Communication

Controlled synthesis and room-temperature pyroelectricity of CuInP2S6 ultrathin flakes

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1. Introduction

Ferroelectric materials exhibit spontaneous electric polarization due to the separation of the positive and negative charges at both sides of the crystal’s surface. Below $T_c$, ferroelectric materials can be polarized by heating or cooling, and electrical charges will be generated at both sides of the crystals [1–9]. Therefore, ferroelectric materials are naturally pyroelectric.\textsuperscript{[10]} Hundreds of pyroelectric and ferroelectric materials have been discovered for the fabrication of electronic and optoelectronic devices [1–3,11–21]. However, the three-dimensional nature of extensively studied ferroelectric oxide lattices limits the growth of high-quality ferroelectric films and heterostructure. Pyroelectric devices at the nanometer or atomic scale is still under exploration. Recently, two-dimensional (2D) metal phosphorus chalcogenide crystals (MDPs, A\textsuperscript{II}B\textsuperscript{III}P\textsubscript{2}S\textsubscript{6} and A\textsuperscript{II}C\textsuperscript{III}P\textsubscript{2}S\textsubscript{6}, in which A\textsuperscript{II} = Ag or Cu, B\textsuperscript{III} = Mn, Cd or Zn, and C\textsuperscript{III} = V, Cr or In) [22–27] have been revisited and serve as an excellent platform to explore the ferroelectricity in the limit of few-atom-thickness. They are van der Waals (vdW) ferroelectric crystals with low surface energy and could be deposited into 2D nanoflakes simply by mechanical exfoliation. Among them, room-temperature ferroelectricity of 2D CuInP2S6 (CIPS) has been reported with a ferroelectric-paraelectric transition temperature of $T_c \sim 320$ K [28–30]. In our previous studies, we have explored the switchable polarization in ultrathin nanoflake (4 nm) and demonstrated non-volatile memory devices by using a CIPS/Si ferroelectric heterostructure [28]. The atomically thin CIPS provide an interesting platform for the investigations of pyroelectric phenomenon, including in-situ pyroelectric surface/interface, pyroelectric process, and the pyroelectric current at the 2D limits.

In this work, by well controlling the appropriate growth time and temperature, we synthesize high-quality CIPS single crystals by using solid state reaction. Raman spectra reveals that the evolutions of Raman peaks are sensitive to the temperature, which are contributed to the change of polar order and ionic conductivity of CIPS induced by varying the temperature. Kelvin probe force microscopy (KPFM) as an in-situ measurement quantitatively provides information on surface...
topography and temperature-dependent surface potential modification simultaneously, characterizing the pyroelectric charges on the surface of CIPS nanoflakes at nanoscale. Further studies reveal that these pyroelectric charges could be harvested and converted into pyroelectric currents, realizing CIPS based vdW nanogenerator.

2. Experimental section

2.1. Sample preparation

High-quality CIPS single crystals were synthesized by solid state reaction as previously reported [31]. Ultrathin CIPS flakes were obtained by mechanical exfoliation from synthetic bulk CIPS crystals onto heavily doped silicon (Si) substrates.
2.2. Characterization of CIPS nano flakes

Thickness of the flakes was identified from their optical contrast and AFM. PFM measurement was carried out by using an Asylum atomic force microscope (Asylum Research MFP-3D). The CIPS samples for transmission electron microscope (TEM) were prepared by dropping the solution with exfoliated CIPS thin flakes onto lacy carbon TEM grids. Z-contrast STEM imaging was done with a Nion UltraSTEM-100, operated at 60 kV, equipped with a fifth order aberration corrector. The convergence angle is set to be ~30 mrad. All Z-contrast STEM images were acquired from the ~86–200 mrad range. Raman spectra were carried out using a confocal Raman system (WITec) with the 532 nm laser excitation. X-ray diffraction patterns (2θ scans) were obtained from CIPS flakes on the SiO₂/Si substrates using an X-ray diffractometer (XRD Shimadzu Thin Film), using Cu-Kα radiation (λ = 1.54050 Å) within a diffraction angle (2θ) from 5° to 30°.

2.3. KPFM measurement

The KPFM measurements were performed using Bruker Dimension Icon Scanning Probe Microscopy with Nanoscope V controller. All the KPFM images were collected in lift mode by using SCM-PIT probes (Pt/Ir coated probes) with a spring constant of ~3 N/m and a resonant frequency of ~75 kHz. During all KPFM measurements, the optimal lift height is 50 nm (the details in Supporting Information, Fig. S7). Heating process was carried out by using TAC thermal applications controller.

2.4. CIPS device fabrication and measurement

The CIPS devices were fabricated by exfoliating the thin flakes of CIPS onto heavily doped silicon substrates. The top electrodes are defined using standard photolithography process followed by thermal evaporation of the Ti/Au (1 nm/10 nm) metal, and lift-off process. Thickness and size of the CIPS flakes and the Ti/Au electrodes were identified from their optical contrast and AFM.

2.5. Peakforce TUNA measurement

All the pyroelectric current measurements of CIPS onto heavily doped silicon substrates. The top electrodes are defined using standard photolithography process followed by thermal evaporation of the Ti/Au (1 nm/10 nm) metal, and lift-off process.

3. Results and discussion

We prepared the single crystals CIPS by chemical vapor transport (CVT) method as shown in Fig. 1a. Mixture elements of Cu, In, P, and S in stoichiometric proportion 1:1:2:6 were mixed as precursors. The air-
stable yellow sample in the form of platelets can be collected in 2 weeks from evacuated silica tubes. Ultrathin CIPS nanoflakes with different thicknesses from 1 to 2 nm to hundreds were mechanically exfoliated on heavily doped Si substrate. They can be identified by color contrast in an optical microscope as shown in the inset of Fig. 1b and Fig. S1. The morphologies and thicknesses of the nanoflakes were identified by atomic force microscopy (AFM) height profiles as shown in Fig. 1b. The atomic structure and quality of the CIPS crystal were further confirmed by atom-resolved scanning transmission electron microscope (STEM) in Fig. 1c and Fig. 2, with the corresponding fast Fourier transform (FFT) patterns and atomic models along the zone axis [312] and [316]. From the STEM image, the metal cations (Cu, In) and P-P occupy the octahedral voids of a sulfur framework. Cu, In and P-P pairs arrange in the triangular motifs and two adjacent monolayers form a complete unit cell because of the alternate site between Cu and P-P pair from one layer to the neighboring one (the bottom part in Fig. 1c). The CIPS is also characterized by using X-ray diffraction patterns (XRD). A set of strong peaks at 13.58 and 27.34 were assigned to [002] and [004] and another set of weak peaks at 18.57, 20.67, and 23.48 were assigned to [012, 111], [11-3] of the CuInP2S6 crystal (ferrielectric phase, space group Cc at T < Tc) with polar Cu sublattice as shown in Fig. 1d [32].

When T < Tc ≈ 320 K, CIPS is pyroelectric (by heating or cooling) with polar Cu+ and In3+ sublattices shifting in antiparallel directions relative to the midplane, and exhibits reversible spontaneous polarizations [24,31–33]. From the Raman spectra of CIPS, major signals are observed at two regions below 400 cm⁻¹, one at 10 < ω < 60 cm⁻¹ and the other at 200 < ω < 300 cm⁻¹, as shown in Fig. 2 and Fig. S3. In the low-energy range, we observe clear signals of Raman peaks at 26, 35, and 47 cm⁻¹ from Fig. 2a, which can be assigned to the CIPS cation translation modes [33]. Interestingly, the positions of all three peaks show temperature dependence: anomalous decreases for the wave-numbers of 26~21 cm⁻¹, 35~32 cm⁻¹ and 47~44 cm⁻¹ (red shift) are observed in the heating process at T < Tc as shown in Fig. 2b. These three peaks are labelled as 1, 2 and 3 in Fig. 2a–c. Moreover, Raman intensities of the spectral lines in Fig. 2c demonstrate that these three low-energy bands appear obvious intensity redistribution with the temperature increase. These low-energy Raman evolutions can be attributed to the copper cation translations. Heating the CIPS nanoflakes leads to copper hopping motions between its twofold equivalent and the off-center intralayer site and the loss of polarity until CIPS transfers into the paraelectric nonpolar phase [25]. Meanwhile, we can find the Raman peaks in the higher-frequency range, assigning to the P2S6 anion deformation modes [33]. Three mainly three peaks at 220, 240 and 270 cm⁻¹ can be found and they are labelled as 4, 5 and 6 in Figs. 2d–2f. These peaks can be ascribed to the P2S6 deformation variations, including P-P pair off-centering, PS3 group twists, P-S distance and S-P-S angle changes. Apparently, there are blue-shifts for the position of peak 4 and 5 by increasing the temperature from 298 K to 328 K and a slight red-shift for peak 6: 226–243 (blue shift) and 263–262 (red shift). Meantime, obviously intensity increase can be seen for peak 6. These observations correspond to the occurrence of a first-order phase transition in CIPS crystal between 298 K and 328 K. Greater copper content leads to an intensity increase for the band at 226 cm⁻¹ and 263 cm⁻¹ and a decrease for the band at 241 cm⁻¹. Therefore, all of the anion deformation modes are closely related to distortions within the sulfur framework occupied by Cu+. These sensitive temperature-dependent evolutions of Raman peaks are originated from the slightly change of the atom positions within the

Fig. 4. Surface potential investigations for CIPS nanoflakes with different temperature. AFM topography (b) and phase (c) images observed by KPFM mode for the CIPS flake in a. (d–h) Corresponding potential mappings with the temperature of 298 K, 303 K, 308 K, 313 K and 318 K, respectively. (i) Average potential variations versus applied temperature calculated from (d–h). Error bars indicate one standard deviation. The scale bar is 2 μm.
CIPS crystal and therefore regulate the polarity of Cu, In and P$_2$S$_6$ sublattices and the CIPS intrinsic ion potential. The changes of the CIPS intrinsic ion potential and charge displacement will regulate the pyroelectric charges on the CIPS surfaces. In order to characterize the charges quantitatively in real-time, the pyroelectric process of ultrathin CIPS nanoflakes is investigated by using Kelvin probe force microscope (KPFM) with lift mode as shown in Fig. 3a [34–37]. Figs. 3b and 3c illustrates the band diagram changes of the sample and tip in KPFM mode. A contact potential difference V$_{\text{CPD}}$ could be explored based on the electrically contacted and work function differences between the sample ($\varphi_{\text{sample}}$) and the tip ($\varphi_{\text{tip}}$). Therefore, we could track V$_{\text{CPD}}$ in this mode and then determine the work function (Fig. S8 and S9), surface potentials and local charge distributions at the surface of scanned samples. For a thick region in CIPS, e.g. ~30 nm in Fig. 3d (also see Fig. S10a), the surface potential will increase as shown in Fig. 3e. Surface potential of CIPS nanoflakes can be quantitatively determined to be ~ 0.2 V higher than one of the silicon substrate, indicating that a lower charge density at the surface of CIPS nanoflakes. By heating the samples from 298 K to 318 K, the corresponding KPFM image are obtained and shown in Fig. 3f, and the surface potential quickly increases from $-0.64$ V to $-0.51$ V. It suggests that CIPS ultrathin flakes remain pyroelectric down to a few nanometers. Temperature-dependent surface potential was observed even for bilayer flake as shown in Fig. 3g-i (the thickness about 1.8 nm as shown in Fig. S10b), which is the thinnest CIPS crystal calculated to have spontaneous polarization and stable pyroelectric phase [28].

Fig. 5. Pyroelectric devices of CIPS nanoflakes. (a) Illustration of PeakForce TUNA setup for simultaneous topography and electrical property mapping. (b) Plots of Z position, Force, and Current as a function of time during one PeakForce Tapping cycle, with critical points including (B) jump-to-contact, (C) peak force, (D) adhesion labelled. AFM topography (c), TUNA current map (heating from 298 K to 318 K) (d) and TUNA current map (cooling from 318 K to 298 K) (e) of CIPS flake is observed by PeakForce TUNA mode. The scale bar in (c-e) is 2 µm. (f) Schematic and (g) AFM image of the vdW CIPS/Si device with 20 nm thick CIPS. (h) The cyclic changes in temperature (298 K–318 K–298 K and 303 K–308 K–303 K), (i) the corresponding differential rate curve, (j) the measured output TUNA current of the vdW CIPS/Si devices as pyroelectric devices under the temperature changes and (k) the TUNA currents with different electrode sizes are observed by using PeakForce TUNA mode.
Based on the above results of the pyroelectric-induced surface potential changes, we further focus on the temperature sensitivity of pyroelectric CIPS nanoflakes. In Fig. 4, the temperature-dependent surface potential are investigated by heating the samples from 298 K to 318 K with interval of 5 K. Fig. 4d–h show the temperature dependent surface potential changes of few-layered CIPS crystal (the thickness ~30 nm as shown in Fig. S7) from −0.639 V (298 K), −0.598 V (303 K), −0.565 V (308 K), −0.537 V (313 K), to −0.519 V (318 K) with the surface potential change ΔV about ~6 mV/K. By heating the CIPS nanoflakes, the spontaneous polarization will be weakened since the Cu, In and P2S6 electric dipoles vibrations are within bigger spread angle due to the higher hopping motions. As a result, the whole magnitude of the spontaneous polarization is decreased, and less charges will be attracted on the surfaces. Therefore, the elevation of surface potential can be observed with a rise of temperature in Fig. 3 and Fig. 4. On the contrary, if the CIPS flake is cooled instead of heated, the spontaneous polarization evolution leads to the increase of attracted charges on the surfaces. The corresponding KPFM images are shown in Fig. S12, and the magnitude of the potential fleetly decreased from −0.513 mV to −0.644 mV. Based on the temperature-dependent surface potential changes, we could quantify the pyroelectric charge displacement and the pyroelectric coefficient, after clarifying the relationship between the charge displacement and surface potential (the details in the supporting information, Eq. S1-S3), are employed to monitor the pyroelectric current generated on CIPS by changing the temperature. An AFM conductive probe with Peakforce TUNA was used for the measurement. By heating the samples from 298 K to 318 K, the TUNA current from 30 nm CIPS flake is ~ −8 pA as shown in Fig. 5d. The flow of electrons is away from the sample so the current by warming is negative in Fig. S14. On the other way, cooling the CIPS flake will enhance the polarization of CIPS and increase the charge density on the surfaces, then the positive current flows towards the sample (~9 pA in Fig. 5e). These results evidence the pyroelectricity in 2D layers of CIPS dielectric permittivity, with the highest dielectric constant εpyro = 380 [22, 29]. Therefore, according to the formulas as shown in Eq. S1−3, the pyroelectric coefficient pCIPS can be obtained up to ~725 µC/m²/K, with the temperature increase, as shown in Fig. 4i.

Conductive AFM (C-AFM, sub-nA up to µA) and tunneling AFM (TUNA, sub-pA up to nA), as powerful techniques for nano-current characterizations (Fig. 5a–b) [38–40], are employed to monitor the pyroelectric current generated on CIPS by changing the temperature. An AFM conductive probe with Peakforce TUNA was used for the measurement. By heating the samples from 298 K to 318 K, the TUNA current from 30 nm CIPS flake is ~ −8 pA as shown in Fig. 5d. The flow of electrons is away from the sample so the current by warming is negative in Fig. S14. On the other way, cooling the CIPS flake will enhance the polarization of CIPS and increase the charge density on the surfaces, then the positive current flows towards the sample (~9 pA in Fig. 5e). These results evidence the pyroelectricity in 2D layers of CIPS, making it a promising energy-harvesting element in vdW heterostructure. We then fabricated the CIPS based vdW devices via stacking CIPS 2D nanostructures, for the first time.

4. Conclusion

In this work, we systematically studied the pyroelectricity in ultrathin CIPS. The phase transition of CIPS was first investigated via thermal evolution of the CIPS Raman spectra, indicating a transition temperature of around 320 K. Through the temperature dependent surface potential and pyroelectric current measurement, we found that, CIPS 2D nanoflakes, even the bilayer CIPS with ~2 nm thickness could generate temporary pyroelectric surface charge induced by slight atom position modifications and the polarization changes with heating or cooling. Taking advantage of pyroelectrical charges, it is the first time that ultrathin CIPS based pyroelectric nanogenerator were demonstrated based on CIPS nanoflakes for harvesting heat energy into pyroelectric currents at the room temperature. These results greatly enrich the self-powered functionalities of 2D ferroelectric materials and opens new possibilities for thermal nanosensors and ultrathin pyroelectric devices.

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Appendix A. Supporting information

Supplementary data associated with this article can be found in the online version at doi:10.1016/j.nanoen.2019.01.085.

References

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