We introduce PyCrystalField, a Python software package for calculating single-ion crystal electric field (CEF) Hamiltonians. This software can calculate a CEF Hamiltonian ab initio from a point charge model for any transition or rare earth ion in either the $J$ basis or the $LS$ basis, perform symmetry analysis to identify nonzero CEF parameters, calculate the energy spectrum and observables such as neutron spectrum and magnetization, and fit CEF Hamiltonians to any experimental data. The theory, implementation, and examples of its use are discussed.

I. INTRODUCTION

A material’s electronic properties—which includes magnetic, optical, and electric qualities—are governed by electron orbitals. Many effects determine an ion’s precise orbital state, but one one of the key effects is crystal electric field (CEF) interactions. These occur when Pauli exclusion and Coulomb repulsion from surrounding atoms shift the relative energies of valence electron orbitals. This can have dramatic effects on an ion’s magnetism, causing single-ion anisotropies, high or low magnetic quantum numbers, or nonmagnetic singlets [1]. To aid in setting up and carrying out such calculations, we introduce PyCrystalField: a python library for calculating and fitting the CEF Hamiltonian of ions.

The theory behind CEF calculations was worked out over 50 years ago [2, 3], but calculating the single-ion Hamiltonian remains a challenge for two reasons. First, the exact Hamiltonian cannot be calculated ab initio. One can use an approximate "point charge model" which neglects higher-order effects [4, 5], but a quantitatively accurate CEF Hamiltonian must be obtained by fitting to data. Second, the complexity of the CEF Hamiltonian depends upon the symmetry of the ion’s environment, causing low-symmetry Hamiltonians to have many independent parameters. Because of this, determining a CEF Hamiltonian can be quite challenging.

Other software has been developed to do CEF calculations, such as FOCUS [6], McPhase [7], SIMPRE [8], or SPECTRE [9]. Useful as they are, these software are limited in either the type of ions they consider—many point-charge models are limited to rare earth ions in the $J$ basis—or the type of data they can fit. PyCrystalField expands the flexibility of CEF software by allowing the user to fit any kind of data: the Python implementation gives the ability to define a custom $\chi^2$ function and fit any variable (including terms in point charge models). It also improves user-friendliness, by automatically building point-charge models from .cif files and fitting to data in just a few lines of code. Finally, PyCrystalField extends the range of point charge modeling to any ion, including to rare earth ions in the $LS$ basis.

II. THEORY

The general CEF Hamiltonian can be written as:

$$\mathcal{H}_{CEF} = \sum_{n,m} B_n^m O_n^m,$$

where $O_n^m$ are the Stevens Operators [2, 3] and $B_n^m$ are multiplicative factors called CEF parameters. $n$ is the operator degree, and is constrained by time-reversal symmetry to be even [10]. $m$ is the operators order, and $-n \geq m \geq n$. These parameters can either be fit to experimental data or calculated from a point-charge model approximation as a starting point of the fit.

A. Point Charge Model

An approximate CEF Hamiltonian can be calculated by treating the surrounding ligands as point-charges and calculating the CEF Hamiltonian from Coulombic repulsion. PyCrystalField calculates $B_n^m$ using the method outlined by Hutchings [3], where

$$B_n^m = -\gamma nm q C_{nm} \langle r^n \rangle \theta_n.$$  

Here, $\gamma nm$ is a term calculated from the ligand environment expressed in terms of Tesseral Harmonics, $q$ is the charge of the central ion (in units of $e$), $C_{nm}$ are normalization factors of the spherical harmonics, $\langle r^n \rangle$ is the expectation value of the radial wavefunction (taken from Ref. [4] for all rare-earth ions), and $\theta_n$ are constants associated with electron orbitals of the magnetic ion. Equation 2 is derived in Appendix A.

B. Neutron Cross Section

Neutron scattering is a common way to measure low-energy (meV range) CEF transitions from one state to another as it can disentangle phonon scattering from crystal field excitations. PyCrystalField calculates the powder-averaged neutron cross section with the dipole approxi-
\[
\frac{d^2\sigma}{d\Omega d\omega} = N (\gamma r_0)^2 \frac{k'}{k} F^2(Q) e^{-2W(Q)} \\
p_n |\langle \Gamma_n^m J_i \rangle|^2 \delta (\hbar \omega + E_n - E_m)
\] (3)

[11], where \(N(\gamma r_0)^2\) are normalization factors, \(k\) and \(k'\) are the incoming and outgoing neutron wavevectors, \(F(Q)\) is the electronic form factor, \(e^{-2W(Q)}\) is the Debye-Waller factor, \(p_n\) is the Boltzmann weight, and \(|\langle \Gamma_n^m J_i \rangle|^2 = \frac{2}{3} \sum_\alpha |\langle \Gamma_m | J_\alpha \Gamma_n \rangle|^2\) is computed from the inner product of total angular momentum \(J_\alpha (\alpha = x, y, z)\) with the CEF eigenstates \(\Gamma_n\).

In practice, the delta function \(\delta (\hbar \omega + E_n - E_m)\) in eq. 3 has a finite width due to the intrinsic energy resolution of the instrument and the finite lifetime of the excited states. By default, PyCrystalField approximates the instrument resolution with a Gaussian profile and the finite lifetimes with a Lorentzian profile, making the effective profile a convolution of the two (PyCrystalField uses a Voigt profile for the sake of computational efficiency). However, the user can specify a custom neutron-peak profile, also with an arbitrary peak-shape dependence on mode energy (This way the user can model asymmetric peaks, where the asymmetry and mode width depend upon energy [12]). With either default or custom peak shapes, PyCrystalField allows the user to specify an arbitrary resolution function giving the FWHM as a function of \(\Delta E\). (A useful approximation for time-of-flight spectrometers can be found in Ref. [13].)

PyCrystalField can calculate 2D (intensity vs \(\Delta E\)) and 3D (intensity vs \(\Delta E\) and \(Q\)) data sets, as shown in Fig. 2. For 3D data sets, the ion and Debye-Waller factor must be specified in order to calculate \(Q\)-dependence. (At the time of writing, the 3D neutron data calculations are only available for rare-earth ions.)

C. Magnetization and Susceptibility

It can be useful to compute susceptibility and magnetization from the CEF Hamiltonian. To do this, PyCrystalField calculates magnetization non-perturbatively as
\[M_n = g_i \langle J_i \rangle, \text{ where } \langle J_i \rangle = \sum \frac{e^{iQ \cdot r}}{Z} |i,J_i\rangle |i\rangle \text{ and } |i\rangle \text{ are the eigenstates of the effective Hamiltonian } \mathcal{H} = \mathcal{H}_{CEF} + \mu_B g \mathbf{B} \cdot \mathbf{J}, \text{ where } \mathbf{B} \text{ is magnetic field. Susceptibility } \chi_{\alpha,\beta} = \frac{\partial M_\alpha}{\partial B_\beta} \text{ is calculated via a numerical derivative of magnetization with respect to field.}

The advantage of the non-perturbative approach is that it can be extended to large magnetic fields and high temperatures without sacrificing accuracy.

D. Intermediate Coupling Scheme

For rare earth ions, the crystal fields are much weaker than spin-orbit coupling and are generally treated as a perturbation to the spin-orbit Hamiltonian, operating on an effective spin \(J\). However, for transition ions (and certain rare earths) where the spin-orbit Hamiltonian \(\mathcal{H}_{SOC} = LS \cdot L\) is of the same magnitude as \(\mathcal{H}_{CEF}\), it is necessary to treat both CEF and spin-orbit coupling non-perturbatively in what is called the "intermediate coupling scheme" where \(\mathcal{H}_{CEF}\) acts only on orbital angular momentum \(L\) [1].

PyCrystalField can do all calculations in either the weak coupling scheme (\(J\) basis) or the intermediate coupling scheme (\(LS\) basis), although in the latter case the ion’s spin-orbit coupling \(\lambda\) must be provided by the user. The advantage of the intermediate scheme is that it can account for intermultiplet transitions, but the disadvantage is that the eigenkets are usually harder to interpret.

In calculating the intermediate-coupling neutron spectrum, \(|\langle \Gamma_m | L_\perp \Gamma_n \rangle|^2 |\langle \Gamma_m | L_\perp + S_\perp \Gamma_n \rangle|^2\) and in calculating magnetization \(M_\alpha = g_i \langle J_\alpha \rangle = \langle L_\alpha + g_e S_\alpha \rangle\). The formula for \(f\)-electron point-charge constants \(\theta_\alpha\) have not been previously published, and are provided in Appendix B.

E. \(g\)-tensor

PyCrystalField can calculate the Lande \(g\)-tensor from an intermediate coupling CEF Hamiltonian. The \(g\) tensor is defined such that
\[
\mathcal{H}_{Zeeman} = \mu_B (\mathbf{B} \cdot \mathbf{L} + g_e \mathbf{B} \cdot \mathbf{S}) = \mu_B \mathbf{B} \cdot \mathbf{g} \cdot \mathbf{J},
\] (4)
where \(\mathbf{B}\) is applied magnetic field, \(\mathbf{L}\) is orbital angular momentum, \(\mathbf{S}\) is spin angular momentum, and \(J\) is total angular momentum. With a doublet ground state \(|\pm\rangle\), the Hamiltonian can be re-written using the Pauli spin matrices. Assuming a magnetic field along \(z\) for simplicity
\[
\mathcal{H}_{eff} = \mu_B B_z \left[ |+|L_z + g_e S_z|+\rangle \langle +|L_z + g_e S_z|+\rangle - |L_z + g_e S_z|\rangle \langle -|L_z + g_e S_z|\rangle \right]
\]
in the \(LS\) basis and
\[
\mathcal{H}_{eff} = \frac{1}{2} \mu_B B_z \left[ \begin{array}{cc} g_{zz} & (g_{zx} - ig_{zy}) \\ (g_{zx} + ig_{zy}) & -g_{zz} \end{array} \right]
\]
in the effective \(J\) basis. Setting these two equations equal to each other and assuming \(B_z\) is small gives the \(g\) tensor values
\[
g_{zz} = 2 \langle +|L_z + g_e S_z|+\rangle, \\
g_{zz} + ig_{zy} = 2 \langle -|L_z + g_e S_z|+\rangle,
\] (5)
and so on for $g_{yy}$, $g_{yx}$, etc. PyCrystalField uses Eq. 5 to compute the $g$ tensor from any intermediate coupling Hamiltonian. For the weak coupling scheme used for rare-earth ions, the equation is the same but with $L_a + g_e S_a$ replaced by $J_a$.

III. IMPLEMENTATION

PyCrystalField is a collection of Python objects and functions which allow the user to build and fit CEF Hamiltonians with just a few lines of code. The workflow is summarized in Fig. 1. The ability to write scripts greatly streamlines the analysis process. It is available for download at https://github.com/asche1/PyCrystalField. Please report bugs to scheieao@ornl.gov.

A. Building CEF Hamiltonian

The user may build a CEF Hamiltonian in two ways: (a) importing a crystal structure for a point-charge model, or (b) specifying the CEF parameters $B_n^m$. For the former case, the user can import the structure from a .cif file or a list of CEF parameters, and then calculate a point charge model or a CEF Hamiltonian respectively. One can fit these models, or directly compute observables like neutron spectrum, susceptibility, etc.

B. Fitting

PyCrystalField allows complete freedom in fitting data. The user must provide a global $\chi^2$ function and PyCrystalField will minimize $\chi^2$ by varying any user-specified variables. The user may fit any set of variables to the data: $B_n^m$, point-charge location (demonstrated in ref. [15]; a similar method is discussed in ref. [16]), effective charge of the point-charges, width of neutron scattering peaks, etc. Any constraints may be imposed. The user may fit to any calculated value: neutron data, magnetization, a list of eigenvalues, $g$-tensor anisotropy, or magnetic susceptibility. In this way, any variable may be fit to any kind of data.

The built-in fit functions are based on the Scipy minimize library, so a variety of methods are available. The PyCrystalField fit function also makes it easy to add more variables to the set of fitted parameters, which is useful for sequentially fitting variables to complex data.

IV. DISCUSSION

A. Examples

In this section we present and discuss three examples to show the performance and capabilities of PyCrystalField.

1. Comparison to Manual Calculation: Yb$^{3+}$ inside a cube of O$^{2-}$ ions

To confirm the accuracy of PyCrystalField, we compare its results to a simple result calculated by hand. We computed the weak-coupling point-charge CEF Hamiltonian for a Yb$^{3+}$ ion surrounded by a cube of O$^{2-}$ ions at
Table I. Calculated CEF parameters for a Yb$^{3+}$ ion inside a cube of O$^{2-}$ ions, by hand via the method in Hutchings [3], and then by PyCrystalField.

| $B_n^{(n)}$ (meV) | By Hand | PyCrystalField |
|------------------|---------|---------------|
| $B_2^4$          | 0.1515035 | 0.1515035     |
| $B_4^4$          | 0.7575173 | 0.7575173     |
| $B_0^{(6)}$      | 0.001604  | 0.001604      |
| $B_6^4$          | -0.0336841 | -0.0336841    |

Figure 2. CEF fits to Nd$_3$Sb$_3$Mg$_2$O$_{14}$ energy and momentum-dependent neutron scattering data from Ref. [17]. (a) shows the data, and (b) shows the fit. The actual fit included three temperatures and three incident energies for a total of nine simultaneously fitted data sets.

PyCrystalField was also used to fit the CEF Hamiltonian of Er triangular lattice delafossites KErSe$_2$ and CsErSe$_2$ [15]. An example fit is shown in Fig. 3. This shows the ability of PyCrystalField to fit multiple temperatures with overlapping peaks. This study also showcases the inherent limitations of CEF fits: two models emerge which fit the data beautifully but have opposite ground state anisotropies. In this case, bulk single-crystal magnetization was used to identify the correct model. This shows that CEF fits to neutron data can be underdetermined, even when fitting to many more peaks than parameters.

3. Ab-initio Ni$^{2+}$ Energy Levels

PyCrystalField can also be used to calculate the ground state energy splitting of an ion in a particular ligand environment. If the ligand positions are well-known, a point-charge model is accurate enough to predict the general character of the ion’s energy spectrum. This is similar to a comparison with a Tanabe-Sugano diagram [1], but with more quantitative accuracy.

An example of this is the calculation of the electron orbital energies of Ni$_2$Mo$_3$O$_8$, which has Ni$^{2+}$ ions surrounded by distorted octahedral and tetrahedral coordination of O$^{2-}$ ions [18]. PyCrystalField allowed for ab initio calculation of the energy spectrum of the valence orbitals in the intermediate coupling scheme. These calculations were used to qualitatively predict magnetic exchanges, single-ion anisotropies, and temperature-dependence in g-factor within Ni$_2$Mo$_3$O$_8$.
B. Uses and Limitations

Before using PyCrystalField, users should be aware of the limitations and pitfalls of crystal field theory. First, the point-charge model is imperfect because the ligand electrons generally have valence p-orbitals, which are anisotropic and cause the effective charge to differ from the idealized point-charge model [19]. Second, the success of fits involving many CEF parameters is highly dependent upon the starting values. There are various approaches to get around this, such as using the point-charge model to generate starting parameters or using a Monte Carlo method to generate many sets of starting parameters. (In the latter case, a comparison to a point-charge model is still helpful to ensure that the CEF parameters are physically sensible.) Third, CEF fits can be underdetermined by neutron scattering data alone [15] and according to Hutchings, where

\[ V(r_n, \theta, \phi) = \sum_{n=0}^{\infty} \sum_{\alpha} r^n \gamma_{n\alpha} Z_{n\alpha}(\theta, \phi), \]

where \( Z_{n\alpha} \) are tesseral harmonics, and

\[ \gamma_{n\alpha} = \frac{4\pi}{(2n+1)} \frac{q_{n\alpha}(\theta, \phi)}{R_n^{n+1}}, \]  

(A1)

summing over \( k \) ligands surrounding the central ion. Recognizing that (according to Hutchings Eq. 5.3):

\[ V(x, y, z) = \sum_{nm} A^m_n \frac{1}{-|e|} f_{nm}^c(x, y, z) \]

and according to Hutchings Eq. 5.5, the Hamiltonian can be written as

\[ \mathcal{H} = -|q| \sum_i V_i(x_i, y_i, z_i) = \sum_i \sum_{nm} A^m_n f_{nm}^c(x, y, z), \]

summing over electrons and where \( q = Ze \). Alternatively, we can write the Hamiltonian in terms of Stevens Operators:

\[ \mathcal{H} = -|Ze| \sum_{n, m} r^n \gamma_{nm} Z_{nm}(\theta_i, \phi_i) \]

\[ = \sum_{k, n, m} A^m_n f_{nm}^c(x_i, y_i, z_i) \sum_{n, m} \frac{(-|e|^2)}{B_n^m} O_n^m \]

\[ \mathcal{H} = \sum_{n, m} B_n^m O_n^m, \]

where \( \theta_n \) is a multiplicative factor which is dependent on the ion \( (\theta_2 = \alpha_J; \theta_4 = \beta_J; \theta_6 = \gamma_J; \) see Table VI in Hutchings). Now, we solve the equations. We can look up \( (r^n) \theta_n \), we just need to find \( A_n^m \).

Because \( A_n^m f_{nm}^c(x_i, y_i, z_i) = -|e|^2 \gamma_{nm} Z_{nm}(\theta_i, \phi_i) \), we should be able to find \( A_n^m \). Now it turns out that, according to Eq. 5.4 in Hutchings, \( C_{nm} \times f_{nm}^c(x_i, y_i, z_i) = r^n Z_{nm}(\theta_i, \phi_i) \), where \( C_{nm} \) is a multiplicative factor in front of the Tesseral harmonics. Therefore,

\[ A_n^m = -\gamma_{nm}^c|e| Ze|C_{nm}. \]

A closed-form expression for the constants \( C \) is very hard to derive, but they are pre-calculated and can be found in "TessHarmConsts.txt".

In the end, we the expression for crystal field parameters \( B_n^m \) is

\[ B_n^m = A_n^m (r^n) \theta_n \]

\[ B_n^m = -\gamma_{nm}^c|e| Ze|C_{nm} \theta_n. \]  

(A2)

We know \( |e|, Z \) (ionization of central ion), and the constants \( C_{nm}, \theta_n \) are given in Hutchings, and \( (r^n) \) are found in Ref. [4] for rare earth ions.

V. CONCLUSION

PyCrystalField is a general-purpose crystal field calculation package which allows the user to consider any type of data, any ion, and any ligand environment. It also allows the user to calculate observable quantities such as neutron scattering and bulk magnetization and susceptibility. It is accurate in reproducing simple point-charge calculations, it can fit highly asymmetric Hamiltonians to complex data sets, and it can make qualitative ground state predictions from a point-charge model.

PyCrystalField simplifies difficult CEF calculations in a way that makes them practical for more of the scientific community. Understanding the single-ion properties in a material is often critical to understanding the behavior of the whole, and CEF calculations are a key part of this.

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Appendix A: Calculation of Point-charge CEF Parameters

We begin with Hutchings [3] Eq. 2.7:

\[ V(r_n, \theta, \phi) = \sum_{n=0}^{\infty} \sum_{\alpha} r^n \gamma_{n\alpha} Z_{n\alpha}(\theta, \phi), \]

where \( Z_{n\alpha} \) are tesseral harmonics, and

\[ \gamma_{n\alpha} = \frac{4\pi}{(2n+1)} \frac{q_{n\alpha}(\theta, \phi)}{R_n^{n+1}}, \]  

(A1)

summing over \( k \) ligands surrounding the central ion. Recognizing that (according to Hutchings Eq. 5.3):

\[ V(x, y, z) = \sum_{nm} A^m_n \frac{1}{-|e|} f_{nm}^c(x, y, z) \]

and according to Hutchings Eq. 5.5, the Hamiltonian can be written as

\[ \mathcal{H} = -|q| \sum_i V_i(x_i, y_i, z_i) = \sum_i \sum_{nm} A^m_n f_{nm}^c(x, y, z), \]

summing over electrons and where \( q = Ze \). Alternatively, we can write the Hamiltonian in terms of Stevens Operators:

\[ \mathcal{H} = -|Ze| \sum_{n, m} r^n \gamma_{nm} Z_{nm}(\theta_i, \phi_i) \]

\[ = \sum_{k, n, m} A^m_n f_{nm}^c(x_i, y_i, z_i) \sum_{n, m} \frac{(-|e|^2)}{B_n^m} O_n^m \]

\[ \mathcal{H} = \sum_{n, m} B_n^m O_n^m, \]

where \( \theta_n \) is a multiplicative factor which is dependent on the ion \( (\theta_2 = \alpha_J; \theta_4 = \beta_J; \theta_6 = \gamma_J; \) see Table VI in Hutchings). Now, we solve the equations. We can look up \( (r^n) \theta_n \), we just need to find \( A_n^m \).

Because \( A_n^m f_{nm}^c(x_i, y_i, z_i) = -|e|^2 \gamma_{nm} Z_{nm}(\theta_i, \phi_i) \), we should be able to find \( A_n^m \). Now it turns out that, according to Eq. 5.4 in Hutchings, \( C_{nm} \times f_{nm}^c(x_i, y_i, z_i) = r^n Z_{nm}(\theta_i, \phi_i) \), where \( C_{nm} \) is a multiplicative factor in front of the Tesseral harmonics. Therefore,

\[ A_n^m = -\gamma_{nm}^c|e| Ze|C_{nm}. \]

A closed-form expression for the constants \( C \) is very hard to derive, but they are pre-calculated and can be found in "TessHarmConsts.txt".

In the end, we the expression for crystal field parameters \( B_n^m \) is

\[ B_n^m = A_n^m (r^n) \theta_n \]

\[ B_n^m = -\gamma_{nm}^c|e| Ze|C_{nm} \theta_n. \]  

(A2)

We know \( |e|, Z \) (ionization of central ion), and the constants \( C_{nm}, \theta_n \) are given in Hutchings, and \( (r^n) \) are found in Ref. [4] for rare earth ions.
PyCrystalField calculates the Hamiltonian in units of meV. $\gamma_{\alpha}$ is in units of $\frac{e^2}{\lambda}$, $C_{nm}$ and $\theta_n$ are unitless, and $(r^n)$ is in units of $(a_0)^n$. This means that $B_m^n$ in the equation written above come out in units of $\frac{e^2}{\lambda}$.

To convert to meV, we first recognize that we have to re-write Hutchings eq. (II.2) with the proper prefactor for Coulomb’s law: $W = \sum_i \frac{1}{4\pi\varepsilon_0} q_i V_i$. Thus, our Hamiltonian becomes

$$H_{CEF} = \sum_{nm} B(exp)_n^m O_n^m = \sum_{nm} \frac{1}{4\pi\varepsilon_0} B(calc)_n^m O_n^m.$$  

Now $\epsilon_0 = \frac{e^2}{2\pi\varepsilon_0}$, so

$$B(exp)_n^m = \frac{1}{4\pi\varepsilon_0} \frac{1}{\gamma_{nm}} \frac{Ze}{C_{nm}} \langle r^n \rangle \theta_n,$$

$$B(calc)_n^m = -\frac{2\alpha\hbar c}{4\pi\varepsilon_0^2} \frac{\gamma_{nm}}{C_{nm}} \langle r^n \rangle \theta_n e^2 Z \frac{a_0^n}{\lambda^{n+1}}.$$

Plugging in the values, we get the equation used by PyCrystalField:

$$B_n^m = 1.440 \times 10^4 (0.5292)^n Z (\gamma_{nm} C_{nm} \langle r^n \rangle \theta_n) \text{ meV}.$$  

**Appendix B: $\theta_n$ in the Intermediate Scheme**

When calculating crystal field levels from the point charge model, the $\theta_n$ constants are listed in Stevens (1952) [2] for the ground states of all the rare earth ions in the $J$ basis. But if we want to look at the state of the rare earth ion in the $LS$ basis (intermediate coupling scheme), we must recalculate $\theta_n$ for each state.

General formulae for first two constants, $\theta_2 = \alpha$ and $\theta_4 = \beta$, are listed in Ref. [20]. However, because they assume that such constants are only necessary for 3d group ions, they do not list $\theta_6$. Therefore, we must derive it following the method of Stevens [2]. We take as an example the Sm$^{3+}$ ion, which has five electrons in its $f$ orbital valence shell and a ground state of Sm$^{3+}$ has $S = 5/2$, $L = 5$, and $J = 5/2$.

We know that

$$V_6^0 = \Sigma (231z^6 - 315z^4r^2 + 105z^2r^4 - 5r^6) = \gamma_6 \alpha \theta_6 O_6^0,$$

where the Stevens operator

$$O_6^0 = 231L_z^6 - (315X - 735)L_z^4 + (105X^2 - 525X + 294)L_z^2 - 5X^3 + 40X^2 - 60X$$

and $X = L(L + 1)$. Now to calculate $\theta_6$, we pick an eigenstate (in this case, $|L = 5, S = 5/2, m_l = 5, m_s = -5/2\rangle$) and calculate the expectation value in terms of the Stevens Operators, and then calculate it in terms of the individual electron wave functions. Then we set the two results equal to each other to find the multiplicative factor necessary to make the Stevens Operator result match the wave function integral.

**Stevens Operators:** This is a straightforward calculation from eq. B2. Letting $G_6^0 = \langle r^6 \rangle \theta_6 O_6^0$,

$$\langle L = 5, m_l = 5 | G_6^0 | L = 5, m_l = 5 \rangle = \theta_6 \langle r^6 \rangle 37800.$$

**Individual electron wave functions:** To carry out this calculation we write $|m_l = 5, m_s = -\frac{5}{2}\rangle$ as a product state of the individual electron wave functions in the valence shell, which have $l = 3, m = 3, 2, 1, 0, -1$ (adding up to $L = 5$). Now we just recompute $V_6^0$ in the basis of individual electrons: $V_6^0 = \gamma O_6^0$, where $m_l = m$ and $L = l$.

$$\langle L = 5, m_l = 5 | V_6^0 | L = 5, m_l = 5 \rangle = \{3, 2, 1, 0, -1\} V_6^0 \{3, 2, 1, 0, -1\}$$

$$= \gamma (180 - 1080 + 2700 - 3000 + 2700) = 900\gamma.$$  

We introduced $\gamma$ to account for the unknown scaling factor. (The similarity to $\gamma_{60}$ is unfortunate, because the $\gamma$s are unrelated. Nevertheless, we follow the notation of Stevens in this appendix.) So now we can relate eq. B3 to eq. B4 so $\beta \langle r^6 \rangle = \frac{900}{37800} \gamma = \frac{4}{42} \gamma$. Now we find $\gamma$ by integrating the wave functions themselves. Let us pick the state $l = 3, m = 3$:

$$\langle l = 3, m = 3 | V_6^0 | l = 3, m = 3 \rangle = 180\gamma$$

$$= \int_0^{2\pi} \int_0^\pi Y_3^3 (231z^6 - 315z^4r^2 + 105z^2r^4 - 5r^6)Y_3^3 \sin \theta d\theta d\phi$$

$$= \frac{-80}{429} r^6.$$  

This means that $\gamma = \frac{-80}{429} r^6 = -\frac{4}{3861} r^6$, so that

$$\theta_6 \langle r^6 \rangle = \frac{1}{42} \gamma = -\frac{4}{42 \cdot 3861} r^6$$

and $\theta_6$ for Sm$^{3+}$ is

$$\theta_6 = \frac{4}{42 \cdot 3861}.$$  

For each ion, we have to calculate $\langle L, m_l | G_6^0 | L, m_l \rangle$ and then calculate $\{3, 2, 1, 0, -1\} V_6^0 \{3, 2, 1, 0, -1\}$. Fortunately, the result $\gamma_6 = \frac{4}{3861} r^6$ holds for all rare earth ions. Thus, we arrive at the general formula for $\theta_6$:

$$\theta_6 = \frac{\{3, 2, 1, 0, -1\} V_6^0 \{3, 2, 1, 0, -1\}}{\langle L, m_l | O_6^0 | L, m_l \rangle} \frac{4}{3861}.$$  

$$\theta_6 = \frac{\{3, 2, 1, 0, -1\} V_6^0 \{3, 2, 1, 0, -1\}}{\langle L, m_l | O_6^0 | L, m_l \rangle} \frac{4}{3861}.$$
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