A Mean Field Model for the Quadrupolar Phases of UPd₃

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UPd₃ is known to exhibit four antiferroquadrupolar ordered phases at low temperatures. We report measurements of the magnetisation and magnetostriction of single crystal UPd₃, along the principal symmetry directions, in fields up to 33 T. These results have been combined with recent inelastic neutron and x-ray resonant scattering measurements to construct a mean field model of UPd₃ including up to fourth nearest neighbour interactions. In particular we find that anisotropic quadrupolar interactions must be included in order to explain the low temperature structures derived from the scattering data.

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I. INTRODUCTION

Low temperature phase transitions in condensed matter systems are usually driven by the cooperative actions of the system’s electronic degrees of freedom. In many cases the exchange interactions between the spins of electrons on neighbouring ions causes them to order below a characteristic transition temperature. However, in lanthanide compounds, the localised 4f-electrons on each ionic site may have non-spherical charge or current distributions, which may be described by a multipole expansion of their electric or magnetic fields. These multipoles may interact, and in certain cases these interactions may be stronger than the spin exchange interactions, driving a phase transition to a multipolar ordered phase. There have been many examples of electric quadrupolar order observed, such as in CeB₆, PrPb, and TmTe, whilst higher order electric multipoles were observed in DyB₂C₂.

In contrast to the lanthanides, there have not been many cases of multipolar order observed in actinide compounds. This is due partly to the larger spatial extent of the 5f wavefunctions, which causes them to become delocalised, and hence invalidates any description of their electric or magnetic fields about a particular point. The insulating actinide oxides UO₂ and NpO₂, however, have been studied extensively, and exhibit electric quadrupolar and high order magnetic multipolar order respectively. In contrast UPd₃ is one of the few metallic actinide compounds which has well localised 5f electrons, and it was one of the first compounds to be shown to exhibit quadrupolar ordering. Anomalies were observed in the heat capacity, thermal expansion, magnetic susceptibility, and electrical resistivity of UPd₃ at low temperatures which are indicative of phase transitions. Polarised neutron diffraction measurements revealed superlattice peaks at Q = (½l0) which appear only in the non-spin-flip channel, and thus can be attributed to a structural transition, from hexagonal to orthorhombic symmetry.

X-ray resonant scattering (XRS) measurements showed that this transition arises from the ordering of the electric quadrupole moment of the 5f² electrons. In addition to this phase transition at T₀ = 7.8 K, there are three further transitions to different antiferroquadrupolar (AFQ) ordered states at T₁₁ = 6.9 K, T₁₋₁ = 6.7 K and T₂ = 4.4 K. Below T₀ a superlattice peak at (½l0) where l is odd is observed, whilst below T₁₋₁ there are additional peaks at (½l0) where l is even. The l odd peaks show that there is antiferroquadrupolar ordering along the c-direction, also denoted as an anti-phase stacking of quadrupoles. The l even peaks show an additional ordering of quadrupole moments in-phase along c.

Measurements of the order parameter using x-ray resonant scattering show that the l odd order is associated mainly with Qₓᵧ quadrupoles, whilst the l even order is associated with Qₓz quadrupoles. The Qₓᵧ ordering is accompanied by a component from the Qₓ₋₁₋₁₋₁ quadrupoles, whilst the Qₓz is accompanied by a Qₓz component. The directions x and z are equivalent to the a and c crystallographic directions and y is perpendicular to both. In addition, there is also an additional ordering of the Qₓᵧ quadrupoles in anti-phase below T₁₁. This sequence of phase transitions is summarised in figure 1. UPd₃ adopts the double hexagonal close packed TiNi₃ crystal structure (space group Dᵣ₄₃̄₃, P63/mmc, no. 194) with lattice parameters a = 5.76 Å and c = 9.62 Å. This means that the nearest neighbour U-U distance, 4.1 Å, is larger than the Hill limit (≈ 3.5 Å) and thus the 5f² electrons are well localised. The U⁴⁺ ions are in the 2a and 2d sites, which have respectively D₃d (3m) and D₃h (6m2) point symmetry.

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which we shall refer to as quasi-cubic and hexagonal. The XRS data shows that the ordering involves primarily the ions on the quasi-cubic site.\textsuperscript{[73]}

Knowledge of the crystal field (CF) interactions is essential in determining the more complex two-ion interactions. In particular it is crucial to know the CF ground state. Moreover, the excited states and the matrix elements of the angular momentum operators, $J_i$, between these and the ground state determine the intensities of excitations observed by inelastic neutron scattering and also to some extent the magnitude of the magnetisation in the ordered phase, as explained by McEwen et al.\textsuperscript{[19]} Thus we shall first consider in section II the CF level scheme deduced from inelastic neutron scattering measurements in the paramagnetic phase.

We then present high field magnetisation and magnetostriction measurements in section III from which the critical fields and magnetic phase diagrams were determined experimentally. Finally, a mean field model of the quadrupolar ordering is constructed in section IV. The order parameters of each of the quadrupolar phases (determined by resonant X-ray scattering) and their transition temperatures, were used to constrain the two-ion quadrupolar interaction parameters for the model, whilst the dipolar exchange parameters were determined from the measured critical fields. The results of the model are then compared to the measured high field data.

Whilst the model which we shall present is empirically based on the measured physical properties of UPd$_3$ rather than directly on measured exchange constants, we note that it is the first attempt to explain comprehensively this fascinating compound with its many competing ordered phases from a microscopic point of view, and hope to stimulate further ab initio studies of the exchange interactions involved. The quadrupolar interactions which drive the many phase transitions in UPd$_3$ are difficult to measure directly, because neutrons couple only to the magnetic dipoles in the system, and the energies are far too low for inelastic X-ray scattering to resolve. However, the wealth of physical property measurements available on UPd$_3$ has encouraged us to try to synthesise this into a mean-field model which explains, to a large extent, these varied measurements. We hope that this may encourage the construction of models to explain the properties of similar quadrupolar (or higher multipolar) ordered compounds thus deepening the understanding of what lies behind these phenomena.

![FIG. 1. (Color Online) The quadrupolar phases of UPd$_3$. Labels below the line indicate order parameter as determined from XRS data. ‘In phase’ indicates the quadrupole moments are aligned in parallel along the c-axis. This corresponds to a superlattice reflection at Q=(1 0 4). ‘Anti-phase’ indicates an anti-parallel alignment along c, and corresponds to a reflection at Q=(1 0 3). After Walker et al.\textsuperscript{[73]}](image)

II. CRYSTAL FIELD INTERACTIONS

The single-ion properties of uranium are generally found to be close to the $LS$-coupling limit.\textsuperscript{[16]} So, in order to simplify the analysis, we shall ignore any mixing with higher order multiplets in determining the crystal field (CF) parameters. As mentioned in section I there are two inequivalent sites for the U$^{4+}$ ions in the crystal structure of UPd$_3$. The different point symmetry of these sites gives rise to different crystal fields, but it happens that both split the 9-fold degenerate $J=4$ ground multiplet into three singlets and three doublets. The energies and wavefunctions of these levels, however, are different for the two sites.

The nature of the ground state may be deduced from single crystal susceptibility measurements and estimates of the magnetic entropy determined from heat capacity measurements. These results indicate a singlet ground state on the hexagonal sites and a doublet on the quasi-cubic site.\textsuperscript{[10]}

The CF split energy levels were determined from previously reported inelastic neutron scattering measurements made on the time-of-flight spectrometer HET at the ISIS Facility, UK\textsuperscript{[1]} and on the triple-axis-spectrometer IN8 at ILL,
Grenoble. We identified magnetic excitations at 4.1, 9.7, 12.3, 16.8, 20.4 and 30 meV. The 16 meV peak exhibits considerable dispersion and is assigned to the transition between the $|J_z = 0\rangle$ singlet ground state and the $|J_z = 1\rangle$ doublet excited state of the hexagonal site ions. Its dispersion was used to determine the exchange interactions between hexagonal site ions. The remaining peaks are assigned to the quasi-cubic site ions.

Henceforth we shall be concerned mainly with the quasi-cubic sites, as the quadrupolar ordering primarily involves the uranium ions on these sites. These sites have trigonal, $3m$ (D$_{3d}$), point symmetry, so that the crystal field Hamiltonian is

$$\mathcal{H}_C = \sum_{k=2,4,6} B_{2k}^O \hat{O}_k^0 + \sum_{k=4,6} B_{2k}^M \hat{O}_k^1 + B_{2k}^Q \hat{O}_k^6$$

(1)

where $B_{2k}^O$ are crystal field parameters and $\hat{O}_k^0$ are Stevens operators. The quantisation (c) axis is taken to be the trigonal axis, which in this case is parallel to c.

From the measured transition energies and with the restriction of a doublet ground state, a crystal field fitting program was used to obtain initial estimates of the CF parameters for the quasi-cubic sites. This program relies on the orthogonality of the spherical harmonic functions from which the CF operators are constructed. It allows one to find a set of CF parameters, $B_{2k}^O$, given the energy levels and wavefunctions produced by the crystal field. The fitting algorithm may thus vary either

$|d_1\rangle = a|4\rangle + b|1\rangle + c|2\rangle$  
$|d_2\rangle = a|-4\rangle - b|-1\rangle + c|2\rangle$

(2)

where for brevity the kets denote states $|J = 4, J_z\rangle$. The singlet wavefunctions have the forms

$|s\rangle = a|3\rangle + c|0\rangle - d|--3\rangle$

$|s\rangle = \frac{1}{\sqrt{2}} (|3\rangle + |--3\rangle)$

(3)

In order to ensure that the $T_0 = 7.8$ K transition is accompanied by only a very small entropy change, as de-
The transition between it and some higher energy state which increases $T$ and increases with decreasing temperature through the doublet ground state $\tilde{J}^0_0$, where $\tilde{J}^0_0$ and $\tilde{J}^0_1$ are large whilst $\langle s|\hat{J}_x|d^2_1\rangle$ is small is thus satisfied if $e \approx 1$, $b^{(2)} \approx 1$ and $b \approx 0$.

These requirements are satisfied by the crystal field parameters in table [1] which yield $b = 0.02$, $b^{(2)} = 0.99$ and $e = 0.92$. The parameters were obtained using a simulated annealing minimisation procedure whereby at each iteration, the algorithm mentioned above was used to refine an initial set of parameters to fit the measured energy levels. These refined parameters are subsequently used to calculate the $b$, $b^{(2)}$ and $e$ matrix elements, from which the simulated annealing ‘energy’ is obtained, and hence minimised. Figure [2] shows the resulting crystal field energy splitting and wave functions for the $U^{4+}$ ions on the quasi-cubic sites.

III. HIGH MAGNETIC FIELD MEASUREMENTS

Single crystals were grown by the Czochralski technique at the University of Birmingham, and cut with faces perpendicular to the orthogonal axes $x$, $y$ and $z$, where $x||a$ and $z||c$. These were used in magnetisation and magnetostriction measurements in fields up to 14 T at Birkbeck College by two of us (KAM and JGP). Subsequently, high field magnetisation measurements were carried out (by JGP) at the Grenoble High Magnetic Field Laboratory, and magnetostriction measurements at the National High Magnetic Field Laboratory, Tallahassee. The magnetostriction was measured using a miniature capacitance dilatometer in which the single crystal samples were mounted with either the $x$, $y$ or $z$ faces parallel to the capacitor plates. The dilatometer could be rotated so that the magnetic field is perpendicular to the capacitor plate allowing the transverse components of magnetostriction to be measured.

Figure [5] shows the magnetisation at several different applied fields, whilst figure [4] collates this and other data to construct the magnetic phase diagrams of UPd$_3$. The data shows the $T_0$ transition increasing in temperature with increasing field which is characteristic of an antiferroquadrupolar transition. In general, three ordered phases can be identified from the data, as the phase between $T_{-1}$ and $T_{+1}$ cannot be distinguished from the magnetisation data.

Figure [5] shows the forced magnetostriction data at 4.2 K, plotted as $\Delta I/I = |I(H, T = 4.2 K) - I(H = 0, T = 4.2 K)|/I(H = 0, T = 4.2 K)$. We measured the longitudinal components of magnetostriction with the field in the $x$, $y$- and $z$-directions, and also the transverse components $\Delta y/y$ and $\Delta z/z$ with applied field parallel to $x$, and $\Delta x/x$ with field parallel to $y$. The measurements were repeated to confirm the reproducibility of the data, and subsequently binned and averaged. In addition, the signal was corrected for artefacts due to eddy currents. The mechanical noise from the magnet cooling system and electrical noise in the leads meant that we obtained a resolution of $10^{-6}$ in $\Delta I/I$.

For $H||x$, the magnetostriction parallel (perpendicular) to the applied field first decreases (increases) until approximately 3 T, then increases (decreases) to about 15 T before...
FIG. 5. (Color Online) The high field longitudinal and transverse magnetostriction of UPd₃ at 4.2 K. Solid vertical lines indicate the observed high field transitions. Blue circles, green squares and red triangles denote the magnetostriction parallel to the x-, y- and z-directions, respectively, for the indicated field directions. Solid lines are the results of the mean-field calculations.

was employed, where the site indices i and j run over nearest- (\( \bar{g} = nn \)) and next-nearest neighbours (\( \bar{g} = nnc \)) within an ab plane and nearest- (\( \bar{g} = nnc \)) and next-nearest neighbours (\( \bar{g} = n3c \)) between planes. \( \mathcal{H}^{ci}_j \) is the crystal field Hamiltonian of the \( \bar{g}^{i} \) quasi-cubic ion given in equation (1) and \( \mathcal{H}^{j}_z \) is the Zeeman Hamiltonian, \(-g_i \mu_B H_i \cdot \mathbf{H} \). The form of the two-ion exchange Hamiltonian was derived by considering isotropic interactions between each pair of neighbours in a local coordinate system defined by an \( \bar{x'} \) axis along the bond, and \( \bar{z'} \) axis along c, and then rotating them into a global coordinate system. In this way, the full hexagonal symmetry of the interactions is satisfied. Such expressions were used to explain the properties of elemental Pr\(_{28}\) and the dispersion of crystal field excitations due to interactions between the hexagonal...

IV. QUADRUPOlar TWO-ION INTERACTIONS AND MEAN FIELD MODEL FOR UPd₃

The ordered quadrupolar structures were calculated from a mean field model, with quadrupolar interactions between the quasi-cubic site ions, using the package McPhase\(^{25,26}\). In order to determine the stable ordered structure, a set of supercells and corresponding wave vectors is generated. From the wave vector a configuration of moments (dipole, quadrupole etc) is generated and used as an initial configuration for a self consistent mean field calculation. For each solution of the mean field iteration the free energy is calculated. The self consistent ordered structure with the lowest free energy is taken to be stable and used for the computation of the physical properties. In this way the phase boundaries between the different quadrupolar ordered phases of UPd₃ were determined in order to construct the phase diagrams of U₁₋ₓPrₓPd₃.

The Hamiltonian:

\[
\mathcal{H} = \sum_i \left( \mathcal{H}^{ci}_j + \mathcal{H}^{j}_z \right) - \frac{1}{2} \left\{ g_{11}^{ij} \sum_{Kj} \left[ \hat{P}_x^i \hat{P}_x^j + \hat{P}_y^i \hat{P}_y^j + \hat{P}_z^i \hat{P}_z^j \right] + \kappa_{11}^{ij} \sum_{Kj} \left[ \cos(2\phi_{ij}) \left( \hat{P}_x^i \hat{P}_y^j - \hat{P}_y^i \hat{P}_x^j \right) + \sin(2\phi_{ij}) \left( \hat{P}_x^i \hat{P}_z^j + \hat{P}_z^i \hat{P}_x^j \right) \right] + g_{10}^{ij} \sum_{Kj} \hat{P}_x^i \hat{P}_z^j \right\} \\
+ \frac{1}{2} g_{21}^{ij} \sum_{Kj} \left[ \hat{Q}_{xz}^i \hat{Q}_{xz}^j + \hat{Q}_{yz}^i \hat{Q}_{yz}^j \right] + \kappa_{21}^{ij} \sum_{Kj} \left[ \cos(2\phi_{ij}) \left( \hat{Q}_{xz}^i \hat{Q}_{xz}^j - \hat{Q}_{yz}^i \hat{Q}_{yz}^j \right) + \sin(2\phi_{ij}) \left( \hat{Q}_{xz}^i \hat{Q}_{yz}^j - \hat{Q}_{yz}^i \hat{Q}_{xz}^j \right) \right] \\
+ g_{22}^{ij} \sum_{Kj} \left[ \hat{Q}_{xy}^i \hat{Q}_{xy}^j + \hat{Q}_{xz}^i \hat{Q}_{xz}^j \right] + \kappa_{22}^{ij} \sum_{Kj} \left[ \cos(4\phi_{ij}) \left( \hat{Q}_{xy}^i \hat{Q}_{xy}^j - \hat{Q}_{xz}^i \hat{Q}_{xz}^j \right) + \sin(4\phi_{ij}) \left( \hat{Q}_{xy}^i \hat{Q}_{xz}^j + \hat{Q}_{xz}^i \hat{Q}_{xy}^j \right) \right] \right\} \\
(4)
\]
sites in UPd$_3$[29].

The Hamiltonian (4) suffices to describe the ordered phases of UPd$_3$. In particular, the anisotropic exchange terms ($\propto \mathcal{K}$) are required since the first (second) order isotropic quadrupolar interactions couple the $\hat{Q}_{xx}$ ($\hat{Q}_{xy}$) and $\hat{Q}_{yz}$ ($\hat{Q}_{xz}$, $\hat{Q}_{z^2}$) operators equally, but the principal order parameters measured by resonant X-ray diffraction are $\hat{Q}_{xx}$ and $\hat{Q}_{xy}$. Thus, the anisotropic terms are required to favour these ($\hat{Q}_{xx}$, $\hat{Q}_{xy}$) interactions over the others ($\hat{Q}_{yz}$, $\hat{Q}_{z^2}$).

In order to stabilise a structure with anti-phase (in-phase) stacking along $c$ we require $\gamma_{nn}^{\text{misc}} < 0$ ($\gamma_{nn}^{\text{misc}} > 0$), whilst the AFQ order in the $ab$ plane requires $\gamma_{nn,nn}^{\text{misc}} < 0$. Thus,

$$\gamma_{nn}^{\text{misc}} = \gamma_{21}^{\text{misc}} + \gamma_{31}^{\text{misc}} < 0$$

$$\gamma_{xy}^{\text{misc}} = \gamma_{22}^{\text{misc}} + \gamma_{32}^{\text{misc}} > 0$$

and

$$\gamma_{nn}^{\text{misc}} > |\gamma_{xy}^{\text{misc}}|$$

$$\gamma_{nn,nn}^{\text{misc}} = \gamma_{21}^{\text{misc}} + \gamma_{31}^{\text{misc}} < 0$$

$$\gamma_{nn,nn}^{\text{misc}} > |\gamma_{xy,xy}^{\text{misc}}|$$

should be satisfied in order to yield two phases with $\hat{Q}_{xx}$ anti-phase ($\hat{Q}_{xy}$ in-phase) order along $c$ at higher (lower) temperatures, as measured. Furthermore, because the $\hat{Q}_{xx}$ ($\hat{Q}_{xy}$) and $\hat{Q}_{z^2}$ ($\hat{Q}_{z^2}$) operators share the same symmetry[30] an ordering of one of these pairs will induce a secondary ordering of the other quadrupole of the pair on the same site. That is, a non-zero expectation value $\langle \hat{Q}_{xx} \rangle$ implies $\langle \hat{Q}_{z^2} \rangle \neq 0$ also (angled brackets denote the thermal expectation values $\langle \hat{O} \rangle = \sum_n \langle n | \hat{O} | n \rangle \exp(-E_n/k_B T)/Z$ where the states $| n \rangle$ are eigenstates of the Hamiltonian (4) and $Z$ is the partition function). This contributes to the effective field acting on an ion and combines with the exchange interaction to reinforce (or suppress) the ordering of some particular quadrupole. The strength of this contribution is dependent on the crystal field wavefunctions, and we found that for UPd$_3$, the $\langle \hat{Q}_{z^2} \rangle$ moment induced by a $\hat{Q}_{xx}$ order is of the same order of magnitude as the primary $\langle \hat{Q}_{xx} \rangle$ moment and acts to reinforce the $\hat{Q}_{z^2}$ ordering (induced $\langle \hat{Q}_{z^2} \rangle > 0$). This means that, unfortunately, in our model there is always a large $\hat{Q}_{z^2}$ moment in disagreement with the measured XRS azimuthal dependence, which indicates a 10-15% contribution.

The other induced moments are generally an order of magnitude weaker than their primary order parameter. They are needed, though, to account for the observation by inelastic neutron scattering of four almost dispersionless excitations below 4 meV at 1.8 K, which arise from transitions between the levels of the ground state doublet on the quasi-cubic sites, whose degeneracy is lifted by the quadrupolar order. As there are four quasi-cubic sites in the ordered unit cell, this implies that the splitting on each site is different, which may only occur if the magnitude of the quadrupolar moments on each site is different. In-phase ordering of the quadrupoles along $c$ (denoted CS for $c$-same in the following for brevity) means that the moments on sites C1-C4 of figure [6] are $\langle \hat{Q}_{C1}^{\text{CS}} \rangle = \langle \hat{Q}_{C2}^{\text{CS}} \rangle = \langle \hat{Q}_{C3}^{\text{CS}} \rangle = \langle \hat{Q}_{C4}^{\text{CS}} \rangle$, whilst anti-phase ordering (henceforth denoted CD, $c$-different) implies that $\langle \hat{Q}_{C1}^{\text{CD}} \rangle = -\langle \hat{Q}_{C2}^{\text{CD}} \rangle = -\langle \hat{Q}_{C3}^{\text{CD}} \rangle = \langle \hat{Q}_{C4}^{\text{CD}} \rangle$. However, since $\langle \hat{Q}_{xy} \rangle = |\langle \hat{Q}_{z^2} \rangle|$ and $\langle \hat{Q}_{xy} \rangle = |\langle \hat{Q}_{z^2} \rangle|$, the combination of $\hat{Q}_{xx}$ CD, $\hat{Q}_{xy}$ CD, $\hat{Q}_{xy}$ CS, $\hat{Q}_{z^2}$ CS ordering imposed by the Hamiltonian [3] and conditions [5] and [6] will result in $\langle \hat{Q}_{C1}^{\text{CD}} \rangle = -\langle \hat{Q}_{C3}^{\text{CD}} \rangle = \langle \hat{Q}_{xy} \rangle$ and $\langle \hat{Q}_{C2}^{\text{CD}} \rangle = -\langle \hat{Q}_{C4}^{\text{CD}} \rangle = \langle \hat{Q}_{xy} \rangle$ so that only two excitations may be expected. Only by including the induced moments, which yields $\hat{Q}_{z^2}$ CD and $\hat{Q}_{xy}$ CS ordering (amongst others), will the moments on each of the sites C1-C4, and thus the splitting of the ground state doublet, be unique. This is illustrated schematically on the right side of figure [6].

Finally, we calculate that for the CF scheme outlined in section [11] $d_{12}[\hat{Q}_{xx}|s]$ and $d_{12}[\hat{Q}_{xy}|s]$ are an order of magnitude lower than $d_{12}[\hat{Q}_{xy}|s]$ and $d_{12}[\hat{Q}_{z^2}|s]$ which means that the exchange parameters $J_{21}$ should be an order of magnitude larger than $J_{22}$ to give similar ordering temperatures for the first and second order quadrupoles as observed [29]. From the above considerations, we first determined the order of magnitude of exchange parameters which result in ordering temperatures below 10 K. Subsequently a simulated annealing and particle swarm optimisation [30] search was carried out to find sets of parameters which yield at least two transitions at $\approx 4$ K and $\approx 8$ K and a splitting of the ground state doublet at 2 K close to the measured values 1.28 meV, 1.68 meV, 2.20 meV and 2.60 meV. Additional criteria, including the requirement that the calculated magnetisation with an in-plane applied field should increase with decreasing temperature, and that there should be non-zero moments of all quadrupoles in the lowest temperature phases were then used to sort the candidate sets of parameters found by the search. We then calculated the magnetic phase diagram for the four best sets of parameters, and selected the set which have phase boundaries most similar to those measured. Finally, the parameters were
refined by hand to better match the transition temperatures and fields.

We found during this procedure that unless the dipolar interactions are included, structures with ordering wavevectors \((\frac{1}{2} \frac{1}{2} 0)\) are favoured over the \((\frac{1}{4} 0 0)\) observed (indexed with respect to the dhcp cell). Thus small values of \(J_{nm}\) and \(K_{nm}\) were included, but not varied in the search procedure. They were subsequently refined by hand along with the quadrupolar interaction parameters to better fit the measured critical fields.

The final parameters are shown in table II. At 2 K, the full mean field model with these parameters yields a splitting of the doublet ground state on ions C1-C4 of 0.69 meV, 1.65 meV, 1.82 meV, 2.08 meV respectively (at \(Q = (\frac{2}{3} \frac{2}{3} 0)\)), which is somewhat lower than the experimentally measured values (1.28 meV, 1.68 meV, 2.20 meV and 2.60 meV). The dispersion of the levels along [001], \(\approx 0.5\) meV, is close to the measured value but the in plane dispersion of \(\approx 0.1\) meV is in stark contrast to the measurements which showed the modes to be almost dispersionless.

Furthermore, the calculated order parameters differ from the XRS measurements: in the model, the dominant order parameters are \(\hat{Q}_{x x y y}\) and \(\hat{Q}_{x z}\) rather than \(\hat{Q}_{x x y y}\) and \(\hat{Q}_{x z}\). The phase denoted AFQ1 in figure 3 has a large \(\hat{Q}_{x x y y}\) moment (ordered in antiphase along \(c\)) inducing a small \((\approx 2\%)\) \(\hat{Q}_{x z}\) moment. In the AFQ2 phase, \(\hat{Q}_{x i}\) moments become ordered, inducing some \(\hat{Q}_{x z}\) quadrupoles; both these quadrupolar moments double below the transition to the AFQ3 phase. Thus although the calculated sequence of ordering agrees with experimental data, the type of quadrupolar order does not. Unfortunately, we found it impossible to stabilise the \(\hat{Q}_{x z}\) order parameter over the \(\hat{Q}_{x x y y}\) order parameter at higher temperatures whilst maintaining a ground state with all quadrupolar moments ordered.

Another discrepancy between the model and experiments is the very low critical fields when \(H|c|\) (seen at the bottom of figure 3) and the magnitude of the \(c\) axis magnetisation which is some three times smaller than measured (figure 3). In principle, this can be altered by increasing the \(J_{nn}\) exchange parameters, however, we found that raising these from the values in table II suppresses the quadrupolar ordering completely, in favour of a dipolar order. Alternatively, the quadrupolar exchange parameters \(J_{nn}\) may be altered, but an increase in the critical field necessitates an increase in the transition temperatures.

Since the exchange interactions are likely to arise from the RKKY mechanism, one expects that it should be long ranged. Thus including interactions further than nearest neighbour may give better agreement with the data. However, this vastly increases the parameter space, and unfortunately we could not obtain good fits with simple analytical forms of the RKKY exchange. A more sophisticated approach, using the measured bandstructure of the Pd-U conduction band, may give better results. This consideration may also apply to the quadrupolar interaction parameters, and may account for the discrepancies between the measured and calculated phase boundaries.

Finally, we note that ultrasound measurements and a symmetry analysis of the XRS data showed that there is a sequence of structural transitions from hexagonal to orthorhombic symmetry at \(T_0\) and from orthorhombic to monoclinic at \(T_{11}\). As the parameters in table II have the symmetry of the high temperature hexagonal structure, it is possible that using temperature dependent exchange parameters which incorporate deviations from hexagonal symmetry proportional to the order parameter may yield a better fit.

The magnetoelastic strain is proportional to the strain derivative of the free energy,

\[
\varepsilon^{\alpha\beta} = -\sum_{\beta} s^{\alpha\beta} \frac{\partial F}{\partial \epsilon^{\beta}} 
\]

where \(s^{\alpha\beta}\) is the elastic compliance and the indices \(\alpha\) and \(\beta\) are the Cartesian directions. Noting that \(F = -k_B T \ln Z\), where the partition function is \(Z = \sum_{\alpha} \exp(-E_{\alpha}/k_B T)\), we thus find that \(\frac{\partial F}{\partial \epsilon^{\beta}} = \langle \frac{\partial E_{\alpha}}{\partial \epsilon^{\beta}} \rangle\) where the angled brackets indicate the thermal expectation value.

In the context of the Hamiltonian developed above, there are two main contributions to the magnetoelastic strain. These arise from the single-ion crystal field and the two-ion exchange interactions, both of which depend on the position of the ions, and are hence coupled to any change in the lattice. These two interactions give rise to crystal field striction and exchange striction respectively. In order to calculate the magnetostriction, we thus have to find expressions for the energy levels \(E_\alpha\) as a function of these two interactions. This is done by expanding the Hamiltonian, equation 4, in a Taylor series and making the harmonic approximation by keeping only the first order term, which yields
The calculated magnetostriction fits the data well in the high field regions, but does not reproduce the low field behaviour, particular for the case where the field is applied along x. It may be that at low fields, some of the measured magnetostriction is due to domain rotation, which is not considered in the mean field model.

V. CONCLUSIONS

We have deduced a mean field model including up to four nearest neighbour dipolar exchange and quadrupolar interactions between the 5f electrons of UPd$_3$ which is in generally good qualitative agreement with a broad range of experimental results. The interactions between electrons on the quasi-cubic sites were deduced from resonant x-ray scattering measurements of the quadrupolar order parameters of each of the four low temperature ordered phases, and from the measured transition temperatures and critical fields. The model was then used to calculate the high field magnetisation and magnetostriction, and the magnetic phase diagram up to 30 T. With an applied magnetic field in the basal plane, the calculations for the x and y directions agree with the data. The calculated magnetisation with the field parallel to z is a factor of 3 too large, however, and the calculated critical field in this direction is much smaller than that measured. Despite this, the z-direction calculated magnetic phase diagrams qualitatively reproduces that measured. In conclusion, we have developed a nearest neighbour mean-field model which reproduces many of the main features observed in UPd$_3$.

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TABLE III. Fitted magnetoelastic parameters.

| $\alpha$ | a | b | c |
|----------|---|---|---|
| $A_{\alpha,x} \times 10^{-5}$ | 26 | -100 |  |
| $A_{\alpha,y} \times 10^{-5}$ | -35 | 85 |  |
| $A_{\alpha,z} \times 10^{-5}$ | -1 | 5 | -30 |
| $A_{\alpha,\alpha} \times 10^{-5}$ | 80 | -50 | -30 |
| $A_{\alpha,x^2-y^2} \times 10^{-5}$ | 5 | -60 | -30 |
| $K_{\alpha,xy} \times 10^{-5}$ | 48 | 20 |  |
| $K_{\alpha,yz} \times 10^{-5}$ | -10 | 30 |  |
| $K_{\alpha,xz} \times 10^{-5}$ | -150 | -45 | 25 |
| $K_{\alpha,x^2-y^2} \times 10^{-5}$ | 1 | 10 | 10 |
| $K_{c,xy} \times 10^{-5}$ | -48 | -14 |  |
| $K_{c,yz} \times 10^{-5}$ | -3 | -40 | 5 |
| $K_{c,xz} \times 10^{-5}$ | 200 | 35 |  |
| $K_{c,x^2-y^2} \times 10^{-5}$ | 8 | -4 |  |

TABLE III. Fitted magnetoelastic parameters.

$$e_{\alpha}^\alpha = -\sum_{kq,i} s^{\alpha\beta} \frac{\partial B_k^q}{\partial \epsilon} \langle O_k^\beta \rangle$$

$$e_{\alpha}^\epsilon = \frac{1}{2} \sum_{\beta ij} s^{\alpha\beta} \frac{\partial f_{ij}}{\partial \epsilon} \langle f_{ij}^\beta \rangle$$

The prefactors $A_{\alpha} = \frac{\alpha}{V} \frac{\partial B_k^q}{\partial \epsilon} \langle O_k^\beta \rangle$ and $K_{\alpha\beta} = \frac{\alpha^2}{V} \frac{\partial f_{ij}}{\partial \epsilon} \langle f_{ij}^\beta \rangle$ may be taken to be independent of field and temperature and can then be fitted to experimental data given the thermal expectation values $\langle O_k^\beta \rangle$ and $\langle f_{ij}^\beta \rangle$ obtained from the mean-field model at different applied magnetic fields.

Considering only the non-zero terms up to rank-2 in the Hamiltonian [4], we thus have:

$$e_{\alpha} = \frac{1}{N} \sum_i \left[ \sum_{\beta} A_{\alpha\beta} \langle Q_k^\beta \rangle + \sum_{\beta} K_{\alpha\beta} \langle Q_k^\beta Q_j^\beta \rangle + \sum_{\beta} K_{\alpha\epsilon} \langle Q_k^\beta Q_j^\epsilon \rangle \right]$$

where $N = 4$ is the number of U$^{4+}$ ions in the magnetic unit cell, and the indices $\alpha = x, y, z$, $\beta = xy, yz, z^2, zx, x^2 - y^2$. In order to reduce the number of parameters in fitting equation [10] to the data, we considered only the quadrupolar exchange interactions between nearest neighbour ions in the c-direction and the basal plane, since the calculated expectation values of $\langle Q_k^\beta Q_j^\beta \rangle \gg \langle Q_j^\epsilon \rangle$. Table III shows the fitted parameters, whilst the calculated magnetostriction is shown as solid lines in figure [5].
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