Bridging lattice-scale physics and continuum field theory with quantum Monte Carlo simulations

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We discuss designer Hamiltonians—lattice models tailored to be free from sign problems (“designed”) when simulated with quantum Monte Carlo methods but which still host complex many-body states and quantum phase transitions of interest in condensed matter physics. We focus on quantum spin systems in which competing interactions lead to non-magnetic ground states. These states and the associated quantum phase transitions can be studied in great detail, enabling direct access to universal properties and connections with low-energy effective quantum field theories. As specific examples, we discuss the transition from a Néel antiferromagnet to either a uniform quantum paramagnet or a spontaneously symmetry-broken valence-bond solid in SU(2) and SU(N) invariant spin models. We also discuss anisotropic (XXZ) systems harboring topological Z2 spin liquids and the XY* transition. We briefly review recent progress on quantum Monte Carlo algorithms, including ground state projection in the valence-bond basis and direct computation of the Renyi variants of the entanglement entropy.

I. INTRODUCTION

Understanding ground states of interacting quantum systems is one of the central defining challenges of condensed matter physics. On the one hand, quantum field theory is a powerful and general framework for studies of...
low-energy properties of complex, strongly correlated and entangled quantum matter [1]. On the other hand, this continuum approach has potential pitfalls due to its generality, and limitations due to its insensitivity to lattice scale physics, making it important to also carry out direct studies of microscopic lattice Hamiltonians. A fruitful direction is to combine the advantages of both methods by analyzing data from unbiased numerical investigations of lattice Hamiltonians with quantum field theoretic predictions of emergent low energy properties. Such connections between the two approaches can have significant synergistic effects in advancing our understanding of quantum many-body phenomena—as they have had and continue to have in classical statistical mechanics, especially in studies of thermal phase transitions [2,3].

Computational many-body research faces important challenges since the quantum models of interest are often beyond the reach of existing algorithms. Two-dimensional (2D) and 3D frustrated quantum spin systems and fermion systems are the main groups of difficult systems, since they suffer from the infamous “sign problem” in quantum Monte Carlo (QMC) simulations [4–6]. The sign problem arises when the weight function in the configuration space constructed using the Euclidean path integral or other mappings to an effective statistical-mechanics problem is not positive definite, hence invalidating the interpretation of the weights as a probability distribution for importance sampling. The approach we advocate and review here is to construct particular “designer Hamiltonians”, which host interesting ground states and quantum phase transitions but still are amenable to large-scale QMC studies without sign problems (i.e., they are designed and “de-signed” to be practically useful).

A common misconception is that interesting designer Hamiltonians should not exist—if a model is sign-problem free it must be trivial or uninteresting. In this article, we will review some recent counter-examples to this pessimistic view, from the area of quantum magnetism. Moreover, advances in QMC methods [7–12] have made it possible to truly approach the low-energy limit of many important models without any approximations (beyond controllable statistical errors), and to compute quantities of current interest in condensed matter physics and quantum information theory, e.g., the fidelity [13], the geometric tensor [14], and the entanglement entropy [15,16]. We will briefly address some of these technical developments as well. We first make some further remarks on field theory and the philosophy of the synergistic use of designer Hamiltonians, and then outline the topics covered in this Review.

A. Field theory and numerical studies of Hamiltonians

The natural starting point for theoretical models of condensed matter systems are microscopic Hamiltonians. An effective field theoretic description of a given Hamiltonian is constructed based on a combination of insights and assumptions on the properties of the Hamiltonian that control the long distance physics according to the renormalization group (RG) approach. These properties most famously include symmetry and dimensionality, but more subtle effects like the nature of the topological defects also play an important role. The fixed points of the field theory control the long distance physics. The RG approach may result in flows to a previously known fixed point, or to some fixed point whose properties are not known. The RG itself often relies on approximations that are not easy to justify, and, if the flow is to some unknown state, it is challenging to characterize it. Often to control the RG flows, analytic expansions around some special simplifying limits are considered. Well known examples are the $\epsilon$ expansion around the upper critical dimension and the $1/N$ expansion in the number of components $N$ (spin components, flavors of fermions, etc.) [17,18]. The extrapolation to the $\epsilon$ or $N$ of interest is often problematic, however, because only low-order expansions are feasible in practice—or worse still, the expansion may not converge at the value of $\epsilon$ or $N$ of primary interest.

One example of the above approach of deriving effective field theories from microscopic Hamiltonians by a limiting expansion is the $(d+1)$-dimensional nonlinear $\sigma$-model description of interacting quantum spins in $d$ dimensions, which was derived by Haldane in the semi-classical limit of large spins $S$ [19,20]. Another example—central to this review article—is the non-compact $CP^{N−1}$ description of the intriguing Néel to valence-bond solid (VBS) “deconfined” quantum-critical point in 2D SU($N$) quantum magnets, which can be justified in the large-$N$ limit [21,22]. Whether or not the theory remains correct down to small $N$, in particular $N = 2$, remains a challenging question.

In view of the uncertainties with analytic methods highlighted above, it is important to test the predictions of field theories in some unbiased way, starting from a microscopic description of the system or phenomenon of interest. In some cases there are exact solutions, e.g., the Bethe Ansatz solution of the $S = 1/2$ Heisenberg chain [23] has been crucial for testing the non-linear $\sigma$-model description of this class of critical spin chains. The exact AKLT state [24] for a special version of the $S = 1$ chain was similarly important in confirming Haldane’s conjecture of the qualitative differences between half-odd integer and integer spins [25]. Exact solutions are rare, however, and mostly limited to 1D models.

Another, more general approach is to study Hamiltonians numerically. Exact diagonalization of the Hamiltonian is possible only for very small systems; currently up to 42 $S = 1/2$ spins [26,27]. White’s density matrix renormalization (DMRG) scheme [28] and related approaches based on matrix-product states [29] have enabled more detailed studies of the low-energy physics of 1D systems, including also “ladders” of several coupled chains [30]. QMC studies are also competitive in 1D in
the absence of sign problems [31–33].

While there has been some progress for 2D systems with DMRG [34] and tensor-product [35–37] states, which are higher-dimensional generalizations of the matrix-product states generated in DMRG [38], in general these methods have not yet reached the point where one can obtain unbiased results for generic nontrivial models. There are encouraging developments, however, of calculations addressing challenging systems and approaching the level of accuracy where definite conclusions can be drawn [33, 40].

Presently, however, the only numerical approach with which one can routinely reach sufficiently large 2D and 3D lattices in a completely unbiased way is QMC simulations [41–42], but, as already noted, they are restricted to systems free from sign problems. The class of models for which the sign problem is either absent or evadable still contains a vast range of Hamiltonians with nontrivial and interesting ground states and quantum phase transitions. To study low-energy emergent properties, the interactions do not necessarily have to correspond in detail to any particular real material—although some times that is also possible [43]. The idea is to design a sign-problem free Hamiltonian in such a way that it contains a particular macroscopic (low-energy) phenomenon of interest—so that universal physics is captured.

Bench-mark results obtained from such designer Hamiltonians, representing various physical phenomena in a prototypical manner, can be very useful experimentally. Here our main focus will be on the theory side. With unbiased large-scale QMC simulations, one can test field theories proposed to capture specific classes of quantum many-body states or quantum phase transitions. Moreover, explorations of designer Hamiltonians can also serve as “experiments” for discovering novel phenomena, and, thereby, stimulate further theoretical developments.

A designer Hamiltonian corresponds to an effective classical statistical-mechanics problem (with a real action), which arises out of the construction of the sampling space in the QMC scheme implemented. In some cases quantum phenomena in d dimensions can be studied using Monte Carlo simulations with the simplest generalization to a classical model with the same global symmetries in d + 1 dimensions [44–46] (e.g., the thermal phase transition of the 2D classical Ising model is in the same universality class as the quantum phase transition of the transverse-field Ising chain [41–47]). Often, however, the effective classical model resulting from the construction of a QMC configuration space, in a complete treatment of a given a quantum Hamiltonian, has unusual degrees of freedom and interactions, unanticipated in the study of classical statistical mechanics. The naive mapping with just d → d + 1 can then miss important and intriguing phenomena special to quantum mechanics [22, 23, 48]. Thus, in our view the approach of directly studying quantum mechanical designer Hamiltonians is an important direction in forming an unbiased picture of quantum criticality and emergent phenomena.

B. Outline of the Review

In this Review, we will focus on some recent examples where it has been possible to study quantum phase transitions in detail by QMC simulations and to make quantitative comparisons of extracted universal quantities with field theoretic results. We also discuss other fascinating quantum phenomena, e.g., emergent gauge fields and entanglement, where QMC studies are playing a critical role in establishing benchmarks and gaining insights that are hard or impossible to extract from the field theories.

In Sec. II we discuss the sign problem and some of the QMC schemes used in large-scale studies of quantum spin systems, including treatments of SU(N) symmetry and recent progress in computing the Rényi entanglement entropy. In Sec. III we give examples of conventional and deconfined quantum-critical points in SU(2) invariant models, and in Sec. IV we show how generalization to SU(N) symmetry can be used to directly connect to large-N studies within field theory. In Sec. V we discuss spin liquid states and associated quantum phase transitions in U(1) symmetric, highly frustrated “XXZ” spin models (or, equivalently, interacting hard-core bosons). We conclude in Sec. VI with some further discussion and an outlook on future prospects and challenges.

II. MODERN QMC METHODS

There is a wide range of QMC methods available for studies of different classes of lattice models [41, 42, 49]. In this Review we focus on spin models, and therefore restrict the discussion to the types of methods most suitable for them. These methods are also often well suited for studies of bosons, where a substantial body of work has been carried out in the past several years, stimulated by ultracold atoms in optical lattices [50–53]. For fermions, there are alternative approaches, based upon auxiliary fields decoupling the interactions and subsequent analytical tracing-out of the fermions [49, 54]. These methods have a much less severe fermion sign problem (in some cases completely avoiding it, e.g., for particle-hole symmetric systems)—with the exception of 1D systems, where the methods discussed here also work well (since the signs from anticommutation normally do not appear in this geometry). We will only give a brief non-technical overview of QMC methods here, with references to more detailed accounts.

A. The sign problem

Solving many-body quantum mechanical problems on a computer is generally exponentially hard (in computing time versus system size) with known algorithms. The statistical mechanics of a d-dimensional quantum Hamiltonian can be rewritten in terms of some classical vari-
ables $C$ in $d + 1$ dimensions, i.e., the partition function ($T > 0$) or the normalization ($T = 0$) can be cast into the form

$$Z = \sum_C W_C. \quad (1)$$

Systems of interest can, for practical, known ways of transforming into this form, have both positive and negative weights $W_C$, invalidating the usual interpretation of $W_C/Z$ as a probability distribution in Monte Carlo simulations. Formally, this difficulty can be side-stepped by not including the sign in the probability, using $P_C \propto |W_C|$, and compensating for this by weighting observables with the sign:

$$\langle O \rangle = \frac{\sum_C W_C O_C}{\sum_C W_C} = \frac{\sum_C O_C \text{Sign}(W_C) |W_C|}{\sum_C \text{Sign}(W_C) |W_C|}$$

$$= \langle \text{Sign} \cdot O \rangle_{|W|}, \quad (2)$$

The problem here is that the numerator and denominator in the last expression both approach zero exponentially in system size and inverse temperature, and, hence, resolving the ratio within statistical noise is exponentially hard. The advantage of importance sampling (time speed-up from exponential to polynomial scaling) is then lost. This predicament is the QMC sign problem.

Here, we consider classes of quantum Hamiltonians for which it is possible to choose a basis in which all the $W_C \geq 0$. These sign-problem free models can generally be simulated in polynomial time by importance sampling, allowing access to large system sizes (with the exception of “glassy” systems and other systems where even classical Monte Carlo simulations scale exponentially). In some cases, sign problems that appear intractable at first-sight can be solved in more sophisticated ways, e.g., in the Meron algorithm [55]. Here we implicitly consider cases where the “de-signing” is essentially automatic, being a direct consequence of the non-positivity of the off-diagonal matrix elements of the Hamiltonian.

**B. $T > 0$ and $T = 0$ QMC methods**

In finite-temperature methods, the goal is to compute thermal averages

$$\langle A \rangle = \frac{1}{Z} \text{Tr} \{ A e^{-\beta H} \}, \quad Z = \text{Tr} \{ e^{-\beta H} \}. \quad (3)$$

where $\beta = 1/T$. As an alternative to studying the ground state in the limit of $T \rightarrow 0$, in a ground-state projector method some operator $P(\beta)$ is applied to a “trial state” $|\Psi_0\rangle$, such that $|\Psi_\beta\rangle = P(\beta) |\Psi_0\rangle$ approaches the ground state when $\beta \rightarrow \infty$. An expectation value

$$\langle A \rangle = \frac{1}{Z} \langle \Psi_\beta | A | \Psi_\beta \rangle, \quad Z = \langle \Psi_\beta | \Psi_\beta \rangle, \quad (4)$$
tends to the true ground state expectation value, $\langle A \rangle \rightarrow \langle 0 | A | 0 \rangle$. For the projector, one can use the imaginary-time evolution operator $P(\beta) = e^{-\beta H}$ or a high power of the Hamiltonian; $P(m) = H^m$. Here $m \propto \beta N$ gives the same rate of convergence for the two choices of projectors for a given system size $N$. This follows from a Taylor expansion of the time evolution operator, which for large $\beta$ is dominated by powers of the order $n = \beta |E_0|$, where $E_0$ is the ground state energy.

**C. Path integrals and stochastic series expansions**

The task is now to rewrite $Z$ in Eqs. (3) or (4) in the form [1], and expectation values as Eq. (2), with classical variables without diagonalizing $H$. In practice, $T = 0$ and $T > 0$ schemes for a given model are often very similar. In both cases, the exponential operator can be treated with path-integral methods—world lines in discrete [54, 55] or continuous [9, 57, 58] imaginary time—starting from a product of “time slice” evolution operators,

$$e^{-\beta H} = \prod_{i=1}^M e^{-\Delta H}, \quad \Delta = \frac{\beta}{M}. \quad (5)$$

Complete sets of states are then inserted between the exponentials (and the trace is further taken at $T > 0$). For small $\Delta$, the matrix elements of the exponentials can be evaluated approximately, giving the form [11] with $W_C$ a product of $M$ matrix elements. In modern methods the limit $\Delta \rightarrow 0$ is taken at the algorithmic level [3, 11, 57] and the Monte Carlo sampling is of paths in continuous imaginary time. Older approaches employed the Suzuki-Trotter decomposition [17] in Eq. (3), which typically leads to an error $O(\Delta^2)$.

An alternative to Eq. (5) is to start from a series expansion [7, 57],

$$e^{-\beta H} = \sum_{n=0}^\infty \frac{\beta^n}{n!} (-H)^n, \quad (6)$$

and insert complete sets of states between each instance of $H$. The power $n$ itself is importance-sampled along with the terms of $H$ (Stochastic Series Expansion; SSE). This expansion is sharply peaked around $\langle n \rangle = -\beta E$, where $E$ is the total internal energy (under the assumption that there is no sign problem), which leads to a computational effort scaling as $\beta N$, which is the same as in the path-integral approaches. Again, $W_C$ in Eq. (1) is a product of matrix elements.

The continuous-time path integral can also be seen as a variant of SSE, where the evolution operator is written in the interaction picture, with $H = H_0 + V$ and expanding only in the perturbation $V$ to some conveniently chosen $H_0$ [57, 60]. The full SSE, where $H_0 = 0$, $V = H$, is normally more efficient for spin systems, while
the continuous-time variant should be better for certain boson systems [61].

Regardless of how the exponential operator is treated, the difference between Eqs. (1) and (4) is essentially only in the boundary conditions in the imaginary time direction; periodic for \( T > 0 \) (due to the trace being taken) and dictated by the nature of the trial state in \( T = 0 \) projections. An equal superposition over all basis states corresponds to fully open boundaries, while other choices of the trial state leads to “biased” boundaries with some weighting of the bra and ket states (depending on the details of the trial state). The projector approach with \( P(\beta) = e^{-\beta H} \) is often called the Path Integral Ground State method (PIGS) in the context of continuous-space systems [62, 63].

There is no sign problem in these approaches for spin systems with non-positive definite off-diagonal matrix elements (for a properly chosen trial state in the case of the \( T = 0 \) approach). For bipartite antiferromagnetic interactions a sublattice rotation can accomplish this, or, equivalently, one can note that for such models the signs are “invisible”, because all non-vanishing terms of the path integral or series expansion have an even number of off-diagonal matrix elements. For further details of sign issues, see, e.g., Refs. [6, 7, 11]. Note that there are no restrictions on the diagonal interactions in the chosen basis, so that one can study, e.g., spin and boson systems in which the potential energy is highly frustrated (as we will discuss below in Sec. V).

### D. Sampling with loop and cluster algorithms

A breakthrough leading to today’s efficient QMC algorithms was the realization by Evertz et al. [8] that the classical Swendsen-Wang (SW) cluster Monte Carlo method [64] could be generalized to loop algorithms (i.e., generating clusters in the form of loops) for vertex models, and that these models are very similar to the effective statistical mechanics problems arising in the transformations of quantum problems discussed above. While the loop algorithm has been adapted to many different models, both within the path-integral [6, 11, 65, 66] and SSE [11] frameworks, they cannot be applied to all cases (in analogy with the SW method, which also has limited applicability). The loop concept has also been further generalized, however, to “worms” [10, 57, 58] and “directed loops” [11, 67], which are essentially loops that can self-intersect during their construction and incorporate the detailed-balance principle in a more general manner. Recently such an algorithm was also formulated without detailed balance [68]. In addition, for some models where loop algorithms and their generalizations are not applicable (e.g., transverse-field Ising models), other efficient generalizations of the SW cluster concept have been developed [69, 70].

### E. SU(2)-symmetric models

A particularly convenient class of trial states in projector QMC studies of SU(2) invariant interactions is the amplitude-product states in the overcomplete valence-bond (singlet pair) basis [71], which were recently generalized to include bond correlations [72]. These total-singlet states incorporate Marshall’s sign rule in a convenient way and also automatically have the appropriate momentum for the ground state of a periodic system (thereby from the outset filtering out a significant fraction of the excited states). The Hamiltonian in this case can typically be expressed in terms of two-spin singlet-projector operators (individual ones or products of two or more of them);

\[
P_{ij} = \frac{1}{4} - S_i \cdot S_j = |s_{ij}| \langle s_{ij} \rangle ,
\]

where \( s_{ij} \) denotes the singlet state of spins \( i \) and \( j \). A power of the Hamiltonian \( H_m \), for fixed \( m \) or in a series expansion, Eq. (9), is expanded into all possible products of these projectors, which act on the ket trial state in (10). In early versions of this scheme, the resulting paths of valence bond states were sampled directly [73, 74], while in more recent formulations the diagonal and off-diagonal parts of the projectors are also sampled, along with spin configurations compatible with the valence-bond configurations of the trial state (which are also sampled) [75]. An example of a configuration in such a simulation is shown in Fig. 1. This configuration space is amenable to the very efficient loop updates discussed above [12]. Apart from the boundary conditions in the “propagation” direction (and the associated sampling of the valence bonds in the trial state), this kind of projector method is in practice implemented in a very similar way as the SSE method for SU(2) models [12].

Evaluation of expectation values (“measurements”) in projector QMC simulations are carried out at the mid-point of the configuration (while in \( T > 0 \) methods averages can be taken over the whole time range, due to the time-periodicity), as indicated in Fig. 1. In the valence-bond basis, when propagating the bra and ket states in the pure valence-bond basis (which corresponds to averaging over all compatible spin configurations), most quantities of interest can be expressed using the loops of the transition graph formed when superimposing the projected bra and ket bond configurations [71, 77]. Estimators for operators \( A \) expressed in the space \( \{L\} \) of transition-graph loops are typically of the form

\[
A(L) = \frac{\langle V_l | A | V_r \rangle}{\langle V_l | V_r \rangle},
\]

where the overlap of the two valence-bond states is given by

\[
\langle V_l | V_r \rangle = 2^{N_v - N/2},
\]

where \( N_v \) is the number of loops and \( N \) the system size. The numerator in Eq. (8) depends on the loop structure,
generalizing the Heisenberg interaction for two spins on opposite sublattices.

Another interaction of interest is an SU(N) invariant interaction between sites having the same representation; the permutation operator, which is defined in the following way by its action on a ket:

$$\frac{1}{N} \Pi_{ij} |\alpha_i\beta_j\rangle \equiv |\beta_i\alpha_j\rangle. \quad (12)$$

We can now relate this operator to the generators $T^a$:

$$\sum_a T^a_i \cdot T^a_j + \frac{1}{N^2} = \Pi_{ij}. \quad (13)$$

This is an SU(N) invariant generalization of the SU(2) Heisenberg interaction for two spins on the same sublattice.

Hamiltonians based the SU(N) operators $P_{ij}$ and $\Pi_{ij}$ are sign-problem free if they come with negative signs, in which case they generalize the SU(2) antiferromagnetic and ferromagnetic Heisenberg exchange, respectively. In terms of the definitions of $P_{ij}$ and $\Pi_{ij}$, the algorithms presented in Sec. II B can be simply generalized from SU(2) to SU(N) by extending the number of “colors” (of the spins and the loops) from 2 to $N$. As a consequence of this, estimators expressed in terms of transition-graph loops are also modified, e.g., in the state overlap Eq. (2) is replaced by $N$. One can even generalize simulations in the pure valence-bond basis to non-integer $N$.

### F. Generalizations from SU(2) to SU(N)

The above methods for SU(2) models can be extended to a certain class of spin models with an SU(N) symmetry on bipartite lattices. First, note again that the SU(2) Heisenberg operator is simply a singlet projector, Eq. (7). In order to generalize this operator to SU(N), we choose models that have an SU(N) spin with a fundamental representation on the A sub-lattice and an SU(N) spin with a conjugate to the fundamental representation on the B sub-lattice. Denoting the $N$ states on each site by $|\alpha\rangle$, where $1 \leq \alpha \leq N$, and the generators of the fundamental representation by $T^a (N \times N$ matrices), the SU(N) singlet of a spin $i$ on the A sub-lattice and $j$ on the B sub-lattice is given by:

$$|s^N_{ij}\rangle = \frac{1}{\sqrt{N}} \sum_{\alpha} |\alpha_i\alpha_j\rangle. \quad (10)$$

We note here that for $N = 2$ and on bipartite lattices, this singlet is equivalent to the standard way of writing the singlet as $|\uparrow\downarrow\rangle - |\downarrow\uparrow\rangle)/\sqrt{2}$, by making the transformation $|\uparrow\rangle \rightarrow |\downarrow\rangle, |\downarrow\rangle \rightarrow -|\uparrow\rangle$ on one of the sublattices. We can use $|s^N_{ij}\rangle$ to rewrite the manifestly SU(N) invariant version of Eq. (4) as an explicitly sign-problem free projection operator,

$$\sum_a T^a_i \cdot T^a_j + \frac{1}{N^2} = |s^N_{ij}\rangle \langle s^N_{ij}| \equiv P_{ij}. \quad (11)$$

For integer $\alpha \geq 1$. Here, $\rho_A$ is the reduced density matrix, $\rho_A = Tr_B \{\rho\}$, and $A$ is a subregion of a lattice system (with $B$ being its complement). The von Neumann entanglement entropy corresponds to the limit $\alpha \rightarrow 1$.

The direct measure of Renyi entropies can not be done in QMC using conventional estimators, however recent work has demonstrated that it is possible to measure $S_\alpha$ for $\alpha \geq 2$ using a replica trick. Here, $\alpha$ copies of the simulation cell are used, with modified periodic boundaries in the “propagation” (or imaginary time) direction. The evaluation of Renyi entropies proceeds in...
formally different ways for $T = 0$ and $T > 0$ QMC methods \cite{13,16}. Namely, for $T = 0$, the calculation of $S_\alpha$ proceeds in analogy with Eq. (4) (where $\Psi$ is the wavefunction of the $\alpha$-times replicated system), with the operator $A$ replaced by a “swap” operator for $\alpha = 2$ \cite{15} or permutation operator for $\alpha \geq 3$ \cite{98}.

The actual simulation procedures are easiest to understand in the context of the projector algorithm discussed in Sec. III B. Here, the second Renyi entropy (for example) is given by the midpoint evaluation of a $\text{SWAP}_A$ operator on two replicas of the system (see Fig. 2), that literally swaps states (spins or valence-bond endpoints) between replicas when they lie in region $A$. Then,

$$S_2 = -\ln \left[ \langle \text{SWAP}_A(L) \rangle \right],$$

where the expectation value is calculated at the midpoint of the configuration, and expressed using the loops of the transition graph $L$ formed when superimposing the projected bra and ket states; Eq. (15). Namely,

$$\frac{\langle V_r \rangle}{\langle V_l \rangle} = 2^{N_{\text{swap}} - N_{\text{e}}},$$

where $N_{\text{swap}}$ is the number of transition-graph loops of the swapped configuration, while $N_{\text{e}}$ is for the configuration before the swap operator is applied (Fig. 2). We refer to Ref. \cite{98} for further details of this particular $T = 0$ projector QMC implementation.

At finite temperature, the explicit evaluation of the $\text{SWAP}_A$ operator can be replaced by an evaluation of the difference in free energies between a replicated system (similar to Fig. 2), and an un-replicated system \cite{16}, using Eq. (15). Although this technique looks quite different formally, both the $T = 0$ and $T > 0$ techniques can be viewed on a similar footing when considered in the context of SW loop or cluster algorithms, much like the unified framework of the QMC itself, discussed above in Sec. III B. In some cases, the implementation for these two different techniques can be made to be practically identical.

III. GROUND STATES AND QUANTUM PHASE TRANSITIONS IN SU(2) MODELS

It has been a matter of debate for a long time whether the non-linear $\sigma$-model can correctly capture a phase transition from the Néel state to a non-magnetic state in two dimensions for $S = 1/2$ spins \cite{20}. Other terms, related to Berry phases of the spins, have to be included in the long wavelength description to obtain a VBS state \cite{99,100}. It was recently proposed that the Néel and VBS states are separated by a “deconfined” quantum-critical (DQC) point \cite{22} described by the *non-compact* CP$^1$ field theory \cite{101} (in contrast to the nonlinear $\sigma$-model, which is equivalent to the *compact* CP$^1$ description, i.e., in the deconfined scenario the Berry phases render the gauge field effectively non-compact at the critical point). This

field theory describes two flavors of spinons interacting with a non-compact U(1) gauge field. The VBS and Néel order parameter form due to confinement and condensation, respectively, of spinons, and therefore cannot be regarded as two separate order parameters at the phase transition.

In this section we first discuss the rather well-studied continuous quantum phase transition between the Néel state and a quantum paramagnet in systems with bimodal Heisenberg couplings forming a static pattern. In this case the paramagnet does not break any symmetries of the Hamiltonian and the transition is believed to fall in the 3D O(3) universality class. A VBS state with spontaneously broken lattice symmetries can be achieved with frustrated interactions, which, however, are sign problematic in QMC simulations. The J-Q model \cite{102} is a designer Hamiltonian circumventing this problem. We will review some of the recent work aimed at characterizing its VBS state and Néel–VBS transition.

A. Néel–paramagnetic transition in dimerized systems

Consider $S = 1/2$ spins on a 2D bipartite lattice (e.g., the simple square lattice), interacting with antiferromagnetic Heisenberg exchange of strength $J_1$, written using the singlet projectors \cite{7},

$$H_1 = -J_1 \sum_{\langle i,j \rangle} P_{ij},$$

FIG. 2. The SWAP$_A$ operator for calculating the Renyi entropy $S_2$, where subsystem $A$ consists of the two top sites. The simulated system is composed of two non-interacting replicas, top (compare to Fig. 1), and bottom, which are sampled independently. Just to the right of the midpoint of the configuration (dashed line) the SWAP$_A$ operator (green vertical bar) connects the A spins in the top replica to the bottom one, as indicated by the colors of the loops 08.
Here \((i,j)\) denotes nearest-neighbor spins. This system has long-range Néel order at \(T = 0\). Now introduce dimerization, by considering pairs (dimers) of spins \([i,j]\) such that each spin belongs exactly to one dimer. In addition to the nearest-neighbor couplings \([17]\), let there be additional intra-dimer couplings of strength \(\lambda\) (i.e., the total intra-dimer coupling is \(J_1 + \lambda\)), such that the Hamiltonian is

\[
H = H_1 - \lambda \sum_{[i,j]} P_{ij}.
\]  

(18)

If \(\lambda > 0\) and \(J_1 \to 0\), the ground state is clearly a product of singlets on the dimers—a quantum paramagnetic state that we will refer to as a valence-bond-liquid (VBL), motivated by the fact that the state breaks no symmetries and is strongly interacting away from the large-\(\lambda\) limit. When \(\lambda = 0\) the system is the standard Heisenberg model \([17]\). There is, thus, a quantum phase transition between Néel and VBL ground states as a function of \(g = (J_1 + \lambda)/J_1\). The dimers can be arranged in different ways, and instead of dimers one can also use larger \(J_1\)-coupled units of an even number of spins, e.g., 4-site plaquettes, or one can use a bilayer, with \(J_1\) and \(J_2\) the inter-and intra-layer couplings, respectively.

1. \(T = 0\) criticality

Based on symmetry arguments alone, the Néel–VBL transition in these dimerized (or polymerized) systems should be in the classical 3D O(3) universality class. Many QMC studies have been devoted to checking the exponents against available results for the classical transition \([102, 108]\). For some of the systems, e.g., bilayers \([106]\) and columnar dimers on the square lattice \([12, 105, 108]\), results have been obtained that rival classical calculations in precision (statistical error bars) and in general the exponents agree very well with each other. In other cases, e.g., with the dimers arranged in a staggered pattern, some studies initially indicated a new universality class \([107]\), while others supported the O(3) transition \([103]\). Later it was realized that these cases, where the dimer pattern lacks inversion symmetry, are associated with certain operators (cubic interactions) in the field-theory formulation which are not present in systems with inversion symmetry \([110]\). These operators are asymptotically irrelevant in the RG sense, but can lead to substantial finite-size corrections that can easily be mistaken for a different universality class.

2. \(T > 0\) scaling

Field-theoretic descriptions using the nonlinear \(\sigma\)-model \([19, 20, 111]\) have given a wealth of predictions also for the \(T > 0\) quantum-critical “fan” extending out into the plane \((g,T)\) from the critical point \((g_c,0)\). Scaling behavior in this extended \(T > 0\) region is an important generic characteristic of quantum phase transitions, in contrast to classical phase transitions with typically very narrow regions of criticality. This aspect of dimerized models has also been investigated in detail in QMC studies \([112, 114]\).

3. Experimental realizations

A Néel–VBL transition of the type discussed above can in principle be experimentally realized in systems consisting of coupled dimers as a function of pressure. The authors are not aware of any material where the phase transition can be crossed in a quasi-2D systems of weakly coupled planes, but a well-studied 3D case is TiCuCl\(_3\) \([112, 116]\). Here neutron scattering shows a transition from a paramagnet to a Néel state at a critical pressure \([117]\), including the analogue of the Higgs boson (longitudinal mode) in the ordered state \([118]\). In this case the values of the couplings and their pressure dependence are not known in detail, but one can still learn a lot from studies of designer Hamiltonians by focusing on universal properties \([119, 123]\).

B. Deconfined Néel–VBS transition in J-Q models

While spontaneously formed VBS states in translationally invariant 2D spin systems have been discussed for more than two decades \([81, 124, 127]\), most of the early work focused on frustrated systems \([125, 127]\), for which it is very difficult to carry out unbiased numerical calculations. The proposal of a DQC point separating the Néel and VBS states motivated detailed numerical studies of this transition, and, thus, construction of suitable designer Hamiltonians amenable to QMC simulations. Numerical work preceding the theory \([128]\) had already indicated the intriguing possibility of continuous magnetic–VBS quantum phase transitions (where standard arguments based on Landau–Ginzburg theory predict generically first-order transitions) in U(1) (quantum XY) models. Tractable SU(2) spin models exhibiting Néel–VBS transitions were lacking, however, until the introduction of the J-Q class of models \([103]\), where \(J\) refers to Heisenberg exchange on a bipartite lattice and \(Q\) is a multi-spin interaction which competes against Néel order but does not generate a sign problem.

The Q interactions are most conveniently expressed in terms of the singlet projectors, Eq. \((7)\), in the simplest case on the 2D square lattice;

\[
H_Q = -Q \sum_{\langle ijkl\rangle} P_{ij} P_{kl},
\]

(19)

where the site pairs \(ij\) and \(kl\) form parallel edges (with both horizontal and vertical arrangements included) on a \(2 \times 2\) plaquette. One can also consider more than two
projectors in the product and denote by $Q_p$ an interaction with $p$ projectors. Fig. 3 illustrates $Q_2$ and $Q_3$ terms leading to columnar VBS ground states. In addition to the number of projectors in the product, they way they are arranged relative to each other on the lattice is also crucial, with different patterns leading to different VBS states that do not support any local dimer fluctuations, while the continuous transitions are into VBSs where such fluctuations exist. Alternatively, one can relate the different behaviors to different types of topological defects in the VBS.

1. $T = 0$ Critical behavior

The critical behavior of both the $J$-$Q_2$ and $J$-$Q_3$ models (with the columnar dimer arrangements in Fig. 3) on the simple 2D square lattice has been analyzed in detail in several different ways. The spin stiffness $\rho_s$ is the quantity of interest to study the destruction of the Neel state by the perturbation $\beta$. It should scale at a critical point as $\rho_s \sim L^{2-d-z}$, where $z$ is the dynamic exponent. The DQC theory is Lorenz-invariant, i.e., $z = 1$. One can test this prediction by graphing $\rho_s(L)$ versus the coupling ratio $J/Q$ for different $L \times L$ lattices (in the ground state or for the inverse temperature $\beta \propto L^\nu$). Such curves should cross at the critical point. The deviations from pure scaling can also be fitted with a conventional multiplicative correction of the form $(1 + a L^{-\omega})$ with large $a$ and small $\omega$.

2. $T > 0$ critical scaling

The quantum-critical “fan” in the $(J/Q, T)$ plane has also been investigated. Initial calculations confirmed that

---

**Fig. 3.** $Q_2$ and $Q_3$ interactions on the square lattice. The bars indicate the locations of singlet projectors $C_{ij}$ on site pairs $ij$. The $Q_2$ and $Q_3$ arrangements in the left and center panels lead to columnar VBS states, while the right $Q_3$ arrangement leads to a staggered VBS. The Hamiltonian contains all unique translations and $90^\circ$ rotations (and in the staggered case also reflections) of the operator patterns.

**Fig. 4.** Finite-size scaling of the spin stiffness of the $J$-$Q_2$ model in the vicinity of its quantum-critical point; from Ref. 76. The calculations were done at inverse temperature $\beta = L$. The standard critical scaling form $\rho_s \sim \text{constant}$ has been modified by a log-correction (with $L_0 = 0.9$) in order to compensate for a weak drift of the crossing points. The red curves, which cross at a single point, the critical point $q_c = (J/Q)_c = 0.0447(2)$, represent a common scaling function for $L \geq 48$. Results for the $J$-$Q_2$ model are graphed in Fig. 4 and discussed next.
the scaling is governed by a $z = 1$ critical point \[36\]. Later studies \[28,114\] noted log-like corrections also here, e.g., while the conventional $z = 1$ scaling implies that the uniform magnetic susceptibility scales linearly in $T$ \[20\], the behavior actually appears to be of the form $\chi \sim T[1 + a \ln(1/T)]$. These corrections, like those at $T = 0$ discussed above, remain an intriguing unexplained aspect of the Néel–VBS transition in the $J$-$Q$ models.

3. Emergent $U(1)$ gauge fluctuations

An important and remarkable aspect of the DQC theory is the emergent conservation law of the gauge flux, due to the irrelevance of monopoles at the critical point. There is a $U(1)$ symmetry associated with this conservation law at the critical point, which is also emergent and not explicitly contained in the $J$-$Q$ model, which only has an $SU(2)$ spin-rotational symmetry, the discrete symmetries of the square lattice and time reversal. The emergent $U(1)$ symmetry can, however, be observed in the fluctuations of the VBS order parameter \[102\]. Naively one would expect these fluctuations to reflect the $Z_4$ symmetry of the VBS (which is dictated by the square lattice), but instead the symmetry group expands to $U(1)$ upon approach to the critical point. The interpretation of this emergent $U(1)$ VBS symmetry is that it reflects the non-compactness of the gauge field (“photon”) of the DQC theory. Here we will review how this emergence is detected in QMC simulations.

Consider the VBS (dimer-dimer) correlation function,

$$C_\alpha(r_{ij}) = \langle B_\alpha(r_i)B_\alpha(r_j) \rangle,$$

(20)

where $r_{ij} = r_i - r_j$ is the spatial separation of the operators $B_\alpha$, $\alpha = \hat{x}, \hat{y}$, which measure the spin correlations on nearest-neighbor bonds in the $\alpha$ direction;

$$B_\alpha(r) = S(r) \cdot S(r+\hat{\alpha}), \quad B_\beta(r) = S(r) \cdot S(r+\hat{\beta}).$$

(21)

We define the expectation value of this operator when averaged over all sites $r = (x,y)$: $b_\alpha = \langle B_\alpha(x,y) \rangle \neq 0$. In a VBS, the expectation value acquires a spatial modulation. With the dimers (stronger bonds) oriented in the $x$ direction, the asymptotic form of the spatially averaged $x$ dimer correlations in a columnar VBS is $C_\alpha(x,y) = b_\alpha^2 + (1 - \xi^2)\frac{Q^2}{D^2 + \alpha e^{-r/\Lambda}}$, where $r = (x^2 + y^2)$, $D$ is the magnitude of the order parameter, and $\alpha$ is a constant. The rate of decay of the oscillating term toward a non-zero value as $r \to \infty$ defines the correlation length $\xi$ (where we suppress the directional dependence of $\xi$, which can be parameterized by $\xi_x$ and $\xi_y$). This is the standard definition of a correlation length, as the length-scale associated with the order parameter. Normally this (the larger of $\xi_x, \xi_y$) is the largest length-scale in the system.

In a symmetry-broken $x$-oriented VBS, we can consider also dimer correlations in the direction $y$ perpendicular to the dimers. The asymptotic form of these correlations is $C_\beta(x,y) = b_\beta^2 + ke^{-r/\Lambda}$, where $k$ is another constant (and again there are really two decay constants: $\Lambda_x$ and $\Lambda_y$). These correlations are, at first sight, not associated with the order parameter, and the length-scale $\Lambda$ would then normally not be referred to as “the correlation length”. However, $\Lambda$ can actually be larger than $\xi$, and this is at the heart of the emergent $U(1)$ symmetry of the VBS.

According to the DQC theory, both lengths diverge as the critical point is approached, and $\Lambda \sim \xi^{1+\epsilon}$ with $\epsilon > 0$. Close to the critical point, where the order parameter $D$ is small and $\Lambda \gg \xi$, on length scales less than $\Lambda$ the system will appear to have both $x$ and $y$ VBS order.

One can also associate $\Lambda$ with fluctuations of the angle of the order parameter. Since the VBS order parameter is a vector, $\mathbf{D} = (D_x, D_y)$, one can define a magnitude $D$ and an angle $\Theta$. $\Lambda$ defines a length scale below which the order parameter exhibits $U(1)$ symmetry—the system not only has both $x$ and $y$ order up to this length scale, but one can think of the angle as fluctuating uniformly, even though it has to take one of the values $n\pi/2$ when the order parameters $D_x$ and $D_y$ of a symmetry-broken state are averaged over regions of size $\gtrsim \Lambda$. There are many interesting consequences of the emergent $U(1)$ symmetry of the VBS order parameter, as further discussed in Ref. \[141\].

This phenomenon of an emergent symmetry higher than that of the order parameter in an ordered state is in fact a general aspect of systems with dangerously irrelevant perturbations. These perturbations reduce the symmetry of a system in such a way that the standard critical exponents at a phase transition into an ordered state are unaffected, but the symmetry broken in the ordered state is reduced \[142,143\]. A prototypical example is the 3D classical XY model with an added on-site potential $h\cos(\theta_i)$ on all sites $i$, where $\theta_i$ is the spin angle and $q$ an integer \[144\]. This model orders at a critical temperature $T_c(h)$, but, regardless of the value of $h$, the critical exponents are fixed at those of the standard XY transition. However, instead of breaking the continuous $U(1)$ symmetry, as the original $h = 0$ model does, now only a $q$-fold ($Z_q$) symmetry is broken. As in the VBS discussed above, there is a length-scale $\Lambda \sim \xi^{1+\epsilon(q)}$ in the ordered state, which is associated with $U(1)$ symmetric fluctuations of a coarse-grained order parameter (magnetization) $\mathbf{m} = (m_x, m_y)$. Finite-size scaling procedures have been developed \[145\] to access the exponent $a(q)$ directly based on the cross-over from $U(1)$ to $Z_q$ symmetry seen in the order-parameter distribution $P(m_x, m_y)$. Such techniques have also been employed to study the VBS fluctuations of the $J$-$Q$ model, as we discuss next.

In simulations of a VBS on a finite lattice, where the $Z_4$ symmetry is not broken, one can study the emergent $U(1)$ and the length-scale $\Lambda$ by accumulating the probability distribution $P(D_x, D_y)$, where $D_x$ and $D_y$ are defined on the whole system in terms of matrix elements of the operators $B_\alpha$ in Eq. \[21\], evaluated for individual configurations using transition-graph loops, as explained in Sec. \[111\] and Fourier transforming at $(\pi,0)$ and $(0,\pi)$ for...
4. Spinon deconfinement

The emergent U(1) length scale discussed above should also be reflected in the confinement of spinons (e.g., the size of a bound state) in the VBS state. By utilizing properties of the valence-bond basis, discussed in Sec. II F, and generalizations to non-zero total magnetization, it is possible to directly access $S = 1$ excitations of VBS states and study their structure in terms of bound or unbound (deconfined) spinons. Such calculations have already been used to characterize deconfined spinons using a designer Hamiltonian with a VBS state in one dimension. Similar calculations for 2D J-Q models should shed further light on the nature of the spinons in the vicinity of the DQC point.

Another approach to confirming spinon excitations, which has already been employed for the 2D J-Q model, is to study the thermodynamic properties at criticality and compare them with predictions for a simple model of a gas of thermally excited bosonic spinons. The results show a remarkable consistency among different quantities which are related through expressions for the spinon gas, thus lending strong support to the deconfinement scenario.

IV. NÉEL-VBS TRANSITION IN SU($N$) SPIN MODELS

In this section we will discuss quantum criticality in SU($N$) spin models on 2D square lattices, as systematic generalizations of the SU(2) systems discussed above. Secs. IV A, IV B, IV C, and IV D describe several studies carried out on a single layer, and Sec. IV E summarizes a recent study on a bilayer.

A. SU($N$) Heisenberg models

A natural extension of the SU(2) Heisenberg model is to spin models with SU($N$) symmetry. Such generalizations were initially introduced to facilitate access to a solvable large-$N$ limit. In the meantime, new physical systems have also motivated an interest in such models at arbitrary finite values of $N$, leading to extensive studies of SU($N$) spin systems. The physical degrees of freedom of a particular spin model depend not only on the global symmetry but also on the chosen representation under which the spins transform. Thus, there are a number of distinct extensions of the SU(2) model to SU($N$), depending on what properties are intended to be retained in the generalization.

In the context of the study of the DQC point at the Néel-VBS transition, the ability of two spins to form a singlet is the most central characteristic required to form a VBS. It is possible to choose SU($N$) models which have this property by working on bipartite lattices and assigning SU($N$) spins that transform as relative conjugate representations on each of the sublattices, as discussed in Sec. II F. In such models two spins on opposite sublattices can form a singlet, allowing for the possibility of a valence-bond solid. The simplest SU($N$) invariant two-site sign-problem free Hamiltonian interaction one can define for such models is given by $H_1$ in Eq. (17), with $P_{ij}$ generalized to the SU($N$) singlet projector defined in Sec. II F. We note again that for $N = 2$ this reduces to the familiar $S = 1/2$ antiferromagnetic Heisenberg model.

The Hamiltonian Eq. (17) has been studied extensively as a function of $N$. On a 1D chain the well known critical Bethe state for $N = 2$ gives way to a VBS ordered state for all $N \geq 3$. The 2D version of $H_1$ was shown in the large-$N$ limit to map onto a quantum dimer model with only a kinetic term. From direct numerical studies of the quantum dimer model, it is known that the kinetic-only model orders into a VBS. This implies that in $H_1$ there must be a transition between the magnetic state at $N = 2$ and the VBS at $N = \infty$. The location of the transition was first determined to lie between $N = 4$ and $N = 5$ by QMC simulations.
B. SU(N) $J_1$-$Q$ model for $N \leq 4$

In the previous section we saw that on the square lattice the Néel-order found for $N = 2, 3, 4$ gives way to VBS order for $N \geq 5$ in the SU($N$) antiferromagnet, Eq. (17). In order to access the quantum phase transition by QMC simulations at a given value of $N \leq 4$ one needs new sign-problem free “designer” couplings that destroy the Néel phase and result in the VBS. The first such term discovered was the $Q$-interaction, Eq. (19), introduced for SU(2) in Sec. [111]. It can be generalized for any $N$, with $P_{ij}$ again the SU($N$) singlet projectors. Since the matrix elements of $P_{ij}$ are all positive, such a $Q$ coupling is free of the QMC sign problem.

It turns out that the $Q$-only model, with the projectors arranged as in Fig. 3 is always VBS ordered, so that for the cases of $N = 2, 3, 4$, the $J_1$-$Q$ model has a Néel-VBS transition [102, 157]. Detailed QMC studies for these $N$ find strong evidence for a continuous transition between the Néel and VBS ground states [156, 158]. As discussed above for the SU(2) case, certain observables show deviations from standard scaling laws for the system sizes studied, that could either be due to conventional corrections to scaling or actual violations of the scaling laws (but they do not resemble violations of the kind that would be expected at a first-order transition [76]). The origin of these corrections remains to be resolved.

C. SU($N$) $J_1$-$J_2$ model for $N \geq 5$

For $N \geq 5$, the $Q$ term strengthens the VBS that is already present in the $J_1$ only model, and hence to study the Néel-VBS transition for large-$N$, a new designer Hamiltonian is needed. To supply such a model, an interaction $J_2$ can be included between sites on the same sublattice which stabilizes the Néel state [54]:

$$H_2 = -J_2 \sum_{\langle i,j \rangle} \Pi_{ij},$$

where the $\Pi_{ij}$ interaction was introduced in Eqs. (12) and (13). In order to see that this interaction favors the Néel state, we note that in the $H_2$ model by itself, the two sublattices are decoupled. The ground state corresponds to having an independent SU($N$) ferromagnet on each sublattice. Turning on a small $J_1 \ll J_2$ interaction will clearly cause the sublattice degeneracy to be lifted, resulting in the two independent ferromagnets to lock into a single antiferromagnetic state. These arguments are true independent of the value of $N$. Thus, for every $N \geq 5$ the VBS state is realized when $J_1 \gg J_2$ and the Néel state is obtained when $J_2 \gg J_1$. Thus, there must be at least one quantum phase transition as the ratio $J_2/J_1$ is tuned. From numerical simulations of the $J_1$-$J_2$ model, there is compelling evidence for a direct transition between the two phases, with no indication of an intervening phase [54].

D. $J_1$-$J_2$-$Q$ model; anomalous dimensions

One can also combine the $J_1$-$J_2$ and $J$-$Q$ model. The phase diagram of the resulting $J_1$-$J_2$-$Q$ model as a function of $N$ is summarized in Fig. 6. So far, the phase boundaries have only been computed in the two perpendicular planes shown in the figures. We here discuss some aspects of the critical behavior for $N \in \{2,3,\ldots,12\}$ obtained in these calculations and relate the results to large-$N$ expansions within the DQC theory.

An important prediction of the DQC scenario [22] at the Néel-VBS transition is that both order parameters are simultaneously quantum critical at the phase transition and that space and time scale in the same way, i.e., the dynamic critical exponent $z = 1$. This implies that both correlation functions decay as Lorentz-invariant power laws,

$$C_{N,V}(r,\tau) \sim \frac{1}{(r^2 + c^2\tau^2)^{(1+\eta_{N,V})/2}}.$$  

where $C_N$ and $C_V$ are the two point correlation functions of the Néel and VBS order parameters. The indices $\eta_{N}$ and $\eta_{V}$ are the so-called anomalous dimensions of the Néel and VBS order parameters. The DQC scenario predicts that the continuum field-theoretic universality of the Néel-VBS critical point is described by the non-compact CP$^{N-1}$ universality. Quantitative estimates for the universal indices that characterize this universality class are available only in the large-$N$ limit [159].

![FIG. 6. Phase Diagram of the SU($N$) symmetric sign-problem free $J_1$-$J_2$-$Q$ model as a function of $N$. Each of the couplings has been introduced in the text: the generalized nearest-neighbor antiferromagnetic exchange $J_1$ in Sec. IV A, the four-spin coupling $Q = Q_2$ in Sec. IV B and Fig. 3 and the generalized ferromagnetic next-nearest-neighbor coupling $J_2$ in Sec. IV C. This SU($N$) antiferromagnet allows for an unbiased study of deconfined quantum criticality at the Néel-VBS transition for each value of $N$.](image)
We note here that as $N \to \infty$, $\eta_N \to 1$ and $\eta_V \to \infty$. Both results are very unusual, since typically $\eta$-exponents are small. For reference, in the $O(N)$ model $\eta \to 0$ in the $N \to \infty$ limit and $\eta \approx 0.037$ for the $O(N=3)$ model.\(^{162}\)

We now turn back to the study of the lattice designer $J_1$-$J_2$-$Q$ Hamiltonian. By analyzing the size dependence of the correlation functions in Eq. (23) at the location of the critical points shown in Fig. 6, it is possible to estimate values for $\eta_N$ and $\eta_V$ as a function of $N$.\(^{160,137}\) Results for $2 \leq N \leq 12$ are summarized in Fig. 7. The lattice Hamiltonians reproduce the “smoking gun” DQC feature that $\eta_N \to 1$ and $\eta_V \to \infty$ in the $N \to \infty$ limit. Furthermore, extrapolations of the data, with the constant terms subtracted off, show quantitative agreement, within a few percent, with the corrections in the large-$N$ expansions, Eq. (24). Note that the correct description for large $N$ is far from trivial, since the continuum theory was not derived from the microscopic Hamiltonians considered here but was posited as a generic theory of the Néel–VBS transition.

The fact that the exponents agree so well lends strong positive support in favor of the DQC scenario for large $N$. Clearly, it would be very interesting to compute further $1/N$ corrections analytically, to test the theory in this quantitative way down to the smallest $N$. We note that the exponent $\eta_N$ for the $SU(2)$ case has been estimated based on classical Monte Carlo simulations of the non-compact $CP^1$ model, and also of the 3D Heisenberg model with “hedgehog” topological defects suppressed (which should lead to the same universality).\(^{101}\) The exponents are in rough agreement with those of the $SU(2)$ $J$-$Q$ model, but it would be very interesting to push these classical simulations to higher precision as well. Another study of a classical model argued to realize the DQC action was found to have a first-order transition.\(^{163}\)

### E. SU($N$) Bilayer

The study of the destruction of SU($N$) magnetic order in the bilayer geometry, illustrated in Fig. 8(a), provides an interesting example to test our understanding of the role of Berry phases at DQC points, since these should cancel between the two square lattice layers. Consider a bilayer in which each layer is described by some combination of the $J_1$-$J_2$ interactions discussed in the previous

![FIG. 8. Physics of the SU($N$) bilayer model. (a) Lattice and couplings. White sites form the A sublattice with SU($N$) spins that transform in the fundamental representation. Black sites on the B sublattice transform as the conjugate to this representation. (b) Phase diagram of the SU($6$) bilayer model as a function of $g_\perp = J_\perp/J_1$ and $g_2 = J_2/J_1$. The black circle at $g_\perp = 0$ is the Néel-VBS quantum critical point. Remarkably, in the bilayer geometry with $g_\perp \neq 0$ this continuous transition becomes first-order (indicated by open circles), due to the cancellation of Berry phases between the layers. The VBL phase at large $g_\perp$ is a featureless paramagnet with singlets predominantly forming between the planes. Cartoons of the basic singlet physics in the symmetry-broken VBS (left) and uniform VBL (right) phases are shown. The SU($N$) Néel-VBL transition is discussed at length in Ref. 164.](image-url)
section and with the following coupling between the layers,

$$H_{\perp} = -J_{\perp} \sum_{[ij]} P_{ij}, \quad (25)$$

where the sum over $[ij]$ is taken between spins vertically next to each other in the bilayer (i.e., $[ij]$ form dimers, as

in Sec. IIIA). For any $N$, the $J_{\perp}$-only model has a

featureless non-degenerate ground state consisting of a product of vertical valence bonds (singlets)—the SU($N$)

generalization of the VBL state discussed in Sec. IIIA.

The introduction of small intra-layer couplings like the

$J_1$-$J_2$ interactions will make the ground state deviate from a simple product state by increasing the amount of entanglement, but these couplings cannot destabilize

the VBL paramagnet, as long as they are small compared to $J_{\perp}$; see Fig. 8(b).

1. Néel-VBL transition

When the value of $J_2/J_1$ places the system within the

Néel phase when $J_{\perp} = 0$, there is a transition from the

Néel state into the VBL for increasing $J_{\perp}$. This transition in the SU(2) case is continuous and belongs to the 3D O(3) universality class, as discussed in Sec. IIIA. For $N > 3$, QMC calculations [164] show that the transition is first-order, however, as also predicted by a mean-field theory. For $N = 3$ the transition appears continuous based on the largest system sizes studied. If this is indeed the case, it would be in the universality class of the compact CP$^2$ model [10]. Further studies of the $N = 3$ model should be carried out to resolve this issue.

2. Néel-VBS transition

A very interesting aspect of the SU($N$) bilayer models is the limit where $J_{\perp}$ is weak and there is a Néel-VBS transition as a function of $J_2/J_1$. It is well known that the Berry phases cancel in the long-wavelength limit between the layers. Nevertheless, both Néel and VBS phases must clearly be stable to a small finite interlayer coupling. What is the fate of the $J_{\perp} = 0$ deconfined Néel-VBS transition in the presence of the bilayer coupling? Remarkably, the QMC simulations show that the transition between the very same Néel and VBS phases becomes first-order in the bilayer geometry. This finding can be understood as a restoration of the Landau paradigm, due to the cancellation of Berry phases which is a relevant perturbation at the deconfined critical point [164]. This is another piece of strong evidence for the correctness of the DQC theory, as well as the more general field-theoretical understanding of the role of Berry phases.

V. SPIN LIQUIDS AND DECONFINEMENT IN U(1) MODELS

One of the main foci of research into designer Hamiltonians is the search, detection and characterization of quantum spin liquid (QSL) phases. That is, although there has been much progress in understanding properties of QSL states from effective field theories, there are very few microscopic models amenable to unbiased calculations that can be argued to exhibit a QSL without controversy. The recent spate of high-profile numerical studies identifying possible QSL candidate Hamiltonians [163, 167] has sparked intense activities on the theory side, highlighting the synergy between analytical theory and numerics. Thus, we are interested in finding the simplest models harboring QSL states, that are also amenable to large-scale simulation by QMC.

Much of the reason for the scarcity of microscopic models is that fundamental ingredients required to give rise to QSL phases, namely geometrical frustration, are also typical harbingers of the sign problem. Despite this, the absence of the sign problem does not fundamentally preclude the existence of spin liquid physics. This is most evident in Hamiltonians with U(1) symmetry—anisotropic spin (or Bose-Hubbard) models specifically designed to capture the essential ingredients needed to promote a QSL phase without causing the sign-problem. A crucial fact in this regard is that there are no limitations on the diagonal couplings, because the sign problem is only caused by the off-diagonal matrix elements in the chosen QMC basis (see Sec. II). Thus, diagonal frustration can be simulated in any form.

Examples of both 2D gapped [168–171] and 3D and gapless [172] QSLs have recently been studied with large-scale QMC simulations of such sign-problem free U(1) models. In this section, we will explore one class of Hamiltonians that contains the simplest type of QSL phase, which can be related to a $Z_2$ gauge theory. We will demonstrate that universal topological properties calculated in QMC can be compared to this effective field-theoretic description. In addition, an associated critical point separating an XY-ferromagnet and a QSL can be demonstrated to be of an exotic “fractionalized” type, based on comparisons of universal quantities calculated from QMC and a related quantum field theory.

A. BFG Hamiltonians with $Z_2$ spin liquid phases

One successful recipe for designing Hamiltonians with QSL phases amenable to QMC study was pioneered by Isakov and co-workers for certain Bose-Hubbard models on the kagome lattice [168, 169, 171]. Models in this class, first proposed analytically by Balents, Fisher and Girvin [172] (BFG), are constructed partly based on their relation to quantum dimer models (QDMs), where a strong local constraint (the number of dimers emanating from a site being fixed) is coupled with a quantum tunneling
that allows the state to fluctuate, while keeping the constraint satisfied. QDM Hamiltonians can be constructed where the groundstate superposition of dimer configurations breaks no symmetry—being a type of QSL (or resonating valence bond) phase, that has an emergent gauge symmetry and related topological order [174].

The recipe to construct QSL phases in XXZ spin (or boson) models is to map the dimer constraint to a plaquette spin configuration on the dual lattice; the dimer may represent an $S^z$ spin configuration on a bond which is frustrated (or unsatisfied). Then, the role of the quantum dimer tunneling moves is done by a spin-exchange processes (which may not be a simple two-body term). Such models have been called cluster-charging models [169], because spin configurations summed over a local cluster (e.g. a lattice plaquette) are penalized when differing from some chosen value. For example, on the kagome lattice (Fig. 9),

$$H_0 = V \sum_{\mathcal{O}} (S^z_{\mathcal{O}})^2,$$  \hspace{1cm} (26)

where

$$S^z_{\mathcal{O}} = \sum_{i \in \mathcal{O}} S^z_i$$  \hspace{1cm} (27)

is a cluster-charging potential term which favors three spin up and three spins down per hexagonal plaquette. This Hamiltonian alone will promote a highly degenerate manifold of groundstate configurations, where each kagome-lattice hexagon satisfies the plaquette constraint but is otherwise disordered. Like all operators which are entirely diagonal in the chosen QMC basis, $H_0$ does not affect the simulation with a sign problem, no matter how the interaction is frustrated, making it a suitable ingredient for a designer Hamiltonian.

In order to promote a QSL phase from this classical degenerate manifold, one desires to add quantum fluctuations to the Hamiltonian such that the cluster-charging constraint is not violated. In this case, a ring-exchange term,

$$H_{\text{ring}} = -K \sum_{i \in \mathcal{R}} (S^+_i S^-_j S^+_k S^-_l + h.c.),$$  \hspace{1cm} (28)

which operates around a “bow-tie” plaquette on the kagome lattice (Fig. 9), promotes quantum fluctuations while preserving the cluster-charging constraint $H_0$. Thus, one expects the kagome lattice model with $H = H_0 + H_{\text{ring}}$ to retain a disordered groundstate with quantum fluctuations—a good candidate for a quantum spin liquid phase. In addition, with $K > 0$, such terms do not have the sign problem for QMC [175].

BFG [173] were the first to show that variations of this kagome-lattice model support the simplest type of $Z_2$ spin liquid—in particular, at an exactly soluble Rokhsar-Kivelson point [131]. Subsequently, using QMC simulations, Isakov and co-workers showed that a variant of the cluster-charging Hamiltonian with different short-range spin exchanges, support a $Z_2$ spin liquid phase. This exchange term can either connect the first, second and third neighbors $(ij)_3$ on the kagome lattice [168, 169]; or it can be nearest-neighbor only, $(ij)_1$ [171]. Again, for $t_{1,3} > 0$ no sign problem exists. In addition to $H = H_0 + H_3$ and $H = H_0 + H_1$, the $Z_2$ spin liquid has been demonstrated on $H = H_1 + H_{\text{ring}}$, where the cluster-charging term is not explicitly required and the Hamiltonian consists of two competing kinetic-energy terms [170], sometimes called the J-K model [173].

In the next section, we explore one procedure by which QMC simulations can positively identify such spin liquid phases. In section 4.3 we examine the XY-ferromagnet (or “superfluid” in the boson language) to QSL phase transition, which is common to these models, as a different example of a deconfined or fractionalized quantum critical point.

B. Identifying the $Z_2$ spin liquid with entanglement

The smoking gun signature for a gapped spin liquid phase could be argued to be the identification of an emergent gauge theory, and the associated topological order [174]. One manifestation is the existence of a ground state degeneracy, which is topological in nature, in the simplest ($Z_2$) case being four-fold on a torus. Previous QMC studies of QSL states have demonstrated the difficulty in clearly identifying this degeneracy, due to the
tendency for tunneling between the equal-energy states on finite-size lattices\cite{165}. Similarly, recent DMRG work studying candidate gapped spin liquid states have been unable to identify the expected topological degeneracy\cite{166, 167}. For example, the\textit{Levin-Wen} construction\cite{179} isolates 2\(\gamma\) from Eq. (31), requiring measurement of the equal time\textit{correlation function} \(\langle \hat{G}(\tau) \hat{G}(0) \rangle\) is modulated not by the physical bosons (which would undergo a quantum phase transition in the 3D XY universality class), but by the fractional spinon fields\cite{183–185}. This leads to the same exponents \(z\) and \(\nu\) as in the ordinary XY critical point. However, the exponent \(\eta\) controlling the equal time correlation function \(\langle \hat{b}^\dagger(0)\hat{b}(x) \rangle\) is modified. In a way similar to the DQC point discussed in Sec. IV.C \(\eta = 1\) in the \(N \rightarrow 1\) limit and is expected to be anomalously large also for \(N = 1\).

Previous estimates for \(\eta\) were obtained from field-theoretic treatments. From an \(1/N\) expansion, Ref. \cite{184} obtains
\[
\eta = 1 + \frac{32}{3\pi^2N},
\]
which gives \(\eta = 2.08\) for \(N = 1\). A more accurate value was obtained from a combination of field theory and Monte Carlo simulations of the correlation of a composite operator in the 3D XY model\cite{186, 187}, leading to \(\eta \approx 1.47(3)\).

The measurement of \(\eta\) in the kagome BFG model is challenging, requiring measurement the equal time Green’s function in real space: \(G(x) \equiv G(\tau = 0, r) = \)

1. The boundary law

Luckily, there is another tool suited for the positive identification of this topological order, through measurements of the Renyi entanglement entropy, introduced in section II.C. The quantities \(S_\alpha\) are well studied in quantum information science; they quantify correlations for a system in a basis-independent way, which makes them suitable in particular for characterizing phases that do not have an explicit broken symmetry (i.e., QSL phases). In fact, some authors have championed the entanglement entropy as a paradigmatic analog to symmetry breaking not having an explicit broken symmetry (i.e., QSL phases).

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of the Hamiltonian $H$ logical entanglement entropy in the scales as $1/L$. Inset: finite-size scaling of one expect another temperature crossover to $2\ln(2)$ (dashed line) identifies the emergent gauge symmetry; kagome lattices [171]. The quantization of the plateau at the XY quantum phase transitions in this model. Namely, since the fractional spinons are bosonic, they can be modeled by a “simulation within a simulation”, achieved by projecting physical bosons at an XY transition to wind strictly in pairs around the space- and time-periodic simulation cell. Since this distribution is universal, it is expected to be the same (up to a non-universal velocity) as the winding number distribution of the physical (composite) bosons in the $XY^*$ transition. Observation of this fact [189] is a striking confirmation that topological properties can still be present even when fractional excitations are gapless. These universal winding number distributions are possible to compute in principle in field theory; hence they are another example of the synergetic connection between and quantum field theory and QMC simulations on designer Hamiltonians.

VI. DISCUSSION

We have presented several examples of the use of judiciously constructed designer Hamiltonians for testing field theories and exploring various quantum many-body phenomena related to quantum-criticality and exotic ground states. Here we discuss some further generic aspects of this approach, followed by a brief discussion of other interesting related studies of systems not covered in this Review, and an outlook on future directions.

A. Designer Hamiltonian, stoquasticity, and the sign problem

The concept of designer Hamiltonians that we have introduced here is different from that of “stoquastic” Hamiltonians [190], which are those for which all off-diagonal matrix elements are non-positive (i.e., sign-problem free) in the standard basis. This is normally the $z$-component basis for spin systems, although a broader definition with respect to an arbitrary basis has also been presented [191], which would seem to indicate that any Hamiltonian could in principle be cast into the stoquastic class. Our criterion is more one of scientific utility than a strict mathematical definition—models that are free from sign problems either automatically (in a practically useful basis), or through some other way of circumventing it (e.g., the Meron algorithm [55] or determinant-based QMC of particle-hole symmetric Hubbard models [49,192,193]). To be considered a designer Hamiltonian, such a model should also represent a prototypical case of some interesting quantum-man body phenomenon which is difficult to access in an unbiased manner in other ways, i.e., the model is designed to shed light on this problem (and must be “de-signed” to be practically useful for studies on large scale).

1. Limitations of designer Hamiltonians

An important fundamental problem is whether it is always possible to construct such designer Hamiltonians, or whether certain types of states are beyond reach in practice because their physical properties are fundamentally tied to the difficulties in circumventing the sign problem.
For instance, the “Bose metal” spin-liquid phase with a Fermi sea of emergent fermions appears to have a complex sign structure that would be difficult to reproduce in a sign-free Hamiltonian in a simple basis [194]. On the other hand, as we have reviewed in Sec. IV A, $Z_2$ spin-liquid ground states have been found in models with U(1) symmetry. An interesting open challenge is to find the $Z_2$ spin liquid (which itself is a broad class if states [195]) in sign-free SU(2) invariant models. It is certainly possible that these states cannot be realized with sign-problem free Hamiltonians in a simple basis, but one can still not exclude that designer Hamiltonians can be constructed for which some useful basis can be found, or for which the sign problem can be solved in some other way. Indeed, the recent discovery of a spin liquid in the Hubbard model on the honeycomb lattice [166] seems to provide an example, although it remains to be seen whether this state really is a $Z_2$ topological state.

2. Computational complexity

Regarding the sign problem itself more broadly, Ref. [196] presented a proof that the sign problem cannot be solved in general. The problem here was defined in terms of constructing a QMC algorithm for which the computational effort scales as a power-law in the system size. A particular problem is considered for which the removal of the negative signs still leaves a problem for which the Monte Carlo sampling is believed to scale exponentially in the system size and, thus, the sign problem was not solved, only recast into a different form. However, the model system chosen in this demonstration was that of a spin glass, and the conclusion reached is equivalent to the statement that the quantum spin glass is no easier to solve than the corresponding classical spin glass. This is hardly a surprising result. The proof does not address QMC solutions of systems that are not directly associated with the well known difficulties of simulating classical frustrated “glassy” systems with complicated energy landscapes. Whether or not the sign problem can be circumvented (at least for representative designer Hamiltonians) for translationally invariant systems such as fermionic Hubbard models without particle-hole symmetry and SU(2) frustrated quantum spins models remains an important open question.

Some non-trivial solutions of sign problems are worth mentioning. In some cases, local basis transformations can be used to render all off diagonal matrix elements non-positive [197]. One can sometimes apply more sophisticated ways to circumvent the problem, e.g., with the Meron algorithm [52] or the newly discovered fermion bag approach [198]. It remains to be seen whether these exiting developments can be applied to a wider range of fermionic and frustrated designer Hamiltonians.

B. Other systems and future prospects

1. Spin models

The models and quantum states we have discussed here only represent a small sample of recent research into fascinating topics in quantum magnetism. Other noteworthy examples include systems exhibiting Bose-Einstein condensation of magnons [199], spin supersolids [200], quadrupolar order [201, 202], and quantum compass models [203]. Here as well, QMC studies of designer Hamiltonians played a crucial role in uncovering novel physics, and unresolved issues will require the kind of synergistic interplay between low-energy theories and QMC simulations that we have emphasized in this Review.

Within the classes of spin models we have discussed here, although the essential physics now has been ascertained, there still remains important work to be done to improve on the quantitative aspects, e.g., the critical exponents. In the case of the studies aiming to connect SU(N) QMC simulations with large-N expansions, summarized in Sec. V D and Fig. 7 it would be very useful to go to still larger lattices, to check the convergence of the computed exponents more precisely. This is important, in particular, in light of the fact there are significant scaling corrections in quantities such as the spin stiffness for $N = 2$. Such corrections could potentially also affect the fitting procedures underlying the exponents graphed in Fig. 7 for small $N$. When higher-order 1/$N$ analytical results eventually become available, it will be important to know the values of the exponents without possible remaining effects of scaling corrections. What is needed here is high-accuracy (small error bars) results for the correlation functions on large lattices. It will also be very interesting to use the methods discussed in Sec. II C to quantify the degree of entanglement in the SU(2) and SU(N) models.

2. Fermions

Another very interesting aspect of the designer Hamiltonians we have discussed here is that they can also be doped with fermions. While this introduces a sign problem, it should be possible to study large lattices doped with a small number of holes, since the magnitude of the sign problem only depends on the number of fermions. This low-density limit of the SU(N) models should already be sufficient for making connections with large-N theories of “holonic” or “strange” metallic states, that have been predicted to arise out of the DQC system upon doping [160].

An interesting conceptual question is whether one can study bosonic systems in which fermions are emergent particles. A positive answer may be suggested by the demonstrated existence of the $Z_2$ fractional phase in a purely bosonic model, as we discussed here in Sec. V B. The finite-temperature structure of the topological en-
tanglement entropy identifies that both a $Z_2$ charge (a spinon) and a $Z_2$ flux (a vison) are realized as excitations in this model $^{171}$. This raises the possibility that, with a suitable “binding” potential $^{204}$ coupling these two, an extension of the model with an emergent fermionic excitation might be created $^{170}$. The ability for QMC to measure the entanglement entropy in a state with an emergent fermion could also have wide implications for the purported application of tensor network algorithms as a “solution” to the sign problem, if such a state were found to violate the boundary law, Eq. (30).

3. Disordered systems and impurities

In this Review we have focused on translationally invariant systems. Systems with disorder, i.e., random couplings, dilution, etc., is another area where designer Hamiltonians can provide unique insights. To mention one example of a spin system for which the standard classical–quantum correspondence fails, in the 2D $S = 1/2$ Heisenberg antiferromagnet diluted with non-magnetic impurities, the interplay of classical percolation and quantum fluctuations lead to low-energy excitations $^{147}$ not captured $^{205}$ within the conventional $d \rightarrow (d + 1)$-dimensional mapping. Other unanticipated effects of dilution on quantum-critical scaling have also been observed that still lack explanation and deserve further study $^{206,210}$. For the designer Hamiltonians discussed in this Review, disorder effects have not yet been studied. One can anticipate interesting behaviors in VBS states, at the Néel–VBS transition, and in the $Z_2$ spin liquids.

Beyond disordered spin models, various manifestations of the “dirty boson” problem remain intriguing and challenging $^{211,213}$, and provide excellent examples of the indispensable role of QMC studies of designer Hamiltonians $^{214}$. An example is the extended hardcore Bose-Hubbard model with random frustrated interactions on a 3D cubic lattice. QMC simulations have been instrumental in discovering that glassiness can coexist with superfluidity, and that quantum fluctuations and random frustration are both crucial ingredients needed to stabilize a “superglass” state $^{215,216}$.

In addition to extensive disorder, single-impurity problems are also interesting and important. Effects of a single missing or added spin have been investigated with QMC in the 2D and 3D Néel states $^{217,219}$ and at conventional quantum-critical points in dimerized models $^{220}$. QMC studies have also been done on vacancies in the VBS state of the 2D $J$-$Q$ model $^{221}$, where one can observe a vortex in the order parameter, and at the SU(2) and SU(3) critical points $^{138,158}$, where impurities provide an interesting way to probe the anomalously large corrections to scaling in these systems. Impurities in 1D chains (with open edges) have also been studied $^{222}$.

In general, impurities and disorder in interacting quantum systems present rich opportunities for further exploring exotic states and phase transitions. Synergistic combinations of low-energy field theories, RG schemes for impurities $^{223}$ and disorder $^{213,224,226}$, and QMC studies of designer Hamiltonians will be needed to make progress here.

4. Non-equilibrium dynamic scaling

A recent development on the methods front is the adaptation of QMC algorithms such as those discussed in Sec. II to non-equilibrium setups $^{14}$. By studying time evolution out of a non-equilibrium state in imaginary time, standard sign-problem free systems can be analyzed in novel ways. For instance, it was found in Ref. $^{14}$ that the dynamic scaling properties when quenching to a quantum-critical point is the same in real and imaginary time, and that such protocols are useful for extracting the dynamic critical exponent. Moreover, the fidelity and full geometric tensor, which characterize the geometrical properties of the state space, can be accessed this way. One can anticipate many fruitful applications of this unconventional QMC approach, which gives access to dynamics beyond the commonly used numerical analytic continuation of QMC-computed equilibrium imaginary-time correlation functions $^{227}$.

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[1] S. Sachdev, Quantum Phase Transitions (Cambridge University Press, 2011).

[2] J. L. Cardy, ed., Finite-Size Scaling (North Holland, 1988).

[3] P. M. Chaikin and T. C. Lubensky, Principles of Condensed Matter Physics (Cambridge University Press, 2000).
O. I. Motrunich and T. Senthil, Phys. Rev. Lett. 89, 277004 (2002).
T. Vojta and J. Schmalian, Phys. Rev. Lett. 95, 237206 (2005).
R. Yu, T. Roscilde, and S. Haas, Phys. Rev. B 73, 064406 (2006).
T. Vojta and J. Schmalian, Phys. Rev. Lett. 95, 237206 (2005).
R. Yu, T. Roscilde, and S. Haas, Phys. Rev. B 73, 064406 (2006).
A. W. Sandvik, Phys. Rev. Lett. 96, 207201 (2006).
T. Roscilde and S. Haas, Phys. Rev. Lett. 99, 047205 (2007).
D.-X. Yao, J. Gustafsson, E. W. Carlson, and A. W. Sandvik, Phys. Rev. B 82, 172409 (2010).
L. Pollet, N. V. Prokof'ev, B. V. Svistunov, and M. Troyer, Phys. Rev. Lett. 103, 140402 (2009).
H. Meier and M. Wallin, Phys. Rev. Lett. 108, 055701 (2012).
S. Iyer, D. Pekker, and G. Refael, Phys. Rev. B 85, 094202 (2012).
V. Gurarie, L. Pollet, N. V. Prokof'ev, B. V. Svistunov, and M. Troyer, Phys. Rev. B 80, 214519 (2009).
K.-M. Tam, S. Geraedts, S. Inglis, M. J. P. Gingras, and R. G. Melko, Phys. Rev. Lett. 104, 215301 (2010).
D. Larson and Y.-J. Kao, arXiv:1202.3908 (2012).
K. H. Höglund and A. W. Sandvik, Phys. Rev. B 70, 024406 (2004).
F. Anfuso and S. Eggert, Phys. Rev. Lett. 96, 017204 (2006).
C.-W. Liu, S. Liu, Y.-J. Kao, A. L. Chernyshev, and A. W. Sandvik, Phys. Rev. Lett. 102, 167201 (2009).
K. H. Höglund, A. W. Sandvik, and S. Sachdev, Phys. Rev. Lett. 98, 087203 (2007).
R. K. Kaul, R. G. Melko, M. A. Metlitski, and S. Sachdev, Phys. Rev. Lett. 101, 187206 (2008).
S. Sanyal, A. Banerjee, and K. Damle, Phys. Rev. B 84, 235129 (2011).
M. Vojta, Phys. Rev. B 85, 115113 (2012).
J. A. Hoyos and T. Vojta, Phys. Rev. Lett. 100, 240601 (2008).
T. Vojta, C. Kotabage, and J. A. Hoyos, Phys. Rev. B 79, 024401 (2009).
E. Altman, Y. Kafri, A. Polkovnikov, and G. Refael, Phys. Rev. B 81, 174528 (2010).
O. F. Syljuåsen, Phys. Rev. B 78, 174429 (2008).