Predicting the Future of Permanent-Magnet Materials

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There are two main thrusts towards new permanent-magnet materials: improving extrinsic properties by nanostructuring and intrinsic properties by atomic structuring. Theory—both numerical and analytical—plays an important role in this ambitious research. Our analysis of aligned hard-soft nanostructures shows that soft-in-hard geometries are better than hard-in-soft geometries and that embedded soft spheres are better than sandwiched soft layers. Concerning the choice of the hard phase, both a high magnetization and a high anisotropy are necessary. As an example of first-principle research, we consider interatomic Mn exchange in MnAl and find strongly ferromagnetic intralayer exchange, in spite of the small Mn-Mn distances.

Index Terms—Magnetic anisotropy, magnetization processes, permanent magnets.

I. INTRODUCTION

PERMANENT magnets [1]–[6] are used in an impressive range of applications, from electromotors and loud-speakers to windscreen wipers, locks, microphones and toy magnets. More recent key developments are applications in computer hard-disk drives, wind generators, and hybrid-car motors. The time of horseshoe magnets has long been over and permanent magnets made from steels are now obsolete, but the high saturation magnetization of Fe 33Co 67 and its pronounced temperature stability remain valued in alnico permanent magnets.

The first true hard-magnetic materials, the T l 10 compound CoPt [7] and the hexagonal ferrite BaFe 12O 19 [8] date back to the mid-20th century, followed by the high-performance rare-earth transition-metal permanent magnets Sm-Co [1], [9] and Nd 2Fe 14B [2], [10], [11]. This has enabled us to enhance the energy product by two orders of magnitude, from about 1 kJ/m 3 around 1900 to 460 kJ/m 3 in this millennium.

The performance of RE-TM intermetallics is difficult to beat, but rare-earth supplies have become a bottleneck in recent years, and there is active research in various directions. First, as emphasized by Skomski and Coey [12], the range of transition-metal-rich rare-earth intermetallics is limited, but improving the energy product to about 1000 kJ/m 3 could be possible by suitable nanostructuring [12], [13]. Second, a fundamental topic is the improvement of permanent-magnet materials, by changing chemical composition and atomic structure.

This paper focuses on the theoretical aspects of the first two questions: What are the optimum hard-soft geometries and phases, and what are the prospects for finding new permanent-magnet compounds?

II. SCIENTIFIC BACKGROUND

The physics of permanent magnets is based on the distinction between properties of atomic origin (intrinsic properties), and properties related to the magnets real structure (extrinsic properties), such as coercivity and energy product.

Intrinsic magnetic properties depend on crystal structure and chemical composition. The spontaneous or saturation magnetization M s and the Curie temperature T C of most magnets are largely determined by the iron-series (or 3d) transition-metal (TM) sublattice. Some of the heavier (4d, 5d, 4f) atoms also carry a magnetic moment m, but their magnetization M s = m/V is diluted by the larger atomic volume V of the heavy elements. The 3d magnetization is largely determined by the spin (S), which is known as the orbital-moment quenching.

Anisotropy means that the magnetic energy depends on the magnetization directions relative to the crystal axes. Simplifying somewhat, the anisotropic electrostatic crystal field modifies the orbital motion of the electrons and affects, via spin-orbit coupling, the spin system [14]. Since the spin-orbit coupling is a relativistic mechanism, the anisotropy is largest for heavy elements, such as rare earths. Most permanent-magnet alloys have uniaxial (hexagonal, tetrahedral or rhombohedral) crystal structures and the easy magnetization axis (c-axis) perpendicular to the basal plane (a-b-plane) [1]–[3].

Rare-earth moments are unquenched, and the spin-orbit coupling ensures that the 4f charge clouds, described by Stevens coefficients, are rigidly coupled to the spin [2], [3]. The corresponding rare-earth anisotropy energy is equal to the electrostatic interaction energy between the 4f ions and the anisotropic crystal field.

By contrast, there are no well-defined rules for the 3d, 4d, and 5d anisotropies as a function of the atomic number (or d-band filling) n. The anisotropy generally oscillates as a function of n, but these oscillations involve complicated Fermi-level-dependent k-space summations [15]. There are crude rules for nearly filled 3d bands [16], [17], but first-principle approaches, such as VASP calculations are much more reliable. Interestingly, such independent-electron calculations cannot be used to predict rare-earth anisotropy, because 4f electrons are highly correlated and the density functional for rare-earth 4f electrons is not known [18]. Methods such as LSDA + U reproduce the correct configuration or charge state, typically R 3+, but become successively less accurate for term, multiplet, and intramultiplet levels. For example, correlations lead to spin-charge separation [18], [19], and even if the charge state is well-known, little can generally be said about the spin direction.

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It is important to keep in mind that a good permanent magnet needs both a high coercivity and a high magnetization. On a Stoner-Wohlfarth-level, the coercivity \( H_c \) is determined by the anisotropy field

\[
H_A = 2K_1/\mu_0M_s
\]

The coercivity can therefore be enhanced by reducing the magnetization, and such magnets have been proposed as permanent magnets, most recently a Rh-substituted \( \varepsilon \)-Fe\(_2\)O\(_3\) [20]. However, the energy product is quadratic in the magnetization, and magnets with a low magnetization cannot be used as permanent magnets, even if the coercivity is very high.

*Extrinsic* properties are usually realized on a scale of several nanometers and on relatively long time-scales, as epitomized by the nonequilibrium character of magnetic hysteresis [21]. There is the general rule that coercivities reach only about 10% to 30% of the anisotropy field (Kronmüller factor \( \alpha = H_c/H_A \)), depending on microstructure. Fig. 1 shows two key mechanisms of coercivity, namely nucleation and pinning. Nucleation-controlled coercivity aims at removing nuclei or "soft spots" and typically implies nearly perfect magnets, without harmful bulk and surface defects. A second mechanism is pinning, where domain walls exist but cannot move due to real-structure imperfections.

Let us consider the nucleation in a soft-phase region and assume that the field \( \mathbf{H} = H_e\mathbf{e}_e \) is parallel to the \( e \)-axis. Starting from saturation, \( \mathbf{M} = M_s\mathbf{e}_s \), nucleation involves a small magnetization component \( \mathbf{M} - M_s\mathbf{e}_e = \mathbf{m} \) [21], [22]. The components \( m_x \) and \( m_y \) of \( \mathbf{m} \) are approximately degenerate, so that we can restrict ourselves to the consideration of \( m = |\mathbf{m}| \) [21]. The analysis starts from the micromagnetic free energy and yields

\[
\mathcal{A} \nabla^2 m + (K(r) - \mu_0M_sH/2) m = 0.
\]

Here the effective lowest-order uniaxial anisotropy constant \( K(r) \) describes the real or defect structure of the magnet and \( \mathcal{A} \) is the exchange stiffness. \( K \) also contains the magnetostatic self-interaction in a rather crude approximation [21]. For homogeneous ellipsoids of revolution, \( K(r) = (K) \), and (2) reproduces the coherent-rotation result of (1).

**III. NANOSTRUCTURED PERMANENT MAGNETS**

Aligned hard-soft nanocomposites continue to be an active research area [23], [24], [26], [27], in spite of the challenging processing requirements. In these structures, the soft-phase material improves the hard-magnetic performance the main phase, sacrificing some anisotropy and coercivity but enhancing magnetization and energy product beyond that of the hard phase [12], [25]. The approach was initially outlined by Kneller and Hawig, who advocated exchange-spring multilayers [13]. Skomski and Coey [12] considered three-dimensional nanostructures and predicted rather accurate upper energy-product limits of about 1000 kJ/m\(^3\). These papers have several precursors, such as earlier work on exchange-spring multilayers [28], [29] and on nucleation fields in inhomogeneous materials [12], [27]–[31]. Actual energy products are always lower than the upper limit, because real-structure imperfections reduce the coercivity [12], [32]. For example, Fig. 2 in [12] shows how the coercivity decreases as the soft regions get bigger. Note that the calculations in [12] and in the present paper assume perfect exchange coupling at the interface. As analyzed in [21], moderately reduced interface exchange has only a very small effect on the interface exchange.

There exist experimental proofs of principle with high magnetization and coercivity in thin films [25], [26], [33]–[36], but the challenge remains to maintain bulk coercivity in the presence of the soft phase. It is well-established that the soft phase of a two-phase system cannot be much larger than twice the Bloch-wall width \( \pi \sqrt{A/K_h} \) of the hard phase, irrespective of dimensionality. Otherwise, it was believed that multilayers [13] and spherical inclusions [12], [31] behave similarly, in spite of the more pronounced micromagnetic localization behavior of one-dimensional structures [21]. The specific involvement of \( \sqrt{A/K_h} \) is indeed a cornerstone of hard-soft exchange coupling, but there are also differences, both quantitatively and qualitatively.

**A. Volume Fraction of the Soft Phase**

How do coercivity and energy product vary as a function of the soft-phase content? In the worst case, a trivially small addition of the soft phase completely ruins the coercivity by creating a harmful nucleus, and the energy product collapses. There are, in fact, thousands of ways to destroy coercivity and energy product. In this subsection, we study two examples of such deterioration mechanisms. By carefully controlling the size and distribution the soft-phase regions, it is possible to limit the coercivity decrease to \( H_c = H_{bh}(1 - f) \), where \( H_{bh} \) is the coercivity of the hard phase and \( f \) is the volume fraction of the soft phase. This corresponds to the limit of perfect exchange coupling [12], where \( K_{R} = K(r) \). Let us assume that the hysteresis loop is rectangular and the hard-phase coercivity \( H_{bh} = \alpha H_A \). The exchange-coupled soft phase enhances the remanence to \( M_r = M_{bh} + f(M_r - M_{bh}) \) but reduces the coercivity. For rectangular loops with \( H_c < M_r/2 \), the energy product \( (BH)_{\text{max}} = \mu_0H_c(M_r - H_c) \), whereas for \( M_r > H_c/2 \), \( (BH)_{\text{max}} = \mu_0M_r^2/4 \). "Poor" hysteresis loops having a straight line from \( M(-H_c) = 0 \) to \( M(0) = M_r \) yield an energy product of \( 1/4\mu_0M_r^2/2(1 + M_r/H_c) \). Fig. 2 shows these energy products for different values of \( \alpha \).

The maxima in Fig. 2(a) depend on the choice of the hard component and are equal to \( 1/4\mu_0M_r^2(1 + (M_r - M_{bh})/\alpha H_A) \). This means that the ratio \( (M_r - M_{bh})/\alpha H_A = (M_r - M_{bh})/K_{bh} [35] \) must be small to achieve high energy products, and there is no point in using semihard materials with relatively small magnetizations and coercivities as hard phases. In fact, the addition of the soft phase *deteriorates* the energy product if the hard-phase coercivity \( \alpha H_A \) is lower than \( M_{bh} - M_r/2 \).
B. Effect of Geometry

Equation (2) has been solved for a variety of geometries [12], [30], [37]. Fig. 3 summarizes the geometries included in the present analysis, namely spheres, embedded plates, cylinders, and thin films on a hard surface. The nucleation field is obtained by starting from (2) and exploiting the boundary conditions at free and clamped surfaces [12], [31], [37].

Fig. 4 shows the nucleation field as a function of soft-phase dimension (radius or layer thickness), assuming a “proper” exchange length $l_s$ of $\sqrt{A/\mu_0 M^2_s} = 2$ nm, $K_1 = 5$ MJ/m$^2$ and $\mu_0 M_h = 1.5$ T. For large soft regions ($R \rightarrow \infty$), the nucleation mode $m(\tau)$ is confined to the center of the soft inclusion. The corresponding magnetization profiles are $\phi(z) = \phi_0 \cos(\pi z/t)$ for plate-like soft inclusions of thickness $t$, $\phi_0 J_0(r/R)$ for cylindrical inclusions of radius $R$, and $2 R \phi_0 \sin(\pi r/2R)/\pi r$ for spherical inclusions of radius $R$. Explicitly, the ratio $H_n/M_s$ is equal to 19.74 $l_s^2/H^2$ (spheres), 11.57 $l_s^2/H^2$ (cylinders), 19.74 $l_s^2/t^2$ (embedded plates), and 4.94 $l_s^2/t^2$ (soft layer with free surface).

In other words, we see that embedded spheres, Fig. 3(a), have the highest coercivity, corresponding to a factor 4 in the nucleation field compared to embedded layers (c) having $t = 2R$. The behavior of embedded soft cylinders is intermediate between spheres and layers. The nucleation fields of soft-magnetic layers with one free surface, Fig. 3(d), have an additional factor 1/4 compared to the embedded layer, or a factor 1/16 compared to embedded spheres.

For very small soft inclusions, the nucleation field approaches the anisotropy field $H_A$ of the hard phase, but due to localization, the approach to the "plateau" $H_n = H_A$ is dimensionality-dependent. This is in close analogy to the quantum-mechanical delocalization of electrons in an inhomogeneous potential $V(r)$ [21] and to the behavior of impurity states in the band gaps of solids of different dimensionalities [38]. Nucleation modes in one-dimensional systems (layers) localize most easily and exhibit a pronounced nucleation-field drop, whereas three-dimensional (spherical) inclusions need a minimum size to be effective, $R = 4$ nm in Fig. 4. Two-dimensional systems (cylinders) form a borderline case, with logarithmically weak localization corrections [38], [39].

C. Other Micromagnetic Approaches

Let us consider alnicos, where elongated soft regions of FeCo, Fig. 5(a), are embedded in a nonmagnetic NiAl matrix [4], both phases crystallizing in the cubic CsCl (B2) structure. The magnetic anisotropy of alnicos is of the shape-anisotropy type, caused by magnetostatic charges at the wires’ ends. Since the energy product $(BH)_{\text{max}}$ never exceeds $\mu_0 M_s^2/4$, where $M_s = \int M_s$, it is advantageous to enhance the volume fraction of the magnetic FeCo phase, the dark area in Fig. 4(a). However, this reduces the coercivity, and the interplay between magnetization increase and coercivity decrease yields a volume fraction $f = 2/3$ at which the energy product reaches a maximum. For highly idealized alnico, this analysis yields $(BH)_{\text{max}} = \mu_0 M_s^2/12$ [40].

What would happen if there was some anisotropy associated with the predominant (001)-type interfaces between the main (FeCo) and matrix (NiAl) phases? Surface and interface anisotropies $K_s$ have values of up to 1 mJ/m$^2$ [1 erg/cm$^2$]. This corresponds to a bulk anisotropy of 5 MJ/m$^2$ (or roughly 0.35 meV per atom), roughly equal to the maximum anisotropy observed in a 3d system (YCo$_5$). The physics behind this similarity is the crystal structure of YCo$_5$, which consists of alternating Co and Y-Co layers, each alternation giving rise to a large ‘interface’ anisotropy.
An approximate energy product is obtained from (3), by using $H_c = 1/2(1 - f)M_s + H_{an}$, where $H_{an}$ is the additional anisotropy-contribution due to the interface anisotropy. If $H_{an}$ is independent of $f$, then the derivative $dH_c/df$ remains unchanged and the energy product is maximized for $H_c = M_s/4$, irrespective of $H_{an}$. The optimum packing fraction $f = 2(1 + 2\Delta H_c/M_s)/3$, yields the dashed line in Fig. 5(c).

The net surface anisotropy, obtained by adding the inequivalent surface contributions $A$ and $B$ in Fig. 5(b), yields an anisotropy-field change of $\Delta H_a = -4K_s/\mu_m M_sw$, where $w$ is the width. Negative interface anisotropy, where the easy axis is parallel to the wires, moves the peak in Fig. 5(b) towards higher volume fractions and higher energy-product values. For a long wire having a square cross section ($w = 20$ nm) and a (001)-type interface, an interface anisotropy of magnitude 1 mJ/m$^2$ translates into substantial anisotropy-field change of up to 0.1 T [1 kOe], comparable to the coercivity of alnico. This is important, because the small coercivity is the bottleneck for the use of an otherwise good permanent-magnet material. Note, however, that the change may have either sign, depending on the sign of $K_s$.

IV. HARD-MAGNETIC COMPOUNDS

The search for new intermetallic compounds has been a long-standing task in magnetism, and the rare-earth intermetallics used at present are highly sophisticated and difficult-to-improve alloys. However, exploring new and modified magnetic phases—such as specialty magnets, including thin-film magnets, and rare-earth-free magnets with energy products between alnico and Nd-Fe-B—remains an important research direction and has attracted renewed interest.

This section discusses some first-principle results by the present authors and by other groups. In our own work, we used a supercell approach, the frozen-core full-potential projected augmented wave (PAW) method as implemented in the Vienna ab-initio simulation package (VASP) [41], and a generalized-gradient approximation (GGA) [41], [42].

A. Iron- and Cobalt-Base Magnets

Iron- and cobalt-based metallic magnets play an important role in magnetism, largely because their magnetization is as high as 2.43 T in Fe$_{63}$Co$_{37}$. Substantial anisotropy, $K_1 = 9.5$ MJ/m$^3$, and a magnetization of $\mu_M = 1.9$ T have been predicted for strongly distorted Fe-Co with $c/a = 1.23$ [43]. Experimental room-temperature anisotropies per strained Fe or Co atom reach about 2.1 MJ/m$^3$, but this value does not account for the large amount of Pt (about 75 vol.%) [44]. In fact, a similar mechanism is responsible for the coercivity of steel magnets, where interstitial carbon yields a tetragonal martensitic lattice distortion [5], and both magnetoelastic and chemical effects contribute to the net anisotropy [45]. Other candidates for nonrare-earth permanent magnets are Zr$_2$Co$_{11}$ [46] and orthorhombic HfCo$_7$ [47], which have magnetizations comparable to Sm-Co, for example 1.09 T for HfCo$_7$. A problem is that the addition of 4$d$ and 5$d$ atoms, such as W, often deteriorates the magnetization [48].

An interesting noncubic high-magnetization phase is Fe$_{11}$N$_2$ [49], where the nitrogen enhances the magnetization by expanding the Fe lattice. The effect on the electronic structure of the Fe is larger than the dilution of the magnetic moment, although magnetization values of 3.0 T have not been convincingly verified [49], and the structures are difficult to prepare.

B. L1$_0$-Ordered Magnets

L1$_0$-ordered alloys such as CoPt, FePt and FePd [7], [5], [50], have long been known in permanent magnetism, but their use has been limited by the price of Pt and Pd. There are a few ferromagnetic Mn alloys with modest magnetizations of less than one tesla, such as MnBi and MnAl [3], [5]. L1$_0$-ordered FeNi (tetrataenite) was originally discovered in meteorites, formed with cooling times in excess of one million years, but is now being explored from the viewpoint of permanent magnetism [51]. Fig. 6 shows the most general L1$_0$ structure, which has the composition AHC$_2$ [52].

In Fe-rich noncubic Pt alloys, a moderate reduction of raw-materials costs can be achieved by substituting Fe and Co for Pt. For example, recently produced Fe-Co-Pt thin-film magnets have a thickness of 20 nm and room-temperature properties of up to $\mu_m H_c = 2.52$ T and $\mu_m M_s = 1.78$ T [26], [35]. In these structures, the compromise between magnetization and coercivity yields a nominal energy product maximum of 510 kJ/m$^3$ for Fe$_{30}$Co$_{23}$Pt$_{33}$.

To understand the magnetic moments in Fe$_{1-x}$Co$_x$Pt, we have performed supercell calculations on various chemically ordered and disordered L1$_0$ derivates [42]. The ordered alloys have the nominal composition Fe$_5$Co$_8$Pt$_{16}$, but differ by their stackings, both vertically and laterally. The disordered alloys are approximated by supercells with 32 atoms and have the nominal compositions Fe$_{15}$Co$_{15}$Pt$_{16}$ ($x = 6.25\%$) and Fe$_{14}$Co$_2$Pt$_{16}$ ($x = 12.5\%$). In Fe$_{15}$Co$_{15}$Pt$_{16}$, all Co sites are
equivalent, but in the Fe\textsubscript{14}Co\textsubscript{2}Pt\textsubscript{16} supercell, there are 7 crystallographically nonequivalent configurations—each of these configurations requires a separate first-principle calculation, followed by configurational averaging.

Both our supercell [42] and previous CPA [53] calculations indicate that the average moment decreases with increasing Co content. This is very different from bcc Fe\textsubscript{1−x}Co\textsubscript{x}, where the moment initially increases with \( x \), from 2.2 \( \mu_B \) (Fe) to 2.4 \( \mu_B \) (Fe\textsubscript{0.5}Co\textsubscript{0.5}). In Fe\textsubscript{4}Co\textsubscript{6}Pt\textsubscript{16}, the effective is Fe-Co moment is 2.42 \( \mu_B \) [42], [54], corresponding to strong 3-D ferromagnetism. Note that perfectly ordered FePt has been predicted to be antiferromagnetic [55]. Our calculations yield a very weak ferromagnetic exchange, 0.005 eV per atom, and the FM-AFM transition is within the margin of error.

**C. Manganese-Based Magnets**

Tripole manganese has a moment of 5 \( \mu_B \) per atom, as compared to 2.2 \( \mu_B \) for Fe. If this moment could be exploited in industrial magnets, it would revolutionize technology far beyond magnetism and open the door for completely new technologies. Furthermore, Mn is a relatively cheap metal, which is of utmost importance in the light of the tight supply and high costs of rare earths. Unfortunately, most manganese compounds are antiferromagnetic, which is typical for elements in the middle of the 3d series. \( L_1_0 \)-ordered (or \( \tau \)-phase) MnAl requires small C additions to become structurally stable [5], but exhibits appreciable intrinsic properties, namely \( \mu_0M_s = 0.75 \) T, \( K_1 = 1.7 \) MJ/m\(^3\) and \( T_c = 650 \) K [3], [50]. Our focus is on exchange interactions between Mn atoms, which largely determine the magnetization and Curie temperature of Mn alloys.

The Slater–Bethe–Néel curve suggests that ferromagnetic interatomic exchange in Mn alloys requires large interatomic distances, with the effect of strongly diluting the magnetization. However, the Slater–Bethe–Néel curve is a crude approximation, and our VASP calculations show that strong ferromagnetic (FM) exchange can exist even for very short Mn-Mn distances. In \( L_1_0 \)-ordered MnAl, the dense-packed Mn sheets exhibit an FM \textit{intralayer} exchange \( J \), whereas the interlayer exchange \( J' \), with enhanced Mn-Mn distances due to the separating Al, is only very weakly ferromagnetic. For the \( L_1_0 \) MnAl alloys, we obtain Mn and Al moments of 2.420 \( \mu_B \) and −0.061 \( \mu_B \), respectively. This means that the coupling between the Mn and Al sublattices is antiferromagnetic.

It is interesting to see how properties change on Fe addition. Fig. 7 shows the spin-resolved density of states (DOS) for \( L_1_0 \) MnFeAl\textsubscript{2}, which crystallizes in the structure of Fig. 6(b). The Fe reduces the net moment, with Fe, Mn and Al atomic moments of 2.461 \( \mu_B \), 1.899 \( \mu_B \), and −0.656 \( \mu_B \), respectively. On the other hand, the Fe substitution enhances the anisotropy constant from 1.77 MJ/m\(^3\) in MnAl to 2.50 MJ/m\(^3\) in MnFeAl\textsubscript{2}.

For MnAl, our first-principle calculations yield \( J = 502 \) K and \( J' = 75 \) K, and an approximate Curie-temperature of 718 K, as compared to the experimental value of about 650 K. For hypothetical \( L_1_1 \)-ordered FeMnAl\textsubscript{2}, the Fe-Mn interlayer coupling is very strong, \( J = 593 \) K, but the Mn-Mn intraplane exchange is negative, \( J = −185 \) K, and the spin structure with the lowest energy is antiferromagnetic.

**V. Conclusion**

In summary, we have investigated several aspects of \( c \)-axis-aligned hard-soft nanostructures and of permanent-magnet alloys. Concerning the hard-soft geometry, there are two rules. First, soft-in-hard geometries are better than hard-in-soft geometries. Second, among the favorable hard-in-soft geometries, embedded soft spheres are best. Embedded layers are worst, whereas embedded cylinders are intermediate. For small feature sizes, the differences are a micromagnetic analog to the dimensionality-dependent localization behavior of impurity states in the electronic band gaps. The hard-phase materials best suited for hard-soft nanostructuring have both a high magnetization and a high anisotropy.

Our first-principle calculations on MnAl show that a strongly ferromagnetic exchange can exist for very short Mn-Mn distances. This is encouraging and calls for future research. Iron addition enhances the anisotropy, from 1.77 MJ/m\(^3\) to 2.50 MJ/m\(^3\) for FeMnAl\textsubscript{2}, but reduces the magnetization by creating antiferromagnetic bonds.

So, let us return to our starting point. Can theory really predict the future of permanent magnetism? Probably not. Can it guide experimentalists to go in meaningful directions and to avoid pitfalls? Yes.

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