CARBON-NANOTUBE GEOMETRIES AS OPTIMAL CONFIGURATIONS

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Abstract. The fine geometry of carbon nanotubes is investigated from the viewpoint of Molecular Mechanics. Actual nanotube configurations are characterized as being locally minimizing a given configurational energy, including both two- and three-body contributions. By focusing on so-called zigzag and armchair topologies, we prove that the configurational energy is strictly minimized within specific, one-parameter families of periodic configurations. Such optimal configurations are checked to be stable with respect to a large class of small nonperiodic perturbations and do not coincide with classical rolled-up nor polyhedral geometries.

1. Introduction

Carbon nanotubes are allotropes of carbon consisting in long, cylindrical, hollow structures including as many as $10^7$ atoms [43]. These are covalently bonded, thus sharing a pair of electrons. The corresponding orbitals are so-called $sp^2$-hybridized, inducing indeed bonds at a given atom to form $2\pi/3$ angles, as happens in alkenes [4]. As a result, the local topology of atomic bonds is hexagonal, for each atom has exactly three active bonds, approximately forming $2\pi/3$ angles. Although evidence of carbon filaments has to be traced back at least to the ’50 [34], a new branch of carbon-nanotube chemistry definitely emerged from the early ’90 with [21] and [2, 22], see the review [32]. Since then, carbon nanotubes have been subject of intensive investigation under a variety of different experimental and computational techniques [24, 20].

The unique trait of carbon nanotubes is their amazing aspect-ratio: structures of a few nanometers in diameter can have lengths up to several centimeters, thus bridging molecular and macroscopic scales. In addition, carbon nanotubes show remarkable mechanical strength [26, 42, 45] and, depending on the topology, high electrical conductivity or semiconductivity [25, 33]. Such exceptional properties make carbon nanotubes interesting for the development of innovative technologies. Carbon nanotubes are presently used in the production of composite materials, coatings, microelectronics, energy-storage devices, and biotechnologies [8].

Carbon nanotubes can be intuitively visualized as the result of the rolling-up of graphene, a two-dimensional carbon sheet where atomic bonds form an ideal hexagonal pattern. More precisely, let the hexagonal lattice $\mathcal{H} := \{pa+qb+rc : p,q \in \mathbb{Z}, r = 0,1\}$ with $a = (\sqrt{3},0)$, $b = (\sqrt{3}/2,3/2)$, and $c = (\sqrt{3},1)$ represent graphene. We may prescribe the topology of the atomic bonds of a nanotube by specifying a vector $(\ell,m)$ for $\ell, m \in \mathbb{N}$, $\ell > 2$ and identifying $x \in \mathcal{H}$ with $x + \ell a + mb$, see Figure 1. This identification actually defines the roll-up, an example of a resulting structure being in Figure 5. Nanotubes are called zigzag for $m = 0$, armchair for $m = \ell$, and chiral in all other cases. We concentrate here on zigzag and armchair topologies, leaving the chiral case aside, for it involves additional intricacies.

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The fine geometry of carbon nanotubes is presently still debated and different models are available. In the zigzag and armchair case, the classical rolled-up \([9, 10]\) model is directly inspired by the above idealized identification and assumes that two of the three bond angles at each atom are exactly \(2\pi/3\). An alternative is the polyhedral model \([6, 7]\), which prescribes the three bond angles at each atom to be equal, see also the generalization in \([27]\).

All the mentioned nanotube models are geometrical in nature for they reside on sets of geometrical postulates. The aim of this note is instead to follow a variational approach by investigating carbon-nanotube geometries within the frame of Molecular Mechanics \([1, 28, 35]\). Atom configurations are modeled as a collection of particle positions, to which a configurational energy is associated. This energy is given in terms of classical potentials and takes into account both attractive-repulsive two-body interactions, minimized at some given bond length, and three-body terms favoring \(2\pi/3\) angles between bonds \([40, 41]\). This simplified, phenomenological approach is obviously very far from the quantum-mechanical nature of actual chemical bonding. Still, it has the advantage of being simple, parametrizable, and of delivering the only computationally amenable option as the dimension of the ensemble scales up, as it is the case of carbon nanotubes.

We single out specific carbon-nanotube geometries by focusing on their stability. This concept is central with respect to the process of molecular structuring, since among the many theoretically possible geometries, only those showing some suitable stability can be expected to be realized. This is formalized here by interpreting stability as the minimality of the configurational energy with respect to small perturbations. This approach is fairly classical, at least in the case of infinite crystals, where a number of different stability criteria has been set forth, essentially by prescribing specific set of admissible perturbations \([14, 15]\). The reader is referred to the discrete-to-continuous analysis in \([12, 13]\) and the recent comparison in \([36]\) of such different stability concepts at the continuum level for free-standing graphene. Our approach here is that of possibly considering all small perturbations, where such smallness is exclusively aimed at preserving the local coordination of the atomic bonds. As a consequence, the stability criterion here addressed turns out to be the most restrictive. The reader is referred to \([18, 39]\).
for related stability analysis for other carbon nanostructures, including carbyne, graphene, the fullerene $C_{60}$, and diamond. Moving from the present analysis, a refined analytic discussion of the stability of carbon nanotubes under moderate stretching has been developed in [17].

Under some classical choice of interaction potentials, we investigate first the minimization of the configurational energy in specific one-parameter classes of periodic zigzag and armchair configurations which include the classical rolled-up [10] and the polyhedral [6] model. Here, the dimensionality of the problem scales down and we can prove that a unique optimal configuration exists. Quite remarkably, this optimal configuration do not coincide neither with the classical rolled-up [10] nor with the polyhedral model [6]. Our analysis thus entails that these classical geometrical models do not admit a variational interpretation and that energy minimality induces a different structure.

The stability of such optimal configurations with respect to general small perturbations has been already computationally ascertained in [30]. Our aim here is to provide some analytical justification of this fact. In particular, we prove stability with respect to a large class of small perturbations including tractions and diameter dilations, among many others. Moreover, we propose an ansatz, namely (5), under which stability can be proved for all small perturbations. Numerical evidence for the validity of such assumption is provided.

This path is of course not new to computational chemists, and numerical investigations of carbon-nanotube geometries at the single-atom level are already available [37, 38, 46]. With respect to the current stand the novelty of our contribution is twofold. On the one hand, we investigate the full structure at once, by dropping the periodicity assumption on perturbations. On the other hand, we obtain rigorous analytical results instead of numerical assessments. Note that a menagerie of different possible choices for energy terms has been implemented in Computational Chemistry codes [3, 5, 19, 31, 44]. By sorting out a set of minimal hypothesis of the configurational energy we are hence contributing to cross-validate these choices in view of their capability of describing stable carbon-nanotube geometries.

The plan of the paper is the following. We devote Section 2 to the specification of the mathematical setting and the statement of the main stability result. The specific geometry of periodic zigzag and armchair configurations is then detailed in Section 3. We analyze energy minimization at single atoms in Section 4 and identify optimal configurations, whose stability is then proved in Section 5 by means of convexity and monotonicity arguments. Eventually, we devote Section 6 to a numerical assessment of ansatz (5).

2. Main result

In this section we introduce the mathematical setting and state the main result of the paper, namely Theorem 2.1. Nanotubes are described as collections of distinct points in $\mathbb{R}^3$ of the form

$$\mathcal{C} := C_n + LZ e_3.$$  

The set $C_n := \{x_1, \ldots, x_n\}$ is called the $n$-cell of the nanotube $\mathcal{C}$, where the points $x_i \in \mathbb{R}^3$ are such that $x_i \cdot e_3 \in [0, L)$. The possibly very large $L > 0$ acts as period of the nanotube along its axis $e_3 := (0, 0, 1)$ and is here introduced for the sake of definiteness only. In particular, such periodicity is immaterial with respect to the fine, local geometry of the nanotube which is here analyzed and it is introduced just to prevent the need of discussing the boundary effects due to finite capping. Note that nanotubes can be as long as $10^6$ times their diameter, so that $L$ would
ideally correspond to such length. In the following, we systematically identify the nanotube $C$ with the pair $(C_n, L)$.

To each nanotube $C$ we associate a configurational energy $E(C)$. This is modeled on the specific phenomenology of $sp^2$ covalent bonding of carbon atoms [4]. In particular, bonds of a specific length (here normalized to 1) and bond angles of $2\pi/3$ and $4\pi/3$ are favored. Correspondingly, the energy $E$ features the sum of a two- and a three-body contribution and reads

$$E(C) = E(C_n, L) := \frac{1}{2} \sum_{(i,j) \in \mathcal{N}} v_2(|x_i - x_j|_L) + \frac{1}{2} \sum_{(i,j,k) \in \mathcal{T}} v_3(\omega_{ijk}),$$

the two terms being described in the following.

The two-body potential $v_2 : \mathbb{R}^+ \to [-1, \infty)$ is assumed to be minimized only in 1 and its minimal value is fixed to be $v_2(1) = -1$, with no loss of generality. Moreover, $v_2$ is asked to be short-ranged, i.e. to be supported in a neighborhood of 1 of the form $(0, \rho)$. For the sake of definiteness, we let $v_2(r) = 0$ for $r > \rho$ with $\rho := 1.1$. By introducing the distance modulo $L$ defined by

$$|x_i - x_j|_L := \min_{z \in \{-1, 0, +1\}} |x_i - x_j + Lze_3|,$$

we say that two points $x_i, x_j \in C_n, i \neq j$, are bonded, or that the bond between $x_i$ and $x_j$ is active, if $|x_i - x_j|_L < \rho$, and we refer to the graph formed by all the active bonds as the bond graph. Let us denote by $\mathcal{N}$ the set of all pairs of indexes corresponding to bonded points, i.e.

$$\mathcal{N} := \{(i,j) : x_i, x_j \in C_n, i \neq j, \text{ and } |x_i - x_j|_L < \rho\}.$$

For $x_j \in C_n$ we also let

$$\mathcal{N}(x_j) := \{i \in \{1, \ldots, n\} : (i,j) \in \mathcal{N}\},$$

and if $i \in \mathcal{N}(x_j)$ we define by $x_i^j$ the point in $\{x_i + Lze_3 : z = -1, 0, +1\}$ such that $|x_i^j - x_j| = |x_i - x_j|_L$.

The three-body potential $v_3 : [0, 2\pi] \to [0, \infty)$ is assumed to be symmetric around $\pi$, namely $v_3(\alpha) = v_3(2\pi - \alpha)$, and to attain minimum value 0 only at $2\pi/3$ and $4\pi/3$. Furthermore, $v_3$ is required to be convex and strictly decreasing in the interval

$$I_\varepsilon := (2\pi/3 - \varepsilon, 2\pi/3),$$

for some given $\varepsilon < \pi/8$, and twice differentiable at each point of $I_\varepsilon$, with continuous second derivative. This entails that $v_3'(2\pi/3) = 0$. Moreover, we additionally assume that $v_3''(2\pi/3) > 0$.

The parameter $\varepsilon$ will be assumed to be fixed throughout the paper. The index set $\mathcal{T}$ in (1) identifies triples of first-neighboring points, namely,

$$\mathcal{T} := \{(i,j,k) : x_i, x_j, x_k \in C_n, i \neq k, i \in \mathcal{N}(x_j) \text{ and } k \in \mathcal{N}(x_j)\}.$$

For $(i,j,k) \in \mathcal{T}$, the bond angle $\omega_{ijk}$ in (1) is the angle formed by the vectors $x_i^j - x_j$ and $x_k^j - x_j$ (counterclockwise oriented with respect to the normal $(x_i^j - x_j) \wedge (x_k^j - x_j)$), see Figure 2.

The above assumptions on $v_2$ and $v_3$ are quite general. In particular, they are satisfied by standard interaction potentials for $sp^2$-bonding in carbon [40, 41]. Note that we do not require $v_2$ to be repulsive at short-range, although this is a fairly classical assumption and, currently, the only frame in which crystallization in the hexagonal lattice has been rigorously proved [11, 16, 29]. Since the energy $E$ is clearly rotation and translation invariant, in the following we will tacitly assume that all statements are to be considered up to isometries.

We aim at characterizing the fine geometry of stable nanotubes, namely those which locally minimize the configurational energy $E$. To this end, we need to introduce specific families of
zigzag and armchair nanotubes, denoted by $\mathcal{F}_i^z$ and $\mathcal{F}_i^a$ in the following, where $\ell \in \mathbb{N}$ stands for the roll-up parameter whose role we discussed in the introduction, see Figure 1. For the sake of convenience we postpone to Section 3 the detailed definition of such families. Let us however anticipate that all nanotubes in $\mathcal{F}_i^z$ (throughout the paper we use the index $s = z$ for zigzag and $s = a$ for armchair) are objective configurations [23], for they can be obtained as orbits of two points under the action of an isometry group of rotations and translations. In particular, nanotubes in $\mathcal{F}_i^a$ all have points disposed on sections orthogonal to their axis, each containing exactly $\ell \in \mathbb{N}$ points, and the topology of their bond graph corresponds to that of the hexagonal lattice $\mathcal{H}$ under the identification $x \mapsto x + \ell a$ for zigzag and $x \mapsto x + \ell(a+b)$ for armchair, as seen in the introduction.

For a given $\ell \in \mathbb{N}$, the collection $\mathcal{F}_i^z$ can be described by a single scalar parameter. As a consequence, when restricted to the specific family $\mathcal{F}_i^s$ of nanotubes, the minimization of the configurational energy is reduced to a scalar problem. Under the above assumptions one can prove that a unique strict minimizer $\mathcal{F}_i^s$ exists in $\mathcal{F}_i^z$. We prove this in Theorem 4.4 by slightly extending the former [30, Thms. 4.3 and 6.1]. Let us mention again that these minimizers do not coincide with the classical rolled-up [10] nor with the polyhedral model [6], see [30, Thms. 4.1-2 and 6.1], which however belong to these families.

The focus of this paper is to prove that the objective nanotubes $\mathcal{F}_i^s$ are stable, namely that they locally minimize the energy with respect to general perturbations, not restricting to $\mathcal{F}_i^z$. In order to illustrate the statement, let $\ell \in \mathbb{N}$ be fixed and denote by $F_n = \{x_1, \ldots, x_n\}$ and $L^s$ the $n$-cell and the period of $\mathcal{F}_i^s$, so that $\mathcal{F}_i = (F_n, L^s)$. We define small perturbations $\mathcal{P}_\eta(\mathcal{F}_i)$ of $\mathcal{F}_i$ as

$$\mathcal{P}_\eta(\mathcal{F}_i) := \{ \tilde{\mathcal{F}} = (F_n, L) : F_n := \{x_1, \ldots, x_n\} \text{ and } L := L^s + \delta L \}$$

with $x_i := x_i^s + \delta x_i$ for $|\delta x_i| < \eta$ and $|\delta L| < \eta$.

The parameter $\eta > 0$ will be always chosen to be small enough so that the topology of the bond graph of a nanotube $\tilde{\mathcal{F}} \in \mathcal{P}_\eta(\mathcal{F}_i)$ coincides with that of $\mathcal{F}_i^s$: in particular, if $\eta$ is small enough we have

$$\mathcal{N}(x_i) = \mathcal{N}(x_i^s), \quad i = 1, \ldots, n. \tag{3}$$

For all nanotubes whose $n$-cell $C_n = \{x_1, \ldots, x_n\}$ fulfills (3), in particular for all perturbations $\tilde{\mathcal{F}} \in \mathcal{P}_\eta(\mathcal{F}_i)$ with small enough $\eta$, we have $\#\mathcal{N}(x_i) = 3$ and we denote by $i_1, i_2, i_3$ the three elements of $\mathcal{N}(x_i)$, for any $i \in \{1, \ldots, n\}$. Moreover, we indicate by $b_i = |x_i-x_{i_3}|$, and $a_i^j = |x_i-x_{i_j}|$, $j = 1, 2$, the lengths of the three bonds at $x_i$. The bond angle (smaller than or equal to $\pi$) between $a_i^1$ and $a_i^2$ will be denoted by $\beta_i$ whereas $\alpha_i^j$ are the bond angles (smaller than or equal to $\pi$) between $b_i$ and $a_i^j$, for $j = 1, 2$, see Figure 3. This notation will be used to highlight the role of the bond $b_i$: this will (approximately) be parallel to the axis for zigzag nanotubes and (approximately) lay on a plane perpendicular to the axis for armchair nanotubes.

![Figure 2. Notation for bonds and bond angles.](image-url)
Given $x_i \in C_n$ with $\mathcal{N}(x_i) = \{i_1, i_2, i_3\}$, the full three-dimensional geometry of its first neighbors can be uniquely determined up to rotations in terms of the bond lengths $(a_1^i, a_2^i, b_i)$ and the bond angles $(\alpha_1^i, \alpha_2^i, \beta_i)$. In fact, the angle $\beta_i$ can be expressed as a function of $\alpha_1^i$, $\alpha_2^i$, and an additional angle $\gamma_s^i$, whose definition differs in the zigzag and in the armchair case. Indeed, we let $\gamma_z^i$ be the (smaller than $\pi$) angle formed by the plane containing $x_i$, $x_{i_1}$, and $x_{i_3}$ and the plane containing $x_i$, $x_{i_2}$, and $x_{i_3}$, and $\gamma_a^i$ is the (larger than $\pi/2$) angle between the line $x_i - x_{i_3}$ and its projection on the plane containing $x_i$, $x_{i_1}$, and $x_{i_2}$ (see Figure 4).

We indicate by $\gamma_\ell$ the internal angle of the regular $(2\ell)$-gon, namely

$$\gamma_\ell := \pi(1 - 1/\ell).$$

Given a nanotube $\tilde{F} \in \mathcal{P}_\eta(F_\ell^s)$, with small enough $\eta$ so that (3) holds, let $\overline{\gamma}$ be the mean of its $\gamma_s^i$ angles, namely

$$\overline{\gamma} := \frac{1}{n} \sum_{i=1}^{n} \gamma_s^i$$

(4)
with \( s \) either \( a \) or \( z \). We note that \( \gamma = \gamma_\ell \) if \( \tilde{F} \in F_\ell \). Indeed, nanotubes in \( F_\ell \) are contained in the surface of a prism (see Section 3 below), whose section is a \((2\ell)\)-gon, and the sum of the \( \gamma_i^s \) is related with the sum of the internal angles of such \((2\ell)\)-gon. On the other hand, \( \gamma \neq \gamma_\ell \) in general.

We are now ready to state our main result.

**Theorem 2.1 (Stability).** Let \( \varepsilon \in (0, \pi/8) \). There exists \( \ell^*_\varepsilon = \ell^*_\varepsilon(v_3, \varepsilon) \in \mathbb{N} \) such that for all \( \ell > \ell^*_\varepsilon \) one can find \( \eta^\varepsilon = \eta^\varepsilon(v_3, \varepsilon, \ell) > 0 \) so that any nontrivial perturbation \( \tilde{F} \in \mathcal{P}_{\eta^\varepsilon}(F_\ell) \) with \( \gamma \leq \gamma_\ell \) satisfies \( E(\tilde{F}) > E(F_\ell) \).

The theorem asserts the local minimality of \( F_\ell \) with respect to all small perturbations with \( \gamma \leq \gamma_\ell \). As already mentioned, this class includes all small perturbations in \( F_\ell \), which indeed fulfill the stronger \( \gamma = \gamma_\ell \). This particularly implies that Theorem 2.1 includes the former analysis in [30], which is restricted to perturbations in \( F_\ell \).

More significantly, a large class of small perturbations which are *not* in \( F_\ell \) fulfill the constraint \( \gamma \leq \gamma_\ell \) as well. In the zigzag case, one has even \( \gamma = \gamma_\ell \) whenever the perturbation preserves the alignment to the nanotube axis of the \( b_i \) bonds. This includes tractions and diameter changes, which may vary section by section of the nanotube. Moreover, sections can be deformed, for instance squeezed to ellipses, and some shear can also be accommodated, all possibly varying section by section. The armchair case is analogous: a large class of relevant perturbations including traction and diameter change fulfill \( \gamma \leq \gamma_\ell \).

Unfortunately not all small perturbations have \( \gamma \leq \gamma_\ell \). One example is obtained by displacing a single point of \( F_\ell \) orthogonally to the axis. The assertion of Theorem 2.1 can be extended to *all* small perturbations under the following assumption: given \( \ell, \eta^\varepsilon \) from Theorem 2.1,

\[
\forall \tilde{F} \in \mathcal{P}_{\eta^\varepsilon}(F_\ell), \exists \tilde{F} \in \mathcal{P}_{\eta^\varepsilon}(F_\ell) \text{ such that } \gamma \leq \gamma_\ell \text{ and } E(\tilde{F}) \geq E(F_\ell),
\]

(5)

where \( \gamma \) is the mean value of the \( \gamma_i^s \) angles of \( \tilde{F} \). The latter has a constructive flavor: for all given small perturbations one is required to exhibit a better energy competitor fulfilling the constraint \( \gamma \leq \gamma_\ell \). We advance two such candidate, energy-reducing constructions and present numerical evidence for (5) in Section 6.

The statement of Theorem 2.1 relates the size of \( \ell \), which roughly corresponds to the diameter of the nanotube, and the parameter \( \varepsilon \), which instead measures the interval of validity of the assumptions on \( v_3 \). The smaller the \( \varepsilon \), namely the weaker the hypothesis on \( v_3 \), the larger the diameter of the nanotube needs to be in order have stability. At the same time, nanotubes of very small diameter may be treated if assumptions on \( v_3 \) are strengthened, namely if \( \varepsilon \) is large enough.

### 3. Zigzag and Armchair Geometries

We devote this section to the description of the geometry of zigzag and armchair nanotubes and, specifically, we define the families \( F_\ell \).

**Zigzag.** For any \( \ell > 3 \), the family \( F_\ell \) is defined up to isometries as

\[
\left\{ \left( r \cos \left( \frac{\pi(2i+k)}{\ell} \right), r \sin \left( \frac{\pi(2i+k)}{\ell} \right), k(1+s) + j \right) \mid i = 1, \ldots, \ell, \ j = 0, 1, \ k \in \mathbb{Z} \right\}
\]

(6)
for some choice of radius \( r \) and step \( 1 + s \) so that
\[
 r \in \left( 0, \frac{1}{2 \sin(\pi/(2\ell))} \right) \quad \text{and} \quad s \in (0, 1)
\]
along with the constraint
\[
s^2 + 4r^2 \sin^2 \left( \frac{\pi}{2\ell} \right) = 1. \tag{7}
\]
Figure 5 illustrates a configuration in \( \mathcal{F}_z^\ell \). As already mentioned, configurations in \( \mathcal{F}_z^\ell \) are objective \([23]\) for they correspond to orbits of two points under translations along \( e_3 \) of \( 2(1 + s) \) and rototranslations in direction \( e_3 \) of angle \( \pi/\ell \) and length \( 1 + s \). The constraint (7) expresses the fact that all bonds have length 1 and we have that \( \mathcal{F}_z^\ell \) is a one-parameter smooth family of configurations: each configuration in \( \mathcal{F}_z^\ell \) is uniquely determined by specifying either \( r \) or \( s \).

Note that for \( r = 0 \) the cylinder is reduced to its axis and for \( s = 0 \) the sections collide.

Nanotubes \( \mathcal{F} \) in \( \mathcal{F}_z^\ell \) have all points lying on the surface of a cylinder with axis \( e_3 \) and radius \( r \) and are organized in planar sections, perpendicular to \( e_3 \), obtained by fixing \( k \) and \( j \) in (6). Each section contains exactly \( \ell \) points are arranged at the vertices of a regular \( \ell \)-gon, and its two closest sections are at distance \( s \) and 1, respectively.

The position of any point in \( \mathcal{F} \) is identified by the triple \( (i,k,j) \) for \( i \in \{1,\ldots,\ell\} \), \( k \in \mathbb{Z} \), and \( j \in \{0,1\} \). The point \((i,k,j)\) is bonded to the three points \((i,k,1)\), \((i,k-1,1)\) and \((i-1,k-1,1)\) for \( j = 0 \), or the three points \((i,k,0)\), from \((i,k+1,0)\) and \((i-1,k+1,0)\) for \( j = 1 \) (along with the convention that \((0,k,j)\) and \((\ell,k,j)\) are identified), since they are at distance 1. The full geometry of such three neighbors to \( x_i \) is hence described by the three angles \( (\alpha^1_i,\alpha^2_i,\beta_i) \), see Figure 3. Equivalently, one can express \( \beta_i \) as a function of \( \alpha^1_i, \alpha^2_i \) and \( \gamma^z_i \) as

\[
\beta_i = \tilde{\beta}^z(\alpha^1_i, \alpha^2_i, \gamma^z_i)
\]

where the function \( \tilde{\beta}^z \) is defined by
\[
\tilde{\beta}^z(\alpha^1, \alpha^2, \gamma) := 2 \arcsin \left( \sqrt{(1 - \sin \alpha^1 \sin \alpha^2 \cos \gamma - \cos \alpha^1 \cos \alpha^2)/2} \right) \tag{8}
\]

In case \( \alpha^1 = \alpha^2 \) we will also use the notation
\[
\beta^z(\alpha, \gamma) := \tilde{\beta}^z(\alpha, \alpha, \gamma) = 2 \arcsin \left( \sin \alpha \sin \frac{\gamma}{2} \right).
\]

Restrictions on the possible choice of the radius \( r \) have to be introduced in order to ensure that bond graph of nanotubes in \( \mathcal{F}_z^\ell \) to locally have the topology of Figure 4. This imposes
to rule out extremal values of the radius $r$. At first, sections of the nanotube should be wide enough to have that second-neighbors stay at distance larger than $\rho = 1.1$. This translates to

$$r > r_\pm := \frac{0.55}{\sin \gamma_\ell}.$$  \hfill (9)

On the other hand, if $r$ is large then the intersectional distance $1+s$ approaches 1, thus activating extra bonds. This calls for the condition

$$r < r_\pm^+ := \frac{\sqrt{0.9975}}{2 \sin(\pi/(2\ell))}$$ \hfill (10)

as already discussed in [30]. We record all relevant facts about the family $\mathcal{F}_\ell^z$ in the following statement.

**Proposition 3.1** (Zigzag geometry). In $F \in \mathcal{F}_\ell^z$ with radius $r \in (r_\pm^-, r_\pm^+)$ all points have exactly three (first-nearest) neighbors, all at distance 1, with one bond $b$ aligned with $e_3$. The two bond angles (smaller than $\pi$) formed by $b$ are equal and have value $\alpha \in (\pi/2, \pi)$. The third bond angle (smaller than $\pi$) is

$$\beta^z(\alpha, \gamma_\ell) = 2 \arcsin \left( \sin \alpha \sin \frac{\gamma_\ell}{2} \right).$$

We refer to [30, Proposition 3.2] for a detailed proof of Proposition 3.1. Let us point out here that, by (7) and the fact that $\sin \alpha = \sqrt{1-s^2} = 2\sin(\pi/(2\ell))$, the family $\mathcal{F}_\ell^z$ is smoothly and uniquely-determined not only by the parameters $r$ and $s$ independently, but also by the angle $\alpha$. In this regard note that the constraints on $\alpha$ corresponding to (10) is

$$\alpha > \alpha_\pm := \arccos(-1/20) \approx 93^\circ$$ \hfill (11)

and the one corresponding to (9) is

$$2 \sin \alpha \sin(\gamma_\ell/2) > \rho.$$

For all $\ell \geq 4$ the latter is implied by the following

$$\alpha < \alpha_\pm^+ := \pi - \arcsin \left( \frac{1.1}{\sqrt{2 + \sqrt{2}}} \right) \approx 143.5^\circ.$$ \hfill (12)

Proposition 3.1 entails in particular that, for all $\alpha \in (\alpha_\pm^-, \alpha_\pm^+)$, a configuration $F_\alpha \in \mathcal{F}_\ell^z$, uniquely defined by its bond angle $\alpha$, features two thirds of the bond angles (smaller than $\pi$) with value $\alpha$ and one third with value $\beta^z(\alpha, \gamma_\ell)$. This allows a simple expression for its energy. Denote its $n$-cell ($n = 4m\ell$ for some $m \in \mathbb{N}$) by $F_n$ and observe that its period $L$ can be computed as

$$L = 2m(1 - \cos \alpha)$$ \hfill (13)

(for $m = 1$ we get the minimal period of the configuration). As all points bring the same contribution to the energy we have

$$E(F_\alpha) = E(F_n, L) = -\frac{3}{2}n + n\mathcal{E}^z(\alpha, \gamma_\ell)$$ \hfill (14)

where

$$\mathcal{E}^z(\alpha, \gamma) := 2v_3(\alpha) + v_3(\beta^z(\alpha, \gamma)).$$
**Armchair.** For all \( \ell \in 2\mathbb{N} \), we introduce the family \( \mathcal{F}_\ell^a \) up to isometries as

\[
\left\{ \left( r \cos \left( \frac{2\pi}{\ell}(2i+k)+q_r,j \right), r \sin \left( \frac{2\pi}{\ell}(2i+k)+q_r,j \right), p k \right) \bigg| i = 1, \ldots, \ell; j = 0, 1, k \in \mathbb{Z} \right\}
\]

(15)

for some choice of radius \( r \) and step \( p \) such that

\[
r \in \left( \frac{1}{2\sin(\pi/\ell)}, \frac{1}{2\sin(\pi/(2\ell))} \right) \quad \text{and} \quad p \in (0,1)
\]

and the following constraint holds

\[
p^2 + 4r^2 \sin^2 \left( \frac{\pi}{\ell} - \frac{q_r}{2} \right) = 1,
\]

(16)

where

\[
q_r := 2 \arcsin \left( \frac{1}{2r} \right).
\]

Points in \( \mathcal{F} \in \mathcal{F}_\ell^a \) lie on the surface of a cylinder with radius \( r \) and axis \( e_3 \) and they are arranged in planar sections, perpendicular to \( e_3 \), obtained by fixing \( k \) and \( j \) in (15). The constraint (16) entails that all bonds have length 1. The collection \( \mathcal{F}_\ell^a \) is a one-parameter smooth family uniquely determined by \( r \) or \( p \). The limit cases \( p = 0 \) (for which we have \( q_r = \pi/\ell \)) and \( p = 1 \) (for which \( q_r = 2\pi/\ell \)) correspond to having the sections on the same plane, and to a prism shape, respectively.

Each of the sections exactly contains \( \ell \) points, arranged at the vertices of two regular \((\ell/2)\)-gons, which are rotated of an angle \( q_r \) with respect to each other. For each section, the two closest sections are both at distance \( p \). Nanotubes \( \mathcal{F}_\ell^a \) are objective [23], for they correspond to the orbit of two points under rotations of \( 4\pi/\ell \) around the axis \( e_3 \) and roto-translations about \( e_3 \) of angle \( 2\pi/\ell \) and step \( p \). By (15) we see that the triple \((i,k,j)\) for \( i \in \{1,\ldots,\ell/2\}, k \in \mathbb{Z}, \) and \( j \in \{0,1\} \) uniquely determines the positions of all points. Note that the point \((i,k,j)\) is bonded with the three points \((i,k,1)\), \((i,k-1,1)\) and \((i-1,k+1,1)\) for \( j = 0 \), or the three points \((i,k,0)\), \((i,k+1,0)\), and \((i+1,k-1,0)\) for \( j = 1 \) (where we identify \((0,k,j)\) and \((\ell/2,k,j)\)), since they are at distance 1. The geometry of such ensemble of three neighbors to \( x_i \) is hence described by the three angles \((\alpha_1^a, \alpha_2^a, \beta_i)\), see Figure 4. Equivalently, one can express \( \beta_i \) as a function of \( \alpha_1^a, \alpha_2^a \) and \( \gamma_i^a \) as

\[
\beta_i = \tilde{\beta}^a(\alpha_1^a, \alpha_2^a, \gamma_i^a)
\]

where the function \( \tilde{\beta}^a \) is defined by

\[
\tilde{\beta}^a(\alpha_1^a, \alpha_2^a, \gamma) := \arccos \left( \frac{\cos \alpha_1^a}{\cos \gamma} \right) + \arccos \left( \frac{\cos \alpha_2^a}{\cos \gamma} \right),
\]

If \( \alpha_1^a = \alpha_2^a \) we will also use the notation

\[
\beta^a(\alpha, \gamma) := \tilde{\beta}^a(\alpha, \alpha, \gamma) = 2 \arccos \left( \frac{\cos \alpha}{\cos \gamma} \right).
\]

As in the case of zigzag geometries, parameters should be additionally constrained in order the bond graph to locally have the topology of Figure 4. Since \( p \) is the distance between two consecutive sections we require \( 2p > \rho \). According to (16) this corresponds to \( 2r \sin(\pi/\ell - q_r/2) < \sqrt{0.6975} \), which yields \( r < r_a^p \), where \( r_a^p \) denotes the unique positive solution of the corresponding equality (notice that the map \( r \mapsto 2r \sin(\pi/\ell - q_r/2) \) is monotone increasing for \( r > 1/2 \), as \( \ell \geq 4 \), and takes values 0 and 1 at extremes of the definition range for \( r \)).
On the other hand, the distance between two nearest points on a section is either $2r \sin(qr/2) = 1$ or $2r \sin(2\pi/\ell - qr/2)$. We impose the latter to be greater than $\rho$ to avoid an extra bond. This requires $r > r^a$, where $r^a$ is the unique positive solution to $2r \sin(2\pi/\ell - qr/2) = \rho$. The armchair version of Proposition 3.1 reads as follows.

**Proposition 3.2** (Armchair geometry). In $F \in \mathcal{F}^a_\ell$ with radius $r \in (r^a, r^a + \rho)$ all points have exactly three (first-nearest) neighbors, all at distance 1, with one bond $b$ orthogonal to $e_3$. The two bond angles (smaller than $\pi$) formed by $b$ are equal and have value $\alpha \in (\pi/2, \pi)$. The third bond angle (smaller than $\pi$) is

$$\beta^a(\alpha, \gamma\ell) = 2 \arccos \left( \frac{\cos \alpha}{\cos \gamma\ell} \right).$$

Analogously as for the zigzag family, by equation

$$\cos \alpha = 2r \cos \gamma\ell \sin \left( \frac{\pi}{\ell} - \frac{qr}{2} \right)$$

one can characterize the element of the family $\mathcal{F}^a_\ell$ via the bond angle $\alpha \in (\pi/2, \gamma\ell)$. The constraint $r < r^a$ is equivalent to

$$\cos^2 \alpha < 0.6975 \cos^2 \gamma\ell,$$

which, for all $\ell \geq 4$, is implied by

$$\alpha < \alpha^a_+ := \arccos(-\sqrt{0.34875}) \approx 126^\circ.$$  

(17)

On the other hand, using the identity $2r \sin(2\pi/\ell - qr/2) = 1 + 4r \cos(\pi/\ell) \sin(\pi/\ell - qr/2)$ and (17), the constraint $r > r^a$ translates to

$$\alpha > \alpha^a_+ := \arccos(-1/20) \approx 93^\circ.$$  

(18)

For all $\alpha \in (\alpha^a_-, \alpha^a_+)$ the energy of the nanotube $F_\alpha \in \mathcal{F}^a_\ell$ has a simple expression, since all points contribute the same amount to it. For $F_\alpha = (F_n, L)$ where $n = 2m\ell$, $m \in \mathbb{N}$, we can compute that

$$L := 2m \left( 1 - \frac{\cos^2 \alpha}{\cos^2 \gamma\ell} \right)^{1/2}$$

and the energy reads

$$E(F_\alpha) = E(F_n, L) = -\frac{3}{2} n + nE^a(\alpha, \gamma\ell)$$

where

$$E^a(\alpha, \gamma) := 2v_3(\alpha) + v_3(\beta^a(\alpha, \gamma)).$$

(21)

4. Angle-energy optimization

The analysis of the previous section entails that the minimization problem for configurations in $\mathcal{F}^a_\ell$ is localized, for all points equally contribute to the energy. As all bond lengths in $\mathcal{F}^a_\ell$ are equal to 1, we reduce ourselves to the study of a scalar functional corresponding to the angle energy of a single point, see (14) and (21). This analysis is performed in this section and, in particular, brings to the identification of the optimal configuration $F^a_\ell$.

Let us start by computing the derivatives of the functions $\beta^a$, which we consider to be defined in $(\pi/2, 3\pi/4) \times (\pi/2, 3\pi/4) \times [3\pi/4, \pi]$. These values are just taken for the sake of definiteness.
and could be differently chosen around 2π/3 for α₁, α², and left to π for γ. Notice in particular that γ_ℓ ∈ [3π/4, π) for ℓ ≥ 4. In this range, we have

\[
\begin{align*}
\partial_\alpha \hat{\beta}^z(\alpha^1, \alpha^2, \gamma) &= -\frac{\cos \alpha^1 \sin \alpha^2 \cos \gamma + \sin \alpha^1 \cos \alpha^2}{\sqrt{1 - (\sin \alpha^1 \sin \alpha^2 \cos \gamma + \cos \alpha^1 \cos \alpha^2)^2}} < 0 \\
\partial_\alpha \hat{\beta}^z(\alpha^1, \alpha^2, \gamma) &= -\frac{\sin \alpha^1 \cos \alpha^2 \cos \gamma + \cos \alpha^1 \sin \alpha^2}{\sqrt{1 - (\sin \alpha^1 \sin \alpha^2 \cos \gamma + \cos \alpha^1 \cos \alpha^2)^2}} < 0 \\
\partial_{\alpha^i} \tilde{\beta}^a(\alpha^1, \alpha^2, \gamma) &= -\frac{2 \sin \alpha^i}{\sqrt{\cos^2 \gamma - \cos^2 \alpha^i}} < 0, \quad i = 1, 2.
\end{align*}
\]

The functions β⁺(α, γ) are smooth on (π/2, 3π/4) × [3π/4, π] and we have

\[
\begin{align*}
\partial_\alpha \beta^z &= \frac{2 \sin(\gamma/2) \cos \alpha}{\sqrt{1 - \sin^2 \alpha \sin^2(\gamma/2)}} < 0, \quad \partial^2_{\alpha \alpha} \beta^z = \frac{2 \sin \alpha \sin(\gamma/2) \cos^2(\gamma/2)}{(1 - \sin^2 \alpha \sin^2(\gamma/2))^{3/2}} < 0, \\
\partial_{\gamma} \beta^z &= \frac{\sin \alpha \cos(\gamma/2)}{\sqrt{1 - \sin^2 \alpha \sin^2(\gamma/2)}} > 0, \quad \partial^2_{\gamma \gamma} \beta^z = \frac{-\sin \alpha \cos^2 \alpha \sin(\gamma/2)}{2(1 - \sin^2 \alpha \sin^2(\gamma/2))^{3/2}} < 0, \\
\partial_\alpha \beta^a &= -\frac{2 \sin \alpha}{\sqrt{\cos^2 \gamma - \cos^2 \alpha}} < 0, \quad \partial^2_{\alpha \alpha} \beta^a = \frac{2 \cos \alpha \sin^2 \gamma}{(\cos^2 \gamma - \cos^2 \alpha)^{3/2}} < 0, \\
\partial_{\gamma} \beta^a &= \frac{2 \cos \alpha \tan \gamma}{\sqrt{\cos^2 \gamma - \cos^2 \alpha}} > 0, \quad \partial^2_{\gamma \gamma} \beta^a = \frac{2 \cos \alpha \cos^2 \gamma - 2 \cos^2 \alpha + \sin^2 \gamma \cos^2 \gamma}{\cos^2 \gamma (\cos^2 \gamma - \cos^2 \alpha)^{3/2}} < 0.
\end{align*}
\]

The values at the extrema of the definition range are

\[
\begin{align*}
\partial_\alpha \beta^z(2\pi/3, \pi) &= -2, \quad \partial^2_{\alpha \alpha} \beta^z(2\pi/3, \pi) = 0, \\
\partial_{\gamma} \beta^z(2\pi/3, \pi) &= 0, \quad \partial^2_{\gamma \gamma} \beta^z(2\pi/3, \pi) = -\sqrt{3}/2, \\
\partial_\alpha \beta^a(2\pi/3, \pi) &= -2, \quad \partial^2_{\alpha \alpha} \beta^a(2\pi/3, \pi) = 0, \\
\partial_{\gamma} \beta^a(2\pi/3, \pi) &= 0, \quad \partial^2_{\gamma \gamma} \beta^a(2\pi/3, \pi) = -2/\sqrt{3}.
\end{align*}
\]

Let us consider first the case in which two of the three bond angles are equal. In particular, we discuss the minimization of E⁺ with respect to α for all given γ. Some care is needed for the choice of the interval of optimization. This has to be specified depending on γ and ε ∈ (0, π/8) from (2) in order to ensure that v₃ is evaluated just in I₁, namely where the assumptions hold. For ε ∈ (0, π/8) and γ ∈ [3π/4, π] we define the two quantities

\[
\sigma^+_\varepsilon(\gamma) := \pi - \arcsin\left(\frac{\sin(\pi/3 - \varepsilon/2)}{\sin(\gamma/2)}\right), \quad \sigma^-\varepsilon(\gamma) := \arccos(\cos \gamma \cos(\pi/3 - \varepsilon/2)).
\]

From the definitions of β⁺ and their monotonicity properties from (23), (24), it is not difficult to check that σ⁺(γ) belongs to (π/2, 3π/4) and that it is the unique solution to the equation β⁺(γ, γ) = 2π/3 - ε on (π/2, 3π/4). Moreover, we have β⁺(α, γ) > 2π/3 - ε if α < σ⁺(γ). The functions σ⁺(γ) are strictly increasing both with respect to ε ∈ (0, π/8) and to γ ∈ [3π/4, π].

By recalling that ε is fixed throughout, one has that σ⁺(γ) is larger than 2π/3 if γ is large enough (i.e., close to π). More precisely, taking into account the strict monotonicity of [3π/4, π] ⊃
γ → σ_ε^*(γ), we define values γ_ε as follows

\[
\gamma_\varepsilon := \begin{cases} 
3\pi/4 & \text{if } \sigma_\varepsilon^*(3\pi/4) \geq 2\pi/3 \\
(\sigma_\varepsilon^*)^{-1}(2\pi/3) & \text{if } \sigma_\varepsilon^*(3\pi/4) < 2\pi/3,
\end{cases}
\]

where (σ_ε^*)^{-1}(·) is the inverse of σ_ε^*(·) on [3π/4, π]. In this way, we obtain σ_ε^*(γ) > 2π/3 if γ ∈ (γ_ε, π). It will hence be important to keep γ in the range (γ_ε, π). Notice that the smaller the ε, the stronger is the restriction on the values of γ. For instance, if ε = π/12, we have γ_ε = 3π/4 and (γ_ε) ∈ (3π/4, 5π/6).

We will also need the quantities σ_0^*(γ), defined as the unique solutions to β^*(·, γ) = 2π/3 on (π/2, 3π/4). These are explicitly given by

\[
\sigma_0^*(γ) := \pi - \arcsin \left( \frac{\sqrt{3}}{2\sin(\gamma/2)} \right), \quad \sigma_0^*(γ) := \arccos \left( \frac{\cos \gamma}{2} \right). \tag{28}
\]

The notation corresponds to the fact that σ_ε^* = lim_{ε→0} σ_ε^*. Notice that γ → σ_0^*(γ) is increasing on [3π/4, π] and converging to 2π/3 as γ ↑ π. Thanks to (23) and (24) we also have that β^*(α, γ) > 2π/3 if α < σ_0^*(γ). The definition of γ_ε and the monotonicity of σ_ε^*(γ) entail that σ_0^*(γ) > 2π/3 - ε for any ε ∈ (0, π/8), as soon as γ ≥ γ_ε.

We can now proceed with the optimization of E^*(·, γ) as follows.

**Lemma 4.1 (Minimality in α).** Given γ ∈ (γ_ε, π), the map α → E^*(α, γ) admits a unique minimizer α^*(γ) on (2π/3 - ε, σ_ε^*(γ)). Moreover, one has that α^*(γ) < 2π/3, β^*(α^*(γ), γ) < 2π/3, ∂_α E^*(α^*(γ), γ) > 0, and α^*(γ) → 2π/3 as γ ↑ π.

**Proof.** Let γ ∈ (γ_ε, π) be fixed. This ensures that σ_ε^*(γ) > 2π/3, by the definition of γ_ε.

First of all, we show that α is not a minimizer of α → E^*(α, γ) if β^*(α, γ) > 2π/3. We recall that σ_0^*(γ) is defined as the unique value in (π/2, 2π/3) such that β^*(σ_0^*(γ), γ) = 2π/3, and that σ_0^*(γ) > 2π/3 - ε. If β^*(α, γ) > 2π/3, the monotonicity of β^* from (23) and (24) shows that 2π/3 - ε < α < σ_0^*(γ), then we have

\[
E^*(α, γ) = 3v_3(α) + v_3(β^*(α, γ)) > 2v_3(σ_0^*(γ)) = 2v_3(σ_0^*(γ)) + v_3(β^*(σ_0^*(γ), γ)) = E_1(σ_0^*(γ), γ),
\]

where we have used the nonnegativity and the strict monotonicity of v_3 and the fact that v_3(2π/3) = 0.

One can similarly prove that there is no minimizer in (2π/3, σ_ε^*(γ)). Indeed, for α in such interval we have β^*(α, γ) ∈ I_ε, from the obvious geometric constraint 2α + β_ε(α, γ) < 2π and from the definition of σ_ε^*(γ). Then, the monotonicity of α → β^*(α, γ) and v_3 entails

\[
E^*(α, γ) = 3v_3(α) + v_3(β^*(α, γ)) > v_3(β^*(2π/3, γ)) = E^*(2π/3, γ).
\]

Let now α ∈ (σ_0^*(γ), 2π/3). Then, we have just shown that β^*(α, γ) ∈ I_ε. Since the composition of v_3 (convex and strictly decreasing on I_ε) and β^* (strictly concave) is strictly convex, it follows that E^*(·, γ) is strictly convex in (σ_0^*(γ), 2π/3). Let us now check that α → E^*(α, γ) is not minimized at the extrema σ_0^*(γ) and 2π/3, and this will conclude the proof. Since σ_0^*(γ) > 2π/3 - ε, then

\[
\partial_α E^*(σ_0^*(γ), γ) = 2v_3'(σ_0^*(γ)) + v_3'(β^*(σ_0^*(γ), γ)) \partial_α β^*(σ_0^*(γ), γ) = 2v_3'(σ_0^*(γ)) < 0,
\]

because v_3'(2π/3) = 0 by assumption. Concerning 2π/3, since β^*(2π/3, γ) < 2π/3, we have

\[
\partial_α E^*(2π/3, γ) = 2v_3'(2π/3) + v_3'(β^*(2π/3, γ)) \partial_α β^*(2π/3, γ) > 0,
\]
thanks once more to the monotonicity of $v_3$ and $\beta^s(\cdot, \gamma)$. The claim is proved, and the unique minimizer $\alpha^*(\gamma)$ belongs to $(\sigma_5^*(\gamma), 2\pi/3) \subset I_\varepsilon$.

Eventually, we have $\beta^s(\sigma_5^*(\gamma), \gamma) < 2\pi/3$ since $\alpha^*(\gamma) > \sigma_5^*(\gamma)$. It is clear from (27) that $\sigma_5^*(\gamma)$ tends to $2\pi/3$ as $\gamma \uparrow \pi$, so that $\alpha^*(\gamma)$ tends to $2\pi/3$ as well. □

Note that Lemma 4.1 generalizes [30, Thms. 4.3 and 6.1] to the case $\gamma \neq \gamma_e$.

We now turn to prove convexity. The following functions angle energies $\tilde{E}^s$ will be used

$$\tilde{E}^s(\alpha, \alpha^2, \gamma) := v_3(\alpha) + v_3(\alpha^2) + v_3(\beta^s(\alpha, \alpha^2, \gamma)).$$

**Lemma 4.2** (Convexity in $\alpha$). There exist $\gamma_s^* \in (\gamma_e^*, \pi)$ such that, for all $\gamma \in (\gamma_s^*, \pi)$, the map $(\alpha^1, \alpha^2) \mapsto \tilde{E}^s(\alpha, \alpha^2, \gamma)$ is strictly convex on $[\sigma_5^*(\gamma), 2\pi/3] \times [\sigma_5^*(\gamma), 2\pi/3]$.

**Proof.** Let $\gamma > \gamma_s^*$ so that we have both $\sigma_5^*(\gamma) > 2\pi/3$ and $\sigma_5^*(\gamma) > 2\pi/3 - \varepsilon$, as previously observed, see (27)-(28). By taking (23) and (24) into account we have that $\alpha \geq \sigma_5^*(\gamma) > 2\pi/3 - \varepsilon$ implies $\beta^s(\alpha, \gamma) \leq 2\pi/3$, and $\alpha \leq 2\pi/3 < \sigma_5^*(\gamma)$ implies $\beta^s(\alpha, \gamma) > 2\pi/3 - \varepsilon$. Thanks to the monotonicity in (22), analogous bounds are found for $\tilde{\beta}^s$ as well. That is, if $\alpha^1, \alpha^2 \in [\sigma_5^*(\gamma), 2\pi/3] \subset I_\varepsilon$, then $\tilde{\beta}^s(\alpha^1, \alpha^2, \gamma) \in (2\pi/3 - \varepsilon, 2\pi/3) = I_\varepsilon$.

In the following $\nabla$ denotes the derivative with respect to $(\alpha^1, \alpha^2)$. The index $s$ as well as the dependence of $\tilde{\beta}^s$ and $\tilde{E}^s$ on the three variables $(\alpha^1, \alpha^2, \gamma)$ will be omitted for brevity. For $(\alpha^1, \alpha^2 \in [\sigma_5^*(\gamma), 2\pi/3]$, we have

$$\nabla^2 \tilde{E}(\alpha^1, \alpha^2, \gamma) = \text{diag}(v_3''(\alpha^1), v_3''(\alpha^2)) + v_3''(\tilde{\beta})(\nabla\tilde{\beta}) \otimes \nabla\tilde{\beta} + v_3''(\tilde{\beta}) \nabla^2\tilde{\beta}.$$ 

A direct computation yields

$$\tilde{E}_{\alpha^1(\alpha^1, \alpha^2, \gamma)} = v_3''(\alpha^1) + \tilde{\beta}_{\alpha^1(\alpha^1, \alpha^2)} v_3''(\tilde{\beta}),$$

$$\tilde{E}_{\alpha^2(\alpha^1, \alpha^2, \gamma)} = v_3''(\alpha^2) + \tilde{\beta}_{\alpha^2(\alpha^1, \alpha^2)} v_3''(\tilde{\beta}),$$

$$\tilde{E}_{\alpha^1(\alpha^1, \alpha^2, \gamma)} = \tilde{\beta}_{\alpha^1(\alpha^1, \alpha^2)} v_3''(\tilde{\beta}) + \tilde{\beta}_{\alpha^1(\alpha^1, \alpha^2)} v_3''(\tilde{\beta}),$$

$$\det \nabla^2 \tilde{E} = \left[ v_3''(\alpha^1) + \tilde{\beta}_{\alpha^1(\alpha^1, \alpha^2)} v_3''(\tilde{\beta}) \right] \left[ v_3''(\alpha^2) + \tilde{\beta}_{\alpha^2(\alpha^1, \alpha^2)} v_3''(\tilde{\beta}) \right]$$

$$+ v_3''(\alpha^1) v_3''(\tilde{\beta}) \tilde{\beta}_{\alpha^1(\alpha^2)} v_3''(\tilde{\beta}) + v_3''(\alpha^2) v_3''(\tilde{\beta}) \tilde{\beta}_{\alpha^2(\alpha^1)} v_3''(\tilde{\beta}) - v_3''(\tilde{\beta}) \tilde{\beta}_{\alpha^1(\alpha^2)} v_3''(\tilde{\beta}) - v_3''(\tilde{\beta}) \tilde{\beta}_{\alpha^2(\alpha^1)} v_3''(\tilde{\beta}) - 2 v_3''(\tilde{\beta}) v_3''(\tilde{\beta}) \tilde{\beta}_{\alpha^1(\alpha^2)} \tilde{\beta}_{\alpha^2(\alpha^1)}$$

where we have exploited the regularity of $v_3$, see Section 2. In particular, we have $v_3''(2\pi/3) = 0$ and $v_3''(2\pi/3) > 0$. We recall that the functions $\beta^s(\alpha^1, \alpha^2, \gamma)$ are smooth and tend to $2\pi/3$ as $(\alpha^1, \alpha^2, \gamma)$ tends to $(2\pi/3, 2\pi/3, \pi)$, see (8). On the other hand, $\sigma_6^*(\gamma) \uparrow 2\pi/3$ as $\gamma \uparrow \pi$, see (28).

Therefore, we can choose $\gamma_s^* \in (\gamma_e^*, \pi)$ such that, for all $(\alpha^1, \alpha^2) \in [\sigma_6^*(\gamma), 2\pi/3) \times [\sigma_6^*(\gamma), 2\pi/3]$ and $\gamma \in (\gamma_s^*, \pi)$, there hold

$$v_3''(\alpha^1) > \frac{1}{2} v_3''(2\pi/3), \quad v_3''(\alpha^2) > \frac{1}{2} v_3''(2\pi/3), \quad |\tilde{\beta}_{\alpha^1(\alpha^2)} v_3''(\tilde{\beta})| < \frac{1}{4} v_3''(2\pi/3)$$

and

$$|v_3''(\alpha^1) \tilde{\beta}_{\alpha^2(\alpha^2)} v_3''(\tilde{\beta}) + v_3''(\alpha^2) \tilde{\beta}_{\alpha^2(\alpha^1)} v_3''(\tilde{\beta})$$

$$+(\tilde{\beta}_{\alpha^1(\alpha^2)} \tilde{\beta}_{\alpha^2(\alpha^2)} - \tilde{\beta}_{\alpha^2(\alpha^1)} \tilde{\beta}_{\alpha^1(\alpha^2)}) v_3''(\tilde{\beta}) - 2 v_3''(\tilde{\beta}) v_3''(\tilde{\beta}) \tilde{\beta}_{\alpha^1(\alpha^2)} \tilde{\beta}_{\alpha^2(\alpha^1)}| < \frac{1}{8} (v_3''(2\pi/3))^2.$$ 

This way we get $4\tilde{E}_{\alpha^1(\alpha^1)} > v_3''(2\pi/3) > 0$ and $8 \det \nabla^2 \tilde{E} > (v_3''(2\pi/3))^2 > 0$ as desired. □
Lemma 4.3 (Monotonicity and convexity in $\gamma$). There exists $\hat{\gamma}^s_s \in (\gamma^s_s, \pi)$ such that the map 

$$
\gamma \mapsto \mathcal{E}^s(\gamma) := \mathcal{E}^s(\alpha^s(\gamma), \gamma) = \min\{\mathcal{E}^s(\alpha, \gamma) \mid \alpha \in (\sigma^s_0(\gamma), 2\pi/3)\}
$$

is strictly decreasing and strictly convex on $(\hat{\gamma}^s_s, \pi)$.

Proof. In the following, we drop the index $s$ from $\beta^s, \mathcal{E}^s, \mathcal{E}^s', \mathcal{E}^s', \gamma^s_s, \sigma^s_0(\gamma)$, and $\alpha(\gamma)$, for the sake of notational simplicity. Let $\gamma \in (\gamma_s, \pi)$ and take derivatives with respect to $\gamma$. Thanks to first order optimality condition for $\alpha(\gamma)$, i.e. $\mathcal{E}_\alpha(\alpha(\gamma), \gamma) = 0$, we have

$$
\mathcal{E}'(\gamma) = \mathcal{E}_\alpha(\alpha(\gamma), \gamma)\alpha'(\gamma) + \mathcal{E}_\gamma(\alpha(\gamma), \gamma) = v_3'(\beta(\alpha(\gamma), \gamma))\beta_\gamma(\alpha(\gamma), \gamma).
$$

As $\beta(\alpha(\gamma), \gamma) < 2\pi/3$ from Proposition 4.1, we have $v_3'(\beta(\alpha(\gamma), \gamma)) < 0$ and we conclude from (23) and (24) that $\mathcal{E}' < 0$ on $(\gamma_s, \pi)$. By taking the derivative with respect to $\gamma$ in the identity $\mathcal{E}_\alpha(\alpha(\gamma), \gamma) = 0$, we get $\mathcal{E}_{\alpha\alpha}(\alpha(\gamma), \gamma)\alpha'(\gamma) + \mathcal{E}_{\alpha\gamma}(\alpha(\gamma), \gamma) = 0$, thus

$$
\alpha'(\gamma) = -\frac{\mathcal{E}_{\alpha\gamma}(\alpha(\gamma), \gamma)}{\mathcal{E}_{\alpha\alpha}(\alpha(\gamma), \gamma)}.
$$

Using the latter expression for $\alpha'(\gamma)$, we find (here $\nabla$ denotes derivation in the couple $\alpha, \gamma$)

$$
\mathcal{E}''(\gamma) = \mathcal{E}_{\alpha\gamma}(\alpha(\gamma), \gamma)\alpha'(\gamma) + \mathcal{E}_{\gamma\gamma}(\alpha(\gamma), \gamma) = \frac{\mathcal{E}_{\alpha\alpha}(\alpha(\gamma), \gamma)\mathcal{E}_{\gamma\gamma}(\alpha(\gamma), \gamma) - \mathcal{E}^2_{\alpha\gamma}(\alpha(\gamma), \gamma)}{\mathcal{E}_{\alpha\alpha}(\alpha(\gamma), \gamma)} = \frac{\det \nabla^2 \mathcal{E}(\alpha(\gamma), \gamma)}{\mathcal{E}_{\alpha\alpha}(\alpha(\gamma), \gamma)}.
$$

The second-order optimality condition for $\alpha(\gamma)$ reads $\mathcal{E}_{\alpha\alpha}(\alpha(\gamma), \gamma) > 0$, as seen in Proposition 4.1. In order to check the positivity of $\mathcal{E}''$, we need to prove that $\det \nabla^2 \mathcal{E}(\alpha(\gamma), \gamma) > 0$. We check this property in a left neighborhood of $\gamma = \pi$. To this aim we compute the general expression of $\det \nabla^2 \mathcal{E}(\alpha(\gamma), \gamma)$ on $(\sigma(\gamma), 2\pi/3) \times (\gamma_s, \pi)$. The computations of second-order derivatives are similar to the ones in Lemma 4.2. We have that

$$
\mathcal{E}_{\alpha\alpha} = 2v_3''(\alpha) + \beta_\alpha^2 v_3'(\beta) + \beta_{\alpha\alpha} v_3'(\beta),
$$

$$
\mathcal{E}_{\gamma\gamma} = \beta_\gamma^2 v_3''(\beta) + \beta_{\gamma\gamma} v_3'(\beta),
$$

$$
\mathcal{E}_{\alpha\gamma} = \beta_{\alpha\gamma} v_3'(\beta) + \beta_{\alpha\gamma} v_3'(\beta).
$$

These entail in particular that

$$
\det \nabla^2 \mathcal{E} = 2v_3''(\alpha)v_3''(\beta)\beta_\gamma^2 + 2v_3''(\alpha)v_3''(\beta)\beta_{\alpha\gamma}
$$

$$
+ v_3'(\beta)v_3''(\beta)\left(\beta_\alpha^2 \beta_{\alpha\gamma} + \beta_{\gamma\gamma}^2 \beta_{\alpha\alpha} - 2\beta_{\alpha\gamma} \beta_{\alpha\gamma}\right)
$$

$$
+ v_3'(\beta)\left(\beta_{\alpha\gamma} \beta_{\gamma\gamma} - \beta_{\gamma\gamma}^2 \right).
$$

Recall now from Proposition 4.1 that if $\gamma \uparrow \pi$ we also have $\alpha(\gamma) \rightarrow 2\pi/3$, and then $\beta(\alpha(\gamma), \gamma) \rightarrow 2\pi/3$ as well. Taking (25)-(26) into account, one finds that $\beta_\alpha^2 \beta_{\alpha\gamma} - 2\beta_{\alpha\gamma} \beta_{\alpha\gamma}$ is negligible with respect to $\beta_{\gamma\gamma}^2 \beta_{\gamma\gamma}$, which has nonzero limit as $\gamma \uparrow \pi$, and that the term involving $\beta_\gamma^2 \beta_{\gamma\gamma}$ dominates the ones involving $v_3''(\beta)$. Indeed, we have $\beta \rightarrow 2\pi/3$ and $|v_3''(\beta)| \ll v_3'(\beta)v_3'(\beta)$,
since \( v'_3(2\pi/3) = 0 \) and \( v''_3(2\pi/3) > 0 \). Therefore, there exists \( \gamma_\varepsilon \in (\gamma_\varepsilon, \pi) \) such that the sign of \( \det \nabla^2 \mathcal{E}(\alpha(\gamma), \gamma) \) in \( (\gamma_\varepsilon, \pi) \) is the same of
\[
2v''_3(\alpha(\gamma))v'_3(\beta(\alpha(\gamma), \gamma))\beta'_3(\alpha(\gamma), \gamma) \\
+ 2v''_3(\alpha(\gamma))v'_3(\beta(\alpha(\gamma), \gamma))\beta_{3\gamma}(\alpha(\gamma), \gamma) \\
+ v'_3(\beta(\alpha(\gamma), \gamma))v''_3(\beta(\alpha(\gamma), \gamma))\beta''_3(\alpha(\gamma), \gamma)\beta_{3\gamma}(\alpha(\gamma), \gamma).
\]
The latter is positive thanks to (23)-(24) and the fact that \( \beta(\alpha(\gamma), \gamma) < 2\pi/3 \) because of \( \beta(\alpha(\gamma), \gamma) < 2\pi/3 \).

Moving from Lemma 4.1 one can prove that the configurational energy \( E \) admits a unique minimizer on the family \( \mathcal{F}_s^\ell \), recovering indeed [30, Thms. 4.3 and 6.1]. In fact, by (14) and (21) such minimization reduces to the minimization of the angle energy of a single point, namely,
\[
\min \{ \mathcal{E}^s(\alpha, \gamma_\varepsilon) : \alpha \in (2\pi/3 - \varepsilon, \sigma^s_\varepsilon(\gamma_\varepsilon)) \}.
\]
In fact, following Section 3 we should require \( \alpha \in (\alpha^s_-, \alpha^s_+) \) from (11)-(12) and (18)-(19). Note nonetheless that
\[
(2\pi/3 - \varepsilon, \sigma^s_\varepsilon(\gamma_\varepsilon)) \subset (\alpha^s_-, \alpha^s_+)
\]
if \( v_2(r) = 0 \) for \( r \geq 1.1 \) and \( \ell \geq 4 \) in the zigzag case or \( \ell \geq 6 \) in the armchair case. Lemma 4.1 entails the following.

**Theorem 4.4 (Minimality in \( \mathcal{F}_s^\ell \)).** Letting \( \ell \geq 4 \) (zigzag) or \( \ell \geq 6 \) (armchair), there exists a unique nanotube \( \mathcal{F}_s^\ell \in \mathcal{F}_s^\ell \) minimizing \( E \) within \( \mathcal{F}_s^\ell \). This corresponds to the unique minimizer of \( \alpha \rightarrow \mathcal{E}^s(\alpha, \gamma_\varepsilon) \) in \( (2\pi/3 - \varepsilon, \sigma^s_\varepsilon(\gamma_\varepsilon)) \).

Before closing this section, let us mention again that the minimizing nanotubes \( \mathcal{F}_s^\ell \) do not coincide with the rolled-up [10] nor with the polyhedral nanotubes [6]. Indeed, these are defined via their corresponding angles \( \alpha^s_\text{pol} = 2\pi/3 \) and by \( \alpha^s_\text{pol} \) solving \( \beta^s(\alpha^s_\text{pol}, \gamma_\varepsilon) = \alpha^s_\text{pol} \). One easily checks that \( \partial_\alpha \mathcal{E}^s(\cdot, \gamma_\varepsilon) \neq 0 \) both in \( \alpha^s_\text{pol} \) and \( \alpha^s_\text{pol} \), see [30, Thms. 4.1-2 and 6.1].

5. **Proof of Theorem 2.1**

We are eventually in the position of proving the assertion of Theorem 2.1. Define \( \ell^s_\varepsilon \) as the smallest integer (larger than 2) such that
\[
\gamma_{\ell^s_\varepsilon} > \max\{\tilde{\gamma}^s_\varepsilon, \tilde{\gamma}^s_\varepsilon\},
\]
where \( \tilde{\gamma}^s_\varepsilon \) and \( \tilde{\gamma}^s_\varepsilon \) appear in Lemma 4.2 and Lemma 4.3. Notice from the proofs of such lemmas that the numbers \( \tilde{\gamma}^s_\varepsilon \) and \( \tilde{\gamma}^s_\varepsilon \) are depending on the specific choice of the potential \( v_3 \). Fix the integer \( \ell > \ell^s_\varepsilon \) from now on. Let moreover \( \eta_\ell^s \) be the largest positive number such that
\begin{itemize}
  \item[i)] the \( n \)-cell \( \{x_1, \ldots, x_n\} \) of any nanotube in \( \mathcal{P}_{\eta_\ell^s}(\mathcal{F}_s^\ell) \) satisfies (3), and moreover
  \item[ii)] the angle \( \gamma^s_i \) at \( x_i \) satisfies \( \gamma^s_i > \max\{\tilde{\gamma}^s_\varepsilon, \tilde{\gamma}^s_\varepsilon\} \), and the angles \( \alpha^1_i, \alpha^2_i \) at \( x_i \) belong to \( (\sigma^s_\varepsilon(\gamma^s_i), 2\pi/3) \), for any \( i = 1, \ldots, n \).
\end{itemize}
The number \( \eta_\ell^s \) gives the smallness of the perturbation and depends on \( v_3, \varepsilon, \) and on \( \ell \).

Let now \( \mathcal{F}_s^\ell \in \mathcal{F}_s^\ell \) be the minimal nanotube from Theorem 4.4 and \( \tilde{\mathcal{F}} \in \mathcal{P}_{\eta_\ell^s}(\mathcal{F}_s^\ell) \) be a nontrivial perturbation, with \( n \)-cell \( \{x_1, \ldots, x_n\} \). In particular, the restrictions on \( \ell \) and \( \eta_\ell^s \)
entail that the topology of $\mathcal{F}_\ell^\gamma$ and that of $\tilde{\mathcal{F}}$ coincide and the energy $E(\tilde{\mathcal{F}})$ can be computed as

$$E(\tilde{\mathcal{F}}) = E(F_n, L) = \frac{1}{2} \sum_{i=1}^{n} \sum_{k=1}^{3} v_2(|x_i - x_{ik}|_L) + \sum_{i=1}^{n} \tilde{E}^s(\alpha_i^1, \alpha_i^2, \gamma_i^s),$$

where $x_{ik}$, $k = 1, 2, 3$, are the three points that are bonded to $x_i$. We recall that $\gamma$ is given by (4) and that we are assuming $\gamma \leq \gamma_\ell$. As $v_2 \geq -1$, we get the bound

$$E(\tilde{\mathcal{F}}) \geq -\frac{3}{2} n + \sum_{i=1}^{n} \tilde{E}^s(\alpha_i^1, \alpha_i^2, \gamma_i^s).$$

By letting $\alpha_i := (\alpha_i^1 + \alpha_i^2)/2$ we have

$$\sum_{i=1}^{n} \tilde{E}^s(\alpha_i^1, \alpha_i^2, \gamma_i^s) \geq \sum_{i=1}^{n} E^s(\alpha_i, \gamma_i^s) \geq \sum_{i=1}^{n} E^s(\alpha_i(\gamma_i^s), \gamma_i^s) = \sum_{i=1}^{n} E^s(\gamma_i^s) \geq n\mathcal{E}^s(\gamma)$$

where we have used Lemma 4.2, the definition of $\mathcal{E}^s(\gamma_i^s)$ as the unique minimizer of $\alpha \mapsto E^s(\alpha, \gamma_i^s)$ from Lemma 4.1, and the definition of $\mathcal{E}^s$ and its convexity from Lemma 4.3 (note that all the angles $\gamma_i^s$, $\alpha_i^1$ and $\alpha_i^2$ belong to the correct range of validity of these properties thanks to the restrictions on $\ell$ and $\eta_\ell^s$). We have obtained,

$$E(\tilde{\mathcal{F}}) \geq -3n/2 + n\mathcal{E}^s(\gamma).$$

As $\mathcal{E}^s$ is decreasing due to Lemma 4.3, from the assumption $\gamma \leq \gamma_\ell$ we deduce that

$$E(\tilde{\mathcal{F}}) \geq -3n/2 + n\mathcal{E}^s(\gamma_\ell) = E(\mathcal{F}_\ell^\gamma).$$

The assertion follows upon noting that equality holds if and only if the perturbation is trivial, since all the convexity and monotonicity properties we used are strict. Note that the proof is independent from the period $L$ of $\tilde{\mathcal{F}}$, which is a small perturbation of the one of $\mathcal{F}_\ell^\gamma$.

6. Numerical evidence for (5)

Theorem 2.1 asserts the stability of $\mathcal{F}_\ell^\gamma$ with respect to all small perturbations fulfilling $\gamma \leq \gamma_\ell$. Under assumption (5) the reach of the result could then be readily extended to all small perturbations. As we are presently not able to present an analytical proof of (5), we resort in recording here some numerical evidence under the specific choice of $v_2$ and $v_3$ given for $f(t) := 1/(2t^{12}) - 1/t^6$ by

$$v_2(r) = \begin{cases} f(r) - f(1.1) & \text{if } 0 < r < 1.1, \\ 0 & \text{otherwise}, \end{cases} \quad v_3(\theta) = 10(\cos \theta + 1/2)^2.$$

These interaction potentials fulfill the general requirements of Section 2. In particular, $v_2$ is a short-ranged, truncated version of the classical Lennard-Jones potential and $v_3$ corresponds to the standard choice in [41]. We argue on the zigzag and the armchair case separately.

Zigzag. Fix $\ell \in \mathbb{N}$ suitably large and let $\mathcal{F}^\gamma_\ell = (F_n^\gamma, L_n^\gamma)$ be the optimal configuration in $\mathcal{F}_\ell^\gamma$. Recall that the $n$-cell $F_n^\gamma$ is characterized by $\alpha^2(\gamma_\ell)$, which is the unique minimum in (29). The reference period is the cell length $L_n^\gamma := 2m(1 - \cos \alpha^2(\gamma_\ell))$, where the integers $m, n, \ell$ are such that $n = 4m\ell$, see (13). As usual, $e_3$ is the axis of the nanotube.

Let now $\tilde{\mathcal{F}} := (F_n, L)$ be a small perturbation of $\mathcal{F}^\gamma_\ell$. Notice that, as in the definition of $\mathcal{P}\gamma(\mathcal{F}_\ell^\gamma)$ in Section 2, we are perturbing the position of the points in the $n$-cell $F_n^\gamma$ and the period $L_n^\gamma$. For every point $x$ in $F_n$, let $x_1, x_2, x_3$ be the points that are bonded to $x$ (modulo $L$). We define the new configuration $\tilde{\mathcal{F}} := (\tilde{F}_n, L)$ by letting its $n$-cell $\tilde{F}_n$ be obtained by rigidly
rotating around the midpoint each bond in $\tilde{F}$ corresponding to a bond in $F_n^a$ parallel to $e_3$, in such a way that the rotated bond is again parallel to $e_3$.

A comparison between the energies $E(\tilde{F})$ and $E(\overline{F})$ is given in Figure 6 with respect to two different reference configurations, namely $\ell = 10$, $m = 2$ ($n = 80$) in the top and $\ell = 20$, $m = 2$ ($n = 160$) in the bottom subfigure. We ran the comparison over $10^6$ random perturbations $\tilde{F}$, $10^2$ of which are plotted in Figure 6. The horizontal axis of the diagram is the mean $\gamma$ of the angles $\gamma_i$ whereas the vertical axis corresponds to the energy (normalized with respect to the number $n$ of atoms). Each random perturbation $\tilde{F}$ is denoted by a cross in the diagram and it is linked via a dotted line to its corresponding configuration $\overline{F}$, indicated by a circle. The decrease of the energy $E(\tilde{F}) \geq E(\overline{F})$ as well as the fact that $\gamma = \gamma_0$ for $\overline{F}$ are apparent, so that (5) is confirmed.

The experiment gives also an illustration of Theorem 2.1, for all random perturbations have a higher energy when compared to the optimal configuration $F_{n}^a$.

**Armchair.** We start from the optimal configuration $F_n^a = (F_n^a, L_n^a)$ corresponding indeed to the unique angle $\alpha^a(\gamma_0)$ realizing the minimum in (29). The period $L_n^a$ is given by (20) for suitable integers $n, m, \ell$ with $\ell$ even and $n = 2m\ell$.

Assume to be given a small perturbation $\tilde{F} = (F_n, L)$ of $F_n^a$. Let us indicate by $b_{ik}$ the bond corresponding to that of $F_n^a$ with endpoints indexed as $(i, 0, k)$ and $(i, 1, k)$ in (15). We define $\overline{F}$ fulfilling (5) by $\overline{F} = T(\tilde{F})$, where the transformation $T$ is the composition of some elementary transformations $T_i$ for $i = 0, \ldots, 3$ defined as follows.

- $T_0$: Rotate each $b_{ik}$ around its midpoint in such a way that the rotated bond lays in an orthogonal plane to $e_3$ and its endpoints are equidistant from $e_3$.
- $T_1$: Rotate each $b_{ik}$ around its midpoint first and then around $e_3$ in such a way that the rotated bond is parallel to the corresponding bond in $F_{n}^a$ and its endpoints are equidistant from $e_3$.
- $T_2$: For each $i$ compute the mean $d_i$ of the distances of the midpoints of the bonds $b_{ik}$ to $e_3$. Translate each $b_{ik}$ by displacing its midpoint in the radial direction with respect to $e_3$ in such a way that the distance of the displaced midpoint from $e_3$ is exactly $d_i$.
- $T_3$: For each $i$ compute the mean $s_i$ of lengths of the bonds $b_{ik}$. Dilate/contract each $b_{ik}$ about its midpoint without rotating it so that the transformed bond has length $s_i$.

Figure 7 and Figure 8 show random perturbations $\tilde{F}$ (crosses) and the transformed configurations $\overline{F} = T(\tilde{F})$ (circles) corresponding to the choices $T \equiv T_2 \circ T_0$ (Figure 7) and $T \equiv T_3 \circ T_2 \circ T_1$ (Figure 8) and the two reference choices for the $n$-cell $\ell = 10$, $m = 4$, $n = 80$ (top), and $\ell = 20$, $m = 4$, $n = 160$ (bottom). Normalized energies with respect to the number of atoms and mean $\gamma$ of the angles $\gamma_i$ are the depicted on the vertical and on the horizontal axis, respectively. We have run the simulation over $10^6$ random perturbation, of which $10^2$ are reported here.

In the case $T = T_2 \circ T_0$ (Figure 7), we have that $E(\overline{F}) \leq E(\tilde{F})$ and that the mean angle $\gamma$ of $\overline{F}$ becomes smaller than $\gamma_0$. In the case $T = T_3 \circ T_2 \circ T_1$ (Figure 8), we have that $E(\overline{F}) \leq E(\tilde{F})$ and $\gamma = \gamma_0$. This last equality could also be proved by means of direct geometric considerations from the definitions of the transformations. In both cases (5) holds.

Note that all perturbations have higher energy with respect to $F_n^a$. 

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Figure 6. Comparison of the energy of a random perturbation $\tilde{F}$ of $F^*_\ell$ with the energy of its corresponding configuration $\mathcal{F}$. We have investigated the two cases $\ell = 10, n = 80$ (above) and $\ell = 20, n = 160$ (below). Energy values (normalized with respect to $n$) are on the vertical axis, while the horizontal axis depicts the mean angle $\gamma$ of each configuration. The energy of each random perturbation $\tilde{F}$ (cross) is higher than the energy of its corresponding configuration $F^*_\ell$ (circle), which satisfies $\gamma = \gamma_{\ell}$. The cross at the bottom corresponds to the minimizer $F^*_{\ell}$. 
Figure 7. Comparison of the energy of a random perturbation $\tilde{F}$ of $F^a_\ell$ with the energy of its transformed configuration $T_2(T_0(\tilde{F}))$. We investigate the two cases $\ell = 10$, $n = 80$ (above) and $\ell = 20$, $n = 160$ (below). Energy values (normalized with respect to $n$) are on the vertical axis, while the horizontal axis depicts the mean angle $\gamma$ of each configuration. The energy of each random perturbation $\tilde{F}$ (cross) is higher than the energy of its transformed configuration $T_2(T_0(\tilde{F}))$ (circle), which satisfies $\gamma \leq \gamma_\ell$. The cross at the bottom corresponds to the minimizer $F^a_\ell$. 
Figure 8. Comparison of the energy of a random perturbation $\tilde{F}$ of $F^*_{a\ell}$ with the energy of its transformed configuration $T_3(T_2(T_1(\tilde{F})))$. We investigate the two cases $\ell = 10, n = 80$ (above) and $\ell = 20, n = 160$ (below). Energy values (normalized with respect to $n$) are on the vertical axis, while the horizontal axis depicts the mean angle $\gamma$ of each configuration. The energy of each random perturbation $\tilde{F}$ (cross) is higher than the energy of its transformed configuration $T_3(T_2(T_1(\tilde{F})))$ (circle), which satisfies $\gamma = \gamma_{\ell}$. The cross at the bottom corresponds to the minimizer $F^*_{a\ell}$. 
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