Deriving Finite Sphere Packings

The Harvard community has made this article openly available. Please share how this access benefits you. Your story matters

<table>
<thead>
<tr>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Published Version</td>
<td>doi:10.1137/100784424</td>
</tr>
<tr>
<td>Citable link</td>
<td><a href="http://nrs.harvard.edu/urn-3:HUL.InstRepos:6098792">http://nrs.harvard.edu/urn-3:HUL.InstRepos:6098792</a></td>
</tr>
<tr>
<td>Terms of Use</td>
<td>This article was downloaded from Harvard University’s DASH repository, and is made available under the terms and conditions applicable to Open Access Policy Articles, as set forth at <a href="http://nrs.harvard.edu/urn-3:HUL.InstRepos:dash.current.terms-of-use#OAP">http://nrs.harvard.edu/urn-3:HUL.InstRepos:dash.current.terms-of-use#OAP</a></td>
</tr>
</tbody>
</table>
DERIVING FINITE SPHERE PACKINGS

NATALIE ARKUS†, VINOTHAN N. MANOHARAN†, AND MICHAEL P. BRENNER†

Abstract. Sphere packing problems have a rich history in both mathematics and physics; yet, relatively few analytical analyses of sphere packings exist, and answers to seemingly simple questions are unknown. Here, we present an analytical method for deriving all packings of $n$ spheres in $\mathbb{R}^3$ satisfying minimal rigidity constraints ($\geq 3$ contacts per sphere and $\geq 3n - 6$ total contacts). We derive such packings for $n \leq 10$ and provide a preliminary set of maximum contact packings for $10 < n \leq 20$. The resultant set of packings has some striking features; among them are the following: (i) all minimally rigid packings for $n \leq 9$ have exactly $3n - 6$ contacts; (ii) nonrigid packings satisfying minimal rigidity constraints arise for $n \geq 9$; (iii) the number of ground states (i.e., packings with the maximum number of contacts) oscillates with respect to $n$; (iv) for $10 \leq n \leq 20$ there are only a small number of packings with the maximum number of contacts, and for $10 \leq n < 13$ these are all commensurate with the hexagonal close-packed lattice. The general method presented here may have applications to other related problems in mathematics, such as the Erdös repeated distance problem and Euclidean distance matrix completion problems.

Key words. packing, sphere, colloid, distance equations

AMS subject classifications. 14, 82

DOI. 10.1137/100784424

1. Introduction. We consider all configurations of $n$ identical impenetrable spheres in $\mathbb{R}^3$ that maximize the number of contacts between the spheres.

Our interest in this problem arose through the following question: can one direct the self-assembly of colloidal particles into any desired structure simply by imposing a spherically symmetric binding specificity on those colloidal particles? Colloidal particles are small (nanometer to micron sized) spherical particles in aqueous solution. In recent years myriad methods have been developed to control the binding of particles to each other, thus causing them to self-assemble into clusters [50, 41, 49, 10, 15, 24]. The interactions between the particles typically have a range much smaller than the particle size; the potential energy of such a cluster is then simply proportional to the number of contacts. Structures with maximum numbers of contacts are thus the minima of the potential energy and correspond to what will form in the thermodynamic equilibrium.

Controlling which structures will form thus becomes a problem of controlling which structures correspond to energetic minima, or equivalently to contact maxima. Although minimal-energy clusters have been catalogued for many different potentials, e.g., the Lennard-Jones potential [21], they had not previously been calculated for hard (impenetrable) spheres. Thus there was no detailed understanding of what structures could self-assemble in this colloidal system.

Received by the editors January 29, 2010; accepted for publication (in revised form) July 27, 2011; published electronically December 20, 2011. This work was supported by the MRSEC program of the National Science Foundation under award DMR-0820484, the NSF Division of Mathematical Sciences under grant DMS-0907985, and DARPA under contract BAA 07-21.


†School of Engineering and Applied Sciences, Harvard University, Cambridge, MA 02138 (narkus@post.harvard.edu, vnm@seas.harvard.edu, brenner@seas.harvard.edu).

1By “spherically symmetric binding specificity” we are referring to a situation in which colloidal particles bind isotropically, but not all colloidal particles can bind to one another. For example, imagine assigning labels to the colloidal particles $A$, $B$, $C$, and so on. Particles sharing common labels will be able to bind to one another, but colloidal particles that do not share a common label cannot bind.
The motivating question of how to control the self-assembly of a colloidal system could thus be broken up into two parts: (i) determine what structures can self-assemble and (ii) determine how to direct the self-assembly of the system such that only the desired structure(s) form. In this paper, we address (i). In [5, 6], we address (ii).

It should be noted that, in addition to self-assembly, the structures of small clusters of atoms or particles bear directly on problems central to materials science and condensed matter physics, including nucleation, glass formation, and the statistical mechanics of clusters [18, 47, 44]. For instance, the first step towards understanding the thermodynamics of a particular cluster system is to calculate the ground states as a function of particle number \( n \) [17].

The structures that can self-assemble in this colloidal system can be determined by solving the mathematical problem of which structures globally or locally maximize the number of contacts between \( n \) spheres in \( \mathbb{R}^3 \), i.e., all structures in which either (i) no additional contacts between spheres can exist, or (ii) a contact must first be broken in order to form an additional contact. We will refer to such structures as packings. We formulate our problem by focusing on enumerating only minimally rigid sphere packings, which we define as packings with at least 3 contacts per particle and at least \( 3n - 6 \) total contacts. Minimal rigidity is necessary, but not sufficient, for a structure to be rigid. We previously detailed the ground states of these packings as well as some of their interesting features in [7]. Here, we present the results and method more completely. The method we introduce combines graph theory and geometry to analytically derive all minimally rigid packings of \( n \) spheres. We perform this enumeration for \( n \leq 10 \) spheres. Due to the large number of packings that must be evaluated, this analytical method is implemented computationally, and near \( n = 10 \) we reach the method’s computational limitations. Finding scalable methods for enumerating packings at higher \( n \) is a significant challenge for the future.

Already by \( n = 10 \), a number of interesting features set in. For \( n \leq 9 \), all minimally rigid packings have exactly \( 3n - 6 \) contacts. The first instance of a nonrigid sphere packing that satisfies minimal rigidity constraints occurs at \( n = 9 \), and more such nonrigid packings arise at \( n = 10 \). The first instance of packings with greater than \( 3n - 6 \) contacts occurs at \( n = 10 \). We discuss the geometrical manner in which these maximum contact packings arise and conjecture that maximum contact packings for all \( n \) in this system must contain octahedra. We provide preliminary evidence for this maximum contact conjecture for \( n \leq 20 \), and we show that the putative maximum contact packings of \( 10 \leq n \leq 13 \) are commensurate with the hexagonal close-packed (HCP) lattice, but that maximum contact packings of \( 14 \leq n \leq 20 \) are not. Furthermore, we show that the number of packings containing the maximum number of contacts is oscillatory with \( n \), and we discuss the origins of these oscillations.

The set of packings we enumerate includes, as a subset, structures previously observed and described in the literature: for example, it includes all minimal-second-moment clusters reported by [48], packings observed experimentally through capillary driven assembly of colloidal spheres [34, 41], as well as the Janus particle structures observed by Hong et al. [29].

This problem is closely related to several unsolved problems in mathematics, such as the Erdős unit distance problem (a.k.a. the Erdős repeated distance problem), Euclidean distance matrix and positive semidefinite matrix completion problems, and 3-dimensional graph rigidity. Thus the method and results presented here may have a direct bearing on these problems.
The organization of this paper is as follows: In the next section we outline our mathematical formulation of the problem and describe our methodology for finding all minimally rigid packings of a fixed $n$. We combine graph theoretic enumeration of adjacency matrices with solving for their corresponding distance matrices. The elements of these distance matrices correspond to the relative distances between spheres in 3-dimensional Euclidean space and thus yield the packings that correspond to those adjacency matrices. Analytical methods for solving such adjacency matrices that scale efficiently with $n$ do not exist. Sections 3 and 4 thus derive a method with improved scaling: Section 3 derives geometrical rules that map patterns in adjacency matrices either (i) to a configuration of spheres, in which case the adjacency matrix is solved for its corresponding distance matrix or matrices; or (ii) to an unrealizable configuration, in which case no real-valued embedding in 3-dimensional Euclidean space of that adjacency matrix exists in which spheres do not overlap. We show how these geometrical rules, combined with adjacency matrix enumeration, can lead to a complete set of minimally rigid packings. Each time a new pattern is encountered, a new geometrical rule must be derived; thus this part of the method requires new derivations at each $n$ and does not scale efficiently.

In section 4, we derive a single geometrical rule (the triangular bipyramid rule) that can solve all iterative adjacency matrices. An iterative adjacency matrix is an $n \times n$ matrix in which all minimally rigid $m < n$ subgraphs also correspond to packings. This part of the method applies to any $n$ and thus scales efficiently for all $n$. Most adjacency matrices at small $n$ are iterative; therefore, this greatly reduces the number of geometrical rules necessary to derive a complete set of packings. Section 5 describes the set of sphere packings we find from our study. We provide analytical formulas for packings up to $n = 7$, and the set of packings for $n = 8, 9, 10$ is included in the supplementary information [8]. Section 6 summarizes notable properties of the packings, including how the number of contacts changes with $n$, the emergence of minimally rigid structures that are not rigid, and the emergence of maximum contact packings that are commensurate with lattice packings. Section 7 summarizes the main roadblocks towards obtaining results at higher $n$ and contains several ideas and conjectures therein, and also discusses extensions to dimensions other than 3, as well as relevance to other problems of mathematical interest. Section 8 provides some concluding remarks.

2. Mathematical formulation. We begin by presenting a two-step method for enumerating a complete set of sphere packings that satisfies minimal rigidity constraints. The set of all packings of $n$ spheres is a subset of all possible configurations of $n$ spheres. Thus, to enumerate a complete set of sphere packings we (i) use graph theory to enumerate all $n$ sphere configurations, and (ii) determine which of those configurations correspond to sphere packings. The sphere packings we consider here correspond to maxima (local or global) in the number of contacts. Provided step (ii) is exact, this method will produce a complete set of packings. However, current analytical methods do not scale efficiently with $n$ and are therefore ill-suited for step (ii). Thus, in sections 3 and 4, we use basic geometry to derive an analytical method that can exactly determine which configurations correspond to sphere packings. Because the number of possible configurations grows exponentially with $n$, this analytical pro-

---

2This method allowed us to enumerate packings up to $n = 8$, while standard methods in algebraic geometry (using the package SINGULAR) allowed us to enumerate up to $n = 7$.

3In other words, either the configuration of spheres cannot form an extra contact (global maxima), or if an extra contact can be formed, it first requires the breaking of an existing contact (local maxima).
cess must be implemented computationally. We focus our search only on those sphere packings satisfying minimal rigidity constraints (≥ 3 contacts per sphere, ≥ 3n − 6 total contacts), because doing this guarantees that if a graph has an embedding in 3-dimensional Euclidean space, it corresponds to a sphere packing, whereas graphs that are not minimally rigid can have 3-dimensional embeddings without corresponding to packings due to a continuous degree of freedom that allows for the formation of another contact.

We also note that spheres can be thought of as points (Figure 1), where points correspond to the centers of spheres, and we measure the distance between spheres as the distance between their centers. Throughout this paper we will use the words sphere, point, and particle interchangeably.

2.1. Graph theory produces the set of possible packings. A configuration of n spheres can be described by an n × n adjacency matrix, A, detailing which spheres are in contact: A_{ij} = 1 if the i-th and j-th particles touch, and A_{ij} = 0 if they do not. A system of n spheres has n(n−1)/2 interparticle distances; the 2 possibilities (touching or not touching) per distance thus leave 2^{n(n−1)/2} different ways of arranging contacts amongst the distances. There are thus 2^{n(n−1)/2} possible adjacency matrices, each of which potentially corresponds to a packing.

Figure 1 shows a packing of 6 particles, both as a sphere packing (Figure 1(a)) and as points connected by line segments (Figure 1(b)). The adjacency matrix corresponding to this packing is shown in Figure 1(c). The set of possible packings can be enumerated by considering all adjacency matrices. For n = 6, there are 2^{15} = 32,768 different adjacency matrices. Table 1 shows that the number of adjacency matrices grows rapidly with n, reaching 3.5184 × 10^{13} by n = 10; however, many of these correspond to the same structure due to particle labeling degeneracy. For example, switching labels 5 and 2 of Figure 1 yields another adjacency matrix corresponding to the same structure.\footnote{Note that this is purely an exercise in switching particle labels; it is not a statement about the symmetry of the structure—we are not saying that particles 2 and 5 have the same contact distribution. The point is that many matrices can lead to the same structure, because how one labels the particles is an arbitrary factor.}

Adjacency matrices corresponding to the same structure are isomorphic to one another—meaning there will exist a permutation of rows and columns that can translate one matrix into the other.\footnote{See [27] for a nice example of such a permutation.}

To generate the complete set of possible packings, we need only enumerate non-isomorphic A’s. Such algorithms exist; examples include nauty and the SAGE pack-
The Growth of Adjacency Matrices with $n$. The number of adjacency matrices (constructed by [42]) decreases rapidly as isomorphism and rigidity constraints are imposed. Iterative and noniterative are defined in the text. The classification of whether an $A$ is iterative or not is here shown after all rules for $n-1$ particles are applied; thus the noniterative column shows $n$-particle noniterative structures only and does not include noniterative structures of less than $n$ particles. Also note that the classification of whether an $A$ is iterative or not is thus sensitive to which geometrical rules are used. The number of iterative and noniterative $A$’s at $n=10$ is different from [7] because some modifications were made to the code since the publication of that paper. Please see the supplemental information [8] for exactly which modifications have been made.

<table>
<thead>
<tr>
<th>$n$</th>
<th>$A$’s</th>
<th>Nonisomorphic $A$’s</th>
<th>Minimally rigid $A$’s</th>
<th>Iterative $A$’s</th>
<th>Noniterative $A$’s</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>2</td>
<td>2</td>
<td>2</td>
<td>1</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>3</td>
<td>8</td>
<td>4</td>
<td>1</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>4</td>
<td>64</td>
<td>11</td>
<td>1</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>5</td>
<td>1,024</td>
<td>34</td>
<td>1</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>6</td>
<td>32,768</td>
<td>156</td>
<td>4</td>
<td>3</td>
<td>1</td>
</tr>
<tr>
<td>7</td>
<td>2,097,152</td>
<td>1,044</td>
<td>29</td>
<td>26</td>
<td>3</td>
</tr>
<tr>
<td>8</td>
<td>268,435,456</td>
<td>12,336</td>
<td>438</td>
<td>437</td>
<td>1</td>
</tr>
<tr>
<td>9</td>
<td>6.8719 · 10$^{10}$</td>
<td>274,668</td>
<td>13,828</td>
<td>13,823</td>
<td>5</td>
</tr>
<tr>
<td>10</td>
<td>3.5184 · 10$^{13}$</td>
<td>12,005,168</td>
<td>750,352</td>
<td>750,258</td>
<td>94</td>
</tr>
</tbody>
</table>

age called nice [42, 3]. The number of nonisomorphic matrices is much smaller but still grows exponentially with $n$. Table 1 shows this growth also; for example at $n=6$ the number of potential structures is 156, and at $n=10$ it is 12,005,168.

The set of $A$’s (potential packings) can be further reduced by imposing rigidity constraints. Most structures with fewer than $3n-6$ total contacts or fewer than 3 contacts per particle will not correspond to a packing because there will exist a continuous degree of freedom through which the structure can form one or more bonds. Rigidity requires (i) there be at least 3 contacts per particle, and (ii) there be at least as many contacts as internal degrees of freedom—thus there must be at least $3n-6$ contacts.\(^6\)

Table 1 shows how imposing minimal rigidity constraints restricts the number of adjacency matrices. For $n \leq 5$ particles, this eliminates all but 1 adjacency matrix, thus identifying a unique packing for each of these $n$: the doublet, triangle, tetrahedron, and triangular bipyramid, respectively (section 5). For $n \leq 4$, all relative distances within these packings are touching and are thus known. However, for $n=5$, the packing contains one unknown relative distance, which must be determined. For $n \geq 6$, more than one minimally rigid $A$ exists, and thus rigidity constraints alone are insufficient to identify the number of sphere packings (or to solve for them, as more than 1 distance in each minimally rigid $A$ is unknown).

2.2. Solving potential packings: Algebraic formulation. To make further progress, we reformulate our problem algebraically. Each adjacency matrix element, $A_{ij}$, is associated with an interparticle distance,

$$r_{ij} = \sqrt{(x_i - x_j)^2 + (y_i - y_j)^2 + (z_i - z_j)^2};$$

\(^6\)Note that in restricting to structures with exactly $3n-6$ contacts, we will also find structures with more than $3n-6$ contacts. This is because when we solve for the packings as outlined below, the solutions can end up having more contacts than are assumed in the algebraic formulation.
which is the distance between particles \( i \) and \( j \) whose centers are located at \((x_i, y_i, z_i)\), \((x_j, y_j, z_j)\), respectively. The distances are constrained by the adjacency matrix as follows:

\[
A_{ij} = 1 \implies r_{ij} = 2r, \tag{1}
\]
\[
A_{ij} = 0 \implies r_{ij} \geq 2r, \tag{2}
\]

where \( r \) is the sphere radius.\(^7\)

For adjacency matrices with \( 3n - 6 \) contacts, this leads to precisely as many equations as unknowns.\(^8\) The task is thus to solve for the \( r_{ij} \geq 2r \) given a particular set of \( r_{ij} = 2r \). The particle configuration encoded by each \( A \) is thus specified by a distance matrix, \( D \), whose elements \( D_{ij} = r_{ij} \).

The fundamental question is to find an efficient, exact method for mapping \( A \to D \). If any \( D_{ij} < 2r \), this implies that particles \( i, j \) overlap; thus any \( D \) with \( D_{ij} < 2r \) is unphysical. Figure 1(d) shows the distance matrix corresponding to an \( n = 6 \) packing. The interparticle distance between each of the particles that are touching is normalized to 1; for this packing, this leaves three distances that need to be determined; \( r_{12}, r_{23}, \) and \( r_{34} \).

In solving \( A \) for \( D \), the following scenarios are possible:

1. Continuous set(s) of real-valued \( D \) correspond to a given \( A \), in which case the structure(s) are not rigid.
2. No real-valued \( D \) exists that solves \( A \), in which case the structure is unphysical.
3. Finite, real-valued \( D \) exist that solve \( A \). In this case, the structure(s) correspond to rigid sphere packing(s) provided that all \( D_{ij} \geq 2r \).

For every nonisomorphic adjacency matrix, whether there exists a corresponding packing requires solving for \( D \) and asking whether the resulting \( r_{ij} \)'s satisfy these constraints.

### 2.3. Limitations of existing solution methods.

The issue now becomes one of solving a system of \( n(n-1)/2 \) constraints (\( 3n - 6 \) equations and \( n(n-1)/2 - (3n - 6) \) inequality constraints). Numerical approaches for solving such a system cannot be guaranteed to converge; for example, Newton’s method requires an accurate initial guess for guaranteed convergence. When a solution does not converge, we do not know whether it is because a solution does not exist or because the initial guess is not sufficiently accurate.\(^9\) Algebraic geometric methods (e.g., Gröbner bases) \(^10\) are effective, but these algorithms do not scale efficiently with \( n \). Our own implementation\(^10\) of this method using the package \textsc{SINGULAR} \(^23\) was only able to solve for structures up to \( n = 7 \).

In the following section, we use another approach and derive a different geometrical method to efficiently solve for all sphere packings given a set of nonisomorphic packings.
minimally rigid $\mathcal{A}$’s. We implement this method up to $n = 10$, at which point we begin to hit some practical roadblocks; these are discussed at the end of the paper, where we outline potential ways to overcome them.

2.3.1. Chiral structures. Before proceeding further, it is worth remarking that structures with different handedness will correspond to the same distance matrix. We can analyze each $\mathcal{D}$ and determine whether it corresponds to a structure that has a chiral counterpart (see section 3.5). We refer to chiral structures as the same packing but different states—thus, a distance matrix having a left- and right-handed counterpart corresponds to 1 packing with 2 different states. Note that according to our definition, a different packing necessarily corresponds to a different state. Thus, the total number of states is equal to the total number of packings plus the total number of chiral counterparts.

3. Geometrical rules solve for sphere packings. We now show how geometrical rules can be used to effectively and analytically solve the class of polynomial equations that are generated by adjacency matrices. We use basic geometry to construct rules associating patterns of 1’s and 0’s in $\mathcal{A}$’s with either a given relative distance, $D_{ij}$, or an unphysical conformation (in which case no $D \geq 2r$ exists). There thus exist two types of rules: 

Elimination rules eliminate an $\mathcal{A}$ as unphysical, and 

distance rules solve an $\mathcal{A}_{ij}$ for its corresponding $D_{ij}$.

3.1. Neighbor spheres and intersection circles. With each sphere of radius $r$, we can associate a neighbor sphere of radius $R = 2r$, whose surface defines where another sphere must lie if it is to touch the original sphere in question (Figure 2(a)). When 2 spheres touch, their neighbor spheres intersect in an intersection circle (Figure 2(b)). The radius of the intersection circle follows from straightforward geometry and is $\frac{\sqrt{3}}{2}R$ (see supplemental information for the derivation [8]).

![Fig. 2. The Neighbor Sphere. (a) A particle and its neighbor sphere. 4 particles are shown touching the center particle, and it is seen that their centers lie on the surface of the particle’s neighbor sphere. (b) 2 touching particles. The associated neighbor spheres of the particles intersect in an intersection circle of radius ($\sqrt{3}/2$)R.](image-url)
Fig. 3. No More than 2 Particles Can Touch 3 Connected Particles. Schematic of 3 linearly connected particles and their associated intersection circles. The center of each intersection circle lies at the midpoint of the line segment connecting the associated points. There can never be more than 2 intersection points of these intersection circles, indicating that no more than 2 particles can touch the same 3 linearly connected particles.

Now we can interpret each $A_{ij} = 1$, in geometrical terms, as an intersection circle between spheres $i$ and $j$. Minimal rigidity constraints imply that each particle is associated with at least 3 intersection circles. Intersection circles can be used to derive geometrical rules because, in general, a packing of $n$ particles involves intersections of intersection circles, and intersections of intersection circles define points in space. A particle touching $m$ other particles will lie at the intersection of those $m$ neighbor spheres. For example, a particle touching the dimer depicted in Figure 2(b) will lie on the circumference of the associated intersection circle. The intersection of $m \geq 3$ neighbor spheres are points—and by defining points in space, basic trigonometry can be used to calculate the distances between those points, thus solving $A$’s for $D$’s.

3.2. Individual geometrical rules. Using intersection circles, we now derive several representative geometrical rules for eliminating and solving adjacency matrices. The supplementary information [8] contains the complete set of rules used to derive the results of this paper.

3.2.1. Rule 1. The simplest rule arises from the fact that intersection circles can intersect in 2, 1, or 0 points, but never in more than 2 points. This geometrical fact implies that any $A$ with the following property is unphysical: any 2 of the set $\{A_{jk}, A_{jp}, A_{kp}\}$ equal 1, and there exist more than 2 $i$’s for which $A_{ij} = A_{ik} = A_{ip} = 1$.

Physically, this implies that no more than 2 spheres can simultaneously touch 3 connected spheres. Three spheres are connected if at least 2 contacts exist between them (Figure 3). This in turn tells us how many identical spheres can mutually touch a trimer: 2. Figure 4 shows an example of an adjacency matrix that is unphysical for this reason: the blue highlighted section shows that spheres 4, 5, 6 make up a trimer; but the purple highlighted section shows that spheres 1, 2, 3 all touch the same trimer. This is impossible given the argument outlined above, and hence this adjacency matrix does not correspond to a packing.

3.2.2. Rule 2. A trimer, a configuration of 3 spheres forming an equilateral triangle, is associated with 3 mutually intersecting intersection circles (Figure 5(a)). These 3 intersection circles intersect at 2 points (shown in red). Here we calculate the distance between these 2 intersection points.
Fig. 4. Example of an Unphysical Adjacency Matrix. (a) An adjacency matrix that is unphysical because it implies more than 2 intersections of intersection circles. The blue (darker) highlights show that particles 4, 5, 6 make up a trimer. The purple (lighter) highlighted part shows that particles 1, 2, and 3 all touch the same trimer, 4, 5, 6. (b) A sphere packing corresponding to this unphysical adjacency matrix (shown in both sphere and point/line representations). For it to be realized, 2 particles must occupy the same point in space.

Fig. 5. The Intersections of 3 Intersection Circles. (a) A trimer and its corresponding intersection circles. The 3 intersection circles mutually intersect at 2 points, shown in red (the two smaller dots within the triangle). (b) The triangles that relate the trimer to one of the points of intersection (red dot at the intersection of the dashed lines). The distance between this point and the center of the triangle is equivalent to half the distance between the 2 points of intersection (here denoted as $h$).

Note that a particle lying at one of the intersection points forms the 4-particle packing (the tetrahedron). Two particles, lying at each intersection point, form the 5-particle packing (the 5-point polytetrahedron). The distance between these 2 intersection points, $h$, is the only distance that is greater than $R$ in the 5-particle packing (Figure 6).

To calculate this distance, we note that the trimer and its associated intersection circles form the set of triangles shown in Figure 5(b) (where the dashed line indicates an out-of-plane triangle). We calculate $a$ by considering the right triangle with sides $\sqrt{3}/2R - a, a, 1/2R$. Trigonometry then implies that $a = R/(2\sqrt{3})$, and $h = 2\sqrt{2/3}R$. 
Fig. 6. The 2 Intersection Points of a Trimer Correspond to the 5-Particle Packing. A 5-particle packing is shown with its point representation overlain. The center triangle of the point representation corresponds to a trimer, and the 2 points that contact the trimer correspond to the 2 intersection points of the trimer’s 3 intersection circles. The 2 intersection points are shown in red (dots touching the ends of the dashed line), and $h$ corresponds to the distance between them.

This implies that the solution to an adjacency matrix corresponding to the 5-particle packing is

$$
\begin{pmatrix}
0 & 1 & 1 & 1 & 1 \\
1 & 0 & 1 & 1 & 1 \\
1 & 1 & 0 & 1 & 1 \\
1 & 1 & 1 & 0 & 0 \\
1 & 1 & 1 & 0 & 0 \\
\end{pmatrix}
\rightarrow
\begin{pmatrix}
0 & 1 & 1 & 1 & 1 \\
1 & 0 & 1 & 1 & 1 \\
1 & 1 & 0 & 1 & 1 \\
1 & 1 & 1 & 0 & 2\sqrt{\frac{2}{3}} \\
1 & 1 & 1 & 2\sqrt{\frac{2}{3}} & 0 \\
\end{pmatrix},
$$

where the right matrix is the corresponding distance matrix, $D$, and without loss of generality we have let $R = 1$. For $n = 5$, there is only 1 nonisomorphic minimally rigid $A$.

We can formalize this construction as a distance rule, which can be used whenever a submatrix of some $A$ has the same structure as the 5-particle packing. Such submatrices can be identified with the following pattern: $A_{ij} = A_{ik} = A_{kj} = 1$, and there exist 2 points $p$ for which $A_{pi} = A_{pj} = A_{pk} = 1$. Whenever this pattern exists, the distance submatrix between the associated points corresponds to $D$ for the 5-particle packing. In particular, this rule solves for the distance between the 2 points $p$; for example, if $p = l, m$, then $D_{lm} = 2\sqrt{\frac{2}{3}} R$.

3.2.3. Rule 3. Another elimination rule follows directly from any distance rule, including Rule 2 derived above. Suppose we determine that for a given pattern of $A_{ij}$, the contact distribution implies that $D_{kp} > R$. If it then happens that $A_{kp} = 1$, then this implies that all of the geometrical constraints cannot be satisfied simultaneously, so that $A$ is unphysical.
Fig. 7. 5 Points on an Intersection Circle. The intersection circle shown in black corresponds to the dimer in the center. 5 points are shown lying on the intersection circle; this corresponds to 5 particles touching the center dimer. The radius of the intersection circle is \((\sqrt{3}/2)R\) (shown as dashed black lines) and connects the center of the dimer (which is the origin of the intersection circle) to points 1–5 on the intersection circle. The arc length swept out by one pair of particles on the intersection circle is \(S\). It can be seen that the 1st and 5th particles nearly touch. The space between them is not big enough to fit another particle, and thus it can be seen that no more than 5 particles can touch a dimer.

For example, if an \(A\) contained the intersection circle construction discussed in the previous section, that would imply that \(D_{lm} = 2\sqrt{2/3}R\), but if the adjacency matrix also stated that \(A_{lm} = 1\), then that \(A\) would be unphysical.

3.2.4. Rule 4. We can derive another set of geometrical rules by finding the maximum number of points that can lie on an intersection circle—this corresponds to the maximum number of spheres that can touch a dimer. Figure 7 shows the dimer (top and bottom spheres), as well as points lying on their intersection circle.

The maximum number of spheres that can lie on an intersection circle is 5, and this can be calculated as follows: we divide the circumference of the entire intersection circle by the arc length swept out by 2 spheres lying a unit distance apart\(^{11}\) (see Figure 7). This arc length is given by \(S = r\theta\), where \(r\) is the radius of the intersection circle, and \(\theta\) is the angle between 2 radial line segments. The law of cosines then implies that

\[
\theta = \cos^{-1}\left(\frac{1}{3}\right),
\]

so that the number of points a distance \(R\) apart that can fit on an intersection circle

\(^{11}\)Without loss of generality, we refer to the distance between two touching spheres, \(R\), as the unit distance.
is given by

$$\frac{2\pi \sqrt{2} R}{\sqrt{3} R \cos^{-1} \left( \frac{1}{3} \right)} \approx 5.1043.$$  

(4)

This indicates that (i) any $A$ implying that more than 5 points lie on an intersection circle, or (ii) any $A$ implying a unit distance between all $m \leq 5$ points lying on an intersection circle, is unphysical. We can identify 5 points lying on an intersection circle by the following adjacency matrix pattern: $A_{ij} = 1$, such that there are 5 points $k$ for which $A_{ik} = A_{jk} = 1$.

To solve for the structure of $m \leq 5$ points lying on an intersection circle, we must compute the distances between the nontouching particles on the intersection circle (Figure 7). Of these distances, we have already calculated that between points 1 and 3 and shown that it is $2\sqrt{2}/3R$ (section 3.2.2). All of these distances can be obtained by the isosceles triangle with equivalent lengths $\sqrt{3}/2R$ (corresponding to the dashed black lines in Figure 7—note that only 2 such lines are shown, but that they exist between the midpoint of the dimer and every point along the intersection circle). The unique length of the isosceles triangle will be the unknown distance, $r_{ij}$, and the angle between the two $\sqrt{3}/2R$ sides connecting particles $i$ and $j$ will be called $\phi_{ij}$. Thus, the unknown distances will all be given by

$$\sin \left( \frac{1}{2} \phi_{ij} \right) = \frac{1}{2} r_{ij} \frac{\sqrt{3}}{2} R,$$

(5)

where

$$\phi_{13} = 2\theta,$$

$$\phi_{14} = 2\pi - 3\theta,$$

$$\phi_{15} = 2\pi - 4\theta,$$

where $\theta$ is given by (3), thus yielding

$$r_{13} = 2\sqrt{\frac{2}{3}} R,$$

(6)

$$r_{14} = \frac{5}{3} R,$$

(7)

$$r_{15} = \frac{4\sqrt{6}}{9} R.$$  

(8)

These calculations apply to any adjacency matrix in which $A_{ij} = 1$, there exist $n$ points, $k$, for which $A_{ik} = A_{jk} = 1$, and there also exist $n-1$ instances where $A_{pq} = 1$ amongst the $n$ points, $k$, where $n = 3, 4, 5$ for $r_{13}, r_{14}, r_{15}$, respectively. Then the distance between the two endpoints of the $n$ particles is given by $D_{pk} = r_{13}, r_{14}, r_{15}$, respectively.

That is for

$n = 2$: if $k = p, q, l$, then $A_{pq} = A_{ql} = 1$, and the distance $D_{pl} = 2\sqrt{2}/3 R$;

$n = 3$: if $k = p, q, l, m$, then $A_{pq} = A_{ql} = A_{lm} = 1$, and the distance $D_{pm} = \frac{5}{3} R$;

$n = 4$: if $k = p, q, l, m, z$, then $A_{pq} = A_{ql} = A_{lm} = A_{mz} = 1$, and $D_{pz} = \frac{4\sqrt{6}}{9} R$.  

Copyright © by SIAM. Unauthorized reproduction of this article is prohibited.
3.3. Using geometrical rules to derive a complete set of sphere packings. The aforementioned geometrical rules can be used, along with the set of nonisomorphic adjacency matrices, to derive a complete set of packings for a given \( n \). To explain more clearly how this is done, we show as an example the derivation for \( n = 7 \) particle packings. For this \( n \), there exist 29 minimally rigid \( \mathcal{A} \)'s, all of which potentially correspond to packings (Table 1).

To these matrices we apply the elimination rules just outlined, as well as those that appear in the supplementary information [8]. This immediately eliminates 24/29 \( \mathcal{A} \)'s as unphysical. Figure 8 shows which of the matrices are eliminated. Table 2 shows which rules are used to eliminate the \( \mathcal{A} \)'s. 17/24 of the matrices are eliminated because they imply more than 2 intersections of intersection circles. Three of the matrices are eliminated because of the relative distance rule for three points on an intersection circle. Two matrices each are eliminated by rules acting on 5 rings and 4 rings, respectively. For the remaining five adjacency matrices, we apply distance rules to the \( \mathcal{A} \)'s to solve for the corresponding \( \mathcal{D} \)'s. Table 3 details which distance rules are

Note that for \( n = 3 \), we have already identified this \( \mathcal{A} \) pattern in section 3.2.2, whereas the \( n = 2, 4 \) structures are new. Also note that, by symmetry, \( r_{13} = r_{24} = r_{35} \), and \( r_{14} = r_{25} \), and that these equivalences are identified by the above patterns in \( \mathcal{A} \).

Fig. 8. Eliminating Adjacency Matrices. There are 29 nonisomorphic adjacency matrices satisfying minimal rigidity constraints for 7 particles. 24 out of the 29 \( \mathcal{A} \)'s are eliminated by geometrical rules, which are shown here as color-coded \( X \)'s—see Table 2 and Appendix A in [8] for the corresponding rules. See Figure 9 for solutions to the 5 physically realizable \( \mathcal{A} \)'s (which correspond to the matrices that appear here without an \( X \)).
Table 2

Elimination Rules Used for 7-Particle Packings. Each rule appears in its own section either in the text or in Appendix A in [8], where the complete set of rules is included. The rule column lists in what section(s) the relevant rules can be found. The X’s in the color column are from top to bottom: red, blue, green, and orange.

<table>
<thead>
<tr>
<th>Color</th>
<th>Unphysical because</th>
<th>Rule:</th>
</tr>
</thead>
<tbody>
<tr>
<td>X</td>
<td>2 or more intersection circles intersect at more than 2 points</td>
<td>A.1 section 3.2.1</td>
</tr>
<tr>
<td>X</td>
<td>All relative distances between 3 points that lie on an intersection circle = R</td>
<td>A.3 sections 3.2.2, 3.2.3, and 3.2.4</td>
</tr>
<tr>
<td>X</td>
<td>A closed 5 ring surrounds a circle of intersection</td>
<td>A.14</td>
</tr>
<tr>
<td>X</td>
<td>2 points on opposite sides of a closed 4 ring touch</td>
<td>A.15</td>
</tr>
</tbody>
</table>

Table 3

Rules Needed to Solve 7-Particle Packings. The rules listed here correspond to distance rules. Rule “A. #” corresponds to rule # in Appendix A in [8]; otherwise the relevant equation and section numbers are listed for rules found within the paper. (Note that rule 4, found in section 3.2.4 (equation (6)), is the same as rules 2 (section 3.2.2) and A.1, rule 4 (equation (7)) is the same as rule A.2, and rule 4 (equation (8)) is the same as rule A.4.) Graphs are numbered in ascending order from left to right and top to bottom as they appear in Figure 8. These graphs correspond to the ones without X’s.

<table>
<thead>
<tr>
<th>Graph number</th>
<th>Rules used:</th>
</tr>
</thead>
<tbody>
<tr>
<td>4</td>
<td>section 3.2.4 eqns (6), (7), and (8)</td>
</tr>
<tr>
<td>8</td>
<td>section 3.2.4 eqns (6) and (7)</td>
</tr>
<tr>
<td>17</td>
<td>section 3.2.4 eqns (6), (7); and A.11</td>
</tr>
<tr>
<td>22</td>
<td>A.7 and A.6</td>
</tr>
<tr>
<td>26</td>
<td>A.3 and A.9</td>
</tr>
</tbody>
</table>

used to solve for the packings corresponding to each A. The analytical solutions for the distance matrices as well as the associated packings are shown in Figure 9; to each A (numbered by the order in which it appears in Figure 8, in ascending order from left to right, and top to bottom, respectively) we apply the rules outlined in Table 3 to analytically solve for the packing.

This is the set of 7-particle sphere packings. Note that the packing corresponding to graph 17 (row 4 from the top) is the only one where distinct left- and right-handed structures are possible; thus it corresponds to 2 distinct states.

3.4. Nonuniqueness of geometrical rules. Note that the geometrical rules described here are not unique in that (i) the rules themselves can be derived in different ways, and (ii) a different set of rules altogether could be derived/applied to solve for the same packings. This is simply one set of rules that works. One example of this is that either Rule 2 or Rule 4 (equation (6)) can be used to determine the unknown distance in a 5-particle packing. Another example of this is that all iterative packings can be solved using the triangular bipyramid rule that will be introduced in section 4 instead of using the aforementioned rules. There are undoubtedly many such examples, and the list of rules just presented were not derived with the goal of conciseness. They are complete in the sense that they allow one to solve all adjacency matrices for the n presented here. Beyond this, however, they can only solve adjacency matrices containing the structure the rule identifies. If an A contains an identifiable structure as well as a nonidentifiable (not previously encountered) structure, then the rules will solve the identifiable part and only partially solve that A for its corresponding D. If an A contains no identifiable structure, then new rules must be derived to solve any part of it. It is because of these latter two cases that we continued to derive new geo-

Copyright © by SIAM. Unauthorized reproduction of this article is prohibited.
metrical rules as we increased in \( n \). The triangular bipyramid rule in section 4 is a rule that we derived in order to have one general rule that could recognize a certain class of adjacency matrix structures—the iterative class—for all \( n \). Because it is a general rule, applicable to all \( n \), its introduction makes the set of rules much more concise.

3.5. Chirality. Once all packings have been derived by solving all \( A \)'s for their corresponding \( D \)'s, we must determine how many states each packing has. If a pack-
ing is chiral, it will have more than one state. This will show up by a packing having a nonsuperimposable mirror image, for example, a packing having different “handedness,” such as distinct left- and right-handed structures.

One can calculate whether a packing is chiral as follows: The automorphism group of a packing, \( \{ \alpha \} \), gives the set of self-isomorphisms, i.e., all possible permutations of the structure into itself. Each element of the automorphism group will thus correspond either to a rotation or to a reflection. Rotations are transformations with determinant \( = 1 \), and reflections are transformations with determinant \( = -1 \). Thus, one can construct the set of all isomorphic graphs, \( \{ D \} \), and construct the automorphism group for any one \( D_i \) within the isomorphic set. If there exists any matrix \( D_j \in \{ D \} \) that is isomorphic to \( D_i \) only through reflections and not through rotations, i.e., if the isomorphism group of \( D_j \) to \( D_i \) corresponds to transformations that all have determinant \(-1\), then the packing \( D \) is chiral.

A property related to chirality is the symmetry number, \( \sigma \). This corresponds to the number of ways that a structure can be rotated into itself. The symmetry number is necessary for calculating the equilibrium probability distribution of packings [5, 43]. The symmetry number of a packing, \( D \), will be equal to the number of transformations within the automorphism group of \( D \) that have determinant 1. Thus, if a packing has no reflections, then the symmetry number will simply equal the size of the automorphism group. If a packing has reflections, then the symmetry number will equal the size of the automorphism group divided by 2 (dividing by 2 will remove all automorphism mappings corresponding to reflections, and not rotations).

Related to both symmetry numbers and chirality are point groups. A point group is a group of symmetry operations which all leave at least one point unmoved. Point groups have been calculated for many structures—and there exist programs that allow one to enter in a set of coordinates and retrieve the point group corresponding to those coordinates [46]. Symmetry numbers and chirality can alternatively be calculated directly from the point group of a structure. For example, compounds in the \( C_m \) point group, where \( C_m \) is the cyclic group consisting of rotations by \( 360^\circ/m \) and all integer multiples (where \( m \) is an integer), are always chiral [25].

Point groups, symmetry numbers, and chirality of packings are included in the lists of packings appearing in section 5 and in the supplementary information [8]. The growth of chiral structures with \( n \) is interesting—surprisingly, over half of all 9-particle packings are chiral—see Table 4.

4. One geometrical rule that solves for all iterative packings: The triangular bipyramid rule. In principle, these types of geometrical rules can be used to derive a complete set of sphere packings for any number of particles, \( n \). However, in practice, the number of rules used here grows too quickly with \( n \) for this to be a practical method: at \( n = 5 \) spheres, only 1 rule is required; 3 rules are needed at \( n = 6 \); 12 rules are needed at \( n = 7 \); and at \( n = 8 \), 14 rules solve \( 435/438 \) minimally rigid nonisomorphic \( \mathcal{A} \)'s. This leaves 3 unsolved \( \mathcal{A} \)'s for which more geometrical rules must be derived; looking ahead at the 13,828 and 750,352 \( \mathcal{A} \)'s that must be solved at \( n = 9, 10 \), respectively, it becomes clear that deriving a rule or set of rules that does not grow significantly with \( n \) is a necessary step. Here, we derive one geometrical rule that can solve one class of packings for any \( n \), thereby greatly reducing the number of rules needed to derive a complete set of \( n \) sphere packings. In section 7.2, we discuss how one geometrical rule can also be used to solve the other class of packings for all \( n \).

Packings can be broken up into two types or classes: iterative and noniterative, or new seeds. Iterative packings are \( n \)-particle packings that are solely combinations...
Table 4

<table>
<thead>
<tr>
<th>n</th>
<th>Packings from [27]</th>
<th>Total packings (current study)</th>
<th>New seeds</th>
<th>Nonrigid packings</th>
<th>Chiral</th>
<th>Total states</th>
</tr>
</thead>
<tbody>
<tr>
<td>2</td>
<td>1</td>
<td>1</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>3</td>
<td>1</td>
<td>1</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>4</td>
<td>1</td>
<td>1</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>5</td>
<td>1</td>
<td>1</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>6</td>
<td>2</td>
<td>2</td>
<td>1</td>
<td>0</td>
<td>0</td>
<td>2</td>
</tr>
<tr>
<td>7</td>
<td>4</td>
<td>5</td>
<td>1</td>
<td>0</td>
<td>1</td>
<td>6</td>
</tr>
<tr>
<td>8</td>
<td>10</td>
<td>13</td>
<td>1</td>
<td>0</td>
<td>3</td>
<td>16</td>
</tr>
<tr>
<td>9</td>
<td>32</td>
<td>52</td>
<td>4</td>
<td>1</td>
<td>28</td>
<td>80</td>
</tr>
<tr>
<td>10</td>
<td>113</td>
<td>262</td>
<td>8</td>
<td>4</td>
<td>201</td>
<td>463</td>
</tr>
</tbody>
</table>

Fig. 10. Iterative Packings. Two examples of iterative packings. (a) A 6-particle polytetrahedron (red) with one particle added to it (blue). This decomposes into a tetrahedron (blue) plus a 6-particle polytetrahedron (red), with a shared triangular base (purple). (b) 2 joined octahedra (one red and one blue, with a shared purple triangular base) forming a 9-particle packing. Color is available online only.

of packings of less than $n$ particles (see Figure 10). New seeds are $n$-particle packings that cannot be constructed solely out of packings of less than $n$ particles; i.e., they contain within them (in part or in whole) an inherently new structure (Figure 11).
Put another way, iterative packings correspond to $A$’s for which all minimally rigid $m \times m$ subgraphs, $m < n$, correspond to packings that have been identified at lower $n$.

4.1. Solving iterative structures. An iterative packing is a polyhedron containing solely packings of less than $n$ particles.\footnote{An iterative $A$ is an $n \times n$ graph composed solely of $m \times m$ ($m < n$) subgraphs, each of which correspond to minimally rigid $A$’s of less than $n$ spheres.} Thus any iterative packing can be decomposed into 2 joined polyhedra (see Figure 10—the red and blue packings are the joined polyhedra). The 2 polyhedra are joined via a common base of particles (shown in purple).\footnote{Given the minimal rigidity constraints we have imposed, this common base will always consist of at least 3 particles.} Because the joined polyhedra are less than $n$-particle packings, all of their intrapolyhedral distances are known from lower-order packings.

Thus, deriving one geometrical rule that can solve for all iterative packings requires solving the following geometrical problem: Given 2 joined polyhedra, where all intrapolyhedral distances are known, derive a general formula for the interpolyhedral distances. Note that the solution to this problem immediately extends to unphysical iterative structures as well, as they are composed of structures of less than $n$ particles, where either (i) one or more of the joined structures is unphysical, or (ii) the particular combination of the structures is unphysical.

The geometrical problem is solved with the following observation: An explicit analytical formula for the distance between any 2 points can be derived if those 2 points can be related to a common triangular base. Let there exist two particles $i,j$ whose interparticle distance, $r_{ij}$, is unknown. If there also exist 3 particles, $k,p,q$, with known interparticle distances ($r_{kp}, r_{kq}, r_{pq}$), and if the distances between $i,j$ and the 3-particle base ($r_{ip}, r_{ik}, r_{jq}, r_{jp}, r_{jk}, r_{jq}$) are also known, then there exists an analytical relationship for the resulting $r_{ij}$. We call this the triangular bipyramid rule because the 5 points $i,j,k,p,q$ together form a (potentially irregular) ditetrahedron or triangular bipyramid (see Figure 12). We show the rule here, while a complete derivation can be found in the supplemental information (Appendix A in [8]).
The triangular bipyramid (or ditetrahedron) constructed in the triangular bipyramid rule. The center triangle $kpq$, shown in red, corresponds to the common 3-particle base. Particles $i$ and $j$ are related to one another through the common base. The distance between $i$ and $j$, $r_{ij}$, shown as the dash-dot blue line, is unknown. $A_1$ corresponds to $\angle jpi$. $A_2$ is the dihedral angle between $\triangle ipk$ and $\triangle jpk$, and $A_3$ is the dihedral angle between $\triangle kpq$ and $\triangle jpk$. Points $i$ and $j$ can either both lie on the same side of the base $kpq$ or each lie on opposite sides of the base (indicated by the dashed lines that can either go into or come out of the plane). If $i, j$ lie on the same side, then $A_1$ is equal to the difference of $A_2$ and $A_3$, and if $i, j$ lie on opposite sides of the base, then $A_1$ is equal to the sum of $A_2$ and $A_3$.

When all distances other than $r_{ij}$ are known, then an explicit analytical formula can be derived to solve $r_{ij}$.

The dihedral angles $A_2$ and $A_3$ of Figure 12 are given by

\[
A_3 = \cos^{-1}\left(\frac{\cos a_3 - \cos b_3 \cos c_3}{\sin c_3 \sin b_3}\right),
\]
\[
A_2 = \cos^{-1}\left(\frac{\cos a_2 - \cos b_2 \cos c_2}{\sin c_2 \sin b_2}\right).
\]

These formulas are essentially obtained using the spherical rule of cosines; see Appendix A in [8] for details.

The dihedral angle $A_1$ will be either the sum or the difference of the dihedral angles $A_2$ and $A_3$ (see Figure 12), depending on whether the points $i, j$ lie on the same or on opposite sides of the base $p, k, q$. If $i, j$ lie on the same side, then $A_1 = |A_2 - A_3|$, and if $i, j$ lie on opposite sides, then $A_1 = A_2 + A_3$.

The angle $a_1$ is then given by

\[
a_1 = \cos^{-1}(\sin c_1 \sin b_1 \cos A_1 + \cos b_1 \cos c_1),
\]

and from the law of cosines, we can then calculate $r_{ij}$:

\[
r_{ij} = \sqrt{r_{ip}^2 + r_{pj}^2 - 2r_{ip}r_{pj} \cos a_1}.
\]
Associated with each $r_{ij}$ we have 2 possible $A_1$ and thus 2 possible solutions (similar, in principle, to one having 2 possible solutions to a quadratic equation).

### 4.2. Applying the triangular bipyramid rule

The triangular bipyramid rule can be used to solve all iterative packings as follows. We first search for subgraphs of $\mathcal{A}$ corresponding to lower-n seeds. The elements of $\mathcal{D}$ corresponding to these lower-order structures are known and inserted appropriately. If $\mathcal{A}$ is iterative, all minimally rigid subgraphs of $m < n$ particles (i.e., $m$ subgraphs with at least $3m - 6$ contacts and at least 3 contacts per particle) will correspond to $m$-particle packings. Once all lower-order seeds are inserted as appropriate, all unknown $r_{ij}$ correspond to the distances between the spheres of different known lower-order subpackings. The triangular bipyramid rule is then applied to each unknown element of $\mathcal{D}$. For each unknown distance, $r_{ij}$, both solutions are potentially stored, as are all possible sets of unknown distances $\{r_{ij}\}$. Along the way, each $r_{ij}$ solution is tested for consistency, and it is always possible that both, 1, or neither solution will be consistent. Once all locally consistent $r_{ij}$ are stored, the resultant $\{r_{ij}\}$ are tested for global consistency.

A solution will be inconsistent, and thus unphysical, for one of the following reasons:

1. It violates the triangle inequality (meaning that no real solution exists—this shows up as the absolute value of the argument of the inverse cosine being greater than 1).
2. One or more distance(s) are less than $R$.
3. Different triangular bases lead to different $r_{ij}$; this indicates that a structure is in conflict with itself. One part of it implies it should have one structure, whereas another part implies a different structure. Such a structure is physically and mathematically inconsistent.

Violation 2 arises within the 5 particles of one triangular bipyramid and thus registers a physical local inconsistency. Violation 1 occurs within individual triangular bipyramids, as each $r_{ij}$ is determined, in which case it registers a local inconsistency; it also occurs over the entire set of triangular bipyramids, once all $\{r_{ij}\}$ have been determined, in which case it registers a global inconsistency. Violation 3 occurs when solutions are consistent within individual triangular bipyramids, and thus locally consistent, but inconsistent within combinations of triangular bipyramids—these solutions are thus **globally inconsistent**. This violation can be checked as follows: Figure 12 shows that the dihedral angle, $A_1$, is given by either the sum or the difference of $A_2$ and $A_3$ if particles $i$ and $j$ lie on opposite sides or on the same side of the triangular base, respectively (to within a $2\pi$ modulation of course). Test all possible 5-particle combinations of triangular bipyramids within the $n$-particle structure, and if there exists a triangular bipyramid that does not satisfy

$$A_1 = \frac{A_2 + A_3}{|A_2 - A_3|} \quad \frac{2\pi - (A_2 + A_3)}{2\pi - |A_2 - A_3|}$$

$^{14}$If $\mathcal{A}$ is not iterative, there will exist unknown $r_{ij}$ that do not correspond to distances between spheres of known subpackings. In this case, (10) will contain at least 1 unknown element on the right-hand side and cannot be applied directly.

$^{15}$In this case, some triangular bipyramids are locally consistent, whereas others are not. All possible triangular bipyramids of a structure need not be tested to solve for all $r_{ij}$; thus it is important to check all bipyramids to ensure global consistency once $\{r_{ij}\}$ have been determined. This violation is related to violation 3, except that here the violation is registered between the angles associated with the line segments, and in violation 3 the violation occurs within the dihedral angles.
the solution is globally inconsistent. (In calculating \( r_{ij} \) in (10), we need not consider the latter two \( A_1 \) solutions, as \( \cos(2\pi - x) = \cos(x) \)).

There is one scenario in which 5 points need not satisfy this global consistency check, and that is when 3 or more of the 5 points lie in a line. In this case, the 3 or more points define a line and not a plane, and thus the dihedral angle is not defined, and (11) cannot be applied. This situation is encountered in certain structures that contain octahedra (for example, see graphs 5416 and 10664 at \( n = 9 \) in the supplementary information [8]). In this scenario, the following global check can be performed: for \( m \) points lying in a line, there must exist the following number of line segments with a given minimum distance:

<table>
<thead>
<tr>
<th>Number of Line Segments</th>
<th>Minimum Length of Line Segment</th>
</tr>
</thead>
<tbody>
<tr>
<td>( m - 1 )</td>
<td>( R )</td>
</tr>
<tr>
<td>( m - 2 )</td>
<td>( 2R )</td>
</tr>
<tr>
<td>( \vdots )</td>
<td>( \vdots )</td>
</tr>
<tr>
<td>( m - (m - 1) )</td>
<td>( (m - 1)R )</td>
</tr>
</tbody>
</table>

(12)

Note that the first constraint, having \( m - 1 \) line segments with a minimum length of \( R \), is automatically satisfied by the fact that we eliminate all solutions containing a distance \( < R \). Up to \( n = 10 \), it turns out that performing this check for \( m = 3 \) alone is sufficient to find all globally inconsistent solutions. We evaluate that 3 points lie in a line by identifying that the angle associated with those 3 points is 0 or \( \pi \). We then check that there is at least one distance amongst the 3 points that is \( \geq 2 \) (where, once again, \( R = 1 \) without loss of generality). If this is not satisfied, then the packing is globally inconsistent and is eliminated. Note that the fact that some dihedral angles may not be defined was overlooked in [7], and this is the primary source of the difference in numbers reported here in Tables 1 and 4 versus the numbers previously reported in [7].

4.3. The growth of new seeds. For new seeds, we have a structure that contains an inherently new polyhedron, and thus some or all intrapolyhedral distances are also unknown; i.e., one or more of the 9 distances within \( \{ r_{ip}, r_{ik}, r_{iq}, r_{jp}, r_{jk}, r_{jq}, r_{pq}, r_{pq} \} \) are unknown. Thus, deriving the equation for \( r_{ij} \), as is done for iterative packings, will yield one equation with more than one unknown. The triangular bipyramid rule, therefore, cannot be applied directly to new seeds, and new geometrical rules must be derived to analytically solve noniterative \( A \)'s.

Using a general rule to solve for iterative \( A \)'s and deriving individual geometrical rules for noniterative \( A \)'s are feasible so long as the noniterative \( A \)'s do not grow too quickly with \( n \). Table 1 shows that this is the case for \( n \leq 9 \), where the number of noniterative \( A \)'s is 5 or less. However, at \( n = 10 \), there are 94 noniterative \( A \)'s.\(^{16}\) To sift through the 94 potential seeds at \( n = 10 \) requires inventing new geometrical rules, and the growth of such rules demonstrates that the method we have described does not scale efficiently with \( n \). In section 7.2, we will discuss a potential extension of the triangular bipyramid rule, which might be able to break this bottleneck, at least computationally. Here, we present the packing results derived from a combination of the triangular bipyramid rule and individual geometrical rules. For \( n \leq 9 \), we

\(^{16}\)Note that the noniterative and iterative \( A \)'s listed in this table are constructed after applying the geometrical rules for \( n \leq 8 \) that appear in the text and in the supplementary information [8]. Thus, this reflects the number of iterative and noniterative \( A \)'s with respect to these geometrical rules, and not the absolute number of iterative and noniterative \( A \)'s.
have analytically solved for all packings. At \( n = 10 \), we analytically solve for all iterative packings, and produce a preliminary list of new seeds, found by solving the noniterative \( A \)'s numerically using Newton iterations.\(^{17}\)

5. The set of sphere packings. Here we present the list of sphere packings derived by this method.\(^{18}\) In principle, the analytical method outlined here will yield a complete set of minimally rigid packings. However, we have not implemented the triangular bipyramid rule symbolically, leading to the following practical issues, which could lead to numerical errors.

**Numerical round-off error.** All calculations involving the triangular bipyramid rule are subject to numerical precision. Our algorithm eliminates many packings by finding situations where the argument of the \( \cos^{-1} \) term is larger than unity; this can also occur erroneously due to round-off error, causing packings to erroneously be recognized as unphysical. Similarly, round-off errors are possible when checking for the consistency of a packing, in checking the equivalence between dihedral angles, or checking that there exists at least one distance greater than or equal to 2 when 3 points lie in a line. Round-off issues could be improved by using general precision libraries such as *gmp* and *mpfr* [1, 2], or altogether avoided by doing all calculations symbolically. Thus, while the analytical method presented here should in principle yield a complete set of sphere packings, practical issues such as these are a source of error.

We present a complete set of sphere packings of \( n \leq 9 \), save round-off error. At \( n = 10 \), we present a complete set of iterative packings and a preliminary list of new seeds. Packings of \( n \leq 7 \) particles are included here, and packings of \( n \leq 8 \leq 10 \) particles are included in supplementary information [8]. All results are summarized in Table 4.

In the list presented here, \( \phi \) corresponds to the point group and \( \sigma \) to the symmetry number. We have included the 2nd moment of each packing, and a “*” appears next to the 2nd moment that corresponds to the minimum of the 2nd moment of \( n \) particles. The “special properties” column denotes whether a structure is convex, a new seed, chiral, or nonrigid. If the special properties column is blank, then that packing contains none of these properties.

\(^{17}\)Our runs of Newton Iterations used random initial guesses (between \(-5\) and \(5\) for the coordinates of the particles). We performed 20 iterations of each initial guess and approximately 150,000 total runs. Every matrix for which a solution was found was found multiple times. We believe this to be a reasonably thorough search and that this preliminary list of new seeds at \( n = 10 \) might be complete. It is worth noting that the preliminary list of new seeds reported here found by Newton iterations is the same as the list reported in [7] found by constructing the noniterative \( A \)'s manually with the construction toy *Geomags*.

\(^{18}\)The number of packings presented here for \( n = 9, 10 \) differs from the number we reported in [7]. This is primarily because our previous code did not run a check to ensure that the dihedral angle was well defined when checking for global consistency. As mentioned in section 4.2, this can occur when the 3 points used to define a plane are collinear, occurring in some of the packings that contain octahedra. In our original code (used for [7]), the dihedral angle check was still being performed in such an instance and erroneously deemed some packings globally inconsistent. It was a personal communication with Rob Hoy, who extended this work in [30], that brought it to our attention that 2 packings were missed in our \( n = 9 \) list. In examining this discrepancy, we discovered that this issue with the dihedral angles was what caused these packings to be missed, and we have since made the relevant correction in the code. This caused 52 packings to be realized at \( n = 9 \) and many more packings to be realized at \( n = 10 \). We also made 2 more modifications to the code, such as further correcting for numerical round-off error, which further caused several more packings to be registered at \( n = 10 \). Please see the supplemental information, Appendix C in [8], for a complete list of the changes that were made to the code.
2 Particle Packings

Packing 1 (Graph 1):

\[ A = \begin{pmatrix} 0 & 1 \\ 1 & 0 \end{pmatrix}, \quad C = \begin{pmatrix} 0 \\ 0 \end{pmatrix} \]

\[ D = \begin{pmatrix} 0 & 1 \\ 1 & 0 \end{pmatrix} R \]

<table>
<thead>
<tr>
<th>2nd Moment</th>
<th>( \phi )</th>
<th>( \sigma )</th>
<th>Special Properties</th>
</tr>
</thead>
<tbody>
<tr>
<td>*0.5R^2</td>
<td>2</td>
<td></td>
<td>New Seed</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Convex</td>
</tr>
</tbody>
</table>

3 Particle Packings

Packing 1 (Graph 1):

\[ A = \begin{pmatrix} 0 & 1 & 1 \\ 1 & 0 & 1 \\ 1 & 1 & 0 \end{pmatrix}, \quad C = \begin{pmatrix} 0 \\ 0 \\ 0 \end{pmatrix} \]

\[ D = \begin{pmatrix} 0 & 1 & 1 \\ 1 & 0 & 1 \\ 1 & 1 & 0 \end{pmatrix} R \]

<table>
<thead>
<tr>
<th>2nd Moment</th>
<th>( \phi )</th>
<th>( \sigma )</th>
<th>Special Properties</th>
</tr>
</thead>
<tbody>
<tr>
<td>*1.0R^2</td>
<td>6</td>
<td></td>
<td>Convex</td>
</tr>
</tbody>
</table>
4 Particle Packings

Packing 1 (Graph 1):

\[
A: \begin{pmatrix}
0 & 1 & 1 & 1 \\
1 & 0 & 1 & 1 \\
1 & 1 & 0 & 1 \\
1 & 1 & 1 & 0
\end{pmatrix}
\quad C: \begin{pmatrix}
0 \\
0 \\
0 \\
-1
\end{pmatrix}
\quad R
\]

\[
D: \begin{pmatrix}
0 & 1 & 1 & 1 \\
1 & 0 & 1 & 1 \\
1 & 1 & 0 & 1 \\
1 & 1 & 1 & 0
\end{pmatrix}
\quad R
\]


<table>
<thead>
<tr>
<th>2nd Moment</th>
<th>(\phi)</th>
<th>(\sigma)</th>
<th>Special Properties</th>
</tr>
</thead>
<tbody>
<tr>
<td>*1.5R²</td>
<td>(T_d)</td>
<td>12</td>
<td>Convex</td>
</tr>
</tbody>
</table>

5 Particle Packings

Packing 1 (Graph 1):

\[
A: \begin{pmatrix}
0 & 0 & 1 & 1 & 1 \\
0 & 0 & 1 & 1 & 1 \\
1 & 1 & 0 & 1 & 1 \\
1 & 1 & 1 & 0 & 1 \\
1 & 1 & 1 & 1 & 0
\end{pmatrix}
\quad C: \begin{pmatrix}
0 \\
0 \\
4/(3\sqrt{3}) \\
-4/3 \\
(2/3)\sqrt{2/3} \\
1/(2\sqrt{3})
\end{pmatrix}
\quad R
\]

\[
D: \begin{pmatrix}
0 & 2\sqrt{3/3} & 1 & 1 & 1 \\
2\sqrt{3/3} & 0 & 1 & 1 & 1 \\
1 & 1 & 0 & 1 & 1 \\
1 & 1 & 1 & 0 & 1 \\
1 & 1 & 1 & 1 & 0
\end{pmatrix}
\quad R
\]


<table>
<thead>
<tr>
<th>2nd Moment</th>
<th>(\phi)</th>
<th>(\sigma)</th>
<th>Special Properties</th>
</tr>
</thead>
<tbody>
<tr>
<td>*2.33333R²</td>
<td>(D_{2h})</td>
<td>6</td>
<td>Convex</td>
</tr>
</tbody>
</table>
6 Particle Packings

Packing 1 (Graph 2):

\[
A = \begin{pmatrix}
0 & 0 & 1 & 1 & 1 & 1 \\
0 & 0 & 0 & 1 & 1 & 1 \\
1 & 1 & 0 & 0 & 1 & 1 \\
1 & 1 & 0 & 0 & 1 & 1 \\
1 & 1 & 1 & 1 & 0 & 1 \\
1 & 1 & 1 & 1 & 1 & 0
\end{pmatrix}
\]

\[
C = \begin{pmatrix}
4/(3\sqrt{3}) \\
1/3 \\
(2/3)\sqrt{2/3} \\
0 \\
-1 \\
-4/9
\end{pmatrix}
\]

\[
R = \begin{pmatrix}
(10/9)\sqrt{3/3} \\
1/(2\sqrt{3}) \\
0 \\
0 \\
1/2 \\
0
\end{pmatrix}
\]

2nd Moment \( \phi \) \( \sigma \) Special Properties

| 3.35185 \( R^2 \) | \( C_{2v} \) | 2 |

Packing 2 (Graph 4):

\[
A = \begin{pmatrix}
0 & 0 & 1 & 1 & 1 & 1 \\
0 & 0 & 1 & 1 & 1 & 1 \\
1 & 1 & 0 & 0 & 1 & 1 \\
1 & 1 & 0 & 0 & 1 & 1 \\
1 & 1 & 1 & 1 & 0 & 0 \\
1 & 1 & 1 & 1 & 0 & 0
\end{pmatrix}
\]

\[
C = \begin{pmatrix}
0 \\
0 \\
0 \\
1 \\
-1 \\
-1
\end{pmatrix}
\]

\[
R = \begin{pmatrix}
0 \\
1/2 \\
0 \\
0 \\
-1/2 \\
-1/2
\end{pmatrix}
\]

\[
P = \begin{pmatrix}
0 & \sqrt{2} & 1 & 1 & 1 & 1 \\
\sqrt{2} & 0 & 1 & 1 & 1 & 1 \\
1 & 1 & 0 & \sqrt{2} & 1 & 1 \\
1 & 1 & \sqrt{2} & 0 & 1 & 1 \\
1 & 1 & 1 & 1 & 0 & \sqrt{2} \\
1 & 1 & 1 & 1 & \sqrt{2} & 0
\end{pmatrix}
\]

2nd Moment \( \phi \) \( \sigma \) Special Properties

| \( ^*3.0 R^2 \) | \( O_h \) | 24 | New Seed Convex |
7 Particle Packings

Packing 1 (Graph 4):

\[
A = \begin{pmatrix}
0 & 0 & 0 & 1 & 1 & 1 & 1 & 1 \\
0 & 0 & 0 & 1 & 0 & 1 & 1 & 1 \\
0 & 0 & 0 & 0 & 1 & 1 & 1 & 1 \\
1 & 1 & 0 & 0 & 0 & 1 & 1 & 1 \\
1 & 0 & 1 & 0 & 0 & 1 & 1 & 1 \\
1 & 1 & 1 & 1 & 1 & 0 & 1 & 1 \\
1 & 1 & 1 & 1 & 1 & 1 & 0 & 1
\end{pmatrix}
\]

\[
D = \begin{pmatrix}
0 & 2\sqrt{\frac{2}{3}} & 2\sqrt{\frac{2}{3}} & 0 & 1 & 1 & 1 & 1 \\
2\sqrt{\frac{2}{3}} & 0 & 4\sqrt{\frac{2}{3}} & 0 & 1 & 1 & 1 & 1 \\
2\sqrt{\frac{2}{3}} & 4\sqrt{\frac{2}{3}} & 0 & \frac{2}{\sqrt{3}} & 0 & 1 & 1 & 1 \\
1 & 1 & \frac{2}{\sqrt{3}} & \frac{2}{\sqrt{3}} & 0 & 1 & 1 & 1 \\
1 & 1 & 1 & 1 & 1 & 0 & 1 & 1 \\
1 & 1 & 1 & 1 & 1 & 1 & 0 & 1
\end{pmatrix}
\]

\[
C = \begin{pmatrix}
0 & 0 & 0 & 0 & 0 \\
4/(3\sqrt{3}) & -1/3 & (3/2)\sqrt{2/3} & 20/(9\sqrt{3}) & -4/9 \\
(10/9)\sqrt{2/3} & 0 & 0 & 0 & 0
\end{pmatrix}
\]

Packing 2 (Graph 8):

\[
A = \begin{pmatrix}
0 & 0 & 0 & 0 & 1 & 0 & 1 & 1 \\
0 & 0 & 0 & 0 & 1 & 0 & 1 & 1 \\
0 & 0 & 0 & 0 & 1 & 0 & 1 & 1 \\
1 & 1 & 0 & 0 & 1 & 1 & 1 & 1 \\
1 & 0 & 1 & 0 & 1 & 1 & 1 & 1 \\
0 & 1 & 1 & 1 & 1 & 1 & 1 & 1 \\
1 & 1 & 1 & 1 & 1 & 1 & 1 & 1
\end{pmatrix}
\]

\[
D = \begin{pmatrix}
0 & 1/2 & 1/2 & 1/2 & 1 & 0 & 1 \\
1/2 & 0 & 1/2 & 1/2 & 1 & 1 \\
1/2 & 1/2 & 0 & 1/2 & 1 & 1 \\
1 & 1 & 1 & 1 & 1 & 0 & 1 \\
2\sqrt{3} & 0 & 1/2 & 1/2 & 1 \\
1 & 1 & 1 & 1 & 1 & 0 & 1
\end{pmatrix}
\]

\[
C = \begin{pmatrix}
0 & 0 & 0 & 0 & 4/(3\sqrt{3}) & 1/3 & (2/3)\sqrt{2/3} & (1/2)\sqrt{2/3} \\
-1 & 0 & 0 & 0 & 0 & 0 & (10/9)\sqrt{2/3} & (1/2)\sqrt{2/3}
\end{pmatrix}
\]

<table>
<thead>
<tr>
<th>2nd Moment</th>
<th>$\phi$</th>
<th>$\sigma$</th>
<th>Special Properties</th>
</tr>
</thead>
<tbody>
<tr>
<td>$4.24868R^2$</td>
<td>$C_{2v}$</td>
<td>2</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>2nd Moment</th>
<th>$\phi$</th>
<th>$\sigma$</th>
<th>Special Properties</th>
</tr>
</thead>
<tbody>
<tr>
<td>$4.47619R^2$</td>
<td>$C_{3v}$</td>
<td>3</td>
<td></td>
</tr>
</tbody>
</table>
Packing 3 (Graph 17):

\[
A = \begin{pmatrix}
0 & 1 & 0 & 1 & 1 & 1 \\
0 & 0 & 1 & 0 & 1 & 1 \\
1 & 0 & 0 & 1 & 0 & 1 \\
1 & 1 & 1 & 0 & 1 & 1 \\
1 & 1 & 0 & 1 & 1 & 1 \\
1 & 1 & 1 & 1 & 1 & 0
\end{pmatrix}, \quad P = \begin{pmatrix}
0 & 2\sqrt{\frac{3}{5}} & 1 & \frac{\sqrt{2}}{3} & 1 & 1 \\
2\sqrt{\frac{3}{5}} & 0 & \frac{\sqrt{2}}{3} & 1 & 1 & 1 \\
1 & \frac{\sqrt{2}}{3} & 0 & \frac{\sqrt{2}}{3} & 1 & 2\sqrt{\frac{3}{5}} \\
\frac{\sqrt{2}}{3} & 1 & \frac{\sqrt{2}}{3} & 0 & 1 & 1 \\
1 & 1 & 2\sqrt{\frac{3}{5}} & 1 & 1 & 0 \\
1 & 1 & 1 & 1 & 1 & 0
\end{pmatrix}, \quad C = \begin{pmatrix}
\frac{4\sqrt{3}}{9} & \frac{2}{3}\sqrt{\frac{3}{5}} & 0 & -1 & 0 & 0 \\
\frac{2}{3}\sqrt{\frac{3}{5}} & \frac{2}{3}\sqrt{\frac{3}{5}} & 1 & \frac{2}{3}\sqrt{\frac{3}{5}} & 1 & 2\sqrt{\frac{3}{5}} \\
0 & 1 & 1 & \frac{2}{3}\sqrt{\frac{3}{5}} & 1 & 1 \\
\frac{2}{3}\sqrt{\frac{3}{5}} & 1 & 1 & 1 & 1 & 0 \\
0 & 1 & 1 & 1 & 1 & 0
\end{pmatrix}
\]

<table>
<thead>
<tr>
<th>2nd Moment</th>
<th>4.64501$R^2$</th>
<th>$C_2$</th>
<th>Special Properties</th>
<th>Chiral</th>
</tr>
</thead>
</table>

Packing 4 (Graph 22):

\[
A = \begin{pmatrix}
0 & 1 & 1 & 1 & 0 & 1 \\
0 & 1 & 1 & 1 & 1 & 1 \\
1 & 0 & 0 & 0 & 1 & 1 \\
1 & 1 & 0 & 0 & 0 & 1 \\
0 & 1 & 1 & 0 & 0 & 1 \\
1 & 1 & 1 & 1 & 1 & 1 \\
0 & 0 & 0 & 0 & 0 & 0
\end{pmatrix}, \quad P = \begin{pmatrix}
0 & \sqrt{\frac{3}{2}} x & 1 & 1 & \sqrt{\frac{3}{2}} x & 1 \\
\sqrt{\frac{3}{2}} x & 0 & \sqrt{\frac{3}{2}} x & 1 & 1 & 1 \\
1 & \sqrt{\frac{3}{2}} x & 0 & \sqrt{\frac{3}{2}} x & 1 & 1 \\
\sqrt{\frac{3}{2}} x & 1 & \sqrt{\frac{3}{2}} x & 0 & \sqrt{\frac{3}{2}} x & 1 \\
1 & 1 & 1 & 1 & 1 & 1 \\
1 & 1 & 1 & 1 & 1 & 1 \\
0 & \sqrt{\frac{3}{2}} x & 1 & 1 & \sqrt{\frac{3}{2}} x & 0
\end{pmatrix}, \quad C = \begin{pmatrix}
0 & 0 & 0 & 0 & 0 & 0 \\
0 & 0 & (1/2)(1/3)(5 + \sqrt{5}) & (1/4)(-3 - \sqrt{5}) & 0 & 0 \\
0 & (1/2)(1/3)(5 + \sqrt{5}) & (1/4)(-3 - \sqrt{5}) & 0 & 0 & 0 \\
(1/9)(\sqrt{2}(5 + \sqrt{5}) + 10(5 + \sqrt{5})) & 0 & 0 & 0 & 0 & 0 \\
(5 + \sqrt{5})^2 / (20\sqrt{2}) & 0 & 0 & 0 & 0 & 0 \\
-(1/10)(5 - \sqrt{5}) & -(1/10)(5 - \sqrt{5}) & -(1/10)(5 - \sqrt{5}) & -(1/10)(5 - \sqrt{5}) & -(1/10)(5 - \sqrt{5}) & -(1/10)(5 - \sqrt{5})
\end{pmatrix}
\]

<table>
<thead>
<tr>
<th>2nd Moment</th>
<th>4.17082$R^2$</th>
<th>$D_{2h}$</th>
<th>Special Properties</th>
<th>Convex</th>
</tr>
</thead>
</table>
6. Properties of packings. Here we highlight some interesting properties of packings.

6.1. New seeds. New seeds are interesting because they are inherently new structures of \( n \) particles. They are also “generating sets”; i.e., once they exist at a given starting \( n = m \), they are propagated iteratively for all \( n > m \). Geometrically, new seeds are inherently global structures, stabilized exactly by the \( n \) particles for which that new seed arises. Iterative packings are geometrically locally stable, in that subsets of less than \( n \) particles within the packing also correspond to packings. Thus, new seeds are unique events for a given number of particles, \( n \). Figure 13 shows all new seeds of \( n \leq 10 \) particles (where the set of new seeds at \( n = 10 \) is putative). The proportion of new seeds to total packings is relatively small for small \( n \), which can be seen in Table 4.

6.2. Rigidity. We have enumerated packings satisfying minimal rigidity constraints; these constraints are necessary but not sufficient for rigidity, and thus we can find nonrigid packings that satisfy these constraints. For these packings, there exists a degree of freedom in which particles can move without breaking or forming additional contacts. The first instance of a nonrigid packing occurs at \( n = 9 \), at which there is one. At \( n = 10 \), there are 4 nonrigid packings: 1 nonrigid new seed, and 3 iterative nonrigid packings that derive from the \( n = 9 \) nonrigid new seed (Figures 14 and 15).

These 10 particle nonrigid packings will iteratively produce at least \( m \geq 1 \) nonrigid packings at \( n = 11 \), and so on. All nonrigid packings enumerated thus far contain at least 2 deformable open square faces. We do not know whether or not at least 2 open square faces are a requisite of nonrigid packings that satisfy minimal rigidity constraints. The open square faces must be “connected” for the extra degree of freedom to exist—in the packings encountered thus far, this manifests itself by the existence of half-octahedra sharing at least 1 vertex.
Fig. 13. New Seeds. All new seeds of $n \leq 10$ particles shown in both sphere and point/line representation. There exists only 1 packing for each $n$ of $n \leq 5$ particles that can each be constructed iteratively from a dimer; thus there exist no new seeds for $n \leq 5$. $n = 6$ is the first instance of a new seed. The set of new seeds reported for $n = 10$ is putative and thus represents a lower bound. New seeds with a * appearing to the right correspond to minima of the 2nd moment out of all packings for that $n$. It can be seen here that, for the packings we have analyzed, when more than 1 packing exists, the minimum of the 2nd moment happens to correspond to a new seed.

Fig. 14. The 9-particle nonrigid new seed (grey) is shown in the top left-hand box. It is composed of two joined 5-point polytetrahedra (red) attached to two joined half-octahedra (blue). The substructures are shown overlain—purple bonds and particles are shared by both substructures, whereas only red or blue bonds and particles belong to the polytetrahedral or octahedral structures alone, respectively. Two representative ways of forming this nonrigid structure are shown in (a) and (b). (a) Two 5-point polytetrahedra are joined by sharing one common point (on bottom). The two polytetrahedra are then fully attached to one another via the remaining 3 bonds first shown in dashed black lines, as potential bonds, and then in solid white lines, as actualized bonds. These bonds form the two connected half-octahedra. (b) 2 particles (red) are attached to the concave side faces of the 2 joined half-octahedra (blue). The 2 red particles form the two 5-point polytetrahedral substructures once they are attached to the joined octahedra.
Fig. 15. Nonrigid Packings of 9 and 10 Spheres. (a) The nonrigid $n = 9$ packing, with nonrigid motion, corresponding to a twisting of the square faces, shown by black arrows. (b) Nonrigid $n = 10$ packings formed iteratively by adding one particle to the nonrigid $n = 9$ seed. The nonrigid motion of these structures is the same as in (a). (c) Nonrigid $n = 10$ seed. Nonrigid motion, corresponding to a twisting of the radially connected square faces, connected by the bottom particle, is shown by the black arrows. This twisting motion can be accomplished by, for example, twisting the top triangle.

Fig. 16. Tree Convergence in a 10-Particle Packing. An example of tree convergence in one of the 25 contact packings of 10 particles. This packing, shown in grey (top panel), can be decomposed into (bottom left panel) the 9-particle nonrigid new seed (red) plus one particle (blue) that rigidifies the structure, or into (bottom right panel) an octahedron (blue) with 2 attached polytetrahedra (red). The red dashed line indicates an “implicit contact,” a contact that automatically forms once the other contacts are in place (this corresponds to the 25th contact). Color is available online only.

6.3. The tree nature of packings. There is a distinct tree nature to packings. New seeds are the origin of a branch in the tree. Iterative packings continue the branch. All $n$-particle iterative packings can be decomposed into combinations of less than $n$-particle packings, and this decomposition is often not unique. When the decomposition is not unique, the branches of the tree converge. Figures 16 and 17 show examples of tree convergence. Figure 16 shows the 2 possible decompositions of one of the 10-particle 25 contact packings. This packing can be formed either by adding 1 particle to the 9 particle nonrigid new seed or by combining two polytetrahedra with a 6-particle octahedron.

Figure 17 shows the tree structure of $2 \leq n \leq 8$ packings. It can be seen that tree convergence occurs from $n = 7$ to $n = 8$, where multiple 7-particle packings produce the same 8-particle packing under the addition of another particle.
Fig. 17. Tree Convergence for $n \leq 8$. An example of tree convergence for $2 \leq n \leq 8$. Packings above which there is no arrow correspond to new seeds and thus to the beginning of a new branch. The arrows then point to the $n$-particle packing or group of packings that form iteratively by adding one particle. It can be seen that even for the iterative case of adding one particle, there is tree convergence from $n = 7$ to $n = 8$ (shown by multiple arrows feeding into the same packing).

6.4. Minima of the 2nd moment. The 2nd moment measures the deviation of particles from their collective center (centroid) and is given by

$$M = \sum_{i=1}^{n} |\mathbf{r}_i - \mathbf{c}|^2 = \sum_{i=1}^{n} (x_i - c_x)^2 + (y_i - c_y)^2 + (z_i - c_z)^2,$$

where $\mathbf{r}_i$ is the $x,y,z$ position of particle $i$, and $\mathbf{c}$ is the centroid, the average $x,y,z$ position over all particles, given by

$$c_x = \frac{1}{n} \sum_{i=1}^{n} x_i$$

and analogously given for $c_y$ and $c_z$.

The minimum of the 2nd moment corresponds to the packing with the smallest $M$. The 2nd moment is listed within the list of packings in section 5 and in Appendix B in [8], and a “*” signifies the minimum of the 2nd moment for each $n$ in Figure 13. We confirm that the minima of the 2nd moment reported by Sloane et al. [48] are correct (they proved the 2nd moment minima for $n \leq 4$, but for $n > 4$ these were putative structures). For $n \leq 10$, each minimum of the second moment corresponds to a new seed, if a new seed exists.
6.5. Ground states and the maximum number of contacts. A fundamental question related to sphere packings is, What is the maximum number of contacts that a packing of \( n \) spheres can have? Not only is this question of mathematical interest in its own right, but it is also of significant physical interest, as such packings correspond to ground states. The number of packings that contain the maximum number of contacts in turn corresponds to the ground state degeneracy. Table 5 shows how the ground state degeneracy changes with \( n \). Interestingly, this relation appears to be oscillatory.

For \( n \leq 9 \), every packing has exactly \( 3n - 6 \) contacts. Thus, at each fixed \( n \), \( n \leq 9 \), all packings have the same potential energy. Table 5 shows that, for \( n \leq 9 \), the ground state degeneracy increases exponentially. But at \( n = 10 \) this trend changes due to a small number of packings that can have \( 25 = 3n - 5 \) contacts (all other 10-particle packings have \( 3n - 6 \) contacts). There exist 3 such packings, each containing octahedra (Figure 18(a)–(c)). These three structures are the ground states at \( n = 10 \).

For \( n \geq 11 \), we conjecture as to what the maximum number of contacts is and provide examples of such structures. The maximum contact packings at \( n = 10 \) arise because it becomes possible to add 4 contacts to minimally rigid 9-particle packings, whereas all other iterative packings of \( n \leq 10 \) spheres are formed by the addition of \( 3m \) contacts to a minimally rigid \( n - m \) sphere packing. All maximum contact packings found thus far correspond to iterative packings. We have not determined whether this is true for all \( n \), but we conjecture that it is, because new seeds tend to contain more empty space and thus fewer contacts. We have found three types of structures that allow for the addition of more than \( 3m \) contacts: (i) \( m \) octahedra, where each pair of octahedra shares one edge (as in Figure 18(c)), (ii) an open square face created by half an octahedra (as shown in blue in Figure 18(a)), and (iii) the concave \( m \) point face created by octahedra sharing 3 edges (as shown in blue in

### Table 5

<table>
<thead>
<tr>
<th>( n )</th>
<th>( 3n - 6 ) Contacts</th>
<th>Ground state packings</th>
</tr>
</thead>
<tbody>
<tr>
<td>2</td>
<td>3</td>
<td>1</td>
</tr>
<tr>
<td>3</td>
<td>3</td>
<td>1</td>
</tr>
<tr>
<td>4</td>
<td>3</td>
<td>1</td>
</tr>
<tr>
<td>5</td>
<td>3</td>
<td>1</td>
</tr>
<tr>
<td>6</td>
<td>3</td>
<td>2</td>
</tr>
<tr>
<td>7</td>
<td>3</td>
<td>5</td>
</tr>
<tr>
<td>8</td>
<td>3</td>
<td>13</td>
</tr>
<tr>
<td>9</td>
<td>3</td>
<td>52</td>
</tr>
<tr>
<td>10</td>
<td>3</td>
<td>3</td>
</tr>
<tr>
<td>11</td>
<td>3</td>
<td>1</td>
</tr>
<tr>
<td>12</td>
<td>3</td>
<td>1</td>
</tr>
<tr>
<td>13</td>
<td>3</td>
<td>7</td>
</tr>
<tr>
<td>14</td>
<td>3</td>
<td>1</td>
</tr>
<tr>
<td>15</td>
<td>3</td>
<td>1</td>
</tr>
<tr>
<td>16</td>
<td>3</td>
<td>1</td>
</tr>
<tr>
<td>17</td>
<td>3</td>
<td>8</td>
</tr>
<tr>
<td>18</td>
<td>3</td>
<td>1</td>
</tr>
<tr>
<td>19</td>
<td>3</td>
<td>1</td>
</tr>
<tr>
<td>20</td>
<td>3</td>
<td>1</td>
</tr>
</tbody>
</table>
Fig. 18. Maximum contact packings for $10 \leq n \leq 20$. We are reasonably confident that the $n = 10$ packings shown here correspond to the maximum contact packings, but maximum contact packings of $n > 10$ are conjectured. (a)–(c) 10-particle packings with $3n - 5 = 25$ contacts. (a) can be formed either by adding one particle (red) to one of the open square faces (blue) or as otherwise detailed in Figure 16. (b) can be formed by adding one particle (red) to the concave 4-particle face created by 2 joined octahedra (blue). (c) is formed by connecting 2 octahedra by one edge. (d) 11-particle maximum contact packing ($3n - 4 = 29$ contacts). This can be formed either by adding one particle (red) to the concave 4-particle face created by the 2 joined octahedra of (b) or by adding one particle to the one remaining open square face of (a). (e) 12-particle maximum contact packing ($3n - 3 = 33$ contacts). This is formed by adding one particle (red) to the concave 4-particle face created by 2 joined octahedra (blue). 13-particle maximum contact packings are constructed iteratively from this packing. (f) 14-particle maximum contact packing ($3n - 2 = 40$ contacts). This is formed by 3 radially connected octahedra (blue) and 2 particles (yellow) added to each of the concave 5-particle faces created by the 3 joined octahedra. (g) 15- and 16-particle maximum contact packings containing $3n - 1$ and $3n$ contacts, respectively. The 15-particle packing corresponds to the addition of only one of the red particles to the concave 4-particle face of (f), and the 16-particle packing includes both red particles. (h) 17-particle 3n contact packing formed by 4 radially connected octahedra (blue) and 2 particles (yellow) connected to the concave 6-particle faces created by the 4 joined octahedra. At 17 particles, 3n contact packings correspond to this packing as well as packings constructed iteratively from (g). (i) 18-, 19-, 20-particle maximum contact packings corresponding to $3n + 1$, $3n + 2$, and $3n + 3$ contacts, respectively. Each of these packings is constructed by adding a particle (red) to one of the concave 4-particle faces created by the joined octahedra—the 18-particle packing is constructed by adding one such particle to (h), 19 by adding 2, and 20 by adding 3. Color is available online only.

Figure 18(d)–(e)). 4-point concave faces are shown, for example, in Figure 18(e); 5-point concave faces in Figure 18(f)–(g); and 6-point concave faces in Figure 18(h)–(i). To each $m$ point concave face it is possible to add one particle with $m$ contacts; this is evidenced, for example, by the red particle with 4 contacts in Figure 18(e) and the yellow particles with either 5 or 6 contacts in Figure 18(f)–(i). All structures leading to maximum contact packings that we have found thus far are related to octahedra, and we conjecture that this will be the case for all $n$. (Interestingly all nonrigid packings found thus far are also related to octahedra, in that they contain the open square faces created by half-octahedral structures; they all contain half-octahedra sharing at least 1 point.)
There are many fewer ways of adding greater than $3n$ contacts to an $n - m$ sphere packing than there are of adding $3m$ contacts, thus leading to a relatively small number of ground state packings when a new maximum number of contacts, as a function of $n$, is reached. Thus, each time a packing with a greater maximum number of contacts, as a function of $n$, is possible, we expect the ground state degeneracy to either decrease or to remain small. When the functional form for the maximum number of contacts remains constant, we expect the ground state degeneracy to grow rapidly, due in large part to the iterative growth of adding a particle with 3 contacts to packings. Table 5, for example, shows that for $n \leq 9$ the ground state degeneracy increases exponentially because all packings have $3n - 6$ contacts. But at $n = 10$ this trend changes because packings with more than $3n - 6$ contacts become possible. At $n = 13$ and at $n = 17$, rapid growth in the ground state degeneracy resumes as the functional form for the maximum number of contacts remains constant, but for all other $n$, the ground state degeneracy either decreases or remains constant, because a new maximum number of contacts, as a function of $n$, is possible.

At low temperatures, we expect the experimentally observable packings to be dominated by the maximum contact packings. Thus, for $n = 10$, and for any $n$ in which either (i) the maximum number of contacts increases as a function of $n$ or (ii) the maximum number of contacts does not remain constant for too long ($\leq 3$ particles), we expect that there are only a small number of packings that will be observable at low temperature. This trend can be seen in the conjectured ground state degeneracy for $9 < n \leq 20$ in Table 5, and we conjecture that a similar low amplitude oscillatory trend might also continue for $n > 20$.

6.6. Lattice structure. The nonrigid new seeds at $n = 9$ and $n = 10$, as well as the maximum contact packings of $n < 13$ are all subunits of the hexagonally close-packed (HCP) lattice, being combinations of face-sharing tetrahedra and octahedra. Additionally, the structure shown in Figure 18(c) is a subunit of either the HCP or face-centered cubic (FCC) lattice. The nonrigid packings are entropically favored, and we thus expect these to form with higher probability at higher temperatures, while the maximum contact packings are energetically favored, corresponding to the structures that will form with higher probability as the temperature $T \to 0$. For $14 \leq n \leq 20$, the maximum contact packings are not HCP subunits (Figure 18(f)–(i)).

Frank predicted [20] that icosahedral short-range order would be a hallmark of liquid structure, and experimental studies have shown local cluster-like order in bulk atomic liquids and glasses [44, 47]. Results from a recent study suggest that structural arrest in condensed phases may be related to geometrical constraints at the scale of a few particles [45]. The propensity for icosahedra [28, 16] in longer-range systems is absent in ours. We have proven that the icosahedron is not the ground state at $n = 12$, nor is an icosahedron with a central sphere the ground state at $n = 13$. A 12-sphere icosahedron has only $3n - 6 = 30$ contacts, and in a 13-sphere icosahedron the outer spheres would not be close enough to interact with each other.

It is possible, and perhaps even likely, that the lattice structures corresponding to ground state packings will be periodic with $n$. For example, although the ground states for $14 \leq n \leq 20$ are not commensurate with HCP, the ground states for a finite range of higher $n$ may be, and may then subsequently return to the lattice structure commensurate with $14 \leq n \leq 20$. Detailing the structures of ground state packings for all $n$, and geometrical patterns contained therein, is a subject of future work. Furthermore, the appearance of crystalline order, such as HCP, at very low $n$ may influence nucleation.
7. Extensions and conjectures.

7.1. The major roadblock for reaching higher $n$. The main roadblock to the analytical enumeration of sphere packings at higher $n$ in the current work is deriving one analytical geometrical rule that can solve for all new seeds. In the next section, we outline a numerical method, based on the triangular bipyramid rule, that is capable of finding all solutions of $A \rightarrow D$ and which can thus solve for all new seeds. However, the implementation of either this numerical scheme or the derivation of an analytical rule would only allow us to enumerate packings of up to about $n = 14$ spheres. This is because the real limitation of the current method arises from the enumeration of minimally rigid, nonisomorphic adjacency matrices. For $n < 10$, the enumeration of such adjacency matrices using nauty [42] takes on the order of seconds. For $n = 10, 11$, this enumeration takes minutes. For $n = 12$, enumerating all minimally rigid, nonisomorphic $A$'s takes approximately 2 hours. Extrapolating, we expect the enumeration at $n = 13, 14$ to take on the order of 2 days and 2 weeks, respectively. Thus, around $n = 14$, we begin to reach the computational limitations of this method, which is due to the enumeration of $A$'s.

Only a very small fraction of adjacency matrices correspond to sphere packings; for example, at $n = 10$, out of the 750,226 $A$'s, only 262 correspond to sphere packings. Thus, the enumeration of all $A$'s really is a brute force and wasteful step. Further advances in enumerating sphere packings will require overcoming this roadblock. In section 7.3, we propose one method that might be able to overcome this limitation.

7.2. Applying the triangular bipyramid rule to new seeds. The triangular bipyramid rule solves for iterative structures but does not work for noniterative ones, which we also showed increase rapidly starting at $n = 10$. Here, we discuss how the triangular bipyramid rule might also be applied to new seeds. In this case, the equations for the unknown interparticle distances, $r_{ij}$, are implicit and thus must be solved numerically.

In a new seed, any triangular bipyramid that contains an unknown distance will contain more than one unknown distance. Thus, $r_{ij}$ cannot be determined one at a time, as with iterative packings; instead we must construct a system of equations to solve for the unknown distances.\textsuperscript{19} Let us consider a general triangular bipyramid, like that shown in Figure 12, but here all 10 distances are potentially unknown (Figure 19). For $m$ unknown $r_{ij}$, we construct $m$ triangular bipyramids to solve for the $r_{ij}$ (see Figure 20, for example). Each triangular bipyramid yields one equation for an unknown distance, thus yielding $m$ equations with $m$ unknowns in total, making the system well defined.

We assign a label, $d_i$, to each of the 10 distances within the triangular bipyramid (Figure 19). Explicit formulas can always be obtained for $d_2$ and $d_3$. Thus, over all $m$ triangular bipyramids, we place each unknown distance in location $d_2$ or $d_3$ at least once. New seeds are inherently global structures; thus the $m$ triangular bipyramids should contain all $n$ points amongst them in order to ensure that the solution space is sufficiently constrained. Also, to avoid redundancy, each triangular bipyramid must contain a unique combination of 5 particles.

The equations for the $r_{ij}$ are derived from the triangular bipyramid property that the dihedral angle $A_1$ must equal either the sum or the difference of the dihedral angles $A_2$ and $A_3$ (as was detailed in section 4 and in Figure 12). This equation

\textsuperscript{19}As can be seen in section 4 or in Appendix A of the supplemental information [8], the equation for an unknown distance is quadratic. Thus, this will be a quadratic system of equations.
is cumbersome, and it as well as its derivation can be found in Appendix A of the supplemental information [8]. As always with the triangular bipyramid rule, due to the 2 possibilities of $A_1$ equaling either the sum or the difference of $A_2$ and $A_3$, each unknown distance has 2 possible solutions.

For each set of $r_{ij}$ that is to be solved, construct initial guesses between the bounds of the triangle inequality and no-overlap constraint, and iterate the initial guess with a step size less than or equal to the minimum difference between different solutions (for rigid structures). There will always exist unknown $r_{ij} \leq 2R$ because each particle has at least 3 contacts.\footnote{This also holds true when each particle has at least 2 contacts.} These are the $r_{ij}$ for which there exists a $k$ satisfying $A_{ik} = A_{jk} = 1$, $A_{ij} = 0$. Thus, first solve the set of $R \leq r_{ij} \leq 2R$. If unknown $r_{ij}$ remain, then solve the set of $r_{ij}$ that now have known triangle inequality bounds, due to the previously solved set of $r_{ij}$; repeat until all $r_{ij}$ have been solved.\footnote{We have tested this method on the new seed $A$'s for $n \leq 8$ and have shown that it works; however, we have not implemented it for up to $n = 14$.}

### The bond breakage conjecture

A packing of $n$ spheres can be formed by (i) taking an $n - m$ sphere packing, breaking a contact (or bond), adding $m$ new spheres, and forming the appropriate contacts to complete the packing, or by (ii) breaking one bond of an $n$ sphere packing and reforming another. From this property, we propose the following theory.

**Bond breakage conjecture.** All packings of $n$ spheres can be obtained by breaking one bond and reforming another in every possible way. For any packing, there exists an $m$ step path, of breaking one contact and reforming another, that will form that packing out of another packing with $3n - 6$ contacts. Each of the $m$ steps will corre-
Fig. 20. A New Seed and the General Triangular Bipyramid. Here we show an example of how to apply the general triangular bipyramid to a new seed in order to determine its unknown distances. This new seed corresponds to the 6-particle octahedron. It has 3 unknown distances, $r_{12}$, $r_{34}$, and $r_{56}$ (dashed black lines). The first triangular bipyramid constructed consists of particles 1, 2, 3, 4, 5 and has unknown distances $d_1$ and $d_2$ corresponding to $r_{34}$ and $r_{12}$, respectively. (Note that $d_3$ to $d_{10} = R$.) The second triangular bipyramid consists of particles 1, 2, 3, 4, 6 and has unknown distances $d_1'$, $d_2'$ corresponding to $r_{12}$, $r_{34}$, respectively. The third triangular bipyramid consists of particles 1, 2, 4, 5, 6 and has unknown distances $d_1''$, $d_2''$ corresponding to $r_{12}$, $r_{56}$, respectively. Note that the first two triangular bipyramids comprise 2 equations and 2 unknowns, and thus are alone sufficient to solve $r_{12}$ and $r_{34}$. Once these 2 distances are known, applying the third triangular bipyramid involves only 1 unknown distance, $r_{56}$, and thus reduces to the same scenario as applying a triangular bipyramid to an iterative packing.

respond to an $n$-particle packing. The end points of the path (i.e., which $3n - 6$ packing one begins with and which packing one ends up with) determine what value $m$ takes. For every packing, there exists at least 1 other packing for which $m = 1$.

This suggests an alternative method for enumerating all packings of $n$ spheres: construct just one $3n - 6$ contact packing of $n$ particles (this can easily be done; simply construct an $n$-particle polytetrahedron, for example), and then break and reform bonds in all possible ways. For each packing, it is important to explore every combination of breaking and reforming a bond, i.e., to go down all paths and not just one path.

We have confirmed this conjecture up to as high as we have enumerated packings ($n = 10$) using the following algorithm:

For every $\mathcal{A}$ that corresponds to a packing
1. For each 1 that appears in the $\mathcal{A}$:
   (a) Swap the 1 with an existing 0. Do this in every possible nonisomorphic way. (This is the mathematical analogue of physically breaking an existing bond and reforming a different bond that was not present in that packing.)
For each new $A$ that is generated by swapping a 1 and a 0 (these are 1 bond away $A$'s, i.e., where $m = 1$):

i. Test for an isomorphism with the $A$'s of all other packings (i.e., all $A$'s other than the one being examined).

ii. If an isomorphism is found, stop the examination of this $A$, as it has been shown that there exists a 1 bond bath between the packing being examined and another packing.\textsuperscript{22}

Thus implementing this algorithm computationally, we have proven that, for each $n \leq 10$, every packing is a 1 bond distance away from at least 1 other packing of the same $n$. We have not proven this for $n > 10$, as we have not enumerated all packings of $n > 10$, but we suspect that this conjecture holds for all $n$.

Mapping out all possible 1 bond distances might be able to shed insight into the kinetic pathways of packings.

To implement the bond breakage conjecture into an improved method for enumerating sphere packings, the bonds must be broken and reformed intelligently, such that unphysical conformations are not explored. If all physical and unphysical conformations were to be explored, then we would simply return to the same computational problem we had with enumerating all nonisomorphic $A$'s. One should be able to break a bond and reform it only as is physically possible by calculating the 1 degree of freedom motion that is left over from the one broken bond. Furthermore, one can take advantage of symmetry to, a priori, not explore redundant (i.e., isomorphic) pathways of breaking and reforming bonds.

7.4. Extensions to other dimensions. The method presented here is, in large part, not dimension specific. The only step which is dimension specific is the set of geometrical rules used to solve $A$ for $D$. However, at least for the triangular bipyramid rule (which solves for most of the packings), the geometrical rules can easily be modified to account for a different number of dimensions, $d$. Once this is done, the same method can be used to solve for sphere packings of $d \neq 3$ dimensions.

7.5. “Lower-dimensional packings.” While a packing of $n$ spheres depends on the dimensionality of its embedding space, there exists a cutoff number, $m$, for which the packing of $n$ spheres remains constant for all $d+i$ dimensions, $i \geq 1$. For this $m$, the $n$ spheres have accessed all dimensions possible to them, and so the embedding space becomes irrelevant. For example, for all $d$, the unique packing of 1 and 2 spheres is the singlet and doublet, respectively. For 3 spheres, the unique packing in 1 dimension is a linear connected chain of 3 spheres, whereas in 2 dimensions it is the equilateral triangle. For $d \geq 2$, however, the unique packing of 3 spheres remains the same; it is always the triangle. For 4 spheres, the unique packing in 2 dimensions is the ditriangle whereas in 3 dimensions it is the tetrahedron. It is generally true that packings of $d+1$ particles remain the same for $d$ or more dimensions.\textsuperscript{23}

7.6. Patterns in adjacencies and distances. Does there exist a signature pattern in the $A$'s or $D$'s that signifies a packing? In other words, is there a pattern

\textsuperscript{22}If one is interested in examining all 1 bond paths that exist between all packings, then this same algorithm can be executed without this termination step to yield all possible 1 bond paths.

\textsuperscript{23}We originally posed this as a question, but one of the reviewers of this paper pointed out that this was true due to the fact that $d+1$ points are always contained in an affine subspace of at most $d$ dimensions (without loss of generality, one point can be taken to be the origin). Thus, the $d+1$ points can only access $d$ dimensions, and considering a higher than $d$-dimensional space will not change the structures that they can form.
in the distribution of adjacencies (i.e., number of contacts per particle and connections therein) and/or the distribution of distances that corresponds to packings. If such a pattern does exist, it could illuminate a more general method for solving for packings. It also might shed light on the spectrum of allowed solutions for the system of quadratic equations corresponding to the adjacency matrices—detailing which do and do not have real valued solutions in \( \mathbb{R}^3 \) satisfying \( D \geq R \).

7.7. Related mathematical problems.

7.7.1. Erdős unit distance problem. The Erdős unit distance problem (a.k.a. Erdős repeated distance problem)\(^{24}\) was posed in 1946 by the Hungarian mathematician, Paul Erdős. It asks what the maximum number of unit (or repeated) distances that can connect \( n \) points in \( d \) dimensions is [19, 9]. This problem is still unsolved. Even in 2 and 3 dimensions, only upper and lower bounds are known [14, 26]. The solution to this problem in 3 dimensions, where the unit distance is also the minimum distance, would answer what the maximum number of contacts in any sphere packing is, thus giving the number of contacts corresponding to the ground state packing(s).

7.7.2. 3-dimensional rigidity. In solving adjacency matrices for both rigid and nonrigid packings that satisfy minimal rigidity constraints in 3 dimensions, this problem is directly related to determining whether a graph is rigid in 3 dimensions. Much work has been done in this field [33, 13, 35, 12, 32, 22], as well as in other dimensions [40]. The existing work on rigidity may help to further inform sphere packings, and the work presented here may in addition be applicable to rigidity theory. In particular, it may allow for the development of a simple method for reading off whether a 3-dimensional graph is rigid or not. By the method presented here, a graph is determined to be nonrigid if there exists a continuum of solutions to \( A \). However, if we can determine a signature pattern that corresponds to all nonrigid (but minimally rigid) \( A \)'s, this would allow for a very simple determination of whether a 3-dimensional graph is rigid.

7.7.3. Solutions to systems of polynomial equations. The method presented here is inherently solving a system of quadratic equations. Thus, it presents an alternative analytical solution to this class of problems. Current standards in the field for analytically solving systems of polynomial equations include Gröbner bases [11]; however, these are time-consuming and thus do not scale efficiently with the number of equations. The method presented here solves a certain class of polynomial equations efficiently for a relatively large number of equations. Is it possible to extend this method in order to more efficiently solve large systems of polynomial equations?

7.7.4. Euclidean distance matrix completion problems. Given a symmetric matrix, \( M \), where only certain elements are specified, the Euclidean distance matrix completion problem is to find the unspecified elements of \( M \) that make \( M \) a Euclidean distance matrix. Euclidean distance matrix and positive semidefinite matrix completion problems are closely linked [36, 37, 38, 39, 31, 4]. In solving adjacency matrices for distance matrices, the method presented here is directly related and potentially directly applicable to the Euclidean distance matrix completion problem and, by extension, to the positive semidefinite matrix completion problem.

\(^{24}\)Without loss of generality, a repeated distance can be called a unit distance, because one can always uniformly rescale all distances such that the repeated distance is the unit distance. Put another way, “a unit” can be given any value—here, the unit is simply given the value of the repeated distance.
8. Concluding remarks. In this work, we present an analytical method for deriving all packings of \( n \) spheres. We carry out this derivation for \( n \leq 10 \), where the set of \( n = 10 \) new seeds is preliminary, and all iterative packings of \( n = 10 \) spheres and all packings of \( n \leq 9 \) spheres are potentially complete, save the numerical round-off error present from implementing this analytical method computationally.

We consider the derivation of these sphere packings to be the first step in directing the self-assembly of spherical colloidal particles, where we have divided this problem up into 2 parts: (i) understanding what the system of colloids can self-assemble, and (ii) deriving a mechanism to control that self-assembly. The derivation of all packings of \( n \) spheres gives us everything that a system of \( n \) colloidal particles can self-assemble, thus taking care of this first step. Future work will detail the second step, which is the derivation of a mechanism that directs the self-assembly of the packings such that only one packing forms.

Beyond the problem of self-assembly, the results reported here are interesting in their own right. We find many interesting properties from the sphere packings enumerated up to \( n = 10 \), as well as from the conjectured maximum contact packings of \( 11 \leq n \leq 20 \). Furthermore, the results are directly related to the physics of colloidal clusters and may have applications to glassy systems and the nucleation of crystals. They are also directly related to unsolved problems in mathematics, such as the Erdős unit distance problem.

Acknowledgments. We thank John Lee for consultations in coding, and for writing code that (i) converted the packing output to LaTeX format for appropriate display in Appendix B in [8], and (ii) interfaced the packing output with the rotational constant calculator website and with the mayavi2 graphing package so that the 10 particle point groups, symmetry numbers, and figures did not have to be generated manually. We also thank David Roach, Noam Elkies, and Marcus Roper for helpful discussions.

REFERENCES


DERIVING FINITE SPHERE PACKINGS


