# Molecular Crystal Global Phase Diagrams III: Sufficient parameter space determination

J. Brandon Keith\* and Richard B. McClurg

Department of Chemical Engineering and Materials Science,

University of Minnesota, Minneapolis, Minnesota

# Abstract

In previous papers [1, 2] we developed a method for constructing global phase diagrams (GPDs) for molecular crystals and partially applied it to identifying reference lattices pertaining to all single component ordered crystal structures of tetrahedral molecules in the Cambridge Structural Database. Here we expand upon these results by outlining a method to derive a representative potential characteristic of all experimental structures in our data set. This is significant because our prior work [1] did not specify the number of parameters needed for GPDs. Although there are suggestions in the literature that thousands of parameters are required to adequately describe tetrahedral molecule intermolecular potentials, we find 15 parameters are sufficient to successfully plot the structures of our test data on molecular crystal GPDs.

<sup>\*</sup>Electronic address: jbrkeith@gmail.com; Also at Department of Physics and Astronomy, Brigham Young University, Provo, Utah

#### I. INTRODUCTION

Supramolecular organic solids and mesophases are of primary interest in pharmaceuticals, electrophotographic processes, organic pigments and dyes, and molecular FET's and dielectrics [3]. To alter their electrical, optical, thermal, and solubility properties it is necessary to design both the molecular and supramolecular structure to produce novel, application-specific materials. While standard quantum chemistry methods are adequate for predicting molecular properties, the weak interactions responsible for determining supramolecular structure are more difficult to design [4, 5].

Previous work [1, 6] has demonstrated the ability to construct global phase diagrams (GPDs) which can be used for materials design by displaying the types of structures obtained from all possible supramolecular potentials. GPDs use an intermolecular potential constructed from a complete set of basis functions for rotational space for all molecules of a particular point group and a set of molecular center-of-mass packings derived in a companion paper [2]. The parameters of the intermolecular potential become axes on GPDs for each packing and the crystal structures are phase regions in the diagrams.

An important benchmark of GPD's utility is to show that experimentally observed structures can be found and their intermolecular potential parameters can be read from the axes. In a companion paper [2] we found our data set consisting of all ordered single-component crystal structures of tetrahedral molecules in the Cambridge Structural Database (CSD) [7] have molecular centers of mass situated on high symmetry space groups, or reference lattices. Our goal in this paper is to determine the number of potential parameters needed to find all experimental phases of this data set on GPDs pertaining to these reference lattices. A small number of each will disprove previous hypotheses that many parameters are needed to describe intermolecular potentials [8] and will enhance the useability of GPDs in materials design. The remainder of this paper is as follows. In Sec. II we review the rotational potential and outline the computational procedure to find potential parameters for each structure. This is done using global optimization techniques among parameter and configurational space. In Sec. III we discuss the reverse-engineered potentials and some aspects of our model. We end with a conclusion about our results.

#### II. REPRESENTATIVE POTENTIAL DETERMINATION

Global phase diagrams require (1) reference lattices consistent with molecular centers of mass in experimental structures and (2) intermolecular potential (IP) parameters to use as independent parameters. In [2] we found the majority of the experimental structures in the CSD of tetrahedral molecules can be classified using only four reference lattices. Since there is little data for other reference lattices, we focus on the set {bcc, fcc, hcp, sc}. Each of these has at least four monomer crystal structures in the data set. In this section we seek to determine a *sufficient* parameter space such that experimental structures are each stable in a separate subspace. Thus we review our choice of IP and define the potential parameters. Then we outline a method for identifying representative potential parameters for each experimental structure. This is done using a low temperature structural limit which is convenient and consistent with the experimental database but is not strictly necessary. Then the IP is truncated to create a finite dimensional IP parameter space. A library of alternative crystal structures is constructed with which to compare experimental structures, and a figure of merit is specified when searching for the potential parameters. This procedure identifies a structure with similar cell shape, molecular center of mass, and molecular orientations as shown in Fig. 1.

## A. Mesoscopic Hamiltonian

Previously [1] we discussed the construction of a nearest neighbor rotational potential for van der Waals molecules. It is a level of abstraction above an atom-atom or site-site potential but retains a firm basis in quantum mechanics [9]. The potential consists of a two center expansion constructed by coupling one center basis functions  $U_{m_{\tau}n_{\sigma}}^{\ell_{i}}$  for pairs of molecules i and j and coupling "matrix"  $J_{m_{\tau}n_{\sigma}m_{\rho}n_{\mu}}^{\ell_{i}\ell_{j}}$ ,

$$V_{\text{or}} = \frac{1}{2} \sum_{ij} \sum_{\ell_i \ell_j m_\tau m_\rho n_\sigma n_\mu} U_{m_\tau n_\sigma}^{\ell_i}(\boldsymbol{\omega}_i) \times J_{m_\tau n_\sigma m_\rho n_\mu}^{\ell_i \ell_j}(\boldsymbol{\nu}, \boldsymbol{\Omega}_{ij}) U_{m_\rho n_\mu}^{\ell_j}(\boldsymbol{\omega}_i).$$

$$(1)$$

The one half avoids overcounting,  $\ell_i, \ell_j \in \mathbb{N}$ , the natural numbers, and  $|\ell_i - \ell_j| \leq \ell \leq \ell_i + \ell_j$ .  $U_{m_\tau n_\sigma}^{\ell_i}(\boldsymbol{\omega}_i)$  are functions of the orientation of the molecule through its Euler angles  $\boldsymbol{\omega}_i$  using the passive convention [10]. They are projected from SO(3) irreducible representations (IR's), also called Wigner functions, and contain both the point group symmetry of the molecule and that of the Wyckoff point in the crystal,

$$U_{m_{\tau}n_{\sigma}}^{\ell_{i}}(\boldsymbol{\omega}_{i}) = \sum_{m_{i}n_{i}} S_{m_{i}m_{\tau}}^{\ell_{i}*} D_{m_{i}n_{i}}^{\ell_{i}}(\boldsymbol{\omega}_{i}) S_{n_{i}n_{\sigma}}^{\ell_{i}}.$$
 (2)

The sparse unitary matrix S provides the linear combinations of Wigner functions which give a particular point group IR symmetry [11]. Specifically, the left multiplication by  $S^{-1}$  in Eq. (2) gives basis functions transforming like Wyckoff point group IR's and the right multiplication by S gives basis functions of the molecular point group IR's. Subscript  $\tau$  is a compound index referring to multiple copies of the Wyckoff point group IR subduced in  $\ell_i$  and  $m_{\tau}$  goes over the dimensions of each IR. Subscript  $\sigma$  is a compound index referring to multiple copies of the molecular point group unit IR subduced in the  $\ell_i$ -th manifold of SO(3) and  $n_{\sigma}$  is its dimension. Point group IR subduction frequencies in spherical harmonics are discussed elsewhere [11] and in Sec. IIB. The symmetry adaption leaves relatively few basis functions since only matrix elements where  $\sigma$  is the unit IR  $(i.e.\ A_1)$  are kept. This gives basis functions in the molecular frame with the full molecular symmetry. Thus the set  $\{U_{m_{\tau}n_{\sigma}}^{\ell_i}\}$  is a complete set of functions taking full advantage of symmetry.

The coupling matrix  $J_{m_{\tau}n_{\sigma}m_{\rho}n_{\mu}}^{\ell_{i}\ell_{j}}$  specifies the angular dependence with respect to molecular centers,

$$J_{m_{\tau}n_{\sigma}m_{\rho}n_{\mu}}^{\ell_{i}\ell_{j}}(\boldsymbol{\nu},\boldsymbol{\Omega}_{ij}) = \sum_{\ell m_{i}mm_{j}} \nu_{\ell_{i},\ell,\ell_{j}}^{n_{\sigma},n_{\mu}}(r_{ij}) \begin{pmatrix} \ell_{i} & \ell & \ell_{j} \\ m_{i} & m & m_{j} \end{pmatrix} \times S_{m_{\tau}m_{i}}^{\ell_{i}} C_{m}^{\ell}(\boldsymbol{\Omega}_{ij}) S_{m_{\rho}m_{j}}^{\ell_{j}},$$

$$(3)$$

The potential coefficients  $\nu_{\ell_i,\ell,\ell_j}^{n_\sigma,n_\mu}(r_{ij})$  are a function of the distance  $r_{ij}$  between molecular centers. The neighbor distance  $r_{ij}$  and orientation  $\Omega_{ij}$  of the intermolecular vector are determined by the reference lattice. Since we consider pairwise interactions only among the equidistant nearest neighbors of the reference lattices, the functions  $\nu_{\ell_i,\ell,\ell_j}^{n_\sigma,n_\mu}(r_{ij})$  are only evaluated at the nearest neighbor distance and are treated as scalars. Thus the coupling matrix  $J_{m_\tau n_\sigma m_\rho n_\mu}^{\ell_i \ell_j}$  contains reference lattice information through  $\Omega_{ij}$  and  $r_{ij}$  as well as pairwise intermolecular potential information through the coefficients  $\boldsymbol{\nu}$ . These coefficients have inversion symmetries that reduce the number of independent values, even for asymmetric molecules. In particular  $\nu_{\ell_i,\ell,\ell_j}^{n_\sigma,n_\mu}$  is zero if  $\ell_i + \ell + \ell_j$  is odd and  $\nu_{\ell_i,\ell,\ell_j}^{n_\sigma,n_\mu} = (-1)^{\ell_i+\ell_j}\nu_{\ell_j,\ell,\ell_i}^{n_\mu,n_\sigma}$  for single component crystals. [12] The full set of  $\nu_{\ell_i,\ell,\ell_j}^{n_\sigma,n_\mu}(r_{ij})$ , denoted  $\boldsymbol{\nu}$ , are the axes of GPD's and the search space for reverse engineering experimental crystal structures.

#### B. Computational Method

Having established translational and rotational components of the intermolecular potential as the parameter space for GPDs, we seek to determine potential parameters  $\nu_{\ell_i,\ell,\ell_j}^{n_\sigma,n_\mu}$  sufficient to produce each experimental crystal structure. Our method has five steps. (1) Define a figure of merit by which to order the crystal structures. (2) Ensure the potential, Eq. (1), is truncated such that it geometrically is able to produce a symmetry change from the assigned reference lattice to the experimental phase. (3) Develop a library of structural types against which this energy may be compared. (4) Determine the energy of an experimental phase as a function of  $\nu$ . (5) Search potential parameter space until a  $\nu$  vector is found which makes the experimental phase energetically minimal relative to the alternatives in the library.

# 1. Figure of Merit

In our previous work [1, 6] linear response theory was used to seek phase transitions as a model crystal is cooled from a disordered plastic crystalline state at high temperature. These transitions are bifurcation points of the free energy surface. Following each transition, the phase was identified by the presence of nonzero thermal averages of rotator functions  $U_{m_{\tau}n_{\sigma}}^{\ell_{i}}(\boldsymbol{\omega}_{i})$  which serve as order parameters. This work examines the low temperature limit of the previous model where low is relative to the temperature  $T_{pt}$  at which the molecules undergo the first phase transition from the reference phase. In this regime the function  $U_{m_{\tau}n_{\sigma}}^{\ell_{i}}$  itself is an adequate approximation to the thermal averages of the rotator functions,

$$\lim_{T/T_{\rm pt} \to 0} \langle U_{m_{\tau}n_{\sigma}}^{\ell_i} \rangle = U_{m_{\tau}n_{\sigma}}^{\ell_i}(\boldsymbol{\omega}_0). \tag{4}$$

Also, the free energy is equal to the potential in that limit,

$$\lim_{T/T_{\text{pt}} \to 0} A = V. \tag{5}$$

Equations (4) and (5) are common approximations used in many crystal structure prediction codes [13] and are further justified by the exclusion of disordered structures in *CSDSymmetry* from which our data set is derived.

## 2. Potential Truncation

The potential in Eq. (1) is a doubly infinite sum over manifolds  $\ell_i$  and  $\ell_j$  that must be truncated for practical application. To determine an adequate truncation of the potential we make use of space group irreducible representations (IR's). Symmetry-breaking mechanisms are classified by space group IR's and an order parameter direction such as (a, b) [14]. The temperature dependent values a and b are given in our model by components of space group IR-adapted basis functions q which are linear combinations of  $U_{m_{\tau}n_{\sigma}}^{\ell_i}(\omega_i)$  [1]. Space group IR distortions of the crystal can be decomposed into point group IR distortions of the distinguishable molecules in the crystal [15]. To determine which IR's  $\tau_w$  of point group w are in a symmetry-breaking space group IR  $\tau_{SG}$ , we calculate their subduction frequencies  $n_{SG}$ 

$$n_{SG} = \frac{1}{|w|} \sum_{g \in w} \chi^{\tau_{SG}}(g)^* \chi^{\tau_w}(g) \quad \forall \ \tau_{SG}$$
 (6)

where |w| is the number of elements g in w.  $\chi(g)^{\tau_{SG}}$  and  $\chi(g)^{\tau_{w}}$  are the traces of the matrix representation of the space group IR and point group IR, respectively. The calculations are easily performed using the ISOTROPY software package [14] using the "show frequency" command. Space group IR and point group IR characters may also be generated.

Using these point groups one may also calculate the number of times each point group IR  $\tau_w$  appears for each manifold of SO(3), or  $\ell_i$  value,

$$n_{SO(3)} = \frac{1}{|w|} \sum_{g \in w} \chi^{\ell_i}(g)^* \chi^{\tau_w}(g) \quad \forall \ \ell_i$$
 (7)

where  $\chi^{\ell_i}$  is the trace of the matrix representation of an IR of SO(3) also called a Wigner rotation matrix. Such subduction frequencies may be easily calculated but are most accessible in standard tables in the literature [11] and are shown for  $O_h$  and  $D_{3h}$  in Table I. As there is no molecular unit IR in the first, second, or fifth manifold, there are no Wyckoff IR rows of  $U_{m_{\pi}n_{\pi}}^{\ell_i}$  so these manifolds are not shown.

If the potential is truncated before the manifold at which a Wyckoff IR forming a symmetry-breaking space group IR is present (i.e. if  $n_{SG} = 0$  in Eq. 6), the desired phase transition cannot occur with that potential. Therefore we calculate minimal SO(3) manifolds necessary to achieve a transition from the assigned reference lattice to an experimental structure. This is shown for the fcc reference lattice in Table II. Point group IR's subduced

in the space group IR and the first manifold or  $\ell_i$  value of SO(3) at which a given Wyckoff IR appears are also shown. For some pathways, such as that of carbon tetraiodide, 225a  $\rightarrow$  121a in Table II, there is a single space group IR  $\Gamma_5^-$  subducing a single point group IR  $T_{2u}$ . Table I shows that this point group IR first occurs in the third manifold of SO(3) when split by an  $O_h$  point group crystal field. Thus  $\ell_i^{\max} \geq 3$  may give the 121a structure. For other pathways, such as that of tetrakis(M<sub>3</sub>-t-Butylimido)-tetraiodo-tetra-indium, 225a  $\rightarrow$  12i, there is more than one Wyckoff IR subduced in a space group IR.  $E_u$ ,  $T_{1u}$ , and  $T_{2u}$  can lead to this transition. As two of them are present at the third manifold (Table I), a potential with  $\ell_i^{\max} \geq 3$  may also produce this transition. For still others, such as that of adamantane, 225a  $\rightarrow$  114a, the pathway is coupled, or composed of the direct sum of two space group IR's. Three different coupled space group IR's decompose into three different point group IR's. One of these pathways uses IR's on the third manifold of SO(3) ( $\ell_i^{\max} \geq 3$ ) while the other two require fourth manifold basis functions. A similar analysis for the bcc, hcp, and sc reference lattices given in the appendix shows symmetry-breaking Wyckoff IR's appear by at least the third or fourth manifold in nearly all cases.

In real phase transitions, IR's on the first non-trivial manifold typically induce symmetry breaking. Second manifold induced symmetry breaking is uncommon and higher manifold induced phase transitions are not generally observed [16]. The reasons for this are different for small and large molecules. For small molecules with only a few atoms (i.e. CH<sub>4</sub>) molecular orbitals tend to form low-energy, slowly varying topologies to lessen kinetic energy contributions in Schrödinger's equation. The result is an IP well represented by smooth, slowly varying basis functions. For large molecules, low manifold contributions to the potential are sufficient to locate attractive/repulsive regions of the potential and the relative magnitudes of attractive configurations, even though these basis functions are not sufficient to represent the fine structure in the potential due to individual atoms [17]. In either case, the most important aspects of the IP are given by slowly varying functions while more rapidly varying functions produce finer details. Thus as a minimal basis set we truncate Eq. (1) at  $\ell_{\text{max}} = 3$  or 4.

Truncating at  $\ell_{\text{max}} = 4$  leaves fifteen coefficients in the potential,  $\nu_{0,3,3}$ ,  $\nu_{0,4,4}$ ,  $\nu_{3,0,3}$ ,  $\nu_{3,2,3}$ ,  $\nu_{3,4,3}$ ,  $\nu_{3,6,3}$ ,  $\nu_{3,1,4}$ ,  $\nu_{3,3,4}$ ,  $\nu_{3,5,4}$ ,  $\nu_{3,7,4}$ ,  $\nu_{4,0,4}$ ,  $\nu_{4,2,4}$ ,  $\nu_{4,4,4}$ ,  $\nu_{4,6,4}$ , and  $\nu_{4,8,4}$ . The coefficient  $\nu_{0,0,0}$  is negligible since it only affects the trivial basis function  $U_{0,0}^0 = 1$  which is isotropic and therefore unimportant in rotational ordering. As the unit IR for  $T_d$  appears once in

the zeroth, third and fourth manifolds,  $n_{\sigma}$  and  $n_{\mu}$  are always  $1_{A_1}$  and have been dropped in the notation for the coefficients in Eq. (3). For space groups whose occupied Wyckoff point group in the reference lattice has the inversion as one of its symmetry operators,  $\nu_{0,x,x}$ cancels in the crystal field for odd values of x. This is the case with the  $\nu_{033}$  coefficient in the bcc, fcc, and sc reference lattices with 14 coefficients but not for hcp with 15.

## 3. Candidate Lattice Library

We seek a set of potential parameters  $\nu$  such that the energy is lower than other structures, whether observed or not. Therefore, a comparative list of structural types should consist of both the experimental structures and a large library of alternate structures. Each alternative structure serves to constrain the representative potential parameters  $\nu$  identified with each experimental structure. In our previous work [1], high symmetry point IR distortions [14, 18] from a reference lattice led to structures classified using their isotropy subgroups (ISs). These subgroups represent the most common types of phase transitions and are minimal in the sense that only one domain of each structure is tested. This compares favorably to crystal structure prediction which generates thousands of multi-domain duplicate structures. Therefore they are convenient for constructing a library of candidate lattices. They are discussed further in Sec. III. As most molecular crystals have one molecule or less per asymmetric part of the unit cell  $(Z' \leq 1)$ , we have discarded ISs that imply more than one occupied Wyckoff point [19]. The library becomes inconveniently large if Z' > 1 is permitted. ISs whose primitive unit cell contains more than eight molecules are also discarded [20].

To minimize the energy of these candidate lattices,  $U_{m_{\tau}n_{\sigma}}^{\ell_i}(\boldsymbol{\omega}_i)$  are placed at one molecular center of the molecular positions of the lattice of each IS and space group operations of the IS are applied to generate basis functions for all other molecules in the Wyckoff orbit. This reduces the number of Euler angles needed to just one set for each candidate lattice. Although these candidate lattices are generated from ISs, after Euler angle minimization they are free to assume whatever structure the minimization algorithm can find provided the basis functions are related by IS space group operations. Thus a candidate lattice has fixed (super) cell parameters and Wyckoff points but variable Euler angles and space group. This results in a generalization of and supergroup to high symmetry isotropy subgroups.

This procedure gives 75 candidate lattices for bcc, 67 for fcc, 19 for hcp, and 105 for sc.

These augment the list of observed structures in our library. The unit cells of the candidate lattices are given in the supplementary material. The unit vectors are given in terms of their reference lattice vectors. Space group operations (of the reference lattice) that relate molecules on distinct lattice points are also given. Minimizing the energy of all candidate lattices gives a lowest energy structure with which to compare the energy of the experimental structure while searching parameter space.

#### 4. Experimental Phase Energies

The potential V in Eq. (1) is a function of Euler angles  $\omega$ , potential parameters  $\nu_{\ell_i,\ell,\ell_i}$ , and intermolecular angle  $\Omega$  (measured with respect to the z-axis in the lab reference frame of molecule i). The intermolecular angles are determined by the reference lattice but the Euler angles are dependent upon the orientation of the molecules on the reference lattice. This is complicated by the fact that the experimental atomic positions in each molecule are slightly distorted due to small strains resulting from embedding the experimental lattice in its reference lattice. We calculate the Euler angles of experimentally observed structures by rotating a rigid tetrahedral molecule in the reference lattice frame to minimize its atomic distances compared to the fractional coordinates of the experimental molecule. The reference lattice molecular positions are also shifted to coincide with the centers of mass of the experimentally determined fractional cell atoms. Thus the Euler angles obtained are a "best fit" to the molecular rotations were the experimental structures exactly proportional to their reference-lattice-derived ideal lattices. These Euler angles are presented in the supplementary material for our reference lattice set. The reference configuration of the molecule is with three-fold axes pointing in the [111],  $[\bar{1}\bar{1}1]$ ,  $[\bar{1}1\bar{1}]$ , and  $[1\bar{1}\bar{1}]$  directions. Typically only one atom lying along each of the three-fold axes was used when determining molecular orientations. Euler angles are not unique due to the  $2\pi$  modulus in the definition of  $\{\alpha, \beta, \gamma\}$ and the point group symmetry of the tetrahedral molecule. Once the Euler angles are determined, the rotator functions  $U_{m_{\tau}n_{\sigma}}^{\ell_{i}}$  may be calculated directly using Eq. (2). These are listed in the supplementary material.  $U_{m_{\tau}n_{\sigma}}^{\ell_{i}}$  is fully symmetric with respect to point group symmetry and is not influenced by non-unique Euler angles. Substituting the  $U_{m_{\tau}n_{\sigma}}^{\ell_i}$  into Eq. (1) yields the interaction potential as a function of the potential parameters  $\nu$ . This gives the energy  $E = V(\nu)$  as a linear polynomial in  $\nu$  such as

$$E = 0.2901 \,\nu_{0,4,4} - 0.0028 \,\nu_{3,0,3} + 0.0036 \,\nu_{3,1,4}$$

$$- 0.1020 \,\nu_{3,2,3} - 0.0425 \,\nu_{3,3,4} - 0.0127 \,\nu_{3,4,3}$$

$$+ 0.0053 \,\nu_{3,5,4} - 0.0006 \,\nu_{3,6,3} + 0.0071 \,\nu_{3,7,4}$$

$$- 0.0055 \,\nu_{4,6,4} - 0.0183 \,\nu_{4,8,4} + 1.2640 \,\nu_{4,0,4}$$

$$+ 0.0109 \,\nu_{4,2,4} + 0.0316 \,\nu_{4,4,4}, \tag{8}$$

for tetrakis(trimethystannyl)silane [CSD structure MEZDIE01], which crystallizes in space group 2.

# 5. Minimum Energy Gap

To find a representative vector  $\boldsymbol{\nu}$  for which each experimental phase is the lowest energy structure (i.e. a phase with an energy such as Eq. (8)), we propose a minimum energy difference algorithm. First we project the  $\boldsymbol{\nu}$  vector onto the unit hypersphere since only relative  $\boldsymbol{\nu}$  magnitudes affect the phase as  $T \to 0$ . Then taking the library of structures discussed previously in this section, we minimize the difference of the energy of the target structure  $E_{target}$  and the minimal energy structure in the collection  $\{E_{lib}(\boldsymbol{\nu}, \boldsymbol{\omega})\}$ . Thus we seek the vector  $\nu_{RP}$  that minimizes

$$\Delta E = E_{target}(\boldsymbol{\nu}) - \min\{E_{lib}(\boldsymbol{\nu}, \boldsymbol{\omega})\}. \tag{9}$$

This gives the point in  $\nu$  space of the *largest* energy gap between  $E_{\text{target}}$  and any other structures in the library and is representative of the family of intermolecular potentials consistent with the target structure. For an exhaustive library,  $\Delta E$  is bounded below by 0 since  $\{E_{lib}(\nu,\omega)\}$  contains the target structure and is minimized with respect to  $\omega$  while the target structure is held at fixed molecular orientation. We call it the representative parameter vector  $\nu_{RP}$ .

While  $E_{\text{target}}$  is based on observed Euler angles  $\omega$  and is only a function of  $\nu$  (i.e. Eq. (8)),  $E_{lib}$  contains a library of structures whose energies are functions of  $\nu$  and  $\omega$ . Since minimizing with respect to  $\omega$  is computationally demanding, we solve Eq. (9) in an iterative manner. First Eq. (9) is minimized for each experimental lattice with  $\{E_{lib}\}$  based only on the energies of the remaining experimental lattices. For each trial  $\nu_{RP}$  found from this

relative minimization, all additional candidate lattices are minimized with respect to their Euler angles at fixed  $\nu$  and the lowest energy solution is appended to the library. Again Eq. (9) is minimized for the energy of each experimental structure with respect to the augmented set to find a new trial RP in  $\nu$ -space and the process iterates until the RP's converge.

Minimization methods using genetic algorithms (differential evolution) and simulated annealing [21] are used on the Euler angle minimization and energy difference minimization, respectively. This is particularly important since the topologies of the energy and energy difference are both nonlinear with many local minima, the latter also having discontinuous derivatives. We reiterate this is a relative energy minimization for  $\Delta E$  in parameter space  $\nu$  and not a simulated annealing with fixed potential and varying temperature as is more commonly done.

We anticipate that the lower manifold basis functions should be more important in the potential because the kinetic energy of molecular orbitals pertaining to these angular momenta is lower. However, the above scheme does not discriminate between  $\nu_{\ell_i,\ell_i,\ell_j}$  corresponding to lower or higher manifolds. To favor lower manifolds the energy difference is minimized for  $\ell_{\text{max}} = 3$  for all experimental structures. Only if a minimum is not found for a given structure are fourth manifold basis functions appended to the parameter space and the process repeated.

## III. DISCUSSION AND CONCLUSION

#### A. Representative Potentials

Calculating potential parameters according to the procedure outlined in Sec. II gives the results in Table III where the potential coefficients are given for experimental structures in the five most prevalent reference lattices. Calculations for other reference lattices such as Aa or A15 are not reported here since they represent a smaller percentage of distinct structures. The difference in energy between the experimental structure and the next lowest energy structure among the candidate lattices is shown in the second column.

For 12 of the 30 structures,  $\Delta E = 0$ . This indicates the target structure is in the library and  $\omega_{\rm exp} \approx \omega_{\rm min}$ . Since  $\Delta E = 0$  throughout the region where the target is stable,  $\nu_{RP}$  is

an arbitrary point in the region and is not necessarily centrally located.

For 11 of the 30 structures,  $\Delta E < 0$ . This indicates the target structure is not in the library. Since  $\Delta E = 0$  on the borders, but  $\Delta E < 0$  in the interior,  $\nu_{RP}$  gives a set of potential parameters where the target structure is relatively stable and is expected to be near the center of the region.

For 5 of the 30 structures,  $0 < \Delta E < 0.01$  and therefore a representative potential for these structures has likely been found. This can be asserted because the potential is composed of products of  $U_{m_{\tau}n_{\sigma}}^{\ell_i}(\omega_i)$ , Clebsch-Gordan coefficients, spherical harmonics, and unit sphere normalized potential coefficients  $\nu$  that are all unitary. This makes the energy and therefore energy difference to be  $\mathcal{O}(1)$ . An energy difference less than this tolerance likely results from a slight mismatch in the Euler angles  $\omega_{\text{exp}}$  and  $\omega_{\text{min}}$  since the minimizer is free to alter the angle to attain the lowest possible energy. Thus  $\nu_{RP}$  is in the correct region but not necessarily centrally located and the target structure is in the library.

For 2 of 30 structures,  $\Delta E > 0.01$ . This indicates the absence of potential parameters consistent with the experimental crystal structure relative to the other structures using the truncation specified. The  $\nu_{RP}$  listed identifies a point in parameter space where the target structure is least metastable compared to the global minimum in the library. The target structure is also usually quite similar to the global minimal structure. This is the case with YEMRIR, for example. It has a positive  $\Delta E$  but is similar to the global minimum structure, space group 132 with molecules at Wyckoff point a. Both have two molecules per unit cell and one of the molecules, that at (0,0,0) in both structures, has the same orientation. The difference is the second molecule which is rotated  $\sim 90^{\circ}$  between the two structures. Reasons for this difference may be related to truncation of the potential after the fourth manifold, lack of molecular displacements and lattice strains in the model, lack of explicit second nearest-neighbor or three-body interactions, or missing quantum effects. Another reason may be that the reported experimental phase is in fact metastable in the limit  $T \to 0$ . Ostwald's step rule supports this, which is that during crystal nucleation and growth a metastable phase frequently crystallizes first which is closest to the liquid and which has the lowest free energy barrier. The crystallite then undergoes a series of metastable modifications until it reaches the thermodynamically stable structure [22]. The experimentally observed structure may have formed during an early stage of nucleation as one of these intermediate structures.

14 of the 30 distinct structures can be located in potential parameter space using only third manifold basis functions while 16 require fourth manifold functions. This is contrary to previous suggestions [8] that many manifolds are needed to accurately describe tetrahedral molecules such as methane and adamantane. However this supports current work [23] showing that fewer manifolds are necessary if the coefficients of the basis functions are evaluated to reflect primarily low energy configurations rather than the common evaluation method that equally weights all configurations. Current work is a proof by demonstration. Although visualization of high dimensional spaces is difficult, software such as GGOBI has made this possible in an interactive way. [24]. Any number of axes may be displayed and spatial relationships between the RP's analyzed.

Interpreting the potential coefficients in Table III is analogous to describing multipole interactions. These multipoles include contributions from all interaction modes and are not limited to electrostatic interactions. In the case of tetrahedral molecules the octopole  $(\ell_i = 3)$  is the first nonzero multipole. Coefficients such as  $\nu_{0,3,3}$  and  $\nu_{0,4,4}$  where  $\ell_i$  or  $\ell_j$  is zero represent an octopole or hexadecapole interacting with the zeroth pole and are the crystal field coefficients. As the basis functions may also be used as quantum basis sets, the  $\nu$  may be calculated ab initio and have been given physical interpretations via symmetry-adpated perturbation theory [9]. For instance, although dispersion and induction forces can be found in all components of  $\nu$ , electrostatic forces contribute only to  $\nu_{\ell_i,\ell_i+\ell_j,\ell_j}^{n_\sigma,n_\mu}$  [25] such as  $\nu_{3,6,3}$ ,  $\nu_{3,7,4}$ , and  $\nu_{4,8,4}$ .

One drawback of using the algorithm in Sec. II B is that, by finding the maximum energy difference between a phase and all others, the RP's of neighboring phases tend to be spread apart. This is because an RP is a vector representing a region in a space, not a unique set of IP parameters. Homologous series of molecules (i.e.  $CF_4$ ,  $CCl_4$ ,  $CBr_4$ ,  $CI_4$ ) are expected to show trends in  $\nu$  space. If members of homologous series have different crystal structures, then they have different RP's. However, different RP's are widely spread by our algorithm. Therefore RP's of homologous series are often widely spaced, even if the molecules are expected to have similar intermolecular potentials. This hides the expected trends among homologues.

Correlations are evident in reference lattice assignments. An example of this is a series of molecules with molecular formula  $C_{16}H_{36}X_4In_4N_4$  where X is Cl, Br, or I. The first two structures, 14e MECKUA and 12i MECKOU, pack in an fcc reference lattice while the

last, 11e MECKIO, packs in a tetragonal reference lattice intermediate between fcc and bcc which is slightly closer to the bcc reference lattice. This indicates at least the translational part of homologue potentials is similar and that small changes in the atomic constituency can slightly alter the intermolecular potential (IP), but significantly alter crystal structure. This is consistent with experience that similar molecules often have very different crystal structures despite similar intermolecular potentials. GPDs acknowledge this by placing the seemingly disparate structures close to one another in IP parameter space.

In [2] we noted that structures 195a and 197a were very similar to higher symmetry structures (215a and 217a). If the molecules remained tetrahedral then the crystal would retain higher symmetry but the reported atomic coordinates indicate a minor molecular distortion reducing the space group symmetry. Assuming that the reported space group is correct, these structures require a symmetry-breaking pathway from sc and bcc with a minimal truncation manifold of  $\ell_i^{\text{max}} = 6$  or 9 (depending on the potential-dependent transition pathway taken). In contrast to these high manifold requirements most pathways require a third or fourth manifold basis set. In view of the success of finding RP's using only the two first manifolds ( $\ell_i^{\text{max}} = 3$  or 4) for all other molecules in the data set, it seems unlikely such a large number of rapidly oscillating basis functions would be required to properly describe the intermolecular potential for these structures. It seems more likely that, barring a Jahn-Teller crystal distortion, the crystal symmetry may have been underspecified when reported to the CSD.

#### B. Extensions and Features of the Methodology

We now discuss some of the ways to improve the model and algorithms. An issue affecting the numerical accuracy of the RP is how large a library of alternate crystal structures is needed to localize the RP in  $\nu$  space. In our previous work [1] we considered just the high-symmetry point isotropy subgroups and in this work have followed suit since these are the most common [18]. Recall that space group IR's are indexed by reciprocal space vectors [26, 27]. There are the same number of k points in reciprocal space as there are unit cells in the crystal and there is a correspondence between k points and supercell patterns in the crystal. Experimental structures are supercells in their reference lattice. As nearly all experimental structures contain a relatively small number of clustered parent lattice

unit cells as sublattices, symmetry-breaking occurs at k vectors corresponding to this small cluster. These are k vectors at high symmetry points in the reference lattice Brillouin zone. If the experimental structure has a unit cell which is large or flattened/elongated, however, k vectors corresponding to this larger or longer/flatter group of parent lattice unit cells will be on high symmetry lines and planes. Symmetry breaking pathways for experimental lattices pertaining to been and fee reference lattices in the appendix show that some experimental structures contain large or non-clustered parent lattice unit cells such as 15f,f,f,f or 152b and their k vectors therefore are from high-symmetry plane and line IR's. Although these cases are less common, such isotropy subgroups could be included in the candidate lattices in Sec. IIB3. Providing such additional structures in the library would place additional constraints on the RP for the observed structures and therefore would further localize the RP for each observed structure at the expense of a much larger library.

In Sec. IIB3 we chose to consider only isotropy subgroups with one occupied Wyckoff point. This is a commonly used simplification. [13] We have also discarded coupled IR isotropy subgroups for the same reason. Our method could be extended to include library structures with multiple Wyckoff points and those from coupled IRs, although the computational demands are much higher because of the larger set of candidate lattices and so we do not pursue it here. The effect would be to further localize the RPs of each phase, again at the expense of a much larger library.

Although tetrahedral molecules are used in the current example to reverse engineer the IP, any molecular point group could be used without a dramatic increase in the number of potential coefficients  $\nu$  in the first two non-trivial manifolds. This is because the absence of IP coefficients in lower manifolds for high-symmetry molecules is offset by the larger number of parameters in higher manifolds. Consider the number of basis functions in the first two non-trivial manifolds of  $I_h$  (the icosahedral group) and  $C_1$  (the point group of no special symmetry). These are the highest and lowest molecular point group symmetries, respectively. The first three manifolds of  $I_h$  containing a totally symmetric molecular representation are the zeroth, sixth, and tenth manifolds. The number of potential coefficients is seven on the sixth manifold,  $\{\nu_{1,0,0}^{1,1}, \nu_{6,2,6}^{1,1}, ... \nu_{6,12,6}^{1,1}\}$ , eleven on the tenth manifold,  $\{\nu_{10,0,10}^{1,1}, \nu_{10,2,10}^{1,1}, ... \nu_{10,20,10}^{1,1}\}$ , and seven for the cross manifolds,  $\{\nu_{6,4,10}^{1,1}, \nu_{6,6,10}^{1,1}, ... \nu_{6,16,10}^{1,1}\}$ . There are also two crystal field coefficients,  $\{\nu_{0,6,6}^{1,1}, \nu_{0,10,10}^{1,1}\}$ , giving a total of 27 potential coefficients for  $I_h$ . For  $C_1$  the first three manifolds are the zeroth, first, and second. The coefficients on the

first manifold are  $\{\nu_{1,0,1}^{n_{\sigma},n_{\mu}}, \nu_{1,2,1}^{n_{\sigma},n_{\mu}}\}$ . Although there are three copies of the totally symmetric molecular representation on the first manifold and therefore the molecular frame indices  $\sigma$  and  $\mu$  in Eq. (1) go over  $\{1,2,3\}$ , we are free to choose the standard orientation for the molecule corresponding to Euler angles  $\{0,0,0\}$ . If it is chosen with the IP major axis parallel to the laboratory z-axis then only one of these is nonzero. This leaves two coefficients  $\{\nu_{1,0,1}^{1,1}, \nu_{1,2,1}^{1,1}\}$ . If the molecular minor axis in the standard orientation is oriented parallel to the laboratory x-axis only four of the  $\sigma$  and  $\mu$  are nonzero in the second manifold. This leaves 48 coefficients on the second manifold,  $\{\nu_{2,0,2}^{n_{\sigma},n_{\mu}}, \nu_{2,2,2}^{n_{\sigma},n_{\mu}}, \nu_{2,4,2}^{n_{\sigma},n_{\mu}}\}$ . With five additional crystal field coefficients  $\{\nu_{1,1,1}^{1,1}, \nu_{0,2,2}^{1,n_{\mu}}\}$  and eight cross manifold coefficients  $\{\nu_{1,1,2}^{1,n_{\mu}}, \nu_{1,3,2}^{1,n_{\mu}}\}$  the total number of coefficients on the first three manifolds of  $C_1$  is 60, an increase of roughly twofold from highly symmetrical  $I_h$ . This shows that although this method has been applied to tetrahedral molecules, it is applicable to other molecular point group symmetries with only a modest change in the number parameters.

Throughout we have implemented a simple direct cutoff truncation scheme in which a doubly infinite summation is cut off at a maximum manifold number  $\ell_i^{\text{max}}$ . An alternative is the manifold-sum cutoff in which we truncate such that  $\ell_i + \ell_j \leq \ell^{\text{max}}$ . This truncation would include smoother functions before more rapidly oscillating ones which is consistent with our expectations of lower-energy electronic contributions to the potential. Also the number of potential parameters increases at a slower rate with this truncation. This is shown in Table IV for both truncation schemes where we compare the cumulative number of parameters for the  $C_1$ ,  $T_d$ , and  $I_h$  point groups at different manifolds. The truncation method used in this study is a square truncation of the double sum while the alternative is a triangular truncation. The manifold sum truncation adds new parameters into the potential more slowly than single manifold truncation. Therefore, the 15 coefficients used here are sufficient but may not be necessary. Further investigation of global phase diagrams with  $\ell$ -sum truncation is needed to test this hypothesis. This is important since lower dimensional GPDs would be easier to construct and to use.

From the foregoing discussion we have seen 15 coefficients are sufficient to reverse engineer our dataset of experimental structures. We have not investigated if a linear combination would be better. Principle component analysis could be used to identify linear combinations of basis functions that better fit molecules. It is possible that less than 15 coefficients are necessary.

#### C. Conclusions

We have shown that a molecular crystal global phase diagram (GPD) with a small number of reference lattices derived in [2] can summarize the experimental data using a modest number of IP parameters. The data set is diverse enough to test the GPD's ability to classify a wide range of space groups using a common intermolecular potential. Just as the van Konynenburg global phase diagram classification based on the simple van der Waals equation of state is nonetheless widely used to classify the phase behavior of real binary mixtures, molecular crystal global phase diagrams may be useful in elucidating phase behavior of a variety of real substances and, in turn, used to develop novel intermolecular potentials and materials.

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TABLE I: Presence of  $O_h$ ,  $D_{3h}$ , and  $D_{2d}$  point group IR's for various manifolds of SO(3).  $O_h$  is the Wyckoff point group of the fcc, bcc, and sc reference lattices.  $D_{3h}$  is the Wyckoff point group of the hcp reference lattice.  $D_{2d}$  is the Wyckoff point group of the A5' reference lattice. The first occurrence of a given IR is shown in bold.

$\ell_i O_h \text{ IR's}$	$D_{3h}$ IR's	$D_{2d}$ IR's
$0   {m A_{1g}}$	$A_1'$	$A_1$
$3 A_{2u} + T_{1u} + T_{2u}$	$A_1^\prime + \boldsymbol{A_2^\prime} + \boldsymbol{E^\prime}^\prime + \boldsymbol{E^{\prime\prime}}^\prime$	$A_1 + \boldsymbol{A_2} + \boldsymbol{B_2} + 2\boldsymbol{E}$
$4 A_{1g} + E_g + T_{1g} + T_{2g}$	$A_1' + A_1'' + A_2'' + 2E' + E''$	$2A_1 + A_2 + B_1 + B_2 + 2E$
$6 A_{1g} + A_{2g} + E_g + T_{1g} + 2T_{2g}$	$2A_1' + A_1'' + A_2' + A_2'' + 2E' + 2E$	$"2A_1 + A_2 + 2B_1 + 2B_2 + 3E$
$7 A_{2u} + \mathbf{E_u} + 2T_{1u} + 2T_{2u}$	$A_1' + A_1'' + A_2' + 2A_2'' + 3E' + 2E$	" $2A_1 + 2A_2 + B_1 + 2B_2 + 4E$
$8 A_{1g} + 2E_g + 2T_{1g} + 2T_{2g}$		
$9 \ \mathbf{A_{1u}} + A_{2u} + E_u + 3T_{1u} + 2T_2$	u	

TABLE II: Group theoretical symmetry-breaking pathways of experimental lattices for the fcc reference lattice. Classified by space group IR and order parameter direction, each pathway shows the point group IR and minimal manifold of SO(3) in Eq. (1) required to achieve it. The order parameter directions are given in an abbreviated form in the notation of Stokes and Hatch [28]. The pathways, space group IRs, order parameter directions, and point group IRs were computed using ISOTROPY. However, transitions belonging to a coupled IR between a high symmetry point and line are currently not a feature of ISOTROPY. These entries have been marked with an asterisk in the table.

Pathway	S. G. IR	OP Dir.	P. G. IR	$\ell^{\mathrm{req'd}}$
$225a\rightarrow152b$	$\Lambda_3$ (k=1/3)	P1	$E_g, T_{1g}, T_{2g}, E_u, T_{1u}, T_{2u}$	3
$225a\rightarrow142a$	$W_3$	P2	$T_{1g}$ , $A_{2u}$ , $E_u$	3
$225a \rightarrow 121a$	$\Gamma_5^-$	P1	$T_{2u}$	3
$225a \rightarrow 114a$	$X_2^- \oplus \Gamma_5^-$	$P1 \oplus P1$	$A_{2u}, E_u \oplus T_{2u}$	3
	$X_3^+ \oplus \Gamma_5^-$	$P1 \oplus P1$	$T_{1g} \oplus T_{2u}$	4
	$X_2^- \oplus X_3^+$	$P1\oplusP1$	$A_{2u}, E_u \oplus T_{1g}$	4
$225a \rightarrow 64d,f$	*			
$225a \rightarrow 15f,f,f,f$	$C_2 \ (k_1 = 1/4, k_2 = 3/4)$	C18,C19	$A_{2g}, E_g, T_{1g}, T_{2g},$	3
			$A_{1u}, E_u, T_{1u}, T_{2u}$	
$225a \rightarrow 15e (REKYUB)$	$L_3^-$	P7	$E_u, T_{1u}, T_{2u}$	3
$225a \rightarrow 14e \text{ (MECKUA)}$	$\Gamma_3^+ \oplus L_3^-$	$\mathrm{P1} \oplus \mathrm{C8}$	$E_g \oplus E_u, T_{1u}, T_{2u}$	4
	$\Gamma_3^+ \oplus X_5^-$	$\mathrm{C1}\oplus\mathrm{S7}$	$E_g \oplus T_{1u}, T_{2u}$	4
	$\Gamma_4^+ \oplus X_5^-$	$P1\oplusC11$	$T_{1g} \oplus T_{1u}, T_{2u}$	4
		$P1\oplusS7$		
	$\Gamma_5^+ \oplus L_3^-$	$C2 \oplus C8$	$T_{2g} \oplus E_u, T_{1u}, T_{2u}$	4
	$\Gamma_5^+ \oplus X_5^-$	$P1\oplusC11$	$T_{2g} \oplus T_{1u}, T_{2u}$	4
		$P1\oplusS7$		
	$L_2^+ \oplus L_1^-$	$P1\oplusP1$	$A_{2g},T_{1g}\oplus A_{1u},T_{2u}$	4
	$L_2^+ \oplus L_3^-$	$P1\oplusP7$	$A_{2g},T_{1g}\oplus E_u,T_{1u},T_{2u}$	4
	$L_2^+ \oplus X_2^-$	$P1\oplusP1$	$A_{2g},T_{1g}\oplus A_{2u},E_{u}$	4
	$L_2^+ \oplus X_3^-$	$P1 \oplus P1$	$A_{2g},T_{1g}\oplus E_u,T_{1u}$	4
	$L_2^+ \oplus X_5^-$	$P1\oplusP1$	$A_{2g},T_{1g}\oplus T_{1u},T_{2u}$	4
	$L_3^+ \oplus L_1^-$	$P7\oplusP1$	$E_g, T_{1g}, T_{2g} \oplus A_{1u}, T_{2u}$	4
	$L_{3}^{+} \oplus L_{3}^{-}$	$P7\oplusP7$	$E_g, T_{1g}, T_{2g} \oplus E_u, T_{1u}, T_{2u}$	4
	$L_3^+ \oplus X_2^-$	$P7\oplusP1$	$E_g, T_{1g}, T_{2g} \oplus A_{2u}, E_u$	4
	$L_3^+ \oplus X_3^-$	$P7 \oplus P1$	$E_g, T_{1g}, T_{2g} \oplus E_u, T_{1u}$	4
	$L_3^+ \oplus X_5^-$	$P7 \oplus P1$	$E_g, T_{1g}, T_{2g} \oplus T_{1u}, T_{2u}$	4
	$L_1^- \oplus L_3^-$	$\mathrm{P1}\oplus\mathrm{C8}$	$A_{1u}, T_{2u} \oplus E_u, T_{1u}, T_{2u}$	4
	$L_1^- \oplus X_2^-$	$P1\oplusP1$	$A_{1u}, T_{2u} \oplus A_{2u}, E_u$	4
	$L_1^- \oplus X_3^-$	$P1\oplusP1$	$A_{1u}, T_{2u} \oplus E_u, T_{1u}$	4
	$L_1^- \oplus X_5^-$	$P1\oplusP1$	$A_{1u}, T_{2u} \oplus T_{1u}, T_{2u}$	4
	$L_3^- \oplus X_2^-$	$P7\oplusP1$	$E_u, T_{1u}, T_{2u} \oplus A_{2u}, E_u$	4
	$L_3^- \oplus X_3^-$	$P7\oplusP1$	$E_u, T_{1u}, T_{2u} \oplus E_u, T_{1u}$	4
	$L_3^- \oplus X_5^-$	$P7\oplusP1$	$E_u, T_{1u}, T_{2u} \oplus T_{1u}, T_{2u}$	4
	$X_2^- \oplus X_5^-$	P1 ⊕ C11	$A_{2u}, E_u \oplus T_{1u}, T_{2u}$	4
		$P1\oplusS7$		
	$X_3^- \oplus X_5^-$	P1 ⊕ C11	$E_u, T_{1u} \oplus T_{1u}, T_{2u}$	4
$225a \rightarrow 14e \text{ (TOHSUE)}$	$\Delta_5 \ (k=1/4)$	C7	$T_{1g}, T_{2g}, T_{1u}, T_{2u}$	3
225a → 12i	$L_3^-$	P2	$E_u, T_{1u}, T_{2u}$	3

TABLE III: RP components for crystals of tetrahedral molecules in the CSD. The  $\ell_i$  truncation value and the presence, absence, or proximity of a global minimum are indicated. The parameters  $\nu_{\ell_i,\ell,\ell_j}$  are indexed according to Eq. (1) and have been mapped to the unit hypersphere.

Identifier	$\Delta \mathrm{E}$	$\nu_{033}$	$\nu_{044}$	$\nu_{303}$	$\nu_{323}$	$\nu_{343}$	$\nu_{363}$	$\nu_{314}$	$\nu_{334}$	$\nu_{354}$	$\nu_{374}$	$\nu_{404}$	$\nu_{424}$	$\nu_{444}$	$\nu_{464}$	$\nu_{484}$
bcc																
$217a~\rm HXMTAM07$	0.000	-	0.000	0.970	0.000	0.230	-0.079	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000
161a TCYMET	0.000	-	0.167	0.012	-0.075	0.166	-0.0823	0.039	0.123	-0.168	0.269	0.067	0.072	-0.403	0.165	0.784
2i MEZDIE01	0.000	-	-0.074	-0.008	-0.191	0.107	-0.234	-0.527	-0.113	-0.198	-0.015	-0.655	-0.162	0.108	0.014	0.309
60c,d YIMWEW	-0.044	-	-0.086	0.114	-0.225	-0.023	0.017	-0.207	-0.404	0.696	0.312	0.095	0.048	-0.167	0.054	0.314
2iii OHABEE	0.004	-	-0.158	-0.002	-0.280	-0.011	-0.006	0.429	0.197	0.416	-0.178	-0.172	0.243	0.494	0.033	0.369
fcc																
121a ZZZKDW $01$	0.000	-	0.000	0.23	0.066	0.32	-0.916	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000
$142a~\mathrm{KUJSIR}$	0.000	-	0.000	-1.	-0.008	-0.004	-0.027	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000
114a ADAMAN08*	0.002	-	0.000	-0.076	0.258	-0.437	-0.858	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000
152b MTRETC10	-0.070	-	-0.202	0.020	-0.060	0.104	-0.144	-0.858	0.121	0.241	-0.214	-0.002	-0.191	-0.021	-0.171	0.048
15e REKYUB	0.002	-	-0.085	0.000	0.095	-0.002	0.001	0.458	-0.226	-0.523	-0.514	-0.191	0.368	0.097	0.034	0.055
12i MECKOU	0.000	-	-0.19	0.007	-0.634	0.044	-0.106	0.172	0.313	0.082	0.372	-0.439	0.109	0.234	-0.058	-0.118
64d,f METHANEIII	-0.239	-	0.087	-0.026	0.014	-0.025	0.016	0.000	-0.024	-0.007	-0.001	0.1	0.067	-0.751	-0.54	-0.346
14e MECKUA	0.014	-	-0.112	-0.018	0.023	0.038	-0.322	-0.819	0.019	-0.233	0.121	0.022	0.084	-0.306	-0.194	0.06
14e TOHSUE	-0.011	-	0.069	-0.005	-0.065	-0.016	-0.163	0.805	0.048	-0.37	0.381	-0.007	-0.1	-0.135	-0.005	0.057
15f,f,f,f CTBROM	-0.185	-	0.041	0.025	-0.053	0.025	-0.062	-0.775	0.352	-0.091	0.458	0.013	0.072	0.074	-0.117	0.153
hcp																
165d DILWIE01	-0.037	-0.772	0.000	0.096	0.303	0.443	-0.326	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000
147d ZIZHIZ	0.000	-0.781	0.000	-0.121	0.198	0.147	-0.56	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000
176h CUCZUV	0.008	0.002	0.000	-0.233	-0.941	0.246	-0.003	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000
14e CARBTC	-0.032	-0.146	0.000	-0.054	0.438	-0.457	0.758	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000
sc																
215a FOHCUA	0.000	_	0.000	0.960	0.000	0.279	-0.017	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000	0.000
120c YEMRIR	0.016	-	-0.102	-0.111	-0.167	-0.439	0.077	0.403	-0.487	-0.078	0.281	0.169	0.056	-0.071	0.449	-0.167

TABLE IV: Comparison of the cumulative number of parameters in potential for a truncation at a given manifold for two truncation strategies and three molecular point groups.

Direct Truncation  $\ell$  sum Truncation

$\ell_i$	$C_1$	$T_d$	$I_h$	$\frac{\ell_i + \ell_j}{}$	$C_1$	$T_d$	$I_h$
0	*	*	*	0	*	*	*
1	3	0	0	1	1	0	0
2	60	0	0	2	6	0	0
3		5	0	3	15	1	0
4		10	0	4		1	0
6			8	6		6	1
10	١		19	10			8

<sup>\*</sup> There is one isotropic basis function for  $\ell_i=\ell_j=0$  in each case, but it does not drive orientational ordering.

TABLE V: Group theoretical symmetry-breaking pathways of experimental lattices from the bcc reference lattice. Classified by space group IR and order parameter direction, each pathway shows the point group IR and minimal manifold of SO(3) in Eq. (1) required to achieve it. The order parameter directions are given in an abbreviated form in the notation of Stokes and Hatch [28].

Pathway	S. G. IR	OP Dir.	P. G. IR	$\ell^{\mathrm{req'd}}$
229a → 217a	$\Gamma_2^-$	P1	$A_{2u}$	3
$229a \rightarrow 161a$	$H_5^- \oplus \Gamma_2^-$	P3 ⊕ P1	$T_{2u} \oplus A_{2u}$	3
	$H_5^- \oplus \Gamma_4^-$	P3 ⊕ P3	$T_{2u} \oplus T_{1u}$	3
	$H_{5}^{-} \oplus H_{4}^{+}$	P3 ⊕ P3	$T_{2u} \oplus T_{1g}$	4
	$H_4^+ \oplus \Gamma_2^-$	P3 ⊕ P1	$T_{1g} \oplus A_{2u}$	4
	$H_4^+ \oplus \Gamma_4^-$	P3 ⊕ P3	$T_{1g} \oplus T_{1u}$	4
	$H_{5}^{-} \oplus H_{2}^{+}$	P3 ⊕ P1	$T_{2u} \oplus A_{2g}$	6
	$H_2^+ \oplus \Gamma_4^-$	$P1 \oplus P3$	$A_{2g} \oplus T_{1u}$	6
	$H_1^- \oplus \Gamma_4^-$	$P1 \oplus P3$	$A_{1u} \oplus T_{1u}$	9
	$H_{1}^{-} \oplus H_{4}^{+}$	$P1 \oplus P3$	$A_{1u} \oplus T_{1g}$	9
$229a \rightarrow 60c,d$ (YIMWEW)	*			
$229a \rightarrow 2i \text{ (MEZDIE01)}$	$N_1^- \oplus \Gamma_4^+$	$P1 \oplus S1$	$A_{1u},E_u,T_{2u}\oplus T_{1g}$	4
	$N_1^- \oplus \Gamma_5^+$	$P1\oplusS1$	$A_{1u},E_u,T_{2u}\oplus T_{2g}$	4
	$N_2^- \oplus \Gamma_4^+$	$P1\oplusS1$	$A_{2u},E_u,T_{1u}\oplus T_{1g}$	4
	$N_2^- \oplus \Gamma_5^+$	$P1\oplusS1$	$A_{2u},E_u,T_{1u}\oplus T_{2g}$	4
	$N_3^- \oplus \Gamma_4^+$	$P1\oplusS1$	$T_{1u}, T_{2u} \oplus T_{1g}$	4
	$N_3^- \oplus \Gamma_5^+$	$P1\oplusS1$	$T_{1u},T_{2u}\oplus T_{2g}$	4
	$N_4^- \oplus \Gamma_4^+$	$P1\oplusS1$	$T_{1u},T_{2u}\oplus T_{1g}$	4
	$N_4^- \oplus \Gamma_5^+$	$P1\oplusS1$	$T_{1u},T_{2u}\oplus T_{2g}$	4
$229a \rightarrow 2iii$	$H_4^-$	S1	$T_{1u}$	3
	H5-	S1	$T_{2u}$	3

TABLE VI: Group theoretical symmetry-breaking pathways of experimental lattices for the hcp reference lattice.

Pathway	S. G. IR	OP Dir.	P. G. IR	$\ell^{\mathrm{req'd}}$
194c → 176h	$K_4$	P1	E'	3
$194c\rightarrow165d$	$A_2$	P3	$A'_{2}, A'_{1}$	3
$194c\rightarrow147d$	$\Gamma_3^+ \oplus \Gamma_2^+$	P1⊕ P1	$A_2^{\prime\prime}\oplus A_2^\prime$	3
	$\Gamma_4^+ \oplus \Gamma_2^+$	P1⊕ P1	$A_1^{\prime\prime}\oplus A_2^\prime$	4
	$\Gamma_4^+ \oplus \Gamma_3^+$	P1⊕ P1	$A_1^{\prime\prime}\oplus A_2^{\prime\prime}$	4
$194c\rightarrow14e\mathrm{CARBTC}$	$M_2^+ \oplus \Gamma_6^+$	P1⊕ P1	$A_2^\prime, E^\prime \oplus E^{\prime\prime}$	3
	$M_2^+ \oplus \Gamma_6^+$	P1⊕ P1	$A_2', E' \oplus E''$	3
	$M_2^+ \oplus \Gamma_3^+$	P1⊕ P1	$A_2', E' \oplus A_2''$	4
	$M_4^+ \oplus \Gamma_3^+$	P1⊕ P1	$A_1^{\prime\prime}, E^{\prime\prime} \oplus A_2^{\prime\prime}$	4
	$M_4^+ \oplus \Gamma_5^+$	P1⊕ C1	$A_1^{\prime\prime},E^{\prime\prime}\oplus E^\prime$	4
	$M_4^+ \oplus \Gamma_6^+$	P1⊕ P1	$A_1^{\prime\prime},E^{\prime\prime}\oplus E^{\prime\prime}$	4
	$M_4^+ \oplus M_2^+$	P1⊕ P1	$A_1^{\prime\prime}, E^{\prime\prime} \oplus A_2^\prime, E^\prime$	4

TABLE VII: Group theoretical symmetry-breaking pathways of experimental lattices for the sc reference lattice.

Pathway	S. G. IR	OP Dir.	P. G. IR	$\ell^{\mathrm{req'd}}$
$221a \rightarrow 215a$	$\Gamma_2^-$	P1	$A_{2u}$	3
$221a\rightarrow120c$	$R_5^- \oplus \Gamma_2^-$	$\mathrm{P}1\oplus\ \mathrm{P}1$	$T_{2u} \oplus A_{2u}$	3
	$R_4^+ \oplus \Gamma_2^-$	$\mathrm{P}1\oplus\ \mathrm{P}1$	$T_{1g} \oplus A_{2u}$	4
	$R_5^- \oplus R_4^+$	$\mathrm{P}1\oplus\ \mathrm{P}1$	$T_{2u} \oplus T_{1g}$	4
	$R_4^+ \oplus \Gamma_3^-$	$\mathrm{P}1\oplus\ \mathrm{P}1$	$T_{1g} \oplus E_u$	7
	$R_5^- \oplus \Gamma_3^-$	$\mathrm{P}1\oplus\ \mathrm{P}1$	$T_{2u} \oplus E_u$	7

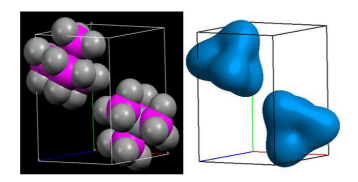


FIG. 1: Comparison of an experimental structure, tetrakis(trimethystannyl)silane [CSD structure MEZDIE01], which crystallizes in space group 2 at Wyckoff point i with arbitrarily-shaped tetrahedral figures whose orientation is determined by orientational energy minimization with molecular center of mass on an ideal bcc reference lattice.