STILL STATES OF BISTABLE LATTICES, COMPATIBILITY, AND PHASE TRANSITION

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Abstract. A two-dimensional bistable lattice is a periodic triangular network of non-linear bistable rods. The energy of each rod is piecewise quadratic and has two minima. Consequently, a rod undergoes a reversible phase transition when its elongation reaches a critical value. We study an energy minimization problem for such lattices. The objective is to characterize the effective energy of the system when the number of nodes in the network approaches infinity. The most important feature of the effective energy is its "flat bottom". This means that the effective energy density is zero for all strains inside a certain three-dimensional set in the strain space. The flat bottom occurs because the microscopic discrete model has a large number of deformed states that carry no forces. We call such deformations still states. In the paper, we present a complete characterization of the "flat bottom" set in terms of the parameters of the network. This is done by constructing a family of still states whose average strains densely fill the set in question.

The two-dimensional case is more difficult than the previously studied case of one-dimensional chains, because the elongations in two-dimensional networks must satisfy certain compatibility conditions that do not arise in the one-dimensional case. We derive these conditions for small and arbitrary deformations. For small deformations a complete analysis is provided.

Key words. Phase transformation in solids, Bi-stable lattice, Eigenstrain, Compatibility conditions.

AMS subject classifications. 70C20, 74N15, 74Q05, 74Q15, 74A50

1. Introduction. This paper suggests a simple discrete model for phase transition theory. In two dimensions, we study deformation of bi-stable lattices (BL), which are periodic triangular networks of bistable elastic rods. A similar network with linearly elastic rods has been used by Cauchy, who formally averaged this assembly and obtained one of the first equations of continuum elasticity [8]. Here, we consider the same assembly, assuming, however, that the rods are bi-stable. The bi-stability models phase transitions: it is assumed that each rod has two equilibrium states that of length \(l\) and \(l(1+s)\), respectively. Here, \(s\) can vary between \(\frac{1}{2}\) and 2, so that a triangle of transformed rods still remains a triangle.

We believe that this bi-stable lattice model is the simplest and natural finite dimensional model for phase transitions in solids. Although it does not have all the features of continuum model, it appears to capture most of the expected features. At the same time this model allows for detailed description of inhomogeneous deformations and does not involve ad hoc assumptions of the continuum theory.

One dimensional chains of bi-stable rods are well investigated [1, 2, 10, 11, 15, 16]. Each bistable element can be in one of the two states (short and long mode) that differ by the equilibrium length. It is assumed for definiteness that initially all rods are in short mode and can transit to the long mode. The state of a chain with some transited elements is characterized by a scalar relative elongation, or one-dimensional strain, that is the normalized difference between the length in the deformed configuration

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and reference length.

Some results on the two-dimensional bi-stable lattices can be found in [12, 13] (explicit construction of certain special solutions, discrete Green functions), and [6, 7] (direct numerical simulations of time-dependent impact fracture problems). Other related problems are treated in the book [14] that also contains a wealth of additional references.

The present work differs from the above mentioned papers both in approach and in scope. It addresses the specific difficulty of the two-dimensional problem – compatibility conditions on the rods’ length. These conditions are automatically satisfied in one dimensional chains. In two-dimensional lattice, the transition of links to a new state generally leads to elastic deformation of all rods. For example, one transformed (elongated) rod does not fit into a triangular lattice of non-transformed (short) links without distorting the lattice. Each such distortion increases the energy of the assembly.

One of the objectives of the paper is to describe special lattice deformations that we call still states. These deformations carry zero forces in each rod, and thus the energy of each still state is zero. In any such state the length of each rod should be either \( l \) or \( l(1 + s) \). Since the number of rods is larger than the number of nodes times the dimension, an arbitrary assignment of lengths may not correspond to an actual lattice deformation. Therefore compatibility conditions must be satisfied at a still state.

We describe the set of average strains of still states. First, we consider the network of rods undergoing arbitrary deformations. We assume that an effective strain tensor is represented by its eigenvalues. In the plane of eigenvalues, the set \( H_L \) of all still states is a nonconvex curvilinear hexagon. We demonstrate structures that densely fill \( H_L \) and derive nonlinear compatibility conditions.

Further, we consider the case of small deformation, linearize the compatibility conditions and develop the theory in more detail. We show that the linearized compatibility conditions are necessary and sufficient for existence of a lattice deformation with a given set of link elongations. In this sense, the conditions are analogous to the compatibility conditions in continuum elasticity. We also describe a sequence of still states whose average strains reach the average strain of any still state by incremental change of the structure starting from the reference configuration. This procedure can describe an evolutionary irreversible process of phase transition, such as the damage distribution. In this case, the initial state corresponds to “undamaged” configuration and structural change – to irreversible damage propagation.

We also construct the set \( D \) of all average strain tensors corresponding to still states and sets of deformed elements that densely fill that set. In the space of entries of the strain tensor, the set \( D \) turns out to be a parallelepiped. We obtain explicit formulas that relate the side lengths and orientation of this parallelepiped to the non-dimensional critical elongation parameter \( s \), and three lattice direction vectors.

Finally, we suggest an equation for the effective energy density as a function of the total strain. It is as follows: if a strain is inside of the set \( D \), then the energy density is asymptotically close to zero as the number of the network nodes goes to infinity. If the strain is outside of this set, the energy density is proportional to the square of the distance between the strain tensor and \( D \).

The paper is organized as follows. Section 2 contains formulation of the problem and derivation of compatibility conditions for arbitrary deformations. Small deformation compatibility conditions are also obtained here. In Section 3 we describe several
explicit constructions of still states for arbitrary deformations, as well as a set of
eigenstrains of all still states. The remainder of the paper deals with the small de-
formation case. In Section 4.1 we formulate the problem. A new formula for average
strain is obtained in Section 4.2. Unlike existing formulas that relate average strain
to nodal displacements, our formula gives an expression in terms of rod elongations.
This allows to relate components of strain to the average elongations along lattice
directions. Such formulas seem to be of independent interest. In section 5 we pro-
vide a complete analysis of the compatibility conditions. In Section 6, we prove a
characterization theorem for $D$. This is done by constructing another family of still
states called stripes. The eigenstrains of the stripes fill $D$ densely, in the sense that
every strain within this set can be approximated by an average strain of some stripe,
with an error of order $N^{-1/2}$ where $N$ is the number of nodes. Section 7 contains a
discussion of the effective energy density. Conclusions are provided in Section 8.

2. The problem.

2.1. Energy of one bistable rod. Consider a bistable elastic rod with the
energy $W_r(x)$ that possesses two equal minima $W_r(l) = W_r(l(1+s)) = 0$. Let us also
assume that $W_r(x)$ is convex outside of the interval $[l, l(1+s)]$, and its second
derivative is positive in a proximity of each minima $x = l$ and $x = l(1+s)$. For
definiteness, we may assume that the dependence of $W_r(x)$ on the length $x > 0$ is
piece-wise quadratic ($W_r^q$) or polynomial ($W_r^p$)

$$W_r^q(x) = \min \left\{ \frac{1}{2} (x-l)^2, \frac{1}{2} C(x-l(1+s))^2 \right\},$$

$$W_r^p(x) = (x-l)^2 (x-l(1+s))^2,$$

(2.1)

where $l$ is the length of the rod in the reference configuration, $l+s$ is the length in
the elongated mode, and $\frac{1}{4} < 1+s < 2$.

The rods are elastic and locally stable in a proximity of the equilibria. The elastic
force $f_r$ in the rod is

$$f_r(x) = \frac{\partial W_r}{\partial x}.$$  

The rod has two equilibrium states $x = l$ and $x = l(1+s)$ of equal energy

$$f_r(x) = 0, \quad W_r(x) = 0 \quad \text{if } x = l, l(1+s).$$

The magnitude of the force monotonically increases with the elongation $l$ in the prox-
imity of equilibria.

There are several equilibrium lengths $x_\alpha$ and $x_\beta$ for every $f_r$ in a proximity of
zero. They are solutions of the equation

$$\frac{\partial W_r}{\partial x} = C$$

2.2. Triangular network. Consider a triangular periodic network made of the
links defined above. For each node, there are six rods joining it with six neighbors.
When the rods are of equal lengths, the network is elastically isotropic, its Poisson
coefficient is equal to $\frac{1}{4}$. The number of nodes in the period is three times less
than the number of rods between them. There are three families of the parallel rods
in the network
When the rods transit to a different state, the network becomes inhomogeneous (each rod can have a different length). We assume that the transition is periodic. Each period consists of \( N \) nodes where \( N \) can be arbitrary large. The network’s energy is the sum of energies stored in all \( 3N \) links. The length of a link can be expressed through the position of its ends, that is the nodes. The nodes are determined by \( N \) pairs of coordinates in a plane, or by \( 2N \) parameters. Hence, the links’ lengths can not be arbitrary (for example, see Figure 2.2). We conclude that they are subject to \( N \) compatibility conditions that we derive now.

### 2.3. Compatibility conditions.

#### 2.3.1. Necessary condition.

There are conditions that constrain the lengths of the rods that can be joined in a hexagonal lattice. Consider an inner node in the lattice. There are six rods that link the node with the neighbors; let us denote the lengths of them as \( a_1, \ldots, a_6 \). Consider also six rods that surround the node forming a hexagon around it; let us denote the lengths of them as \( b_1, \ldots, b_6 \). These rods form six
triangles joined at the node. The angles at the node will be denoted by $\phi_1, \ldots, \phi_6$.

The lengths of the listed twelve rods cannot be arbitrary. The sum of angles of the six triangles $a_1 a_2, b_1, \ldots$ is equal to $2\pi$,

$$\sum_{i=1}^{6} \phi_i = 2\pi. \quad (2.2)$$

To express this condition in terms of the lengths, we use trigonometry. Cosine of each angle $\phi_i$ is expressed through the lengths of the links as

$$\cos(\phi_i) = \frac{a_i^2 + a_{i+1}^2 - b_i^2}{2a_i a_{i+1}}, \quad i = 1, \ldots, 6. \quad (2.3)$$

Here, $a_7 = a_1$. Therefore, the lengths of rods are constrained as follows

$$\sum_{i=1}^{6} \arccos \left( \frac{a_i^2 + a_{i+1}^2 - b_i^2}{2a_i a_{i+1}} \right) = 2\pi. \quad (2.4)$$

The number of these constraints is equal to the number of inner nodes in the structure.

**2.3.2. Linearized compatibility conditions.** In a linearized case when the lengths of the rods are close to $l$, constraints (2.4) are simplified. Assuming that

$$a_i = l(1 + \kappa_{a_i}), \quad b_i = l(1 + \kappa_{b_i}), \quad i = 1, \ldots, 6 \quad (2.5)$$

we write $\phi_i$ as the function of $\kappa_{a_i}, \kappa_{a_{i+1}}, \kappa_{b_i}$:

$$\phi_i = \arccos \left( \frac{(1 + \kappa_{a_i})^2 + (1 + \kappa_{a_{i+1}})^2 - (1 + \kappa_{b_i})^2}{2(1 + \kappa_{a_i})(1 + \kappa_{a_{i+1}})} \right).$$

Linearizing near

$$\begin{pmatrix} 
\kappa_{a_i} \\
\kappa_{a_{i+1}} \\
\kappa_{b_i} 
\end{pmatrix} = \begin{pmatrix} 
0 \\
0 \\
0 
\end{pmatrix}$$

we find

$$\sum_{i=1}^{6} \phi_i = 2\pi + \frac{\sqrt{3}}{3} \left( \sum_{i=1}^{6} \kappa_{a_i} + \sum_{i=1}^{6} \kappa_{a_{i+1}} - 2\sum_{i=1}^{6} \kappa_{b_i} \right)$$

Substituting this expression into (2.2), we obtain an elegant linearized compatibility condition for lattices

$$\sum_{i=1}^{6} \kappa_{a_i} = \sum_{i=1}^{6} \kappa_{b_i}. \quad (2.6)$$

It states that the sum of the elongations of the spokes that come out of a node is equal to the elongation of the hexagonal rim around this node.

**3. Eigenstrains and still states.**

**3.1. Definitions.**
3.1.1. Still configurations. In this section we consider still configurations. These are deformed states with zero force in each rod. Because of the force-elongation dependence, this requires the length of each rod to be either \( l \) or \( l(1 + s) \). To simplify presentation, in this section we scale \( l = 1 \) and denote \( l(1 + s) = a \). Given a collection of such rod lengths, we call a configuration still if it is geometrically compatible, that is the ends of the link of the length one and \( a \) can meet in the nodes. The examples of still configurations are given below in Figure 3.4. This section contains somewhat informal, but explicit description of the set of (average) eigenstrains of all still states corresponding to arbitrary (not necessarily small), deformations. We recall that eigenstrain is a generic name given to various non-elastic strains, such as strains due to thermal expansion, phase transformation, initial strains and so on.

Notice that the problem is reduced to a geometric problem of tiling of the plane with triangles of four types with the lengths of the sides being equal to \((1, 1, 1)\), \((1, 1, a)\), \((1, a, a)\), and \((a, a, a)\), respectively.

Any still configuration can be split into parts that are also still configurations. Each configuration is characterized by a symmetric second-rank eigenstrain tensor which shows the change of the initial shape of a periodic cell. Assume that a unit square element of periodicity of an initial (non-transformed) network contains a very large number of triangles. After deformation, it changes its shape. Asymptotically, this element is close to a parallelogram. The eigenstrain tensor describes the shape of it: the lengths of its sides and the angle between them.

3.1.2. Eigenstrain of still states. Generally, a deformation due to transformation is followed by elastic deformation of all links due to compatibility conditions \([2.4]\). Therefore, eigenstrain of the composite network is not a convex combination of eigenstrains of its components that are, in principle, incompatible and lead to additional distortions on the boundaries of components. The eigenstrain of a compatible configurations of still states is an affine function (the average) of the eigenstrains of the phases. It is independent of the structure of the mixture

\[
E_{\text{mixture}} = \sum_{i} \mu_{i} E_{i}, \quad \sum_{i} \mu_{i} = 1, \quad \mu_{i} = k_{i} N, \quad i = 0, \ldots, N
\]

(3.1)

where \( \mu_{i} \) is the volume fraction of the phase with eigenstrain \( E_{i} \). The only role of the structure is the selection of compatible states.

Next, we describe the variety of still states.

3.2. The homogeneous still states. In the initial network, there are three families of codirected rods. The homogeneous still states are the states in which all the rods in one family are in either “short” or “long” mode.

3.2.1. Classification of homogeneous states. There are eight \(2^3 = 8\) homogeneous states that consist of periodic arrays of equal triangles and their reverses.

1. Initial state \( S_{\alpha} \) with the lengths of the links equal to \((1, 1, 1)\). The elementary square of the state (the eigenstrain) is assumed to be an identity matrix,

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

(3.2)

2. The dual state \( S_{\beta} \) with the lengths of the links equal to \((a, a, a)\). The unit square element \( E_{\alpha} \) of the initial state is transformed to

\[
E_{\beta} = \begin{pmatrix} a & 0 \\ 0 & a \end{pmatrix}
\]

(3.3)
and it is still proportional to an identity matrix. We call $E_\beta$ the eigenstrain of the state $S_\beta$.

3. State $S_\gamma$ with the lengths of the links equal to $(a, a, 1)$ that consists of isosceles triangles. The square element $E_\alpha$ of the initial state is transformed to

$$E_\gamma = \begin{pmatrix} \tan \gamma & 0 \\ \sqrt{3} & 1 \end{pmatrix}$$

(3.4)

Here, $\gamma$ is the angle between the base of the isosceles triangle and its side, $\gamma$ is expressed through $a$ as

$$a = \frac{1}{2 \cos \gamma}$$

$E_\gamma$ is the eigenstrain of the state $S_\gamma$.

4. Together with $S_\gamma$, there are two twin states, $S'_\gamma$ with the lengths of the links equal to $(a, 1, a)$ and $S''_\gamma$ with the lengths of the links equal to $(1, a, a)$. The eigenstrains of these states are obtained from $E_\gamma$ by rotating it to the angle of $\pm 60^\circ$, respectively.

$$E'_\gamma = R^T E_\gamma R \quad E''_\gamma = R E_\gamma R^T$$

(3.5)

where $R$ is the $60^\circ$ rotation matrix.

5. State $S_\delta$ with the lengths of the links equal to $(1, 1, a)$ that consists of isosceles triangles. The square element $E_\alpha$ of the initial state is transformed to

$$E_\delta = \begin{pmatrix} \frac{2 \sin \delta}{\sqrt{3}} & 0 \\ 0 & a \end{pmatrix} = \begin{pmatrix} \frac{2 \sin \delta}{\sqrt{3}} & 0 \\ 0 & 2 \cos \delta \end{pmatrix}$$

(3.6)

Here, $\delta$ is the angle between the base of the isosceles triangle and its side, $\delta$ is expressed through $a$ as

$$a = 2 \cos \delta$$
The angles $\delta$ and $\gamma$ are related as
\[
\cos \gamma \cos \delta = \frac{1}{4}
\]

6. State $S_\delta$ is also accompanied by the two twin states, $S'_\delta$ with the lengths of the links equal to $(a, 1, 1)$ and $S''_\delta$ with the lengths of the links equal to $(1, a, 1)$. The eigenstrains of these states are obtained from $E_\gamma$ by rotating it by the angle of $60^\circ$, and $-60^\circ$, respectively.

\[
E'_\delta = R^T E_\delta R \quad E''_\delta = R E_\delta R^T
\]

(3.7)

Remark 3.1. Pairs $S_\alpha$, $S_\beta$ and $S_\gamma$, $S_\delta$ are dual: each of them is transferred into another by mutually replacing rods of the length 1 and $a$.

3.2.2. Vertices of the eigenstrain states set. Consider a set $\mathcal{D}$ of all possible eigenstrains. In a three-dimensional space where coordinates are entries of the eigenstrain tensor, this set is a bounded region. The eigenstrains of the homogeneous states correspond to the vertices of $\mathcal{D}$. Indeed, there, the elongation of the each family of the rods is extreme, that is the deformation in the directions of the rods’ families is extreme. The rods cannot be elongated (contracted) in these directions without an elastic deformation. Therefore, there exist three trial matrices $E_{tr_1}$, $E_{tr_2}$ and $E_{tr_3}$ of deformation that correspond to impossible change of eigenstrain in the sense that

\[
E_z + \epsilon E_{tr_i}, \quad z = \alpha, \ldots, \beta, \ i = 1, 2, 3
\]
does not correspond to any still state, for any $\epsilon > 0$.

Consider, for example, state $S_\alpha$. Set of impossible deformations $E_{tr}$ contains a cone of matrices of the form

\[
E_{tr_i}^\alpha = \sum_{i=1}^{3} d_i t_i t_i^T
\]

(3.8)

\[
d_i < 0, \quad t_1 = \left(\begin{array}{c} 1 \\ 0 \end{array}\right), t_3 = \frac{1}{2} \left(\begin{array}{c} 1 \\ \sqrt{3} \end{array}\right), t_3 = \frac{1}{2} \left(\begin{array}{c} -1 \\ \sqrt{3} \end{array}\right),
\]

here $d < 0$ is arbitrary. These deformations require contraction of one of the rods. In the state $S_\alpha$ such contraction is impossible. The impossible deformations form a cone in the eigenstrain space. Similar cones can be established for all homogeneous states because the structure cannot be extended (compressed) in the direction of long (short) rods. Below we describe the two- and three-phase composites of still states. The eigenstrain of any still state composite is a convex combination of eigenstrains of its phases, therefore, the two-phase composites correspond to edges of $\mathcal{D}$ and two-phase composites – to its phases. Set $\mathcal{D}$ is a polyhedron.

We also describe a two-dimensional set $\mathcal{H}_L$ of eigenvalues $(\Lambda_A, \Lambda_B)$ of eigenstrains. This set is also bounded and symmetric (includes pairs $(\Lambda_A, \Lambda_B)$ and $(\Lambda_B, \Lambda_A)$). The eigenstrains of the homogeneous states correspond to vertices of $\mathcal{H}_L$, and the differently oriented homogeneous still states $S_\gamma$, $S'_\gamma$ and $S''_\gamma$ (and $S_\delta$, $S'_\delta$ and $S''_\delta$) correspond to the same pairs of eigenvalues – vertices of $\mathcal{H}_L$. However, the eigenvalues of a two-phase composite are convex combinations of eigenvalues of components only if their vertices are coaxial tensors (the eigen-axes are codirected). Otherwise, they are located along a curve that join the vertices. Therefore, $\mathcal{H}_L$ is a symmetric curvilinear hexagon.
3.3. Laminate structures. There are still states composed from several basis homogeneous states. We call two still states $E_y$ and $E_z$ are called laminate compatible if lengths of their rods along the layers are the same. In terms of eigenstrains, the states are laminate compatible if a vector $\tau$ (the tangent) exists such that

$$\tau^T (E_y - E_z) \tau = 0$$  \hspace{1cm} (3.9)$$

that is, matrices $E_y$ and $E_z$ have the same tangent component. All pairs of homogeneous states except $S_\alpha$ and $S_\beta$ can be laminated together, along the side of a common self-strain, if the states are allowed to rotate freely. We also remark that threads in laminate-compatible structures may possess kinks, as in Figure 4. The continuity of the angles is not required for compatibility of a lattice, in contrast with continuum mechanics.

3.3.1. Structures from an isotropic and anisotropic phases. States $S_\alpha$ and $S_\delta$ are laminate compatible. The eigenstrains of mixture are

$$E_{\alpha\delta} = \mu E_\alpha + (1 - \mu) E_\delta, \hspace{1cm} \mu = \frac{k_\alpha}{k_\alpha + k_\delta}$$

where $\mu$ is the volume fraction of phase $S_\alpha$, $k_\alpha$ is the thickness of the layer, and $k_\delta$ is the thickness of the layer of phase $S_\delta$. The laminate is shown in Figure 3.4.

In addition, the eigenstrains $E_\alpha$ and $E_\beta$ are isotropic and therefore their eigenvectors are codirected with eigenvalues of any phase they are mixed with. A structure of one of them and another phase leads to average of eigenvalues $\lambda_1, \lambda_2$ of eigenstrain. The eigenvalues $\lambda^x_{\alpha\delta}, \lambda^y_{\alpha\delta}$ of $E_{\alpha\delta}$ are located at the straight line between eigenvalues of $E_\alpha$ and $E_\delta$.

$$\lambda^x_{\alpha\delta} = \mu \lambda^x_\alpha + (1 - \mu) \lambda^x_\delta, \hspace{1cm} \lambda^y_{\alpha\delta} = \mu \lambda^y_\alpha + (1 - \mu) \lambda^y_\delta$$

Similar laminates can be built for $E_{\alpha\gamma}$, $E_{\beta\delta}$, $E_{\gamma\beta}$, and $E_{\delta\gamma}$. The eigenvalues of these structures span the intervals between the eigenvalues of the homogeneous states.

3.3.2. Laminates from two anisotropic phases $S_\delta$ and $S_\gamma$. Consider the compatibility of two anisotropic states $S_\delta$. Each of them consists of "long" and "short" rods. Assume for definiteness, that tangent $\tau$ to laminates corresponds to the line of "short" links. This line is an axis of symmetry for the state $E_\gamma$ that consists of two families of long and one family of short rods. In orthogonal coordinates $\tau, \nu$, the eigenstrain $E_\gamma$ is diagonal, its eigenvectors are codirected with $\tau$ and $\nu$. 

![Fig. 3.2. Examples of laminates](image)
The state $E_\delta$ (that consists of two families of short and one family of long rods) can be made compatible with $E_\gamma$ if it is rotated so that its $\tau\tau$ line consists of short rods. In other words, tensor of eigenstrain $E_\delta$ must be rotated to a proper angle so that its tangent entry becomes compatible with $E_\gamma$, $|E_\delta|_{\tau\tau} = 1$, see compatibility conditions (3.3). The rotated matrix is computed from the additional conditions that trace and determinant are independent of rotation. The rotated eigenstrain becomes

$$E_{\delta(\text{rotated})} = \begin{pmatrix} \frac{2}{\sqrt{3}} \sin(b) + 2 \cos(b) - 1 & \rho \\ \rho & 1 \end{pmatrix}$$

where $\rho^2 = \frac{1}{3}(2\cos(b) - 1)(2\sin(b)\sqrt{3} - 3)$. Eigenstrain $E_{\delta\gamma}$ of the mixture is still an average of those in the phases according to (3.1).

$$\mu E_{\delta(\text{rotated})} + (1 - \mu) E_{\gamma} = \begin{pmatrix} \mu \left( \frac{2}{\sqrt{3}} \sin(b) + 2 \cos(b) - 1 \right) + (1 - \mu) \tan\gamma \mu \rho \\ \mu \rho \end{pmatrix}$$

where $\mu$ is the volume fraction, as before.

Eigenvalues $\Lambda_{\delta\gamma}$ of a composite $E_{\delta\gamma}$ are parametrized by $\mu \in [0, 1]$. They depend on $\mu$ nonlinearly. In the eigenvalue plane, they are located at an arch that spans $\Gamma_{\delta}$ and $\Lambda_{\gamma}$, not at an interval between them.

3.3.3. Two-phase twins. Besides these structures, there are two-phase twins: the laminates of differently oriented phase $E_\delta$ or $E_\gamma$. These structures preserve the trace of $E_\delta$ and $E_\gamma$, respectively, and their eigenstrain is as follows

$$E_{\delta(\text{twin})} = \begin{pmatrix} \frac{2}{\sqrt{3}} \sin(b) + 2 \cos(b) - 1 & (1 - 2\mu)\rho \\ (1 - 2\mu)\rho & 1 \end{pmatrix}$$

where $\mu = k/N$, $k \in [-N, N]$. One could check that the eigenvalues of these twins are never isotropic, Figure 3.2(b).

Together, all listed structures form a region $\Omega$ in the plane of eigenvalues of $E$. The boundary has three components corresponding to structures $S_{\alpha\delta}$ (an interval), $S_{\delta\gamma}$ (an arch), and $S_{\gamma\beta}$ (an interval) and four vertices, see Figure 3.3.

3.3.4. Three-phase laminates. There exist three-phase still structures with eigenstrains that densely cover the domain $\Omega$ bounded by the eigenstrains of two-phase structures. Curvilinear triangular domain $\alpha, \delta, \gamma$ is covered with laminates from three phases, obtained by laminating the isotropic phase $S_\alpha$ with the laminate $S_{\delta\gamma}$. The tangents to laminates correspond to the sides of unit lengths in all three components. The eigenstrain $E_{\alpha\delta\gamma}$ is

$$E_{\alpha\delta\gamma} = \nu E_{\alpha} + \mu E_{\delta(\text{rotated})} + (1 - \mu - \nu) E_{\gamma}$$

where

$$\mu = \frac{k}{N}, \quad k = 0, \ldots, N, \quad \nu = \frac{p}{N}, \quad p = 0, \ldots, N - k,$$

The corresponding eigenvalue pairs cover the part of $\Omega$ that lies outside the rectangle, three of whose vertices are $\alpha, \eta, \beta$ in Figure 3.3.
3.4. Hexagon-triangles-strips. The central triangle $\alpha, \eta, \beta$ can be covered by mosaic assembly from elements of $S_\alpha, S_\beta$ and differently oriented elements $S_\delta, S'_\delta, S''_\delta$ shown in Figure 3.4. Notice that $S_\alpha$ and $S_\beta$ are incompatible, and the assembly uses anisotropic elements $S_\delta$ between them. The assembly is as follows.

1. First, three parallelograms from $S_\delta, S'_\delta$ and $S''_\delta$ are formed of the sizes $k, n_1$, $k, n_2$, and $k, n_3$, respectively. The links of the unit length form the base of each parallelogram and the links of the length $a$ form the sides (Figure 3.4 (a)).

2. Second, these parallelograms are joined by the edges with the equilateral triangle with the side $ka$ in between, forming the structure shaped as Y (Figure 3.4 (b)).

3. Third, the obtained figure is copied and translated. The copies are joined by edges, leaving empty triangles and hexagons with the parallel sides of $n_1, n_2$ and $n_3$ unit-length elements.

4. Finally, the triangular parts are filled with $S_\delta$ and the hexagon parts with $S_\alpha$. The obtained periodic assembly covers the whole plane (Figure 3.4 (c)).

The obtained periodic assembly covers the whole plane.

The periodicity element consists of

- three parallelograms from $S_\delta, S'_\delta$ $S''_\delta$ respectively, of $2n_1k, 2n_2k$, and $2n_3k$ triangles each,
- two triangles composed of $S_\gamma$ of $k(k + 1)$ triangles each.
- one hexagon containing $S_\alpha$ of $2(n_1n_2 + n_2n_3 + n_3n_1)$ triangles.

Thus the element consists of

\[ N = k(k + 1 + n_1 + n_2 + n_3) + n_1n_2 + n_2n_3 + n_3n_1 \]

pairs of triangles.

The eigenstrain $E_{\text{hex}}$ of the assembly is a sum of eigenstrains of its pieces,

\[ E_{\text{hex}} = \frac{D}{N} \quad (3.11) \]
where

\[ D = k(k + 1)E_a + n_1 kE_\delta + n_2 k'E_\delta' + n_3 k'E_\delta'' + (n_1 n_2 + n_2 n_3 + n_3 n_1)E_\beta \quad (3.12) \]

\[ E_{\text{hex}} = \frac{k(k + 1)E_a + n_1 kE_\delta + n_2 kE'_\delta + n_3 kE''_\delta + (n_1 n_2 + n_2 n_3 + n_3 n_1)E_\beta}{k(k + 1 + n_1 + n_2 + n_3) + n_1 n_2 + n_2 n_3 + n_3 n_1} \]

3.4.1. Asymptotics. In particular, \( E_{\text{hex}} \) is isotropic if \( n_1 = n_2 = n_3 = n \). Then

\[ E_{\text{hex}} = \epsilon_{is} I, \]

\[ \epsilon_{is} = \frac{k(k + 1) + \frac{3}{2} n k \text{ Tr}(E_\delta) + 3n^2 a}{k(k + 1) + 3n + 3n^2} \quad (3.13) \]
When $n \to \infty$ and $\frac{k}{n}$ varies in $[0, \infty]$, the isotropic deformation $\epsilon_{i\delta}$ varies from one to $a$.

The described structures degenerate into two-phase structures $E_{\delta\beta}$ and $E_{\alpha\delta}$. If $n_1 \gg n_2, n_3, k$, then

$$E_{\text{hex}} \to \frac{kE_{\delta} + (n_2 + n_3)E_{\beta}}{k + n_2 + n_2 + n_3} = E_{\delta\beta}$$

If $n_2 \ll n_1, k$ and $n_3 \ll n_1, k$, we have

$$E_{\text{hex}} \to \frac{k(k + 1)E_{\alpha} + n_1 kE_{\delta}}{k(k + 1) + n_1 k} = E_{\alpha\delta}$$

3.5. Covering of the whole domain $\Omega$. Together, the hexagonal and three-phase-laminate structures densely fill the whole curved hexagon of eigenvalues of still states. The two types of covers are met on the line $\alpha, \gamma$ when $k_{h} \to 1$ in hexagons and $\mu \to 0$ in three-phase laminates. $\Lambda_{x}\Lambda_{y}\Lambda_{z}$. The covering is non-unique: For instance, one can replace $S_{\alpha}$ with $S_{\beta}$ and $S_{\delta}$ with $S_{\gamma}$ everywhere (complementary or dual replacement) and obtain another complete coverage. The density of the coverage is easy to estimate through the explicitly given parameters $k, p$. The distance from an eigenstrain of a discrete $N \times N$ periodic structure to the arbitrary point of the domain is of the order of $N^{-1}$, as follows from the above explicit formulas.

4. Small deformations.

4.1. Linearized elongations. In the remainder of this paper we consider the case of small deformations which means that rod elongations are linearized near the reference configuration. The linearized relative elongation $\kappa_{ij}$ of the rod $(i, j)$ can be written as

$$\kappa_{ij} = q_{ij} \cdot \frac{u_j - u_i}{l}, \quad (4.1)$$

where $q_{ij}$ is the unit direction from node $i$ to node $j$, $u_i$ and $u_j$ are the displacements of nodes $i$ and $j$, respectively.

Consider a triangular periodic network of $N$ nodes, where $N$ is finite, even though it can be arbitrary large. We also suppose that the nodes are arranged into a hexagonal shape containing $n$ nodes along a side (see Figure 5.1 for an example).

Factoring out $l^2$ in (2.1) and using nodal displacements instead of the rod length, we write the energy of a rod $(i, j)$ as

$$w(\kappa_{ij}) = \frac{C l^2}{2} \min \{ \kappa_{ij}^2, (\kappa_{ij} - s)^2 \}, \quad (4.2)$$

where, $s > 0$ is a dimensionless parameter characterizing the critical relative elongation. The graph of the dependence $w(\kappa_{ij})$ is shown on figure 4.1 (Compare with (2.1), where $x$ denotes the actual rod length).

The displacements $u_k$ in the equilibrium state are found by minimizing the total energy of the network:

$$W(u_1, \ldots, u_N) = \sum_{(i,j)} w(\kappa_{ij}) = \sum_{(i,j)} w(q_{ij} \cdot \frac{u_j - u_i}{l})$$
4.2. **Average strain.** An effective homogeneous deformation state of the network is characterized by an average (linearized) strain tensor $\mathbf{E}$. In this section we obtain a formula for the average strain in terms of elongations $\kappa_{ij}$.

Denote the number of nodes in the network by $N$. Given a set of displacements of nodes $u_1, \ldots, u_N$ we define a continuum deformation $\mathbf{u}(x)$ that coincides with $\mathbf{u}_i$ at each $x_i$. This can be done by interpolating. In the present case, the lattice forms a triangulation of the physical domain $\Omega$. Therefore it is easy to construct a piecewise linear interpolant, using finite elements. After this is done we obtain a continuous function $\mathbf{u}(x), x \in \Omega$ such that

- $\mathbf{u}(x_i) = \mathbf{u}_i$ for $i = 1, \ldots, N$ and $x_i$ being the position of node $i$;
- $\mathbf{u}(x)$ is linear on every elementary triangle of the lattice.

To $\mathbf{u}(x)$ we associate strain tensor $\mathbf{\varepsilon}$:

$$\mathbf{\varepsilon} = \frac{\nabla \mathbf{u} + (\nabla \mathbf{u})^T}{2}$$

Average strain $\mathbf{E}$ is defined by the formula

$$\mathbf{E} = \frac{1}{|\Omega|} \int_{\Omega} \mathbf{\varepsilon}(\mathbf{x}) d\mathbf{x}, \quad (4.3)$$

where $\Omega$ is the domain of $\mathbf{u}$ and $|\Omega|$ is

$$|\Omega| = \int_{\Omega} dx$$

There exist formulas for $\mathbf{E}$ in terms of displacements $\mathbf{u}_i$ (see [4], [9]). However, for our purposes it is better to work with the relative elongations $\kappa_{ij}$ because we would often prescribe elongations, rather than displacements. To use standard formulas one would have to solve the system (5.1) (see the next section) that relates elongations and displacements. We choose a more direct route that involves relating three average elongations along the lattice directions with the three independent components of the average strain.

Rewrite (4.3) as

$$\mathbf{E} = \frac{1}{|\Omega|} \int_{\Omega} \mathbf{\varepsilon}(\mathbf{x}) d\mathbf{x} = \frac{1}{|T|} \sum_{\Delta \in \mathbf{T}} \mathbf{\varepsilon}_{\Delta},$$
where $T$ is the set of elementary triangles of the network, $|T|$ is the number of elementary triangles in the network, and $\varepsilon_\Delta$ is the strain tensor on elementary triangle $\Delta$:

$$
\varepsilon_\Delta = \frac{\nabla u_\Delta + (\nabla u_\Delta)^T}{2} = \frac{1}{|\Omega_\Delta|} \int_{\Omega_\Delta} \varepsilon(x) dx
$$

Since the average strain tensor $E$ is a $2 \times 2$ symmetric matrix, it has three independent components:

$$
E = \begin{bmatrix}
a & b \\
b & c
\end{bmatrix}
$$

Consider three lattice direction vectors $q_1$, $q_2$, and $q_3$ (see figure 4.2(a)):

$$
q_1 = \frac{1}{2} \begin{bmatrix} 2 \\ 0 \end{bmatrix}, \quad q_2 = \frac{1}{2} \begin{bmatrix} 1 \\ \sqrt{3} \end{bmatrix}, \quad q_3 = \frac{1}{2} \begin{bmatrix} -1 \\ \sqrt{3} \end{bmatrix}.
$$

We can find the components of $E$ using the values $q_r \cdot (E q_r)$, $r = 1, 2, 3$. Hence, consider

$$
q_r \cdot (E q_r) = \frac{1}{|T|} \sum_{\Delta \in T} q_r \cdot (\varepsilon_\Delta q_r) = \frac{1}{|T|} \sum_{\Delta \in T} q_r \cdot (\nabla u_\Delta q_r).
$$

Using linearity of $u$ on triangle $\Delta$ for edge $(i, j)$ of the triangle we find

$$
q_{ij} \cdot (\varepsilon_\Delta q_{ij}) = q_{ij} \cdot (\nabla u_\Delta q_{ij}) = q_{ij} \cdot (\frac{u_j - u_i}{l}) = \kappa_{ij}
$$

Given a triangle $\Delta$ with vertices $h, i, j$ (see figure 4.2(b)) define vector $k_\Delta \in \mathbb{R}^3$ as:

$$
k_\Delta = \begin{bmatrix}
\kappa_{hj} \\
\kappa_{hi} \\
\kappa_{ij}
\end{bmatrix}
$$
Using this notation we rewrite (4.4) as a linear system for finding the components $a, b, c$ of $E$ given elongations $\kappa_{ij}$:

$$Q \begin{bmatrix} a \\ b \\ c \end{bmatrix} = \frac{1}{|T|} \sum_{\Delta \in T} k_{\Delta},$$

(4.5)

where the matrix $Q$ is given by

$$Q = \begin{bmatrix} q^2_{1,1} & 2q_{1,1}q_{1,2} & q^2_{1,2} \\ q^2_{2,1} & 2q_{2,1}q_{2,2} & q^2_{2,2} \\ q^2_{3,1} & 2q_{3,1}q_{3,2} & q^2_{3,2} \end{bmatrix} = \begin{bmatrix} 1 & 0 & 0 \\ \frac{1}{\sqrt{3}} & \frac{1}{\sqrt{3}} & \frac{1}{2} \\ -\frac{1}{\sqrt{3}} & \frac{1}{1} & \frac{1}{2} \end{bmatrix}$$

(4.6)

and function $m: \mathbb{R}^3 \to \mathbb{R}^{2 \times 2}$ by

$$m(x) = x_1 \begin{bmatrix} 1 & 0 \\ 0 & 0 \end{bmatrix} + x_2 \begin{bmatrix} 0 & 1 \\ 1 & 0 \end{bmatrix} + x_3 \begin{bmatrix} 0 & 0 \\ 0 & 1 \end{bmatrix}$$

Function $m$ is simply a linear 1-1 mapping from the space of three dimensional vectors to the space of $2 \times 2$ symmetric matrices.

Solving (4.5) we find average strain $E$:

$$E = \frac{1}{|T|} m \left( Q^{-1} \sum_{\Delta \in T} k_{\Delta} \right),$$

(4.7)

where

$$Q^{-1} = \begin{bmatrix} 1 & 0 & 0 \\ 0 & \frac{2}{3} & -\frac{1}{3} \\ -\frac{1}{3} & \frac{1}{3} & \frac{1}{3} \end{bmatrix}$$

Further, we will rewrite the sum over triangles in (4.7) as the sum over edges (this is just the change in indexing). Denote all edges of the array by $E$, all boundary edges of the array by $E_B$ and all non-boundary edges of the array by $E_I$ ($E_I = E \setminus E_B$). Then

$$\sum_{\Delta \in T} k_{\Delta} = 2 \sum_{e \in \mathcal{E}} k_e - \sum_{e \in \mathcal{E}_B} k_e,$$

where vector $k_e \in \mathbb{R}^3$ has the components defined by

$$(k_e)_r = \begin{cases} \kappa_e, & \text{if edge } e \text{ is parallel to } q_r \\ 0, & \text{else} \end{cases}, \quad r = 1, 2, 3$$

Next, equation (4.7) can be rewritten as

$$E = \frac{2}{|T|} m \left( Q^{-1} \sum_{e \in \mathcal{E}} k_e \right) - \frac{1}{|T|} m \left( Q^{-1} \sum_{e \in \mathcal{E}_B} k_e \right).$$

Denote the average elongation by $\bar{k}$:

$$\bar{k} = \frac{1}{|E|} \sum_{e \in E} k_e.$$

(4.8)
Then we rewrite (4.7) and obtain the desired formula for the average strain in terms of average elongations:

\[ E = 2 \frac{|E|}{|T|} m(Q^{-1}\bar{k}) - \frac{1}{|T|} m \left( Q^{-1} \sum_{e \in E_B} k_e \right). \] (4.9)

This formula is used extensively in the remainder of the paper. The first term in the right hand side contains contributions from all interior edges of the lattice, while the second term contains the contributions from the boundary edges only. As the number of nodes increases, the second term becomes a small perturbation of the first. This becomes clear from the following estimate

\[ \left\| m \left( \frac{1}{|T|} Q^{-1} \sum_{e \in E_B} k_e \right) \right\|_F \leq \sqrt{2} \frac{|E_B|}{|T|} \|Q^{-1}\|_2 \max_{e \in E_B} ||k_e||_2. \] (4.10)

Here \( \| \cdot \|_F \) denotes the Frobenius norm of a matrix, and \( \| \cdot \|_2 \) is the Euclidean norm. Note that the ratio

\[ \frac{|E_B|}{|T|} \]

does not approach zero as the number of nodes approaches infinity.

5. Compatibility conditions. In the next two sections, we study the linearized version of the problem of finding still states. The linearized formulation is as follows.

Equations (4.1) relating node displacements with link elongations can be concisely written as

\[ RU = \kappa, \] (5.1)

where \( U \) is a tuple of displacements \( u_i \), say \( (u_1, \ldots, u_N) \), and \( \kappa \) is a tuple of \( \kappa_{ij} \). The number of unknowns in (5.1) is \( 2N \) (since each \( u_i \) has two independent components).

The number of equations is larger, approximately \( 3N \) (see Lemma 5.1 below for the exact count). Therefore, we have an overdetermined system (the number of unknowns is less than the number of equations). As a consequence, (5.1) may be unsolvable for some \( \kappa \). The values of \( \kappa \) for which (5.1) is solvable are called admissible.

5.1. Compatibility conditions characterize range of \( R \). In this section we show that compatibility conditions (2.6) are necessary and sufficient for solvability of (5.1). Necessity is clear from the derivation of (2.6), so we focus on sufficiency.

Suppose that we have the hexagonal array with a side of \( n \) points (see figure 5.1).

For every hexagon consisting of 6 vertices we formulate the condition (2.6): the sum of elongations of the outer edges equals to the sum of elongations of the inner edges. For example, for figure 5.1 the equation will be

\[ \kappa_{12} + \kappa_{23} + \ldots + \kappa_{56} + \kappa_{61} = \kappa_{71} + \kappa_{72} + \ldots + \kappa_{75} + \kappa_{76} \]

The collection of all such equations will be called hexagonal equations.

Next, we need a preliminary result giving the exact count of the number of nodes, edges and hexagonal equations. The following lemma is used later to carry out proofs by induction.

**Lemma 5.1.** The total number of nodes \( N \), the number of edges \( E \), and the number of hexagonal equations \( M \) depend on \( n \) as follows.

\[ N(n) = 3n^2 - 3n + 1, \quad E(n) = 9n^2 - 15n + 6, \quad M(n) = 3n^2 - 9n + 7. \]
Fig. 5.1. Example of the network with $N = 19$, $n = 3$. A hexagon that forms a hexagonal equations shown in bold

**Proof.** We will prove the lemma using mathematical induction.

For $n = 2$ we have a hexagon that has 7 vertices, 12 edges, and 1 hexagonal equation. Indeed,

$$N(2) = 3 \cdot 2^2 - 3 \cdot 2 + 1 = 7, \quad E(2) = 9 \cdot 2^2 - 15 \cdot 2 + 6 = 12, \quad M(2) = 3 \cdot 2^2 - 9 \cdot 2 + 7 = 1$$

Thus, the formulas are true for $n = 2$.

Suppose that given formulas are true for $n = k$. Let’s prove that they are true for $n = k + 1$.

When we increase $n = k$ by 1 we “wrap” the array by a layer of points. Therefore, we add $6k$ vertices, we allow $6(k - 1)$ new hexagonal equations, and add $6(3k - 1)$ edges. Thus, the number of vertices $N^*$ for the size $n = k + 1$ is:

$$N^* = N(k) + 6k = 3k^2 - 3k + 1 + 6k = 3(k + 1 - 1)^2 + 3k + 1$$

$$= 3(k + 1)^2 - 6(k + 1) + 3 + 3k + 1 = 3(k + 1)^2 - 3(k + 1) + 1,$$

the number of equations $M^*$ is

$$M^* = M(k) + 6(k - 1) = 3k^2 - 9k + 7 + 6(k - 1) = 3(k + 1 - 1)^2 - 3k + 1$$

$$= 3(k + 1)^2 - 6(k + 1) + 3 - 3k + 1 = 3(k + 1)^2 - 9(k + 1) + 7,$$

and the number of edges $E^*$ is

$$E^* = E(k) + 6(3k - 1) = 9k^2 - 15k + 6 + 18k - 6 = 9(k + 1 - 1)^2 + 3k$$

$$= 9(k + 1)^2 - 18(k + 1) + 9 + 3k = 9(k + 1)^2 - 15(k + 1) + 6,$$

Since $N^* = N(k + 1)$, $M^* = M(k + 1)$, and $E^* = E(k + 1)$, the formulas are true. \[\blacksquare\]

Observe that

$$2N - 3 = E - M.$$

We know that for the mapping $R : \mathbb{R}^{2N} \to \mathbb{R}^E$ we have $\dim(\ker R) = 3$. This follows from the well known results on graph rigidity, in particular theorems on the first-order rigidity of triangulations (see e.g. [3] and references therein). Thus,

$$\dim(\text{im} \, R) = 2N - 3$$
The hexagonal equations can be concisely written in the form

$$Z\kappa = 0,$$  \hspace{1cm} (5.2)

where $Z$ is a matrix, $Z : \mathbb{R}^E \to \mathbb{R}^M$; $i$-th row of this matrix corresponds to $i$-th equation.

**Lemma 5.2.** Matrix $Z$ has full rank $M$.

**Proof.** We will restate the lemma in the form: all hexagonal equations are linearly independent.

We will prove the lemma in its new form using mathematical induction on $n$. For $n = 2$ we have $M = 1$. One equation is linearly independent.

Suppose, the lemma is true for $n = k$. Let’s prove it for $n = k + 1$.

Increasing $n$ by 1 we wrap a hexagon of side size $k$ with a layer of points. Doing so we add some number of hexagonal equations to those that we had in the original hexagon. The hexagonal equations of the original hexagon are linearly independent. Each new hexagonal equation contains an edge that is present in no other equation. Thus, all equations are linearly independent.

Since matrix $Z$ has rank $M$, we find $\dim(\ker Z) = E - M$. Thus,

$$\dim(\text{im } R) = \dim(\ker Z)$$  \hspace{1cm} (5.3)

Necessity of hexagonal equations implies that each admissible $\kappa$ satisfies them. This fact can be stated as follows: for any $\kappa \in \text{im } R$ we have $Z\kappa = 0$. This in turn implies

$$\text{im } R \subset \ker Z.$$  \hspace{1cm} (5.4)

**Theorem 5.3.** For any vector $\kappa \in \mathbb{R}^E$ satisfying $Z\kappa = 0$ there is a unique (up to translation and rotation) vector $u \in \mathbb{R}^{2N}$ satisfying $Ru = \kappa$.

**Proof.** The conditions (5.3) and (5.4) imply $\text{im } R = \ker Z$.

Thus, for any $\kappa \in \mathbb{R}^E$ we can find a vector $u \in \mathbb{R}^{2N}$. If we neglect rigid rotations and translations of the whole lattice (they form the null space of $R$ by the well known results on rigidity of triangulations, [3] and references therein), then this vector is unique. \[\square\]

**6. Still states and small deformation eigenstrains.** In the small deformation case, a still state is a collection of nodal displacements $u_i$ satisfying equations (5.1) with the right hand side $\kappa$ of special form. The components $\kappa_{ij}$ can take only two values: 0 and $s/2$, where $s/2$ is the critical elongation from (4.2). If $\kappa_{ij} = 0$ we call the corresponding edge $(i, j)$ short, otherwise the edge is called long. To construct a still state, one could choose a length for each edge (long or short). The resulting configuration is accepted if the resulting triangles form a tessellation. Otherwise the configuration is rejected. In the case of small deformations, an admissible elongation vector $\kappa$ should lie in the range of the matrix $R$ from (5.1). Equivalently, such $\kappa$ must be a solution of the hexagonal equations (5.2).

Denote the set of all still states by $U$. To characterize $U$, one needs to solve the following problem.

Given $\kappa$ with $\kappa_{ij} \in \{0, s\}$, find $U$ realizing that $\kappa$, that is, find $U$ such that $RU = \kappa$. 19
This is possible only if $\kappa$ satisfies hexagonal equations from Section 5. Complete characterization of $\mathcal{U}$ seems very difficult, and is not addressed in this paper. The reason for the difficulty is the lack of a convenient structure. It is easily checked that $\mathcal{U}$ is not a linear space, or even a convex set. Therefore, a description of this set cannot be obtained by linear algebra methods.

6.1. An algorithm of adding still states. In this section we characterize the set of strains that can be well approximated by the average strains of still states. An average strain of a still state can be conveniently described by a triple of concentrations of long edges parallel to the lattice directions. In this section we propose an explicit construction that furnishes a large number of still states. Concentrations of these states densely fill a certain unit cube in the concentration space. Our construction is based on two observations. First, one needs a binary operation that produces new still states by combining two already known still states. Second, one needs a simple "building block", that is, a still state that could be easily combined with its slightly modified (e.g. translated and rotated) replicas. In view of the linear structure of $\text{im } R$, the convenient operation is summation. However, summation of two still states does not always produce a still state. An additional condition that ensures that the sum of still states remains a still state is non-overlapping of the long edges (see Lemma 6.1 below): if a particular edge is long in the first state, then it should be short in the second state, and vice versa.

From this we deduce that the building block should have low concentrations of long edges, and the placement of these edges should be localized as highly as possible. If both of these requirements are satisfied, one can add together shifted copies of the building block to generate new still states with higher concentrations.

Lemma 6.1. The sum $\kappa_1 + \kappa_2$ of two still states $\kappa_1$ and $\kappa_2$ is a still state provided $\kappa_1 \cdot \kappa_2 = 0$.

Proof. Since $\kappa_1$ and $\kappa_2$ are still states, their components satisfy

$$(\kappa_1)_{ij} \in \{0, s\}, \quad (\kappa_2)_{ij} \in \{0, s\}$$

for all connected nodes $i$ and $j$. Moreover, the condition of orthogonality $\kappa_1 \cdot \kappa_2 = 0$ says that for each certain pair $(i, j)$ the values $(\kappa_1)_{ij}$ and $(\kappa_2)_{ij}$ cannot equal $s$ simultaneously. Thus,

$$(\kappa_1)_{ij} + (\kappa_2)_{ij} \in \{0, s\}$$

Also, since $\kappa_1$ and $\kappa_2$ are still states, we have

$$Z\kappa_1 = 0, \quad Z\kappa_2 = 0,$$

where $Z$ is the matrix of the hexagonal system (5.2). By linearity $Z$,

$$Z(\kappa_1 + \kappa_2) = Z\kappa_1 + Z\kappa_2 = 0$$

Since the necessary conditions hold, the sum $\kappa_1 + \kappa_2$ is a still state.

Notice that Lemma 6.1 states that sum

$$\kappa_1 + \kappa_2 + \ldots + \kappa_m$$

is a still state, provided $\kappa_i, \ i = 1, \ldots, m$ have non-overlapping long edges.
For a still state in (4.10) we have \(|k_2| \leq s\). Thus

\[
\left\| \text{m} \left( \frac{1}{T} Q^{-1} \sum_{e \in E_B} k_e \right) \right\|_F \leq \sqrt{2|E_B|/|T|} ||Q^{-1}||_2 s \tag{6.1}
\]

**Theorem 6.2.** For any \(\alpha_i \in [0, 1], i = 1, 2, 3\), and any \(n \geq 3\) there exists a still state \(\kappa^*\) such that its concentrations \(\alpha_i^*, i = 1, 2, 3\), satisfy

\[|\alpha_i - \alpha_i^*| < \frac{1}{n}\]

**Proof.** To prove the theorem we will use special still states called *stripes*. These states have \(k_{ij} = 0\) everywhere except for one stripe that has non-zero elements in one direction (figure 6.1). We can see that these stripes are still states because their components are either 0 or \(s\) and they satisfy the hexagonal equations: each hexagon has either all zero elongations or one non-zero inner elongation and one non-zero outer elongation.

![Diagram of stripes](image)

**Fig. 6.1.** 6.1(a), 6.1(b), and 6.1(c) are “stripe” still states. 6.1(d) is a still state composed of 6.1(a), 6.1(b), and 6.1(c). Solid line corresponds to \(k = s\), dotted line is \(0\).
Observe, that for two different stripes \( \kappa_1 \) and \( \kappa_2 \) we have \( \kappa_1 \cdot \kappa_2 = 0 \). In other words, the long edges of \( \kappa_1 \) do not overlap the long edges of \( \kappa_2 \), and thus \( \kappa_1 + \kappa_2 \) is a still state (figure 6.1(d)).

We have 3 different groups of stripes, \( \mathcal{G}_1 \), \( \mathcal{G}_2 \), and \( \mathcal{G}_3 \), one for each direction: one group of horizontal stripes and two groups of rotated stripes. The number of stripes in each group is \( 2(n - 1) \).

Next, define the vectors \( \mathbf{d}_r \), \( r \in \{1, 2, 3\} \) as follows. The dimension of \( \mathbf{d}_r \) is \( E \) (the number of edges), and the components are equal to one if the corresponding edge is parallel to \( \mathbf{q}_r \) (a lattice direction vector), and equal to zero otherwise. It is easy to see that if \( \kappa \in \mathcal{G}_r \) for some \( r \in \{1, 2, 3\} \) then \( \kappa \cdot \mathbf{d}_r < 2n \), \( r = 1, 2, 3 \), since each “stripe” has no more than \( 2n \) long edges. Also, since each stripe has long edges only in \( r \)-th direction, we get

\[
\kappa \cdot \mathbf{d}_r = \frac{E}{3} \alpha_r, \quad \kappa \cdot \mathbf{d}_t = 0 \text{ for } t \neq r.
\]

This implies

\[
\alpha_r < \frac{6n}{E} = \frac{6n}{9n^2 - 15n + 6} < \frac{1}{n} \text{ for } n \geq 3.
\]

First, we explain the main idea of the proof. Consider \( i \)-th group (stripes of one certain direction). If we add all stripes of this group then we’ll get \( \alpha_i = 1 \). For the still state \( \kappa = 0 \) we have \( \alpha_i = 0 \). Therefore, adding stripes one by one we increase \( \alpha_i \) from 0 to 1 with steps less than \( n^{-1} \). Hence, for any \( x \in [0, 1] \) there exists a step when \( |\alpha_i - x| < n^{-1} \). Finally, since different directions are independent from each other, we can run this algorithm for \( i = 1, 2, 3 \) and at the end add all still states together to get what we need. Next, we present the technical details.

For any subset \( S \) of set \( \mathcal{G}_i \) we will define functions \( \hat{\kappa}(S) \) and \( \hat{\alpha}_j(S) \) by

\[
\hat{\kappa}(S) = \sum_{\kappa \in S} \kappa \quad \text{and} \quad \hat{\alpha}_j(S) = \sum_{\kappa \in S} \frac{3}{E} \kappa \cdot \mathbf{d}_j = \frac{3}{E} \hat{\kappa}(S) \cdot \mathbf{d}_j.
\]

Since \( \hat{\kappa}(\mathcal{G}_i) = \mathbf{d}_i \), we have \( \hat{\alpha}_i(\mathcal{G}_i) = 1 \) and \( \hat{\alpha}_j(\mathcal{G}_i) = 0 \) for \( j \neq i \). Also, we define \( \hat{\kappa}(\emptyset) = 0 \) and \( \hat{\alpha}_j(\emptyset) = 0 \), \( j = 1, 2, 3 \).

From the corollary of lemma [6.1][1] it follows that for any \( S \subset \mathcal{G}_i \) the value of function \( \hat{\kappa}(S) \) is a still state with concentrations \( \hat{\alpha}_j(S) \).

Suppose we are given values \( \alpha_i \in [0, 1], i = 1, 2, 3 \). Let’s define sets \( \mathcal{H}_i \), \( i = 1, 2, 3 \), as

\[
\mathcal{H}_i = \{ S \subset \mathcal{G}_i : \hat{\alpha}_i(S) \leq \alpha_i \}
\]

Having defined \( \mathcal{H}_i \) we define values \( S^*_i, i = 1, 2, 3 \), by

\[
S^*_i = \text{argmax}_{S \in \mathcal{H}_i} \hat{\alpha}_i(S)
\]

Since the sets \( \mathcal{G}_i \) are finite, the sets \( \mathcal{H}_i \) are finite and the maximum exists. Therefore, the definition of \( S^*_i \) is consistent.

Now if for some \( i \in \{1, 2, 3\} \) we have \( \mathcal{G}_i \setminus S^*_i = \emptyset \) then \( S^*_i = \mathcal{G}_i \) and \( \hat{\alpha}(S^*_i) = 1 \).

From the definition of \( S^*_i \) we have

\[
\hat{\alpha}_i(S^*_i) \leq \alpha_i
\]
Since $\alpha_i \leq 1$, we get $\hat{\alpha}_i(S_i^*) = \alpha_i$.

If for some $i \in \{1, 2, 3\}$ we have $G_i \setminus S_i^* \neq \emptyset$ then there exists $k \in G_i \setminus S_i^*$. From the definition of function $\hat{\alpha}$ we get

$$\hat{\alpha}(S_i^* \cup \{k\}) = \hat{\alpha}(S_i^*) + \hat{\alpha}(\{k\}) > \hat{\alpha}(S_i^*)$$

And, since $S_i^*$ has maximal value in $H_i$, we have $S_i^* \cup \{k\} \notin H_i$. Hence,

$$\alpha_i < \hat{\alpha}_i(S_i^* \cup \{k\})$$

Now using

$$\hat{\alpha}_i(S_i^*) \leq \alpha_i < \hat{\alpha}_i(S_i^* \cup \{k\})$$

and

$$\hat{\alpha}_i(S_i^* \cup \{k\}) - \hat{\alpha}_i(S_i^*) = \hat{\alpha}_i(\{k\}) < \frac{1}{n}$$

we get

$$0 \leq \alpha_i^* - \hat{\alpha}_i(S_i^*) < \frac{1}{n}$$

Finally, for the still state $\kappa^*$ defined by

$$\kappa^* = \hat{\kappa}(S_1^* \cup S_2^* \cup S_3^*)$$

we have

$$|\alpha_i^* - \alpha_i| < \frac{1}{n}$$

Let us define a set $D$:

$$D = \{ sm(Q^{-1}x) : x \in \mathbb{R}^3, 0 \leq x_r \leq 1, r = 1, 2, 3\}$$

where $Q^{-1}$ is the inverse of the matrix given by Eq. (4.6). Set $D$ is a convex polygon in the space of $2 \times 2$ symmetric matrices, since it is an image of a cube under a non-singular linear transformation. The components of the elements of $D$ form a set shown on figure 6.2.

**Theorem 6.3.** For any $E \in D$ there exists a still state of the hexagonal array with a side of $n \geq 3$ points such that average strain $E^*$ of this still state satisfies

$$||E^* - E||_F \leq s||Q^{-1}||_2 \frac{8}{n-1}$$

**Proof.** Since $E \in D$, there exists $\alpha \in \mathbb{R}^3, 0 \leq \alpha_r \leq 1$, such that

$$E = sm(Q^{-1} \alpha)$$

Using theorem 6.2 we can claim that for $n \geq 3$ there is a still state with the concentration values $\alpha^*$ such that

$$||\alpha^* - \alpha||_2 \leq \frac{\sqrt{3}}{n}$$
Fig. 6.2. The cube and its image under $Q^{-1}$

Hence, using (4.9) we find
\[
\|E^* - E\|_F = \left\| \frac{2|E|}{|T|} m(Q^{-1}k^*) - \frac{1}{|T|} m(Q^{-1} \sum_{e \in E_B} k_e^*) - E \right\|_F
\]
\[
= \left\| \frac{2s |E|}{3 |T|} m(Q^{-1} \alpha^*) - \frac{1}{|T|} m(Q^{-1} \sum_{e \in E_B} k_e^*) - s m(Q^{-1} \alpha) \right\|_F
\]
\[
\leq \sqrt{2} ||Q^{-1}||_2 \left\| \frac{2s |E|}{3 |T|} \alpha^* - s \alpha \right\|_2 + \left\| m \left( \frac{1}{|T|} Q^{-1} \sum_{e \in E_B} k_e^* \right) \right\|_F
\]

Next,
\[
\left\| \frac{2s |E|}{3 |T|} \alpha^* - s \alpha \right\|_2 \leq s ||\alpha^* - \alpha||_2 + s \left| \frac{2 |E|}{3 |T|} \right| ||\alpha^*||_2
\]
\[
\leq \frac{\sqrt{3}}{n} s + s \left| \frac{2 |E|}{3 |T|} \right| \sqrt{3}
\]

Further, using
\[
|T| = 6(n-1)^2, \quad |E| = 9n^2 - 15n + 6, \quad |E_B| = 6(n-1)
\]
and estimation (6.1) we find
\[
\|E^* - E\|_F \leq \sqrt{2} ||Q^{-1}||_2 \left( \frac{\sqrt{3}}{n} s + s \left| \frac{2 |E|}{3 |T|} \right| \sqrt{3} \right) + \sqrt{2} |E_B| ||Q^{-1}||_2 s
\]
\[
= \sqrt{2} s ||Q^{-1}||_2 \left( \frac{\sqrt{3}}{n} + \sqrt{3} \frac{n-1}{3(n-1)} + \frac{1}{n-1} \right)
\]
\[
\leq s ||Q^{-1}||_2 \frac{8}{n-1}
\]

Let us now discuss the significance of the Theorem 6.3 for linking the microscopic (lattice-level) and macroscopic (continuum) models. First, it is of interest to understand how geometry of $D$ depends on the microstructure. We observe that $D$ is the
image of a unit cube under a linear mapping defined in (6.2). The unit cube and
the mapping \( m \) are both independent of the lattice geometry and particulars of the
microscopic force definition. The two microstructure-dependent quantities in (6.2) are
the relative (non-dimensional) critical elongation \( s \) and the matrix \( Q^{-1} \). The actual
critical elongation is \( sl \), where \( l \) is the equilibrium edge length. Passing to the \( \Gamma \)-limit
requires proper scaling of \( l \) with \( N \), a typical scaling being \( l \sim N^{-1/2} \). In the most
natural setting, \( sl \) should scale as \( l \). If that is the case, then \( s = O(1) \) as \( N \to \infty \).
The set \( D \) in that case is independent of \( N \), and \( s \) is a non-dimensional constant of
the problem that determines susceptibility of the material to phase transition. The
diameter of \( D \) increases linearly with \( s \). The matrix \( Q^{-1} \) determines orientation of \( D \)
in the strain space and provides the explicit dependence of \( D \) on the lattice geometry.
Indeed, by definition (4.6), \( Q \) depends only on the products of components of the
lattice direction vectors \( q_r, r = 1, 2, 3 \).

If \( s = O(1) \), then \( D \) is the "flat bottom" of the macroscopic energy density
because, by Theorem 6.3, any strain in this set can be approximated by a still state
eigenstrain and the error of this approximation vanishes in the limit \( N \to \infty \). Since
the energy of every still state is zero, and \( D \) is fixed as \( N \to \infty \), the conclusion follows.

7. Effective energy of the network. In the previous section, we described
the set \( D \) of all strains that corresponds to zero effective energy. Here, we suggest the
effective energy equation that corresponds to strains outside this set.

The average elastic energy \( W \) of the network without the transition is described
by Cauchy formula [8]

\[
W(\varepsilon) = \frac{2C}{3} \left[ \text{Tr}(\varepsilon^2) - \frac{1}{4} (\text{Tr} \varepsilon)^2 \right],
\]

where \( \varepsilon \) is the average strain of the network. Assume first that the transition state is
fixed in one of the still states \( E \in D \). In other words, assume that no Long-to-Short
and Short-to-Long transition are allowed. Let us denote the total average strain as \( e \).
This strain consists of an eigenstrain \( E \) and elastic strain \( \varepsilon \):

\[
e = E + \varepsilon.
\]

The network is elastically inhomogeneous. However, we will assume for simplicity
that the elastic properties stay constant. We neglect the effect of density variation
in the transition which is reasonable for small elongations \( s \). Under this assumption,
the average elastic energy of the network is

\[
W(\varepsilon) = W(e - E).
\]

Minimizing the energy over all still state strains \( E \), we conclude that, for each fixed
\( n \), the energy density \( J_n \) can be written as follows

\[
J_n(e) = \min_{E \in D} W(e - E) = O(n^{-1}) \quad \text{if} \quad e \in D. \tag{7.1}
\]

Assume now that \( e \not\in D \). The density becomes

\[
J_n(e) = \min_{E \in D} W(e - E) \tag{7.2}
\]

Notice that \( J \) is a convex linear combination of the energies of homogeneous states
that correspond to the vertices of \( D \). This follows from convexity of \( D \) and \( W(\varepsilon) \).
Motivated by this, we suggest a representation for the effective energy density $J$ in the limit $n \to \infty$ (Γ-limit). By the well known argument [5], using density of piecewise linear deformations, it is enough to consider only linear deformations (equivalently, constant effective strains). For these strains,

$$J(e) = \begin{cases} \min_{E_i} W(e - E_i), & \text{if } e \notin D, \\ 0, & \text{if } e \in D. \end{cases} \quad (7.3)$$

Here $E_i$ are the eigenstrains corresponding to the corners of $D$.

8. Conclusions. We studied the discrete model of solid-solid phase transitions. The model consists of a two-dimensional triangular periodic lattice of nonlinear bistable rods. In order to characterize the effective energy of the lattice, we obtained the description of the set of the average eigenstrain tensors corresponding to so called still states, e.g. microstructure deformations that carry no forces. We also proposed several explicit constructions of such states for small and large deformations.

The compatibility conditions for rod elongations were derived. In the small deformation case, these conditions are necessary and sufficient for the existence of a deformation realizing a given set of elongations. To characterize the states with zero effective energy, we proposed a construction of a special class of still states. The eigenstrains of these special patterns densely cover the set $D$ of all eigenstrains corresponding to still states. We showed that the set $D$ for our network is a parallelepiped in 3D space of independent components of the strain tensor. The orientation and side lengths of $D$ were explicitly described in terms of the parameters of the underlying microscopic discrete model: the side lengths are proportional to $s$ and orientation is determined by the components of the lattice direction vectors via the matrix $Q^{-1}$. Components of the inverse matrix $Q$ are defined via components of the direction vectors according to (4.6).

The above results suggest the following properties of the energy density:

- if a strain is inside of $D$, then the energy density is asymptotically close to zero as the number of the lattice nodes approaches infinity;
- If the strain is outside of this set, the energy density is proportional to the square of the distance between the strain and $D$.

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