

Polynomials for crystal frameworks and the rigid unit mode spectrum

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To each discrete translationally periodic bar-joint framework \mathcal{C} in \mathbb{R}^d we associate a matrix-valued function $\Phi_{\mathcal{C}}(z)$ defined on the d -torus. The rigid unit mode spectrum $\Omega(\mathcal{C})$ of \mathcal{C} is defined in terms of the multi-phases of phase-periodic infinitesimal flexes and is shown to correspond to the singular points of the function $z \rightarrow \text{rank } \Phi_{\mathcal{C}}(z)$ and also to the set of wave vectors of harmonic excitations which have vanishing energy in the long wavelength limit. To a crystal framework in Maxwell counting equilibrium the determinant of $\Phi_{\mathcal{C}}(z)$ is defined and gives rise to a unique multi-variable polynomial $p_{\mathcal{C}}(z_1, \dots, z_d)$. For ideal zeolites the algebraic variety of zeros of $p_{\mathcal{C}}(z)$ on the d -torus coincides with the RUM spectrum. The matrix function is related to other aspects of idealised framework rigidity and flexibility and in particular leads to an explicit formula for the number of supercell-periodic floppy modes. In the case of certain zeolite frameworks in dimensions 2 and 3 direct proofs are given to show the maximal floppy mode property (order N). In particular this is the case for the cubic symmetry sodalite framework and other idealised zeolites.

Key words: Crystal framework, rigidity operator, crystal polynomial, rigid unit mode.

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1. Introduction

Let \mathcal{C} be a mathematical crystal framework, by which we mean a connected structure in the Euclidean space \mathbb{R}^d consisting of a set \mathcal{C}_e of framework edges, representing bars or bonds, with a corresponding set \mathcal{C}_v of framework points (vertices), representing joints or atoms, such that \mathcal{C}_e is periodic with respect to a discrete translation group \mathcal{T} of isometries of \mathbb{R}^d , with \mathcal{T} of full rank. We consider mainly the case $d=2,3$ together with the locally finite assumption that \mathcal{C}_e is generated by the translations of a finite set of edges. Such a geometric bar-joint framework \mathcal{C} can serve as a model for the essential geometry of the disposition of atoms and bonds in a material crystal \mathcal{M} . In this case the vertices have atomic identifiers, such as H, He, Li, B, ... , and the chosen edges may correspond just to the strong bonds. The identification of strongly bonded molecular units, such as SiO_4 and TiO_6 , imply a polyhedral net structure for \mathcal{C} and in particular

aluminosilicate crystals and zeolites provide in this way a fascinating diversity of tetrahedral nets in which every vertex is shared by two tetrahedra.

Material scientists are interested in the manifestation and explanation of various forms of low energy oscillation and excitation modes. Of particular interest are the rigid unit modes (RUMs) in crystalline materials, the low energy (long wavelength) modes of oscillation related to the relative motions of rigid units, such as the SiO_4 tetrahedral units in quartz. The wave vectors of these modes are observed in neutron scattering experiments and have been shown to correlate closely with those for the modes observed in computer simulations with periodic networks of rigid units. In both the experimental measurements and in the simulations the background mathematical model is classical lattice dynamics and the rigid unit mode wave vectors are observed where phonon dispersion curves display markedly low energy. There is now a considerable body of literature tabulating the rigid unit mode wave vectors of various crystals and it has become evident that the primary determinant in a material \mathcal{M} is the geometric structure of an associated abstract framework \mathcal{C} . This was outlined in the seminal paper of Giddy, Dove, Pawley and Heine [7]. See also Swainson and Dove [23], Hammond et al [9], [10] and Dove et al [6], This experimental work shows that the wave vectors of RUMs often lie along lines and planes in reciprocal space. However, for many materials the wave vectors also lie on more mysterious curved surfaces. See also the recent computer assisted analysis of Wegner [24].

In the sequel we develop a mathematical theory of rigid unit modes in idealized crystal frameworks. As we shall demonstrate, this is essentially a linear first order theory and one can side-step lattice dynamical formulations that relate to higher energy phonons and their dispersion curves. In fact in Definition 8 we define the RUM spectrum $\Omega(\mathcal{C})$ of an idealised crystal framework \mathcal{C} , with given translation group, as the set of multi-phases for which there exists a nonzero phase-periodic infinitesimal flex. This form of the spectrum was first given in Owen and Power [18] as a byproduct of the analysis of square-summable infinitesimal flexes. Mapping the d -torus to the unit cube in \mathbb{R}^d by taking logarithms gives the usual wave vector parametrisation space for RUMS used by crystallographers. The spectrum $\Omega(\mathcal{C})$ leads naturally to a definition of the RUM dimension $\dim_{\text{rum}} \mathcal{C}$, which takes integer values from 0 to d and which gives a measure of the infinitesimal flexibility of \mathcal{C} . In the interesting special case of a crystal framework in Maxwell counting equilibrium, such as, for example, a tetrahedral net framework derived from an idealised zeolite, the spectrum $\Omega(\mathcal{C})$ is determined as the zero set of a multi-variable polynomial $p_{\mathcal{C}}(z_1, \dots, z_d)$ defined on the d -torus. This polynomial may vanish identically, which corresponds to the case $\dim_{\text{rum}}(\mathcal{C}) = d$, and for $d = 2, 3$ this is also known as "order N". (See Theorem 2.) This property occurs for example in the case of the cubic form sodalite framework \mathcal{C}_{SOD} , as we prove below in Section 6 by infinitesimal analysis. Interestingly, Kapko et al [13] have recently conducted a simulation analysis to determine the extent of this property in idealized zeolites.

The infinitesimal flex perspective is useful for several reasons. Firstly it brings into play the fairly well-established theory of infinitesimal rigidity for finite bar-joint frameworks and this is of significance for local flexibility. On the other hand the consideration of general infinitesimal flexes in infinite bar-joint frameworks gives a route to understanding and predicting the appearance of linear components (lines, planes, hyperplanes etc) observed experimentally in RUM wave vector sets. In addition, the first order infinitesimal flexibility properties of a crystal framework

\mathcal{C} are implicit in the infinite rigidity matrix $R(\mathcal{C})$ of \mathcal{C} , and for phase-periodic flexibility this simplifies to the consideration of a finite function matrix $\Phi_{\mathcal{C}}(z)$ defined on the d -torus. This matrix function, which we also refer to as the symbol function of \mathcal{C} (borrowing terminology from Hilbert space operator theory) also arises naturally from square-summable flex perspectives and may be a useful tool more generally, as we speculate below. When the matrix is square the *crystal polynomial* $p_{\mathcal{C}}(z)$ of \mathcal{C} is defined as a natural normalisation of its determinant.

In the development we give definitions, theorems, proofs and illustrative examples all of which lie within a mathematical theory of infinite bar-joint frameworks. While the focus is on rigidity and flexibility properties related to the disposition of the bonds, rather than their strengths, the theory has the potential to inform applied analysis and simulations.

In Sections 2 and 3 we give examples of crystal frameworks and various spaces of infinitesimal flexes. In Sections 4, 5 and 6 we define the matrix function $\Phi_{\mathcal{C}}(z)$, the RUM spectrum $\Omega(\mathcal{C})$, the polynomial $p_{\mathcal{C}}(z)$ and give connections with phase-periodic flexes, floppy modes and low energy harmonic excitations. In the final section we give determinations of $\Omega(\mathcal{C})$ in a range of examples. In particular we give a two-dimensional zeolite whose floppy modes are of order N .

2. Crystal frameworks: terminology and examples.

Let $G = (V, E)$ be a simple graph, finite or countable, with vertices $V = \{v_1, v_2, \dots\}$, and $E \subseteq V \times V$ a countable set of edges, and let p_1, p_2, \dots be a sequence of points in the Euclidean space \mathbb{R}^d , with $p_i \neq p_j$ if (v_i, v_j) is an edge. Then the pair (G, p) , with $p = (p_1, p_2, \dots)$ is said to be a *bar-joint framework* in \mathbb{R}^d with framework points, or joints, p_i and framework edges, or bars, given by the line segments $[p_i, p_j]$ between p_i and p_j when (v_i, v_j) is an edge in E . In all our examples in fact, the framework points are distinct.

An *isometry* of \mathbb{R}^3 is a distance-preserving map $T: \mathbb{R}^3 \rightarrow \mathbb{R}^3$. A *full rank translation group* \mathcal{T} is a set of translation isometries $\{T_k: k \in \mathbb{Z}^3\}$ with $T_{k+l} = T_k + T_l$ for all k, l , $T_k \neq I$ if $k \neq 0$, and such that the three *period vectors*

$$a = T_{\gamma_1}0, \quad b = T_{\gamma_2}0, \quad c = T_{\gamma_3}0,$$

associated with the generators $\gamma_1 = (1, 0, 0)$, $\gamma_2 = (0, 1, 0)$, $\gamma_3 = (0, 0, 1)$ of \mathbb{Z}^3 are not coplanar. Full rank translation groups in \mathbb{R}^d are similarly defined.

The following definition follows the formalism of Owen and Power [18] which emphasises the periodic partitioning of the points and edges of \mathcal{C} by translates of a geometrical *motif*, this being a pair consisting of a finite set of framework vertices together with a finite set of framework edges.

DEFINITION 1. A *crystal framework* $\mathcal{C} = (F_v, F_e, \mathcal{T})$ in \mathbb{R}^d , with full rank translation group $\mathcal{T} = \{T_k: k \in \mathbb{Z}^d\}$ and motif (F_v, F_e) , is a countable bar-joint framework with framework points $p_{\kappa, k}$, for $1 \leq \kappa \leq t, k \in \mathbb{Z}^d$, such that

(i) F_v is a finite set of framework vertices, $\{p_{\kappa, 0}: 1 \leq \kappa \leq t\}$ in \mathbb{R}^d , and F_e is a finite set of framework edges,

(ii) for each κ and k the point $p_{\kappa, k}$ is the translate $T_k p_{\kappa, 0}$,

- (iii) the set \mathcal{C}_v of framework points is the union of the disjoint sets $T_k(F_v)$ for $k \in \mathbb{Z}^d$,
- (iv) the set \mathcal{C}_e of framework edges is the union of the disjoint sets $T_k(F_e)$ for $k \in \mathbb{Z}^d$.

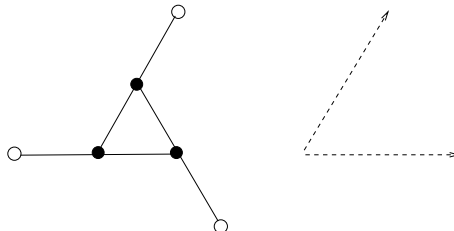


Figure 1. Motif and period vectors for the kagome framework, \mathcal{C}_{kag} .

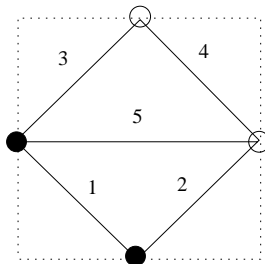


Figure 2. A five-edged motif for the squares framework, \mathcal{C}_{sq} .

In Figures 1 and 2 motif choices are shown for the kagome framework \mathcal{C}_{kag} and the squares framework \mathcal{C}_{sq} , where the filled vertices indicate the points of F_v and where the translation group is determined by the period vectors. Thus \mathcal{C}_{kag} is the well-known framework of pairwise corner-connected congruent equilateral triangles in regular hexagonal arrangement, while \mathcal{C}_{sq} is a translationally periodic framework of corner-connected rigid square units.

One can similarly identify motifs for other well-known basic frameworks, such as (i) the grid framework $\mathcal{C}_{\mathbb{Z}^d}$ in \mathbb{R}^d with $\mathcal{C}_v = \mathbb{Z}^d$ and \mathcal{C}_e equal to the set of line segments between nearest neighbours, and (ii) \mathcal{C}_{tri} and (iii) \mathcal{C}_{hex} , by which we denote the two crystal frameworks in the plane associated with regular triangle and hexagon tilings respectively.

In the examples we employ a mnemonic notational convenience with, typically, \mathcal{C}_{xyz} with all lower case letters indicating a planar framework, \mathcal{C}_{Xyx} indicating a 3D framework, and \mathcal{C}_{XYZ} indicating a 3D framework which derives in a well-defined way from the zeolite with name XYZ. For example we write \mathcal{C}_{Oct} to denote the basic regular octahedron net framework in three dimensions formed by corner-connected congruent octahedra with maximal cubic symmetry. Also \mathcal{C}_{SOD} , defined

below, derives from the cubic form of the zeolite sodalite, while the companion framework C_{RWY} derives from the sodalite RWY. These conventions are useful, for example, when discussing subframeworks lying in vector subspaces (slices).

The following definition is convenient.

DEFINITION 2. *A crystal framework \mathcal{C} in \mathbb{R}^d is said to be in Maxwell counting equilibrium if $d|F_v| = |F_e|$ for some, and hence every, motif. If $d|F_v| < |F_e|$ then \mathcal{C} is said to be edge rich while if $d|F_v| > |F_e|$ then \mathcal{C} is said to be edge sparse.*

We now define a number of illustrative crystal frameworks in dimensions 2 and 3 and in Section 6 we compute their RUM spectra. Of particular interest with regard to rigidity and flexibility are the 4-regular (4-coordinated) frameworks in 2D and the 6-regular frameworks in 3D, examples of which are provided by idealized zeolites in the sense of Definition 3.

Graphene and diamond bar-joint frameworks, $\mathcal{C}_{\text{gra}}, \mathcal{C}_{\text{Dia}}, \mathcal{C}_{\text{Dia}}^2$. The usual visualisation of graphene is as a two-dimensional hexagonal bond-node network of carbon atoms with the geometry of \mathcal{C}_{hex} . However \mathcal{C}_{hex} is edge sparse and this image is not suggestive of the strength of the material. If we view the $C-C-C$ angles as rigid, or, equivalently, if we also view *second* nearest neighbours as bonded, then this leads to the edge rich crystal framework, $\mathcal{C}_{\text{hex}}^2$ say, implied by Figure 3. In the motif shown we may take two of the edges of one of the equilateral triangles to determine period vectors and a corresponding translation group \mathcal{T} .

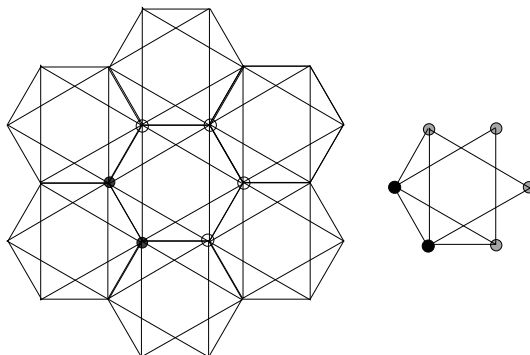


Figure 3. Part of $\mathcal{C}_{\text{gra}} = \mathcal{C}_{\text{hex}}^2$ and a choice of motif.

This crystal framework is of interest in its own right and we also write it as \mathcal{C}_{gra} when viewed as a bar-joint framework in \mathbb{R}^2 . It may be assembled or decomposed in a number of ways to reveal substructure and in particular it may be constructed as a fusion of two congruent crystal subframeworks as follows. Let $\mathcal{C}_{\text{tri}}^+$ be obtained from the triangular framework \mathcal{C}_{tri} by adding bars, in which alternate triangles have three extra short bars added, connecting the triangle joints to a new joint at the centroid of the triangle. Note that these added centroid joints are in natural one to one correspondence with the joints of \mathcal{C}_{tri} by a small translation. Then

\mathcal{C}_{gra} is congruent to the framework formed from the join of two copies of $\mathcal{C}_{\text{tri}}^+$, one of which is rotated by π , and where the copies are connected by identifying the centroids of one copy with the non-centroid joints of the other, together with identification of the resulting double edges.

Similarly, crystalline diamond is usually indicated pictorially by a face-centred unit cell, with 14 C atoms at face centres and corners, plus 4 internal C atoms, and nearest neighbour connectivity. Again, the implied 4-coordinated edge sparse bar-joint framework, \mathcal{C}_{Dia} say, does not of itself impart a sense of rigidity. It is natural for us to consider, once again, the derived first-and-second-nearest neighbour framework, and to take this as the definition of an associated bar-joint framework, which we denote $\mathcal{C}_{\text{Dia}}^2$. This too may be understood, or defined, in various constructive ways. For one such construction, echoing the graphene framework decomposition, note that there is a bipartite red-blue colouring of the nodes of \mathcal{C}_{Dia} with face atoms red and internal atoms blue say. The extra edges of $\mathcal{C}_{\text{Dia}}^2$ are either blue-blue or red-red. The red-red determined subframework we refer to as the tetrahedron framework \mathcal{C}_{Tet} . Adding to this framework the blue-red edges of \mathcal{C}_{Dia} gives a framework we call $\mathcal{C}_{\text{Tet}}^+$ (created by centroid addition). It follows that $\mathcal{C}_{\text{Dia}}^2$ is a join of two copies of $\mathcal{C}_{\text{Tet}}^+$ (with reflected orientation), the join being effected by centroid/noncentroid identification, as before.

The cubic sodalite framework \mathcal{C}_{SOD} . The crystal framework \mathcal{C}_{SOD} in three dimensions is built from 4-rings of tetrahedra in a way which echoes the crystal structure of the cubic form of the zeolite sodalite.

The following general definition is convenient.

DEFINITION 3. *An ideal (or mathematical) zeolite in two (resp. three) dimensions is a crystal framework \mathcal{C} in the plane (resp. \mathbb{R}^3) consisting of congruent triangles (resp. congruent tetrahedra), each pair of which intersect disjointly or at a common vertex, or edge, (or face,) and is such that every vertex is shared by two triangles (resp. tetrahedra).*

We remark that in databases material zeolite frameworks are most frequently indicated as a network of "T atoms" corresponding to tetrahedral centres, each of which is 4-coordinated with neighbouring T atoms. This contrasts with the rigid unit view here of a tetrahedral net framework implied by the positions of O atoms as vertices.

The 4-ring building units of \mathcal{C}_{SOD} are oriented in the high symmetry arrangement indicated in Figure 4. Six such rings may be placed on (the outside of the) six faces of an imaginary cube so that the contact vertices sit on the midpoints of the edges of the cube. This gives a finite bar-joint framework consisting of six regular 4-rings connected together to form a finite bar-joint framework which we call the sodalite cage framework. With unit edge length for the tetrahedra the cube has sidelength $1 + \sqrt{2}$, while the three orthogonal period vectors (determining unit cell geometry) have length $2 + \sqrt{2}$.

A motif for the framework can be given using the set F_e of edges in three pairwise-connected pairwise orthogonally oriented 4-rings of the sodalite cage. The images of the edges of F_e under the action of the associated isometry group \mathcal{T} are essentially disjoint and generate the crystal framework \mathcal{C}_{SOD} . For an appropriate

set F_v an examination of the positioning of F_v in the sodalite cage shows that one must take the vertices of F_e except for 9 redundant exterior vertices.

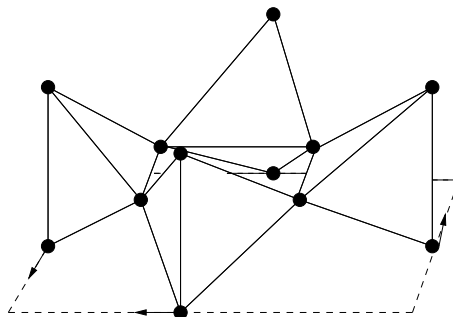


Figure 4. The top 4-ring of the sodalite cage.

The kagome net framework $\mathcal{C}_{\text{Knet}}$. We give two specifications of the *kagome net* framework in three dimensions. Firstly, it may be constructed in a layered manner. Form upward tetrahedral rigid unit frameworks on alternate triangles of a two-dimensional kagome framework lying in the xy -plane. Similarly, form downward tetrahedra on the other triangles and thereby create a layer framework of pairwise connected tetrahedra. Parallel copies of such layers can be joined at their exposed joints together to fill space, creating, unambiguously, the crystal framework we denote as $\mathcal{C}_{\text{Knet}}$.

Alternatively, $\mathcal{C}_{\text{Knet}}$ is a translationally periodic bar-joint framework with period vectors formed by three edges of a regular parallelapiped, with pairwise angles of $\pi/3$. Each parallelapiped contains two tetrahedral rigid units located at opposite "acute" corners of the parallelapiped and with edge length half that of the parallelapiped edges. The planar slices of $\mathcal{C}_{\text{Knet}}$, determined by each pair of period vectors, give copies of \mathcal{C}_{kag} .

The frameworks $\mathcal{C}_{\text{star}}$ and \mathcal{C}_{oct} . The kagome framework can be viewed as arising from the connection of translates of a regular 6-pointed star. There are analogous frameworks, $\mathcal{C}_{\text{star}}$ and \mathcal{C}_{oct} , arising from similar tilings using a regular 4-pointed star and an 8-pointed star respectively. Figure 6, in the final section, indicates the (primitive) star template for $\mathcal{C}_{\text{star}}$, while Figure 5 indicates tiling templates for four 2D zeolite crystal frameworks with implied translation groups. The third framework (with exterior angle $8\pi/12$) is a copy of $\mathcal{C}_{\text{star}}$, with a modified orientation and translation group, while the second and fourth examples give \mathcal{C}_{oct} in different orientations, with choices of translation group.

A motif (F_v, F_e) for \mathcal{C}_{oct} may be provided with F_v the set of four boundary vertices (indicated as solid vertices in the fourth template) plus the eight internal vertices (of the octagon), and with F_e the set of all 24 edges of the template. Evidently, there is a smooth periodic edge-length-preserving deformation linking these frameworks which is parametrised by the exterior angle. This may be viewed

as an example of the idealisation of displacive phase transitions in materials and in Section 6 we see how the RUM spectrum evolves in this example.

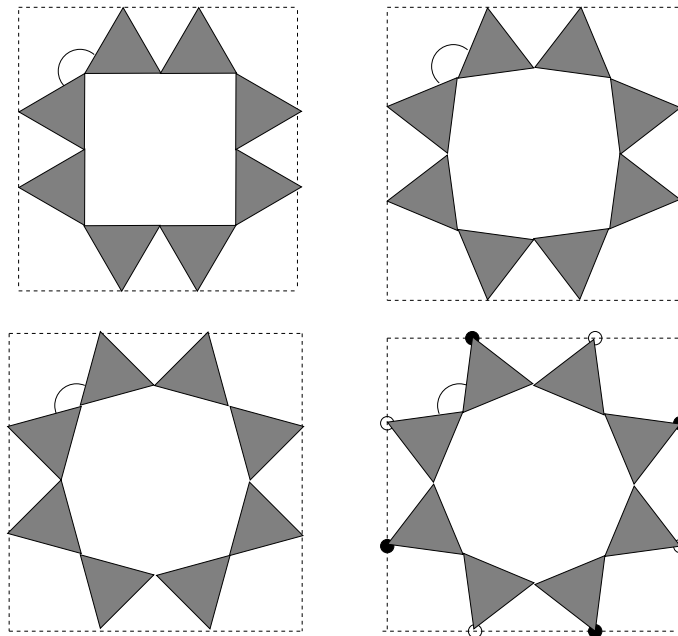


Figure 5. Templates for 2D zeolite frameworks, with exterior angles $10\pi/12$, $9\pi/12$, $8\pi/12$, $7\pi/12$.

Further examples. Simple but informative examples of 3D zeolite frameworks can be built from 2D zeolite frameworks in various ways by layer constructions. With \mathcal{C}_{oct} for example, embedded in the x, y plane of \mathbb{R}^3 , we may add bars and joints to obtain alternately upward and downward pointing tetrahedral units and so create a layer framework. These layers may be joined consecutively at their exposed points to fill \mathbb{R}^3 and thereby so create an associated ideal zeolite framework $\tilde{\mathcal{C}}_{\text{oct}}$. Similarly one can view $\mathcal{C}_{\text{Knet}}$ as the framework $\tilde{\mathcal{C}}_{\text{kag}}$.

We also note that interesting and diverse examples of mathematical crystal frameworks are implied by various tilings and periodic nets. For an account of three-periodic nets and connections with crystal chemistry see Delgado Friedrichs, O’Keeffe and Yaghi [3], [4]. Such an (unlabelled) net, in any dimension, may be defined as a pair (N, P) , where N the union of the edges of a crystal framework whose pairs of edges only intersect at framework vertex points, and where P is the subset of these points.

3. Infinitesimal flexibility and rigidity.

We now define various flexes which act on the entire infinite crystal framework in a locally infinitesimal manner. The definition is the same as that for a finite bar-joint framework.

DEFINITION 4. *An infinitesimal flex of a finite or countable bar-joint framework (G, p) is a vector $u = (u_i)$, with each component u_i a vector in \mathbb{R}^d , such that for each edge $[p_i, p_j]$*

$$\langle p_i - p_j, u_i \rangle = \langle p_i - p_j, u_j \rangle.$$

Regarding the u_i as velocity vectors this asserts that for each edge the components of the endpoint velocities in the edge direction are in agreement. This is equivalent to the assertion that an infinitesimal flex is a velocity vector $v = (v_i)$ for which the distance deviation

$$|p_i - p_j| - |(p_i + tu_i) - (p_j + tu_j)|$$

of each edge is of order t^2 as the time parameter t tends to zero.

We will not be concerned particularly with *continuous flexes*, which are also called finite flexes or finite (edge-length-preserving) deformations. For such flexes each framework point undergoes a continuous motion $p_{\kappa,k}(t)$ such that edge lengths are preserved for all values of time t in some range. However we note that, as for a finite framework, the derivative $u = p'(0) = (p'_i(0))$ of a differentiable continuous flexes $p(t) = (p_i(t))$ provides an infinitesimal flex u .

In the case of a crystal framework in \mathbb{R}^d a velocity vector is doubly-indexed sequence v of vectors $v_{\kappa,k}$ in \mathbb{R}^d regarded as instantaneous velocities applied to the frameworks vertices $p_{\kappa,k}$, and it is convenient to consider the vector space of all velocity sequences, written as a direct product, namely

$$\mathcal{H}_{atom} = \prod_{\kappa,k} \mathbb{R}^d.$$

Thus, a real infinitesimal flex u for the crystal framework \mathcal{C} is a velocity vector u in \mathcal{H}_{atom} such that

$$\langle u_{\kappa,k} - u_{\tau,l}, p_{\kappa,k} - p_{\tau,l} \rangle = 0$$

for each framework edge $[p_{\kappa,k}, p_{\tau,l}]$. In particular the set of all infinitesimal flexes forms a vector subspace, \mathcal{H}_{fl} say, of \mathcal{H}_{atom} . Also each nontrivial isometry of \mathbb{R}^d gives rise to a one-dimensional vector subspace of \mathcal{H}_{fl} .

The *rigidity matrix* $R(\mathcal{C})$ of \mathcal{C} is a real infinite matrix defined as in the finite framework case.

DEFINITION 5. *The rigidity matrix $R(\mathcal{C})$ of the crystal framework \mathcal{C} in \mathbb{R}^3 has rows labelled by the edges $e = [p_{\kappa,k}, p_{\tau,l}]$ and columns labelled by the framework point coordinate indices (κ, x, k) , (κ, y, k) , (κ, z, k) . The row for edge e takes the form*

$$[\cdots 0 \quad (p_{\kappa,k} - p_{\tau,l}) \quad 0 \cdots 0 \quad (p_{\tau,l} - p_{\kappa,k}) \quad 0 \cdots]$$

where the entry $(p_{\kappa,k} - p_{\tau,l})$ indicates that the three coordinates of this vector lie in the columns for (κ, x, k) , (κ, y, k) , (κ, z, k) .

The definition of $R(\mathcal{C})$ for $d = 2, 4, 5, \dots$, and also for general countably infinite bar-joint frameworks, is essentially the same. We remark that one may take the

view that $R(\mathcal{C})$ is $1/2J(\mathcal{C})$ where $J(\mathcal{C})$ is the generalised Jacobian, evaluated at the $p_{\kappa,k}$, for the infinite quadratic equation system

$$|q_{\kappa,k} - q_{\tau,l}|^2 = d_e^2,$$

where the equations, labelled by the edges, are in the coordinate variables of the points $q_{\kappa,k}$, and where the constants d_e are the given lengths of the edges e of \mathcal{C} .

It is natural to consider various linear transformations that derive from $R(\mathcal{C})$. To this end let

$$\mathcal{H}_{bond} = \prod_{e \in \mathcal{C}_e} \mathbb{R} = \prod_{e \in F_e, k \in \mathbb{Z}^d} \mathbb{R}$$

be the space of real sequences $w = (w_{e,k})_{e \in F_e, k \in \mathbb{Z}^d}$ labelled by the framework edges. Then $R(\mathcal{C})$ gives a linear transformation $R: \mathcal{H}_{atom} \rightarrow \mathcal{H}_{bond}$. Indeed, each row of R has at most $2d$ nonzero entries and the image $R(u)$ is given by the well-defined matrix multiplication $R(\mathcal{C})u$. As for finite frameworks one has the following elementary proposition.

PROPOSITION 1. *The infinitesimal flexes of the crystal framework \mathcal{C} are the velocity vectors in \mathcal{H}_{atom} that lie in the nullspace of the linear transformation $R(\mathcal{C})$.*

Let us introduce notation for the natural "basic sequences" of \mathcal{H}_{atom} and \mathcal{H}_{bond} . Write ξ_x, ξ_y, ξ_z for the standard coordinate basis of \mathbb{R}^3 , $\xi_x = (1, 0, 0)$ etc., and for $\sigma \in \{x, y, z\}$ write $\xi_{\kappa,\sigma,k}$ for the position indicator vector in \mathcal{H}_{atom} with

$$(\xi_{\kappa,\sigma,k})_{\kappa',k'} = \delta_{\kappa,\kappa'} \delta_{k,k'} \xi_\sigma,$$

where $\delta_{\kappa,\kappa'}$ is the Kronecker delta. While \mathcal{H}_{atom} does not have countable vector space dimension its subspace of finitely nonzero sequences has the set $\{\xi_{\kappa,\sigma,k}\}$ as a vector space basis. However, the set is a generalised *product type basis* for \mathcal{H}_{atom} in the sense of the next definition. In particular we may define the infinitesimal unit translation flex u_x in the x direction as the well-defined infinite sum

$$u_x = \sum_{\kappa,k} \xi_{\kappa,x,k}.$$

Similarly we may write $\eta_{e,k}$ for the basic sequence in \mathcal{H}_{bond} which is zero but for the value 1 for the coordinate position e, k .

DEFINITION 6. *Let (G, p) be a countably infinite bar-joint framework in \mathbb{R}^d . A product type basis for a subspace \mathcal{M} of the velocity space \mathcal{H}_{atom} of (G, p) is a countable set $\mathcal{S} = \{w^1, w^2, \dots\}$ of vectors in \mathcal{M} such that,*

(i) *every vector u in \mathcal{M} has a unique representation*

$$u = \sum_{n \in \mathbb{N}} \alpha_n w^n, \quad \alpha_n \in \mathbb{R},$$

(ii) *for each index k only a finitely many elements w^n of \mathcal{S} have nonzero k th component w_k^n .*

(In view of (ii) an arbitrary infinite sum of the form in (i) is well-defined.) The basic grid framework $\mathcal{C}_{\mathbb{Z}^d}$ has evident nonzero infinitesimal flexes u that act only on linear subframeworks (copies of $\mathcal{C}_{\mathbb{Z}}$ in \mathbb{R}^2). One can show that a set, \mathcal{S}_d of representatives of all such flexes, is a product type basis for \mathcal{H}_{\square} . In fact, it is

possible to identify product type bases for the vector space of all infinitesimal flexes for many other basic crystal frameworks (Power [20]). Two examples are \mathcal{C}_{kag} and the 3D crystal framework \mathcal{C}_{Oct} for example.

Some natural special classes of infinitesimal flexes of a crystal framework \mathcal{C} with translation group \mathcal{T} which will concern us are the following:

- (i) *strictly periodic infinitesimal flexes* u , with the periodicity condition $u_{\kappa,k} = u_{\kappa,0}$ for all $k \in \mathbb{Z}^d$,
- (ii) *supercell-periodic flexes* u , with the periodicity condition $u_{\kappa,k} = u_{\kappa,0}$, for all k in a subgroup $r_1\mathbb{Z} \times \cdots \times r_d\mathbb{Z}$ for some positive integers r_1, \dots, r_d ,
- (iii) *local infinitesimal flexes* u , for which $u_{\kappa,k} = 0$ for all but finitely many values of κ, k .

Note the elementary fact that if there exists a local flex for \mathcal{C} then this framework is rich in supercell-periodic flexes. Indeed if u is such a local infinitesimal flex and if $k \rightarrow \alpha_k$ is any supercell-periodic coefficient sequence then the sum

$$w = \sum_k \alpha_k T_k u,$$

is well-defined supercell-periodic infinitesimal flex.

In Section 5 we turn attention to complex scalar infinitesimal flexes which are phase-periodic, the real and imaginary parts of which provide real infinitesimal flexes. It is such phase-periodic flexes that are closely allied to rigid unit mode wave vectors. They lead naturally to the formulation of a matrix-valued function associated with \mathcal{C} and \mathcal{T} and we describe this association in the next section.

We remark that there is an important class of infinitesimal flexes that we do not consider here which do not feature in fixed lattice rigid unit mode analysis. These are the affinely periodic infinitesimal flexes which allow, roughly speaking, an infinitesimal adjustment of period vectors. Such adjustment may reflect infinitesimal lattice contraction for example. For more on such flexibility and associated finite motions see, for example, Borcea and Streinu [1], Malestein and Theran [15], Owen and Power [18], Power[19] and Ross et al [21].

If a connected bar-joint framework (G, p) , finite or infinite, has no infinitesimal flexes other than rigid motion flexes then it is said to be *infinitesimally rigid*. Perhaps the simplest way in which this occurs is when (G, p) is *sequentially infinitesimally rigid* (Owen and Power [18]) in the sense that it is the union of an increasing sequence of infinitesimally rigid finite frameworks. This is evidently the case for the edge rich frameworks \mathcal{C}_{tri} , \mathcal{C}_{gra} , \mathcal{C}_{Tet} and $\mathcal{C}_{\text{Dia}}^2$. In particular it follows from the definitions below that the RUM spectra of these frameworks are trivial. On the other hand overconstrained frameworks such as these are rich in infinitesimal self-stresses in the following sense.

Following terminology for finite frameworks we may define the notion of a *self-stress* $w = (w_e)_{e \in \mathcal{C}_e}$ for a crystal framework \mathcal{C} . This is an assignment of scalars to edges such that for every framework point $p_{\kappa,0}$ in F_v (and hence for every framework point) the finite vector sum

$$\sum_{\tau, l: e=[p_{\kappa,0}, p_{\tau,l}] \in \mathcal{C}_e} w_e (p_{\kappa,0} - p_{\tau,l}),$$

taken over all edges incident to $p_{\kappa,0}$, is equal to zero. This is a companion notion to that of an infinitesimal flex and indeed w is a self-stress if and only if w lies in the nullspace of the transpose matrix $R(C)^T$.

4. The matrix function $\Phi_{\mathcal{C}}(z)$.

A matrix-valued function, or *symbol function*, for \mathcal{C} is associated with a choice of periodicity group \mathcal{T} and a given motif (F_v, F_e) as follows.

Write $z = (z_1, \dots, z_d)$, with $z_i \in \mathbb{C}$, $|z_i| = 1$, to denote general points in the d -torus. Also, write z^k for the monomial function $z \rightarrow z^k$ from \mathbb{T}^d to \mathbb{C} . Since $z_i^{-k} = \bar{z}_i^k$ for points on the circle \mathbb{T} we may think of general monomials z^δ as products of the z_i or \bar{z}_i with just non-negative powers.

It is convenient to define the edge vector v_e of the directed edge $e = [p_{\kappa,k}, p_{\tau,l}]$ as $v_e = p_{\kappa,k} - p_{\tau,l}$ and to write $v_{e,\sigma}$, for $1 \leq \sigma \leq d$, for the coordinates of v_e .

DEFINITION 7. Let \mathcal{C} be a crystal framework in \mathbb{R}^d with motif sets

$$F_v = \{p_{\kappa,0} : 1 \leq \kappa \leq |F_v|\}, \quad F_e = \{e_i : 1 \leq i \leq |F_e|\}.$$

Then $\Phi_{\mathcal{C}}(z)$ is the matrix-valued function on \mathbb{T}^d with rows labelled by the edges $e = [p_{\kappa,k}, p_{\tau,l}] \in F_e$ and with columns labelled by pairs κ, σ . As a matrix of scalar function the entries are given by

$$(\Phi_{\mathcal{C}}(z))_{e,(\kappa,\sigma)} = v_{e,\sigma} \bar{z}^k,$$

$$(\Phi_{\mathcal{C}}(z))_{e,(\tau,\sigma)} = -v_{e,\sigma} \bar{z}^l,$$

if $\kappa \neq \tau$, while for a reflexive edge, with $\kappa = \tau$,

$$(\Phi_{\mathcal{C}}(z))_{e,(\kappa,\sigma)} = v_{e,\sigma} (\bar{z}^k - \bar{z}^l).$$

The other entries are equal to the zero function.

Different motifs for $(\mathcal{C}, \mathcal{T})$ give matrix functions that are equivalent in a natural way. Indeed, replacement of a motif edge (resp. vertex) by a \mathcal{T} -equivalent one results in the multiplication of the appropriate row (resp. columns) by a monomial. Thus in general two motif matrix functions $\Phi(z)$ and $\Psi(z)$ satisfy the equation

$$\Psi(z) = D_1(z)A\Phi(z)BD_2(z),$$

where $D_1(z), D_2(z)$ are diagonal matrix functions with monomial functions on the diagonal and where A, B are permutation matrices, associated with edge and vertex relabelling.

The next two examples and those we consider later occur in two and three dimension and in this case we simply write (z, w) and (z, w, u) respectively for general points of \mathbb{T}^2 and \mathbb{T}^3 .

Example (a). The motif for \mathcal{C}_{sq} implied by Figure 2 has F_v equal to the ordered set $\{(1/2, 0), (0, 1/2)\}$ and $F_e = \{e_1, \dots, e_5\}$. Here the period vectors, given by the sides of the parallelogram unit cell, are scaled with unit length. It follows that the

matrix function for \mathcal{C}_{sq} is

$$\frac{1}{2} \begin{bmatrix} 1 & -1 & -1 & 1 \\ -1 & -1 & \bar{z} & \bar{z} \\ \bar{w} & \bar{w} & -1 & -1 \\ -\bar{w} & \bar{w} & \bar{z} & -\bar{z} \\ 0 & 0 & -2 + 2\bar{z} & 0 \end{bmatrix}.$$

If the final row of $\Phi_{\mathcal{C}_{\text{sq}}}(z, w)$ is deleted then one has the matrix function for the realisation of the square grid framework, $\mathcal{C}_{\mathbb{Z}^2}$, when rotated by $\pi/4$. This framework is in Maxwell counting equilibrium and so the matrix function is square and we may compute

$$\det \Phi_{\mathcal{C}_{\mathbb{Z}^2}}(z, w) = 4 (\bar{w} - \bar{z}) (\bar{w}\bar{z} - 1).$$

Example (b). With a choice of labeling for the motif in Figure 1, with period vectors of length one, the matrix function $\Phi_{\text{kag}}(z, w)$ of the kagome framework \mathcal{C}_{kag} takes the form given by

$$\Phi_{\text{kag}}(z, w) = \frac{1}{4} \begin{bmatrix} -2 & 0 & 2 & 0 & 0 & 0 \\ 0 & 0 & 1 & -\sqrt{3} & -1 & \sqrt{3} \\ -1 & -\sqrt{3} & 0 & 0 & 1 & \sqrt{3} \\ 2 & 0 & -2z & 0 & 0 & 0 \\ 0 & 0 & -1 & \sqrt{3} & \bar{z}\bar{w} & -\sqrt{3}\bar{z}w \\ \bar{w} & \sqrt{3}\bar{w} & 0 & 0 & -1 & -\sqrt{3} \end{bmatrix}.$$

In this case the determinant is equal to a constant multiple of

$$\bar{z}\bar{w}(z-1)(w-1)(z-w).$$

For a different motif for the given translation group this determinant would change by a monomial factor.

Polynomials for crystal frameworks. Let \mathcal{C} be a crystal framework in \mathbb{R}^d with a given isometry group \mathcal{T} . If \mathcal{C} is in Maxwell counting equilibrium then we may form the polynomial $\det(\Phi_{\mathcal{C}}(z))$ of the matrix function associated with a particular motif. This is a polynomial in the coordinate functions z_i and their complex conjugates \bar{z}_i , and is possibly identically zero. In the nonzero case we remove dependence on the motif and formally define the *crystal polynomial* $p_{\mathcal{C}}(z_1, \dots, z_d)$, associated with the pair \mathcal{C}, \mathcal{T} and a lexicographic monomial ordering, as the product $\alpha z^\gamma \det(\Phi_{\mathcal{C}}(z))$ where the multi-power γ and the scalar α are chosen so that

- (i) $p_{\mathcal{C}}(z)$ is a linear combination of nonnegative power monomials,

$$p_{\mathcal{C}}(z) = \sum_{\alpha \in \mathbb{Z}_+^d} a_\alpha z^\alpha,$$

- (ii) $p_{\mathcal{C}}(z)$ has minimum total degree, and
 (iii) $p_{\mathcal{C}}(z)$ has leading monomial with coefficient 1.

It is natural to order monomials lexicographically, so that, for example, the monomial function $z_1^2 z_2$ has higher multi-degree than $z_1 z_2^3$. In this way one defines the leading term of a multivariable polynomial. (See also the discussion in Cox, Little and O'Shea [2] for example.)

It follows that the crystal polynomial for the kagome framework and the (primitive case) translation group, as above, is

$$p_{kag}(z, w) = (z - 1)(w - 1)(z - w),$$

with lexicographic order $z > w$. Also, for the grid framework $\mathcal{C}_{\mathbb{Z}^2}$ and the non-axial translation group given above we see from the form of the determinant that

$$p_{\mathbb{Z}^2}(z, w) = (z - w)(zw - 1).$$

For the grid frameworks $\mathcal{C}_{\mathbb{Z}^d}$ it is in fact more natural to take the standard axial translation group \mathcal{T} and a minimal motif which consists of a single vertex and d edges, one for each axial direction. For this pair \mathcal{C}, \mathcal{T} the crystal polynomial is simply

$$(z_1 - 1)(z_2 - 1) \dots (z_d - 1).$$

5. Rigid Unit Modes and $\Phi_{\mathcal{C}}(z)$.

We first show how $\Phi_{\mathcal{C}}(z)$ arises as a family of matrices parametrised by points z in the d -torus where the matrix for $z = \omega$ determines the possible existence of infinitesimal flexes which are periodic modulo the multi-phase $\bar{\omega}$.

Let $\mathcal{K}_{atom}, \mathcal{K}_{bond}$ be the complex scalar versions of the vector spaces $\mathcal{H}_{atom}, \mathcal{H}_{bond}$. Write \mathcal{K}_a^ω for the complex vector subspace space of complex velocity vectors $v = (v_{\kappa, k})$ such that $v_{\kappa, k} = \omega^k v_{\kappa, 0}$ for $\kappa \in F_v, k \in \mathbb{Z}^d$. This is a finite-dimensional subspace of \mathcal{K}_{atom} of dimension $d|F_v|$.

Similarly let $\mathcal{K}_b^\omega \subset \mathcal{K}_{bond}$ be the subspace of the bond vector space of complex sequences $w = (w_e)_{e \in \mathcal{C}_e}$ which are phase-periodic in this way for the phase ω . Note that the rigidity matrix $R(\mathcal{C})$ provides a linear transformation $R^\omega : \mathcal{K}_a^\omega \rightarrow \mathcal{K}_b^\omega$ by restriction. Indeed, with $d = 3$, let $\gamma_i, 1 \leq i \leq 3$, denote the usual generators for \mathbb{Z}^3 and let W_i and U_i be the shift transformations on \mathcal{K}_{atom} and \mathcal{K}_{bond} respectively, with

$$\begin{aligned} W_i : \xi_{\kappa, \sigma, k} &\rightarrow \xi_{\kappa, \sigma, k + \gamma_i}, \\ U_i : \eta_{e, k} &\rightarrow \eta_{e, k + \gamma_i}. \end{aligned}$$

Then we have the commutation relations

$$R(\mathcal{C})W_i = U_i R(\mathcal{C}), \quad 1 \leq i \leq 3,$$

and the identities $W_i u = \bar{\omega}_i u$, for $u \in \mathcal{K}_a^\omega$, and $U_i v = \bar{\omega}_i v$, for $v \in \mathcal{K}_b^\omega$. Thus for u in \mathcal{K}_a^ω ,

$$U_i(R(\mathcal{C})u) = R(\mathcal{C})(W_i u) = R(\mathcal{C})(\bar{\omega}_i u) = \bar{\omega}_i R(\mathcal{C})u$$

and so $R(\mathcal{C})u \in \mathcal{K}_b^\omega$.

Let $\{\xi_{\kappa, \sigma} : \kappa \in F_v, \sigma \in \{x, y, z\}\}$ be the natural basis for the column vector space $\mathbb{C}^{3|F_v|}$. Write $\xi_{\kappa, \sigma}^\omega$ for the displacement vectors in \mathcal{K}_a^ω which "extend" the basis

elements $\xi_{\kappa,\sigma}$. Formally, in terms of Kronecker delta symbol, we have

$$(\xi_{\kappa,\sigma}^\omega)_{\kappa',k} = \delta_{\kappa,\kappa'} \omega^k \xi_{\kappa,\sigma}.$$

Similarly let $\eta_e, e \in F_e$, be the standard basis for $\mathbb{C}^{|F_e|}$ and write $\eta_e^\omega, e \in F_e$, for the natural associated basis for \mathcal{K}_b^ω , with

$$(\eta_e^\omega)_{e',k} = \omega^k \delta_{e,e'}.$$

THEOREM 1. *Let \mathcal{C} be a crystal framework in \mathbb{R}^d with matrix function $\Phi_{\mathcal{C}}(z)$ and let $\omega \in \mathbb{T}^d$. Then the scalar matrix $\Phi_{\mathcal{C}}(\bar{\omega})$ is the representing matrix for the linear transformation $R^\omega : \mathcal{K}_a^\omega \rightarrow \mathcal{K}_b^\omega$ with respect to the natural bases $\{\xi_{\kappa,\sigma}^\omega\}$ and $\{\eta_e^\omega\}$.*

Proof. Let \tilde{u} be a velocity vector in \mathcal{K}_a^ω determined by $u \in \mathbb{C}^{|F_v|}$ as above. Let e in F_e be an edge of the form $[p_{\kappa,k}, p_{\tau,l}]$ and let $\langle \cdot, \cdot \rangle$ denote the bilinear form on \mathbb{C}^d . Then, from the definition of the rigidity matrix $R(\mathcal{C})$, the $(e, 0)^{th}$ entry of $R(\mathcal{C})\tilde{u}$ in \mathcal{K}_b^ω can be written as

$$\begin{aligned} (R(\mathcal{C})\tilde{u})_{e,0} &= \langle v_e, \tilde{u}_{\kappa,k} \rangle + \langle -v_e, \tilde{u}_{\tau,l} \rangle \\ &= \langle v_e, \omega^k u_\kappa \rangle + \langle -v_e, \omega^l u_\tau \rangle \\ &= \langle \omega^k v_e, u_\kappa \rangle + \langle -\omega^l v_e, u_\tau \rangle. \end{aligned}$$

This agrees with $(\Phi_{\mathcal{C}}(\bar{\omega})u)_e$, both in the case $\kappa \neq \tau$ and in the reflexive case $\kappa = \tau$ and the theorem follows. \blacksquare

In particular the strictly periodic (one-cell-periodic) (real or complex) infinitesimal flexes are determined by the (real or complex) vectors in the nullspace of the real matrix $\Phi(1, \dots, 1)$. This *periodic rigidity matrix* has rows carrying entries from the vectors $v_e, -v_e$ in the case of nonreflexive edges of the motif, with reflexive edges contributing zero rows.

The terminology of the following definition is justified in the next section.

DEFINITION 8. *The rigid unit mode spectrum (RUM spectrum) of the crystal framework \mathcal{C} in \mathbb{R}^d , with translation group \mathcal{T} , is the set $\Omega(\mathcal{C})$ of points $\omega = (\omega_1, \dots, \omega_d)$ in \mathbb{T}^d for which there is a nonzero vector u in \mathcal{K}_a^ω which is an infinitesimal flex for \mathcal{C} .*

From the theorem we have

$$\Omega(\mathcal{C}) = \{\omega \in \mathbb{T}^d : \ker \Phi(\bar{\omega}) \neq \{0\}\}$$

and so from a commutative algebra perspective this set is a real algebraic variety.

The *rigid unit modes* themselves are the nonzero infinitesimal flexes giving rise to points in the RUM spectrum and the *mode multiplicity function* is the integer-valued function defined on $\Omega(\mathcal{C})$ by $\mu(\omega) = \dim \ker R^\omega$. We may write $\Omega(\mathcal{C}, \mathcal{T})$ if we want to make explicit the dependence \mathcal{T} . Additionally we define the *primitive RUM spectrum* of a general crystal framework \mathcal{C} to correspond to the choice $(\mathcal{C}, \mathcal{T})$ with \mathcal{T} chosen in the maximal sense, corresponding to the translation group for a primitive unit cell. Note that if one doubles the period vectors, and hence the unit cell that they subtend, then the new RUM spectrum

is obtained simply as the range of the old spectrum under the argument doubling map $(w_1, w_2, w_3) \rightarrow (w_1^2, w_2^2, w_3^2)$. In particular it follows that the real dimension $\Omega(\mathcal{C})$ as a real algebraic variety is independent of the choice of translation group, and we refer to this dimension as the *rigid unit mode dimension* of \mathcal{C} , denoted $\dim_{\text{rum}}(\mathcal{C})$.

While we have given the multi-phase form of the RUM spectrum the convention in material science is to indicate such a spectrum in two and three dimensions as the set of (reduced) wave vectors $\mathbf{k} = (\mathbf{k}_1, \mathbf{k}_2, \mathbf{k}_3)$ in the unit cube $[0, 1]^3$ obtained by taking logarithms coordinatewise. For a simple crystal framework $(\mathcal{C}, \mathcal{T})$ this often consists of a union of a finite number of points, lines and planes in $[0, 1]^d$, and possibly all of $[0, 1]^d$. In this case we say that $\Omega(\mathcal{C})$ is *standard*. Otherwise, borrowing terminology from Dove et al [6], we say that the RUM spectrum is *exotic*.

For a framework in Maxwell counting equilibrium $\Omega(\mathcal{C})$ is the zero set of $p_{\mathcal{C}}(z)$. For the kagome framework, for example, the polynomial is $(z - 1)(w - 1)(z - w)$ and we obtain the set which is the union of the three curves on \mathbb{T}^2 defined by $z = 1$, $w = 1$ and $z = w$. In terms of wave vectors this translates to the union of the three parametrised lines $(0, \alpha)$, $(\alpha, 0)$ and (α, α) .

When \mathcal{C} is edge rich, with $|F_e| > d|F_v|$ then one may instead form the finite family of polynomials of the $d|F_v| \times d|F_v|$ submatrices of $\Phi_{\mathcal{C}}(z)$. Then the RUM spectrum will be contained in the intersections of the zero sets of these polynomials on the torus.

The RUM spectrum will generally carry symmetries reflecting the point group symmetries of the crystal framework. Even so the point group may be trivial and the following rather theoretical inverse problem may well have an affirmative answer.

Problem. Let $q(z, w)$ be a polynomial with real coefficients with $q(1, 1) = 0$. Is there a crystal polynomial $p(z, w)$ whose zero set on the 2-torus is the same as that for $q(z, w)$?

Floppy modes and their asymptotic order. In applications the term *floppy mode* often refers to rigid unit flexibility and oscillation within a large supercell and there is interest in the asymptotic order as the supercell dimensions tend to infinity. In particular, a so-called *order N crystal* or periodic structure is one for which the asymptotic order agrees with the order of the number of atoms in the supercell, which is of order n^3 in an $n \times n \times n$ supercell of a 3D crystal. We now formalise this terminology in the direction of infinitesimal flexes and indicate connections with the RUM spectrum.

DEFINITION 9. Let \mathcal{C} be a crystal framework in \mathbb{R}^d with translation group $\mathcal{T} = \{T_k : k \in \mathbb{Z}^d\}$.

(i) A *floppy mode*, or *n -fold periodic floppy mode*, of \mathcal{C} is a nonzero real vector $u = (u_{\kappa, k})$ in the nullspace (kernel) of $R(\mathcal{C})$ which is supercell-periodic for the subgroup $n\mathbb{Z} \times n\mathbb{Z} \times n\mathbb{Z}$ for some n ; $u_{\kappa, k} = u_{\kappa, 0}$ for all k in this subgroup.

(ii) ν_n is the dimension of the real linear space of real n -fold periodic floppy modes.

(iii) A crystal framework \mathcal{C} in \mathbb{R}^3 is of order N^α for floppy modes, where $\alpha = 0, \frac{1}{3}, \frac{2}{3}$ or 1, if $\nu_n \geq Cn^{3\alpha}$ for all n for some $C > 0$, while there is no such

constant for the power $\alpha + \frac{1}{3}$. In particular \mathcal{C} is said to be of order N if $\nu_n \geq cn^3$ for some constant $c > 0$.

Since the real and imaginary parts of a complex infinitesimal flex are real infinitesimal flexes it follows that $\nu_n \leq \dim \ker R_n(\mathcal{C}) \leq 2\nu_n$ where $R_n(\mathcal{C})$ is the rigidity matrix for n -fold periodicity viewed as a complex vector space linear transformation. Thus, in considerations of asymptotic order we may more conveniently consider the complex scalar case. The matrix $R_n(\mathcal{C})$ is the rigidity matrix for strict periodicity relative to the subgroup \mathcal{T}' of \mathcal{T} corresponding to the index subgroup $n\mathbb{Z} \times \dots \times n\mathbb{Z}$. Accordingly it is given as the periodic rigidity matrix associated with a motif for the n -fold supercell. Such a motif can be taken simply as the union of n^d translates of the given motif. More conveniently, it is possible to explicitly block diagonalise $R_n(\mathcal{C})$ (as a complex vector space transformation) as a direct sum (even an orthogonal direct sum for natural inner product) of the matrices $\Phi(\omega)$ as ω ranges over the set of points, \mathbb{T}_n say, with coordinates ω_j of the form $e^{2\pi k_j/n}$, where $0 \leq k_j < n$ are integers. This then gives the following counting formula for floppy modes:

$$\dim \ker R_n(\mathcal{C}) = \sum_{0 \leq k_i < n, 1 \leq i \leq d} \dim \ker(\Phi_{\mathcal{C}}(\omega^k))$$

where $\omega^k = (e^{2\pi i k_1/n}, \dots, e^{2\pi i k_d/n})$. (This answers a question posed by Simon Guest). In fact an elementary direct proof of this formula follows from the fact that nonzero vectors u, v from distinct nullspaces $\ker \Phi_{\mathcal{C}}(\omega^k)$ are linearly independent, on the one hand, and that, on the other hand, by the usual averaging arguments, any n -fold periodic flex may be decomposed as a sum of pure frequency n -fold periodic infinitesimal flexes. By "pure frequency" we mean phase-periodic in each coordinate for some n^{th} root of unity (depending on the coordinate).

It is of interest then to consider the rational subset of the RUM spectrum corresponding to floppy modes, namely

$$\Omega_{\text{rat}}(\mathcal{C}) := \bigcup_{n=1}^{\infty} (\Omega(\mathcal{C}) \cap \mathbb{T}_n^d)$$

and to ask:

To what extent do the asymptotics of floppy modes determine the RUM dimension ?

In the case of order N (the maximal order) there is a close connection as we show below. Also, if we suppose that $\Omega(\mathcal{C})$ has dimension t with t less than d and that $\Omega(\mathcal{C})$ is *rationally parametrised*, in the sense that there are rational coefficient real polynomials $g_1(s), \dots, g_d(s)$, in variable s_1, \dots, s_t , such that

$$\Omega(\mathcal{C}) = \{(\exp(2\pi i g_1(s)), \dots, \exp(2\pi i g_d(s))) : s \in [0, 1]^t\} \subseteq \mathbb{T}^d,$$

then it follows that $\Omega_{\text{rat}}(\mathcal{C})$ is dense in $\Omega(\mathcal{C})$ and that the n -fold periodic floppy modes are of order n^t . But in the general case one should expect examples where the rational points of the RUM spectrum are not dense. It would be of theoretical interest to identify some such examples in which, for example, a curved RUM spectrum only contains a finite number of irrational wave vectors.

For the proof of Theorem 2 we note the following lemma.

LEMMA 1. (i) Let \mathcal{C} be a d -dimensional crystal framework with motif set (F_v, F_e) and RUM spectrum $\Omega(\mathcal{C}) \subseteq \mathbb{T}^d$. Then

$$d - 1 + |\Omega(\mathcal{C}) \cap \mathbb{T}_n^d| \leq \dim \ker R_n(\mathcal{C}) \leq d|F_v||\Omega(\mathcal{C}) \cap \mathbb{T}_n^d|$$

where $|F_v|$ is the number of vertices in the partition unit cell and where \mathbb{T}_n^d is the discrete torus determined by n^{th} roots of unity.

(ii) If $\dim \ker R_n(\mathcal{C}) \geq cn^\alpha$ for some $c > 0, \alpha > 0$, then $\dim(\Omega(\mathcal{C})) \geq \alpha$.

Proof. (i) The counting formula implies the second inequality since $\dim \ker(\Phi_{\mathcal{C}}(\omega) \leq d|F_v|$ for all ω . Also, if $\omega^k \in \Omega(\mathcal{C}) \cap \mathbb{T}_n^d$ then $\dim \ker \Phi_{\mathcal{C}}(\omega^k) \geq 1$, while for wave vector $\mathbf{k} = (0, 0, 0)$ we have $\dim \ker(\Phi_{\mathcal{C}}(1, \dots, 1)) \geq d$, since there are certainly d linearly independent translation infinitesimal flexes. Thus the first inequality follows.

(ii) follows from (i) since for any algebraic variety Ω , if the dimension is less than the integer α then the cardinality of $\Omega \cap \mathbb{T}_n^d$ is at most of order $n^{\alpha-1}$. ■

It can be shown by direct linear algebra, as we now sketch, that if a crystal framework has order N then there exists a local infinitesimal flex. The idea of the proof is to consider the components, $P(u_k)$ say, of a basis u_1, u_2, \dots of the vector space of n -fold supercell flexes, where these components correspond to the "boundary velocity space" of the supercell motif (F_v^n, F_e^n) . More precisely, the boundary space is the span of the basis element corresponding to vertices of edges in F_e^n which are not in F_v^n . The boundary vector space has dimension of order n^{d-1} and so if n is large enough then (by order N) there exists a linear dependence between the vectors $P(u_k)$. Thus the component $P(u)$, for the vector u which is the corresponding linear combination of the u_k , is zero. This infinitesimal flex u , vanishes on the boundary, since $P(u) = 0$, and this readily gives rise to a local infinitesimal flex.

THEOREM 2. With the notation above the following assertions are equivalent for a crystal framework \mathcal{C} in \mathbb{R}^d .

- (i) \mathcal{C} has a local floppy mode.
- (ii) \mathcal{C} is of order N .
- (iii) $\dim_{\text{rum}}(\mathcal{C}) = d$.
- (iv) $\Omega(\mathcal{C}) = \mathbb{T}^d$.

Proof. We have indicated how (i) follows from (ii) above. To see that (ii) follows from (i) note that if u is a nonzero local infinitesimal flex and ω is a multi-phase in \mathbb{T}^d then the sum

$$v = \sum_{k \in \mathbb{Z}^d} \omega^k T_k u$$

is a phase-periodic infinitesimal flex and it is nonzero for almost every ω . The lemma shows that (ii) and (iii) are equivalent, and the equivalence of (iii) and (iv) follow since $\Omega(\mathcal{C})$ is a real algebraic variety in \mathbb{T}^d . ■

Square-summable flexes. Departing from periodicity and phase periodicity it is natural to enquire to what extent a crystal framework \mathcal{C} might be resistant to flexes whose velocities diminish to zero at infinity. With this in mind write \mathcal{K}_a^2 and

\mathcal{K}_b^2 for the Hilbert spaces of square summable sequences in \mathcal{K}_{atom} and \mathcal{K}_{bond} . Thus $u = (u_{\kappa,k}) \in \mathcal{K}_a^2$ is such that the sum of the squares of the Euclidean norms $|u_{\kappa,k}|$ is finite. Then it is elementary to show that $R(\mathcal{C})$ determines a bounded Hilbert space operator from \mathcal{K}_a^2 to \mathcal{K}_b^2 . This operator intertwines the shift transformations, as before, although now these transformations are unitary operators on Hilbert spaces. Identifying square-summable sequences with square-integrable functions in a standard way one obtains unitary equivalences between \mathcal{K}_a^2 and $L^2(\mathbb{T}^d) \otimes \mathbb{C}^{|F_v|}$ and between \mathcal{K}_b^2 and $L^2(\mathbb{T}^d) \otimes \mathbb{C}^{|F_e|}$. With careful spatial identifications one can identify the corresponding unitary transform $U_b R(\mathcal{C}) U_a^*$ of the operator $R(\mathcal{C})$ with a multiplication operator between these matrix-valued function spaces, where the multiplying function is $\Phi_{\mathcal{C}}(\bar{z})$. In this way the matrix function for \mathcal{C} and its translation group appears naturally from the point of view of square-summable velocity sequences. For more details see Owen and Power [18] where other operator-theoretic considerations are given.

More speculatively, it would be of interest to investigate other possible roles of the matrix function, particularly with regard to approximate flexes and quantitative issues. For example for the 3D framework \mathcal{C} we may define the non-negative scalar function λ on \mathbb{T}^3 with

$$\lambda : (z_1, z_2, z_3) \rightarrow \lambda_{\min}(\Phi_{\mathcal{C}}(z_1, z_2, z_3)^* \Phi_{\mathcal{C}}(z_1, z_2, z_3))$$

where $\lambda_{\min}(A)$ denotes the smallest eigenvalue of the positive operator A . In particular, when the spectrum is trivial, that is, equal to the singleton set $\{(1, 1, 1)\}$ (corresponding to the acoustic modes), the function is nonvanishing except at this point and could be viewed as an measure of RUM resistance for the purposes of comparisons.

6. RUMs and low energy phonons.

In the traditional, less idealised, mathematical models for crystalline dynamics the atoms oscillate harmonically (under the harmonic hypothesis). The bond strengths are finite and a dynamical matrix embodying them governs the modes and wave vectors of phonon excitations. We show how the RUM spectrum $\Omega(\mathcal{C})$ arises as the set of wave vectors \mathbf{k} of the harmonic excitations of \mathcal{C} which induce vanishing bond distortion in their low frequency (long wavelength, low energy) limits.

Suppose that \mathcal{C} is a crystal framework in \mathbb{R}^d , with motif data (F_v, F_e, \mathcal{T}) and suppose that the vertices $p_{\kappa,k}$ undergo a standard wave motion,

$$p_{\kappa,k}(t) = p_{\kappa,k} + u_{\kappa,k}(t), \quad \kappa \in F_v, k \in \mathbb{Z}^d,$$

where $u_{\kappa,k}(t)$ represents the local oscillatory motion of atom κ in the translated unit cell with label $k \in \mathbb{Z}^3$. Following standard formula-simplifying conventions, the framework point motions take values in \mathbb{C}^d , the case of real motion being recoverable from real and imaginary parts. (See Dove [5].) Thus we have

$$u_{\kappa,k}(t) = u_{\kappa} \exp(2\pi i \mathbf{k} \cdot k) \exp(i\alpha t)$$

where $\mathbf{u} = (u_{\kappa})_{\kappa \in F_v}$ is a fixed vector in $\mathbb{C}^{|F_v|}$, where \mathbf{k} is the wave vector and where α is the frequency.

Considering the distortion $\Delta_e(t)$ for the edge $e = [p_{\kappa,k}, p_{\tau,k+\delta(e)}]$ we have

$$\begin{aligned}\Delta_e(t) &:= |p_{\kappa,k}(t) - p_{\tau,k+\delta(e)}(t)|^2 - |p_{\kappa,k}(0) - p_{\tau,k+\delta(e)}(0)|^2 \\ &= 2\text{Re}\langle p_{\kappa,k} - p_{\tau,k+\delta(e)}, u_{\kappa,k}(t) - u_{\tau,k+\delta(e)}(t) \rangle \\ &\quad + 2\text{Re}\langle p_{\kappa,k} - p_{\tau,k+\delta(e)}, u_{\kappa,k}(0) - u_{\tau,k+\delta(e)}(0) \rangle \\ &\quad + \epsilon(\mathbf{u}, \mathbf{k}, k, \alpha t)\end{aligned}$$

where

$$\epsilon(\mathbf{u}, \mathbf{k}, k, \alpha t) = |u_{\kappa,k}(t) - u_{\tau,k+\delta(e)}(t)|^2 - |u_{\kappa,k}(0) - u_{\tau,k+\delta(e)}(0)|^2.$$

First note that in any finite time period $[0, T]$ the difference quantities $\epsilon(\mathbf{u}, \mathbf{k}, k, \alpha t)$ tends to zero uniformly, for all $t \in [0, T]$ and all k in \mathbb{Z}^3 , as the frequency α tends to zero. This follows readily from the fact that for any θ the quantity

$$|\sin(\alpha t + \theta) - \sin(\alpha t)|^2 - |\sin(\theta) - \sin(0)|^2$$

tends to zero uniformly for $t \in [0, T]$ as α tends to zero.

For the other terms for $\Delta_e(t)$ note that

$$\begin{aligned}& 2\text{Re}\langle p_{\kappa,k} - p_{\tau,k+\delta(e)}, u_{\kappa,k}(t) - u_{\tau,k+\delta(e)}(t) \rangle \\ &= 2\text{Re}[e^{-\alpha t - 2\pi i \mathbf{k} \cdot k} \langle p_{\kappa,k} - p_{\tau,k+\delta(e)}, u_{\kappa} - \omega^{\delta(e)} u_{\tau} \rangle] \\ &= 2\text{Re}[e^{-\alpha t - 2\pi i \mathbf{k} \cdot k} \langle p_{\kappa} - p_{\tau, \delta(e)}, u_{\kappa} - \omega^{\delta(e)} u_{\tau} \rangle]\end{aligned}$$

which is zero, irrespective of t , if $(\omega^k u_{\kappa})$ is an infinitesimal flex of the framework.

It follows that we have proven the implication (i) implies (ii) in the following proposition and in fact the converse assertion follows from the same equations. The theorem underlies the correspondence of the points in $\Omega(\mathcal{C})$ with the wave vectors of RUM phonons that arise in simulations.

THEOREM 3. *Let \mathcal{C} be a crystal framework, with specified periodicity, and let \mathbf{k} be a wave vector with point $\omega \in \mathbb{T}^3$. Then the following assertions are equivalent.*

- (i) $(\omega^k u_{\kappa})_{\kappa,k}$ is a nonzero phase-periodic infinitesimal flex for \mathcal{C} .
- (ii) For the vertex wave motion

$$p_{\kappa,k}(t) = p_{\kappa,k} + u_{\kappa} \exp(2\pi i \mathbf{k} \cdot k) \exp(i\alpha t),$$

and a given time interval, $t \in [0, T]$, the bond length changes

$$\delta e(t) = |p_{\kappa,k}(t) - p_{\tau,k+\delta(e)}(t)| - |p_{\kappa,k}(0) - p_{\tau,k+\delta(e)}(0)|,$$

tend to zero uniformly, in t and e , as the wavelength $2\pi/\alpha$ tends to infinity.

In the last two decades the RUM spectra of frameworks associated with specific material crystals have been derived by experiment and by simulation using lattice dynamics. Some of the results of this approach can be found in Giddy et al [7], Hammond et al [9], [10], Dove et al [6] and Swainson and Dove [23]. In particular the programme CRUSH has been used for this purpose and the method here reflects principle (ii) in the theorem above. Indeed in the simulations a double limiting process is used (the split atom method) in which each shared vertex (often an oxygen atom) is duplicated, for each rigid unit, and connected by bonds

of zero length and increasing strength, tending to infinity. In this setting the RUM wave vectors coincide with those for which the long wavelength limits have vanishing energy, and so they can be identified and counted.

7. Determinations of RUM spectra.

The rigid unit mode spectrum is now determined for a variety of basic crystal frameworks. Also we emphasise an infinitesimal flex method for the identification of lines and planes of wave vectors. The spectrum is of standard type for the frameworks $\mathcal{C}_{\mathbb{Z}^d}$, \mathcal{C}_{sq} , $\mathcal{C}_{\text{star}}$, \mathcal{C}_{kag} , $\mathcal{C}_{\text{Knet}}$, \mathcal{C}_{Oct} and \mathcal{C}_{SOD} , while for the 2D zeolite \mathcal{C}_{Oct} it is a union of four closed curves.

Consider once again the basic grid framework $\mathcal{C}_{\mathbb{Z}^2}$ in the plane with motif consisting of a single vertex, $F_v = \{p_\kappa\}$, and two edges. Examining all edges it becomes evident that there exists an infinitesimal flex u supported on the x -axis, with $u_{\kappa, (k_1, 0)} = (1, 0)$ for all $k_1 \in \mathbb{Z}$. Using all the parallel translates $T_{(0, k_2)}u$ of u , we may define a phase-periodic velocity vector v in $\mathcal{K}_{\text{atom}}$,

$$v = \sum_{k_2 \in \mathbb{Z}} \omega_2^{k_2} T_{(0, k_2)} u,$$

where ω_2 is a fixed point in \mathbb{T} . Note that v is well-defined and

$$R(\mathcal{C})v = R(\mathcal{C}) \sum_{k_2 \in \mathbb{Z}} \omega_2^{k_2} T_{(0, k_2)} u = \sum_{k_2 \in \mathbb{Z}} \omega_2^{k_2} R(\mathcal{C})T_{(0, k_2)} u = \sum_{k_2 \in \mathbb{Z}} \omega_2^{k_2} T_{(0, k_2)} R(\mathcal{C})u = 0.$$

Thus v is an infinitesimal flex, phase-periodic for the point $(1, \omega_2)$ in \mathbb{T}^2 and so $(1, \omega_2)$ lies in the RUM spectrum. In the language of wave vectors the RUM spectrum contains the line of wave vectors $(0, \alpha)$. By symmetry the line $(\alpha, 0)$ is also included. Similar arguments apply to the kagome lattice which also has linearly localised infinitesimal flexes. (See also [8], [11] for example.)

More generally, suppose that a crystal framework \mathcal{C} has a nonzero infinitesimal flex u which is

- (a) *band limited*, in the sense of being supported by a set of framework vertices within a finite distance of a direction axis for \mathcal{T} , and
- (b) *periodic*, or more generally, *phase-periodic* in the direction axis direction.

By (a) one can form a sum analogous to that above, using the complementary axis direction(s), to obtain a well-defined phase-periodic infinitesimal displacement which, by translational invariance and linearity, is an infinitesimal flex. If ω_1 is the phase in (b) then we deduce that $\{\omega_1\} \times \mathbb{T}^{d-1}$ is contained in $\Omega(\mathcal{C})$.

Thus, for the grid framework $\mathcal{C}_{\mathbb{Z}^3}$ in three dimensions one deduces from the evident line-localised infinitesimal flexes that there are three surfaces, $z = 1$, $w = 1$ and $u = 1$, in $\Omega(\mathcal{C}_{\mathbb{Z}^3})$. In general a line-localised flex of this type leads directly to a hyperplane of wave vectors in the RUM spectrum.

Similar observations hold for plane-localised flexes. In three dimensions, for example, such a flex, which is assumed to be "in-plane phase-periodic", leads to a line of RUM wave vectors. This is the case for \mathcal{C}_{Oct} , considered below, and the RUM spectrum here is the union of these planes.

Example (a): The regular 4-ring framework $\mathcal{C}_{\text{star}}$. This 2D zeolite is defined by translates of the regular 4-ring of equilateral triangles in Figure 6. It is sufficiently simple that one can deduce its RUM spectrum and its crystal polynomial $p_{\text{star}}(z, w)$ from infinitesimal arguments.

For a motif we may take F_e to consist of the edges of the 12-edged star and take F_v to be the set of four vertices of the square together with the westward and southern vertex. The four edges in the motif incident to the external vertices (north and eastward) provide four rows of the 12 by 12 matrix $\Phi_{\mathcal{C}}(z, w)$ each of which carries simple monomials (either z or w or their conjugates). Thus $p_{\mathcal{C}}(z, w)$ has total degree at most 4. One can identify band-limited infinitesimal flexes as indicated in Figures 6 and 7. Here the top and bottom vertices of each are fixed and there is horizontal periodic extension to a band-limited infinitesimal flex. In the former case there is two-step horizontal periodicity while in the latter case there is strict horizontal periodicity although the band is two cells wide.

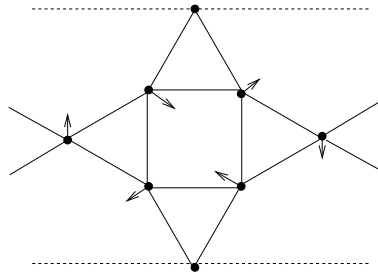


Figure 6. A 2-cell-periodic band-limited flex of $\mathcal{C}_{\text{star}}$.

From the discussion above the first band-limited flex shows that the phase $(-1, \omega_2)$ lies in $\Omega(\mathcal{C})$ for all $\omega_2 \in \mathbb{T}$. By symmetry $(\omega_1, -1) \in \Omega(\mathcal{C})$ for all $\omega_1 \in \mathbb{T}$. The second band-limited flex shows that $\{1\} \times \mathbb{T}$ lies in $\Omega(\mathcal{C}_{\text{star}})$ and hence so too does $\mathbb{T} \times \{1\}$ by symmetry. Thus $\Omega(\mathcal{C})$ contains the set

$$(\{1\} \times \mathbb{T}) \cup (\mathbb{T} \times \{1\}) \cup (\{-1\} \times \mathbb{T}) \cup (\mathbb{T} \times \{-1\})$$

and so $p_{\mathcal{C}}(z, w)$ must be divisible by the irreducible factors $z - 1, w - 1, z + 1, w + 1$. Since $p_{\mathcal{C}}(z, w)$ has total degree at most 4 it follows that either p vanishes identically or

$$p_{\mathcal{C}}(z, w) = (z - 1)(w - 1)(z + 1)(w + 1).$$

In fact the former case does not hold. One can see this, thematically, by demonstrating that there are no local flexes or one may compute $\det \Phi(1/3, 1/3) \neq 0$. Thus the RUM spectrum is precisely the fourfold union above.

Example (b): The 2D zeolite framework \mathcal{C}_{oct} . There are no local or band-limited infinitesimal flexes evident for the regular octagon framework and the so the expectation is that the RUM spectrum is trivial or a union of proper curves.

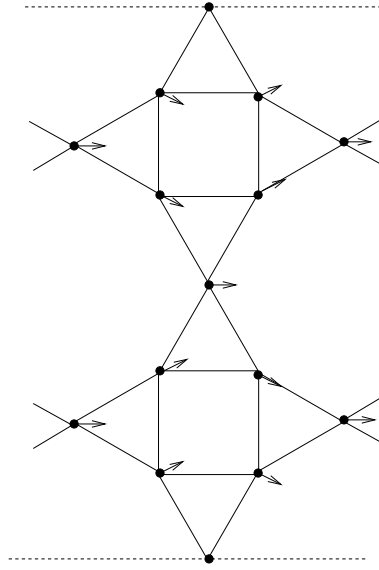


Figure 7. A 1-cell-periodic band-limited flex of $\mathcal{C}_{\text{star}}$.

Returning to the 2D zeolites of Figure 5 the third of these, with external angle $8\pi/12$, is equal to $\mathcal{C}_{\text{star}}$, although with a different translation group, \mathcal{T}' say, for which the old period vectors are rotated by $\pi/4$ and scaled by the factor $\sqrt{2}$. In view of this rotation it follows that

$$\Omega(\mathcal{C}_{\text{star}}, \mathcal{T}') = \{(w, w), (w, -w) : w \in \mathbb{T}\}.$$

In terms of reduced wave vectors this corresponds to the subset of the unit square $[0, 1]^2$ given as the union of the two diagonals.

As we have noted earlier, the 4-pointed star framework is related to its 8-pointed star companion \mathcal{C}_{oct} by a continuous flex. It follows that the 24 by 24 symbol matrix function $\Phi_{\text{star}}(z_1, z_2)$ for the former (for \mathcal{T}') is naturally "continuously connected" to the symbol function $\Phi_{\text{oct}}(z_1, z_2)$ by an explicit continuous path $t \rightarrow \Phi_t(z_1, z_2)$. This in turn provides a set-valued map which we refer to as the *RUM spectrum evolution* for this (periodicity-preserving) flex:

$$t \rightarrow \Omega(\Phi_t(z)).$$

When this is made explicit by computation the octagon framework has exotic spectrum as indicated in Figure 8 and evolves towards the cross-shaped spectrum of $\mathcal{C}_{\text{star}}$ under the continuous flex.

In fact one can obtain the RUM spectrum of the octagon framework completely analytically, although with some significant algebraic complexity, as follows.

Note first that a motif for \mathcal{C}_{oct} is formed by the 24 edges of the 8-ring for F_e with F_v obtained by omitting four boundary vertices as, for example, in Figure 5. There are 8 edges with external vertices and each contributes a row to Φ_{oct}

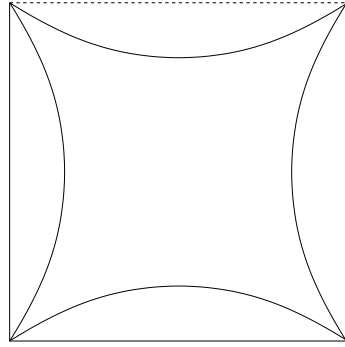


Figure 8. The curved wave vector spectrum of \mathcal{C}_{oct} .

with a simple monomial and so it follows that $p_{\text{oct}}(z, w)$ has degree 8 at most. The 24×24 function matrix $\Phi_{\text{oct}}(z)$ is sparse and the (at most) four nonzero functions in each row may be conveniently normalised by dividing by the magnitude of the x -coordinate difference for that row. The magnitudes of the nonzero nonunit entries are then the tangents of the angles $k\pi/24$, for $k = 1, 3, 5, 7, 9, 11$, all of which lie in the field extension $\mathbb{Q}(\sqrt{2}, \sqrt{3})$. The crystal polynomial can be computed and admits an explicit factorisation as the product

$$p_{\text{oct}}(z, w) = p_1(z, w)p_2(z, w),$$

where

$$p_1(z, w) = z^2w - (\sqrt{3} + \sqrt{2})zw^2 + 2(\sqrt{3} + \sqrt{2} - 1)zw - (\sqrt{3} + \sqrt{2})z + w,$$

$$p_2(z, w) = z^2w - (\sqrt{3} - \sqrt{2})zw^2 + 2(\sqrt{3} - \sqrt{2} - 1)zw - (\sqrt{3} - \sqrt{2})z + w.$$

Each of the factors is responsible for two of the four closed curves that comprise the RUM spectrum.

Returning to the as yet unconsidered 2D zeolite of Figure 5 (the first framework indicated, with an "8-ring of triangles encircling a square") we remark that one can also show, by band-limited infinitesimal flex analysis, that it has standard RUM spectrum, being the subset of the unit square $[0, 1]^2$ given as the union of the axes.

Each of these 2D zeolites has a 3D zeolite companion obtained by the layer construction. The companion $\tilde{\mathcal{C}}_{\text{oct}}$ for \mathcal{C}_{oct} also has exotic RUM spectrum and in fact by earlier arguments contains the surface of points (z, w, u) in \mathbb{T}^3 with (z, u) in $\Omega(\mathcal{C}_{\text{oct}})$ and u any point of \mathbb{T} .

The two-dimensional crystal framework deformation implied by Figure 5 is an example of a *finite flex* and continuous and smooth flexes such as these serve to model flexibility considerations for zeolites and other micro-porous materials. These finite motions usually take place with an associated contraction and increase in rigid unit density. See for example the collapsing mechanisms of Kapco et al [14] and the flexibility window determinations in Kapco et al [12].

Example (c): A 2D zeolite with order N . Figure 9 shows a unit cell for a 2D zeolite, $\mathcal{C}_{\text{bowtie}}$ say, which is of order N . (This resolves an existence question posed by Mike Thorpe.) To see this property one can verify that there is an

infinitesimal flex of the enclosed finite framework which assigns zero velocities to the six boundary vertices and a nonzero vertical velocity to the central vertex. Thus the entire framework has a local infinitesimal flex and so the RUM spectrum is all of \mathbb{T}^2 .

We remark that in general it need not be the case that an order N crystal framework has a local RUM internal to a unit cell. For example, one could take a new motif and unit cell in which the central vertex is shifted to the boundary and in this case one has to consider a threefold supercell before a local RUM appears.

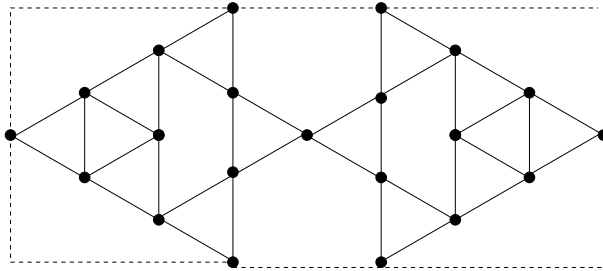


Figure 9. A unit cell which defines $\mathcal{C}_{\text{bowtie}}$.

Example (d): The kagome net framework $\mathcal{C}_{\text{Knet}}$ and its polynomial. The RUM spectrum of the 3D kagome net framework can be derived from that of the 2D kagome framework. The spectrum of the latter is the zero set on the torus \mathbb{T}^2 determined by the crystal polynomial, which an earlier computation showed was equal to $(z - 1)(w - 1)(z - w)$. One can derive this from infinitesimal flex analysis as follows. It is elementary to show that there is no local infinitesimal flex and so $p_{\text{kag}}(z, w)$ is necessarily nonzero. There are line-supported infinitesimal flexes in the directions of the period vectors a_1 and a_2 so it follows from the discussion above that $(z - 1)$ and $(w - 1)$ are factors. Let u be a (similar) infinitesimal flex supported on a line in the direction $a_1 - a_2$ and consider the infinitesimal flexes

$$v = \sum_{k \in \mathbb{Z}} \omega^k T_{(k,0)} u$$

for $\omega \in \mathbb{T}$. In view of the triangular symmetry of \mathcal{C}_{kag} in fact this flex is phase-periodic for the phase (ω, ω) and follows that $(z - w)$ is necessarily a factor of $p_{\text{kag}}(z, w)$. One can see, without calculation, that the total degree of this polynomial is at most 3 and so the derivation is complete.

Moving up a dimension, a phase-periodic flex of \mathcal{C}_{kag} , with phase (ω_1, ω_2) say, induces a "layer-limited" infinitesimal flex of the kagome net framework $\mathcal{C}_{\text{Knet}}$. Thus, for all $\omega \in \mathbb{T}$ there is an infinitesimal flex of $\mathcal{C}_{\text{Knet}}$ with phase $(\omega_1, \omega_2, \omega)$. Similar assertions hold for the other two translation group planes. The crystal polynomial $p_{\text{Knet}}(z, w, u)$ has total degree at most 6 and must vanish on the six planes $z - 1 = 0, w - 1 = 0, u - 1 = 0, z - w = 0, w - u = 0, z - u = 0$. It

follows that

$$p_{Knet}(z, w, u) = (z - 1)(w - 1)(u - 1)(z - w)(w - u)(z - u),$$

for the monomial order with $z > w > u$.

The polynomial above was also obtained in Wegner [24] and Owen and Power [18] by direct calculation.

The kagome net framework and its deformations feature as the tetrahedral net framework for a range of materials and their phases. It is the framework for β -cristobalite, for example, while a particular deformation gives the framework for tridymite. This was the first material for which curved surfaces of RUMs were observed. (Dove et al [6].)

Example (e): Sodalite and \mathcal{C}_{SOD} . The framework \mathcal{C}_{SOD} has a symbol function with 72 rows and columns. Indeed, it is in Maxwell counting equilibrium, being a 3D zeolite crystal framework, and the motif set F_e consists of the edges of three 4-rings of tetrahedra. We prove that \mathcal{C}_{SOD} is of order N . Specifically we show, by infinitesimal flex geometry, that there is a nonzero infinitesimal flex v of the finite sodalite cage framework such that all the outer vertices are fixed by v . That is, $v_{\kappa,\delta} = 0$ if $p_{\kappa,\delta}$ is any of the 24 outer vertices of the cage. The "outer fixed" sodalite cage framework has 36 free vertices with 108 degrees of freedom while there are 144 constraining edges. Despite this considerable over-constraint there is sufficient symmetry to allow in a proper infinitesimal flex.

We shall show that an individual 4-ring, R_1 say, of the sodalite cage has an infinitesimal flex, $v^{(1)}$ say, which fixes the quadruple of outer vertices, that is, the upper vertices in Figure 4. In this flex the quadruple of vertices on the imaginary square have equal magnitude velocity vectors which are directed towards two opposing corners of the imaginary cube. (See the flex arrows in Figure 4). Taking $v^{(1)}$ so that these vectors have magnitude 1 it follows that $v^{(1)}$ is determined up to sign and that this sign may be specified by labeling the cube corners "a" and "r" for their attracting and repelling sense. Note that one can label the eight corners of the imaginary cube in this manner so that no like labels are adjacent. In this case the implied flexes $v^{(1)}, \dots, v^{(6)}$ of the six 4 rings of the sodalite cage have equal displacement vectors at common vertices. This consistency shows that there is an infinitesimal flex of the entire sodalite cage in which the outer vertices are fixed, as required.

It remains to show that there is the stated flex of the 4-ring R_1 . To this end let p_1, p_2 be two non-opposite top vertices of R_1 with intermediate vertex p_3 , let p_1, p_3, p_4 be the vertices of an inward facing face of a tetrahedron of R_1 with vertices p_1, p_3, p_4, p_5 , so that the lower vertex p_5 is a cube-edge midpoint. There is a unique "inward and upward" displacement velocity u_3 of the intermediate vertex p_3 which has unit length and is such $(u_1, u_2, u_3) = (0, 0, u_3)$ is a flex for the two edges $[p_1, p_3], [p_3, p_2]$. The displacement vector u_3 induces a unique displacement vector u_5 which is in the direction of the cube edge and is such that

$$\langle u_5 - u_3, p_5 - p_3 \rangle = 0.$$

The triple $u_1 = 0, u_3$ and u_5 now determine the infinitesimal motion of the tetrahedron, with flex vector u_4 for p_4 . However, the reversal (sign change) of u_3 induces the reversal of u_5 so it is clear from the symmetric position of the tetrahedron that u_4 must be the unique unit norm "outward and downward" flex

at p_4 . Continuing around the ring it follows that R_1 has the desired infinitesimal flex.

One can apply similar constructive flex arguments to other zeolite frameworks and of course to any zeolite crystal framework which contains a sodalite cage as above, such as \mathcal{C}_{LTA} . Also we note (as do Kapko et al [12]) that \mathcal{C}_{RWY} is derived from \mathcal{C}_{SOD} by replacing each tetrahedron by a rigid unit of four tetrahedra. Thus the same infinitesimal flex geometry applies and \mathcal{C}_{RWY} has order N .

Example (f): Perovskite, \mathcal{C}_{sq} and \mathcal{C}_{Oct} . Consider the integer translation group \mathcal{T} and the determination of the framework through the primitive motif (F_v, F_e) where

$$F_v = \{0, 1/2, 1/2), (1/2, 0, 1/2), (1/2, 1/2, 0)\} = \{p_{\kappa,0} : 1 \leq \kappa \leq 3\}$$

and where F_e consists of the twelve framework edges between the centres of adjacent faces of the unit cube $[0, 1]^3$. Thus the vertices of $V(F_e) \setminus F_v$ have the form p_{i,γ_i} , where $\gamma_1 = (1, 0, 0)$, $\gamma_2 = (0, 1, 0)$, $\gamma_3 = (0, 0, 1)$. The framework is therefore edge rich and the matrix function $\Phi_{Oct}(z)$ is 12 by 9.

The framework \mathcal{C}_{Oct} is a 3D analogue of the 2D squares framework \mathcal{C}_{sq} and may be obtained from it by a layer construction and a discarding of redundant edges internal to the octahedra. Thus infinitesimal flexes of the 2D squares lattice imply plane-localised flexes for \mathcal{C}_{Oct} . This observation can be made the basis for an infinitesimal flex analysis determination of the RUM spectrum. A more algebraic approach is possible as follows.

Performing row operations on $\Phi_{\mathcal{C}_{sq}}(z)$, as given in Section 2, we see that (\bar{z}, \bar{w}) is a point of the RUM spectrum if and only if the equivalent matrix

$$\Psi(z, w) = \begin{bmatrix} 1 & -1 & -1 & 1 \\ 0 & -2 & z-1 & z+1 \\ 0 & 0 & 1-wz & 1-wz \\ 0 & 0 & 0 & -2z+2w \\ 0 & 0 & -2+2z & 0 \end{bmatrix}$$

has rank less than 4. This occurs if and only if

$$\begin{bmatrix} 1-wz & 1-wz \\ 0 & -2z+2w \\ -2+2z & 0 \end{bmatrix}$$

has rank equal to 0 or 1. The rank is 0 if and only if $z = w = 1$, corresponding to the two-dimensional space of rigid motions with phase $(1, 1)$, and the rank is 1 if and only if $z = w = -1$. Thus

$$\Omega(\mathcal{C}_{sq}) = \{(1, 1), (-1, -1)\}.$$

The infinitesimal flex for the phase $(-1, -1)$ is the one for which the rigid units, in this case squares with diagonals, rotate infinitesimally in alternating senses.

The alternating rotation flex of \mathcal{C}_{sq} induces a plane-localised flex of \mathcal{C}_{Oct} in each of the framework planes $x = 1/2, y = 1/2, z = 1/2$. It follows that $\Omega(\mathcal{C}_{Oct})$ contains the three sets of phases,

$$\mathbb{T} \times \{-1\} \times \{-1\}, \quad \{-1\} \times \mathbb{T} \times \{-1\}, \quad \{-1\} \times \{-1\} \times \mathbb{T}.$$

That the spectrum is no more than the union of these sets and the singleton $(1, 1, 1)$ can be seen from a row analysis of the 12 by 9 function matrix $\Phi_{\mathcal{C}_{Oct}}(z)$

in the same style as the argument for \mathcal{C}_{sq} . Thus, in wave vector formalism, the RUM spectrum of the octahedral net \mathcal{C}_{Oct} is the set of lines

$$(\alpha, 1/2, 1/2), \quad (1/2, \alpha, 1/2), \quad (1/2, 1/2, \alpha)$$

together with the acoustic wave vector $(0, 0, 0)$.

The corner connected octahedron net crystal framework \mathcal{C}_{Oct} is associated with cubic perovskites, such as SiTO_3 , and RUM distributions have been determined experimentally, Giddy et al [7], Dove et al [6].

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