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A microspectroscopic insight into the resistivity switching of individual Ag–TCNQ nanocrystals†

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We investigate the resistivity switching in individual Ag–TCNQ wires with on/off-ratios of up to 10³. Raman and soft X-ray absorption microspectroscopy studies disclose reverse charge transfer. Quantification of the fraction of neutral TCNQ within the switched material yields values up to 22.3%. These findings expedite the understanding of the switching process in Ag–TCNQ nanowires.

(7,7',8,8')-Tetracyanoquinodimethane (TCNQ) is a widely-known organic electron acceptor and readily forms charge transfer complexes with a wide range of metals, such as potassium, nickel, copper and silver. The molecule is thereby reduced to the stable radical anion and forms needle-shaped crystallites with the positively charged metal ion.^{1–6} Cu–TCNQ and Ag–TCNQ are the most prominent representatives of this class of materials: both materials have been intensely studied due to their exceptional electrical, magnetic and optical properties. Among these, the electrically bistable resistivity switching is of specific interest in molecule-based electronics, as the intrinsic resistivity of the naturally semi-conducting materials can be electrically switched at a certain threshold voltage.^{3,7–11} This renders TCNQ charge transfer salts promising candidates for storage devices.^{9,12–14} The resistivity switching is linked to a structural phase transition which originates from the reverse electron transfer partly forming neutral Ag⁰ atoms and TCNQ⁰ molecules, which act as dopants in the charge transfer salt according to eqn (1).^{7,9}



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For the implementation of Ag–TCNQ in device fabrication, it is crucial to connect and address individual crystallites on the nanoscale. Controlling the resistivity switching Ag–TCNQ nanocrystals effectively requires comprehensive understanding of the underlying mechanism. To date, the switching behaviour has only been investigated either in bulk materials or in thin films using spectroscopic methods. For instance, evidence for the coexistence of TCNQ anions and neutral TCNQ molecules has been found by Raman spectroscopy of Cu–TCNQ and Ag–TCNQ films and by X-ray diffraction showing two different phases of Ag–TCNQ crystals.^{3,8,12} These investigations are methodically limited to the bulk phase instead of locally probing individual crystals and their electronic structure. However, it is well known that physical processes on small scales do not essentially reflect those in the bulk materials due to boundary conditions.^{15,16} Individual Ag–TCNQ crystallites have been merely characterized electrically and imaged independently using scanning and transmission electron microscopy (TEM).^{9,12,17–19}

Within this work, we report on the resistivity switching and the underlying charge transfer mechanism in individual Ag–TCNQ crystallites in detail. Two complementary spectroscopic nanoprobbers, *i.e.*, near-edge X-ray absorption fine structure (NEXAFS) microspectroscopy and confocal Raman microspectroscopy, are employed to monitor changes in the spectroscopic fingerprints *in operando*. The dissimilar density of unoccupied states of the neutral TCNQ molecule and the respective radical anion thereby enables stoichiometric quantification of the reverse charge transfer upon switching within individual nanowires.

Results and discussion

Electrical characterization of individual Ag–TCNQ nanowires

There are basically two ways to prepare Ag–TCNQ crystallites which can be individually addressed using a microscope, namely the wet-chemical synthesis^{4,20} and a reaction from the vapour phase.²¹ Length and diameter distributions as well as morphologies and orientation can be tailored by varying the



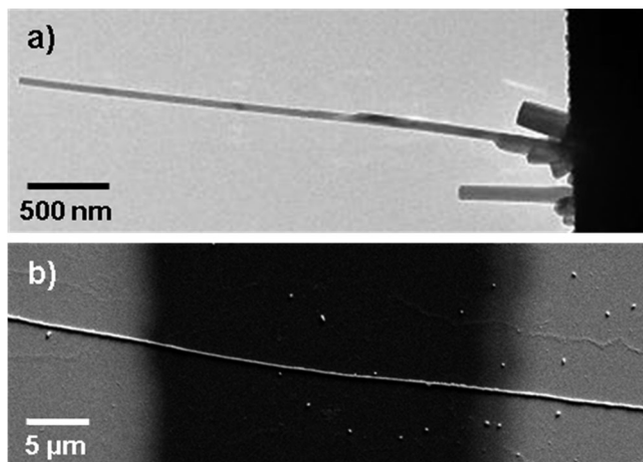


Fig. 1 Individual Ag–TCNQ wires for microscopic and spectroscopic investigations *in operando*. (a) Transmission electron micrograph of an Ag–TCNQ nanowire ($d = 41$ nm) on a gold support, which can be contacted with a tungsten tip in the TEM. (b) Scanning electron micrograph of a crystallite ($d = 0.35$ μm) contacted with gold electrodes (the brighter areas on both sides).

experimental parameters (see the ESI† for a detailed description of the sample preparation). Fig. 1a shows a TEM image of an upright standing nanowire on a gold substrate prepared *via* the gas phase reaction. The wires prepared using this method can be contacted by the tip of a scanning tunnelling microscope and electrically characterized inside the TEM. Fig. 1b shows an Ag–TCNQ wire prepared in solution and subsequently contacted with two gold pads as electrodes. These crystallites are larger in diameter (up to several micrometres) but exhibit comparable switching behaviour, and are thus utilized as model systems for Raman and NEXAFS spectroscopy studies in this work.

Fig. 2 shows the typical I – V characteristics of an individual Ag–TCNQ wire contacted inside the TEM indicating resistivity switching. Upon sweeping the voltage from 0 V to -10 V, the resistivity changes from a high resistance state to a low resistance state below -8 V. The subsequent, reverse voltage sweep shows a hysteresis-type behaviour, maintaining the low resistance state. Sweeping further to positive values, the current increases again, yet is more restrained than in the negative regime until it reaches comparable currents for forward and reverse sweeps at approx. 5.5 V. Upon taking a closer look to the on/off-ratios, the switching is evident. From negative voltages up to +2 V, the on/off-ratio well exceeds 10^2 (maximum value at -0.3 V: 1.15×10^3), whereas it drops to values below 10 above 3.5 V. Thus, the on-state is induced by the negative voltage, and the material is brought back to its off-state by application of the positive voltage. The current is thereby limited to 10^{-7} A in order to protect the crystal from electrical breakdown (see below). Note that especially contributions from contact resistance in the experimental setup decrease the effective potential at the crystal itself; absolute values for the applied voltage have thus to be considered with care.

Probing the reverse charge transfer using Raman spectroscopy

A fundamental measure to probe and distinguish the state of TCNQ anions and neutral TCNQ molecules is their vibrational

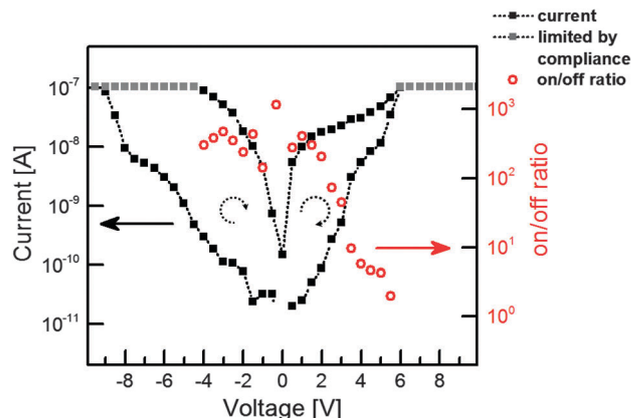


Fig. 2 Current–voltage characteristics of an individual Ag–TCNQ crystallite. Negative and positive dual voltage sweeps have been conducted: 0 V \rightarrow -10 V \rightarrow 10 V \rightarrow 0 V (black, squares). The on/off-ratios well exceed 10^2 (red, open circles). The on/off-ratios in the regime with limited current flow by the chosen compliance value (grey squares) cannot be determined and are therefore not shown.

structure (see the ESI† for reference spectra from the pristine Ag–TCNQ and the neutral TCNQ species). Raman spectroscopy and theoretical considerations on TCNQ, Na–TCNQ and Cu–TCNQ^{3,22,23} allow us to assign the bands in the TCNQ and Ag–TCNQ spectra to distinct features. The bands at 1208 rel. cm^{-1} and 1605 rel. cm^{-1} represent vibrational modes within the TCNQ molecule, which are barely affected by the reduction to TCNQ⁻ (ν_3 and ν_5 modes, a_g symmetry).²³ The TCNQ⁰ vibration at 1457 rel. cm^{-1} (C=C stretching mode, ν_4 , a_g symmetry) is red-shifted to 1388 rel. cm^{-1} by the uptake of an electron into the lowest unoccupied (π^*)-molecular orbital (LUMO), weakening the respective bonds due to antibonding effects. The electron is hereby located at one of the carbon atoms between the cyan groups and the quinone ring.²⁴

Thus, we recorded the Raman spectra of individual Ag–TCNQ wires with various voltages applied to track the reverse charge transfer qualitatively (see Fig. 3). Whereas the spectral fingerprint

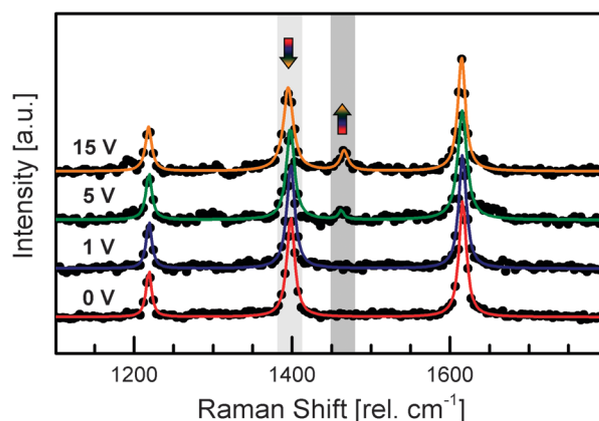


Fig. 3 Reverse charge transfer probed using Raman spectroscopy. Raman spectra of an individual Ag–TCNQ wire under operation. With an increase in negative voltage (absolute values shown), the typical band for neutral TCNQ arises at 1457 rel. cm^{-1} whereas the vibration of the negative molecule at 1388 rel. cm^{-1} is diminished.



remains unchanged at 1 V, the vibrational band at $1457 \text{ rel. cm}^{-1}$, typical for neutral TCNQ, starts to increase at 5 V and appears more pronounced at 15 V, whereas the typical vibrational mode for the negatively charged molecule at $1388 \text{ rel. cm}^{-1}$ decreases. This change is a direct spectroscopic proof that reverse charge transfer takes place when the resistivity state of an individual Ag-TCNQ wire is switched. The evolution of the characteristic band of the neutral TCNQ during voltage-induced resistivity switching demonstrates that a remarkable amount of the materials undergoes reverse charge transfer, forming the neutral species within the crystallite. However, an absolute quantification of affected molecules based on Raman spectroscopy is problematic even with the clear shift observed, due to dissimilar polarizability tensors of neutral and negatively charged molecules.²⁵

Quantification by NEXAFS spectroscopy

Complementary to probing the vibrational structure, NEXAFS spectroscopy provides a powerful tool for investigating the chemical state of TCNQ. Both species, Ag-TCNQ and neutral TCNQ, are oriented equally with respect to the X-ray polarization to exclude effects which arise from linear X-ray absorption dichroism. Whereas Raman spectroscopy discloses a specific feature disparate for TCNQ^- and TCNQ^0 , both the NEXAFS spectra of the reference materials show several strong and overlapping C $1s \rightarrow \pi^*$ absorption bands between 283 eV and 289 eV (Fig. 4a). The intensities and energy positions of the peaks show sufficient differences to distinguish the electronic states of TCNQ, especially the resonances of neutral TCNQ at 288.0 eV and 289.6 eV.^{26,27}

The neutral TCNQ fraction in the Ag-TCNQ wire was derived as a function of applied voltage by linear combination (Fig. 4b, see ESI†). In this particular experiment, we obtain an increase in the TCNQ^0 fractions up to $15.5 \pm 4.9\%$ (7.5 V, Fig. 4c). Most interestingly, the neutral TCNQ fraction remains in this range when the voltage is switched off ($12.3 \pm 4.6\%$) for five minutes, and increases further after 10 V has been applied for 1 min ($22.3 \pm 9.0\%$). Thus, NEXAFS confirms the findings from Raman spectroscopy that resistivity switching is accompanied by reverse charge transfer from the radical TCNQ anion to silver. Moreover, the quantification of this effect becomes possible using thickness-normalized spectra, yet with non-negligible uncertainty. Minor geometric modifications in the TCNQ moieties upon switching affect the absorption probability, which we consider in the experimental error. Nevertheless, the high percentages exceed those of typical dopants in semiconductors by far, indicating that the mechanism for resistivity switching is not a pure doping effect, but is more likely related to the accompanying structural rearrangement with inherent changes in the band structure.²⁸ This structural change remains, however, concealed due to damage by electrons (in TEM) preventing a detailed electron diffraction analysis in the switched state.

Electrical breakdown and failure mechanism

High current densities lead to electrical breakdown of the nanowires. Typical currents exceed 10^{-7} to 10^{-6} A when the failure sets in. Substantial thinning and mechanical rupture of the Ag-TCNQ wires upon this process are revealed using TEM

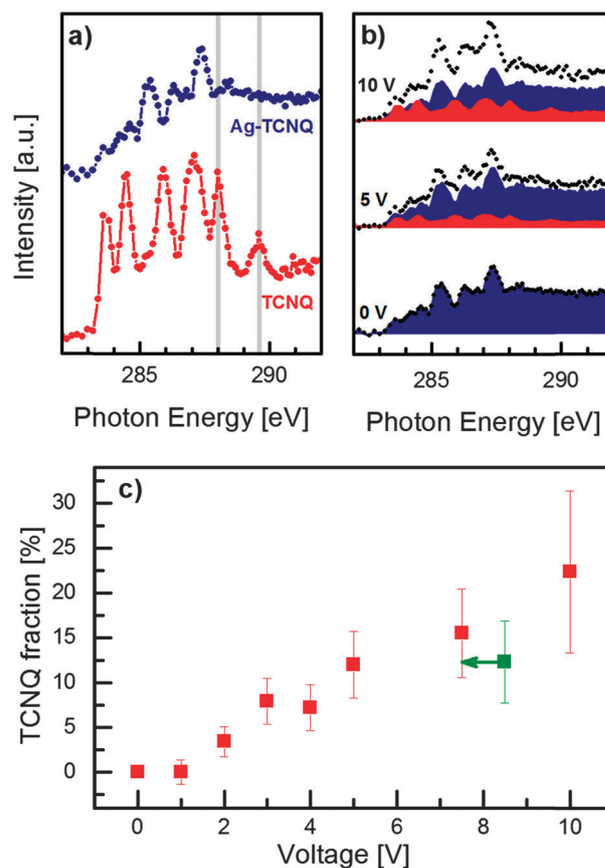


Fig. 4 Quantification of the reverse charge transfer using NEXAFS spectroscopy. (a) Reference C K-edge NEXAFS spectra of TCNQ (red) and Ag-TCNQ (blue) recorded from the respective microcrystals. The typical resonances for TCNQ at 288.0 eV and 289.6 eV are indicated in grey. (b) Exemplary NEXAFS spectra of Ag-TCNQ (shown in Fig. 1b), (black dots) with different voltages applied. The underlying areas show the deconvolution of the Ag-TCNQ (blue) and neutral TCNQ contributions (red). (c) Fraction of neutral species (TCNQ^0) as derived from spectral deconvolution. Note that the green data point corresponds to 7.5 V after five minutes of relaxation.

(see ESI†). We also observe the decomposition of the Ag-TCNQ crystallites upon too intense laser irradiation, accompanied by condensation of the neutral TCNQ around the crystal (see ESI†). These findings can be attributed to an excessive temperature increase by current-induced/resistive heating. With the reverse charge transfer during switching in the formation of the neutral TCNQ species, these extremely volatile molecules²⁹ sublime and then condense in close vicinity of the crystallite, which leads to the observed thinning of the crystallites and finally to electrical breakdown. The experiments have therefore been performed with great care. The microspectroscopic imaging techniques employed here offer immediate inspection and control.

Conclusions

The ability to contact and switch Ag-TCNQ crystallites with diameters down to tens of nanometres is a key step in advancing towards the implementation of this material into storage devices.



Within this study, we demonstrate the reversible resistivity switching in individual, contacted Ag–TCNQ wires with on/off-ratios up to 10^3 . The perspective to grow well-defined structures^{13,30} renders the materials appealing for the fabrication of structured arrays of switchable metal–organic nanowires.

The use of microspectroscopic probes with high lateral resolution offers unique insights for investigating the resistivity switching *in operando*. Confocal Raman microscopy reveals the evolution of the vibrational band typical for neutral TCNQ molecules within the metal–organic salt upon voltage application. Essentially, the switching is accompanied by a remarkable reverse charge transfer from TCNQ[−] to the silver cations. These results confirm the mechanistic model that a mixed-valence state of the metal–organic TCNQ salt is an inherent precondition for the transition from the semiconducting high resistance state towards metallic conductivity.²⁸ Moreover, NEXAFS spectroscopy enables the quantification of the neutral fraction as a function of the applied potential, yielding up to 22.3%. These values exceed the typical amounts for dopants in semiconductors by orders of magnitude. In light of these findings, an explanation for enhanced conductivity by structural rearrangements with inherent changes in the band structure²⁸ seems plausible.

The potential benefit of this effect leading to switching of the resistance states is, however, limited by the heating and subsequent decomposition of the active Ag–TCNQ wires. This applies especially to crystallites with poor thermal coupling for sufficient heat dissipation. For the fabrication of respective electronic devices, encapsulation of the electrically active materials in an insulating, thermally conductive matrix to ensure structural stability of the contacted Ag–TCNQ moieties is therefore highly recommended.

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