Wafer scale synthesis of organic semiconductor nanosheets for van der Waals heterojunction devices

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Organic semiconductors (OSC) are widely used for consumer electronic products owing to their attractive properties such as flexibility and low production cost. Atomically thin transition metal dichalcogenides (TMDs) are another class of emerging materials with superior electronic and optical properties. Integrating them into van der Waals (vdW) heterostructures provides an opportunity to harness the advantages of both material systems. However, building such heterojunctions by conventional physical vapor deposition (PVD) of OSCs is challenging, since the growth is disrupted due to limited diffusion of the molecules on the TMD surface. Here we report wafer-scale (3-inch) fabrication of transferable OSC nanosheets with thickness down to 15 nm, which enable the realization of heterojunction devices. By controlled dissolution of a poly(acrylic acid) film, on which the OSC films were grown by PVD, they can be released and transferred onto arbitrary substrates. OSC crystal quality and optical anisotropy are preserved during the transfer process. By transferring OSC nanosheets (p-type) onto prefabricated electrodes and TMD monolayers (n-type), we fabricate and characterize various electronic devices including unipolar, ambipolar and antiambi bipolar field-effect transistors. Such vdW p-n heterojunction devices open up a wide range of possible applications ranging from ultrafast photodetectors to conformal electronics.

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Although the rest of the OSC film is still sticking to the Si wafer at one side, it remains intact without rupture or cracking. A photograph of an intermediate state of the release of a 50 nm thick pentacene film from a 3-inch Si wafer is shown in Fig. 1g. Finally, by adding more water, the PAA dissolves completely and the whole mechanically stable OSC nanosheet floats on the water surface, Fig. 1c. A photograph of such a 50 nm thick pentacene nanosheet is shown in Fig. 1h. After the organic film is released from the PAA template, the transfer of the OSC onto the target substrate resembles other meniscus-based transfer schemes for all-organic heterojunctions, which instead require soluble molecules. Further details on the PAA dissolution and a time-lapse video of the process are provided in Supplementary Notes 1 and in Supplementary Video 1. Next, the glass support with the floating OSC nanosheet on the wafer is immersed into a water bath (Fig. 1d) and the floating OSC nanosheet is taken by the target substrate, Fig. 1e. Note that transfer of the nanosheet is particularly favorable on hydrophilic target substrates. However, the transfer onto hydrophobic target substrates such as Si wafers after cleanroom processing is also possible. To this end, the edge of the target Si wafer needs to establish a physical contact with the floating OSC nanosheet to enable vdW adhesion as shown schematically in Fig. 1e. Once the adhesion is established, the target wafer with the OSC nanosheet is taken out of the water and dried in air. We found out that positioning the wafer almost vertically allows excess water to drain out easily and dry faster, Fig. 1f. A 50 nm pentacene nanosheet transferred in this way on a 2-inch wafer is shown in Fig. 1i.

**Structural and optical characterization of transferred organic nanosheets**

Morphology and crystallinity of the OSC nanosheets were characterized before and after transfer by AFM and X-ray diffraction. For 50 nm pentacene and DNTT nanosheets, PAA promotes the formation of large crystalline grains with lateral grain size dimensions of ~6 µm and ~3 µm, respectively (see Fig. 2a, d). These grains are much larger than those obtained by the direct growth on oxide surfaces such as SiO2 and Al2O3, or on TMDs, see Supplementary Fig. 3 for a comparison. After transfer, the grain morphology of the OSC nanosheets is preserved as verified by AFM, see Fig. 2b, e and Supplementary Fig. 4. X-ray diffraction further confirms that PAA promotes OSC nanosheet growth in a single crystallographic phase, i.e., before transfer, there is only one series of Bragg peaks for both materials (see black curves in Fig. 2c, f). The Bragg reflections of pentacene change after transfer (see red curve in Fig. 2c). Pentacene is known to exhibit different polymorphs such as thin film and Campbell bulk phase which can be converted in response to stress. Apparently, the transfer induces a stress sufficient to convert some fraction of the thin-film phase into Campbell bulk phase. Such stress might originate from the curvature of the water meniscus during the release of the OSC nanosheet. Stress-induced phase transitions have not been reported for DNTT. In agreement, the crystallographic phase of the DNTT nanosheets is preserved after transfer.

It is well-known that the optical reflection of pentacene grains is strongly polarization dependent for some specific wavelengths, a phenomenon called Davydov splitting. In order to probe for...
Davydov splitting, we raster scan the transferred pentacene nanosheet in confocal geometry with a linear polarized laser beam (635 nm) (see Supplementary Notes 4). Indeed, we found out that the reflection of individual grains depends on the polarization direction of the laser, which is indicated by the yellow arrow in Fig. 2g, h. Each grain reflects only one of the two polarization states efficiently, as verified by a false-colour superposition map in Fig. 2i. This observation confirms that the transferred pentacene nanosheets preserves their specific optical properties.

**Transport characteristics of transferred organic nanosheets**

Next, the electronic transport properties of transferred OSC nanosheets were investigated in the inverted coplanar field-effect geometry, i.e., bottom-contact and bottom-gate FETs, as shown in Fig. 3a (see also Supplementary Fig. 5). A highly doped Si substrate is used as a global bottom-gate electrode and an ALD grown 33 nm alumina (Al2O3) film serves as the gate dielectric. A self-assembled monolayer (SAM) of n-tetradecylphosphonic acid was used to passivate the alumina film. Bottom-contacts were defined by shadow masks, which we use to evaporate a structured Ti adhesion layer and Au contacts. Finally, a 50 nm DNTT nanosheet is transferred on the wafer. The device characterization is shown in Fig. 3b. The DNTT device exhibits an on/off ratio of up to $10^5$ at $V_D = -5.0$ V, a threshold voltage of $-0.63$ V, and a little hysteresis of 217 mV (see Supplementary Fig. 6 for further transistor characteristics). The extracted mobility for DNTT is $0.16 \text{ cm}^2\text{ V}^{-1}\text{s}^{-1}$ and the subthreshold swing is 289 mV per decade. A further improvement of the extracted OSC mobility might be possible by bottom-contact modification. These values show that the OSC/gate dielectric junction is structurally and energetically well-ordered after the nanosheet transfer, i.e. we see only mild signatures of the charge traps that typically lower device performance in OSC bottom-contact devices. In addition, transferred nanosheet FETs with top-contacts show similar transfer characteristics as bottom-contact devices suggesting that the well-known Au bottom-contact problems were resolved by transfer (Supplementary Fig. 7).

Next, we compare the device with a transferred nanosheet with a similar device fabricated via direct PVD of the DNTT. This device yields drain currents, which are more than one order of magnitude lower compared to the transferred DNTT nanosheet device.
heterojunction by transfer of the OSC nanosheets onto ML MoS$_2$ nanosheets. We tested two different device configurations resulting in ambipolar and antiambipolar FETs, see Fig. 4a, c. To this end, high-quality ML MoS$_2$ single crystals$^{26}$ were synthesized on thermally oxidized Si wafers by chemical vapor deposition (CVD) (see Supplementary Notes 6). The MoS$_2$ crystals were transferred by a PMMA assisted transfer process$^{27}$ and Au/Ti source/drain electrodes were patterned. Next, a 50 nm DNTT nanosheet was transferred (see Supplementary Fig. 9a). Figure 4b shows the transfer curves of such a device under ambient conditions demonstrating the characteristic drain current V-shape of ambipolar FETs, see also Supplementary Figure 10. The current in the ambipolar region can be expressed as the sum of an electron and hole current$^{28}$:

$$I_{\text{d}} = \frac{W C_G}{2L} \left\{ \mu_e (V_G - V_{\text{th},e})^2 + \mu_h (V_D - (V_G - V_{\text{th},h}))^2 \right\}$$  

(1)

The drain voltage $V_D$ affects the threshold voltage of one of the two currents, (cf. Eq. 1), here the hole current, and in turn the transfer curve shifts in response to the drain voltage, cf. Fig. 4b. Furthermore, the two asymptotic branches of the drain voltage allow us to model the saturation behavior of the ambipolar device to determine electron and hole mobilities $\mu_e$, $\mu_h$ and threshold voltages $V_{\text{th},e}$, $V_{\text{th},h}$. The DNTT/MoS$_2$ vdW heterojunction device shows rather balanced mobilities of $\mu_e = 0.18$ cm$^2$/V$\cdot$s$^{-1}$ and $\mu_h = 0.25$ cm$^2$/V$\cdot$s$^{-1}$. So far, similar balanced mobilities have only been reported for organic single crystals placed on 2D materials$^{10}$. Such balanced mobilities are important in applications where the device operates in bipolar or ambipolar mode, as needed for, e.g., inverters and oscillators$^{29}$. We performed drain bias dependent ambipolar FET measurements to resolve the nature of the conducting channel. We find that both channels (p and n) can be switched off in the ambipolar FET geometry via the gate, a typical behavior for a semiconductor (Supplementary Fig. 10a, b).

The drain current dip in the transfer curve provides additional information about the subthreshold behavior of the device. According to Eq. 1, the ambipolar current drops to zero for drain voltages smaller than the electron and hole threshold voltage difference. Here, we measure a finite current of $I_{\text{d}} = 100$ pA for $V_D = \pm 5.0$ V in the minimum. This current can be explained by subthreshold behavior of ML MoS$_2$ and DNTT; i.e., we can model current in the minimum with a ML MoS$_2$ subthreshold swing of 2.31 V per decade, see Supplementary Notes 7 for further analysis. The subthreshold swing allows to estimate the density of ML MoS$_2$ subthreshold traps to $N_{\text{sub}} = 9.16 \times 10^{12}$ eV$^{-1}$ cm$^{-2}$ assuming a capacitance per area of $C_i = 38.4$ nF per cm$^2$. Similar values are found for DNTT.

The geometric design of the ambipolar device allows that current flows in the two junction materials without crossing the two materials. In order to enforce current flow across the heterojunction, we employ now the antiambipolar geometry, Fig. 4c. In this geometry, the electrodes are exclusively in contact with one material only. Here, the source is connected to a ML MoS$_2$ and drain to a 50 nm pentacene nanosheet. The overlap region of the two materials, i.e., the p-n junction, is 15 $\mu$m along the channel length. Technically, this configuration was realized with help of an insulating SU-8 polymer coating on one contact, and suited placement of the second contact (see Fig. 4c and Supplementary Fig. 9b, 11). In this antiambipolar geometry, the drain current minimum of the ambipolar geometry changes to a drain current maximum. This peak in drain current is the fingerprint of the antiambipolar geometry, confirming that indeed a functional p-n heterostructure junction formed between a 2D material and a transferred OSC. The signal to noise ratio of the drain current maximum is up to $10^5$, cf. Fig. 4d, exceeding previously reported values for this materials combination$^{10}$.

**Transport characteristics of van der Waals heterostructures**

Now, we demonstrate the vdW heterostructure devices made of OSC nanosheets and ML MoS$_2$. We assemble vdW p-n heterojunction by transfer of the OSC nanosheets onto ML MoS$_2$ nanosheets. We tested two different device configurations resulting in ambipolar and antiambipolar FETs, see Fig. 4a, c. To this end, high-quality ML MoS$_2$ single crystals$^{26}$ were synthesized on thermally oxidized Si wafers by chemical vapor deposition (CVD) (see Supplementary Notes 6). The MoS$_2$ crystals were transferred by a PMMA assisted transfer process$^{27}$ and Au/Ti source/drain electrodes were patterned. Next, a 50 nm DNTT nanosheet was transferred (see Supplementary Fig. 9a). Figure 4b shows the transfer curves of such a device under ambient conditions demonstrating the characteristic drain current V-shape of ambipolar FETs, see also Supplementary Figure 10. The current in the ambipolar region can be expressed as the sum of an electron and hole current$^{28}$:

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**Transport characteristics of van der Waals heterostructures**

Now, we demonstrate the vdW heterostructure devices made of OSC nanosheets and ML MoS$_2$. We assemble vdW p-n
Au/TiSiO\textsubscript{2} by an O\textsubscript{2} (10 sccm) plasma treatment at 50 W for 5 min. An approx. 50 nm thick layer of pentacene or DNTT was immediately loaded into an ultra-high vacuum (UHV) chamber. The OSC transfer method can be used to transfer OSC nanosheets to the improvement of the subthreshold behavior. The OSC transfer can be patterned by e-beam lithography while the antiambipolar FET was patterned by photolithography. OSC nanosheets for FETs were evaporated with a rate of 0.1 Å per s at room temperature for pentacene and DNTT films respectively. In summary, we have developed a method for wafer-scale synthesis and transfer of organic semiconducting nanosheets with thickness down to 15 nm, demonstrated for the example of DNTT and pentacene. The nanosheets can be transferred onto arbitrary substrates by a water-assisted transfer protocol. Unipolar devices fabricated via OSC transfer outperform prepared conventionally by PVD DNTT bottom-contact FETs by more than one order of magnitude in drain current. Fabrication of ambipolar and antiambipolar devices with OSC and ML MoS\textsubscript{2} was demonstrated and such devices reach the performance of heterojunction devices previously built from organic single crystals, outperforming also OSC films deposited by PVD directly on MoS\textsubscript{2}. Such vdW heterojunction devices have potential application in optoelectronics as ultrafast photodetectors or light-emitting devices. Further progress in this direction relies on the improvement of the subthreshold behavior. The OSC transfer method can be used to transfer OSC nanosheets to flexible or curved substrates for applications in large area photodetection (see Supplementary Fig. 14), biomedical sensing applications\textsuperscript{30}, and mechanical sensing as in artificial skin devices\textsuperscript{31}.

**METHODS**

**Organic semiconductor deposition**

Si wafer with SiO\textsubscript{2} oxide surface were sonicated 5 min + 10 min in acetone and in isopropanol then 10 min DI at 60 °C. The SiO\textsubscript{2} surface was activated by an O\textsubscript{2} (10 sccm) plasma treatment at 50 W for 5 min. An approx. 50 nm thick layer of PAA (volume fraction 2.5%, filtered by 0.22 µm pore-sized syringe filter) was spin-coated (60 s at 4000 rpm) onto the Si wafer. The wafers were immediately loaded into an ultra-high vacuum (UHV) chamber and up to 50 nm thick pentacene or DNTT films were evaporated at a base pressure of middle 10\textsuperscript{-8} mbar. The OSC films were evaporated with a rate of 0.1 Å per s at room temperature for pentacene and with a rate of 0.1 Å per s at 60 °C for DNTT.

**AFM and X-ray measurements**

AFM images are recorded by a Bruker Dimensional Icon. The deposition rates and substrate temperatures for the nanosheets used in AFM measurements are 0.02 Å per s at room temperature and 0.1 Å per s at 60 °C for pentacene and DNTT respectively. For X-ray measurements an in house X-ray setup was used with Mo source and a monochromatic beam in reflection geometry. The deposition rates and substrate temperatures for the nanosheets used in X-ray measurements are 0.1 Å per s at room temperature for both pentacene and DNTT films, respectively.

**Device fabrication**

The shadow polyimide mask for patterning organic FETs was manufactured by CADILAC Laser GmbH. For vdW heterostructure FETs, SiO\textsubscript{2} is used as gate dielectric and its thickness is 90 nm for ambipolar DNTT/ML MoS\textsubscript{2} FET and 300 nm for antiambipolar pentacene/ML MoS\textsubscript{2} FET. The ambipolar FET was patterned by photolithography while the antiambipolar FET was patterned by e-beam lithography. OSC nanosheets for FETs were evaporated with 0.1 Å per s at room temperature for pentacene and 60 °C for DNTT films.

**Electrical characterization**

A probe station in dark ambient conditions was used for organic FETs and DNTT/ML MoS\textsubscript{2} ambipolar FET where pentacene/ML MoS\textsubscript{2} antiambipolar FET was measured under vacuum.

**DATA AVAILABILITY**

The data that support the findings of this study are available from the corresponding author upon reasonable request.

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AUTHOR CONTRIBUTIONS

The research project was initiated by B.N. and A.T. S.B.K., A.G., A.T., and B.N. developed the OSC transfer technique. S.B.K. pursued the growth and transfer of organic nanosheets including their characterization with help from F.A.E.N., Z.G. and A.G. carried out the preparation of TMDs. S.B.K. and U.H fabricated OSC/TMD devices. S.B.K. characterized the unipolar and ambipolar FETs while E.N. and A.G. measured the antiambipolar FET. S.B.K. and B.N. analyzed the FET characteristics and wrote the manuscript. All authors contributed to the final form of the manuscript.

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COMPETING INTERESTS

An institutional patent application titled "Method of transfer of organic semiconductor films to a substrate and electronic devices made therefrom" with the German Patent and Trade Mark Office (GPTO) was filed by FSU Jena with the official file number "10 2021 107 057.0." Inventors (share equal part, in alphabetical order): Antony George, Sirri Batuhan Kalkan, Bert Nickel, Andrey Turchanin.

ADDITIONAL INFORMATION

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