Spin Crossover in a Vacuum-Deposited Submonolayer of a Molecular Iron(II) Complex

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Supporting Information

ABSTRACT: Spin-state switching of transition-metal complexes (spin crossover) is sensitive to a variety of tiny perturbations. It is often found to be suppressed for molecules directly adsorbed on solid surfaces. We present X-ray absorption spectroscopy measurements of a submonolayer of [FeII(NCS)2L] (L: 1-{6-[1,1-di(pyridin-2-yl)ethyl]-pyridin-2-yl}-N,N-dimethylmethanamine) deposited on a highly oriented pyrolytic graphite substrate in ultrahigh vacuum. These molecules undergo a thermally induced, fully reversible, gradual spin crossover with a transition temperature of $T_{1/2} = 235(6)$ K and a transition width of $\Delta T_{10} = 115(8)$ K. Our results show that by using a carbon-based substrate the spin-crossover behavior can be preserved even for molecules that are in direct contact with a solid surface.

SECTION: Physical Processes in Nanomaterials and Nanostructures

Since its first observation in 1931 and rationalization using ligand-field theory in the 1950s, the spin-crossover (SCO) phenomenon has been intensely researched and discussed, particularly with regard to potential applications in information storage and processing. Most of the known SCO compounds are transition-metal complexes, especially 3d4–3d5 metal ions in pseudo-octahedral coordination, with a predominance of iron(II) in N6 environments. Among these, (pyridine)bis(thiocyanato-κN)iron(II) complexes constitute an exceptionally well-studied class. Assemblies of such molecules have been synthesized and characterized in bulk, nanoscale, and surface phases. Typical methods for preparing the latter comprise Langmuir–Blodgett techniques, spin coating, and vacuum deposition, leading to thin or ultrathin films. Because the spin state of a molecule influences a multitude of its physical and chemical properties, it can be monitored using a wide variety of methods, for example, IR, Mössbauer, NMR, Raman, UV/vis, and X-ray absorption spectroscopies (XAS), conductometry, dielectrometry, diffractometry, refractometry, and susceptometry.

Recently, an electron-induced spin-state switching of [FeII(bpb)2(phen)] (bpb: bis(1H-pyrazol-1-yl)borate, phen: 1,10-phenanthroline) in the second molecular layer deposited on Au(111) in ultrahigh vacuum (UHV) was observed by differential tunneling spectroscopy at 5 K, but the molecules of the first layer could not be switched. The remarkably high sensitivity to the environment, which is typical of SCO compounds, often suppresses the switching behavior of molecules in direct contact with a substrate. Similarly, [FeII(NCS)2(phen)2] on Cu(100) does not show an SCO transition in the first molecular layer, whereas isolated [FeII(NCS)2(phen)2] molecules on an interfacial layer of CuN on Cu(100) may be switched between their high-spin (HS) and low-spin (LS) state by the tunneling current at 4.6 K. Oligonuclear “beads” in chains of [FeII(L′)2](BF4)2 (L′: 2,6-di(1H-pyrazol-1-yl)-4-(thiocyanatomethyl)pyridine), after self-assembly on highly oriented pyrolytic graphite (HOPG) from acetonitrile solution, do show spin-state interconversion, as detected by current-imaging tunneling spectroscopy (CITS).

Here we report on the thermally induced, reversible spin-state switching of submonolayers of the molecular complex [FeII(NCS)2,L] (L: 1-{6-[1,1-di(pyridin-2-yl)ethyl]-pyridin-2-yl}-N,N-dimethylmethanamine) obtained by vacuum deposition onto an HOPG surface. Complex 1 has a novel N4 chelate ligand designed to contain both imine and aliphatic amine N donors and two thiocyanato-κN ligands. HOPG was chosen as substrate because it promises only weak interaction, possibly leaving the molecular properties and thus the SCO behavior largely unaffected.

Submonolayer coverages of 1 were obtained by thermal deposition at ~510 K from a tantalum Knudsen cell onto an HOPG substrate under UHV conditions and measured by X-
ray absorption spectroscopy (XAS). To characterize the adsorption of 1 on the HOPG surface, we have performed low-energy electron diffraction (LEED) experiments. Figure 2 shows LEED images of the clean HOPG surface (a), 1.2 monolayer (ML) of 1 on HOPG directly (b) and after 10 min of measuring with the electron beam at the same position (c). The ring-shaped diffracted electron intensity in panel a demonstrates that the graphite surface is highly oriented, while containing crystallites of different azimuthal orientation. The disappearance of the diffraction pattern in panel b proves that the molecules wet the surface and form a molecular layer because the electrons diffracted by the HOPG substrate are strongly damped by the adsorbate. If 1 would form crystallites, then most parts of the surface would remain uncovered, and the ring-shaped diffraction pattern would be only slightly damped. Even though we cannot rule out that a minor part of the molecules may already occupy second monolayer positions while the first monolayer is not yet complete, we can conclude that 1 grows layerwise on HOPG and that no crystallites are formed. The partial reappearance of the substrate diffraction pattern in panel c is interpreted as the result of electron beam damage and subsequent partial desorption of fragments of the molecules.

Fe-L_{2,3} XA spectra give direct access to detailed information on the electronic structure of the iron(II) ions, allowing for straightforward identification of the HS and LS state and their interconversion with temperature. Figure 3 shows the isotropic Fe-L_{2,3} X-ray absorption spectra of 0.8 ML of 1 on HOPG measured at 300 (black) and 75 K (red). Strong temperature-dependent changes of the Fe XAS signal are evident. At room temperature, the Fe-L_{3} edge displays a double-peak feature at photon energies of 707.8 and 708.5 eV. At 75 K, the Fe-L_{3} edge is shifted by ~1.1 eV to higher photon energy and slightly compressed. These changes are accompanied by a change of the relative intensity of the Fe-L_{3} (705 to 715 eV) and L_{2} (718 to 726 eV) edge. The branching ratio, defined as the ratio of the integrated L_{3} intensity and the intensities of the L_{1} and the L_{2} edges taken together, is reduced from 0.74(2) at 300 K to 0.64(2) at 75 K. Such a reduction is typical of an HS-to-LS transition and reflects the decrease in 3d spin–orbit coupling energy due to the vanishing magnetic moment in the LS state.\textsuperscript{17,18} The shapes of the Fe-L_{2,3} XA spectra of both the HS and the LS state closely resemble those of [Fe\textsuperscript{II}(NCS)\textsubscript{2}(phen)]\textsubscript{2},\textsuperscript{19} indicating that the ligand field of the Fe center is very similar for the two systems. The spin transition of 0.8 ML of 1 on HOPG appears to be nearly complete at low temperatures, and the spectra at 300 and 75 K represent almost pure HS and LS states, respectively. The isotropic Fe-L_{3} XA spectra shown in Figure S1 of the Supporting Information were recorded during heating the sample from 75 to 300 K. The spectra display a gradual transition between the two spectra presented in Figure 3. Figure 4 shows the fraction of HS molecules plotted over the temperature. For the determination of the HS fraction, each Fe-L_{3} XA spectrum was fitted as a linear combination of the spectrum at 300 and at 75 K. Because the temperature-dependent spectral change is saturating at low temperatures, it can be assumed that the spectrum at 75 K represents a pure LS spin state. At 300 K, the transition to the HS state is not complete. However, the maximal LS contribution can be estimated by subtracting fractions of the spectrum at 75 K shown in Figure 2 from the one at 300 K and must be smaller than 20% because the resulting intensities cannot be smaller.
formation. As expected for an SCO transition, the spin of the Fe centers of 1 on HOPG can be switched repeatedly and in a fully reversible manner between HS and LS by altering the temperature (Figure S3 of the Supporting Information).

In conclusion, a thermally induced, reversible SCO transition has been demonstrated for a submonolayer of [FeII(NCS)2L] molecules thermally deposited in UHV on an HOPG surface. The HS and LS spin states could be clearly identified by means of XAS at the Fe-L2,3 edges. The SCO transition is nearly complete and shows a very gradual behavior between 90 and 300 K, indicating that the transition acts on the single-molecule level with a low degree of cooperativity. Our result may serve as proof of principle that the quenching of switching of surface-mounted molecular units can be overcome by using carbon-based surfaces. This opens up new possibilities to realize the vision of highly sensitive SCO molecules serving as tools for creating molecular spintronic devices on surfaces. It remains interesting to see if unquenched switching behavior can also be found on graphene surfaces.

### EXPERIMENTAL SECTION

Ligand L was synthesized by Eschweiler–Clarke methylation of the corresponding primary amine, \{6-[1,1-di(pyridin-2-yl)-ethyl][pyridin-2-yl]methylamine,22 following a standard literature procedure.23 It was purified by column chromatography and subsequently reacted with [Fe(NCS)2(py)4]24 (py: pyridine) in methanol to give pure [Fe(NCS)2L] (1) in a yield of 73% (see Supporting Information). Solid-state variable-temperature magnetic susceptibility measurements were performed using a Quantum Design MPMS-XLS SQUID magnetometer operating at 0.1 T. Diamagnetic corrections for the sample and the sample holder were applied.

LEED images were taken using an Omicron Spectaeed system with a spot size of the electron beam of ∼0.5 mm in diameter and a sample current of ∼0.8 μA during the measurement. We purchased 10 × 10 × 2 mm3 SPI-1 high-grade HOPG substrates exhibiting a mosaic angle of 0.4 ± 0.1° from Structure Probe. A clean HOPG surface was obtained by cleaving an HOPG substrate at a pressure of 10−6 mbar by means of a carbon tape. The quality of the HOPG surface was checked by the angle dependence of the C π* resonance at 285.4 eV. Successful preparations typically showed a ratio of 1:100 between the resonance intensity for vertically s- and horizontally p-polarized X-rays at 20° grazing incidence. Cleavage was repeated if the ratio was larger. I was deposited at ∼510 K from a tantalum Knudsen cell at 5 × 10−9 mbar onto the substrate held at room temperature. Coverages were determined by using a quartz microbalance and the absolute Fe-L2,3 XAS intensity. After deposition, the sample was heated to 350 K to desorb volatile nitrogen-containing fragments produced during the deposition process. These fragments are likely L, which decoordinates at 460 K in high vacuum (Figure S4 of the Supporting Information). After desorption, only intact molecules of compound 1 remain on the surface, as concluded from the observation that the spin transition is complete at low temperatures. XAS measurements were performed at the beamline UE56/2-PGM1 at BESSY II at a pressure of 5 × 10−10 mbar. Fe-L2,3 isotropic XA spectra were recorded at the magic angle (54.7°) between the X-ray wave vector and the surface using linearly p-polarized X-rays. At this angle, the XAS resonance intensities are independent of the orientations of the molecular orbitals.25 The energy resolution was set to 300 meV at a photon flux of ∼1013 photons s−1 cm−2.

![Figure 4](image-url) Fraction of HS molecules (squares) as a function of temperature and fit of a model based on noninteracting molecules.
XA spectra were acquired in total-electron-yield mode by recording the sample drift current as a function of photon energy. The XAS signal of the sample was normalized to the one of a gold grid upstream to the experiment that was recorded in parallel. Subsequently, the spectra were normalized to the substrate signal approximated by a linear function. Time-dependent X-ray-induced irreversible modifications of the Fe-L edge structure lead to apparent reduction of the HS contribution of 20% within 1 h of continuous illumination at room temperature with three times the flux density used for recording the spectra (Figure S5 of the Supporting Information). Consequently, the measuring time for each spectrum was limited to 3 min before a new position on the sample was chosen. On this time scale, the spectra at room temperature and at low temperatures showed no time-dependent variations so that chemical modifications of the molecules as well as an X-ray-induced formation of crystallites can be ruled out.

**REFERENCES**


