

Cite this: *J. Mater. Chem. C*, 2021,
9, 10875

Conformationally rigid molecular and polymeric naphthalene-diimides containing C₆H₆N₂ constitutional isomers†

Vincenzo Mirco Abbinante,^a Gonzalo García-Espejo,^a Gabriele Calabrese,^b Silvia Milita,^{*b} Luisa Barba,^c Diego Marini,^b Candida Pipitone,^d Francesco Giannici,^d Antonietta Guagliardi^{*e} and Norberto Masciocchi^{ib}^{*a}

Organic thin films based on naphthalenediimides (NDIs) bearing alkyl substituents have shown interesting properties for application in OLEDs, thermoelectrics, solar cells, sensors and organic electronics. However, the polymorphic versatility attributed to the flexibility of alkyl chains remains a challenging issue, with detrimental implications on the performances. Aryl analogues containing C₆H₆N₂ constitutional isomers are herein investigated as one of the possible way-out strategies. The synthesis of molecular and polymeric species is described, starting from naphthaleneteracarboxyldianhydride with isomeric aromatic amines and hydrazine. The materials are fully characterized by spectroscopy, thermal and structural X-ray diffraction methods, both as bulk powders and thin films, revealing a rich structural landscape. Depending on the stereochemistry of the branching aryls, the compounds show a variety of parallel stacking of the NDI cores, and high structural stability upon heating, up to 560 °C in the polymeric form. Thin films prepared by spin coating from organic solvent solutions and studied by grazing-incidence X-ray diffraction exhibit a high degree of crystallinity indicating the intrinsic tendency of these molecules to self-assemble in an ordered fashion without the need for any post-processing technique. In line with other NDI-based diimides, UV-vis spectroscopy indicates the optical band gaps falling in the visible region (2.87–3.02 eV). DFT calculations reveal a significant lowering of the frontier orbital energies of the hydrazido derivative. Beyond solution processing, the high thermal stability and the absence of polymorphic forms of these materials suggest that sublimation-based routes for films and device preparation can also be followed.

Received 4th February 2021,
Accepted 16th June 2021

DOI: 10.1039/d1tc00564b

rsc.li/materials-c

1. Introduction

Organic molecular semiconductors are a class of compounds that have attracted a great deal of attention, in the form of thin films, for appealing applications in photovoltaics, optoelectronics, computing and sensing¹ and, more recently, low-power thermoelectric generators.² These compounds greatly benefit from easy

chemical functionalization, rapid solution processability of pristine molecular entities, simple deposition of thin films and superb flexibility.³ After several years of fundamental studies, they have recently been incorporated into large commodities,⁴ and rapidly filled the market as flexible light panels, rollable solar cells, cell phone displays and battery-free artificial retina, to mention a few.⁵

Among the most investigated species, a number of studies focused on symmetrically disubstituted naphthalene diimide (NDI)⁶ and perylene diimide (PDI)⁷ molecules, where the presence of π - π stacked aromatic cores, in optimized geometrical arrangements, enhances important functional properties, such as electron conductivity above all.⁸ In particular, NDI compounds (having a smaller aromatic core than the PDI congeners) possess two strong electron-withdrawing imide groups enabling several NDI-based systems to achieve relatively low LUMO energies, excellent ambient stability, and promising charge-transport properties. Nonetheless, very slight changes in the nature or position of the side groups may result in a big

^a Dipartimento di Scienza e Alta Tecnologia & INSTM, Università dell'Insubria, via Valleggio 11, 22100 Como, Italy. E-mail: norberto.masciocchi@uninsubria.it

^b Istituto per la Microelettronica e Microsistemi, Consiglio Nazionale delle Ricerche, via Gobetti 101, 40129 Bologna, Italy. E-mail: milita@bo.imm.cnr.it

^c Istituto di Cristallografia @ Elettra Synchrotron, Consiglio Nazionale delle Ricerche, c/o Area Science Park, 34139 Basovizza, Trieste, Italy

^d Dipartimento di Fisica e Chimica, Università di Palermo, viale delle Scienze, Ed. 17, 90128 Palermo, Italy

^e Istituto di Cristallografia & To.Sca.Lab., Consiglio Nazionale delle Ricerche, via Valleggio 11, 22100 Como, Italy

† Electronic supplementary information (ESI) available: Details of the 2D-GIXD, structural and spectroscopic characterization. See DOI: 10.1039/d1tc00564b

difference in molecular packing and in material properties, as observed, for example, in many of the NDI-based species discussed in a previous paper.⁹ Indeed, once isolated as powders and films, alkyl chain derivatives show a remarkable structural flexibility, witnessed by a rich polymorphic behavior and reversible crystal-to-crystal and crystal-to-mesophase transformations, with detrimental implications for the applications/devices.¹⁰ On the other hand, while the performances of these materials are heavily dependent on the supramolecular assembly of the constituent molecules in the solid state (mostly in thin films),¹¹ predicting/devising the best performing geometry presently remains a challenging task. Stimulated by these considerations, we extended our interest to less flexible residues. Inspired by the earlier reports on amino-bearing residues in fullerenes, polyfluorenes and PDI derivatives,¹² where the presence of the amino group was not found prejudicial to materials properties, here we focus our attention on NDI-based solids bearing amino- (or hydrazido)-substituted benzenes, where the presence of aromatic rings is expected to block, to some extent, the mobility of the N-linked moieties, preventing “easy” polymorphic transformations under working conditions.

The choice of *aromatic* residues (here, in the form of amino-substituted phenyls) is also encouraged by the increased conjugation which can extend off the NDI core, enhancing the electron mobility. To the best of our knowledge, studies on pendant amino groups for NDI are scarce and mostly limited to their use as fluorescent probes¹³ and in cancer therapy.¹⁴ Although, admittedly, NH₂ groups are prone to oxidation and molecular degradation (more so in solution than in the solid phase), our studies show that the species presented here are stable in air up to (at least) 280 °C. Thus, amino-group reactivity seems not to be a worrisome issue.

One appealing, and so far underexplored, complementary aspect is the impact of the constitutional isomers considered in the present work on the NDI structural landscape, *i.e.* that of molecular entities possessing the same chemical formula but with a slightly different atom connectivity in their periphery. It is noteworthy that these NDI-materials sharing constitutional isomers are not the result of non-stereo-/regio-selective syntheses in which it is difficult to separate mixtures (as in the commercially available PDI-Br₂¹⁵ and N1400¹⁶ dyes); instead, by using predefined aminoaromatic residues, they are prepared as pure chemical entities with unique crystallographic phases.

While having the same chemical composition (as polymorphs do), crystal forms determined by the presence of *constitutional* isomers possess distinct physical and chemical properties from those one would expect in (conformationally different) polymorphic forms, with additional upstream control and greater stability. Similar considerations would apply as well to the case of *configurational* stereoisomers, pointing to the importance of understanding and addressing the selective formation of solid organic materials of isomeric species in organic electronics,¹⁷ as much as it is considered in pharmaceutical and pigment industries.¹⁸ Notable examples in these industrially relevant fields are the polymorphism of rubrene for efficient OLEDs,¹⁹ stereoisomerism of amphetamine²⁰ and, for what matters

here, the constitutionally different Pigments Red 194 and Orange 43 molecules.²¹ It is noteworthy that an additional degree of freedom, that is the occurrence of structurally induced *mesomeric* forms in organic semiconductors, has also been recently unveiled.²²

In the present work, the synthesis of a number of NDIs is presented, based on C₆H₆N₂ isomers (see Chart 1). Their rich structural landscape is unveiled and deeply discussed based on

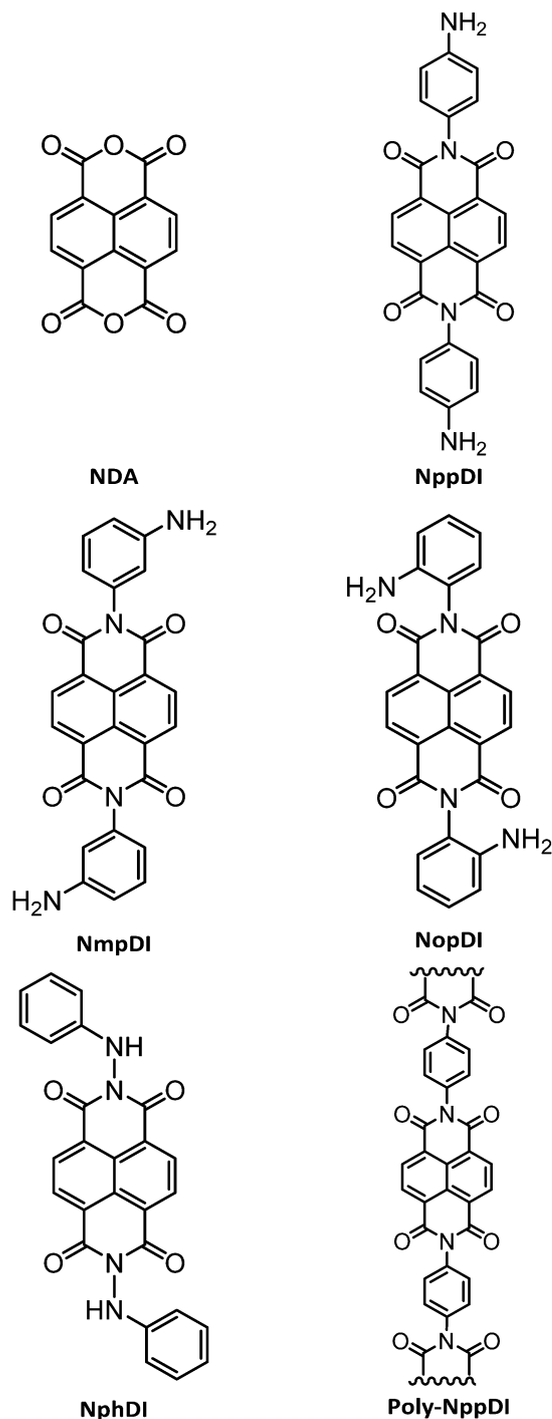


Chart 1 Molecular sketches and labeling of the species discussed in this work.

the combination of multiple structural characterization methods including X-ray powder diffraction, spectroscopic and thermal analyses, DFT calculations, and variable temperature diffraction methods. UV-Vis reflectance spectra on powders and grazing-incidence X-ray diffraction analysis of spin-coated films are presented with a general discussion of the properties and the possible processability routes of these materials.

2. Experimental section

(a) Materials

Reagents and solvents were purchased from Merck (*p*-phenylenediamine, >98%; *m*-phenylenediamine, 99%; *o*-phenylenediamine, >98%; phenylhydrazine, >97%; benzoic acid, >99%; methanol, 99.8%; dimethylformamide, 99.8%; toluene, 99.8%; dimethylacetamide, >99%; *m*-cresol, 99%), BLD (1,4,5,8-naphthalenetetracarboxylic acid monoanhydride, 98%) and Riedel-de-Haën (triethylamine, 99%) and used without further purification. All reactions were performed under a N₂ atmosphere.

(b) Syntheses of the materials

1,4,5,8-Naphthalenetetracarboxylic dianhydride (NDA). NDA was prepared using the synthetic method reported in ref. 23. Into a 500 mL three-necked flask, 25 g (0.087 mol) of 1,4,5,8-naphthalenetetracarboxylic acid monoanhydride, 50 mL of DMA and 100 mL of dry toluene were added under a N₂ atmosphere. The mixture was kept at 111 °C and stirred at this temperature for 2 h. Then, the flask was cooled to 10 °C, and stirred overnight. Filtration gave 23.5 g (0.087 mol, quantitative yield) of 1,4,5,8-naphthalenetetracarboxylic dianhydride as a yellow solid. ¹H-NMR (400 MHz, DMSO-d₆, 25 °C): δ 8.71 (bs, 4H). Calc. for C₁₄H₄O₆ (%): C, 62.7; H 1.5; Exp. C 62.5, H 1.8. *m/z* (ESI⁺): 299.0 (M + CH₃OH - H)⁻, calc. for C₁₅H₇O₇: 299.2.

1,4,5,8-Naphthalenetetracarboxylic *p*-aminophenylenedimide (NppDI). NppDI was prepared following the already published procedures.^{24,25} To a round bottom flask equipped with a stirrer and condenser, 1,4,5,8-naphthalenetetracarboxylic dianhydride (1 g, 3.73 mmol, 1 eq.), 1,4-phenylenediamine (4.033 g, 37.3 mmol, 10 eq.) and then 67 mL of anhydrous MeOH were added under a N₂ atmosphere. The reaction mixture was heated at the refluxing temperature for 20 h. The mixture was then cooled to rt, filtered and the brown solid was suspended in 50 mL of MeOH for 1 h. The suspension was filtered and dried under vacuum. NppDI was obtained as a brownish solid in 78% yield (1.310 g, 2.90 mmol). ¹H-NMR (400 MHz, DMSO-d₆, 25 °C): δ 8.69 (s, 4H), 7.02 (d, *J* 8.4 Hz, 4H), 6.67 (d, *J* 8.4 Hz, 4H), 5.30 (s, 4H); ¹³C-NMR (100 MHz, DMSO-d₆, 25 °C): δ 163.21, 148.77, 130.40, 129.09, 126.96, 126.56, 123.24, 113.68. Calc. for C₂₆H₁₆N₄O₄ (%): C, 69.6; H 3.6, N 12.5; Exp. C 68.9, H 3.8, N 12.1. *m/z* (ESI⁺): 447.3 (M - H)⁻, calc. for C₂₆H₁₅N₄O₄: 447.4.

1,4,5,8-Naphthalenetetracarboxylic *m*-aminophenylenedimide (NmpDI). To a round bottom flask equipped with a stirrer and condenser, 1,4,5,8-naphthalenetetracarboxylic dianhydride

(300 mg, 1.118 mmol, 1 eq.), 1,3-phenylenediamine (1.209 g, 11.18 mmol, 10 eq.) and 20 mL of anhydrous MeOH were added under a N₂ atmosphere. The reaction mixture was heated at the refluxing temperature for 20 h. The mixture was then cooled to rt and another 15 mL of MeOH were added leaving the mixture under stirring for an additional one hour. The suspension was filtered and NmpDI was obtained as a light brownish solid in 83% yield (417 mg, 0.929 mmol). ¹H-NMR (400 MHz, DMSO-d₆, 25 °C): δ 8.76 (s, 2H), 7.22 (t, *J* 7.6 Hz, 1H), 6.74 (d, *J* 8 Hz, 1H), 6.64 (s, 1H), 6.59 (d, *J* 7.6 Hz, 1H), 5.34 (s, 2H); ¹³C-NMR (100 MHz, DMSO-d₆, 25 °C): δ 162.77, 149.47, 136.14, 130.37, 129.16, 126.94, 126.65, 115.85, 114.17, 113.87. Calc. for C₂₆H₁₆N₄O₄ (%): C, 69.6; H 3.6, N 12.5; Exp. C 68.9, H 3.8, N 12.4. *m/z* (ESI⁺): 447.3 (M - H)⁻, calc. for C₂₆H₁₅N₄O₄: 447.4.

1,4,5,8-Naphthalenetetracarboxyl-*o*-aminophenylenedimide (NopDI). To a round bottom flask equipped with a stirrer and condenser, 1,4,5,8-naphthalenetetracarboxylic dianhydride (300 mg, 1.118 mmol, 1 eq.); 1,2-phenylenediamine (1.209 g, 11.18 mmol, 10 eq.) and 20 mL of anhydrous MeOH were added under a N₂ atmosphere. The reaction mixture was heated at the refluxing temperature for 16 h. The mixture was then cooled to rt and another 15 mL of MeOH were added, leaving the suspension under stirring for 1 h. The mixture was filtered and impure NopDI was obtained as a brownish solid (325 mg). The procedure was also repeated by changing the reaction time from 16 to 4 h but, also in this case, the formation of the by-products could not be avoided. Further shortening the reaction time to 1 h only led to incomplete NDA conversion, together with the formation of a mixture of the condensation products.

1,4,5,8-Naphthalenetetracarboxyl-phenylhydrazinedimide (NphDI). NphDI was prepared following an already published procedure.²⁶ Phenylhydrazine (2.199 mL, 22.371 mmol, 3 eq.) was transferred in a 250 mL round bottom flask, equipped with a stirrer and condenser, containing 1,4,5,8-naphthalene-tetracarboxylic dianhydride (2 g, 7.457 mmol, 1 eq.) in 100 mL of anhydrous DMF under a N₂ atmosphere. The reaction mixture was heated at 140 °C for 6 h and then cooled to rt. DMF was removed under vacuum and the reaction crude was dissolved in 100 mL of EtOAc and washed with 150 mL of water. The organic layer was washed with a brine solution (1 × 100 mL). EtOAc was then evaporated under vacuum and the residue was suspended in water, filtered, and washed with water, acetone and then with CH₂Cl₂. NphDI was obtained in 65% yield (2.18 g, 4.861 mmol) as a light brownish powder. ¹H-NMR (400 MHz, DMSO-d₆, 25 °C): δ 8.83 (s, 2H), 8.81 (s, 1H), 7.24 (m, 2H), 6.89 (m, 3H); ¹³C-NMR (100 MHz, DMSO-d₆, 25 °C): δ 162.16, 147.02, 131.03, 128.83, 126.98, 119.63, 112.62. Calc. for C₂₆H₁₆N₄O₄ (%): C, 69.6; H 3.6, N 12.5; Exp. C 69.4, H 4.2, N 11.9. *m/z* (ESI⁺): 447.3 (M - H)⁻, calc. for C₂₆H₁₅N₄O₄: 447.4.

Poly-1,4,5,8-naphthalenetetracarboxyl-*p*-phenylenedimide (poly-NppDI). *p*-Phenylenediamine (60.4 mg, 0.559 mmol, 1 eq.) was placed in a two-neck round bottom flask equipped with a stirrer and condenser under a N₂ atmosphere. Then, 2.5 mL of *m*-cresol, and subsequently triethylamine (TEA, 120 μL, 0.866 mmol, 1.55 eq., dropwise) were added. The reaction mixture was kept at 160 °C for 5 min and then 1,4,5,8-

naphthalenetetracarboxylic dianhydride (150 mg, 0.559 mmol, 1 eq.), benzoic acid (273.08 mg, 2.236 mmol, 4 eq.) and another 4.2 mL of *m*-cresol were added. The reaction was left under stirring at 175 °C for 15 h and then at 195 °C for an additional 3 hours. The mixture was then cooled to room temperature, filtered and washed with methanol, H₂O, acetone, and CH₂Cl₂ and dried in air, obtaining 176 mg of a yellow powder. Due to the very limited solubility in DMSO, very weak NMR signals were measured: ¹H-NMR (400 MHz, DMSO-d₆, 25 °C): δ 9.19 (s, 2H), 8.82 (s, 1H), 8.77 (s, 1H), 8.51 (s, 1H), 7.64 (d, 2H), 7.03 (m, 2H), 6.69 (s, 1H), 6.56 (m, 1H). Calc. for C₂₀H₈N₂O₄ (%): C, 70.6; H 2.4, N 8.2; Exp. C 66.3, H 2.9, N 6.9.

(c) Physico-chemical characterization

UV-Vis spectroscopy. The UV-Vis reflectance spectra of the powders were acquired in the 200–800 nm range, using a Shimadzu UV-2600 spectrophotometer. BaSO₄ was used as the reflectance standard and the measurements were performed on pelletized powders. The Kubelka–Munk function $F[R]$ was calculated using the reflectance spectrum $R(E)$, where E is the photon energy, using the $F[R] = (1 - R)^2/2R$ relationship. Taking $F[R]$ as a representative of the sample absorbance spectrum, extrapolation of the linear portion of the $(F[R]*E)^2$ vs. E on the E axis provided experimentally accessible optical direct band gap values.

NMR characterization. Solution ¹H and ¹³C (APT) NMR spectra were recorded in d⁶-DMSO at 400 and 100 MHz, respectively, using a Bruker Avance 400 spectrometer (Fig. S1–S4, ESI†). ¹H and ¹³C NMR data are reported as chemical shifts (in ppm, internal TMS standard), multiplicity (s = singlet, bs = broad singlet, d = doublet, t = triplet, and m = multiplet), coupling constants (in Hz) and integration. The poor solubility of our materials (see Table S1, ESI†) prevented NMR data collection in other organic solvents, and, similarly, limited the choice of experimental conditions for other solution studies, such as voltammetric measurements.

MS characterization. Mass spectra (Fig. S5–S8, ESI†) were recorded on a Thermo Scientific, Q-Exactive Plus System using 1 mg mL⁻¹ of centrifuged CH₃OH solutions.

Thermal analyses. Thermogravimetric (TG) and differential scanning calorimetric (DSC) traces were acquired from 30 to 900 °C (with a scan rate of 10 °C min⁻¹) using a STA 409 PC Luxx[®] analyser (Netzsch) under a nitrogen flow or in air and alumina sample holders equipped with a pierced lid (Fig. S9, ESI†).

Electrochemical characterization. The samples were dissolved in DMSO (typically 0.15 mg mL⁻¹) and 0.1 M NBut₄PF₆ was added as the electrolyte. Sweeps at 0.1 V s⁻¹ rate were performed in the –1.5 to +1.5 V interval. Ag/AgCl was used as the reference electrode, taken at 4.4 eV below the zero vacuum-level (Fig. S10, ESI†).

X-ray powder diffraction (PXRD). PXRD measurements were performed using a Bruker AXS D8 Advance diffractometer in Bragg–Brentano θ : θ geometry, equipped with a Lynxeye position sensitive detector. DS: 0.5°; generator setting: 40 kV, 40 mA; Ni-filtered Cu-K α radiation, λ = 1.5418 Å. PXRD data for

structure solution were collected in the 3–105° 2θ range for all samples (3–75° for poly-NppDI), sampling at 0.02°, with scan time lasting approximately 16 h.

Variable temperature PXRD analysis. Thermodiffractometric experiments were performed from 30 to 540 °C (580 °C for poly-NppDI). Powdered batches were deposited in the hollow space of an aluminium sample holder of a custom-made heating stage (Officina Elettrotecnica di Tenno, Ponte Arche, Italy). Diffractograms were acquired in air, in the most significant (low-angle) 3–30° 2θ range, under isothermal conditions in 20 or 30 °C steps. Since in powder diffraction experiments the samples are in direct contact with air and some thermal drifts/gradients are present, the accurate phase transition temperatures are calibrated using TG measurements.

(d) Preparation of thin films

The NDI derivatives were used as obtained from the washing and purification processes after their respective synthesis. For each compound, the solvent and the concentration of the solution were chosen close to the saturation limit. The least soluble NppDI derivative was dissolved in *N*-methyl-2-pyrrolidinone (NMP) and acetone (volume ratio 3:7) at a concentration of 1 mg mL⁻¹, while the NmpDI and NphDI solids were dissolved in *N,N*-dimethylformamide (DMF), with concentrations of 17 and 30 mg mL⁻¹, respectively. Thin films were prepared by spin-coating 20 μ L of solution onto 1 × 1 cm² silicon substrates covered with a 300 nm-thick thermal silicon oxide layer. Before spin coating, the SiO₂/Si substrates were cleaned sequentially for 5 minutes in acetone, ethanol, isopropyl alcohol and deionized water, and blow-dried with nitrogen gas. Finally, the substrates were etched by Ar/O₂ plasma for 15 minutes (flux ratio of 3:1), aiming at improving their surface wettability.

Though NMP and DMF are solvents not commonly used for the deposition of thin organic films for device fabrication (chloroform and chlorobenzene being much more widely employed), reports on thin film processing with less volatile solvents (surprisingly including also methanesulfonic acid) have appeared.²⁷ Additionally, the high thermal stability and the absence of polymorphic phases of these molecular solids, later discussed, suggest that sublimation processing of thin films and devices is a further viable option.

For each film, the spin coating conditions were tuned to optimize the film homogeneity and reproducibility. The NppDI solution was stirred and heated up to 170 °C for 48 h before being centrifuged at 10⁴ rpm for 5 minutes to separate the suspended powders from the solution. The NppDI films were spin-coated at 1000 rpm for 30 s followed by a treatment at 5500 rpm for 30 s. For variance, the NmpDI and NphDI solutions were heated up to ca. 130 °C for 10 minutes until a complete dissolution of the powders was achieved. The NmpDI films were spin-coated at 1500 rpm for 30 s, while the NphDI films were spin-coated at 1000 rpm for 30 s, followed by 2500 rpm for 30s. All films were left to dry at room temperature. Each sample has been investigated both as fabricated and after thermal annealing in air at 180 °C for 10 minutes.

(e) X-ray structural studies

Ab initio crystal structure solution from X-ray diffraction data. PXRD structure solution of the NppDI, NmpDI, NphDI and poly-NppDI phases was performed using the TOPAS-R²⁸ software. Standard peak search methods followed by profile fitting allowed the accurate estimation of the low-angle peak position. These values, through the SVD indexing algorithm,²⁹ provided primitive orthorhombic or monoclinic cells [$a = 26.12$, $b = 5.16$, $c = 7.83$ Å, $\beta = 101.4^\circ$, and $\text{GOF}(20) = 16.7$ for NppDI; $a = 31.96$, $b = 8.53$, $c = 7.44$ Å, and $\text{GOF}(23) = 19.9$ for NmpDI; $a = 20.94$, $b = 5.39$, $c = 9.28$ Å, $\beta = 96.1^\circ$, $\text{GOF}(29) = 32.1$ for NphDI; $a = 10.74$, $b = 12.64$, $c = 5.02$ Å, and $\text{GOF}(10) = 20.1$ for poly-NppDI]. Space group determination through the analysis of systematic absences after cell reduction indicated, for the four phases, $P2_1/c$, $Pbca$, $P2_1/n$ and $Pnmm$, respectively, later confirmed by successful structure solution and refinement. Density considerations enabled the determination of the molecular symmetry, the Z' value and the content of the asymmetric unit. Structure solution of the four molecular crystals was performed using the Monte Carlo/simulated annealing technique using a rigid model (flexible at the imide link) described by the Z-matrix formalism with standard geometrical parameters.

The paucity of the diffraction peaks in the poly-NppDI PXRD trace, their breadth and severe overlap, did not enable (semi)-automatic structure solution. A few competitive structural models were built through sensible stereochemical considerations and tested against the experimental pattern, eventually validating a set of collinear alternate copolymers of NDI/C₆H₄ fragments as the best model. The accuracy of the refined metrical and structural descriptors is less than ideal. Nevertheless, the material periodicity and its main stereochemical features are (reasonably) well addressed by the present model.

The final refinements were eventually carried out by the Rietveld method, maintaining the rigid bodies introduced at the structure solution stage (see Fig. 1 and Fig. S11, ESI†) The background was modelled by a polynomial function of the Chebyshev type and the peak profiles were described by the Fundamental Parameters Approach³⁰ and a common (refinable) isotropic thermal factor was attributed to all atoms. March-Dollase correction for preferred orientation³¹ was applied in the form of $g(hkl)$ and quoted below. Fractional atomic coordinates and crystal structure details were deposited with the CCDC (CSD Codes 2059769–2059772).† Specific sample-dependent issues are discussed in the Results section.

Crystal data for NppDI. C₂₆H₁₆N₄O₄, $f_w = 448.43$ g mol⁻¹, orthorhombic, $P2_1/c$, $a = 7.8185(4)$, $b = 5.1566(2)$, $c = 25.727(2)$ Å, $\beta = 95.918(3)^\circ$, $V = 1031.7(1)$ Å³, $Z = 2$, $\rho_{\text{calc}} = 1.444$ g cm⁻³, $\mu(\text{Cu-K}\alpha) = 8.2$ cm⁻¹, $g(010) = 1.16$; R_p and R_{wp} , 0.082 and 0.109 respectively, 6–105° 2θ range. $R_{\text{Bragg}} = 0.047$.

Crystal data for NmpDI. C₂₆H₁₆N₄O₄, $f_w = 448.43$ g mol⁻¹, orthorhombic, $Pbca$, $a = 32.012(9)$, $b = 8.5412(8)$, $c = 7.443(1)$ Å, $V = 2035.0(7)$ Å³, $Z = 4$, $\rho_{\text{calc}} = 1.464$ g cm⁻³, $\mu(\text{Cu-K}\alpha) = 8.4$ cm⁻¹, $g(100) = 1.22$; R_p and R_{wp} , 0.072 and 0.102 respectively, 5–105° 2θ range. $R_{\text{Bragg}} = 0.038$.

Crystal data for NphDI. C₂₆H₁₆N₄O₄, $f_w = 448.43$ g mol⁻¹, monoclinic, $P2_1/n$, $a = 20.956(2)$, $b = 5.3945(3)$, $c = 9.2901(6)$ Å,

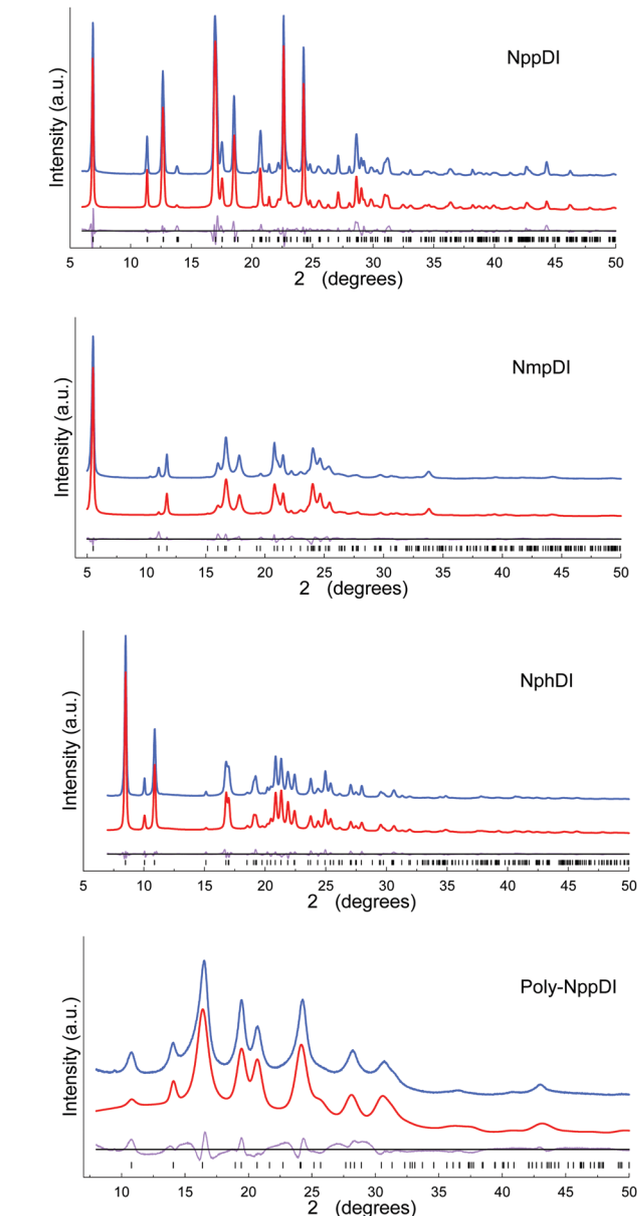


Fig. 1 Final Rietveld refinement plots, with difference plot and peak markers at the bottom. The observed and calculated curves (blue and red traces, respectively) have been vertically shifted for clarity.

$b = 96.132(5)^\circ$, $V = 1044.2(1)$ Å³, $Z = 2$, $\rho_{\text{calc}} = 1.426$ g cm⁻³, $\mu(\text{Cu-K}\alpha) = 8.2$ cm⁻¹, $g(100) = 1.13$; R_p and R_{wp} , 0.048 and 0.067 respectively, 7–105° 2θ range. $R_{\text{Bragg}} = 0.033$.

Crystal data for poly-NppDI. [C₂₀H₈N₂O₄]_n, $f_w = n \times 340.30$ g mol⁻¹ (n ca. 8), orthorhombic, $Pnmm$, $a = 10.82(2)$, $b = 5.04(1)$, $c = 12.58(2)$ Å, $V = 685(3)$ Å³, $Z = 2$, $\rho_{\text{calc}} = 1.65$ g cm⁻³, $\mu(\text{Cu-K}\alpha) = 9.8$ cm⁻¹, R_p and R_{wp} , 0.062 and 0.072 respectively, 8–70° 2θ range. $R_{\text{Bragg}} = 0.025$.

(f) Grazing incidence X-ray diffraction (GIXD) studies

GIXD measurements were performed at the XRD1 beamline of the ELETTRA synchrotron radiation facility in Trieste (Italy). $\lambda = 1.00$ Å; beam size: 200 × 200 μm². 2D-GIXD patterns were

collected by using a 2 M Pilatus silicon pixel X-ray detector (DECTRIS Ltd) positioned perpendicular to the incident beam, 350 mm away from the sample. Incidence angles of the X-ray beam with the film surface of $\alpha_i = 0.05^\circ$ and 0.1° were chosen to probe the uppermost film layers (the X-ray penetration depths are *ca.* 4 and 8 nm, for $\alpha_i = 0.05^\circ$ and 0.1° respectively), as well as $\alpha_i = 0.2^\circ$ to probe the full film thickness (penetration depth: *ca.* 30 μm). During the measurements, the samples were slowly rotated around the substrate normal (180° in 60 s), to average signals originating from laterally inhomogeneous film regions. Both as-prepared and annealed films were investigated to study the effect of thermal annealing on film crystallinity and mosaicity.

The visualization, analysis of the 2D-GIXD patterns and intensity profile extraction were performed using GIDVis software.³²

(g) Quantum-chemical calculations

Ground state geometry optimization and HOMO–LUMO energy calculations for NphDI, NmpDI and NppDI were performed with the Gaussian16 package,³³ using the nonlocal hybrid Becke three-parameter Lee–Yang–Parr (B3LYP) functional^{34,35} and the 6-31G(p,d) basis set. The experimentally determined crystal structure of each compound was used as a starting point for geometry optimization. The method and the chosen basis set were validated against the already published LUMO value of NphDI and HOMO/LUMO values of fluorinated analogues,³⁶ with absolute differences of less than 0.1 eV.

3. Results and discussion

(a) Synthesis

The syntheses of the NppDI^{24,25} and NphDI²³ molecules were already reported, though their characterization was limited to standard spectroscopic and (for NphDI) electrochemical methods. In the present work, we have adopted the synthetic approach of ref. 24 for what concerns NppDI, whereas some modifications of the procedure reported in ref. 26 were applied to NphDI, regarding changes in the work-up procedure (see details in the Experimental section). In particular, instead of dissolving the extracted crude material in THF and subsequently adding water to crystallize the product at 4°C overnight, the pure product was obtained upon washing the extracted reaction crude material with water, acetone and then CH_2Cl_2 , thus improving the product yield from 43%²⁶ to 65%.

The syntheses of the NDI cores branched with *m*- and *o*-substituted diaminobenzenes (NmpDI and NopDI), not present in the scientific literature, were performed in strict similarity with the protocol used for the *p*-regioisomer. NmpDI was easily recovered as a pure solid in high yields (>80%). The synthesis of the NopDI molecule, attempted in several ways, mostly modifying temperature solvents and reaction times, proved to be very difficult. Indeed, two ring-closure adducts, known as Pigment Orange 43³⁷ and Pigment Red 194,³⁸ were systematically detected

in the reaction mixture (see Chart S1, ESI†). Not being possible to isolate a single (pure) crystal phase of NopDI, its full structural and analytical characterization was abandoned. It is worth noting that such *o*-aminophenylene condensation has been used in the synthesis of BBL, or poly(benzimidazobenzophenanthroline), a soluble polymer somewhat reminiscent of both NopDI and poly-NppDI, which has recently attracted the attention of many for its excellent performances in n-channel OFETs.^{27c}

Finally, aiming at enhancing the thermal stability and the conductive properties of this class of materials, the polymerization capability of the NppDI molecule was explored through the synthesis of a polymeric species, poly-NppDI. A one-pot reaction using a 1:0.8 NDA:*p*-diamino-benzene molar ratio proved to be unsuccessful (a complex mixture of products was obtained). Co-polymerization with NDA was also tried by reacting NDA and NppDI monomers overnight in 1:0.8 ratio in refluxing toluene, but this attempt also failed.

However, taking inspiration from polymerization reactions of NDA with a series of aromatic diamines,³⁹ suitable conditions for the obtention of the sought copolymer were unearthed. Employing *m*-cresol as the solvent (a liquid with a high boiling point and solvent power) and with the crucial usage of two additives (excess benzoic acid – 4 eq. – and TEA – 1.5 eq.), poly-NppDI was synthesized with a 93% yield. The role of these additives can be explained by benzoic acid coordination to the NDA oxygen atoms, making carbonyl groups more susceptible to the nucleophilic attack, and by TEA-induced avoidance of proton transfer from the benzoic acid to *p*-diaminobenzene. Quantitative yields were obtained when employing 1 eq. of *p*-diaminobenzene as the unique nucleophile and heating at 175°C for 15 h and then at 195°C for an additional 3 hours. poly-NppDI appeared to be a nanosized material by X-ray diffraction analysis. Its structural analysis (later discussed) confirmed the formation of the sought material, though the average chain length is limited to a few (alternating) NDA/*p*-diaminobenzene monomers.

(b) Crystal structures

The crystal structures of the NppDI, NmpDI and NphDI molecular phases contain symmetrically disubstituted NDI cores, lying about inversion centers in monoclinic or orthorhombic space groups, with $Z' = 0.5$. Their molecular structures and crystal packings are shown in Fig. 2, and the relevant stereochemical features are shown in Table 1, where the structural analogues 1,4,5,8-naphthalenetetracarboxyl-*p*- CH_3 -benzylidimide (NhbdI) and *p*- CF_3 -benzylidimide (NfbDI) are also included.⁴⁰ As detailed in the Experimental Section, the structural models presented here make use of rigid body descriptions of the stiff portions of the molecules. Accordingly, reliable structural information is mostly confined in the conformational freedom (the orientation of the branching residuals), in the intermolecular effects (the stacking periodicity) and in the degree of overlap of parallel, adjacent, NDI cores. With reference to the stacking periodicity, while for aliphatic substituents the π – π stacking distances were typically around 3.4 Å or lower⁹ (very similar to the interlayer interaction in graphite, 3.35 Å),⁴¹

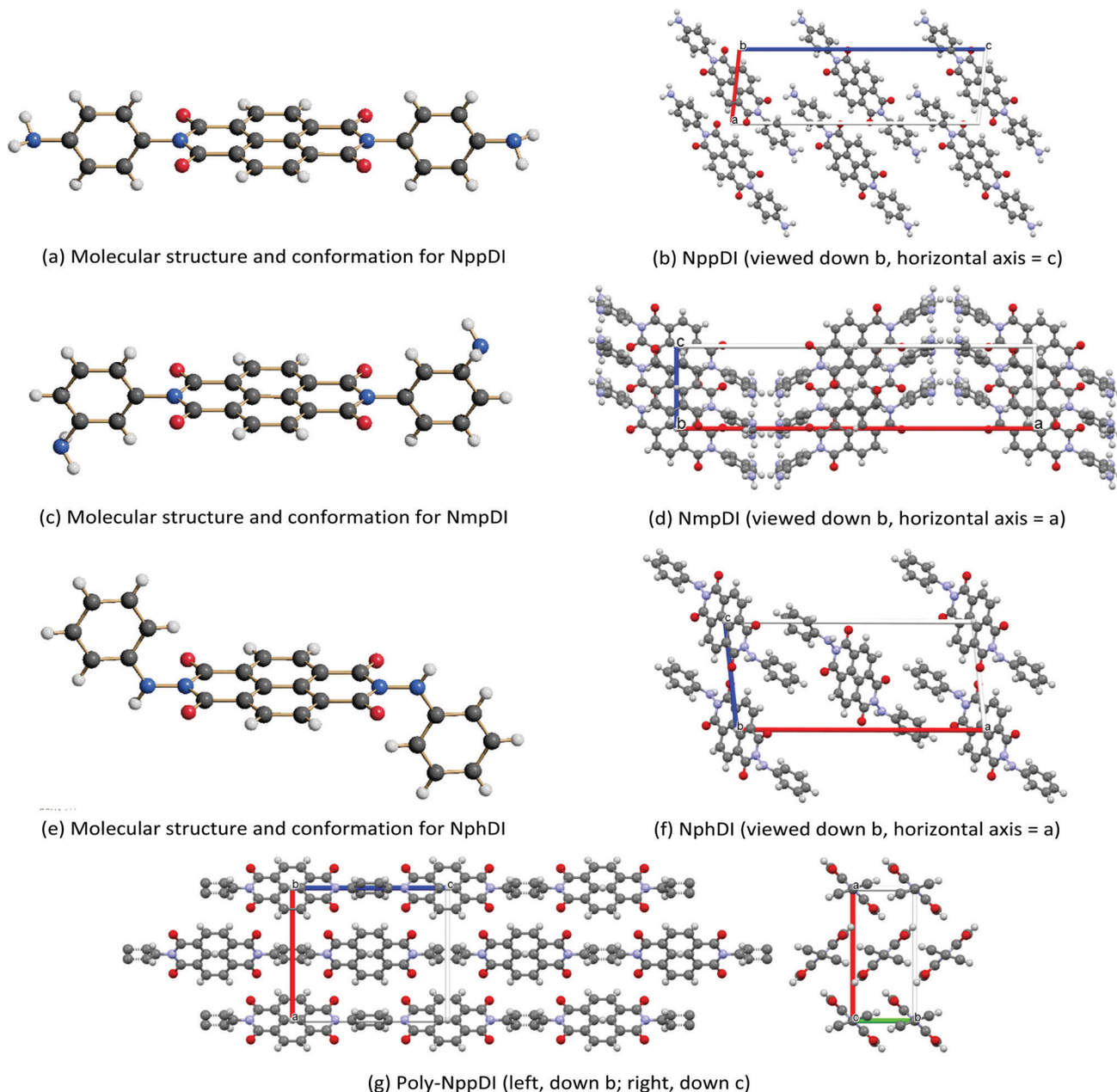


Fig. 2 Schematic drawings of the molecular structure and crystal packings of (a and b) NppDI, (c and d) NmpDI and (e and f) NphDI, together with a note of their orientations. In (g), the polymeric poly-NppDI structure is shown. Color codes: C in grey, N in blue, O in red and H in white. Graphics realized by Schakal⁴² (a, c and e) and Mercury⁴³ (b, d, f and g).

in the present case this value increases to 3.5 Å or above for NppDI monomeric and polymeric derivatives, whereas parallel stacking does not occur in the monoclinic crystal structure of NmpDI. In the latter case, the aromatic cores, stretching out in the approximate [102] direction, stack in herringbone geometry, tilted away one from the other by *ca.* 48° (see Fig. S12, ESI†). Further comparative geometrical descriptors of NppDI *vs.* NmpDI are the tilt angles of the aromatic cores *vs.* the lamellar stacking (100) [63.2° *vs.* 29.7°] and the thickness of the molecular slabs [<11.5 *vs.* 16.0 Å], witnessing the more sprawled disposition of the NDI cores in NppDI. Only the

phenylhydrazine derivative (NphDI) shows a short(er) NDI-core interaction.

As per the relative slip of the aromatic core, for all crystal phases studied here, the stereochemical descriptors (which in aliphatic systems and in NhbdI and NfbDI as well, have found to cluster into two specific portions of the χ , ψ plot)⁹ are more widely spread here (see Fig. 3). These findings indicate that the most relevant interactions driving the formation of the stacks are not anymore assignable to the NDI cores; instead, they must be attributed to the interlocking of the rigid (aromatic) branches. The geometrical degree of overlap (not the orbital

Table 1 Relevant geometrical features for the crystal phases studied in this work. τ is the NDI-R torsional angle, d is the interplanar distance of parallelly stacked NDI-cores, χ and ψ are the stereochemical descriptors defined in ref. 9, and H-bonds are the shortest intermolecular H-bond contacts. E_{gap} corresponds to the optical band gap energies derived as described in the Experimental section

Species	τ , °	d , Å	χ , °	ψ , °	H-Bonds, Å	DoO (eqn (1))	E_{gap} , eV
NppDI	82.6	3.63	46.7	55.9	3.11 (N··N) 3.22 (N··O)	0.46	2.87
NmpDI	78.4	no π - π	—	—	None	—	3.01
NphDI	84.7	3.38	49.2	64.9	3.06 (N··N)	0.63	2.94
NhbDI ⁴⁰	80.0	3.54	67.1	46.2	None	0.40	n.a.
NfbDI ⁴⁰	78.6	3.40	73.7	39.8	None	0.58	n.a.
Poly-NppDI	63.0	3.50	90.0	44.0	None	0.69	2.90
Aliphatic NDI	—	3.27–3.42	68.4–80.5	42.2–53.4	—	0.62–0.72	

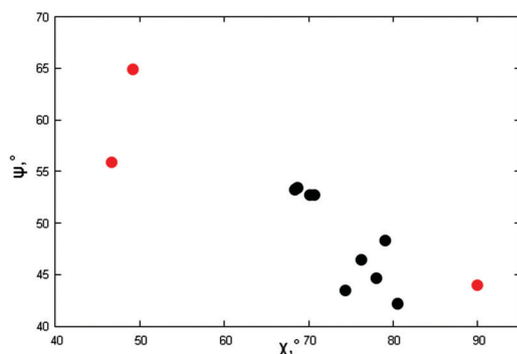


Fig. 3 Stereochemical descriptors χ, ψ for the π - π stacking of the crystal phases of this work (red dots, values from Table 2) and the aliphatic analogues (black dots) reported in ref. 9.

one) defined by:

$$\text{DoO} = \sqrt{1 - \cos^2\chi - \cos^2\psi} \quad (1)$$

measures the complete (DoO = 1.0) or partial ($0 < \text{DoO} < 1.0$) superimposition of neighboring NDI cores along the direction orthogonal to the molecular plane. The values in the range 0.63–0.72 are typical for aliphatic substitutions and are also found for the molecular NphDI and polymeric poly-NppDI crystals. In contrast, in the molecular NppDI and NhbDI species (DoO < 0.46), the NDI cores overlap to a minimal extent; these compounds must be considered definite outliers in the set of symmetrically substituted aliphatic and aromatic NDIs.

Two of these materials, NmpDI and poly-NppDI, require a special set of comments. NmpDI crystallizes forming extended slabs running in the bc plane (thus normal to a), weakly bound by hydrogen bonds of the N-H··N type. Given that the m -positioned $-\text{NH}_2$ residue was found to be crystallographically disordered in two positions roughly with a 1 : 3 ratio, this brings about a lack of ideal periodicity and the occurrence of a (possibly conditional) stereochemical local preference, with localized defects of the 2D type. This hypothesis is also partially confirmed by the less-than-ideal Rietveld fit shown in Fig. 1, possibly manifesting the slight misorientation of each molecule in the crystal from the average one, *i.e.* smoothing out the structural misfits.

Finally, poly-NppDI is a rare example of a crystalline polymer of NDI-based materials. Its extreme insolubility probably limits

the growth of microstructured materials, and very broad peaks are observed in the PXRD trace. Thanks to the knowledge of the stereochemical feature of the NppDI molecular counterpart, it was possible to define a sensible structural model using symmetry restraints and geometrical considerations, later refined against the PXRD data using the standard Rietveld technique.

Our result revealed the occurrence of *ca.* 10 nm chains of elliptical section (suggesting a sequence of 8 NDI/1,4-phenylenes along the c axis and an average molecular weight of 2700 Dalton), packed in the crystal in a herringbone motif and a pseudo-hexagonal lattice (Fig. 2d), reminiscent of smectic E liquid crystalline phases. The fact that these chains are not extremely long is also witnessed by the less-than-ideal elemental analysis values reported in the Experimental section, where the C/N molar ratio is slightly above 11 (and should be 10 in an infinite chain), as if the chains were terminated by NDA, and not by p -phenylenediamine residues.

(c) Thermal stability

The thermal stabilities of the di- and poly-imide species were studied using the TG plots (Fig. 4) and the structural variations occurring upon heating were followed by variable-temperature X-ray experiments (Fig. 5). For NppDI, NmpDI, NphDI and poly-NppDI bulk samples, isolated in powdered form, the PXRD patterns were acquired from room temperature up to the decomposition temperature, measured by TG to occur near 300 °C for NphDI and at much higher temperatures for the other materials. Comparing TG and PXRD data, no

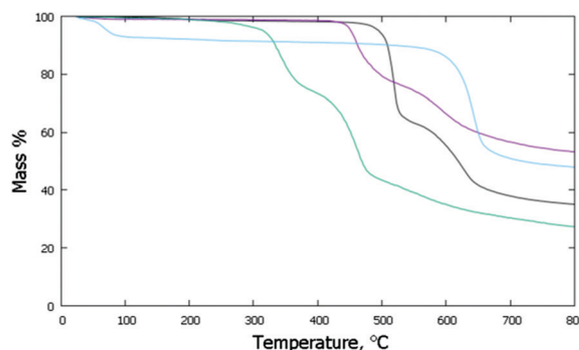


Fig. 4 TG traces collected under N_2 at a scan rate of $10^\circ \text{ min}^{-1}$: NppDI in black; NmpDI in magenta; NphDI in green; poly-NppDI in blue.

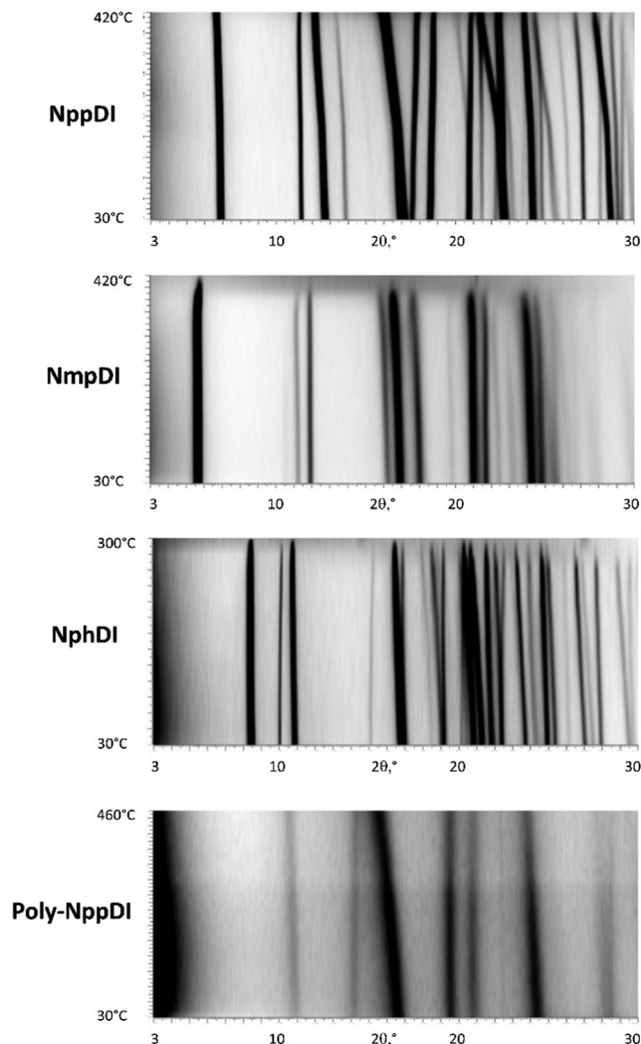


Fig. 5 Variable temperature XRD scans (2θ in the 3–30° range), from RT (bottom) to 420, 420, 300, and 460 °C, respectively.

polymorphic changes upon heating were observed. This is in striking difference with many aliphatic congeners, where the occurrence of several crystalline and liquid crystalline phases was the rule more than the exception.⁹

The observed mass variations are not easy to interpret. In all three molecular compounds a double decomposition step is observed, accounting for a total of 52 to 61 w% loss. These values are in line with the elimination of the imidic branches

and partial (and differential) decomposition (to CO or CO₂) of the carbonyl residues. Of much simpler description is the TG curve of the polymeric poly-NppDI phase: the observed 41.2 w% loss is easily reproduced by the elimination of two C₂O₂N fragments (40.9%), leaving behind only condensed polyaromatic hydrocarbons.

The coefficients of thermal expansion were determined using the cell parameter variations derived from the structure-less Le Bail fitting using the linear approximation:

$$p(T) = p(T_0)[1 + \kappa_p(T - T_0)] \quad (2)$$

where $p = a, b, c, (\beta)$ or V ; T and T_0 are the actual and initial temperatures and κ_p is the pertinent linear (or volumetric) thermal expansion coefficient. The entire set of measured datapoints could be reproduced by the mathematically equivalent $\Delta p/p_{RT} = \kappa_p \Delta T$; and the resulting κ_p values and their uncertainties are shown in Table 2.

The κ_p values, spatially visualized in Fig. 6, are highly anisotropic, in line with the crystal symmetries of these materials. This is particularly true for poly-NppDI, where two nearly null coefficients are observed along the **b** and **c** axes, at the expenses of a very large value measured along **a**.

Additionally, the very high decomposition temperature of poly-NppDI powders (onset near 615 °C) closely matches that of films of the 2,3,6,7 regioisomer⁴⁴ (measured by T_d^5 , the 5% weight loss temperature – 590 °C in N₂ and 580 °C in air), confirming the extreme thermal stability of aromatic polyimide (co)polymers.⁴⁵ The oxidative stability of all species was also established by performing TG/DSC measurements in air (see Fig. S9, ESI†).

(d) Spectroscopic analysis

The powdered materials appear light brown in color and, as expected, show “absorbance” spectra, spanning the wide 400–600 nm interval (Fig. 7). Such behavior is in line with other NDI-based diimides, though a wider band is observed here. Since most of the studied species (NphDI excluded, for the presence of an sp³ hybridized N atom) may manifest extended charge delocalization onto the aromatic branching residues, the occurrence of several accessible states with low-energy separation is in agreement with the observed broad absorption band falling in the visible region.

The steep increase observed in proximity to 400 nm, on the verge of the violet-UV range, has been interpreted by the

Table 2 TG inflection points and the corresponding mass loss, and the linear and volumetric thermal expansion coefficients (10⁻⁶ K⁻¹), calculated by linear regression of the PXRD data acquired in the 30–300 °C range

Species	TG inflection, °	Mass loss, %	κ_a	κ_b	κ_c	κ_β	κ_V	DLD ^a
NppDI	520.0 (624.2 ^b)	35.2 (60.9 ^b)	51(4)	-34(1)	51(2)	79(5)	88(7)	0.0126
NmpDI	460.3 (594.2 ^b)	22.0 (51.9 ^b)	37(3)	19(1)	55(2)	—	112(3)	0.0068
NphDI	334.8 (464.9 ^b)	25.4 (57.4 ^b)	26(2)	104(4)	11(2)	-46(1)	150(2)	0.0123
Poly-NppDI	643.0	41.2	109(3)	-4(3)	-1(3)	—	104(6)	0.0113

^a Note: DLD is the degree of lattice distortion, or the spontaneous strain, that is the square root of the sum of squared eigenvalues of strain tensor calculated for the 30–300° range, divided by 3. ^b Second inflection point and the total mass change after the second event.

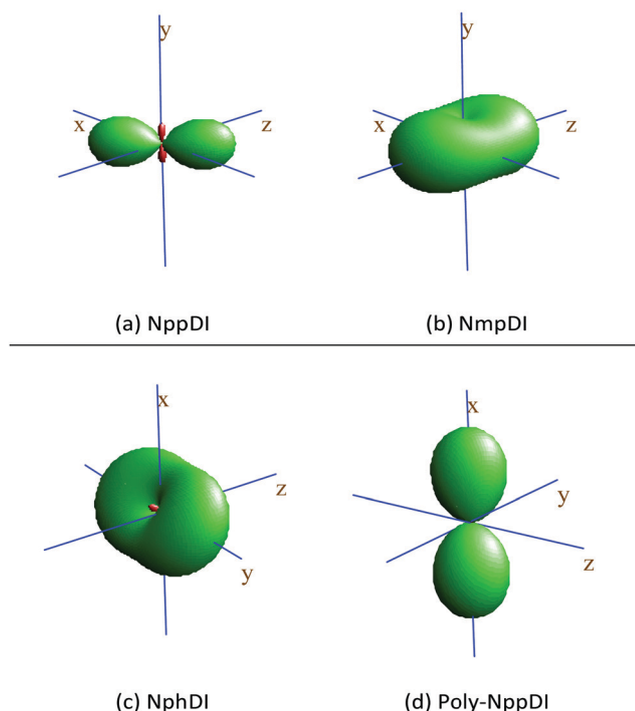


Fig. 6 Visualization of the thermal expansion tensors, calculated by comparing the lattice metrics at RT and at 300 °C. Positive lobes in green, (tiny) negative ones in red.

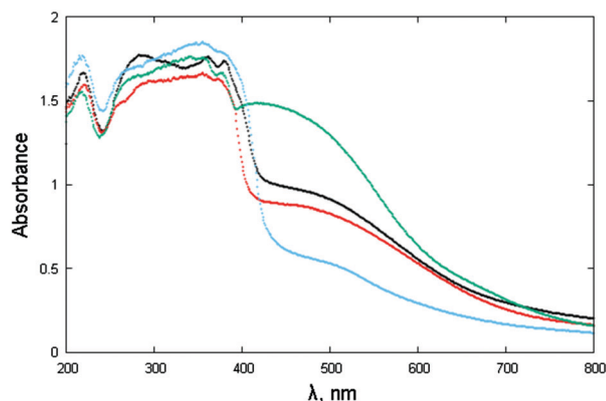


Fig. 7 Absorbance spectra (in the form of $F[R(E)]$ through Kubelka–Munk transformation of the diffuse reflectance spectra) of NppDI (black), NmpDI (red), NphDI (green) and poly-NppDI (blue).

occurrence of a direct bandgap, the actual values of which, falling in the narrow 2.87–3.01 eV range and derived as illustrated in Fig. S13 and S14 (ESI[†]), are reported in Table 1. The lowest values (2.87 and 2.90 eV) are found for the *p*-diaminophenylene derivatives, the molecular (NppDI) and polymeric (poly-NppDI) ones, respectively. Such observations, which cannot be explained by the crystal packing features discussed above and quantified in Table 1, speak for a slightly higher extent of conjugated bond overlap in these molecules. Slightly larger direct band gap energies are found for NmpDI

(not manifesting π - π stacking in the solid) and NphDI (where significant bond conjugation through the N–N bond can easily be excluded). As reference values, for markedly unconjugated residues, the typical optical bandgap falls near 3.0 eV.⁴⁶

(e) Electronic structure

As reported in Fig. 8, the HOMO and LUMO, derived from DFT calculations, are mostly localized on the side aromatic rings and on the naphthalene-diimide aromatic core, respectively. In NppDI and NmpDI, the HOMO levels are aligned (at *ca.* –5.6 eV); since they differ only for the (remote enough) position of the amino group on the side rings, this has negligible influence on the HOMO energy. Since the polyaromatic core is the same in both compounds (see the optimized geometries in Fig. S15 (ESI[†]), coordinates are provided in Tables S2–S4, ESI[†]), the energies of the LUMO differ by less than 0.1 eV (–3.15 and –3.23 eV). This finding can likely be ascribed to the difference in dihedral angles (reported in Table S5, ESI[†]), affecting a partial, though minor, electronic conjugation. On the other hand, in NphDI, the energies of both HOMO/LUMO are lowered (–5.95/–3.63 eV, in line with those reported in ref. 26) and favor $1e^-$ reduction to the corresponding radical anion, useful for sensing and for the generation of *n*-type organic semiconductors. These results suggest that the HOMO/LUMO energy is sensitive to nitrogen substituents only when they are directly linked to the core rings. Experimentally measured optical band gaps (measured in the solid) deviate by *ca.* 0.4–0.6 eV from the DFT-calculated molecular ones (known to vary depending on the basis-set used⁴⁷), an effect which was already observed in a large variety of organic semiconductors,⁴⁸ and also in strictly related NDI-based molecules and congeners (up to *ca.* 0.7 eV).⁴⁹

To experimentally corroborate our DFT simulations, cyclic voltammetric measurements of NppDI, NmpDI and NphDI (in DMSO) were performed (see Fig. S10, ESI[†]) and the data

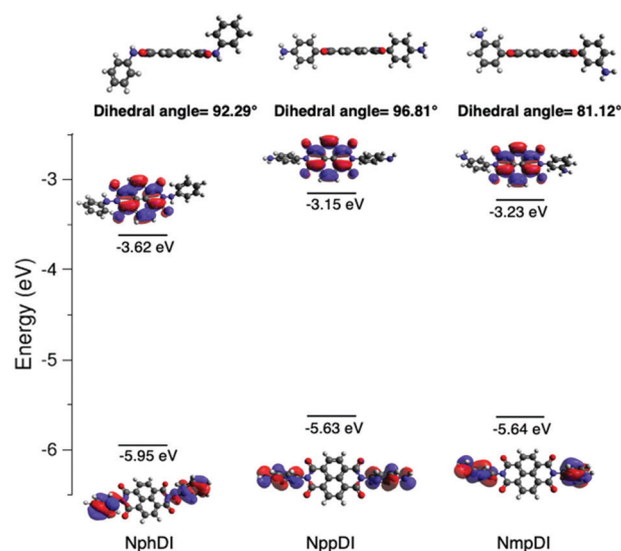


Fig. 8 Energy scheme and the HOMO and LUMO isosurfaces for NphDI, NppDI, and NmpDI (left to right).

were analyzed following the procedure used for similar compounds (*n*-dopable conjugated NDI-based polymers).⁵⁰ Similarly to these polymers, our molecular compounds show two separate reduction steps (*ca.* 0.5 eV apart, see Table S6, ESI[†]), but with slightly lower electrochemical band gaps ($\Delta E_{g_{el}}$, *ca.* 1.4 eV vs. *ca.* 2.0 eV in ref. 50). Such lowering is mostly attributed to the changes of the absolute energy values of the HOMOs, while those of the LUMOs are still located near -4 eV. Thus, LUMO-dependent reduction or partial *n*-doping is a favored process also in our molecular species synthesis.

(f) Thin film analysis

For the three molecular NDI derivatives, both the as-deposited and thermally annealed thin films have been investigated. The thin film structures, in terms of crystal phase, degree of crystallinity and texture (mosaicity) have been determined by the analysis of the 2D-GIXD measurements.⁵¹ As anticipated, these images have been collected just below and slightly above the critical angle for total reflection, and their mutual comparison enables the retrieval of information on the variation of the film structure along its thickness.⁵²

Due to the extremely poor solubility of NppDI (providing approximately $10\times$ more dilute solutions than NmpDI and NphDI), the as-deposited (too thin) films did not provide measurable diffracted signals, and only few and weak signals appeared in the 2D-GIXD images after annealing (see Fig. S16, ESI[†]). Although the paucity of information does not allow a complete description of the NppDI film structure, the observed signals can be ascribed to the strongest reflections of the bulk phase, suggesting that the

molecular arrangement in the film remains unchanged and that no distinct thin-film structure is formed.

Fig. 9 shows the 2D-GIXD images for the annealed NmpDI (a) and NphDI (b) thin films, together with the corresponding integrated intensity profiles along the q_z (Fig. 9b and e) and q_{xy} (Fig. 9c and f) directions. The 2D-GIXRD images of both films exhibit several intense Bragg spots having a perfect symmetry with respect to the central vertical line, overlapped with minor arc-shaped intensity contributions, the latter being more extended and pronounced for the NmpDI film than for NphDI (Fig. 9a and d). These features indicate the coexistence of a majority of crystallites/domains highly oriented along one specific direction with respect to the substrate surface (sharper $h00$ signals), and a minority of less, or even randomly, oriented crystallites (arc-shaped signals). For both films, all the reflections have been indexed by using the monoclinic crystallographic structure of the RT-stable bulk phases and by assuming the (100) texturing (Fig. S17 and S18a, ESI[†]), as confirmed by the q_z integrated intensity profiles (Fig. 9b and e), where only $h00$ reflections are observed. The molecular orientations associated with this texturing are schematically reported in Fig. 9a and b. Interestingly, though sharing the same texturing, dictated by the stacking of molecular layers (arranged in slabs), the molecular contacts on the substrate surface distinctly differ with the angle between the aromatic cores and the surface being 83.7° and 40.8° for NmpDI and NphDI, respectively. However, in addition to peaks of the $h00$ class, the q_{xy} integrated intensity profiles (Fig. 9c and f) show the presence of further peaks, witnessing the presence of largely misoriented crystallites.

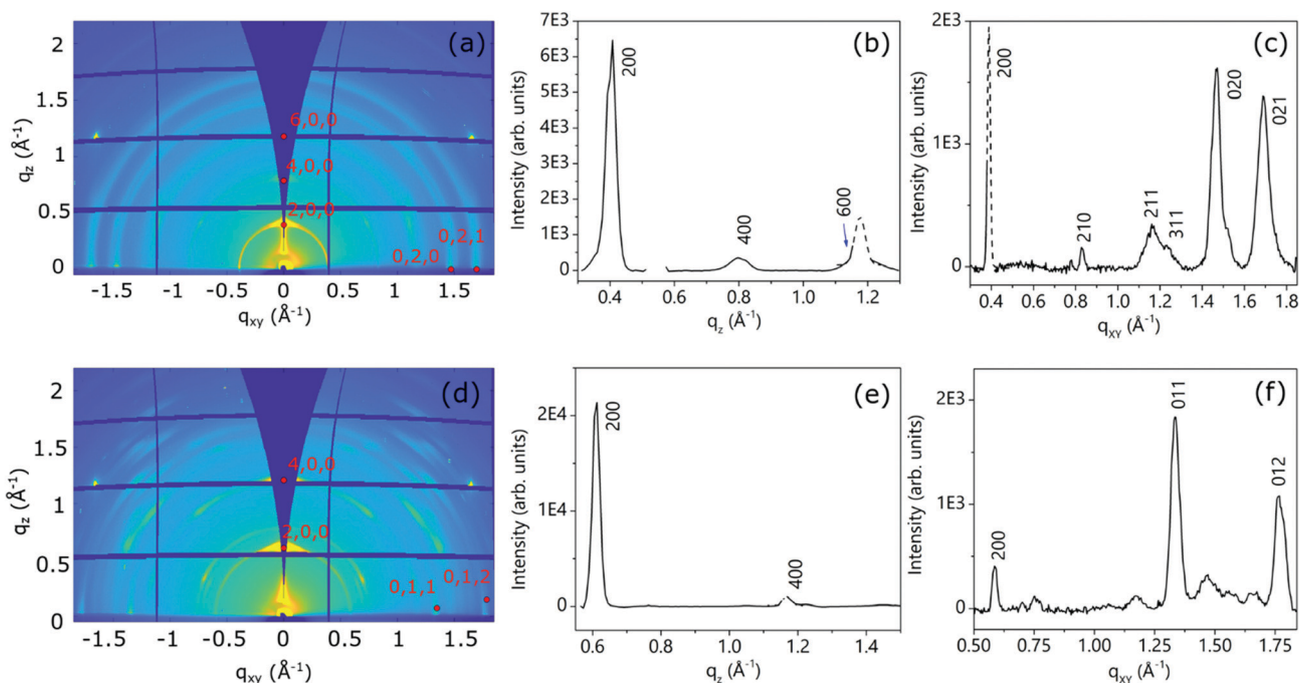


Fig. 9 2D-GIXD images of annealed (a) NmpDI and (d) NphDI thin films recorded for $\alpha_i = 0.1^\circ$. The corresponding integrated intensity profiles along q_z , [(b) and (e)], and along q_{xy} , [(c) and (f)] are reported. The dashed lines in b, c and e are Gaussian fits to the experimental data at the positions where the detector cuts hamper collection of the diffracted intensity.

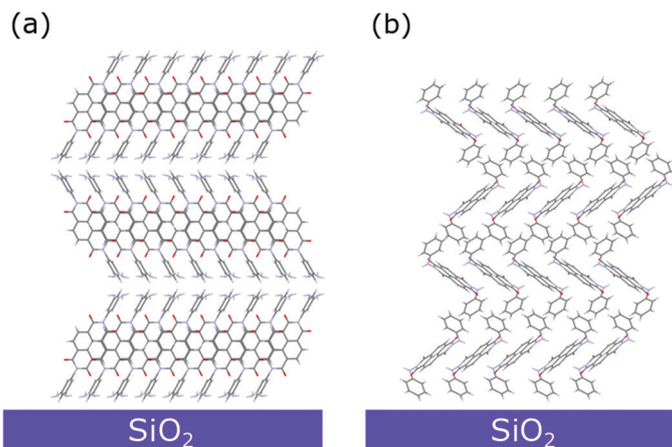


Fig. 10 Sketch of the dominant molecular arrangement of the (a) NmpDI and (b) NphDI thin films on the substrate surface, as retrieved from GIXD analysis.

A closer look into the 2D-GIXD images of the NphDI film shows that all reflections are azimuthally misplaced with respect to those expected for an exact $h00$ texturing (Figure S18a, ESI[†]), as already observed in the 2D GIXD images of other highly ordered organic films⁵³ (Fig. 10).

However, the experimental positions of the reflection maxima can be properly reproduced by tilting by *ca.* 3° the (100) crystal planes with respect to the substrate surface (Fig. S18b, ESI[†]). This tilting only marginally affects the angle between the aromatic cores and the substrate surface (lowered to *ca.* 40.1°).

The mosaicity of the oriented crystallites/domains, *i.e.* their angular deviation with respect to the main orientation, is estimated from the analysis of the experimental azimuthal profile of the strong 200 reflection (see Fig. S19, ESI[†]). The very small value of the determined mosaicity, *ca.* $\pm 2^\circ$ and $\pm 3.5^\circ$ for NmpDI and NphDI, respectively, is a direct measure of the strong orientation of the domains (*i.e.*, of the associated molecules).

The 2D GIXD patterns collected below and above the critical angle for total reflection (Fig. S20 and S21, ESI[†] for NmpDI and NphDI, respectively), probing only the first 4 nm close to the air interface or the entire film thickness, allowed us to establish the homogeneity of the molecular arrangement across the film thickness. Regardless of the probed depth, the films show the same texturing, as evidenced by consistence of the extracted q_z profiles, and the same degree of mosaicity, given by the almost invariance of the width of azimuthal profiles. Interestingly, the as-deposited films exhibit a high degree of crystallinity (slightly lowered upon annealing), indicating the extreme propensity of these molecules to self-assemble in an ordered fashion on the substrate surface without the need for any postprocessing technique (Fig. S22 and S23, ESI[†]).

4. Conclusions

In this paper we presented the isolation and the thorough characterization of powders and thin films of new molecular

crystals containing the NDI cores, substituted by different isomeric, constitutionally different, N-containing aromatic residues. The complete structural analysis demonstrated the similarity of the bulk and film structures, and the morphological texturing of the latter. In search for NDI-based organic solids not showing the polymorphic variability of the alkyl-chains analogues, in the present work we report species which exhibit high thermal stability and absence of polymorphic transitions up to melting. Both properties make the sublimation-route as a viable option for films and device preparation, in addition to solution-based methods, thus compensating the limited solubility properties of our systems. The structural, spectroscopic and thermal analyses presented in this work, together with the determination of the anisotropy of thermal expansion coefficients, add further knowledge on how NDI cores approach, and overlap, in the search for optimized performances (*e.g.*, electron and thermal conductivity, and Seebeck coefficient), which are at the heart of thermoelectric generators based on aromatic-polyimides.⁵⁴

DFT calculations indicated a significant lowering of the frontier orbital energies of the hydrazido derivative (NphDI), fostering its use as an electronic acceptor. Such an effect can be further, and easily, enhanced by adding electron-withdrawing groups (such F or CF_3 residues) on the aromatic rings in the molecular periphery. Work can be anticipated in the direction of exploring the synthesis and structure of NphDI fluorinated congeners, in search for soluble, filmable and dopable systems with improved electronic performances, electron mobility and rich(er) carrier concentrations. Indeed, the preliminary results indicate that polyfluorinated NDIs possess significantly enhanced solubilities in organic solvents commonly used for film deposition (acetonitrile, DMF and THF – but not chlorinated ones) and energy levels and redox properties suitable for *n*-doping.

Finally, we have also prepared and characterized an organic nanosized material, stable well above 500°C and containing a 100% ordered 1:1 copolymer (of admittedly short chain length), together with its structure determination from the severely broadened XRD trace.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

This project was partially supported by MIUR (PRIN-2017, Project 2017L8WW48, HY-TEC). We thank G. Chita for assistance during the synchrotron X-measurements at Elettra. The courtesy of G.B. Giovenzana (Università del Piemonte Orientale) for providing MS data is also acknowledged.

Notes and references

- W. Brütting and C. Adachi, *Physics of Organic Semiconductors*, Wiley, NY (USA), 2012.
- (a) Y. Chen, Y. Zhao and Z. Liang, *Energy Environ. Sci.*, 2015, **8**, 401–422; (b) M. Culebras, C. M. Gómez and A. Cantarero, *Materials*, 2014, **7**, 6701–6732; (c) L. M. Cowen, J. Atoyo, M. J. Carnie, D. Baran and B. C. Schroeder, *ECS J. Solid State Sci. Technol.*, 2017, **6**, N3080–N3088.
- M. M. Hussain and N. El-Atab, *Handbook of Flexible and Stretchable Electronics*, CRC Press, Boca Raton, FL (USA), 2019.
- M. Caironi and Y.-Y. Noh, *Large Area and Flexible Electronics*, Wiley VCH, Weinheim (Germany), 2015.
- L. A. Francis and K. Iniewski, *Novel Advances in Microsystems Technologies and Their Applications*, CRC Press, Boca Raton, FL (USA), 2014.
- M. Al Kobaisi, S. V. Bhosale, K. Latam, A. M. Raynor and S. V. Bhosale, *Chem. Rev.*, 2016, **116**, 1685–11796.
- C. Li and H. Wonneberger, *Adv. Mater.*, 2012, **24**, 613–636.
- (a) H. Dong, C. Wang and W. Hu, *Chem. Commun.*, 2010, **46**, 5211–5222; (b) A. Nowak-Król, K. Shoyama, M. Stolte and F. Würthner, *Chem. Commun.*, 2018, **54**, 13763–13772.
- S. Milita, F. Liscio, L. Cowen, M. Cavallini, B. A. Drain, T. Degousée, S. Luong, O. Fenwick, A. Guagliardi, B. C. Schroeder and N. Masciocchi, *J. Mater. Chem. C*, 2020, **8**, 3097–3112.
- (a) M. Dharmarwardana, R. P. Welch, S. Kwon, V. K. Nguyen, G. T. McCandless, M. A. Omary and J. J. Gassensmith, *Chem. Commun.*, 2017, **53**, 9890–9893; (b) P. M. Alvey, J. J. Reczek, V. Lynch and B. L. Iverson, *J. Org. Chem.*, 2010, **75**, 7682–7690; (c) J. J. Reczek, K. R. Villazor, V. Lynch, T. M. Swager and B. L. Iverson, *J. Am. Chem. Soc.*, 2006, **128**, 7995–8002; (d) T. He, M. Stolte, C. Burschka, N. H. Hansen, T. Musiol, D. I. Kalblein, J. Pflaum, X. Tao, J. Brill and F. Würthner, *Nat. Commun.*, 2015, **6**, 5954; (e) M. Diebold, E. Christ, L. Biniak, L. Karmazin, B. Heinrich, C. Contal, S. Ghosh, P. J. Mesini and M. Brinkmann, *J. Mater. Chem. C*, 2019, **7**, 13120–13129; (f) X. Jiao, S. Maniam, S. J. Langford and C. R. Mc Neill, *Phys. Rev. Mater.*, 2019, **3**, 013606.
- (a) R. Pfattner, M. Mas-Torrent, I. Bilotti, A. Brillante, S. Milita, F. Liscio, F. Biscarini, T. Marszalek, J. Ulanski, A. Nosal, M. Gazicki-Lipman, M. Leufgen, G. Schmidt, L. W. Molenkamp, V. Laukhin, J. Veciana and C. Rovira, *Adv. Mater.*, 2010, **22**, 4198–4203; (b) S. Galindo, A. Tamayo, F. Leonardi and M. Mas-Torrent, *Adv. Funct. Mater.*, 2017, **27**, 1700526; (c) D. Gentili, M. Gazzano, M. Melucci, D. Jones and M. Cavallini, *Chem. Soc. Rev.*, 2019, **48**, 2502–2517.
- (a) Z.-G. Zhang, H. Li, B. Qi, D. Chi, Z. Jin, Z. Qi, J. Hou, Y. Li and J. Wang, *J. Mater. Chem. A*, 2013, **1**, 9624–9629; (b) T. Akiyama, *Bull. Chem. Soc. Jpn.*, 2019, **92**, 1181–1199; (c) T. de Roo, J. Haase, J. Keller, Ch. Hinz, M. Schmid, D. V. Seletskiy, H. Cölfen, A. Leitenstorfer and S. Mecking, *Adv. Funct. Mater.*, 2018, **14**, 2714–2719; (d) B. Russ, M. J. Robb, B. C. Popere, E. E. Perry, C.-K. Mai, S. L. Fronk, S. N. Patel, T. E. Mates, G. C. Bazan, J. J. Urban, M. L. Chabinye, C. J. Hawker and R. A. Segalman, *Chem. Sci.*, 2016, **7**, 1914–1919.
- F. Doria, A. Oppi, F. Manoli, S. Botti, N. Kandath, V. Grande, I. Manet and M. Freccero, *Chem. Commun.*, 2015, **51**, 9105–9108.
- H. Wei, M. Lv, X. Duan, S. Li, Y. Yao, K. Wang, P. Zhang, X. Li and H. Chen, *Med. Chem. Res.*, 2014, **23**, 2277–2286.
- F. Würthner, V. Stepanenko, Z. Chen, C. R. Saha-Möller, N. Kocher and D. Stalke, *J. Org. Chem.*, 2004, **69**, 7933–7939.
- L. Ferlauto, F. Liscio, E. Orgiu, N. Masciocchi, A. Guagliardi, F. Biscarini, P. Samori and S. Milita, *Adv. Funct. Mater.*, 2014, **24**, 1090–1099.
- O. D. Jurchescu, D. A. Mourey, S. Subramanian, S. R. Parkin, B. M. Vogel, J. E. Anthony, T. N. Jackson and D. J. Gundlach, *Phys. Rev.*, 2009, **80**, 0852012009.
- (a) A. S. Raw, M. S. Furness, D. S. Gill, R. C. Adams, F. O. Holcombe and L. X. Yu, *Adv. Drug Delivery Rev.*, 2004, **56**, 397–414; (b) G. Goldbeck, E. Pidcock and C. Groom, *Solid Form Informatics for Pharmaceuticals and Agrochemicals: Knowledge-Based Substance Development and Risk Assessment*, CCDC, 2012 www.ccdc.cam.ac.uk/support-and-resources/ccdcresources/Solid_Form_Informatics.pdf; (c) J. Bernstein, *Polymorphism in Molecular Crystals*, Clarendon Press, Oxford, 2002.
- D. W. Armstrong, K. R. Rund, U. B. Nair and G. L. Raid III, *Current*, 1996, **15**, 57–61.
- S. Bergantin and M. Moret, *Cryst. Growth Des.*, 2012, **12**, 6035–6041.
- J. Mizuguchi, *J. Phys. Chem. B*, 2004, **108**, 8926–8930.
- K. C. Lee, G. S. Ryu, S. Chen, G. Kim, Y. Y. Noh and C. Yang, *Org. Electron.*, 2016, **37**, 402–410.
- K. Tagami, A. Saitoh, K. Sekido, M. Sekiya and M. Nishi, *US Pat.*, US10065969B2, 2018.
- C. R. DeBlase, K. Hernández-Burgos, J. M. Totter, D. J. Fortman, D. dos, S. Abreu, R. A. Timm, I. C. N. Diógenes, L. T. Kubota, H. D. Abruña and W. R. Dichtel, *Angew. Chem., Int. Ed.*, 2015, **54**, 13225–13229.
- S. A. Bhat, C. Das and T. K. Maji, *J. Mater. Chem. A*, 2018, **6**, 19834–19842.
- M. R. Ajayakumar, P. Mukhopadhyay, S. Yadav and D. Ghosh, *Org. Lett.*, 2010, **12**, 2646–2649.
- See for example: (a) Do-K Kim, H. Lee, X. Zhang, J.-H. Bae and J. Park, *Micromachines*, 2019, **10**, 727–736; (b) L. Ruan,

- D. Zhang, J. Tong, J. Kang, Y. Chang, L. Zhou, G. Qin and X. Zhang, in *Ferroelectrics and Their Applications*, ed. H. Irzaman, Intechopen, London, UK, 2018, ch. 8; (c) F. S. Kim, C. H. Park, Y. Na and S. A. Jeneke, *Org. Electr.*, 2019, **69**, 301–317.
- 28 TOPAS-R, V. 3.0, 2005, Bruker AXS, Karlsruhe, Germany.
- 29 A. A. Coelho, *J. Appl. Crystallogr.*, 2003, **36**, 86–95.
- 30 R. W. Cheary and A. Coelho, *J. Appl. Crystallogr.*, 1992, **25**, 109–121.
- 31 W. A. Dollase, *J. Appl. Crystallogr.*, 1986, **19**, 267–272.
- 32 B. Schrode, S. Pachmajer, M. Dohr, C. Röthel, J. Domke, T. Fritz, R. Resel and O. Werzer, *J. Appl. Crystallogr.*, 2019, **52**, 683–689.
- 33 M. J. Frisch, G. W. Trucks, H. B. Schlegel, G. E. Scuseria, M. A. Robb, J. R. Cheeseman, G. Scalmani, V. Barone, G. A. Petersson, H. Nakatsuji, X. Li, M. Caricato, A. V. Marenich, J. Bloino, B. G. Janesko, R. Gomperts, B. Mennucci, H. P. Hratchian, J. V. Ortiz, A. F. Izmaylov, J. L. Sonnenberg, D. Williams-Young, F. Ding, F. Lipparini, F. Egidi, J. Goings, B. Peng, A. Petrone, T. Henderson, D. Ranasinghe, V. G. Zakrzewski, J. Gao, N. Rega, G. Zheng, W. Liang, M. Hada, M. Ehara, K. Toyota, R. Fukuda, J. Hasegawa, M. Ishida, T. Nakajima, Y. Honda, O. Kitao, H. Nakai, T. Vreven, K. Throssell, J. A. Montgomery, Jr., J. E. Peralta, F. Ogliaro, M. J. Bearpark, J. J. Heyd, E. N. Brothers, K. N. Kudin, V. N. Staroverov, T. A. Keith, R. Kobayashi, J. Normand, K. Raghavachari, A. P. Rendell, J. C. Burant, S. S. Iyengar, J. Tomasi, M. Cossi, J. M. Millam, M. Klene, C. Adamo, R. Cammi, J. W. Ochterski, R. L. Martin, K. Morokuma, O. Farkas, J. B. Foresman and D. J. Fox, *Gaussian 16, Revision C.01*, Gaussian, Inc., Wallingford CT, 2016.
- 34 A. D. Becke, *J. Chem. Phys.*, 1993, **98**, 5648–5652.
- 35 P. J. Stephens, F. J. Devlin, C. F. Chabalowski and M. J. Frisch, *J. Chem. Phys.*, 1994, **98**, 11623–11627.
- 36 M. R. Ajayakumar and P. Mukhopadhyay, *Chem. Commun.*, 2009, 3702–3704.
- 37 J. Mizuguchi, *Zeit. Kristallogr. NCS*, 2002, **218**, 137–138.
- 38 J. Mizuguchi, *Zeit. Kristallogr. NCS*, 2002, **218**, 139–140.
- 39 K. Miyatake, H. Zhou, T. Matsuo, H. Uchida and M. Watanabe, *Macromolecules*, 2004, **37**, 4961–4966.
- 40 H. E. Katz, T. Siegrist, J. H. Schön, C. Kloc, B. Batlogg, A. J. Lovinger and J. Johnson, *ChemPhysChem*, 2001, **3**, 167–172.
- 41 P. Trucano and R. Chen, *Nature*, 1975, **258**, 136–137.
- 42 E. Keller, *Chem. Zeit*, 1980, **14**, 56–60.
- 43 C. F. Macrae, I. Sovago, S. J. Cottrell, P. T. A. Galek, P. McCabe, E. Pidcock, M. Platings, G. P. Shields, J. S. Stevens, M. Towler and P. A. Wood, *J. Appl. Crystallogr.*, 2020, **53**, 226–235.
- 44 M. Hasegawa and S. Horii, *Polym. J.*, 2007, **39**, 610–621.
- 45 See, e.g., (a) A. S. Mathews, H. Kim and C.-S. Ha, *Macromol. Res.*, 2007, **15**, 114–128; (b) ed. K. L. Mittal, *Polyimides and other high temperature polymers: synthesis, characterization, and applications*, CRC Press, Boca Raton, FL, 2009, vol. 5.
- 46 N. B. Kolhe, R. Nandini Devi, S. P. Senanayak, B. Jancy, K. S. Narayan and S. K. Asha, *J. Mater. Chem.*, 2012, **22**, 15235–15246.
- 47 A. Dittmer, R. Izsák, F. Neese and D. Maganas, *Inorg. Chem.*, 2019, **58**, 9303–9315.
- 48 J. C. S. Costa, R. J. S. Taveira, C. F. R. A. C. Lima, A. Mendes and L. M. N. B. F. Santos, *Opt. Mater.*, 2016, **58**, 51–60.
- 49 A. Insuasty, S. Carrara, L. Tang, C. Forsyth, C. F. Hogan, C. R. McNeill and S. J. Langford, *ChemPlusChem*, 2019, **84**, 1638–1642.
- 50 P. Ledwon, D. Ovsianikova, T. Jarosz, S. Gogoc, P. Nitschke and W. Domagala, *Electrochim. Acta*, 2019, **307**, 525–535.
- 51 J. Lee, J. L. Hernandez, I. Pelse, J. R. Reynolds and C. Yang, *J. Mater. Chem. C*, 2018, **6**, 10532–10537.
- 52 W. Porzio, C. Scavia, L. Barba, G. Arrighetti and S. Milita, *Eur. Polym. J.*, 2011, **47**, 273–283.
- 53 (a) S. Milita, M. Servidori, F. Cicoira, C. Santato and A. Pifferi, *Nucl. Instr. Meth. B*, 2006, **246**, 101–105; (b) S. Milita, C. Santato and F. Cicoira, *Appl. Surf. Sci.*, 2006, **252**, 8022–8027.
- 54 K. Feng, H. Guo, J. Wang, Y. Shi, Z. Wu, M. Su, Z. Zhang, J. H. Son, H. Y. Woo and X. Guo, *J. Am. Chem. Soc.*, 2021, **143**, 1539–1552, and references therein.