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Supramolecular Chalcogen-Bonded Semiconducting Nanoribbons at Work in Lighting Devices

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Abstract: This work describes the design and synthesis of a π -conjugated telluro[3,2- β][1]-tellurophene-based synthon that, embodying pyridyl and haloaryl chalcogen-bonding acceptors, self-assembles into nanoribbons through chalcogen bonds. The ribbons π -stack in a multi-layered architecture both in single crystals and thin films. Theoretical studies of the electronic states of chalcogen-bonded material showed the presence of a local charge density between Te and N atoms. OTFT-based charge transport measurements showed hole-transport properties for this material. Its integration as a p-type semiconductor in multi-layered Cu^I-based light-emitting electrochemical cells (LECs) led to a 10-fold increase in stability (38 h vs. 3 h) compared to single-layered devices. Finally, using the reference tellurotellurophene congener bearing a C–H group instead of the pyridyl N atom, a herringbone solid-state assembly is formed without charge transport features, resulting in LECs with poor stabilities (<1 h).

Introduction

Since the first observation in the 1960s^[1] and conceptualization in the 2000s,^[2] chalcogen-bonding interactions (ChBIs, X–Ch $\cdots$ Y)^[3] have raised great interest among chemists to gain control on the conformation of π -conjugated polymers,^[4] conceive new catalysts,^[5] receptors for anion recognition,^[6] and molecular tectons for crystal

engineering.^[7] In particular, in molecular crystals, ChBIs have been used to program discrete and polymeric architectures in the solid state.^[8] Prior art in the field includes the use of benzo-2,1,3-chalcogenodiazoles to engineer infinite supramolecular ribbons.^[9] In these architectures, the Ch and N atoms engage in multiple ChBIs, forming 2Ch-2N squares recognition motifs. When mounted as walls on a resorcin-[4]arene scaffold, the benzo-2,1,3-chalcogenodiazole units engage in 16 intramolecular ChBIs, templating the formation of a self-assembled capsule.^[10] Other solid-state architectures include kinked supramolecular polymers,^[11] in which 2-substituted benzo-1,3-chalcogenazoles are wired through single ChBIs. Heteromolecular solid-state polymers were also obtained by exploiting simultaneous chalcogen- and halogen-bonding interactions^[12] using the chalcogenazolo[5,4- β]pyridine (CGP) unit developed by our group,^[13] known to persistently form non-covalent 2Ch-2N hexagon motifs.^[14] However, the exploitation of chalcogen-bonded soft materials in optoelectronics has so far remained underdeveloped,^[15] with the exclusion of few examples in non-linear optics^[16] and in organic thin-film transistors (OTFTs).^[17] Capitalizing on the strong orbital contribution ($n^2_{(Y)} \rightarrow \sigma^*_{(Ch-X)}$),^[18] one could envisage to exploit ChBIs to conceive supramolecular semiconductors in which the molecular units are electronically coupled.^[19] Building on this idea, this work describes the first example of the design of a π -conjugated molecular synthon that, by embodying two doubly chalcogen-bonded units, assembles into supramolecular nanoribbons at the solid-state (Scheme 1).

In thin films, the ribbon would π -stack in a multi-layered architecture, ultimately giving rise to a p-type semiconduct-

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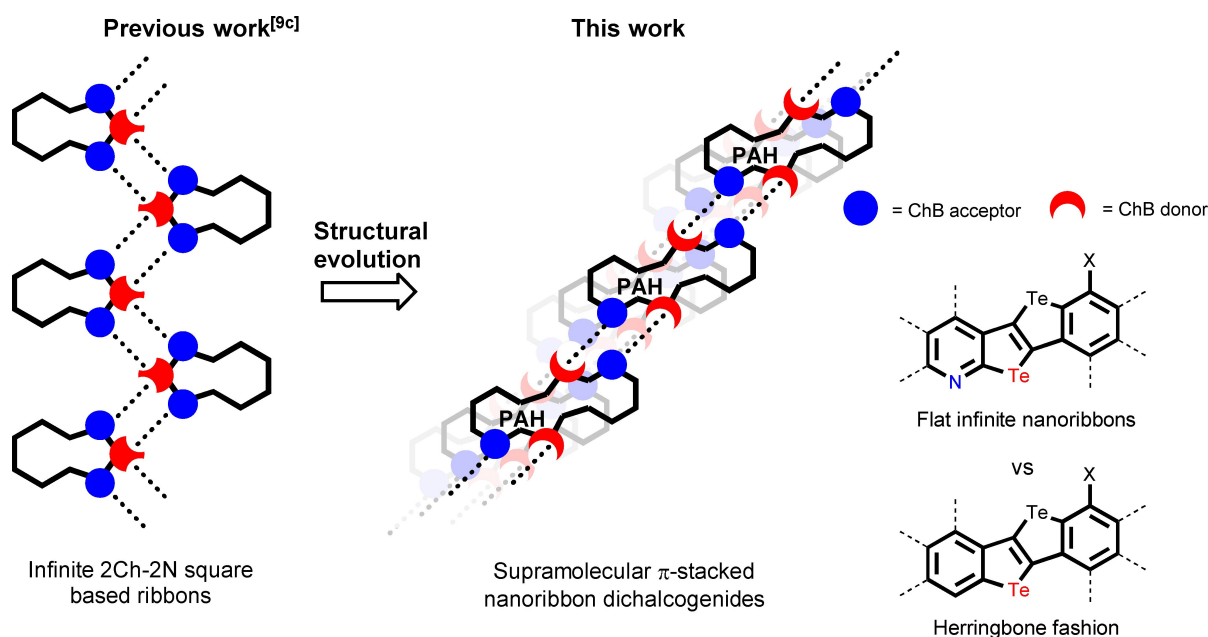
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Scheme 1. Schematic representation of supramolecular chalcogen-based nanoribbons held by ChBIs in the solid-state.

ing material as depicted by charge-transport measurement with TFTs. This was used to decouple hole injection/transport and exciton formation in multi-layered red-emitting Cu^I-complex-based light-emitting electrochemical cells (LECs), realizing 10-fold increased stabilities compared to reference devices with traditional hole-transport layers.^[20]

Results and Discussion

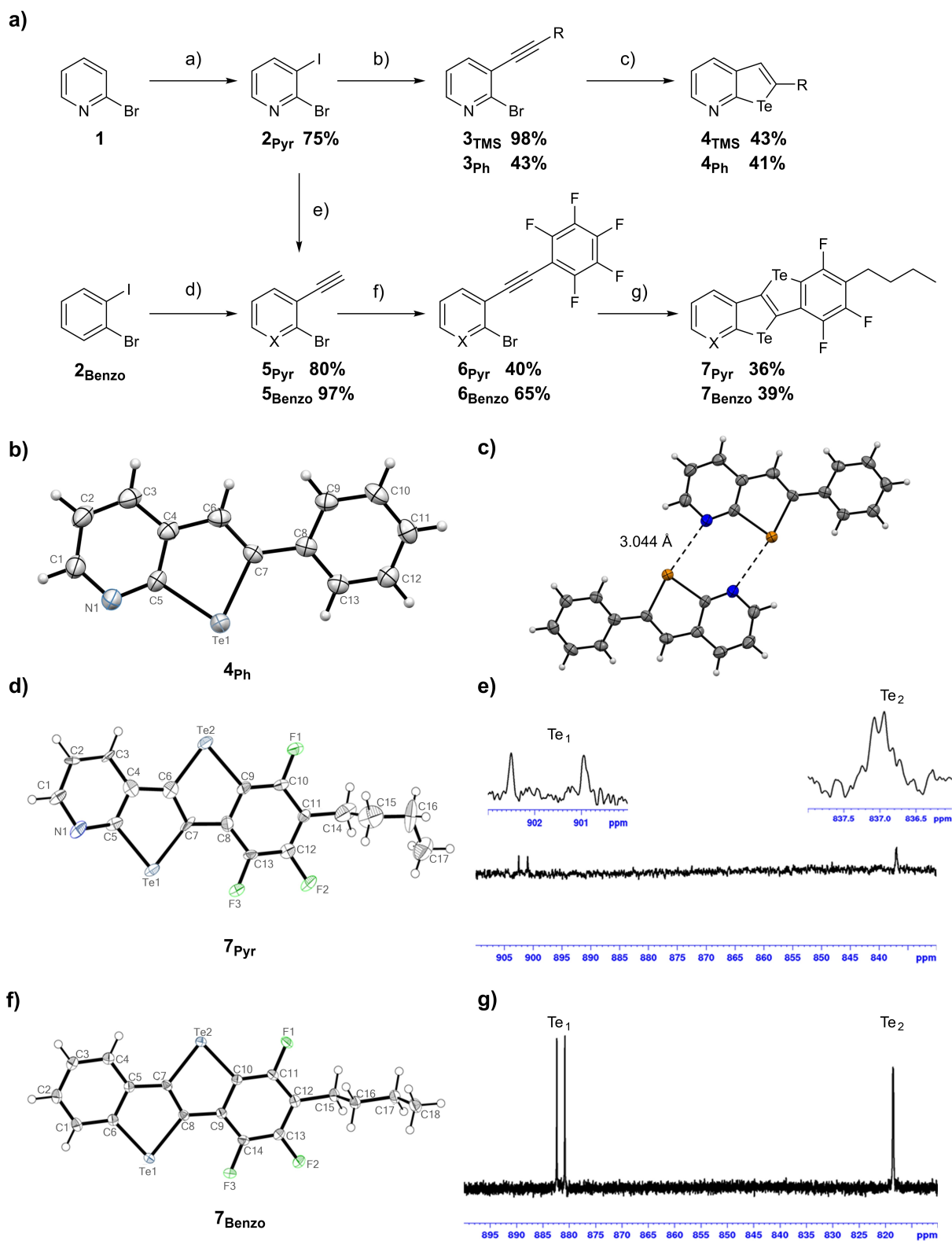
Design and Synthesis

Chalcogen-bonded ribbons have been previously obtained with benzo-1,2,5-telluradiazole synthons exploiting the two Te-centered σ -holes to engage in bifurcated ChBIs.^[21] In our approach, we considered the use of π -conjugated scaffolds containing two chalcogen-bonding recognition units at their extremities (Scheme 1).^[22] Capitalizing on the persistent recognition behavior of the Te-CGP unit, we conjectured that any polycyclic aromatic framework encoding chalcogen-bonding donor Te and acceptor N atoms in a second nearest neighboring (SNN) topology,^[13] *i.e.* a peripheral telluropheno[5,4- β]pyridine-like substructure, can undergo in-plane self-assembly. Anticipating scarce solubility and processability of any derivative exposing multiple SNN-type recognition units,^[23] we envisaged a π -conjugated scaffold that would include heteromolecular chalcogen-bonding recognition units. One moiety would develop strong 2Ch-2N hexagons, while the second one would establish weak interactions, to self-assemble into supramolecular ribbons at the solid state (Scheme 1). Building on the telluro[3,2- β][1]-tellurophene bicycle as the chalcogen-bond donor framework,^[24] we designed a self-assembling polycyclic aromatic module that includes both a Te/N SNN-type and a Te/halogen recognition pairs (Scheme 1b).

At the synthetic planning level, this design guided us to contemplate an organometallic-mediated Te-insertion reaction, followed by a one-pot intramolecular cyclization as the key synthetic step to form the telluro[3,2- β][1]-tellurophene core from the relevant disubstituted alkyne derivative bearing both pyridyl and haloaryl moieties. A reference module bearing a C–H functionality at the place of the N atom, which is unable to undergo ribbon-like assembly, has been also designed.

Taking advantage of the methodologies developed for preparing benzotellurophenes,^[25] our synthetic efforts started with the investigations of the intramolecular cyclization reaction to obtain pyridinotellurophene derivatives from the corresponding 2-bromo-3-alkynylpyridine (Scheme 2a). Selective deprotonation of commercially available 2-bromopyridine **1** using LDA in THF at -95°C , followed by the slow addition of I₂, gave access to 2-bromo-3-iodopyridine **2_{pyr}** in high yield.^[26] Sonogashira-type cross-coupling reaction of **2_{pyr}** with the relevant ethynyl derivative in the presence of [Pd(PPh₃)₂Cl₂] and CuI, gave molecules **3_{TMS}** and **3_{ph}** in excellent and good yields, respectively.^[27] Treatment of the bromo-containing alkynes with a trialkyl magnesiate (prepared in situ by mixing *i*-PrMgCl with *n*-BuLi at 0°C)^[28] followed by the addition of freshly-ground elemental Te powder led to pyridinotellurophenes **4_{TMS}** and **4_{ph}** in 43 % and 41 % yields, respectively, through Te-insertion and consecutive intramolecular cyclization (Scheme 2a).

X-ray analysis of a single crystal of **4_{ph}** obtained by slow evaporation of a CHCl₃ solution confirmed the presence of the bicyclic tellurophenopyridine moiety. Each bicycle arranges in non-covalent 2Ch-2N hexagonal dimers through the formation of double ChBIs ($d_{\text{Te}\cdots\text{N}} = 3.044 \text{ \AA} < \text{sum of vdW radii} = 3.67 \text{ \AA}$, $\text{C}_7\text{--Te}_1\cdots\text{N}_1 = 169.8^{\circ}$; Scheme 2c), validating our hypothesis, for which the tellurophenopyridine



Scheme 2. a) Synthetic pathway to **4** and **7**. Reagents and conditions: a) 1. LDA, THF, -95°C , 1 h; 2. I_2 , THF, -95°C to r.t., 4 h; b) $\text{R}\equiv$, 5 mol% $[\text{Pd}(\text{PPh}_3)_2\text{Cl}_2]$, 10 mol% CuI, NEt_3 , 1,4-dioxane, r.t., 1.5 h; c) 1. $i\text{-PrBu}_2\text{MgLi}\cdot\text{LiCl}$, THF, 0°C , 1 h; 2. Te^0 , r.t., 3 h; 3. EtOH, r.t., overnight; d) 1. TMSA, 5 mol% $[\text{Pd}(\text{PPh}_3)_2\text{Cl}_2]$, 10 mol% CuI, NEt_3 , 1,4-dioxane, r.t., 1.5 h; e) the same conditions as those used for (d); f) for **6_Pyr**: 5 mol% $[\text{Pd}(\text{PPh}_3)_2\text{Cl}_2]$, 10 mol% CuI, $\text{C}_6\text{F}_5\text{I}$, $i\text{-PrNH}_2$, toluene, 80°C , 3 h; for **6_Benzo**: 5 mol% $[\text{Pd}(\text{PPh}_3)_2\text{Cl}_2]$, 10 mol% CuI, $\text{C}_6\text{F}_5\text{I}$, NEt_3 , 40°C , overnight; g) the same conditions as those used for (c); ORTEP representation of a single molecule of b) **4_Ph**, d) **7_Pyr** and f) **7_Benzo** drawn with 50% displacement ellipsoid;^[32] c) crystal structure of **4_Ph** showing the dimer formation in the solid state; 189.5 MHz ^{125}Te NMR spectrum in C_6D_6 of (e) **7_Pyr** and g) **7_Benzo**.

scaffold undergoes non-covalent dimerization similarly to the CGP-type motifs.^[13,14] To prepare the target tellurotellurophene module, we conjectured that highly electron-deficient perfluorinated aryl substituents could be used as favorable functionalities to induce an intramolecular S_NAr cyclization reaction.^[29,30] Thus, the synthesis of a disubstituted alkyne derivative bearing both a pyridyl and a pentafluoro aryl moiety was planned. TMS-deprotection of **3**_{TMS} with K_2CO_3 in a 2:1 mixture of MeOH and CH_2Cl_2 followed by Sonogashira cross-coupling reaction with iodopentafluorobenzene in the presence of $[PdCl_2(PPh_3)_2]$ and CuI led to the disubstituted alkynyl derivative **6**_{Pyr} in 40 % yield. Derivative **6**_{Pyr} was converted into bis tellurotellurophenes **7**_{Pyr} by Te-insertion followed by a double intramolecular cyclization reaction upon addition of *i*-PrBu₂MgLi-LiCl in THF and of elemental Te (Scheme 2a). Notably, an unexpected alkylation substitution reaction occurred at the 4-position of the pentafluoroaryl moiety.^[31] A similar synthetic pathway was also exploited to prepare congener **7**_{Benzo}. Bis-substituted precursor **5**_{Benzo} was prepared in 97 % yield starting from 2-bromiodobenzene **2**_{Benzo} by Sonogashira cross-coupling with TMS-acetylene (TMSA) followed by TMS-deprotection. Subsequent cross-coupling reaction of iodopentafluorobenzene with **5**_{Benzo} gave **6**_{Benzo} in 65 % yield. Te-insertion in the presence of *i*-PrBu₂MgLi-LiCl gave the tellurotellurophene derivative **7**_{Benzo} in 39 % yield. All compounds were fully characterized by 1H , ^{19}F , ^{13}C and ^{125}Te (for **7**_{Pyr} and **7**_{Benzo}—Schemes 2e and 2g) NMR spectroscopy, IR, HR-mass spectrometry, TGA (Figures S32, S33) and X-ray diffraction analysis (Schemes 2b–f—see Supporting Information).^[32] In particular, ^{125}Te NMR spectra of **7**_{Pyr} and **7**_{Benzo} were measured in C_6D_6 to probe the presence of two chemically-non-equivalent chalcogen nuclei in the telluro-[3,2-*b*][1]-tellurophene core, namely Te_1 and Te_2 (Schemes 2e and g). While a significant “through-space” internuclear spin-spin ^{19}F - ^{125}Te coupling constant ($^4J_{^{19}F-^{125}Te}$) value of 290 Hz was observed for the Te_1 resonances, a weak through-bond $^3J_{^{19}F-^{125}Te}$ value of about 26 Hz was measured

for the Te_2 nuclei in both molecules, confirming the in-plane molecular asymmetry.^[33]

Photophysical and Redox Properties

The absorption spectra of derivatives **7**_{Pyr} and **7**_{Benzo} were recorded in CH_2Cl_2 at r.t. (Figure 1). Both compounds exhibit well-structured absorption bands with two peaks centered at ≈ 330 and 380 nm associated to $\epsilon_{max} \approx 10000\text{ M}^{-1}\text{cm}^{-1}$. Although both molecules are not emissive in CH_2Cl_2 at r.t. in both aerated and deaerated conditions, they revealed significant emissive signals in $CHCl_3/EtOH$ (1:1 v/v) glassy matrix at 77 K (Table S2, Supporting Information).^[11a] The emission profiles of **7**_{Pyr} and **7**_{Benzo} depict identical well-structured envelopes with maxima at ≈ 552 , 570 and 615 nm.

Notably, they exhibit only phosphorescence emission with excited state lifetimes of $\approx 700\text{ }\mu\text{s}$. This suggests a quantitative intersystem crossing (ISC) process induced by the presence of the Te atom (see below).^[11a,34]

Cyclic voltammetry (CV) experiments in CH_2Cl_2 and 1,1,2,2-tetrachloroethane (TCE) were performed to unravel the redox properties of **7**_{Pyr} and **7**_{Benzo} (all potential values are given vs. Fc/Fc^+). Both compounds exhibited *quasi*-reversible redox behaviors (Table S3 and Figure S34, Supporting Information). The CV behavior was similar in different solvents, with the redox values measured in TCE being anodically shifted with respect to those in CH_2Cl_2 . Moreover, a second irreversible oxidation peak (*ca.* 0.5 V) appeared for both molecules after multiple scans, suggesting the occurrence of side reactions and/or decomposition (Figures S32e and S32f in the Supporting Information). Finally, thin films of **7**_{Pyr} and **7**_{Benzo} showed *quasi*-reversible oxidation events at +0.79 V and +0.87 V, respectively (scan rate of 100 mV s^{-1} ; Figure S35, Supporting Information). Upon repetitive CV scans, the current intensity increased due to a partial charging of the films. This led to a slight shift ($\approx 10/20\text{ mV}$) in the oxidation potential. No other

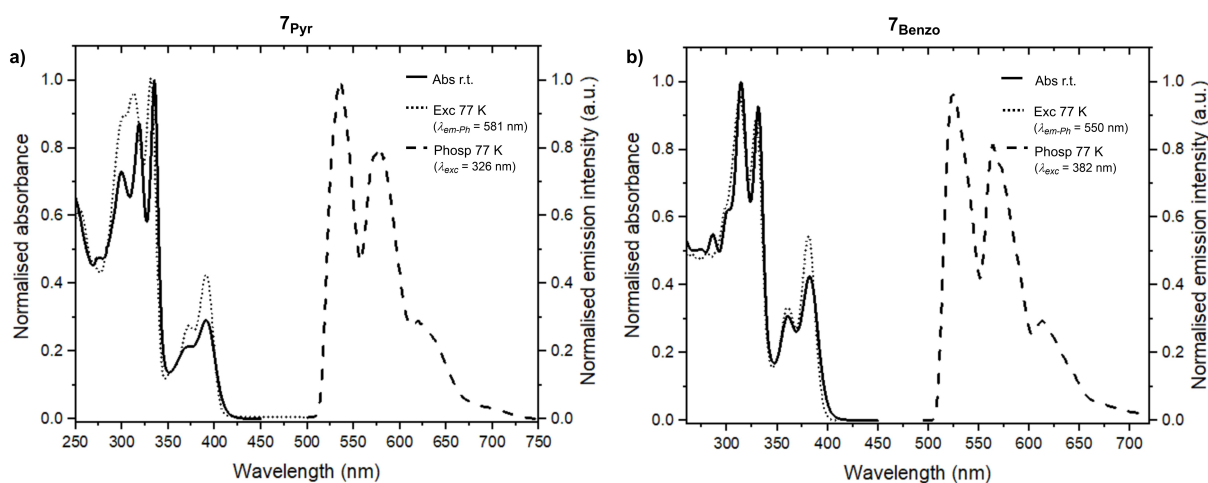


Figure 1. Normalised absorbance (solid line), excitation (dotted line) and phosphorescence (dashed line) spectra of solutions of **7**_{Pyr} (a) and **7**_{Benzo} (b). Absorption was measured in CH_2Cl_2 at r.t., phosphorescence and excitation in $CHCl_3/EtOH$ (1:1 v/v) glassy matrix at 77 K.

redox event was detected after 20 scans, suggesting that these compounds are electrochemically stable in thin films.

Solid-State Organization

Next, we studied the structural properties and the organization of solids made of **7_{Pyr}** and **7_{Benzo}**. The X-ray analysis of single crystals of **7_{Pyr}** is marked by the presence of chalcogen-bonded molecules, each developing into supramolecular ribbons (Figure 2a) held together through two types of ChBIs. While the proximal N and Te₁ atoms ($d_{\text{Te1}\cdots\text{N}}=2.961\text{ \AA}$ < sum of vdW radii = 3.67 Å, $\text{C}_7\text{-Te}_1\cdots\text{N}_1=172.6^\circ$) undergo double ChBIs through the typical doubly-bonded 2Ch-2N hexagon recognition motif of the CGP unit,^[13,14] the Te₂ atom establishes a weak ChBI with a F atom of a neighboring molecule ($d_{\text{Te2}\cdots\text{F2}}=3.295\text{ \AA}$ < sum of vdW radii = 3.53 Å, $\text{C}_9\text{-Te}_2\cdots\text{F}_2=172.6^\circ$). Each ribbon is organized into π - π stacks ($d_{\pi-\pi}=3.390\text{ \AA}$; Figure S36b in the Supporting Information), ultimately leading to a two-dimensional brickwork-like organization (Figure 2b). Crystals of **7_{Benzo}** showed a different molecular packing organization, with only the Te₂ atom engaging in bifurcated intermolecular ChBIs with two neighboring F atoms ($d_{\text{Te(a)}\cdots\text{F3}}=3.259\text{ \AA}$, $\text{C}_{10}\text{-Te}_2\cdots\text{F}_3=164.7^\circ$; $d_{\text{Te(b)}\cdots\text{F2}}=3.489\text{ \AA}$, $\text{C}_7\text{-Te}_2\cdots\text{F}_2=168.9^\circ$, both ChBIs < sum of vdW radii = 3.53 Å; Figure 2c). These bifurcated interactions force the molecules to adopt a herringbone organization stabilized through intermolecular C-H $\cdots\pi$ contacts ($d_{\text{C}\cdots\pi}=2.860\text{ \AA}$, Figure 2d).

To further characterize the materials, we have investigated the local chemical and electronic environment

nearby the chalcogen atoms by mean of solid-state ¹²⁵Te NMR spectroscopy (Figure 3).^[35] The simulation (red line) provides isotropic chemical shift (δ_{iso} , marked with * in Figure 3), the principal component (δ_{11} , δ_{22} and δ_{33}), span (Ω) and skew (κ) values for both the chalcogen nuclei (Table S1 and comment in the Supporting Information). Building on the seminal ¹²⁵Te ssNMR studies by Bryce and co-workers,^[35b] the different values shown by Te₁ and Te₂ nuclei for isotropic chemical shift δ_{iso} (898.5 vs. 845.5 ppm), the largest principal component δ_{11} (1854.7 vs. 2118.3 ppm), the shielded δ_{33} (251.6 vs. 181.2 ppm) and the span values (1603.2 vs. 1937.1 ppm) easily suggest that the Te₁ atom engages in stronger and more directional ChBIs than the Te₂ nucleus due to the formation of the double 2Ch-2N motif. To validate our interpretation, we performed ¹²⁵Te ssNMR measurements with solids of reference **7_{Benzo}**. Here, the different trend of the Ω values for Te₁ and Te₂ (*i.e.*, 2079.9 vs. 1825.4 ppm, respectively) confirms the X-ray observations, for which only weak ChBIs are observed in solids of **7_{Benzo}**.

Computational Studies

To support our hypothesis regarding the bonding contribution of the strong N $\cdots$ Te ChBIs to the electronic properties of the supramolecular materials, the organization of **7_{Pyr}** and **7_{Benzo}** was investigated within the density functional theory (DFT) framework, starting from the experimental cell parameters of the crystal structure (Supporting Information for the calculation methods). *Ab initio* structural optimiza-

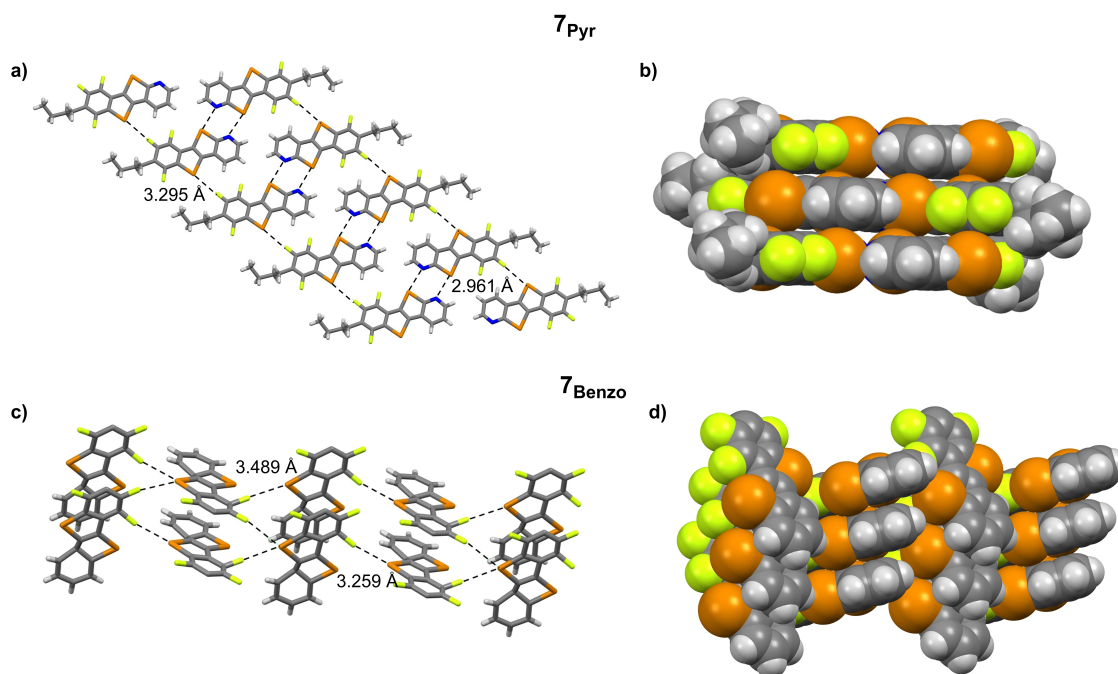


Figure 2. Left: X-ray stick representation of a) the supramolecular nanoribbon developed by **7_{Pyr}**, and c) the herringbone organization of **7_{Benzo}** (*n*-butyl chains omitted for clarity). ChBIs are highlighted with dashed lines. Right: space-fill representation of the crystal packing of b) **7_{Pyr}** and d) **7_{Benzo}**. Crystallization solvents: CHCl₃ and toluene for **7_{Pyr}**, petroleum ether for **7_{Benzo}**. Space groups: *Pbca* for **7_{Pyr}**, *P2₁/c* for **7_{Benzo}**.^[32]

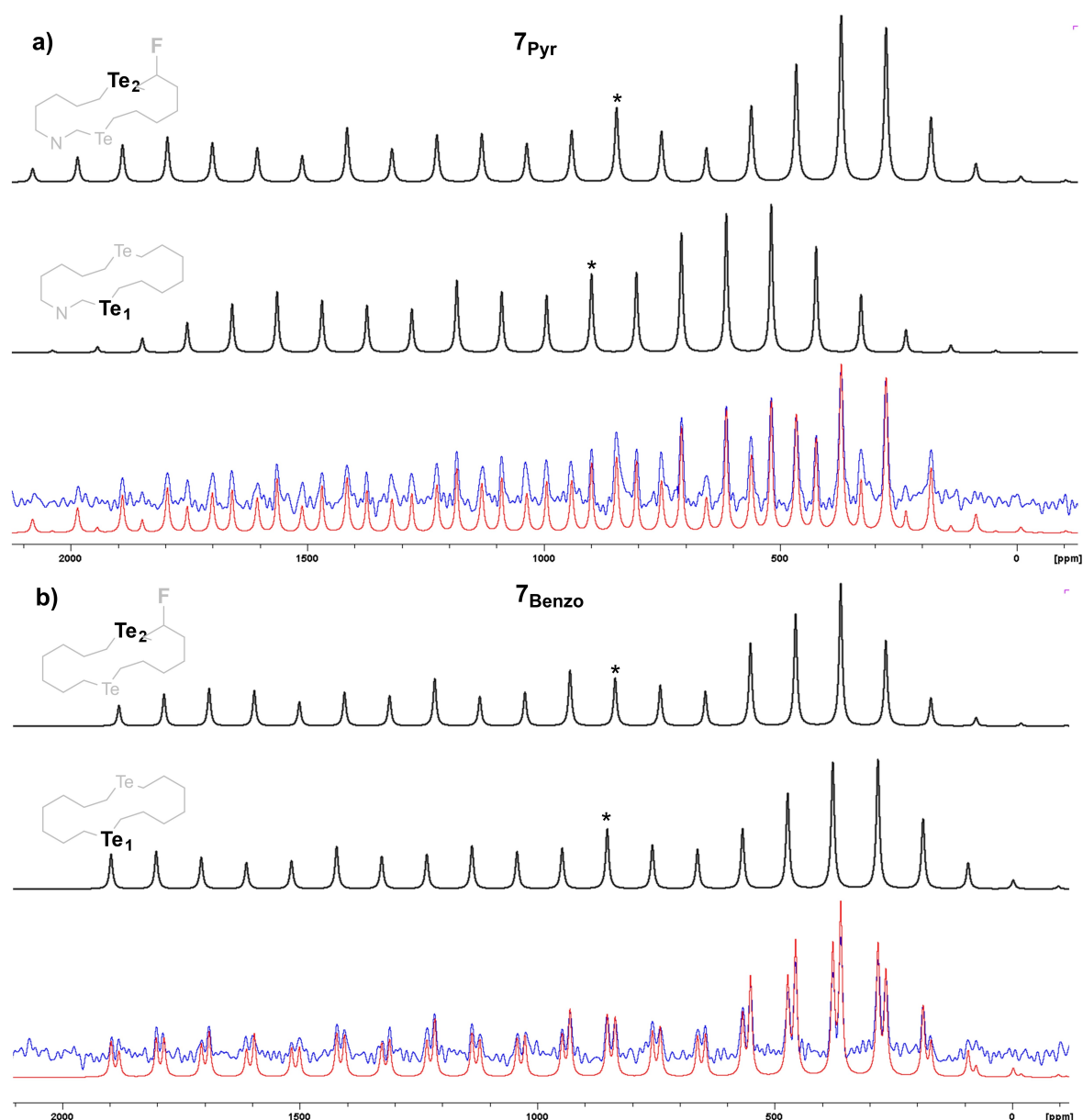


Figure 3. Experimental (blue lines) and simulated (red lines) ^{125}Te ssNMR spectra of a) 7_{Pyr} and b) 7_{Benzo} . Black spectra correspond to the simulated spectra of Te₁ and Te₂ atoms, with the isotropic peaks marked as black asterisks.

tion has been performed considering a chalcogen-bonded monolayer for 7_{Pyr} structure, while a bilayer structure has been considered for 7_{Benzo} to model every Te...F interaction. Te...N and Te...F intermolecular distances have been determined, as well as the interlayer π - π contact, being in good agreement with the experimental values (Figures S38a and S41a). The Partial Density of States (PDOS) has been evaluated (Figure 4). For both structures, we found that the highest states of the valence band are mainly based on Te p orbitals, whereas C p orbitals compose the lowest states of the conduction band.

When investigating the local electronic states at Γ point for 7_{Pyr} , a local charge density is found between Te and N atoms (Figure 5a), located at -3.56 eV below the Fermi energy. On the other hand, when the local electronic states

at Γ point were calculated for 7_{Benzo} (using the same isosurface value as for 7_{Pyr}), no localized charge density was observed between Te and F atoms (Figure 5b).

In order to complete the theoretical study, first-principles simulations were also performed for π - π stacked multi-layers of 7_{Pyr} and 7_{Benzo} structures. In both cases, analogous results were obtained for the multi-layered system of 7_{Pyr} when the local electronic states at Γ point were investigated, showing the same type of Te...N interactions located at -3.71 eV. In accordance with the bilayer case, no localized charge density is found between Te and F atoms in 7_{Benzo} multi-layered structure.

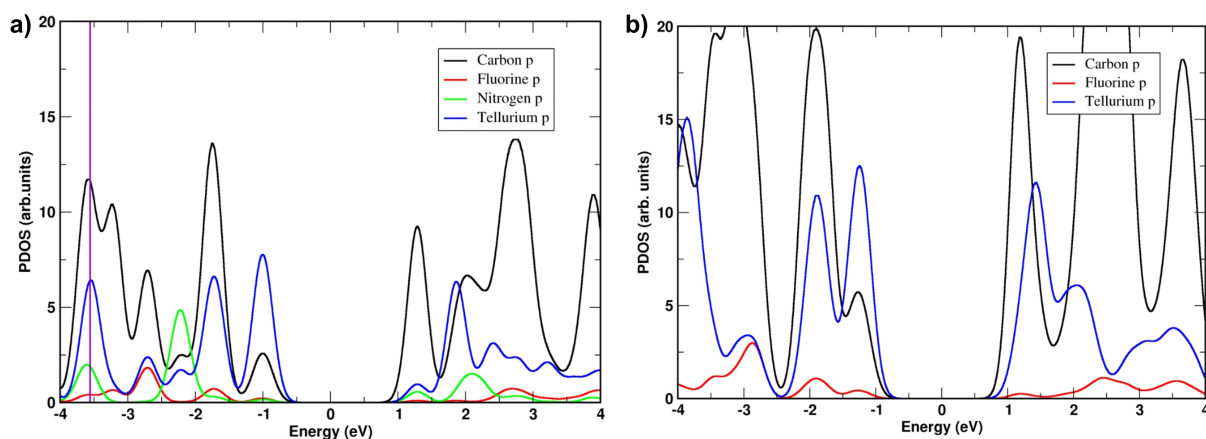


Figure 4. Partial Density of States (PDOS) of a) 7_{Pyr} and b) 7_{Benzo} structures at the DFT-GGA level of theory. The Fermi energy has been normalized to zero. The purple line in (a) indicates the energy at which the Te...N local charge density has been extracted.

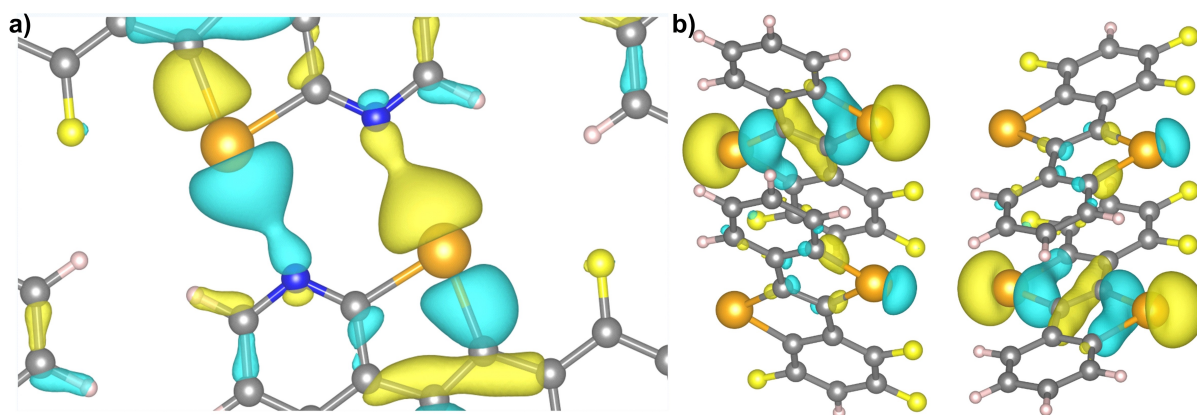


Figure 5. Local charge densities for a) 7_{Pyr} and b) 7_{Benzo} structures, calculated at -3.56 and -3.74 eV, respectively. The isosurface value is chosen to be 0.001 eV Bohr⁻³. The positive (negative) values of the electronic wavefunctions are depicted in yellow (blue). C, Te, N, F and H atoms are respectively represented by grey, orange, blue, yellow and white spheres.

Thin Film Organization

In view of the use of 7_{Pyr} and 7_{Benzo} in LECs (see below), we focused our gaze on the morphological characterization of thin 7_{Pyr} - and 7_{Benzo} -containing films. They were prepared by either spin-coating or drop-casting techniques onto quartz, glass, silicon wafers and ITO/poly(3,4-ethylenedioxythiophene)polystyrene sulfonate (PEDOT:PSS) coated glass (*i.e.*, similar to those used in devices, see below). To get deeper insights into the molecular arrangements of the thin films, we complemented the characterization (for the macroscopic morphological characterization see Polarized Light Microscopy (PLM) and Scanning Electron Microscopy (SEM) measurements in Section 7 of the Supporting Information) with Grazing Incidence X-ray Diffraction (GIXRD) using synchrotron radiation. In line with the observations provided by PLM and SEM, the morphology of thin 7_{Pyr} -films is substrate-independent (Figure 6). In Figure S45a, depicting the GIXRD pattern of a highly ordered 7_{Pyr} -film spin-coated on a silicon wafer, one can notice the presence of intense and high-order reflections

$[002n]$ ($n \in \{0,1,2,\dots\}$) in the out-of-plane direction. This demonstrates a preferential orientation of molecules 7_{Pyr} along the c -axis in edge-on configuration (Figure 6e and Figure S45c). This structural hypothesis is further supported by the full GIXRD indexing of the main reflections calculated from the crystal structure. Moreover, the lattice constant derived from the GIXRD reflection $[004]$ $c = 44.12(2)$ Å corresponds to the crystal lattice constant within the experimental error bar.

The intense reflection $[025]$ is the fingerprint for periodic π - π stacking ($d_{\pi-\pi} = 3.39$ Å) as observed in the single crystal X-ray structure, as well as visualized in the view perpendicular to the $[025]$ reflection direction (Figure S45d). The same finding is observed in the spin-coated film (Figures 6a,c), but with higher disorder. The corresponding drop-casted film on ITO/PEDOT:PSS covered glass (Figure S45b, Supporting Information) shows additionally random orientation. This supports the idea for which the organization of 7_{Pyr} -films is similar to that measured in the crystal structure, with the ChBIs ruling the solid-state arrangement also in thin films. On the contrary, spin-coated

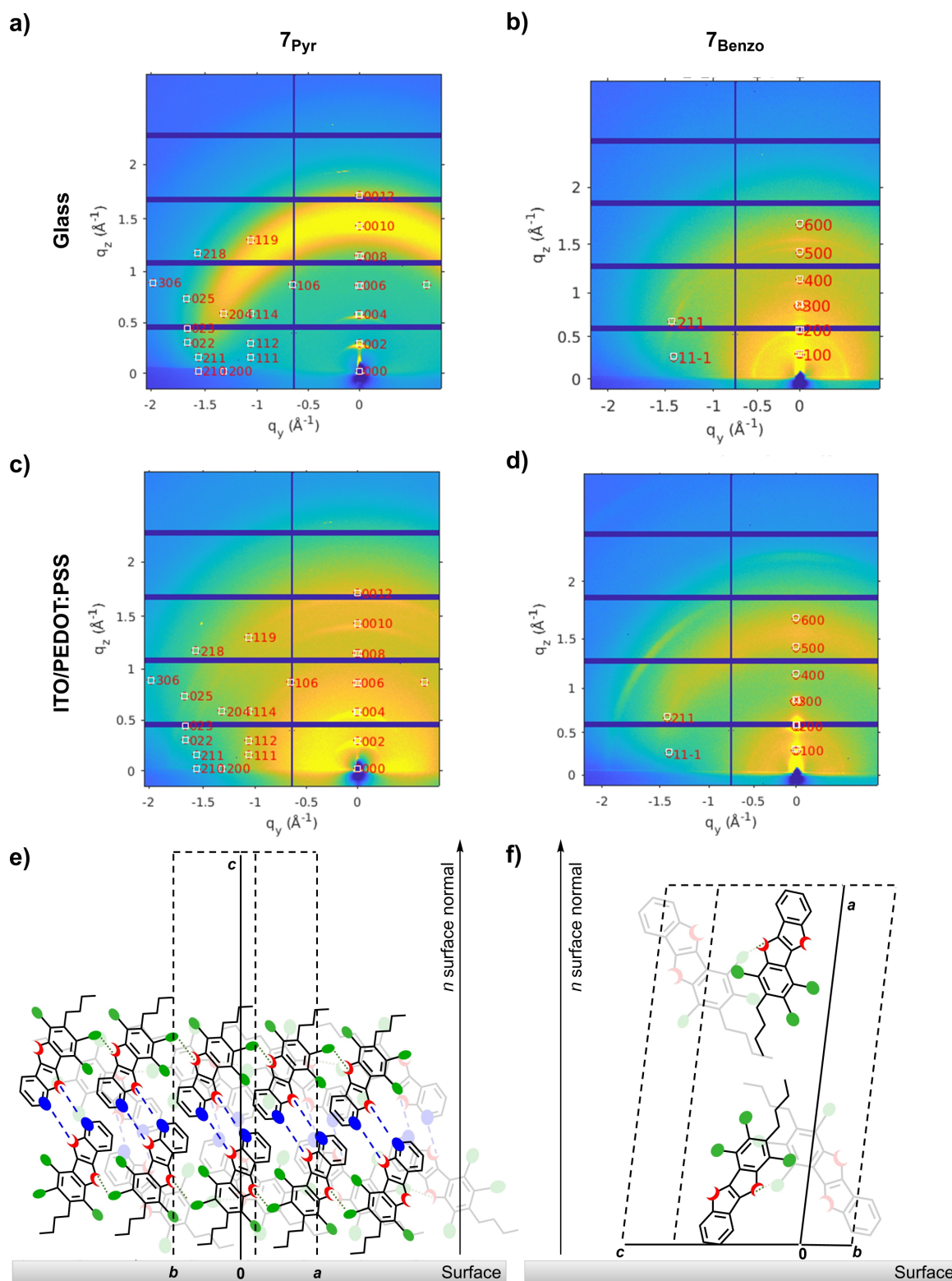


Figure 6. On the left, 2D-GIXRD patterns of glass spin-coated with thin films of a) 7_{Pyr} and b) 7_{Benzo} ; ITO/PEDOT:PSS covered glass with films of c) 7_{Pyr} and d) 7_{Benzo} ; schematic representation of the molecules on the surface for e) 7_{Pyr} and f) 7_{Benzo} . The indices of the main GIXRD reflections calculated from the crystal structure has been overlaid with their appropriate out-of-plane orientation and in-plane random orientation.

7_{Benzo}-films show a lower degree of order than those obtained from **7**_{Pyr} (Figure 6b,d). The 2D-GIXRD patterns suggest the presence of both ordered lamellar stacks along the reciprocal out-of-plane direction [100] arranged in a head-to-tail fashion (as shown in Figure 6f and Figure S46) and partly randomly-oriented molecular distribution on the substrate. Only the high intensity reflections [211] and [111] are visible in the in-plane direction. The simulated GIXRD pattern from the crystal structure was overlaid with a modified lattice parameter *a* (determined from the out-of-plane diffraction peaks) of 26.56 Å compared to the 22.937 Å from the crystal structure.

Thin-Film Transistor Fabrication

To shed light onto the charge transport properties of **7**_{Pyr} and **7**_{Benzo}, and to gain insight into the role of the order at the supramolecular level within the assemblies obtained by solution processing, we prepared and characterized thin-film transistors in a bottom-gate top-contact configuration. In full accordance with the previously presented results, both optical microscopy (Figure S47a,b) and atomic force microscopy (Figure S47c) revealed that molecule **7**_{Pyr} self-assembles into ordered structures whereas **7**_{Benzo} forms disordered clusters. More specifically, **7**_{Pyr} self-assembles into crystals with a thickness of 40 nm and lateral size of several tens of μm, whereas **7**_{Benzo} forms disordered aggregates with a lateral size of a few μm and a thickness exceeding the hundreds of nm scale. The electrical characterization of the thin-film transistors was carried out by recording transfer curves. While both materials revealed the absence of electron transport, a major difference has been observed in terms of hole mobility. Thin films of **7**_{Benzo} displayed a lack of hole transport whereas **7**_{Pyr}-based crystals exhibited a hole transport behavior. A representative transfer curve is displayed in Figure S47g. It exhibits an excellent $I_{\text{on}}/I_{\text{off}}$ ratio of 10^5 and a low off current, indicating the low defect states in the vicinity of HOMO. The extracted hole mobility of **7**_{Pyr} was calculated as $8.8 \times 10^{-5} \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$. While the device is obviously unoptimized for a specific application and there is room for enhancement of the device performance, e.g. via optimization of interfaces of **7**_{Pyr} with the dielectric gate and the metallic electrodes, this result represents the first demonstration of a semiconducting characteristic of supramolecular architecture held together *via* chalcogen bonds.

Implementation in LECs

Considering the ribbon-type, chalcogen-bond-driven solid-state organization, the p-type semiconducting properties and the lack of electroluminescence (Supporting Information, Figure S48), we used thin films of **7**_{Pyr} as hole-transport layers to build LECs.^[20a,c,36] **7**_{Pyr}-containing films were integrated as hole-transport layers to decouple charge injection/transport and exciton formation in LECs with Cu^I-complexes. These emitters are notoriously known to provide

poorly stable LECs under device operation.^[20b,37] A 10-fold enhancement in stability for blue-, yellow- and red-emitting LECs without affecting brightness and efficiency has recently been reported implementing a 10–20 nm thick hole-transport layer (*i.e.*, 4,4'-bis(*N*-carbazolyl)-1,1'-biphenyl or **CBP**), in multi-layered LECs.^[37] However, **CBP** layers tend to easily form micrometer-size aggregates upon device fabrication and storage (*e.g.*, either under ambient or inert conditions), compromising the device self-stability.^[37b,38] Moreover, **CBP**-LECs show a significant bluish green electroluminescence that limits their use in terms of device chromaticity.^[36] **7**_{Pyr}-films are very robust with respect to morphology and fabrication, while they feature a similar electronic energy levels to those of **CBP** (Figure S49).

In view of the above lines, proof-of-concept LECs combining **7**_{Pyr}-, **7**_{Benzo}- or **CBP**-films as hole-transporters and [Cu(dcbq)(Xantphos)]PF₆ (**Cu1**; dcbq: 4,4'-diethylester-2,2'-biquinoline and Xantphos: 4,5-bis(diphenylphosphino)-9,9-dimethylxanthene) as the red emitter were fabricated with the architectures: i) ITO/PEDOT:PSS (70 nm)/**7**_{Pyr} or **7**_{Benzo} (15 nm)/**Cu1** (80 nm)/Al (90 nm) as targeted devices (Figure 7), and ii) ITO/PEDOT:PSS (70 nm)/**CBP** (15 nm)/**Cu1** (80 nm)/Al (90 nm) and ITO/PEDOT:PSS (70 nm)/**Cu1** (80 nm)/Al (90 nm) as reference devices. They were driven at the optimized pulsed current of 20 mA.^[20b] As expected, **7**_{Pyr}-containing LECs outperformed the reference single-layered LECs,^[20b] showing similar efficiency values (*ca.* 0.18 lm/W), but one order of magnitude enhanced stability (3 h *vs.* 38 h for single- and multi-layered LECs, respectively). Both devices showed the same deep-red electroluminescence centered at 663 nm, associated to x/y CIE color coordinates of 0.65/0.32, stable over the entire device lifespan (Figure 7b). Thus, electron-hole recombination efficiently occurs at the interface **7**_{Pyr}/**Cu1**, while hole injection/transport occurs firstly in the chalcogen-bonded layer and it is further transferred to the **Cu1** layer. As such, the formation of oxidized products (*i.e.*, Cu^{II} complex species) is significantly reduced, leading to a significant enhancement of the device stability. At last, we compared the performances of **7**_{Pyr}-based with those integrating the benchmark **CBP**- and reference **7**_{Benzo}-film. In line with the I–V assays (see above), reference **7**_{Benzo} multi-layered LECs showed a dramatic loss of stability ($t_{1/2} < 1$ h) at pulsed 20 mA (Figure 7c). In stark contrast, Figure 7 shows that similar irradiances and stabilities were noted for **7**_{Pyr}- and **CBP**-LECs, while the efficiency is slightly enhanced (0.18 *vs.* 0.15 lm W⁻¹).

Conclusion

Herein we have reported on the design and synthesis of two telluro[3,2-β][1]-tellurophene π-conjugated modules, **7**_{Pyr} and **7**_{Benzo}, which undergo programmed self-assembly at the solid-state through ChBIs. While **7**_{Pyr} was designed to undergo ribbon-like organization through the formation of two double ChBIs, congener **7**_{Benzo} was edited in such a way that only herringbone-type architectures could be formed at the solid-state through weak ChBIs as displayed by single-

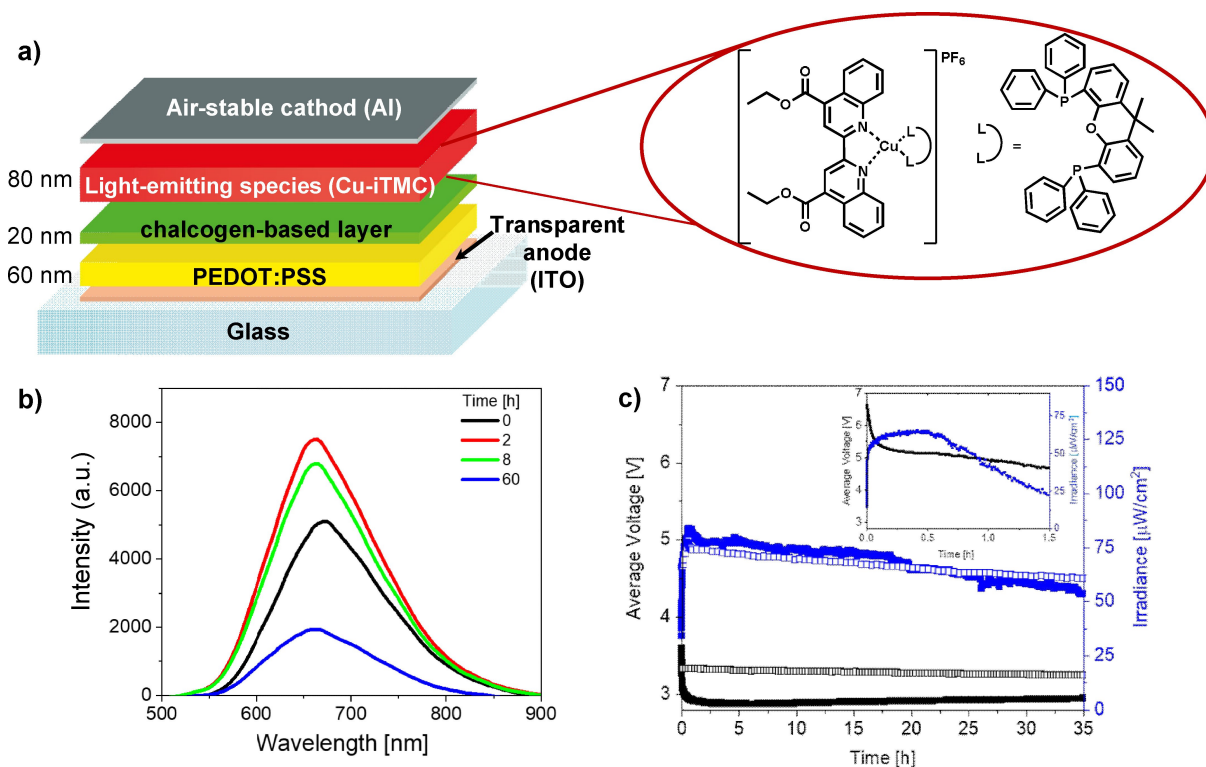


Figure 7. a) Schematic representation of the multi-layered LECs together with the chemical structure of **Cu1**. b) Electroluminescence spectra over time of ITO/PEDOT:PSS (60 nm)/**7Pyr** (15 nm)/**Cu1** (80 nm)/Al at pulsed 20 mA. c) Average voltage and luminance vs. time of devices with **7Pyr**/**Cu1** (full squares), **CBP**/**Cu1** (empty squares), **7Benzo**/**Cu1** (inset) active layers at pulsed 20 mA.

crystal X-ray analysis. A computational investigation suggested that both **7Pyr** and **7Benzo** electronic structures were characterized by Te *p* orbitals in the highest states of the valence bands, and by C *p* orbitals in the lowest states of the conduction band but, only for **7Pyr**, a local charge density was found between Te and N atoms. OTFT-based charge transport measurements of single crystals of **7Pyr** showed hole-transport properties, whereas reference materials with **7Benzo** showed no measurable mobility values. 2D-GIXRD investigations showed that thin films containing **7Pyr** display consistent molecular organization with that observed in single crystals. However, films containing **7Benzo** depicted only short-range order and no π -stacking arrangements. Notably, the structural properties of films of **7Pyr** revealed to be substrate-independent and the same as those measured in single crystals.

Finally, it was demonstrated for the first time the use of chalcogen-bonded compounds in multi-layered LECs as a hole transport layer to decouple transport and emission using red-emitting Cu^I complexes. LECs containing **7Pyr** displayed a stability enhancement (*ca.* 40 h) when compared to devices integrating **7Benzo** (<1 h), with performances similar to those obtained when traditional hole transport layers like CBP are used. Notably, the lack of electroluminescence signal is an interesting feature that paves the way towards its versatile implementation in LECs.

This work describes for the first time the design and preparation of a chalcogen-bonded supramolecular semi-

conductor and its use in solid-state lighting devices, establishing such intermolecular interactions as an effective supramolecular tool for designing functional materials, in general, and non-electroluminescent hole transporters as effective strategy to enhance LECs, in particular.

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Conflict of Interest

The authors declare no conflict of interest.

Data Availability Statement

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

Keywords: Chalcogen Bond • Crystal Engineering • Organic Semiconductor • Supramolecular Architectures

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- [31] ¹H NMR studies of the reaction mixture quenched after 50 minutes (Figure S30, Supporting Information), *i.e.* before the addition of freshly grounded Te, displayed the presence of the alkylated intermediate only. This suggests that the alkylation reaction have already occurred within few minutes after the treatment of **6_{Benzo}** with the organomagnesium reagent. Furthermore, ¹⁹F NMR studies (Figure S31, Supporting Information) allowed to unambiguously determined the position of the *n*-butyl chain on the perfluorinated ring.
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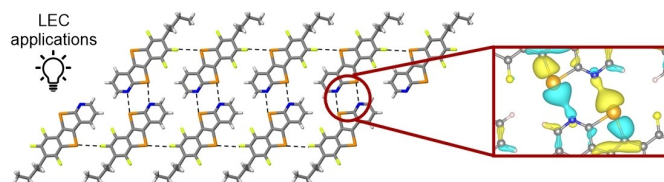
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Research Articles

Crystal Engineering

D. Romito, E. Fresta, L. M. Cavinato,
H. Kählig, H. Amenitsch, L. Caputo,
Y. Chen, P. Samorì, J.-C. Charlier,
R. D. Costa,* D. Bonifazi* — **e202202137**

Supramolecular Chalcogen-Bonded Semi-
conducting Nanoribbons at Work in Light-
ing Devices



Chalcogen-bonded p-type semiconducting ribbons have been engineered through the self-assembly of tailored π -conjugated modules. Theoretical studies of the electronic states showed the presence of a local charge density

between Te and N atoms. Implementation of the ribbons as hole-transport layers in light-emitting electrochemical cells gave rise to devices with performances similar to those obtained with state-of-art semiconductors.