Annealing a Magnetic Cactus into Phyllotaxis

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The appearance of mathematical regularities in the disposition of leaves on a stem, scales on a pine-cone and spines on a cactus has puzzled scholars for millennia; similar so-called phyllotactic patterns are seen in self-organized growth, polypeptides, convection, magnetic flux lattices and ion beams. Levitov showed that a cylindrical lattice of repulsive particles can reproduce phyllotaxis under the (unproved) assumption that minimum of energy would be achieved by 2-D Bravais lattices. Here we provide experimental and numerical evidence that the Phyllotactic lattice is actually a ground state. When mechanically annealed, our experimental “magnetic cactus” precisely reproduces botanical phyllotaxis, along with domain boundaries (called transitions in Botany) between different phyllotactic patterns. We employ a structural genetic algorithm to explore the more general axially unconstrained case, which reveals multijugate (multiple spirals) as well as monojugate (single spiral) phyllotaxis.

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I. INTRODUCTION

Symmetrical morphologies and regular patterns in living organisms (Fig.\textsuperscript{1}) have been credited with originating the idea of beauty, the notion of art as an imitation of nature, and humanity’s first mathematical inquiries \cite{1-4}. The fascinating symmetrical patterns of organs in plants, called phyllotaxis \cite{1,3}, were known to the Romans (Pliny) and ancient Greeks (Theofrastus), while early recognitions are found in sources as ancient as the Text of the Pyramids \cite{1}. Leonardo da Vinci \cite{5}, Andrea Cesalpino, and Charles Bonnet \cite{6} studied phyllotaxis in the modern era. Kepler proposed that the Fibonacci sequence \{1, 2, 3, 5, 8…\}, where each term is the sum of the two preceding ones \cite{7}, describes these phyllotactic patterns.

A discipline that thrived on multidisciplinary interactions \cite{8}, phyllotaxis found its standard mathematical description when August and Louis Bravais \cite{9} introduced the point lattice on a cylinder to represent the dispositions of leaves in 1837 (see Fig.\textsuperscript{2}), thirteen years before August’s seminal work on crystallography \cite{10}. Unfortunately botanists neglected the work of the Bravais brothers, and it wasn’t until Church rediscovered it eighty years later that more progress was achieved in the field \cite{11}.

The geometrical description of cylindrical phyllotaxis relies, in the simplest case, on the phyllotactic lattice introduced by the Bravais brothers \cite{1,3,9}. It consists of a so-called generative spiral of divergence angle $\Omega$. We can visually decompose the resulting lattice in crossing spirals that join nearest neighbors, as in Fig.\textsuperscript{2}, which botanists call parastichies. It is a fundamental observation (made first by Kepler) that the numbers $n$, $m$ of crossing parastichies needed to cover the lattice are consecutive terms of the standard Fibonacci sequence, or less frequently the variants obtained by changing the second term, also called Lucas numbers: 1, 3, 4, 7, 11… and 1, 4, 5, 9… often referred to as second and third phyllotaxis. From that, one can prove that the divergence angle of the generative spirals in plants assumes values close to \cite{2,12}

$$\Omega_p = \frac{360\degree}{(\tau + p)}$$

where $p = 1, 2, 3$ denotes first, second or third phyllotaxis and $\tau = (1 + \sqrt{5})/2$ is the golden ratio. For more than one generative spiral (“multijugate” phyllotaxis), parastichies share a common divisor $(n, m) = (kn', km')$, $k$ being the number of generative spirals \cite{2,14}. Not unlike domain boundaries in crystals, plants show kinks between domains, called transitions by botanists \cite{1,13}.

In the last 50 years, phyllotactic patterns have been seen or predicted outside of botany: polypeptide chains \cite{14,15}, tubular packings of spheres \cite{16}, convection cells \cite{17}, layered superconductors \cite{18}, self-assembled microstructures \cite{19}, and cooled particle beams \cite{20,21}. While it is still debated whether such systems might shed light on botanical phyllotaxis, the occurrence of such mathematical regularities outside of botany is fascinating and leads to generalizations that – unlike quasistatic botany – allows for dynamics \cite{22}.

In a groundbreaking work Levitov recognized phyllotaxis in vortices of layered superconductors \cite{18}. He next described how phyllotactic patterns represent states of minimal energy of a cylindrical lattice (that is of a lattice with cylindrical boundary conditions) of mutually repelling objects, the repulsion mimicking the interactions between spines, leaves, or seeds in plant morphology \cite{23,24}. Yet such a constraint to a lattice is absent both in botany and in the physical systems to which this energetic model might apply, such as adatoms or low-density electrons on nanotubes and ions or dipolar...
molecules in cylindrical traps.

Following up on earlier work that focused on the dynamics of rotons and solitons in physical phyllotactic systems \[22\], we provide here a detailed experimental and numerical demonstration that Levitov’s constraint is not necessary, and that the lowest energy states of repulsive particles in cylindrical geometries are indeed phyllotactic lattices. In addition, we describe the experimental and numerical generation of multijugate phyllotaxis, static kink-like domain boundaries between different phyllotactic lattices, and unusual disordered yet reflection-symmetric structures that may be a static relic of soliton propagation.

We show that when a “magnetic cactus” of magnets (spines) equally spaced on co-axial bearings (stem) with south poles all pointing outward is annealed, it precisely reproduces botanical phyllotaxis. When studied numerically via a structural genetic algorithm, the fully unconstrained case reveals both multijugate and mono jugate phyllotaxis. In addition, we describe the experimental and numerical demonstration that Levitov’s constraint is not necessary, and that the lowest energy states of repulsive particles in cylindrical geometries are indeed phyllotactic lattices. In addition, we describe the experimental and numerical generation of multijugate phyllotaxis, static kink-like domain boundaries between different phyllotactic lattices, and unusual disordered yet reflection-symmetric structures that may be a static relic of soliton propagation.

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In section II we describe the statics of repulsive particles in cylindrical geometries. In section III we detail the experiment on the magnetic cactus. In Section IV we discuss the more general case of multijugate phyllotaxis.

II. PHYLLOTAXIS OF REPULSIVE PARTICLES IN CYLINDRICAL GEOMETRIES

In this section we will recall Levitov’s model \[23, 24\] and some of our own findings \[22\]. Following Levitov, let us assume that the lowest energy configuration for a set of particles with repulsive interactions, confined to a cylindrical shell of radius \( R \), is a helix with a fixed angular offset \( \Omega \) between consecutive particles and a uniform axial spacing \( a \), as in Fig. 1 (this so far unproved ansatz will be investigated later both numerically and experimentally). For a generic pair-wise repulsive interaction \( v_{ij} \) between particles \( i \) and \( j \), the energy of the helix is

\[
V = \frac{1}{2} \sum_{i \neq j} v_{ij}.
\]

Since the lattice structure is defined by \( \Omega \), we can write \( V(\Omega) \).

In Fig. 3 we plot \( V(\Omega) \) for various values of the ratio \( a/R \): for specificity we employed a dipole dipole interaction \( v_{ij} = p_i \cdot p_j / r_{ij}^3 - 3(p_i \cdot r_{ij})(p_j \cdot r_{ij}) / r_{ij}^5 \), repulsive at the densities considered here. However, the following considerations only depend upon geometry and therefore apply to a vast range of reasonably behaving, long range repulsive interactions.

When \( a/R \gg 1 \), the angle \( \Omega = \pi \) maximizes distance between neighboring particles and therefore \( V(\Omega) \) has a minimum in \( \pi \). The angle between second nearest neighbors along the helix is \( 2\pi \), which means that they face each other. And thus, as the density increases, interaction between the facing second nearest neighbors becomes predominant, and \( \Omega = \pi \) is not a minimum for \( V(\Omega) \) anymore. If we shift the helical angle from \( \pi \), the repulsive interaction between second nearest neighbors is reduced, with minimal penalty from nearest neighbors. In terms of \( V(\Omega) \), that means a local maximum \( \Omega = \pi \).

This argument can be iterated for every commensurate winding that allows particles separated by \( j \) neighbors to face each other. As density increases further, the angles \( 2\pi/3 \) and \( 4\pi/3 \) also become unfavorable due to third-neighbour interactions. Any commensurate spiral of divergence angle \( \Omega = 2\pi i/j \) with \( i, j \) relatively prime corresponds to a configuration where each particle faces each \( j \)th neighbor. For every \( j \) there will be a value of \( a/R \) low enough such that \( \Omega = 2\pi i/j \) is a local maximum,
which we call a peak of rank \( j \).

The proliferation of peaks for increasing linear density is shown in Fig 3. We can see that at high density, peaks of equal rank are nearly degenerate; that is natural, since their principal defining energetic contribution arises from particles facing each other at a distance \( ja \). The minima also become more nearly degenerate as the density increases. That can be explained intuitively, since for angles incommensurate to \( \pi \) each particle “sees” the others as incommensurately smeared around the cylinder, and is therefore embedded in a nearly uniform background charge from the other particles. The degenerate energy of the ground state can be well approximated by an uniform continuum distribution \( \epsilon_0 \), whereas the energy of a peak of order \( j \) will be \( V(2\pi i/j) \simeq v(ja) + \epsilon_0 \), where \( v(r) \) is the energy of two particles facing at a distance \( r \): for our dipole interaction \( v(ja) = p^2/a^2 j^2 \).

The first step to calculate the degeneracy of our system at a given density, is to find the corresponding maximum rank of the peaks. As all of the peaks of the same rank have the same energy, and appear in the spectrum together, we can focus on the emergence of \( 2\pi/J \). For \( a/R \ll 1 \), this new peak will emerge when the distance between particles separated by \( ja \) equals that of particles separated by \( 2\pi R/J \). Therefore one finds for the maximum rank

\[
J = \left\lceil \sqrt{\frac{2\pi R}{a}} \right\rceil,
\]

which as expected only depends on purely geometrical parameters. A little more tricky is to compute the degeneracy, given \( J \). The set of all the peaks has the cardinality of the class of all the fractions \( i/j \), with \( i, j \) coprime and \( j \leq J \). This can be considered as the disjoint union of other classes, called Farey classes of order \( j \), defined as follows: \( P_j = \{ \Omega = 2\pi i/j \} \) for \( i, j \) coprime and \( i \leq j \), i.e. all fractions in lowest terms between \( 0 \) and \( 1 \) whose denominators do not exceed \( j \).

The union of all Farey classes up to a certain order \( J \) has the cardinality of the set of peaks for a spectrum of maximum rank \( J \). Now, the cardinality of \( P_j \) is know from number theory to be Euler’s totient function, \( \phi(j) \). Therefore, the degeneracy \( D \) of the energy minima for a system with a maximum peak rank \( J \) is

\[
D = \sum_{j=1}^{J} \phi(j) = \frac{3}{\pi^2} J^2 + O(J \log J),
\]

which, from Eq. 2 scales as \( D \sim 2R/a \).

Finally, we recall that the order \( j_1, j_2 \) of the peaks bracketing a minimum relates to its structure in a straightforward way: the helix corresponds to a rhombic lattice where each particle has its nearest neighbors at axial displacements of \( \pm a j_1, \pm a j_2 \) and second nearest neighbors at \( \pm a (j_1 + j_2) \) or \( \pm a (j_1 - j_2) \). Also, \( j_1 \) and \( j_2 \) give the number of crossing secondary spirals (parastichies) needed to cover the lattice by connecting nearest neighbors.

For completeness, let us now follow Levitov, and consider the adiabatic evolution of our system as the linear density is increased. As new sets of maxima and minima emerge, the true minimum goes through a series of quasi-bifurcations, the consequence of an elusive symmetry whose explanation goes beyond our scope. Suffice it to say that the system evolves quasi-statically from one of these optimal \( \Omega \) to another as \( R/a \) increases, asymptoting to the golden angle \( \Omega_1 = 2\pi/(\tau + 1) \) \( \tau = (1 + \sqrt{5}) /2 \), ubiquitous in botany, as each minimum is bracketed by peaks whose ranks, because of the Farey tree structure described above, are consecutive elements of the Fibonacci sequence. Occasional “wrong turns” at later stages, will not shift the convergence too far from the golden angle, yet the Fibonacci structure would be lost. However if one or two consecutive wrong turns happen at the second or second and third bifurcations the system will converge to the alternative angles of second or third phyllotaxis, given by Eq. 1.

We have only surveyed so far spiraling lattices generated by a single helix. A straightforward generalization gives multijugate phyllotaxis, when two or more elements grow at the same axial coordinate. This case, which Levitov does not explore, can be easily mathematically reduced to monojugate case, by considering two or more replicas of the phyllotactic lattice as in Fig 2. In our experimental realization we restrict ourselves to the monojugate phyllotaxis, and explore multijugate only numerically.

### III. A MAGNETIC CACTUS

There is a long history of experimental reproductions of phyllotactic patterns. Recently, Doady et al. described phyllotaxis in terms of dynamical systems and then veri-
A. Experimental Apparatus

We built a magnetic cactus by mounting permanent magnets (spines) on stacked co-axial bearings (a stem) which are free to rotate about a central axis, as in Fig. 4. All the magnets point outward, to produce a repulsive interaction between all magnet pairs. To avoid effects of gravity, the apparatus rests in the vertical position, and is non-magnetic. We built two different versions, the second with magnets twice as long as the first, as to have a larger effective radius which gives three rather than two stable structures.

We employed cylindrical permanent iron-neodymium magnets, 1.2 cm long and 0.6 cm in diameter. They are mounted on fifty aluminum rings of 2.2 cm outer diameter, each affixed to a non-magnetic radial ball bearing (acetal/silicon, Nordex) as in Fig. 4. These unit rings are evenly spaced on an aluminum rod in a stacked structure of 39.9 cm axial length.

At static equilibrium we measure separation angle between each magnet element, by rotating the cactus until a magnet element aligns with the reference wires. A telescope and a vertical viewing slit accompanied by two vertical reference wires assist in data acquisition.

B. Annealing

By measuring the dipole-dipole interaction between an individual magnet pair, we can reconstruct the curve of the lattice energy \( V(\Omega) \) as a function of the angular offset \( \Omega \) between magnets. We find that the first arrangement, with short magnets, admits two minima, given by the angles of Eq. 1 for \( p = 1, 2 \). The second arrangement, with long magnets, has three minima corresponding to the angles of Eq. 1 for \( p = 1, 2, 3 \), one of which (\( p = 3 \)) is a weak metastable minimum. These divergence angles of stable helices are very close to those predicted by phyllotaxis, of Eq. 1, and are all accessible by experimental procedure described below.

Before every data acquisition, the cactus is disordered and then athermally annealed into a low-energy state. The protocol involves repeatedly winding the bottom-most magnet to generate an ever-tightening spiral, until an explosive release of energy disorders the lattice. Next, an independent external magnet is oscillated in small circular motions near randomly chosen points while the cylinder as a whole is slowly rotated, to further randomize magnet orientations. After 10-30 second of mechanical annealing through applied vibrations, the system consistently enters a robust ordered state which does not anneal further on experimental timescales.

C. Results

Figure 5 reports the experimental results for both arrangements by plotting the measured angle between con-
FIG. 5: Top: A 3-D rendering of the experimental data and the corresponding Bravais lattice of the magnetic cactus annealed in a spiral configuration of divergence angle $\Omega_1$ and parastichies (2,3) (blue and red dashed lines), Fibonacci numbers. The bottom two panels show the experimentally measured angular offsets $\omega$ between successive magnets for magnetic cacti with short (middle) and long (bottom) magnets, plotted versus magnet index, which simply counts the number of magnets along the axis. Flat regions are perfect spirals while steps are boundaries between different phyllotactic domains. The dotted lines give the phyllotactic angles $\Omega_1$, $\Omega_2$, $\Omega_3$ and $2\pi-\Omega_2$ defined in the text, whereas the dashed lines are minima of the magnetic lattice energy (insets) calculated by interpolating the measured pair-wise magnet-magnet interaction. Data reproduced from Ref. [22].

FIG. 6: A kink between domains of first and second phyllotaxis. From top to bottom: a 3-D rendering, and its unwound 2-D Bravais lattice from numerical simulations. Below, the same kink plotted as angle increments between successive magnets for the experimental system (crosses) and numerical simulation (circles). The more narrow (short-magnet) cactus self-organizes into the spirals with divergence angles precisely reproducing those of first phyllotaxis, $\Omega = \Omega_1$, and second phyllotaxis, $\Omega = \Omega_2$, as in Eq. (1). When the results are represented in a 2-D lattice, as in the top of Fig. 5, parastichies can be drawn. As parastichial numbers for $\Omega = \Omega_1$ we find the Fibonacci numbers (2,3), and for $\Omega = \Omega_2$, the Lucas numbers (3,4), as seen also in botany. The larger-radius system also forms first and second Phyllotaxis helices, as well as limited domains of third phyllotaxis [with $\Omega = \Omega_3$ and Lucas numbers (1,4)], bracketed by domains of second phyllotaxis. The insets of Fig. 5 show the magnetic interaction energy $V(\Omega)$ of the lattice obtained by interpolating measured values for the pair-wise magnet-magnet interaction, plotted as a function of divergence angle $\Omega$. As we can see, local minima correspond to phyllotactic angles. Figure 5 also shows that in many instances the system fragments into two or three distinct domains whose domain walls always share a common parastichy, as seen in botany [1, 13], and as expected in physics for a quasi-one-dimensional degenerate system. We have computed numerically one such transition via dynamical simulations in a velocity-Verlet algorithm, in the following way: we start from a crude static step-like kink as an initial condition, and allow it to radiate energy in the form of phonons waves until it stabilizes in a kink with superimposed vibrations; we then average this configuration over time, to remove these residual oscillations. When the result is
used as new initial conditions, it proves to be a static kink. Fig. 7 reports our numerical results for a kink in a system whose size and interaction reproduces the physical realization of the magnetic cactus, along with the experimental data for such a kink. The match is essentially perfect, indicating that the dissipative (i.e. frictional) forces neglected in our model do not significantly affect the static configurations. We apply the same numerical procedure to calculate a kink in a system of larger degeneracy, among domains which are absent in our physical realization. We use a smaller $a/R$ ratio and a different interaction between particles (ideal dipole instead of physical dipole). The result shown in Fig. 7 reproduces the same qualitative shape of the previous, lower degeneracy case. Similar kinks are present also in a fully unconstrained cactus, one in which the particles can move along the axis, and are found in early generations of our structural genetic algorithms (see below). Finally, these kinks can travel as novel topological solitons, with a rich phenomenology that is explained elsewhere [22, 29].

Finally, in the system with longer magnets, we occasionally found intriguing yet hard-to-interpret configurations that contain two nearly reflection symmetric domain boundaries. Fig. 8 reports two such configurations, measured in independent experimental runs. Although we do not have a firm explanation for these structures, we speculate that they form as frozen-in soliton waves that initiated symmetrically at both ends of the structure, upon release of the wound-up elastic energy during initial preparation. Indeed an analytical, continuum theory for phyllotactic solitons which we have developed recently, and which explains results of the dynamical simulations also supports the existence of similar frozen-in pulses [29].

IV. FULLY UNCONSTRAINED CACTUS: STRUCTURAL GENETIC ALGORITHM

Our experimental apparatus is not fully unconstrained: the axial coordinates of the dipoles are fixed, and so only the azimuthal movement is allowed. While this is an huge improvement toward the original helical constraint of Levitov, many (most) physical systems that could manifest phyllotactic patterns do not possess such a lesser azimuthal constraint. To corroborate and extend our experimental results to a completely unconstrained system, we seek the energy minimum in a set of repulsive
particles that can move axially as well as angularly on a cylindrical surface, via a non-local numerical optimization. To this purpose, we developed a structural genetic algorithm.

A. Genetic Algorithm

A genetic algorithm is a method of optimization that mimics evolution to find the absolute minimum in a function which shows a large number of metastable minima. The coordinates of the energy functions are called genes, and a set of genes is a particular specification of value for those variables. The routine typically starts with a set of “parents”, or specific points in the domain of the energy function. At each stage of the routine, parents “mate” to produce children via exchange of genes: a subset of the coordinates of each of the two configurations are swapped, therefore generating new points in the energy domain, called children. Each of those children is then locally relaxed to a minimum via a local search. The new population of parents and children undergoes genetic selection and only the fittest (the lowest energy ones) form a new population.

There are many different implementations of this general idea: care is taken not to lose genetic diversity during selection, to avoid a population of almost identical replicas: that is usually achieved with a more or less skilled genetic selection, which might retain less genetically fit individuals, and often by introducing mutations in the form of random alteration of the gene sequence, which would hopefully prevent the routine from getting stuck around a metastable region. Choice of parameterization of the structures (genes) and mating (crossover) is crucial to the performance of the algorithm.

About fifteen years ago, Deaven and Ho [30] introduced a so-called structural genetic algorithm, which proved particularly efficient in minimizing the energy of physical structures, as it allows for physical intuition in defining the genes and mating procedure. With it, they found the C_{60} fullerene structure as a ground state of 60 carbon atoms interacting with suitable atomic potentials [30] and solved the celebrated Thomson problem of repulsive charges on a sphere [31], a task quite similar to ours.

B. Our Algorithm

In our implementation we use a population of 10 members. Each member represents a configuration of 101 particles on the cylinder: more explicitly, the genetic structure of each member \( P_k, k = 1, \ldots, 10 \), of the population is a set of variables, or \( P_k = \{ \theta_i, z_i \}_{i=1}^{101} \) which specifies, in cylindrical coordinates, the positions of the particles composing its structure. The particles interact via a pair-wise inverse quadratic repulsion \( V = V_o(r_o/r)^2 \), where \( r \) is the three dimensional distance between particles; we introduce a confining potential in the form of an external axial square-well of width \( L \), which sets the length of the cylinder, and hence the density. The choice of the pairwise interaction is not fundamental, as long as it is long ranged, repulsive, and well behaved [22]. Our particular choice simply speeds up the computation.

We generate the first population randomly. At each step, we randomly couple mates, and exchange their genes employing the following mating procedure: we order the genes by increasing axial coordinates \( z_1 < z_2 < \cdots < z_{101} \) and swap the first \( 1 < n < 101 \) genes, where \( n \) is a random number, between randomly selected parents. The children obtained in this way are then relaxed to a stable structure via a standard conjugate gradient algorithm. We then prepare the new generation by selecting the lowest energy individuals in the population of parents and children, yet making sure that the energy difference between members does not fall below a certain threshold, to preserve genetic diversity: when new children cannot produce a new population of 10 in accordance with the energy threshold, we introduce mutations by randomly altering a certain number of members.

C. Numerical Results

During the structural evolution, the earliest populations contain metastable disordered states. Members of intermediate populations show kinks between domains of different divergence angle, configurations which are also seen experimentally. After fifteen to twenty generations, the algorithm typically converges to a single crystalline domain.

Figure 9 reports the energy of the fittest member of the population in every generation. Early generations return very high energy configurations corresponding to disordered metastable states. Intermediate generations show populations of phyllotactic domains separated by kinks between. Finally, the algorithm converges to a single crystalline domain in the bulk (deformations at the boundaries simply accommodate the system to the confining potential).
fit structure progressively decreases in energy in intermittent steps separated by plateaux. The final converged results form well-defined two-dimensional cylindrical crystals away from boundaries.

Figure 10 shows the crystalline structure to which the algorithm converges, for $R = 1.65L/N$, a single spiral with $\Omega = \Omega_1$, as defined in Eqn. (1), corresponding to first phyllotaxis with parastichies $(1, 2)$. A plot of $\Delta z = z_{i+1} - z_i$-returns the value $L/N$ in the bulk, which implies a single generative spiral. This choice of $RN/L$ corresponds to a density close to that of our experimental apparatus.

**V. MULTIJUGATE PHYLLOTAXIS**

For highly degenerate systems the genetic algorithm returns configurations with more than one generative spiral, corresponding to what in botany is called multijugate phyllotaxis [2]. We have seen before that helices make cylindrically symmetric lattices. On the other hand, every cylindrically symmetric lattice can be decomposed into a suitable number of equispaced generative spirals [2, 9]. That is accomplished by discretizing the cylinder along its axis into equally spaced rings and then assigning at each ring $n$ sites, equally spaced and separated by a $2\pi/n$ angular shift. As before, each ring is shifted consecutively by a divergence angle $\Omega$. The case $n = 2$ is shown at the top of Fig. 11. The case $n = 1$ is shown in our experimental arrangement of Fig. 1.

By decomposing the $n$-jugate cylindrical lattice into $n$ lateral replicas of single-spiral lattice, as in Fig. 2, the reader is easily convinced that multijugate phyllotaxis reduces to the previously described monojugate case. All the considerations above apply, provided that one now takes the periodicity to be $2\pi/n$, and the distance between rings to be $na$ (with, as before, $a = L/N$). It follows that a $n$-jugate configuration will have local maxima in $2\pi i/n$, $i = 1 \ldots n$ and, following the discussion of section II, one finds that there will be other local maxima corresponding to angles $2\pi/n \times i/j$ when $j \leq [J/n]$, and $J$ is the maximum rank given by Eq. 2.

Note now that if two multijugate lattices of jugation $n, n'$ have a peak in the commensurate angle $2\pi i/j$, then the energy of the peak is the same, as is shown in Fig. 11 bottom, which compares the plots of the energy of such an arrangement for different values of $n$. In fact both configurations correspond to particles facing each other after $j/n$ and $j/n'$ rings, and therefore at the same distance $na \times j/n = n'a \times j/n' = ja$, independent of $n$ or $n'$. For small $n$ the minima in the energy of $n$-jugate config-
urations essentially degenerate with the monojugate one previously explored. For large $n$ they have higher energy. If $a/R$ is small enough, the threshold is $n > J$, as interaction between particles on the same ring become comparable to those facing in the minimal monojugate peak.

VI. CONCLUSION

We have studied the lowest energy configurations of repulsive particles on cylindrical surfaces, both experimentally and numerically. We have found that they correspond to the spiraling lattices seen in the phyllotaxis of living beings, both monojugate and multijugate. By establishing experimentally and numerically that phyllotactic point lattices are ground states in the very general geometric scenario of unconstrained repulsive particles on cylinders, we have opened the study of phyllotaxis to a much wider range of annealable physical systems where the particles could be electrons, adatoms, ions, dipolar molecules, nanoparticles, etc. constrained by external potentials.

Unlike plants, these multifarious, non-biological Phyllotactic systems could access various degrees of dynamics, providing new phenomenology well beyond that available to over-damped, adiabatic botany. We have reported elsewhere [22] on the dynamical richness of this physical phyllotaxis, including classical rotons and a large family of novel, inter-converting topological solitons.

[1] I. Adler, D. Barabe, R. V. Jean, Ann. Bot. 80 231-244 (1997).
[2] R. V. Jean, Phyllotaxis: A Systemic Study in Plant Morphogenesis (Cambridge Univ. Press, Cambridge, 1994).
[3] http://maven.smith.edu/~phyllo/ provides an overview and useful applets.
[4] N. Grew, The anatomy of plants (Johnson Reprint Corp., New York, 1965).
[5] E. MacCurdy, The notebooks of Leonardo da Vinci (Braziller, New York, 1955).
[6] C. Bonnet, Recherches sur l'usage des feuilles dans les plantes (E. Luzac, fils, Göttingen and Leiden, 1754).
[7] L. E. Sigler, Fibonacci's Liber Abaci (Springer-Verlag, New York, 2002).
[8] V. Jean, B. Denis, Multidisciplinarity: a key to phyllotaxis in Symmetry in Plants 619-653 (World Scientific, 1998).
[9] L. Bravais, A. Bravais Annales des Sciences Naturelles Botanique 7 42-110, 193-221; 291-348 8 11-42 (1837).
[10] A. Bravais, J. École Polytech. 19 1-128 (1850).
[11] A. H. Church, On the Relation of Phyllotaxis to Mechanical Laws. (Williams and Norgate, London, 1904).
[12] R. V. Jean, D. Barabe, Ann. Bot. 88 173-186 (2001).
[13] A. Frey-Wyssling, Nature 173 596 (1954).
[14] S. F. Albdurur, K. A. Laki, J. Theor. Biol. 104 599-603 (1983).
[15] R. O. Erickson, Science 181 705-716 (1973).