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# Hidden orders in spin-orbit entangled correlated insulators

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## ABSTRACT

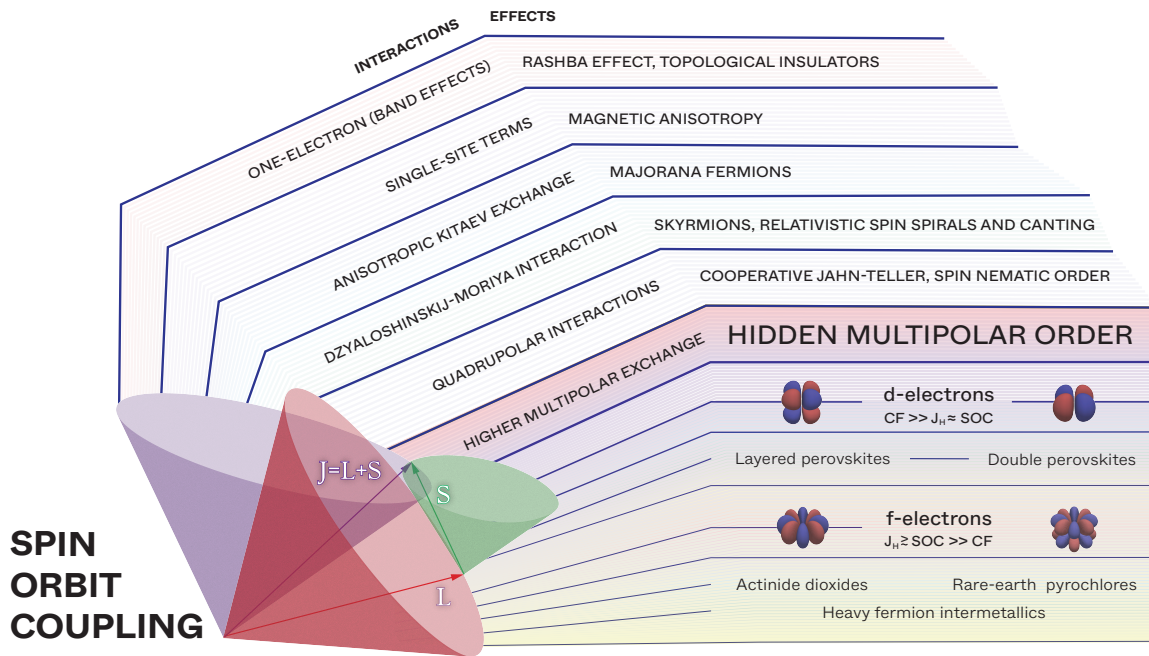
In many materials, ordered phases and their order parameters are easily characterized by standard experimental methods. "Hidden order" refers to a phase transition in which an ordered state emerges without such an easily detectable order parameter, despite clear thermodynamic evidence of the transition. The underlying mechanisms for these unconventional states of matter stem from spin-orbit coupling, which intertwines inter-site exchange, classical electromagnetic interactions, and electron-lattice effects. This physics is elusive to experimental probes and beyond traditional theories of insulating magnetism, requiring sophisticated methodologies for its exploration. In this Review, we survey exotic hidden-order phases in correlated insulators, particularly focusing on the latest progress in material-specific theories and numerical approaches. The relevant degrees of freedom in these phases are local high-rank multipole moments of magnetic and charge density that emerge from spin-orbit entangled correlated shells of heavy  $d$  and  $f$  electron ions and interact on the lattice via various mechanisms. We discuss approaches to modelling hidden orders in realistic systems via direct ab initio calculations or by constructing low-energy many-body effective Hamiltonian. We also describe how these new theoretical tools have helped to uncover driving mechanisms for recently discovered multipolar phases in double perovskites of heavy transition metals, and how they have proved instrumental in disentangling the role of various interactions in "traditional"  $f$ -electron multipolar materials like actinide dioxides. In both cases, material-specific theories have played a key role in interpreting and predicting experimental signatures of hidden orders.

## [H1] Introduction

Spin-orbit coupling (SOC) and electronic correlation are two fundamental interactions that substantially influence the properties of many-body electronic systems. SOC in solids is a relativistic interaction that entangles the electron spin with its orbital angular momentum in a crystal field environment. SOC is strongly enhanced for elements with a large atomic number<sup>1,2</sup> and is primarily responsible for effects such as Rashba coupling<sup>3,4</sup>, the spin Hall effect<sup>5,6</sup>, and topological spin textures such as skyrmions<sup>7</sup>. Electronic correlation is the repulsive interaction between electrons, which is particularly intense in strongly correlated materials with localized electronic states, typically partially filled  $3d$  orbitals, and leads to phenomena such as Mott insulating behavior and quantum criticality.<sup>8-11</sup>

The synergistic interplay of large SOC and strong electronic correlation can give rise to new phenomena distinct from conventional electronic and magnetic states,<sup>12-18</sup> such as spin-orbit coupled Mott insulators<sup>19</sup>, insulator-to-metal transitions driven by the combined action of SOC and magnetism<sup>20,21</sup>, exotic magnetic states such as quantum spin liquids<sup>22</sup>, valence-bond glasses<sup>23</sup>, high-rank multipolar orders<sup>24</sup>, skyrmions<sup>25,26</sup>, non-collinear Dzyaloshinskii-Moriya interactions<sup>27-29</sup>, non-trivial topological Mott systems<sup>30</sup> and Rashba effects<sup>4</sup>. These effects are typically observed in systems containing heavy  $5d$  transition metals or heavy  $4f$ - $5f$  elements, which constitute an extensively studied class of materials and still represent a frontier in condensed matter research.<sup>31-36</sup> A compendium of the

most relevant phases is provided in **Table 1** and pictorially represented in **Fig. 1** together with the one-body and many-body interactions that are at their origin.



**Figure 1. Spin-orbit entangled phenomena.** Interplay of spin-orbit coupling with band and correlation effects induces unusual interactions (left side), which, in turn, generate unconventional phenomena (right side), including hidden multipolar orders. Characteristic energy scales and key families of materials hosting hidden orders are listed for *d*- and *f*-electron systems. Representative multipolar order parameters are shown for both classes of materials: quadrupolar ( $K = 2$ ) and octupolar ( $K = 3$ ) moments for *d* systems and hexadecapolar ( $K = 4$ ) and triakontadipolar ( $K = 5$ ) moments for *f* systems.

A common characteristic of spin-orbit entangled correlated insulators is the formation of unconventional spin-orbital phases, often developing in geometrically frustrated lattices. In conventional magnetic materials, a phase transition to a well-defined pattern of local magnetic moments is typically associated with an order parameter detectable experimentally by both thermodynamic and spectroscopic probes such as magnetic susceptibility, specific heat, and neutron scattering measurements. In contrast, in a correlated spin-orbit coupled background, the strong entanglement of spin and orbital moments can lead to spontaneous symmetry breaking driven by a complex order parameter that, though clearly observed in thermodynamic probes, is challenging to characterize by standard magnetic measurements and neutron spectroscopy. The resulting order is typically referred to as ‘hidden’, a term introduced in the early 2000s for the unknown ordered state observed in the heavy-fermion metal  $\text{URu}_2\text{Si}_2$ <sup>37–39</sup>. Since then, hidden-order phases (**Box 1**) have been reported for different types of correlated insulators with localized and partially filled *f* and *d* spin-orbital manifolds.<sup>40–71</sup>

In conventional magnetic phase transitions, the order parameter is associated with distinct ordering patterns of the electronic spin dipole moments in a crystal. These emergent orders are typically induced by a Heisenberg-type effective Hamiltonian describing spin-spin interactions, and they can be investigated by ab initio electronic structure theory using density functional theory (DFT) and dynamical mean-field theory (DMFT) techniques. This well-established approach facilitates a quantitative comparison of computed intersite dipolar interactions with experimental data, allowing for a material-specific analysis.<sup>72</sup> In spin-orbit entangled correlated materials, the interplay of spin and unquenched orbital degrees of freedom may lead to the activation of higher-rank multipole

interactions, which can be responsible for the onset of hidden orders<sup>73</sup>. Compared to dipolar couplings, evaluating multipolar intersite interactions is more complex due to the necessity of accounting for tensorial operators in the extended multipolar Hamiltonian. Equally challenging is the extraction of high-rank coupling from experimental measurements, which are often "blind" to multipolar hidden orders.<sup>74–79</sup> Nevertheless, pioneering experiments have provided insights in deciphering the microscopic nature of hidden orders using different techniques: X-ray magnetic circular dichroism (XMCD) has been employed to inspect the ferroic order of the magnetic octupole in  $\text{Mn}_3\text{Sn}$ <sup>80</sup>, and a variety of tools including neutron and X-ray diffraction (XRD), muon spin resonance ( $\mu\text{SR}$ ), inelastic neutron scattering, resonant and non-resonant elastic X-ray scattering (REXS) and nuclear magnetic resonance (NMR) were used to elucidate the octupolar order in Os-based double perovskites<sup>35,52,63</sup> and  $\text{UPd}_3$ <sup>81</sup> as well as the rank-5 (trikontadipolar, the name referring to 32 lobes of this multipole) order in  $\text{NpO}_2$ <sup>82,83</sup>.

Unraveling and understanding hidden orders is challenging, demanding advanced theories and computational methodologies. These efforts can uncover novel quantum states of matter, advancing the boundaries of materials research.

In this Review, we provide an overview of recent *ab initio* quantitative approaches for studying the complex hidden multipolar orders in correlated insulators. We detail how these methods can offer essential insights into material-specific properties, which are crucial for a microscopic understanding of non-conventional order parameters. We begin by reviewing the fundamental theoretical aspects of multiplet physics and the formalism necessary to construct low-energy microscopic Hamiltonians. Next, we present computational approaches to derive these Hamiltonians from first principles, focusing specifically on multipolar DFT and DMFT frameworks. We then demonstrate how these theories have clarified the role of multipolar physics in two main classes of materials: "traditional" *f*-electron Mott insulators<sup>73</sup> and *5d* double perovskites<sup>84,85</sup>. Very complex low-energy Hamiltonians involving couplings between high-rank multipolar moments have been fully derived from first principles for prototypical "hidden-order" *f*-electron systems<sup>34,67,86</sup>. The double perovskite family has recently garnered significant attention as a rich playground for discovering novel high-rank multipolar quantum phases and unraveling the coupling of hidden orders with phonons<sup>33,87</sup> and vibronic effects<sup>35,88–90</sup> and the role of charged defects, polarons and doping<sup>31</sup>, providing knowledge that can be transferable to other quantum materials. Although this Review focuses primarily on theoretical and computational aspects, we discuss direct comparisons with experimental observations of multipolar orders as a necessary validation for the proposed numerical protocols.

We note that ideas and symmetry organization principles similar to those discussed in this Review also apply to materials with exotic magnetic orders such as toroidal order<sup>91</sup> or the recently proposed altermagnetic order, which can occur in systems with larger unit cells<sup>92,93</sup>. In such materials, while the net magnetic dipole moment within the unit cell integrates to zero, one can nevertheless find emergent higher-rank multipoles constructed out of the local dipole moments. These include quadrupolar symmetry breaking, which is akin to nematic order, time-reversal breaking octupolar order, which may be viewed as a *d*-wave ferromagnet, or even multipolar variants of multiferroic materials.<sup>94–97</sup>

## [H1] Theories and methods

This section provides an overview of theoretical approaches to multipolar interactions and orders in realistic materials. We start by briefly introducing the concepts of multipolar moments (and the corresponding operators) acting on a localized atomic shell in a correlated insulator. Various mechanisms providing coupling between multipoles at different sites are also briefly reviewed together with model approaches designed to derive those couplings from tight-binding hopping parameters.

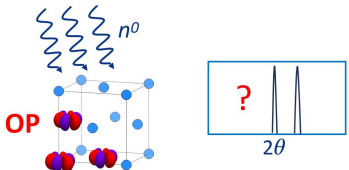
We then focus on material-specific theoretical tools for modeling multipolar phases and extracting experimental signatures of multipolar moments. *Ab initio* total-energy DFT-based methods are extensively used to study conventional magnetic orders<sup>146</sup>. Generalizations of DFT functionals in conjunction with the  $+U$  correction have been elaborated to treat multipolar orders. An alternative route is to construct, from DFT or its combination with DMFT, a many-body effective Hamiltonian (MBEH) encoding low-energy multipolar physics. We review newly developed "force-theorem" approaches to correlated insulators that derive the MBEH from linear-response

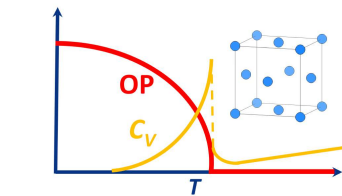
Phases of matter and exotic effects in spin-orbit entangled correlated materials		
Year of Report	Type of phase	Materials
1958 <sup>98</sup> , 1960 <sup>28,99</sup>	Dzyaloshinskii-Moriya interaction	Fe <sub>2</sub> O <sub>3</sub> , SrFeO <sub>3</sub> <sup>100</sup> Sr <sub>2</sub> IrO <sub>4</sub> <sup>101</sup> , polar and frustrated magnets <sup>102</sup> , multiferroics <sup>103,104</sup> , 2D magnets <sup>105</sup> , 2D Van der Waals <sup>106</sup>
		Reviews <sup>102,104,105,107,108,108-110</sup>
1970 <sup>24</sup> ,1996 <sup>111</sup>	Multipolar (hidden) order	NpO <sub>2</sub> , UO <sub>2</sub> <sup>24</sup> (actinide dioxides <sup>73,82</sup> ), CeB <sub>6</sub> <sup>112,113</sup> , URu <sub>2</sub> Si <sub>2</sub> <sup>37,44,111</sup> , R <sub>3</sub> Pd <sub>20</sub> X <sub>6</sub> (R=Ce, Pr: X=Si, Ge) <sup>114</sup> , PrX <sub>3</sub> (X=Pb,Mg) <sup>115,116</sup> , Sr <sub>2</sub> VO <sub>4</sub> <sup>41,47</sup> , Ba <sub>2</sub> MOsO <sub>6</sub> <sup>42,63,66</sup> , Sr <sub>2</sub> RuO <sub>4</sub> <sup>87,117</sup> , Sr <sub>2</sub> IrO <sub>4</sub> <sup>49</sup> , Rare-earth pyrochlores <sup>36,48,59</sup> , further heavy fermions <sup>118</sup> , Mn <sub>3</sub> Sn <sup>80</sup> , magnetoelectric oxides (Ca <sub>3</sub> Ru <sub>2</sub> O <sub>7</sub> <sup>119</sup> , Cr <sub>2</sub> O <sub>3</sub> & Fe <sub>2</sub> O <sub>3</sub> <sup>71</sup> )
		Reviews <sup>16,59,61,73,120,121</sup>
1973 <sup>122</sup>	Quantum spin liquid	Candidates <sup>123</sup> : Triangular organics <sup>124,125</sup> , Kitaev materials <sup>19,126</sup> , Rare-earth pyrochlores <sup>43</sup> , Kagomé Herbertsmithite <sup>127</sup>
		Reviews <sup>59,123,125,128,129</sup>
2009 <sup>19</sup>	Spin-orbit coupled Mott insulator	Sr <sub>2</sub> IrO <sub>4</sub> (Layered) <sup>19,130</sup> , Na <sub>2</sub> IrO <sub>3</sub> (Honeycomb) <sup>131</sup> , BaIrO <sub>3</sub> <sup>132,133</sup> , NaOsO <sub>3</sub> <sup>20,21</sup> , other 4d and 5d oxides <sup>134</sup>
		Reviews <sup>12,14,16,85,134</sup>
2010 <sup>135</sup>	Non-trivial topological phases	SbB <sub>6</sub> <sup>135</sup> , SrIrO <sub>3</sub> <sup>136</sup> , Ce <sub>3</sub> Bi <sub>4</sub> Pd <sub>3</sub> <sup>137</sup> , Ce <sub>2</sub> Au <sub>3</sub> In <sub>5</sub> <sup>138</sup>
		Reviews <sup>30</sup>
2010 <sup>3,139</sup>	Rashba effects in correlated systems	SrTiO <sub>3</sub> <sup>139</sup> , KTaO <sub>3</sub> <sup>140</sup> , SrNbO <sub>3</sub> <sup>141</sup> , RETM <sub>2</sub> Ir <sub>2</sub> Si <sub>2</sub> <sup>4,142,143</sup> (RE=rare earth; TM=transition metal)
		Reviews <sup>4,144,145</sup>

**Table 1. Phases and behaviours originating from the coexistence of spin-orbit coupling and electronic correlation. Historical overview and exemplary materials.**

of the DFT (+U/+DMFT) total energy functional, as well as another DFT+DMFT-based approach that extracts the multipolar MBEH from the dynamical susceptibility. A different class of methods – cluster MBEH derivation utilizing quantum-chemistry or exact diagonalization methods within a strong-coupling perturbative perspective – is considered afterwards. Theoretical approaches to electron-lattice coupling and the Jahn-Teller effect in multipolar systems are also briefly reviewed, as well as methods for solving the MBEH and for modeling experimental responses of multipolar orders.

## Box 1. Hidden order

- **Ordered phases and order parameters.** Second-order phase transitions in solids involve the emergence of an ordered phase at a specific temperature, with a symmetry lower than that of the high-T disordered phase. This low-T ordered phase is characterized by a non-zero of a certain physical observable, known as the *order parameter* (OP), which gradually vanishes as the temperature approaches the transition point, as shown in the schematics. The order parameters can be usually directly measured due to their interactions with either external probes, e. g., neutron diffraction can detect superstructural peaks due to an antiferromagnetic OP.The diagram shows a lattice of atoms with red and blue arrows representing magnetic moments. A wavy line labeled  $n^0$  is above the lattice. To the right, a diffraction pattern shows several sharp peaks on a  $2\theta$  axis.
- **Hidden-order phases.** In certain cases, experimental observations reveal clear thermodynamic signatures of a phase transition, such as the  $\lambda$ -type anomaly in specific heat  $C_V$  shown in the schematics, without any detectable spontaneous symmetry breaking. These scenarios pose long-standing challenges in solid-state physics and are referred to as *hidden order* transitions.The diagram shows a lattice of atoms with red and blue spheres. A wavy line labeled  $n^0$  is above the lattice. To the right, a diffraction pattern shows a question mark and two sharp peaks on a  $2\theta$  axis.
- **Multipolar orders.** In a common class of hidden orders, the OPs are high-rank local-moment degrees of freedom (multipole moments). These systems are classified by the rank of their multipolar order parameters and are typically parity-even, with the exception of magnetic and electric toroidal moments, which are parity-odd (see Supplemental Material):
  - *Quadrupolar orders* The simplest case is the ordering of quadrupoles (rank 2), which are, in a quantum treatment, represented by symmetrized quadratic polynomials of angular momenta. Since quadrupolar moments interact directly with the lattice and are a source of magnetic anisotropy, quadrupolar orders have been characterized in various materials by studying anisotropic susceptibility and magnetoelastic effects.
  - *High-rank multipolar orders.* Octupolar (rank 3) and higher-rank orders arise from electronic coupling between multipoles rather than due to a coupling by the lattice. They are extremely challenging to detect. Promising techniques for identifying such hidden orders include methods sensitive to time-reversal symmetry and to the splitting of ionic energy levels: X-ray magnetic circular dichroism, neutron and X-ray diffraction, muon spin resonance, resonant and non-resonant elastic X-ray scattering, and nuclear magnetic resonance. *Magnetoelectric hidden order.* Magnetoelectric materials exhibit a net change in magnetization in response to an applied electric field or vice versa. The leading, linear contribution to the magnetoelectric response—requiring the simultaneous breaking of both spatial and time-reversal symmetries—is closely associated with the presence of magnetoelectric multipoles. These are odd-parity, second-order multipoles arising from the magnetization density.
- **Multipolar systems.** "Hidden-order" phases have been found in insulators, e. g. actinide dioxides or double perovskites of heavy transition metals, magnetoelectric materials, as well as in heavy-fermion intermetallics, see Table 1 for selected references. Multipolar systems share key features that serve as preconditions for "hidden order":
  - Strong electronic correlations
  - High lattice symmetry
  - Strong spin-orbit coupling
  - Large spatial separation between magnetic ions
  - Geometrical frustration with respect to conventional antiferromagnetism



### Box 1. Hidden order (continued)

- **Ab initio modelling.** A major difficulty lies in determining the true spatial symmetry of the multipolar order, even when associated local symmetry breaking is revealed by experiment. This challenge underscores the critical role of theoretical methods and models in exploring hidden-order phases in solid-state systems. In recent years, several *ab initio* methodologies have been developed to investigate multipolar order in correlated systems and their underlying microscopic quantum interactions.

## [H2] The ground-state multiplet and its moments

Local multipolar degrees of freedom are hosted by strongly correlated, typically  $d$  or  $f$ , shells in Mott insulators. The local electronic Hamiltonian for such a shell reads

$$H_{\text{loc}} = H_{\text{CF}} + H_{\text{SO}} + H_{\text{C}}, \quad (1)$$

where the terms on the right-hand side are the crystal field, SOC and on-site Coulomb repulsion (C). With charge fluctuations suppressed due to a large value of the on-site Coulomb repulsion parameter  $U$ , the correlated shell has a well-defined integer ground-state occupancy. Other local energy scales – Hund’s rule coupling  $J_H$ , crystal field splitting, SOC constant  $\xi_{\text{SO}}$  – are, as a rule, much larger than the relevant energy scale of intersite coupling. The highest known transition temperature for the onset of a purely "hidden" order without any observable conventional dipole moments, about 50 K, is probably observed in  $d^2$  double perovskites of Os<sup>63</sup>. This energy scale is to be compared with the magnitude of  $J_H$  (0.2-1 eV) and  $\xi_{\text{SO}}$  (several hundreds meV) in heavy transition metals and  $f$ -electron materials<sup>16,147–150</sup>. Hence, a well-separated ground-state multiplet (GSM) of the local Hamiltonian can be defined for a given shell; only those GSM states are thus relevant for low-temperature orders.

The choice for the GSM depends on the problem at hand. In  $f$ -electron materials the crystal field is much weaker than  $\xi_{\text{SO}}$  ( $\sim J_H$ ). However, the relevant crystal field splitting is still generally substantially larger than the magnitude of intersite coupling, and only the lowest crystal field level is relevant for the corresponding order. In this situation, it is useful to introduce the concept of orbital "pseudo-spin" operators, a fictitious angular momentum encoding the crystal field ground-state manifold<sup>73</sup>.

In contrast, in correlated insulators of heavy ( $4d$  and  $5d$ ) transition metals, the crystal field term in equation (1) is typically larger than  $J_H$ . The crystal field thus defines a relevant manifold (such as  $t_{2g}$  orbitals in cubic perovskites and double perovskites with  $d$  shell occupancy  $N_d \leq 6$ ), which is in turn split by the action of spin-orbit and smaller crystal field terms<sup>16</sup>. In the case of  $t_{2g}$  systems, it is convenient to introduce a pseudo angular-momentum operator  $\ell = 1$  whose matrix elements within this manifold are the same as those of the physical orbital momentum  $l = 2$  with the inverse sign<sup>151</sup>. The SO term for the full  $d$  shell,  $\xi_{\text{SO}} \sum_i \hat{\mathbf{l}}_i \hat{\mathbf{s}}_i$ , transforms into  $-\xi_{\text{SO}} \sum_i \hat{\boldsymbol{\ell}}_i \hat{\mathbf{s}}_i$  acting within the  $t_{2g}$  space. The flip of the SOC sign results in the inversion of the atomic third Hund’s rule, with the shell total angular momentum  $J = L + S$  or  $L - S$  for the less than or more than half-filled shell, respectively<sup>16</sup>.

The GSM with occupancy  $N$  and effective  $J$  is fully determined by the corresponding  $N$ -electron density matrix  $\rho$ ; thus, the value of any operator  $A$  acting within the GSM is given by  $\text{Tr}[\rho A]$ . Apart from its usual matrix representation, the density matrix operator can be equally well defined by its multipole expansion<sup>73,152</sup>

$$\rho = \langle O_{\bar{K}}^Q \rangle O_{\bar{K}}^Q, \quad (2)$$

where  $O_{\bar{K}}^Q$  with  $K \leq 2J$  are normalized,  $\text{Tr} \left[ O_{\bar{K}}^Q O_{\bar{K}'}^{Q'} \right] = \delta_{\bar{K}\bar{K}'} \delta_{QQ'}$ , Hermitian spherical tensor operators of rank  $K$  and projection  $Q$  constructed from the standard multipolar spherical tensors<sup>153</sup> akin to spherical harmonics<sup>73</sup>. Alternatively, the degrees of freedom within the GSM space can also be defined as polynomials of angular momentum operators (Stevens operators) or as products of pseudo-spin-1/2 operators. An expansion similar to that in equation (2) can also be introduced for the one-electron density matrix of a given correlated shell using one-electron multipoles  $\hat{D}_{\bar{K}}^q$ .

The states belonging to a single  $J$ -multiplet have the same parity, leading to only parity-even – charge (even  $K$ ) or magnetic (odd  $K$ ) – multipoles contributing to the expansion in equation (2). Two additional kinds of (parity-odd)

multipoles – magnetic and electrical toroidal moments – need to be included in the multipolar expansion of a general electronic system<sup>154</sup>, that is, in the case of either an atomic cluster or hybridized orbitals on a single atomic site. These multipoles play a crucial role in the magnetoelectrical response<sup>91</sup>. One-electron operators for such multipoles have been recently derived<sup>155,156</sup> and classified by their symmetry<sup>157,158</sup>; the corresponding toroidal order parameters have been proposed in a number of correlated *d* and *f*-electron systems<sup>157</sup>. Recently, magnetic multipoles have been utilized in the cluster-multipole expansion, enabling the decomposition of arbitrary magnetic configurations into magnetic multipole components. This approach facilitates the systematic and efficient generation of magnetic configurations, which can be screened in high-throughput studies to identify the magnetic ground-state phase.<sup>159,160</sup>

The experimental determination of the nature of multipolar order parameters in spin-orbit entangled materials presents a substantial challenge. Quadrupolar moments exhibit strong coupling to phonons with the same symmetry, and can be studied using polarization-resolved Raman spectroscopy, which can probe the symmetry of crystal field multiplets as well as the relevant phonon modes as a function of temperature.<sup>161–164</sup> Quadrupolar ordering also leads to weak local lattice displacements, which may be measured using high-resolution synchrotron X-ray diffraction, thus allowing one to infer the nature of quadrupolar ordering.<sup>60</sup> Higher-order multipolar symmetry breaking, such as octupolar ordering, could be probed using a judicious combination of magnetic field and strain.<sup>165,166</sup> Crudely, we may view octupolar order as a product of dipolar and quadrupolar orders, so that applying a magnetic field can pin the dipolar order, allowing one to detect the emergent quadrupolar signature via lattice response. Conversely, the application of a symmetry-tailored strain field can reveal the octupolar order via an emergent dipolar signal. Polarized neutron scattering,<sup>167,168</sup> resonant X-ray scattering<sup>35,73</sup>, unconventional magnetoelectric responses,<sup>169</sup> second-harmonic generation,<sup>170,171</sup> and nuclear magnetic resonance studies<sup>52,172,173</sup> may also be used to infer partial symmetry information for higher multipolar orders, but their complete characterization remains a complex task.

We detail various definitions of multipolar operators and their connection to the classical multipolar expansion of the charge and magnetic densities in the Supplemental Material. Intersite coupling mechanisms between multipolar moments are outlined in **Box 2**.

## Box 2. Intersite interactions between multipole moments

The local multipolar moments introduced in the text interact on the lattice through various mechanisms, including purely electronic interactions—such as exchange and classical electromagnetic interactions—as well as electron-lattice coupling. These couplings were initially uncovered and intensively investigated in more conventional magnetic and orbital-ordered  $3d$  systems<sup>174–179</sup>. In the context of high-rank multipole moments in spin-orbit insulators, the relative importance of those coupling terms may change as we discuss below while introducing most important coupling mechanisms.

- *Intersite exchange interactions (IEI)*. In strongly-correlated systems, direct exchange due to correlated orbitals on neighboring sites overlapping is usually small, and kinetic exchange due to correlated hopping dominates. In complex systems like double perovskite, a significant contribution due to hopping through non-magnetic cations is revealed by ab initio analysis<sup>66</sup>. In  $f$ -electron materials, the CF splitting is weak and one starts with deriving symmetry-allowed coupling between one-electron hopping terms and the full  $J$ -multiplet<sup>180, 181</sup>. The resulting formulae are quite cumbersome and semi-empirical superexchange is often used together with the assumption that dipolar superexchange dominates over multipolar one.<sup>182</sup> The latter assumption is not of general validity as shown by recent direct ab initio calculations of superexchange in rare-earth oxides and nitrides<sup>34, 86</sup>.

Whatever the exchange mechanism is, the low-energy Hamiltonian for a chosen GSM reads

$$H_{\text{IEI}} = \sum_{\langle ij \rangle} \sum_{KK'} V_{KK'}^{QQ'}(\mathbf{R}_{ij}) O_K^Q(i) O_{K'}^{Q'}(j), \quad (3)$$

in terms of the multipolar operators  $O_K^Q$  (2) and the intersite exchange interactions (IEI)  $V_{KK'}^{QQ'}(\mathbf{R}_{ij})$  for the bond  $\langle ij \rangle$  linking two corresponding correlated sites. The second sum in (3) is over the repeated  $K$  and  $Q$  indices. The IEI constant  $\hat{V}$  for a given bond is thus a  $(2J_i + 1)^2 \times (2J_j + 1)^2$  matrix. The bond point-group and time-reversal symmetries nullify some matrix elements<sup>73</sup> in  $\hat{V}$ , though non-zero couplings are still numerous for systems with large effective  $J$ <sup>34, 66, 67, 86</sup>.

- *Classical electromagnetic coupling*. The classical dipole-dipole ( $K = K' = 1$ ) magnetic coupling  $\sum I_{11}^{QQ'}(\mathbf{R}_{ij}) g_i O_1^Q(i) g_j O_1^{Q'}(j)$  is well known to be important in rare-earth magnets with large on-site dipole moments and gyromagnetic ratios  $g$  as well as weak exchange couplings, where it may provide a leading contribution to the two-site magnetic anisotropy<sup>182</sup>. Since classical electromagnetic interactions decay as  $\sim r^{-K-K'-1}$ , the higher-rank magnetic (odd  $K$ ) coupling are expected to be small. In contrast, electrostatic interaction between quadrupoles ( $K=2$ ) has been suggested to play an important role in some systems, in particular, due to a large spacial extend of  $5d$  orbitals, in multipolar  $5d$  double perovskites<sup>42, 183</sup>. Its magnitude remains, however, poorly constrained by ab initio calculations or experiments<sup>184</sup>.
- *Electron-lattice (EL) coupling*. In accordance with the Jahn-Teller (JT) theorem<sup>185</sup>, the on-site JT term  $g_{\text{JT}} O_K^Q(i) q_Q(i)$  that linearly couples electric (even  $K$ ) operators with a local lattice distortion mode  $q_Q$  of the same symmetry must be present in the Hamiltonian whenever the GSM has a non-Kramer's (non-time-reversal related) degeneracy<sup>186</sup>. The local distortions on different sites are then coupled elastically by the lattice resulting in an effective lattice-mediated intersite interaction between electric multipoles<sup>177, 179</sup>. In practice, this cooperative JT effect in strongly-correlated systems always involve quadrupoles ( $K=2$ ); relevant hexadecapoles ( $K=4$ ) of the same symmetry are expected to give a small contribution. The shape of quadrupoles is affected by SO entanglement resulting in qualitative changes in the JT potential energy surface between weak to strong SO limit<sup>187, 188</sup>. In high-symmetry (cubic) multipolar systems, the EL coupling is generally important and can give a leading contribution to intersite interactions<sup>32, 65, 89</sup>. In lower-symmetry systems with the GSM that is formally a singlet or a Kramer's doublet, a pseudo-JT effect<sup>117, 189</sup> may arise due to inter-multiplet mixing induced by  $q_Q$ .

## Box 2. Intersite interactions between multipole moments (continued)

Overall, the low-energy many-body effective Hamiltonian (MBEH) thus reads

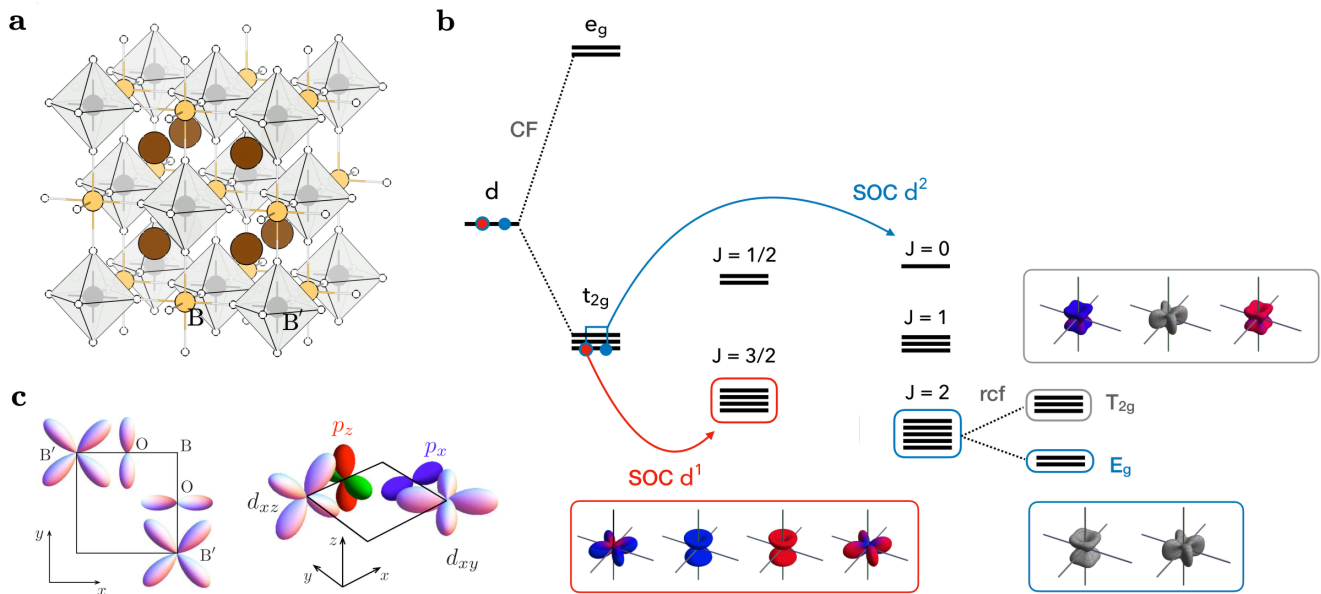
$$H = H_{\text{IEI}} + H_{\text{EM}} + H_{\text{EL}} + \sum_i H_{1s}(i), \quad (4)$$

where the classical electromagnetic term  $H_{\text{EM}}$  has the same form as the IEI term (3). The electron-lattice term  $\hat{H}_{\text{EL}}$  can be written differently depending whether ionic degrees of freedom are kept explicitly (thus JT dynamics is included) or integrated out. In the latter case the EL term also takes the same bi-linear form (3) with  $K, K' = 2$ . Finally,  $\hat{H}_{1s}$  contains all the remaining single-site terms that are active within the GSM like its CF splitting.

### [H2] Model superexchange Hamiltonians

Qualitative features of the multipolar orders, in particular in  $d$ -electron materials, have been intensively studied by the traditional method of deriving intersite exchange interactions from tight-binding Hamiltonians. This framework, initiated by the classical work of Philip Anderson<sup>174</sup>, revealed the importance of superexchange through the  $p$ -orbitals of surrounding ligands. The tight-binding (Goodenough-Kanamori)<sup>175,176</sup> rules relating the transition metal-ligand-transition metal bond geometry with the sign of the resulting superexchange between transition metal spins have been intensively used to analyze coupling in correlated magnetic insulators. The orbital (Kugel-Khomskii) superexchange<sup>178</sup> needs to be simultaneously included. These approaches have been generalized to the case of strong SOC entangling orbital and spin inter-site exchange<sup>16,19,42,183</sup>.

In particular, this approach was employed to elucidate low-temperature orders in spin-orbit double perovskites (Fig.2a)<sup>42,183</sup>. First, the spin-orbit GSM is established by focusing on the  $t_{2g}$  manifold, separated from the excited  $e_g$  one by a large octahedral crystal field (Fig. 2b). In the absence of SOC, the superexchange coupling within this manifold comprises separate  $t_{2g}$ -orbital and Heisenberg spin terms. Following well-established tight-binding



**Figure 2. Levels structure, ground state multiplets and hopping paths in  $d^1$  and  $d^2$  spin-orbit double perovskites.** (a) The conventional unit cell of  $A_2BB'O_6$  cubic double perovskites. (b) The level splitting on the  $d^1$  and  $d^2$  shells of magnetic  $B'$  sites due to the octahedral crystal field and spin-orbit coupling. Electron density distribution for the ground-state multiplet orbitals is shown after Ref.<sup>190</sup>, the spin-up/spin-down densities are indicated by red-blue color. (c) Hopping paths included in the superexchange model of Chen *et al.*<sup>42</sup> (after Ref.<sup>42</sup>).

rules for  $dp$  hopping<sup>175,176,179</sup>, the leading hopping term is assumed to be due to  $\pi$ -hopping between  $t_{2g}$  and an appropriate ligand- $p$  orbital lying in the same plane  $\alpha$  (such as an  $xy$  electron along the  $y$  direction hopping to the  $p_x$  orbital, **Fig. 2c**)<sup>42</sup>. From the "diagonal" process between  $t_{2g}$  orbitals of the same kind one obtains strong antiferroic spin and ferroic orbital superexchange

$$H_{\text{dr}}^{\alpha} = J_{\text{SE}} \sum_{\langle ij \rangle \in \alpha} \left( S_{i\alpha} S_{j\alpha} - \frac{1}{4} n_{i\alpha} n_{j\alpha} \right),$$

where  $n_{i\alpha}$  is the occupancy operator for the corresponding site and orbital,  $J_{\text{SE}}$  is the superexchange coupling constant. Hopping between two orthogonal  $t_{2g}$  orbitals along two orthogonal bonds (**Fig. 2c**) leads, due to Hund's rule on the ligand- $p$  shell, to a weaker superexchange of the opposite sign with respect to the "direct" term.

In the limit of strong SOC, well justified for heavy  $5d$  transition metal ions, this spin-orbital Hamiltonian can be projected onto the spin-orbit GSM. In the case of  $d^1$  systems, the GSM is the  $J_{\text{eff}} = 3/2$  quadruplet shown in **Fig. 2b**, which, in accordance with equation (2), can host dipoles ( $K = 1$ ), quadrupoles ( $K = 2$ ), and octupoles ( $K = 3$ ). The time-odd spin operators are projected onto superpositions of time-odd  $J = 3/2$  dipoles and octupoles, while  $n_{i\alpha}$  become time-even  $J = 3/2$  quadrupoles of  $e_g$  symmetry. Hence, the  $t_{2g}$  charge and magnetic degrees of freedom project onto the corresponding degrees of freedom of the spin-orbit GSM; moments of opposite symmetry under time reversal do not couple, as expected.

A similar analysis was carried out for  $d^2$  double perovskites<sup>183</sup>, with the projection onto the  $J_{\text{eff}} = 2$  GSM of  $d^2$  ions. In this case, however, it is crucial<sup>63</sup> to take into account the "remnant" crystal field splitting of the  $J_{\text{eff}} = 2$  manifold by the cubic crystal field (**Fig. 2b**). The lower  $E_g$  manifold does not carry any dipole  $K = 1$  moments. This can be shown either by a group-theory analysis or by explicitly projecting the dipole-moment operators onto the  $E_g$  space, where they become zero. The  $E_g$  doublet is thus described by three moments – two quadrupoles and an octupole – that project onto  $x$ ,  $z$ , and  $y$  pseudo-spin-1/2 operators of the  $E_g$  space. Hence, by using such symmetry analysis one can immediately show that in the limit of dominating remnant cubic crystal field only multipolar phases can be realized in  $d^2$  double perovskites.

The approach used in Refs.<sup>42,183</sup> has been intensively employed to explain experimental trends in spin-orbit double perovskites. Despite its success, its predictions on the relative magnitude of various multipolar intersite exchange interactions (IEIs) are based on a rather simplified tight-binding picture and not always supported by recent ab initio calculations. For example, the treatment used in Ref.<sup>42</sup> predicts no superexchange coupling between  $J = 3/2$   $t_{2g}$  quadrupoles in  $d^1$  double perovskites, while this coupling is found to be the leading term in the  $d^1$  double perovskite  $\text{Ba}_2\text{MgReO}_6$  by first principles methods (see **Fig. 3** and the section on materials). Hence, we now turn to ab initio methodologies for multipolar ordering.

## [H2] Total energy DFT-based methods

Total-energy DFT-based approaches do not attempt to construct the low-energy MBEH (equation 4) explicitly. Instead, they treat all electronic degrees of freedom on equal footing and rely on the well-known precision and robustness of DFT<sup>191</sup> to capture the subtle contribution of hidden orders in the total energy. In this framework, an effective MBEH can be constructed *a posteriori* from the computed dependence of the total energy on order parameters. Strong electronic correlations are not properly described by standard DFT with local or semi-local treatment of exchange and correlations; hence, on-site Coulomb interactions need to be treated explicitly, by adding the Hubbard  $U$  term for the correlated shell or employing more advanced corrections such as hybrid functionals<sup>192,193</sup> and the self-interaction correction methods<sup>194</sup>.

[H3] *DFT+U*. The simplest and well-tested approach of this type, *DFT+U*<sup>195</sup>, consists of treating a shell with a local Hubbard interaction within the Hartree-Fock approximation, producing a static orbital-dependent contribution to the DFT one-electron potential and the corresponding correction to the total energy. Once *DFT+U* iterations are converged, the multipolar moments can be calculated from the one-electron density matrix  $\rho^{1el}$  using one-electron multipolar operators  $\hat{D}_k^{q73,74,156,196,197}$  (which are defined in the Supplemental Material).

This framework has been intensively applied to multipolar orders in  $f$ -electron materials<sup>118</sup>. Its main difficulty stems from the dependence of DFT+U on the initial guess for  $\rho^{1el}$ , since the DFT+U total energy exhibits multiple local minima in the space of possible order parameters<sup>198–200</sup>. Hence, to obtain a correct multipolar order one generally needs to preset it by a correct initial guess for  $\rho^{1el}$  on all sites in the magnetic unit cell<sup>118,201,202</sup>. Moreover, since DFT+U always converges to a certain total-energy minimum, it is not possible to estimate the total energy dependence on a continuous change in the order parameters. The straightforward DFT+U also has difficulty treating the higher-temperature orders that only partially lift the GSM degeneracy (such as quadrupolar orders, which preserve Kramer's degeneracy), because the Hartree-Fock approach representing the state by a single Slater determinant cannot describe multiplet physics.

New DFT+U-based approaches that overcome or mitigate these limitations have recently been proposed. In particular, constrained versions of DFT+U<sup>101,203</sup> have been devised to evaluate the total energy upon continuous evolution of the order parameters (dipole moments) in their phase space. To fix the moment of a given site  $i$  along the predefined direction  $\mathbf{M}_i^0$  an energy penalty

$$E = E_0 + \sum_i \gamma [\mathbf{M}_i - \mathbf{M}_i^0 (\mathbf{M}_i^0 \cdot \mathbf{M}_i)]^2, \quad (5)$$

where  $\gamma$  is the penalty energy parameter, is added to the DFT+U total energy. The corresponding "penalty field" then appears in the one-electron Kohn-Sham potential acting to rotate the moment  $\mathbf{M}_i$  towards the chosen direction. While the "penalty" energy contribution is unphysical, it becomes negligibly small once  $\gamma$  is sufficiently large to force the moment to align along  $\mathbf{M}_i^0$ . This approach has been first applied to calculate the MBEH of the  $d^5$   $J_{\text{eff}} = 1/2$  iridate  $\text{Sr}_2\text{IrO}_4$ <sup>101</sup> and, subsequently, to estimate multipolar intersite interactions in  $\text{UO}_2$ <sup>204</sup> and in the  $J_{\text{eff}} = 3/2$  double perovskite  $\text{Ba}_2\text{NaOsO}_6$ <sup>65</sup>. In  $\text{Sr}_2\text{IrO}_4$  and  $\text{Ba}_2\text{NaOsO}_6$  the magnetic order is planar; the magnitude of the total-energy contribution from IEIs between various multipoles (equation 3) is parameterized as a function of the ordered moments' canting angle  $\phi$  in the  $xy$  plane. This method does not enable full separation of all bilinear contributions in equation (3), since multipoles of different  $K$  with the same  $Q$  exhibit the same  $\phi$ -dependence. However, the method can identify multipolar Dzyaloshinskii-Moriya IEIs that arise due to inversion-symmetry breaking by lattice distortions<sup>65</sup>.

Another version of DFT+U<sup>79</sup> in the spirit of standard constrained DFT<sup>205</sup> adds the term  $\sum_{ikq} s_{kq}^i (\hat{w}_{kq}^i - \tilde{w}_{kq}^i)$  to the DFT(+U) functional for a chosen subset of sites  $i$  and one-electron multipole operators<sup>74</sup>  $\hat{w}_{kq}^i$  with target expectation values  $\tilde{w}_{kq}^i$ . The Lagrange multiplier  $s_{kq}^i$  is in fact a staggered multipole field inducing corresponding on-site moments. Its contribution to the total energy is then subtracted to obtain the physical total energy for the induced multipolar order.

The constrained DFT+U methods<sup>79,101,203</sup> described above cannot treat purely electric (such as quadrupolar) orders, as they rely on magnetic polarization to obtain a non-degenerate ground state, which can be represented by a single Slater determinant, on correlated shells. A recent development<sup>206</sup> addresses this limitation through an approach similar to the DFT+U treatment for non-spin orbit correlated paramagnetic systems with local moments<sup>207,208</sup>. Namely, atomic multiplet physics is mimicked by a static magnetic disorder within a large supercell. The random orientation of the moments is enforced by the penalty term in equation (5). While time-reversal symmetry is broken locally at each site, it is globally preserved due to the random orientations of the moments, enabling the evaluation of quadrupole moments in a paramagnetic environment.

### [H3] DFT+DMFT.

A more advanced treatment of electronic correlations is provided by combining DFT with DMFT<sup>209–213</sup>. In this framework, a single correlated shell is embedded into a self-consistent electronic bath representing the rest of system; this "quantum impurity problem" is then solved beyond the static Hartree-Fock approximation, with local correlations fully encoded by a dynamical single-site potential: DMFT self-energy  $\Sigma_i(\omega)$ . Solving the quantum impurity problem becomes increasingly more complex and time-consuming with increasing degeneracy of the correlated shell, reducing its symmetry and lowering the temperature, which corresponds exactly to the regime where multipolar orders in spin-orbit insulators arise. Direct DMFT calculations of multipolar ordered phases are

thus very rare<sup>214</sup>. The quadrupolar phase of the d<sup>1</sup> double perovskite Ba<sub>2</sub>ReMgO<sub>6</sub> has been recently studied by DFT+DMFT<sup>215</sup>. In this case, also especially hard for DFT+U, direct DFT+DMFT calculations of the ordered phase were prohibitively computationally expensive. Hence, the relative stability of competing orders was estimated by applying suitable small staggered quadrupolar fields in the symmetry-unbroken high-temperature phase; the transition temperatures were then extracted from Curie-Weiss fits of corresponding static susceptibilities.

## [H2] Force-theorem methods

Force-theorem approaches are based on a general DFT "force theorem"<sup>216</sup> that states that the change of DFT total energy upon a small perturbation is given by the corresponding change in the one-electron band energy only. The magnetic force theorem (MFT)<sup>217,218</sup> then evaluates the change in DFT grand thermodynamic potential  $\Omega$  in an ordered magnetic state upon an infinitesimal tilting  $\delta\Theta$  of two moments at the sites  $i$  and  $j$  away from the ground state direction. The resulting expression for intersite coupling in an effective classical Heisenberg model involves one-electron propagators between sites  $i$  and  $j$  as well as the on-site DFT exchange field on the two sites. The MFT formula can be straightforwardly generalized to DFT+DMFT<sup>219</sup> with the DFT exchange field substituted by the difference between DMFT spin-up and spin-down self-energies. MFT approaches of different flavors<sup>218,220,221</sup> have been widely used to investigate magnetic systems<sup>72</sup> and its formalism generalized to treat Dzyaloshinskii-Moriya IEIs<sup>222</sup>. Its generalization to multipolar IEIs is, however, not straightforward. First, an ordered state for a given set of multipolar moments needs to be obtained before the MFT can be applied, which is a daunting task for correlated insulators. Second, multipolar operators do not have any obvious classical counterparts, and their effective Hamiltonian should naturally be quantum.

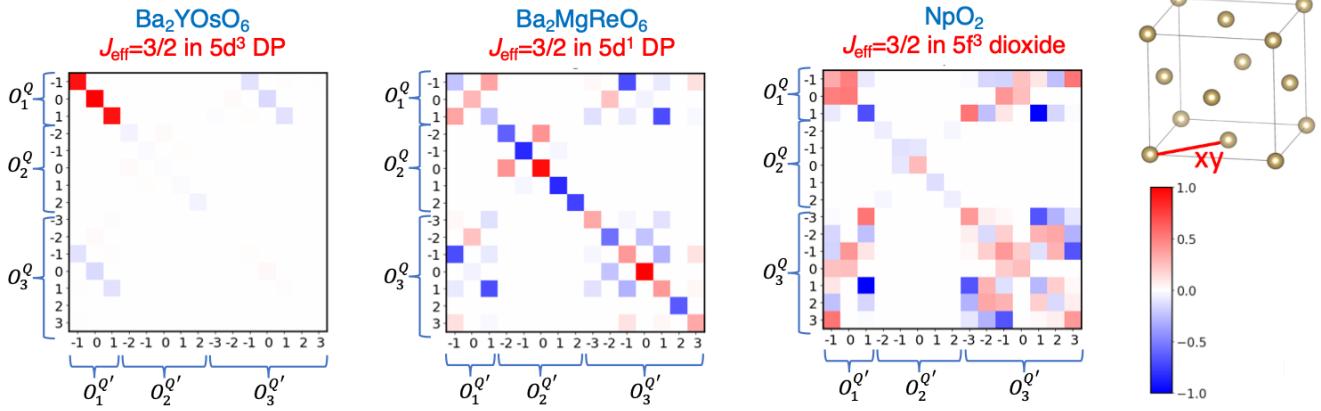
The MFT was generalized to multipolar orders within DFT+U<sup>76,77</sup>. This approach consists in starting with an ordered state and calculating the change in the DFT+U band energy upon simultaneous flipping of two multipole moments  $\langle O_K^Q(i) \rangle$  and  $\langle O_{K'}^Q(j) \rangle$  on neighboring sites  $i$  and  $j$  while freezing the converged DFT+U Kohn-Sham potential. The method can in principle extract the IEIs for all the moments that are consistent with the single Slater-determinant ground state. As other DFT+U based techniques, this method has difficulties with time-even orders only partially lifting the GSM degeneracy, such as the purely quadrupolar order in the case of a half-integer  $J_{\text{eff}}$ .

[H3] *Force theorem in Hubbard-I.* A reasonable starting point for treating correlated insulators can often be obtained within a quasi-atomic Hubbard-I (HI)<sup>223</sup> approximation. This DFT+HI approach<sup>211</sup> is a substantial simplification over the general DFT+DMFT framework, because the hopping between the effective bath and quantum impurity is neglected. The DMFT impurity problem then reduces to diagonalizing a local Hamiltonian for a single correlated shell subjected to external (crystal or exchange) fields, which contribute to the one-electron level positions of the local Hamiltonian. The atomic Green's function and self-energy are then evaluated. DFT+HI contains all single-shell multiplet effects and produces a rather good description of the electronic spectrum in local-moment paramagnets<sup>224-227</sup>. The straightforward DFT+HI is not suitable for ordered phases, because it neglects kinetic exchange, which is contained in electronic hopping from the bath.

The "kinetic energy" term of the DFT+DMFT grand thermodynamic potential  $\Omega$ <sup>212,228</sup>, however, still contains all electron hopping terms through Kohn-Sham Hamiltonian  $H_{\text{KS}}$  even when the HI approximation is used for the DMFT self-energy. The force theorem in Hubbard-I (FT-HI)<sup>75</sup> utilizes this fact to extract IEIs from the paramagnetic (symmetry unbroken) DFT+HI electronic structure. Namely, one considers fluctuations corresponding to the simultaneous appearance of small moments within the GSM of two sites in an otherwise paramagnetic environment and, similar to the MFT, evaluates the response of  $\Omega$ . Introducing trace-conserving density matrix fluctuations  $\rho_{M_1 M_2}$  within the GSM and neglecting, in the force-theorem spirit, their impact on the Kohn-Sham potential, the following formula was derived<sup>75</sup> for the matrix elements of the quantum MBEH (equation 3):

$$\langle M_1^i M_3^j | \hat{H}_{\text{IEI}} | M_2^i M_4^j \rangle = \text{Tr} \left[ G_{ij} \frac{\delta \Sigma_j}{\delta \rho_{M_3 M_4}^j} G_{ji} \frac{\delta \Sigma_i}{\delta \rho_{M_1 M_2}^i} \right], \quad (6)$$

where  $i$  and  $j$  are site labels. The FT-HI structure is very similar to the MFT one, but uses the paramagnetic intersite propagators  $G_{ij}$  together with the on-site "vertices", which are derivatives of the HI self-energy over fluctuations



**Figure 3. Intersite exchange interaction (IEI) matrices for three  $J_{\text{eff}}=3/2$  SO insulators calculated by the force theorem in Hubbard-I (FT-HI) method. Left.** The  $d^3$  double perovskite (DP)  $\text{Ba}_2\text{YOsO}_6$ . In this case there is no spin-orbit entanglement within the  $J_{\text{eff}}=3/2$  ground state (GS), and the dominating IEI are antiferroic dipole-dipole Heisenberg terms as expected for a pure spin- $3/2$  system. Perturbative admixture of excited multiplets by SO gives rise to weaker multipolar IEI, in particular, to IEI between dipoles ( $K=1$ ) and octupoles ( $K=3$ ) that induce a large gap in the magnetic excitation spectrum of  $\text{Ba}_2\text{YOsO}_6$ <sup>231</sup>. **Middle.** The  $d^1$  DP  $\text{Ba}_2\text{MgReO}_6$  exhibits a spin-orbit entangled  $J_{\text{eff}}=3/2$  GS. The states within this multiplet feature a strongly anisotropic charge distribution i. e. large quadrupole (QP) moments. The QP-QP ( $K=2$ ) IEI thus dominate in this system, leading to a competition between ferroic  $t_{2g}$  and antiferroic  $e_g$  electronic QP orders<sup>35,90</sup>. **Right.** The  $f^3$  dioxide  $\text{NpO}_2$ . Proliferation of non-zero IEI in this case stems from complex shapes of  $f$ -electron orbitals leading to many non-zero contributions to superexchange. The time-odd (dipole-octupole) IEI are seen to dominate. These degrees of freedom within  $\text{Np } J_{\text{eff}}=3/2$  are in fact downfolded high-rank magnetic moments of the physical  $J=9/2$  atomic GS of  $\text{Np } f^3$ ; the dipole-octupolar order in the  $J_{\text{eff}}$  space of  $\text{NpO}_2$  thus corresponds to a rank-5 (trikontadipole) order of physical multipoles<sup>67,82</sup>. All IEI matrices are plotted for the  $\mathbf{R} = [1/2, 1/2, 0] \equiv xy$  nearest-neighbor bond of the face-centered cubic (fcc) lattice shown in the top right corner.

calculated with analytical formulas<sup>75</sup>. Once all MBEH matrix elements for a given bond  $ij$  are computed, they are transformed to the form of interaction between multipoles (equation 3) by using the orthonormality properties of multipolar operators  $O_K^Q$ .

The FT-HI method can thus calculate the IEI matrix (equation 3) for all multipoles starting from a single self-consistent DFT+HI calculation for a paramagnetic phase. Its applicability is limited to Mott insulating phases, where the FT-HI formula (equation 6) reproduces the lowest-order in hopping/ $U$  contribution to superexchange<sup>75</sup>. The mapping to a multipolar MBEH also requires a specific choice of the phases for GSM states  $|M\rangle$ , that is, a proper pseudo-spin basis should be constructed<sup>229</sup>.

The method has been extensively applied to multipolar  $d$  and  $f$  electron compounds. The IEI matrices calculated by FT-HI for several  $J_{\text{eff}}=3/2$  systems with magnetic fcc sublattice are displayed in **Fig. 3**; a clear connection between the nature of the ground state multiplet and the structure of IEIs can be observed. All IEI matrices in **Fig. 3** exhibit no coupling between charge quadrupoles ( $K=2$ ) and magnetic dipoles/octupoles, because the IEI Hamiltonian calculated in the paramagnetic state must be invariant under time reversal. Zero matrix elements within various  $KK'$  blocks can be understood from a group theory analysis<sup>73</sup>, as demonstrated, for example, for a  $d^1$  double perovskite<sup>90</sup>. **Fig. 3** shows purely electronic IEIs; the electron-lattice coupling can also be mapped into an effective interaction between quadrupoles<sup>177,178,230</sup> which is particularly important in the case of  $d^1$  double perovskites, as discussed in the section on materials. Since electron-lattice quadrupole-quadrupole terms obey the same symmetry as electronic IEI ones, the electron-lattice quadrupole-quadrupole blocks feature the same non-zero matrix elements.

Various other applications of the FT-HI are discussed in the section on materials.

## [H2] Susceptibility-based approaches

Second-order phase transitions can be detected by the divergence of the general static susceptibility  $\chi_K^Q(\mathbf{q}) = \langle O_K^Q(\mathbf{q}) O_K^Q(-\mathbf{q}) \rangle$  for a given primary order parameter  $\langle O_K^Q \rangle$  at the ordering  $\mathbf{q}$ -vector in the Brillouin zone.

In the DMFT framework, the general static susceptibility  $\chi(\mathbf{q}) = \sum_{n,n'} X(i\omega_n, i\omega_{n'}, \mathbf{q})$  is given by the two-electron Green's function, which is a function of two fermionic Matsubara frequencies and four  $(m, \sigma)$  indices labeling the correlated-shell orbitals. This Green's function is obtained by solving a Bethe-Salpeter equation<sup>209,232</sup> :

$$\hat{X}^{-1}(\mathbf{q}) = \hat{X}_{\text{imp}}^{-1} - \hat{X}_{\text{imp},0}^{-1} + \hat{X}_0^{-1}(\mathbf{q}), \quad (7)$$

where the subscripts "imp" and "0" label the impurity and bare (single bubble) susceptibilities, respectively, and the hats label matrices in frequency indices. The bare susceptibilities are diagonal in the  $nn'$  indices, while the two-particle Green's function  $\hat{X}_{\text{imp}}$  is a large non-diagonal matrix with the rank rapidly increasing with the number of Matsubara points included and correlated shell degeneracy. Though the impurity susceptibilities can be efficiently calculated by some many-body methods<sup>233</sup>, solving the Bethe-Salpeter equation is numerically very heavy, especially for large shell degeneracies and in the low-temperature regime relevant to multipolar orders.

To tackle this problem, a single-value decomposition of the impurity susceptibility was performed<sup>234</sup>, keeping only the leading eigenvalue  $\hat{X}_{\text{imp}}(i\omega_n, i\omega_{n'}) \propto \phi_0(i\omega_n)\phi_0^*(i\omega_{n'})$  and showing that this approximation is valid for leading fluctuation channels in one and two-band Hubbard models in the strong-coupling limit. The susceptibility is then rewritten in a random-phase approximation (RPA)-like form,  $\chi^{-1}(\mathbf{q}) = [\chi_{\text{imp}}^{-1} - I(\mathbf{q})]$ , where the IEI reads  $I(\mathbf{q}) = \text{Tr}_n |\phi_0(i\omega_n)|^2 \left( \hat{X}_{\text{imp},0}^{-1}(i\omega_n) - \hat{X}_0^{-1}(\mathbf{q}, \omega_n) \right)$ . The method was applied<sup>235</sup>, in conjunction with DFT+HI, to multipolar interactions in CeB<sub>6</sub>.

## [H2] Cluster methods

Cluster methods consider a many-electron problem on finite-size clusters of correlated sites with its parameters (hopping between the sites and their local Hamiltonian) derived in an ab initio way for a continuous system. Subsequently, within a strong-coupling perspective suitable for correlated insulators, the single-site terms are treated exactly, while the hopping within the cluster is typically included perturbatively to obtain the IEI between the cluster's sites. To derive the kinetic exchange IEI, a pair of correlated sites is typically considered.

A generalization of this traditional approach to superexchange<sup>236,237</sup> to the case of multipolar interactions<sup>86,238,239</sup> includes all active orbitals on correlated sites into the cluster Hamiltonian that reads

$$H_{\text{cl}} = \sum_{il} H_{\text{loc}}^{il} + H_{\text{C}}^{\text{int}} + H_t = H_0 + V$$

where local terms (Eq. 1) are included for active orbitals  $l$  on each correlated site  $i$ , and  $H_{\text{C}}^{\text{int}}$  and  $H_t$  are the intersite Coulomb interaction and hopping terms within the cluster, respectively. On the right-hand side, those terms are regrouped into those that contribute to the splitting within the GSM of correlated shells ( $V$ ) and those that do not ( $H_0$ ). Subsequently, by projecting  $H_{\text{cl}}$  to the manifold of GSM,  $P_0 = \otimes_i (\sum_M |iJM\rangle \langle iJM|)$ , the MBEH is evaluated. It includes crystal field splitting within the GSM as well as various intersite terms. In particular, the kinetic exchange is evaluated within the second-order perturbation theory

$$H_{\text{KE}} = P_0 H_t \left( \sum_{s \notin \text{GSM}} P_s \frac{1}{E_0 - H_0} P_s \right) H_t P_0,$$

where the projection  $P_s$  runs over all states outside of the GSM manifold, and  $E_0$  is the GSM eigenenergy within  $H_0$  (which is the same for all states within the GSM manifold). In addition to the kinetic exchange, other contributions like direct exchange are also included.

Another approach<sup>240</sup>, which is suitable for small clusters, performs an exact diagonalization of  $H_{\text{cl}}$  followed by a unitary (Schrieffer-Wolff<sup>241</sup>) transformation  $\mathcal{R}_{P \rightarrow P_0}$  from the low-energy subspace  $P$  of the dimension  $((2J_{\text{eff}} + 1)^2)$  of a two-site cluster to the GSM subspace  $P_0$  of two disconnected sites defined as above. The resulting Hamiltonian

within the  $P_0$  subspace then becomes  $H_{\text{eff}} = P_0 \mathcal{R}_{P \rightarrow P_0} H_{\text{cl}} \mathcal{R}_{P \rightarrow P_0}^\dagger P_0$ . This Hamiltonian  $H_{\text{eff}}$  contains on-site and IEI terms; the latter can be again transformed to the multipolar form (equation 3). Several similar methods based on exact diagonalization<sup>242–244</sup> also perform a mapping between non-interacting space  $P_0$  and  $P$  by expanding  $P$  into the basis of pseudo-spin states  $P_0$ .

## [H2] Approaches to electron-lattice interactions

The electron-lattice term in (Eq. 4) can provide a crucial contribution to the energetics of multipole orders<sup>32,35,89</sup>. It arises due to the Jahn-Teller coupling between an electronic quadrupole operator and local distortions of the same symmetry (both labeled by their irreducible representation  $\Gamma$  and projection  $Q$ ). The electron-lattice term consists of single-site and intersite contributions<sup>177</sup>:

$$H_{\text{EL}} = H_{\text{harm}} + H_{\text{JT}} + H_{\text{EC}} = \sum_{i\Gamma Q} \left( \hbar\omega_\Gamma (\hat{n}_{\Gamma Q}^i + 1/2) - g_\Gamma O_2^{\Gamma Q}(i) q_{\Gamma Q}(i) \right) + \sum \Phi_{\Gamma\Gamma'}^{QQ'}(ij) q_{\Gamma Q}(i) q_{\Gamma'Q'}(j), \quad (8)$$

where  $\Phi_{\Gamma\Gamma'}^{QQ'}(ij)$  is the intersite elastic coupling, also termed the force constant matrix between irreducible local distortion modes on correlated sites  $i$  and  $j$ , and the summation over all repeated indices is implied in the last term. Hence, to define the electron-lattice interaction one needs to determine all so-called vibronic parameters, that is, the local force constant matrix  $\Phi_{\Gamma\Gamma'}^{QQ'}(ii)$  that determines the single-site oscillator strength  $\omega_\Gamma$ , the intersite one  $\Phi_{\Gamma\Gamma'}^{QQ'}(ij)$ , and the Jahn-Teller couplings  $g_\Gamma$ .

In the context of multipolar order in spin-orbit double perovskites, a framework for single-site terms in the electron-lattice Hamiltonian was developed<sup>245</sup> based on evaluating the adiabatic potential energy surface, that is, the electronic energy as a function of fixed ionic coordinates  $\{q_{\Gamma Q}\}$ , by ab initio DFT or quantum chemistry methods. This potential energy surface was calculated at a grid in the  $\{q_{\Gamma Q}\}$  space and then fitted with the expected form<sup>186</sup> for  $t_{2g}$  electrons on a transition metal ion coupling to all  $e_g$  and  $t_{2g}$  symmetry distortions of a ligand octahedron surrounding it. Going beyond the linear Jahn-Teller coupling in equation (9), higher-order Jahn-Teller couplings were included.

Alternative and computationally less demanding approaches<sup>32,35</sup> are based on DFT or DFT+HI calculations for continuous systems to extract the linear Jahn-Teller coupling only. In particular, the DFT total energy as well as the quadrupoles of the correlated shell were calculated<sup>35</sup> upon the  $\mathbf{k}$ -space distortion mode amplitude  $q_{\mathbf{k}\theta}$ , where  $\theta$  labels irreducible distortion modes at a given  $\mathbf{k}$  point. By writing the total energy as the sum of a bilinear Jahn-Teller term and a quadratic elastic term<sup>246,247</sup>, Jahn-Teller couplings and corresponding elastic constants are obtained from a fit of total energy vs  $q_{\mathbf{k}}^\theta$ .

Jahn-Teller couplings in DFT+HI were calculated by a similar approach of applying small distortion modes<sup>32</sup>. The splitting of the GSM levels that are degenerate in the undistorted structure is due to the Jahn-Teller term  $g_\Gamma q_{\Gamma Q} O_2^{\Gamma Q}$ . Since the GSM eigenvalues and states are directly computed in the HI approach, the magnitude of the coupling constant can be simply extracted from a linear fit of the splitting versus distortion amplitude. The electronic quadrupole IEI cannot contribute to this splitting within the HI approximation, since the electron hopping inducing them is neglected while solving the impurity problem. The spurious self-interaction contribution to the Jahn-Teller splitting is eliminated by charge-density averaging within GSM<sup>248</sup>. The force constant matrix was evaluated by standard DFT methods<sup>249</sup> and then projected into the space of irreducible displacement modes  $q_{\Gamma Q}(i)$ <sup>32</sup>.

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## [H2] Solving the many-body effective Hamiltonian and calculating properties

The quantum ab initio MBEH (equation 4) for the system at a given volume that includes electronic and, possibly, electron-lattice terms, needs to be solved to obtain ordered phases as a function of temperature and external fields (such as an applied magnetic field). While there is a vast array of advanced methods to solve quantum spin-1/2 models, especially in low dimensions<sup>254–256</sup>, the choice is much more limited for multipolar orders. The size of the Hilbert space rapidly increases for  $J_{\text{eff}} > 1/2$ ; moreover, the IEI Hamiltonian (equation 3) typically has rather low symmetry, unlike typical spin-1/2 models, where such symmetries (such as total  $\langle S_z \rangle$  conservation) are employed to partition the Hilbert space into subblocks. Mapping the full multipolar MBEH into a classical model is generally also not possible.

Hence, multipolar MBEHs are typically solved using a generalized single-site mean-field approximation<sup>182,252</sup>. The intersite terms in equation (3) are mean-field decoupled, that is, products of the fluctuation terms are neglected. This leads to the mean-field Hamiltonian  $\sum_{\alpha K Q} \left( O_K^Q(\alpha) - 1/2 \langle O_K^Q \rangle_{\alpha} \right) V_{KQ}^{\text{MF}}(\alpha)$ , where each site  $\alpha$  in the magnetic unit cell interacts with the Weiss field  $V_{KQ}^{\text{MF}}(\alpha)$  produced by the rest of systems. This mean-field problem is solved self-consistently to obtain the set of order parameters  $\{\langle O_K^Q \rangle\}$  and the free energy  $F_{\text{MF}}$  for a set of possible ordered structures identifying the one minimizing  $F_{\text{MF}}$  at a given temperature.

The electron-lattice term (equation 9) can be mapped into effective lattice-mediated quadrupole-quadrupole coupling<sup>177,178,257</sup>, but this treatment neglects dynamical Jahn-Teller effects. Alternatively, the mean-field approximation can be extended to electron-lattice terms<sup>258</sup> by an analogous mean-field decoupling of the intersite elastic term in eq. 9. A vibronic single-site basis that consists of the electronic wavefunction and local phonon modes is introduced<sup>186</sup> and the Jahn-Teller single-site problem in electronic and elastic mean field is solved by exact diagonalization<sup>245</sup>. When the Jahn-Teller couplings  $g$  are sufficiently weak or strong, weak or strong-coupling

Theories and methods to explore multipolar magnetism and hidden orders		
Properties	Theory	Materials
<i>IEI from the ordered phase</i>	Magnetic Force Theorem <sup>76</sup> Constrained DFT + U <sup>101</sup>	LaMnO <sub>3</sub> <sup>222</sup> , UO <sub>2</sub> <sup>76,77</sup> UO <sub>2</sub> <sup>204</sup> , Ba <sub>2</sub> NaOsO <sub>6</sub> <sup>65</sup>
<i>IEI from the paramagnetic phase</i>	Cluster perturbation theory <sup>86,238,239</sup> Exact diagonalization <sup>46,240</sup>  Force Theorem in Hubbard-I <sup>75</sup>   DMFT susceptibility in strong coupling limit <sup>234</sup>	UO <sub>2</sub> <sup>238</sup> , NdN <sup>86</sup> Ba <sub>2</sub> MOsO <sub>6</sub> ( <i>M</i> = Ca, Mg, Zn) <sup>240</sup> Ba <sub>2</sub> NaOsO <sub>6</sub> <sup>65</sup> UO <sub>2</sub> <sup>250</sup> , NpO <sub>2</sub> <sup>67</sup> , PrO <sub>2</sub> <sup>34</sup> , Ba <sub>2</sub> MgReO <sub>6</sub> <sup>90</sup> , Ba <sub>2</sub> YMO <sub>6</sub> ( <i>M</i> = Os, Ru) <sup>231</sup> Ba <sub>2</sub> MOsO <sub>6</sub> ( <i>M</i> = Ca, Mg, Zn) <sup>66</sup>  CeB <sub>6</sub> <sup>235</sup>
<i>Constrained multipoles</i>	Polymorphous DFT+U <sup>207,251</sup> DFT + HI + Constrained DFT + U <sup>78,79</sup> Constrained DFT + U	Ba <sub>2</sub> MgReO <sub>6</sub> <sup>206</sup> Ba <sub>2</sub> MOsO <sub>6</sub> ( <i>M</i> = Ca, Mg, Zn) <sup>78</sup> KCu <sub>1-x</sub> Zn <sub>x</sub> F <sub>3</sub> <sup>79</sup>
<i>Multipolar susceptibilities</i>	Classical Monte Carlo <sup>70,240</sup> DFT + DMFT <sup>35</sup>	Ba <sub>2</sub> Ca <sub>1-δ</sub> Na <sub>δ</sub> OsO <sub>6</sub> <sup>70</sup> Ba <sub>2</sub> MgReO <sub>6</sub> <sup>35,215</sup>
<i>Electron-lattice interactions</i>	Cluster <sup>35,229</sup> DFT, DFT + Hubbard-I <sup>90</sup>	d <sup>1</sup> Double perovskites <sup>89,229</sup> , Ba <sub>2</sub> MgReO <sub>6</sub> <sup>3</sup> Ba <sub>2</sub> MgReO <sub>6</sub> <sup>90</sup>
<i>Multipolar contributions to E<sub>H</sub> and E<sub>x</sub></i>	DFT+U <sup>74</sup>	Actinides <sup>74</sup>
<i>Excitation spectra</i>	Mean field + RPA <sup>182,252</sup> +multipolar form-factors <sup>66,253</sup>	Ba <sub>2</sub> MOsO <sub>6</sub> ( <i>M</i> = Ca, Mg, Zn) <sup>66</sup> , Ba <sub>2</sub> YMO <sub>6</sub> ( <i>M</i> = Os, Ru) <sup>231</sup>

**Table 2. Theoretical approaches to multipolar magnetism and hidden orders in materials.** For each target property (left column) we list theoretical approaches developed for its calculation (middle column). Applications of each approach to specific materials are listed in the right column.

perturbative approaches can be employed<sup>32,89</sup>.

As in the case of conventional magnetic orders, the mean-field approximation leads to a systematic overestimation of transition temperatures. The overestimation factor depends on the system's effective dimensionality and lattice geometry. In particular, overestimation by about a factor of two was observed in a number of 3D multipolar systems studied with a FT-HI Hamiltonian solved by mean-field<sup>34,67,231,250</sup>. In those cases, the magnetic sublattice is fcc and thus frustrated with respect to antiferromagnetism. Alternatively, a classical model can be introduced for a subset of multipoles, such as quadrupoles or octupoles, and solved by classical Monte Carlo<sup>70,259</sup>.

Once a stable order is identified, single-site excitation within the GSM space can also be calculated from mean-field eigenstates using the Fermi golden rule,  $\sum P_i |\langle f | \mathcal{V} | i \rangle|^2 \delta(\omega - E_f - E_i)$ , where  $P_i$  is the thermal occupation probability for the initial mean-field state  $i$ ,  $\mathcal{V}$  is an appropriate scattering operator for a given probe (such as Raman<sup>260</sup> or resonant inelastic X-ray scattering (RIXS)<sup>261</sup>). To evaluate  $\mathbf{q}$ -dependent excitations measured, for example, by inelastic electron scattering (INS) or RIXS, a generalized RPA can be employed<sup>182,252</sup> to obtain generalized lattice susceptibility. Subsequently, the fluctuation-dissipation theorem is used to obtain from it the relevant scattering cross-section. Appropriate form-factors for scattering from multipoles can be obtained by expanding, within the GSM, the corresponding scattering operator in multipolar operators  $O_K^Q$ <sup>66,253,262</sup>.

Finally, electronic excitations in a wide energy range can be affected by the ordering of the low-energy degrees of freedom that are included in the MBEH. These effects are outside of the GSM space and, hence, cannot be obtained from the MBEH alone. Instead, a mean-field ordered state found by solving the MBEH can be utilized to provide a proper initial guess for all electron DFT+U or DFT+DMFT calculations. A scheme was implemented within DFT+U to initialize the one-electron density matrix  $\rho^{1el}$  in accordance with a multipolar order derived from a FT-HI MBEH. Once an appropriate  $\rho^{1el}$  is constructed, it is used as an initial guess for DFT+U<sup>78</sup>. This typically results into DFT+U converging to the local minimum in total energy corresponding to a given multipolar order; its electronic spectrum is then calculated.

A summary of representative results obtained using the approaches discussed in this section is provided in **Table 2**.

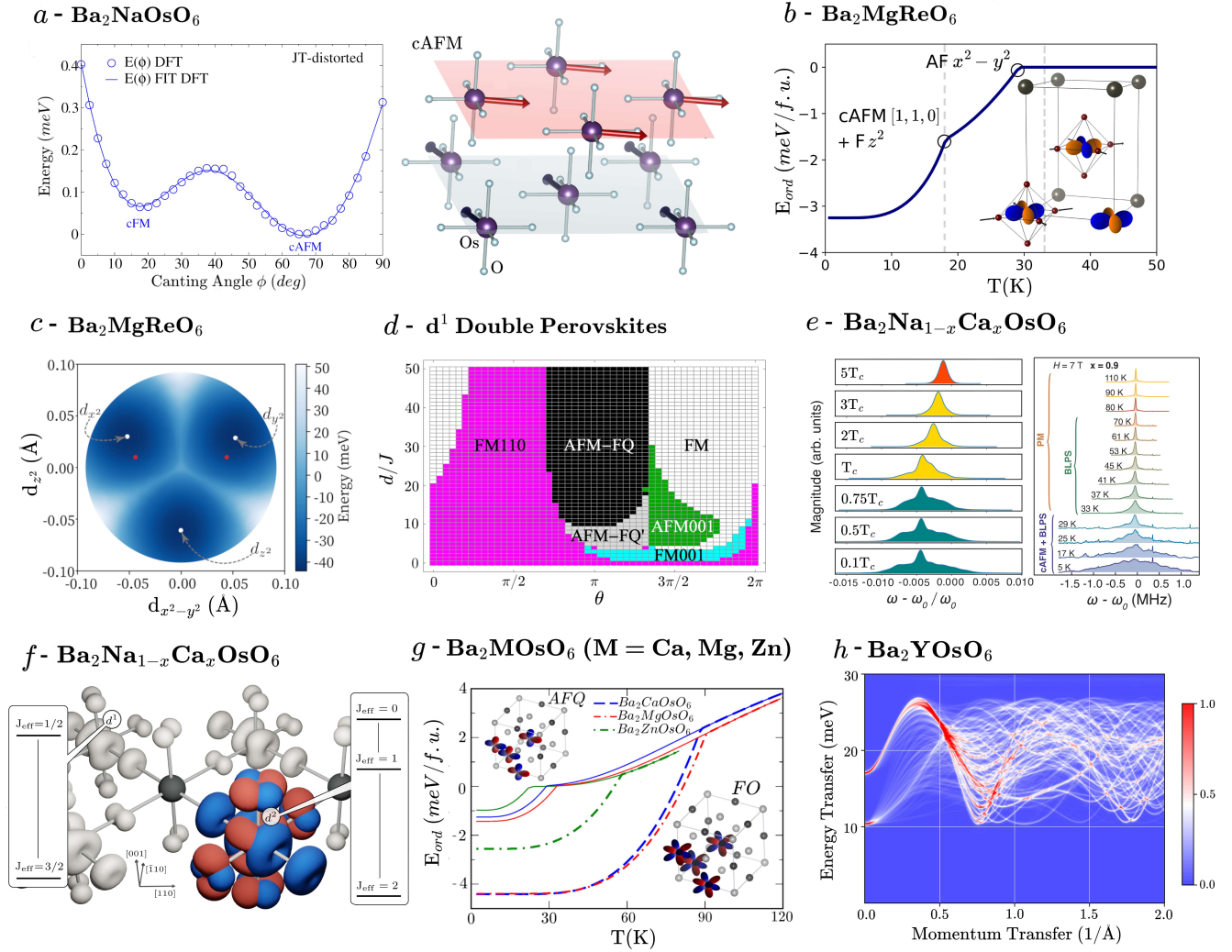
## [H1] Materials

After reviewing theories and methods for studying multipolar hidden orders, we now turn to a survey of the emergence of multipolar phases in materials, focusing on two main materials classes:  $5d$  double perovskites and  $f$ -electron systems.

### [H2] $d$ -electrons: Double Perovskites

The emergence of multipolar degrees of freedom relies on having atomic orbital degeneracy, electron interactions, and strong spin-orbital entanglement. These conditions are met in oxides of heavy  $4d/5d$  transition metals with partially filled  $t_{2g}$  orbitals. Due to the larger spread of the  $4d/5d$  atomic wavefunctions, which tends to favor metallicity, the route to access localized multipole moments is to explore crystals where the heavy  $4d/5d$  transition metal ions are spaced farther apart than in typical perovskite materials. Ordered double perovskites with the chemical formula  $A_2BB'O_6$  provide ideal platforms for such phenomena. Here, B is occupied by an electronically inert ion, and B' by magnetically active  $4d/5d$  ions. In these compounds, the B and B' sites form two sublattices of a cubic lattice, resulting in a B'-B' spacing that is  $\sqrt{2}$  times greater than in single perovskites. Moreover, the relatively large on-site Hubbard interaction  $U$ , in the range of 2–4 eV, combined with substantial SOC strength of a few hundred meV, further facilitates the development of multipolar phases. Consequently, these perovskite compounds are commonly classified as "multipolar Mott insulators."<sup>42,52,60,65,66,263</sup> Indeed, various compounds in this class, featuring electron fillings such as  $d^1$ ,  $d^2$ , and  $d^3$ , have been experimentally and theoretically shown to exhibit clear evidence of distinct multipolar degrees of freedom.

Early studies utilizing RIXS spectra in conjunction with single-site or cluster exact diagonalization successfully extracted the effective Hund's coupling and SOC in these systems. These investigations provided insights into the ground-state and excited-state multiplet structures, laying the groundwork for the identification of multipolar phases in this class of materials<sup>149,150,264,265</sup>.



**Figure 4. Multipolar properties of d-electron systems computed with different methods** **a** | Energy as a function of the canting angle for the 5d<sup>1</sup> double perovskite (DP) Ba<sub>2</sub>NaOsO<sub>6</sub> in the Jahn-Teller (JT) distorted phase with constrained DFT + U<sup>65</sup> (left panel); graphical representation of the canted anti-ferromagnetic (cAFM) phase (right panel)<sup>65</sup>. **b** | Mean field ordering energy vs temperature for the 5d<sup>1</sup> DP Ba<sub>2</sub>MgReO<sub>6</sub> with low-energy effective Hamiltonian including both superexchange and vibronic interactions, as computed with force theorem in Hubbard-I (FT-HI) and DFT(+HI) methods, respectively. The anti-ferro quadrupolar AF $x^2 - y^2$  phase is shown in the inset<sup>90</sup>. **c** | The potential energy surface of the ReO<sub>6</sub> octahedron in the space of  $E_g$  JT normal modes in 5d<sup>1</sup> DP Ba<sub>2</sub>MgReO<sub>6</sub> calculated by quantum chemistry methods<sup>35</sup>. The result is compared with the experimental static octahedral distortions plotted as red dots.<sup>35</sup> **d** | Spin-vibronic phase diagram of 5d<sup>1</sup> DPs as a function of magnetic ( $J$ ) and vibronic ( $d$ ,  $\theta$ ) parameters as obtained within a mean-field approximation<sup>89</sup>. **e** | Computed <sup>23</sup>Na nuclear magnetic resonance (NMR) spectra displaying the shift and broadening of the NMR peak while lowering temperature for lightly Na doped 5d<sup>2</sup> DP Ba<sub>2</sub>CaOsO<sub>6</sub> (left panel)<sup>70</sup>; experimental NMR spectra for comparison with Na concentration  $x = 0.9$  (right panel)<sup>266</sup>. **f** | The magnetic density of the 5d<sup>2</sup> polaron coexisting with the charge density of the 5d<sup>1</sup> sites the Ca doped 5d<sup>1</sup> DP Ba<sub>2</sub>NaOsO<sub>6</sub> as computed in DFT+U<sup>31</sup>. **g** | Mean field ordering energy vs temperature for the 5d<sup>2</sup> DPs Ba<sub>2</sub>MOsO<sub>6</sub> (M = Ca, Mg, Zn) obtained from a many-body effective Hamiltonian computed with DFT+HI and FT-HI. The ferro-octupolar/anti-ferro quadrupolar (FO/AFQ) are graphically represented in the insets<sup>66</sup> **h** | The spherically-averaged inelastic neutron scattering (INS) intensity for the 5d<sup>3</sup> DP Ba<sub>2</sub>YOsO<sub>6</sub> computed for the ground-state 2q AFM phase from FT-HI intersite exchange interactions within the random-phase approximation.<sup>231</sup>

### [H3] $d^1$

$d^1$  double perovskites are well-studied and include  $\text{Ba}_2\text{LiOsO}_6$ ,<sup>267–269</sup>  $\text{Ba}_2\text{NaOsO}_6$ ,<sup>267–269</sup>  $\text{Ba}_2\text{YMoO}_6$ ,<sup>270</sup>  $\text{Ba}_2\text{CaReO}_6$ ,<sup>271,272</sup>  $\text{Ba}_2\text{CaOsO}_6$ ,<sup>271</sup>  $\text{Ba}_2\text{MgReO}_6$ ,<sup>60,273,274</sup> and  $\text{Ba}_2\text{ZnReO}_6$ .<sup>273</sup> While early work on many of these nominally cubic double perovskites with a  $J = 3/2$  ground state multiplet reported a single low-temperature magnetic transition into weakly ordered antiferromagnetic or ferromagnetic phases with small ordered moments, more recent work has revealed that there is a higher-temperature transition at which the crystal point group symmetry is subtly broken, splitting the  $J = 3/2$  quartet into two Kramers doublets. This weak crystal symmetry lowering occurs either through local distortions, as seen in  $\text{Ba}_2\text{NaOsO}_6$  via NMR and nuclear quadrupole resonance experiments as well as DFT calculations,<sup>52,65,275–278</sup> or through global symmetry breaking, as seen via high-resolution X-ray diffraction measurements of tiny lattice displacements in  $\text{Ba}_2\text{MgReO}_6$ .<sup>35,60,274,279</sup> Such symmetry breaking has been proposed to arise from ordering of quadrupolar degrees of freedom  $\vec{Q} = (J_x^2 - J_y^2, 3J_z^2 - J^2)$  due to orbital charge repulsion.<sup>42,184</sup> However, more recent work assigned a more central role to the lattice degrees of freedom to explain the symmetry-breaking pattern seen in  $\text{Ba}_2\text{MgReO}_6$ , invoking a cooperative spin-orbit Jahn-Teller distortion of phonon modes coupled to these quadrupolar moments.<sup>32,35,90,190,280,281</sup> In certain materials, such as  $\text{Ba}_2\text{YMoO}_6$ ,<sup>23,282</sup> which shows no sign of a magnetic transition down to the lowest temperature, dynamic Jahn-Teller fluctuations might prevent magnetic ordering, leading to exotic spin liquid states, although a valence bond glass has also been proposed as a possible explanation for these observations.

**Figure 4** presents examples of predicted multipolar properties. Magnetically constrained non-collinear DFT successfully identifies the canted antiferromagnetic phase with a canting angle of  $\phi = 66^\circ$  as the ground state of Jahn-Teller-distorted  $\text{Ba}_2\text{NaOsO}_6$  (**Figure 4a**), in excellent agreement with NMR measurements<sup>52,65</sup>. The vibronic mechanisms underlying the antiferroic quadrupolar hidden order in  $\text{Ba}_2\text{MgReO}_6$  have been elucidated using a low-temperature many-body effective Hamiltonian (equation 4) incorporating electronic and electron-lattice coupling mechanisms<sup>90</sup>. This Hamiltonian is derived from ab initio calculations by combining FT-HI superexchange interactions with DFT(+HI) elastic and Jahn-Teller couplings. This approach reproduces the two phase transitions observed in thermodynamics, XRD and REXS experiments<sup>35,60,273</sup>, which correspond to the breaking of cubic symmetry and the subsequent emergence of an antiferroic quadrupolar (AFQ) phase and a canted AFM phase (**Figure 4b**).

The role of electron-lattice coupling in  $\text{Ba}_2\text{MgReO}_6$  and the regime of Jahn-Teller problem relevant for this compound are the subjects of ongoing debate. In particular, a weak-coupling Jahn-Teller regime and a subtle interplay of electron-lattice and superexchange interactions were predicted<sup>90</sup> to stabilize the observed AFQ order. In contrast, other work<sup>89</sup> assumed a strong Jahn-Teller regime, that is, the  $\text{ReO}_6$  Jahn-Teller system tunneling between the minima of the Jahn-Teller potential energy surface. This potential energy surface with respect to local Jahn-Teller distortion computed by quantum-chemical approaches<sup>35</sup> is shown in **Fig. 4c**; it reveals three degenerate minima, in qualitative consistency with the experimental distortion amplitudes<sup>60</sup>. Using the strong Jahn-Teller-regime approximation<sup>89</sup> various quadrupole orders were obtained, as well as a global diagram of magnetically ordered states in  $d^1$  double perovskites (**Fig. 4d**). In this picture, the emergence of various ordered phases is mainly driven by the cooperative effects of spin-orbit coupling and electron-lattice interactions. Moreover, only  $R\ln 2$  magnetic entropy seems to be recovered by heating  $\text{Ba}_2\text{MgReO}_6$  up from zero to the AFQ transition point<sup>281,283</sup>; this should be contrasted with the value of  $R\ln 4$  expected upon recovering the  $J = 3/2$  degeneracy in the paramagnetic phase. None of the approaches discussed above can account for this missing entropy in  $\text{Ba}_2\text{MgReO}_6$ .

### [H3] $d^2$

For the  $d^2$  double perovskites  $\text{Ba}_2\text{CaOsO}_6$ ,<sup>271</sup>  $\text{Ba}_2\text{MgOsO}_6$ ,<sup>273</sup> and  $\text{Ba}_2\text{ZnOsO}_6$ ,<sup>273</sup> thermodynamic measurements reveal a single magnetic phase transition, with  $T_c \sim 30\text{-}50\text{K}$ , which was attributed to the onset of an antiferromagnetic order, although muon spin resonance ( $\mu\text{SR}$ ) oscillations found puzzling evidence for unusually weak internal fields. While earlier theoretical studies had suggested that the  $5d^2$  configuration could form a robust  $J = 2$  moment,<sup>183,184</sup> more recent theoretical work<sup>63,263</sup> argued that this five-fold multiplet gets split, even in a cubic environment, to yield a non-Kramers  $E_g$  doublet ground state  $\{|0\rangle, |2\rangle + |-2\rangle\}$  and an excited  $T_{2g}$  magnetic triplet. This splitting arises from SOC-induced mixing of  $t_{2g}\text{-}e_g$  orbitals, while it is absent in models projected from the outset onto the

$t_{2g}$  orbitals. This non-Kramers doublet acts as a pseudospin-1/2 degree of freedom, with the pseudospin operators  $(\tau_x, \tau_z) \sim (J_x^2 - J_y^2, 3J_z^2 - J^2)$  encoding quadrupoles, while  $\tau_y \sim \text{Sym}[J_x J_y J_z]$  encapsulates an octupolar moment (with ‘Sym’ denoting symmetrization; these Stevens operators are the same, up to a normalization, as the operators  $O_2^0, O_2^2$ , and  $O_3^{-2}$  introduced in the Supplemental Material). This scenario, together with the assignment of the observed phase transition to ferro-ordering of  $\tau_y$ , was shown to capture the entropy, magnetic susceptibility, and weak  $\mu\text{SR}$  internal fields.<sup>63,263,284</sup> An ab initio electronic MBEH derived for these systems by DFT+HI and FT-HI approaches<sup>66</sup> exhibited dominant ferro-octupolar exchange, with subdominant quadrupolar interactions; these results were supported by later calculations using a multiorbital tight-binding model with interactions and SOC.<sup>240,285</sup> **Figure 4e** shows the temperature dependence of the total energy for the  $d^2$  series  $\text{Ba}_2\text{ROsO}_6$  ( $R = \text{Ca}, \text{Mg}, \text{Zn}$ ) obtained by mean-field solution of the MBEH<sup>66</sup>, highlighting the greater stability of the ferromagnetic octupolar order compared to the competing antiferromagnetic quadrupolar phase.

The competition between quadrupolar and octupolar order in these systems is highly susceptible to local and global strains that can break the crystalline point group symmetry.<sup>240</sup> Indeed, strong coupling of this non-Kramers doublet to phonons might favor local Jahn-Teller distortions and quadrupolar order,<sup>259</sup> potentially explaining the absence of an octupolar ordering transition in  $\text{Ba}_2\text{CdOsO}_6$ .<sup>273,286</sup> This competition is also revealed in NMR experiments<sup>266</sup> on  $\text{Ba}_2\text{Na}_{1-x}\text{Ca}_x\text{OsO}_6$ . Chemical substitution of  $\text{Na}^+$  with  $\text{Ca}^{2+}$  on the  $B$  site induces an effective electron transfer to the Na site<sup>277</sup>, accompanied by local structural modifications that alter the nature of the multipolar ground state<sup>31,266</sup>. Surprisingly, the system remains insulating throughout the entire doping range. The Mott gap is protected by the formation of Jahn-Teller polarons, which generate additional mid-gap SOC levels leading to the unusual coexistence of  $J = 3/2$  and  $J = 2$  states<sup>31</sup> (**Figure 4f**). In the  $x \sim 1$  regime, dilute impurities act as local probes of  $d^2$  multipolar order but also induce local strains that can locally pin quadrupoles and generate parasitic dipole moments tied to the octupolar ordering, providing an explanation for the observed NMR lineshapes (**Figure 4g**)<sup>70</sup>.

### [H3] $d^3$

Turning to the  $d^3$  double perovskites,  $\text{Ba}_2\text{YRuO}_6$ <sup>287–289</sup> and  $\text{Ba}_2\text{YO}_6$ <sup>149,290</sup>, though the half-filled  $t_{2g}$  configuration could be expected to quench the orbital angular momentum and suppress SOC effects, the observation of spin gaps in inelastic neutron scattering (INS)<sup>288,290</sup> and RIXS in  $\text{Ba}_2\text{YO}_6$ <sup>149</sup> suggests that this scenario is incorrect. Indeed, ab initio IEI Hamiltonians (equation 3) derived by the FT-HI method for  $\text{Ba}_2\text{YO}_6$  and  $\text{Ba}_2\text{YRuO}_6$  exhibit, apart from expected AFM Heisenberg couplings, also substantial dipole-octupole IEIs<sup>231</sup>. These terms scale as the square of spin-orbit coupling. They lift the continuous symmetry of the Hamiltonian, opening an excitation gap, as shown<sup>231</sup> by direct calculations of INS spectra of  $\text{Ba}_2\text{YO}_6$  (**Fig. 4h**) and  $\text{Ba}_2\text{YRuO}_6$  from the mean-field ground state of the IEI Hamiltonian. The theoretical excitation gap and shape of the INS spectra are in good qualitative agreement with the measurements<sup>289,290</sup>.

The same dipole-octupole IEIs are also predicted<sup>231</sup> to stabilize a non-collinear double- $\mathbf{q}$  magnetic structure in both compounds. At the same time, non-coplanar triple- $\mathbf{q}$  spin textures in  $\text{Ba}_2\text{YRuO}_6$  have been proposed<sup>289</sup> on the basis of INS measurements. The competition between various non-collinear orders stabilized by multipolar effects hence remains to be clarified.

### [H3] Vacancy-ordered double perovskites

We finally turn to the vacancy-ordered halide double perovskites  $\text{A}_2\text{MX}_6$ , a recently explored class of materials in the same category as  $\text{A}_2\text{BB}'\text{O}_6$  but with oxygen replaced by ligand halides  $X$  ( $=\text{Cl}, \text{Br}, \text{I}$ ) and B site ions replaced by vacancies. The set of  $d^1$  compounds  $\text{Cs}_2\text{MX}_6$  ( $\text{M}=\text{Ta}, \text{Nb}; \text{X}=\text{Cl}$ ),<sup>56,291,292</sup> shows evidence of weak moments,  $R\ln 4$  entropy, and ordering consistent with a  $J = 3/2$  ground state multiplet. Experiments on the  $d^2$  compounds  $\text{A}_2\text{WCl}_6$  ( $\text{A} = \text{Cs}, \text{Rb}, \text{CH}_3\text{NH}_3$ )<sup>293</sup> find evidence of strong deviations from the classic Kotani formula for the magnetic susceptibility, which has been theoretically ascribed, via exact diagonalization calculations, to the formation of a non-Kramers doublet and the resulting spin gap;<sup>294</sup> this scenario has been confirmed through more detailed cluster calculations.<sup>295</sup> The predicted ferro-octupolar order<sup>294</sup> in  $\text{A}_2\text{WCl}_6$  is yet to be experimentally confirmed. An important quantitative difference between these systems and the oxide double perovskites is their much narrower bandwidth, which results in weaker exchange interactions and lower transition temperatures for symmetry breaking.

There is recent evidence from RIXS on the  $d^5$  compound  $\text{K}_2\text{IrCl}_6$  that strong coupling to phonons results in a clear vibronic character of the  $J = 3/2$  excited states.<sup>296</sup> This suggests that coupling to the lattice may play an important role also in the vacancy-ordered halide double perovskites.

## [H2] $f$ -electron systems

$f$ -electron systems have been a traditional avenue to explore multipolar phases starting from the discovery<sup>297–299</sup> of a hidden-order phase transition in  $\text{NpO}_2$  in the 1960s, with multipolar orders identified and characterized in more than a dozen compounds<sup>73, 118, 121</sup>.

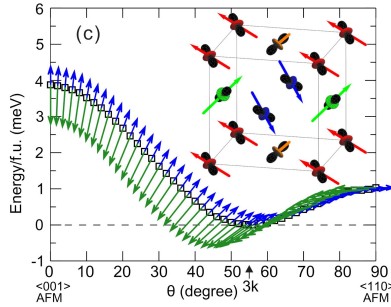
These systems have several distinctive features compared to their  $d$ -electron counterparts. The SOC is very strong, especially in the  $5f$ -electronic shell of actinides. Together with a pronounced spatial localization of  $f$ -electron atomic states, this leads to the SOC dominating over crystal-field effects. The latter can be considered as a perturbation splitting the energy levels of the ionic ground-state  $J$ -multiplet<sup>182</sup>. Unlike in  $d$ -electron materials, the crystal-field potential does not substantially reduce the  $f$ -shell orbital moment; hence, physics of the multipolar interactions, including electron-lattice ones, emerges rather often. Another important consequence of the strong localization and large SOC is that  $f$ -shells often keep their multiplet structure in solids and can be treated in the framework of standard crystal-field theory even in the metallic state. In this case, a well-localized  $f$ -electrons subsystem coexists with  $s$ - $d$  metallic conduction bands. This situation occurs in elemental rare-earth metals and their inter-metallic compounds. Therefore, multipolar effects can play an essential role even in metallic systems. Multipolar interactions are suggested to be behind the iconic hidden-order phase of  $\text{URu}_2\text{Si}_2$ <sup>214</sup> and to mediate superconductivity in  $\text{PrV}_2\text{Al}_{20}$ <sup>300, 301</sup>. It is generally well established that metallic rare-earth-based compounds, such as  $\text{DyB}_2\text{C}_2$ <sup>302, 303</sup>,  $\text{CeB}_6$ <sup>112</sup>,  $\text{CeAg}$ <sup>304</sup>,  $\text{TmCd}$ <sup>305</sup>, and  $\text{TmZn}$ <sup>306</sup> harbor hidden quadrupolar orders. Since quadrupolar moments of  $J$ -multiplets directly couple to the lattice via magneto-elastic coupling, ferroic and more complex antiferroic quadrupolar orders in rare-earths and actinides materials<sup>307</sup> can be identified and characterized by lattice strains and local atomic distortions below the quadrupolar phase transition. Early examples of the antiferroic quadrupole order detected in this way include measurements in  $\text{TbGa}_3$ <sup>308</sup> and  $\text{PrPb}_3$ <sup>309</sup>. Interestingly, spin-liquid phases associated with localized  $f$ -electrons multipolar degrees of freedom have also been reported for  $\text{Gd}_3\text{Ga}_5\text{O}_{12}$ <sup>310</sup> and  $\text{Ce}_2\text{Sn}_2\text{O}_7$ <sup>64</sup>.

## [H3] Actinide and praseodymium dioxides

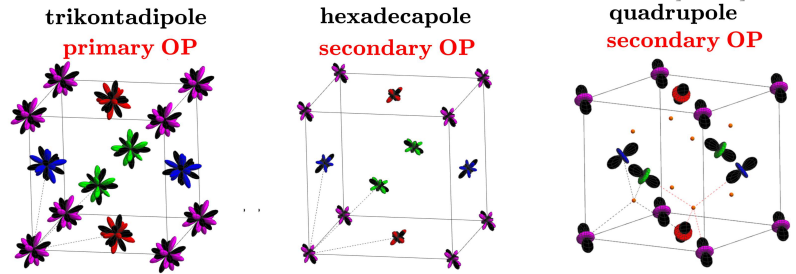
Actinide and praseodymium dioxides are prototypical cases of multipolar order and have been extensively investigated for decades by a combination of experiment, models and DFT calculations<sup>73, 118</sup>. We emphasize recent developments brought about by advanced ab initio methodologies.

Perhaps the most influential example of a magnetic system in which quadrupolar degrees of freedom play a fundamental role is  $\text{UO}_2$  with rutile crystal structure<sup>73</sup>. The U ions feature a spherically symmetric ground-state crystal-field triplet, which is well separated from excited crystal field levels. They occupy a geometrically frustrated fcc sublattice. For any set of antiferro dipolar IEIs, three different AFM structures – single, double and triple- $\mathbf{q}$  – have exactly the same energy and ordering temperature. Although  $\text{UO}_2$  exhibits conventional dipolar magnetic ordering at low temperatures, the secondary quadrupolar order parameter is believed to be the main reason for the stabilization of the observed<sup>311</sup> complex non-collinear  $3\mathbf{q}$  AFM structure with respect to competing non-collinear  $2\mathbf{q}$  and collinear  $1\mathbf{q}$  AFM orders. Several mechanisms have been proposed to explain the stabilization of the  $3\mathbf{q}$  magnetic structure by the secondary quadrupolar order parameter: via indirect quadrupolar exchange interactions mediated by phonons associated with oxygen vibrations<sup>312</sup>; purely electronic quadrupolar direct IEIs<sup>238</sup>; and due to static distortions of the oxygens in the magnetic ground state caused by interactions with quadrupolar degrees of freedom<sup>313</sup>. The effects of quadrupole-quadrupole IEIs in  $\text{UO}_2$  were investigated<sup>76</sup> using the DFT+U mean-field theory. Mapping magnetically constrained DFT total energies onto the model introduced in ref.<sup>238</sup> provided an ab initio confirmation of the important role of quadrupole-quadrupole interactions in establishing the  $3\mathbf{q}$  phase.<sup>204</sup> The corresponding non-collinear magnetic energy surface is displayed in **Fig. 5a**. However, the relative stability of the  $3\mathbf{q}$  ground state with respect to the other competing AFM phases is largely overestimated, a known issue in DFT<sup>65</sup>. This scenario has been confirmed by FT-HI calculations of the full IEI Hamiltonian,<sup>250</sup> predicting quadrupole-quadrupole

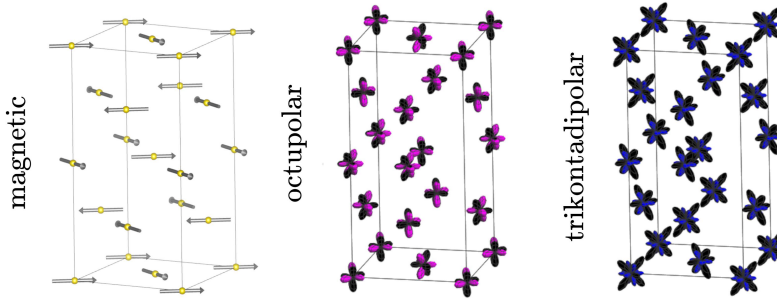
a -  $\text{UO}_2$



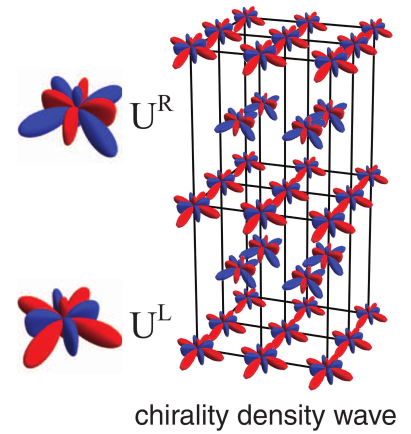
b -  $\text{NpO}_2$



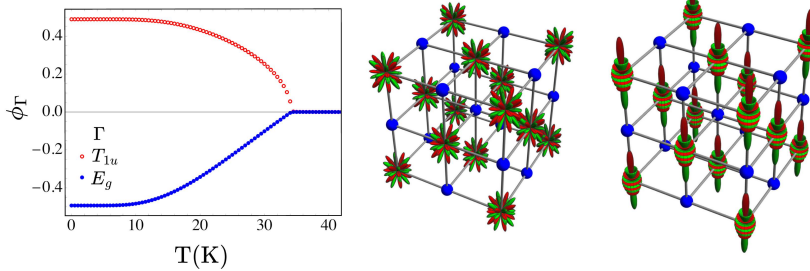
c -  $\text{PrO}_2$



d -  $\text{URu}_2\text{Si}_2$



e -  $\text{NdN}$



**Figure 5. Multipolar orders of f-electron systems computed with different methods** **a** | Total energy of  $\text{UO}_2$  vs magnetic canting angle along the  $1\mathbf{k}$  antiferromagnetic (AFM)  $\langle 001 \rangle \rightarrow 3\mathbf{k}$  AFM  $\rightarrow 1\mathbf{k}$  AFM  $\langle 110 \rangle$  transformation pathway calculated by constrained DFT+U<sup>204</sup>. Inset shows  $3\mathbf{k}$  quadrupolar and magnetic orders predicted from intersite exchange (IEI) Hamiltonian derived with the force theorem in Hubbard-I (FT-HI) method<sup>250</sup>. **b** | Non-zero order parameters (OP) in the "hidden-order" phase of  $\text{NpO}_2$  predicted from FT-HI IEI Hamiltonian<sup>67</sup>. The primary OP is a rank-5 (trikontadipolar) antiferromagnetic (left); charge hexadecapole (middle) and quadrupole (right) are secondary OPs. The quadrupole OP is scaled up by a factor of 4 to be visible. **c** | Dipole magnetic structure of Jahn-Teller-distorted  $\text{PrO}_2$  (left), which is induced by "hidden" octupole (middle) and trikontadipole (right) OPs. These OPs have been derived from an effective Hamiltonian comprising DFT+HI crystal-field splitting and FT-HI intersite exchange<sup>34</sup> **d** | Chiral hexadecapolar OP proposed<sup>161</sup> for the "hidden-order" phase of  $\text{URu}_2\text{Si}_2$  on the basis of DFT+dynamical mean-field theory (DMFT) calculations<sup>214</sup>. **e** | The primary and secondary OP vs  $T$  as well as the largest rank-7 (middle) and rank-9 (right) contributions into the primary OP in  $\text{NdN}$ <sup>86</sup>; the IEI Hamiltonian in this work was calculated by cluster quantum chemistry methods.

IEIs that favor the  $3\mathbf{q}$  order of both magnetic moments and quadrupoles (inset in **Fig. 5a**). However, the stabilization energy for this structure with respect to the predicted  $2\mathbf{q}$  one is extremely small (of the order of 0.01 meV). This casts doubt on the purely electronic quadrupole-quadrupole origin of the  $3\mathbf{q}$ -structure stabilization. The static distortions of the oxygen sublattice are also unlikely to stabilize the  $3\mathbf{q}$  structure, because the gain due to the magneto-elastic energy of the static Jan-Teller distortion is fully compensated by the elastic energy. Hence, such distortions cannot affect the ordering temperature of the competing magnetic phases, as it was shown almost 50 years ago for the  $1\mathbf{q}$  structure<sup>314</sup>. Therefore, the principal mechanism to resolve the magnetic frustration in  $\text{UO}_2$  still remains to be uncovered. Taking into account the importance of this material for nuclear technology,  $\text{UO}_2$  remains of high interest as an archetypal system for studying the physics of the multipolar order<sup>73</sup>.

High-rank multipolar IEIs can play a significant role also in the case of a conventional magnetic dipole order in an  $f$ -electron magnetic insulator. This situation occurs in  $\text{PrO}_2$ , where the experimentally observed<sup>315</sup> complex non-collinear AFM order (**Fig. 5b**) cannot be obtained from any physically reasonable choice of dipolar IEI,<sup>316</sup> suggesting the importance of multipolar couplings<sup>73</sup>. The full IEI matrix between nearest-neighbor  $J = 5/2$   $\text{Pr}^{4+}$  ions calculated by the FT-HI approach<sup>34</sup> exhibits leading couplings between high-rank multipoles. The main contribution to the magnetic ordering stems from those high-rank multipoles, with the corresponding contribution due to the conventional dipolar de Gennes IEIs being about three orders of magnitude lower. Hence, the experimentally observed magnetic order is entirely due to non-dipolar IEIs and arises as a slave order to the complex antiferromagnetic ordering of the octupoles and trikontadipoles<sup>34</sup> (**Fig. 5b**), to which it is entangled by the crystal-field potential. Recent calculations of multipolar IEIs<sup>34,86</sup> reveal their importance in the conventional magnetism of  $f$ -electron insulators. The standard paradigm<sup>182</sup> that the observed magnetic order is due to crystal field (or magnetic anisotropy) and relativistic de Gennes dipolar terms can be thus violated in many cases<sup>34</sup>.

The same ab initio FT-HI approach has also been applied to the purely multipolar order in the oldest known hidden-order material,  $\text{NpO}_2$ <sup>67</sup>. The calculated multipolar IEIs predict a complex antiferromagnetic trikontadipolar order together with small secondary quadrupolar and hexadecapolar order parameters (**Fig. 5c**). The magnetic dipolar moments cancel each other exactly on each Np site, in agreement with experiment<sup>317</sup>. Moreover, the predicted mutual orientation of the quadrupolar moments on different Np sites is such that the magnetoelastic coupling with the Np  $f$ -electrons subsystem is fully compensated for all oxygen atoms. As a result, the cubic lattice structure of  $\text{NpO}_2$  is preserved in the ordered phase. The predicted quadrupole structure also explains the lowering of the  $\text{NpO}_2$  symmetry detected by NMR measurements<sup>318</sup>. Indeed, the quadrupole lobes of neighboring Np either all point towards O (for two oxygen sites along one of the cube diagonals) or all point away from it (for six other oxygen sites in the unit cell, **Fig. 5c**). Hence, the local environment for these two sets of oxygen sites becomes different in the ordered phase, as detected by NMR probes. The structure of primary and secondary order parameters in  $\text{NpO}_2$  was previously inferred from the results of a battery of experimental probes<sup>82,82,83,318,319</sup>; it has been fully reproduced using the ab initio FT-HI Hamiltonian. It is also shown<sup>67</sup> that the volume dependence of the multipolar IEIs leads to spontaneous volume-striction effects, similarly to the ordinary two-site exchange magnetostriction effect in magnetic materials<sup>307</sup>. This multipolar magnetostriction accounts for a tiny uniform volume contraction observed in  $\text{NpO}_2$  below the hidden-order transition<sup>320</sup>.

### [H3] Rare-earth mononitrides

The importance of multipolar IEIs was also demonstrated for rare-earth mononitrides  $\text{RN}$ , where  $R$  is a rare-earth  $3+$  ion. They form a simple rock-salt structure and exhibit rather conventional ferromagnetic or antiferromagnetic orders<sup>321</sup>.

Nevertheless, kinetic IEIs obtained for  $R_2\text{N}_2$  complexes<sup>239</sup> using a cluster strong-coupling approach exhibit multipolar terms. This approach was subsequently applied to derive the full IEI interaction matrix between the  $\text{Nd}^{3+}$   $J = 9/2$  ground state multiplets in  $\text{NdN}$ . High-rank multipolar IEI contributions up to the 7th and 9th rank were shown to provide an important contribution to the thermodynamics of the  $\text{NdN}$  ferromagnetic phase; this contribution is comparable to that of the usual magnetic dipolar IEI<sup>86</sup>. The observed ferromagnetic phase in  $\text{NdN}$  is thus revealed to be a complex multipolar order (**Fig. 5d**) exhibiting substantial high-rank contributions to the order parameters together with the dipolar component.

### [H3] Heavy-fermion intermetallics

Ab initio theories of multipolar orders still find limited application in  $f$ -electron metals compared to  $f$ -electron insulators. DFT+U and DFT+DMFT approaches generalized to handle multipolar moments<sup>40,322</sup> have, however, been employed to treat those systems<sup>118,214</sup>. In general, DFT+U-based methods need to be properly constrained by occupation matrix control<sup>118</sup>. The predictive power of these schemes is still limited, because the proper symmetry breaking needs to be introduced from the onset. The space of possible multipolar order parameters is often too vast to explore all possibilities with such total energy calculations. Further advances in this area will be important to tackle outstanding problems like the true nature of the hidden order in the canonical URu<sub>2</sub>Si<sub>2</sub> compound, for which exotic chiral hexadecapolar orders are proposed (Fig. 5e),<sup>161,214</sup> as well as a number of alternative multipolar order parameters<sup>323</sup>, and an interplay between the multipolar order and the Kondo effect<sup>324</sup>.

### [H2] Magnetoelectric systems

We conclude the materials section by briefly touching on magnetoelectric systems<sup>91,155,325,326</sup>. Magnetoelectric materials exhibit a coupling between magnetic and electric properties, whereby an applied electric field induces a linear magnetization, and conversely, an applied magnetic field generates a linear electric polarization. In these systems, antisymmetric spin-orbit coupling associated with the simultaneous breaking of spatial inversion and time-reversal symmetries can induce odd-parity multipoles like second-order (quadrupole) moments of the magnetization density—a departure from the even-parity multipole orders hosted by single  $d$  or  $f$ -electron shells discussed thus far. Magnetoelectric multipoles, which have been briefly introduced in the section about methods, are central to the fundamental mechanisms of magnetoelectric coupling and have been extensively studied both theoretically<sup>71,119,156,327–330</sup> and experimentally<sup>80,331–336</sup>. The prototypical magnetoelectric material in which the connection between local magnetoelectric multipolar order and the atomic-scale magnetoelectric response has been studied is Cr<sub>2</sub>O<sub>3</sub><sup>337</sup>. This compound also exhibits an antiferroic arrangement of magnetoelectric multipoles, representing a distinct form of hidden order<sup>71</sup>. Moreover, fluctuating parity-odd multipoles have been proposed as potential order parameters in hidden-order phases of unconventional superconductors<sup>301</sup>, and may also play a role in the enigmatic hidden order phase observed in cuprates<sup>338</sup>.

### [H1] Conclusions and Future Perspectives

As emphasized in this Review, a vast space of competing order parameters and interactions in multipolar systems calls for approaches to construct their realistic low-energy effective Hamiltonians in an unbiased way. The progress in modeling insulating multipolar phases stems from recently formulated efficient ab initio methods that enable parameter-free, material-specific descriptions of hidden multipolar orders. Despite these successful applications, several open questions and challenges remain, requiring continued efforts.

Compared to insulating systems, the scope of ab-initio theories for metallic multipolar phases, such as  $f$ -electron heavy-fermion systems, remains more limited. The reason is that the force theorem<sup>75</sup> and cluster approaches<sup>86,239</sup> developed for high-rank intersite exchange interactions are based on a strong-coupling perturbative picture that becomes generally invalid in the metallic state. The conventional magnetic force theorem for symmetry-broken phases can be applied,<sup>72,339</sup> but it lacks a proper generalization to high-rank multipolar couplings. This limits its applicability not only for heavy-fermion systems, but also for multipolar intermetallics with localized  $f$  shells, where the kinetic exchange mechanism is enabled by the on-site Hund's rule coupling between localized and conduction electrons. In this respect, one may notice the promising development of extracting multipolar IEs from the full DMFT susceptibility<sup>234</sup>, as discussed, and its application to CeB<sub>6</sub><sup>235</sup>.

Unlike purely spin-dipole moments, multipolar orders such as the quadrupolar order can linearly couple to phonons, leading to new material functionalities via strain-tuning or driven phonons. This connects with the emerging area of “straintronics”<sup>340,341</sup> and “phononics”<sup>33,87,342</sup> in bulk oxides and 2D materials, enabling the use of strain or phonons to read and write information into multipolar degrees of freedom.

The coupling of phonons with doping-injected electrons and holes allows for charge localization, the breaking of crystal symmetry<sup>70</sup> and the formation of polarons<sup>31,343</sup>. This can lead to the coexistence of different high-rank

multipolar degrees of freedom, as reviewed for the case of  $J = 3/2$  and  $J = 2$  states in  $\text{Ba}_2\text{Na}_{1-x}\text{Ca}_x\text{OsO}_6$ <sup>31</sup>. The interaction between different types of local high-rank multipoles within the same system remains unexplored. Understanding these interactions as well as the interaction of polarons with the underlying hidden order could lead to the discovery of novel and exciting physics and will provide insights into the evolution of hidden phases upon doping<sup>266</sup>.

In the context of dilute impurities, isolated  $d$ -orbital ions or actinide  $f$ -orbital ions embedded at high-symmetry sites in cubic crystals may be viewed as single defects that support many sharp low-energy multiplets hosting a variety of multipolar moments. Certain multiplets, for example, have no dipole moments but only possess quadrupole and octupole moments. Such multiplets can thus act as sensors of electric field gradients, which couple linearly to quadrupoles, analogous to how an ordinary magnetic field couples to spin dipoles. At the same time, since quadrupole and octupole moments couple relatively weakly to actual magnetic fields, at  $\mathcal{O}(B^2)$  or  $\mathcal{O}(B^3)$  respectively, they are relatively immune to stray magnetic field noise. Embedded multipolar ions might also act as strain sensors, because their multiplet degeneracies can be broken in specific patterns by externally applied strain fields, which can couple linearly to the quadrupole moments. Multipolar defect centres can thus provide a rich new class of states for quantum sensing or for application as multipolar quantum bits, potentially extending the capabilities of existing systems such as nitrogen vacancy centers in diamond<sup>344–346</sup>.

Another intriguing aspect is the evolution of intersite exchange interactions from the bulk to the surface<sup>192, 347</sup>. Microscopic Hamiltonian approaches have demonstrated that the breaking of crystal symmetry at surfaces in multipolar materials can pin quadrupolar orders<sup>240</sup>, potentially giving rise to novel physics at crystal surfaces, interfaces, or within nanoparticles<sup>348, 349</sup>. This remains a largely unexplored field of research, requiring combined experimental and theoretical efforts. In particular, atomically resolved, surface-sensitive experimental techniques—such as (spin-dependent) scanning tunneling microscopy, atomic force microscopy, and ARPES—paired with first-principles atomistic simulations are crucial to unravel the surface geometry, identify possible reconstructions, and distinguish the electronic and magnetic properties.

Finally, the integration of artificial intelligence algorithms and data-driven approaches into many-body and materials physics<sup>350</sup> is driving a transformative shift in the field. Machine learning (ML) strategies have recently been employed to accelerate and optimize the study of magnetic phases within first-principles calculations<sup>351–357</sup>, as well as effective spin Hamiltonians and Monte Carlo simulations<sup>358–361</sup>. A range of ML architectures, including ML-based magnetic interatomic potentials<sup>351–354</sup>, deep neural networks<sup>355, 360</sup>, reinforcement learning<sup>358, 361</sup>, and Bayesian optimization<sup>357</sup>, has been applied to explore collinear and non-collinear magnetic phases, magnetic potential energy landscapes, and spin-spin interaction parameters. Although still in its early stages, this field is rapidly advancing and is expected to mature sufficiently to address high-rank intersite exchange interactions, (pseudo)spin excited states, temperature effects and spin-lattice unconventional interactions. By mapping ab initio theories onto efficient ML frameworks, these approaches have the potential to greatly enhance the exploration of complex magnetic interactions, such as those reviewed here, ultimately advancing our understanding of quantum magnetism and unconventional hidden orders.

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### Author contributions

C.F. conceived the initial idea for the review and coordinated its writing. L.V.P., C.F., and A.P. outlined the content. The initial draft was written by C.F. (Introduction), L.V.P. (Theory and Methods), A.P. (*d*-electron materials), and S.K. (*f*-electron materials). L.C. and D.F.M. contributed to drafting, compiled the tables, and designed the major figures. All authors participated in the final editing of the text and figures.

### Competing interests

The authors declare no competing interests.