In-plane and out-of-plane excitonic coupling in 2D molecular crystals

Understanding the nature of molecular excitons in low-dimensional molecular solids is of paramount importance in fundamental photophysics and various applications such as energy harvesting, switching electronics and display devices. Despite this, the spatial evolution of molecular excitons and their transition dipoles have not been captured in the precision of molecular length scales. Here we show in-plane and out-of-plane excitonic evolution in quasi-layered two-dimensional (2D) perylene-3, 4, 9, 10-tetracarboxylic dianhydride (PTCDA) crystals assembly-grown on hexagonal boron nitride (hBN) crystals. Complete lattice constants with orientations of two herringbone-configured basis molecules are determined with polarization-resolved spectroscopy and electron diffraction methods. In the truly 2D limit of single layers, two Frenkel emissions Davydov-split by Kasha-type intralayer coupling exhibit energy inversion with decreasing temperature, which enhances excitonic coherence. As the thickness increases, the transition dipole moments of newly emerging charge transfer excitons are reoriented because of mixing with the Frenkel states. The current spatial anatomy of 2D molecular excitons will inspire a deeper understanding and groundbreaking applications of low-dimensional molecular systems.

Excitons in molecular solids mediate efficient light harvesting in photosynthesis complexes and important organo-electronic applications such as photovoltaics and light-emitting diodes. They also present a myriad of intriguing photophysical phenomena such as Davydov splitting (DS) associated with J/H aggregates, coherence-induced superradiance, mixing of Frenkel and charge-transfer excitons, and singlet fissions. Despite their small footprints spanning a few unit cells, molecular excitons are greatly affected by intermolecular interactions through Kasha-type Coulombic and intermolecular CT couplings. Geometric arrangements are crucial in both interactions in that the former is governed by the long-range coupling among transition dipoles, whereas the latter is by the short-range orbital overlap. Notably, the CT coupling is ubiquitous in π-stacked systems, differentiates the J/H-aggregate behaviors, limits the coherence of Frenkel excitons (FE), and facilitates the singlet fission. Despite many experimental breakthroughs and progress, however, it remains unclear how the evolution of Frenkel-CT mixing can be revealed by other than differing energetics. By exploiting the capability of polarized spectroscopy that directly maps the transition dipoles, more direct proof for the mixing and its spatial progression should be within experimental reach.

For such an exploration, two-dimensional molecular crystals (2DMCs) can be an ideal system. In principle, the above interactions can be tuned by varying intermolecular arrangements, dielectric environments, and the extent of coupled units. In bulk crystals, however, the two former parameters are thermodynamically fixed and can hardly be modified except for polymorphic transitions. Whereas the number of molecular units under excitonic coupling can be purposefully varied by forming oligomers with chemical bonds, it is not applicable to the molecular crystals formed via van der Waals (vdW) interactions, either. In contrast, variable-thickness 2DMCs epitaxially grown on crystalline templates can provide crucial controls over the...
Results and discussion

Structural and optical characterization of 2D PTCDA crystals

Single and few-layer 2D PTCDA crystals (nLP) shown in Fig. 1a were grown on few-layer hBN (nLBN) by physical vapor deposition (PVD) at elevated temperatures. The AFM height image (Fig. 1a) and profile (Fig. 1b) showed a flat layered structure with an interlayer spacing of 0.31 ± 0.08 nm (more examples with optical micrographs are given in Supplementary Fig. 1). As described in Methods, crystalline substrates for growth and excess thermal energy were crucial in obtaining ordered structures. The crystalline nature of the samples was revealed by electron diffraction, as shown in the selected area electron diffraction (SAED) image of 2LG/1LP/nLBN (Fig. 1c), where the encapsulation with bilayer graphene (2LG) led to greatly enhanced stability against electron-induced damages (see Methods). Analysis of multiple samples verified that the rectangular unit cell determined by the diffraction method corresponds to (102)-plane unit cell of bulk crystals. Then, m and n spanning the 2D unit cell (Fig. 1d) are equivalent respectively to b and [201] of bulk: m = 1.256 ± 0.018 (1.220 ± 0.030) nm, n = 1.971 ± 0.017 (1.972 ± 0.008) nm for 1LP (3LP). As summarized in Supplementary Table 1, m of 1LP was ~6% longer than that of bulk, but the deviation was reduced for 3LP, which suggests convergence toward the bulk value with increasing thickness. In addition, the fact that interlayer spacing in 2D PTCDA is very close to that of bulk...
corroborates that 1L\(D\) corresponds to a single (102) plane\(^{29}\). As shown below, DS in PL spectra indicated that two-basis molecules are arranged in a herringbone configuration as in bulk crystals\(^{30}\). The orientation of each basis molecule within the unit cell could be determined by polarization-resolved absorption spectra as described in Supplementary Fig. 2. The simulated diffraction pattern in Supplementary Fig. 3 also showed that 2D PTCDA is closer to the \(\alpha\) polymorph, preferred for higher growth temperature, rather than \(\beta\)\(^{19}\) (see Supplementary Fig. 4 for their structural difference). Note that the unit cells in two neighboring layers are displaced against each other along \(n\) as depicted in Fig. 1d. The formation of the 2D crystalline layered structure is assisted by the interplanar \(\pi-\pi\) interaction\(^{32,33}\) and in-plane hydrogen bonding\(^{34}\). We also note there is a specific orientational registry between \(n_{LP}\) and underlying hBN: the \(n\) axis of 1L\(D\) and 3L\(D\) formed 10.6 ± 0.1° with the armchair direction of hBN (Supplementary Fig. 3h). This indicates that the crystalline nature of 2D PTCDA is supported by substrate-mediated epitaxial growth. In contrast, graphene allows two different rotational polytypes (Supplementary Fig. 3b, c). More restrictive binding on hBN than graphene was also reported for tetracene molecules\(^{35}\).

In addition to the topographic measurements, the average thickness of PTCDA samples could be determined by optical contrast (Methods), which is proportional to the degree of absorption for thin samples supported on transparent substrates\(^{36}\). Figure 1e showed a good resolution for thickness and linearity up to 6 layers. The spectral changes in the room-temperature PL spectra (Fig. 1f) were largely consistent with previous studies on bulk crystals and thin films\(^{32,33}\) whereas monolayer samples present localized FEs with a substantial vibronic progression, thicker samples are dominated by CT excitons denoted as \(C_{T\text{high}}\) and \(C_{T\text{low}}\), which are coupled with lattice phonons to varying degrees\(^{32,33}\). CT excitons have been modeled to reside in 1D molecular stacks under the assumption that the in-plane excitonic coupling is negligible compared to the interplanar \(\pi-\pi\) interactions\(^{9,20,36}\). Although the low-energy PL peaks are mostly of vibronic components, where \(v\) refers to the vibrational quantum number of the electronic ground state and the pair of LDS and UDS corresponds to the 0-0 transition. The origin of the peak at 2.21 eV (marked with asterisks) is unclear. e Polar graph of PL intensity for LDS and UDS peaks, where the data were fitted with the square of cosine. d Polar graph of differential reflectance (DR) for LDS and UDS peaks, for the latter of which an angle-independent constant was subtracted (see Supplementary Fig. 2 for complete DR spectra). e Orientations of molecular and DS transition dipole moments determined with polarized spectroscopy and electron diffraction.

**Fig. 2** Optical anisotropy and epitaxial growth of 1L PTCDA. a Scheme of Davydov splitting in an oblique PTCDA dimer described by ground (\(|\psi_1\rangle\)) and excited-state wavefunctions (\(|\psi_2\rangle\)). LDS and UDS are the superpositions of \(|\psi_1\rangle\) and \(|\psi_2\rangle\), resulting from the Coulombic interactions (\(J_{\text{Coul}}\)) between molecular transition dipoles (oblique arrows). b LDS and UDS-polarized PL spectra of 1L\(D\) on hBN obtained in a parallel configuration, where the polarizations of the excitation beam and PL signals were aligned with \(\mu_{\text{LDS}}\) (upper) and \(\mu_{\text{UDS}}\) (lower), respectively. Complete sets of spectra (Supplementary Fig. 5) were globally fitted with multiple Gaussian functions for 0-\(v\) vibronic components, where \(v\) refers to the vibrational quantum number of the electronic ground state and the pair of LDS and UDS corresponds to the 0-0 transition. The origin of the peak at 2.21 eV (marked with asterisks) is unclear. e Polar graph of PL intensity for LDS and UDS peaks, where the data were fitted with the square of cosine. d Polar graph of differential reflectance (DR) for LDS and UDS peaks, for the latter of which an angle-independent constant was subtracted (see Supplementary Fig. 2 for complete DR spectra). e Orientations of molecular and DS transition dipole moments determined with polarized spectroscopy and electron diffraction.
corresponding to 2JCoul will be governed by mutual orientations and measurements, respectively. The only experimental EDS, 37m eV for a complete set of spectra. In Fig. 2b presenting LDS and UDS-polarized spectra, the 0-0 transition led to two prominent DS peaks with polarized spectra, the 0-0 transition led to two prominent DS peaks. The scheme in d depicts an energy inversion in DS and the associated change in Boltzmann-type populations with decreasing temperature. Coherence number (Nc) of ILp determined by comparing linewidths following: @c = \sigma_{\text{mono}} / \sigma_{\text{pol}} and \( \sigma \) = \( \sqrt{T} \), where \( \sigma_{\text{mono}} / \sigma_{\text{pol}} \) is FWHM of 0-0 peaks for monomeric (2D) PTCDA and T is temperature, respectively. PL spectra of monomeric solutions are given in Supplementary Fig. 8. Dotted lines in e and f are linear fits, whereas those in d and f are guides to the eye. Error bars in e and f indicate standard deviations obtained from multiple samples.

and \( \varphi_p \). Under a point-dipole approximation, the split energy \( (E_{\text{DS}}) \) corresponding to \( 2\mu_{\text{pol}} \) will be governed by mutual orientations and displacement between the two transition dipole moments that are represented by the red (blue) arrow pairs in Fig. 2a. Their transition dipole moments (\( \mu_{\text{LDS}} \) and \( \mu_{\text{UDS}} \)) are expected to be orthogonal to each other, no matter what angle the transition dipole moments of the two bases (\( \mu_{\text{L}} \) and \( \mu_{\text{H}} \)) form. To determine the orientations of the DS dipoles, we obtained polarized PL spectra of ILp in a parallel configuration by rotating excitation polarization (see Supplementary Fig. 5 for a complete set of spectra). In Fig. 2b presenting LDS and UDS-polarized spectra, the 0-0 transition led to two prominent DS peaks with \( E_{\text{DS}} \) of 36 meV. Their polar intensity graph in Fig. 2c verified their orthogonality (90.2 ± 1.3 deg for multiple samples). LDS still showed significant signals at the intensity minimum unlike LDS, the origin of which is not clear. The DS of 0–1 and 0–2 peaks could not be discerned because of their large spectral widths and overlap. Notably, the DS has not yet been observed in any form of PTCDA solids directly by emission or absorption spectroscopy. In addition to the lacking availability of single crystals of sufficient quality and size, dominant CT emission and extremely high absorption impede the conventional measured measurements, respectively. The only experimental \( E_{\text{DS}} \) of 37 meV for \( \alpha \)-PTCDAs, was determined by ellipsometry.

Correlated measurements of electron diffraction and polarized PL (Supplementary Fig. 2) allowed us to determine the orientation of \( \mu_{\text{LDS}} \) and \( \mu_{\text{UDS}} \) with respect to the crystallographic coordinates of 2D PTCDA. As shown in Fig. 2e for ILp, \( \mu_{\text{LDS}} \) was found to make an angle of 21.5° with the \( n \) axis. We note that this information can be used in determining the molecular orientation in 2D PTCDA because molecular transition dipoles are aligned along the long molecular axis. Assuming that \( \mu_{\text{L}} \) and \( \mu_{\text{H}} \) are identical in their magnitudes, \( \mu_{\text{DS}} \) (\( \mu_{\text{LDS}} \)) is parallel (perpendicular) to the bisector of the angle (\( \theta_{\text{LH}} \)) formed by \( \mu_{\text{L}} \) and \( \mu_{\text{H}} \) (Fig. 2e). As \( \theta_{\text{LH}} \) governs the relative magnitude of the DS dipoles, it could be determined by comparing the absorption by LDS and UDS (see Supplementary Fig. 2 for details). The scheme in Fig. 2e depicts the molecular arrangement of 1LP determined by the diffraction, polarized PL, and absorption measurements. The fact that \( \theta_{\text{LH}} \approx 89.2° \) indicates that the two bases are aligned almost perpendicularly to each other, unlike bulk crystals with \( \theta_{\text{LH}} \approx 96° \).

In Fig. 3, we show a distinctive temperature dependence of the DS peaks of ILp. Whereas the emission from LDS (UDS) of J (H) dimers is dipole-forbidden, both DS transitions of oblique dimers found in 2D PTCDA and tetracene44 are allowed. Because of rapid internal conversion, the emission from UDS is insignificant unless \( E_{\text{DS}} \) is close to or smaller than the thermal energy like PTCDA (Fig. 2a). For example, 2D tetracene showed negligible UDS intensity because of its large \( E_{\text{DS}} \) (95 meV) which hinders the thermal population of UDS. The two sets of polarized PL spectra of ILp (Fig. 3a, b) clearly show that LDS and UDS redshift with decreasing temperature. Remarkably, the rate of redshift is 2.7 times larger for UDS than LDS as shown in Fig. 3c (see...
Kasha crucial insight into the nature of excitonic coupling in 1LP. According to induced dephasing, the prediction in Fig. 4a is qualitatively consistent with decreasing temperature, more DS dipoles undergo coherent coupling, which can be approximated as the double-cell and N-cell cases in a. The delocalized coupling leads to excitonic bands for LDS (red dotted band) and UDS (blue dotted band). Only the uppermost (lowermost) state is optically accessible for the LDS (UDS) band, where \( k \) represents the crystal momentum. A 2D arrangement of molecular transition dipoles determined for 1LP and used for the electrostatic calculations shown in c. Four type-II dipoles (red and blue arrows) in the first shell of the reference dipole at the center (black arrow; type-I) orient in opposite directions for LDS (red arrows) and UDS (blue arrows) coupling, whereas all type-I dipoles including eight in the second shell (gray arrows) remain unchanged. Type-I dipoles in the two orange quadrants interact repulsively with the reference dipole. Energy of the reference transition dipole moment interacting with another delocalized over neighboring type-II dipoles. Coherence length is defined as the number of type-II shells surrounding the reference dipole: the first and second shells contain 4 and 12 type-II dipoles, respectively. We used 6.93 Debye for the monomeric transition dipole moment and \( \varepsilon = 1 \) for simplicity.

Supplementary Fig. 6 for more data and Supplementary Fig. 7 for complete spectra), which leads to the inversions in the peak energies (Supplementary Fig. 7) and intensity ratio (Fig. 3c, d). The spectral narrowing (Fig. 3e) accompanied by the increase of their intensities suggests that their excitonic decay is mainly governed by superradiance, an accelerated spontaneous emission from coherently coupled or delocalized excitons. The degree of delocalization will be discussed later using the molecular coherence number, \( N_e \), which was extracted from the linewidth (Fig. 3f), which leads to the inversions in the peak energies (Fig. 3e).

The temperature dependence of both DS peaks in Fig. 3 provides a crucial insight into the nature of excitonic coupling in 1LP. According to Kasha’s exciton model, the excited state of a herringbone-configured PTCDA dimer splits into LDS and UDS (Fig. 4a), and their average energy is slightly lower than that of the excited monomer. The former is induced by the Coulombic interaction between two transition dipoles, whereas the latter is due to differing vdW interaction between ground-state and excited molecules. The long-range Coulombic interaction can be further extended to 2D arrays of unit cells containing two bases, where the number of unit cells represents the degree of exciton delocalization. For example, the inter-dipole interaction will split LDS and UDS into two states for double cells and N states for N cells (Fig. 4a). Because only the uppermost (lowermost) state is optically allowed for the LDS (UDS) band, the optical transition energy increases (decreases) for LDS (UDS) with increasing N. As excitons are more delocalized at a lower temperature because of reduced phonon-induced dephasing, the prediction in Fig. 4a is qualitatively consistent with the differing shift rates of the DS peak energies (Fig. 3c). For a more realistic description, one may need to consider the influence of structural disorder that can be present in samples. Disorder limits the degree of delocalization and induces a redshift in excitonic states under short-range potential fluctuations.

To account for the long-range Coulombic interaction among transition dipole moments of excitons delocalized in 2D crystals, we performed a simple electrostatic calculation under the point-dipole approximation. Using the structural information revealed by the diffraction and polarized spectroscopy, 2D arrays of transition dipoles for 1LP were generated as schematically given in Fig. 4b. The reference dipole (black arrow; basis I) at the center interacts attractively and repulsively with four type-II bases (first-shell neighbors) under LDS (red arrow) and UDS (blue arrow) coupling, respectively. Figure 4c shows that the energy for UDS (LDS) coupling decreases (increases) with increasing the number of interacting dipoles represented by the coherence length, which is defined as the number of type-II shells interacting with the reference dipole in Fig. 4b. As \( L_c \) is equivalent to length, it is proportional to \( N_e \). We note that UDS is stabilized by -10 meV when \( L_c \) increases sufficiently and this value is one-third of the change in LDS energy as temperature decreased from 298 K to 77 K (Fig. 3c). The opposite \( L_c \)-dependences of LDS and UDS captured in Fig. 3c provides a qualitative justification for the observed energy inversion despite the limitations of the simple model. Quantitative description can be obtained by considering exciton-phonon interactions and non-dipolar short-range interactions.

We note that the significant redshift found for 2D PTCDA differs from the behavior of bulk polyacene crystals. Negligibly small temperature dependence of the excitonic energy of the bulk system is due to the almost complete cancellation of two competing factors: thermal expansion and exciton-phonon coupling. The sole contribution of the former is also insignificant for the current system assuming compliant thermal expansion: despite its negative thermal expansion coefficient (TEC), few-nm-thick hBN shrinks with decreasing temperature when supported on amorphous quartz substrates with positive TEC because hBN complies with the thermal expansion of underlying substrates because of the vdW bonding between the two. Similarly, 2D PTCDA is likely to comply with hBN because of their epitaxial registry. Then, the TEC of amorphous quartz predicts that the substrate-induced thermal expansion of 2D PTCDA by cooling from 298 to 79 K is \( -0.02 \% \), for which many-body calculations' predicted sub-meV redshift for polyacene molecular crystals. In case 2D PTCDA...
does not comply with hBN, however, its contraction induced by the cooling may reach up to 1% based on its bulk TEC. Then, the contraction-induced redshift can be nonnegligible and may contribute to the observed redshift.

The superradiant behavior of 2D PTCDA shown in the spectral narrowing and redshift indicates that both DS excitons reside in multiple molecules, the number of which can be equated to \(N_c\). Because \(N_c\) is inversely proportional to \(\sigma\), it can be determined by comparing LDS and CT peaks with monomeric PTCDA dissolved in solvents as shown in Fig. 3. For a similar dielectric environment as PTCDA crystals with a dielectric constant (\(\varepsilon\)) of 3.6, chloroform (\(\varepsilon = 4.81\)) and toluene (\(\varepsilon = 2.38\)) were used as solvents. As shown in Supplementary Fig. 8, monomeric P.T. spectra exhibited prominent vibrational progressions like 1LP but with twice larger linewidth. Figure 3f shows that \(N_c\) increases from -3 for 298 K to -7 for 77 K. Viewing the simplicity of the model in Fig. 4, the estimated change in \(N_c\) needs to be considered semi-quantitatively: at 298 K, a given excited molecule (basis I) interacts mainly with neighboring molecules in the first shell (basis II), which leads to DS. At a lower temperature, the excitonic wavefunction becomes more delocalized beyond the first shell. Whereas \(N_c\) of -4 and -14 have recently been obtained at 77 K for single-layered brick-wall type PTCDA films, the in-plane excitonic coupling within bulk (102) plane had been considered negligible.

Out-of-plane coupling via Frenkel-CT mixing in few-layered PTCDA crystals

In this section, we show how CT excitons are modified by mixing with FE states using multilayer PTCDA crystals. CT excitons often dominate or affect the excited state of molecular aggregates with significant \(\pi-\pi\) interactions. When CT states are energetically close to Frenkel states, their coupling can be significant enough to affect their transition energies and transition dipole moments. Using momentum-resolved electronic energy loss spectroscopy, for example, Knupfer et al. observed an energy-dispersive state from PTCDA crystals and derived the presence of Frenkel-CT coupling. Whereas mixing-induced energetic variation is known for some other representative systems, the effects on transition dipole moments have not been directly observed. In this regard, thickness-controlled 2D PTCDA is a good model system to unravel the progression of Frenkel-CT mixing. Previous studies established that \(\pi-\pi\) interactions are so dominant that bulk PTCDA crystals can be viewed as a non-interacting assembly of 1D \(\pi\)-stacked PTCDA columns. Figure 5 shows the polarized PL spectra obtained for various thicknesses at 77 K. Unlike 1LP, thicker crystals exhibited a few broad bands including CT high and CT low, which are located at slightly higher energies than 298 K (see Supplementary Fig. 9 for complete angle-resolved polarized spectra for 1LP, 6LP). We note that the 0-0 emission, mostly originating from LDS at 77 K (Supplementary Fig. 9), is still preserved in multilayers, whereas the other vibronic bands are not discernible because of the intense CT emissions. As shown in Supplementary Fig. 9, the LDS emission of multilayers is also plane-polarized. This fact suggests that the 0-0 emission of multilayers originates from FEs localized at the surface layers and subsequently not coupled to CT states. In contrast, the CT emissions exhibited a non-Malus behavior with distinct dependence on thickness. As shown in Fig. 5b, c and Supplementary Fig. 9, the polar graphs of the CT peaks for 2~3LP are elliptical unlike that of LDS, which suggests that each CT peak consists of transitions from multiple states. A similar observation could be made for 2LP~6LP samples (Supplementary Fig. 9). Remarkably, the transition dipole moments (\(\mu_{\text{CT}}\)) of CT states approach \(\mu_{\text{LDS}}\) in orientation with an increasing thickness (Supplementary Fig. 9). Assuming that \(\mu_{\text{CT}}\) is aligned along the major axis of ellipses, the angle (\(\theta_\text{T-LDS}\)) between \(\mu_{\text{CT}}\) and \(\mu_{\text{LDS}}\) was determined (Fig. 5d): \(\theta_\text{T-LDS}\) decreased from 67° at 2LP to 19° at 6LP.

As depicted in Fig. 5e, the thickness-dependent reorientation of \(\mu_{\text{CT}}\) is attributed to the varying degree in Frenkel-CT mixing. For qualitative understanding, we adapted a perturbative mixing model to multilayer PTCDA crystals as given in Supplementary Note 1. In essence, observed CT states are viewed as a mixed state of
unperturbed CT (CT*) and FE states (Fig. 5e), and can be described in a perturbation approach. For \( n \)-stacked dimers representing 2Ls shown in Fig. 5f, the mixing generates symmetric (+) and antisymmetric (−) wavefunctions (Equation S3 of Supplementary Note 1). Whereas the transition dipole moment (\( \mu_{\text{CT}} \)) of the former vanishes because of symmetry, that of the latter is modified from \( \mu_{\text{CT}}^e \) by \( \sqrt{2(\kappa - \lambda)} \mu_{\text{FE}} \), where \( \kappa \) and \( \lambda \) can be related to hole and electron transfer integrals, respectively. When expanded to \( n \)-mers for NLs, the degree of FE-CT mixing may decrease because of the increasing energy difference between two constituent unperturbed states (Supplementary Note 1). This suggests that the contribution of FE to CT state decreases with increasing thickness, which is schematically depicted in Supplementary Fig. 10a: For 1LP, \( \mu_{\text{LDS}} \) for thick layers. Notably, the transition dipoles \( \mu_{\text{LDS}} \) and \( \mu_{\text{CT}} \) are oriented along the in-plane axis (3D description is given in Supplementary Note 1). This suggests that the contribution of FE to CT state decreases with increasing thickness, which is schematically depicted in Supplementary Fig. 10a: For 1LP, \( \mu_{\text{LDS}} \) for thick layers.

We note that the model analysis not only supports the experimental observation in Fig. 5d but also provides valuable insight into the nature of the Frenkel-CT mixing. Specifically, the current results visualized the orientational progression of Frenkel-CT mixing by mapping transition dipoles with respect to crystallographic axes, which is unprecedented to our knowledge. It is also notable that the experimental \( \theta_{\text{CT-LDS}} \) approached its theoretical limit of \( \sim 21.5^\circ \) (Supplementary Fig. 11) with increasing thickness to 6L. This sets a rough thickness limit for the bulk behavior in regard to FE-CT mixing. Finally, thickness-dependent mixing can also be utilized in modulating exciton functions and applications in various transport devices.

In this work, we described the spatial evolution of the intralayer and interlayer excitonic couplings in 2D PTCDA crystals using polarization-resolved variable-temperature spectroscopies. 2D PTCDA grown on hBN crystals by physical vapor deposition enabled fine controls over the thickness and crystallographic orientation. Electron diffraction patterns combined with polarized spectra led to the crystallographic details including molecular orientations of two herringbone-arranged bases. For 1L, the 0-0 emission pair Davydov-split by Kasha-allowed and forbidden, respectively, with increasing thickness, which was successfully explained with a Frenkel-CT mixing model. This work will lead to a deeper understanding of molecular excitons bound in low-dimensional organic materials and thus contribute to the better design of molecular structures with novel functions.

### Methods

#### Growth of 2D PTCDA crystals

Mono- and few-layer PTCDA crystals were grown on high-quality inorganic 2D crystals by the self-limited epitaxy. Briefly, few-layer hBN and graphene were mechanically exfoliated on amorphous quartz and 285 nm SiO\(_2\)/Si substrates from hBN powder (Momentive Performance Materials) and natural graphite (NGS Naturgraphite GmbH). PTCDA powder (TCI, >98%) and exfoliated samples were placed at the center and downstream of a tube furnace (ThermoFisher, Lindberg/Blue M), respectively. After the quartz tube was purged with a high-purity Ar gas for 30 min at 500 mL/min, the flow rate was reduced to 100 mL/min. Subsequently, the tube was heated to 330 °C at a rate of -22 °C/min and the temperature was maintained for 15-30 min. Sublimated PTCDA molecules were found to deposit as 2D crystals on hBN and graphene. The average thickness was controlled by varying the deposition time and location within the tube.

#### PL spectroscopy

The employed micro-spectroscopy setup has been described elsewhere. Briefly, solid-state laser beams of 457.8 and 514.3 nm (Cobolt, Twist, and Fandango) were focused onto samples within a spot size of ~1 μm using a microscope objective (×40, NA = 0.60). The signals were collected using a CCD camera (Princeton Instruments, PyLon) with a spectrometer (Princeton Instruments, SP2300). All measurements were performed in the ambient conditions unless noted otherwise. The average power was maintained below 2 μW to avoid unwanted photoinduced degradation.

#### Temperature-dependent PL spectroscopy

A tunable pulsed source (Coherent, Chameleon Ultra II and Compact OPO) was used for variable-temperature PL measurements. Excitation beams of 515 nm were focused with an objective (×50, NA = 0.55) onto a 1-μm focal spot on samples. The light source was operated at a repetition rate of 80 MHz generating pulses with 140-fs duration. Samples placed in a liquid-N\(_2\) cryostat (Linkam, HMS600-PEB4) could be cooled down to 77 K. PL signals were collected by a Czerny-Turner spectrometer (Andor, SR303i) equipped with an EMCCD camera (Andor, DU971P). For polarization-resolved experiments, an analyzing polarizer was placed in front of the spectrometer to select the polarization component parallel to that of the plane-polarized excitation beam. The polarization vector of the excitation beam could be rotated within the sample plane by rotating an achromatic half-wave plate, which was mounted on a rotational stage (Thorlabs, KPRM2E/M) and placed between the objective and tube lens. The average power was maintained below 0.2 μW to avoid unwanted photoinduced degradation.

#### Optical contrast and differential reflectance

Optical absorption of 2D PTCDA on transparent substrates could be measured using its differential reflectance (DR), defined as \( (R - R_0)/R_0 \), where \( R \) and \( R_0 \) are the spectral reflectance of sample and bare substrate, respectively. Optical contrast, essentially DR, was similarly determined using the green-channel signals of optical micrographs.

#### AFM measurements

The topographic details of samples were studied with an atomic force microscope (Park Systems, XE-70). Height and phase images were obtained in a noncontact mode using Si tips with a nominal radius of 8 nm (MicroMasch, NSC-15).

#### TEM sample preparation and measurements

PTCDA on 2D crystals (hBN or graphene) was covered with few-layer graphene flakes before transferring to TEM grids. Then, the resulting samples of graphene/PTCDA/hBN (or graphene)/SiO\(_2\)/Si were overlaid with polydimethylsiloxane (PDMS) film. When soaked in ethanol/DI water mixtures (ethanol 50%, DI water 50%) for 2 h, the 2D hetero layers of the samples were released from SiO\(_2\)/Si substrates and transferred to the PDMS film. The heterostructures on PDMS were dried under ambient conditions and transferred again to holey Si TEM grids (Norcada Inc.) by stamping via a micromanipulator under optical microscope observation. SAED acquisition was performed with a JEOL ARM 200 F equipped with image and probe aberration.
correctors operated at 80 kV. SAED simulations were performed by CrystalDiffract (CrystalMaker Software Ltd.).

Data availability
The authors declare that all data supporting the findings of this study are available within the paper and its supplementary information file or available from the corresponding author upon request.

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Author contributions
S.R. conceived the project. D.K. and S.R. designed the experiments. D.K. prepared samples, performed spectroscopy experiments, and analyzed the data. J.P., J.L., and H.C.C. established the growth method of PTCDA crystals. S.L. and K.K. conducted the SAED measurements and analysis. D.K. and S.R. wrote the manuscript with contributions from all authors.

Competing interests
The authors declare no competing interests.

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