Hidden order and multipolar exchange striction in a correlated f-electron system

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The nature of order in low-temperature phases of some materials is not directly seen by experiment. Such “hidden orders” (HOs) may inspire decades of research to identify the mechanism underlying those exotic states of matter. In insulators, HO phases originate in degenerate many-electron states on localized f or d shells that may harbor high-rank multipole moments. Coupled by intersite exchange, those moments form a vast space of competing order parameters. Here, we show how the ground-state order and magnetic excitations of a prototypical HO system, neptunium dioxide NpO2, can be fully described by a low-energy Hamiltonian derived by a many-body ab initio force theorem method. Supercritical interactions between the lowest crystalline quadruplets of Np4+ ions induce a primary noncollinear order of time-odd rank 5 (triakontadipolar) moments with a secondary quadrupole order preserving the cubic symmetry of NpO2. Our study also reveals an unconventional multipolar exchange striction mechanism behind the anomalous volume contraction of the NpO2 HO phase.

Strongly correlated electrons | hidden order | ab initio | linear response | effective Hamiltonians

Spontaneous symmetry breaking, or a phase transition, in extended systems is associated with emergence of a macroscopic order parameter, which is the statistical average over some physical observable. In some systems, the onset of order is clearly observed in the behavior of thermodynamic functions; however, the order parameter is not detectable by standard probes like neutron scattering or magnetic susceptibility measurements. In metals, such phenomena are typically associated with strongly correlated heavy-fermion behavior, as in the case of enigmatic URu2Si2 (1–3). In correlated insulators, hidden order (HO) phases typically originate in high-rank multipolar degrees of freedom on localized f- and d-electron shells. It was realized a long time ago (4) that in correlated magnetic insulators with strong spin-orbit coupling, apart from the ordinary Heisenberg interaction between localized spins, there may also exist intersite interactions coupling higher-order spin operators (magnetic multipoles). If those interactions are essentially large, this might lead to a new state of matter—a multipolar order without any associated magnetic order (5–7). Such purely multipolar orders have been observed in various f-electron (2, 8–11) and transition metal-correlated systems (12–14). Owing to numerous competing order parameters and small relevant energy scales, quantitative description of the HO represents a formidable challenge for theory. First-principles approaches to HO are typically restricted to simulations of a few likely phases inferred experimentally and do not attempt to explore the full space of possible multipolar orders (1, 15).

Difficulties of identifying the physical order parameter in a vast phase space of possible HOs are exemplified by the case of cubic NpO2 (8). A sharp second-order phase transition at \( T_D \approx 26 \text{K} \) was detected in NpO2 more than a half-century ago (16), with no evidence for underlying magnetic order and structural transformations (17–19), apart from a small anomalous contraction of the cubic unit cell volume observed (20) below \( T_D \). At the same time, NMR measurements (21) detect two inequivalent oxygen sites in the unit cell below \( T_D \), due to lowering of the cubic symmetry from high-temperature \( Fm\overline{3}m \) to \( Pn\overline{3}m \) by a longitudinal order of Np quadrupoles. However, a primary quadrupolar order parameter initially suggested (22) is excluded since muon spin-rotation measurements detect a nonzero magnetic density (23). Moreover, the crystal-field (CF) ground-state (GS) quadruplet \( \Gamma_8 \) is split in the HO phase into two singlets and a doublet, suggesting a time-odd primary order parameter (19, 24, 25). There is a multitude of possible high-rank odd order parameters realizable within the \( J = 9/2 \) ground-state multiplet of Np4+ \( (J = 3, 5, 7; \text{i.e., octupolar, triakontadipolar, etc.}) \). A lot of effort has been directed over the last two decades (21, 23, 24, 26–29) to identify a possible primary order able to reconcile various experimental facts. In particular, assumption of a triakontadipolar antiferromagnetic (AF) 3k order of the \( \Gamma_5 \) symmetry was shown (24, 25, 30) to lead to the best agreement with X-ray scattering and inelastic neutron scattering (INS) spectra. The same hypothesis is also strongly supported by estimates of the relative strength of odd \( \Gamma_5 \) multipole moments on the Np f3 shell in the presence of CF splitting (24, 31).

Although there is substantial experimental evidence to support the 3k rank 5 order in NpO2, the mechanism of its formation is still not well understood. The simplest possible form of the superexchange Hamiltonian, consisting of diagonal nearest-neighbor (NN) interactions between three \( \Gamma_5 \) triakontadipoles and between three dipole moments, has been employed in

Significance

Second-order phase transitions in solids occur due to spontaneous symmetry breaking with an order parameter continuously emerging from the disordered high-temperature phase. In some materials, the phase transitions are clearly detected in thermodynamic functions (e.g., specific heat), but the microscopic order parameters remain “hidden” from researchers, in some cases for decades. Here, we show how such hidden-order parameters can be unambiguously identified and the corresponding ordered phase fully described using a first-principles many-body linear response theory. Considering the canonical “hidden-order” system neptunium dioxide, we also identify an unconventional mechanism of spontaneous multipolar exchange striction that induces an anomalous volume contraction of the hidden-order phase in NpO2.
analysis of low-temperature susceptibility and INS data (24, 25). The full structure of this Hamiltonian cannot be extracted from experiment due to a large number of possible superexchange interactions (SEIs). The measured CF splitting of 55 meV between the $\Gamma_8$ ground state and first exited CF level (8, 9, 32) is much larger than $T_0$, suggesting SEI between $\Gamma_8$ states on neighboring Np ions as the origin of its exotic ordered phase. Such low-energy superexchange Hamiltonian has not been so far derived theoretically. Previous theoretical density functional theory (DFT) + U studies have confirmed the stability of a triakontadipolar order in NpO$_2$ (15, 33); however, they imposed an initial symmetry breaking consistent with the 3k rank 5 order inferred experimentally.

In this work, we apply a first-principles many-body framework to the problem of "hidden" multipolar orders in correlated insulators as exemplified by NpO$_2$. It consists of evaluating the full low-energy superexchange Hamiltonian from an ab initio description of the symmetric paramagnetic phase. We start with charge self-consistent DFT + dynamical mean-field theory (DMFT) (34–36) calculations for paramagnetic NpO$_2$ treating Np 5f within the quasaiatomic Hubbard-I (HI) approximation (37) (this method is abbreviated below as DFT + HI) to obtain its electronic structure and the composition of a CF-split Np 5f$^0$ shell. A force theorem within HI (FT-HI) approach (38) is subsequently employed to derive SEIs between the calculated CF $\Gamma_8$ ground-state quadruplets. In summary, we perform an ab initio electronic structure calculation of a complete superexchange Hamiltonian for high-rank magnetic multipoles in an f-electron crystalline material. By solving this Hamiltonian, we find a 3k rank 5 primary magnetic multipolar order (MMO) accompanied by a secondary longitudinal quadrupole order. The calculated time-odd splitting of $\Gamma_8$ and magnetic excitation spectra are in a good agreement with experiment. The lattice contraction effect (20) is shown to be not related to quadrupole ordering as suggested before (39) but rather, to stem from the volume dependence of leading time-odd SEIs. This multipolar "exchange spring" effect is quite unique and has not been, to our awareness, discussed previously in the literature. Overall, we show that within our first-principles scheme, which treats all competing order parameters on the equal footing, high-rank multipolar orders in correlated insulators can be captured both qualitatively and quantitatively.

**Results**

**CF Splitting and Superexchange Hamiltonian.** We start with evaluating the Np CF splitting in paramagnetic NpO$_2$ as discussed above, this splitting determines the space of low-energy states forming the MMO. The CF splitting of the Np 5f$^5$ ground-state multiplet $J = 9/2$ calculated by DFT + HI is shown in Fig. 1A. The ground-state $\Gamma_8$ quartet is separated from another excited $\Gamma_8$ quartet by 68 meV, in agreement with the experimental range for this splitting, 30 to 80 meV, inferred from INS measurements (19, 32, 40). The broad experimental range for the excited $\Gamma_8$ energy is due to the presence of a dispersive phonon branch in the same range [this overlap has been a major source of difficulties for the phenomenological analysis of NpO$_2$ in the framework of effective CF model model (8)].

Our calculated CF corresponds to $x = -0.54$ parameterizing the relative magnitude of the four- and six-order contributions to the cubic CF (41). Our values are in good agreement with $x = -0.48$ inferred from INS measurements (19) and analysis of CF excitation energies along the actinide dioxide series (31). The CF-level energies calculated within DFT + HI are

![Fig. 1. Np 5f on-site splitting and Np–Np intersite interactions in NpO$_2$.](https://doi.org/10.1073/pnas.2025317118)
also in good agreement with previous DFT + DMFT calculations by Kolorenč et al. (42) employing an exact diagonalization approach to Np 5f. The calculated wave functions (WFs) of the CF ground-state Γ₈ quartet (SI Appendix, Table S1) feature a small admixture of the excited J = 11/2 and 13/2 multiplets.

The space of CF GS Γ₈ quadruplet is conveniently represented by the effective angular momentum quantum number Jₐf = 3/2, with the Γ₈ WFs labeled by the corresponding projection M = −3/2,..., +3/2 and the phases of WFs chosen to satisfy the time-reversal symmetry (SI Appendix, Table S1). The on-site degrees of freedom within the GS quadruplet are (pseudo-)dipole, quadrupole, and octupole moments defined for Jₐf = 3/2 in the standard way (8). Hence, the most general form for a superexchange coupling between Γ₈ quadruplets on two Np sites reads

\[ \sum_{KK'Q} V^{QQ'}_{KK'}(R) \hat{O}_K^{Q}(R_0) \hat{O}_K^{Q'}(R_0 + R). \]  

where \( \hat{O}_K^{Q}(R_0) \) and \( \hat{O}_K^{Q}(R_0 + R) \) are the real spherical tensor operators (8) for J = 3/2 of rank K = 1, 2, or 3, −K ≤ Q ≤ K, acting on the sites R₀ and R₀ + R, respectively. \( \hat{V}^{QQ'}_{KK'}(R) \) is the SEI that couples them, and to the translational invariance, \( \hat{V}^{QQ'}_{KK'} \) depends only on the intersite lattice vector \( R \).

We employed the FT-HI approach (38) to evaluate all interactions \( \hat{V}^{QQ'}_{KK'}(R) \) from the converged DFT + HI NpO₂ electronic structure for the first Np coordination shells. Only NN SEIs are significant, with longer-distance ones being more than an order of magnitude smaller. All SEIs \( \hat{V}^{QQ'}_{KK'}(R) \) for a given bond R form a matrix, designated below as the SEI matrix, with the rows and columns labeled by the moments K and K' Q' on the sites R₀ and R₀ + R, respectively. The NN SEI matrix \( \hat{V}(R) \) is graphically represented in Fig. IB (SI Appendix, Table S2) using a local coordinate system with the quantization axis z[R]. This 15 × 15 matrix is of a block-diagonal form since the interactions between time-even and time-odd moments are zero by symmetry. It can thus be separated into the dipole–dipole (DD), quadrupole–quadrupole (QQ), octupole–octupole (OO), and dipole–octupole (DO) blocks. Despite this simplification, the SEI matrix \( \hat{V} \) in principle contain 70 distinct elements. The number of distinct nonzero matrix elements in \( \hat{V} \), while reduced by the cubic symmetry to 38, remains rather large.

Our calculations predict the largest values for the diagonal DD x × x (AF, 1.6 meV) SEI. However, the off-diagonal OO zyg to \( y(x^2 − 3yz) \) (ferro, −1.5 meV) and DO z to \( z^2 \) (AF, 0.95 meV) couplings are of about the same magnitude as the DD x × x one. Overall, the calculated \( \hat{V} \) matrix shown in Fig. 1B features many nonnegligible DD, OO, and DO interactions of a comparable magnitude. The QQ interactions are weaker, reflecting the secondary nature of the quadrupole order. Our calculations thus predict a complex and frustrated superexchange in NpO₂, which may give rise to multiple competing time-odd orders. Therefore, as has been previously noted (8, 25), extracting a full superexchange Hamiltonian of NpO₂ from experimental (e.g., INS) data is virtually impossible due to a large number of parameters entering into the fit of an excitation spectra. The same difficulty is encountered by ab initio approaches based on total energy calculations for symmetry broken phases (15, 33, 43), which require a large number of very precise calculations to extract multiple nonnegligible matrix elements of \( \hat{V} \), with the magnitude of 0.5 meV and above. Within the present framework, all interactions are extracted from a single ab initio calculation for paramagnetic NpO₂.

Ordered State of NpO₂. The calculated superexchange Hamiltonian for NpO₂ reads

\[ H_{SE} = \frac{1}{N} \sum_{R \in \text{NN}} \hat{O}(R_0) \hat{V}(R) \hat{O}(R_0 + R), \]  

where \( \hat{R} \) runs over all NN bonds in the Np face-centered cubic (fcc) sublattice and \( N \) is the number of Np sites; we also introduced the obvious vector notation \( \hat{O} = [O_{\uparrow}, O_{\downarrow}, \ldots, O_{\uparrow}] \) for multipole tensors. The SEI matrix \( \hat{V}(R) \) in the local frame \( \hat{R} \) is (Eq. 1, see also Fig. 1B) rotated to align the corresponding Np NN bond. We solved [2] numerically within the mean-field (MF) approximation (44), obtaining a second-order transition at \( T_\text{N} = 37 \text{K} \), in good agreement with its experimental value of 26 K taking into account the usual MF overestimation of ordering temperatures. The numerical results were verified by a linearized (MF) theory, derived by a first-order expansion of MF equations in the order parameters \( \hat{O}(\hat{R}) \) (Methods).

The resulting GS order of NpO₂ in the \( J_{\text{eff}} = 3/2 \) space consists of a primary (pseudo-)DO order combined with secondary (pseudo-) quadrupole one (SI Appendix, Table S3 lists the values of all \( J_{\text{eff}} \) order parameters; see also SI Appendix, Fig. S1). The pseudodipole order is a complex 3k planar AF structure, with four inequivalent single-cubic sublattices forming two pairs with different moment magnitude; the moments of those two pairs are aligned along the \((1,1,0)\) and \((3,1,0)\) directions in the fcc lattice, respectively. The origin of this inclined AF structure of pseudodipoles is in the DO SEI, similar to physical AF magnetic orders found in some materials with large spin-orbit coupling that are likely induced by higher-order magnetic multipolar interactions. The pseudotetrapoles is also oriented in nonsymmetrical directions. The secondary pseudotetrapole order is of a 3k type, which we analyze in detail below.

We subsequently mapped the moments calculated in the \( J_{\text{eff}} = 3/2 \) space into the observable multipolar moments that are defined in the physical \( J = 9/2 \) space of Np 5f³ GS multiplet (Methods). The calculated physical multipolar order of NpO₂ is displayed in Fig. 2. Notice that observable moments up to \( K = 7 \) can exist on an f-electron shell (8); we show the largest primary (odd) and secondary (even) order parameters as well as the physically important quadrupole order. All nonzero multipole moments are listed in SI Appendix, Table S4. The physical dipole magnetic moments are found to completely vanish since their contribution into the \( \langle O_3 \rangle \) and \( \langle O_4 \rangle \) pseudodipole \( J_{\text{eff}} = 3/2 \) moment is exactly canceled by that due to the \( \langle T_5 \rangle \) and \( \langle T_7 \rangle \) pseudotetrapoles. The primary order parameter is of rank 5 (triaxialdipole); the physical octupole moment is an order of magnitude smaller, and the magnitude of rank 7 moments is about 1/3 that of triaxialdipoles, in agreement with previous estimates for the relative contribution of those multipoles into the MMO order in NpO₂ (24). The magnetic triaxialdipole on different sublattices are oriented in four different directions (forming mutual angles corresponding to the angles between the cube’s main diagonals), thus structured similarly to the 3k–AFM dipole order in UO₂ (Fig. 2A). The secondary order is dominated by hexadecapole (rank 4) (Fig. 2B). The ordered quadrupole moments (Fig. 2C) are roughly twice smaller. The quadrupole order is directly related to the pseudotetrapole one since the \( \Gamma_5 \) (or \( \Gamma_{25} \)) pseudotetrapoles directly map into the physical ones, apart from swapping \( xy \leftrightarrow yz \) and \( x^2 + y^2 \leftrightarrow xxz \). The resulting physical quadrupole order can be represented, in the space of \( \Gamma_5 \) quadrupoles \([O_{xx}, O_{xy}, O_{xz}] \) by four directions \([111], [111], [111], \) and \([111]\) for four inequivalent Np sublattices \([0,0,0], [1/2,1/2,0], [1/2,0,1/2], \) and \([0,1/2,1/2]\), respectively. These quadrupoles can be depicted as \( \langle O_{3z} \rangle \) moments with the principal axes \( z \) along the corresponding direction at each given...
site (Fig. 2C). One sees that the ordered quadrupoles on the four Np sites forming the tetrahedron around each oxygen can either have their principal axes directed along the Np–O bonds toward the central O or form the same angle of 70.5° with respect to those bonds. In both cases, the tetrahedron symmetry is preserved. The first case is realized for two O along one of the principal cubic diagonals, while the second one is found for six others, resulting in the lowering of NpO$_2$ symmetry to Pn3$_m$ from Fm3$_m$ without any distortion of the cubic structure. This longitudinal 3k quadrupole order, previously proposed on the basis of resonant X-ray scattering (30) and subsequently confirmed by the splitting of $^{17}$O-NMR spectra in ordered NpO$_2$ (21), is thus predicted by our ab initio superexchange Hamiltonian (2).

Exchange Splitting and Magnetic Excitations. Having obtained the MMO of NpO$_2$, we subsequently calculated its excitation spectra, which have been previously measured, in particular, by INS (19, 25).

The MMO lifts the degeneracy of CF GS $\Gamma_5$ quartet; the resulting exchange splitting calculated from the ab initio superexchange Hamiltonian (2) is depicted in Fig. 1 A. Right. The ground state is a $\Gamma_5$ singlet with the first excited doublet found at 6.1 meV above the GS $\Gamma_5$ singlet and the second excited level, singlet, located at 12.2 meV. The calculated position of first excited doublet is in excellent agreement with the location of a prominent peak in INS spectra at about 6.4 meV (19) in the ordered phase; another broad excitation was observed in the range of 11 to 18 meV (25) (see below).

Previously, an exchange splitting of the $\Gamma_8$ was obtained in ref. 24 assuming a diagonal uniform SEI between $\Gamma_5$ triakontadipole. The value of this SEI was tuned to reproduce the relative magnitude of the principal momenta indicates their relative magnitude. Black dashed lines are the primitive lattice translations of the original Np cubic face-centered sublattice, which, in the ordered phase, connects inequivalent Np sublattices. (C) Longitudinal 3k order of slave quadrupole moments, which are scaled on the plot by a factor of four. Np–O bonds for two inequivalent O sites (small orange balls) in ordered NpO$_2$ are indicated by dashed lines. One sees that in the Np tetrahedron around each O, all ordered Np quadrupoles either are directed along the Np–O bonds or form the same angle with those bonds. This longitudinal 3k quadrupolar order preserves the cubic positions of oxygen sites in the ordered phase of NpO$_2$, in contrast to the transverse 3k quadrupole order in UO$_2$. 

ref. 24. The $S(q, E)$ structure is richer along other directions showing multiple branches in the $E$ range from 4 to 8 meV. The experimental INS spectrum $S(q, E)$ has not been measured so far due to lack of large single-crystal samples (8); hence, our result represents a prediction for future experiments. The calculated spherically averaged INS cross-section $\int S(q, E)dq$ is compared in Fig. 3B with that measured (25) on powder samples at the same $|q|=2.5$ Å$^{-1}$. The theoretical INS spectrum exhibits prominent peaks at around 6 meV and at about 12.5 meV, corresponding to the transition from the $\Gamma_5$ GS to the excited doublet and singlet, respectively.

The theoretical low-energy feature agrees very well with the measured INS spectra, after experimental broadening is taken into account (Fig. 3B). The high-energy feature, however, is clearly split in experiment into two broad peaks centered at about 12 and 16 meV. In order to understand whether the relative weights of the low- and high-energy features are captured in the theory, we employed the same analysis as Magnani et al. (25). Namely, we evaluated, as a function of the momentum transfer, the ratio of high-energy feature spectral weight to that of the low-energy one. The calculated ratio is in excellent agreement with experiment up to $|q|=2.5$ Å$^{-1}$. As noticed in ref. 25, a phonon contribution to INS appears below 18 meV for large $|q|$, thus rendering the separation of magnetic and phonon scattering less reliable for $|q|>2$ Å$^{-1}$. The splitting of high-energy peak was clearly observed at all measured $|q|$; it was not reproduced by the simplified semiempirical SEI employed by Magnani et al. (25). They speculated that this splitting might stem from complex realistic Np–Np SEIs, which they could not determine from experiment. In the present work, we determined the full superexchange Hamiltonian for the GS CF $\Gamma_5$ quadruplet. Hence, the fact that the splitting of INS high-energy feature is still not reproduced points out to its origin likely being a superexchange coupling between the ground-state and first excited $\Gamma_5$ quadruplets (a significant contribution of the very high-energy $\Gamma_5$ CF doublet is unlikely). This interquadruplet coupling can be in principle derived using the present framework; we have not attempted to do this in the present work.

Alternatively, lattice-mediated interactions might be also considered as the origin of the high-peak splitting. Those interactions couple time-even moments (i.e., the quadrupoles within the $J_{QG}$ space). However, the QQ coupling is rather expected to impact the shape of the low-energy peak in Fig. 3B because the corresponding lowest on-site excitation in the ordered phase
the volume contraction and volume constants calculated for NpO$_2$ elastic energies. To that end, we adopted the elastic constants below, this is not the case, and the volume contraction in NpO$_2$ cannot be attributed to the conventional volume magnetostriction high-rank magnetic multipoles.

Rather stems from the volume-dependent SEI coupling high-rank quadrupole order coupling to the lattice. As we show below, this is not the case, and the volume contraction in NpO$_2$ rather stems from the volume-dependent SEI coupling high-rank magnetic multipoles.

In order to make a quantitative estimation for this anomaly, we evaluated the volume dependence of NpO$_2$ ordering and elastic energies. To that end, we adopted the elastic constants calculated for NpO$_2$ in the framework of the DFT + U approach (45): $C_{11} = 404$ GPa, $C_{12} = 143$ GPa. The corresponding parabolic volume dependence of elastic energy $E_{\text{elas}} = 1/3 (C_{11}/2 + C_{12}) \epsilon^2 \equiv K \epsilon^2$, where $\epsilon = (V - V_0)/V_0$ is the volume contraction and $V_0$ is the NpO$_2$ equilibrium volume (45), is depicted in Fig. 4. The dependence of MMO energy vs. volume was obtained by calculating the SEIs at a few different volumes and then evaluating the MF order and ordering energy vs. volume expansion or contraction. The superexchange ordering energy remains linear vs. $\epsilon$ in a rather large range ($\epsilon = \pm 1\%$); its dependence upon $\epsilon$ for the relevant range of small $\epsilon$ is thus easily obtained.

As is seen in Fig. 4, the superexchange contribution shifts the equilibrium volume in the ordered state toward smaller volumes. The negative slope for the ordering energy vs. volume is expected as the SEIs become larger with decreasing Np–Np distance. Thus, the multipolar SEIs act as springs (scheme in Fig. 4, Inset) pulling Np atoms closer as the order parameters increase below $T_0$. Our approach is thus able to qualitatively capture this very small in magnitude subtle effect: The calculated spontaneous multipolar exchange striction is 0.023\% (Fig. 4) at zero $T$ as compared with the experimental estimate of 0.018\% (20, 39).

We also performed the same calculations suppressing the QQ order parameter, obtaining only a minor change, by about 0.5\%, in the slope of MMO energy vs. volume. Hence, the secondary quadrupole order plays virtually no role in the anomalous volume contraction. The physical origin of this effect is the volume dependence of leading, time-odd SEIs.

To analyze the temperature dependence of the anomalous contraction, one may recast the linear in volume MMO energy (Fig. 4) into a general form of $K_{\text{sec}} \xi_T^2(T) + K_{\text{pr}} \xi_T^2(T)$, where $\xi_T(T)$ and $\xi_{\text{sec}}(T)$ are primary and secondary order parameters, respectively, and $K_{\text{pr}}$ and $K_{\text{sec}}$ are the slopes of volume dependence for the corresponding contributions to MMO energy. The temperature dependence of the anomalous volume contraction is thus given by that of squares of the order parameters, $\xi_T^2(T)$ and $\xi_{\text{sec}}^2(T)$. As shown in SI Appendix, Fig. S2, secondary quadrupole $\xi_{\text{sec}}^2(T)$ exhibits a smooth evolution across $T_0$ and rather slowly increases vs. decreasing $T$, while primary time-odd $\xi_T^2(T)$ features a discontinuity in the slope at $T_0$, as expected, with a rapid growth for $T < T_0$, reaching about 60\% of total magnitude at $3/4 T_0$. This behavior of $\xi_T^2(T)$ is in a perfect agreement with the shape of temperature dependence of the volume anomaly (20), thus confirming that it is induced directly by the time-odd primary order.

Fig. 3. INS spectra in ordered NpO$_2$. (A) INS cross-section $S(q, E)$ with the momentum transfer $q$ along a high-symmetry path in the fcc BZ. The special points are $\Gamma = [0,0,0]$, $X = [1,0,0]$, $W = [1,1,0]$, $K = [\frac{11}{2}, \frac{11}{2}, \frac{11}{2}]$, in units of $2\pi/\alpha$. (B) Powder (spherically averaged) INS cross-section vs. energy transfer $E$ for $|q| = 2.5$ Å$^{-1}$. The theoretical spectra were broadened with the Gaussian resolution function of 1.5 meV. The experimental points are from Magnani et al. (25). (C) Ratio of the spectral weights of the low-energy (around 6-meV) and high-energy (10- to 20-meV) features vs. momentum transfer in the powder INS spectra (B). The experimental points are from Magnani et al. (25). Theoretical points from the same work are calculated with a semiempirical superexchange Hamiltonian (in the text).
Discussion

In conclusion, we have applied an advanced ab initio framework to the HO phase of neptunium dioxide NpO₂. Our framework is based on the density functional + DMFT (DFT + DMFT) in conjunction with a quasistatic approximation to local correlations on Np 5f. Its crucial part is a force theorem method (38) that we employ to calculate SEIs between all multipole moments of the Np f³ lowest CF manifold. From the resulting superexchange Hamiltonian, we derive all order parameters of the HO phase, its ordering temperature, magnetic excitations, and volume effect. In fact, numerous properties of the NpO₂ HO phenomenon that have been painstakingly determined in experiments over several decades—absence of conventional magnetic order, the CF-level scheme, the primary triadecapole order and secondary longitudinal 3k quadrupole order, the singlet–doublet–singlet exchange splitting of the CF ground state, the two-peak structure of the magnetic excitations, and volume effect. In practice, numerous properties of the NpO₂ HO phenomenon that have been painstakingly determined in experiments over several decades—absence of conventional magnetic order, the CF-level scheme, the primary triadecapole order and secondary longitudinal 3k quadrupole order, the singlet–doublet–singlet exchange splitting of the CF ground state, the two-peak structure of the magnetic excitations spectra—are reproduced by our calculations that contain essentially no adjustable parameters. Therefore, the present scheme is shown to provide a full, parameter-free, and quantitatively correct description of superexchange in complex realistic correlated insulators. On the basis of our theory, we may also identify the features that are not stemming from the intersite coupling between the ground-state CF levels. For example, the splitting of the high-energy peak in NpO₂ INS spectra likely stems from an SEI with excited CF levels. We also uncover an unconventional mechanism for the anomalous volume contraction observed in ordered NpO₂, which is induced not by the secondary quadrupole order coupling to the lattice as previously assumed but rather, by the volume dependence of leading time-odd SEIs.

This first-principles methodology—a dynamical MF treatment of symmetric paramagnetic phase combined with the force theorem extraction of the full complex intersite exchange responsible for the spontaneous symmetry breaking—can be applied to a wide range of rare earth, actinide, and heavy-transition metals correlated systems (2, 8, 9, 46), in which the interplay of a large spin-orbit coupling with crystalline environment gives rise to a large degeneracy of the CF ground state and high-rank multipole moments. HOs, stemming from coupling between those moments, can be predicted and their interplay with various parameters—external or chemical pressure, applied field, lattice distortions—identified, thus opening up an avenue for theoretical search of new exotic phases of matter.

Methods

DFT + Hi First-Principles Calculations. Our charge self-consistent DFT + DMFT calculations using the Hi approximation for Np 5f, abbreviated as DFT + Hi, were carried out for the CaF₂-type cubic structure of NpO₂, with the experimental lattice parameter a = 5.434 Å. We employed the Wien-2k full-potential code (47) in conjunction with "TRIQS" library implementations for the DMFT cycle (36, 48) and Hi. The spin-orbit coupling was included in Wien2k within the standard second-variation treatment. The Brillouin zone (BZ) integration was carried out using 1,000 k points in the full BZ, and the local density approximation (LDA) was employed as DFT exchange correlation potential.

The Wannier orbitals representing Np 5f states were constructed by the projective technique of refs. 49 and 50 using the Kohn–Sham bands enclosed by the energy window [−2.04; 2.18] eV around the Kohn–Sham Fermi energy; this window thus encloses all Np 5f-like bands. The use of a narrow window enclosing only the target (5f-like) band for the construction of local orbitals results in a so-called "extended" Wannier basis. By employing such a basis within DFT + Hi, one may effectively include the contribution of hybridization to the CF splitting on localized shells, as discussed in ref. 51. The same choice for the local 5f basis was employed in our previous DFT + Hi study (52) of UO₂, resulting in a good quantitative agreement of the calculated CF splitting with experiment.

The on-site Coulomb interaction between Np 5f was specified by the Slater parameter U₀ = 4.5 eV and the Hund's rule coupling J₅ = 0.6 eV; the same values were previously employed for UO₂ (52). The double-counting correction was computed using the fully localized limit (53) with the atomic occupancy of Np 5f shell (54). The DFT + DMFT charge self-consistency was implemented as described in ref. 55. In our self-consistent DFT + Hi calculations, we employed the spherical averaging of the Np 5f charge density, following the approach of Delange et al. (56), in order to suppress the contribution of LDA self-interaction error to the CF. The DFT + Hi calculations were converged to 5 μRy in the total energy.

CF and SEIs. The self-consistent DFT + Hi calculations predict a CF split ⁴I₉/₂ atomic multiplet to have the lowest energy, in agreement with Hund’s rules for an f⁶ shell; the calculated spin-orbit coupling λ = 0.27 eV. The CF splitting of the ⁴I₉/₂ multiplet predicted by these calculations is shown in Fig. 1A,
and the corresponding CF WFs are listed in SI Appendix, Table S1. The cubic CF parameters—\( A_0(0^2) = -152\) meV, \( A_0(1^2) = 5A_0(0^2) \), \( A_0(2^2) = 32.6\) meV, and \( A_0(3^2) = 21A_0(0^2) \)—were extracted by fitting the converged DFT + Hi one-electron Si-level positions (56).

The states of the CF ground-state quadruplet \( \Gamma_8 \) were labeled by projection \( M \) of the pseudoangular quantum number \( J_M = 3/2 \) as specified in SI Appendix, Table S1. We subsequently employed the FT-HI method (38) to evaluate all SEIs between the \( J_M = 3/2 \) quadruplet for several first Np–Np coordination shells. Previously, the FT-HI method was applied to systems with conventional magnetic primary order (52, 57). Within this method, matrix elements of intersite coupling \( V^{\alpha\beta}_{\mathbf{R}} \) for the Np–Np bond \( \mathbf{R} \) read

\[
\langle M_i M_j | V^\alpha_{\mathbf{R}} | M_i M_k \rangle = \text{Tr} \left[ \frac{\delta_{\mathbf{R}_0 + \mathbf{R}, \mathbf{R}_0}}{\mathbf{G}_{\mathbf{R}_0} \mathbf{M}_{\alpha \beta} \mathbf{G}_{\mathbf{R}_0} \mathbf{M}_{\alpha \beta}} \right],
\]

where \( \rho_{M_i M_j}^{\alpha\beta} \) is the corresponding element of the \( J_M = 3/2 \) density matrix on site \( \mathbf{R}_0 \), \( \mathbf{M}_{\alpha \beta} \) is the derivative of atomic (Hi) self-energy \( \Sigma_{\mathbf{R}_0}^{\alpha\beta} \) over a fluctuation of the \( \mathbf{M}_{\alpha \beta} \) element, and \( \mathbf{G} \) is the intersite Green’s function for the Np–Np bond \( \mathbf{R} \) evaluated within the DFT + Hi. After all matrix elements (3) are calculated, they are transformed to the couplings \( V_{\mathbf{R}^\alpha\beta}^{\gamma\delta} \) between onsite moments (1) as follows:

\[
V_{\mathbf{R}^\alpha\beta}^{\gamma\delta} = \sum_{\mathbf{M}_i \mathbf{M}_j \mathbf{M}_k \mathbf{M}_l} \langle \mathbf{M}_i \mathbf{M}_j | V^\alpha_{\mathbf{R}} | \mathbf{M}_k \mathbf{M}_l \rangle \langle \mathbf{M}_k \mathbf{M}_l | V^\beta_{\mathbf{R}} | \mathbf{M}_i \mathbf{M}_j \rangle.
\]

where \( \mathbf{M}_{\alpha \beta} \) is the \( \mathbf{M}_{\alpha \beta} \) matrix element of the real spherical tensor defined in accordance with equation 10 in Santini et al. (8). The SEI matrix \( V \) shown in Fig. 18 was subsequently obtained by rotating calculated \( V \) by 45° about the [010] axis, thus aligning one of the NN bonds with \( Z \).

**MF Calculations and Analysis of Order Parameters.** We solved the obtained superexchange Hamiltonian (2) using the numerical MF package MCFPHASE (44) including all 1k structures up to \( 4 \times 4 \times 4 \) unit cells. We have also verified the numerical solution by an analytical approach. Namely, the MF equations read

\[
\langle \hat{O}_\mu \rangle = \frac{1}{Z} \text{Tr} \left[ \hat{O}_\mu \exp(-i \hat{H}_{\text{MF}}) \right],
\]

where \( H_{\text{MF}} = \sum_{\mathbf{R}_0} \hat{O} \mathbf{V}_{\mathbf{R}_0} \hat{O} (\mathbf{R}_0) \) is the MF Hamiltonian for sublattice \( a, \mathbf{V}_{\mathbf{R}_0} \) is the SEI matrix between sublattices \( a, b \) and \( \mathbf{V} = \{ \mathbf{V} = \{ \sum_{\mathbf{R}_0} \mathbf{V}_{\mathbf{R}_0} \} \} \) with \( \mathbf{R}_0 \in a \), \( Z = \text{Tr} \left[ \exp(-i \hat{H}_{\text{MF}}) \right] \), and \( \beta = 1/T \). Expanding the RHS of (5) to the linear order in \( \langle \hat{O} \rangle \) and allowing for four inequivalent simple cubic Np sublattices, which is a minimal number needed to cover all possible ordered states on fcc lattice with an NN coupling (58), one obtains a system of 60 linear equations. Solutions of the linearized MF equations unambiguously identify primary order parameters. With both numerical and analytic MF approaches, we obtained the highest ordering temperature \((T = 37 K)\) and lowest free energy for the primary DO order displayed in SI Appendix, Fig. S1.

This order in the \( J_M = 3/2 \) space was subsequently mapped into the physical \( J = 9/2 \) space. The \( J_M \) density matrix on the inequivalent site \( a \) reads

\[
\rho_{J_M}(\mathbf{R}) = \sum_{\mathbf{R}_0} \rho_{J_M}(\mathbf{R}_0) \exp(-i \hat{H}_{\text{MF}}),
\]

where \( \rho_{J_M}(\mathbf{R}_0) \) are the corresponding pseudo-Jmoments. This density matrix is upfolded into the one in the \( J = 9/2 \) space, neglecting small contributions of the excited \( J = 11/2 \) and \( 13/2 \) multiplets and renormalizing the \( \Gamma_8 \) CF states accordingly, as \( \rho_{J_M}(\mathbf{R}_0) \) are the corresponding CF wavefunctions, \( \langle \hat{O} \rangle \) the CF GS basis, with WFs written as columns in the order of \( M = -J_M \ldots J_M \). The physical \( J = 9/2 \) moments are then calculated in the standard way.

\[
\langle \hat{O}(\mathbf{R}) \rangle_a = \text{Tr} \left( \rho_{J_M}(\mathbf{R}) \hat{O} \right).
\]

**Calculations of Dynamical Susceptibility.** In order to calculate the dynamical magnetic susceptibility \( \chi_{\alpha\beta}(q, \omega) \), we implemented a general RPA approach (59). Namely, the general susceptibility matrix in the \( J_M = 3/2 \) space reads

\[
\chi_{\alpha\beta}(q, \omega) = \sum_{\mathbf{R}} \langle \hat{O}(\mathbf{R}+\mathbf{q}) \hat{O}^\dagger(\mathbf{R}) \rangle_a \chi_{\alpha\beta}(q, \omega) = \sum_{\mathbf{R}} \langle \hat{O}(\mathbf{R}+\mathbf{q}) \hat{O}^\dagger(\mathbf{R}) \rangle_a \chi_{\alpha\beta}(q, \omega),
\]

where \( \chi_{\alpha\beta}(q, \omega) \) is the local bare susceptibility, \( \hat{O} \) is the Fourier transform of SEI matrices \( \mathbf{V} \), the bar ... designates a matrix in combined \( [a, \alpha] \) indices, where a labels inequivalent sublattices, and \( \alpha = [\delta, \delta] \) labels multiples. The local susceptibility \( \chi_{\alpha\beta}(q, \omega) \) for the inequivalent site \( a \) is calculated from the MF eigenvalues \( E^a \) and eigenstates \( |\psi^a\rangle \)

\[
\chi_{\alpha\beta}(q, \omega) = -\sum_{\omega \neq E^a} \langle \psi^a | \hat{O}(\mathbf{R}+\mathbf{q}) \hat{O}^\dagger(\mathbf{R}) | \psi^a \rangle \chi_{\alpha\beta}(q, \omega),
\]

where \( \chi_{\alpha\beta}(q, \omega) \) labels four eigenvalues and eigenstates of the MF Hamiltonian \( \hat{H}_{\text{MF}} \) on the site \( a \) and \( \chi_{\alpha\beta}(q, \omega) \) is the corresponding Boltzmann weight.

After the \( J_M \) susceptibility matrix \( \chi_{\alpha\beta}(q, \omega) \) is calculated, it is “upfolded” to the physical \( J = 9/2 \) space similarly to the \( \rho_{J_M}(\mathbf{R}) \) density matrix as described above. The magnetic susceptibility \( \chi_{\alpha\beta}(q, \omega) \) is given by the DD blocks of the upfolded \( \chi_{\alpha\beta}(q, \omega) \) summed over the sublattice indices.

**Data Availability.** All relevant data are available within the manuscript and SI Appendix. Raw computational data have been also deposited in the publicly accessible Materials Cloud Archive (10.24435/materialscloud:12-7qg60).
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