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The Role of Symmetry Breaking in the Structural Trapping of Light-Induced Excited Spin States

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Light-Induced Excited Spin State Trapping (LIESST) data are reported for seven isostructural solvate salts from the iron(II)/2,6-di(pyrazol-1-yl)pyridine family. A complicated relationship between their spin-crossover $T_{1/2}$ and $T(\text{LIESST})$ values may reflect low-temperature thermal and light-induced symmetry breaking, which is shown by one of the compounds but not by two others.

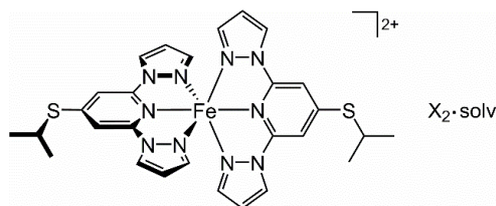
Spin-crossover (SCO) compounds^{1,2} are versatile molecular switches for use in multifunctional materials, macroscopic devices and nanoscience.^{2,3} SCO transitions can be induced by a range of stimuli including temperature, pressure and visible irradiation.^{1,4} Light induced SCO is most often measured as a photo-conversion of a low-spin compound to a metastable high-spin state at low temperatures. This is the Light-Induced Excited Spin State Trapping (LIESST) effect.⁵ The sample can only reconvert to its low-spin ground state upon heating above the activation barrier to its relaxation process, which typically lies below 150 K.

Mean-field theory of the propagation of SCO transitions through solid lattices predicts an inverse relationship between the thermodynamic SCO temperature $T_{1/2}$ and the lifetime of the photoinduced metastable state,⁶ Some years ago an empirical relationship of this type was indeed proposed in different families of compounds (eq 1):

$$T(\text{LIESST}) = T_0 - 0.3T_{1/2} \quad (1)$$

where $T(\text{LIESST})$ is the relaxation temperature of the kinetically trapped spin state^{7,8} and T_0 reflects the rigidity of the metal ion coordination sphere.^{8,9} Although there is often a degree of scatter in $T_{1/2}$ vs $T(\text{LIESST})$ plots, eq 1 is a useful predictor of $T(\text{LIESST})$ for many types of SCO materials. This includes complex salts derived from $[\text{Fe}(\text{bpp})_2]^{2+}$ ($\text{bpp} = 2,6\text{-di}\{\text{pyrazol-1-yl}\}\text{pyridine}$), which often show good agreement to eq 1 with $T_0 \approx 150$ K.^{10,11}

We recently reported six compounds of general formula $[\text{Fe}L_2][\text{BF}_4]_2 \cdot \text{solv}$ ($\mathbf{1}[\text{BF}_4]_2 \cdot \text{solv}$, Scheme 1).¹² This is a rare family of isostructural SCO materials,¹³ which facilitates studies of structure: function relationships underlying their SCO behaviour. We have expanded the series with the perchlorate salts $\mathbf{1}[\text{ClO}_4]_2 \cdot \text{solv}$, and report a photomagnetic and photocrystallographic study on these isostructural compounds that reveals the relationship between structure and $T(\text{LIESST})$ in unprecedented detail.



Scheme 1. Compound $\mathbf{1}X_2 \cdot \text{solv}$ ($X^- = \text{BF}_4^-$ or ClO_4^- ; $\text{solv} = \text{MeNO}_2, \text{MeCN}, \text{Me}_2\text{CO}, \text{H}_2\text{O}$ or sf [solvent-free]).

Complexation of $\text{Fe}[\text{ClO}_4]_2 \cdot 6\text{H}_2\text{O}$ by 2 equiv L in the appropriate solvent affords $\mathbf{1}[\text{ClO}_4]_2 \cdot \text{MeCN}$, $\mathbf{1}[\text{ClO}_4]_2 \cdot \text{MeNO}_2$ and $\mathbf{1}[\text{ClO}_4]_2 \cdot \gamma\text{Me}_2\text{CO}$ ($\gamma \approx 0.7$) after the usual work-up. $\mathbf{1}[\text{ClO}_4]_2 \cdot \gamma\text{Me}_2\text{CO}$ is converted to $\mathbf{1}[\text{ClO}_4]_2 \cdot \text{H}_2\text{O}$ in single-crystal-to-single-crystal fashion, when stored *in vacuo* at 290 K for 24 hrs and then exposed to air. Solvent-free $\mathbf{1}[\text{ClO}_4]_2 \cdot \text{sf}$ was also prepared *in situ*, by annealing crystals of $\mathbf{1}[\text{ClO}_4]_2 \cdot \gamma\text{Me}_2\text{CO}$ on the diffractometer (ESI †). The $\mathbf{1}[\text{ClO}_4]_2 \cdot \text{solv}$ and $\mathbf{1}[\text{BF}_4]_2 \cdot \text{solv}$ ¹² compounds are high-spin, isostructural and phase-pure at room temperature ($P2_1/c$, $Z = 4$), while all except $\mathbf{1}[\text{ClO}_4]_2 \cdot \text{MeCN}$ exhibit SCO upon cooling without a crystallographic phase change (Fig. 1 and ESI †). The SCO temperature and cooperativity vary between the compounds, but the high-spin state is consistently stabilised when $X = \text{ClO}_4^-$ compared to $X = \text{BF}_4^-$ for each solvent. That might reflect expansion of the crystal lattice by the larger ClO_4^- anion, which would favour the larger high-spin cations.¹⁴ Any fraction of the samples that is high-spin near 100 K remains frozen in below that temperature.^{15,16} Poising $\mathbf{1}[\text{ClO}_4]_2 \cdot \text{MeNO}_2$ at 102 K for 80 mins leads to a slow reduction in $\chi_M T$, until the sample is fully low-spin and the warming branch of the transition proceeds monotonically (ESI †). That confirms the kinetic origin of the low-temperature spin-state trapping, and the apparent SCO hysteresis, in that material. Such kinetic effects arise when thermal trapping of the high-spin state occurs at a similar temperature to $T_{1/2}$ in an SCO material.¹⁵

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† Electronic Supplementary Information (ESI) available: experimental procedures and characterisation data; crystallographic experimental data, Figures and Tables; magnetic susceptibility data for all the compounds, measured under the same conditions as in ref. 12; kinetic studies of thermal SCO in two $\mathbf{1}[\text{ClO}_4]_2 \cdot \text{solv}$ samples; and a Table of the $T(\text{LIESST})$ data plotted in Fig. 2. See DOI: 10.1039/x0xx00000x

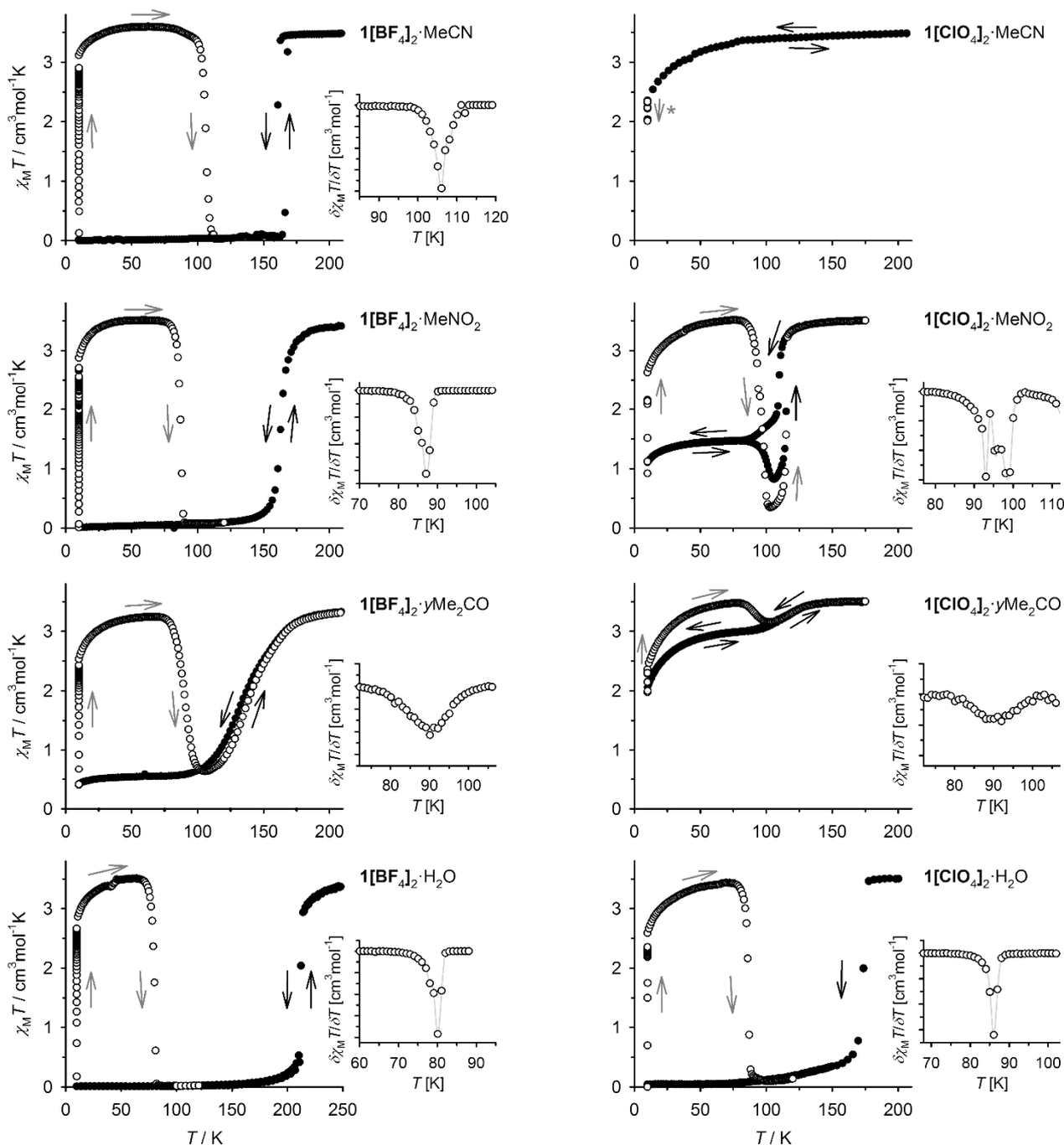


Figure 1. Temperature dependence of magnetic susceptibility data for eight $1\mathbf{X}_2$ -solv compounds before irradiation (●, black arrows), and during the $T(\text{LIESST})$ measurement (○, grey arrows). The samples were cooled to 10 K, irradiated at that temperature ($\lambda = 510$ nm), then rewarmed in the dark. Scan rate 0.4 Kmin^{-1} . The insets show the first derivatives of the relaxation curves, with data points linked by spline curves for clarity. The starred compound is high-spin at 10 K, and was irradiated at $\lambda = 980$ nm in a reverse-LIESST experiment.¹⁷

Seven freshly prepared $1\mathbf{X}_2$ -solv samples showed essentially quantitative low \rightarrow high-spin photoconversion upon irradiation at 510 nm at 10 K (Fig. 1). After equilibration, warming the samples in the dark showed the high-spin forms are long-lived until ca 80 K, where thermal relaxation to their thermodynamic low-spin states took place.¹⁷ The $T(\text{LIESST})$ curves are mostly monotonic but of differing abruptness, with samples exhibiting the least cooperative thermal SCO ($1[\text{BF}_4]_2 \cdot \gamma\text{Me}_2\text{CO}$ and $1[\text{ClO}_4]_2 \cdot \gamma\text{Me}_2\text{CO}$) showing the most gradual LIESST relaxation. An exception is $1[\text{ClO}_4]_2 \cdot \text{MeNO}_2$, whose relaxation is split into three closely spaced components.

Although other explanations are possible,¹⁸ this stepped relaxation may reflect crystallographic phase changes occurring during the LIESST relaxation process (see below).¹⁹

The $T(\text{LIESST})$ values, from the minima of the $\delta\chi_M T / \delta T$ curves, are typical for complexes of this type (Table 1).¹¹ However, a plot of $T_{1/2}$ vs $T(\text{LIESST})$ for these data can be interpreted in two ways (Fig. 2). At first glance, all the compounds lie on the same $T(\text{LIESST})/T_{1/2}$ line except $1[\text{BF}_4]_2 \cdot \text{MeCN}$, whose $T(\text{LIESST})$ is ca 20 K higher than for $1[\text{BF}_4]_2 \cdot \text{MeNO}_2$ despite their similar $T_{1/2}$ values (Table 1). However, comparison of these data with the literature shows an

Table 1. Thermal SCO and LIESST properties of $1[\text{BF}_4]_2\text{-solv}$ and $1[\text{ClO}_4]_2\text{-solv}$, with a temperature ramp of 0.4 Kmin^{-1} . (Figs. 1 and 2). Data for the BF_4^- salts differ slightly from those in ref. 13, which were measured at a faster scan rate of 5 Kmin^{-1} (ESI †).

	$T_{1/2}\downarrow / \text{K}$	$T_{1/2}\uparrow / \text{K}$	cooperativity	$T(\text{LIESST}) / \text{K}$
$1[\text{BF}_4]_2\text{-MeCN}$	160	167	abrupt	106
$1[\text{ClO}_4]_2\text{-MeCN}$	HS ^a	–	–	–
$1[\text{BF}_4]_2\text{-MeNO}_2$	171	–	gradual	87
$1[\text{ClO}_4]_2\text{-MeNO}_2$	102	115	gradual, incomplete	93,96,99
$1[\text{BF}_4]_2\cdot\gamma\text{Me}_2\text{CO}$	131	–	gradual, incomplete	91
$1[\text{ClO}_4]_2\cdot\gamma\text{Me}_2\text{CO}$	<100 ^b	–	gradual, incomplete	ca 90 ^c
$1[\text{BF}_4]_2\cdot\text{H}_2\text{O}$	212	214	abrupt	81
$1[\text{ClO}_4]_2\cdot\text{H}_2\text{O}$	173	–	abrupt	86

^aHS = high-spin between 3-300 K. ^bOnly 20 % of the SCO transition occurs before the remaining high-spin fraction is frozen in below 100 K. ^cNot included in Fig. 2.

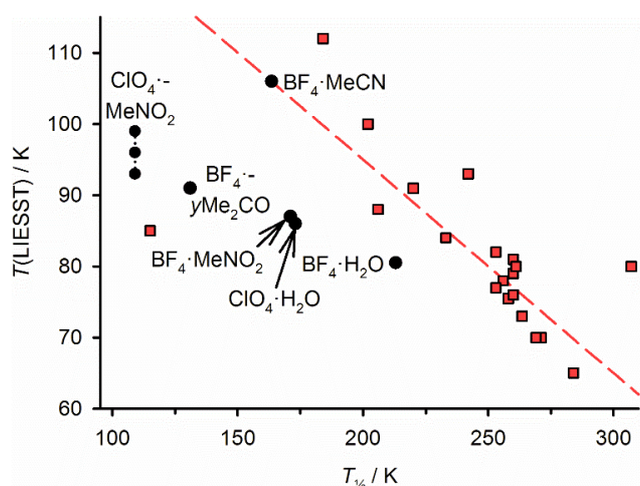


Figure 2. Plot of $T_{1/2}$ vs $T(\text{LIESST})$ for the compounds in this work (black circles; Table 1), and from previously published compounds from our laboratory (red squares; ESI†). The dashed line shows eq 1 with $T_0 = 155\text{ K}$, close to the $T_0 = 150\text{ K}$ correlation that was originally proposed for this family of compounds.^{10,11}

alternative picture. Only $1[\text{BF}_4]_2\text{-MeCN}$ and $1[\text{BF}_4]_2\text{-H}_2\text{O}$ lie within experimental error of the trend expected from our previous measurements. The other compounds show reduced $T(\text{LIESST})$ values, deviating increasingly from eq 1 as $T_{1/2}$ is lowered (Fig. 2). Crystallographic studies on three of the materials shed light on these differences. Unexpectedly, $1[\text{BF}_4]_2\text{-MeNO}_2$ undergoes a symmetry-breaking phase transition upon cooling from 100 K (phase 1; $P2_1/c$, $Z = 4$) to 15 K (phase 2; $P2_1/c$, $Z = 12$), involving a tripling of the unit cell b dimension. Three unique low-spin cations in the asymmetric unit, labelled ‘A’, ‘B’ and ‘C’, are grouped into layers parallel to (010). Individual layers contain either A-type molecules or alternating B and C types, with the layers arranged as A–(B/C)–(B/C)–A–(B/C)–(B/C) down the b axis (Fig. 3). Irradiation at 660 nm at 15 K transforms the crystal to a new high-spin phase (phase 3; $P2_1$, $Z = 4$), whose unit cell dimensions resemble phase 1 but which lacks the c glide plane. The ‘A’ and ‘B’ cation sites in this phase are grouped into corrugated layers along (001) (Fig. 4). The cations in phases 2 and 3 have similar metric parameters to the corresponding spin states of phase 1. The symmetry breaking is reflected in changes to the orientations of the *isopropyl* groups, anions and solvent molecules (ESI †).

In contrast, $1[\text{BF}_4]_2\text{-H}_2\text{O}$ (at 20 K) and $1[\text{BF}_4]_2\text{-MeCN}$ (at 85 K) both retain phase 1 before and after irradiation; the high-spin form

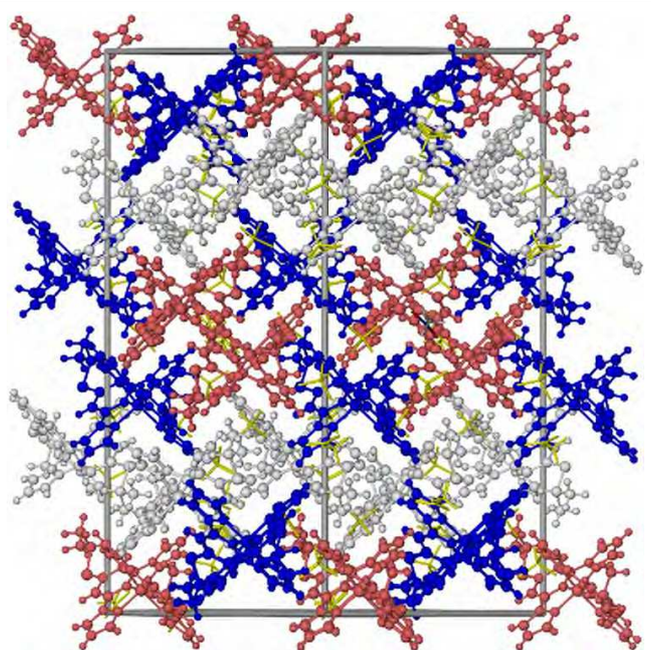


Figure 3. Packing diagram of low-spin phase 2 of $1[\text{BF}_4]_2\text{-MeNO}_2$ along the (101) crystal vector, with the b axis vertical. Cations A, B and C are coloured white, blue and red, respectively, while the anions and solvent (yellow) are de-emphasised for clarity.

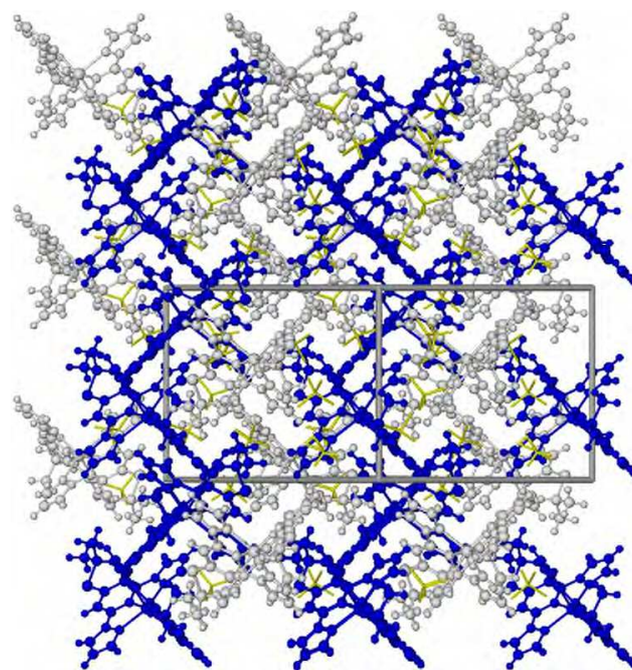


Figure 4. Packing diagram of high-spin phase 3 of $1[\text{BF}_4]_2\text{-MeNO}_2$, in the same view as Fig. 3. Cations A and B are coloured white and blue, respectively. Other details as for Fig. 3.

of $1[\text{BF}_4]_2\text{-MeCN}$ was also characterised at 15 K, again adopting phase 1. Hence $1[\text{BF}_4]_2\text{-H}_2\text{O}$ and $1[\text{BF}_4]_2\text{-MeCN}$, which align more closely with eq 1 ($T_0 = 150\text{ K}$, Fig. 2), show no evidence for symmetry breaking under these conditions. The isothermal low-spin→high-spin photoconversion of both phase 1 compounds results in an expansion of the unit cell a axis and a contraction of b and β . The reduction in β is much larger for $1[\text{BF}_4]_2\text{-MeCN}$ at 85 K, causing an unusual 0.2 % contraction of the unit cell volume in its

high-spin state. In contrast, $1[\text{BF}_4]_2 \cdot \text{H}_2\text{O}$ and $1[\text{BF}_4]_2 \cdot \text{MeNO}_2$ both undergo a more typical expansion of their normalised unit cell volume during photoexcitation experiments (ESI †).

The light-induced high-spin state of $1[\text{BF}_4]_2 \cdot \text{MeNO}_2$ (phase 3) has reduced crystallographic symmetry, and thus a lower entropy, than its thermodynamic high-spin state (phase 1). That should shift the (theoretical) $T_{1/2}$ of phase 3 to a higher temperature than phase 1,²⁰ leading to a lower $T(\text{LIESST})$ for phase 3 as observed (eq 1).²¹ This symmetry-breaking entropy change is unlikely to be electronic in origin, since the coordination geometries of the C_1 -symmetric iron centers are similar in each phase. Rather, it predominantly reflects a reduction in vibrational entropy through a lifting of lattice phonon degeneracy, associated with the loss of the crystallographic glide plane and inversion center in phase 3. Attempts to access phase 1 of $1[\text{BF}_4]_2 \cdot \text{MeNO}_2$ by photoirradiation, for comparison with phase 3, have thus far been unsuccessful.

In conclusion, isostructural $1[\text{BF}_4]_2 \cdot \text{solv}$ and $1[\text{ClO}_4]_2 \cdot \text{solv}$ exhibit a complex relationship between $T_{1/2}$ and $T(\text{LIESST})$. Most of the compounds exhibit a linear $T_{1/2}$ vs $T(\text{LIESST})$ dependence, with a reduced slope compared to eq 1 (Fig. 2). Hence, a generalisation of eq 1 can be applied to this subset of compounds (eq 2).

$$T(\text{LIESST}) = T_0 - aT_{1/2} \quad (2)$$

The data in Table 1 (omitting $1[\text{BF}_4]_2 \cdot \text{MeCN}$) are best fit by $T_0 = 108 \text{ K}$ and $a = 0.13$ (ESI †). Moreover, $T(\text{LIESST})$ for $1[\text{BF}_4]_2 \cdot \text{MeCN}$ and $1[\text{BF}_4]_2 \cdot \text{MeNO}_2$ differ by 20 K, despite their almost identical $T_{1/2}$ values (Table 1). That can be explained by the thermodynamic consequences of a series of thermal and light-induced symmetry-breaking transitions, which are undergone by $1[\text{BF}_4]_2 \cdot \text{MeNO}_2$ but not $1[\text{BF}_4]_2 \cdot \text{MeCN}$ or $1[\text{BF}_4]_2 \cdot \text{H}_2\text{O}$. This clearly demonstrates the impact of crystallographic phase changes on $T(\text{LIESST})$,²¹ at least in these two compounds. Other compounds that deviate unexpectedly from eqs 1 or 2 may also exhibit unresolved structural chemistry in the LIESST experiment.

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Conflicts of interest

There are no conflicts to declare.

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