Competition between magnetic frustration and spin dimensionality in the ground state of the classical antiferromagnetic $n$-vector model with arbitrary $n$

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A new method to characterize the strength of magnetic frustration is proposed by calculating the dimensionality of the absolute ground state of the classical nearest-neighbor antiferromagnetic $n$-vector model with arbitrary $n$, describing interactions between spins that reside on the vertices of frustrated molecules. Spins having more than three components may minimize the energy more effectively against frustration by allowing bigger angles between nearest-neighbors, leading to ground states in higher than three-dimensional spin space. Platonic solids in three and four dimensions, which have equivalent sites and consist of only one type of polygon, have lowest-energy configurations in a number of spin dimensions equal to their real-space dimensionality. The same is true for the Archimedean solids, which have equivalent sites but consist of more than one type of polygon. Fullerene molecules and geodesic icosahedra, whose sites are not equivalent, can produce ground states in as many as five spin dimensions. It is also found that the introduction of interactions in the next higher dimension does not necessarily lower the energy immediately, but it typically takes a finite coupling for this to happen. Frustration is also characterized by the maximum value of the ground-state energy when the exchange interactions are allowed to vary, which reveals symmetry patterns in the magnetic behavior and that symmetry and connectivity can play a more important role than molecular size in alleviating frustration.

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I. INTRODUCTION

Strongly-correlated electron interactions can very often be described by effective Hamiltonians involving only spin degrees of freedom [1, 2]. A Hamiltonian describing interactions between localized spins is the nearest-neighbor antiferromagnetic Heisenberg model, which has been considered on various lattices and clusters. A single spin is located at each vertex of the underlying structure and only nearest-neighbors interact. It is of special interest when the underlying structure’s topology is frustrated, meaning that due to competing interactions not all nearest-neighbor spin pairs are simultaneously antiparallel in the ground state when the spins are classical [3, 4].

In the nearest-neighbor antiferromagnetic Ising model the spins interact only along a single direction in spin space, making it maximally anisotropic [5]. The dimensionality of the interactions can be increased to two and three in a controlled way, by allowing the spins to interact in the added spin dimensions with increasing strength. The corresponding models are the XY in two dimensions and the anisotropic Heisenberg, or XXZ, in three. When the interaction components become equal in all two or three spin directions one gets respectively the XX model and the isotropic XXX Heisenberg model. Frustration manifests itself in all these cases but is more pronounced in the XXX limit, where magnetic anisotropy plays no role. Then any deviation from antiparallel nearest-neighbor classical spins in the zero-magnetic-field ground state and any magnetization or susceptibility discontinuities in an external field are solely due to the frustrated connectivity of the structure that hosts the spins.

In the case of bipartite structures, which lack frustration, nearest-neighbors point at antiparallel directions in spin space in the ground state. The ground-state energy is minimal and the spin configuration is collinear already at the Ising limit. Perhaps the simplest structures associated with frustration are polygons with an odd number of vertices with edges that correspond to exchange interactions of the same strength. The smallest member of this family is the triangle, for which in the classical Ising ground state not all three spin pairs can be simultaneously antiparallel, demonstrating the effect of frustration. Going to the XY model frustration is relaxed by allowing the spins to interact in a plane, and the lowest-energy configuration becomes coplanar and its energy is continuously lowered with increasing coupling in the second spin direction. This holds up to the XX limit, where the ground-state energy assumes its minimum value and does not decrease any further if interaction in the third spin direction is introduced. The energy per bond is then equal for any of the three bonds. Analogous results hold for any member of the family of the polygons with an odd number of vertices, even though frustration gets weaker with size [6]. The dimensionality of the lowest-energy configuration in spin space is equal to the real-space dimensionality of the polygon.

A basic question for molecules including at least one type of odd-membered polygon is if the assembly of more than one frustrated unit further strengthens frustration, by increasing the ground-state energy per bond and making the lowest-energy spin configuration noncoplanar. For the icosidodecahedron, an Archimedean solid with vertex-sharing triangles, this is not true, as the ground-
state energy is simply the sum of the energies of the individual triangles and the corresponding configuration is coplanar \[n\]. The same is true for the ground-state energies of two other Archimedean solids, the truncated tetrahedron and the truncated icosahedron, however the corresponding spin configurations are noncoplanar \[11\, 12\]. Their frustrated units are triangles and pentagons respectively, which are isolated from one another. On the other hand for the icosahedron, a Platonic solid with edge-sharing triangles \[13\], not only is the lowest-energy configuration three-dimensional, but also the corresponding energy per bond is higher than the one of an isolated triangle \[9\, 14\, 15\]. A similar result holds for another Platonic solid, the dodecahedron, which is formed from edge-sharing pentagons \[16\].

In this paper magnetic frustration is characterized by calculating the spin-space dimensionality of the absolute ground state of the nearest-neighbor classical \(n\)-vector model, when it describes interactions between spins with \(n\) components residing on vertices of frustrated molecules. For \(n = 1\) one gets the Ising model, for \(n = 2\) the XY model, and for \(n = 3\) the Heisenberg model, but \(n\) is allowed to be greater than three until the ground-state energy does not decrease any more. In this way the spins are given more space for nearest neighbors to attempt to direct themselves in an antiparallel fashion, not being restricted by the three dimensions of the Heisenberg model but only by the frustrated topology of the underlying structure. The ground-state energy is monitored from the Ising limit up to the dimension where it assumes its absolute minimum. This is done in a continuous way, allowing the detection of dimensionality windows where the lowest energy remains constant before it starts to decrease again. The point-group symmetry of the molecule determines the number of independent exchange interactions, with edges connected by symmetry operations taken to correspond to the same interaction strength. A second way with which the strength of magnetic frustration is characterized is by calculating the maximum possible ground-state energy, when there is more than one independent exchange interaction and these are allowed to vary in the number of spin dimensions of the absolute ground state. In this paper molecules having up to four unique exchange interactions are considered.

The task to investigate the properties of the \(n\)-vector model when \(n > 3\) has been mostly undertaken in the continuum case \[17\, 27\]. Here Platonic solids in three and four spatial dimensions are considered, as well as Archimedean solids, fullerene molecules \[28\], and geodesic icosahedra \[29\, 30\]. These molecules include at least one type of odd-membered polygon.

It is found that for the icosahedral Platonic solids and their fourth-dimensional real space analogues (Table \[I\]) the spin-space dimensionality of the lowest-energy configuration is equal to the real-space dimensionality of the molecule, showing a direct correlation between spin space and real space. Frustration is stronger than the one of the isolated polygon these molecules are made from, since the ground-state energy per bond for them is higher. On the other hand the truncated dodecahedron and the rhombicosidodecahedron, which are Archimedean solids (Table \[II\]), have three-dimensional ground states and frustration is not stronger than the one at the level of the isolated polygons they are made of. However in the case of the snub dodecahedron the assemblage of the individual polygons increases the maximum possible frustration.

For the fullerene molecules (Table \[III\]) the lowest-energy configuration spin-space dimensionality is typically 3 or 4, with the ones with icosahedral symmetry belonging to the former case. Frustration is minimal for them, as the maximum ground-state energy does not exceed the ground-state energy of an isolated pentagon. This demonstrates a direct correlation between magnetic behavior and symmetry and shows that frustration does not necessarily decrease with the size of the fullerenes, but rather there is a specific symmetry that minimizes it. Molecules with nonisolated pentagons and \(T_d\) symmetry have ground states in \(n = 5\). \(T_d\) fullerenes share patterns of common magnetic behavior not only in the dimensionality of the absolute ground state, but also in the correlations of the maximally frustrated ground state. Fullerenes more generally show that the maximum ground-state energy per bond does not necessarily decrease with the number of vertices, showing that symmetry is also important in determining the strength of frustration, as was also pointed out earlier for the icosahedral members of the family.

The smallest geodesic icosahedron, the pentakis dodecahedron, develops its absolute ground state in \(n = 4\) (Table \[IV\]). Its maximum ground-state energy per bond equals the ground-state energy per bond of the icosahedron, even though it develops in four dimensions. The ground state of the next bigger member of the family, the pentakis icosidodecahedron, develops in \(n = 5\), and its maximum possible energy per bond value is higher than the one of the icosahedron. A bigger member of the family, the hexapentakis truncated icosahedron, has also a five-dimensional ground state. Its maximum energy value is achieved when two of the exchange interactions are zero and is also equal to the ground-state energy per bond of the icosahedron, also existing in three dimensions. The maximum ground-state per bond of the geodesic icosahedra also shows that frustration does not necessarily decrease with the number of vertices, with the pentakis icosidodecahedron having the highest one. Since these molecules have the same spatial symmetry, what makes the difference is their connectivity.

A common characteristic of the lowest-energy configuration in a spin-space dimension \(n\) less than the one of the absolute ground state, is that it typically does not change before the interactions that start to develop in the next higher dimension assume a minimum strength. Unlike odd-membered polygons which are simple frustrated structures, the ground-state energy does not continuously decrease as the interaction in the new dimension is getting stronger. The minimum interaction strength
required to lower the ground-state energy typically in- 
creases with dimensionality, showing that the lowest-
energy configuration is becoming more stable with n. 
Similarly, less strongly frustrated ground states for a 
specific n require a stronger minimum interaction strength 
in the new dimension in order to lower their energy.

The plan of this paper is as follows: in Sec. I1 the 
n-vector model is introduced, and in Secs. III to V111 its 
ground state for different families of molecules is calcu-
lated. Finally Sec. XX presents the conclusions.

II. MODEL

N classical spins $s_i, i = 1,\ldots,N$ of unit magnitude 
are considered, defined in an n-dimensional spin space. 
A spin is mounted on each of the N vertices of the 
different molecules under consideration. The spins in-
teract according to the Hamiltonian $H_n$ of the nearest-
neighbor n-vector model, with two interacting spins con-
ected by an edge of the molecule. The exchange interac-
tions are taken to obey the molecular symmetry, with two 
edges connected by a symmetry operation of the mole-
cular point group corresponding to the same interaction 
strength. An interaction between nearest neighbors i and 
j is taken equal to $J_{ij}$ in $n − 1$ of the spin directions, and 
is scaled with $\alpha_n$ in the remaining one:

$$H_n = \sum_{<ij>} J_{ij}(\sum_{\sigma=1}^{n-1} s_i^\sigma s_j^\sigma + \alpha_n s_i^n s_j^n)$$ (1)

The brackets indicate that interactions are limited to 
nearest neighbors. $J_{ij}$ is nonnegative and $0 < \alpha_n \leq 1$, as 
the dimensionality of spin interactions $d = n−1+\alpha_n$ goes 
from $n−1$ to n in spin space. For bipartite structures 
nearest neighbors point in antiparallel directions in the 
ground state, while for frustrated ones the topology of 
the molecule determines their relative orientation. Here 
the lowest-energy configuration of Hamiltonian (1) as $n$ is 
varied is of interest. The lowest possible dimensionality 
is the Ising limit $n = 1$, and n is allowed to increase until 
the ground state does not change any further. When 
$n = 2$ and $0 < \alpha_2 < 1$ the Hamiltonian is the one of the 
XY model, and at $n = 2$ and $\alpha_2 = 1$ of the XX model. 
For $n = 3$ and $0 < \alpha_3 < 1$ one gets the XXZ model, and 
for $n = 3$ and $\alpha_3 = 1$ the XXX model.

The number of independent exchange interactions 
grows with molecular size and decreasing symmetry. A thorough investigation of interaction space has been made 
for molecules with up to four unique interactions. 
These are parametrized so that the sum of the squares 
of their magnitudes equals one. If their number is two 
they are parametrized by a polar angle $\omega$, and their values 
are equal to the corresponding x and y components. 
If it is three a polar $\theta$ and an azimuthal angle $\phi$ do 
the parametrization, with the three different interactions 
equal to the corresponding x, y, and z components.

The calculations were done numerically [15, 31, 40]. 
Each spin $s_i$ being a classical unit vector in n dimen-
sions is defined by $n−2$ polar angles varying from 0 to 
$\pi$ and an azimuthal angle varying from 0 to $2\pi$. A ran-
don initial configuration of the spins is selected and each 
angle is moved opposite its gradient direction, until the 
lowest energy configuration is reached. Repetition of this 
procedure for different initial configurations ensures that 
the global energy minimum is found.

III. POLYGONS WITH AN ODD NUMBER OF 
VERTICES

The lowest-energy configuration of model (1) on bipar-
tite structures, which lack any frustration, is of the Néel 
type, with nearest-neighbor spins pointing in antiparal-
lel directions. It is already the ground state at the Ising 
limit of Hamiltonian (1).

A simple family of frustrated structures are polygons 
with an odd number of vertices. In the simplest case 
all polygon edges are equivalent and $J_{ij} \equiv J$ in Hamil-
tonian (1), with each spin having two nearest neigh-
bors. In the Ising-limit ground state neighboring spins 
are antiparallel, except from a single parallel pair due 
to the odd total number of spins. Frustration is getting 
weaker with N as the number of antiparallel nearest-
neighbor pairs increases. As interaction in the second 
direction is introduced by allowing $\alpha_n$ to be nonzero, spin-
space anisotropy and consequently frustration are getting 
weaker. This results in a two-dimensional lowest-energy 
configuration in contrast to the unfrustrated case, and a 
continuously decreasing energy per bond and net magneti-
ization with increasing d (Fig. II). At the XX limit the 
angle between nearest-neighbors equals $\frac{\pi}{d−1}$ \cite{10}, while the net magnetization becomes zero. Further increasing 
$d$ in the Hamiltonian does not alter the ground state, and 
the dimensionality of the ground state in spin space 
equals the real-space dimensionality of the polygons.

IV. PLATONIC SOLIDS

A. Icosahedron

The icosahedron belongs to the class of Platonic solids 
\cite{13}, which are convex regular polyhedra with equivalent 
vertices that consist of only one type of polygon. Their 
edges are also equivalent and each $J_{ij} \equiv J$ in Hamil-
tonian (1). The icosahedron consists of 20 triangles, has 
12 vertices, and belongs to the $I_h$ symmetry group, the 
point group with the largest number of symmetry op-
ervations \cite{41}. At the XXX limit the ground state has 
been found to be noncoplanar \cite{9, 14, 15}. The triangles 
share edges resulting in stronger frustration in compar-
ison with an isolated triangle, as the ground-state nearest-
neighbor correlation increases from $−\frac{1}{2}$ in the latter case 
to $−\frac{\sqrt{5}}{5} = −0.44721$. Fig. IIa) shows the evolution of
the ground-state energy per bond as \( d \) varies from 1 to 3. At the Ising limit it equals \(-\frac{3}{4}\), as in the case of an isolated triangle. Unlike the triangle and the other odd-membered polygons though (Fig. 1(a)), the lowest-energy Ising configuration does not immediately change when \( \alpha_2 \) becomes nonzero, remaining the same up to \( \alpha_2 = \frac{\sqrt{5}}{2} \). This can also be seen in the plot of the nearest-neighbor correlations as a function of \( \alpha \) (Fig. 1(b)). The lowest-energy configuration becomes then noncollinear with several unique nearest-neighbor correlation values, until the XX limit where nearest-neighbors are antiparallel or at angles \( \frac{\pi}{2} \) and \( \frac{3\pi}{4} \) with each other, and the ground-state energy per bond equals \(-\frac{5}{4}\). The XX limit ground state is also protected against \( \alpha_3 \) as it does not change up to \( \alpha_3 = 0.47178 \), where it becomes noncoplanar. Finally at the XXX limit the nearest-neighbor correlations become the same for every pair. The ground state does not change as \( d \) is further increased, and \( d = 3 \) is the spin-space dimensionality for which all nearest-neighbor correlations become equal and obey the symmetry of the icosahedron. The total spin equals zero for any value of \( d \).

### V. PLATONIC SOLIDS IN FOUR SPATIAL DIMENSIONS

The icosahedron and the dodecahedron analogues in four spatial dimensions are the 600-cell and the 120-cell respectively [42]. The 600-cell has 120 vertices, 720 edges, and 1200 triangular faces. Fig. 3 shows the evolution of the ground-state energy of Hamiltonian (1), with nearest-neighbor correlations becoming equal exactly at \( d = 4 \). This is determined by the more strongly frustrated triangles as in the case of Platonic solids. The value of \( d = 4 \) is 0.62348, less than \( \alpha_3 \), again unlike the three-dimensional Platonic solids. The ground-state magnetization per spin has discontinuities in its value and also in its derivative as a function of \( d \).

The four-dimensional analogue of the dodecahedron, the 120-cell, has 600 vertices, 1200 edges, and 720 pentagonal faces. Its ground-state energy per bond evolution from \( d = 3 \) to 4 is shown in Fig. 3. The average ground-state nearest-neighbor correlation when \( d = 3 \) equals \(-0.29098 \), and at \( d = 4 \) it is equal to \(-\frac{\sqrt{5}}{4} = -0.30902 \) for every bond, significantly higher than the value of the icosahedron. This shows that frustration is getting quite stronger when going from three to four-dimensional spatial dimensions, and that the dimensionality of the ground-state spin configuration follows the spatial dimensionality of the molecule (Table II), with nearest-neighbor correlations becoming equal exactly at \( d = 4 \). The value of \( \alpha_4 = 0.62348 \), less than \( \alpha_3 \), again unlike the three-dimensional Platonic solids. The ground-state magnetization per spin has discontinuities in its value and also in its derivative as a function of \( d \).

### VI. ARCHIMEDEAN SOLIDS

The Archimedean solids are convex uniform polyhedra that have identical vertices like the Platonic solids, but are formed by more than one polygon. The icosidodecahedron is made of triangles and pentagons, with like polygons sharing vertices and unlike polygons sharing edges, and has \( I_h \) symmetry and identical edges. The ground-state energy per bond of Hamiltonian (1) equals the one of an isolated triangle, with the lowest-energy configuration similarly being coplanar, showing that it is determined by the more strongly frustrated triangles [10].
The truncated tetrahedron and the truncated icosahedron, which have \( T_d \) and \( I_h \) symmetry respectively, have been shown to have ground states three-dimensional in spin space for any ratio of their two symmetrically independent exchange constants \[11, 12\]. They achieve the lowest-possible energy allowed by the connectivity of the frustrated polygon they include, the triangle and the pentagon respectively, which is the sum of energies of isolated frustrated polygons and antiparallel interpolygon bonds.

The ground states of the truncated dodecahedron and the rhombicosidodecahedron, which also have \( I_h \) symmetry and two symmetrically independent exchange constants, are also three-dimensional and achieve the lowest-possible ground-state energy allowed by their odd-membered polygons for any relative value of the exchange constants. In the former case the energy is the sum of energies of isolated triangles and antiparallel intertriangle bonds, while in the latter of isolated triangles and pentagons. Consequently they achieve the maximum possible energy when only the triangle bonds are nonzero, with the corresponding configuration two-dimensional in spin space (Table \[I\]).

The snub dodecahedron has chiral icosahedral symmetry \( I \), lacking a center of inversion. Unlike the aforementioned Archimedean solids it has edge-sharing frustrated units of the same type, which are triangles. It has three symmetrically independent edges, which divide it into pentagons, triangles, and dimers, since not all triangle bonds are symmetrically equivalent. Figure \[G\] shows the reduced ground-state energy per bond \( \frac{E_g}{\sum_{<ij>} J_{ij}} \) when the relative strengths of the corresponding three unique exchange interactions are parametrized by a polar angle \( \theta \) and an azimuthal angle \( \phi \). The \( z \)-component corresponds to the exchange interaction of the pentagon edges, the \( x \) to one of the triangles, and the \( y \) to the one of the dimers. The lowest-energy configuration is again three-dimensional in spin space, with the nearest-neighbor correlations assuming three independent values, corresponding to the unique exchange interactions. The total spin of the ground state is zero. For \( \theta = 0 \) only the pentagon bonds are nonzero and the ground-state energy is the sum of energies of isolated pentagons. For \( \theta = \frac{\pi}{2} \) only the triangle and dimer bonds are nonzero. As \( \phi \) increases from 0 to \( \frac{\pi}{2} \) the energy goes from the sum of energies of isolated triangles to the sum of energies of isolated dimers. For \( 0 < \theta < \frac{\pi}{2} \) the ground-state energy is determined by the competition of the three different exchange constants. The competition between the triangular and pentagonal bonds increases the energy for smaller \( \phi \) when \( \theta \neq \frac{\pi}{2} \), while for higher \( \phi \) the dimer bonds, which link different pentagons, are getting stronger and lower the ground-state energy.

Frustration has a peak when the lowest energy becomes maximum. This occurs when \( \theta = 0.379280\pi \) and \( \phi = \frac{3\pi}{7} \), where the triangle and dimer interactions are both equal to 0.65686 while the pentagonal is equal to 0.37023. The nearest-neighbor correlations are equal to -0.4863181 for every bond (Table \[II\]), higher than the corresponding value for an isolated triangle but lower than the value for the icosahedron. For \( \theta \) not very close to \( \frac{\pi}{2} \) and \( \phi = 0.481\pi \), where the dimer bonds are strong and the triangle bonds weak, minimization of the pentagonal energy is favored and the reduced energy per bond is equal to the one of an isolated pentagon. The equality occurs for slightly different values of \( \phi \) as \( \theta \) is varied.

Archimedean solids have lowest-energy configurations which are at most three-dimensional (Table \[II\]), and frustration is not stronger than the one at the level of an isolated odd-membered polygon. The exception is the snub dodecahedron, which achieves a maximally frustrated ground state with an energy per bond higher than the one of an isolated triangle. Simultaneously, and even though there are three geometrically distinct edges in the molecule, the maximally frustrated ground state achieves the same energy for any bond in the molecule.

**VII. FULLERENE MOLECULES**

Fullerene molecules are allotropes of carbon that consist of 12 pentagons and \( \frac{2\pi}{5} - 10 \) hexagons \[23\]. The polygons share edges and their vertices are three-fold coordinated. Members of the family are the dodecahedron (Sec. \[VI\]) and the truncated icosahedron (Sec. \[VI\]). The pentagons are the source of frustration, and the molecules are further characterized by having neighboring (nonisolated) pentagons or not. A three-dimensional ground state has been found to be a typical feature of fullerene molecules when spins mounted on their vertices interact according to the XXX model and all exchange interactions are equal \[12, 23, 32, 37, 53\]. Frustration even results in a finite ground-state magnetization for many of the molecules. On average frustration decreases with the number of vertices as the number of hexagons increases, but a more precise characterization of frustration, especially for molecules with the same number of vertices, can be done if \( n \) in Hamiltonian \[I\] is allowed to be greater than 3 and the exchange interactions can take arbitrary values. The interaction parameter space has been investigated for fullerene molecules that have up to four symmetrically independent exchange interactions.

\( I_h \)-symmetry molecules bigger than the dodecahedron have pentagons not neighboring one another. For the molecules with 80 (chamfered dodecahedron) and 180 (truncated pentakis dodecahedron) vertices and the \( I \)-symmetry molecule with 140 (hexpropello dodecahedron) the lowest-energy configuration is three-dimensional. The corresponding average energy per bond varies between the one of an isolated pentagon and that of antiparallel spins, meaning that the maximum possible energy occurs when only the pentagons bonds are nonzero and it is two-dimensional in spin space (Table \[I\]). This shows that for icosahedral symmetry frustration is never stronger than the one of an isolated pentagon, as has already been found for the truncated icosahedron \[12\],
and demonstrates a strong correlation between symmetry and magnetic behavior. It has also been found that the icosahedral fullerene molecules have the same classical magnetization response and are also the ones that support magnetization discontinuities for quantum spins 31, 35, 43.

More specifically, for \( N = 80 \) there are two symmetrically unique types of edges, the first between same-pentagon spins and the second between a pentagon and a nonpentagon spin. The reduced ground-state energy per bond is shown in Fig. 7 as a function of \( \omega \), with \( \tan \omega \) the relative strength of the two exchange interactions and for \( d \) ranging from 1 to 3. Also shown is the lower bound for the energy, which is the sum of the ground-state energies of the corresponding number of isolated pentagons and hexagons. Unlike the truncated icosahedron 12, the chamfered dodecahedron does not attain this lower bound, which explains why the intrapentagon and pentagon-nonpentagon nearest-neighbor correlations vary with \( \omega \) and are not constant and equal to \(-\frac{2}{\sqrt{3}}\) and -1 respectively. However the maximum frustration does not exceed the one of an isolated pentagon, with the reduced ground-state energy per bond never higher than \(-\frac{2}{\sqrt{3}}\). This is also true for the icosahedral molecules with \( N = 140 \) and 180, which have four symmetrically unique types of edges, showing that the icosahedral clusters achieve minimal frustration. Frustration does not have to do with the size of the molecule and the fact that the number of unfrustrated hexagons increases with \( N \), but rather with molecular symmetry.

For 24 vertices and \( D_{5h} \) symmetry 28, 44 the ground state of Hamiltonian 11 is in general three-dimensional (Fig. 8) and has zero magnetization. The maximum energy per bond equals -0.7606899 when the exchange interactions in the top and bottom hexagon equal \( J_1 = 0.369747 \), the ones between spins in the top or bottom hexagon and spins in the middle ring equal \( J_2 = 2J_1 \), and the ones between spins in the middle ring equal \( J_3 = 0.562526 \), with the ground state developing then in two spin dimensions. The \( J_1 \) and \( J_2 \) bonds have an energy equal to \(-\frac{1}{\sqrt{3}}\), which equals the ground-state energy per bond, while in the middle ring the energy alternates in value between -1 and \( 1 - \frac{1}{\sqrt{3}} \), so that the average also equals \(-\frac{1}{\sqrt{3}}\).

For 28 vertices and \( T_d \) symmetry 28, 44 the ground state is four-dimensional (Fig. 9). Maximum frustration occurs when same-hexagon exchange interactions are equal to \( J = 0.362873 \), interhexagon interactions to \( 2J \), and the rest of the pentagon-only interactions to 1.61070\( J \). All nearest-neighbor correlations are equal to -0.7843647, and the magnetization per spin equals 0.0599442. The exchange interaction values for the maximum energy per bond are not very different from the ones of the 24-vertices cluster, and similarities can also be detected between the \( \theta \) and \( \phi \) dependence in Figs 8 and 9 with the number of hexagons much less than the one of pentagons for the two clusters.

For 30 vertices and \( D_{3h} \) symmetry 28, 44 the lowest-energy configuration is three-dimensional. Its energy is maximized when the exchange interactions that connect the lower with the upper part of the molecule are zero, dividing it in two parts with respect to its mirror plane. The interaction within the top and bottom pentagons equals 0.73198, between these pentagons and the hexagons 0.59805, and hexagon spins within the upper or the lower side of the molecule interact with bond strengths equal to 0.32641. Again these values are similar to the two fullerene molecules examined before. All nearest-neighbor correlations equal -0.7707600, a value higher that the one of the \( T_d \) molecule with the lower number of 28 vertices. This demonstrates that even though the number of vertices increases frustration does not get weaker, according to the maximum possible ground-state energy criterion. The corresponding magnetization per spin for each half of the molecule equals 0.122080.

For 36 vertices and \( D_{6h} \) symmetry 28, 49 the lowest-energy configuration is two-dimensional. The maximally frustrated ground state occurs again when the exchange interactions that connect the lower with the upper part of the molecule are zero, resulting in two independent parts with respect to the molecule’s mirror plane. The exchange interactions in the top and bottom hexagon have half the strength of the ones that connect them to the inner hexagons, and they have equal strength with the inner-hexagon interactions. Each bond has an energy equal to the energy per bond of an isolated pentagon, demonstrating that this molecule is minimally frustrated. The magnetization per spin for each half of the molecule equals.

For 40 vertices and \( T_d \) symmetry 28 the lowest-energy configuration is five-dimensional. The maximally frustrated ground state is three-dimensional, and occurs when interactions that only belong to hexagons are zero, splitting the molecule in four parts with ten spins each. Interactions that belong to both a pentagon and a hexagon are equal to \( J = 0.52746 \), and pentagon-only interactions have a strength equal to 1.61070\( J \), similarly to the \( N = 28 \) molecule with the same symmetry. Furthermore, all nearest-neighbor correlations are equal to the value of the ones in the maximally frustrated configuration of that molecule, demonstrating the link between symmetry and magnetic properties. The magnetization per spin for each of the four separate parts of the molecule equals 0.0479013. Again and according to the maximum possible ground-state energy criterion, this molecule is more strongly frustrated than the \( D_{6h} \) fullerene with the smaller number of 36 vertices, showing the importance of symmetry for frustration.

When the number of symmetrically independent exchange interactions is greater than four, the lowest-energy configuration of Hamiltonian 11 has been calculated for all \( N \leq 36 \) molecules and for higher symmetry molecules with \( N \leq 100 \) when all exchange interactions are equal, and it is typically three or four-dimensional in
spin space. Along with the \( N = 40 \) molecule with \( T_d \) symmetry, the other exception is the \( N = 100 \) molecule with the same symmetry and also nonisolated pentagons, which has a five-dimensional ground state. Table II lists the ground-state energy of Hamiltonian (1) for both molecules with increasing \( n \). The absolute minimum is achieved for \( n = 5 \), while the minimum \( \alpha_n \) required to lower the ground-state energy when adding a new dimension increases with \( n \), as was mostly the case for the Platonic solids.

Fullerene molecules demonstrate that even though pentagons are less frustrated than triangles, their assemblage with hexagons leads to the formation of ground states up to five spin dimensions (Table I), higher even than the four of the four real-space dimensional Platonic solids. What distinguishes fullerenes from Platonic and Archimedean solids is that their sites are not equivalent, and as \( N \) increases so does the number of symmetrically unique vertices. On the other hand the icosahedral fullerenes are minimally frustrated, pointing to the importance of high symmetry for the magnetic properties, especially since they have also been found to behave non-trivially in a magnetic field both for classical and quantum spins \([31, 35, 43, 45]\). This in agreement with the icosahedral and the four-dimensional Platonic solids forming ground states in a number of spin dimensions equal to their dimensionality in real space, which is also true for the icosahedral Archimedean solids. These results show that frustration in fullerene molecules does not necessarily decrease with \( N \) and the number of hexagons, but symmetry plays an important role as well, as is the case with the minimally frustrated \( D_{6h} \) fullerene with \( N = 36 \). It was also found that in order to achieve maximum energy the exchange interaction values of different molecules are similar in value.

VIII. GEODESIC ICOSAHEDRA

The geodesic icosahedra are polyhedra derived from the icosahedron \([29, 30]\). They have icosahedral symmetry and are duals of fullerene molecules. Twelve of their vertices have five and the rest six nearest-neighbors and they only include triangles. The pentakis dodecahedron is the dual of the truncated icosahedron. It is derived from the dodecahedron by adding a vertex at the center of each one of its 12 pentagons, and has \( N = 32 \). There are two unique types of edges, one corresponding to the dodecahedron edges and the other to the edges linking a pentagon vertex with the vertex at the pentagon’s center. The two corresponding exchange constants are parametrized as \( \cos \omega \) and \( \sin \omega \). The zero-field ground state has been calculated for \( d = 3 \), along with the magnetization response both at the classical and quantum level \([40]\). Figure 11 shows the reduced ground-state energy per bond of Hamiltonian (1) as a function of \( \omega \) for \( d = 1 \) to 4. For \( \omega = 0 \) the ground state of the dodecahedron is three-dimensional \([10]\). Introducing the second exchange interaction further enhances frustration and the reduced energy per bond increases, and eventually the ground state develops a four-dimensional structure in spin space at \( \omega = 0.10414\pi \). The maximum ground-state energy is achieved for \( \omega = \tan^{-1} \frac{\sqrt{5} + 1}{4} \) and equals \( -\frac{\pi}{4} \) for any bond (Table III), as in the ground state of the icosahedron, with the corresponding magnetization per bond equal to 0.088251. This is also the maximum possible ground-state energy for spins located at the vertices of an isolated pentagon with an extra spin located at its center, which is three-dimensional, when the exchange strength with the spin at the center equals \( \sqrt{\frac{\sqrt{5} + 1}{2}} \) times the intrapentagon interaction. At \( \omega = 0.33926\pi \) the lowest-energy configuration becomes collinear.

Figure 11 shows the ground-state energy of Hamiltonian (1) as a function of \( \omega \) away from the collinear regime for values of \( d \) which are in general noninteger. Focusing between \( d = 1 \) and 2, for the ranges of \( \omega \) where the ground-state energy and consequently frustration are maximum, a weak \( \alpha_n \) is sufficient to further lower the energy from its \( d = 1 \) value, while for \( \omega \)-ranges of weaker frustration the \( d = 1 \) lowest-energy configuration is more robust and the corresponding \( \alpha_n \) value bigger. Going from \( d = 2 \) to 3 the corresponding \( \alpha_n \) values are much stronger, and they get even stronger when going from \( d = 3 \) to 4. This shows that the lowest-energy configurations are becoming more robust with increasing \( d \), a conclusion that has been typically drawn for the Platonic solids.

The ground-state nearest-neighbor correlations are shown in Fig. 12. They obey the symmetry of the molecule, with edges connected by symmetry operations corresponding to the same correlation value. They are constant for lower \( \omega \) where the ground-state is three-dimensional, and for higher \( \omega \) where it is collinear. In between they vary with \( \omega \) and become equal at the point of maximum frustration.

The next bigger geodesic icosahedron is the pentakis icosidodecahedron. It is derived from the icosidodecahedron by introducing a vertex at the center of each one of its 12 pentagons, and has \( N = 42 \). The number of geometrically distinct edges is again two, one corresponding to the edges of the icosidodecahedron and the other to the edges linking a pentagon vertex with the vertex at the pentagon’s center. The exchange constants are again parametrized as \( \cos \omega \) and \( \sin \omega \) respectively. Figure 13 shows the reduced ground-state energy per bond of Hamiltonian (1) plotted as a function of \( \omega \) for \( d = 1 \) to 5. At \( \omega = 0 \) the lowest-energy configuration of the icosidodecahedron is two-dimensional. Then for finite \( \omega \) the lowest-energy configuration develops in \( d = 5 \) dimensions, with the maximum energy per bond occurring at \( \omega = 0.256725\pi \), being equal to \(-0.4403875\) for both unique bonds (Table IV). This value is higher than the corresponding one for the pentakis dodecahedron. At \( \omega = 0.34593\pi \) the ground-state becomes four-dimensional, and at \( \omega = 0.41416\pi \) the spins become col-
Figure 14 shows the ground-state nearest-neighbor correlations as a function of $\omega$. Again there is a unique correlation for each unique exchange interaction in Hamiltonian (1), and the two correlations become equal at the point of maximum frustration. At the discontinuity the total spin jumps from zero to a finite value. The stronger frustration of the pentakis icosidodecahedron than the pentakis dodecahedron is also visible at the $\omega = 0$ limit, since it reduces to the icosidodecahedron and not to the dodecahedron, with the latter less strongly frustrated.

The hexapentakis truncated icosahedron has 92 vertices and four symmetrically independent edges. The maximum ground-state energy equals $-\frac{\sqrt{5}}{2}$ (Table III) and occurs when the only nonzero exchange interactions are between the 5-fold coordinated spins and their nearest neighbors, and among these nearest neighbors themselves, with the ratio of the two equal to $\frac{\sqrt{5}+1}{2}$. This is the three-dimensional state that has the maximum ground-state energy for an isolated pentagon with a spin at its center.

The geodesic icosahedra have five and six-fold coordinated vertices. Each one of the former resides at the center of a pentagon, and forms with it a structure that achieves a maximum ground-state energy per bond of $-\frac{\sqrt{5}}{2}$, which equals the ground-state energy of the icosahedron $[3, 5, 5]$. The pentakis dodecahedron achieves this maximum in four spin dimensions, while the pentakis icosidodecahedron has a higher maximum possible energy than $-\frac{\sqrt{5}}{2}$ in five spin dimensions. On the other hand, the hexapentakis truncated icosahedron achieves the maximum when the pentagons with a spin at their center are isolated from the rest of the cluster, which only requires three dimensions in spin space. What distinguishes this molecule from the two smaller ones is that the pentagons with a spin at their center are isolated from one another. These results show that frustration as determined from the maximum ground-state energy criterion does not necessarily weaken with the increase in the number of vertices, which makes the twelve five-fold coordinated vertices much less in number than the six-fold ones. The specific connectivity of each cluster is important, with the pentakis icosidodecahedron having the maximum possible ground-state energy per bond.

**Table I: Different families of molecules with one symmetrically independent exchange interaction for which the ground state of Hamiltonian (1) was calculated. The real-space dimensionality of the molecules is listed, along with the number of vertices $N$, the point-group symmetry, the ground-state dimensionality in spin space $n$, and the ground-state energy per bond $E_\text{bond}$.**

| molecule                        | real-space dimensions | $N$ | point-group symmetry | $n$ | $E_\text{bond}$  |
|---------------------------------|-----------------------|-----|----------------------|-----|------------------|
| Polygons with an Odd Number of Vertices |
| triangle                        | 2                     | 3   | $D_4$                | 2   | $\frac{1}{4}$  |
| pentagon                        | 2                     | 5   | $D_5$                | 2   | $-\frac{\sqrt{5}+1}{4}$  |
| Platonic Solids                 |
| icosahedron                     | 3                     | 12  | $I_h$                | 3   | $-\frac{\sqrt{5}}{2}$  |
| dodecahedron                    | 3                     | 20  | $I_h$                | 3   | $-\frac{\sqrt{5}}{2}$  |
| Four-Dimensional Platonic Solids |
| 600-cell                        | 4                     | 120 | $H_4$, [3,3,5]       | 4   | $-\frac{\sqrt{13}+1}{2}$  |
| 120-cell                        | 4                     | 600 | $H_4$, [3,3,5]       | 4   | $-\frac{\sqrt{13}+1}{2}$  |

**IX. CONCLUSIONS**

Magnetic frustration has been characterized by the dimensionality of the absolute ground state of the $n$-vector model, by allowing $n$ to take arbitrary values with the spins mounted on the vertices of different molecules being more than three-dimensional. Molecules of high symmetry such as Platonic solids and Archimedean solids (Table II) have been found to form ground states in a number of spin-space dimensions equal to their real-space dimension. When there is more than one unique type of vertex, as in the case of fullerene molecules and geodesic icosahedra, the ground state can develop in as many as five spin-space dimensions, in order to optimize nearest-neighbor interactions. Frustration is also characterized by the maximum ground-state energy per bond when there are more than one symmetrically independent exchange interactions and they are allowed to vary. Typically the nearest-neighbor correlations are then equal, unless frustration does not exceed the one at the level of the maximally frustrated polygon of the molecule, as in the case of the $I_h$-symmetry fullerenes. This second way of characterizing frustration also demonstrates the existence of symmetry patterns within the same family of molecules. Furthermore, increasing the number of sites does not necessarily weaken frustration as is expected, but the molecular point-group symmetry also plays an important role, as well as the specific connectivity of each molecule if they are of the same symmetry. It is also found that going from a dimension $n$ to the next higher one by switching on the interactions in the new dimension, it typically takes a finite strength of the latter to lower the ground-state energy.

The study of the $n$-vector model with arbitrary $n$ allows a more precise characterization of the frustration introduced by the molecular connectivity. This is because the energy minimization is not constrained by the requirement that the spins can only be up to three-dimensional, allowing them to minimize the nearest-neighbor interactions more efficiently. The method introduced here can reveal more symmetry patterns if the computational resources are available to study molecules with more than four symmetrically independent exchange interactions. Furthermore, the $n$-vector model can be taken as an effective model, for example in the case of spin-orbit interactions.
TABLE II: Different families of three-dimensional molecules with more than one symmetrically independent exchange interaction for which the ground state of Hamiltonian (1) was calculated for the whole range of their exchange interactions parameter space. The number of vertices of the molecules $N$ is listed, along with the point-group symmetry, the number of symmetrically independent exchange interactions, the ground-state dimensionality in spin space $n$, and the maximum possible reduced ground-state energy per bond \( \frac{E_g}{\sum_{<ij>} J_{ij}} \) along with the corresponding dimensionality in spin space $n_{max}$.

| molecule                        | $N$ | point group | unique inter. | $n$ \( \sum_{<ij>} J_{ij} \) | $n_{max}$ |
|--------------------------------|-----|-------------|---------------|---------------------------|-----------|
| Archimedean Solids             |     |             |               |                           |           |
| truncated dodecahedron          | 60  | $I_h$       | 2             | 3                         | $-\frac{1}{2}$ | 2         |
| rhombicosidodecahedron          | 60  | $I_h$       | 2             | 3                         | $-\frac{1}{2}$ | 2         |
| snub dodecahedron               | 60  | $I$         | 3             | 3                         | -0.4863181 | 3         |
| Fullerene                       |     |             |               |                           |           |
| chamfered dodecahedron          | 80  | $I_h$       | 2             | 3                         | $-\frac{\sqrt{5}+1}{4}$ | 2         |
| hexpropello dodecahedron        | 140 | $I$         | 4             | 3                         | $-\frac{\sqrt{5}+1}{4}$ | 2         |
| truncated pentakis dodecahedron  | 180 | $I_h$       | 4             | 3                         | $-\frac{\sqrt{5}+1}{4}$ | 2         |
| -                               | 24  | $D_{6d}$    | 3             | 3                         | -0.7606899 | 2         |
| -                               | 28  | $T_d$       | 3             | 4                         | -0.7843647 | 4         |
| -                               | 30  | $D_{5h}$    | 4             | 3                         | -0.7707600 | 3         |
| -                               | 36  | $D_{5h}$    | 4             | 2                         | $-\frac{\sqrt{5}+1}{4}$ | 2         |
| -                               | 40  | $T_d$       | 4             | 5                         | -0.7843647 | 3         |
| Geodesic Icosahedra             |     |             |               |                           |           |
| pentakis dodecahedron           | 32  | $I_h$       | 2             | 4                         | $-\frac{\sqrt{5}}{5}$ | 4         |
| pentakis                        | 42  | $I_h$       | 2             | 5                         | -0.4403875 | 5         |
| icosidodecahedron               |     |             |               |                           |           |
| hexapentakis                    | 92  | $I_h$       | 4             | 5                         | $-\frac{\sqrt{5}}{\sqrt{3}}$ | 3         |

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TABLE III: Ground-state energy of Hamiltonian (1) for different \( n \) at the isotropic limit \( \alpha_n = 1 \) and minimum \( \alpha_{n,\text{min}} \) required to lower the ground-state energy in \( n \) dimensions for the \( T_d \)-symmetry nonisolated-pentagon fullerenes with \( N = 40 \) and 100. All the exchange interactions are equal. The corresponding magnetizations are zero.

| \( n \) | \( E_{\text{bond}} \) (\( \alpha_n = 1 \)) | \( \alpha_{n,\text{min}} \) | \( E_{\text{bond}} \) (\( \alpha_n = 1 \)) | \( \alpha_{n,\text{min}} \) |
|------|-----------------|----------------|-----------------|----------------|
| 1    | -0.8265611766   | 0.48794        | 0.47175         |
| 2    | -0.8298861510   | 0.98118        | 0.95423         |
| 3    | -0.8298861626   | 0.9999998      | 0.999995        |
| 4    | -0.8298861626   | 0.9999998      | 0.999995        |
| 5    | -0.8298861626   | 0.9999998      | 0.999995        |

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FIG. 1: (a) Ground-state energy per bond $E_{g}/N$ for Hamiltonian (1) for a triangle (black solid line), a pentagon (red dashed line), a heptagon (green long-dashed line), and a nonagon (blue dot-dashed line) as a function of the dimensionality $d$ of the interactions in spin space. (b) Corresponding magnetization per spin $M_{g}/N$.

FIG. 2: Ground-state energy $E_{g}$ per bond for Hamiltonian (1) as a function of the dimensionality $d$ of the interactions in spin space for (a) the icosahedron, and (b) the dodecahedron.
FIG. 3: Ground-state nearest-neighbor correlations \((\vec{s}_i \cdot \vec{s}_j)_g\) for Hamiltonian (1) as a function of the dimensionality \(d\) of the interactions in spin space for (a) the icosahedron, and (b) the dodecahedron.

FIG. 4: (a), (b), and (c): ground-state energy \(E_g\) per bond and (d), (e), and (f): ground-state magnetization per spin \(M_g/N\) for Hamiltonian (1) as a function of the dimensionality \(d\) of the interactions in spin space for the 600-cell.
FIG. 5: Ground-state energy $E_g$ per bond for Hamiltonian (1) as a function of the dimensionality $d$ of the interactions in spin space for the 120-cell.

FIG. 6: Reduced ground-state energy per bond $\frac{E_g}{\sum_{<ij>} J_{ij}}$ for Hamiltonian (1) in $d = 3$ for different values of $\theta$ as a function of $\phi$ for the snub dodecahedron. $\theta$ and $\phi$ control the relative strength of the three symmetrically independent exchange constants, with the $z$-component the exchange interaction of the pentagon edges, the $x$ the one of the triangles, and the $y$ the one of the dimers.
FIG. 7: Reduced ground-state energy per bond \[ E_g / \sum_{ij} J_{ij} \] for Hamiltonian \( H \) as a function of \( \omega \) for the chamfered dodecahedron \( (N = 80) \), with \( \cos \omega \) the intrapentagon and \( \sin \omega \) the pentagon-nonpentagon interaction. The dimensionality of spin space \( d \) ranges from 1 to 3. "No frustration" shows the sum of reduced ground-state energies of isolated pentagons and hexagons. The intrapentagon and pentagon-nonpentagon nearest-neighbor correlation functions are also plotted.

FIG. 8: Reduced ground-state energy per bond \[ E_g / \sum_{ij} J_{ij} \] for Hamiltonian \( H \) in \( d = 3 \) for different values of \( \theta \) as a function of \( \phi \) for the 24-site fullerene molecule. \( \theta \) and \( \phi \) control the relative strength of the three symmetrically independent exchange constants, with the \( z \)-component the interactions in the top and bottom hexagon, the \( x \) the interaction between spins in the middle ring, and the \( y \) the interaction between spins in the top or bottom hexagon and spins in the middle ring.
FIG. 9: Reduced ground-state energy per bond $E_g/\sum_{ij} J_{ij}$ for Hamiltonian (1) in $d = 4$ for different values of $\theta$ as a function of $\phi$ for the 28-site $T_d$ fullerene molecule isomer. $\theta$ and $\phi$ control the relative strength of the three symmetrically independent exchange constants, with the $z$-component the same-hexagon interactions, the $x$ pentagon-only interactions, and the $y$ interhexagon interactions.

FIG. 10: Reduced ground-state energy per bond $E_g/\sum_{ij} J_{ij}$ for Hamiltonian (1) as a function of $\omega$ for the pentakis dodecahedron. The dodecahedron-edge interaction equals $\cos \omega$ and the pentagon vertex with the pentagon’s center interaction equals $\sin \omega$. The dimensionality of spin space $d$ ranges from 1 to 4. The vertical lines show the $\omega$ values where the dimensionality of the ground state changes from 3 to 4 and then to 1.
FIG. 11: Reduced ground-state energy per bond \( \frac{E_g}{\sum_{ij} J_{ij}} \) for Hamiltonian (1) as a function of \( \omega \) for the pentakis dodecahedron. The dodecahedron-edge interaction equals \( \cos \omega \) and the pentagon vertex with the pentagon’s center interaction equals \( \sin \omega \). The dimensionality of spin space \( d \) ranges from 1 to 4 in steps of \( \frac{1}{10} \), increasing from top to bottom, with integer-\( d \) values corresponding to straight lines, and noninteger to dashed lines. Noninteger values of \( d \) may coincide with integer values, especially with increasing \( d \). The vertical lines show the \( \omega \) values where the dimensionality of the ground state changes from 3 to 4 and then to 1.

FIG. 12: Ground-state nearest-neighbor correlations \( (\vec{s}_i \cdot \vec{s}_j)_{g} \) for Hamiltonian (1) as a function of \( \omega \) for the pentakis dodecahedron. The dodecahedron-edge interaction equals \( \cos \omega \) and the pentagon vertex with the pentagon’s center interaction equals \( \sin \omega \). The vertical lines show the \( \omega \) values where the dimensionality of the ground state changes from 3 to 4 and then to 1.
FIG. 13: Reduced ground-state energy per bond $\frac{E_g}{\sum J_{ij}}$ for Hamiltonian (1) as a function of $\omega$ for the pentakis icosidodecahedron. The icosidodecahedron-edge interaction equals $\cos \omega$ and the pentagon vertex with the pentagon’s center interaction equals $\sin \omega$. The dimensionality of spin space $d$ ranges from 1 to 5. The vertical lines show the $\omega$ values where the dimensionality of the ground state changes from 5 to 4 and then to 1.

FIG. 14: Ground-state nearest-neighbor correlations $(\vec{s}_i \cdot \vec{s}_j)_g$ for Hamiltonian (1) as a function of $\omega$ for the pentakis icosidodecahedron. The icosidodecahedron-edge interaction equals $\cos \omega$ and the pentagon vertex with the pentagon’s center interaction equals $\sin \omega$. The vertical lines show the $\omega$ values where the dimensionality of the ground state changes from 5 to 4 and then to 1.