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Room Temperature Ferroelectricity in an Organic Cocrystal

Ren A. Wiscons[†], N. Rajesh Goud[†], Joshua T. Damron, and Adam J. Matzger*

Abstract: Ferroelectric materials exhibit switchable remanent polarization due to reversible symmetry breaking under an applied electric field. Previous research has leveraged temperature-induced neutral-ionic transitions in charge-transfer (CT) cocrystals to access ferroelectrics that operate through displacement of molecules under an applied field. However, displacive ferroelectric behavior is rare in organic CT cocrystals and achieving a Curie temperature (T_C) above ambient has been elusive. Here a cocrystal between acenaphthene 2,3,5,6-tetrafluoro-7,7,8,8-tetracyanoquinodimethane presented that shows switchable remanent polarization at room temperature ($T_C = 68^{\circ}$ C). Raman spectroscopy, X-ray diffraction, and solid-state NMR spectroscopy indicate the ferroelectric behavior is facilitated by AN rotation, deviating from conventional design strategies for CT ferroelectrics. These findings highlight the relevance of non-CT interactions in the design of displacive ferroelectric cocrystals.

Crystalline charge transfer (CT) complexes formed between neutral molecular components demonstrate a broad range of fractional electron transfer that predicates applications in most fields of electronics.[1] With benefits offered by solution processability, flexibility, reduced toxicity, and ease of tunability, synthesis, and purification, organic electronic materials are promising alternatives to traditional inorganics. [1g, 2] Of particular interest is the application of CT cocrystals as displacive ferroelectrics in data storage. [2c, 3] The exchangeable nature of the CT pair permits rapid screening of various molecular combinations relative to single-component materials; this is advantageous in the field of ferroelectric materials, which show and statistically disfavored crystallographic requirements. Taking advantage of this design feature, several organic CT ferroelectrics have been reported, [1d, 1e, 2, 4] although most of materials that have been discovered to date only exhibit ferroelectricity below room temperature.

Conventional design strategies towards displacive ferroelectric cocrystals focus on tailoring the CT gap in 1D chains of alternating π -electron donor (D) and acceptor (A) molecules to achieve a transition between neutral (DADA...) and ionic ($D^+A^-D^+A^-...$) solid phases (NI transition)^[5] through a reversible Peierls-type lattice instability at the Curie temperature, T_C . [1b-e, 1i, 2-4, 6] The Peierls distortion can manifest as a molecular

deformation (Figure 1a) or displacement (Figure 1b) that polarizes the CT chain in the ionic/ferroelectric phase. [1e, 3] However, design strategies that leverage electronic instabilities modeled in 1D CT chains are limited in their ability to predict interchain interactions that may prevent structural transitions. We argue that the paucity of reported cocrystals that exhibit room temperature displacive ferroelectricity is due, in part, to design strategies that consider CT interactions as the driving force for the paraelectric-ferroelectric transition, while assuming that interchain interactions are non-participatory.

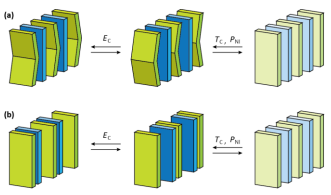


Figure 1. Two mechanisms of displacive ferroelectric transitioning in CT cocrystals: (a) molecular deformation and (b) molecular displacement (adapted from Horiuchi *et al.*)^[1e]. The polarization direction can be switched under a coercive electric field, E_C , and the ferroelectric-paraelectric transition can be thermally induced at the T_C or the NI transition pressure, P_{NI} .

Herein, a room-temperature displacive ferroelectric CT cocrystal formed between acenaphthene (AN) and 2,3,5,6tetrafluoro-7,7,8,8-tetracyanoguinodimethane (F₄TCNQ) presented. A record high T_C of 68 °C allows room temperature ferroelectricity in AN-F₄TCNQ; the T_C corresponds to an activated by rotational motion in AN that decouples AN-F4TCNQ dimers. Weak electronic coupling between AN and F4TCNQ, as measured by Raman spectroscopy, supports the claim that the mechanism of the ferroelectric-paraelectric transition is primarily structural. No formal NI transition occurs at the T_C although this material displays switchable remanent polarization, prompting reevaluation of conventional redox potential-focused design strategies. AN-F₄TCNQ highlights the importance of interchain interactions and molecular dynamics in the design of CT cocrystals that undergo ferroelectric transitions and provides critical insights to improve the diversity and reliability of subsequent ferroelectric design strategies.

AN and F_4TCNQ (Figure 2a) were selected as cocrystallization partners because they display a minimal calculated difference in position of the HOMO and LUMO energy levels for isolated gas-phase D and A molecules. This approach is consistent with current strategies towards CT cocrystal

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design. $^{[1f,\ 1h]}$ Cocrystals of AN-F₄TCNQ were grown by solvent evaporation from an equimolar acetonitrile solution (see Supporting Information, SI). Thermal cycling by differential scanning calorimetry (DSC) revealed a reversible endotherm upon heating at 341.1 K (68.1 °C) and an exotherm at 335.0 K (62.0 °C) upon cooling. Single-crystal XRD (SCXRD) structures were obtained at 85 K, 293 K (20 °C), and 348 K (75 °C). At both 85 K and 293 K, the cocrystal solves in the polar Pc space group, whereas the 348 K crystal structure becomes centrosymmetric (P2₁/c). AN-F₄TCNQ adopts a mixed-stack motif parallel to the a-axis (Figure 2c). Interplanar spacings reveal that the molecules are dimerized $(D^+A^- D^+A^-...)$ in the two low temperature structures with alternating long and short π -stacking distances (3.415 Å and 3.369 Å at 293 K measured D centroid to A plane), while the high-temperature crystal structure is differentiated by a decoupling of the DA dimers, evenly spacing the D and A molecules (3.461 Å), and a 180° rotational disorder of AN. Despite this rotational disorder, 3D crystallinity is maintained in the $P2_1/c$ phase with close interchain contacts formed between C-F and C=N on neighboring F4TCNQ molecules (Figure 2d). These contacts are present in the Pc (at 293 K) and $P2_1/c$ phases at distances between 3.05 and 3.07 Å... Furthermore, the interchain contacts are strengthened through C≡N···H-C interactions^[7] that are present in all three crystal structures at an interaction distance of 2.46 Å at 293 K (normalized hydrogen positions, Figure 2e).

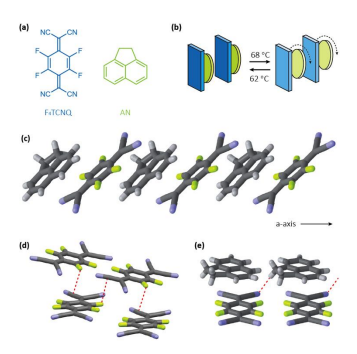


Figure 2. (a) Chemical structures of F₄TCNQ (blue) and AN (green); (b) illustration summarizing the NI transition between the low-temperature Pc (left) and high-temperature $P2_1/c$ (right) phases with the dashed arrow representing the high temperature rotational disorder; (c) 1D chains of alternating AN and F₄TCNQ molecules along the a-axis from the room temperature crystal structure; (d) chains of C-F····C=N interactions formed between neighboring F₄TCNQ molecules; (e) view highlighting the C=N···H-C interactions.

Given that the phase change in AN-F₄TCNQ is accompanied by a reversible symmetry change (noncentrosymmetric to

centrosymmetric), the room-temperature ferroelectricity was investigated by measuring polarization hysteresis parallel to the CT stacking direction (a-axis) on single crystals (see SI). Polarization hysteresis loops measured directly show dielectric leakage that becomes more dramatic upon scanning to lower frequencies. Positive-up-negative-down (PUND) measurements reveal that two components (dielectric and ferroelectric) contribute to the overall polarization intensity and, at a frequency optimized for the ferroelectric component, the dielectric component contributes to 95% of the overall intensity and the ferroelectric component the other 5% (see SI). The dielectric component is likely associated with rotation of AN under an applied electric field due to its dipole moment, which is consistent with the increased motion of AN observed in the crystal structure of the high-temperature P2₁/c phase. Remanent polarization hysteresis loops were measured using a double triangular waveform voltage to remove the effects of nonhysteresis character (see SI). Room temperature ferroelectricity in AN-F₄TCNQ was confirmed and the remanent polarization (Pr) was measured up to 0.08 \pm 0.01 nC/cm² (see SI). By comparison to the model displacive ferroelectric formed between tetrathiafulvalene (TTF) and p-chloranil (CA), the Pr of AN-F₄TCNQ is small: however, AN-F₄TCNQ represents remarkable progress for displacive ferroelectric cocrystals as it displays ferroelectric behavior above room temperature, while TTF-CA is only ferroelectric below 81 K. Electronic coupling between AN and F4TCNQ was further characterized by calculating the ionicity (ρ) of the cocrystal during the ferroelectric-paraelectric transition. Ionicity values were calculated from Raman spectra[1h, 8] obtained between 30 °C and 80 °C, revealing a minor shift in ionicity between 65 and 70 °C associated with the transition (see SI). The ρ of the ionic phase was calculated at 0.3 e, which decreases to 0.2 e upon transitioning to the $P2_1/c$ phase. The $\Delta \rho$ for this material is 0.1 e (about one fourth that of TTF-CA), implicating that a true NI transition[8b] is not crossed.

By assuming a double-well potential model for a ferroelectric transition, [9] the $T_{\rm C}$ can be used to estimate the energy difference between the ferroelectric and paraelectric states. Participation of non-CT interactions altered during the transition can be probed by comparing the $T_{\rm C}$ of non-deuterated and perdeuterated analogues of ferroelectric materials. For example, TTF-CA exhibits a $T_{\rm C}$ 2-5 K lower than d_4 TTF-CA, [1e, 10] which was attributed to lengthening, destabilization, and symmetrization of interchain C-H···O hydrogen bonds upon transitioning to the paraelectric (neutral) phase. [11]

Investigation into the ferroelectric-paraelectric transition mechanism for AN-F₄TCNQ began with synthesis of d_{10} AN-F₄TCNQ. The DSC trace revealed a reversible endotherm at 69.2 °C upon heating and an exotherm at 62.0 °C upon cooling. The 1.1 °C increase in $T_{\rm C}$ upon deuteration is in the opposite direction of what would be expected for a change in redox potential due to the reduced hyperconjugative stabilization afforded by deuterium. This observation indicates that non-CT interactions contribute to the ferroelectric-paraelectric transition mechanism (*vide infra*).

Examples of flip-flop^[12a] and plastic^[12b] ferroelectricity in CT metal-organic systems and antiferroelectric-paraelectric transitions in organic crystals^[12c] have suggested the possibility

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of forming ferroelectrics in which structural transitions are controlled by activated vibrational and/or rotational modes. Such a mechanism would be consistent with the AN motion identified by SCXRD and dielectric leakage observed during polarization measurements. Evidence of dynamic molecular rotation was collected by solid-state ¹H magic-angle-spinning nuclear magnetic resonance (MAS-NMR) spectroscopy between 30 and 80 °C. Above 60 °C, the aromatic and non-aromatic proton environments in AN can be differentiated in the 1D 1H spectra (Figure 3a); however, with a decrease in temperature, the methylene signal shifts downfield, appearing as a shoulder on the aromatic peak, while the aromatic signal shows progressive broadening upon cooling. The sharpening upon heating from 30 to 60 °C suggests that the transition is preceded by continuous structural changes and confirms that the AN rotational disorder observed by SCXRD is dynamic. The ¹H T₁ relaxation data collected between 30 and 80 °C is most suitably fit with a bicomponent T₁ curve (see SI). Both components show a decrease in relaxation time upon heating to 60 °C, which is likely a product of two effects: a decrease in CT and interruption of C≡N···H-C interactions due to rotational motion. At temperatures beyond 60 °C, an increase in the T₁ is measured, caused by a freeing of AN motion, pushing the dynamics past the T₁ minimum and lengthening the T_1 time. At and above the T_C , this effect becomes more dramatic (Figure 3b).

The MAS-NMR studies prompted a targeted investigation of the C≡N···H-C interaction during the ferroelectric-paraelectric transition. Classical C≡N···H-O H-bonds show -ΔH° between 2-5 kcal mol^{-1 [13]} (likely lower in AN-F₄TCNQ due to the weaker donation), too low of an energetic barrier to independently prevent AN rotational motion at room temperature. The barrier to rotation is likely a convolution of the C≡N···H-C interaction and lattice reorganization enthalpies to accommodate AN rotation. The relative significances of these effects to the structural phase transition were studied by powder-XRD (PXRD) and confirmed by SCXRD (see SI). The 20 shift in the reflections associated with the (010) and (002) interplanar spacings were monitored between 20 °C and 80 °C as these planes describe crystallographic expansions/contractions due to C≡N···H-C interactions and rotation-induced lattice deformation, respectively (Figure 3c). As shown in Figure 3d, the reflection corresponding to the (010) plane does not significantly shift during the transition, indicating that destabilization and elongation of the C≡N···H-C interaction is not a dominant contributor to the structural transition observed by XRD. Contrastingly, the {002} interplanar spacings dramatically expand \sim 5 °C before the $T_{\rm C}$. Activation of AN rotation is consistent with the preferential interplanar expansion between the {002} planes, which would account for additional enthalpic contributions to the barrier preventing ferroelectric-paraelectric transitioning at room temperature.

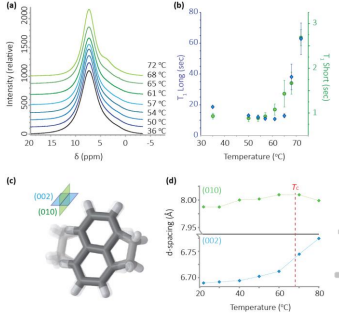


Figure 3. (a) 1D MAS-NMR 1H spectra collected between 30 and 80 $^{\circ}$ C; (b) T_1 relaxation time for the long (blue) and short (green) components as a function of temperature; (c) orientation of AN relative to the (002) and (010) Miller planes; (d) d-spacing expansion between the (010) and (002) Miller planes with increasing temperature.

It has been shown that AN-F4TCNQ undergoes a space group change from Pc to P21/c at 68 °C associated with a decrease in ionicity, consistent with a ferroelectric-paraelectric phase transition, despite weak electronic coupling between AN and F4TCNQ. Activation of AN rotational motion was found to accompany the phase transition as shown by MAS-NMR and PXRD. Reversible remanent polarization of AN-F4TCNQ was measured, confirming that this material is ferroelectric at room temperature although the mechanism of transition is primarily structural. Given these findings, we suggest inclusion of interchain interactions and molecular dynamics that facilitate displacement events responsible for polarization switching in future design of ferroelectric CT cocrystals. Shifting from electronically-driven to structure-driven mechanisms allows a greater diversity of materials that undergo lattice-facilitated ferroelectric-paraelectric transitions to be accessed that may not be predicted by a 1D Peierls distortion model.

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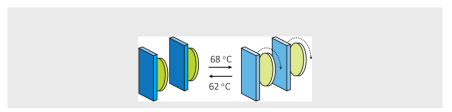
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The organic charge-transfer cocrystal AN-F₄TCNQ shows switchable remanent polarization at room temperature. The ferroelectric Curie temperature was measured at 68 °C by differential scanning calorimetry, prompting further electronic and structural characterization of AN-F₄TCNQ that dynamic motion of AN contributes to the high-temperature polarization switching.

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