# An Fe (III) wheel with a zwitterionic ligand: the structure and magnetic properties of $\left[\mathrm{Fe}(\mathrm{OMe})_{2}(\text { proline })\right]_{12}\left[\mathrm{ClO}_{4}\right]_{12}$ 

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The structure and magnetic properties of a dodecanuclear iron(III) wheel are reported, featuring a bridging zwitterionic ligand.

Cyclic metal cages featuring paramagnetic ions have been heavily studied since the report by Lippard and co-workers of the first "ferric wheel". ${ }^{1}$ The majority are either octa- or deca-nuclear; the metal $\cdots$ metal vectors in the former are normally bridged by one $\mu_{2^{-}}$ ligand (e.g. fluoride) and two carboxylates, ${ }^{2}$ in the latter the metal $\cdots$ metal edges are normally bridged by two $\mu_{2}$-ligands (e.g. methoxide) and one carboxylate. ${ }^{1,3}$ More recently dodecanuclear wheels have been reported featuring the latter bridging motif, but in both cases the wheels become slightly "buckled", i.e. the metal cores are no longer planar. ${ }^{4,5}$ There are also reports of wheels where the edges are asymmetrically bridged. ${ }^{6}$ We have been investigating complexes with a bridging L-prolinato ligand, ${ }^{7}$ which can become zwitterionic, ${ }^{8}$ and here demonstrate that when it features in a "ferric wheel", a planar dodecanuclear wheel results which carries a $12+$ charge.
$\left[\mathrm{Fe}(\mathrm{OMe})_{2} \text { (proline) }\right]_{12}\left[\mathrm{ClO}_{4}\right]_{12} \mathbf{1}$ is formed from the reaction of hydrated iron(II) perchlorate ( 6 mmols ) with L-proline ( 8 mmols ) in methanol ( 150 ml ). The solution was stirred for ten minutes and crystals of $\mathbf{1}$ grew after standing for several days at room temperature. The yield was $34 \% . \dagger$ Caution. Perchlorate salts of metal complexes are potentially explosive and should be handled with care.
The structure $\ddagger$ of the cation (Fig. 1) shows twelve Fe (iiI) centres, with each $\mathrm{Fe} \cdots \mathrm{Fe}$ contact bridged by two methoxides and one proline. The molecule has a six-fold symmetry axis, and the cage is planar making the non-crystallographic symmetry $S_{12}$.
Previously reported "ferric wheels" containing this bridging motif are either decanuclear and planar, ${ }^{1}$ with the $\mathrm{Fe} \cdots \mathrm{Fe} \cdots \mathrm{Fe}$ angle $c a .144^{\circ}$, or dodecanuclear and slightly puckered. ${ }^{4,5}$ Here the $\mathrm{Fe} \cdots \mathrm{Fe} \cdots \mathrm{Fe}$ angle is $146^{\circ}$ at $\mathrm{Fe}(1)$ and $154^{\circ}$ at $\mathrm{Fe}(2)$ - giving the average $150^{\circ}$ required for a planar dodecagon. The $\mathrm{Fe}-\mathrm{O}$ distances are normal: $\mathrm{Fe}-\mathrm{O}$ (methoxide), 1.974 to 2.009 (5); $\mathrm{Fe}-\mathrm{O}$ (proline), 2.042 to $2.081(5) \AA$, and bond angles are not unusual. Therefore the planar dodecagon is consistent with typical $\mathrm{Fe}-\mathrm{O}$ bond lengths and normal bond angles.
The high charge requires the presence of twelve perchlorate anions per wheel. These lie between the wheels, forming H-bonds from the oxygen atoms of the perchlorates to the protonated N atoms of the proline ligands (Fig. 2). As H-bonds are formed to wheels above and below each perchlorate the result is to form a H bonded stack of $\left\{\mathrm{Fe}_{12}\right\}$ wheels. The $\mathrm{O} \cdots \mathrm{N}$ distances in the H -bonds vary from 2.821 to $2.867(6) \AA$.

The magnetic behaviour§ of $\mathbf{1}$ shows the expected antiferromagnetic exchange between the Fe (III) centres. The room temperature value of $\chi_{\mathrm{m}} T$ is $c a .35 \mathrm{~cm}^{3} \mathrm{~K} \mathrm{~mol}^{-1}$ (where $\chi_{\mathrm{m}}$ is the molar magnetic susceptibility), which is lower than the calculated value for twelve Fe (III) centres $\left(52.5 \mathrm{~cm}^{3} \mathrm{~K} \mathrm{~mol}^{-1}\right.$ for $\left.g=2.0\right)$, suggesting a strong exchange interaction is operational. At low temperature there is evidence for the presence of a paramagnetic
impurity; the magnetic fit was therefore restricted to data measured above 20 K .
This behaviour has been modelled for a ring of twelve $S=5 / 2$ spins, derived from the method developed by Fisher for classical spins. ${ }^{10}$ The best fit of the data (Fig. 3) for the Hamiltonian $H=$ $-J . S_{i} S_{i+1}\left(i \leq 12, S_{13} \equiv S_{1}\right)$, gives $J=-15.8 \mathrm{~cm}^{-1}$, and $g=1.99$. We also calculated the exchange interaction using density functional theory (DFT)II. An exchange interaction was derived by DFT for a dimeric Fe (III) fragment containing the bridging unit found in 1, i.e. two methoxides and a proline, with terminal $\mathrm{H}_{2} \mathrm{O}$ molecules used to complete the Fe coordination spheres. This gave an exchange interaction of $-13.9 \mathrm{~cm}^{-1}$; this value was then used in a Monte Carlo simulation of the magnetic data with an excellent fit. It is very reassuring that the two methods of fitting the magnetic data reach a similar value.
This value of between -14 and $-16 \mathrm{~cm}^{-1}$ is more antiferromagnetic than the exchange interaction of $-9.6 \mathrm{~cm}^{-1}$ found for decanuclear ferric wheels, ${ }^{1}$ or that of $-10.9 \mathrm{~cm}^{-1}$ found in a $\left\{\mathrm{Fe}_{12}\right\}$ ring. ${ }^{4}$ Similarly in a $\left\{\mathrm{Fe}_{2}\right\}$ complex with two bridging MeO and a $\mathrm{O}_{2} \mathrm{CMe}$ bridge $J$ was found to be $-10.1 \mathrm{~cm}^{-1} .{ }^{13}$ Our value is in moderate agreement with proposed correlations between the exchange interaction and either the average ${ }^{14}$ or smaller $\mathrm{Fe}-\mathrm{O}-\mathrm{Fe}$ angle ${ }^{15}$ in di(alkoxo)-bridged Fe (III) complexes where there is no bridging carboxylate. Here these angles are 99.3 and $99.1^{\circ}$ respectively, which would give values of either -12.0 or -12.4


Fig. 1 The structure of the cation of $\mathbf{1}$ in the crystal. All unlabelled atoms are C-atoms.
$\mathrm{cm}^{-1}$ depending on the correlation used. The electronic character of the carboxylate may be causing this slight increase in the magnitude of the exchange due to a zwitterion being involved here,


Fig. 2 The H -bonding of $\mathrm{ClO}_{4}$ anions to $\left\{\mathrm{Fe}_{12}\right\}$ wheels in $\mathbf{1}$ viewed perpendicular to the plane of the wheels. Fe and N atoms and atoms in $\mathrm{ClO}_{4}$ units highlighted.


Fig. 3 The variable temperature behaviour of $\chi_{\mathrm{m}}(\triangle)$ and $\chi_{\mathrm{m}} \mathrm{T}(\bigcirc)$ for $\mathbf{1}$.
rather than the more conventional carboxylates used previously. The change is very slight and the predominant superexchange path probably remains through the methoxide ligands.

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## Notes and references

$\dagger$ Anal. Calcd (Found) for $\mathrm{C}_{84} \mathrm{H}_{180} \mathrm{Cl}_{12} \mathrm{Fe}_{12} \mathrm{O}_{96}$ : C, 25.29(24.96); H, 4.65 (4.52); N, 4.21 (4.12)\%.
$\ddagger$ Crystal data for $\mathrm{C}_{84} \mathrm{H}_{180} \mathrm{Fe}_{12} \mathrm{O}_{48} .12 \mathrm{ClO}_{4}, \mathbf{1}$ : hexagonal, $P 622$, $a=$ 25.0265(17), $c=7.7474(11) \AA, V=4202.3(7) \AA^{3}, M=3990, Z=1$ (the molecule lies on a six-fold axis and a two-fold axis), $T=150.0(2) \mathrm{K}, R 1=$ 0.0722. Data collection, structure solution and refinement used SHELXTL. 9 Full details have been deposited and will be published later. CCDC 222686. See http://www.rsc.org/suppdata/cc/b3/b312947k/ for crystallographic data in .cif or other electronic format..
§ Variable temperature magnetic measurements on $\mathbf{1}$ in the region 1.8-325 K were made using a SQUID magnetometer (Quantum Design) with samples sealed in gelatine capsules in a 100 G field. The data have been adjusted for the diamagnetism of the sample using Pascal's constants. The data were fitted to: $\chi_{\mathrm{m}} T=\chi_{\text {freeion }} T\left(1+\sum_{n=1,11}\left(2 u^{n}\right)+u^{12}\right)$ where $u$ is the Langevin function $J S(S+1) / T(J$ expressed in Kelvin).
If The DFT calculation used Gaussian 98, ${ }^{11}$ with the hybrid B3LYP functional together with Ahlrich's TZV basis set. This has been found to give good numerical estimates of $J$-values in previous work. ${ }^{12}$

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