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## **Designing Isotropic Interactions for Self-Assembly of Complex Lattices**

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We present a direct method for solving the inverse problem of designing isotropic potentials that cause self-assembly into target lattices. Each potential is constructed by matching its energy spectrum to the reciprocal representation of the lattice to guarantee that the desired structure is a ground state. We use the method to self-assemble complex lattices not previously achieved with isotropic potentials, such as a snub square tiling and the kagome lattice. The latter is especially interesting because it provides the crucial geometric frustration in several proposed spin liquids.

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Self-assembly [1], the process through which simple entities spontaneously form complex structures, is a phenomenon of fundamental interest and a compelling tool for future nanoscale fabrication. As technology is pushed to ever smaller length scales, direct fabrication becomes increasingly challenging and material design may turn towards tailoring interactions to cause constituents to order into the desired patterns without directed external influence. This vision has been the subject of intense experimental work during the past decades, but there is still a relative lack of theoretical results beyond pure simulationbased work.

What theoretical work exists can be understood as trying to solve an "inverse" statistical mechanics problem, finding interactions that cause self-assembly into a target structure or configurations with desired properties. Various schemes for approaching this problem have been proposed [2,3] and several isotropic potentials self-assembling into low-coordinated lattices have been reported, e.g., square, honeycomb, simple cubic, and diamond lattices [4]. The focus is on isotropic interactions, partly due to theoretical tractability but also motivated by the recent years' remarkable improvement in both understanding and design capabilities of such interactions in especially colloidal and nano systems [5,6]. By combining and tuning forces including, but not limited to, van der Waals, electrostatic, magnetic, and steric forces as well as forces of purely entropic origin, it is possible today to construct complex effective interactions. With such a profusion of possibilities, the forward statistical mechanics' prescription of sweeping the parameter space to give a complete picture of the possible structures becomes computationally infeasible, and the inverse formulation increasingly appealing.

In this Letter we present a new method for designing potentials for targeted self-assembly based on an energy spectrum approach. The method is direct in contrast to iterative or optimization approaches more commonly used to solve inverse problems. We show how interactions that cause particles to organize into the kagome lattice can be obtained. The kagome lattice is a much sought after topology due to its highly frustrated geometry [7,8], possibly resulting in spin liquid ground states in several materials [9,10] as well as in the corresponding Heisenberg model [11]. The self-assembly of nanoscale [12,13] and colloidal [14] kagome lattices through directed interactions has caused considerable interest, but whether this target structure is achievable via isotropic potentials has until now been an open question (cf. the negative result in [15]). To demonstrate the generality of our method, we also apply it to the honeycomb lattice and a snub square tiling [16].

We consider self-assembly governed by energy minimization where an interaction is designed with certain features so that the target structure is a stable low energy state, which should typically be reached when the system self-assembles from random initial configurations. Our method differs from previous ones in that we use a reciprocal representation of the lattice to construct a theoretical potential in a way that guarantees the target structure to be the unique ground state. The potential is then systematically simplified into a more realizable form. In general a lattice structure is defined by the small |**k**| region of the Fourier spectrum while the large **|k|** region only contributes with higher resolution of the distribution defining the particles. The kagome lattice for instance is clearly resolved (Fig. 1) when expressed as the sum of the Fourier modes of the (weighted) reciprocal lattice vectors up to length  $2|\mathbf{k}_p|$ , where  $\mathbf{k}_p$  is a primitive reciprocal lattice vector. This makes the reciprocal perspective a natural approach for targeted self-assembly of lattice structures, as opposed to real-space methods focusing on the pair correlation function [4].

In two dimensions the potential energy for a configuration with particle density  $\rho(\mathbf{r})$  can be expressed in reciprocal space as

$$E = \int d\mathbf{r}_1 d\mathbf{r}_2 \rho(\mathbf{r}_1) V(|\mathbf{r}_1 - \mathbf{r}_2|) \rho(\mathbf{r}_2) = \int dk |\hat{\rho}(k)|^2 \hat{V}(k),$$
(1)

where  $|\hat{\rho}(k)|^2 dk = \int_{|\mathbf{k}|=k} d\mathbf{k} |\hat{\rho}(\mathbf{k})|^2$ ,  $\hat{\rho}(\mathbf{k})$  is the standard Fourier transform of  $\rho(\mathbf{r})$ , and  $\hat{V}(k)$  is the radial Fourier (Hankel) transform of V(r) [17],



FIG. 1 (color online). The kagome lattice in real space (a) as described by its first few reciprocal lattice vectors (b). Structure factors for the included reciprocal vectors are shown in black and primitive lattice vectors  $\mathbf{k}_p$  as gray arrows. The kagome lattice (illustrated in black, with basis and lattice vectors in white) is indeed determined by this limited part of the reciprocal lattice.

$$\hat{V}(k) = 2\pi \int r dr V(r) J_0(rk), \qquad (2)$$

where  $J_0(x)$  is a Bessel function. If  $V(\mathbf{r}_1, \mathbf{r}_2) = V(|\mathbf{r}_1 - \mathbf{r}_2|)$  is interpreted as a linear operator, then Eq. (1) defines a diagonalization of  $V(\mathbf{r}_1, \mathbf{r}_2)$ . We therefore refer to  $\hat{V}(k)$  as the energy spectrum of the potential [18].

We use the power spectrum of the target lattice  $|\hat{\rho}(k)|$ and Eq. (1) to construct a potential that guarantees that the target is a ground state in the following way. Consider a potential V(r) whose energy spectrum is chosen positive,  $\hat{V}(k) \ge 0$ , and has zeros only at the frequencies coinciding with the reciprocal lattice points  $G_i$  of the Bravais lattice describing the target lattice,  $\hat{V}(|\mathbf{G}_i|) = 0$  for all *i*. This design guarantees that the target lattice is a ground state with E = 0. However, any lattice with reciprocal points being a subset of  $G_i$ , i.e., with arbitrary basis (as well as reciprocal lattices with opposite chirality), is also a ground state. This degeneracy can be broken in favor of the target lattice by introducing small perturbations of the energy spectrum at its zeros and using the fact that there are amplitude differences in the peaks of  $\hat{\rho}(\mathbf{G}_i)$  defining the target basis. The principle behind the choice of perturbations is that negative perturbations of the energy spectrum at frequencies where  $|\hat{\rho}(k)|$  of the target structure is large, and positive perturbations where  $|\hat{\rho}(k)|$  is small, exclude lattice bases with structure factors other than the target's as ground states.

In many cases, such as for the lattices considered in this Letter, it is possible to exclude the complete set of alternative bases and analytically construct a potential with the target lattice as the unique ground state. There are two characteristics of  $\hat{\rho}(\mathbf{k})$  we need to consider in these cases. For a closed system with *n* particles  $\hat{\rho}(\mathbf{0}) = n$  so  $\hat{V}(0)$  is irrelevant to the ground state configuration. More importantly, the mass provides a maximum for the amplitude  $|\hat{\rho}(\mathbf{k})| = |\sum_{j=1}^{n} e^{i\mathbf{k}\cdot\mathbf{r}_j}| \leq n$ , with equality for certain  $\mathbf{k} = \mathbf{G}_i$  if the basis of  $\mathbf{r}_j$  satisfies  $\mathbf{k} \cdot \mathbf{r}_j = 2\pi m + \phi$ , *m* being an integer and  $\phi$  an arbitrary phase. From Eq. (1) it

follows that a negative perturbation of the energy spectrum at a frequency where  $|\hat{\rho}(k)|$  is maximal limits the bases of the degenerate ground states to the finite number of configurations fulfilling the maximum amplitude constraint  $\mathbf{k} \cdot \mathbf{r}_j = 2\pi m + \phi$ . Smaller negative perturbations at the next highest peaks, or positive perturbations at the lowest peaks, further reduce the degeneracy. The hierarchy of perturbations then singles out the target lattice as the unique ground state.

We apply this design scheme to the kagome lattice. Consider an energy spectrum with zeros corresponding to the hexagonal Bravais lattice with primitive vectors  $\{(2, 0), (1, \sqrt{3})\}$ , which encompasses the kagome lattice [basis {(0, 0), (1, 0),  $(1/2, \sqrt{3}/2)$ ]. The maximum amplitude constraint is fulfilled at the reciprocal points  $|\mathbf{G}_{i}| = \frac{4\pi}{\sqrt{3}}$  only if the basis is a subset of {(0, 0), (1, 0),  $(1/2, \sqrt{3}/2), (3/2, \sqrt{3}/2)$ . Any subset of three of this set is a basis for the kagome lattice, assuming a density of approximately three particles per primitive cell. Including a hard core repulsion makes  $\mathbf{r}_{i} = \mathbf{r}_{l}, j \neq l$ , unfeasible without energetically benefiting any configuration nor affecting the kagome lattice. Therefore a negative perturbation of  $\hat{V}(k)$  at  $k = \frac{4\pi}{\sqrt{3}}$  combined with a hard core repulsion breaks the degeneracy in favor of the kagome lattice which becomes the unique ground state. An energy spectrum constructed by following these prescriptions is shown in Fig. 2(a).

A typical energy spectrum fulfilling the above constraints results in a highly complex potential, not realizable in experimental systems. However, the properties of the energy spectrum that select the ground state are far from a complete specification. This has dual consequences in that, on one hand, there is considerable leeway in varying experimental realizations for a target system, but, on the other hand, the prescription leaves a fairly large space of possible designs to be searched. While having no effect on the ground states, these choices can greatly affect the realizability of the potential as well as the extent to which a system of particles interacting with the potential can achieve the ground state under realistic dynamics. Reasonable requirements on the designed potentials include smoothness and restrictions on their range. To understand how this can be achieved we note that the Fourier relation in Eq. (2) implies that there exist versions of Heisenberg's uncertainty principle relating the smoothness of the potential in real space to a screening in reciprocal space, and vice versa. Several general principles for how the power spectrum should be designed follow from this observation. Restricting the power spectrum to the low  $\mathbf{k}$  region gives smoother potentials, evident in Figs. 2(a) and 2(b). Notice how the first few minima of the power spectrum are sufficient to determine the lattice periodicity. A screening of the power spectrum provides further smoothing of the potential [19]. Conversely, to get a short-ranged potential the spectrum



FIG. 2 (color online). A spectral method for targeted self-assembly. By designing an energy spectrum  $\hat{V}(k)$  (a) with minima at the peaks of  $\hat{\rho}$  and taking its Fourier transform we arrive at a potential V(r) (b) that causes particles to self-assemble into a target lattice. (c) Time evolution of a self-assembling kagome lattice. The number of grains diminishes by random walk of their boundaries, a process faster for smaller systems (d) but where some local defects are still present as the dimensions of the system is not fine-tuned to fit the lattice.

should be smooth; in Fig. 2(a) this is achieved using third orders splines as interpolations between the extreme points.

To test our method we perform Monte Carlo simulations with the constructed interaction potentials to confirm that they reliably and in a positive density range self-assemble into their respective target lattices. All simulations start in a random disordered state and we consider only local moves in the Monte Carlo updates. We do not fine-tune the dimensions of the simulated systems, therefore some deformations of the lattices are necessary for them to fit. The simulations confirm that the designed potentials are robust to such complications.

Figure 2(c) illustrates the relaxation process for the physically most interesting structure, the kagome lattice. The snapshots of the time evolution show how the lattice assemble locally into grains with boundaries that diffuse and slowly annihilate. The system size in Fig. 2(c) is chosen large enough to illustrate this process but makes complete relaxation too computationally demanding. In Fig. 2(d) we show the global ground state for a smaller system with the same potential. It should be noted that the kagome lattice is significantly more complex than lattices previously achieved through targeted self-assembly with isotropic interactions.

To demonstrate the range of our method, we also assemble a snub square tiling with isosceles triangles [16], shown in Fig. 3(a). This structure is even more complex than the kagome lattice since it has a square Bravais lattice and requires more perturbations to single out the target basis. We use perturbations at four points in the energy spectrum, three positive perturbations where  $|\hat{\rho}(|\mathbf{G}_i|)|^2 = 0$ , and one negative where  $|\hat{\rho}(|\mathbf{G}_i|)|^2 = n$ .

For completeness we also show how our method applies to the previously self-assembled honeycomb lattice [4]. It is significantly simpler, but physically interesting especially in its graphene incarnation [20]. Since the power spectrum has fewer features [ $\hat{\rho}(\mathbf{G}_i) \in \{2, 1\}$ ], even a relatively simple potential can exhibit the crucial properties for self-assembly of the honeycomb lattice; see Fig. 3(b).

All examples above utilize key features of the lattices in reciprocal space, namely, that the power spectra reaches its theoretical maximal limit at the reciprocal lattice sites where  $|\mathbf{G}_i| = k$  for some k. If this criterion is not fulfilled, as is the case for several Archimedean tilings like the equilateral snub square, one cannot reduce the set of possible basis configurations to a finite number (discrete positions) and the choice of perturbations is not as straightforward. However, allowing for competing perturbations



FIG. 3 (color online). Isotropic potentials that cause particles to self-assemble into snub square (a) and honeycomb (b) lattices.

expands the group of achievable lattices. To evaluate a basis with respect to a given set of perturbations it is sufficient to calculate the structure factors of the reciprocal points coinciding with the perturbation frequencies. This is computationally efficient and therefore an iterative method could be used to find the necessary perturbations for this type of lattice. Note that we cannot exclude that there can exist a basis where the uniqueness of the ground state cannot be enforced by any perturbation.

In general, proving that a certain configuration is the ground state of a given potential is a very hard problem. In fact, the exact nature of the ground state is not rigorously known even for simple interactions such as the Lennard-Jones potential [21]. In this Letter we have described a direct method to design potentials for targeted self-assembly of lattices, a problem usually approached using iterative methods involving repeated relaxations of the system [2,3]. From our construction follows the somewhat counterintuitive observation that it is actually simpler to find a potential with a given configuration as a ground state than to determine the ground state(s) of a given potential.

To conclude, we have demonstrated that there exist isotropic potentials that produce complex lattices such as the kagome. Previous work addressing targeted lattice formation has focused on the structure of the potential in real space, e.g., [2,3], whereas our approach focuses on the energy spectrum of the potential. We believe our method to be better suited for targeted self-assembly of lattices since their translational invariance is more naturally expressed in reciprocal space.

The ability to design interactions that self-assemble into complex lattices is interesting in itself, but this remains a theoretical observation if the potentials are too complicated to be feasible except in computer simulations. By analyzing how smoothing and screening affect the energy spectrum we can systematically simplify the potentials and thereby design interactions that should be possible to implement in experimental systems. A complementary approach to achieve realizability could be to only consider potentials that can be spanned as a linear combination, or some other parametrization, of a set of basis potentials that can be controlled in an experimental system. Our method can be applied directly in this context by analyzing the parametrized potential in reciprocal space according to the design principles we have introduced. Exploring the possibilities and limitations of combinations of standard potentials is currently work in progress.

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$$\Psi[\Delta_k]\hat{V}(k) = \int r dr V(r)\Psi(-r^2)J_0(kr)$$

with any function  $\Psi$  for which the integral converges. By choosing, e.g.,  $\Psi[X] = e^{aX}$ , one see that a Gaussian smoothing of the spectrum corresponds to a screening of the potential and vice versa.

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