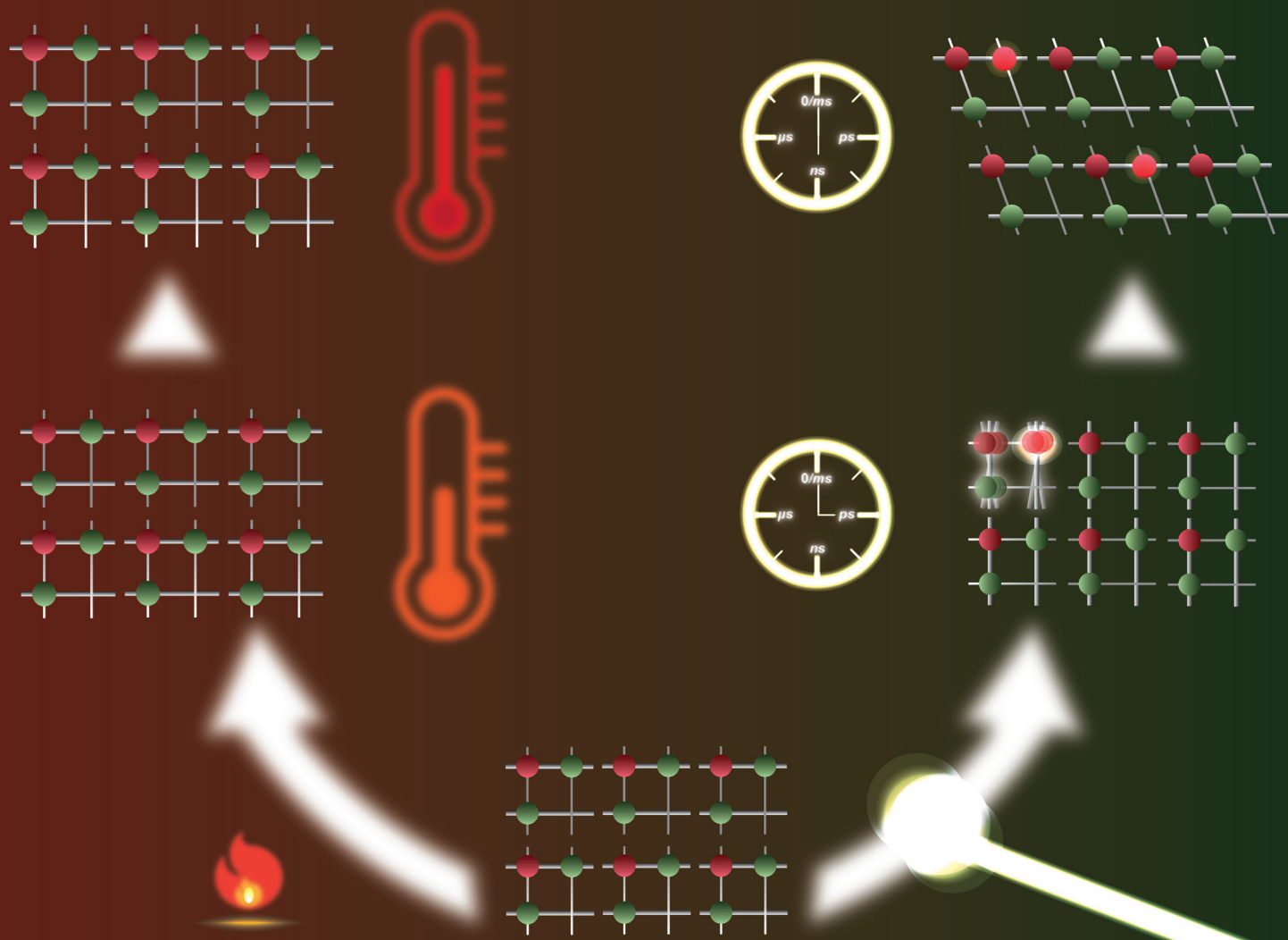


Dalton Transactions

An international journal of inorganic chemistry

rsc.li/dalton



ISSN 1477-9226

PAPER

Jose de Jesus Velazquez-Garcia, Simone Techert *et al.*
Structural dynamics of a thermally silent triiron(II) spin
crossover defect grid complex

Cite this: *Dalton Trans.*, 2023, **52**, 12224

Structural dynamics of a thermally silent triiron(II) spin crossover defect grid complex†

 Jose de Jesus Velazquez-Garcia,^a *a Krishnayan Basuroy,^b ^a
 Darina Storozhuk,^a Joanne Wong,^b Serhiy Demeshko,^b Franc Meyer,^b ^b
 Robert Henning^c and Simone Techert^b *a,d

The structural evolution of spin crossover (SCO) complexes during their spin transition at equilibrium and out-of-equilibrium conditions needs to be understood to enable their successful utilisation in displays, actuators and memory components. In this study, diffraction techniques were employed to study the structural changes accompanying the temperature increase and the light irradiation of a defect [2 × 2] triiron(II) metallogrid of the form [Fe₃L^H₂(HL^H)₂](BF₄)₄·4MeCN (**FE3**), L^H = 3,5-bis(6-(2,2'-bipyridyl))pyrazole. Although a multi-temperature crystallographic investigation on single crystals evidenced that the compound does not exhibit a thermal spin transition, the structural analysis of the defect grid suggests that the flexibility of the grid, provided by a metal-devoid vertex, leads to interesting characteristics that can be used for intermolecular cooperativity in related thermally responsive systems. Time-resolved photocystallography results reveal that upon excitation with a ps laser pulse, the defect grid shows the first two steps of the out-of-equilibrium process, namely the photoinduced and elastic steps, occurring at the ps and ns time scales, respectively. Similar to a previously reported [2 × 2] tetrairon(II) metallogrid, **FE3** exhibits a local distortion of the entire grid during the photoinduced step and a long-range distortion of the lattice during the elastic step. Although the lifetime of the pure photoinduced high spin (HS) state is longer in the tetranuclear grid than in the defect grid, suggesting that the global nuclearity plays a crucial role for the lifetime of the photoinduced species, the influence of the co-crystallising solvent on the lifetime of the photoinduced HS state remains unknown. This study sheds light on the out-of-equilibrium dynamics of a thermally silent defect triiron SCO metallogrid.

Received 1st July 2023,
Accepted 4th August 2023

DOI: 10.1039/d3dt02067c

rsc.li/dalton

Introduction

Spin crossover (SCO) complexes are switchable molecules capable of reversibly changing their spin state from a low-spin (LS) to a high-spin (HS) state under the influence of various external stimuli, such as temperature, pressure, electric field or light.^{1–5} Most of the vast number of SCO compounds are mononuclear, Fe^{II}-centered complexes where the LS state corresponds to $S = 0$ ($t_{2g}^6e_g^0$ configuration, singlet) and the HS

state to $S = 2$ ($t_{2g}^4e_g^2$, quintet).^{6–9} However, recent progress in supramolecular chemistry has allowed for the synthesis of a variety of discrete polynuclear SCO Fe^{II} complexes.^{10,11} [2 × 2] Fe₄^{II} metallogrids are an example of the latter type of complexes, in which four metal ions and four ditopic ligand strands give a matrix-like structure with up to four potentially addressable sites arranged in a square.^{12–17} This type of complex is appealing because it offers the potential for intramolecular cooperativity through the strong linking of the metal centres, which can be used in molecular logic devices, such as nanoscale memory devices and quantum automata, among others.¹⁸

In a series of publications, some of us have reported a family of metallogrid complexes with the general formula [Fe_XL^R₄], where $X = 2, 3, 4$, $L = R$ -3,5-bis{6-(2,2'-bipyridyl)}pyrazolate and $R = H$, methyl.^{19–22} This rational design approach has allowed for the customisation of various SCO properties, such as transition temperature, number of SCO steps and cooperative behaviour. For instance, the parent complex [Fe₄L^H₄](BF₄)₄·MeCN (**1**) undergoes an incomplete multistep thermal transition from [4HS] to [2LS–2HS] in the 300K to 50K range.

^aDeutsches Elektronen-Synchrotron DESY, Notkestr. 85, 22607 Hamburg, Germany.
E-mail: jose.velazquez@desy.de, simone.techert@desy.de

^bInstitut für Anorganische Chemie, Georg-August-Universität Göttingen,
Tammannstraße, 4, 37077, Göttingen, Germany

^cCenter for Advanced Radiation Sources, The University of Chicago, Argonne
National Laboratory, 9700 South Cass Ave, Lemont, Illinois, 90439, USA

^dInstitut für Röntgenphysik, Georg-August-Universität Göttingen, Friedrich-Hund-
Platz 1, Göttingen, 37077, Germany

† Electronic supplementary information (ESI) available. CCDC 2270541–2270546, 2270556–2270560, 2270561–2270563, 2270566–2270568, 2270570–2270574, 2270763–2270767. For ESI and crystallographic data in CIF or other electronic format see DOI: <https://doi.org/10.1039/d3dt02067c>



By introducing a methyl substituent to the C⁴ position of the pyrazolate ring, a metallogrid complex [Fe₄L^{Me}₄](BF₄)₄·2MeCN (**FE4**) was obtained. The stronger ligand field in **FE4** stabilises the LS fraction, resulting in a di-mixed-spin configuration [HS–LS–HS–LS] over a wider temperature range, from 7 to 250 K. Above 250 K, the complex undergoes a single gradual SCO and a change of configuration to [1LS–3HS]. Furthermore, the removal of one metal centre in **1** leads to the formation of a trinuclear Fe^{II} grid, also known as ‘defect grid’, of the form [Fe₃L^H₂(HL^H)₂](BF₄)₄·4MeCN (**FE3**), as shown in Fig. 1. The defect grid has the unique combination of strongly elastically coupled metal ions and a structurally soft metal-devoid vertex, which gives rise to a cooperative transition from [2LS–1HS] to [1LS–2HS] upon loss of the lattice solvent. Although this spin transition of **FE3** at equilibrium conditions has been reported previously,²² there is no information about the out-of-equilibrium dynamics of this grid.

The presence of solvent molecules in the lattice of the SCO complexes has varying effects on the spin transition. Previous studies have demonstrated that small solvent molecules can induce,^{23,24} alter^{25,26} or suppress^{27–31} the thermal spin transition of SCO complexes. Although some reports have demonstrated a photoinduced transition following light irradiation at low temperatures in iron complexes, where thermal spin transition is assisted by the desolvation process,^{29,30} there is no clear understanding of the effects of the solvent in the photo-

induced spin transition or the out-of-equilibrium dynamics of such complexes.

In response to a fs laser pulse, crystals of mononuclear SCO compounds exhibit complex out-of-equilibrium dynamics involving three consecutive steps: photoinduced, elastic and thermal steps, occurring at characteristic time-scales of picoseconds (ps), nanoseconds (ns) and microseconds (μs), respectively.^{32–36} The initial photoinduced step involves local conversion from LS to HS states *via* intersystem crossing through metal-to-ligand charge transfer (MLCT) excitation; this photo-switching process is proportional to the density of laser excitation, where one photon switches one metal centre from the LS to the HS state, accompanied by a structural reorganisation at the molecular scale. The second step, known as the ‘elastic step’, is associated with an additional conversion to the HS state and the volume expansion of the lattice at the material scale, which is not instantaneous and requires propagation of strain waves³⁷ at the speed of sound (~2000 ms⁻¹). Finally, the third step, referred to as the thermal step, occurs on the μs time scale when the average temperature of the crystal increases due to heat diffusion.

In our previous works,^{38,39} we employed time-resolved photocrystallography to investigate the out-of-equilibrium dynamics of **FE4**. Our findings revealed that the out-of-equilibrium dynamics triggered by a ps laser pulse follow the same sequence as observed in mononuclear SCO complexes. However, unlike mononuclear complexes, **FE4** exhibits two distinct types of elastic distortions occurring at different time scales. The first type is a short-range distortion, observed during the ps time scale, that propagates over the entire Fe₄ grid complex. This distortion results from the rearrangement of the coordination sphere of the photo-switching ion and the constant feedback between strongly linked metal ions. The second type takes place during the ns time scale and is a long-range distortion caused by the anisotropic expansion of the lattice, which is commonly observed in the dynamics of mononuclear SCO materials. The structural analysis of **FE4** demonstrated that the long-range distortion prevails over the short-range distortion, leading to the largest deformation of both the entire grid and the coordination sphere of each metal ion in the ns time scale.

However, the aforementioned studies do not provide information on the out-of-equilibrium dynamics of SCO complexes, where the solvent silences the thermal SCO, or in SCO defect grids. Therefore, this work aims to investigate the out-of-equilibrium structural dynamics of the thermally silent **FE3**. Since the structure of **FE3** at 133 K, as well as the magnetic properties and physicochemical characterisation, have been reported in a previous publication,²² this work focuses on exploring its out-of-equilibrium structural dynamics during the first 1 ns after light excitation, as well as the structural changes upon heating the grid from 100 K to 310 K. The multi-temperature structural analysis reveals that while **FE3** lacks a thermal spin transition, the flexibility of the grid, facilitated by a metal-devoid vertex, offers intriguing characteristics that may support intermolecular cooperativity in other thermally

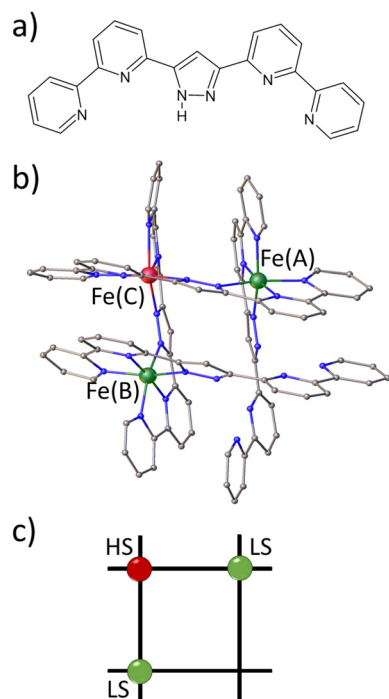


Fig. 1 (a) Pyrazole-bridged compartmental ligand HL. (b) Molecular representation of the [Fe₃L₄]⁴⁺ grid. Fe(A) and Fe(B) ions are in the LS state, and Fe(C) ions are in the HS state. Counter ions, solvent molecules and hydrogen atoms are omitted for clarity. (c) Schematic representation of the grid in **FE3**.



responsive systems. More interestingly, we show that upon excitation, **FE3** presents the first two steps of the out-of-equilibrium process, as well as the two types of elastic distortions observed in **FE4**. A comparison between the **FE3** and **FE4** grids demonstrates that the photo-switched HS state relaxes more rapidly in **FE3** than in **FE4**.

Methods

Synthesis and steady-state spectroscopy

The **FE3** metallogrid and its ligands were synthesised following the procedure reported in the literature.²²

UV-vis spectra were collected at room temperature on a Varian Cary-5E UV-vis Spectrometer. All data were collected at a scan rate of 0.2 nm s⁻¹ over the wavelength range of 200–800 nm with an interval size of 0.5 nm. The absorption spectrum of **FE3** was collected on 10 mm cuvettes, using a reference spectrum of acetonitrile, which was also used to dissolve the compound.

X-ray diffraction

Single-crystal X-ray diffraction experiments were performed from 100 K to 310 K (the maximum temperature possible before crystal decay became unacceptable). The X-ray data were collected in phi scan mode on undulator synchrotron radiation with $\lambda = 0.6199 \text{ \AA}$ at P11 beamline in PETRA III, DESY, Hamburg, Germany. Indexing of the X-ray diffraction pattern, unit cell refinement and spot integration were performed using the XDS program package.⁴⁰ The crystal structures were solved with the SHELXT⁴¹ structure solution program within the Olex2 software package.⁴² Subsequently, the structures were refined against F² isotropically, followed by full matrix anisotropic least squares refinement by SHELXL.⁴³ For all structures, hydrogen atoms were fixed geometrically, in idealised positions, and allowed to ride with the respective C or N atoms, to which each was bonded, in the final cycle of refinement. Crystal data and structure refinement parameters for the 100 K dataset are provided in Table S1,† while the selected bond lengths and bond angles are given in Table S2.† The asymmetric unit cell, thermal ellipsoid plot and packing of molecules down all crystallographic axes at 100 K are given in Fig. S1–S5.

Time-resolved pink Laue crystallography

Time-resolved X-ray diffraction (TRXRD) measurements were performed at the BioCARS beamline at the Advanced Photon Source, Argonne National Laboratory, IL, USA. TRXRD data were collected at 15KeV with a Rayonix MX340-HS detector. Pulses from a Ti:sapphire laser tuned to 390 nm wavelength were used as a pump source and set perpendicular to the φ -rotation axis. The time delay between the laser pump and the X-ray probe was set by varying the arrival times of the picosecond laser pulses with respect to the synchrotron X-ray pulses. Delay times (dt) between the laser pump and the X-ray probe were selected at 200 ps, 500 ps, 700 ps and 1 ns. A refer-

ence measurement for each crystal was taken at the negative delay time (dt < 0) of –500 ps. Laser power was set at 2.0 (mJ mm⁻²) per pulse. Crystallographic data were acquired at 100 K since the lifetime of the locally excited molecules is larger at low temperatures. The latter temperature was limited by the equipment capability. The recorded data sets for each crystal covered a 180° scan with 1° step.

Since crystals of **FE3** showed a fast decay under laser exposure, a typical data collection strategy of 5 laser-OFF/ON cycle used in our previous works^{38,39,44} was not recommended. Therefore, a strategy of one laser-ON at dt < 0 followed by all laser-ON at dt was used for each crystal. The measurement was repeated in five different crystals for each compound. Crystals used in this study had approximate dimensions of 90 × 50 × 30 μm³.

The pink Laue diffraction images were processed with the software package Precognition/Epinorm,⁴⁵ using variable elliptical integration for all data sets. After integration, the dt and dt < 0 data sets were scaled together, but the repetitive and symmetry-related reflections from both data sets were merged separately. These data sets were used to obtain the photodifference maps and the final structural models at each delay time. For all data sets, initial models of the crystal structures were taken from those obtained from the monochromatic X-ray diffraction experiment at 100 K. Subsequently, the structures were refined using the SHELXL program within the Olex2 software package. Thermal ellipsoid plots (50% of probability) for all time-resolved data sets are provided in ESI (Fig. S6–S10).†

Modification of the coordination environment

The Octadist⁴⁶ program was used to characterise the coordination geometry of the metal ions in each compound, in particular the ⟨Fe–N⟩ bond length distance and the angular distortion parameter (θ). The angular distortion parameter is the sum of the deviations from 60° of the 24 N–Fe–N angles, six per pseudo three-fold axis, measured on a projection of opposite triangular faces of the FeN₆ octahedron, orientated by superimposing the face centroids [see Fig. S11†]. For comparison, continuous symmetry measurements⁴⁷ calculated using the SHAPE program⁴⁸ were also used to characterise the change in symmetry and shape of the metal coordination sphere. The calculation of $S(\text{Oh})$ and $S(\text{itp})$, which compare the coordination sphere with respect to a perfect octahedron and trigonal prism, respectively, were done for all crystallographically independent metal ions from structures at 100 K, 250 K and 310 K (see ESI† for more information). Both θ and $S(\text{Oh})$ parameters are small and closer to 0 when the metal centre is in LS state, while $S(\text{itp}) \gg 0$, and the opposite is shown for metal centres in HS state.

Analysis of temperature increase due to laser exposure

An estimate of the temperature change between the dt and dt < 0 data sets was obtained from the well-known photo-Wilson plots.^{49,50} The plots for all time-resolved data are shown in Fig. S13.† The slope of the plot corresponds to twice the difference of the Debye–Waller factor ($2\Delta B^{\text{dt}-\text{dt}<0}$), associated with



the isotropic atomic motion (for more information see the temperature difference section in ESI†). Therefore, measuring ΔB between the dt and $dt < 0$ data sets provides an estimate of the temperature increase due to laser exposure through the associated increase in thermal motion. Values of $\Delta B^{dt-dt<0} \approx 0$ indicate a non-detectable global heating of the sample, while values greater than zero suggest an increase in temperature of the crystal.

Results

Multi-temperature structural analysis

Structures at initial and final conditions. Fig. 1b and 1c show the structure of the cation in FE3 and its schematic representation, respectively. For clarity, Fe^{II} ions are labelled by their corresponding corner on the grid, with LS Fe^{II} ions occupying corners A and B and the HS ion occupying corner C. To differentiate between the two LS ions, corner A is assigned to the LS Fe^{II} with the higher distortion in its coordination sphere at 100 K.

Table 1 provides the geometrical characteristics of the FeN₆ octahedra in FE3 at both 100 K and 310 K. A more detailed structural description of the compound is provided in Table S3.† The structural analysis confirms that the FE3 defect grid maintains its $P\bar{1}$ space group across the temperature range. At both temperatures, four acetonitrile molecules were observed cocrystallising in the lattice, from which two were refined as disordered over two positions (Fig. S1†). As reported in the literature,²² the asymmetric unit of the compound reveals that all metal ions are crystallographically independent with a molecular configuration of [LS–HS–LS], which was deduced from analysing the coordination sphere of each metal ion (and in accordance with the results of ⁵⁷Fe Mössbauer spectroscopy²²). The analysis of the coordination geometry demonstrates that LS and HS ions exhibit typical values of $\langle \text{Fe–N} \rangle$ bond lengths, with LS atoms between 1.95 and 1.97 Å and HS atoms around 2.18 Å. A further difference between the metal centres is the distortion of the {FeN₆} coordination environment from an ideal octahedron. The less distorted FeN₆ octahedron is found for the LS ions, Fe(A) and Fe(B) with $\theta = 301\text{--}314^\circ$, $S(\text{Oh}) = 2.1\text{--}2.3$, while the HS ion, Fe(C), displays a greater distortion of the coordination sphere with $\theta = 570^\circ$ and $S(\text{Oh}) = 6.0$.

Metal ion coordination geometry. The multi-temperature study presented here provides insight into the temperature

Table 1 Geometry changes in the coordination sphere of the metal centres in FE3 with the temperature increase

Metal	100 K			310 K		
	$\langle \text{Fe–N} \rangle$ (Å)	θ (°)	$S(\text{Oh})$	$\langle \text{Fe–N} \rangle$ (Å)	θ (°)	$S(\text{Oh})$
Fe(A)	1.966	306.1	2.256	1.974	313.5	2.332
Fe(B)	1.959	301.9	2.183	1.968	310.5	2.284
Fe(C)	2.183	577.3	6.140	2.188	575.7	6.160

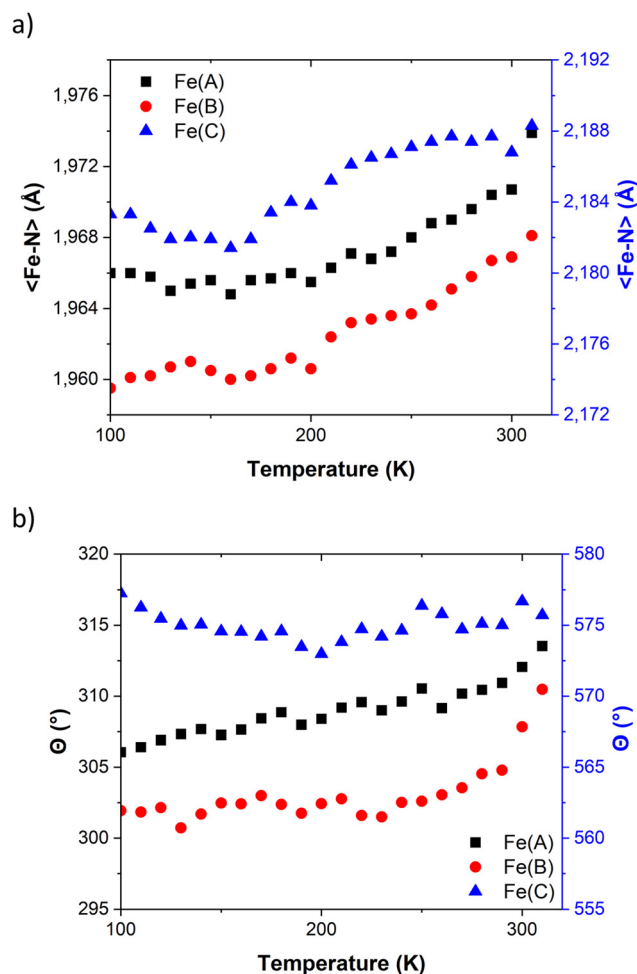


Fig. 2 Temperature dependence of (a) $\langle \text{Fe–N} \rangle$ bond length distances and (b) angular distortions (θ) in FE3. Values for Fe(A) and Fe(B) share the same axis.

dependence of the $\langle \text{Fe–N} \rangle$ bond lengths and the distortion of the coordination sphere, as measured by the θ parameter, for each metal ion in the FE3 defect grid. As shown in Fig. 2a, the average metal–ligand bond length expands by less than 0.01 Å for all three metal ions. It is worth noting that Fe(A) exhibits a noticeable increase in the $\langle \text{Fe–N} \rangle$ bond lengths around 260 K, which indicates the beginning of the LS \rightarrow HS transition for that ion. However, this is attributed to the loss of the co-crystallisation solvent rather than a feature of the solvated sample, as stated above (see Introduction).

The temperature dependence of θ (Fig. 2b) indicates that below 270 K there is no notable change in distortion of the {FeN₆} octahedrons as the temperature increases. However, following the solvent loss and a potential small LS \rightarrow HS transition, a slight change in the angular distortion of the Fe(A) and Fe(B) ions is observed. Notably, the Fe(B) ion exhibits a comparatively larger change in θ , suggesting that it may undergo the most significant distortion during the spin transition.

Molecular geometry. The metal devoid vertex plays an important role in the thermal expansion of the FE3 grid. As depicted



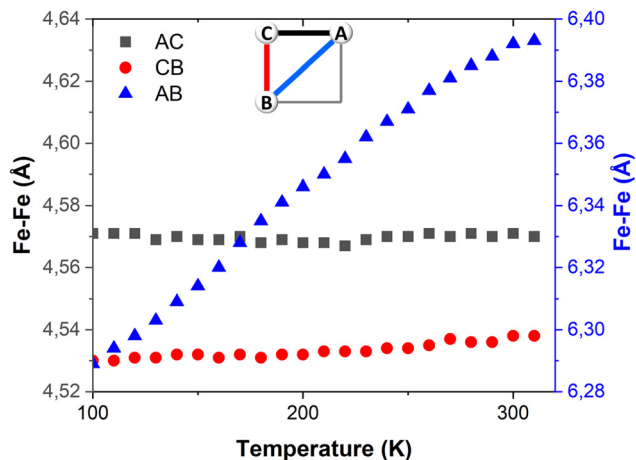


Fig. 3 Temperature dependence of the Fe–Fe distances in FE3. Distances AC and CD share the same axis.

in Fig. 3, the temperature dependence of the Fe–Fe distances shows that the thermal expansion of the parallelogram in FE3 is primarily driven by the separation between ligands at the metal devoid vertex. In contrast, the Fe–Fe distance between strongly linked pyrazolato-bridged metal ions remains relatively constant, showing a negligible impact on the overall grid thermal expansion.

Disorder. One important feature to consider during the multi-temperature analysis is the increase in disorder with temperature. Two types of disorder are observed: the static disorder of the solvent molecules and the dynamic disorder of the pyridyl group. Two solvent molecules were refined as disordered over two positions, which is observed at 100 K (Fig. S1†). As the temperature continues to rise, accurately modelling the disordered solvent molecules becomes more challenging, especially beyond 270 K, when solvent evaporation is expected to occur. In the case of the dynamic disorder, the rotation of the pendant pyridyl ring at the metal-devoid site initiates around 180 K and increases gradually until the end of the multi-temperature experiment (Fig. S13†). Since the disordered solvent molecules are found in the proximity to the rotating pyridyl ring (Fig. S14†), it is possible that the increase of disorder in the solvent and the evaporation of it contribute to the larger rotation of the pyridyl group as the temperature rises.

Intermolecular interactions. The Hirshfeld surface^{51,52} approach, utilizing fingerprint representations, was employed to analyse and summarize the intermolecular interactions within the FE3 grid. Hirshfeld surfaces were calculated using CrystalExplorer 17.5 software.⁵³ In Table S3,† two-dimensional fingerprint plots of (di, de) points from all contacts contributing to the Hirshfeld surface analysis are presented for all atoms in the FE3 grid at temperatures of 100 K, 250 K, and 310 K. The fingerprint plots in the table provide valuable insights into the dominant intermolecular interactions in the crystal packing. Notably, H...H contacts are the most significant, constituting between 45.1% and 47.7% of the overall

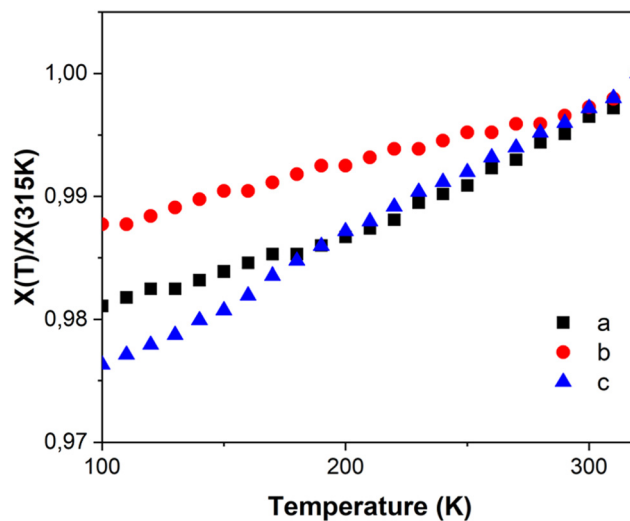


Fig. 4 Relative temperature dependence of unit cell parameters of FE3.

crystal packing. Other important interactions and their respective contributions include H...F (25.9–27.6%), C...H (9.8–10.3%), H...N (6.3–6.7%), H...C (4.4–4.7%), C...C (1.7–2.2%), and C...N (1–1.7%). While the contribution of each contact remains relatively stable across temperatures, noteworthy changes in the distances of H...N, H...C, and H...H interactions occur as the temperature increases from 100 K to 250 K or 310 K. These changes can be attributed to solvent disorder and the rotation of the pyridyl group, as mentioned earlier. C...C interactions, which are primarily associated with π ... π interactions, they exhibit weaker strengths within the FE3 grid. This is evident from the substantial separation between π - π planes, measuring 4.2–4.8 Å at 100 K (see Table S3†). Moreover, it is important to note that the rotation of the pyridyl group at 250 K and 310 K leads to an increased separation between π - π planes up to 5.1 Å.

Unit cell. Although the FE3 cation shows a larger expansion than the defect-free grid FE4, this does not have a significant impact on the expansion of the unit cell parameters. Fig. 4 displays the temperature dependence of the unit-cell dimensions, which exhibit an anisotropic behaviour with no abrupt modification. A quasi-linear expansion of all unit cell parameters in FE3 is observed with the increase in temperature. This suggests that there is no significant effect due to any spin transition or by the rotation of pyridyl groups or a sudden positional change of counter ions and solvent molecules as observed by other systems.^{54,55}

Out-of-equilibrium process

Optical measurements. Fig. 5 displays the UV-vis absorption spectrum of FE3 in an acetonitrile solution at room temperature, covering the 200–800 nm range. The spectrum reveals similar features to the previously reported spectrum of the defect-free FE4 grid. The bands below 350 nm are attributed to ligand-based $\pi \rightarrow \pi^*$ transitions, while the bands spanning the 450–700 nm range are assigned to MLCT transitions of mixed



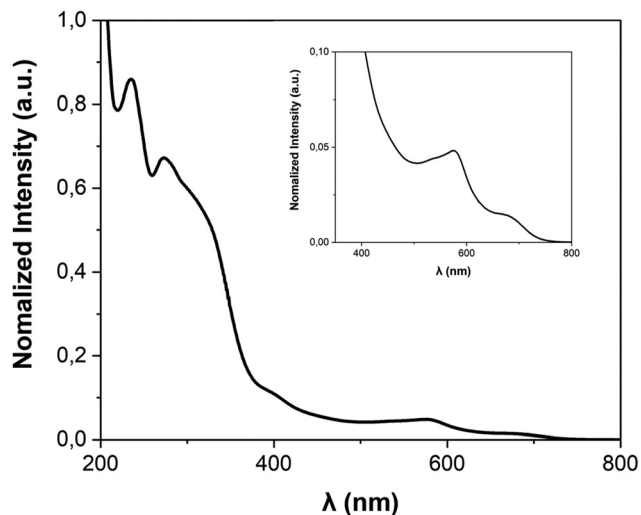


Fig. 5 UV-vis spectrum of FE3 grid in acetonitrile. (inset) Enlargement of the 350–800 nm range.

singlet–triplet character. As with FE4, the photocrystallographic experiment for FE3 employed an irradiation wavelength at the absorption tail of the MLCT band (390 nm) to avoid excessive photon absorption at the surface of the crystal that hinders laser penetration.

Photodifference maps. The light-induced changes are illustrated in photodifference maps of the form $F_{\text{obs}}^{\text{dt}} - F_{\text{obs}}^{\text{dt}<0}$ based on all independent reflections with $I/\sigma(I) > 3$ (Fig. 6). To highlight the Fe displacements, isosurfaces are drawn between ± 0.08 and 0.18 e \AA^{-3} . The molecular response of the FE3 defect

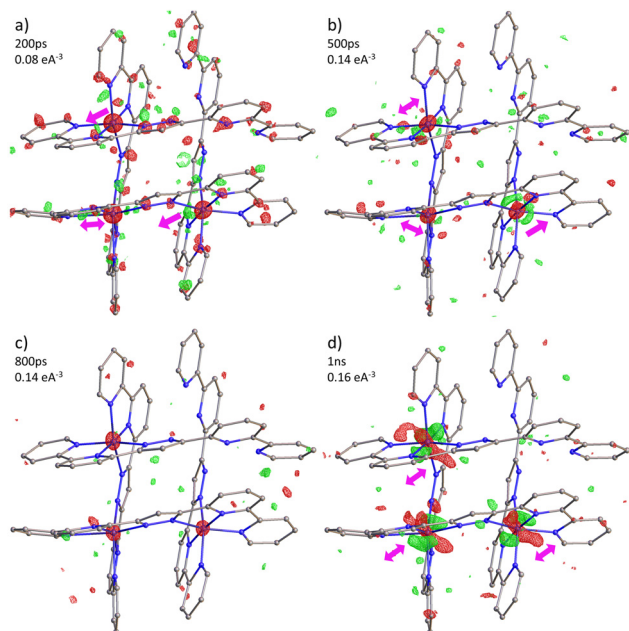


Fig. 6 Photodifference maps of FE3 during (a–c) the photoinduced step and (d) the elastic step. Isosurfaces (green positive, red negative) and delay times are shown in each figure. Arrows give an approximate direction of the electron density shift.

grid on excitation at any time is illustrated by the shift of electron density of all iron ions. Similar to the FE4 grid, the photodifference maps of the defect grid before 800 ps suggest a continuous change in the direction of the electron density due to a constant reorganisation of the grid. Unlike the FE4, where only two metal ions were crystallographic-symmetry independent, here, it is possible to observe the individual movement of all metal ions in the grid. The maps show that each iron ion moves in a different direction at delay times of 200 ps and 500 ps, suggesting that this constant arrangement is not caused by lattice expansion or by thermal motion. The photodifference maps at 800 ps suggest that most of the photo-induced step is almost over, similar to what was found in FE4 at 220 K. Finally, the photodifference maps at 1 ns exhibit an electron shift of all metal ions towards the 'a' axis, suggesting a movement due to the anisotropic thermal expansion of the lattice along such axis and the beginning of the elastic step.

Structural response. The temporal evolution of the average Fe–N bond length of all Fe^{II} ions and the variation of the trigonal distortion parameter are presented in Fig. 7. The results in the graphs agree with the photodifference maps showing two structural responses at different times. The first is the expansion in $\langle \text{Fe–N} \rangle$ bond length of both Fe(A) and Fe(B) ions, which is attributed to the local LS \rightarrow HS photoswitching in the photoinduced step. This step has a duration of less than 800 ps, after which both ions relax to their original states. Since the grid has returned mostly to its ground state configuration, the second expansion of the $\langle \text{Fe–N} \rangle$ bond lengths of Fe(A) and Fe(B) is attributed to the elastic step and is caused by the anisotropic thermal expansion of the lattice, as suggested by the photodifference map at 1 ns delay time.

Although the elongation of the $\langle \text{Fe–N} \rangle$ bond lengths in both ions, which originally are in the LS state, may suggest a transition of the entire grid from a 2LS–1HS to a 3HS configuration, we consider an alternative scenario more likely where only one metal ion in the LS state switches to the HS state. If Fe(A) and Fe(B) ions have a similar probability to transit to a HS state upon excitation, the change in configuration from 2LS–1HS to 1LS–2HS in the crystal would result in a mixture of molecules with only Fe(A) or Fe(B) as the transition ion. Since crystallography provides an average representation of all molecules in the crystal, the expansion of the $\langle \text{Fe–N} \rangle$ bond lengths will be observed in both ions in the asymmetric unit, regardless of whether the structural changes belong to different molecules.

The variation of the trigonal distortion parameter in the time domain for the transiting metal ions Fe(A) and Fe(B) is shown in Fig. 7b. The figure reveals a slight increase in the distortion of the coordination sphere of both transiting metal ions during the photoinduced step. Notably, the Fe(B) ions exhibit a more pronounced distortion, consistent with the observations from the multi-temperature study. As observed in our previous studies on the FE4 grid, the distortion of the coordination spheres of both transiting atoms are larger during the elastic step than in the photoinduced step, as observed in the value of $\Delta\theta$ at 1 ns delay time.



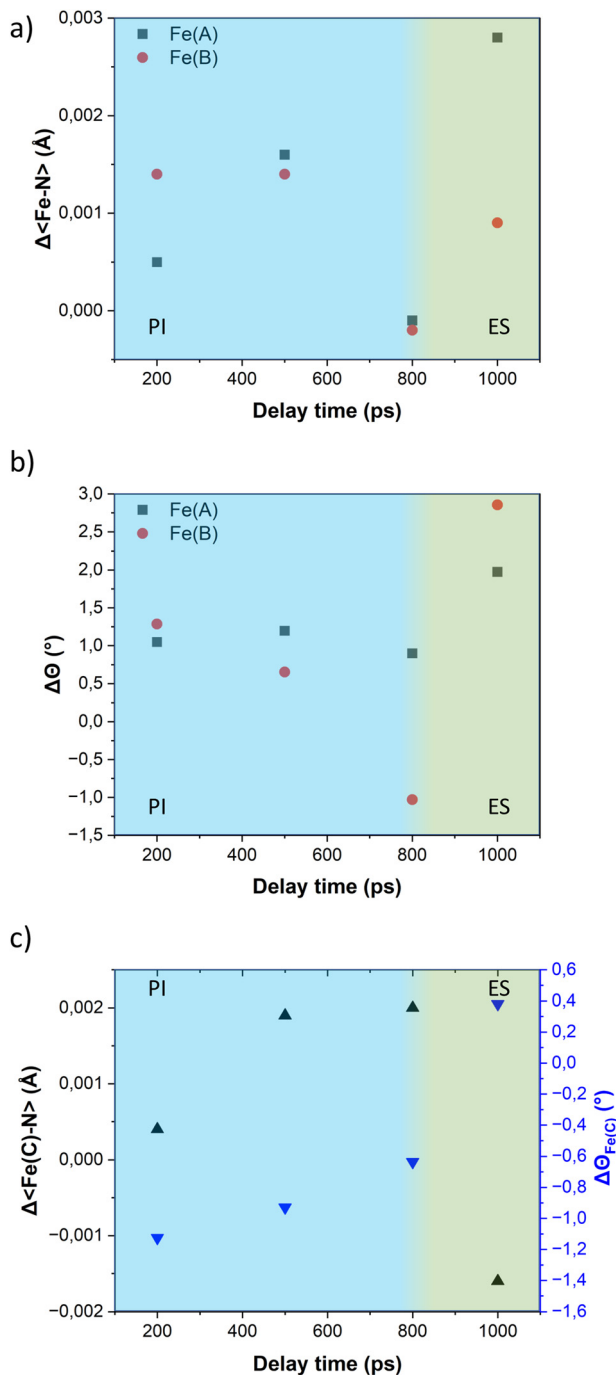


Fig. 7 Time-delay dependence of (a) $\Delta\langle\text{Fe-N}\rangle$; (b) $\Delta\theta$ of the Fe(A) and Fe(B) ions; and (c) those from the Fe(C) ion. Photoinduced (PI) and elastic (ES) steps are shown in blue and green background, respectively.

Similar to the observations for the FE4 grid, the photo-switching at the Fe(A) and Fe(B) ions of the FE3 grid leads to changes in the coordination sphere of the adjacent metal ion in the HS state, Fe(C). Fig. 7c displays the expansion of the $\langle\text{Fe-N}\rangle$ bond length of Fe(C) ion and the concomitant change in the distortion of its coordination geometry following light irradiation. However, it should be noted that the changes are not related to any spin transition since the Fe(C) ion remains

in the HS state throughout the experiment. Instead, these changes in its coordination sphere are attributed to a knock-on effect from the structural reorganisation of the whole grid following the spin transition in the neighbouring ions.

Photoconversion and thermal factor. In the time domain, the increase in population of the HS ions leads to an expansion of the $\langle\text{Fe-N}\rangle$ of the Fe(A) and Fe(B) ions. Therefore, the variation of the HS fraction can be estimated based on the time variation of the $\langle\text{Fe-N}\rangle$ according to the expression:

$$\% \Delta X_{\text{HS}} = \frac{\langle\text{Fe(A)-N}\rangle^{\text{dt}} - \langle\text{Fe(A)-N}\rangle^{\text{dt} < 0}}{\langle\text{Fe(A)-N}\rangle_{\text{HS}} - \langle\text{Fe(A)-N}\rangle_{\text{LS}}} + \frac{\langle\text{Fe(B)-N}\rangle^{\text{dt}} - \langle\text{Fe(B)-N}\rangle^{\text{dt} < 0}}{\langle\text{Fe(B)-N}\rangle_{\text{HS}} - \langle\text{Fe(B)-N}\rangle_{\text{LS}}} \quad (1)$$

where $\langle\text{Fe-N}\rangle_{\text{HS}}$ and $\langle\text{Fe-N}\rangle_{\text{LS}}$ are the average metal-ligand bond lengths of the transiting ions in the HS and the LS state, respectively. Since the $\langle\text{Fe-N}\rangle_{\text{HS}}$ value is unknown, the denominators in the equation can be replaced by 0.2 Å, the maximum bond-length expansion expected for both ions. Consequently, the variation of the HS state can be obtained by:

$$\% \Delta X_{\text{HS}} = \frac{\Delta\langle\text{Fe(A)-N}\rangle + \Delta\langle\text{Fe(B)-N}\rangle}{0.2 \text{ \AA}} \quad (2)$$

where the $\Delta\langle\text{Fe-N}\rangle$ is the variation of the average metal-ligand bond length of the transiting ions at delay times dt with respect to the reference measurement dt < 0.

Table 2 presents the change in the HS-state molecular fraction from 200 ps to 1 ns, calculated from the expansion of the $\langle\text{Fe-N}\rangle$ bond length of the Fe(A) and Fe(B) ions. The estimated population of the excited state during the photoinduced step (dt < 800 ps) is in the 1–1.5% range, while ~2% is estimated for the delay time in the elastic step (dt > 800 ps). The change in $\% \Delta X_{\text{HS}}$ is slightly larger in the elastic step than in the photoinduced step as typically reported in monometallic materials, where photoconversion of the elastic and thermal steps tends to overcome the photoinduced step.³² At 800 ps, most of the molecules relax to their ground state shown by the low value of the $\% \Delta X_{\text{HS}}$ and in agreement with the photo-difference maps.

In the studied range of time, the LS \rightarrow HS transition occurs without any significant energy exchange with the environment, as observed in the time-delay dependence of the isotropic thermal factor, $\Delta B^{\text{dt}-\text{dt} < 0}$. Table 2 shows that the values of $\Delta B^{\text{dt}-\text{dt} < 0}$ are close to zero and comparable with those reported for the FE4 grid in the photoinduced step and beginning of the elastic step. This suggests that heating of the crystal is not detectable within the measured range and that the two

Table 2 Change in the HS-state molecular fraction and ΔB values from photo-Wilson plots

	200 ps	500 ps	800 ps	1 ns
$\% \Delta X_{\text{HS}}$	1.0 (13)	1.5 (13)	-0.2 (12)	1.9 (13)
$\Delta B^{\text{dt}-\text{dt} < 0}$	0.00 (6)	0.07 (3)	0.00 (2)	0.06 (3)



increases in the % ΔX_{HS} , one before and one after 800 ps, are solely attributed to the photoinduced step and elastic step, respectively.

Discussion

One of the most ambitious goals of synthetic chemists in the SCO field is the rational design of SCO systems with tailored properties. With the aim of enhancing the SCO properties, *e.g.* abrupt, hysteretic and multistep switching, the design of SCO complexes has explored various strategies, including varying the number of metal centres (ranging from mononuclear to polymeric) and adopting diverse geometries, such as squares, cages and grids. One potential approach has been the utilisation of $[2 \times 2]$ grids formed by highly constrained ligands, which promote the communication between metal centres. Tetranuclear grids are rigid molecules, which restricts the overall movement of the architecture in favour of the communication between the linked metal centres. While such communication may lead to intramolecular cooperativity,⁵⁶ there is no clear evidence suggesting that $[2 \times 2]$ grid-like arrangements promote intermolecular cooperativity. However, defect metalogrids with a metal-devoid flexible vertex may present a solution to this problem.

The present multi-temperature crystallographic analysis sheds light on two interesting characteristics of such defect grids. Firstly, the **FE3** defect grid displays a larger thermal expansion in its geometrical dimension compared to the more constrained **FE4** grid, primarily due to the reduced restraint provided by the metal-devoid site. Secondly, the rotation of the pyridyl rings at this site offers a potential avenue for introducing long-range elastic interaction capable of modifying the unit cell expansion, as observed in other compounds.⁵⁵ Although **FE3** does not exhibit thermal SCO, these two characteristics of defect grids hold promise for promoting intermolecular cooperativity in other thermally responsive SCO grids. Thus, defect grids with one metal-devoid vertex and communicating metal ions may present an alternative strategy to enhance both the intermolecular cooperativity, by increasing the flexibility of the grid, and intramolecular cooperativity, provided by the strong linking of the metal ions. However, it is indispensable to consider other factors that can affect the SCO and the cooperativity, such as the lattice type, counter ions and solvent effects.

Solvent silencing is a well-known phenomenon that occurs in SCO complexes where the solvent delays the spin transition to higher temperatures or prevents it from happening. Numerous SCO complexes, including the **FE3** grid, have been reported to be silenced by the presence of a specific solvent molecule within the crystal lattice.^{22,27–31} However, the out-of-equilibrium dynamics in this type of complexes remains largely unexplored. In this study, we observed that while the solvent suppresses the thermal-induced SCO in **FE3**, two out of the three consecutive steps of the out-of-equilibrium process are observed following light irradiation.

In line with the observation in mononuclear Fe^{II} and Fe^{III} complexes, as well as in the tetranuclear **FE4** grid, excitation of the **FE3** grid with a ps laser pulse initiates a multistep process, with each step occurring at distinct time scales.^{32–36} First, the light absorption leads to a local LS \rightarrow HS transition in a fraction of molecules, resulting in the conversion of one of the Fe^{II} ions from the LS to the HS state. The spin flip is accompanied by the reorganisation of the $\{\text{FeN}_6\}$ coordination sphere of the transiting ion, which triggers a short-range elastic distortion that propagates through the grid due to the tightly linked metal ions. This distortion develops within the first few hundreds of picoseconds after light irradiation, and it is evidenced by the distinct shift in electron density observed in the photo-difference maps at 200 ps and 500 ps. Notably, this molecular reorganisation resembles the behaviour observed in the **FE4** grid, reinforcing the idea that the strong linking of metal ions leads to an elastic metal-to-metal communication during the spin transition. Subsequently, the elastic step takes place when the lattice heating, resulting from the energy dissipation, induces the coherent propagation of strain waves. This leads to lattice expansion and crystal deformation at the material scale, causing all metallic ions to move in the direction of the propagating strain waves, as observed in the photodifference maps at 1 ns. The internal pressure generated at this step leads to the second LS \rightarrow HS switching, as reported in Table 2. Although the dynamics beyond 1 ns are not reported in this study, this second step typically lasts until the beginning of the μs time scale, after which it is followed by the thermal step when the energy distribution reaches a global temperature rise on the crystal. However, since the **FE3** grid is thermally silent, this step is expected not to be present. Although the suppression of the thermal step in the out-of-equilibrium dynamics of **FE3** could be perceived as a limitation, it provides valuable insights about the tailoring of the response of the SCO materials upon light excitation. Given that the spin switching during the elastic and thermal steps usually prevails over the pure photoinduced spin flip,³² the knowledge of how to limit these steps by chemical engineering is a valuable resource.

Comparing the lifetime of the photoinduced HS state in the **FE3** defect grid with that observed in the **FE4** grid suggests that the metastable HS state resulting from pure photo-switching exhibits a longer lifetime in **FE4** than in **FE3**. While our previous report³⁹ indicated that the photoinduced step in **FE4** lasts approximately 800 ps, similar to **FE3**, those studies were conducted at a significantly higher temperature of 220 K, compared to 100 K used in this work. As the lifetime of the metastable states is known to be longer at lower temperatures, the duration of the photoinduced HS state in the **FE4** at 100 K is estimated to be considerably longer than 800 ps (>2 ns), as suggested by our first studies on the **FE4** grid.³⁸ This observation aligns with time-resolved X-ray spectroscopic studies performed in solution, where the parent grid **1** exhibited a longer lifetime of the photoexcited state (~ 210 ns) compared to the **FE3** grid (~ 123 ns).^{57,58} Thus, the results here reinforce the idea that global nuclearity of the architecture plays a pivotal role in tailoring the lifetime of the photoinduced



species. However, it is important to consider that the presence of the solvent in both grids may also influence the lifetime of the pure photoinduced HS state, and therefore, more research is needed to understand its influence.

It is worth noting that the present study has been limited to a few data points within the first nanosecond after laser irradiation of a defect grid, whose thermal transition from a 2LS–1HS to a 1LS–2HS state is silenced by the presence of the co-crystallising acetonitrile solvent. Further investigations are necessary to obtain a detailed picture of the structural changes over extended timescales ranging from picoseconds to microseconds. Additionally, studies in a solvent-free grid and with other solvents would provide valuable information about the influence of the solvent on the lifetime of the metastable state and the structural changes occurring in each step of the out-of-equilibrium dynamics. This also helps to differentiate between the effects of the solvent and the metal-devoid site in the grid. A comprehensive analysis of all factors influencing the out-of-equilibrium dynamics of oligomeric SCO complexes would provide crucial insights for designing new SCO materials with tailored properties.

Conclusions

The rational design of spin crossover systems with diverse nuclearity and topology has emerged as a powerful approach to enhance cooperative effects and expand the variety of SCO properties, considering their potential application in nanoscale memory devices and quantum automata, among others. To effectively engineer SCO materials with tailored properties, it is crucial to develop a comprehensive understanding of factors influencing the spin transition in solid state, both at equilibrium and out-of-equilibrium conditions. Some significant factors include the number and arrangement of metal centres as well as the presence of solvent within the crystal lattice. In this study, we presented the structural analysis of a defect $[2 \times 2]$ metallogrid with one metal-devoid vertex, where the co-crystallising solvent silenced the thermal spin transition while allowing for the photoinduced transition at out-of-equilibrium conditions.

Although the studied system **FE3** does not exhibit thermal SCO, its structural flexibility, facilitated by the metal-devoid site, imparts attractive characteristics that can potentially favour the SCO in related systems, that is the large thermal expansion of the overall structure and rotation of the pyridyl rings at the metal-devoid site. These features are attractive for introducing long-range elastic interactions that can favour intermolecular cooperativity in other defect grids whose SCO is not affected by the solvent. Despite the solvent induced silencing of the thermal SCO transition, **FE3** undergoes two of the three steps of the out-of-equilibrium process. Upon irradiation with a ps laser pulse, the **FE3** defect grid exhibits the photoinduced and elastic steps occurring at ps and ns time scales, respectively. Each step introduces an increase in the fraction of the HS state and a different reorganisation. The

structural reorganisation of **FE3** during the out-of-equilibrium process is similar to that of the previously reported **FE4** grid, involving a short-range distortion appearing from the continuous molecular rearrangement of the grid during the photoinduced step, triggered by LS \rightarrow HS photo-switching of only one metal ion, and a long-range distortion caused by the anisotropic deformation of the lattice during the elastic step.

Compared to **FE4**, the pure photoinduced HS state in **FE3** relaxes back to its original LS state in ~ 800 ps at 100 K, while in **FE4**, it takes longer than 2 ns at 100 K and ~ 800 ps at 220 K. This observation supports previous time-resolved X-ray spectroscopic findings in solution and suggests that global nuclearity plays a crucial role in the lifetime of the photoinduced species. However, the influence of the co-crystallising solvent on the lifetime of the photoinduced HS state remains unknown and requires further research with other solvents and solvent-free grids. Understanding the solvent effect at each step of the out-of-equilibrium process is essential for controlling the properties of SCO materials with bi- and multi-stability at equilibrium and out-of-equilibrium conditions, laying the foundations for their practical applications.

Author contributions

JJVG drafted the manuscript. JJVG, KB and DS performed the time-resolved crystallographic experiment with assistance from RH in beamline operation at APS. ST and FM conceived the study. FM and SD designed the molecules. JW synthesised the material. All authors provided input and agreed on the final manuscript.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

This research used resources of the Advanced Photon Source, a US Department of Energy (DOE) Office of Science User Facility operated for the DOE Office of Science by Argonne National Laboratory under contract no. DE-AC02-06CH11357. Use of BioCARS was also supported by the National Institute of General Medical Sciences of the National Institutes of Health under grant number P41 GM118217. Time-resolved set-up at Sector 14 was funded in part through a collaboration with Philip Anfinrud (NIH/NIDDK). The content is solely the responsibility of the authors and does not necessarily represent the official views of the National Institutes of Health. Portions of this research were carried out at the light source PETRA-III at DESY, a member of the Helmholtz Association (HGF). We would like to thank P11 staff for assistance in using beamline P11. The current work has been funded by the Deutsche Forschungsgemeinschaft (DFG, German Research Foundation) – 217133147/SFB 1073, projects B06, C02. HG-



recruitment, HG-Innovation “ECRAPs”, HG-Innovation DSF/DASHH and CMWS.

References

- P. Gütllich, A. Hauser and H. Spiering, *Angew. Chem., Int. Ed. Engl.*, 1994, **33**, 2024–2054.
- P. Gütllich and H. A. Goodwin, *Spin Crossover in Transition Metal Compounds I-III*, Springer Berlin Heidelberg, Berlin, Heidelberg, 2004.
- A. Bousseksou, G. Molnár, L. Salmon and W. Nicolazzi, *Chem. Soc. Rev.*, 2011, **40**, 3313.
- E. A. Bousseksou, *C. R. Chim.*, 2018, **21**, 1055–1300.
- K. S. Murray, *Spin-Crossover Materials*, John Wiley & Sons, Ltd, 2013, pp. 1–54.
- H. A. Goodwin, in *Spin Crossover in Transition Metal Compounds I*, ed. P. Gütllich and H. A. Goodwin, Springer Berlin Heidelberg, Berlin, Heidelberg, 2004, pp. 59–90.
- Y. Garcia, V. Niel, M. C. Muñoz and J. A. Real, in *Spin Crossover in Transition Metal Compounds I*, ed. P. Gütllich and H. A. Goodwin, Springer Berlin Heidelberg, Berlin, Heidelberg, 2004, pp. 229–257.
- J. R. Thompson, R. J. Archer, C. S. Hawes, A. Ferguson, A. Wattiaux, C. Mathonière, R. Clérac and P. E. Kruger, *Dalton Trans.*, 2012, **41**, 12720.
- A. Lennartson, A. D. Bond, S. Piligkos and C. J. McKenzie, *Angew. Chem., Int. Ed.*, 2012, **51**, 11049–11052.
- R. W. Hogue, S. Singh and S. Brooker, *Chem. Soc. Rev.*, 2018, **47**, 7303–7338.
- J. Olguín and S. Brooker, *Spin-Crossover Materials*, John Wiley & Sons, Ltd, 2013, pp. 77–120.
- M. Ruben, J. Rojo, F. J. Romero-Salguero, L. H. Uppadine and J.-M. Lehn, *Angew. Chem., Int. Ed.*, 2004, **43**, 3644–3662.
- J. G. Hardy, *Chem. Soc. Rev.*, 2013, **42**, 7881.
- A. Stadler, *Eur. J. Inorg. Chem.*, 2009, **2009**, 4751–4770.
- M. Ruben, J.-M. Lehn and P. Müller, *Chem. Soc. Rev.*, 2006, **35**, 1056–1067.
- L. N. Dawe, T. S. M. Abedin and L. K. Thompson, *Dalton Trans.*, 2008, 1661–1675.
- L. N. Dawe, K. V. Shuvaev and L. K. Thompson, *Chem. Soc. Rev.*, 2009, **38**, 2334.
- K. S. Kumar and M. Ruben, *Coord. Chem. Rev.*, 2017, **346**, 176–205.
- B. Schneider, S. Demeshko, S. Dechert and F. Meyer, *Angew. Chem., Int. Ed.*, 2010, **49**, 9274–9277.
- B. Schneider, S. Demeshko, S. Dechert and F. Meyer, *Inorg. Chem.*, 2012, **51**, 4912–4914.
- B. Schneider, S. Demeshko, S. Neudeck, S. Dechert and F. Meyer, *Inorg. Chem.*, 2013, **52**, 13230–13237.
- M. Steinert, B. Schneider, S. Dechert, S. Demeshko and F. Meyer, *Angew. Chem., Int. Ed.*, 2014, **53**, 6135–6139.
- G. J. Halder, C. J. Kepert, B. Moubaraki, K. S. Murray and J. D. Cashion, *Science*, 2002, **298**, 1762–1765.
- W. Vreugdenhil, J. H. V. Diemen, R. A. G. D. Graaff, J. G. Haasnoot, J. Reedijk, A. M. V. D. Kraan, O. Kahn and J. Zarembowitch, *Polyhedron*, 1990, **9**, 2971–2979.
- D. Sertphon, P. Harding, K. Murray, B. Moubaraki, S. Neville, L. Liu, S. Telfer and D. Harding, *Crystals*, 2019, **9**, 116.
- W. Phonsri, P. Harding, L. Liu, S. G. Telfer, K. S. Murray, B. Moubaraki, T. M. Ross, G. N. L. Jameson and D. J. Harding, *Chem. Sci.*, 2017, **8**, 3949–3959.
- I. C. Berdiell, R. Kulmaczewski, N. Shahid, O. Cespedes and M. A. Halcrow, *Chem. Commun.*, 2021, **57**, 6566–6569.
- T. Charytanowicz, J. J. Zakrzewski, K. Dziedzic-Kocurek, S. Chorazy and B. Sieklucka, *J. Appl. Phys.*, 2021, **129**, 143902.
- S. Marcén, L. Lecren, L. Capes, H. A. Goodwin and J.-F. Létard, *Chem. Phys. Lett.*, 2002, **358**, 87–95.
- A. Djemel, O. Stefanczyk, C. Desplanches, K. Kumar, R. Delimi, F. Benaceur, S. Ohkoshi and G. Chastanet, *Inorg. Chem. Front.*, 2021, **8**, 3210–3221.
- M. Hostettler, K. W. Törnroos, D. Chernyshov, B. Vangdal and H.-B. Bürgi, *Angew. Chem., Int. Ed.*, 2004, **43**, 4589–4594.
- E. Collet, N. Moisan, C. Baldé, R. Bertoni, E. Trzop, C. Laulhé, M. Lorenc, M. Servol, H. Cailleau, A. Tissot, M.-L. Boillot, T. Graber, R. Henning, P. Coppens and M. B.-L. Cointe, *Phys. Chem. Chem. Phys.*, 2012, **14**, 6192.
- M. Lorenc, J. Hébert, N. Moisan, E. Trzop, M. Servol, M. Buron-Le Cointe, H. Cailleau, M. L. Boillot, E. Pontecorvo, M. Wulff, S. Koshihara and E. Collet, *Phys. Rev. Lett.*, 2009, **103**, 028301.
- H. Cailleau, M. Lorenc, L. Guérin, M. Servol, E. Collet and M. Buron-Le Cointe, *Acta Crystallogr., Sect. A: Found. Crystallogr.*, 2010, **66**, 189–197.
- M. Lorenc, C. Balde, W. Kaszub, A. Tissot, N. Moisan, M. Servol, M. B.-L. Cointe, H. Cailleau, P. Chasle, P. Czarnecki, M. L. Boillot and E. Collet, *Physical Review B: Solid State*, 2012, **8**.
- R. Bertoni, M. Lorenc, H. Cailleau, A. Tissot, J. Laisney, M.-L. Boillot, L. Stoleriu, A. Stancu, C. Enachescu and E. Collet, *Nat. Mater.*, 2016, **15**, 606–610.
- S. Techert, *J. Appl. Crystallogr.*, 2004, **37**, 445–450.
- J. de J. Velazquez-Garcia, K. Basuroy, D. Storozhuk, J. Wong, S. Demeshko, F. Meyer, R. Henning and S. Techert, *Dalton Trans.*, 2022, **51**, 6036–6045.
- J. de J. Velazquez-Garcia, K. Basuroy, D. Storozhuk, J. Wong, S. Demeshko, F. Meyer, R. Henning and S. Techert, *Dalton Trans.*, 2022, **51**, 17513–17922.
- W. Kabsch, *J. Appl. Crystallogr.*, 1993, **26**, 795–800.
- G. M. Sheldrick, *Acta Crystallogr., Sect. A: Found. Adv.*, 2015, **71**, 3–8.
- O. V. Dolomanov, L. J. Bourhis, R. J. Gildea, J. A. K. Howard and H. Puschmann, *J. Appl. Crystallogr.*, 2009, **42**, 339–341.
- G. M. Sheldrick, *Acta Crystallogr., Sect. C: Struct. Chem.*, 2015, **71**, 3–8.



- 44 K. Basuroy, J. de J. Velazquez-Garcia, D. Storozhuk, R. Henning, D. J. Gosztola, S. Thekku Veedu and S. Techert, *J. Chem. Phys.*, 2023, **158**, 054304.
- 45 J. Z. Ren, *Precognition User Guide Ref. Tutor*, Renz Res. Inc., Westmont IL USA, 2006.
- 46 R. Ketkaew, Y. Tantirungrotechai, P. Harding, G. Chastanet, P. Guionneau, M. Marchivie and D. J. Harding, *Dalton Trans.*, 2021, **50**, 1086–1096.
- 47 H. Zabrodsky, S. Peleg and D. Avnir, *J. Am. Chem. Soc.*, 1992, **114**, 7843–7851.
- 48 M. Llunell, D. Casanova, J. Cirera, P. Alemany and S. Alvarez, *SHAPE Program Version 21 Barc*, 2003.
- 49 M. S. Schmøkel, R. Kamiński, J. B. Benedict and P. Coppens, *Acta Crystallogr., Sect. A: Found. Crystallogr.*, 2010, **66**, 632–636.
- 50 I. I. Vorontsov and P. Coppens, *J. Synchrotron Radiat.*, 2005, **12**, 488–493.
- 51 J. J. McKinnon, A. S. Mitchell and M. A. Spackman, *Chem. – Eur. J.*, 1998, **4**, 2136–2141.
- 52 M. A. Spackman and D. Jayatilaka, *CrystEngComm*, 2009, **11**, 19–32.
- 53 M. J. Turner, J. J. McKinnon, S. K. Wolff, D. J. Grimwood, P. R. Spackman, D. Jayatilaka and M. A. Spackman, *CrystalExplorer17*, University of Western Australia, 2017.
- 54 P. Guionneau, *Dalton Trans.*, 2014, **43**, 382–393.
- 55 W. Guo, N. Daro, S. Pillet, M. Marchivie, E. Bendeif, E. Tailleux, K. Chainok, D. Denux, G. Chastanet and P. Guionneau, *Chem. – Eur. J.*, 2020, **26**, 12927–12930.
- 56 E. M. Zueva, E. R. Ryabikh and S. A. Borshch, *Inorg. Chem.*, 2011, **50**, 11143–11151.
- 57 M. A. Naumova, A. Kalinko, J. W. L. Wong, S. Alvarez Gutierrez, J. Meng, M. Liang, M. Abdellah, H. Geng, W. Lin, K. Kubicek, M. Biednov, F. Lima, A. Galler, P. Zalden, S. Checchia, P.-A. Mante, J. Zimara, D. Schwarzer, S. Demeshko, V. Murzin, D. Gosztola, M. Jarenmark, J. Zhang, M. Bauer, M. L. Lawson Daku, D. Khakhulin, W. Gawelda, C. Bressler, F. Meyer, K. Zheng and S. E. Canton, *J. Chem. Phys.*, 2020, **152**, 214301.
- 58 M. A. Naumova, A. Kalinko, J. W. L. Wong, M. Abdellah, H. Geng, E. Domenichini, J. Meng, S. A. Gutierrez, P.-A. Mante, W. Lin, P. Zalden, A. Galler, F. Lima, K. Kubicek, M. Biednov, A. Britz, S. Checchia, V. Kabanova, M. Wulff, J. Zimara, D. Schwarzer, S. Demeshko, V. Murzin, D. Gosztola, M. Jarenmark, J. Zhang, M. Bauer, M. L. Lawson Daku, W. Gawelda, D. Khakhulin, C. Bressler, F. Meyer, K. Zheng and S. E. Canton, *J. Phys. Chem. Lett.*, 2020, **11**, 2133–2141.

