Polymorphism and magnetic properties of Li$_2$MSiO$_4$ (M = Fe, Mn) cathode materials

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Transition metal-based lithium orthosilicates (Li$_2$MSiO$_4$, M = Fe, Ni, Co, Mn) are gaining a wide interest as cathode materials for lithium-ion batteries. These materials present a very complex polymorphism that could affect their physical properties. In this work, we synthesized the Li$_2$FeSiO$_4$ and Li$_2$MnSiO$_4$ compounds by a sol-gel method at different temperatures. The samples were investigated by XRPD, TEM, $^7$Li MAS NMR, and magnetization measurements, in order to characterize the relationships between crystal structure and magnetic properties. High-quality $^7$Li MAS NMR spectra were used to determine the silicate structure, which can otherwise be hard to study due to possible mixtures of different polymorphs. The magnetization study revealed that the Néel temperature does not depend on the polymorph structure for both iron and manganese lithium orthosilicates.

Polyanion framework compounds based on PO$_4$ or SiO$_4$ structural units are now under intense study for the application as cathode materials in lithium–ion batteries. All of these materials are characterized by low costs and toxicity, are environmentally friendly, and highly safe$^1$. Along with the well-known phospho-olivine, lithium orthosilicates appear especially promising because they can afford more than one electron reversible exchange per transition metal atom, so increasing the overall cathode capacity. In fact, Li$_2$MnSiO$_4$ can reach the capacity of 333 mAh g$^{-1}$, while Li$_2$FeSiO$_4$ could deliver 166 mAh g$^{-1}$ for the extraction of one Li ion$^2$. However, the low electronic conductivity of silicates has to be overcome in order to reach the theoretical capacity and different approaches have been tried to improve their electrochemical performances, e.g. by mixing Fe and Mn$^{3-6}$, by doping with Cr$^7$, V$^8$, Mg$^9$, Zn, Cu, and Ni$^{10}$, by adding a proper carbon-coating$^{11}$ or by preparing composites with carbon nanotubes$^{12}$.

Another critical feature of the orthosilicates is their rich polymorphism with numerous, different crystal structures that could be stabilized depending on the synthesis conditions$^{13}$. Usually, the monoclinic $P2_1/n$ and the orthorhombic $Pmn_2_1$ or $Pmnb$ space groups are reported for both the Li$_2$MnSiO$_4$ and Li$_2$FeSiO$_4$ compounds. The differences among these structures are mainly due to different arrangements of the cation tetrahedra, and local-order probes such as solid-state Magic Angle Spinning Nuclear Magnetic Resonance (MAS NMR) are very useful to identify the obtained polymorphs and also mixtures of them$^{14}$. This rich polymorphism, with the associated small transition energies, is one of the factors affecting the long-term cyclability of these materials. We recently showed by density-functional theory (DFT) calculations that, under delithiated conditions, Li$_2$MnSiO$_4$ can transform from the $Pmn_2_1$ or $Pmnb$ polymorphs to the electrochemically weakest $P2_1/n$ one, and this mechanism may gradually lead to electrochemical and structural collapse$^{15}$.

The dependence of Li$_2$FeSiO$_4$ electrochemistry on structure was also recently investigated$^{16,17}$, while information regarding polymorphism and magnetic properties is still lacking. Some works report about the antiferromagnetic (AFM) magnetic ordering temperature of Li$_2$MSiO$_4$ (M = Co, Mn, Fe) and the magnetic measurements were mainly used to probe the presence of ferro or ferri-magnetic impurities in the samples$^{18,19}$. In this work we report the results of the magnetic and spectroscopic study of Li$_2$MSiO$_4$ (M = Fe, Mn), prepared via sol-gel synthesis at different temperatures to obtain different polymorphs. The magnetic properties are investigated by SQUID magnetometry and their relationships with the polymorph structure are discussed. The combined use of X-ray powder diffraction (XRPD) and $^7$Li MAS NMR allowed us to individuate the phases in the samples. TEM
microscopy coupled with Selected Area Electron Diffraction (SAED) was also used as a helpful tool for the assignment of the correct structure to the synthesized materials.

**Results**

**XRPD and Rietveld results.** Figures 1a and b show, for comparison, the XRPD patterns of the Li₂FeSiO₄ and Li₂MnSiO₄ samples, respectively. The Li₂FeSiO₄ samples present low amount of different impurities, which can be identified as Fe₃O₄ (JCPDS card No. 19-0629), Li₃Fe₅O₈ (JCPDS card No. 74-1754) and FeO (JCPDS card No. 89-0687). Besides, other differences in the patterns can suggest the presence of different silicate polymorphs. For the Li₂MnSiO₄ samples the main impurities are Li₂SiO₃ (JCPDS card No. 29-0828) and MnO (JCPDS card No.75-1090) and again some pattern differences can be ascribed to the formation of different polymorphs. The Fe samples show a higher degree of purity than the Mn ones (see also Table 1), suggesting that under these synthesis conditions iron silicates can be obtained more easily than the Mn ones.

The Rietveld refinements were performed on the basis of the structural models present in the literature, that take into account the existence of different polymorphs. The results of the Rietveld refinements are reported in Table 1 together with the discrepancy factors. Rwp and S are satisfactory and suggest a good quality of the structural refinement. Different synthesis conditions allowed the stabilization of different polymorphs, as evidenced both for the Li₂FeSiO₄ and Li₂MnSiO₄ compounds. In fact, for the Fe compound synthesized at low temperature the monoclinic P2₁/n s.g. was obtained (see Rietveld refinement in Figure 1c), while at 900 °C the Pmnb space group provided a good description of the structure of the sample. The s.g. assignment depends on some typical reflections, i.e. the peak at about 20/20, distinctive of the Pmnb s.g., and the peak at 31.8°/29 that otherwise pertains to the P2₁/n s.g. In addition, the P2₁/n structure presents three peaks in the range 20–23.5°/29, while only two peaks are found in the same region for the orthorhombic form.

For the Mn compounds, a co-presence of polymorphs was found for the Mn-900, including Pmnb and P2₁/n phases. In case of Mn-650, the Pmn₂₁ structure was found as a single phase.

**NMR results.** The NMR spectroscopy of these materials is traditionally challenging since the paramagnetic trivalent ions induce very large shifts and shift anisotropies, resulting in spectra that are very broad and difficult to excite using standard NMR methods. However, the combined use of ultra-fast MAS (60 kHz)
and sophisticated pulse sequences enabled us to obtain extremely high-quality broadband $^7$Li spectra with a short experiment time of about 16 minutes per spectrum. Figure 2 reports the full $^7$Li MAS NMR spectra of the Mn-650 and Fe-650 samples (parts a and b, respectively). Both spectra are characterized by broad spinning-sideband manifolds which are due to the hyperfine interactions between the unpaired electrons of the transition metal ion and the observed nucleus. The unpaired electrons of the tetrahedrally-coordinated high-spin Mn$^{2+}$ ion are present in the configuration $e_g^3t_{2g}^3$, giving an isotropic $g$-tensor whose value is equal to the free-electron $g$-value $g_e$. In this case, the isotropic contribution to the paramagnetic shift is due to the Fermi-contact interaction (and is referred to as the Fermi-contact shift), and the spinning sidebands arise from the spin-dipolar interaction between the unpaired electrons and the nucleus. The interactions giving rise to the paramagnetic $^7$Li shift for Li$_2$FeSiO$_4$ are more complicated, as the high-spin Fe$^{2+}$ electron configuration of $e_g^3t_{2g}^3$ gives an anisotropic $g$-tensor due to spin-orbit coupling. In addition to the Fermi-contact interaction, the isotropic shift now has a contribution from the pseudo-contact shift, which is due to the coupling of the $g$-anisotropy to the spin-dipolar term in the hyperfine coupling tensor. For a detailed description of the Hamiltonian terms describing these spectra see refs 22,23.

In addition, the $g$-anisotropy can be used to rationalize the difference in the linewidths of the sidebands in each spectrum. The increased broadening in the spectrum of Fe-650 relative to Mn-650 is entirely inhomogeneous and arises from the anisotropic bulk magnetic susceptibility (ABMS) due to the $g$-anisotropy in the former material. It should be noted that the differences in linewidth cannot be explained by the differences in the homogeneous contributions, such as spin-spin relaxation. We measured the $^7$Li homogeneous decay time constants $T_2^*$ for both samples, which were found to be 497 μs and 720 μs for Mn-650 and Fe-650, respectively. The corresponding homogeneous linewidth 442 Hz for Fe-650 is therefore found to be lower than the value of 640 Hz for Mn-650. The linewidth in the spectrum of Fe-650 is therefore dominated by the inhomogeneous contribution.

Figure 3 shows an expansion of the spectral region containing the isotropic peaks, which were identified using the recently-published adiabatic magic-angle turning (aMAT) experiment, together with their best-fits whose results are reported in Table 2. In order to check the sensitivity of our approach, we performed different best-fits by including, beside the isotropic peaks, also up to three couples of spinning sidebands, but the overall agreement did not change significantly, and the ratios among the different components change of about 10–15%.

The spectrum of Mn-650 is characterized by the presence of three peaks at 2.5 ppm, −98.1 ppm and −120.2 ppm (Figure 3a). The downfield peak is attributed to the Li$_2$SiO$_3$ diamagnetic impurity, and accounts for 9.8% of the total observed lithium, in reasonable agreement with the 15.6% obtained by the Rietveld analysis (see Table 1). The peaks upfield are attributed to the $P_{mn21}$ (−98.1 ppm) and $P_{mn2}$ (−120.2 ppm) polymorphs, respectively, in agreement with the literature assignments. The $P_{mn2}$ polymorph accounts for about 10% of the overall active phase. We stress here that further Rietveld refinements performed by using both $P_{mn2}$ and $P_{mn2}$ polymorphs gave us only a marginal improvement of the fit for the Mn-650 sample with respect to that reported in Table 1 (see Discussion). On the other hand, the reflections of $P_{mn2}$ and $P_{mn2}$ polymorphs are very similar. The spectrum of Fe-650 (Figure 3b) is characterized by two peaks at −8.7 ppm and −52.0 ppm, which can be attributed at the two Li sites of the $P2_1/n$ polymorph, in good agreement with the assignment of Sirisopanaporn et al. By considering the spinning sidebands, and therefore

**Table 2**: $^7$Li MAS NMR parameters obtained by spectral best fit

| Sample | Chemical shift (ppm) | Anisotropy (ppm) | Percentage (%) |
|--------|----------------------|-----------------|----------------|
| Mn-650 | Li$_2$SiO$_3$ | 2.5 | 112 | 9.8 |
|        | $P_{mn21}$ | −98.1 | 954 | 8.2 |
|        | $P_{mn2}$ | −120.2 | 36 | 82.0 |
| Fe-650 | $P2_1/n$ Li$_1$ | −8.7 | −1750 | 56.0 |
|        | $P2_1/n$ Li$_2$ | −52.0 | 790 | 44.0 |
the different shift anisotropies, the best-fitted ratio between the two sites is 56:44. No evidence of diamagnetic impurities or other Li-containing active phases is observed, in agreement with the Rietveld refinement.

**SEM and TEM.** SEM analysis showed that for the samples synthesized by sol-gel at 650°C the morphology was constituted by aggregates of small rounded particles, whereas at 900°C the samples appeared to be constituted by large agglomerates of fused particles. SEM micrographs of the as-prepared samples are shown in Figure S1 (ESI).

In order to verify the polymorph assignments obtained by XRPD and NMR, TEM analysis was also performed. However, the measurements to investigate the crystalline structure were performed only on the Fe-900 specimen (Figure 4a), because the samples synthesized at 650°C showed crystallite aggregation (see Figure S2). From the morphological point of view, the sample comprises rounded particles with diameters ranging from approximately 0.2 to 0.5 μm. In general, the particles were too thick to give enough contrast for High Resolution Electron Microscopy. In order to obtain information about the crystallographic structure, therefore, we performed Selected Area Electron Diffraction (SAED) on single grains after orienting them along some low order Zone Axis (ZA). A representative SAED is reported in Figure 4b. To check which space group was responsible for the diffraction patterns many image simulations were performed using the JEMS software. All the ZAs were calculated for both the space groups P21/n and Pmnb. The best result, obtained for the Pmnb space group with the crystal oriented along [011], is reported in Figure 4c, where the simulated electron diffraction pattern is superimposed to the experimental one.

**Magnetic characterization.** Figure 5a shows the temperature dependence of the magnetization (M/H vs. T curves) for the Fe-900 and Fe-650 samples in zero field cooling (ZFC) and field cooling (FC) regimes, obtained by applying a 10000 Oe magnetic field. Both the samples undergo the paramagnetic to antiferromagnetic transition, with Neél temperature TN ≈ 20 K, typically observed for the lithium iron silicate. Deviations from the usual Curie-Weiss behaviour emerge from Figure 5a: a markedly higher M value pertains to the Fe-650 sample, with respect to the Fe-900 one, in the whole investigated temperature range, while a clear-cut separation between ZFC and FC curves is observed, for temperatures lower than about 50 K, only for the Fe-900 sample. The field dependence of the magnetization, investigated at different temperatures, allowed us to verify the origin of these peculiar features. Figure 5b shows the M vs. H curves for Fe-900 and the fit of the linear part of each curve; experimental curves at room temperature, together with their linear fit, for Fe-900 and Fe-650 are instead compared in the inset.

**Figure 4** | (a) TEM micrograph and (b) SAED of the specimen Fe-900; (c) The same experimental pattern (SAED), as reported in Figure b, together with simulated Electron Diffraction pattern; the Miller indexes are also indicated. The drawing makes use of the kinematical theory of electron diffraction. The intensity of the spots is just proportional to their structure factor (bright red spots correspond to higher structure factors while black spots correspond to lowest structure factors). The Double diffraction spots are also indicated in yellow.

**Figure 5** | (a) ZFC and FC temperature dependence of molar magnetization at 10000 Oe for Fe-900 (black line) and Fe-650 (red line) samples. (b) M vs H curves for Fe-900 at different temperatures. The linear fit of the high field region is also shown. Inset: comparison between room temperature M vs H curves of Fe-900 (black symbols) and Fe-650 (red symbols) samples, reported with their linear fits (black lines). (c) ZFC and FC M vs T curves at 10000 Oe for Mn-900 (black), and Mn-650 (red) samples. Inset: FC curves in the low temperature region.
The non-linear M(H) behaviour evident for Fe-900 at 5 K for low magnetic fields suggests the presence of a ferromagnetic-like extrinsic contribution, with a value of magnetization at null field, $M_0(H = 0)$, obtained by extrapolating the linear behaviour at high fields and corresponding to the residual magnetization of such ferromagnetic-like ordered phase, of about 45 emu/mol. This value decreases with increasing the temperature, becoming negligible at r.t. A linear behaviour is indeed generally observed for T ≈ 50 K and H ≈ 500 Oe. The non-null $M_0(H = 0)$ contribution plausibly arises from the impurity phase disclosed by XRPD data, i.e., Li$_3$Fe$_5$O$_8$, for which a ferro- or ferri-magnetic behaviour can be reasonably supposed. This assumption is well supported by quantum-mechanical calculations$^{29}$, which, taking into account magnetic interactions, foresee an arrangement of Fe atoms, in the structure of Li$_3$Fe$_5$O$_8$, compatible with both ferromagnetic and ferrimagnetic orderings, being the ferrimagnetic arrangement energetically favoured. Incidentally, we remark that the $M_0(H = 0)$ value here obtained at 5 K corresponds to about 1 Bohr magneton per unit formula, indeed supporting a ferrimagnetic ordering for this phase. The small amount of Li$_3$Fe$_5$O$_8$ in the sample can also be responsible of the separation between ZFC and FC M vs. T curves (see Figure 5a).

A M(H) non-linear behaviour at low magnetic fields is instead always detected for the Fe-650 sample, with a $M_0(H = 0)$ value ranging between 135 and 140 emu/mol, in the whole T range. This value corresponds to the additive contribution to the M vs. T curve pertaining to the Fe-650 sample, shown in Figure 5a. We can undoubtedly attribute this additive contribution to the ferrimagnetic Fe$_5$O$_8$ phase, detected by XRPD in this sample in very small amount, whose $T_N$ value is much higher than r.t.$^{30}$. Taking into account the $M_0$ value of this phase (92 emu/m at r.t.$^{30}$), the amount of Fe$_3$O$_4$ in Fe-650, as revealed by the saturation magnetization values, results to be in very good agreement with the XRPD data (Table 1). No sign of the antiferromagnetic (AF) transition of the small amount of FeO phase (T$_N$ ≈ 186–198 K$^{30}$), disclosed by XRPD, is instead detectable in the M vs. T curve of Fe-650.

In order to avoid the contribution of saturated magnetic phases, Curie-Weiss constants for our lithium iron silicates have been inferred considering the $1/\chi_{mol}(T)$ values obtained from the slopes of the linear part of the M vs. T curves at different temperatures ($\chi(T) = dM(T)/dH, T = 300, 200, 100, 50, 5 K$) in the T range corresponding to the linear paramagnetic region.

For Fe-650 the obtained Curie constant leads to an effective magnetic moment, $\mu_{eff}$, for the iron silicate very near to 5 $\mu_B$, thus consistent with a spin only contribution of Fe$^{2+}$ in the high spin configuration. Besides, a Weiss constant, $\theta$, of about −35 K has been obtained. We remark that these values, and the related $T_N$ value, are in very good agreement with those reported in the literature for this compound$^{1}$. A higher $\mu_{eff}$ value has instead been obtained for Fe-900 ($\mu_{eff} \approx 5.15$), which could imply, for example, the contribution, in addition to the one of divalent iron ions, from Fe ions with higher oxidation state. Nevertheless, this should not be the case because a $T_N$ value of 20 K in this compound is related to magnetic interactions between Fe$^{2+}$ spins only, while the coexistence of Fe$^{3+}$ and Fe$^{2+}$ ions should give rise to AF ordering with a $T_N$ value higher than 20 K$^{31}$. On this basis, an extrinsic contribution can be instead invoked to explain the higher estimated $\mu_{eff}$ value for Fe-900, as, for example, the one due to the small amount in the sample of the ferrimagnetic Li$_3$Fe$_5$O$_8$ phase not fully saturated in the whole temperature range considered to estimate Curie and Weiss constants. Moreover, it must be taken into account that even a small amount of ferromagnetic-like impurity phase can enhance the $\theta$ value of the sample. Indeed, for Fe-900 a higher $\theta$ value ($\theta = −60$ K) has been estimated with respect to Fe-650, against the unchanged $T_N$ value of the iron silicate phase.

In view of the small amount of well-characterized impurity phases in the samples, magnetization data attest the good quality and stoichiometry of the lithium iron silicate phases.

The magnetic characterization of Li$_2$MnSiO$_4$ samples is affected by the presence, together with the silicate, of a great amount of spurious phases, both Li-/Si- based compounds, diamagnetic, and manganese oxide, characterized by a paramagnetic-to-antiferromagnetic transition with a T$_N$ value (112–115 K$^{30}$) falling inside the investigated temperature range. Figure 5c reports the ZFC and FC temperature dependence of the molar magnetization at 10000 Oe for the two investigated samples. Both of them clearly display the typical Li$_3$MnSiO$_4$ paramagnetic to antiferromagnetic transition with Néel temperature T$_N$ ≈ 12 K$^{32}$. A small, but not negligible difference between the T$_N$ values is evident in the enlargement of the FC curves, reported in the inset of Figure 5c. In particular, a lower value (=11 K) is detected for the sample synthesized at the lower temperature. No sign of the AF transition related to the manganese oxide is detectable in the M vs. T curves at the foreseen T$_N$ value even for Mn-650, for which a remarkable MnO amount is indeed disclosed by XRPD.

The Curie-Weiss constants have been extracted from 1/\chi_{mol}(T) values in the high temperature paramagnetic region. The contribution of the MnO impurity phase to the overall paramagnetic behaviour of the samples has been taken into account to try to estimate the $\mu_{eff}$ values for the main lithium manganese silicate phase. For Li$_3$MnSiO$_4$ in Mn-900, $\mu_{eff}$ is found to be near to 6 $\mu_B$, consistently with a spin only contribution of Mn$^{3+}$ in high spin configuration. A 0 value of about −35 K has been obtained for this sample. These values are in fair agreement with those reported in literature for the manganese lithium silicate$^{33}$.

For the Mn-650 sample a higher $\theta$ value has been obtained with respect to Mn-900, consistently with the presence of a much higher MnO amount, for which the antiferromagnetic interactions are stronger with respect to the silicate phase. Besides, a $\mu_{eff}$ value appreciably lower than 6 $\mu_B$ has been inferred in this case and, although this estimate can be strongly affected by the low purity of the sample, it seems however to suggest the presence, together with divalent manganese ions, of an appreciable amount of Mn ions with higher oxidation states in the Li$_3$MnSiO$_4$ phase.

**Discussion**

The combined use of XRPD and $^7$Li MAS NMR allowed us to recognize the stable polymorphs for the lithium iron/manganese silicates synthesized at the different temperatures. The main difference between the two series of samples concerns the possibility of obtaining a single phase for Li$_2$Fe$_2$SiO$_4$ independently of the synthesis temperature, while Li$_2$MnSiO$_4$ was obtained as a single polymorph only at 650°C. Besides, in the case of Li$_3$MnSiO$_4$ we observe the stabilization of the Pnnm polymorph at 900°C in agreement with the findings of Gummock et al.$^{14}$, who recently discussed the formation of the Pnnm structure for the Li$_2$MnSiO$_4$ material synthesized at 900°C, with a different connectivity of LiO$_4$ and MnO$_4$ tetrahedra with respect to the Li$_2$FeSiO$_4$ Pnnm polymorph.

We remark that $^7$Li MAS NMR was chiefly used to check that the polymorphs found by XRD Rietveld analysis were correct, in particular for the samples which where found to be single phase. A very good agreement was found for the Fe-650 sample, while some discrepancies were observed in the case of Mn-650. At first, the Rietveld refinement was carried out by considering only the Pnnm$_2$ structure but, on the basis of the NMR results, the Pnnm one was also considered afterwards. In fact, a slight improvement of the fitting was obtained (R$_{wp}$ = 11.6 vs. 12.9), and the amount of the Pnnm$_2$ phase was 15%, in reasonable agreement with the about 10% value obtained by NMR. At the light of these results, we believe that XRPD cannot easily discriminate between so similar phases, such as the Pnnm$_2$ and Pnnm polymorphs, and that the parallel use of a powerful local probe such as $^7$Li MAS NMR is mandatory.

Once the crystal structures of the samples have been precisely determined, the relationship between the polymorphic form and
the physical properties was investigated. In particular, we focused our attention on the magnetic properties. We recall that the antiferromagnetic ordering in Li$_2$MnSiO$_4$ comes from the long-range M-O-Li-O-M interactions. Our data, in addition to what reported in the literature, do support the hypothesis that the structural differences between the different polymorphs is too small to affect the interaction paths to such an extent that the $T_N$ value may change. Indeed, as already discussed, the same $T_N$ value was detected for the lithium iron silicate phase in Fe-650 and Fe-900, which are characterized by the monoclinic $P2_1/n$ and the orthorhombic $Pmn2_1$ polymorphs, respectively. Moreover, the same $T_N$ value was also detected for the orthorhombic $Pmn2_1$, lithium iron silicate $^{31,32}$. Then, a clear correlation cannot be found between polymorphism and magnetic features of Li$_2$FeSiO$_4$. Rather, the $T_N$ value of the lithium iron silicate can be influenced by the coexistence between Fe ions with different oxidation states.$^{33}$ Also the results shown above for the Li-Mn silicates are in agreement with these hypotheses: the small $T_N$ shift from 12 K (i.e. from the value commonly reported in literature for stoichiometric Li-Mn silicates $^{33}$) is indeed observed for the sample for which the coexistence of Mn$^{2+}$ ions with manganese ions with different oxidation states has been disclosed.

In conclusion, Li$_2$FeSiO$_4$ and Li$_3$MnSiO$_4$ compounds were synthesized by a sol-gel route, and the polymorphic form and impurity phases were investigated by using XRPD, $^{34}$ Li MAS NMR and magnetization measurements. The magnetic analysis was applied to obtain information not only on the general presence of ferro- or ferrimagnetic impurities, but also to verify their nature and quantify them. Besides, the study of the silicate magnetic features allowed us to rule out the dependence of the $T_N$ on the polymorphism.

A new procedure of 7Li NMR spectra collection was applied, based on the combined use of very fast MAS rotation and sophisticated pulse sequences. The clear advantage of using $^{7}$Li with respect to the less abundant $^{6}$Li, and obtaining similar quality or even better spectra through Hydrothermal-Assisted Sol-Gel Process Batteries and Energy Storage.

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Author contributions
M.C.M. performed the magnetic measurements with related data analysis and wrote the magnetic section. C.F., A.J.P., G.P. and P.M. collected and analysed the NMR spectra. P.C. performed TEM analysis. M.B., S.F., D.C. performed XRPD analysis. S.F. performed the synthesis. M.B. and S.F. wrote the manuscript and all authors reviewed the manuscript.

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