Atomic Fermi gas in the trimerised kagomé lattice at the filling 2/3

B. Damski\(^1,2\), H.-U. Everts\(^3\), A. Honecker\(^3\), H. Feermann\(^1\), L. Santos\(^4\) and M. Lewenstein\(^1,5,*\)

(1) Institut für Theoretische Physik, Universität Hannover, Appelstr. 2, D-30167 Hannover
(2) Institute of Physics, Jagiellonian University, Poland
(3) Institute for Theoretical Physics, TU Braunschweig, Mendelssohnstr. 3, D-38106 Braunschweig
(4) Institut für Theoretische Physik III, Universität Stuttgart, Pfaffenwaldring 57 V, D-70550 Stuttgart
(5) ICFO-Institut de Ciències Fotòniques, Jordi Girona 29, Edifici Nexus II, E-08034 Barcelona, Spain

We study low temperature properties of a spinless interacting Fermi gas in the trimerised kagomé lattice. The case of two fermions per trimer is described by a quantum spin 1/2 model on the triangular lattice with couplings depending on the bond directions. Using exact diagonalisations we show that the system exhibits non-standard properties of a quantum spin-liquid crystal, combining a planar antiferromagnetic order with an exceptionally large number of low energy excitations.

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One of the most fascinating recent trends in the physics of ultracold gases concerns atomic gases in optical lattices, where strongly correlated systems may be realized. Such systems offer an "atomic Hubbard toolbox" \(^1\) to simulate various sorts of Hubbard models, and to study phenomena known in condensed matter physics in an unprecedentedly controlled manner. To name just few examples, atomic lattice gases may serve to study various spin models \(^2\), to simulate high \(T_c\) superconductivity \(^3\), to investigate a variety of quantum disordered systems \(^4\), or to process quantum information \(^5\). Seminal experiments of Ref. \(^6\) have stimulated a great interest in experimental studies of atomic lattice gases (cf. Ref. \(^7\)).

Particularly fascinating in this context is the possibility of studying quantum frustrated antiferromagnets, which lie at the heart of modern quantum magnetism \(^8\). Recently we have proposed how to create ideal and trimerised kagomé optical lattices, and have studied physics of various quantum gases in such lattices \(^9\). A Fermi-Fermi mixture with half filling for both species in the limit of strong interspecies coupling behaves in the original kagomé lattice, the problem of two fermions per trimer is described by a quantum spin 1 \(\frac{1}{2}\) model on the triangular lattice with couplings depending on the bond directions. Using exact diagonalisations we show that the system exhibits non-standard properties of a quantum spin-liquid crystal, combining planar antiferromagnetic order with an exceptionally large number of low energy excitations, and a small (if any) gap. For the effectively antiferromagnetic coupling the quantum results agree very well with the classical results indicating antiferromagnetic planar order and a gapped spectrum.

The experimental realization of the considered system requires a creation of trimerised kagomé lattice, using superlattice techniques as shown in Ref. \(^10\). The spinless interacting Fermi gas can then be formed, for instance, in a Bose-Fermi mixture, in the strong coupling limit, when bosons form a Mott insulator (MI), and fermions together with 0, 1, ..., bosons (bosonic holes) form fermionic composites \(^11\). Alternatively, one could use a gas of polarised ultracold dipolar fermions, that interact via repulsive dipolar potential.

The spinless interacting Fermi gas in the trimerised kagomé lattice is described by the extended Fermi-Hubbard Hamiltonian

\[
H_{FH} = -\sum_{\langle ab \rangle} (t_{ab} f^\dagger_a f_b + \text{h.c.}) + \sum_{\langle ab \rangle} U_{ab} n_a n_b, \quad \text{where} \, \langle a, i \rangle \, \text{referring to intra-trimmer indices and} \, i \, \text{numbering the trimers. The} \, t_{ab} \, \text{and} \, U_{ab} \, \text{take the values} \, t \, \text{and} \, U \, \text{for intra-, and} \, t' \, \text{and} \, U' \, \text{for inter-trimer hopping,} \, n_a = f^\dagger_a f_a, \, \text{and} \, f_a \, \text{is the fermionic annihilation operator. The sites in each trimer are enumerated as in Fig. \(^11\). We denote the 3 different intra-trimer modes by} \, f^{(i)}(i) \equiv (f_{1,i+} + f_{2,i+} + f_{3,i+})/\sqrt{3} \, \text{(zero momentum mode), and} \, f^{(i)}_{\pm} = (f_{1,i+} \pm z_{\pm} f_{2,i+} + z_{\pm} f_{3,i+})/\sqrt{3} \, \text{(left and right chirality modes)}, \, \text{where} \, z_{\pm} = \exp(\pm 2\pi i/3). \]

In the limit of weak coupling between the trimers of the original kagomé lattice, the problem of two fermions per trimer (filling 2/3) becomes equivalent to a quantum magnet on a triangular lattice with couplings that depend on the bond directions as described by the Hamiltonian

\[
H_{\text{trimer}} = \frac{J}{2} \sum_{i=1}^{N} \sum_{j=1}^{6} s_i (\phi_{i\to j}) s_j (\tilde{\phi}_{j\to i}), \tag{1}
\]
where \( N \) denotes number of trimers, \( J = 4U'/9 \), and the nearest neighbours are enumerated as in Fig. 1. In Eq. 1 we have \( s_i(\phi) = \cos(\phi) s_x^{(i)} + \sin(\phi) s_y^{(i)} \), where the spin-1/2 operators \( s_x^{(i)}, s_y^{(i)} \) are defined as: \( s_x^{(i)} = (f_+^{(i)} f_-^{(i)} + f_-^{(i)} f_+^{(i)})/2, s_y^{(i)} = -i(f_+^{(i)} f_-^{(i)} - f_-^{(i)} f_+^{(i)})/2 \). The angles \( \phi \) are: \( \phi_{i\rightarrow j} = 0 \), \( \phi_{i\rightarrow j} = 2\pi/3 \), \( \phi_{i\rightarrow 3} = -2\pi/3 \), \( \phi_{i\rightarrow 2} = 2\pi/3 \), \( \phi_{i\rightarrow 4} = -2\pi/3 \), \( \phi_{i\rightarrow 4} = 2\pi/3 \). This Hamiltonian has previously appeared in the context of a block-spin approach to the Heisenberg KAF [13, 14]. The main purpose of that approach has been to find the origin of the exponentially large number of low-lying singlets that had been found in numerical studies of the kagomé antiferromagnet [15, 16]. From Refs. [13, 14] it also follows that \( H_{\text{trimer}} \) describes the physics of the trimerised KAF in a magnetic field that drives this system into the plateau region at 1/3 of the saturation magnetisation. We stress that the Hamiltonian \( H_{\text{trimer}} \) to be studied in this Letter describes a physically feasible situation.

Let us begin by discussing the classical theory of the model (1), which describes the large spin limit. In addition to being translationally invariant, the model of Eq. 1 is invariant under the point group of order 6, \( Z_6 = Z_3 \times Z_2 \), where the generator of \( Z_3 \) (order 3) is the combined rotation of the lattice by the angle \( 4\pi/3 \), and of the spins by the angle \( 2\pi/3 \), while the generator of \( Z_2 \) (order 2) is the spin inversion in the \( x-y \) plane. We remark that our model possesses no continuous spin rotational symmetry. There exist three ordered classical states with small unit cells that are compatible with this point-group symmetry of the model: a ferromagnetic state, and two 120° Néel type structures with left (Fig. 1b) and right (Fig. 2a) chiralities. The energies per site of these states are \( E_{\text{class}} = -3S^2 J/4 \) and \( E_{\text{class}} = 3S^2 J/2 \), where the subscripts “right” and “left” refer to chiralities. Hence, for \( J > 0 \) the state with left-handed chirality will be the ground state (GS). For \( J > 0 \) the situation is more complex: the states with right-handed chirality and the ferromagnetic state are degenerate ground states.

To understand further the nature of classical ground states we have done a numerical analysis of the 12-spin cell by fixing the direction of every spin to \( n\pi/3 \) \((n = 0 \cdots 5)\), and checking the energies of the resulting 612 configurations. This analysis has revealed that for \( J < 0 \) there are 6 ground states (\( Z_6 \) symmetry of Fig. 1) each of them exhibiting the left chirality Néel order. For \( J > 0 \) the results are dramatically different: there are 240 degenerate classical GSs in this case among them 6 pure right chirality Néel states and 6 purely ferromagnetic states. For an illustration, we show in Fig. 3 two ordered GSs with very large unit cells (Figs. 3a, 3b) together with their parent states (Figs. 3c, 3f). As will be seen below the large number of degenerate classical GSs finds its analogue in a large density of low-lying excitations of the quantum version of Eq. 1.

To describe the physics of spinless fermions on a trimerised optical kagomé lattice at filling 2/3 we need to consider the model, Eq. 1, for spin 1/2, i.e. in the extreme quantum limit. Questions to be answered for this case are: (i) Is the GS of the model Eq. 1 an ordered state, or is it a spin liquid either of type I, i.e. a state without broken symmetry, with exponentially fast decaying spin-pair correlations and a gap to the first excitation, or of type II, i.e. a kagomé-like GS again without broken symmetry, with extremely short ranged correlations, but with a dense spectrum of excitations adjacent to the GS; (ii) What are the thermal properties of our system? After all, the model can only be realized at finite, albeit low temperatures.

To answer the above questions we have performed exact diagonalisation of the Hamiltonian 1 for \( N = 12, 15, 18, 21 \), and 24 spins using the ARPACK routines 18. To simplify the calculations we block-diagonalised the Hamiltonian 1 by exploiting its translational symmetries thereby reducing the dimensions of the matrices that had to be diagonalised from \( 2^N \times 2^N \) to \( \approx 2^N/N \times 2^N/N \). Despite all these efforts, studies of larger systems require the use of massive computer resources. Fortunately, the
the large manifold of classical GSs by quantum effects implies the breaking of the translational and of the point group of our model Eq. 1 but there is no continuous symmetry that the ordered GS could break. Therefore, the standard expectation would be that the excitations have a gap of order of J. Instead, we find that the system has an exceptionally large number of low energy excitations (see Fig. 4). For instance, for N = 21 in the energy interval 0.1J/2 there are about 800 excited states. Most of them support the spin order of the GS so that this order persists at finite temperatures.

The analysis of the results for different N’s are compatible with an exponential increase of the number of low-energy states with the system size N similarly as in the case of the S = 1/2 KAF 15, 16. For the KAF Mila has been able to explain this high density of low energy states by associating them approximately to dimer coverings of an effective triangular lattice with uncorrelated products of nearest-neighbour pair states 14. His method fails here, because the low-lying states of our model must certainly be highly correlated. On account of the breaking of the discrete symmetries of our model by the Néel order, one expects the ground state of the infinite system to be sixfold degenerate. For finite systems this degeneracy is lifted. Nevertheless, we expect to find six low-lying states in the gap below the lowest excited state. In view of this scenario the inspection of the lower panel of Fig. 4 suggests that the gap, if any, is smaller than 10−2J/2. The appearance of this very small energy scale is completely unexpected and puzzling. Obviously, the answer to the questions (i) and (ii) above, is that the GS is ordered, and that the order survives at low T. The smallness of the gap and the large density of low-energy states, however, resemble very much the behaviour of a quantum spin liquid of type II. For these reasons we propose to term our system a quantum spin-liquid crystal. We remark in this context, that the specific heat of

### TABLE I: Spin correlations for N = 21 and 24 spins

| N       | J        | \( J/N \) |
|---------|----------|-----------|
| 21      | 1.0      | 2.0       |
| 24      | 1.0      | 2.0       |

FIG. 3: (colour online) (a) Right chirality Néel configuration; (b) localised defect in configuration (a); (c) ferromagnetic configuration; (d) line defect in configuration (c). Open arrows present spins determined by the boundary conditions for the 12-spin cell. Defects are marked by red dashed contours.

The selection of a GS with 120° planar Néel order from

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One can then increase 20 to the observation of the lattice. For instance one can start with a slower than the final 1/J lattice; by repeated measurement of the energy measure the energy of the system simply by opening the TABLE II: Spin-spin planar correlations for \( J < 0 \) as a function of distance 1, \( \ldots \), \( \sqrt{7} \) in lattice units.

\[ \begin{array}{|c|c|c|c|} 
\hline
\text{ } & 1 & \sqrt{3} & 2 & \sqrt{7} \\
\hline
N=21 & -0.134 & 0.237 & -0.117 & -0.116 \\
N=12 & -0.137 & 0.251 & -0.125 & \text{ } \\
\hline
\end{array} \]

The above results for \( J > 0 \) contrast dramatically with those for \( J < 0 \), summarised in Table II. In the latter case we deal with the standard quantum antiferromagnet with 120° Néel order and left chirality [19] (Fig. 1b). The spectrum is gapped, and the classical spin-spin correlations approximate the quantum correlations remarkably well. The gap is of the order \( |J|/2 \) in this case (\( N = 12, 18, 21 \)), meaning that there are at most a few states with energies substantially below \(|J|/2\) for \( J < 0 \), as opposed to the huge number for \( J > 0 \) (Fig. 4).

The observation of physics described in this Letter requires achieving low, but not unrealistic \( T \simeq 10 \text{mK} \) to 100nK (c.f. Ref. [12]). In experiments \( N \) could vary from \( \simeq 20 \) to \( \simeq 1000 \). The low energy states may be prepared by employing adiabatic changes of the degree of trimerisation of the lattice. For instance one can start with a completely trimerised lattice; the filling \( \nu = 2/3 \) may be achieved then by starting with \( \nu = 1 \), and eliminating 1 atom per trimer using, for instance, laser excitations. One can then increase \( t \) and \( U \) slowly, on the time scale slower than the final 1/J (\( \approx \text{seconds} \)). Alternatively, one could start with \( \nu \approx 2/3 \) in the moderately trimerised regime. As in Ref. [6], the inhomogeneity of the lattice due to the trapping potential, would then allow to achieve the Mott state with \( \nu = 2/3 \) per trimer in the centre of the trap. Nearly perfect 2/3 filling can be achieved by loading a BEC of molecules formed by 2 fermions into a triangular lattice, generating an MI state, adiabatically transforming the lattice to a trimerised kagomé one, “dissociating” the molecules by changing the scattering length to negative values, and by finally optically pumping the atoms into a single internal state. Preparing \( \nu = 2/3 \) might involve undesired heating (due to optical pumping), which can be overcome by using laser, or phonon cooling afterwards (cf. [20]). Note that the imperfections of \( \nu \) can be described by a "\( t - J \"-kind of model, and are of interest themselves.

After state preparation it should then be possible to measure the energy of the system simply by opening the lattice; by repeated measurement of the energy \( E(T) \) at (definite) finite temperatures one would get in this way an access to the density of modes, i.e. could compare the results with Fig. 4. From such measurements one could infer about the existence of the gap \( E_{\text{gap}} \), since if \( E_{\text{gap}} \) is large enough, \( E(T) \) becomes \( T \)-independent for \( kT \leq E_{\text{gap}} \). Various other correlations could be measured using the methods proposed in Ref. [21]. In order to measure planar spin correlations, one has, however, to lift the degeneracy of the \( f_\pm \) modes, e.g. by slightly modifying the intensity of one of the superlattices forming the trimerised lattice. This should be done on a time scale faster than the characteristic time scales of other interactions, so that the state of the system would not change during the measurement. In such a case one can use far off resonant Raman scattering (or scattering of matter waves) to measure the dynamic structure factor, which is proportional to the spatio-temporal Fourier transform of the density-density correlations. At frequencies close to the two photon Raman resonance between the \( f_\pm \) modes, only \( f_+ - f_- \) transitions contribute to the signal, and hence such measurement yields the desired information about the correlations \( \langle f_+^{(i)} f_+^{(j)} f_-^{(j)} f_-^{(i)} \rangle \), and the spin correlations of Fig. 2.

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