to the flake geometry. We treat the flakes as infinitesimally thin disks where the disk edge has a negligible contribution to the flake area when compared to the faces of the disk. The most stable configuration of the flake at the interface, where \(E_d\) is maximized and the interfacial energy is minimized, is with the flake faces parallel to the interface, maximizing \(A_t\). For this system, \(A_t = A_{hw} = A_{ph} = A_0\) where \(A_t\) is the area of one face of the disk. The minimum energy for detachment for 2D flakes given by eq 2 therefore simplifies to

\[
E_d = \gamma_{hw} A_t (1 \pm \cos \theta)
\]

(5)

where the sign in the parentheses is positive when \(\theta > 90^\circ\) and negative when \(\theta < 90^\circ\).

Applying eq 5 to our experimental system, \(\gamma_{hw} = 50.5\) m J m\(^{-2}\), \(A_t = 0.95\) mm\(^2\) (Supporting Information, Figure S1), and \(\theta = 99.7^\circ\) (determined by the sessile drop method (Supporting Information, Figure S4)), we find that \(E_d\) is \(9.70 \times 10^6\) kJ T. This value is almost identical to the value calculated using eqs 4a and 4b for the rounded disk model (\(a = 0.55\) \(\mu\)m, \(b = 1\) nm), validating our simplified approach for 2D materials at liquid/liquid interfaces and confirming that the contribution of flake edges to the reduction in interfacial energy at the interface is indeed negligible. By comparison, \(E_d\) for a spherical particle of equal total surface area is \(6.19 \times 10^5\) kJ T, as calculated using eq 3. Thus, the stability of confinement at the liquid/liquid interface is greater for disklke particles and flakes of 2D materials than for spherical particles, further increasing the driving force for assembly.

**Liquid/Liquid Film Assembly.** Figure 2b provides a schematic illustration of the three-step process for film assembly and transfer onto a substrate. The dispersed 2D material is injected into the upper hexane phase, and the flakes assemble at the liquid/liquid interface driven by the reduction in interfacial energy, as discussed above. The high energy required to detach flakes from the interface, as described by eq S1, ensures minimal overlap between adjacent flakes during the assembly process. The IPA used to disperse the flakes causes a decrease in the water/hexane interfacial tension at the point of injection,\(^{40}\) hence, the resulting local interfacial tension gradient sweeps the 2D material to the opposite side of the interface. This effect is evident at the macroscopic scale, as shown in Figure 2a (see also Video S1 in the Supporting Information). The interfacial tension gradient compresses the individual nanoflakes into a tightly packed, tiled film, with enhanced edge-to-edge contact and connectivity of the flakes within the thin film (high-magnification SEM image available in the Supporting Information, Figure S2). The assembled film of densely tiled (jammed) 2D flakes is stable at the liquid/liquid interface for at least 2 days, with no mixing of the 2D material with either bulk phase observed. This can be attributed to the poor solubility of MoS\(_2\) materials in both hexane and water, and its stability at the interface of hexane and water. Finally, the film is transferred onto a substrate by manually lifting it through the film parallel to the surface, analogous to the Langmuir–Schaeffer coating method (Figure 2b).\(^{41,42}\)

Hence, this liquid/luid film assembly method is applicable to many 2D materials, and a variety of substrates can be coated in this way. Additional benefits of the injection method discussed here over previously reported liquid/liquid methods are improved control over the amount of material added to the interface and a reduced number of processing steps without the need for further sonication or an inducing agent to facilitate film formation.
Characterization of Films on Substrates. Figure 3 shows representative SEM and AFM images of MoS$_2$ films deposited using: (d) liquid/liquid interface assembly, (e) spin coating, and (f) spray coating. Line scans from the AFM images (d–f) corresponding to the dashed yellow lines. The colored regions indicate the z height in terms of coverage (c) in flake monolayers. The black dotted line corresponds to the height of the substrate. All scale bars = 5 μm.

Figure 3. SEM images of MoS$_2$ thin films using: (a) liquid/liquid interface assembly; (b) spin coating; and (c) spray coating. AFM images of MoS$_2$ thin films deposited using: (d) liquid/liquid interface assembly, (e) spin coating, and (f) spray coating. (g–i) Line scans from the AFM images (d–f) corresponding to the dashed yellow lines. The colored regions indicate the z height in terms of coverage (c) in flake monolayers. The black dotted line corresponds to the height of the substrate. All scale bars = 5 μm.

Characterization of Films on Substrates. Figure 3 shows representative SEM and AFM images of MoS$_2$ films on a Si wafer (90 nm SiO$_2$) formed via liquid/liquid interfacial assembly, spin coating, and spray coating with similar mean coverage. Films formed by spray and spin coating contain stacked/overlapped flakes and a large fraction of uncovered area. The liquid/liquid interface assembly method, however, confines flakes to a single assembly plane. This leads to minimal flake/flake overlap and clear edge-to-edge contact between the majority of the flakes (Figure 3a,d) (high-magnification SEM image available in the Supporting Information, Figure S2); the resulting film is therefore continuous and well packed, with an average coverage of 89% (AFM, Figure 3a) to 91% (SEM, Figure 3b), similar to the predicted packing fraction for maximally jammed binary hard disks (0.78−0.91).43−48 The z-height data in the AFM images can also be extracted and processed to provide a quantitative measure of the distribution of flakes on the substrate via the assignment of coverage values.

Coverage is a dimensionless discrete variable, c, giving the number of objects that intersect a line perpendicular to a surface at a given point. The mean coverage, $\overline{c}$, is given by the

$$\overline{c} = \frac{\sum_{i=1}^{n} c_i}{n}$$

where $c_i$ is the coverage of the $i^{th}$ object and $n$ is the total number of objects.
number of objects per unit area multiplied by the area of one object. A visual representation of these concepts is given in Figure 4a. For top-down deposition, such as spin coating and spray coating, the arrival of material at the surface can be treated as a series of independent random events, and thus, given the finite size of the flakes, overlap of the arriving material is possible. In this case, the probability, \( P(c' \mid c) \), that a point on the surface has a coverage, \( c' \), when the whole surface has a mean coverage \( \bar{c} \), is given by the Poisson distribution:

\[
P(c' \mid c) = \frac{e^{-\bar{c}} \bar{c}^{c'}}{c!} \quad \text{for} \quad c = 0, 1, 2, 3, \ldots
\]  

(6)

When film assembly occurs at an interface, if the concentration of the flake dispersion is sufficiently dilute, there will be minimal overlap and face-to-face contact of the flakes with each confined to the 2D interface on injection. The flakes at the interface are then compressed into a tightly packed film via an interfacial tension gradient, resulting from the material injection, as previously discussed. We also note that capillary/electrostatic forces may also compress the film over a short range, but such effects are expected to be slight for particles of thickness <100 nm. If compression forces are large enough, this may lead to localized buckling or overlap such as that shown in Figure 4b, despite the 2D constraint of the interface. Further compression and roughening of the film may also occur as part of the assembled film transfer process. The expected final mean coverage for 2D films assembled at interfaces in this case is, thus, \( \bar{c} \approx 1 \). If the flakes were truly confined to the interface, forbidding overlap from occurring, then we would expect \( \bar{c} < 1 \) because perfect tiling of irregular flakes is not possible.

Thus, from our consideration of the mechanisms leading to the formation of a film through the addition of individual 2D material flakes, we would expect that the top-down deposition methods will show a range of film thicknesses over a given area, described by the Poisson distribution. In contrast, films produced by interfacial assembly should show a more singular distribution of thicknesses dominated by a thickness defined by the expected dimensions of a single flake.

The histogram of AFM data given in Figure 4c, for the films produced by liquid/liquid assembly, spin coating, and spray coating, provides an experimental measure of the film coverage probability distribution. This shows the fractional areas of the resulting films with coverages 0, 1, 2, etc. (analysis of AFM data for four 20 \( \times \) 20 \( \mu \)m\(^2\) areas at different locations on each film). The bins for the histogram data were chosen based on average flake height measurements (Figure 1b), which found that MoS\(_2\) flakes had an average height of 2 nm, with a minimum flake height of 1 nm. The height of a point on a film surface can be roughly equated to the coverage at that point by dividing by the height of a single flake. Therefore, any pixels in AFM film scans with a measured height <0.94 nm were treated as uncovered (\( c = 0 \)), and increasing coverages were binned with bin widths of 2 nm from that value to correspond to an additional flake layer, i.e., 1–3 nm is \( c = 1 \), 3–5 nm is \( c = 2 \), and so on. It should be noted that this method introduces some error by miscounting the coverage for overlaps and subsurface holes in the film, but such shortcomings should be relatively small in number.

From Figure 4c, the film produced using the liquid/liquid interface indeed shows the expected almost singular distribution of film thicknesses. The total covered area is 0.87 (1 minus area of zero coverage) with the majority of that area only covered by a single layer of flakes (0.71). This is further evidenced by the X-ray diffraction (XRD) pattern for the liquid/liquid assembled film that displays no 002 peak, which suggests that minimal stacking has occurred (Supporting Information, Figure S5). The measured value for the total covered area is close to that found from the analysis of SEM images (0.89) and agrees well with theoretical packing fractions for maximally jammed systems of hard disks in 2D planes (0.78–0.906). Nonzero values for coverages of 2 and greater show evidence that some flake overlap occurs, possibly due to the compressive mechanisms suggested earlier. By comparison, the coverage distributions for the top-down methods (spray and spin coating) are much broader with a greater proportion of the film area exhibiting coverages of 2 or more and an increased proportion of uncovered area (\( c = 0 \)).

To test if the coverage distributions for the top-down methods describe a stochastic flake deposition process, as we would expect, they must be compared to a Poisson distribution. Figure 4d–f shows separate plots of the coverage histograms for the liquid/liquid, spin, and spray-coated films, given in Figure 4c, respectively. Each data set is fitted to a Poisson distribution for the appropriate mean coverage, shown by a solid line. The fit to a Poisson distribution is good for spin-coated and reasonably good for spray films (\( R^2_{adj} = 0.92 \) and 0.85, respectively). This validates our description of these processes as a series of random flake deposition events at the interface. As expected, the fit is poor for the liquid/liquid assembled film (\( R^2_{adj} = 0.23 \)) clearly differentiating the film production mechanism used in this case.

Thin-Film Transistors. Thin-film transistors were manufactured using the liquid/liquid interface assembled MoS\(_2\) film as the channel material. The devices demonstrate good repeatability (device repeatability test data is available in the Supporting Information, Figure S3). The electronic performance of an example FET is shown in Figure 5c,d, with peak mobility \( \mu = 0.73 \) cm\(^2\) V\(^{-1}\) s\(^{-1}\) and on/off ratio = 10\(^5\). This is comparable to previously reported CVD-grown FETs and FETs assembled using pristine mechanically cleaved few-layer MoS\(_2\) single crystals with a back gate geometry and SiO\(_2\) as the gate dielectric. The devices also demonstrate superior properties to most solution-processed MoS\(_2\) FETs reported in the literature (Figure 6).

However, the mobility value for our device is reduced in comparison to previously reported work using the same MoS\(_2\) flake preparation method and a spin-coating film preparation method. This discrepancy is likely due to a number of factors. We expect that around an order of magnitude mobility enhancement may be achievable by reducing coulomb scattering effects via the addition of a high-\( \kappa \) dielectric coating, and further enhancement may be achievable by sulfur vacancy passivation via TFSI superacid surface treatment. We also note that our device testing was conducted under ambient conditions, and around a 4-fold improvement to mobility can be expected by testing under vacuum as in the previous report.

We ascribe the relatively high device performance of these simple FETs to: decreased contact resistance of the edge-to-edge contacts between flakes in the film, minimal heterojunction barriers due to narrow nanoflake thickness distribution and minimal stacking, and increased gate field modulation resulting from the ultrathin and flat nature of the film produced via liquid/liquid interface assembly. As we have shown, such high-quality films are easily produced via this...
method, with minimal parameter optimization, resulting in films with less flake overlap than conventional spin-coating or spray-coating methods. Therefore, by combining the benefits of liquid/liquid interface assembly of thin films with the optimization of device performance via the methods outlined above, vastly improved device characteristics could be achieved in future studies.

**CONCLUSIONS**

Densely packed and predominantly single-layer films of flakes of 2D MoS$_2$ have been produced via assembly at the interface between two immiscible liquids, driven by gradients in interfacial tension. This is shown to be consistent with simple models for particle assembly at the liquid/liquid interface, which have been adapted for 2D material systems with negligible thickness. On comparison with films made using spray- or spin-coating deposition methods, the liquid/liquid assembled films show greater packing density (reduced uncovered area), an increased proportion of edge-to-edge contacts (reduced flake overlap), a greater area covered by a single-flake thickness ($c = 1$), and thus a reduced film roughness (a narrow distribution of coverages). The films have 13% uncovered area, consistent with random tiling models, and are on average 2 nm thick.

Statistical analysis of coverage data, measured using AFM, for films produced by liquid/liquid assembly and two other common, top-down film assembly methods, spin coating and spray coating, highlighted the differences in the deposition mechanisms. The distribution of flake coverage for top-down methods is well described by a Poisson distribution suggesting the stochastic nature of flake placement, whereas the liquid/liquid method shows a more singular distribution of coverages, indicative of flake confinement at the liquid/liquid interface.

Transistors were assembled using liquid/liquid assembled thin films of MoS$_2$ nanosheets as the channel material. The resulting devices display good mobility and on/off ratio with minimal surface treatment, exhibiting equivalent performance to previously reported mechanically exfoliated single-crystal MoS$_2$ transistors with matching device layouts. These properties compare well with those of solution-processed and CVD-grown MoS$_2$-based transistors reported in the literature while requiring minimal postprocessing steps. We attribute this improved performance to the ultrathin and flat nature of the film and the increased proportion of edge-to-edge contacts between flakes achieved by this liquid/liquid assembly method.

The process is scalable, with the potential to be adapted to roll-to-roll processing and could be applied to other 2D, 1D, and 0D materials. Furthermore, ultrapure and degassed solvents can be used as the upper and lower phases to allow the technique to be used in the assembly of air-sensitive materials such as BP and InSe, and we are currently exploring these possibilities.

![Figure 5](https://example.com/figure5.png)

**Figure 5.** (a) Device layout for the prepared FETs. (b) Optical microscopy image of the thin film coating the source/drain electrodes of an assembled FET. Scale bar = 20 μm. (c) Transfer and (d) output curves for an assembled FET with a channel length of 10 μm. $V_{dd} = 0.5$ V for the transfer curve.

![Figure 6](https://example.com/figure6.png)

**Figure 6.** Electrical performance of the FET produced in this work (green star) compared with literature data for FETs manufactured using single mechanically cleaved few-layer MoS$_2$ crystal (blue symbols) CVD-grown MoS$_2$ thin films (black symbols) and solution-processed thin films (red symbols). Data source indicated adjacent to each plotted point. Full data is in the Supporting Information, Table S1.10,55,57,62−69.
**MATERIALS AND METHODS**

**Exfoliation of MoS2.** Thin flakes of MoS2 were prepared using the electrochemical method previously described by Lin et al. This uses the intercalation of bulky tetraheptylammonium (THA) ions to expand the layers within a molybdenite crystal. Since fewer ions are inserted in comparison to the smaller ions used in some intercalation experiments (e.g., Li+), the 2H semiconducting phase of MoS2 is retained. A molybdenite crystal (SPI supplies, West Chester, PA 19381-0856) was used as the cathode, and a graphite strip (Gee Graphite Ltd., Dewsbury, U.K.) as the anode in a two-electrode electrochemical cell. A solution of tetraheptylammonium bromide (THAB) (Sigma, Dorset, U.K., ≥99.0%) in acetonitrile (Sigma, ≥99.5%) (40 mL, 12.5 mg mL−1) acts as the electrolyte. The [THA+ ions were inserted into the layers of the molybdenite crystal under a potential difference of −10 V for 1 h. The expanded molybdenite crystal was washed thoroughly with ethanol (Sigma, 96%) before being sonicated in a 200 mg mL−1 poly(vinylpyrrolidone) (PVP) (Sigma, 30 000 mw) in dimethylformamide (DMF) (Sigma, 99.8%) solution using an ultrasonic bath (P120H, Elmasonic, Singen, Germany) 100 W, for 30 min at 30% power. The material formed a bright green dispersion after the sonication step.

The exfoliated dispersion was separated according to nanoflake size using a cascade centrifugation (1–16, Sigma, Osterode am Harz, Germany). Centrifugation speed of 5900 and 13 200 relative centrifugal force (RCF) for 20 min per step were used to separate nanoflakes of different sizes. After each sedimentation step, the solid sedimented material was redispersed in isopropyl alcohol (IPA) (>99.5%, Sigma) and then washed three times by pelleting in a centrifuge and redispersion in IPA. The material pelleted at 13 200 RCF was used in film assembly and the manufacture of the devices herein. XRD patterns for the MoS2 dispersion, as exfoliated and after size selection, are available in the Supporting Information (Figure S5).

**Liquid/Liquid Thin-Film Assembly.** Thin films were assembled at the interface between a lower phase of hexane (Sigma, ≥99%), density 655 kg m−3, and a lower phase of deionized water, density 995 kg m−3, in a 50 mL glass beaker. The tip of a glass Pasteur pipette containing around 0.4 mL of the dispersion of nanoflakes in IPA (Sigma), density 786 kg m−3, was inserted into the hexane layer. As the dispersion has a higher density than hexane, the IPA dispersion slowly falls from the tip of the pipette to merge with the hexane/water interface. The hexane exchanges with IPA in the closed system until all of the material from the dispersion assembles into a film parallel to the interface; this takes approximately 2 min, depending on the size of the interface and the 2D material used. All of the IPA is eventually exchanged for hexane, completing the transfer of the nanoflake material to the interface. The progress of film deposition can easily be seen by the naked eye (Supporting Information, Video S1); furthermore, the addition of the dispersion can be arrested when the interface is seen to be filled with a monolayer of nanoflakes. The resulting film of the 2D material can then be coated onto a variety of substrates by placing a substrate aligned parallel to and then lifting it away using polyether ether ketone (PEEK) tipped tweezers to select three points to level the plane of the substrate. The z-height value (relative to the leveled substrate) for each pixel in the scan area was used to generate the histograms in Figure 4.

**Flake and Film Characterization.** Flake thicknesses were measured using an atomic force microscope (AFM) (Dimension FastScan, Bruker, Billerica, MA). AFM scans were performed using Antimony (n)-doped Si TESPA-V2 probe tips (Bruker). Scanning electron microscopy (SEM) images of the liquid/liquid assembled films on substrates were obtained using an Ultra S5 FEG SEM (Zeiss, Oberkochen, Germany). XRD patterns were obtained using a Rigaku SmartLab X-ray diffractometer (Rigaku, Tokyo, Japan). UV−vis spectroscopy was performed using a Lambda 25 spectrometer (Perkin Elmer, Waltham, MA).

Lateral flake areas were measured using manipulation of an SEM image of nanoflakes with the Fiji software. First, the SEM image was thresholded to maximize the contrast of the flake area against the substrate area. Next, the “analyze particles” function was used to count the area of each of the particles in the image (n = 714) and calculate the mean lateral flake area. A histogram of the flake areas generated by this analysis is given in the Supporting Information, Figure S1b.

The AFM images were processed and manipulated using the Gwyddion software. Scans were leveled using average plane fitting. The individual scan lines were then flattened using a median of averages. Any large scars caused by impurities in the film were cropped from the final scan area. The substrate plane was then flattened using the three-point method by manually selecting three locations where the substrate was visible (in thick films, where the uncovered substrate cannot be clearly seen, the film can be scratched away using polyether ether ketone (PEEK) tipped tweezers to select three points to level the plane of the substrate). The z-height value (relative to the leveled substrate) for each pixel in the scan area was used to generate the histograms in Figure 4.

**Poisson distributions were fitted to AFM histograms (Figure 4d–f) using the Origin software (OriginPro), Version 2015, OriginLab Corporation, Northampton, MA). The averaged data points with standard deviation were fitted using the Levenberg–Marquardt fitting algorithm and statistical weighting.**

**Three-Phase Contact Angle Measurement.** The three-phase contact angle at the water, hexane, and MoS2 contact line was found using the sessile drop method. A droplet of deionized water was placed on the surface of a MoS2 crystal (SPI supplies), which was submerged in hexane (99%, Sigma). The three-phase contact angle was recorded using a drop shape analyzer (DSA100, Kruss, Hamburg, Germany).

**Device Manufacture and Testing.** To fabricate transistors, n-doped silicon substrates, OFET test chip, and 230 nm SiO2 (IPMS Fraunhofer, Dresden, Germany), with prefabricated interdigitated electrodes (ITO 10 nm, Au 30 nm), were first washed in acetone, water, and IPA with sonication for 10 min with each solvent. The substrates were then coated with the MoS2 thin film using our liquid/liquid assembly method. The film was heated in a tube furnace (MTF, Carbolute, Hope, U.K.) in an Ar atmosphere for 2 h at 200 °C. The devices were tested immediately after removal from the furnace with no surface treatment. Some devices were also soaked in deionized water overnight after the heat treatment to remove any retained (THAB) from the exfoliation step. The n-doped silicon acts as a back gate, while the thin film of MoS2 nanosheets acts as the channel. The interdigitated electrodes are the source and drain for the back gate/back contact (BG/BC) device (Figure 5a).

Device testing was performed using a source meter (2400 series, Keithley, Solon, OH) and a probe station (P4E, Everbeing, Hsinchu City, Taiwan).
The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acsami.0c03794.

Bond number calculation for MoS\(_2\) particles at the interface, demonstrating that capillary forces dominate gravitational forces; thickness and lateral size data for the MoS\(_2\) dispersion; high-magnification SEM image of the MoS\(_2\) liquid/liquid assembled thin film; transfer curves of three repeat FET devices; drop watcher image of the three-phase contact angle among MoS\(_2\) water, and hexane interfaces; XRD patterns for liquid/liquid interface assembled thin films of MoS\(_2\) flakes; table providing literature values used to produce main text (Figure 6) (PDF)

Liquidd/liquid assembly method (Video) (MP4)

**AUTHOR INFORMATION**

**Corresponding Author**

Brian Derby — Department of Materials, University of Manchester, Manchester M13 9PL, UK; orcid.org/0000-0001-5753-0166; Email: brian.derby@manchester.ac.uk

**Authors**

Joseph Neilson — Department of Materials, University of Manchester, Manchester M13 9PL, UK; orcid.org/0000-0002-2472-4869

Michael P. Avery — Department of Materials, University of Manchester, Manchester M13 9PL, UK; orcid.org/0000-0002-2154-1354

Complete contact information is available at: https://pubs.acs.org/10.1021/acsami.0c03794

**Notes**

The authors declare no competing financial interest.

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