**INTRODUCTION**

Following the discovery of graphene, the appearance of 2D transition metal dichalcogenides (TMD) significantly broadened the knowledge in the field of 2D materials, as well as opening potential optoelectronic applications. Besides this, vertical stacks of two TMD monolayers (ML) forming a bilayer demonstrate exciting optoelectronic properties, not present in individual MLs\(^1\). Of two TMD monolayers (ML) forming a bilayer demonstrate potential optoelectronic applications. Besides this, vertical stacks of two TMD monolayers (ML) forming a bilayer demonstrate exciting optoelectronic properties, not present in individual MLs\(^1\). In such TMD bilayers, the two constituent MLs may possess different crystal directions creating in their overlapping region a moiré superlattice. The relative direction between the two MLs is called the twist-angle.

Photoluminescence (PL) studies on MoS\(_2\) TMD bilayers with different twist-angles reveal an interlayer electronic coupling, which corresponds to an indirect bandgap recombination which varies with twist-angle\(^2\). Recently, it was also discovered that the moiré periodic potential in twisted MoS\(_2\) bilayer can modify the properties of phonons in the respective ML constituents to generate Raman modes related to moiré phonons\(^3\). More recent studies revealed the presence of moiré excitons in twisted TMD homo- and hetero-bilayers\(^4\,^5\). The moiré pattern in the crystal symmetry of a twisted bilayer can be controlled through the rotation of the adjacent layers. In this context, the twist-angle is regarded as a new degree of freedom, enabling tuning of the physical properties of the TMD superlattices. Consequently by tuning the twist-angle in real-space, the change in the moiré pattern and consequently to the moiré periodic potential, one can control the interlayer coupling in order to obtain the desired superlattice properties. Obviously, the precise characterization of the twist-angle in moiré superlattices is essential for a global understanding and quality control in such 2D material systems, as well as precise tuning of the respective vdW devices’ performance.

Although TEM is the most commonly used technique to atomically reconstruct twisted TMD bilayers\(^6\), it requires tedious sample transfer on TEM grids, which is incompatible with most 2D materials fabrication and characterization techniques. Apart from being technically challenging, this procedure might eventually distort the relative lattice direction and alignment. Twisted TMD bilayers have also been imaged using atomic force microscopy (AFM)\(^7\), but this requires direct contact with the active area of the bilayer, thus entailing the risk of damaging the sample. While scanning electron microscopy (SEM) techniques do not generally suffer from these limitations, conventional SEM techniques used for crystallographic imaging rely on the detection of backscattered primary electrons, which is not enough to probe mono- or bilayer materials\(^8\).

As far as the optical techniques are concerned, up to date, the estimation of twist-angle in 2D TMD bilayers has been based either on simple optical microscopy observations, or on the production of SHG signals\(^5\,^6\). However, the approaches reported to date do not exhibit high accuracy and cannot image the twist-angle over extended bilayer areas. Having a tool that spatially resolves, with high-resolution and minimally invasively, the twist-angle in large area vdW heterostructures, would be therefore of great importance in the quality characterization of such structures. In this work, we demonstrate such a technique based on the areal imaging of polarization-resolved SHG (P-SHG) signals from TMD superlattices complemented with theoretical modeling that predicts the SHG signals interference from the respective overlapping areas of twisted-bilayers.

2D TMD materials like WS\(_2\) MLs belong to the D\(_{3h}\) point symmetry group with broken inversion symmetry along the armchair direction. This lack of inversion symmetry in the TMD ML results in coherent SHG signals, when an intense field is incident on the 2D material\(^9\,^{10}\). The non-centrosymmetry which creates the SHG signals originates from the honeycomb lattice of the WS\(_2\) 2D crystal, because of the alternating S and W atoms (top view in...\(^1\)Institute of Electronic Structure and Laser, Foundation for Research and Technology-Hellas, Heraklion Crete, Greece. \(^2\)Electron Microscopy for Materials Science (EMAT) University of Antwerp, Antwerp, Belgium. \(^3\)NANOlab Center of Excellence, University of Antwerp, Antwerp, Belgium. \(^4\)Department of Materials Science and Technology, University of Crete, Heraklion Crete, Greece. \(^5\)Department of Physics, University of Crete, Heraklion Crete, Greece. \(^6\)Email: sopsilo@iesl.forth.gr; stratak@iesl.forth.gr

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Optical versus electron diffraction imaging of Twist-angle in 2D transition metal dichalcogenide bilayers

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bilayer atoms configuration of Fig. 1a). The positions of the alternating S and W atoms define the broken symmetry axis of the crystal, which lies in the armchair direction. In contrast, the alternating S and S or W and W atoms define the zig–zag direction of the crystal.

The generated SHG from a WS₂ ML (crystal class D₃h) is described by its corresponding susceptibility tensor, $\chi^{(2)}$. In our approach, we rotate the direction of the linear polarization $\varphi$ of the excitation field and we detect the SHG component parallel to the X-axis ($\zeta = 0(°)$ in Fig. 1b). Then, the recorded SHG from a WS₂ ML is given by (a four-leaved rose-like, polar diagram)$^2$

$$I_{\text{ML}}^{\varphi} = |A\cos(3\theta_1 - 2\varphi)|^2,$$  

(1)

where $A = E_0^2\varepsilon_0\chi^{(2)}$, with $\varepsilon_0$ being the dielectric constant, $E_0$ the amplitude of the excitation field and $\theta_1 \in [0(°)–60(°)]$, defines the armchair direction of the ML modulo 60(°). This means that the armchair directions with $\theta_1 = k \times 60(°)$ in Eq. (1) (where k is an integer) will provide the same P-SHG polar diagram.

When two TMDs, i.e. two WS₂ MLs, are stacked to form a bilayer, the SHG signals from each ML interfere and the total SHG intensity from the WS₂/WS₂ superlattice is described by

$$I_{\text{ML-BL}}^{\varphi} = A^2[\cos(3\theta_1 - 2\varphi) + \cos(3\theta_2 - 2\varphi)]^2,$$  

(2)

where $A = 2\varepsilon_0\chi^{(2)}E_0^2$.

Now, one can use the concept of effective armchair direction $\theta_{\text{eff}}$ in the overlapping region of the two WS₂ monolayers and express the total SHG intensity produced by the 2 MLs as$^7$

$$I_{\text{BL}}^{\varphi} = |A_{\text{eff}}\cos(3\theta_{\text{eff}} - 2\varphi)|^2,$$  

(3)

where

$$A_{\text{eff}} = 2\varepsilon_0\chi^{(2)}E_0^2 \cos \frac{3}{2}\delta,$$  

(4)

where $\delta = \theta_1 - \theta_2$ is the twist-angle, between the MLs and $\theta_{\text{eff}} = \frac{\theta_1 + \theta_2}{2}$.

As a result, the P-SHG modulation emerging from a bilayer region consisting of two WS₂ MLs, at twist angle $\delta = \theta_1 - \theta_2$, behaves as if it was the P-SHG modulation of a single ML with armchair direction $\theta_{\text{eff}}$. The $\theta_{\text{eff}}$ can be extracted experimentally from the P-SHG polar obtained from the bilayer region. In the above, the excitation field propagates along the Z-axis and the $x_1$ armchair direction of WS₂ ML-1 is at angle $\theta_1$ with respect to the X-axis (Fig. 1b). While the second WS₂ ML-2 has its armchair direction lying in the $x_2$ direction, at angle $\theta_2$ with respect to the X-axis (Fig. 1b).

Note in Eq. (4), that the SHG intensity from the twisted bilayer depends on the twist-angle $\delta$, being maximum for $\delta = 0(°)$ and zero for $\delta = 60(°)$. When the two WS₂ MLs are perfectly aligned, e.g. for $\delta = 0(°)$, we have the AA stacking sequence (S or W atoms in one layer lie respectively above the S or W atoms of the second layer (S–S, W–W in Fig. 1a). In this case, the total SHG signal from the bilayer is the result of constructive interference, resulting in SHG intensity four times larger than that of the ML (for $N$ number of MLs the produced SHG signal is analogous to $N^2$). In the case of AB stacking sequence ($\delta = 60(°)$) the S or W atoms in one layer lie, respectively, above the W or S atoms of the second layer (S–W, W–S in Fig. 1c). In this case, centrosymmetry is restored and the SHG signal from the bilayer vanishes (SHG = 0).

The above considerations, refer to the ideal cases of complete constructive or destructive interference in AA and AB bilayer stacking sequences, respectively. However, deviations from these ideal stacking sequences, can occur$^{14}$. A direct consequence of such departures from the stacking sequences AA and AB is the incomplete constructive or destructive interference of the SHG

### Fig. 1 SHG signals originating from WS₂/WS₂ bilayers.

- **a** Schematic representation of the top view in the atomic configuration of a 2D WS₂ bilayer, for the AA stacking sequence. Atoms of W–W or S–S are on top of each other and the SHG signals depend quadratically on the number $N$ of the MLs (SHG~$N^2$).
- **b** Coordinate system of the experimental configuration used for P-SHG imaging. $\theta_1$ and $\theta_2$ denote the armchair directions of ML-1 and ML-2, respectively.
- **c** Atomic configuration for the AB stacking sequence. Alternating W–S atoms are on top of each other and the SHG signals cancel (SHG = 0), because centrosymmetry is restored.
- **d** Block diagram of the experimental setup used for P-SHG imaging. HWP: half-wave plate, GM: galvanometric mirrors, M: mirror, O: objective; SP: sample plane, C: condenser, SPF: short pass filter, BPF: bandpass filter, A: analyzer, PMT: photomultiplier tube.

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from different MLs. In this case, the P-SHG signal modulation depends on the twist-angle between the MLs as shown in Fig. 2.

The graphical representation of Eq. (1) and the corresponding visualization in a polar diagram (presented in Fig. 2) demonstrates a fourfold symmetry of the P-SHG intensity modulation that rotates for different armchair directions \( \theta \). Thus, each armchair direction corresponds to a characteristic fourfold symmetric (four-leaved rose-like) polar diagram. Consequently, we can calculate \( \theta \) by fitting Eq. (1) to experimentally retrieved P-SHG signal intensities with analyzer parallel to the \( X \)-axis and different linear excitation directions \( \phi \). The experimental configuration of a fixed analyzer parallel to the \( X \)-axis used here (\( \zeta = 0^\circ \) in Fig. 1b), is much simpler to that of a rotating analyzer parallel to the rotating linear excitation polarization (\( \zeta = \varphi \) in Fig. 1b), used previously.

The green polar diagrams in Fig. 2 correspond to \( \theta_1 = 0^\circ \) (created using Eq. (1)), while the blue polar diagrams correspond to \( 0^\circ \leq \theta_2 \leq 110^\circ \) with a step of 10\(^\circ\). The red polar diagrams correspond to the product of interference P-SHG in the bilayer region (created using Eq. (2)).

By comparing Fig. 2a, with Fig. 2b, we note the effect of the modulo of 60\(^\circ\) in the calculation of the armchair direction of individual MLs. That is, individual MLs with armchair directions \( \theta^\circ \pm k60^\circ \), where \( k \) is an integer, will provide the same P-SHG polar diagrams. This implies that P-SHG measurements can calculate the armchair direction of an individual ML in the range \( 0^\circ \leq \theta \leq 60^\circ \). Nevertheless, in the case of the bilayer, the SHG signals originate from constructive or destructive interference due to the atomic phase matching between the individual MLs (Fig. 2).

Adapted, as we note in Fig. 2, the SHG signals originating from the bilayers regions with twist-angles of 10\(^\circ\) and 110\(^\circ\), 20\(^\circ\) and 100\(^\circ\), 30\(^\circ\) and 90\(^\circ\), 40\(^\circ\) and 80\(^\circ\), are of equal SHG intensities. Nevertheless, the P-SHG interference polar diagrams from the bilayers' regions (red lines in Fig. 2) for the twist-angles of 10\(^\circ\) and 110\(^\circ\), 20\(^\circ\) and 100\(^\circ\), 30\(^\circ\) and 90\(^\circ\), 40\(^\circ\) and 80\(^\circ\) are different, thus P-SHG is able to identify and discriminate twist-angles that produce similar SHG intensities from the bilayer regions.
RESULTS AND DISCUSSION

PHG measurements

In order to create the WS$_2$/WS$_2$ bilayer, two WS$_2$ monolayers were produced by mechanical exfoliation and were stacked with dry stamping on a Si$_3$N$_4$ support-grid (see Methods). This allows direct comparison between P-SHG and 4D STEM. The excitation source used for the SHG experiments is an fs oscillator, at 1030 nm and repetition rate in the order of MHz, which is adequate to excite non-linear signals like SHG (see Fig. 1b and Methods for the P-SHG microscope).

In Fig. 3a, a CCD image of two individual WS$_2$ MLs (ML-1 and ML-2), forming a WS$_2$/WS$_2$ bilayer on the supporting TEM grid, is shown. One can identify two different types of regions, created by the above stacking, which produce different SHG signals (Fig. 3b, c). Specifically, we identify regions where only one of the two ML-1, and ML-2, WS$_2$ is present, as well as the region where the two MLs spatially overlap and create the WS$_2$/WS$_2$ bilayer. We note that, in the absence of an analyzer in the detection path, the SHG intensity from the bilayer region is lower than the SHG from the ML regions.

In order to calculate the twist-angle of the bilayer, we utilize high-resolution P-SHG measurements with a step of $\varphi = 1^\circ$. Figure 3d presents the respective P-SHG images for fixed analyzer ($\kappa = 0^\circ$) and varying direction of the excitation linear polarization for $\varphi \in [0^\circ)-360^\circ)$. The corresponding polar diagrams (Fig. 3e, f) obtained from two points of interest (POIs 1,2 in Fig. 3c), one for each ML, are fitted to Eq. (1) in order to calculate the armchair direction of the MLs. Then, by using the effective armchair $\theta_{eff}$ obtained from the bilayer (POI 3 in Fig. 3c), we can calculate the twist-angle in the superlattice (using the experimentally retrieved polar of Fig. 3g). We find that $\theta_1 = 44.69^\circ$, $R^2 = 0.92$ in the ML-1 region and $\theta_{eff} = 6.88^\circ$, $R^2 = 0.92$ in the bilayer region.

In previous reports, calculation of twist-angle is performed by measuring the armchair direction of the individual MLs, outside the bilayer region and then deducing their twist angle in their overlapping area. If we follow this strategy we obtain $\delta = \theta_1 - \theta_2 = 44.69^\circ - 29.17^\circ = 15.52^\circ$. Nevertheless, as we see in Fig. 2a, a twist-angle of $15.5^\circ$ should have resulted to constructive SHG interference, i.e. the SHG signal in the bilayer region should have been bigger than the SHG signal of the MLs (Fig. 3e–g). This is not the case for our experimental data, therefore we should follow Fig. 2 and use the modulo $60^\circ$ in the calculation of $\theta_1$ and obtain, $\theta_1 = 44.64^\circ + 60^\circ = 104.64^\circ$. This enables us to unequivocally determine the twist-angle $\delta = \theta_1 - \theta_2 = 104.64^\circ - 29.22^\circ = 75.42^\circ$. We could also use the mean $\theta_{eff}$ acquired from ROI-3 in the bilayer region and Eq. (5) to retrieve the twist-angle as $\delta = \theta_1 - \theta_2 = 2(\theta_1 - \theta_{eff}) = 2(44.64^\circ - 6.96^\circ) = 75.36^\circ \pm 0.55^\circ$.

4D STEM measurements

4D STEM was employed to measure the crystal directions of the individual monolayers, as well to independently estimate the twist-angle in the WS$_2$/WS$_2$ bilayer region seen in Fig. 3a. The schematic representation of the application of the 4D STEM method for studying WS$_2$ monolayers is shown in Fig. 5a. The microscope settings are presented in the Methods section. In particular, the bilayer region that has been analyzed by P-SHG is scanned with 256 × 256 probe positions with a step size of 25 nm. A virtual dark field image (VDF), calculated by summing the intensity of diffraction spots in each probe position over the selected virtual aperture, is shown in Fig. 5b. The inner and outer radii of the virtual aperture are chosen to select the second-order diffraction spots of the WS$_2$ structure (Fig. 5c). From the VDF intensity distribution, we can identify four distinct regions corresponding to the silicon nitride support-gd (dark contrast) (mean diffraction pattern is shown in Fig. 5c), two monolayers (Fig. 5b, e) and an overlapped region with a bilayer (Fig. 5f). At the edges of the monolayers there are regions with increased intensity due to their folding (white arrows in Fig. 5b).

To calculate a direction map of the region, a custom-made peak finding routine was used. Once peak positions are found, two reciprocal vectors can be fitted to describe all diffraction spots defining the relative crystal direction in every probe position. This algorithm works only for diffraction patterns from monolayers. For the bilayer region, four vectors are fitted to each acquired pattern when more than six peaks are detected inside the virtual aperture area. The relative direction map of the region with a bilayer area (white square in Fig. 5b) is shown in Fig. 6a. The tilt angle of every diffraction pattern for two sheets is presented in the histogram Fig. 6d. The twist-angle between the two monolayers was calculated with subpixel accuracy by fitting a Gaussian function for each peak on the histogram and was found to be $15.5^\circ \pm 0.3^\circ$. These results are in good agreement with the experiments on the same area using the all-optical P-SHG imaging microscopy technique, presented above.

However, for the direction determination only the positions of the diffraction spots were used without taking into account their intensity. By looking at the diffraction spot symmetry, one might think that WS$_2$ crystal have six-fold symmetry along the [001] direction, however, due to dynamic electron scattering in combination with the non-centrosymmetry of the crystal, Friedel's
Fig. 3  P-SHG microscopy of WS$_2$/WS$_2$ bilayer on a TEM grid. 

**a** Wide field microscopy using the CCD camera of the microscope. We note two individual MLs of WS$_2$ (ML-1 and ML-2), overlapping in a bilayer WS$_2$/WS$_2$ region. 

**b** SHG imaging of the same region seen in a., without the use of the analyzer in the detection path (see Fig. 1b, d). This ensures that all of the produced SHG signals will be acquired in the same image. Scale bar indicates 5 $\mu$m. 

**c** A zoom of the SHG image seen in (b). Three pixels-of-interest (POI) are chosen. One from the WS$_2$ ML-1, one from the WS$_2$ ML-2 and one from the WS$_2$/WS$_2$ bilayer region. Scale bar shows 1 $\mu$m. 

**d** P-SHG images of the region seen in c. for fixed analyzer ($\zeta = 0^{\circ}$), indicated by a blue double arrow and rotating linear excitation polarization ($\phi \in [10^{\circ} - 90^{\circ}]$, with step 10$^{\circ}$), indicated by a red double arrow. 

**e**, **f** Experimentally retrieved P-SHG polar diagrams for the POIs 1,2, respectively. The polarization rotates with $\phi \in [0^{\circ} - 360^{\circ}]$, and step 1$^{\circ}$ and the analyzer is fixed at $\zeta = 0^{\circ}$. The solid lines (green for ML-1 and blue for ML-2) are the fittings of Eq. (1) to the P-SHG data. This resulted to armchair directions $\theta_1 = 44.69^{\circ}$, $R^2 = 0.92$ and $\theta_2 = 29.17^{\circ}$, $R^2 = 0.96$ for ML-1 and ML-2, respectively, where $R^2$ denotes the quality of the fitting. 

**g** Experimentally retrieved P-SHG data for POI 3 in the WS$_2$/WS$_2$ bilayer region using the same rotating polarization $\phi \in [0^{\circ} - 360^{\circ}]$, with step 1$^{\circ}$ and fixed analyzer at $\zeta = 0^{\circ}$ as in (e, f). The solid red line is the fitting of Eq. (3) to the P-SHG data and $\theta_{\text{eff}} = 6.88^{\circ}$, $R^2 = 0.92$ is the result of the fitting.
law is violated resulting in a symmetry reduction to three-fold symmetry and a nonequivalent angular range from 0° to 120° instead of 0° to 60°. Figure 6(See Figure) shows diffraction patterns of the first and second monolayer together with their azimuthal intensity profiles of the first order spots (Fig. 6(d)) clearly showing 3-fold symmetry. This allows us to identify whether the twist angle is 15.5° or 75.5° (15.5° + 60°) by comparing to simulated diffraction patterns shown in Fig. 6(e, f). The simulation is performed using the Multem software where the experimental conditions such as convergence angle and electron beam energy are chosen to closely resemble the experimental setup (see Methods section). A noticeable difference between the simulated and experimental patterns (Fig. 6(h)) is caused by the presence of the Si3N4-support film, which gives rise to an isotropic background signal. As can be seen in the simulated patterns the intensity of the first order spots changes azimuthally from two low, two high for the twist angle 15.5°, while it has an alternating low-high character for the 75.5° angle. The annular intensity profile of the first order spots of the experimental pattern (Fig. 6(i)), shows a clear signature of a 75.5 ± 0.3(°) twist angle and rules out the 15.5 ± 0.3(°) hypothesis. This result is in excellent agreement with the P-SHG data presented above.

In terms of precision, the STEM data demonstrates 0.3(°) rotational precision compared to 0.55(°) for the P-SHG optical result with a significantly higher spatial resolution but with the downside of a technique which is far less attractive to employ in an inline production environment as compared to the all optical setup proposed in this paper.

Although P-SHG is fast, minimally invasive and can cover large areas, it cannot offer the atomic resolution of the STEM technique. In addition, P-SHG requires breaking of the symmetry and therefore vanishes in intrinsically centrosymmetric systems (such as graphene) or in systems where centrosymmetry is restored e.g. due to the number of layers (i.e., 2H-stacked TMDs with even number of layers) or e.g to homo-bilayers with twist-angle of 60°. For the case of encapsulated TMDs (e.g. between hBN layers) the collected SHG signal from the overlapping region may include additional contributions (as in the case of hBN which is a polar material), and an additional measurement in an uncovered region is needed to unequivocally determine the relative crystal orientation of the individual layers comprising the bilayer.

Considering that the electronic and optical properties of 2D TMD bilayers can be tuned by changing their twist-angle, a robust and minimally invasive tool that can provide spatially resolved determination of the twist-angle, would be of great importance in research, production and large-scale characterization of 2D TMD bilayers. Here, we have used all-optical, laser raster-scanning, P-SHG imaging microscopy to precisely map the twist-angle in large areas of overlapping WS2/WS2 stacked monolayers and we benchmarked the results against 4D STEM electron microscopy. It is found that the twist-angle of WS2/WS2 bilayer obtained using P-SHG mapping is in excellent agreement, either in value and in...
precision, with that obtained using 4D STEM. It is additionally revealed that, given that the produced SHG signal from a bilayer is the vectorial addition of the SHG signals of the individual monolayers, the intensity modulation of the P-SHG signal can be used to deduce unequivocally the armchair direction. This is also in excellent agreement with 4D STEM microscopy analysis.

While STEM provides significantly higher spatial resolution, the P-SHG compensates for this with a wide range of advantages including: no need for vacuum, wide field of view, rapid data acquisition, significantly lower cost and instrument size, and most importantly its capability to work on TMD bilayers deposited on substrates without the need to transfer the films to a TEM grid. Our setup provides an accurate and robust all-optical twist-angle mapping of 2D TMD bilayers. Importantly, the technique is non-destructive, paving the way or directly correlating local twist-angle values with electronic properties, which is crucial for the development and scaling up of vdW bilayer devices with precisely controlled functionality.

**METHODS**

**WS₂/WS₂ bilayer fabrication on a TEM grid**

Polydimethylsiloxane films (PDMS) were fabricated from 10:1 mixing ratio (Sylgard 182 Silicone Elastomer Kit) with heat cure at 80 °C for two hours. High quality WS₂ bulk crystals (HQ Graphene) were mechanically exfoliated directly on the aforementioned PDMS films. The films were placed on typical microscope glass slides using standard protocol. Monolayers of these crystals were realized under an optical microscope. In order to produce the bilayer, at first, a glass slide with a WS₂ monolayer was mounted on a XYZ micromechanical stage under a coaxially illuminated microscope and transferred on a silicon nitride (Si₃N₄) support-grid using viscoelastic stamping. Finally, another WS₂ monolayer was stamped on the previous one in a partial overlapping manner allowing for comparative P-SHG and 4D STEM imaging.

![Image of WS₂ Monolayer and Bilayer](image-url)
Custom-built P-SHG microscope

SHG imaging was performed in the forward-detection geometry using a custom-built laser raster-scanning microscope. As shown schematically in Fig. 1d, a diode-pumped Yb:KGW fs oscillator (1027 nm, 90 fs, 76 MHz, Pharos-SP, Light Conversion, Lithuania), was inserted into a modified, inverted microscope (Zeiss Axio Observer Z1, Germany), after passing through a pair of silver-coated galvanometric mirrors (6215H, Cambridge Technology, UK). A motorized rotation stage (M-060.DG, Physik Instrumente, Karlsruhe, Germany), holding a zero order λ/2 wave plate (QWPO-1030-10-2, CVI-Laser, USA) was used to rotate the direction of the excitation linear polarization. Then, the beam was reflected on a silver mirror at 45° (PFR10-P01, ThorLabs, Germany), placed at the turret box of the microscope, just before the objective (Plan Apochromat 40 × 1.3NA, Zeiss, Germany). The SHG signals were collected from a high-numerical aperture (1.4NA) condenser lens (Zeiss) and guided into a photomultiplier tube (PMT) detector (H9305-04, Hamamatsu, Japan) using another silver mirror (CM1-P01, ThorLabs, Germany) at 45°. The polarization extinction ratio was 28:1. In front of the PMT, a home-built mount was holding a bandpass filter (FF01-514/3-25, Semrock, USA) and a short pass filter (FF01-680/SP-25, Semrock, USA), appropriate for SHG imaging. After the filters, a film polarizer (LPVIS100-MP, ThorLabs) was inserted just in front of the PMT to measure the anisotropy of the SHG signals due to the rotation of the excitation linear polarization. Coordination of PMT recordings with the galvo-mirrors movements and with all the motors, as well as the image formation, was performed using LabView (National Instruments, USA). All of our SHG images were of 500 × 500 pixels and approximately 1.1 s was required for each image to be recorded. This resulted in pixel dwell time of approximately 4.4 μs.

4D STEM measurements

In order to determine local information on the single-layer direction and the bilayer twist angle of 2D materials, a scanning transmission electron microscope ThermoFisher Scientific (FEI) Titan X-Ant-EM was used. The electron microscope was operated in microprobe STEM mode at 300 kV at a convergence semi-angle α of 1 mrad resulting in a probe size of 1.2 nm in diameter. Local information was obtained by scanning the electron probe over the sample and acquiring an electron diffraction pattern at every probe position on a Medipix3 hybrid pixel direct electron detector (Quantum Detectors Merlin) with a camera length of 115 mm and exposure time of 5 ms. In comparison to the SHG pixel dwell time of 4.4 μs, the 5 ms are three orders of magnitude slower. This detector offers a high frame rate and high efficiency enabling the detection of individual electrons without dark or read-out noise and offering 24 bit dynamic range.
REFERENCES

1. He, J., Hummer, K. & Franchini, C. Stacking effects on the electronic and optical properties of bilayer transition metal dichalcogenides MoS2, MoSe2, WS2, and WSe2. *Phys. Rev. B* **89**, 075409 (2014).

2. Liu, K. H. et al. Evolution of interlayer coupling in twisted molybdenum disulfide bilayers. *Nat. Commun.* **5**, 4966 (2014).

3. Lin, M. L. et al. Moiré phonons in twisted bilayer MoS2. *ACS Nano* **12**, 8770 (2018).

4. Jin, C. et al. Observation of moiré excitons in WSe2/WS2 heterostructure superlattices. *Nature* **567**, 76–80 (2019).

5. Seyler, K. L. et al. Signatures of moiré-trapped valley excitons in MoSe2/WSe2 heterobilayers. *Nature* **567**, 66–70 (2019).

6. Alexeev, E. M. et al. Resonantly hybridized excitons in moire superlattices in van der Waals heterostructures. *Nature* **567**, 81–86 (2019).

7. Westen, A. et al. Atomic reconstruction in twisted bilayers of transition metal dichalcogenides. *Nat. Nanotechnol.* **15**, 592–597 (2020).

8. Rosenberger, M. R. et al. Twist angle dependent atomic reconstruction and Moiré patterns in transition metal dichalcogenide heterostructures. *ACS Nano* **14**, 4550–4558 (2020).

9. Schwartz, A. J., Kumar, M., Adams, B. L. & Field, D. P. Electron Backscatter Diffraction in Materials Science (Springer, 2000).

10. Castellanos-Gomez, A. et al. Deterministic transfer of two-dimensional materials and their heterostructures via imaging of their non-linear optical properties (MIS 5050340)” J. V acknowledges funding from FWO G093417N ("Compressed sensing enabling low dose imaging in transmission electron microscopy") from the Flanders Research Fund, EU. J.V. and N.G. acknowledge funding from the European Union under the Horizon 2020 programme within a contract for Integrating Activities No 823717—ESTEEM3. J.V. N.G. and A.O. acknowledge funding through a GOA project "Solarpaint" of the University of Antwerp.

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COMPETING INTERESTS

The authors declare no competing interests.

ADDITIONAL INFORMATION

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