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Z₃ topological order in the quantum dimer-pentamer model

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We introduce the quantum dimer-pentamer model (QDPM) on the square lattice. This model is a generalization of the square lattice quantum dimer model as its configuration space comprises fully-packed hard-core dimer coverings as well as dimer configurations containing pentamers, where four dimers touch a vertex. Thus in the QDPM, the fully-packed, hard-core constraint of the quantum dimer model is relaxed such that the local dimer number at each vertex is fixed modulo 3, resulting in an exact local $Z_3$ gauge symmetry. We construct a local Hamiltonian for which the Rokhsar-Kivelson (RK) equal superposition state is the exact ground state and has a 9-fold topological degeneracy on the torus. Using Monte Carlo calculations, we find no spontaneous symmetry breaking in the RK wavefunction and that its dimer-dimer correlation function decays exponentially. By doping the QDPM RK state with a pair of monomers, we demonstrate that $Z_3$ electric charges are deconfined. Additionally, we introduce a $Z_3$ magnetic string operator that we find decays exponentially and shows no signatures of magnetic vortex condensation and with correlations. These numerical results suggest that the ground state of the QDPM is a dimer liquid with $Z_3$ topological order.

I. INTRODUCTION

Strongly interacting quantum many-body systems are known to support a variety of exotic quantum phases of matter with no classical analogue. Among these are topologically ordered quantum liquids; their non-trivial topological order can not be identified by a local order parameter as these phases of matter don’t break conventional symmetries¹,². Both because of the exotic nature of topological phases and their potential use for quantum information processing³,⁴, there is great interest in the development of theoretical models which display these phases and could provide candidate experimental systems to constructively engineer topological order⁵–¹¹.

A variety of exactly soluble lattice models are known to support ground states with topological order that ranges from the simplest variety, $Z_2$ topological order⁴,¹², to those with non-Abelian anyonic excitations¹³,¹⁴; such models generally involve complex many-body interactions, and thus it is desirable to find simpler models supporting topological order that are closer to what is accessible experimentally. Among the simplest classes of models displaying topological quantum liquid phases are those with local geometric constraints; the constraints inhibit the formation of symmetry breaking local order and given sufficient dynamics, may lead to a symmetric liquid ground state. The quantum dimer model (QDM) describing dimer degrees of freedom living on the links of a lattice is one of the most basic locally constrained models¹⁵,¹⁶; the hard-core, fully-packed QDM has a local constraint that requires that exactly one dimer touches each vertex.

While on non-bipartite two-dimensional lattices, the QDM has a topologically ordered ground state¹⁷,¹⁸, on the square lattice there is only a symmetric liquid ground state at an isolated critical quantum point¹⁹,²⁰. If the hard-core constraint of the square lattice QDM is relaxed such that only the parity of the number of dimers touching each vertex is fixed, the simplest non-trivial dynamics leads to a $Z_2$ topologically ordered ground state⁴,¹². The existence of topologically ordered ground states in these models is intimately related to the local constraint which imposes a local gauge symmetry in the Hilbert space; the topological order that arises is directly related to a discrete gauge theory with the appropriate local gauge constraint²¹.

Given that these models with dimer degrees of freedom and local constraints posses the simplest non-trivial local Hilbert space at each link, they provide a potentially simpler route to engineering exotic phases. The local constraints can be enforced by a local potential energy penalty and the minimal dynamics often arise at the lowest non-trivial order in perturbation theory from a related spin model²². We will refer to such models with dimer degrees of freedom on links and local constraints as gen-
eralized quantum dimer modes (GQDMs). Previous work has demonstrated that GQDMs on the square lattice support gapped $Z_2$ topologically ordered phases as well as gapless quantum critical points. On non-bipartite lattices, GQDMs support $Z_2$ topological order\cite{1,2} as well as doubled-semion phases\cite{24,25}. Additionally, recent work on a trimer resonating valence bond state suggests that it may be described by a $Z_3$ gauge theory\cite{26}. A full characterization of all exotic phases that can exist within this framework is thus desirable.

In this work we extend this paradigm to a new GQDM on the square lattice, what we term the quantum dimer-pentamer model (QDPM). In the QDPM, the local constraint is relaxed from that of the QDM to allow either 1 or 4 dimers to touch each vertex, as shown in figure Fig. 1. Correspondingly, the QDPM extends the Hilbert space of the QDM to include both fully-packed hard-core dimer configurations, as well as those with pentamers—four dimers touching a vertex. We define a local Hamiltonian that provides the simplest non-trivial dynamics that preserve this local constraint. Consequently, the Hamiltonian has an exact local $Z_3$ symmetry and conserves a $Z_3$ topological winding number; thus the QDPM has a 9-fold topological ground state degeneracy on the torus.

Additionally, we present a numerical study of the ground state of the QDPM at an exactly soluble point. Using Monte Carlo calculations, we explicitly demonstrate the absence of symmetry breaking order and the exponential decay of correlation functions. Additionally, we define a $Z_3$ magnetic string operator, whose correlations display no evidence of magnetic vortex condensation. Finally, by studying systems doped with a pair of monomer defects, we demonstrate that monomers are deconfined in the QDPM. Thus we argue that the ground state of the QDPM is a $Z_3$ topologically ordered dimer liquid.

This paper is organized as follows. In Sec. II we provide relevant background for the QDM and related models. In Sec. III we present the details of the QDPM. Sec. IV provides the results of a Monte Carlo study of the ground state of the QDPM at its exactly soluble point. Finally, Sec. V discusses potential future studies of the QDPM.

II. BACKGROUND

A. Generalized quantum dimer models

Here we will provide background about generalized quantum dimer models on the square lattice. We consider systems with Ising-like dimer degrees of freedom on each link, such that each link is either unoccupied or occupied by a single dimer. Given an Ising spin $\sigma_\ell$ on each link $\ell$ we define the number of dimers on the link $n_\ell \equiv 1/2(1 + \sigma_\ell^z)$, where $\sigma_\ell^z$ are the associated Pauli matrices. The Hilbert space is spanned by an orthonormal basis of dimerizations of the square lattice that satisfy a local constraint at each vertex. We define a $U(1)$ charge $Q_\nu$ at each vertex $\nu$ of the lattice in terms of the number of dimers touching $\nu, n_\nu \equiv \sum_{\ell \in \nu} n_\ell$:

$$Q_\nu \equiv \pm (n_\nu - c_p)$$

where $c_p$ is a constant, and we define the even (odd) sector of the constraint to have $c_p = 2$ ($c_p = 1$); the $+(-)$ sign corresponds to vertices on the $A(B)$ sublattice (see Tab. I). A local dimer number constraint requires $Q_\nu$ to vanish for all vertices; a dimer parity constraint requires $(Q_\nu \mod 2)$ to vanish (see Fig. 2).

The Hamiltonian of a GQDM is of the form

$$H_{\text{GQDM}} = - \sum_\nu t_\nu T_\nu + \sum_\nu v_\nu V_\nu$$

(1)

where $T_\nu$ are off-diagonal kinetic energy operators, $V_\nu$ are diagonal potential energy operators, $t_\nu$ and $v_\nu$ parametrize the strength of these terms, and each $\nu$ represents a set of neighboring links. We will limit our discussion to $t_\nu > 0$. $T_\nu$ is constructed to generate resonances between two distinct allowed dimerizations of $\nu$, $c_\nu$ and its compliment $\bar{c}_\nu$, and is written as:

$$T_\nu = |c_\nu\rangle \langle \bar{c}_\nu| + h.c.$$  

in the simplest case, $T_\nu$ involves the minimum non-trivial dimer dynamics that preserve the constraint. Similarly, the potential energy term provides an energy cost for states with these local configurations:

$$V_\nu = |c_\nu\rangle \langle c_\nu| + |\bar{c}_\nu\rangle \langle \bar{c}_\nu|.$$  

Therefore we may define a Hamiltonian for a GQDM using Eq. (1) and defining a set of dimer resonances
define an oriented dimer flux of the links of the square lattice such that they point from one entangled path to another. The ground state is gauge invariant and magnetic string operators generate defects and computing their spatial correlation function; these magnetic vortices should be gapped excitations. The condensation of magnetic vortices can drive a phase transition from a topologically ordered phase to a conventionally ordered phase, where the magnetic string correlation can be used as a test for deconfinement of electric charges. On surfaces with non-trivial topology, we may define the topological loop operators

\[ W_X (\alpha) \equiv S_{\Gamma X} (\alpha) \]

where \( \Gamma_X \) is a topologically non-trivial closed loop that winds once about the \( x \) axis; we define \( W_Y \) similarly. Since \( W_X (\alpha) \) and \( W_Y (\alpha) \) commute with the Hamiltonian, we may use their eigenvalues to label the ground states.

B. Fixed dimer number constraints

The two unique fixed-number constraints on the square lattice lead to the QDM in the odd sector (\( n_v = 1 \)) and the quantum full-packed loop model (QFPLM) in the even sector (\( n_v = 2 \)). For configurations that satisfy these number constraints, \( Q_v \) strictly vanishes, and thus all states in the Hilbert space are invariant under \( G_v \) for \( 0 \leq \alpha_v \leq 2\pi \) in Eq. (3); this represents a local \( U(1) \) gauge symmetry. The Hamiltonian for these models gives dynamics to
parallel dimers on the plaquettes of the lattice:

\[ H_{QDM}(t, v) = H_{QFPLM} \left( \{(t, v)\} \right) \]  

where we have chosen \( t_c = t, v_c = v, \forall \mathcal{D} \). In this case of a fixed number constraint, the Hamiltonian conserves \( Q_v \) and therefore the dimer flux through any closed loop such that:

\[ [\Phi_\Gamma, H_{QDM}] = 0. \]

We can define two conserved topological winding numbers \( \Phi_X \) and \( \Phi_Y \), which are the dimer flux through topologically non-trivial loops that wind around each axis of the lattice when the system has periodic boundary conditions (see Fig. 3). On a lattice of dimensions \( L_x \times L_y \), the eigenvalues of \( \Phi_X \) and \( \Phi_Y \) are integers in the range

\[ -L_x/2 \leq \Phi_X \leq L_x/2, \]

and similarly for \( \Phi_Y \), thus there are \( O(L_xL_y) \) distinct topological sectors.

At the Rokhsar-Kivelson point \( t = v \), we may write \( H_{QDM} \) in the form of Eq. (2) by defining projection operators \( h_v \):

\[ h_v \equiv \left( \frac{|c_v\rangle - |\bar{c}_v\rangle}{\sqrt{2}} \right) \left( \frac{\langle c_v| - \langle \bar{c}_v|}{\sqrt{2}} \right). \]

Since the plaquette flip term in Eq. (7) is believed to be ergodic in each of topological sector \( (\Phi_X, \Phi_Y) \) on the square lattice, each topological sector corresponds to a unique RK ground state where \( \Omega \) is labeled by \((\Phi_X, \Phi_Y)\).

Thus the RK points of the QDM and QFPLM have an extensively degenerate ground states.

Defects in these models violate the local dimer number constraint and thus carry non-zero U(1) electric charge. Tab. I lists all the possible defects and their associated U(1) charges in QDM and QFPLM. When a defect lives on sublattice A, it has the opposite charge that the same defect on sublattice B has. Note that in the QDM, the maximum charge is \( \pm3 \).

The QDM and QFPLM on the square lattice have isolated liquid ground states at the RK point surrounded by symmetry broken phases.\(^{15,16,19,20,33,34}\) The RK point of these models have been shown to have power-law decaying dimer-dimer and monomer-monomer correlation functions and gapless excitations.\(^{19,35–49}\) The algebraic decay of monomer correlations is indicative of the logarithmic confinement of electric charges in these models. Thus, such fixed number constrained models on bipartite lattices are expected to have an isolated unstable gapless critical quantum liquid points with logarithmically confined defect and an extensive topological degeneracy. These features of fixed number constrained models are consistent with the effective U(1) gauge theory description of the RK point;\(^{21,50,51}\) in 2+1D pure U(1) gauge theories are confining and thus the lack of an extended deconfined topologically ordered phase near the RK point.\(^{52}\)

### C. Fixed dimer parity constraints

We now relax the local constraint to a parity constraint such that only \( Q_{v}^{2} \equiv (Q_v \mod 2) \) is required to vanish; the resulting GQDM are the even and odd sectors are the even & odd Ising gauge theories (IGT) with constraints \( n_v \in \{0, 2, 4\} \) and \( n_v \in \{1, 3\} \), respectively. In these cases, the local constraint requires \( \alpha_v \in \{0, \pi\} \) in Eq. (3) and therefore the gauge symmetry is reduced from \( U(1) \) to \( Z_2 \). The set of allowed dimer resonances \( \mathcal{C}_{IGT} \) is

| type   | \( n_v \) | configuration | \( Q_{v}^{22} \) | \( Q_{v}^{23} \) |
|--------|----------|--------------|----------------|----------------|
| monomer | 0        | –            | -1             | 1              |
| dimer   | 1        | –            | 0              | 0              |
| trimer  | 2        | –            | 1              | 1              |
| tetramer| 3        | –            | 2              | 0              |
| pentamer| 4        | –            | 3              | 1              |

TABLE I. Local dimer configurations where \( n_v \) dimers touch a vertex; only one representative of symmetry related configurations is shown. \( Q_v, Q_v^{22}, \) and the \( Q_v^{23} \) are the associated \( U(1), Z_2 \) and \( Z_3 \) charges on the A sub-lattice in the odd sector theories (QDM, odd IGT, QDPM). Defects on the B sub-lattice have opposite charge, with the exception of \( Q_v^{22}, \) as there is only one type of non-trivial \( Z_2 \) charge.
expanded accordingly:

\[ C_{\text{IGT}} \equiv \{(c_{\square,i}, \bar{c}_{\square,i})\} \]

\[ = \{(\mathbf{r}, \mathbf{z}), (\mathbf{z}, \mathbf{r}), (\mathbf{u}, \mathbf{n}), (\mathbf{n}, \mathbf{u})\} \]

where we implicitly include symmetry related resonances. We will limit our discussion to \( v_1 = 0 \) such that the corresponding Hamiltonian is

\[ H_{\text{IGT}}(t) \equiv H_{\text{GQDM}}(C_{\text{IGT}}, \{(t, 0)\}) \]

In this case the equal superposition RK wavefunction is the exact ground state and occurs in the absence of a potential energy term, as we can define a projection operator on each plaquette

\[ h_{\square} \equiv \frac{1}{2}(1 - \sum_i |c_{\square,i} \rangle \langle c_{\square,i}| + \text{h.c.}) \]

such that \( H_{\text{IGT}} \) takes the form of Eq. (2) (up to a constant), and the RK state is the ground state for all values of \( t \).

The additional dynamics of \( H_{\text{IGT}} \) vs. \( H_{\text{GQDM}} \) no longer conserve the dimer flux \( \Phi_\Gamma \) through a closed curve; only \( \Phi_\Gamma \mod 2 \) is preserved by \( H_{\text{IGT}} \). Consequently, to define the string and loop operators that commute with \( H_{\text{IGT}} \) (for closed strings) we fix \( \alpha = \pi \) in Eq. (6):

\[ S_{\gamma}^{Z_3} \equiv S_{\gamma}(\pi), \quad W_{X}^{Z_3} \equiv W_{X}(\pi), \]

and \( W_{Y}^{Z_3} \) is defined similarly. Thus the \( W_{X}^{Z_3} \) and \( W_{Y}^{Z_3} \) have two eigenvalues \( \pm 1 \), and the topological sectors on the torus can be labeled as \( \Omega = (\pm 1, \pm 1) \); consequently the extensive degeneracy of the fixed number constrained models has been reduced to a finite ground state degeneracy in the fixed parity models.

Defects violating the dimer parity constraint in the IGT’s have \( Q_{\gamma}^{Z_3} = 1 \), as listed in Tab. I; we identify these defects as \( Z_2 \) electric charges on the corresponding vertices. Note that there is only one non-trivial charge in \( Z_2 \) gauge theory, thus there is no distinction between the charges of defects living on the two substraticles.

The ground states of \( H_{\text{IGT}} \) are fourfold degenerate on the torus, and have exponentially decaying dimer-dimer and monomer-monomer correlations and a finite energy gap; such features demonstrate the deconfined topologically ordered nature of the ground state of the IGTs.

We may understand the emergence of the \( Z_2 \) topologically ordered phase of IGTs from the QDM and QF-PLM from the following picture. The additional vertex configurations that are allowed in the IGTs relative to the models with a dimer number constraint have \( U(1) \) charge \( Q_v = \pm 2 \) (see Tab. I). It is known that when a \( U(1) \) gauge field is coupled to a dynamic charge \( q \) matter field, this results in a \( Z_q \) gauge theory when the matter field condenses\(^3\); as in the IGT’s we have added dynamic charge \( 2 \) defects to the underlying \( U(1) \) symmetric model, and we may expect a \( Z_2 \) gauge theory to result. We note that pentamers are charge \( \pm 3 \) objects in the QDM (see Tab. I); thus below we investigate whether introducing pentamers into the QDM leads to an effective \( Z_3 \) gauge theory, as suggested in Ref. 54 (see Appendix A for a review of \( Z_3 \) gauge theory).

### III. THE QUANTUM DIMER PENTAMER MODEL

We define the Hilbert space of the QDPM to be a GQDM where vertices are constrained to be touched by a single dimer or the center of a pentamer, such that \( n_v \in \{1, 4\} \). For defect-free configurations in the QDPM, \( Q_v \) only vanishes modulo 3; thus we may define a \( Z_3 \) charge,

\[ Q_v^{Z_3} \equiv (Q_v \mod 3) \]

which strictly vanishes at each vertex, representing the local constraint. All states in the QDPM are invariant under an exact local \( Z_3 \) gauge symmetry given by Eq. (3) with \( \alpha_v \in \{0, \pm 2\pi/3\} \). Thus introducing pentamers in the QDM reduces the gauge symmetry from \( U(1) \) to \( Z_3 \).

The Hamiltonian of the QDPM is defined by Eq. (1) including both plaquette flips as well as local dimer resonations that create and destroy pairs of pentamers, and allow single pentamers to translate on the same sublattice; these local dynamics involving pentamers are illustrated in Tab. II. Note that pentamers give dynamics to “staggered” regions with no flippable plaquettes that are frozen in the QDM (see Fig. 4 for an illustration of this); thus we expect the QDPM to have weaker correlations than the QDM, where the dynamics are more constrained. We may write \( H_{\text{QDPM}} \) as the following:

\[ H_{\text{QDPM}} \equiv H_{\text{GQDM}} + H_{\text{pen}} \]

where \( H_{\text{pen}} \) only involves the pentamer resonances in Tab. II and those related by symmetry:

\[ H_{\text{pen}} \equiv H_{\text{GQDM}}\left(\{(c_{v_1}, \bar{c}_{v_1}), (c_{v_2}, \bar{c}_{v_2}), (c_{v_3}, \bar{c}_{v_3})\},\right.\]

\[ \left.\{(t_1, v_1),(t_2, v_2),(t_3, v_3)\}\right) \]

Because of the terms involving pentamers, \( H_{\text{QDPM}} \) conserves \( Q_v^{Z_3} \) rather than \( Q_v \), and thus \( \Phi_\Gamma \) does not commute with the Hamiltonian for closed loops \( \Gamma \). The appropriate string and loop operator corresponds to Eq. (6) with \( \alpha = 2\pi/3 \):

\[ S_{\gamma}^{Z_3} \equiv S_{\gamma}(2\pi/3), \quad W_{X}^{Z_3} \equiv W_{X}(2\pi/3) \]

and similarly for \( W_{Y}^{Z_3} \). With these definitions, closed string operators \( S_{\Gamma}^{Z_3} \) and the topological loop operators...
TABLE II. The pairs of local dimer & pentamer configurations $c_\nu \leftrightarrow \bar{c}_\nu$ generate minimal local dynamics that create/annihilate pairs $(\nu_1)$ of pentamers and translate single pentamers $(\nu_2 \& \nu_3)$. Note that pentamers always stay on the same sublattice which is associated with their $U(1)$ charge.

| $\nu$  | $c_\nu$  | $\bar{c}_\nu$ |
|--------|--------|-----------|
| $\nu_1$ | $+$    | $-$       |
| $\nu_2$ | $+$    | $-$       |
| $\nu_3$ | $+$    | $-$       |

W$^{Z_3}_X$ and W$^{Z_3}_Y$ commute with $H_{QDPM}$.

With the exception isolated "frozen" configurations (e.g. pentamers occupying all vertices), our numerical simulations (discussed below in Sec. IV) suggest that the dynamics of the $H_{QDPM}$ access a substantial region of configuration space in each topological sector $\Omega$ that are labeled by the topological loop operators:

$$\Omega = \left(W^{Z_3}_X, W^{Z_3}_Y\right).$$

The exact zero energy ground state at the RK point, where $t_1 = v_i, i \in \{1,2,3\}$, is the RK state, where the superposition is taken to be over all states in the topological sector. Since the $W^{Z_3}_X$ and $W^{Z_3}_Y$ have eigenvalues $\{1, e^{i2\pi/3}, e^{-i2\pi/3}\}$, the RK point of the QDPM is therefore 9-fold degenerate on the torus.

Defects in the QDPM include monomers, trimers, and tetramers. As these dimer configurations violate the local constraint at the vertex, these defects carry a non-vanishing $Z_3$ electric charge $Q^{Z_3}_{\nu} = \pm 1$, where the sign depends on the sub-lattice and type of defect. Tab. I lists the $Z_3$ charge for these defects on one sublattice; defects of the same type carry opposite charge when living on opposite sublattices.

We have demonstrated that the ground state at the RK point of the QDPM has an exact $Z_3$ gauge symmetry and a corresponding 9-fold topological degeneracy on the torus. We emphasize that despite the exact $Z_3$ local gauge symmetry of the QDPM, there is not an exact mapping to a $Z_3$ gauge theory on the square lattice, as the QDPM has Ising degrees of freedom on the links, rather than the 3 dimensional local Hilbert space of a $Z_3$ gauge theory (see Appendix A for details of $Z_3$ gauge theory). It remains to address whether this RK state is in the deconfined $Z_3$ topologically ordered phase of a $Z_3$ gauge theory, or a confined symmetry broken or critical state, which we discuss below.

IV. NUMERICAL STUDY OF THE RK GROUND STATE OF THE QDPM

To determine the nature of the ground state at the RK point of the QDPM, we have Monte Carlo sampled $|\Psi_{RK}\rangle$, using local updates corresponding to the dimer resonances shown in Tab. II and single plaquette flips, as defined in Eq. (7). Below we present the results of calculations on $L \times L$ square lattices with periodic boundary conditions on systems of linear dimension up to $L = 128$.

A. Topological sectors

We explicitly demonstrate that the extensive topological degeneracy of the QDM is reduced to a finite degeneracy in the QDPM by computing the histogram of $U(1)$ topological winding numbers $\Phi_X$ and $\Phi_Y$, which is shown in Fig. 5. The different colors represent calculations that were initialized to different winding sectors ($\Phi_X, \Phi_Y$), and we find that $\Phi_X$ and $\Phi_Y$ are not strictly conserved (as they are in the QDM), but are only conserved modulo 3. The histogram $P(\Phi)$ has a Gaussian shape with a width that increases with system size; this is expected as $\Phi = \pm L/2$ correspond to ordered configurations and thus there are fewer dimer configurations near these limits. Although we cannot rule out the possibility of a variety of isolated “frozen” configurations (e.g. pentamers occupying all sites) which are not accessed by the dynamics described in Tab. II, we find that the increased dynamics of the QDPM connect $U(1)$ topological sectors that are disconnected in the QDM; in particular we note that the QDPM gives dynamics to “staggered” dimer configurations that are frozen in the QDM (see Fig. 4). For the remainder of the paper we present results in the $(0,0)$ topological sector, but we see no qualitative dependence between topological sectors for any of the results presented here. Thus we find that the QDPM has a finite 9-fold degeneracy on the torus, where the topological sectors are labeled $(W^{Z_3}_X, W^{Z_3}_Y)$, in contrast with the extensive degeneracy of the QDM.
B. Dimer density correlations

We will now present a numerical study of the dimer density correlations in the RK ground state of the QDPM. In Fig. 6 (a) we plot the bare dimer-dimer density correlation function \( \langle n_\alpha n_\tau \rangle \) on a \( L = 32 \) lattice; here the origin \( 0 \) is represented by the orange link, and \( r \) varies over all other links of the lattice. These bare dimer correlations display only short range correlations with no obvious symmetry breaking. In Fig. 6 (b) we show the equal-time dimer-dimer structure factor,

\[
S_d(k) = \frac{1}{L^2} \sum_{i,j} e^{-i \mathbf{k} \cdot \mathbf{r}_{ij}} \langle n_\alpha(r) n_\tau(r) \rangle
\]

where we restrict the sum to horizontal links, for a \( L = 64 \) lattice. Fig. 6 (c) shows \( S_d(k) \) for a particular path through the Brillouin zone. The dimer structure factor displays broad maxima at \( k = (0, \pm \pi) \) and \( k = (\pm \pi, \pm \pi) \). The lack of rotational symmetry of \( S_d(k) \) is due to the fact that we have isolated horizontal dimers; the correlations between vertical dimers display the same behavior, rotated by \( \pi/2 \). We have plotted the finite size scaling of the intensity of the broad maximum of \( S_d \) in Fig. 7, which we find to vanish in the thermodynamic limit; these maxima are therefore consistent with finite size effects due to short distance correlations. In Appendix B we additionally have plotted order parameters that specifically test for rotational and translational symmetry breaking; in all cases we find no symmetry breaking in the thermodynamic limit.

Additionally we have studied the nature of the decay of the connected dimer-dimer correlation function,

\[
C_d(r) \equiv \langle n_\alpha n_\tau \rangle - \langle n_\alpha \rangle \langle n_\tau \rangle
\]

where \( r \) is the distance between the links \( 0 \) and \( r \). In Fig. 8, \( C_d \) between parallel links is plotted as a function of separation, for \( r \) perpendicular to the orientation of
the dimers for a $L = 64$ lattice. We find that $C_d$ decays exponentially with distance having a correlation length $\chi \approx 1$. Additionally, $C_d$ displays a sub-lattice oscillation that is consistent with the finite-size maximum in $S_d$ at $(0, \pi)$. Thus we find that the RK point of the QDPM is a symmetric dimer liquid with exponentially decaying dimer correlations, in contrast with the power-law decaying correlations of the QDM.

C. $Z_3$ magnetic string correlations

To investigate the $Z_3$ gauge theory description of the QDPM RK ground state, we study the $Z_3$ magnetic string correlation function as defined in (6) & (11):

$$C_M (r_{pp'}) \equiv \left\langle \exp \left[ \frac{2\pi}{3} i \left( \Phi_{p p'} - \Phi_{p' p} \right) \right] \right\rangle,$$

where $p$ and $p'$ are two plaquettes, $r_{pp'}$ is the displacement between $p$ and $p'$, and $\gamma_{pp'}$ is any open string on the dual lattice connecting $p$ and $p'$. In Fig. 9 (a), we have plotted $|C_M|$ for a $L = 32$ lattice. Note that while $C_M$ depends on the choice of the background dimerization which fixes the gauge, $|C_M|$ is independent of the background dimerization. We find only short range magnetic string correlations that display no obvious symmetry breaking.

Additionally, we have computed the magnetic string structure factor

$$S_M (k) \equiv \frac{1}{L^2} \sum_{p,p'} e^{-i k \cdot r_{pp'}} C_M (r_{pp'})$$

and plot the magnitude $|S_M|$ in Fig. 9 (b). We have chosen a columnar background dimerization comprising horizontal background dimers stacked in columns along the $y$-axis, as shown in Fig. 3; this choice breaks the translational symmetry of the lattice and thus requires doubling the unit cell the $x$-direction. While $|C_M|$ displays no obvious symmetry breaking, there are peaks in $|S_M|$ at $(0,0) & (0,\pi)$. We have plotted the finite-size scaling of these peaks in Fig. 7 which demonstrates that these peaks vanish in the thermodynamic limit. We have additionally studied $S_M$ using different background dimerization and find that the location of the finite-size peaks depend on the gauge choice; however in all cases we find that these peaks vanish in the thermodynamic limit. The vanishing of the $(0,\pi)$ peak demonstrates the restoration of rotational and translational symmetries in the thermodynamic limit, and the additional vanishing of the $(0,0)$ peak demonstrates the lack of magnetic vortex condensation, which is a mechanism for destroying topological order and breaking lattice symmetries.

Fig. 10 shows the decay of $|C_M|$ along one of the axes of the dual lattice which we find decays exponentially with distance, with a correlation length $\chi \approx 3$. These exponentially decaying magnetic string correlations are therefore consistent with a symmetric dimer liquid with no magnetic vortex condensation.

D. Monomer deconfinement

To study the confinement of $Z_3$ electrically charged defects in the QDPM, we have Monte Carlo sampled the RK ground state doped with a single pair of monomers.
We also have demonstrated that monomer defects are deconfined in the QDPM, in contrast with the critical confinement of monomers in the QDM. Finally, we find that the $Z_3$ magnetic string correlations decay exponentially and display no evidence of $Z_3$ vortex condensation; this is consistent with $Z_3$ vortices being gapped quasiparticles excitations above the QDPM ground state. These results suggest that the low energy physics of the QDPM is described by a $Z_3$ gauge theory with a $Z_3$ topological ordered ground state. Thus we have demonstrated that systems with Ising degrees of freedom and local constraints can lead to $Z_N$ topological order for $N > 2$, providing the a new stable phase of matter beyond the commonly seen $Z_2$ topological spin liquid.

Future work could directly probe the existence of a finite gap above the RK point by studying the imaginary time dynamics of correlations functions using Monte Carlo methods similar to those used in here.\textsuperscript{47,48} Such an approach could be used to explore if the low energy excitations are best described by $Z_3$ magnetic vortices.\textsuperscript{31,55,56} Additionally, one could conclusively demonstrate the existence of $Z_3$ topological order by computing the topological entanglement entropy in the QDPM ground state using Monte Carlo methods.\textsuperscript{13,57,58}

A finite gap above the RK point suggests that a stable topologically ordered phase should exist surrounding the RK point. While the Monte Carlo techniques used in this manuscript are limited to the RK point, the phase diagram near the RK point can be explored in future work using quantum Monte Carlo methods, as the QDPM has no sign problem. Away from the RK point, the condensation of $Z_3$ magnetic vortices could drive a continuous phase transition out of the topologically ordered phase, analogous to vison condensation in $Z_2$ spin liquids.\textsuperscript{30–32,59,60} If such a continuous transition exists in the QDPM it may provide access to a new universality class of quantum transitions.\textsuperscript{61}

Finally, given the stability of the topologically ordered

\begin{equation}
\text{FIG. 10.} \text{ The magnitude of the } Z_3 \text{ magnetic string correlation } |C_{m1}| \text{ along one axis of the dual lattice of a } L = 64 \text{ lattice. The line corresponds to an exponential fit with correlations length } \chi \approx 3.
\end{equation}

\begin{equation}
\text{FIG. 11.} \text{ The monomer-monomer correlation function } \langle m_0 m_r \rangle \text{ along a coordinate axis in the QDPM RK ground state doped with a single pair of monomers on lattices up to } L = 64. \text{ The horizontal lines correspond to } L^{-4} \text{ for the system size corresponding to the color.}
\end{equation}
phase, it may be possible to realize the QDPM in a more microscopically realistic spin model with two body interactions. The local constraints of the QDM can be realized in two-body spin-1/2 and Bose-Hubbard models, and the QDM Hamiltonian can arise perturbatively in the constrained low energy Hilbert space. Thus future work, it may be possible to engineer the QDPM in the low energy subspace of an experimentally realizable two-body model, providing a route to the experimental realization of a $Z_3$ topologically ordered spin liquid.

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Appendix A: $Z_3$ Gauge theory on the square lattice

Here we will review a string-net implementation of $Z_3$ gauge theory on the square lattice. We define a three-dimensional Hilbert space on each link, corresponding to the presence or absence of an oriented string segment that carries $Z_3$ electric flux, which are eigenstates of the Hermitian $Z_3$ electric flux operator $E_{vv'}$

\[
E_{vv'}(\mathbf{v},\mathbf{v}') = 0, \quad (A1)
\]

\[
E_{vv'}(\mathbf{v},\mathbf{v}') = +1, \quad (A2)
\]

\[
E_{vv'}(\mathbf{v},\mathbf{v}') = -1. \quad (A3)
\]

At each vertex we define a local unitary gauge transformation $G_v$, $G_v \equiv \prod_{v'} P_{vv'}^{1}$

where the product is over all nearest neighbor vertices $v'$, and we have defined the unitary operator

\[
P_{vv'}^{1} \equiv e^{i2\pi/3E_{vv'}}.
\]

We consider eigenstates of $G_v$ to have a definite $Z_3$ electric charge at vertex $v$. The charge-free sector is the gauge invariant subspace where

\[
G_v(\psi) = |\psi\rangle \quad \forall v.
\]

This gauge invariant subspace corresponds to states where the “string flux” vanishes modulo 3 at each vertex, thus including vertices where three inward or three outward strings meet at a vertex. For eigenstates of $G_v$ with nonzero electric charge

\[
G_v|\psi\rangle = e^{\pm2\pi/3i}|\psi\rangle
\]

which corresponds to the presence of a $Z_3$ charge of $\pm 1$ at vertex $v$. A vertex with a positive (negative) charge has a net outward (inward) string flux of 1 modulo 3, or net inward (outward) flux of 2. Fig. 12 shows examples of charged and charge free string configurations.

To provide dynamics, we define a unitary operator electric flux ladder operator $Q^l_{vv'}$ that cyclically increments the electric flux:

\[
Q^l_{vv'}|\mathbf{v},\mathbf{v}'\rangle = |\mathbf{v},\mathbf{v}'\rangle, \quad (A4)
\]

\[
Q^l_{vv'}|\mathbf{v},\mathbf{v}'\rangle = |\mathbf{v},\mathbf{v}'\rangle, \quad (A5)
\]

\[
Q^l_{vv'}|\mathbf{v},\mathbf{v}'\rangle = |\mathbf{v},\mathbf{v}'\rangle. \quad (A6)
\]

The simplest gauge invariant Hamiltonian is then

\[
H_{Z_3} = -J \sum_p \left( Q^l_{v_i v_{i+1}} Q^l_{v_{i+1} v_{i+2}} Q^l_{v_{i+2} v_{i+3}} Q^l_{v_{i+3} v_{i+4}} + h.c. \right)
\]

where $v_i^p$ represent the four vertices in the plaquette $p$, ordered cyclically by nearest neighbors.

We can define two oriented open string operators which commute with $H_{Z_3}$, everywhere except at the endpoints. The electric string operator $S_e$, which acts on an open, oriented string on the lattice $\gamma_{vv'}$ with endpoints $v, v'$, and creates a $+1 (-1)$ charge at vertex $v (v')$, is defined as

\[
S_e(\gamma_{vv'}) \equiv \prod_{l \in \gamma_{vv'}} Q^l_{v_l v_{l+1}}
\]

where the product is over all links $l$ in $\gamma_{vv'}$, $v_l$ and $v_{l+1}$ are the two vertices in $l$, ordered according to the orientation of $\gamma_{vv'}$ (see Fig. 12). The magnetic string operator $S_m$ is defined by an open, oriented string on the dual lattice and creates a magnetic vortex-antivortex pair on plaquettes at the endpoints of $\gamma_{pp'}$

\[
S_m(\gamma_{pp'}) \equiv \prod_{l \in \gamma_{pp'}} P^l_{v_l v_{l+1}}
\]

where the product is over all links $l$ that cross $\gamma$ and $v_l$ ($v_{l+1}$) is the vertex to the left (right) of the string as $\gamma$ is traversed from $p$ to $p'$.

Note that $S_e$ and $S_m$ do not commute if their corresponding strings intersect—this underlies the non-trivial statistics of electric charges and magnetic vortices. Consider the magnetic string operator $S_{mn}(\Gamma)$ defined by an oriented closed loop $\Gamma$ on the dual lattice. This operator corresponds to the creation of a vortex-antivortex pair, winding of the vortex around $\Gamma$, and annihilation of the vortex pair. A state with definite electric charge inside $\Gamma$ will be an eigenstate of $S_{mn}(\Gamma)$, with eigenvalue $e^{i2\pi/3q}$, where $q = 0, \pm 1$ is the net $Z_3$ electric charge (modulo 3) inside $\Gamma$. Thus for $\Gamma$ winding around an isolated electric charge, we see that electric charges and magnetic vortices are anyons with relative braiding statistics with a statistical phase of $e^{i2\pi/3}$.

Additionally, these string operators give rise to a topo-
logical degeneracy in systems of non-trivial topology. For example, consider the charge free sector on a torus. We can consider two topologically non-trivial closed loops $\Gamma_X \& \Gamma_Y$ which have a single winding around the $x$ or $y$ directions of the torus. Since $S_m(\Gamma_X)$ commutes with $H_{Z_3}$, we can label the energy eigenstates by the eigenvalues of $S_m(\Gamma_X)$, and similarly for $\Gamma_Y$: $\{1, e^{\pm 2\pi i/3}, e^{-\pm 2\pi i/3}\}$. Thus $H_{Z_3}$ has a 9-fold ground state degeneracy on the torus, and the ground states subspace can be labeled by the $Z_3$ electric flux winding about each axis of the torus.

**Appendix B: Dimer order parameters QDPM RK ground state**

To explicitly test for the possibility of translational and rotational symmetry breaking corresponding to the features in $S_d$ at the Brillouin zone edges, we define several order parameters which would capture such potential symmetry breaking. For rotational symmetry breaking we define:

$$M_{\text{rot}} \equiv \frac{2}{L^2} (N_v - N_h), \quad \text{(B1)}$$

where $N_v$ ($N_h$) is the total number of vertical (horizontal) dimers; note that $\langle M_{\text{rot}}^2 \rangle \neq 0$ implies there is rotational symmetry breaking. To capture translational symmetry breaking by one lattice site in a direction perpendicular to the majority of the dimers we define:

$$M_\perp \equiv \frac{8}{L^2} \sum_\Box (-1)^r \theta (M_{\text{rot}}) \Pi_\Box^h \left[ (-1)^y \theta (-M_{\text{rot}}) \Pi_\Box^h \right], \quad \text{(B2)}$$

where $\theta(x)$ is the Heaviside function, and $\Pi_\Box^h (\Pi_\Box^v)$ is one if the plaquette $\Box$ has two horizontal (vertical) dimers, and zero otherwise. Similarly, we define $M_\parallel$ to be given by Eq. (B2) with $r_x$ and $r_y$ swapped between terms. Additionally, we define order parameters that indicate “columnar” and “staggered” dimer order:

$$M_{\text{stag}}^2 \equiv \frac{4}{L^4} \left[ \left( \sum_{\ell_h} (-1)^{r_x + r_y} n_d (\vec{r}_{\ell_h}) \right)^2 + \left( \sum_{\ell_v} (-1)^{r_x + r_y} n_d (\vec{r}_{\ell_v}) \right)^2 \right], \quad \text{(B3)}$$

$$M_{\text{col}}^2 \equiv \frac{4}{L^4} \left[ \left( \sum_{\ell_h} (-1)^{r_x} n_d (\vec{r}_{\ell_h}) \right)^2 + \left( \sum_{\ell_v} (-1)^{r_y} n_d (\vec{r}_{\ell_v}) \right)^2 \right], \quad \text{(B4)}$$

where the sums over $\ell_h$ ($\ell_v$) are taken over all horizontal (vertical) links. In Fig. 13 we have plotted the finite-size scaling of these order parameters; we find in all cases that while finite size systems display symmetry breaking due to finite-size effects, these order parameters all vanish in the thermodynamic limit, as demonstrated by the extrapolation shown in Fig. 13.
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