Partial flux ordering and thermal Majorana metals in higher-order spin liquids

Tim Eschmann, Vatsal Dwivedi, Henry F. Legg, Ciarán Hickey, and Simon Trebst

Institute for Theoretical Physics, University of Cologne, 50937 Cologne, Germany

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In frustrated quantum magnetism, chiral spin liquids are a particularly intriguing subset of quantum spin liquids in which the fractionalized parton degrees of freedom form a Chern insulator. Here we study an exactly solvable spin-3/2 model, which harbors not only chiral spin liquids but also spin liquids with higher-order parton band topology—a trivial band insulator, a Chern insulator with gapless chiral edge modes, and a second-order topological insulator with gapless corner modes. With a focus on the thermodynamic precursors and thermal phase transitions associated with these distinct states, we employ numerically exact quantum Monte Carlo simulations to reveal a number of unconventional phenomena. This includes heightened thermal stability of the ground state phases, the emergence of a partial flux ordering of the associated $Z_2$ lattice gauge field, and the formation of a thermal Majorana metal regime extending over a broad temperature range.

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I. INTRODUCTION

The emergence of topological phases from local constraints induced by competing interactions in frustrated quantum magnets has fascinated researchers for decades [1,2]. In a groundbreaking conceptual work, Kalmeyer and Laughlin [3] in the late 80s put forward the formation of a bosonic analog of the fractional quantum Hall state in what has since been termed a chiral spin liquid. First envisioned as resonating valence bond (RVB) ground states of geometrically frustrated Heisenberg antiferromagnets and relevant to high-temperature superconductivity [4,5], such chiral spin liquids have remained elusive for many years. In the past decade, however, tremendous progress has been achieved in firmly establishing chiral spin liquids as ground states of microscopic models [6–11] and their experimental observation in quantized thermal Hall measurements [12,13]. On the theoretical side, conceptual insight has been gained from the unambiguous numerical detection of chiral spin liquids via the calculation of modular matrices [14,15] from the ground state entanglement structure [16] of a number of kagome models [10,11]. Analytically, Kitaev showed that a time-reversal symmetry breaking magnetic field gives rise to a non-Abelian chiral spin liquid in his eponymous model [6]. The advent of Kitaev materials [17] has since produced a direct experimental observation of this state in measurements of a half-quantized thermal Hall effect in the spin-orbit entangled Mott insulator $\alpha$-RuCl$_3$ [12].

The physical mechanism underlying the formation of a chiral spin liquid can be elegantly formulated using Wen’s parton construction [18]. At low temperatures, the original spin degrees of freedom fractionalize into a parton coupled to an emergent lattice gauge field [2]. For the Kitaev model, this parton decomposition is typically done [6] in terms of Majorana fermions coupled to a $Z_2$ gauge field—thereby casting the original interacting spin model to a free (Majorana) fermion problem with a static $Z_2$ gauge order (at zero temperature). One can then resort to the tools of topological band theory to classify the band structure of the emergent Majorana fermions, and thereby classify the fundamental topological properties of the spin liquid state. For the case of the Kitaev honeycomb model, this perspective results in an understanding of the field-induced topological spin liquid as the formation of a Chern insulator (in the Majorana band structure) with all bands carrying a nontrivial Chern number $|\nu|=1$. Such a state—a Chern insulator of the emergent Majorana fermions—is gapped in the bulk, but exhibits chiral gapless modes at its boundary that are topologically protected, as depicted in Fig. 1(b). That such a Majorana Chern insulator indeed forms in the Kitaev material $\alpha$-RuCl$_3$ is substantiated by the observation of a half-quantized thermal Hall conductance, which is direct evidence for Majorana fermions (and not conventional electrons) carrying the thermal edge currents. Subsequent measurements [13] of the anomalous character of the quantized thermal Hall effect, which arises even without any perpendicular magnetic field component, have brought verification that the observed state is indeed arising from the formation of topological Chern bands (and not the formation of Landau levels).

In parallel, the field of topological band theory has been developed [21]. There one distinguishes strong topological

$^{†}$Present address: Department of Physics, University of Basel, Klinikgasse 82, CH-4056 Basel, Switzerland.
insulators (TIs), whose topological features are protected by

time reversal and/or charge conjugation symmetries, from
crystalline TIs whose topology is endowed by certain lattice

symmetries. A particular example of such crystalline TIs are

second-order topological insulators, an instance of higher-

order topology [22]. An $n$th-order TI in $d$ spatial dimensions

exhibits $(d - n)$-dimensional topologically protected gapless

modes that are localized at the intersection of $n$ boundary

planes, while the boundaries of codimension less than $n$

remain fully gapped. A second-order TI in two spatial di-

mensions thus exhibits topologically protected corner modes,

i.e., zero-dimensional gapless modes at the intersection of two

boundaries giving rise to a corner. Such higher-order topology

can also play out in the context of a quantum spin liquid—

with the itinerant fractionalized parton degrees of freedom

forming a nontrivial topological band structure. The example

of a second-order spin liquid with topologically protected

Majorana corner modes has been discussed in the context

of an exactly solvable spin-3/2 generalization of the Kitaev

model [23]. Within this same framework, the chiral spin liquid

can be thought of as a first-order spin liquid, since a Chern

insulator can be considered an example of a first-order topo-

logical insulator.\(^2\)

In this manuscript, we study the thermodynamic precursors

and symmetry-breaking thermal phase transitions leading to

the formation of a family of spin liquid ground states which

exhibit the full range of parton band topology, second-order,

conventional (first-order), and trivial topology, in a general-

ized Kitaev model. We employ sign-problem free quantum

Monte Carlo simulations in the parton basis [24]. These

numerically exact calculations allow us to track the fractiona-

lization of the original spin degrees of freedom, the formation

of gauge order, and the spontaneous breaking of time-reversal

symmetry upon entering the different flavors of spin liquid

ground states. Our main results include (i) the observation

that the thermal stability of the chiral spin liquid in our model

is enhanced by almost an order of magnitude in comparison

with other chiral spin liquid models, with the highest transition

temperatures reaching about 1/10 of the bare coupling

strength; (ii) the emergence of partial flux order in an inter-

mediate temperature range, accompanied by a characteristic

three-peak signature in the specific heat; and (iii) the for-

mation of a gapless phase at finite temperatures that is best

described as a thermal Majorana metal.

Our discussion of these results in the remainder of the

manuscript is structured as follows. In Sec. II, we briefly

introduce a generalized spin-3/2 Kitaev model, its $\Gamma$-matrix

representation, and the underlying five-coordinated Shastry-

Sutherland lattice. In discussing its analytical solution at zero

temperature, we also introduce the parton basis relevant to our

sign-free QMC simulations to explore the thermodynamics at

finite temperatures. The formation of a conventional (first-

order) chiral spin liquid is discussed in Sec. III. Our main

results on thermal stability, partial flux ordering, and thermal

metal formation are all discussed in detail here. In Sec. IV,

we then turn to the formation of a second-order spin liquid,

whose zero-temperature properties we previously discussed in

Ref. [23]. Our focus here is on its thermodynamic properties.

We conclude with an outlook in Sec. V.

\section*{II. THE SHASTRY-SUTHERLAND KITAEV MODEL}

We start our discussion with a brief review of the general-

ization of the Kitaev model to the Shastry-Sutherland lattice

[23,25], its fundamental (lattice) symmetries, the formation

of spin liquid ground states of various levels of topology, and its

numerical representation in sign-free quantum Monte Carlo

simulations.

\subsection*{A. The model: spin-3/2 and Gamma matrices}

The Kitaev honeycomb model is the paradigmatic example

of an exactly solvable quantum spin liquid model. The model

consists of spin-1/2 degrees of freedom on the sites of a

honeycomb lattice interacting via bond-dependent Ising

interactions. By representing the spin operators in terms of

Majorana fermions, the model can be reduced to a nearest-

neighbor hopping model of noninteracting fermions coupled
to a static $\mathbb{Z}_2$ gauge field. The Kitaev model and its exact

solution can be straightforwardly generalized to other lattices

with an odd coordination number, $z = 2n - 1$, wherein the

local “spins” are decomposed into $2n$ Majorana fermions.

Here, we study such a generalization of Kitaev’s hon-
neycomb model to the pentacoordinated Shastry-Sutherland

lattice, previously introduced in Refs. [23,25] (see Fig. 2). The

lattice is most well-known for the orthogonal dimer model,

which was solved by Shastry and Sutherland [26] and serves

as an effective low-temperature model for the transition metal

oxide SrCu$_2$(BO$_3$)$_2$ [27]. The generalized Kitaev model is

\footnote{Note that, though all of the spin liquid ground states spontaneously
break time-reversal symmetry, we reserve the term “chiral spin li-
quid” for those that exhibit chiral edge states (or, more technically,
those that possess a nonzero chiral central charge).}
described by the Hamiltonian
\[
\mathcal{H} = -\sum_{(j,k)\gamma} J^\gamma_j \Gamma^\gamma_j \Gamma^\gamma_k,
\]
where \(\gamma = 1, \ldots, 5\) labels the bond direction and we have a set of five \(4 \times 4\) \(\Gamma\)-matrices on the lattice sites representing spin-3/2 degrees of freedom. The horizontal and vertical (blue) bonds carry a coupling \(J\) (which we set to unity), and the diagonal (red) bonds a coupling \(J_d\). For a suitable choice of these parameters, the resulting higher-order spin liquid system is shown to possess a topologically nontrivial ground state.

To solve this model exactly, we represent the \(\Gamma\) matrices on each site in terms of six Majorana operators \(c_j\) and \(b_j^\dagger\) by setting \(\Gamma^\gamma_j = ib_j^\dagger c_j\). The “bond Majoranas” \(b_j\) are recombined into bond operators \(u_{jk} = ib_j^\dagger b_k\). Since all \(u_{jk}\) commute with the Hamiltonian, they can be replaced by their eigenvalues \(u_{jk} \pm 1\). We are thus left with a hopping model of Majorana fermions \(c_j\) coupled to a static \(\mathbb{Z}_2\)-gauge field \(u_{jk}\), described by the Hamiltonian
\[
\mathcal{H} = \frac{i}{2} \sum_{j,k} J^\gamma_{jk} u_{jk}^\gamma c_j c_k.
\]

The spectrum of \(\mathcal{H}\) must be invariant under \(\mathbb{Z}_2\) gauge transformations, and can thus depend only on the \(\mathbb{Z}_2\) fluxes \(W_p\). Associated with plaquettes \(p\), defined as
\[
W_p = \prod_{(j,k)\in p} (-iu_{jk}),
\]
where, as a convention, the product here is taken with a clockwise orientation. The Shastry-Sutherland lattice has two types of elementary plaquettes, viz, square plaquettes with flux \(W_\Box = \pm 1\), and triangular plaquettes with flux \(W_\triangle = \pm i\).

### B. Sign-problem free sampling of flux configurations

To determine the ground state, one primary task is to find the flux configuration that minimizes the total energy. This is generally a nontrivial matter and one that can almost never be resolved in an analytically exact fashion—with the honeycomb Kitaev model being the most notable exception, for which a theorem by Lieb [28] can be invoked. For the general case one can, however, rely on a numerically exact treatment by performing quantum Monte Carlo (QMC) sampling of the gauge field (while keeping track of all Majorana physics) [23,24]. Such a QMC approach is possible without encountering a sign problem [29] in the spin decomposition introduced above, which allows to treat the gauge field as a classical degree of freedom and incorporating the free Majorana fermions via an exact diagonalization to compute the associated statistical weights. This sign-free QMC approach should be contrasted with recent implementations of sign-free fermion models that are based on an antunitary symmetry [30,31] such as particle-hole symmetry (intrinsically in Majorana fermion systems), time-reversal symmetry and variations thereof [32–36].

In more technical terms, in such a sign-problem-free QMC approach the \(\mathbb{Z}_2\) gauge degrees of freedom \(u_{jk}\) are sampled as classical Ising variables, with the Majorana quantum physics entering in the weights of the Markov chain sampling. For a fixed gauge configuration, the noninteracting Majorana Hamiltonian [Eq. (2)] can be numerically diagonalized, a computation which scales as \(N^3\), where \(N\) is the number of lattice sites [24,37]. All thermodynamic observables can then be obtained without needing to introduce an additional imaginary-time dimension or other nontrivial mapping schemes for the quantum problem. A comprehensive discussion of the technicalities associated with this QMC approach to the Shastry-Sutherland Kitaev model was given in Ref. [23] and is summarized in Appendix A.

Performing such a quantum Monte Carlo (QMC) simulation readily demonstrates that the ground state flux order corresponds to \(W_\Box = -1\) for all square plaquettes. However, though \(W_\Box = W_\triangle^2\), the value of \(W_\triangle\) itself is not fixed. The two possibilities \(W_\triangle = \pm i\) are related by time-reversal symmetry (TRS) and degenerate in energy. At zero temperature, the ground state must thus spontaneously break TRS by selecting either \(W_\triangle = i\) or \(W_\triangle = -i\) for all triangular plaquettes. This spontaneous breaking of time-reversal symmetry is generally true for any Kitaev-type model on a lattice containing plaquettes with an odd number of bonds, as first noted by Kitaev [6], and later elucidated within a concrete spin-1/2 model by Yao and Kivelson [7]. This is precisely what is seen in our thermodynamic QMC data for finite temperatures, as we will discuss in detail below.
C. Symmetries and the ground-state phase diagram

On a conceptual level, the possible ground-state phases of the (Majorana) Hamiltonian can be inferred from the tenfold classification of topological insulators and superconductors [38,39]. The Majorana Hamiltonian, by design, obeys a particle-hole symmetry (PHS) which squares to +1, while time-reversal symmetry is broken spontaneously, as discussed earlier. Thus this Majorana Hamiltonian belongs to symmetry class D. In two spatial dimensions, this symmetry class D allows for a Z invariant, viz., a Chern number, signaling the possibility of the formation of topological order. Such a topological state is precisely the chiral spin liquid (with a gapless chiral edge mode) discussed in the introduction.

If we further restrict to the case where all the “square lattice” bonds are of equal strength (which we set to \( J = 1 \)), the only remaining parameter is \( J_d \), the strength of the diagonal “dimer” bonds of the lattice. As a function of varying \( J_d \) the Majorana band structure exhibits two gapped phases: a topological phase with Chern number \( C = \pm 1 \) for \( J_d < 2\sqrt{2} \) and a trivial one for \( J_d > 2\sqrt{2} \). The two gapped phases are separated by a gap closing at \((\pi, \pi)\), as illustrated in the lower panel of the schematic phase diagram in Fig. 3. The \( C = \pm 1 \) phase is a chiral spin liquid with a chiral Majorana edge mode and bulk Ising anyon topological order. On the other hand, the \( C = 0 \) phase, while still spontaneously breaking TRS, is fully gapped and possesses the same Abelian topological order as the toric code. We refer to this phase as a “trivial” spin liquid.

III. TRIVIAL AND CHIRAL SPIN LIQUIDS

Coming to the actual results of our finite-temperature analysis of the Shastry-Sutherland Kitaev model, we first concentrate on the thermodynamic behavior above the phase transition from the topological (first-order) spin liquid (with Chern number \( v = \pm 1 \)) to the trivial spin liquid (with Chern number \( v = 0 \)).

A. Thermodynamics

A central quantity distinguishing the various finite-temperature phases of our model is the specific heat \( C_v(T) \) and its characteristic multipeak structure, as illustrated in Fig. 4. For the closely related Kitaev spin liquids, it is well established [24] that one finds two well-separated peaks in the specific heat—a smooth high-temperature peak, indicating a thermal crossover, corresponding to the (local) fractionalization of spins and a second low-temperature peak associated with the freezing of the \( Z_2 \) gauge field. We observe a similar two-peak structure of the specific heat in our model for a broad range of parameters \( J_d \gtrsim 1 \) as well. Plotted in Fig. 4 is a characteristic \( C_v(T) \) trace in comparison with data for the Kitaev honeycomb model for similar system sizes. Both systems show a shallow high-temperature crossover around the value of the elementary coupling strength, in our case \( T \sim 2 \), whose shape and height are essentially independent of the system size—indicating a purely local crossover. In fact, this is where the spin fractionalization happens and the system releases precisely half of its entropy [23,24]. A more pointed distinction is found in the low-temperature peak—this...
is a sharp peak for the model at hand (that sharpenes with increasing system size), while it is a more shallow feature in the Kitaev honeycomb model. This is an immediate reflection of the fact that in the model at hand there is a true phase transition occurring at this lower temperature—the spontaneous breaking of time-reversal symmetry upon entering the chiral spin liquid regime, while in contrast the Kitaev honeycomb model exhibits a finite-temperature crossover at this lower temperature scale at which, for a given system size, the $Z_2$ gauge field freezes into its ground-state configuration. While the latter is a true phase transition in three spatial dimensions [40], it remains a thermodynamic crossover in two spatial dimensions [41,42].

The key distinction between these model systems is that the Shastry-Sutherland lattice is nonbipartite and exhibits elementary (triangular) plaquettes with an odd number of bonds. For the emergent Majorana fermions, this results in an am-

mentary (triangular) plaquettes with an

The latter is a true phase transition in three spatial dimensions [40], it remains a thermodynamic crossover in two spatial dimensions [41,42].

The key distinction between these model systems is that the Shastry-Sutherland lattice is nonbipartite and exhibits elementary (triangular) plaquettes with an odd number of bonds. For the emergent Majorana fermions, this results in an ambiguous situation in which they can pick up a phase $e^{i\pi/2}$ upon hopping around such a triangular plaquette, endowing the latter with a flux $W_\triangle = \pm i$ as discussed in the previous section. By spontaneously breaking time-reversal symmetry, one of the two possible signs is chosen, with the system simultaneously undergoing an Ising-type phase transition. Such a time-reversal symmetry breaking thermal phase transition was first observed in the Yao-Kivelson model (on a decorated honeycomb lattice) [43] at a temperature scale $T_c \sim 10^{-2}J$. This is also the typical temperature scale for thermal phase transition in 3d Kitaev models. In comparison, the critical temperature scale of the model at hand is elevated, see Fig. 4, with its maximum close to one-tenth of the Kitaev coupling (at $J_d \sim 1.2$). One might speculate that this enhanced transition temperature is a reflection of the higher coordination number of the Shastry-Sutherland lattice ($z = 5$) in comparison to conventional Kitaev models on tricoordinated lattice geometries. This idea, however, does not hold up when further generalizing our model to a seven-coordinated lattice (by placing additional diagonal bonds on the lattice), which has a transition temperature of the same order of magnitude as the Shastry-Sutherland case.

B. Partial flux ordering

Upon closer inspection, the formation of flux order at low temperatures turns out to be slightly more intricate. As noted above, the spontaneous breaking of time-reversal symmetry is intimately connected with the flux ordering of the triangular plaquettes. With two such triangular plaquettes constituting a single square plaquette, this also implies ordering for the latter. The corresponding flux satisfies $W_\Box = W_\triangle \cdot W_\triangle = +1$, independent of the actual assignment of the triangular plaquettes. Thus an ordering of the 3-plaquettes implies ordering of at least half the 4-plaquettes. The converse is, however, not true; we can have ordering of the 4-plaquettes while the 3-plaquettes remain disordered, resulting in a partial flux ordering. This is exactly what we observe for a limited parameter range $0 < J_d \lesssim 1$, where the 4-plaquette fluxes order at a higher temperature $T' \neq T_c$ for the 3-plaquette fluxes. This is illustrated in Fig. 5 where we plot the phase diagram of our model by color coding the flux of the 3-plaquettes (top panel) and 4-plaquettes (bottom panel) as function of $J_d$. A schematic rendering of the intermediate partial flux ordering is provided in Fig. 6.

The evolution of this partial flux ordering with varying coupling strengths can be seen in the sequence of data sets for vertical cuts through the phase diagram provided in Fig. 7. The top row shows the gauge contribution to the specific heat (i.e., omitting the Majorana contribution resulting in the higher temperature crossover). In the partial flux ordering regime $0 < J_d \lesssim 1$, one finds that the low-temperature specific heat peak actually splits into two parts—with the lower temperature peak indicating the true thermal phase transition associated with time-reversal symmetry breaking and 3-plaquette flux ordering, see also the medium row of panels. While this transition quickly moves to zero temperature

\footnote{Such a $\pi$-flux ground state for square plaquettes is also generally in line with the expectation from Lieb’s theorem on ground-state flux assignments in bipartite lattices with certain mirror symmetries [28], tough it does not strictly apply to the lattice geometry at hand.}
FIG. 6. Partial flux ordering. At $T_c \sim 0.1 J_d$, all triangle plaquettes order into one of the two homogeneous flux configurations (a). For $J_d \lesssim 1$, the square plaquettes assume an ordered $\pi$-flux configuration at $T''$ (b). In this regime, the triangle plaquettes remain disordered, assuming pairwise fluxes $W_\Delta = \pm i$ (darker/lighter yellow).

as $J_d \to 0$, there remains a signature in the specific heat around $\mathcal{O}(1/10)$, which upon closer inspection is the thermal crossover associated with the ordering of the remaining half of the 4-plaquettes, see the lower row of panels.

Taking a step back, we conclude that the emergence of a partial flux ordering is, for a limited range of parameters, a precursor phenomenon to the formation of a low-temperature topological chiral spin liquid.

C. Thermal Majorana metal

The thermodynamic signatures discussed so far—the finite-temperature, time-reversal symmetry breaking phase transition as well as the emergence of a partial flux ordering and its associated thermal crossover—are both closely connected to the underlying lattice gauge theory. From a more conceptual perspective, these aspects are an interesting variation to the gauge physics that has been intensely studied in the context of two- and three-dimensional Kitaev models [24,40]. We now turn to additional thermodynamic aspects of our model at hand, which are genuinely rooted in the physics of the emergent Majorana fermions. The most notable feature here is the formation of a thermal metal regime above the transition to the topological chiral spin liquid as illustrated in the schematic phase diagram of Fig. 3, and as evidenced in the numerical observations of Fig. 8. This thermal metal regime, principally located in the intermediate temperature regime $T_c \lesssim T \lesssim T'$ (i.e., between spin fractionalization and the time-reversal symmetry breaking transition), does not extend over the entire parameter space of our model, but marks a relatively sharp transition for a critical value of $J_d$ that is considerably shifted in comparison with the zero-temperature transition between the chiral and trivial spin liquids. We rationalize these numerical observations using analytical arguments based on the analytical (self-consistent) Born and T-matrix approximations along with a numerical computation using transfer matrices. We thus explain how the entire

FIG. 7. Thermodynamics. For different cuts through the phase diagram of Fig. 5, we show (a) the specific heat $C_{\text{eff}}(T)$ ($\mathbb{Z}_2$ gauge field contribution), (b) the average 3-plaquette flux $|W_\Delta|$, and (c) and the average 4-plaquette flux $W_\Box$ for various values of $J_d$. The low-temperature peak in $C_{\text{eff}}(T)$ indicates a thermal phase transition (dashed lines), which is associated with the ordering of the fluxes $W_\Delta = \pm i$ on the triangular plaquettes of the Shastry-Sutherland lattice. The difference between the transition temperature for the 3-plaquettes and the 4-plaquettes in the left column indicates an intermediate partial flux order.
gapless thermal metal phase emanates from the quantum critical point between the chiral and trivial spin liquid at zero temperature.

1. Numerical observations

In our numerics, the existence of two distinct regimes above the low-temperature chiral spin liquid phases is most evident in calculations of the average Chern number $|\nu|$ of the Majorana band structure encountered when sampling (and averaging over) the different gauge configurations for a given temperature. The behavior of this average Chern number as a function of $J_d$ is plotted in Fig. 8(a), which immediately reveals three distinct regimes. For a horizontal cut in the lowest temperature regime, the average Chern number jumps from $|\nu| = 0$ in the trivial phase to $|\nu| = 1$ in the topological phase, as expected. Unanticipated is probably the behavior for a horizontal cut in the intermediate-temperature regime, where our simulations also reveal a relatively sharp (vertical) boundary at $J_d \approx 1.9$, at which the average Chern number mimics the low-temperature behavior, jumping from $|\nu| = 0$ for large couplings $J_d \gtrsim 1.9$, to $|\nu| \approx 0.6$ for $J_d \lesssim 1.9$ (see also the scans of $|\nu|$) shown in Figs. 19 and 20 of the Appendix). This clearly separates two different thermodynamic regimes, which seemingly persist not only up to the fractionalization crossover scale but beyond into the paramagnetic regime extending all the way up to infinite temperatures.

One important aspect in interpreting this numerical result, in particular the nonquantization of the average Chern number for $J_d \lesssim 1.9$, is to ask whether the Majorana band structure actually remains gapped—a prerequisite for the proper calculation of a Chern number—also at finite temperatures. In fact, this is precisely what distinguishes the two regimes: while the system remains gapped for $J_d \gtrsim 1.9$, it becomes gapless for smaller couplings. That the phase for $J_d \lesssim 1.9$ is in fact gapless can be shown in a straightforward manner by computing the gap in the Majorana spectrum averaged over flux configurations sampled with a uniform distribution, or equivalently, at infinite temperature. As shown in the top panel of the schematic phase diagram of Fig. 3, the gap in the Majorana spectrum indeed vanishes for small $J_d$ and slowly opens only for $J_d \gtrsim 1.9$, as compared to the scenario of a gap closing and reopening for $J_d = 2\sqrt{2} \approx 2.8$ in the zero-temperature case. The Chern number results are thus only well-defined in the gapped regimes of the phase diagram, $J_d \gtrsim 1.9$ and $T < T_c$. For detailed scans of the average Chern number we refer to Appendix B.

Having established the principal gapless character of the finite-temperature spectrum for $J_d \lesssim 1.9$, a more meaningful quantity to calculate is the effective number of states accessible to the system at a given temperature

$$N_0(T) = -\int_0^\infty dE \rho(E) \partial_E n_F(E, T),$$

where $\rho(E)$ is the Majorana density of states and $n_F(E, T)$ is the Fermi function, whose derivative has a peak at $E \approx 0$ of width $\sim T$. Plotting $N_0(T)$ as a function of $J_d$ for various temperatures in Fig. 9, we indeed see that the number of available state becomes finite for small $J_d$, indicating a gapless state, and vanishes only near the transition point $J_d \approx 1.9$, indicating a truly gapped state for larger $J_d$.

We note in passing that the partial flux ordered phase also appears to be gapped, as $N_0(T) \to 0$ in the corresponding parameter regime, i.e., for $J_d \lesssim 0.3$ and $T \lesssim 0.04$. This gap opens at the crossover to the thermal metal phase, as indicated by the increase in $N_0(T)$. This is also indicated in the Chern number plot of Fig. 8(a), where a narrow blue stripe above the partially flux-ordered phase indicates the thermal crossover points associated with the ordering of square plaquettes. Thus, for $J_d < 1$, the Kitaev Shastry-Sutherland system starts in the gapped chiral spin liquid ground state at $T = 0$, undergoes a thermal phase transition into a gapped partial flux-ordered
phase, and the gap finally collapses at the crossover to the intermediate temperature phase.

2. Symmetry-class considerations

To interpret these numerical observations, it is useful to remind oneself of the symmetry class classification of free fermion systems \cite{44,45} and its application to topological states of matter \cite{38,39}. When considering the Majorana fermion representation of our model, the system naturally exhibits particle-hole symmetry (due to the self-conjugation of Majorana fermions) and broken time-reversal symmetry, which is implied by the nonbipartite geometry of the lattice as discussed earlier. This puts our model into symmetry class D.

In the tenfold way classification of topological states of matter \cite{38,39}, this symmetry class allows, in two spatial dimensions, for the formation of both a trivial and a topological insulator (with vanishing and integer Chern number, respectively). In addition, it is well established that in the presence of disorder this symmetry class allows for the formation of a gapless “thermal metal” phase \cite{46–51}.

In the Kitaev Shastry-Sutherland model, all three of these phases are found to be realized. In the low-temperature regime, where time-reversal symmetry is spontaneously broken and the gauge field is ordered, we realize the two “clean” phases of symmetry class D—the trivial insulator (with Chern number $\nu = 0$) for $J_d > 2\sqrt{2}$ and the topological insulator (with Chern number $\nu = \pm 1$) for $J_d < 2\sqrt{2}$. Going to the intermediate-temperature regime, i.e., above the ordering transition at $T_c$, a more subtle situation arises: Here the $\mathbb{Z}_2$ gauge field does not exhibit any order, effectively providing a static disorder potential for the Majorana system. And while time-reversal symmetry (TRS) is not broken on a global level, which strictly speaking does not put the system into symmetry class D anymore, TRS is broken for any given static disorder configuration: Each particular flux configuration must have a fixed value of flux $(\pm i)$ across the 3-plaquettes, and hence breaks time-reversal symmetry. The ensemble of Majorana Hamiltonians thus always corresponds to the symmetry class D at any temperature, even if the true state of the Kitaev model is itself time-reversal invariant. This allows the system to principally form the thermal metal phase of symmetry class D.

This thermal metal phase is indeed realized in the intermediate temperature regime for $J_d \lesssim 1.9$, i.e., in the parameter regime that our numerical experiments indicated as gapless. The very nature of the thermal metal phase can be probed explicitly via a calculation of the low-energy density of states $\rho(E)$ of the Majorana fermions, which is supposed to show a characteristic “ringing” \cite{46}. For $E \to 0$, the latter takes the universal form

$$\rho(E) = \alpha + \frac{\sin(2\pi \alpha EL^2)}{2\pi EL^2}$$

with a single fitting parameter $\alpha$. This is indeed what we find upon numerical inspection, as shown in Fig. 10 for a parameter set deep in the thermal metal regime ($J_d = 1.5$ and $T = 1.86$). Similar behavior has first been reported for the honeycomb Kitaev model in a magnetic field in a broad temperature regime above the thermal crossover into its chiral spin liquid ground state \cite{52}.

3. Analytical perspective

The essential physics of the formation of the thermal metal phase can be understood analytically by analyzing our Majorana hopping model with uncorrelated $\mathbb{Z}_2$ flux disorder.\(^5\) Explicitly, this implies that arbitrary disorder configurations can be obtained by starting from the ground state flux configuration and flipping each plaquette flux with a probability $p$, independent of all other fluxes. This disorder density $p$ then roughly corresponds to the temperature in the full Kitaev model. In particular, $p = 0$ is the ground state ($T = 0$), while $p = 1/2$ is the state with completely random fluxes ($T = \infty$).

We analytically study the effect of this flux disorder by computing the self-energy as a power series in the disorder density—which corresponds to a series in the number of scattering events—using the self-consistent Born approximation (SCBA) as well as the self-consistent T-matrix approximation (SCTA). We find that the net effect of the disorder is a renormalization of the bare coupling strength $J$ (which we have set to 1 so far), so that the new phase boundary occurs at

$$J_d = 2\sqrt{2} J_{\text{cut}}(p) \approx 2\sqrt{2} (1 - \Sigma_p(0, k_0)),$$

where $k_0 = (\pi, \pi)$ is the gap closing point at $J_d = 2\sqrt{2}$ in the clean limit (see Appendix C for a detailed derivation). In Fig. 11, we plot this phase boundary as a function of the disorder density. We find that for $p \lesssim 0.25$, the disorder-induced renormalization of $J$ leads to a continuous shift of the phase

\(^5\)For the full Kitaev model at finite temperature, the disorder is far from uncorrelated, since the probability of a disorder realization is the Boltzmann weight corresponding to all the vison excitations required to arrive at it from the ground state flux configuration.
The fraction of plaquettes with zero flux, so that \( y \) (SCBA) and T-matrix (SCTA) approximations. The laid with the phase boundaries obtained from self-consistent Born channels for the Majorana hopping model on a cylinder, over-quantum critical point. Shown is the effective number of gapless jumps abruptly from \( J \) diagram w.r.t. temperature (Fig. 8), where we see the phase boundary the Ising transition, this entire range is compressed in the phase boundary between the trivial and topological gapped phases.

The most intriguing result of this computation is that the entire thermal metal phase can be thought of as emerging from the renormalization of the quantum critical point at \( J_d = 2\sqrt{2} \). This can be understood by a simple scaling argument: The leading term in the perturbation series for \( \Sigma_p(0, k_0) \) corresponding to scattering events from \( n \) impurities scales as \( (4pF)^n \). Thus, for small \( p \), we neglect the \( n > 1 \) terms to write \( J_{\text{eff}} \) as a linear function of \( p \). Within this “noncrossing” approximation, Eq. (6) has a single solution for a given \( p \), leading to a single gapless point for each disorder density which we can approximate using the self-consistent Born (SCBA) and T-matrix approximations (SCTA). On the other hand, for \( p \gtrsim 1/4 \), we can no longer ignore the higher-order diagrams, so that Eq. (6) has, in fact, an infinite number of solutions, accounting for an entire gapless region in the phase space, which is precisely the thermal metal phase.

Since the perturbative approach is no longer useful in the high disorder density regime, we complement it with a numerical computation of the transmission coefficient. In particular, we compute the disorder-averaged generalized transfer matrix [53] for the Majorana model on a cylinder geometry with \( L_y \gg L_x \). The eigenvalues of this transfer matrix are related to the inverse localization lengths \( \xi \), which can be used to compute the transmission coefficient as \( g = \sum_{i=1}^{2L_y} \sech^2(L_x/\xi_i) \). This can be thought of as the effective number of gapless channels, and is proportional to conductance for complex fermions [54].

This approach effectively captures the contributions from all \( n \)-impurity scattering events, and thus yields a reliable estimate of the phase boundary also for large disorder densities. In Fig. 11, we plot \( g \) as a function of \( J_d \) and \( p \), overlaid with the phase boundaries obtained from SCBA/SCTA. We see a qualitative agreement between the numerical and analytical computations in the low density regime \( p \ll 1/4 \), verifying the effective renormalization of \( J \). For \( p \gtrsim 1/4 \), we clearly see the spreading of the phase boundary into a gapless region extending up to \( J_d \approx 1.8 \), which is also in good agreement with the results obtained using QMC for the full Kitaev model (Fig. 8).

4. Physical interpretation

In closing our discussion of the thermal metal, we need to address the question whether this thermal metal is a legitimate physical phase of our spin model or a mere artefact of our calculations relying on a Majorana decomposition. The relevance of this question acutely presents itself in the observation that the putative thermal metal regime seems to possibly extend beyond the intermediate-temperature range all the way up to infinite temperatures, as possibly suggested by the calculation of the average Chern number plotted in Fig. 8.

The answer to this question is twofold. It sensitively depends on whether the Majorana fermions correspond to actual physical degrees of freedom of the system (after spin fractionalization) or whether they remain mere mathematical objects that can always be invoked in a spin decomposition. This fine distinction is well known from the solution of the honeycomb Kitaev model [6], whose intricacy lies precisely in the fact that the spin decomposition “becomes real” in the low-temperature phase and describes the emergent fractionalized degrees of freedom. It is also exactly this distinction which distinguishes the intermediate-temperature regime from the high-temperature paramagnet in our model. The intermediate-temperature regime is defined as the temperature regime sandwiched between the fractionalization crossover at \( T' \) and the low-temperature gauge ordering transition at \( T_c \), in which the emergent fractionalized degrees of freedom of itinerant Majorana fermions moving in the background of a static, but still disordered \( Z_2 \) gauge field, form. In this regime, our line of arguments outlined above fully applies and we conclude that in this temperature regime the thermal metal is a true physical phase.

The high-temperature paramagnet, on the other hand, is different. Here the Majorana decomposition of the spin operators is a mathematical possibility, but there is no deeper physical meaning associated with it (or any other spin decomposition using, e.g., complex fermions or other parton degrees of freedom). As such our numerical observation of a finite average Chern number and other class D physics in the Majorana representation is a direct reflection of the choice of decomposition, but not physically relevant.

We return to the question of how the thermal metal regime in the intermediate-temperature regime can be probed in the

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6We contrast this to a similar effect in topological insulators with Anderson disorder, where disorder leads to an enhancement of the topological phase. This difference is due to the purely imaginary disorder matrix in the present case, in contrast to a purely real one for topological insulators.
IV. SECOND-ORDER SPIN LIQUID

The physics of the Kitaev Shastry-Sutherland model becomes even richer when considering a *staggering* of the plaquette couplings, which has been shown [23] to induce another variant of spin liquid ground state. This “second-order” spin liquid (SOSL) exhibits a Majorana band structure in the ground state that, akin to a second-order topological insulator (SOSL) exhibits a Majorana band structure in the other variant of spin liquid ground state. This “second-order” symmetry-enriched are a manifestation of the formation of a second-order spin liquid. Specifically, in doing so, we will concentrate on a representative choice of the lattice (indicated in Fig. 12).

In the following, we will turn to the thermodynamics accompanying the formation of a symmetry-enriched topological order that is protected by two mirror symmetries of the lattice (indicated in Fig. 12).

In doing so, we will concentrate on a representative choice of coupling parameters deep in the SOSL regime. Specifically, we introduce the plaquette staggering $\delta J = 0.7$ (resulting in weakly and strongly coupled plaquettes with coupling strength $J - \delta J$ and $J + \delta J$, respectively) and vary the relative coupling of the diagonal versus plaquette bonds $J_{d}/J$. At zero temperature, the phase diagram again consists of two phases [23]: a chiral spin liquid for $J_{d}/J > 2\sqrt{2}$ and a second-order spin liquid phase for $J_{d}/J < 2\sqrt{2}$.

A. Thermodynamics

The principle thermodynamic signatures above the formation of this SOSL are similar to what we have seen for the more conventional spin liquids discussed in the previous section: The specific heat $C_v$ shows a three-peak structure (a). While the high-temperature crossover is associated with spin fractionalization and the low-$T$ phase transition with spontaneous breaking of time-reversal symmetry, the intermediate crossover indicates a partial flux-ordering of square plaquettes (b), which is a consequence of staggered bond couplings $J \pm \delta J$. This choice of coupling generates a hierarchy between the square plaquettes of the lattice, which results in different ordering temperature scales.

FIG. 12. Kitaev Shastry-Sutherland mode with staggered plaquette couplings. The dashed/solid lines correspond to staggered couplings $J \pm \delta J$, respectively, which generates a hierarchy among the square plaquettes: A quarter of the squares possess either only weak (green) or strong (blue) couplings on the edge bonds, while the remaining half of plaquettes has two weak and two strong bonds (cyan). The dotted gray lines denote the two mirror axes.

FIG. 13. Signatures of a three-step thermodynamic transition. Partial flux ordering in the SOSL phase (here, $J = 0.9$ and $\delta J = 0.4$). The specific heat $C_v$ shows a three-peak structure (a). While the high-temperature crossover is associated with spin fractionalization and the low-$T$ phase transition with spontaneous breaking of time-reversal symmetry, the intermediate crossover indicates a partial flux-ordering of square plaquettes (b), which is a consequence of staggered bond couplings $J \pm \delta J$. This choice of coupling generates a hierarchy between the square plaquettes of the lattice, which results in different ordering temperature scales.
range of our QMC simulations. In this limit, where the coupling \( J - \delta J \) on half of the lattice bonds approaches 0, it is expected that the transition temperature rapidly decreases to lower temperature scales [23].

### B. Partial flux ordering

The emergence of a partial flux ordering in an intermediate temperature range is again a precursor phenomenon for the formation of spin liquid ground states that, in the presence of staggered plaquette couplings, comes in even more variations – see Fig. 15. This is readily illustrated by measurements of the three- and four-plaquette fluxes, overlaid as color coding in the thermal phase diagrams of Fig. 14, which reveal a partially flux ordered regime in the parameter range \( J_\delta / J \lesssim 6.7 \) and above the thermal phase transition \( T'' > T > T_c \). However, in this case, the pattern of partial ordering is very different to the one previously discussed in Sec. III for the original model. While there, we observed a region in which the square plaquettes were fully ordered, and the triangular plaquettes remained disordered, here the partial flux order is characterized by partial order of the square plaquettes, with the triangular plaquettes again remaining disordered. This can be clearly seen in the thermal phase diagram with Fig. 14(a) showing the three-plaquette flux \( |W_3| = 0 \) and Fig. 14(b) showing the square-plaquette flux \( -0.75 \lesssim W_\Box \lesssim -0.25 \) within the partial flux ordered regime.

What is the reason for this new behavior? When looking at the model with staggered couplings, \( J \pm \delta J \), we see that the lattice is now composed of three different kinds of square plaquettes: (i) one quarter are “strong plaquettes,” which contain a diagonal \( J_\delta \) bond and four “strong” bonds with coupling \( J + \delta J \), (ii) one quarter are “weak” plaquettes, which contain a diagonal \( J_\delta \) bond and four “weak” bonds with coupling \( J - \delta J \), (iii) while the remaining half are “mixed” plaquettes, which do not contain any diagonal bond and are made up of two “strong” bonds and two “weak” bonds. This hierarchy of couplings, and thus vison gaps, for the three different kinds of square plaquettes is precisely what underlies the emergence of the partial flux ordering.

If we consider a lattice of \( N_p \) total square plaquettes, the behavior seen within the partially flux-ordered regime can be explained as follows. Starting from the lowest temperatures, for \( 0 < T < T_c \), all square plaquettes are in an ordered \( \pi \)-flux phase, with \( W_\Box = -1 \) for all \( N_p \) square plaquettes. At \( T_c \) the triangular plaquettes become disordered, triggering the recovery of time-reversal symmetry, and, also at \( T_c \), the \( N_p/4 \) “weak” square plaquettes similarly become disordered. This loss of \( N_p/4 \) plaquettes explains the drop of \( W_\Box \) from \(-1\) to \(-3/4\) at \( T_c \), shown in the lower panel of Fig. 13. Further increasing the temperature within the regime \( T_c < T < T'' \), the \( N_p/2 \) “mixed” plaquettes gradually disorder for higher temperatures, resulting in a smooth change of \( W_\Box \) from \(-3/4\) just above \( T_c \) to \(-1/4\) just below \( T'' \), see again the lower panel of Fig. 13. Finally, at \( T'' \), the remaining \( N_p/4 \) “strong” plaquettes disorder, resulting in a fully disordered flux state with \( W_\Box = 0 \) for all plaquettes for \( T > T'' \).

### C. Thermal Majorana metal

Turning to signatures of Majorana physics in the thermodynamic behavior, we find evidence for the formation of a thermal metal regime also for the staggered model, similar to what we discussed extensively for the original Kitaev Shashtry-Sutherland model in Sec. III C above. The numerical evidence
emergent Majoranas. Below the thermodynamic crossover where these fractionalized degrees of freedom (locally) come to live, the gauge physics manifests itself through a sequence of (partial) flux ordering transitions, with the transition into the ground-state manifold being accompanied by (global) time-reversal symmetry breaking. The latter is also mandated (already on a local level) by the Majorana physics for non-bipartite lattice geometries.

Another striking manifestation of the interplay of Majorana physics and the $\mathbb{Z}_2$ gauge structure is the formation of a thermal metal regime in an intermediate temperature range where the gauge field is intrinsically disordered. Notably, this gapless phase emerges only for a limited parameter regime of the Kitaev Shastry-Sutherland model, with a sharp transition to a gapped regime. This principle distinction might also make the thermal metal phase observable in experimental studies, e.g., in heat transport measurements which have been demonstrated, both theoretically [55–57] and experimentally [12,13,58,59], to be sensitive probes of the Majorana physics. The gapless versus gapped character of the intermediate-temperature phases should also reflect itself in four-spin correlation functions that probe the algebraic versus exponential decay of the bond-energy bond-energy correlations.

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APPENDIX A: QUANTUM MONTE CARLO SIMULATION

The quantum Monte Carlo (QMC) simulation method for Kitaev models was introduced in Ref. [24], with an extensive review provided in a recent PhD thesis [60]. While originally implemented for spin-1/2 Kitaev models, the approach can readily be generalized to the higher-order spin model considered here, as first discussed in Ref. [23]. The principal idea of the Monte Carlo scheme is to sample the (classical) $\mathbb{Z}_2$ gauge field configurations $\{u_{jk}\}$, while obtaining the statistical weight of each configuration via an exact diagonalization of the Majorana Hamiltonian (2). The latter is performed according to the canonical transformations introduced on Kitaev’s original honeycomb model [6]. Explicitly, the Majorana representation of the higher-order spin model allows to write the partition function of the system as a double trace

$$Z = \text{tr}_{\{|u_{jk}\}} \text{tr}_{\{c_l\}} e^{-\beta \mathcal{H}} = \text{tr}_{\{u_{jk}\}} e^{-\beta F(|u_{jk}|)} ,$$

where the free energy $F(|u_{jk}|)$ in a fixed $\mathbb{Z}_2$ gauge field configuration can be calculated from the eigenspectrum $\{\epsilon_{jk}\}$ of the Majorana Hamiltonian,

$$F(|u_{jk}|) = -T \sum_{\lambda} \ln \left(2 \cosh \left(\frac{\beta \epsilon_{jk}}{2}\right)\right) .$$

The latter step requires the explicit summation over all fermionic Fock states. Here, the distinction between physical

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**FIG. 16.** Thermal metal. The color coding shows the average Chern number $|\nu|$ across the thermal phase diagram of the staggered Kitaev Shastry-Sutherland model. For the low-temperature ground-state phases, the Chern number is $|\nu| = 1$, in the topological spin liquid phase and, as expected, $|\nu| = 0$ in the second-order spin liquid (SOSL). In the intermediate temperature regime, we find that the average Chern number vanishes in the partially flux-ordered phase in the temperature range $T_c < T < T^{\nu}$ for the respective range of couplings. Above $T^{\nu}$ (and $T_c$ for large $J_d$, respectively) the average Chern number is finite, but not quantized around $|\nu| \approx 0.8$. The black data points in indicate the respective transition temperatures of the system as a function of the coupling ratio $J_d/J$, extracted from specific heat traces akin to the ones shown in Fig. 13.

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V. SUMMARY

One of the most intriguing phenomena associated with the formation of quantum spin liquids is the emergence of novel, fractionalized quantum mechanical degrees of freedom—a quasiparticle, generally referred to as a parton, coupled to the gauge field of a deconfined lattice gauge theory. For the spin-3/2 Kitaev Shastry-Sutherland model studied in this manuscript, these emergent degrees of freedom are itinerant Majorana fermions and a $\mathbb{Z}_2$ lattice gauge field, with all spin liquids coming in the form of varying levels of topology—trivial, first-, and second-order, akin to the classification of higher-order topological insulators.

With a focus on the thermodynamic signatures of this fractionalization, we have observed in our numerically exact (sign-free) quantum Monte Carlo simulations characteristic fingerprints of both the underlying gauge physics and the
and unphysical Fock states can be neglected in the thermodynamic limit $N \to \infty$ ($N$ being the number of spins), since these states only differ in terms of their fermionic parity [61]. Thus, with the argument first presented in Ref. [62], unphysical states only contribute deviations of the order $1/N$.

APPENDIX B: AVERAGE CHERN NUMBER

To characterize the Majorana band structures in the intermediate-temperature regime, we have numerically calculated an average Chern number $\langle |v| \rangle$ in our Monte Carlo simulations. A single measurement contributing to this average is done for a given $\mathbb{Z}_2$ gauge field configuration (which effectively acts as a disorder potential for the Majorana fermions in this intermediate temperature regime). The calculation is performed according to the method for non-Abelian Berry connections, which was introduced in Ref. [63]. Here, the real-space $\mathbb{Z}_2$ gauge field configuration $\{u_{ij}\}$ of the size $L = 10$ ($N = 400$ sites) is used for the construction of a supercell in momentum space, described by the Hamiltonian $\mathcal{H}_k(\{u_{ij}\})$.

The Brillouin zone of the system is discretized with a mesh of $10^2$ $k$ points. For each pair of nearest neighbor $k$ points $(k_i, k_i + q_j)$, the U(N) gauge variable

$$U_{q_i} = \frac{\det(\psi^\dagger(k_i)\psi(k_i + q_j))}{\det(\psi^\dagger(k_i)\psi(k_i + q_j))}$$

is calculated, where $\psi$ is the $N/2 \times N/2$ matrix that is composed of the multiplet $|\mu_1(k)\rangle \ldots |\mu_{N/2}(k)\rangle$ of eigenstates, belonging to the lower half of eigenvalues $\epsilon_1, \ldots, \epsilon_{N/2}$ of $\mathcal{H}_k$. Here, the relaxed gap opening condition $\epsilon_m \neq \epsilon_n$ for $n \leq N/2$ and $m > N/2$ has to be fulfilled. From the U(N) gauge variables $U_{q_i}$, we calculate a (gauge-invariant) plaquette field strength

$$F_{12}(k_i) = \log \left( \frac{U_{q_1}(k_i)U_{q_2}(k_i + q_1)}{U_{q_1}(k_i)U_{q_2}(k_i)} \right),$$

which is defined on the square plaquettes of the discretized Brillouin zone, and for which

$$-\pi < \frac{1}{i} F_{12}(k_i) \leq \pi.$$ 

The (numerical) Chern number of the band structure can now be expressed by the sum over all plaquette field strengths $F_{12}(k_i)$ of the lattice,

$$\tilde{\nu} = \frac{1}{2\pi i} \sum_i F_{12}(k_i),$$

such that $\tilde{\nu} \in \mathbb{Z}$ and $\tilde{\nu} \to \nu$ for a sufficiently fine discretization of the Brillouin zone. Note that the condition $\tilde{\nu} \in \mathbb{Z}$ follows from the way the plaquette field strength (B2) is constructed, but the integer value only has a physical meaning if the band structure is gapped. While this is the case for the ground state phases of the Shastry-Sutherland Kitaev system and the high-temperature phase at $J_d > 0$ (Fig. 8), the different integer results for the gapless high-temperature phase at $J_d < 0$ are nonphysical, and a mere “breakdown” indicator for the gapless phase—see the Chern number histograms in Fig. 17 and the band structures in Fig. 18. The same is true for the high-temperature results shown in Fig. 16.

Detailed scans of the average Chern number calculations are shown in Figs. 19 and 20.

FIG. 17. Chern number histograms for two different temperatures at the parameter point $J_d = 1.5$ (in phase diagram 8). In the intermediate-temperature regime, the occurrence of integer Chern number results is an artifact, since here, the system is in a gapless phase.

FIG. 18. Band structures near $E = 0$ for different parameters values $J_d$ at $T = \infty$ (in phase diagram 8), showing the transition from the gapless to the gapped high-temperature phase.
APPENDIX C: DISORDER-INDUCED RENORMALIZATION

The Shastry-Sutherland lattice is described by a four-site unit cell, periodic with translation vectors \( a_1 = \hat{x} \) and \( a_2 = \hat{y} \). Thus the Shastry-Sutherland-Kitaev model, after a Majorana decomposition, can be written in the momentum space as

\[
\mathcal{H}_B(k) = \begin{pmatrix} 0 & J' \gamma & -J \sigma_x e^{-ik_x} & J' \gamma \\ -J' \gamma & 0 & J \sigma_x e^{ik_x} & -J' \gamma \\ J \sigma_x e^{-ik_x} & -J \sigma_x e^{ik_x} & 0 & J' \gamma \\ J' \gamma & J' \gamma & -J' \gamma & 0 \end{pmatrix},
\]  

(C1)

where we set \( J_x = J_y = J + \delta J \), \( J'_x = J'_y = J - \delta J \) and \( J_z = J_d \). In the following, we restrict to the case of \( \delta J = 0 \).

1. Vison excitations as additive disorder

The Bloch Hamiltonian of Eq. (C1) corresponds to the ground state flux sector with \( \pi \)-flux through each 4-plaquette. Flux-disordered configurations, corresponding to one of the four inequivalent disorder configurations is obtained by flipping each \( J \) bonds, which can be taken as the four intracell hoppings. We think of these vison excitations as an additive disorder and write it schematically as \( \mathcal{H} = \mathcal{H}_0 + \sum_{\alpha} U^{(\alpha)}(r) \), where \( U^{(\alpha)}(r) \) is an impurity potential matrix at position \( r \) and \( \alpha \) indicates the matrix structure corresponding to one of the four inequivalent vison excitations. For instance, such a matrix for flipping the 1–2 bond can be written as

\[
X^{12} = 2iJ \begin{pmatrix} 0 & -1 & 0 & 0 \\ 1 & 0 & 0 & 0 \\ 0 & 0 & 0 & 0 \\ 0 & 0 & 0 & 0 \end{pmatrix},
\]  

(C2)

and \( X^{23} \), \( X^{34} \), and \( X^{41} \) take a similar form, which can be compactly written as \( [X^{ab}]_{ij} = 2iJ(\delta_\alpha \delta_\beta - \delta_\alpha \delta_\beta) \). A typical disorder configuration is obtained by flipping each \( J \) bond with probability \( p \). For small \( p \), the disorder corresponds to a set of localized scatterers, which can be analyzed via perturbation theory.

The average effect of disorder is encoded by the self-energy matrix \( \Sigma_p(\omega, k) \), that appears in the definition of the disorder-averaged Green’s function:

\[
\mathcal{G}(\omega, k) \equiv \langle \langle \omega - \mathcal{H}(k) \rangle^{-1} \rangle_{\text{dis}} = [\omega - \mathcal{H}_0(k) - \Sigma_p(\omega, k)]^{-1},
\]  

(C3)

where \( \langle . \rangle_{\text{dis}} \) represents averaging over the position and bond direction of the disorder. In the following, we are only interested in the self-energy at \( \omega = 0 \). Owing to the particle-hole symmetry, the entries in the self-energy—as in the real-space Hamiltonian—must be purely imaginary. We can further identify its matrix structure by noting that the perturbation series above consists of terms of the form \( \mathcal{H}_0^{-1}(k)X^a \mathcal{H}_0^{-1}(k) \), where \( X^a \) is one of the disorder matrices defined above. Given the form of \( \mathcal{H}_2 \), such a product has only two nonzero components at the same position as \( X^a \), \( \mathcal{v} \), at indices \( ab \) and \( ba \). Since each type of disorder is equally likely, the disorder-averaged self-energy takes the form

\[
\Sigma_p(0, k) = i\Sigma_p(0, k) \begin{pmatrix} 0 & -1 & 0 & -1 \\ 1 & 0 & -1 & 0 \\ 0 & 1 & 0 & -1 \\ 1 & 0 & 1 & 0 \end{pmatrix},
\]  

(C4)

FIG. 19. Average chern number \( |\langle \nu \rangle| \) for different horizontal (constant temperature) cuts in phase diagram 8.

FIG. 20. Average chern number \( |\langle \nu \rangle| \) for different vertical (constant \( J_d \)) cuts in phase diagram 8.
where $\Sigma_p(0, k) \in \mathbb{R}$. Since the coefficient (matrix) of $J$ in the Bloch Hamiltonian takes precisely this form for any $k_z = k_w$, the effect of disorder is essentially a renormalization of $J$. In particular, at the quantum critical point $J_d = 2\sqrt{2}J_0$, the spectrum consists of a single Dirac point at $k_0 = (\pi, \pi)$, so that we write $J \rightarrow J_{\text{eff}} = J - \Sigma_p(0, k_0)$. This can also be interpreted as renormalizing the mass of this Dirac fermion, and the phase boundary corresponds to the new $J_d$ for which this mass vanishes.

To compute the disorder averaged self-energy, we expand it in powers of $p$ as

$$
\Sigma_p(\omega, k) = p \left( \begin{array}{c} \xi \\ \eta \\ \eta^* \end{array} \right) + p^2 T^{(2)} + O(p^3),
$$

(C5)

where $T^{(n)}$ has the same matrix structure as $\Sigma_p$ (Eq. (C4)) with coefficient $iT^{(n)}$. Physically, this corresponds to an expansion in the number of scattering events [64]. Note that we only have terms with an even number of scattering events, since those with an odd number cancel after disorder averaging. The self-consistent version of Eq. (C5) is obtained by simply replacing the bare Green’s functions with the full disorder-averaged Green’s functions [64].

2. The Born and T-matrix approximations

At low disorder densities, we can truncate the perturbation series of Eq. (C5) using the T-matrix approximation to compute the disorder-averaged self-energy at linear order in $p$, i.e., only including terms corresponding to a single impurity scattering, so that $\Sigma_p(0, k) \approx p T^{(1)}$. This can be further simplified by taking the Born approximation to the full T-matrix $T^{(1)} \approx T^{(1)}_{\text{Born}}$, wherein we retain only the first Feynman diagram of Eq. (C5). The self-consistent versions of these approximations can be calculated by replacing the bare Green’s function in Eq. (C5) with their disorder-averaged counterparts (which contains $\Sigma_p(\text{iso}, k)$). Setting $J = 1$ throughout this subsection, the renormalization of the coupling constant $J$ for all these approximations take the form

$$
J_{\text{eff}}(p) = 1 - p T^{(1)}, \quad T^{(1)} > 0.
$$

(C6)

The system is now gapless when $J_d = 2\sqrt{2}J_{\text{eff}}(p)$, which yields the phase boundary on the $J_d$–$p$ plane. The net effect of the disorder is a shift of the phase boundary towards smaller $J_d$.

The validity of the approximations discussed above is, at first sight, unclear. This is because our system is gapped and so the usual small parameter for a Fermi sea, $1/(k_F \ell)$—with $\ell$ being the mean free path—is not well defined. Nonetheless, the results for the phase boundary obtained using these approximations agree qualitatively with the numerical results obtained from a transfer matrix calculation, as shown in Fig. 11. Furthermore, they also suffice to explain the origin of a “thermal metal” phase (Fig. 8) and predict its location in the phase diagram.

The basic mechanism for the appearance of the thermal metal is the breakdown of the T-matrix approximation as $p$ is increased and the interference effects from by multiple-impurity scattering become relevant. More precisely, since the disorder strength (i.e., the prefactor in $X_{\text{iso}}$) is 2 and each impurity must have an even number of scattering events, the leading order term for an $n$-impurity scattering scales as $(4p)^n$. These terms can be neglected only when $4p \ll 1$. However, as the disorder density increases to $p \sim 1/4$, the diagrams at all orders in Eq. (C5) become relevant, so that the coefficients of $p^n$ in $\Sigma_p(0, k_0)$ must be taken into account for all $n$. This means that for each value of $J_d$, the equation $J_d = 2\sqrt{2}J_{\text{eff}}(p)$ can have an infinite number of solutions for $p$. Thus, once this density is reached, it is possible for the phase boundary to become a region on the $J_d$–$p$ of the phase diagram and a disorder induced metal can be formed.

Using the various approximations to the T-matrix, we can estimate the coupling constant for which the thermal metal appears by computing the value of $J_{\text{eff}}(p)$ for which the phase boundary hits the line $p = 1/4$, i.e., as $J_d = 2\sqrt{2}(1 - T^{(1)})/4$. It turns out that $T^{(1)}_{\text{Born}}$ can be computed analytically for $J_d = 2$, since the corresponding integral can be evaluated using methods for rational trigonometric functions. In fact, for this special value of the coupling constant, we find that

$$
T^{(1)}_{\text{Born}} = 8 \int \frac{d^2k}{(2\pi)^2} G_0(0, k) = 4 - 2\sqrt{2},
$$

(C7)

which exactly satisfies the condition for the phase boundary obtained above. From the Born approximation, we therefore obtain an entirely analytic estimate of the critical value of $J_d$ where the thermal metal emerges, which is already fairly close to the numerically observed value (see Fig. 11). Better approximations can be obtained by including more diagrams in the perturbative expansion of Eq. (C5) and/or by using the self-consistent version of this expansion. For example, the self-consistent T-matrix estimates the position of the transition as $J_d \approx 1.8$, which is comparable to that found by the transfer matrix technique (Fig. 11) and quantum Monte Carlo (Fig. 8).

APPENDIX D: VISION GAPS

To complement the discussion of the phase diagram in Fig. 3 of the main text we here report on the relation between the transition temperatures and the vision gaps of the system (summarized in Fig. 21). Corresponding to the two sets of elementary plaquettes in the Shastry-Sutherland lattice, we
can distinguish two kinds of vison excitations. A pair of visons on the triangle plaquettes is created by flipping a diagonal $J_d$ bond, whereas flipping a horizontal or vertical bond generates a pair of visons on the square plaquettes. Figure 21(a) shows the values of the critical temperature $T_c$ as a function of the triangle-plaquette vison gap $\Delta_5$. Figure 21(b) shows the values of the square-ordering transition temperature, which is $T''$ for $J_d/J < 0.9$ and $T_c$ for $J_d/J \geq 0.9$, as a function of the square-plaquette vison gap $\Delta_5$.

We can determine a pronounced linear correlation between $T_c$ and $\Delta_5$ for a wide range of values $\Delta_5 \leq 0.5$, which corresponds to coupling parameter values $J_d/J > 1.5$ (orange and blue data points). For the partial-flux order limit $J_d/J < 1$, there is apparently also a linear correlation between the square-ordering temperature $T''$ and $\Delta_5$ with a different (negative) slope (red data points). This implies that a larger vison gap corresponds to a lower transition temperature $T''$ in this limit. For $1 \leq J_d/J \leq 1.5$, where $\Delta_5$ is the largest, the data points are too close to each other to determine a functional relationship. It is here that the $T''-\Delta_5$ curve "U turns" after the gap $\Delta_5$ reaches its largest value.

For the 3-plaquettes, we see a linear correlation between $T_c$ and $\Delta_5$ for the trivial phase $J_d/J > 2.8$ and parts of the chiral phase $J_d/J \geq 2.3$, where $\Delta_5$ has its maximum value. The pronounced "U turn" of the $T_c-\Delta_5$ curve thereby corresponds to moving $J_d$ to lower values. For $J_d/J \leq 1$, there is again a linear correlation between both quantities.

We can state that the sections with a linear correlation between the transition temperature and the vison gap are consistent with results from 3D Kitaev systems [40]. However, the "U turn" behavior, which is witnessed for both the $T_c-\Delta_5$ and the $T''-\Delta_5$ curves, suggests that the relation between transition temperature and gap is not a simple, global linear function $T(\Delta) = m\Delta$. Instead, the slope $m$ is changed in different parameter regions, whereas in the region of extremal $\Delta_5$ values, there is no linear correlation at all. Nonetheless, it can be stated that a general correlation between both quantities is verified by these results.
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