Magnetic anisotropy in trigonal planar Fe(II) bis(trimethylsilyl)amido complexes of the type [Fe{N(SiMe$_3$)$_2$}$_2$L]—experiment and theory†

Tilmann Bodenstein*† and Andreas Eichhöfer Ø *

Systematic ac (alternating current) magnetic investigations on four new trigonal planar high-spin Fe$^{2+}$ complexes [Fe{N(SiMe$_3$)$_2$}$_2$L] reveal that complexes which comprise a phosphine or arsine type ligand (L = PPh$_3$, PMe$_3$ and AsPh$_3$) display slow magnetic relaxation at temperatures below 8 K under applied dc (direct current) fields, whereas a complex with a phosphine oxide ligand (L = OPPh$_3$) does not. Accordingly, the parameters characteristic for magnetic anisotropy, derived both from dc magnetic measurements and quantum chemical calculations, reveal distinct differences for these two types of complexes. Extensive ab initio calculations of multi-reference wave function type were performed on the four new complexes listed above and the related reported ones with L = py, thf and PCy$_3$ in order to get a reasonable description of the local electronic states involved in the magnetic relaxation. These calculations confirm that strong spin–orbit effects generate the magnetic anisotropy of complexes with L = PPh$_3$, PMe$_3$, AsPh$_3$ and PCy$_3$. On the other hand, the complexes with L = OPPh$_3$, py and THF exhibit only small spin–orbit splittings, consistent with the fast relaxation found experimentally.

Introduction

Single-ion molecule magnet behavior, which was first observed in lanthanide-containing complexes,$^{1,2}$ and meanwhile also in several transition metal complexes,$^{3–6}$ may once enable the utilization of such molecules as the smallest possible units in spin-based computational or data storage devices.$^7$ However, the task to control slow relaxation of the magnetization as a desired property in these molecules is still a challenge.

Maximization of the anisotropy of the magnetic moment for a respective molecule will in principle increase its energy barrier which determines the Orbach relaxation process.$^8$ Successful strategies based upon magneto-structural correlations have been developed to achieve these goals.$^{3–6}$ In this respect, the most promising properties of 3d transition metal complexes have been found recently for coordinatively unsaturated compounds. For example, the linearly coordinated cobalt complex [Co(C(SiMe$_3$)$_3$Naph)$_2$] displays a record spin reversal barrier of 450 cm$^{-1}$ in a zero applied direct current field due to the realization of an unusual maximum of the orbital momentum of $L = 3$ arising from a non-Aufbau ground state.$^9$ A similarly high barrier has also been described for a NHC complex of cobalt(u), [[sIPr]Co(u)NDmp] [sIPr = 1,3-bis(2',6'-diisopropylphenyl)-4,5-dihydro-imidazol-2-ylidene; Dmp = 2,6-dimesitylphenyl] with a value of 413 cm$^{-1}$, although the orbital momentum is thought to amount to a value of $L = 2$ only.$^{10}$ In comparison, the two-coordinate complexes of iron(i) and iron(u), [Fe(C(SiMe$_3$)$_3$)$_3$]$^-$ and [Fe(C(SiMe$_3$)$_3$)$_3$]$^{2-}$, display slow magnetic relaxation without application of an external dc field with smaller effective spin reversal barriers of $U_{\text{eff}} = 226$ cm$^{-1}$ and 146 cm$^{-1}$, respectively. In view of the same $L = 2$ ground state like the aforementioned cobalt NHC complex, this difference most probably originates from smaller spin–orbit coupling constants in the iron compounds.

Related investigations have been performed on trigonal planar iron(u) complexes [Fe{N(SiMe$_3$)$_2$}$_2$(L)] (L = PCy$_3$, $^{13}$ thf, N(SiMe$_3$)$_2$, $^{14}$ PCyp$_3$). For this type of complexes one usually expects spatially non-degenerate ground states with $L = 0$. However, small ligand field splittings could in principle allow for a coupling to low lying excited states, and the recent findings suggest an influence of the type of ligand L on the dc and
ac magnetic properties of such complexes. Concerning the understanding of the magnetic properties of such compounds, quantum chemical studies based on wave function based methods have been shown to be able to predict and describe electronic properties of open-shell ion complexes quite accurately (for reviews, see for example ref. 16 and 17) and may be used as a guideline for experimentalists for designing novel complexes with desired magnetic anisotropy.

Herein, we report on the experimental and theoretical characterization of seven trigonal planar \( \text{Fe}^{2+} \) bis(trimethylsilyl)amido complexes of the type \([\text{Fe}\{\text{N(SiMe}_3\}_2\}_2(L)]\) with a focus on the influence of the ligand \( L \) on their electronic and magnetic properties.

**Results and discussion**

Table 1 introduces the molecular formulas and the numbering scheme of the complexes 1–7 which are considered in this work. The synthesis and structural as well as magnetic data have already been published for 6 \(^{14,18,19}\) and 7 \(^{13,14}\) in preceding papers whereas for 5 only an experimental procedure has been published so far.\(^{20}\)

**Synthesis and structure**

The new complexes 1–4 were synthesized in good yields by the reaction of \([\text{Fe}\{\text{N(SiMe}_3\}_2\}_2\]) with the respective ligand either in heptane (1, 3 and 4) or in a solvent free reaction (2) upon heating in accordance to earlier papers on related complexes.\(^{18,19}\) 5 was synthesized prior by sublimation of \([\text{Fe}\{\text{N(SiMe}_3\}_2\}_2\] (\(\text{py}\)) but no single crystal XRD has been performed.\(^{20}\) We obtained single crystals of 5 from reactions of \([\text{Fe}\{\text{N(SiMe}_3\}_2\}_2\] and 1 equiv. of pyridine in pentane according to Scheme 1. However, powder XRD and elemental analysis indicate that the material isolated in this way is not pure. Apart from that, geometrical parameters of 1–5 are similar to the related and known complexes 6,\(^{18,19}\) 7,\(^{13}\) and \([\text{Fe}\{\text{N(SiMe}_3\}_2\}_2(\text{PCyp}_3)]\).\(^{15}\) Shortest interatomic distances between the metal atoms in the crystal lattices are in the range between 891 and 1018 pm (1: 978, 2: 891, 3: 961, 4: 951, 5: 1018 pm).

The measured powder patterns of 1–4 show a good agreement with the calculated ones based on the single crystal data (Fig. S6–S8†) which proves the crystalline purity of the compounds.

![Scheme 1](image)

**Table 1**  List of complexes \([\text{Fe}\{\text{N(SiMe}_3\}_2\}_2L\] (1–7)

| No. | Ligand \( L \) |
|-----|---------------|
| 1   | \( \text{PPh}_3 \) |
| 2   | \( \text{PMe}_3 \) |
| 3   | \( \text{AsPh}_3 \) |
| 4   | \( \text{OPPh}_3 \) |
| 5   | \( \text{py} \) |
| 6   | \( \text{thf} \) |
| 7   | \( \text{PCy}_3 \) |

![Fig. 1](image)
The values of \( \chi T \) for 1–3 display distinct downturns below 100 K (Fig. 2 and 3). This deviation from the ideal Curie behavior is in the absence of close Fe⋯Fe contacts attributable to magnetic anisotropy which is indicative of a significant zero-field splitting and g-tensor anisotropy resulting from the pseudo-trigonal crystal field. Magnetic anisotropy for these three complexes is also indicated by the room temperature values of \( \chi T \) (1: 4.14, 2: 4.27, 3: 4.35 cm\(^3\) mol\(^{-1}\) K) which are much larger than the theoretical spin-only value of 3 cm\(^3\) mol\(^{-1}\) K for one Fe\(^{2+}\) ion (high-spin, \( S = 2 \)). Comparably large values for \( \chi T \) at room temperature have been reported for 7 (\( \chi T = 4.10, 5.12 \) cm\(^3\) mol\(^{-1}\) K). \(^{13,14} \)

In contrast, 4 does not display such an early deviation from the ideal Curie behavior and a distinctly lower value for \( \chi T \) at room temperature (3.63 cm\(^3\) mol\(^{-1}\) K) comparable to those of 5 (3.51 cm\(^3\) mol\(^{-1}\) K)\(^{20} \) and 6 (3.70 cm\(^3\) mol\(^{-1}\) K). \(^{14} \) Also, the value of \( \chi T \) at 2 K for 4 (1.55 cm\(^3\) mol\(^{-1}\) K) differs distinctly from those observed for 1–3 (3.30, 3.44, 3.71 cm\(^3\) mol\(^{-1}\) K). In agreement, the curves of the field dependence of the magnetization (M) of 1–3 on the one, and 4 on the other hand are also different (Fig. S11 and S12†), suggesting either different ground state properties for these complexes or distinct different magnetic anisotropies or both.

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**Table 2 Structural parameters (atomic distances [pm], bond angles [°], shortest intermolecular Fe⋯Fe distances [pm] and deviation of the iron atom from the trigonal plane [pm]) in 1–7**

|            | Fe–N   | Fe–P/As/O/N | N–Fe–N | N–Fe–P/As/O/N | Shortest Fe⋯Fe | \( \Delta \) plane |
|------------|--------|------------|--------|---------------|----------------|------------------|
| 1          | 192.6, 192.5(2) | 253.1(1)   | 135.7(8) | 109.08(6), 115.15(6) | 978             | 2.9(1)          |
| 2          | 192.3, 194.6(2) | 247.5(1)   | 139.03(8) | 109.11(8), 111.85(8) | 891             | 2.9(1)          |
| 3          | 191.7, 192.0(2) | 263.0(1)   | 141.0(1)  | 106.2(1), 111.7(1)   | 961             | 3.5(1)          |
| 4          | 192.9, 193.3(2) | 197.6(2)   | 136.48(9) | 110.12(9), 113.38(9) | 951             | 1.2(1)          |
| 5a         | 192.4, 193.1(4) | 210.1(4)   | 141.7(2)  | 109.14(11), 109.14(11)| 1018             | 3.1(1)          |
| 6b         | 191.8, 191.8(4) | 211.5(4)   | 144.14(16)| 108.75(16), 107.03(15)| 0               |                |
| 7          | 192.7, 192.9(2) | 261.9(12)  | 128.11(2) | 115.72(2), 116.15(2) | 924             | 1.8(1)          |

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"This work. †From ref. 19. ‡From ref. 13."
In order to probe this magnetic anisotropy further, we modeled the \( \mu_{\text{eff}} \) vs. \( T \) and \( M \) vs. \( H \) curves at different temperatures simultaneously using least-squares fits using the program PHI.\(^{21} \) The model includes both axial (\( D \)) and rhombic (\( E \)) ZFS terms as well as Zeeman interactions with an anisotropic treatment of \( g \) (eqn (S1))\(^{19} \) with \( g_x = g_y = g_z \). The best sets of parameters are listed in Table 3 and shown as solid green lines in Fig. 2, 3 and S11, S12.\(^{\dagger} \) Complexes 1–3 display anisotropic \( g \) parameters and large negative \( D \) values, whereas fits of the anisotropic \( g \)-factor of 4 resulted in less anisotropic \( g \)-values as well as a less negative \( D \) (\( D = -15.4 \) \( \text{cm}^{-1} \)). The lower increase of the magnetization curves \( (M_S \text{ vs. } H) \) in 4 (most visible at low \( T \) and \( H < 2 \) \( T \), Fig. S12\(^{\dagger} \)) is reflected in the large \( E/D \) ratio obtained in the fit. This lifts the ground state degeneracy present in 1–3, 7 as confirmed by the \( ab \) \( initio \) calculations (see Theory section). We note that the data of 4 can in principle also be fitted by a set of parameters comprising a positive value of \( D = +12.7 \) \( \text{cm}^{-1} \) \( (g_x = g_y = 2.17, g_z = 2.10, \ E = \pm 4.35, \ \text{TIP} = 0.34 \times 10^{-2} \text{cm}^{-2} \text{mol}^{-1} \)) but with a worse goodness-of-fit factor \( R = 0.214 \), compared to \( R = 0.006 \) for the first option. Related negative values of \( D \) have been observed in [(IPr)Fe(N(SiMe\(_3\))\(_2\)] (IPr = 1,3-bis(diisopropylphenyl)imidazol-2-ylidene) \( (D = -18.2 \) \( \text{cm}^{-1} \)) and [[IMes]Fe(N(SiMe\(_3\))\(_2\)] \( (D = -23.3 \) \( \text{cm}^{-1} \)).\(^{22} \) 6 \( (D = -3.0 \text{ cm}^{-1}); \) 7 \( (D = -33 \text{ cm}^{-1}); \) and [FeN(SiMe\(_3\))\(_2\)(PCyp\(_3\))] \( (D = -38 \text{ cm}^{-1}); \)\(^{15} \) A positive \( D \) value has been recently reported for the trigonal planar iron(II) complex [FeN(SiMe\(_3\))\(_2\)] \( (D = +9.9 \text{ cm}^{-1}); \)\(^{14} \)

In order to probe the dynamic magnetic behavior further, ac measurements were performed in the 1.8–10 \( K \) range using a 3.0 Oe ac field, oscillating at frequencies between 1 and 1500 Hz (Fig. 4–6 and S13–S22\(^{\dagger} \)). In the absence of an external dc field, the out-of-phase components of the ac susceptibility (\( \chi'' \)) of 1–4 have much lower intensity than the in-phase component (\( \chi' \)) and display no maximum, indicating that spin lattice relaxation is faster than the timescale of the experiment. With the application of a static dc field, the intensity of \( \chi'' \) is significantly enhanced for 1–3, but not for 4. This effect can in 1–3 be ascribed to a lifting of degeneracy of the electronic states by the dc field which impedes relaxation processes \( \text{via} \) Quantum Tunneling (QTM).\(^{23,24} \) In contrast, the absence of slow relaxation of magnetization for complex 4 is in line with a smaller magnitude of \( D \) and less axial anisotropy expressed by its \( g \)-factors (Table 3) being not sufficient to establish a significant intrinsic barrier for reversal of the magnetization. This is similar to the behavior of 6.\(^{14} \) In addition, the fitted value of the rhombic ZFS parameter \( E = \pm 5.0 \) is quite high. Quantum tunneling due to transversal fields expressed by \( E \) is among others thought to be one of the reasons for lowering the theoretical spin reversal barrier especially in non-Kramers ions through mixing of the \( M_S \pm 1 \) states.\(^{25,26} \)

Both, the temperature and field dependence of the relaxation time provide an experimental probe for the processes responsible for spin reversal in a magnetic system. The magnetic relaxation times for 1–3 where therefore assessed by measuring the frequency dependencies at different fields (Fig. S13–S18\(^{\dagger} \)) and temperatures (Fig. 4–6 and S19\(^{\dagger} \)), and were subsequently extracted (from these measurements) by fits to eqn (S2) and (S3).\(^{\dagger} \)

For 1–3, the field dependence of the inverse relaxation time at 2 K displays a curvature feature with a minimum around 1500 Oe and a negative slope for smaller and a positive for

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**Table 3** Results of the fittings of the dc magnetic data (simultaneous treatment of \( J \) \( T \) vs. \( T \) and \( M \) vs. \( H \) plots at different temperatures, Fig. 2, 3 and S11, S12\(^{\dagger} \)) of 1–4, 6 and 7 by the PHI program (eqn (S1))\(^{21,21} \)

| \( g_x \) | \( g_y \) | \( g_z \) | \( D \) \( \text{[cm}^{-1}] \) | \( E \) \( \text{[cm}^{-1}] \) | TIP \( \text{[cm}^{-2} \text{mol}^{-1}] \) | \( R \) \( \times 10^{-3} \) |
|---|---|---|---|---|---|---|
| 1 | 2.08 | 2.50 | -39.6 | \( \pm 0.22 \) | 0.96 \( \times 10^{-3} \) | 18.9 |
| 2 | 2.12 | 2.71 | -37.9 | \( \pm 4.22 \) | 0.175 \( \times 10^{-3} \) | 2.7 |
| 3 | 2.08 | 2.65 | -42.3 | \( \pm 0.52 \) | 0.47 \( \times 10^{-3} \) | 9.6 |
| 4 | 1.98 | 2.25 | -15.4 | \( \pm 5.0 \) | 1.06 \( \times 10^{-3} \) | 0.61 |
| 6\(^{\dagger} \) | 2.07 | 2.28 | -20 | \( \pm 4.0 \) | 0.7 \( \times 10^{-3} \) | 0.25 |
| 7\(^{\dagger} \) | 2.14 | 2.61 | -33 | \( \pm 3.4 \) | 3.15 |
larger fields (Fig. S14, S16, S18†). Such a behaviour was for example also observed in the low field region (<3000 Oe) of dilute samples of Cu²⁺ in (NH₄)₂Zn(SO₄)₂·6H₂O27 and in recent investigations on linear coordinated molecular Fe²⁺ complexes,11 and is commonly attributed to quantum tunneling effects. Approximate fits to these data of 1–3 can be obtained using eqn (S4),† where the first term represents the direct process, while the second corresponds to the zero-field tunneling process, accordingly to a procedure recently used for linear Fe(II) complexes.11 The resulting curves are shown as green lines in Fig. S14(a), S16(a) and S18(a)† and the respective parameters displayed in Table S2.†

Concerning temperature dependence, plots of the logarithmic relaxation times of 1–3 vs. the reciprocal temperature show two different regions with an approximately linear behavior at higher temperatures (Fig. 4(b)–6(b)). In order to describe the energy transfer between the magnetic spin system and the lattice vibrations, different possible processes are considered:28,44 A direct process of a resonant phonon, a Raman and an Orbach process describing multiphonon processes via real and virtual intermediate states, respectively. In addition, quantum tunneling may play an important role at zero dc fields. Firstly, the thermal variation of τ at higher temperatures was fitted to an Arrhenius expression 29 (eqn (S5)† where τ₀ is a preexponential factor and Uₑff is the principal energy barrier for reversing the magnetization direction, leading to slightly differing values of Uₑff for 1–3 (Table 4) and relaxation times in the order of 10⁻⁶ s, typical for an Orbach process in such compounds. However, the observed energy barriers are by far smaller than the calculated ones (U ~ S²|D| for integer spin) which is a commonly observed phenomenon and could be assigned to the presence of quantum tunneling

Fig. 4 (a) Temperature dependence of the out-of-phase χ'' component of the ac magnetic susceptibility at H_{dc} = 1500 Oe at different frequencies for 1 (solid lines correspond to the fits to a distribution of single relaxation processes (eqn (S2) and (S3))†; (b) relaxation time (τ) versus the inverse temperature (T⁻¹) for 1. Red and green lines represent fits according to an Arrhenius law (eqn (S5))† and to eqn (S6),† respectively.

Fig. 5 (a) Temperature dependence of the out-of-phase χ'' component of the ac magnetic susceptibility at H_{dc} = 1500 Oe at different frequencies for 2 (solid lines correspond to the fits to a distribution of single relaxation processes (eqn (S2) and (S3))†; (b) relaxation time (τ) versus the inverse temperature (T⁻¹) for 2. Red and green lines represent fits according to an Arrhenius law (eqn (S5))† and to eqn (S6),† respectively.
Effects.\(^{30}\) Secondly, the whole temperature dependence of \(\tau\) showing a strong curvature feature down to low temperature was fitted by eqn (S6)\(^{\dagger}\) including contributions from direct, Raman, quantum tunneling and Orbach relaxation mechanisms. Starting values for the Orbach process \(\left( \text{U}_{\text{Orb}} \text{ and } \tau_{\text{Orb}} \right)\) were taken from the ‘Arrhenius fits’ and the parameters \(B_1\) and \(B_2\) of the quantum tunneling term were taken as fixed parameters from fits of the field dependence of the relaxation time according to eqn (S4)\(^{\dagger}\) (Fig. 4(b)–6(b)).

These fits also result in similar parameter sets (Table 4) for \(1\)–\(3\). The values for \(\text{U}_{\text{Orb}}\) and \(\tau_{\text{Orb}}\) for the Orbach process change only moderately compared to those of the ‘Arrhenius Fits’, and the parameters \(A_{\text{Dir}}\) and \(C_{\text{Ram}}\) for the direct and Raman relaxation process are of similar magnitude for all three compounds. We note that in the case of \(2\), a reasonable fit could not be obtained including the low temperature data. Therefore, only values of \(\tau\) for temperatures above 2.6 K were considered. The broadening of the \(\chi''\) vs. \(\nu\) curve of \(2\) at low temperature (Fig. 5a) might be indicative for a second relaxation pathway or the occurrence of a phonon bottleneck effect.\(^{31}\)

The Cole–Cole/Argand plots \(\chi''\) vs. \(\chi'\) (see Fig. S20–S22\(^{\dagger}\)) show semicircular isotherms. Values of the distribution coefficient \(a\) were found to decrease upon heating which additionally supports the presence of other relaxation processes at lower temperatures.

Theory

In order to investigate the electronic and magnetic structures, multi reference \(ab\) initio calculations (see Experimental section for details) were performed for \(1\)–\(7\). All states corresponding to the \(^{5}\text{D}\) state of the free \(\text{Fe}^{2+}\) \((d^{6}, \text{high-spin})\) ion are considered, differing mainly in the form of the doubly occupied d-orbital. The results of these calculations are summarized in Tables 5–7 and Fig. 7 as well as Tables S3–S9.\(^{\dagger}\)

The scalar-relativistic \(\text{CAS(12,13)SCF}\) and \(\text{CAS(12,13)PT}\) calculations reveal quasi-degenerate ground manifolds for the phosphine and arsine complexes \(1\)–\(3\) and \(7\), in the following denoted type A (Table 5 and S3). On the other hand, the scalar-relativistic ground state of the other complexes \(4\)–\(6\) with an \(\text{OPPh}_3\), pyridine and thf ligand respectively (denoted type B) is, especially in \(4\), well isolated from the first excited state. As the mixing of the two first states anticorrelates with the energy gap, one has to expect large spin–orbit coupling

![Fig. 6](image)

(a) Temperature dependence of the out-of-phase \(\chi''\) component of the ac magnetic susceptibility at \(H_{\text{ac}} = 1500\) Oe at different frequencies for \(3\) (solid lines correspond to the fits to a distribution of single relaxation processes (eqn (S2) and (S3))

(b) Relaxation time \(\tau\) versus the inverse temperature \(T^{-1}\) for \(3\). Red and green lines represent fits according to Arrhenius laws (eqn (S5)\(^{\dagger}\)) and eqn (S6),\(^{\dagger}\) respectively.

**Table 4** Magnetic relaxation parameters\(^a\) derived from fittings of the ac data (Fig. 4–6 and S19)\(^{\dagger}\) either to a pure Orbach process (eqn (S5)\(^{\dagger}\)) or to a combination of an Orbach and Raman relaxation process (eqn (S6)\(^{\dagger}\)).

| \(H_{\text{dc}}\) [Oe] | \(U_{\text{eff}}\) [cm\(^{-1}\)] | \(\tau_{0}\) [s] | \(U_{\text{eff}}\) [cm\(^{-1}\)] | \(\tau_{0}\) [s] | \(A_{\text{Dir}}\) [s\(^{-1}\) K\(^{-1}\)] | \(C_{\text{Ram}}\) [s\(^{-1}\) K\(^{-1}\)] |
|-----------------|-----------------|----------------|-----------------|----------------|----------------|----------------|
| \(1\)            | 1500            | 21.4(4)        | 2.4(1) \times 10^{-6} | 20.5(2)        | 9.7(7) \times 10^{-6} | 12.2(3)        |
| \(2^\prime\)     | 1500            | 23.7(3)        | 2.2(1) \times 10^{-6} | 26.2(9)        | 6(1) \times 10^{-6} | 41.0(1)        |
| \(3\)            | 1500            | 17.9(2)        | 3.4(2) \times 10^{-6} | 14.2(2)        | 2.0(1) \times 10^{-5} | 15.6(6)        |
| \(7^\dagger\)    | 600             | 16.0(3)        | 1.6(2) \times 10^{-6} | 17.1(16)       | 7(3) \times 10^{-7} | 53.2(11)       |

\(^a\) Parameters: external magnetic field \(H_{\text{dc}}\), effective energy barrier \(U_{\text{eff}}\), relaxation time \(\tau_{0}\), parameters for direct and Raman relaxation process \(A_{\text{Dir}}\) and \(B_{\text{Ram}}\).\(^{\dagger}\) The parameters \(B_1\) and \(B_2\) of the quantum tunneling term in eqn (S6)\(^{\dagger}\) were taken as fixed values from fittings of the field dependence of \(\tau\) according to eqn (S4)\(^{\dagger}\) (Fig. S14(a), S16(a) and S18(a)).\(^{\dagger}\) The temperature dependence of \(\ln(\tau^{-1})\) for \(2\) was fitted only with data points in a region from 2.6 to 8.8 K due to the increasing broadening of the \(\chi''\) peaks at \(T < 2.8\) K.\(^{\dagger}\) Data from ref. 14.
complexes are mixed heavily. From the evaluation of the con-spin di
from the SO-RASSI energies (Fig. 7, Table 6 and S4
whereas in
between the ground state and first excited state in
restrictions on the excitation manifold, see Table S3.
Table 5 Spin-free MS-CAS(12,13)PT2 energies E (in cm
E with respect to the ground state
Table 6 SO-RASSI energies (in cm
Table 7 Effective spin Hamiltonian parameters (eqn (2) and (3)) calculated for the lowest Pseudo Spin Multiplet (S = 2) from RASSI wave functions (CAS SCF state(12,13), MSPT2) \(^{a, b}\)

\[
\begin{align*}
g_\lambda &\quad 0.189 &\quad 0.192 &\quad 2.88 &\quad -54.27 &\quad \pm 0.75 \\
g_\sigma &\quad 0.93 &\quad 0.93 &\quad 2.89 &\quad -49.39 &\quad \pm 0.62 \\
g_\pi &\quad 0.87 &\quad 0.92 &\quad 3.04 &\quad -56.77 &\quad \pm 0.73 \\
D &\quad 1.99 &\quad 2.14 &\quad 2.31 &\quad +12.46 &\quad \pm 3.82 \\
E &\quad 1.96 &\quad 2.09 &\quad 2.63 &\quad -30.81 &\quad \pm 3.14 \\
5a &\quad 1.97 &\quad 2.11 &\quad 2.58 &\quad -27.42 &\quad \pm 3.48 \\
5b &\quad 1.98 &\quad 2.11 &\quad 2.51 &\quad -23.67 &\quad \pm 3.61 \\
7 &\quad 1.90 &\quad 1.95 &\quad 2.77 &\quad -52.41 &\quad \pm 1.17 \\
\end{align*}
\]

\(^{a}\) For more results from comparative calculations and concerning restrictions on the excitation manifold, see Table S3. \(^{b}\) Experimental values from ref. 14.

between the ground state and first excited state in 1–3 and 7 whereas in 4, this effect should be very small. Indeed, this different influence of spin–orbit coupling can also be seen from the SO-RASSI energies (Fig. 7, Table 6 and S4†) where type B complexes show a clear energy gap after the first five spin–orbit coupled states whereas the first ten states in type A complexes are mixed heavily. \(^{32}\) From the evaluation of the con-

\textbf{Table 6} SO-RASSI energies (in cm$^{-1}$) of the lowest 11 states of 1–7 based on CAS(12,13)/SCF wave functions and MS-CAS(12,13)PT2 energies (Fig. 7)

\begin{tabular}{cccccccc}
1 & 2 & 3 & 4 & 5a & 6 & 7 \\
1 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
2 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\
3 & 0 & 136 & 146 & 152 & 27 & 81 & 142 \\
4 & 140 & 151 & 157 & 53 & 101 & 83 & 148 \\
5 & 208 & 232 & 243 & 56 & 127 & 99 & 222 \\
6 & 539 & 486 & 490 & 1728 & 797 & 1004 & 523 \\
7 & 591 & 554 & 565 & 1736 & 827 & 1025 & 585 \\
8 & 627 & 588 & 593 & 1752 & 841 & 1038 & 619 \\
9 & 750 & 722 & 734 & 1786 & 927 & 1104 & 750 \\
10 & 751 & 723 & 735 & 1788 & 927 & 1105 & 731 \\
11 & 4264 & 4053 & 4429 & 3482 & 3722 & 3742 & 3621 \\
\end{tabular}

\textbf{Fig. 7} SO-RASSI energies of the first ten states of 1–7 based on CAS(12,13) SCF wave functions and MS-CAS(12,13)PT2 energies (Table 6 and S4†).
reported previously, 6 does not show slow relaxation of the magnetization in related ac experiments. So far, we did not succeed to obtain pure samples of 5 what prevents us to perform related ac measurements. In view of the slightly larger D and slightly smaller E than in 6, it is difficult to predict the magnetic ac properties of 5.

In case of type A complexes, an analysis using an S = 2 spin Hamiltonian may be less meaningful considering the mixing of the lowest CAS states. However, even in this case, an analysis of the lowest two states can be done using an effective S = 1/2 spin Hamiltonian\(^{14}\) \(H(S = 1/2) = g_z^\text{eff}S_zB_z + \Delta S_z\) where \(\Delta\) models the zero-field splitting between the two states in \(H(S = 1/2)\) and is related to the D and E values of \(H(S = 2)\) (eqn (4)) by \(\Delta = 2D(-1 + \sqrt{1 + 3(E/D)^2}) \approx 3E^2/D\) assuming \(D < 0\). \(g_z^\text{eff}\) is an effective \(g\)-factor with \(8 < g_z^\text{eff} \leq 12.\)\(^{37}\) Though reproducing the low-lying spectrum, \(H(S = 1/2)\) cannot be used to interpret magnetic relaxation in terms of electronic states because of the symmetry properties of the zero-field term. Extracted \(g\)-factors for pseudo spins \(S = 2\) and \(S = 1/2\) models, together with main magnetic axes can be found in Table S7.† The main magnetic axes for the lowest pseudo spin multiplet (both \(S = 2\) and \(S = 1/2\)) is oriented for 1–7 in the molecular plane defined by the two nitrogen atoms and the ligand L (along x in Fig. 1, Table S7†). Following ref. 37, the spin-only contribution to \(g_z^\text{eff}\) amounts to \(25g_z^\text{eff} = 8.0092\). Consequently, large orbital contributions are found in type A complexes with \(3 > 2 \geq 1 > 7\) and smaller contributions in type B complexes with \(5a > 5b > 6 > 4\), implying a higher degree of quenching of angular momentum in the latter. The same trend can be found for \(S = 2\), where the orbital contributions to \(g\) are most pronounced in \(z\)-direction, with \(g_z^\text{eff}\) being collinear to \(g_z^\text{eff}\), with smaller deviation from \(g_z^\text{eff}\) in \(x\) - and \(y\)-direction. A very similar trend has been found in a related Fe\(^{3+}\) chloro-\(β\)-diketiminate complex.\(^{37}\)

Following ref. 37, the splitting \(\Delta\) between \(d_{xz}\) and \(d_{yz}\) orbitals may be computed from the effective \(g\)-factor at \(S = 1/2\) using a spin–orbit coupling constant \(\lambda = (436.2\ \text{cm}^{-1})/4\) (Landé-rule, value taken from NIST\(^{35}\)) by eqn (1)

\[
g_z^\text{eff}(\Delta) = 4g_z^\text{eff} + 2\sqrt{3} \cos \left(2\tan^{-1}\left(\frac{4\sqrt{3}\lambda}{\Delta e} - \frac{\sqrt{4\sqrt{3}\lambda^2}}{\Delta e} + 1\right)\right)
\]

(1)

The relation between \(g_z^\text{eff}\) and the splitting is plotted in Fig. S29.† The values found using this simple method amount to 412, 339, 3020, 818, 954, 1160, and 607 cm\(^{-1}\) for 1, 2, 4–7, whereas the value for 3 is undetermined. These values are similar to the values computed with quantum chemical methods (Table S8†).

The complicated nature of electron correlation in multireference-type computations makes it difficult to deduce general rules that can be used for the prediction of electronic and magnetic properties, respectively. However, natural orbitals and energies may be used to construct effective one-particle models that can be useful to a certain extent.\(^{3,36}\) Therefore, effective d-orbital splittings have been computed from the CAS(12,13)SCF wave functions in order to investigate the energetic position of the \(d_{xz}\) orbital relative to the others (note that the complexes lie in the \(xy\)-plane with the ligand L pointing along the \(y\)-axis; Fig. 1). The computed splittings for type A (1–3 and 7) and type B (4–6) complexes are shown in Fig. 8. Due to many-body interactions, it is not unexpected that these values do not reproduce the classical trigonal-planar splitting derived for the d\(^3\) case.\(^{37,41}\) However, the findings from above are confirmed, i.e. upon ‘filling’ the levels with six electrons, type B complexes possess a doubly occupied \(d_{xz}\) orbital yielding a non-degenerate ground state, whereas in type A complexes, \(d_{yz}\) and \(d_{xz}\) orbitals are close in energy with \(d_{yz}\) being lower. Hence, the d-orbital splittings can be used to interpret the low-energy spectrum and thus the magnetic anisotropy in these types of complexes. In order to do that, we computed the d-orbital splittings at different levels of theory by three different methods (Table S8†),\(^{39}\) each of which accounts for electronic effects to a different extent: (1) CAS(5,5)SCF (ROHF) computations on \(d^5\) (\(\text{Mn}(n)\)) derivatives of the complexes include leading interactions between ligands and the d-shell, (2) CAS(6,5)CI computations include d–d correlation effects on top of the orbitals generated by method (1), (3) state average CAS(6,5)SCF computations in which the orbitals of method (2) are allowed to relax. The results are then compared to the CAS(12,13)SCF results shown above, as well as to well-established time-dependent density functional theory results which are known to yield good results in cases with non-degenerate states (complexes 4–6).\(^{14}\)

![Fig. 8 Splitsings of the d-orbital of (a) 1–3 and 7 (type A) and (b) 4–6 (type B) computed from CAS(12,13)SCF wave functions and energies (Table S8†). The grey areas mark the distribution of energies found in the different complexes.](image-url)
Using these methods, the ‘bare’ splitting of d-orbitals is obtained from method (1) by eliminating multi-reference correlation effects in the d-shell going from the 3D ground manifold of Fe(u) to the spatially non-degenerate 6S ground state of Mn(u). These results already reflect the findings of the CAS(12,13)SCF reference calculations with respect to the orbital occupation in the ground state, but in the case of type A complexes do not show the quasi-degeneracy. This is improved by considering d-d correlation and orbital relaxation effects in methods (2) and (3), respectively. For complexes 4 and 6, the results from method 1 are slightly better than the ones obtained with method (3) although the latter is considered superior. However, the differences are generally small and all methods reproduce the findings of the reference calculations qualitatively. The wave functions from method 3 where used to further investigate the mechanistic origin of the differences in type A and type B complexes (see below).

Using the combination of the analysis of the spin–orbit states and the d occupation in the spin-free states, the difference of type A and type B complexes may be summarized as follows, keeping in mind the limitations of ligand-field theory. The spin-free ground states of type A and B complexes are significantly different. In type B complexes, the d_{z^2} orbital is doubly occupied in the ground state, the next states are separated by more than 900 cm\(^{-1}\) and their spin–orbit coupling is weak. In consequence, the energy range of the first five spin–orbit states is small inhibiting single molecular magnet behavior in combination with a large E/D value and a rather isotropic g-factor. In contrast for type A complexes, the spin-free ground state is quasi-degenerate (with an energy difference <350 cm\(^{-1}\)) due to the quasi-degeneracy of the d_{z^2} and d_{xy} orbitals. For these complexes spin–orbit coupling is large resulting in ten spin–orbit states at low energies which mainly consist of the first two spin-free quintet states. The first two spin–orbit states are degenerate and separated by ~150 cm\(^{-1}\) from the next states enabling single molecular magnetism in combination with a large negative D value and strong g-factor anisotropy.

Chemically, a major difference is that the ligands in type A complexes comprise phosphine and arsine moieties while the coordinating atoms in type B complexes are oxygen and nitrogen. In ref. 11, the bonding mechanisms of σ-donor methanide, amide and alkoxide ligands in linearly coordinated Fe(u) complexes has been studied in detail, revealing complicated mechanisms, involving 3d–s mixing, x-bonding anisotropy and orbital mixing of ‘σ-π type’. Due to its π-acceptor capabilities via σ*-orbitals involving the second ligand shell,\(^{40,41}\) a proper ligand field consideration of type A complexes can be considered to be even more complicated in terms of the orbitals involved. However, since the compounds investigated are fairly symmetric (see Table 2), one may distinguish bonding interactions using symmetry arguments: in terms of the Stevens operator equivalents,\(^ {42}\) an ideal trigonal planar crystal field splitting is induced by the axially symmetric terms O_{0}^6 and O_{q}^6.\(^ {43}\) The other terms can thus in principle be assigned to distortions from the ideal D_{3h} symmetry. The ligand field splitting of the atomic 5D ground manifold of Fe(u) may therefore be described by means of the effective Hamiltonian (eqn (2))

$$\hat{H}_{CF} = \sum_{k=0,4,8} \sum_{q=-k}^{k} B_{q}^{0} O_{q}^{0}$$

(2)

where the coefficients B_{q}^{0} are real and the Stevens operators O_{q}^0 transform as the tesseral harmonics.\(^ {44}\) Assuming that bonding interactions can be described roughly by means of effective electromagnetic fields characterized by a certain symmetry type denoted by (k,q) (k = 0, 2, 4, ..., q = −k, ..., k) and respective ‘strength’ B_{q}^{0}, the metal–ligand interactions may be decomposed according to eqn (2).

In order to investigate the relative strength of these interactions, effective model Hamiltonians describing the splitting of the atomic 5D state in complexes 1–7 where constructed using CAS(5,6)SCF energies and wave functions, and subsequently decomposed to give the coefficients B_{q}^{0} in the unit cm\(^{-1}\). The results are summarized in Table S9.\(^ {†}\) In these computations, the only significant differences distinguishing type A from type B complexes are observed in the B_{q}^{0} coefficients. This term describes an interaction potential that has the shape of a d_{z^2–xy} orbital in the xy-plane pointing towards the y-axis. Thus, it has in principle the correct symmetry to describe ‘σ-type’ interactions between the iron atom and the ligand L. On the other hand, the coefficients B_{q}^{0} and B_{q}^{0} which may be related to the respective ‘σ-type’ bonding in the yz and xy planes, respectively, do not display such obvious differences and are by far overpowered by the ‘σ-type’ term B_{4}. Note, however, that all interactions contributing to a symmetry splitting induced by a term (k,q) are collected in B(k,q), i.e. π-bondings may indirectly contribute to terms of σ-symmetry and vice versa.

In order to check the impact of the B_{q}^{0}, B_{q}^{0}, and B_{q}^{0} terms on the ground state symmetry, test simulations have been performed keeping the totally symmetric terms B_{0}^{0} and B_{0}^{0} fixed at their average values (779 cm\(^{-1}\) for B_{0}^{0} and 9 cm\(^{-1}\) for B_{0}^{0}) while varying (a) B_{2}^{0} and B_{2}^{0} and (b) B_{3}^{0} and B_{3}^{0} independently in the range found in complexes 1–7 (−1000 cm\(^{-1}\) ≤ B_{2}^{0} ≤ 0 cm\(^{-1}\), −200 cm\(^{-1}\) ≤ B_{3}^{0} ≤ 150 cm\(^{-1}\)). The resulting surfaces are depicted in Fig. S30.\(^ {†}\) The transition between a doubly occupied d_{z^2} and d_{xy} ground state configuration is, given the average values of B_{2}^{0} and B_{3}^{0}, found to be at around B_{2}^{0} = −555 cm\(^{-1}\) with less negative values corresponding to a doubly occupied d_{z^2} orbital in the ground state of the Fe(u) complexes, which is consistent with the B_{2}^{0} values found for the complexes (type A: −597, −471, −537, −521 cm\(^{-1}\), type B: −278, −375, −350, −363 cm\(^{-1}\)).

From this symmetry analysis, we conjecture that ‘σ-type’ interactions, i.e. integrals of B_{q}^{0}-symmetry dominate the differences in the ligand field splitting of the atomic 3D ground manifold of Fe(u) in the complexes 1–7. Effectively, B_{2}^{0} is connected to a stabilization of the d_{z^2} orbital in comparison to d_{x^2} and d_{y^2}. The stabilization is larger in type A than in type B complexes and induces a change in the ground state occupation to a quasi-degenerate ground state. Whether this
change is really attributable to differences in the σ-bonds of the different ligands L or whether it is connected to π-backbonding indirectly changing B_{2g} cannot be distinguished finally. Furthermore, due to the fact that the operator coefficients computed depend on the choice of the quantization axis (see Experimental section) employed in the computational framework, we cannot exclude that other mechanisms contribute significantly to the electronic structure in the complexes investigated.

Summarizing discussion and conclusion

In this paper, we present a combined experimental and computational analysis of the electronic structure and magnetic properties of a series of trigonal planar iron(II) complexes \([\text{Fe} \{(\text{N(SiMe}_3)_2\}_2\_L\} \_2\] \((L = \text{PPPh}_3, \text{PCy}_3, \text{AsPh}_3, \text{OPPh}_3, \text{thf}, \text{py}, \text{PCy}_3)\). In terms of their experimental dynamic magnetic properties, the complexes can be divided into two classes: type A with ligands \(L = \text{PPPh}_3, \text{PCy}_3, \text{AsPh}_3, \text{PCy}_3\) showing slow magnetic relaxation and those of type B \(i.e. L = \text{OPPh}_3, \text{thf}\) which relax faster than the time scale of the experiment \((\nu = 1500 \text{ s}^{-1})\).

Extensive quantum chemical \textit{ab initio} calculations allow for a more refined picture of the origin of the experimental findings. In all complexes, the first two spin–orbit states are degenerate or quasi-degenerate with an uniaxial g-tensor obtained with pseudospin \(S = \frac{1}{2}\). However, the spin-free electronic origins of these states are quite different: type A complexes reveal quasi-degenerate spin-free scalar-relativistic ground manifolds, whereas the ground state of the type B complexes (including \(L = \text{py}\)) is well isolated from the first excited state, indicating different importance of spin–orbit interaction in these types. This difference is mirrored by the SO-RASSI energy spectrum, where type B (second order) complexes show a substantial energy gap after the first five spin–orbit coupled states (the gap \(\Delta E_S\) increases for the complexes in the order \(L = \text{py}, \text{thf}, \text{OPPh}_3\)). Their ground state wave functions are dominated by the first spin-free quintet state, whereas in the first ten states of type A (first order) complexes the first two spin-free quintet states are heavily mixed.

As already pointed out in an earlier paper,\textsuperscript{14} this different magnetic behavior seems to be related to the heuristic σ/π donor/acceptor strength of the ligands and its effects on the splitting of the d-orbitals. Accordingly, three cases can be distinguished: (1) trigonal planar \([\text{Fe} \{(\text{N(SiMe}_3)_2\}_2\_L\} \_2\] complexes with pure σ donor ligands L like \(L = \text{OPPh}_3\) give rise to a low-lying, isolated, doubly occupied \(d_z\) orbital. As evident from the experiments and quantum chemical calculations, this type of complexes comprises only second order spin–orbit coupling and thus a small magnetic anisotropy of the g tensors (obtained with \(S = 2\)) together with a small \(D\) value.

Furthermore, a large calculated value of \(E\) rationalizes the fast relaxation observed in this complex. (2) In complexes with ligands \(L = \text{thf, py}\) which, in addition to σ-donor properties, also possess weak π-acceptor capabilities, the \(d_z\) orbital is still lowest in energy. However, the experimental and computational data indicate stronger spin–orbit interaction, leading to an increased g-factor anisotropy and larger \(D\) values compared to the complex with the pure σ-donor ligand \(L = \text{OPPh}_3\). Despite this, the relaxation of \([\text{Fe} \{(\text{N(SiMe}_3)_2\}_2\_\text{thf}\}]\) is still fast (not observable in ac measurements up to \(\nu = 1500 \text{ s}^{-1}\) even under applied dc fields). (3) The third case is depicted by type A complexes where \(L\) possess stronger π-acceptor capabilities. Here, the calculations reveal that the \(d_{z^2}\) orbital becomes lowest in energy (doubly occupied in the ground state), being almost degenerate with the slightly higher \(d_{x^2}\) orbital (singely occupied). Thus, strong orbital contributions to the angular momentum arising from spin–orbit coupling of first order type are induced in these complexes which means that a \(S = 2\) pseudo-spin description of the magnetic anisotropy tensors has to be taken with care. However, there is still an energy gap after the first five spin–orbit states whose energies are reasonably well reproduced with a second order spin Hamiltonian (Table S10\textsuperscript{f}). While these considerations overall rationalize the magnetic properties, a quantitative assessment with respect to the σ/π acceptor/donor properties of the ligands \(L\) remains challenging as a symmetry analysis of the ligand field interaction shows.

Concluding, in type A complexes \((L = \text{PPPh}_3, \text{PCy}_3, \text{AsPh}_3, \text{PCy}_3)\), the quasi-degenerate spin-free relativistic ground state leads to an axial magnetic anisotropy via first order spin–orbit coupling, enabling the observation of single-ion magnet behavior, whereas the second order spin–orbit coupling in type B complexes \((L = \text{OPPh}_3, \text{py}, \text{thf})\) induces transversal fields resulting in a faster relaxation.

Experimental section

Synthesis

Standard Schlenk techniques were employed throughout the syntheses using a double-manifold vacuum line with high-purity dry nitrogen (99.9994%) and an MBraun glovebox with high-purity dry argon (99.999%). The solvent heptane was dried over LiAlH\textsubscript{4}, diethyl ether over sodium-benzophenone and both distilled under nitrogen. Li[N(SiMe\textsubscript{3})\textsubscript{2}], PCy\textsubscript{3} (Cy = cyclohexyl, C\textsubscript{6}H\textsubscript{11}), PPh\textsubscript{3}, AsPh\textsubscript{3}, OPPh\textsubscript{3} and anhydrous FeCl\textsubscript{2} were purchased from Aldrich. Li[N(SiMe\textsubscript{3})\textsubscript{2}] was distilled prior to use. \([\text{Fe} \{(\text{N(SiMe}_3)_2\}_2\_\text{PCy}_3]\] \textsuperscript{45} and PMe\textsubscript{3} \textsuperscript{46} were synthesized according to a literature procedure.

\[\text{Fe} \{(\text{N(SiMe}_3)_2\}_2\_\text{PCy}_3\} \_2\] \textsuperscript{7} was synthesized according to a procedure recently published by us.\textsuperscript{14}

Compounds \([\text{Fe} \{(\text{N(SiMe}_3)_2\}_2\_\text{L}\} \_2\] \((L = \text{PPPh}_3 \_1, \text{AsPh}_3 \_3, \text{OPPh}_3 \_4)\) can be synthesized by a procedure analogous to that of \([\text{Fe} \{(\text{N(SiMe}_3)_2\}_2\_\text{thf}\} \_2\] \((0.2 \text{ g}, 0.27 \text{ mmol})\) and two equivalents of the corresponding ligand \((1): \text{PPPh}_3 \_0.139 \text{ g}, 0.53 \text{ mmol}\), 3: AsPh\textsubscript{3} \_0.162 \text{ g}, 0.53 \text{ mmol}\) and 4: OPPh\textsubscript{3} \_0.148 \text{ g}, 0.53 \text{ mmol}\) were dissolved with 2 ml of heptane in a

\textsuperscript{5} Magnetic data have not been measured for \([\text{Fe} \{(\text{N(SiMe}_3)_2\}_2\_\text{py}\}]\) as it could not be isolated purely.
Schlenk tube and heated by an oil bath for five minutes to give a clear pale green solution (1: 100 °C, 3: 85 °C, 4: 100 °C). After cooling down to rt the amount of solvent was reduced by half under a reduced pressure and the reaction solutions stored in a freezer (−42 °C). If the compounds did not crystallize after three days we ‘shock-freeze’ the solutions with liquid N₂, slightly warmed them till everything is dissolved again and put them in the freezer again. In this way 1, 3 and 4 can be obtained as pale green crystals or crystalline precipitates. For their isolation the supernatant solutions were disposed and the crystalline residues in the case of 1 and 4 washed two times with 3 ml of −70 °C cold pentane to give total yields of 1: 0.24 g (72%) and 4: 0.28 g (81%). In the case of 3 washing was not possible due to its high solubility in pentane even at low temperatures (yield: 0.27 g (75%)).

1 C₁₀H₁₅Fe₃PSi₆ (638.90): calcld C 56.4, H 8.1, N 4.4
found C 56.9, H 8.0, N 4.3.

2 (C₂H₅)₂Fe₃PSi₆ (654.90): calcld C 55.0, H 7.9, N 4.3
found C 54.5, H 7.8, N 4.2.

3 C₁₀H₁₅Fe₃AsSi₆ (682.90): calcld C 52.8, H 7.5, N 4.1
found C 53.1, H 7.7, N 4.2.

4 C₁₀H₁₅Fe₃OPSi₆ (654.90): calcld C 56.1, H 8.1, N 4.3
found C 54.5, H 7.3, N 4.5.

[Fe(N(SiMe₃)₂)₂(PMe₃)] (2) can be synthesized by a direct solvent-free reaction of [Fe(N(SiMe₃)₂)₂]₀ (0.3 g, 0.8 mmol) and a slight excess of PMe₃ (0.130 g, 1.7 mmol). The reaction mixture was carefully heated to 70 °C forming a green clear melt. After cooling to rt excess PMe₃ was removed under reduced pressure to give 2 as a solid, pale green crystalline residue.

(2) C₁₅H₄₅Fe₃PSi₆ (452.69): calcld C 39.8, H 10.0, N 6.2
found C 39.3, H 10.0, N 6.3%

Crystallography

Due to the extreme air and moisture sensitivity of the compounds crystals suitable for single crystal X-ray diffraction were selected in perfluoroalkylether oil in a glove box and transferred rapidly under argon atmosphere to the diffractometer equipped with an Oxford Cryosystem. Single-crystal X-ray diffraction data of 1–5 were collected using graphite-monochromatised Mo-Kα radiation (λ = 0.71073 Å) on a STOE IPDS II (Imaging Plate Diffraction System). Raw intensity data were collected and treated with the STOE X-Area software Version 1.39. Data for all compounds were corrected for Lorentz and polarisation effects.

Based on a crystal description a numerical absorption correction was applied for 1–5. The structures were solved with the direct methods program SHELXS of the SHELXTL PC suite programs, and were refined with the use of the full-matrix least-squares program SHELXL. Molecular diagrams were prepared using Diamond. In 1–5 all Fe, As, N, O, P, Si, and C atoms were refined with anisotropic displacement parameters whilst H atoms were computed and refined, using a riding model, with an isotropic temperature factor equal to 1.2 times the equivalent temperature factor of the atom which they are linked to. In 1–4 some of the C atoms of methyl groups were refined isotropic with a split model of site disorder.

Physical measurements

C, H, S elemental analyses were performed on an ‘Elementar vario Micro cube’ instrument.

UV-Vis absorption spectra of 1–4 in C₆D₆ were measured on a PerkinElmer Lambda 900 spectrophotometer in quartz cuvettes.

Zero-Field-Cooled temperature dependent susceptibilities were recorded for 1–4 in dc mode using a MPMS-III (Quantum Design) SQUID magnetometer over a temperature range from 2 to 300 K in a homogeneous 0.1 T external magnetic field. The magnetization curves were measured on the same instrument up to a dc field of 7 T. The ac susceptibility measurements have been performed using a MPMS-XL (Quantum Design) SQUID magnetometer with an oscillating ac field of 3 Oe and ac frequencies ranging from 1 to 1500 Hz. The samples were contained in gelatine capsules filled in a glove box under argon atmosphere owing to the high degree of moisture and oxygen sensitivity of the compounds. The samples were transferred in sealed Schlenk tubes from the glove box to the magnetometer and then rapidly transferred to the helium-purged sample space of the magnetometer. The data were corrected for the sample holder including the gelatine capsule and for diamagnetism using Pascal’s constants.

Details about the simulations are given in the ESI.

Quantum chemical calculations

Electronic and magnetic structures of the complexes 1–5 were studied by ab initio methods. For comparison, calculations were also performed on the published compounds [Fe₃(N(SiMe₃)₂)₂] (L = thf (6)) and [PCy₃ (7)]. All calculations were performed on the crystal structures in gas phase using the MOLCAS program package in version 8.0.15-06-18.

Multi reference wave function models were employed to account for the complicated nature of electron correlation in 1–7. Orbitals were optimized using the complete active space self-consistent field (CASSCF) method, energy corrections were obtained using second order perturbation theory with a complete active space self-consistent field reference function (CASPT2) and spin–orbit coupling was accounted for by using the restricted active space state interaction (RASSI) method in conjunction with the atomic-mean field approximation (AMFI) for the spin–orbit operator (SO-RASSI). Scalar relativistic effects were accounted for in the orbital optimization by means of decoupling of one-electron operators (X2C) using

CCDC 1902301 (1), 1902303 (2), 1902305 (3), 1902304 (4) and 1902302 (5) contain the supplementary crystallographic data for this paper.
the relativistic ANO-type basis sets (ANO-RCC) of triple zeta quality for iron\textsuperscript{56} and the atoms of the first coordination shell\textsuperscript{57} and of double zeta quality for the remaining atoms.\textsuperscript{58}

Two active spaces have been investigated for orbital optimization (1) 3d-shell of the iron centers (CAS(6,5)) and (2) 3d-shell of iron plus a second 3d′ shell with ligand orbital admixtures, as well as the three lone pairs located on the atoms constituting the first coordination sphere pointing towards the iron center (CAS(12,13)). In all cases, the orbitals were optimized using an average density, constructed from the five lowest quintet states with equal weights. Additionally, a CAS(12,13)CI was performed using the CAS(6,5)SCF orbitals in order to distinguish between the effects stemming from orbital optimization and configuration interaction expansion, respectively.

On top of the CASSCF wave functions, single- as well as multi-state CASPT2 calculations where performed using the “afreeze” option to freeze certain inactive orbitals as described in the MOLCAS documentation\textsuperscript{29} with thresholds 0.1 and 0.01 (see Table S3\textsuperscript{†}). In all CAS(12,13) computations, the lowest spin-free states of the CAS(12,13) calculations are spanned by the 100 spin-states of the CAS(6,5)-manifold, i.e. 5 quintets, 45 triplets and 50 singlets. The states arising from different occupation patterns are much higher in energy, with an energy gap for quintets states with equal weights. Additionally, a CAS(12,13)CI using an average density, constructed from the five lowest states, was computed from the five lowest quintet states were shifted by the SS- and MS-CASPT2 correlation energies (eqn (3))\textsuperscript{28,44}

\[
H = \mu_B \sum_{a} B_{\alpha} g_{\alpha} S_{\alpha}
\]

and (4), respectively.

\[
H = D(S_z^2 - 1/3S(S + 1)) + E(S_x^2 - S_y^2)
\]

Because of the quasi-degeneracy of the first two SO-RASSI states, g-factors were also obtained using a pseudo-spin of $S = \frac{1}{2}$ in eqn (3).

In order to account for dynamic correlation effects, the diagonal elements of the spin–orbit CI matrix which are associated to the components of the five quintet states were shifted by the SS- and MS-CASPT2 correlation energies $E_{\text{corr}}(i)$ ($i = 1, \ldots, 5$) (eqn (7)).\textsuperscript{60} The energies of the remaining 95 states where shifted by an arithmetic average correlation energy, i.e. $E_{\text{corr}}(f) \approx 1/5 \sum_{i=1}^{5} E_{\text{corr}}(i)$ for all $j > 5$. Note that all results are based on CASSCF wave functions.

In order to interpret the \textit{ab initio} spectrum and compare to experimentally derived values, the parameters of phenomenological spin Hamiltonians where computed from the \textit{ab initio} data using the SINGLE\_ANIS0 modules documented in detail in ref. 61. These Hamiltonians are used to model the input spectrum in terms of fictitious spins and matrix-valued parameters, e.g. $H_{\text{ZFS}} = SDS$ where $S$ is a spin (vector) operator and $D$ a symmetric $3 \times 3$ matrix which describes the second-order zero-field splitting of a multiplet with pseudo spin $S$. In the case of large spins $S$, i.e. $2S + 1$ degrees of freedom, other (spin-orbit) coupling schemes, or generally in order to increase the accuracy, $H_{\text{ZFS}}$ may be augmented with higher rank terms, e.g. $D'S^4$.

While being in principle able to describe any system of 2$S + 1$ states, a spin Hamiltonian model is physically valid only if there is a clear correspondence between the pseudo-spin and the wave functions of the true Hamiltonian. In lanthanide systems, $S$ usually corresponds to the atomic multiplet. In the case of the second-order interaction modeled by $H_{\text{ZFS}}$ in transition metal complexes, this requires the ligand-field split multiplet to be sufficiently isolated from other states. Then, $D$ describes the splitting induced by second-order spin–orbit and first order spin–spin coupling and may be computed using perturbation theory arguments.\textsuperscript{1,3,34,44} However, in some cases such a low-rank analysis can be performed anyways, and the spectrum of $H_{\text{ZFS}}$, may serve as a marker for assessing the validity of the model.

Magnetic experiments are often conducted at cryogenic temperatures. Therefore, only the lowest states contribute to the properties observed. It is therefore not necessary to reproduce the entire spectrum to high accuracy, but rather the lowest states. Furthermore, adding higher-rank terms promotes overparametrization of the problem, in particular if non-collinear tensors are used at different orders, but give little physical insight on the other hand.

If not mentioned otherwise, the spin Hamiltonian parameters were computed using an effective spin $S = 2$, justified by the energy gap observed after the first five SO-RASSI states. The spin Hamiltonians are given in their main axes by eqn (3) and (4), respectively.

Further CAS computations for analyzing the bonding situation have been carried out using the Bochum suite of \textit{ab initio} programs.\textsuperscript{62,63} In these calculations, the metal atom was equipped with a de2-TZVPP(-f) basis,\textsuperscript{64,65} while the remaining atoms where described by a def2-SVP basis.\textsuperscript{1,4,64} For the computation of magnetic susceptibilities, all 210 microstates where considered in the Boltzmann-averaging.

The Kohn–Sham time dependent density functional theory (TDDFT) calculations where performed with the TURBOMOLE program package.\textsuperscript{66} In these calculations, all atoms where equipped with def2-TZVPP bases.\textsuperscript{64,65} In these calculations, the B3-LYP functional\textsuperscript{67} was used together with a grid of size m3 and the RI-J approximation.

D-orbital splittings where derived from energies and occupations of state-specific natural orbitals as follows: using the state-average CAS orbitals, state-specific density matrices are constructed and diagonalized to give the natural orbitals. Due
to the Full CI nature of the CAS method, the corresponding energies of the states are invariant under this transformation. In the case of ROHF (CAS(5,5)SCF) and TDDFT computations, orbital energies and Jacobian eigenvalues have been used, respectively. The assignments to metal 3d-orbitals have been made by inspecting the orbitals used in the above calculations.

The Hamiltonian decompositions into contributions from irreducible tensor operators have been computed from the CAS energies $E_n$ and wave functions $|n\rangle$ using the program package recently developed in Karlsruhe and Kaiserslautern.\textsuperscript{68}

In this method, the effective Hamiltonian $H_{\text{eff}}$ contains the operator (with $L = 2$ and a quantization axis coinciding with the molecular $z$-axis, see Fig. 1) in the basis of the lowest five CAS states $|n\rangle$, i.e.

$$H_{\text{eff}} = \sum_{n,M} |S,M\rangle \langle S,M'| \langle S,M'| H_{\text{eff}} |S,M\rangle$$

was then used to compute the coefficients $B_{kq}^j$ (eqn (2)) by projecting the effective Hamiltonian onto the matrix representation $O_{ij}^k$ of the Stevens operator equivalents $O_{ij}^{\alpha\beta}$ (see eqn (S8)–(S11)) and then tracing over all pseudo spin functions as described in ref. 61:

$$B_{kq}^j = \frac{\text{tr}(H_{\text{eff}} O_{ij}^k)}{\text{tr}(O_{ij}^k O_{ij}^k)}. \quad (5)$$

**Author contributions**

T. Bodenstein: quantum chemical calculations, A. Eichhöfer: synthesis and characterization.

**Conflicts of interest**

The authors declare no financial interests.

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**References**

1. N. Ishikawa, M. Sugita, T. Ishikawa, S. Koshihara and Y. Kaizu, *J. Am. Chem. Soc.*, 2003, 125, 8694–8695.

2. D. N. Woodruff, R. E. P. Winpenny and R. A. Layfield, *Chem. Rev.*, 2013, 113, 5110–5148.

3. S. Gómez-Coca, D. Aravena, R. Morales and E. Ruiz, *Coord. Chem. Rev.*, 2015, 289–290, 379–392.

4. G. A. Craig and M. Murrie, *Chem. Soc. Rev.*, 2015, 44(8), 2135–2147.

5. A. K. Bar, C. Pichon and J.-P. Sutter, *Coord. Chem. Rev.*, 2016, 308, 346–380.

6. J. M. Frost, K. L. M. Harriman and M. Murugesu, *Chem. Sci.*, 2016, 7, 2470–2491.

7. J. M. Zadrožný, D. J. Xiao, M. Atanasov, G. J. Long, F. Grandjean, F. Neese and J. R. Long, *Nat. Chem.*, 2013, 577–581.

8. C. P. Finn, R. Orbach and W. P. Wolf, *Proc. Phys. Soc.*, 1961, 77, 261–268.

9. P. C. Bunting, M. Atanasov, E. Damgaard-Møller, M. Perfetti, I. Crassee, M. Orliš, J. Ovegaard, J. van Slageren, F. Neese and J. R. Long, *Science*, 2018, 362(6421), 1–9.

10. X.-N. Yao, J.-Z. Du, Y.-Q. Zhang, X.-B. Leng, M.-W. Yang, S.-D. Jiang, Z.-X. Wang, Z.-W. Ouyang, L. Deng, B.-W. Wang and S. Gao, *J. Am. Chem. Soc.*, 2017, 139, 373–380.

11. M. Zadrožný, M. Atanasov, A. Bryan, C.-Y. Lin, B. D. Rekken, P. P. Power, F. Neese and J. R. Long, *Chem. Sci.*, 2013, 4, 125–138.

12. M. Atanasov, M. Zadrožný, J. R. Long and F. Neese, *Chem. Sci.*, 2013, 4, 139–156.

13. P.-H. Lin, N. C. Smythe, S. I. Gorelsky, S. Maguire, N. J. Henson, I. Korobkov, B. L. Scott, J. C. Gordon, R. T. Baker and M. Murugesu, *J. Am. Chem. Soc.*, 2011, 133, 15806–15809.

14. A. Eichhöfer, Y. Lan, V. Mereacre, T. Bodenstein and F. Weigend, *Inorg. Chem.*, 2014, 53, 1962–1974.

15. C. G. Werncke, L. Vendier, S. Sabo-Etienne, J.-P. Sutter, C. Pichon and S. Bontemps, *Eur. J. Inorg. Chem.*, 2017, 1041–1046.

16. M. Atanasov, D. Aravena, E. Suturina, E. Bill, D. Maganas and F. Neese, *Coord. Chem. Rev.*, 2015, 289–290, 177–214.

17. J. P. Malrieu, R. Caballol, C. J. Calzado, C. de Graaf and H. Nojiri, *Proc. Phys. Soc.*, 2013, 177(5110), 429–492.

18. R. A. Andersen, K. Faegri, J. C. Green, A. Haaland, M. F. Lappert, A.-P. Leung and K. Rydral, *Inorg. Chem.*, 1988, 27, 1782–1786.

19. M. M. Olmstead, P. P. Power and S. C. Shoner, *Inorg. Chem.*, 1991, 30, 2547–2551.

20. A. Panda, M. Stender, M. M. Olmstead, P. Klavins and P. P. Power, *Polyyhedron*, 2003, 22, 67–73.

21. N. F. Chilton, R. P. Anderson, L. D. Turner, A. Soncini and K. S. Murray, *J. Comput. Chem.*, 2013, 34, 1164–1175.

22. R. A. Layfield, J. J. McDouall, M. Scheer, C. Schwarzmaier and F. Tuna, *Chem. Commun.*, 2011, 47, 10623–10625.

23. D. Gatteschi, R. Sessoli and J. Villain, *Molecular Nanomagnets*, Oxford University Press, Oxford, 2006.

24. S. Koizumi, M. Nihei, T. Shiga, M. Nakano, H. Nojiri, R. Bircher, O. Waldmann, S. T. Ochsenbein, H. U. Güdel, F. Fernandez-Alonso and H. Oshio, *Chem. – Eur. J.*, 2007, 13, 8445–8453.
37 H. Andres, E. L. Bominaar, J. M. Smith, N. A. Eckert, R. A. Lindh, J. Autschbach, Relativistic Methods for Calculating Electron Paramagnetic Resonance (EPR) Parameters, in Handbook of Relativistic Quantum Chemistry, ed. W. Liu, Springer, Berlin Heidelberg, 2019.

35 NIST Atomic Spectra Database (ver. 5.6.1), https://www.nist.gov/pml/atomic-spectra-database.

19 Regarding the quantum chemical methods employed, one has to distinguish between effects stemming from the self-consistent field (SCF) relaxation of the orbitals and from changing the flexibility of the configuration interaction (CI) wave function in terms of explicitly interacting states.

20 D. G. Gilheany, Chem. Rev., 1994, 94, 1339–1374.

21 J. E. Huheey, E. A. Keiter and R. L. Keiter, Anorganische Chemie, Prinzipien von Struktur und Reaktivität, hrsg. v. R. Steudel, bearb. v. F. Breher, M. Finze, D. Johrendt, M. Kaupp, H.-J. Lunk, U. Radius, U. Schatzschneider, 5te Auflage, 2014, Kap. 15, S. 797, Walter de Gruyter GmbH & Co. KG Berlin/Boston.

22 K. W. H. Stevens, Proc. Phys. Soc., London, Sect. A, 1952, 65(3), 209–215.

23 J. S. Griffith, The theory of transition-metal ions, Cambridge University Press, Cambridge, 1971, ISBN 978-0521051507.

24 A. Abragam and B. Bleaney, Electron Paramagnetic Resonance of Transition Ions, Oxford University Press, 1970.

25 D. Lj. Broere, I. Coric, A. Brosnahan and P. L. Holland, Inorg. Chem., 2017, 56(6), 3140–3143.

26 W. Wolfsberger and H. Schmibaur, Synth. React. Inorg. Met.-Org. Chem., 1974, 4(2), 149–156.

27 A. C. De Vroome, F. F. Lijphart and N. J. Poulis, Physica, 1970, 47, 458–484.

28 R. L. Carlin and A. J. Duyneveldt, Magnetic Properties of Transition Metal Compounds, Springer, Berlin, Heidelberg, New York, 1977.

29 R. J. Glauber, J. Math. Phys., 1963, 4, 294–307.

30 D. E. Freedman, W. H. Harman, T. D. Harris, G. J. Long, C. J. Chang and J. R. Long, J. Am. Chem. Soc., 2010, 132, 1224–1225.

31 J. Soeteman, L. Bevaart and A. J. van Duyneveldt, Physica, 1974, 74, 126–134.

32 F. Neese, J. Am. Chem. Soc., 2006, 128(31), 10213–10222.

33 R. Boča, Coord. Chem. Rev., 2004, 248, 757.

34 H. Bolvin and J. Autschbach, Relativistic Methods for Calculating Electron Paramagnetic Resonance (EPR) Parameters, in Handbook of Relativistic Quantum Chemistry, ed. W. Liu, Springer, Berlin Heidelberg, 2019.

35 NIST Atomic Spectra Database (ver. 5.6.1), https://www.nist.gov/pml/atomic-spectra-database.

36 S. Gómez-Coca, E. Cremades, N. Aliaga-Alcalde and E. Ruiz, J. Am. Chem. Soc., 2013, 135, 7010–7018.

37 H. Andres, E. L. Bominaar, J. M. Smith, N. A. Eckert, P. L. Holland and E. J. Münck, Am. Chem. Soc., 2002, 124, 3012.

38 A. L. Companion and M. A. Komorynskyl, J. Chem. Ed., 1964, 41, 257.

39 Regarding the quantum chemical methods employed, one has to distinguish between effects stemming from the self-consistent field (SCF) relaxation of the orbitals and from changing the flexibility of the configuration interaction (CI) wave function in terms of explicitly interacting states.