Detailed Characterization of a Nanosecond-Lived Excited State: X-ray and Theoretical Investigation of the Quintet State in Photoexcited [Fe(terpy)(2)](2+)

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Published in:
The Journal of Physical Chemistry Part C

Link to article, DOI:
10.1021/acs.jpcc.5b00557

Publication date:
2015

Document Version
Publisher's PDF, also known as Version of record

Link back to DTU Orbit

Citation (APA):
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ABSTRACT: Theoretical predictions show that depending on the populations of the Fe 3d_{xy}, 3d_{xz}, and 3d_{yz} orbitals two possible quintet states can exist for the high-spin state of the photoswitchable model system [Fe(terpy)2]2+. The differences in the structure and molecular properties of these 5B2 and 5E quintets are very small and pose a substantial challenge for experiments to resolve them. Yet for a better understanding of the physics of this system, which can lead to the design of novel molecules with enhanced photoswitching performance, it is vital to determine which high-spin state is reached in the transitions that follow the light excitation. The quintet state can be prepared with a short laser pulse and can be studied with cutting-edge time-resolved X-ray techniques. Here we report on the application of an extended set of X-ray spectroscopy and scattering techniques applied to investigate the quintet state of [Fe(terpy)2]2+ 80 ps after light excitation. High-quality X-ray absorption, nonresonant emission, and resonant emission spectra as well as X-ray diffuse scattering data clearly reflect the formation of the high-spin state of the [Fe(terpy)2]2+ molecule; moreover, extended X-ray absorption fine structure spectroscopy resolves the Fe–ligand bond-length variations with unprecedented bond-length accuracy in time-resolved experiments. With ab initio calculations we determine why, in contrast to most related systems, one configurational mode is insufficient for the description of the low-spin (LS)—high-spin (HS) transition. We identify the electronic structure origin of the differences between the two possible quintet modes, and finally, we unambiguously identify the formed quintet state as 5E, in agreement with our theoretical expectations.
Switchable molecular compounds have significant potential as very high-density devices in the areas of data storage systems, molecular switching, and display devices. Promising candidates include transition metal compounds, in particular octahedral FeII complexes, which can exist either in a low-spin (LS) or a high-spin (HS) state, depending on parameters such as temperature or pressure. Light can also induce a spin-state transition in many of these compounds, typically at low temperatures, thus creating a molecular switch in the “on” position. In order to improve on the performance of these systems, it is essential to characterize the HS state thoroughly, particularly in compounds where some sort of anomaly occurs; understanding how we can modify the properties and the stability of the excited HS state can lead to developments toward applications at room temperature.

The full dynamics of the LS–HS switching has long been described by a simple and rather complete theoretical framework, suggested by A. Hauser and co-workers. This theory describes the transitions along a single configurational coordinate (SCC), typically the breathing mode of the molecule (i.e., a symmetric stretching along the Fe–ligand bonds), which was found to be valid for all complexes with monodentate, bidentate, and often even polynuclear ligands. However, experiments and theory suggested that the tridentate [Fe(terpy)2]2+ (terpy: 2,2′,6′,2″-terpyridine) molecule behaves differently in several respects, the most important being that the SCC model for the HS–LS transition fails, and the switching mechanism involves (at least) two vibrational modes (configuration coordinates). In addition to this anomaly, the nature of the HS state was unclear. Due to the axial compression exerted by the terpy ligands, the symmetry is lowered, and thus the octahedral 5T2 is split into two quintet states of different symmetry with terms 3B2 and 3E. These states differ in the configuration of the dxy subshell and the molecular geometry, particularly the Fe–N bond lengths. A very recent extended X-ray absorption fine structure (EXAFS) study made an attempt to determine the structure of the populated quintet state, but instead of resolving the structure it could only address its distortion due to limited signal-to-noise conditions. On the basis of this it was concluded that the structure for the 3E is in somewhat better agreement with the experimental data than that of the 3B2, although this proposed structure contradicts the energetics arising from the density functional theory (DFT) models applied by these authors, which suggest 3B2 as the more stable quintet state.

In this paper, we revisit the electronic and molecular structure of the quintet state of [Fe(terpy)2]2+ in aqueous solution with advanced experimental and theoretical techniques. We investigate the reasons for and the consequences of the failure of the SCC model (and thus the necessity for a description with two modes) in the molecular geometry and electronic structure. By combining X-ray absorption and emission spectroscopies (XAS and XES, respectively) and X-ray diffuse scattering (XDS) in a single pump–probe setup at the megahertz (MHz) repetition rate, we acquire simultaneously time-resolved structural and electronic structural data with unprecedented quality in the picosecond time domain. We prepare the quintet state of [Fe(terpy)2]2+ with intense, 10 ps, green (532 nm) laser pulses leading to metal-to-ligand charge-transfer (MLCT) excitations that subsequently relax into the HS state on a subpicosecond time scale. We exploit XES and XAS X-ray probes to identify the formation of the quintet state and determine its lifetime; we also employ resonant X-ray emission spectroscopy (RXES) at the 1s → 3d excitations to gain further insights into the 3d states. With EXAFS and XDS we determine the structure of the molecule in the quintet state and its surrounding solvent cage. The results are compared to those obtained on photoexcited [Fe(bipy)3]2+, which has been characterized by a wide range of experimental and theoretical techniques and which follows the SCC model (the formation of its photoinduced quintet state was shown to be accompanied by a 0.2 Å elongation of all Fe–N bonds). Moreover, it has served to calibrate coarsely the arsenal of different X-ray tools used in this work. Indeed, optical tools to date suffer from the clear identification of the excited state in such compounds, as this requires an unambiguous observable. The UV–vis range is quite useful in detecting ultrafast changes, but the overlapping bands due to different excited states hampered in most cases a clear identification of the excited-state electronic and (even more) molecular structure. The X-ray tools employed here seek to fill this knowledge gap, and exploiting laser-pump–X-ray-probe techniques at MHz repetition rates allows us to gain new insights into both the electronic and geometric structure changes due to its unprecedented signal quality, as we demonstrate in this work.

## EXPERIMENTAL AND COMPUTATIONAL METHODS

### Experimental Details.

The laser-pump X-ray-probe experiments were performed at sector 7ID-D of the Advanced Photon Source (APS) and at beamline ID26 of the European Synchrotron Radiation Facility (ESRF). Both setups utilized amplified laser systems with adjustable repetition rates in the MHz range, thus increasing the duty cycle of the pump–probe studies by orders of magnitude compared to others that use conventional amplified laser systems with kilohertz (kHz) rates. The experimental strategy combined different time-domain X-ray tools into one single setup, which enabled simultaneous measurements of X-ray diffuse scattering (XDS) together with nonresonant X-ray emission spectroscopy (XES) and also variants of X-ray absorption spectroscopy (XAS): (i) high-energy-resolution fluorescence detection (HERFD) and (ii) total fluorescence yield (TFY) XAS. The former was possible owing to the implementation of a secondary X-ray crystal spectrometer that energy-resolved the emitted fluorescence. This spectrometer also permitted us to record resonant XES (RXES) to better resolve the 1s pre-edge and thus exploit the improved sensitivity to the 3d orbitals (details further below). Time-resolved XDS was recorded in the forward direction up to a Q-value of around 4 Å−1 with a MHz-gateable, single-photon-counting area detector. This strategy of combined tools had been implemented and tested on the photoinduced spin transition of aqueous [Fe(bipy)3]2+, whose investigation is supported by the analysis of the [Fe(bipy)3]2+ data. The experiments at 7ID-D (APS) utilized 24-bunch mode (in top-up mode with a constant 102 mA ring current) with a 6.52 MHz X-ray pulse repetition rate. X-rays were monochromatized with a diamond (111) double-crystal monochromator. An amplified laser system (Duetto from Time-Bandwidth) generating 10 ps pulses at its second harmonic, 532 nm, was used for laser excitation. It was synchronized to the storage ring radio frequency system and operated at a repetition rate of 3.26 MHz, half that of the X-ray rate. The time delay between laser and X-ray pulses could be electronically adjusted in step sizes of 5 ps minimum duration. Details of this time-resolved setup are given

DOI: 10.1021/acs.jpcc.5b00557

in refs 14 and 30. The X-ray beam was focused by a pair of Kirkpatrick–Baez mirrors to 8 × 7 μm², (H × V) on the sample, and the laser beam overlapped this spot with a larger spot size (100 × 80 μm², H × V). The sample consisted of a free-flowing liquid jet of 13 mM aqueous [Fe(terpy)₂]²⁺ solution (total volume 200 mL). The liquid near the nozzle output was a flat sheet, with thickness 0.1 mm, which was oriented at about 45° to both laser and X-ray beams. This geometry permitted simultaneous recording of XDS data (collected in forward scattering direction) and XES data (collected sideways at 90° scattering angle, along the X-ray polarization vector, via the analyzer crystal). For these measurements the incident X-ray probing energy was tuned to 8.2 keV, far above the Fe K edge. The XDS experiments were performed with a Pilatus 100 K detector gated and synchronized to the bunch frequency as described in our previous MHz work. Following masking, azimuthal integration, and scaling, difference signals were constructed and averaged for each time delay and then corrected for multiple-pump–pulse contributions as described in ref 14. Considering the jet speed (6 m/s), the 80 μm spot illuminated by the laser travels only about 2 μm before the next laser pulse arrives 307 ns later. Therefore, it is obvious that the same sample volume is hit many times in such a high repetition rate experiment. Nevertheless, the studied [Fe(terpy)₂]²⁺ complex is very stable, and no degradation was found in any of the signals with exposure. The lifetime of the excited state is 2 orders of magnitude shorter than the separation between the X-ray pulses; therefore, the spectroscopic data are not affected by the previous excitation pulses. On the other hand, the energy transfer to the solvent leads to a temperature increase in the water, which affects the X-ray scattering data. This has been circumvented by using the scattering signal recorded at a Δt = −200 ps time delay as the background. The increased solvent temperature due to the previous laser shots is present in this water signal, and thus it serves as an appropriate reference, as explained in detail in our previous work on [Fe(bipy)₂]³⁺.  

The Kα XES, (1s2p) RXES, and HERFD-XAS data were collected using a 10 cm diameter, spherically bent (R = 1 m), single-crystal Ge(440) wafer analyzer that focused fluorescence onto a MHz-gated scintillator coupled to a photomultiplier. A bent Si(531) crystal with similar dimensions was used to record the Kβ spectra; the energy resolution was 1 eV for both setups. For recording HERFD-XANES (X-ray absorption near-edge structure), the selected emission energy was kept constant (at the maximum of the Kα₁ line), while in the case of RXES both the energy of the incident beam (Ω) and the fluorescence energy (α) were scanned. The recorded RXES intensity is displayed on a 2D map with one axis being the incident energy and the other their indices, which quantify bond strengths utilizing the actual computed wave function of the molecule, and degenerate frequencies were found for the 5B₂ state. Since this could indicate that the obtained 5B₂ structure is a transition state, we performed single-point computations along the potential energy surface, vibrational frequencies were calculated as second derivatives of the electronic energy. They were found to be positive for the 1A₁ and 5E states confirming that these geometries corresponded to true minima. However, two negative and degenerate frequencies were found for the 5B₂ state. Since the 5B₂ geometry also corresponds to a true minimum.

There is a DHXES intensity line 5B₂ with D₂d symmetry is susceptible to the Jahn–Teller (JT) effect. To estimate the contribution of this effect, geometry optimizations were repeated with the BP86/TZP method, by applying D₂d and C₂ point group symmetries with the ADF2013.01 code. (Note that in C₂ symmetry the correct notation for the JT-distorted 5E state would be 5B, as seen in the correlation table in the Supporting Information (SI). However, in order to avoid possible confusion between 5B and 5B₂ term symbols, we decided to keep the 5E notation throughout this paper.) Potential energies of excited states have been calculated using time-dependent DFT (TD-DFT) at the B3LYP/ TZVP level, which was found to perform well for describing spin-state switching for a series of Fe compounds. Since the ability of DFT to provide accurate spin-state energetics of Fe complexes is limited, the 5B₂ and 5E potentials have been evaluated with an ab initio multireference second-order perturbation theory, the CASPT2 method (Complete Active Space Second-Order Perturbation Theory). These computations were performed with the MOLCAS7.6
program package\textsuperscript{43--45} and their detailed description is given elsewhere.\textsuperscript{12} The potentials were calculated along two different lines, one which connects the $^1A_g$ and $^3E$ minima and the other the $^1B_g$ and $^3E$ minima. The $^1B_g$–$^3E$ line is almost orthogonal to the $^1A_g$–$^3E$ line (cf. Figure 9).

A first-principles approach based on DFT in the generalized gradient approximation (GGA) was adopted to model the X-ray absorption near-edge spectra.\textsuperscript{46} The Fe K-edge XANES was calculated in two steps: first the charge density with a 1s core-hole on the absorbing atom was computed, and then the XANES spectrum was calculated for the electric dipole ($1s \rightarrow p$) transitions only, using the broadening parameters given in ref 47. These two steps were performed with the PWscf\textsuperscript{48} and XSpectra\textsuperscript{49} packages, respectively, of the Quantum ESPRESSO suite of codes.\textsuperscript{50} These codes use plane-wave basis sets, pseudopotentials, and periodic boundary conditions; the [Fe(terpy)$_2$]$^{2+}$ molecule was included in a cubic cell large enough to avoid interactions between molecules belonging to neighboring cells.\textsuperscript{51,52} A 110 Ry cutoff energy was used for the plane-wave expansion; the charge density with a core-hole on the iron atom was determined at the Γ point; and a $2 \times 2 \times 2$ Monkhorst–Pack grid was chosen for the calculation of the XAS spectrum of the LS ground state ($3 \times 3 \times 3$ for the photoinduced HS state). Spin-polarized calculations were performed in the case of the photoinduced HS state ($S = 2$). Norm-conserving pseudopotentials\textsuperscript{53} in the Kleinman–Bylander\textsuperscript{54} form were used, and the parameters for their generation are given in ref 52.

The absorption cross-section of RXES is given by the Kramers–Heisenberg equation\textsuperscript{55} and can be simulated using multilet calculations based on the Ligand Field Multiplet (LFM) theory.\textsuperscript{58} The 1s$^2p^5$$^3d^4$ RXES process for [Fe(terpy)$_2$]$^{2+}$ was modeled assuming electric quadrupole transitions from the initial state $1s^2p^53d^4$ to the intermediate state $1s^2p^63d^4$, followed by an electric dipole emission to the final state $1s^2p^53d^4$. The individual absorption and emission transition matrix elements were first calculated using the method developed by Thole,\textsuperscript{56} in the framework established by Cowan\textsuperscript{57} and Butler,\textsuperscript{58} and then combined in a postprocessing step. Details about these calculations can be found in refs 55, 59, 60, and 61. The calculations were performed assuming an octahedral approximation ($D_{6h}$ symmetry), with a crystal field parameter $\Delta_D = 2.0$ eV for the LS state and 1.3 eV for the HS state; the transition lines were then convolved with a Gaussian, accounting for (some of) the experimental broadening ($0.3$ eV) and a Lorentzian accounting for the core-hole lifetime broadening ($1.2$ eV for the intermediate state and $0.2$ eV for the final state). The Slater integrals that describe the electronic interactions were scaled down to 80% of their atomic values to account for covalency.

\section*{RESULTS AND DISCUSSION}

\textbf{Formation and Decay of the Quintet State.} Potential energy surfaces (PESs) are the key to understanding the photochemistry and photochemistry of molecular systems. It has been found recently that the PES of Fe$^{41}$ or Ru$^{41}$ compounds with a MN$_6$ core (M = Fe or Ru) can be described reasonably well using DFT and TD-DFT\textsuperscript{12,62,63} providing a means to qualitatively examine the de-excitation mechanism.\textsuperscript{59} Therefore, we have completed our previous calculations\textsuperscript{12} on [Fe(terpy)$_2$]$^{2+}$ by adding the potential energies for the singlet and triplet metal-to-ligand charge transfer (MLCT) states. The most relevant ones of them are plotted together in Figure 1 with those of the lowest singlet and quintet as well as three of the triplet metal-centered states, along a combined coordinate based on the $R$(Fe–N$_\text{terpy}$) bond length and the NNN angle of the donor atoms of the terpy ligands. This coordinate connects the LS and HS ($^3E$) minima, which was found to be relevant to the description of this system.\textsuperscript{11,12} The potential energy curves show very similar structure and crossings to those of [Fe(bipy)$_3$]$^{2+}$\textsuperscript{12,26} Therefore, similar de-excitation pathways for both complexes after the excitation into the MLCT may be expected. Although the metal-centered quintet ($^3E$) crosses the $^1$MLCT band somewhat close to its minimum, the coupling between these states was found negligible in a recent theoretical work on [Fe(bipy)$_3$]$^{2+}$,\textsuperscript{51} and it was suggested that the system relaxes via the metal-centered triplet states before the intersystem crossing to the quintet.\textsuperscript{26} This theoretical expectation has been proven experimentally very recently by femtosecond-resolved XES.\textsuperscript{27} In [Fe(terpy)$_2$]$^{2+}$ de-excitation via the triplet states seems highly likely given that the potential curve of the $^3B_g$ state crosses the $^3$MLCT band at its minimum. Strong coupling between them would make relaxation to this metal-centered triplet state rapid. Verification of this de-excitation pathway in [Fe(terpy)$_2$]$^{2+}$ requires more involved theoretical or experimental evidence; moreover, one should keep in mind that the true triplet minima do not lie on this combined coordinate, as will be obvious in the final section of this discussion. Nevertheless, based on the mentioned crossing and the almost identical structure of potential energy curves to those of [Fe(bipy)$_3$]$^{2+}$, the system is very likely to follow a decay path that includes triplets. A possible sequence is indicated by the arrows in Figure 1. From the intermediate lifetimes determined for [Fe(bipy)$_3$]$^{2+}$,\textsuperscript{26,27} it is understood that relaxation to the quintet state takes place on the subpicosecond time scale, and based on the similarities and the possibly even better positioned quintet states, one can assume this is true for [Fe(terpy)$_2$]$^{2+}$ as well.

Having argued that the photocycle of [Fe(terpy)$_2$]$^{2+}$ in many respects resembles that of [Fe(bipy)$_3$]$^{2+}$, it is evident that
the spectra to LS/HS references and was found to be photoinduced HS fraction can be determined from XES by comparing the signature to each change from the LS to HS state. In order to verify the formation of the HS species and quantify its yield, XAS and XES spectra of an aqueous solution of [Fe(terpy)$_2$]$^{2+}$ were taken before and 80 ps after light excitation. The TFY-XANES spectra (Figure 2a) show significant light-induced variations, the largest ones being the intensity increase of the 7125 eV B feature and the intensity decrease of the 7142 eV D feature. These changes are similar to those observed for the thoroughly studied [Fe(bipy)$_3$]$^{2+}$ complex, confirming the formation of the photoinduced HS state. In the XES spectra (Figure 2b,c), for both the K$\alpha$ and K$\beta$ emission lines the spectral variations associated with the formation of the photoinduced HS state are evident and fully consistent with previous studies. The variations of both the K$\alpha$ and the K$\beta$ spectra are essentially identical to those of the [Fe(bipy)$_3$]$^{2+}$ complex which is expected since the XES line shapes of 3d transition metal ions reflect practically only the spin state. The first coordination shells of Fe in both complexes are distorted octahedra, and XES spectra are not sensitive to the other relevant differences in symmetry.

The laser ON XAS and XES spectra (Figure 2, red line) represent the laser-excited state of the [Fe(terpy)$_2$]$^{2+}$ complex, which is a combination of both LS and HS contributions. To infer the photoinduced HS fraction (\(\gamma\)), we used an approach that utilizes XES spectral differences between LS and HS references, referred to as the integrated absolute difference (IAD) approach. The typical photoinduced HS fraction observed during our experiments at 80 ps delay was found to be \(\gamma = 40\%\), although at high laser power 60% could also be achieved.

The formation and decay of the photoinduced HS state can be followed by monitoring the intensity of an appropriate spectral or scattering feature at different time delays (\(\Delta t\)). From the XANES of [Fe(terpy)$_2$]$^{2+}$ the intensity of the B feature at 7125 eV (which presents the largest variation at the spin-state transition) was used for this purpose; its time evolution is displayed in Figure 3.

Figure 3. Time evolution of the intensity of the 7125 eV (B) feature of the [Fe(terpy)$_2$]$^{2+}$ XANES spectrum (Figure 2a), reflecting a quintet lifetime of \(\tau_{HS} = 2.61 \pm 0.01\) ns.

The corresponding fit of the kinetics uses a model consisting of an exponential decay and a broadening by a Gaussian pulse, which is described in the SI. The lifetime of the photoinduced HS state is \(\tau_{XAS}^{HS} = 2.61 \pm 0.01\) ns. This value is fully consistent with the lifetime of \(\tau_{II}^{HS} = 2.54 \pm 0.13\) ns obtained with the optical probe for aqueous [Fe(terpy)$_2$]$^{2+}$.

Structure of the Photoinduced Quintet State. Having verified the formation of the HS state and determined its lifetime, we make an attempt at determining its structure. The theoretical molecular geometry is available from DFT calculations and the ground state matches excellently the available experimental data, as can be seen in Table 1. For the possible quintet structures, the DFT models are very similar to each other; their corresponding Fe–N bond lengths differ at the few picometer level. This presents a great challenge to the experimental efforts that aim to determine the structure in order to identify which HS state is populated. The most distinct difference between the possible quintet states is the anisotropic distribution of Fe–N bond lengths in the first coordination shell, which is present in the case of the \(^5E\) quintet (leading to small but appreciable differences between axial and equatorial bonds), whereas it is lifted in the \(^5B_2\) state. As will be shown in the following section, the very high quality of the picosecond EXAFS spectra presented here was decisive to resolve these very small structural
Fe axially compressed. This leads to an anisotropic distribution of theoretical structure of the LS singlet state, the molecule is shorter axial Fe−N bond lengths in the range 0.004 Å and 0.01 Å, in excellent agreement with the DFT-calculated bond lengths listed in Table 1.

Table 1. Most Relevant Structural and Bonding Parameters of the LS and HS States of [Fe(terpy)2]^{2+}

<table>
<thead>
<tr>
<th></th>
<th>LS ((^{1}A_{1}))</th>
<th>HS ((^{3}E))</th>
<th>HS ((^{3}B_{2}))</th>
</tr>
</thead>
<tbody>
<tr>
<td>R(Fe−N(_{ax}))</td>
<td>1.891(5) Å</td>
<td>1.886 Å</td>
<td>2.103 Å</td>
</tr>
<tr>
<td>R(Fe−N(_{eq}))</td>
<td>1.988(10) Å</td>
<td>1.985 Å</td>
<td>2.198 Å</td>
</tr>
<tr>
<td>NNN angle</td>
<td>102.8(3)°</td>
<td>102.6°</td>
<td>108.4°</td>
</tr>
<tr>
<td>BAB</td>
<td>1.969</td>
<td>0.39</td>
<td>0.33</td>
</tr>
<tr>
<td>R(Fe−N(_{ax}))</td>
<td>2.20 Å</td>
<td>0.34</td>
<td>0.34</td>
</tr>
</tbody>
</table>

“*The experimental data are taken from ref 66. The calculations were carried out at the DFT B3PW91/TZVP level (for the \(^{3}E\) at the relaxed JT-distorted geometry). BAB denotes the Mayer bond order index, a powerful parameter quantifying the strength of chemical bonds between atoms A and B (based on MO coefficients and overlaps).”37,39,67

Resolving the Quintet Structure with EXAFS Spectroscopy.

We have measured the EXAFS spectra of the [Fe(terpy)2]^{2+} complex in its ground LS and photoexcited HS state (at a time delay of 80 ps) with outstanding statistical quality. The spectra allowed us to verify experimentally the DFT-predicted molecular geometries for both spin states. Moreover, the bond length accuracy of the HS spectrum, unprecedented in time-resolved experiments, permitted us to determine with high confidence which of the two possible quintet states (\(^{5}E\) and \(^{5}B_{2}\)) is populated after the light excitation.

The conventional curve-fitting approach of Fourier-transformed EXAFS spectra with standard software packages has been used. A detailed description of EXAFS data reduction procedures, fitting approach, and methods used for statistical evaluation of fit errors is available in the SI.

The Fourier-transformed (FT) EXAFS spectrum of the LS ground state is shown in Figure 4a together with the fit function. In this case we have started with the DFT-optimized \(^{1}A_{1}\) geometry as input structure and refined it during the fitting process. The inset in Figure 4a shows the corresponding \(k^2\)-weighted EXAFS spectrum in \(k\)-space together with the fit curve. (The spectra in this figure appear shifted, as they have not been phase corrected.) The first peak observed in the FT-EXAFS spectrum in Figure 4a corresponds to the first coordination shell, which contains solely contributions from single-scattering paths involving axial and equatorial N atoms (N\(_{ax}\) and N\(_{eq}\) respectively) and the central Fe atom. The peaks in the range between 2 and 4 Å correspond to the second and further coordination shells around the Fe\(^{3+}\) ion, as well as multiple scattering paths from the first coordination shell. In the theoretical structure of the LS singlet state, the molecule is axially compressed. This leads to an anisotropic distribution of Fe−N bond lengths in the first coordination shell, with the axial Fe−N bonds about 0.1 Å smaller than the equatorial ones (cf. Table 1). The obtained EXAFS fit is in full agreement with this prediction, and we can unambiguously distinguish between the shorter axial Fe−N\(_{ax}\) bonds and the expanded equatorial bonds. The refined experimental bond lengths are R(Fe−N\(_{ax}\)) = 1.874 ± 0.004 Å and R(Fe−N\(_{eq}\)) = 1.969 ± 0.004 Å, in very good agreement with the DFT-calculated bond lengths listed in Table 1.

The EXAFS spectrum of the photoexcited [Fe(terpy)2]^{2+} molecule measured at 80 ps time delay is presented in Figure 4b. The spectrum was obtained using the data reduction method described in the SI. The expected elongation of Fe−N bonds (together with the expansion of outer-lying coordination shells) is clearly visible from the shift of all main peaks present in the FT spectrum toward higher \(k\)-values. Similarly to Figure 4a, the inset shows the photoexcited EXAFS spectrum in \(k\)-space with the corresponding best-fit result.

To ensure an unbiased determination of the excited-state structure, a range of starting parameters was used that covered the structures of the two proposed quintet states (using the corresponding theoretical models) as initial guesses. As derived from the DFT, the anisotropy affecting the Fe−N bond lengths is preserved in the HS excited state only for the \(^{3}E\) structure, whereas the structure of the \(^{3}B_{2}\) state has fairly similar Fe−N bond lengths for all six N atoms. Independent of the initial guess, in all cases the fit converged to a structure with substantially distinct Fe−N\(_{ax}\) and Fe−N\(_{eq}\) bonds lengths, clearly indicating that the anisotropic bond distribution is preserved in the photoexcited state and that we can distinguish between the two possible excited states. Furthermore, a rigorous statistical evaluation of the fit results based on \(\chi^2\)-analysis is discussed in detail in the SI and undoubtedly excludes the \(^{3}B_{2}\) quintet state due to the lack of any substantial differences between its Fe−N bonds within the first coordination shell and supports the \(^{3}E\) quintet state as the only possible structural model of the HS state. The refined Fe−N bond lengths are R(Fe−N\(_{ax}\)) = 2.08 ± 0.02 Å and R(Fe−N\(_{eq}\)) = 2.20 ± 0.01 Å, in excellent agreement with the theoretical model for the \(^{3}E\) state (cf. Table 1), which allows us to unambiguously assign its structure to the dynamically measured HS state.

It is worth mentioning that the inclusion of outer coordination shells (beyond Fe−N single scattering contributions in the 1−2 Å range in Figure 4) was possible given the superior EXAFS data quality obtained in the experiment for both the LS ground and photoexcited HS state spectra. In both cases, the statistical
goodness of the fit, evaluated using the $\chi^2$ test (see the SI for more details), significantly improved when extending beyond the first coordination shell and also by taking into account some of the multiple scattering contributions from outer-lying C atoms. For both LS and HS spectra, only very minor changes to the second and third coordination shells of DFT-optimized geometries were needed for fits to converge to minimum $\chi^2$ values (further details can be found in the SI).

**Structural Information from XDS.** The X-ray scattering signal from a (dilute) solution usually contains only a very small contribution from the solute. However, calculating the difference signal $\Delta S(Q, \Delta t) = S(Q, \Delta t) - S(Q, -\infty)$ effectively removes the otherwise dominating, but almost constant, scattering signal from the solvent. This procedure thus highlights the scattering signature of the structural dynamics of the solute and its environment, the solvent cage, and the structural changes in the surrounding bulk solvent. The analysis approach applied here is fully analogous to the one presented in ref 14, where the analogous [Fe(bipy)$_3$]$^{2+}$ compound was investigated.

Such difference signals acquired at six time delays from $\Delta t = 100$ ps to $\Delta t = 25$ ns measured on the [Fe(terpy)$_3$]$^{2+}$ solution are shown in Figure 5a. Qualitatively, the characteristic negative feature around $Q = 0.8$–1.2 Å$^{-1}$ corresponds to the immediate (on the 100 ps time scale) appearance of the bond-elongated HS structure, and the oscillatory feature around $Q = 2.2$ Å$^{-1}$ arises from temperature ($T$) and density ($\rho$) changes in the bulk solvent, as described in detail in ref 14 for the [Fe(bipy)$_3$]$^{2+}$ complex and in ref 69 for the PtPOP complex. Figure 5b shows the experimental $\Delta S(Q, \Delta t = 1$ ns) as well as two simulated signals $\Delta S_{\text{model}} = \gamma \times \Delta S_{\text{Solute+Cage}} + \Delta T \times \Delta S_{\text{AT}} + \Delta \rho \times \Delta S_{\rho}$, where $\gamma$ is the photoexcitation fraction and $\Delta S_{\text{Solute+Cage}}$ has been calculated for each of the two proposed HS structures $\beta^E$ and $\beta_2$. The $\Delta T \times \Delta S_{\text{AT}}$ and $\Delta \rho \times \Delta S_{\rho}$ contributions describe the changes in scattering due to bulk-solvent heating and density changes, respectively, and are determined in a separate experiment.70,71 Figure 5c displays each of these three contributions to the model difference signal at $\Delta t = 1$ ns, where the magnitudes of the individual contributions are comparable. Separate fits were carried out for each time delay and for both putative HS states, yielding essentially similar results.

The time evolution of the magnitude of $\gamma$, the photoexcited fraction for the HS state, is shown in the inset of Figure 5a. Fitting $\gamma$ in analogy with the fit of the XAS signal described above, we find $\gamma_{\text{XDS}} = 2.7 \pm 1.5$ ns, which is in reasonable agreement with the XAS-derived value; the larger uncertainty arises from the fewer time-domain data points in the XDS data set. The time evolution of $\Delta T$ and $\Delta \rho$ reflects the dynamics of the energy transfer to the solvent. In general, they follow what was observed in the analysis of [Fe(bipy)$_3$]$^{2+}$, although the density increase is somewhat smaller in magnitude in the present case ($\sim 0.1$ vs $\sim 0.4$ kg m$^{-3}$) and is at the limit of detection. The temperature increase is also observed to be lower ($\sim 0.3$ vs $\sim 0.6$ °C), in agreement with the lower concentration of excited-state solutes depositing energy to the solvent through the nonradiative decay processes. Finally, it is evident from Figure 5b that the expected difference signals for the two putative structures are very similar and that they fit the acquired data almost equally well.

In conclusion, the XDS results confirm the plausibility of the DFT structural models, but the present data set does not allow us to recognize the tiny differences of the quintet structures in the variations of the solute signal. Moreover, the time evolution of $\Delta T$ and $\Delta \rho$ provides invaluable insights into the solvent dynamics around the excited solute.

**Structure and Electronic Structure from XANES.** K-edge XANES relates to the p-projected unoccupied electronic density of states of the absorbing atom. Since this is largely influenced by the coordination and bonding, this technique is usually rather sensitive to structural changes around the absorbing atom. For the laser-induced quintet state of [Fe(terpy)$_3$]$^{2+}$, the population of the antibonding $e_g^*$ orbitals leads to dramatic rearrangements in the molecular and electronic structure which, as has been shown in Figure 2a, are clearly evident in the near-edge region. The structural differences between the two quintet states, however, are considerably smaller, and distinguishing them with XANES is more challenging. Yet the electronic structure of the $\beta^E$ and $\beta_2$ states differs, and one expects that this would yield differences in their XANES spectra. With the objective of obtaining further insights into the nature of the quintet state while at the same time assessing the capabilities of XANES to reveal electronic and structural information when very small changes are involved, we compare experimental XANES spectra with theoretical spectra obtained from DFT calculations.

The experimental XANES spectra of the ground and laser-excited ($\Delta t = 80$ ps) states were recorded in both TFY and HERFD (taken at the maximum of the Kot peak) modes. These spectra are shown in Figure 6. HERFD-XANES usually leads to improvement in the energy resolution when compared to the conventional detection modes.28 In the current experiment,
Figure 6. Experimental and calculated Fe K-edge XANES spectra of (molecular) \([\text{Fe(terpy)}_2]^{2+}\). With the experimental data, both detection modes (TFY: thick lines and HERFD: dotted lines) are presented; for the theoretical spectrum (thin line), only the electric dipole contribution is plotted. (a) LS ground state; the theoretical spectrum was calculated using the \(^1A\) structure. (b) Partially excited \([\text{Fe(terpy)}_2]^{2+}\) (Laser ON, \(\Delta t = 80\) ps); the theoretical spectrum was constructed as a linear combination of the calculated LS (60%) and HS (40%) spectra. (Given the similarity of the two theoretical HS spectra, the reconstruction was done only for the \(^5E\) structure for clarity.) (c) Photoinduced HS state of \([\text{Fe(terpy)}_2]^{2+}\); the plot shows the experimental spectra reconstructed from the laser ON and LS measurements (utilizing the population from XES) and the theoretical spectra calculated for both possible HS states, \(^5E\) (thin line) and \(^5B_2\) (dashed line). (d) Transient XANES obtained as the difference of the laser ON—LS (ground state) spectra.

Although the HERFD spectra appear somewhat sharper and better resolved, the effect is not dramatic. This can be explained by the fact that the relatively broad energy resolution of the spectrometer (∼0.8 eV) could not improve too much further on the good resolution obtained already in the TFY-XANES due to the narrow energy bandwidth of the incident beam (∼0.5 eV).

Theoretical modeling of K-edge XANES based on the single electron picture has become reliable, and its combination with experiment grew a powerful tool in electronic structure analysis. The theoretical spectra were obtained with such a well-established, reliable approach that takes into account the 1s core hole, utilizing the Quantum ESPRESSO program packages. Electric dipole transitions from the 1s state to the unoccupied p-states were calculated, and DFT (BP86/TZVP) optimized structures for the LS and HS states were utilized. Comparing first the spectra of the LS ground state in Figure 6a it is clear that the DFT-derived spectrum agrees very well with the experiment. The energy positions and the relative intensities of the major spectral features, labeled B, C, and D, are well reproduced, including the high-energy shoulder of peak C (which is resolved only in HERFD-XANES). The pre-edge peak labeled A consists of mostly quadrupolar transitions and so is not expected to be properly accounted for by the current electric dipole calculation.

Comparisons to the laser-excited (\(\Delta t = 80\) ps) experimental spectra are shown in Figure 6b, c, and d. In Figure 6b the measured laser ON spectra, which contain a mixture of LS ground state and HS excited state contributions, are compared with a superposition of LS and HS theoretical spectra. Here the calculated HS component arises solely from the \(^5E\) state. The proportion of HS contribution is taken to be 40% based on the excitation fraction derived from the XES measurement. The calculated spectrum is seen to again reproduce the key spectral features, peaks B, C, and D. For a comparison that is independent of the LS and HS population fractions, the differences between the laser ON spectra and the LS ground state spectra are shown in Figure 6d. Here the theoretical HS—LS difference, which has been scaled to match the intensity of the measured curves, can be seen to mimic the shape of the experimental difference spectra. The agreement in both of these comparisons validates the compatibility of the DFT calculated HS state structure with the experimental observations.

To test whether XANES could distinguish between the two possible HS states and perhaps confirm the EXAFS assignment, theoretical spectra for the \(^5E\) and \(^5B_2\) states were compared to experimental spectra for the HS state that have been reconstructed from the laser ON spectra assuming a 40% HS state fraction. These are shown in Figure 6c. The calculated spectra are seen to be very similar, making a definitive assignment of the experimental spectra to one or the other state difficult. Comparing the ratio of the B and C peak intensities might lead to the opposite conclusion, which warns us that this approach might be far-stretched, as the calculated intensities are apparently not accurate enough for such conclusions in the present case.

The primary electronic structure differences between the \(^5E\) and \(^5B_2\) quintets manifest themselves in the occupation of the 3d orbitals, which brings us to the discussion of the variation of the experimental pre-edge region (peaks A, A′, and A″) upon excitation. This part of the spectrum stems from 1s → 3d transitions. Being sensitive to the occupation of the 3d orbitals, this shall provide further information on the electronic structure.
Indeed, as is evident from Figure 6, this region shows large variations after excitation. However, in conventional XANES these peaks are usually weak and overlapping, and they are also partly obscured by the tail of the main K edge; therefore, they are often poorly resolved. In order to overcome these difficulties and obtain better resolved spectra, we have recorded 1s2p3/2 RXES, the combination of XANES in the preedge region, and the (1s2p) XES around the Kα1 peak. This is presented in the next section.

**Resolving the XANES Pre-Edges with RXES: Further Insights into the 3d Electron States.**

Exploring the pre-edge region of 1sXAS with RXES can reveal more details about the 3d-related part of the electronic structure. We performed 1s2p RXES on light-excited [Fe(terpy)2]2+ at a time delay of Δt = 80 ps. These data are plotted in the top row of Figure 7 next to the ground-state spectrum. The difference between the light-excited and ground-state spectra is shown along with the (fully converted) HS spectrum that was reconstructed from the difference spectrum assuming the population obtained from XES.

FeII in (close-to) octahedral environments is expected to have a single pre-edge resonance when in the LS state and three resonances when in the HS state.55,75 This has been confirmed in several experiments including 1s2p RXES of an FeII compound at thermal spin-state switching.64 These resonances are clearly present in the spectra in Figure 7, and the increase in resolution and information content with respect to XANES is evident when these features are compared to peaks A, A′, and A″ in Figure 6.

Calculated RXES spectra are shown in the middle row of Figure 7. While a single-electron approach usually cannot give an accurate description for 1s2p RXES spectra, multiplet theory can do that for both LS and HS states.55 For octahedral complexes the nature of the pre-edge transitions is electric quadrupolar due to symmetry reasons. The FeN6 cores in [Fe(terpy)2]2+ and [Fe(bipy)3]2+ are distorted octahedra but at a level of distortion that makes the quadrupolar description sufficient to model the spectra. The calculations assumed LS and HS FeII ions in octahedral crystal fields, and the results reproduced the single LS and three HS resonances as can be seen in Figure 7. Multiplet interactions in the RXES final state of the ground-state system lead to small intensity peaks that appear vertically, at different energy transfer values, in the plotted RXES spectra. In the measured LS spectrum the expected (vertical) multiplet structure is weak but obviously present near the single resonance at around 7113.5 eV. In addition, a diagonal feature is observed (at ca. Ω = 7115.5 eV incident energy, Ω − ω = 710.5 eV energy transfer) which is not present in the calculation. This feature must reflect a p-type contribution as a result of orbital mixing (i.e., covalency). In fact, this mixing is expected to bring in a dipolar character, which is seen in the XANES-calculated dipolar contribution to the pre-edges in Figure 6. Upon laser excitation two more peaks (at ca. Ω = 7113 eV, Ω − ω = 706 eV, and ca. Ω = 7116 eV, Ω − ω = 711 eV) appear, while the intensity of the main peak drops. The difference of these spectra together with the known HS population allows us to reconstruct the 1s2p.

**Figure 7.** Experimental Fe 1s2p RXES spectra, from aqueous solutions of [Fe(terpy)2]2+ (top row) and [Fe(bipy)3]2+ (bottom row), showing the ground state, the laser-excited state at 80 ps time delay, their difference, and the reconstructed HS spectra. The theoretical spectra (middle row) were obtained from ligand-field multiplet calculations in an approximate O₃ local symmetry, and there the “Laser ON (partly HS)” panel was constructed by the superposition of 40% HS and 60% LS. The spectral intensity is plotted above the plane spanned by the incident energy and the difference between the incident and emitted energy, the energy transfer. The energy scaling is equal for both axes and identical for all the plots in order to aid the comparison.
RXES spectrum of the HS state of \([\text{Fe(terpy)}_2]^{2+}\), which shows a striking similarity to the calculated one.

We have also studied the 1s2p RXES spectra of the benchmark system \([\text{Fe(bipy)}_3]^{2+}\) which has a somewhat different symmetry \((D_3h)\) and less geometrical constraints from the ligands as they are bidentate. The measured spectra are shown in the bottom row of Figure 7. Despite the somewhat worse energy resolution caused by a twice-larger incident bandwidth, it is apparent that the spectra are very similar to those of the \([\text{Fe(terpy)}_2]^{2+}\). The almost identical difference spectra indicate that the nature of the variations in the two complexes are essentially the same.

The crystal field multiplet calculations nicely reproduce the overall spectral shapes and the difference spectra. Intra-atomic electron–electron interactions together with a crystal-field splitting are therefore the main contributions to the spectra. The calculations neglect orbital mixing effects and only consider electric quadrupole transitions; as a result spectral features may be missing on the high-energy side, and the spectral intensities cannot be fully reproduced.

The chemical sensitivity of the K absorption pre-edge is very different from the main edge. While the main edge spectral shape is dominated by the coordination shells, the pre-edge reflects electron–electron interactions within the d-shell as well as crystal-field effects and orbital mixing. The striking similarity of the difference spectra in the two compounds shows that intra-atomic d–d interactions in the HS state dominate the spectral shape. The 1s2p RXES planes thus visualize the dramatic change of the electronic levels when the molecules are in their short-lived photoexcited state. However, the reasonably good description by the multiplet calculations, an iconic approach utilizing \(O_h\) symmetry, points toward the need for more involved calculations. Distinguishing between \(5E\) and \(5B_2\) in RXES would require full multiplet calculations including dipole and quadrupole transitions with molecular wave functions that describe the whole complex. This is currently not possible with sufficient accuracy. Angular-dependent RXES could help to identify the exact 3d configuration in single crystals; however, this is hardly possible in solution.

**Theoretical Insights into the Electronic Structure.** Although the \([\text{Fe(terpy)}_2]^{2+}\) molecule has already been investigated theoretically,\(^\text{11,12}\) a few details relevant to the characteristics of the quintet states have been overlooked or at least not discussed in due detail. These include the relation between the structural differences and the occupation of the 3d orbitals in the two quintet states \(5E\) and \(5B_2\), the effects of the Jahn–Teller (JT) distortion on the \(5E\), and the electronic structure origin of the two modes needed to describe the spin-state transition. Furthermore, the previous results raise disturbing questions: is it hopeless to distinguish the electronic structure differences of two possible quintets with X-ray spectroscopy? Why is the \(5E\) state populated when almost all DFT calculations predict \(5B_2\) as being more stable? Here we complement the referred works by performing the necessary analysis using TD-DFT and CASPT2 and provide answers to all of the above-mentioned issues.

In order to unveil the variations in the electronic structure that cause the structural and bonding differences between the \(5E\) and \(5B_2\) states, we compare the relevant molecular orbitals (MOs) obtained with spin-unrestricted DFT. To examine this in detail, in Figure 8 we present the highest occupied and the lowest unoccupied (Kohn–Sham) MOs. (A more extended version of this MO diagram is plotted in the SI.) Not surprisingly, these orbitals at the HOMO–LUMO gap have substantial Fe 3d character. The most interesting and relevant molecular orbitals at the gap are those with a dominant contribution from the atomic Fe \(d_{xy}\), \(d_{xz}\), and \(d_{yz}\) orbitals of \(\beta\) spin (i.e., spin down). Only one spin-down electron is available for these three orbitals. Which of these orbitals the electron occupies determines the atomic and electronic structural differences.

In the simplest approximation of ligand-field theory, the Fe \(d_{xy}\), \(d_{xz}\), and \(d_{yz}\) atomic orbitals are nonbonding. However, this is altered in such a covalent molecule where the atomic orbitals are mixed to some extent with ligand orbitals. In the bottom of Figure 8 3D representations show the actual composition of these MOs for the \(5B_2\) state. Here the \(5b_1\) orbital is populated, which still has nonbonding \(d_{yz}\) character; the atomic orbital...
contribution (mixing coefficient) for the Fe d_{xy} is 84.2%, while those for all nitrogen orbitals are below 1%. Nevertheless, it is apparent that the d_{xz} and d_{yz} orbitals are mixed with in-phase p-type orbitals from the axial nitrogens (58.5% Fe(3d), 6.24% N(p)). The resulting 2-fold-degenerate 32e pair of MOs have a ($\pi$-)bonding character. However, this has no effect on the properties of the 5B_2 state since these MOs are unoccupied. In the 5E state (when constraining the symmetry to D_{2d} as in ref 11), the energy order of the 5B_2 and 32e orbitals is swapped, and the 32e orbital pair is populated by a single electron. These orbitals bond to the axial nitrogens, as we just discussed, which shortens the Fe-N_{ax} bond lengths. (This orbital mixing and the corresponding charge transfer to the ligand via the bonding is usually referred to as backbonding or back-donation.)

Without constraints on the symmetry, the JT effect removes the degeneracy of the 32e orbitals in the 5E state and lowers the symmetry to C_2. The highest occupied orbital becomes the 63b, which has the main contribution from the Fe d_{xy} atomic orbital, and thus the backbonding to the axial nitrogens remains effective. The enhanced bonding is clearly reflected in the Mayer bond order indices reported in Table 1 and provides a natural explanation for the geometry differences of the quintet states. Moreover, the energies of the levels of the quintet MOs become strikingly similar after applying the JT distortion to the 5E; the frontier orbitals have almost identical energies. The LUMO pair 64b and 70a (l_{xz},l_{yz}) mimics the 32e (l_{xz},l_{xy}) pair of 5B_2, and the even higher-energy unoccupied orbitals show high resemblance for the 5B_2 and 5E (C_2) states (see SI). Such similarities of the electronic structure readily explain why the presented spectroscopy data are so similar for the two quintet states.

The question remains why the 5E state is populated, contradicting most DFT results that favor the 5B_2 state as more stable.\(^{11,13}\) The energy difference between the quintet states is rather small, and on such a scale the DFT calculations show a large scatter that depends on the choice of the exchange-correlation functional.\(^{12}\) Therefore, we have performed CASPT2 calculations that provide more accurate energy differences. As can be seen in Figure 9(b), the two quintet minima are separated along a line that is almost orthogonal to the line connecting the LS and quintet states. The energy difference and the relative location of the potential wells belonging to the 5E and the 5B_2 states are best appreciated along this coordinate. This is plotted in Figure 9(c). The energy difference is 41 meV with the 5E state lower in energy. Moreover, as the 5E state is expected to split due to the Jahn–Teller effect, we can expect the energy to be even lower. (Note that the JT effect only affects the angle between the planes of the two terpy ligands, which changes by 4°, while the bond lengths and other angles stay practically the same.)\(^{12}\) The combination of the energy difference from CASPT2 and the JT energy lowering obtained from DFT suggests that the distorted 5E can be as much as 150 meV lower in energy than the 5B_2 state. As the transfer to the quintet(s) takes place in the first picosecond after light excitation and the measurements are performed at a delay $\geq 80$ ps by which time the excess energy is transferred to the solvent, we expect to find the system in the 5E, the more stable quintet state. Yet, since geometry optimization is practically impossible at the CASPT2 level on a such a large system with numerous electrons, one should keep in mind that this expectation might be biased.

Finally, we address the origin of the breakdown of the SCC model. In most octahedral Fe\(^{II}\) compounds that undergo spin-state transitions a single configuration coordinate is sufficient as a reaction coordinate, typically the breathing mode of the molecule which is proportional to the metal–ligand distance. This has been exploited in a very successful quantitative approach proposed by A. Hauser that describes well the HS–LS relaxation times for most complexes but not Fe(terpy)\(^{2+}\)\(^{10,11}\). Given the slower than expected relaxations (and apparently higher barrier) found for Fe(terpy)\(^{2+}\)\(^{12}\) in different matrices, it was suggested that the SCC-based model is not applicable to this tridentate complex, and the involvement of an extra mode, probably the ligand bending measured by the NNN angle, is required, which increases the barrier height. Many details of this have not been brought to light; in a recent paper we found that the NNN angle of the ligand and the R(Fe–N_{ax}) bond lengths are promising candidates to describe the reaction coordinates.\(^{12}\)

\[ e^{-\frac{(z^2)}{2}} \]
Starting with the description of the electronic structure of the $^{1}A_{g}$ ground state, which has no unpaired electrons, we note that the spin-restricted KS MOs have identical energies for $\alpha$ and $\beta$ electrons, as is observed on the frontier orbitals of this state in Figure 9a. The tridentate terpyridine has geometrical constraints as the N atoms of its three linked pyridine rings in this planar ligand cannot take up ideal octahedral positions around the Fe center. Due to this, the energy of the molecular orbitals that have a dominant contribution from the 3d orbitals will have a relevant split in energy, depending on whether they are oriented parallel or orthogonal to the molecular axis that goes through the nitrogens of the central pyridine rings and the Fe$^{II}$ ion. This splitting is small for the $d_{x^2}$, $d_{y^2}$, and $d_{z^2}$ nonbonding or weakly backbonding orbitals but high for the $d_{x^2-y^2}$ and $d_{xy}$, $d_{xz}$, $d_{yz}$-based orbitals, which originate from the octahedral $e_{g}^\ast$ antibonding orbitals as seen in Figure 9a. In addition to the geometrical constraints, the backbonding between the axial N and the Fe contributes to the axial distortion of the molecule as well. The stronger bonding along the axis is again clearly identified in the Mayer bond order indices, as seen in Table 1.

To reach the quintet states, both antibonding $e_{g}^\ast$ orbitals need to be populated with a spin-up electron. However, it is revealing to examine what would happen to the molecular geometry if only one of these orbitals is populated. The $e_{g}^\ast (lz^2-xy^2)$ MO is based on the Fe $d_{x^2-y^2}$ atomic orbital mixed with $p$-orbitals of the four equatorial nitrogens; therefore, populating this antibonding orbital causes an expansion in the molecule in this direction, opening up the NNN angle of the ligand, without affecting the axial $R(Fe−N_{eq})$ bond length. Indeed, the transition to the $^{3}A_{1}$ state is almost perfectly described by the NNN opening, as can be seen in Figure 9. This suggests that the opening of the NNN angle of the ligands is one of the relevant modes needed for the LS−HS transition. The $e_{g}^\ast (lz^2)$ antibonding orbital has large mixing of the Fe $d_{x^2}$ with the $p$-orbitals of the axial N atoms and smaller mixing with those of the equatorial N atoms. Therefore, the population of this MO leads to an increase in the lengths of the Fe−N$_{ax}$ bonds and a smaller weakening of the equatorial Fe−N$_{eq}$ bonds. This results in the increase of the axial $R(Fe−N_{eq})$ bond length without substantial effect on the NNN angle of the ligands because the pull on the side rings at the equatorial nitrogens by the iron caused by axial elongation is compensated by the simultaneous weakening of the Fe−N$_{eq}$ bonds. This change corresponds to the second coordinate needed for the transition.11,12 The optimized geometry of the triplet with the populated $d_{y^2}$-based orbital, $^{3}B_{1}$, differs from that of the ground state almost only in $R(Fe−N_{eq})$, as shown in Figure 9. (Note that the geometry optimization of the $^{3}B_{1}$ could only be obtained by forcing the occupation of the $e_{g}^\ast (lz^2)$ MO with a symmetry constraint on the orbital populations.) Consequently, we can identify that these population changes combined with the geometrical constraints of the terpy ligand explain the need for the two-mode description of the switching and the breakdown of the SCC model in this molecule. The validity of relating the two necessary configuration coordinates to electronic structure effects is further supported if we add up the two vectors pointing from the singlet ground state to the triplet states in this coordinate system. Each of these vectors represent the structural variations caused by adding one electron to an $e_{g}^\ast$ orbital. Therefore, we can expect that their sum approximates well the real quintet geometry, if the relation between structure and orbital population variations is valid. The resulting position (marked by a hollow cross in Figure 9) lies indeed fairly close to the actual positions found for the quintet states with the geometry optimizations. The mismatch is comparable to the distance between the $^{5}E$ and $^{3}B_{2}$ states, which differ in the populations of t$_{2g}$-like orbitals. Accordingly, the two modes based on the opening of the NNN angle and the variation of the $R(Fe−N_{eq})$ bond lengths grasp rather well the main structural changes, and these seem to clearly correlate with the populations of the $e_{g}^\ast$-like orbitals. However, remaining smaller-scale variations in the electronic structure are also reflected in the molecular structure. Consequently, further refinement of these modes or involvement of further ones is needed to obtain a complete and fully satisfactory description of this system.

## CONCLUSIONS

The challengingly small differences in the molecular geometry and electronic structure of the $^{3}B_{2}$ and $^{5}E$ quintets were addressed utilizing a variety of time-resolved X-ray techniques. The minuscule explorable differences provided a demanding testing ground for the techniques, enabling assessment of their ability to discern small changes that occur in short-lived transient states. The formation and decay of the high-spin state was followed with all applied techniques. XANES, XES, and RXES spectra, despite their high quality and clear variations, could not reveal which of the two quintets is realized. XDS, which captured both the light-induced conversion to HS and the behavior of the solvent in the vicinity of the molecule, was also unable to distinguish between the two quintets. Excellent quality EXAFS spectra were able to resolve the bond-length variations accurately, leading to unambiguous identification of $^{5}E$ as the formed quintet state which is in agreement with our theoretical predictions. Applying theoretical calculations (DFT, TD-DFT, and CASPT2), we were able to explain the electronic structure origin of the differences in the properties of the two possible quintet states. We identified that back-bonding in the $^{5}E$ state is responsible for the structural differences between the quintet states. We revealed why the $^{5}E$ state is the more stable one and also checked how this state varies upon the Jahn–Teller effect. The failure of spectroscopy to distinguish the quintets was readily explained by the striking similarity of their molecular orbital diagrams. Finally, we determined why one configurational coordinate is not sufficient for the description of the LS−HS transition in [Fe(terpy)$_2$]$^{2+}$.

Due to the anomalous behavior of [Fe(terpy)$_2$]$^{2+}$ among Fe$^{II}$ complexes that undergo photoinduced LS to HS transitions, the identification of the quintet state produced after light excitation represents a milestone. Our work should inspire studies of the dynamics on the way to the quintet state, during the first picosecond after light excitation, using X-ray free electron lasers. This will lead to a more complete understanding of the photocycle of this system, which can motivate the design of novel molecules with better photoswitching performance.

## ASSOCIATED CONTENT

### Supporting Information

Further details on the relation between the states in the different symmetry groups, the lifetime fitting, the EXAFS analysis, and the calculated molecular orbital diagrams. This material is available free of charge via the Internet at http://pubs.acs.org.

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DOI: 10.1021/acs.jpcc.5b00557
■ ACKNOWLEDGMENTS

This project was supported by the European Research Council via contracts ERC-StG-259709 and ERC-AdG-2011.618485, the "Lendület" (Momentum) Program of the Hungarian Academy of Sciences, and the European XFEL. K.H., M.N., T.B. van D., K.K., and H.L. acknowledge support from the Danish National Research Foundation’s Centre for Molecular Movies and DANSATT. C.B., W.G., A.G., A.B., and T.A. acknowledge funding by the Centre of Ultrafast Imaging (Hamburg), by the EU-funded Cluster of Research Infrastructures for Synergies in Physics (CRISP) project, and the Deutsche Forschungsgemeinschaft via SFB925. A.B., N.Z., G.V., D.C., and A.J. acknowledge the French Ministry of Foreign Affairs (MAE) and the French Ministry of Higher Education and Research (MESR) for funding through Balaton project no. 27875YK, as well as the Hungarian and French Intergovernmental S&T Cooperation Program (TET11FR-XTHEOEXP). Z.N. acknowledges support from the Bolyai Fellowship of the Hungarian Academy of Sciences. J.U. acknowledges continued funding from the Knut and Alice Wallenberg Foundation. D.C. and A.J. acknowledge GENCI (Grand Equipement National de Calcul Intensif), which allows access to the HPC resources of IDIRIS under the allocation 2013-100172. Use of the Advanced Photon Source, an Office of Science User Facility operated for U.S. Department Of Energy Office of Science by Argonne National Laboratory, was supported by the U.S. DOE under contract no. DE-AC02-1984. The authors declare no competing financial interest.

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Supporting information for the paper “Detailed characterization of a nanosecond-lived excited state: X-ray and theoretical investigation of the quintet state in photoexcited [Fe(terpy)$_2$]$^{2+}$”

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1. Correlation table for the symmetry groups of $[\text{Fe(terpy)}_2]^{2+}$

Table S1: Correspondence between the notation of the relevant electronic states for the point group symmetries used in the paper.

<table>
<thead>
<tr>
<th>O$_h$ point group</th>
<th>D$_{2d}$ point group</th>
<th>C$_2$ point group</th>
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<td>$^1$A$_1$</td>
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<td>$^1$A$_2$</td>
<td></td>
</tr>
<tr>
<td></td>
<td>$^1$E</td>
<td></td>
</tr>
<tr>
<td>$^3$T$_{1g}$</td>
<td>$^3$A$_2$</td>
<td></td>
</tr>
<tr>
<td></td>
<td>$^3$E$_1$</td>
<td></td>
</tr>
<tr>
<td>$^3$T$_{2g}$</td>
<td>$^3$B$_1$</td>
<td></td>
</tr>
<tr>
<td></td>
<td>$^3$E$_2$</td>
<td></td>
</tr>
<tr>
<td>$^5$T$_{2g}$</td>
<td>$^5$B$_2$</td>
<td></td>
</tr>
<tr>
<td></td>
<td>$^5$E</td>
<td></td>
</tr>
<tr>
<td>$^5$E$_g$</td>
<td>$^5$B$_1$</td>
<td></td>
</tr>
<tr>
<td></td>
<td>$^5$A$_2$</td>
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</table>

2. XAFS data quality

The unprecedented quality of our time-resolved X-ray absorption data is demonstrated...
in Fig. S1. The data was collected during 5 hours, in a repetitive 1 bunch laser ON / 4 bunch laser OFF pattern. This means that only 1 hour was effectively used to record laser-excited data, accumulating $7 \times 10^8$ counts. The data has been summed up and normalized without any smoothing.

3. **Lifetime fitting**

In the Fe 1$s$ XAS, as seen in Fig. S1, the 7125 eV feature shows the largest variation at the spin transition. The time evolution of the photoexcited sample was measured by recording the intensity of this feature. For each time delay setting, the acquisition alternated between measuring the signal with and without the laser excitation. These are shown in Fig. S2. The data have been corrected to eliminate artefacts emerging from the laser phase shifter and the corrected ratio has been fitted with Eq. S1.

Figure 3 of the paper displays the time evolution of the intensity of this signal feature, which corresponds to the formation and decay of the HS state. The width of the rise centered at $\Delta t = 0$ corresponds to the X-ray probe pulse width ($\approx 80$ ps), while the later exponential decay is governed by the lifetime of the HS state. This time evolution is fitted by an expression of the form
Figure S2: Time evolution of the intensity of the 7125 eV feature of the [Fe(terpy)$_2$]$^{2+}$ XANES spectrum with and without laser excitation as a function of the time delay between the laser and the X-ray pulses.

\[ I(\Delta t) = I(t - t_0) = \gamma_0 \int_{-\infty}^{\infty} \frac{1}{\sigma \sqrt{2\pi}} e^{-y^2/2\sigma^2} H(t - t_0 - y)e^{-(t - t_0 - y)/\tau} dy + C \quad (S1) \]

derived from a Gaussian broadening of an exponential decay starting at $\Delta t = 0$. $\tau$ stands for the lifetime of the HS state, $t_0$ is the time of the laser excitation, and $H$ is the heaviside step function, and $C$ is the background.

Table S2: Parameters for fitting Eq. S1 to the delay-dependent 7125 eV XANES intensity.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
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<tbody>
<tr>
<td>$\tau$</td>
<td>2.612 ± 0.006 ns</td>
</tr>
<tr>
<td>$\sigma$</td>
<td>0.0323 ± 0.0004 ns</td>
</tr>
<tr>
<td>$t_0$</td>
<td>-0.0075 ± 0.0003 ns</td>
</tr>
<tr>
<td>$\gamma_0$</td>
<td>0.4157 ± 0.0004</td>
</tr>
<tr>
<td>$C$</td>
<td>0.0009 ± 0.0004</td>
</tr>
</tbody>
</table>

* FWHM: 76.2 ± 0.9 ps

A large correlation was only observed between the lifetime and the background: \text{correl}(\tau, C) = -0.82.

4. EXAFS data reduction and fitting
This section describes the details of the EXAFS data reduction and subsequent fitting procedure for both LS and HS spectra, which are shown in Fig. 4 of the paper.

As explained in the Theoretical Methods section of the paper, Density Functional Theory (DFT) has been applied to obtain the molecular structures of both the LS $^1A_1$ ground state (Fig. S3A) and two possible lowest energy HS states, namely $^5B_2$ (Fig. S3B) and $^5E$ (Fig. S3C) quintets. (We have used Mercury CSD 2.0 software\(^1\) for plotting the DFT-optimized structures shown in Figs. S3 and S4.)

Note that the $^5E$ undergoes a Jahn-Teller distortion, and its symmetry lowers to $C_2$. In this point group the correct notation for $^5E$ would be $^5B$, but we keep the higher symmetry notation ($^5E$) in order to avoid confusion with the state $^5B_2$.

Due to the steric constraints imposed by terpyridine ligands, the N atoms of pyridine rings cannot be accommodated in ideal octahedral positions around the Fe atom, which results in a distorted geometry for the LS $^1A_1$ ground state with two different types of coordinating nitrogen atoms: 4 N atoms lie in the equatorial plane ($N_{eq}$) and the remaining 2 N atoms ($N_{ax}$) are on the molecular axis (Fig. S4A). The axial bond lengths $R(Fe-N_{ax})$ are about 0.1 Å shorter than the equatorials ($R(Fe-N_{eq})$), thus the molecule is axially compressed. For the $^5E$ quintet state, DFT calculations predict a fairly similar bond elongation for both axial and equatorial directions, preserving the
LS axial compression. On the contrary, for the $^5\text{B}_2$ quintet state the bond elongation in the axial direction is expected to be 0.05 Å larger than in equatorial direction thus the Fe–N bond lengths become almost uniform, as can be seen in Fig. S4B.

Figure S4: The effect of geometrical constraints, due to the nature of tridentate terpyridine, is shown for $^1\text{A}_1$ (A) and $^5\text{B}_2 / ^5\text{E}$ geometries (B). The red and blue circles in (A) mark the positions of equatorial and axial N atoms, respectively. The red arrows in (B) indicate the structural distortion of Fe-ligand bonds (bond expansion) in the two possible quintet states.

Using structural models for ground and both possible excited states (ES), EXAFS spectra have been simulated and fitted using FEFF$^3$ and IFEFFIT$^4$ codes, respectively. The calculations of scattering amplitudes and phases were carried out using the FEFF8.20 code. The atomic potentials were calculated self-consistently within the muffin-tin approximation and the Hedin-Lundqvist self-energy model was chosen as an exchange correlation potential for both the fine structure and the atomic background.

The experimental EXAFS spectrum of the ground state (GS) was reduced using the Athena program (part of IFEFFIT package). The data reduction consisted of normalization and removal of the atomic background $\mu_0$ using a cubic spline function. Once reduced, the GS EXAFS spectrum was fitted with the Artemis software (also part of IFEFFIT) using the scattering amplitudes and phases calculated by FEFF8.20 for
A$_1$ geometry (Fig. S3A). All fits were performed in $R$-space using $k^2$ weighting and the results are summarized in Table S3. We have used identical $k$ ranges for Fourier transforms of both LS and HS spectra (2.5–10.5 Å$^{-1}$) and identical $R$ ranges (1-3.8 Å) for both fits presented in Fig. 4 of the paper. This resulted in the same number of statistically independent data points $N_i = (2\Delta k\Delta R)/\pi$, where $\Delta k$ and $\Delta R$ are the ranges in $k$- and $R$-space, respectively. The fit evaluation was done using a standard statistical $\chi^2$ function and the confidence limits for fit parameters were estimated using the error estimation method described in Refs 5,6. One should note that the number of degrees of freedom $\nu = N_i - P$, where $P$ is the maximum number of parameters used in the fit, was different for LS and HS fits and the corresponding values are reported in Table S3.

Table S3: EXAFS fit results for the LS and photoexcited HS states of [Fe(terpy)$_2$]$^{2+}$ in solution.

<table>
<thead>
<tr>
<th></th>
<th>LS (A$_1$)</th>
<th>HS</th>
</tr>
</thead>
<tbody>
<tr>
<td>$R$(Fe–N$_{ax}$)</td>
<td>1.874(4) Å</td>
<td>2.08(2) Å</td>
</tr>
<tr>
<td>$R$(Fe–N$_{eq}$)</td>
<td>1.969(4) Å</td>
<td>2.20(1) Å</td>
</tr>
<tr>
<td>$S_0^2$</td>
<td>0.78(4)</td>
<td>0.78(4)</td>
</tr>
<tr>
<td>$\sigma^2(R$(Fe–N))</td>
<td>$2.6(7) \times 10^{-3}$ Å$^2$</td>
<td>$2.6(7) \times 10^{-3}$ Å$^2$</td>
</tr>
<tr>
<td>$E_0$</td>
<td>$-5.8(4)$ eV</td>
<td>$-4.0(7)$ eV</td>
</tr>
<tr>
<td>$N_i$</td>
<td>14</td>
<td>14</td>
</tr>
<tr>
<td>$\nu$</td>
<td>6</td>
<td>9</td>
</tr>
<tr>
<td>$\chi^2$</td>
<td>165</td>
<td>315</td>
</tr>
<tr>
<td>Reduced $\chi^2$</td>
<td>28</td>
<td>35</td>
</tr>
<tr>
<td>$R$-factor</td>
<td>0.1 %</td>
<td>1.1 %</td>
</tr>
</tbody>
</table>

We have optimized the change in Fe–N bond lengths for axial (Fe–N$_{ax}$) and equatorial (Fe–N$_{eq}$) N atoms by one value, next to the change in second-shell Fe–C bond distances and the change in additional further lying single and multiple scattering shells of C atoms (not shown in the table).

We have constrained the changes in Fe–N bond lengths for two $\Delta R$ values: one for the axial (Fe–N$_{ax}$) and one for the equatorial (Fe–N$_{eq}$) bonds, while using one parameter for the change in second-shell Fe–C bond distances and the one for the change in
additional further lying single and multiple scattering shells of C atoms (not shown in the table).

For each coordination shell we have obtained a Debye-Waller (DW) factor and we have used a global value of the energy threshold \( E_0 \) and amplitude reduction factor \( S_0^2 \) for all scattering contribution of the GS spectrum, which results in a maximum number of fit parameters \( P = 8 \). The fit was done by optimizing all variables simultaneously. The results for the GS spectrum show a very good agreement with the geometry of the LS \(^1\text{A}_1\) state with only small corrections to the Fe–N\(_{\text{ax}}\) and Fe–N\(_{\text{eq}}\) bond lengths. We have not observed any significant parameter correlations for multi-shell fitting. Therefore we concluded that the DFT-predicted atomic structure of the singlet LS state provides a good structural model of the GS [Fe(terpy)]\(^{2+}\) complex.

![Figure S5: Square residuals between the experimental excited state EXAFS data and \(^5\text{E}\) and \(^5\text{B}_2\) model geometries of the HS quintet state as a function of Fe–N\(_{\text{ax}}\) (left) and Fe–N\(_{\text{eq}}\) (right) bond distances. All other EXAFS fit parameters were fixed during the analysis. The dotted vertical lines indicate the predicted axial and equatorial Fe–N bond lengths as obtained by DFT.](image)

The quintet EXAFS spectrum was reconstructed from the transient difference spectrum and the LS spectrum using a relation described in Ref. 7 and the excited HS state fraction derived from XES measurements (see main text for details). In order to minimize the number of systematic errors introduced by data normalization and
reduction procedures, we have used the GS $\mu_0$ function to normalize and remove the background from the ES spectrum using a Matlab code. Fitting of the ES EXAFS spectrum was carried out similarly to the GS case, and we have fixed DW factors and $S_0^2$ in order to minimize the free parameter space in the fitting procedure. The $E_0$ parameter was constrained in the fit to vary in a limited range, as it is strongly correlated with Fe–N bond distance changes. Fits using both $^5B_2$ and $^5E$ atomic coordinates as initial parameters were performed in $R$-space, with the same $k^2$-weighting as for the GS spectrum and using the same number of statistically independent data points, i.e. $N_i = 14$. As mentioned earlier, only the energy shift $E_0$ and the changes in bond lengths, for Fe–N$_{ax}$ and Fe–N$_{eq}$ separately, as well as small corrections to the second shell Fe–C distances and to all longer scattering paths, were optimized, which resulted in $P = 5$. The summary of the most relevant fit results including statistical parameters used in the estimation of the fit errors is presented in Table S3.

We have performed fitting using both $^5B_2$ and $^5E$ input geometries independently as initial guesses. The fit results converged to an identical HS state model, which was very close to the DFT-predicted $^5E$ structure. Therefore we conclude that the measured ES EXAFS spectrum resembles closely the $^5E$ geometry of the aqueous [Fe(terpy)$_2$]$^{2+}$ obtained by DFT, independent of the initial structural model used for FEFF8 simulations and with only minor changes in the relevant Fe–N$_{ax}$, Fe–N$_{eq}$ bond distances and for higher coordination shells. The observed red-shift of $\Delta E_0 = -1.8$ eV is also in agreement with what has been observed earlier for similar spin-transition complexes, which reflects the expansion of the Fe–ligand bonds.$^8,9$

Since the structural differences between $^5B_2$ and $^5E$ geometries are very subtle, only due to the very high quality of our EXAFS data and a precise reconstruction of the HS EXAFS spectrum (using a very accurate measure of the photoexcited HS fraction derived from XES spectrum) was it possible to distinguish both models with statistically relevant accuracy. As mentioned earlier, standard $\chi^2$ tests were also used to
evaluate and compare the statistical goodness of the EXAFS fits as a function of the most relevant structural parameters, namely Fe–N\textsubscript{ax} and Fe–N\textsubscript{eq} bond distances. Here, we have varied the Fe–N\textsubscript{ax} bond lengths in small steps of 0.02 Å and the Fe–N\textsubscript{eq} bond lengths in steps of 0.01 Å and let the rest of the molecular structure relax with DFT.

With fixed values for all remaining fit parameters (including $E_0$ and DW factors), we calculated the corresponding reduced $\chi^2_r = \chi^2/\nu$ residual using the experimental and the calculated EXAFS spectrum of both $^5\text{B}_2$ and $^5\text{E}$ models. These results are shown in Fig. S5. As can be seen in the figure, in both cases a minimum $\chi^2_r$ value is reached; however, it always leads to significantly smaller values for the $^5\text{E}$ starting geometry, as compared to $^5\text{B}_2$. For comparison, we have plotted the DFT-optimized values for Fe–N\textsubscript{ax} and Fe–N\textsubscript{eq}, which are represented by vertical lines in Figs. S5A and S5B, respectively. We observe a consistent result by comparing the DFT-calculated Fe–N bond distances with the best-fit value, as the $^5\text{E}$ geometry is closest for both axial and equatorial N positions to the minimum $\chi^2_r$ positions obtained in our analysis.

5. **Diagrams of Kohn-Sham spin-unrestricted molecular orbitals**
Figure S6: A molecular orbital diagram of the quintet states of [Fe(terpy)$_2$]$^{2+}$, a more detailed version of Fig. 8 in the main text. $\alpha$ orbital symbols are left, $\beta$ are right of the orbital energy levels; those containing relevant $3d$-contribution are noted in parenthesis). Both the highly symmetric (D$_{2d}$), and Jahn-Teller distorted (C$_2$) states are shown for the $^5E$ (similarly to Fig. 8.)
Figure S7: Energy diagram of the DFT-calculated molecular orbitals with Fe p-orbital contributions of [Fe(terpy)$_2$]$^{2+}$ for the two quintet states; blue and yellow denotes $\alpha$ (spin-up) and $\beta$ (spin-down) orbitals, respectively. The apparent similarity for the structure and density of the unoccupied state levels in the two states suggest that their absorption spectra must be very similar.
References


