Zigzag Magnetic Order in the Iridium Oxide Na$_2$IrO$_3$

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We explore the phase diagram of spin-orbit Mott insulators on a honeycomb lattice, within the Kitaev-Heisenberg model extended to its full parameter space. Zigzag-type magnetic order is found to occupy a large part of the phase diagram of the model, and its physical origin is explained as due to interorbital $t_{2g} - e_g$ hopping. The magnetic susceptibility, spin wave spectra, and zigzag order parameter are calculated and compared to the experimental data, obtaining thereby the spin coupling constants in Na$_2$IrO$_3$ and Li$_2$IrO$_3$.

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In the quest for materials with novel electronic phases, iridium oxide Na$_2$IrO$_3$ came into focus recently [1–7] due to theoretical predictions [8,9] that this system may host Kitaev model physics and the quantum spin Hall effect. Na$_2$IrO$_3$ is an insulator with a sizable and temperature independent optical gap $\approx 0.35$ eV [7], and shows Curie-Weiss type susceptibility [1,6] with moments corresponding to an effective spin one-half Ir$^{4+}$ ion with a $t_{2g}$ configuration [10]. These facts imply that Na$_2$IrO$_3$ is a Mott insulator with well-localized Ir moments.

Collective behavior of local moments in Mott insulators is governed by three distinct and often competing forces: (i) orbital-lattice (Jahn-Teller (JT)) coupling, (ii) virtual hopping of electrons across the Mott gap resulting in exchange interactions, and (iii) relativistic spin-orbit coupling (see Ref. [11] for extensive discussions). The corresponding energy scales $E_{JT}$, $J$, and $\lambda$ vary broadly depending on the type of magnetic ions and chemical bonding [12]. When $\lambda > (E_{JT}, J)$, as often realized for Co, Rh, and Ir ions in an octahedral environment, local moments acquire a large orbital component which may result in a strong departure from spin-only Heisenberg models [8,11]. The direct observation of large spin-orbit splitting $3\lambda/2 \sim 0.6–0.7$ eV in insulating iridates Sr$_2$IrO$_4$ [13], Sr$_3$Ir$_2$O$_7$ [14], and Na$_2$IrO$_3$ [15] made it certain that $\lambda > (E_{JT}, J)$. Thus, the low-energy physics of Na$_2$IrO$_3$ is governed by interactions among the spin-orbit entangled Kramers doublets of Ir ions.

It is also established now [3–5] that Ir moments in Na$_2$IrO$_3$ undergo antiferromagnetic (AF) order at $T_N \approx 15$ K. The fact that $T_N$ is much smaller than the paramagnetic Curie temperature ($\approx 125$ K) [6] and spin-wave energies [4] implies that the underlying interactions are strongly frustrated. This is natural in the so-called Kitaev-Heisenberg (KH) model [16] where long range order is suppressed by the proximity to the Kitaev spin-liquid (SL) state. However, the observed “zigzag” magnetic pattern [ferromagnetic (FM) zigzag chains, AF coupled to each other] came as a surprising challenge to this simple and attractive model. To resolve the “zigzag puzzle”, a number of proposals, ranging from various modifications of the KH model [4,6,17–19] to a complete denial [20] of a local moment picture in Na$_2$IrO$_3$, have been put forward.

In this Letter, we show that the zigzag order is in fact a natural ground state (GS) of the KH model, in a previously overlooked parameter range. Next, we identify the exchange process that supports a zigzag-phase regime. Further, we calculate spin-wave spectra, the ordered moment, and magnetic susceptibility of the model in the zigzag phase, and find a nice agreement with experiment. This lends strong support to the KH model as a dominant interaction in Na$_2$IrO$_3$ and related oxides.

The model.—Nearest-neighbor (NN) interaction between isospin one-half Kramers doublets of Ir$^{4+}$ ions, coupled via $90^\circ$-exchange bonds, reads as follows (the exchange processes are described later):

$$\mathcal{H}^{(\gamma)}_{ij} = 2KS^\gamma_iS^\gamma_j + JS_i \cdot S_j.$$ (1)

Here, $\gamma(= x, y, z)$ labels 3 distinct types of NN bonds of a honeycomb lattice [16] of Ir ions in Na$_2$IrO$_3$, and spin axes oriented along the Ir-O bonds of IrO$_6$ octahedron. The bond-dependent Ising coupling between the $\gamma$ components of spins is nothing but the Kitaev model [21], and the second term stands for the Heisenberg exchange.

Let us introduce the energy scale $A = \sqrt{K^2 + J^2}$ and the angle $\phi$ via $K = A \sin \phi$ and $J = A \cos \phi$; the model (1) takes then the following form:

$$\mathcal{H}^{(\gamma)}_{ij} = A(2 \sin \phi S^\gamma_i S^\gamma_j + \cos \phi S_i \cdot S_j).$$ (2)

We let the “phase” angle $\phi$ vary from 0 to $2\pi$, uncovering, thereby, additional phases of the model that escaped attention previously [16], including its zigzag ordered state which is of a particular interest here.

It is instructive to introduce, following Refs. [11,16], 4 sublattices with the fictitious spins $\hat{S}$, which are obtained from $S$ by changing the sign of its two appropriate
components depending on the sublattice index. This transformation results in the $\hat{S}$ Hamiltonian of the same form as (1), but with effective couplings $\tilde{K} = K + J$ and $\tilde{J} = -J$, revealing a hidden $SU(2)$ symmetry of the model at $K = -J$ (where the Kitaev term $\tilde{K}$ vanishes). For the angles, the mapping reads as $\tan \varphi = -\tan \phi - 1$.

**Phase diagram.**—In its full parameter space, the KH model accommodates 6 different phases, best visualized using the phase-angle $\varphi$ as in Fig. 1(a). In addition to the previously discussed [16,22,23] Néel-AF, stripy-AF, and SL states near $\varphi = 0$, $-\pi$, and $-\frac{\pi}{2}$, respectively, we observe 3 more states. First one is “AF” ($K > 0$) Kitaev spin-liquid near $\varphi = \frac{\pi}{2}$. Second, the FM phase broadly extending over the third quadrant of the $\varphi$ circle. The FM and stripy-AF states are connected [see Fig. 1(a)] by the 4-sublattice transformation, which implies their identical dynamics. Finally, near $\varphi = \frac{3}{4} \pi$, the most wanted phase, zigzag AF, appears occupying almost a quarter of the phase space. Thanks to the above mapping, it is understood that the zigzag and Néel states are isomorphic, too.

In particular, the $\varphi = \frac{3}{4} \pi$ zigzag state is identical to the Heisenberg-AF state of the fictitious spins [24].

To obtain the phase boundaries, we have diagonalized the model numerically, using a hexagonal 24-site cluster with periodic boundary conditions. The cluster is compatible with the above 4-sublattice transformation and $\varphi \leftrightarrow \phi$ mapping. As seen in Fig. 1(b), the second derivative of the GS energy $E_{GS}$ with respect to the $\varphi$ well detects the phase transitions. Three pairs of linked transition points are found: $\approx (88^\circ, 92^\circ)$ and $(-76^\circ, -108^\circ)$ for the spin liquid-order transitions around $\pm \frac{\pi}{4}$, and $(162^\circ, -34^\circ)$ or the transitions between ordered phases.

The transitions from zigzag-AF to FM, and from stripy-AF to Néel-AF are expected to be of first order by symmetry; the corresponding peaks in Fig. 1(b) are indeed very sharp. The spin liquid-order transitions near $\varphi = -\frac{\pi}{2}$ lead to wider and much less pronounced peaks, suggesting a second- (or weakly first-) order transition [16]. On the contrary, liquid-order transitions around $\varphi = \frac{\pi}{2}$ show up as very narrow peaks; on the finite cluster studied, they correspond to real level crossings. The nature of these phase transitions remains to be clarified [25].

While at $J = 0$ (i.e., $\varphi = \pm \frac{\pi}{2}$) the sign of $K$ is irrelevant [21], the stability of the AF- and FM-type Kitaev spin liquids against $J$ perturbation is very different; the SL phase near $\frac{\pi}{2}$ ($-\frac{\pi}{2}$) is less (more) robust. This phase behavior is related to a different nature of the competing ordered phases: for the $\frac{\pi}{2}$ SL, these are highly quantum zigzag and Néel states, while the SL near $-\frac{\pi}{2}$ is sandwiched by more classical (FM and “fluctuation free” stripy [16]) states which are energetically less favorable than the quantum SL state.

**Exchange interactions in Na$_2$IrO$_3$.**—Having fixed the parameter space ($K > 0$, $J < 0$) for the zigzag phase, we turn now to the physical processes behind the model (1). Exchange interactions in Mott insulators arise due to virtual hoppings of electrons. This may happen in many different ways, depending sensitively on chemical bonding, intra-ionic electron structure, etc. The case of present interest (i.e., strong spin-orbit coupling, $t_{2g}^5$ configuration, and 90°-bonding geometry) has been addressed in several papers [8,11,16,26]. There are the following four physical processes that contribute to $K$ and $J$ couplings.

**Process 1:** Direct hopping $t'$ between NN $t_{2g}$ orbitals. Since no oxygen orbital is involved, 90° bonding is irrelevant; the resulting Hamiltonian is $H_1 = I_1 \mathbf{S}_i \cdot \mathbf{S}_j$ with $I_1 \approx (\frac{3}{4} t')^2/U$ [16]. Here, $U$ is the Coulomb repulsion between $t_{2g}$ electrons. Typically, one has $t' t < 1$, when compared to the indirect hopping $t$ of $t_{2g}$ orbitals via oxygen ions.

**Process 2:** Interorbital NN $t_{2g} - e_g$ hopping $\tilde{t}$. This is the dominant pathway in 90° bonding geometry since it involves strong $t_{pd\sigma}$ overlap between oxygen-2$p$ and $e_g$ orbitals; typically, $\tilde{t}/t \sim 2$. The corresponding Hamiltonian is [11]

\[
E_{GS} = \sum_{\langle i,j \rangle} t_{ij} \mathbf{S}_i \cdot \mathbf{S}_j + \sum_i J_i \mathbf{S}_i \cdot \mathbf{S}_i + \mathcal{H}_2 + \mathcal{H}_3.
\]
\[ H_2^{(y)} = I_3(2S_i^z S_j^z - S_i \cdot S_j). \] \tag{3} 

This is nothing but the model (1) with \( K = -J = I_2 > 0 \), i.e., at its \( SU(2) \) symmetric point \( \varphi = \frac{3}{4} \pi \) inside the zigzag phase; see Fig. 2. For the Mott-insulating iridates (as opposed to charge-transfer cobaltates \([11]\)), we estimate \( I_2 \approx \frac{\hbar}{2}(i/U)^2J_H \), where \( U \) is the (optically active) excitation energy associated with \( t_{2g} - e_g \) hopping, and \( J_H \) is Hund’s interaction between the \( t_{2g} \) and \( e_g \) orbitals. The physics behind this expression is clear: \((i/U)^2\) measures the amount of \( t_{2g} \) spin which is transferred to the NN \( e_g \) orbital; once arrived, it encounters the “host” \( t_{2g} \) spin and has to obey the Hund’s rule.

For its remarkable properties, the Hamiltonian \( H_2 \) (3) deserves a few more words. On a triangular lattice, it shows the “J” part of a doped \( t-J \) model, it leads to an exotic pairing \([11,28]\).

**Process 3:** Indirect hopping \( t \) between NN \( t_{2g} \) orbitals via oxygen ions. This gives rise to the Kitaev model \( H_3^{(y)} = -I_3S_i^z S_j^z \), with \( I_3 \approx \hbar^2/(2iU)(J_H/U) \) \([8]\), where \( J_H \) is Hund’s coupling between \( t_{2g} \) electrons. This process supports \( \varphi = -\frac{\pi}{2} \) SL state; see Fig. 2.

**Process 4:** Mechanisms involving \( pd \) charge-transfer excitations with energy \( \Delta_{pd} \). Two holes may meet at an oxygen and experience Coulomb \( U_p \) and Hund’s \( J_H^{\parallel} \) interactions, or cycle around a \( \text{Ir}_2\text{O}_2 \) plaquette (Fig. 2). The resulting Hamiltonian \( H_4 \) has the form of \( H_2 \) (3). The coupling constant \( I_4 \approx \hbar^2/(2\Delta_{pd} + U_p - J_H^{\parallel}) - \frac{1}{\Delta_{pd}} \) is negative \([29]\), supporting the stripy-AF state not observed in \( \text{Na}_3\text{IrO}_3 \).

Putting things together, we observe that it is the inter-orbital \( t_{2g} - e_g \) hopping \( H_2 \) process that uniquely supports zigzag order in \( \text{Na}_3\text{IrO}_3 \). This implies in general that multi-orbital Hubbard-type models, when applied to iridates with 90°-bonding geometry, must include \( e_g \) states as well, even though the moments reside predominantly in the \( t_{2g} \) shell.

Up to this point, we neglected trigonal field splitting \( \Delta \) of the \( t_{2g} \) level due to the \( c \) axis compression present in \( \text{Na}_3\text{IrO}_3 \). This approximation is valid as long as \( \Delta \) is much smaller than the spin-orbit coupling \( \lambda \approx 0.4 \) eV \([13,15,30]\) and seems to be justified, since the recent \( \text{ab initio} \) calculations \([20]\) suggest that \( \Delta \approx 75 \) meV only \([31]\).

We have also examined the longer-range couplings, using the hopping matrix of Ref. \([20]\), and found that the second-NN interaction has the form of (3) (as previously noticed Refs. \([32,33]\)), while the third-NN coupling is of the AF-Heisenberg type \([\text{the corresponding coupling constants are } \frac{1}{2}(t_{2g}^2/U)]\). The second (third)-NN interaction would oppose (support) zigzag order; however, we believe that these couplings are not significant in \( \text{Na}_3\text{IrO}_3 \) because the hoppings \( t_2 \) and \( t_3 \) are small \([34]\).

We do not attempt here to evaluate the parameters involved in \( H_1 - H_4 \); \( \text{ab initio} \) calculations as in Ref. \([35]\) might be more useful in this regard. Instead, having obtained a zigzag order in our model (1) and identified the physical process driving this order, we turn now to the experimental data. The \( J \) and \( K \) values in \( \text{Na}_3\text{IrO}_3 \) and \( \text{Li}_2\text{IrO}_3 \) will be extracted below from analysis of the neutron scattering and magnetic susceptibility data.

**Spin waves in the zigzag phase.**—Consider a single domain zigzag state, e.g., with FM chains running perpendicular to \( z \)-type bonds. Following Ref. \([4]\), we introduce a rectangular \( a \times b \) magnetic unit cell \( (\sqrt{3}a_0 \times 3a_0) \) in terms of hexagon-edge \( a_0 \); see Fig. 1(a)], and define the \( ab \)-plane wave vector \( \mathbf{q} \) in units of \((h, k)\) as \( \mathbf{q} = (\frac{2\pi}{3}h, \frac{2\pi}{3}k) \). Standard spin-wave theory gives four dispersive branches:

\[
\omega_{1,2}^2(h, k) = \left[ K^2 + (K + J)^2 \right] c_h^2 - KJ(1 - s_h s_k)
\]

\[
\pm |(K + J)c_h| \sqrt{2K - J}^2 - (2Ks_h - J s_k)^2,
\]

and \( \omega_{3,4}(h, k) = \omega_{1,2}(-h, -k) \), with \( c_h = \cos \pi h \), \( s_h = \sin \pi h \), and \( s_k = \sin \pi k \). If \( K = -J \), i.e., at the \( \varphi = \frac{3}{4} \pi \) point of hidden \( SU(2) \) symmetry, two branches are degenerate \((\omega_1 = \omega_2)\) and become true Goldstone modes. Away from this special point, the small magnon gap is expected to open by quantum effects not considered here. For \( \mathbf{q} \) with \( h = k \), the dispersions (4) simplify to \( \omega_1(h, h) = \sqrt{2K(2K + J)}c_h \) and \( \omega_2(h, h) = \sqrt{2}Jc_h \), revealing two different energy scales in the magnon spectra set by \( K \) and \( J \) couplings.
While the bandwidth of the lowest dispersive mode (set by $J$) is already known to be about 5–6 meV [4], we are not aware of the high energy magnon data to estimate $K$ in Na$_2$IrO$_3$. We have therefore examined (see below) the magnetic susceptibility data [1,6], and obtained $(J, K) \approx (-4.0, 10.5)$ meV that well fit the susceptibility as well as the neutron scattering data [4]. With this, we predict the magnon spectra for Na$_2$IrO$_3$ shown in Fig. 3. The lowest dispersive ($J$) mode is as observed [4], indeed. However, mapping out entire magnon spectra is highly desirable to quantify the Kitaev term $K$ directly.

Magnetic susceptibility.—We have calculated the uniform magnetic susceptibility $\chi(T)$ of the model (1) on 8- and 14-site clusters by exact diagonalization, and on 24-site cluster using the finite-temperature Lanczos method [36,37]. The parameters are varied such that $J = A \cos \varphi$ is consistent with the neutron data [4] while $\varphi$ stays within the zigzag sector of Fig. 1(a); this strongly narrows the possible $K$ window. For the data fits, we let the $g$ factor of the Ir$^{4+}$ ion deviate from 2 (due to the covalency effects [10]), and include the $T$-independent Van Vleck term $\chi_0$. The result for $J = -4.0$ meV, $K = 10.5$ meV, $g = 1.78$, $\chi_0 = 0.16 \times 10^{-3}$ cm$^3$/mol fits the Na$_2$IrO$_3$ data nicely (Fig. 4); deviations occur at low temperatures only, when correlation length exceeds the size of the cluster used. The fit is quite robust: similar results can be found for small only variations, locating Na$_2$IrO$_3$ near $\varphi = 111 \pm 2^\circ$ of the model phase diagram Fig. 1(a). The spin couplings obtained are reasonable for the 90$^\circ$-exchange bonds (as expected [8,11]), they are much smaller than in 180$^\circ$-bond perovskites [13,14]). The magnitude of the Van Vleck term also agrees with our estimate $\chi_0 \approx \frac{g^2}{3 \pi} \mu_B^2 N_A \approx 0.2 \times 10^{-3}$ cm$^3$/mol for the Ir$^{4+}$ ion, considering spin-orbit coupling $\lambda \approx 0.4$ eV [13,15,30].

Dominance of the Kitaev term ($2K/J \approx 5$ in Na$_2$IrO$_3$) implies strong frustration hence enhanced quantum fluctuations; this explains the reduced ordered moment $m \approx 0.22 \mu_B$ [5]. With the $J$, $K$, and $g$ values above, we calculated the leading order spin-wave correction to $m$ and obtained $m \approx 0.33 \mu_B$ [38].

For the sake of curiosity, we have also fitted the $\chi(T)$ data of Li$_2$IrO$_3$ [6], a sister compound of Na$_2$IrO$_3$. Acceptable results have been found for the angle window $\varphi = 124 \pm 6^\circ$; a representative plot for $J = -5.3$ meV, $K = 7.9$ meV, $g = 1.94$, $\chi_0 = 0.14 \times 10^{-3}$ cm$^3$/mol is shown in Fig. 4. It is worth noticing that the value of $J$, which controls the bandwidth of the softest spin-wave mode (see Fig. 3), appears to be similar in both compounds. This may explain why they undergo magnetic transition at similar $T_N \approx 15$ K, despite very different high temperature susceptibilities.

To conclude, we have clarified the origin of zigzag magnetic order in Na$_2$IrO$_3$ in terms of nearest-neighbor Kitaev-Heisenberg model for localized Ir moments. The model well agrees with the low-energy magnon and high temperature magnetic susceptibility data. A general implication of this work is that the interactions considered here should hold a key for understanding the magnetism of a broad class of spin-orbit Mott insulators with 90$^\circ$-exchange bonding geometry, including triangular, honeycomb, and hyperkagome lattice iridates.

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[24] After initial submission of this work, we became aware of the recent derivation [Y. Yu, L. Liang, Q. Niu, and S. Qin, Phys. Rev. B 87, 041107(R) (2013)] of the KH model and zigzag phase from a single band Hubbard model with the spin- and link-dependent NN-hoppings. However, such hoppings are not present in the hexagonal iridates $A_2IrO_3$ [8,9].
[29] The results for $I_2$ of Refs. [8,11,26] were incomplete.
[31] Typically, “noncubic” corrections to the interactions between Kramers doublets scale as $(\Delta/\lambda)^2$ [28], which is about 0.01 if $\Delta = 75 \text{ meV}$. $\theta$ is given by $\tan(\theta) = 2\sqrt{2}(\lambda + 2\Delta)/(\lambda + 2\Delta)$. The case of $\Delta > \lambda$ can be excluded also on the grounds that, in this limit, the interactions become bond-independent and support either Ising-FM or xy-AF states [11], instead of zigzag order observed.
[34] The second-NN hopping $t_2 = 75 \text{ meV}$ is about 4 times less than NN hopping $t$ via oxygen, and $t_2 = 30 \text{ meV}$ [20].
[37] We used $M = 100$ Lanczos steps and $N_d = 1024$ random sampling vectors. The values of $\chi$ and their statistical error are presented in Fig. 4 in the form of $3\sigma$ intervals estimated by taking many sets of the sampling vectors.
[38] Hybridization of the Ir-5$d$ and O-2$p$ orbitals on antiferromagnetic bonds [5], as well as the higher order quantum corrections may further reduce $m$. 

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