Monopole ordered phases in dipolar and nearest-neighbours Ising pyrochlore: from spin ice to the “all-in–all-out” antiferromagnet

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We study Ising pyrochlores by means of Monte Carlo simulations. We cover a set of exchange constants ranging from the frustrated ferromagnetic case (spin-ice) to the fully-ordered “all-in–all-out” antiferromagnet in the dipolar model, reinterpreting the results –as in an ionic system– in terms of a temperature vs. magnetic charge density phase diagram. In spite of its spin nature and the presence of both double and single non-conserved magnetic charges, the dipolar model gives place to a phase diagram which is quite comparable with those previously obtained for on-lattice systems of electric charges, and on spin ice models with conserved number of single magnetic charges. The contrast between these systems, to which we add results from the nearest-neighbours model, put forward other features of our phase diagram –notably, a monopole fluid with charge order at high monopole densities that persists up to arbitrarily high temperatures– that can only be explained taking into account construction constraints forced by the underlying spin degrees of freedom.

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I. INTRODUCTION

Describing a complex material in terms of low-lying particle-like collective excitations –quasi-particles– is one of the key approaches in condensed matter physics. Phonons, magnons, electrons and holes in semiconductors, are some of the better known examples of these excitations. But quasi-particles form a very rich set, including topological excitations and fractionalisation. In a first approximation, these excitations are considered as non-interacting, with all the complexity of the system hidden in the quasi-particles themselves. The next layer of description, where interactions between quasi-particles are included, leads to a great variety of behaviours: from anharmonic effects in crystals to the stabilization of higher hierarchies of order such as magnon binding2 or magnon-mediated heavy fermion superconductivity.3 Recently, a new kind of fractional point-like topological excitation has been proposed theoretically4 and evidence of its existence found experimentally5–7 in the spin ice compounds. These new quasi-particles are sources of magnetic field and interact via a Coulomb-like potential, whence their name magnetic monopoles. This approach allows for a very effective description of the thermodynamics, the dynamics and the out of equilibrium behaviour of spin-ice systems.4,8–15

The magnetic properties of spin ice materials can be described by classical magnetic moments in a pyrochlore lattice, occupying the vertices of corner-sharing tetrahedra. They behave at low temperatures as Ising-like spins16 $\mu_i = \mu_S e_\parallel$ with $S_i = \pm 1$, pointing along the ⟨111⟩ directions $\hat{e}_i$ (Fig. 1). The magnetic interactions of exchange and dipolar origin –of strengths $J$ and $D$, respectively– are well accounted for by the dipolar spin ice model (DSIM) Hamiltonian:

$$\mathcal{H} = \frac{D}{T} \sum_{\langle ij \rangle} S_i S_j +$$

$$+ a^3 \sum_{\langle ij \rangle} \left( \frac{\hat{e}_i \cdot \hat{e}_j}{|\mathbf{r}_{ij}|^3} - \frac{3(\hat{e}_i \cdot \mathbf{r}_{ij})(\hat{e}_j \cdot \mathbf{r}_{ij})}{|\mathbf{r}_{ij}|^5} \right) S_i S_j \tag{1}$$

where $T$ is the temperature, $a$ is the lattice spacing, $\langle ij \rangle$ means that the sum is carried over nearest neighbours, $|\mathbf{r}_{ij}|$ is the distance between spins $i$ and $j$ and $D = \mu_0 \mu^2 / (4\pi a^3)$.

The nearest neighbours spin ice model (NNSIM) Hamiltonian is derived from the DSIM by keeping only the nearest neighbour contributions of the dipolar interaction:

$$\frac{\mathcal{H}_{nn}}{T} = \frac{J_{nn} + D_{nn}}{T} \sum_{\langle ij \rangle} S_i S_j \tag{2}$$

with $J_{nn} = J/3$ and $D_{nn} = 5D/3$, i.e. an effective exchange interaction of strength $J_{\text{eff}} = J_{nn} + D_{nn}$. For the spin ice materials, $J_{\text{eff}}$ in (2) is positive ($J_{nn}/D_{nn} > -1$) and imposes the spin-ice rule (named after Bernal and Fowler’s ice rules17): two spins should point in and two out of a tetrahedron. This rule can be translated into field theory language as a divergence free condition, which gives rise to a “Coulomb phase”.18 Following Ref.
beauty and simplicity of the monopolar scenario introduced by Castelnovo and collaborators. A recent contribution addresses this same issue in a wider scope. Excluding double charges in the dumbbell model (which — unlike the previous approach — takes magnetic charges and not spins as their interacting simple entities), they show that a Coulomb phase can still be defined beneath a crystal of magnetic single charges. One drawback of these approaches is that the explicit omission of double defects is somewhat contrived: in real materials, the limit \( J_{\text{eff}}/T \to 0 \) implies the proliferation of both single and double excitations. Furthermore, within the CDSIM model, the ordered ground state expected for negative \( J_{\text{eff}}/T \) is not allowed.

In this paper we return to the usual DSIM in order to address these shortcomings in the previous analysis and to extend it in order to include the antiferromagnetic case. Building up on previous results by den Hertog et al., we show that when examining using the framework of monopolar excitations both the DSIM and the CDSIM lead to the same physics, and in particular to very similar phase diagrams. Since within the NNSIM one would naively expect no monopole-monopole interaction, one would think charge degrees of freedom to play no role. In spite of this, we will see that the phase diagram obtained for the NNSIM model can be reinterpreted in terms of effective nearest-neighbours interactions between double charges. We will also show that these effective interactions (neither dipolar nor exchange in origin, but arising from correlations imposed by the internal degrees of freedom of the charges) also affect the phase diagram in the presence of dipolar interactions.

**Simulation details**

We performed Monte Carlo simulations with single spin-flip Metropolis algorithm, using Ewald summations to take into account the long-range interactions. We used a conventional cubic cell for the pyrochlore lattice, which contains 16 spins, and simulated systems with \( L \times L \times L \) cells. Thermodynamic data were collected by starting at high temperatures and cooling very slowly, for different values of \( J_{\text{nn}}/D_{\text{nn}} \) (as this ratio will be negative throughout this work, we will usually refer to its absolute value). Typically, we needed \( 10^4 \) Monte Carlo steps for equilibration and \( 2 \cdot 10^4 \) for averaging at each temperature for \( L = 4 \), but we used up to \( 10^5 \) steps for bigger lattices. First order transition points deserve to be mentioned separately; there, full equilibration was only achieved after longer times (up to \( 5 \cdot 10^5 \) Monte Carlo steps) for sizes below \( L = 5 \).
II. CHARGE DEGREES OF FREEDOM IN THE DIPOLAR MODEL

Fig. 2 shows the monopole density $\rho$ as a function of temperature for $L = 4$. We define $\rho$ as the number of single charges per unit tetrahedron, thus counting double monopoles as the superposition of two single charges of the same sign in a tetrahedron. In a similar way to Fig. 2 of Ref. 23, where only single charges were considered, we can distinguish two families of curves according to their limit as $T \to 0$. The first type, which tend to $\rho = 0$ (with $|J_{nn}/D_{nn}| \leq 0.911$), were characterised as spin ices by determining their residual entropy, via numerical integration of the specific heat divided by temperature. The second type are curves that tend to $\rho = 2$ when $T \to 0$, meaning that they reach a state in which all the tetrahedra are occupied by double monopoles (i.e. the non-frustrated antiferromagnetic phase). For certain values of $|J_{nn}/D_{nn}|$ in the latter group of curves, the monopole density per tetrahedron suffers a sudden change at low temperatures from $\rho \approx 0$ to $\rho \approx 2$. It is straightforward to associate the appearance of this zinc-blende structure to the presence of Coulomb-like interactions between monopoles. However, later we will show that this phase (which is no other than the “all-in-all-out” phase mentioned on the title and in the introduction) can also be stabilized in the NNSIM, where these interactions are absent.

The curves in Fig. 3 (upper panel) represent the molar specific heat $C$ as a function of temperature for different values of $|J_{nn}/D_{nn}|$. It is possible to distinguish two very different regimes: some of the curves (those with $|J_{nn}/D_{nn}| \leq 0.911$) reach $\rho = 0$ when $T \to 0$, while the rest approach $\rho = 2$. Among the latter it is easy to notice that the jump between low and high density becomes smoother as $|J_{nn}/D_{nn}|$ increases. As $|J_{nn}/D_{nn}|$ increases, correlated with the jumps we noticed in $\rho$. With the development of a zinc-blende structure in mind, we also present our results of the double monopole staggered density $\rho^S_d$, defined as the average of the modulus of the total magnetic charge due to double monopoles in up tetrahedra per sublattice site per unit charge. A nearly zero value of this quantity implies that there is no symmetry breaking between the up and down tetrahedra sublattices, while $\rho^S_d \approx 1$ is the result of a staggered ordering of the magnetic charges: positive and negative double monopoles alternating in the sublattices, occupying all the tetrahedra. In Fig. 3 (lower panel) we observe a jump between these two states, which is sudden and step-like for low values of $|J_{nn}/D_{nn}|$ and slowly changes into continuous as that ratio increases. It is interesting to note that these step-like jumps are correlated to the ones in $\rho$: the system becomes dense and charge-ordered suddenly and simultaneously, just as it happens in the crystallisation transition of ionic fluids. On the other hand, the continuous developing of both the staggered density and the number density at higher $|J_{nn}/D_{nn}|$ indicates the transition to a phase in which the local density is homogeneous (fluid-like) but charge-ordered.

To confirm the existence of these transitions and determine their order we performed finite size analysis over values of $J_{nn}/D_{nn}$ representative of the two behaviours. In Fig. 4 we present our results of the molar specific heat.
for \( J_{nn}/D_{nn} = -0.919 \). The previous suggestion of a first order transition taking place for these values of \( J_{nn}/D_{nn} \) is backed up by the fact that the value of the specific heat and the double monopole susceptibility \( \chi_d^{\rho} \) (defined as the fluctuations of the corresponding staggered density over the temperature) at their maximum are proportional to the volume of the system (Fig. 4, inset). We also studied \( J_{nn}/D_{nn} = -1.064 \), in which the double monopole susceptibility (Fig. 5), as well as the specific heat and the double monopole staggered density (not shown), evolve with the size of the system as in a second order phase transition. The critical exponents are consistent with the three-dimensional Ising universality class (Fig. 5, inset). Since this line of second order transitions becomes first order, there must be a tri-critical point of the Blume-Emery-Griffiths universality class.

A nice way to summarise all the previous results while emphasising the importance of charge degrees of freedom is to construct a phase diagram in terms of the monopole density and the temperature (Fig. 6). The colour map corresponds to the interpolated value of the double monopole staggered density \( \rho_S^d \) as a function of temperature and monopole number density (for \( L = 4 \)). The graph was obtained by combining a set of more than thirty \( \rho_S^d \) vs. \( T \) curves (of which those shown on Fig. 3, bottom, constitute a subset) with \( \rho \) vs. \( T \) curves (like those on Fig. 2). The filled circles represent the location of the maximum on the specific heat for various values of \( J_{nn}/D_{nn} \) (see Fig. 3, top). Note that the vertical asymptote observed for this curve at high temperature corresponds to the limit in which antiferromagnetic exchange dominates over the dipolar interactions. The white dome (drawn by estimating the region where discontinuities would be observed for the infinite system) represents a forbidden region in parameter space in which, in the thermodynamic limit \( L \to \infty \), the system cannot stay homogeneous. A system with \( \rho = 0.4 \) at \( T/D_{nn} \approx 0.22 \) is exactly at the edge of the dome; if cooled down, it would separate into two phases occupying different parts of the material. On one hand, we would get a low-density monopole fluid, in which most of the tetrahedra are empty. On the other hand, a phase with a high double monopole density would crystallise into the zinc-blende ionic structure. In a tri-critical point around \( \rho = 1.17 \) and \( T/D_{nn} \approx 0.34 \) the first order transition becomes second order, separating the monopole fluid and a staggered charge fluid in which local density is homogeneous but negative and positive charges display a tendency to occupy different sublattices, thus breaking the symmetry without phase separation. It is interesting to note that both this staggered charge fluid and the crystal phase are more natural, double charge analogs of the single monopole ordered fluid and crystal found in the slightly artificial CDSIM (see Fig. 4 of Ref. 22). The apparent differences between this phase diagram and that presented on Fig. 1 of Ref. 24 (of which we give our own version on Fig. 9) should not mask the fact that they both describe the same physics.

On the other hand, despite the great similarity between the phase diagram of Fig. 6, and that for the CDSIM, several differences arise. Since we used a single spin flip algorithm and did not fix the number of monopolar defects, the Melko-Gingras-den Hertog first order transition\(^{25}\) at \( T/D_{nn} = 0.077 \) to an “ordered vacuum” of magnetic charges\(^{22}\) is not visible. Also, Fig. 6 exhibits a re-entrant behaviour of the monopole fluid at temperatures just above \( T_l/D_{nn} \). Finally, one would expect that a disordered phase should always be found...
at high temperatures ($T/D_{nn} \gg 1$) for any charge density, when entropic forces overcome monopole attraction. While this is true both for a system of real Coulomb charges in a lattice and for the CDSIM, we do not see the line of second order transitions joining $\rho = 2$ in the present case. Instead, this line reaches an asymptotically vertical behaviour, parallel to the temperature axis. This fact, which seems to point to an infinitely high interaction energy between charges, could hardly be explained by the monopolar Hamiltonian on its own. These last two cases deserve special attention, and will be analysed in depth in the next sections.

III. COMPARISON BETWEEN THE DIPOLAR AND THE NEAREST-NEIGHBOURS MODEL

In order to better understand the limits of the magnetic charges picture and the energetics of the models we have introduced, we also studied the NNSIM (Eq. (2)) for different ratios $J_{nn}/D_{nn}$. Similar to the previous case, we found a peak in the specific heat and the double monopole susceptibility, and a steep rise in the double monopole staggered density as a function of temperature (not shown), suggesting a phase transition. In this case the behaviour of these quantities is typical of a second order transition for each value of $J_{nn}/D_{nn}$ studied, with no delta-like peaks or discontinuities.

In Fig. 7 we show the peak in the the molar specific heat for $J_{nn}/D_{nn} = -1.064$ and different sizes of the system. The inset shows that its value at the critical temperature—as well as the order parameter $\rho_d^S$ and its fluctuations $\chi_d^S$—evolves with size as a power law. As expected, we determined that the transition again belongs to the three-dimensional Ising universality class. Furthermore, since only one energy scale is present in this model (see Eq. (2)), the $T$ vs. $\rho$ phase diagram shown in Fig. 8 does not depend on $T/D_{nn}$.

FIG. 6. Temperature vs. monopole density phase diagram with dipolar (long-range) interactions. At low temperatures the system undergoes a first order phase transition (white dome), where a low density gas and a crystal coexist. The transition becomes second order (dot-dashed line) at a tricritical point near $\rho_t \approx 1.17$, $T_t/D_{nn} \approx 0.34$. The diagram is overlaid on top of an interpolated contour plot of $\rho_d S$ for $L = 4$. The vertical dotted line at $\rho_t \approx 1.4$ indicates the critical density of the geometric transition in the NNSIM (see Sec. IV A).

FIG. 7. In the case of the nearest neighbour model, the specific heat displays finite size effects characteristic of a second order phase transition for $J_{nn}/D_{nn} = -1.064$. Inset: again, the values of $\rho_d^S(T_c(L))$, $C(T_c(L))/R$ and $\chi_d^S(T_c(L))$ evolve as power laws with the size of the system and present good correspondence with the behaviour expected for the three dimensional Ising model (grey lines).

FIG. 8. Temperature vs. monopole density phase diagram with nearest-neighbours interactions. The second order phase transition (dot-dashed line) between the gas and the staggered charge fluid occurs at $\rho_c \approx 1.4$ at all temperatures, hence we call it a geometric phase transition. The diagram is overlaid on top of an interpolated contour plot of $\rho_d^S$ for $L = 4$.

The dependence of the critical temperature on the rel-
The strength of the monopole picture, that allows not only a qualitative but also a quantitative understanding of the phase diagram, seems at first sight to be weakened by a plain fact: no Coulomb attraction between oppositely charged monopoles is expected in the NNSIM (note that we continue calling the excitations “monopoles” despite the fact that they have lost their main quality as charges). While it is simple to understand why the spins would order in the alternating “all-in–all-out” configuration when only a negative nearest-neighbours interaction $J_{\text{eff}}$ is considered (Fig. 9), it is much harder to rationalise, limiting ourselves to the monopole picture, why non-attracting monopoles would experience any sort of staggered ordering.

This simple puzzle can be solved by taking into account the correlations between monopoles, which transcend any Coulomb-like interactions. It is not energetic disfavour, but construction constraints that prevent more than one double charge in a single diamond site. In the same way, the underlying spin configuration (“all-in” or “all-out”) forbids two double monopoles with the same double charge to be placed in adjacent tetrahedra. A similar reason makes more probable to find single monopoles with the opposite charge around any double monopole. Provided there are enough double charges in a diamond lattice, this correlation—quite equivalent to an infinite repulsion between double charges of the same sign at nearest neighbour sites—will induce staggered ordering on the system, irrespective of the system temperature.

In order to test this point quantitatively, we have simulated a system of nominal “plus” and “minus” hard spheres in the diamond lattice within the Grand Canonical Ensemble. In analogy to the NNSIM, we included no interactions between spheres, but a constraint was imposed forbidding two like spheres to be placed in neighbouring sites. Figure 10 shows the staggered charge density $\rho_S$ for the spheres as a function of total average sphere density $\rho$ for different system sizes—a cubic unit cell of side $L$ is again implied. The sudden increase in $\rho_S$ reflects the charge-like order being stabilised in the system as a consequence of sphere correlations. A previously known example of this kind of symmetry breaking in hard-sphere systems can be found in binary mixtures with a radius ratio of $R_A/R_B \approx 0.4$ and 0.76, where $R_A$ ($R_B$) is the radius of the small (large) spheres. Driven by differential excluded volume effects, these systems present NaCl and CsCl structures, respectively, in which every $A$ particle is surrounded only by $B$ particles, and vice versa, resembling the staggered ordering of our non-interacting double “monopoles”.

Once more, finite size scaling of the order parameter and its fluctuations (Fig. 10, inset) allowed us to identify the three dimensional Ising universality class. The

![FIG. 9. $T/D_{nn}$ vs. $J_{nn}/D_{nn}$ phase diagram for both dipolar and nearest-neighbours spin ice models. Dotted lines represent crossovers, while solid and dot-dashed lines stand for first and second order phase transitions, respectively. The effect of the dipolar interactions is clearly seen: the system orders at the alternating charge configuration even for values of $J_{\text{eff}} > 0$ because the Coulomb attraction between monopoles of opposite charge makes it energetically favourable to create and order these particles. Melko-Gingras-den Hertog first order transition $^{25}$ is not visible because we used a single spin-flip algorithm that mimics the real material and thus freezes at low temperatures.](image-url)
extrapolated critical density \( \rho_c(L \to \infty) = 0.964 \pm 0.004 \) is near the critical value found for \( \rho_c \) in Fig. 8 for the NNSIM. The differences between these two critical monopole concentrations can be explained realizing that our simple sphere model does not consider single monopoles, which add extra charge with smaller correlations. Considering then that even the NNSIM involves an effective contact interaction between monopoles, we can now understand the independence of \( \rho_c \) with temperature observed in Fig. 8 as a result of a construction constraint.

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**FIG. 10.** Staggered charge density (main figure) and its fluctuations (inset) vs. particle density for a system of “plus” and “minus” non-interacting hard spheres within the Grand Canonical Ensemble. Finite size scaling of these quantities is consistent with the three dimensional Ising universality class\(^{27} \) (not shown).

This reasoning gains in depth when we contemplate its implications to the dipolar model studied before. While the low temperature region of the phase diagram in Fig. 6 is, as discussed, obviously dominated by the Coulomb attraction between monopoles, the ordered phase seen for \( T/D_{nn} \gg 1 \) at high monopole density can now be trivially explained in terms of the aforementioned charge correlations. Indeed, the vertical asymptote we noticed in Fig. 6 shows the equivalence between the DSIM and NNSIM in the limit of high \( T/D_{nn} \).

As mentioned, correlations beyond charge interactions are much smaller for single charges. Indeed, using our conserved-monopole algorithm\(^{22} \) for nearest neighbours spin interactions we have proved that a system of single monopoles remains charge-disordered, even in the limit of \( \rho = 1 \). This explains the observation of a high temperature limit for the stability of the fluid of single monopoles observed in Ref. 22.

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**B. Discussion: re-entrance near the tri-critical point**

Figure 6 suggest the re-entrance of the charge-disordered phase near the dome. Indeed, we have observed that this behaviour persists for bigger lattices, of which we have explored up to \( L = 5 \). Though not in a conclusive way, this suggests that this behaviour is not the mere consequence of the tri-critical fluctuations. As in the previous discussion, we propose an explanation for the re-entrant behaviour –which is not observed in the Blume-Emery-Griffiths model\(^{26} \) nor in lattice models of real charges\(^{29,33} \) nor in the CDSIM\(^{22} \) – which transcends the mere charge-like degrees of freedom.

A clue of the entropic origin of the re-entrance can be obtained by noticing that the charge-ordered phase extends its region of stability on increasing temperature, reaching lower densities. We have checked that along the transition line the reduction in \( \rho \) is related with an increase in the number of empty (“2-in/2-out”) tetrahedra, while keeping an approximately constant concentration of single-charge monopoles. This mechanism, which necessarily decreases the number of double-defects, is very efficient in increasing the entropy of the system. This entropic contribution –which can be related to internal degrees of freedom of the charges– can outweigh the expected decrease in energy, reducing the free energy of the staggered phase. The balance is no longer possible for low enough densities, and the previously observed charge-like behaviour\(^{22,29,33} \) is recovered, with \( \rho_c \) increasing with temperature.

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**C. Charge order with spin disorder: best chance within an Ising pyrochlore material**

We have already mentioned the impossibility of stabilising in a real Ising pyrochlore material a perfect single-monopole crystal at zero magnetic field. As stated in Ref. 23, such a crystal would imply the existence of charge order coexisting with a Coulomb phase. We believe that the staggered charge-ordered phase near the re-entrant region is the closest we can get in this system to this situation. At \( T/D_{nn} \approx 0.57 \) and \( \rho \approx 1.1 \) (see Fig. 1 for a snapshot of a part of the system at these values) almost 50% of the diamond lattice sites are occupied by single-defects, while only 30% are double monopoles. This leftmost transition point, in which the staggered charge-order is impending, is quantitatively compatible with the phase diagram obtained for the CDSIM (since only single charges are allowed in this model, a double monopole in the DSIM must be equated to a single monopole in the CDSIM, then the total density \( \rho = 1.1 \) in the first model corresponds to \( \rho \approx 0.5 + 0.3 = 0.8 \) in the latter). Indeed, Fig. 4 in Ref. 22 shows that for 80% occupation staggered order is established near \( T/D = 5/3 \) \( T/D_{nn} \approx 0.95 \), plainly consistent with the value \( T/D_{nn} \approx 0.57 \) we find for the unrestricted DSIM. This coincidence implies that
double monopoles for this level of dilution contribute to the staggered-charge order similarly to single monopoles. An important consequence of this fact is that a neutron scattering measurement of an Ising pyrochlore material with $J_{nn}/D_{nn} \approx -0.947$ at $T \approx 0.57 D_{nn}$ (so as to have $\rho \approx 1.1$ inside the ordered phase, but with a significant fraction of single-charged monopoles) should give a structure factor quite similar to that simulated in Fig. 4 of Ref. 23. This pattern shows signs of spin fragmentation, combining Bragg peaks from an “all-in–all-out” structure with the pinch points which characterise the underlying Coulomb phase.

V. CONCLUSIONS

The path we have taken in this work has been two-fold. Firstly, we have shown how the monopole picture arises naturally as a tool to reinterpret and understand the physics of dipolar Ising pyrochlore systems, encompassing both spin ice and antiferromagnetic materials. The magnetic phase diagram for classical Ising pyrochlores calculated by den Hertog and collaborators, 24 separating the “all-in–all-out” antiferromagnet from the spin ice systems in terms of the effective exchange constant values, has been recast into temperature vs. density, in analogy to the phase diagrams in molecular systems. Notably, in spite of its underlying spin nature and the presence of four types of different non-conserved magnetic charges (plus and minus, single and double), the dipolar model gives rise to a phase diagram which is quite comparable with those previously obtained for on-lattice systems of electric charges, 29 and on spin ice models with conserved number of single magnetic charges. 22 Thinking just on monopoles and their interactions made it simple to justify quantitatively the extent to which the antiferromagnetic phase gets into the region of ferromagnetic first neighbours effective exchange interaction, by evaluating the Madelung energy of a double monopole crystal. This plainly exemplifies how the power of the picture does not limit itself to the qualitative understanding of the phases present. Within the nearest-neighbours model, the stabilisation of the antiferromagnetic phase was interpreted within the monopole picture in terms of the proliferation of monopoles with no Coulomb interactions but which implicitly force a nearest-neighbour exclusion condition between like types. This exclusion condition mimics an attraction/repulsion between like/different charges, maintaining the idea of monopole as a useful concept even within the NNSIM.

We also encountered certain aspects on the phase diagrams which cannot be explained in terms of simple charges. Like the discovery of internal degrees of freedom in particles previously thought as indivisible building blocks, these findings are far from making the monopole picture less interesting. Among these peculiarities we found a re-entrance of the disordered fluid, and a staggered charge-ordered fluid phase which—for high enough densities—can be stable at temperatures arbitrarily much higher than the energy scale characterising charge interactions. This second fact could be understood in terms of construction constraints—the exclusion condition we referred to in the previous paragraph— inherent to the spin nature of the excitations. The local constraint (as opposed to Coulomb-like monopole attraction) is the predominant correlation mechanism between monopoles at high temperatures and high monopole density, and the only interaction for the NNSIM. We explained the re-entrance noticing that lowering the density of double charges induces a reduction in the energy, but a boost in this energy’s degeneracy. A final remark is the identification near the re-entrance of a good candidate for a state where charge order can coexist with a Coulomb phase 23 (an Ising pyrochlore material with $J_{nn}/D_{nn} \approx -0.947$, at $T/D_{nn} \approx 0.57$). The scattering pattern of this state should show signs of spin fragmentation, combining Bragg peaks from an “all-in–all-out” structure with the pinch points which characterise the underlying Coulomb phase. 23

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It should be stressed that the right part of this diagram corresponds to a single phase. The use of different names for two regions of it—which follows Ref. 22—does not imply that a different symmetry is broken, but intends to emphasise their distinct dynamical properties and the fact that, while the double monopole crystal may coexist with a low density charge-disordered fluid, the staggered charge fluid (composed of both single and double monopoles) is homogeneous and occupies the whole space.