Computational Studies of Quantum Spin Systems

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Abstract. These lecture notes introduce quantum spin systems and several computational methods for studying their ground-state and finite-temperature properties. Symmetry-breaking and critical phenomena are first discussed in the simpler setting of Monte Carlo studies of classical spin systems, to illustrate finite-size scaling at continuous and first-order phase transitions. Exact diagonalization and quantum Monte Carlo (stochastic series expansion) algorithms and their computer implementations are then discussed in detail. Applications of the methods are illustrated by results for some of the most essential models in quantum magnetism, such as the $S = 1/2$ Heisenberg antiferromagnet in one and two dimensions, as well as extended models useful for studying quantum phase transitions between antiferromagnetic and magnetically disordered states.


Keywords: Quantum spin system, antiferromagnet, valence-bond solid, quantum phase transition, finite-size scaling, quantum Monte Carlo, exact diagonalization, Lanczos method

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1. INTRODUCTION

One of the primary goals of theoretical physics is to provide the simplest models capturing various complex physical phenomena. In condensed matter physics, as well as in statistical mechanics more broadly, spin systems often serve in this role. As the Ising model and other idealized classical spin models have been invaluable in forming our understanding of thermal phase transitions and critical phenomena, so are various quantum spin models now instrumental in developing a theoretical framework for exotic quantum many-body states and quantum phase transitions, i.e., phase transitions driven by quantum fluctuations (controlled by some tunable interaction parameter) at temperature $T = 0$ \[1, 2\]. With their many possible ordered and disordered ground states and different types of excitations arising from them, quantum spin systems also provide rich opportunities to study other manifestations of collective quantum behavior \[3, 4\].

Although often used as simplified prototypical model systems for various phenomena, and not always intended for describing fully all the details of specific real materials, quantum spin systems have also been very successful in explaining quantitatively the antiferromagnetic properties of a variety of Mott insulators with localized electronic spins. A prime example of this is provided by the undoped parent compounds of the high-temperature superconductors and other related quasi-two-dimensional and quasi-one-dimensional copper oxides. Their experimentally measured magnetic response functions can be remarkably well reproduced by the $S = 1/2$ Heisenberg model on two-dimensional (2D) planes \[5, 6\], isolated chains \[7, 8\], and “ladders” \[9\] consisting of two or more coupled chains. In agreement with model calculations, 2D layered systems exhibit an exponentially divergent correlation length as $T$ is lowered (until ordering sets in below some critical temperature due to 3D couplings or anisotropies), while chain and ladder compounds exhibit only short-range (power-law or exponentially decaying) correlations. In addition to cuprates, many other inorganic and organic antiferromagnets also show similarly good agreement between theory and experiments \[10\].

A prominent research theme in contemporary condensed matter physics is to model and explain \textit{magnetically disordered ground states} of 2D or quasi-2D materials with non-uniform or frustrated (competing) antiferromagnetic interactions \[11, 12, 13, 14\]. Quantum phase transitions in 2D spin systems challenge the classical Ginzburg-Landau framework \[15\] for understanding and classifying phase transitions based on order parameters, as exemplified by the recent theory of “deconfined” quantum critical points \[2, 16\], which separate antiferromagnetic (Néel) and non-magnetic valence-bond solid (VBS) ground states \[17\]. In a field-theory proposed to describe this quantum phase transition (the non-compact CP$^1$ model), deconfined spinons (collective $S = 1/2$ degrees of freedom) are the “elementary particles”, out of which the two order parameters can be formed due to condensation (in the Néel state) or confinement (in a VBS state) \[16\]. Apart from the interest in such unusual phase transitions in condensed matter physics, there are also intriguing connections to deconfinement in gauge theories in particle physics \[18\]. Interacting quantum spins have also recently become interesting in the context of ultra-cold atoms in optical lattices \[19, 20\], as well as in quantum information theory \[21\]. Fundamental many-body concepts such as entanglement entropy \[22\] are currently explored in various ground states of quantum spin systems \[23\].

Exact solutions of quantum spin systems are very rare beyond one dimension, where
there are several important cases (enough to fill a whole encyclopedia [24], in fact). In
two dimensions there are also some examples [25, 26], but normally analytical calculations
rely on approximations or assumptions that cannot be rigorously justified. Purely
computational studies of model hamiltonians are therefore also essential. Unbiased nu-
merical results are important for testing theories and analytical calculations (in par-
icular, continuum field theories for the low-energy physics). Moreover, numerical “simula-
tions” can also in their own right serve as laboratories for exploration and discovery, and
may thus stimulate further theoretical and experimental developments.

In classical statistical physics, almost any model can be studied in detail using Monte
Carlo or molecular dynamics simulations (although there are also challenging classical
systems, e.g., ones with very slow, “glassy” dynamics [27]). The situation is different
in quantum mechanics. There are still enormous hurdles limiting computational studies
of generic quantum spin hamiltonians, especially ones with frustrated interactions and,
going beyond pure spin models, strongly correlated fermion systems. Devising efficient
practically useful numerical algorithms for these types of systems is one of the greatest
challenges in theoretical physics. Thanks to a series of significant developments over the
past couple of decades, large-scale computational studies have already become possible
for some important classes of quantum lattice models. Very large 1D systems can be
studied using the density matrix renormalization group (DMRG) method [28, 29] or
related methods formulated using matrix-product states [21, 30]. Quantum Monte Carlo
(QMC) methods with loop-cluster updates [31, 32, 33] can be used to study a wide
range of spin and boson models in any number of dimensions, typically on lattices with
up $10^4$ sites or more in the ground state, and much larger still at elevated temperatures.
In addition to breakthroughs in efficient algorithms, the impressive improvements in
computer performance have of course played an important role in recent progress, too.
The most dramatic gains have, however, been achieved as a result of better algorithms,
and there is reason to believe that this will continue to be the case in the future as well.

Topics. Two classes of computational methods will be discussed in these lecture
notes: Numerical (exact) diagonalization and quantum Monte Carlo simulation. Exact
diagonalization methods will be developed primarily for 1D systems, followed by some
discussion of extensions to 2D square-lattice systems. The use of symmetries for block-
diagonalization will be developed and used in both complete diagonalization ($T > 0$
calculations) and with the Lanczos method (for obtaining the ground state and low-
energy excitations in a given symmetry sector). QMC simulations based on the series
expansion of the partition function (stochastic series expansion; SSE) will be developed
for $T > 0$ calculations (and applied also in the limit $T \to 0$). Computer programs written
in close correspondence with the pseudocodes are available on-line [34].

Beyond describing the technical aspects of the numerical methods, an integral goal of
these lecture notes is to introduce the most essential quantum spin models (hamiltonians)
and to present some of their physical properties from a computational perspective. While
the discussion is largely self-contained as far as the algorithms and implementations are
concerned, the physics of the systems is for the most part discussed in a rather “light”
fashion, in the form of elementary calculations (e.g., spin-wave theory) and qualitative
descriptions illustrated by numerical results. Connections to complementary analytical
approaches (e.g., predictions based on field theories) are also pointed out, with key
references for further study. The topics range in maturity from well established basics to very recent and ongoing research on exotic quantum phase transitions.

Outline. The quantum spin models to be discussed in the subsequent sections are first introduced in Sec. 2, along with a brief summary of various types of ground states and quantum phase transitions. Classical phase transitions, Monte Carlo simulations, and finite-size scaling techniques are reviewed Sec. 3 in order to set the stage for quantum-mechanical finite-lattice calculations and data analysis. Exact diagonalization techniques and their applications to 1D spin systems are discussed in Sec. 4. Basic properties of the Heisenberg chain and its extension with frustrated interactions are illustrated with numerical results (including the frustration-driven quantum phase transition into a dimerized VBS state). Extensions of the methods to 2D systems are also summarized, and used to study the low-energy states (quantum-rotor states) of small antiferromagnetic systems. Sec. 5 begins with a general discussion of path integrals, followed by the alternative series-expansion formulation of quantum statistical mechanics, on which the SSE QMC method is based. The SSE method is then developed in detail for the \( S = 1/2 \) Heisenberg model. Illustrative results for chains, ladders, and 2D planes are presented, including a study of quantum-criticality in dimerized 2D systems. Applications of the SSE method to “J-Q” models with four- and six-spin interactions are also discussed, and the Néel–VBS transitions occurring in these systems as a function of the strength of the multi-spin interactions are studied. Sec. 6 concludes with a brief survey of other recent works related to the topics of the lecture notes.

2. QUANTUM SPIN MODELS, THEIR GROUND STATES AND QUANTUM PHASE TRANSITIONS

In solid-state physics, quantum spin hamiltonians describe the effective magnetic interactions between localized electronic spins. As such, they can be derived starting from the full problem of interacting electrons [35, 36, 37]. Here we will not discuss their relationships with real materials in detail, but just introduce some well established models with Heisenberg couplings and other, similar interactions. To motivate and prepare for the quantitative numerical calculations in the later sections, in this introductory part we will first survey some of the possible types of ground states and quantum phase transitions. We will discuss the nature of the quantum fluctuations in spin systems from different perspectives, including spin wave theory and singlet pairing (valence bonds).

The Heisenberg exchange is the most important spin-spin interaction and forms the starting point for understanding many materials and phenomena in quantum magnetism. Two spins are coupled according to the hamiltonian

\[
H_{ij} = J_{ij} \mathbf{S}_i \cdot \mathbf{S}_j = J_{ij} (S^x_i S^x_j + S^y_i S^y_j + S^z_i S^z_j).
\]

(1)

Often this pair interaction is summed over only nearest-neighbor sites \((i, j)\), but longer-range interactions can also be included. The type of ground state, the nature of the excitations, and the finite-temperature properties of a system with Heisenberg interactions depend strongly on the underlying lattice. The dimensionality plays a crucial role. According to the Mermin-Wagner-Hohenberg theorem [38, 39], a continuous symmetry
of a quantum system with short-range interactions, here the global SU(2) spin rotation symmetry, can be broken neither at $T \geq 0$ in one dimension nor at $T > 0$ in two dimensions. This normally rules out magnetic order in 1D Heisenberg models (unless there are long-range interactions, in which case the theorem does not apply—we will discuss an example of this in Sec. 4.3.3), but in two dimensions the ground state can be magnetic, i.e., $\langle S_i \rangle \neq 0$ (with all these vector expectation values parallel in a ferromagnet and alternating between two opposite directions in a bipartite antiferromagnet). Beyond dimensionality, the microscopic details of the lattice and the coupling strengths $J_{ij}$ (e.g., uniform, modulated in some periodic pattern, or in some random, disordered way) are also decisive, and many different types of ground states and quantum phase transitions can be realized. Some of these states and transitions are still not very well understood and subjects of ongoing research.

Apart from a brief review of spin wave theory for general $S$, we will in these lecture notes focus on the simplest case of $S = 1/2$ spins, corresponding to individual uncompensated electronic spins. This is often the most interesting case, as $S \to \infty$ is the classical limit, and $S = 1/2$, thus, maximizes the effects of quantum fluctuations (although in some cases, $S = 1$ or higher can actually lead to even larger quantum effects, e.g., in the case of the “Haldane state” of the $S = 1$ chain [26]). We will only consider antiferromagnetic interactions, $J_{ij} > 0$ in (1), which from a theoretical perspective are more interesting than ferromagnetic couplings. Antiferromagnetic interactions in strongly-correlated systems are also more prevalent in nature. A much broader area of quantum many-body physics can be entered by also allowing anisotropies in spin space, i.e., different $x$, $y$, and $z$ coupling strengths in (1). In addition to the relevance of such anisotropies in many real magnetic materials, the mapping between $S = 1/2$ spins and “hard-core” bosons makes such models interesting also for other reasons. The spin-isotropic Heisenberg interaction can be considered the essence of quantum magnetism, however, and we will focus almost exclusively on this case here.

**Section Outline.** After discussing the antiferromagnetic (Néel) state and the nature of its quantum fluctuations based on spin wave theory in Sec. 2.1, two important classes of non-magnetic states—spin liquids and valence-bond solids—will be discussed in Sec. 2.2. In Sec. 2.3 the special properties of 1D systems are briefly reviewed. Sec. 2.4 introduces the extended quantum Heisenberg models and the types of quantum phase transitions that we will study in much more detail in the later sections, in applications of the various computational methods.

### 2.1. The Néel state and its quantum fluctuations

On the 2D square lattice (and other bipartite 2D and 3D lattices with uniform interactions) the ground state of the Heisenberg model with only nearest-neighbor interactions is antiferromagnetically (Néel) ordered, with neighboring spins being oriented, on average, in an antiparallel (staggered) fashion. Note that a maximally ordered antiferromagnetic state, e.g., $|\uparrow \downarrow \uparrow \downarrow, \ldots \rangle$ on a chain (or a checkerboard pattern of $\uparrow$ and $\downarrow$ spins on the 2D square lattice), is not an eigenstate of the Heisenberg hamiltonian, whereas a
fully polarized ferromagnetic state, e.g., $|↑↑↑⟩$, is an eigenstate (and, in the case of ferromagnetic interactions, has the minimum energy). This can be easily seen with the pair interaction (1) written as

$$H_{ij} = J_{ij}(S_i^x S_j^x + S_i^y S_j^y + S_i^z S_j^z) = J_{ij} [S_i^z S_j^z + \frac{1}{2}(S_i^+ S_j^- + S_i^- S_j^+)].$$

When acting on the perfect Néel state, the raising and lowering operators flip pairs of spins, causing local defects, whereas they destroy the perfect ferromagnetic state. The antiferromagnetic order must therefore always be reduced by quantum fluctuations, whereas the fully polarized ferromagnetic state is the ground state also in the presence of the off-diagonal interactions. The amount of magnetic order (if any) remaining in the true ground state of a system with antiferromagnetic interactions depends sensitively on details of the lattice and the interactions included in the Hamiltonian.

Note again that the magnetic order parameter of a system with Heisenberg interactions (e.g., the magnetization of a ferromagnet or the sublattice magnetization of an antiferromagnet) is a vector in spin space. The sublattice magnetization operator is

$$m_s = \frac{1}{N} \sum_{i=1}^{N} \phi_i S_i,$$

where $\phi_i = \pm 1$ is the staggered phase factor, e.g., on the 2D square lattice $\phi_i = (-1)^{x_i+y_i}$, where $x_i$ and $y_i$ are the lattice (integer) coordinates of site $i$. In a Néel state the expectation value $\langle m_s \rangle = m_s^x \hat{x} + m_s^y \hat{y} + m_s^z \hat{z}$ is non-zero in the thermodynamic limit. The order can form—the spin-rotational symmetry can be broken—in any direction in spin space. For convenience one normally associates the $z$ spin components with this direction, so that the staggered magnetization $\langle m_s \rangle = |\langle S_i^z \rangle|$. In a finite system a non-zero magnitude $m_s$ of the Néel order can form, even though the direction of the vector remains fluctuating over all angles and, thus, $\langle m_s \rangle = 0$. In a calculation for a finite system one should therefore detect the presence of order using a quantity which is independent of the direction, e.g., $\langle m_s^2 \rangle$ or $|\langle m_s \rangle|$. We will see many examples of finite-lattice calculations later. First, let us discuss some of the basic properties of the symmetry-broken Néel state in more detail.

### 2.1.1. Spin wave theory

The Néel state and its excitations in 2D and 3D systems can be understood within a simple linear spin wave theory (discussed in more detail in many standard works, e.g., the review article by Manousakis [6] and the book by Mattis [36]). Such a calculation starts from a maximally ordered state of staggered spins, which is the exact ground state in the classical limit where the spin magnitude $S \rightarrow \infty$. This is regarded as a vacuum state, on top of which quantum effects are included systematically by adding bosons, representing the deviations of the spins from $|S_i^z| = S$, in such a way as to obtain a good approximation to the ground state of the system for finite $S$.

The relationship between spins and bosons for $S = 1/2$ is illustrated in Fig. 1. The reason for using this mapping is that it is easier to carry out calculations with the bosons, due to their simpler commutation relations. Linear spin wave theory corresponds to non-interacting bosons and is, by construction, exact for $S \rightarrow \infty$. Results for finite $S$ can be
FIGURE 1. Correspondence between $S = 1/2$ spins and bosons, here illustrated for a 1D chain. The fully staggered reference state (left) is the vacuum for bosons (with – indicating zero boson occupation). For each spin flipped with respect to the reference state there is a boson (•) at the corresponding site (right). For $S = 1/2$ the boson occupation numbers are 0 or 1 on each site, while for arbitrary spins they are 0, …, $2S$. Linear spin wave theory corresponds to non-interacting bosons, and the occupation number constraints are not enforced. Higher-order calculations in $1/S$ gradually restore the constraints.

systematically improved by including interactions in the form of an $1/S$ expansion. Here we just outline the lowest-order calculation.

If we neglect the constraint that the boson number $n_i$ for each site should be within the range $0, \ldots, 2S$ (the physical subspace), we can use the following simple (lowest order in $1/S$) mapping between the spin operators in the original Hamiltonian (2) and boson creation and destruction operators $a_i^+$ and $a_i$ (and the number operators $n_i = a_i^+ a_i$):\(^1\)

\[
\begin{align*}
  i & \in \uparrow \text{ sublattice} : & S^+_i &= S - n_i, \\
  & & S^-_i &= \sqrt{2S} a_i, \\
  j & \in \downarrow \text{ sublattice} : & S^+_j &= n_j - S, \\
  & & S^-_j &= \sqrt{2S} a_j.
\end{align*}
\]

(4)

It is useful to look at Fig. 1 to understand this mapping—apart from the obvious way in which the off-diagonal operators can affect the states, one just has to be careful with the different factors associated with boson creation/destruction and spin raising/lowering (discussed in standard quantum-mechanics texts). The factors in (4) are correct for $n_i \ll S$, but note that they are also exact for $S = 1/2$ in the physical subspace. In principle, one can write down more complicated expressions that are formally correct for any $S$ and $n_i$ in the physical subspace (and also automatically decouple the physical and unphysical subspaces), but these are only relevant when going beyond linear spin wave theory.

We now transform the terms in the Hamiltonian (2) using the mapping (4). Because we are considering a bipartite system, where the two sites $i$ and $j$ are always on different sublattices, we obtain the off-diagonal term

\[
\frac{1}{2} (S^+_i S^-_j + S^-_i S^+_j) \rightarrow S (a_i a_j + a_i^+ a_j^+),
\]

(5)

and the diagonal interaction is

\[
S^+_i S^-_j \rightarrow -S^2 + S(n_i + n_j) - n_i n_j.
\]

(6)

spin wave theory should formally be regarded as a large-$S$ calculation. In the lowest-order calculation we should therefore discard the interaction term $n_i n_j$ in (6), because it is a factor $1/S$ smaller than the non-interacting contributions from (5) and (6). Let us

\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*\*

1 Often two species of bosons, $a_i$ and $b_j$, are used, corresponding to the two sublattices. In momentum space, the use of a single boson species implies that we are here working in the full Brillouin zone of the original lattice with $N$ momenta, instead of one with $N/2$ momenta corresponding to a two-site unit cell.
for definiteness consider the two-dimensional square lattice with \( N = L^2 \) sites. We then have the effective hamiltonian
\[
H = -2NS^2 J + 4SJ \sum_{i=1}^{N} n_i + SJ \sum_{\langle ij \rangle} (a_i a_j + a_i^+ a_j^+).
\] (7)

Here it should be noted that we consider a finite system but assume that the symmetry is broken (the direction of the staggered magnetization has been locked) which can be strictly true only in the limit \( N \to \infty \) (unless some symmetry-breaking field is added to \( H \)). This is fine, however, because we will anyway take the limit \( N \to \infty \) at the end.

The boson hamiltonian (7) can be easily diagonalized (i.e., written in terms of number operators). To construct this solution, we first Fourier transform,
\[
a_k = N^{-1/2} \sum_{\mathbf{r}} e^{i \mathbf{k} \cdot \mathbf{r}} a_{\mathbf{r}}, \quad a_{\mathbf{r}} = N^{-1/2} \sum_{\mathbf{k}} e^{-i \mathbf{k} \cdot \mathbf{r}} a_k,
\] (8)

where the real-space operators have been labeled by their lattice coordinate vectors \( \mathbf{r} \) instead of just the site index \( i \), and the momentum \( \mathbf{k} \) is in the first Brillouin zone of the square lattice (i.e., the reciprocal square lattice of \( N \) sites). The hamiltonian is then
\[
H = -2NS^2 J + 4SJ \sum_{\mathbf{k}} n_k + 2SJ \sum_{\langle \mathbf{i} \mathbf{j} \rangle} \gamma_k (a_k a_{-\mathbf{k}} + a_k^+ a_{-\mathbf{k}}^+),
\] (9)

where, with the lattice constant set to 1 (i.e., \( k_x, k_y = n2\pi/L, n \in \{0, \ldots, L-1\} \)),
\[
\gamma_k = \frac{1}{2} [\cos(k_x) + \cos(k_y)].
\] (10)

The next step is to carry out a Bogolubov transformation to boson operators mixing the original \( +\mathbf{k} \) and \( -\mathbf{k} \) operators;
\[
\alpha_{\mathbf{k}} = \cosh(\Theta_k) a_{\mathbf{k}} + \sinh(\Theta_k) a_{-\mathbf{k}}^+,
\] (11)

which has the inverse
\[
a_{\mathbf{k}} = \cosh(\Theta_k) \alpha_{\mathbf{k}} - \sinh(\Theta_k) \alpha_{-\mathbf{k}}^+.
\] (12)

It is easy to verify that the operators \( \alpha_{\mathbf{k}} \) obey the standard bosonic commutation relations for any \( \Theta_k \). The trick is to choose these “angles” for each \( \mathbf{k} \) such that all operators of the form \( \alpha_{\mathbf{k}} \alpha_{-\mathbf{k}} \) and \( \alpha_{\mathbf{k}}^+ \alpha_{-\mathbf{k}}^+ \) cancel out in the hamiltonian (9). This is the case if
\[
\frac{2 \cosh(\Theta_k) \sinh(\Theta_k)}{\cosh^2(\Theta_k) + \sinh^2(\Theta_k)} = \gamma_k.
\] (13)

The Bogolubov-transformed hamiltonian (the spin wave hamiltonian) is then diagonal in the occupation numbers;
\[
H = E_0 + \sum_{\mathbf{k}} \omega_k \alpha_{\mathbf{k}}^+ \alpha_{\mathbf{k}},
\] (14)

where, after some algebra making use of Eq. (10), the constant can be written as
\[
E_0 = -2SJ \sum_{\mathbf{k}} \frac{\gamma_k^2}{1 + \sqrt{1 - \gamma_k^2}} - 2NS^2 J,
\] (15)
and the dispersion relation in (14) for the spin wave states $\alpha^+_k |0\rangle$ is given by

$$\omega_k = 4SJ \sqrt{1 - \gamma_k^2}. \quad (16)$$

For momenta close to $(0, 0)$ and $(\pi, \pi)$, this dispersion is linear, $\omega_k = ck$ and $\omega_k = c|\langle \pi, \pi \rangle - k|$, respectively, with velocity (the spin wave velocity) $c = 2\sqrt{2S}J$.

The ground state $|0\rangle$ of (14) is the vacuum for spin waves, where the energy is just $E_0$ given by (15). The sum can be evaluated numerically, most easily by a straight-forward summation over the momenta on large lattices, and extrapolating $E_0/N$ to $N \to \infty$ (or converting the sum divided by $N$ to an integral, the numerical evaluation of which gives the $N = \infty$ value directly). The result is $E_0/JN = -0.65795$ for $S = 1/2$.

Note that while the ground state does not contain any Bogolubov $\alpha$-bosons (spin waves), it does contain some amount of the original $\alpha$-bosons. The sublattice magnetization is directly related to the density of these bosons, which is uniform and can be computed at any site or averaged over the sites:

$$\langle m_s \rangle = S - \langle 0|a_i^+ a_i|0\rangle = S - \frac{1}{N} \sum_{i=1}^N \langle 0|a_i^+ a_i|0\rangle. \quad (17)$$

Using the Bogolubov transformation, this becomes

$$\langle m_s \rangle = S - \frac{1}{N} \sum_k \sinh^2(\Theta_k). \quad (18)$$

In the most interesting case of $S = 1/2$, this evaluates to $\langle m_s \rangle = 0.3034$, or $\approx 61\%$ of the “classical” value $1/2$. Thus, the quantum effects (zero-point fluctuations, represented by the presence of bosons) reduce, but do not destroy, the long-range order.

In principle it is not clear whether spin wave theory should be reliable for small $S$. There has been much discussion of this issue, but the method does in fact give a good description of the 2D Heisenberg model on the square lattice. As we will see later, unbiased QMC calculations give results for $E_0$ and $\langle m_s \rangle$ differing from those quoted above by only $1 - 2\%$. This can be traced, a posteriori, to the true value of $\langle m_s \rangle$ being quite large (i.e., spin wave theory can be expected to be accurate when the density of bosons is low). In cases where the true sublattice magnetization is small or vanishes, spin wave theory normally does not work that well, even when going to higher orders in $1/S$ (which increases the complexity of the calculation very significantly [40, 41, 42]).

### 2.1.2. Destruction of the Néel order

When going beyond bipartite lattices with uniform interactions, or by supplementing the Heisenberg model with additional interactions (e.g., including more than two spins) the quantum fluctuations can become so significant that the ground state loses its long-range Néel order (retaining only short-range antiferromagnetic correlations). There are several other possible types of ordered and disordered ground states, some of which have
no classical counterparts. Much of the current interest in quantum spin models is related
to the existence of non-magnetic states and quantum phase transitions between them
and the Néel state [2]. This is also the main theme of the numerical calculations to be
discussed in these lecture notes. One long-standing motivation for studying such transi-
tions stems from the cuprate high-\(T_c\) superconductors, the undoped parent compounds
of which are antiferromagnets corresponding closely to weakly coupled Heisenberg lay-
ers (many properties of which can be understood based on a single layer) [6]. In these
systems the magnetic order is destroyed upon doping by mobile charge carriers. This is
a very challenging electronic many-body problem, where computational studies are also
playing an important role [43, 44, 45]. While the full solution of the high-\(T_c\) problem
will of course require more complicated models (perhaps some variety of the t-J or Hub-
bard model), some generic aspects of the physics close to a quantum phase transition
out of the Néel state can, however, be understood based on spin-only models [46].

Apart from the cuprates and related antiferromagnetic systems, there are also many
materials with non-uniform or frustrated spin interactions [11, 12, 13], which can lead
to non-magnetic low-temperature states. Many of these states, and the possible quantum
phase transitions between them, are still not well understood. It is therefore useful
to search for and study prototypical quantum spin models that realize various types of
ground states and quantum phase transitions. Studies of quantum phase transitions also
have a broader context of understanding “exotic” manifestations of quantum mechanics
at the collective many-body level [2]. There are even interesting connections with par-
ticle physics—close analogies exist between phase transitions in supersymmetric gauge
theories in 2+1 dimensions and “deconfined” quantum-critical points that may separate
the 2D Néel and VBS ground states [18].

2.2. Spin liquids and valence-bond solids

Within linear spin wave theory, a non-magnetic state corresponds to boson density
\(\langle n_i \rangle = S\). This kind of state is, however, not a good representation of actual non-magnetic
ground states of Heisenberg and related quantum spin models, because it does not
contain any correlation effects. To discuss more interesting non-magnetic states, it is
useful to first look at the quantum fluctuations from a different perspective.

For two isolated spins \(i, j\) (a dimer, \(N = 2\)), the ground state of the \(S = 1/2\) antiferro-
magnetic Heisenberg hamiltonian (1) is the singlet;

\[
|\phi_{ij}^s\rangle = \frac{|\uparrow_i\downarrow_j\rangle - |\downarrow_i\uparrow_j\rangle}{\sqrt{2}}.
\] (19)

Although the two spins in such a singlet are always perfectly anti-correlated (entangled),
the individual spins are strongly (maximally) fluctuating, and there is no static spin
order; \(\langle S_i \rangle = \langle S_j \rangle = 0\). In contrast, the perfect Néel states for \(N = 2\), \(|\uparrow_i\downarrow_j\rangle\) and \(|\downarrow_i\uparrow_j\rangle\), are
product states (i.e., of the form \(|\phi_i\rangle \otimes |\phi_j\rangle\)) with no fluctuations (and no entanglement—
loosely speaking, the degree of entanglement corresponds to the deviations from a
product state). Note that for \(N = 2\) the (eigen) energy of the singlet is \(-3J_{ij}/4\), whereas
the expectation value of the energy in the Néel states is much higher, \(-J_{ij}/4\) (and the
FIGURE 2. Valence-bond states in two dimensions. The thick lines represent singlets. The arrows show a spin configuration compatible with the valence bonds (i.e., the spins on each bond are antiparallel—the valence-bond state is a superposition of all such spin configurations). A valence-bond state with only short bonds (or any superposition of such states), as shown here in the extreme case of all bonds of length one, has no magnetic order in two dimensions. There can, however, be order in the valence bonds. (a) and (b) represent typical configurations of a disordered RVB spin liquid and a columnar VBS, respectively.

states are not eigenstates). The tendency of interacting spins to entangle by forming pairwise singlets to minimize the energy remains in multi-spin systems, but when \( N > 2 \) a spin cannot simultaneously form pure singlet pairs with all its neighbors. The system can instead be thought of as a superposition of different singlet pairings. No pair is then in a pure singlet, and the energy contribution from each interaction \( H_{ij} \) is therefore always higher than the singlet energy \(-3J_{ij}/4\). This can be regarded as a reduction of quantum fluctuations (leading to less than maximal two-spin entanglement) for \( N > 2 \), bringing the state (or, more correctly, the density matrix) of each interacting pair closer to a product state. A state with antiferromagnetic long-range order, breaking the rotational invariance of the hamiltonian, can form in the thermodynamic limit if there are enough fluctuations in the singlet pairings (and note that, from the perspective of singlets, the Néel state has larger fluctuations than, e.g., a state of fixed singlet pairings—to be precise, fluctuations should always be specified with respect to some reference state). If the system is one-dimensional, or if the interactions are strongly frustrated (competing), or if the lattice geometry and couplings favor the formation of singlets in a specific pattern (e.g., in a system of weakly coupled dimers), antiferromagnetic order may not be present in the ground state when \( N \rightarrow \infty \).

Valence-bond states. The above intuitive picture of a state of fluctuating singlets can in fact be made rigorous. Any singlet state can be expanded in basis states that are products of singlet pairs, or valence bonds. Denoting by \((i, j)\) a singlet of spins \(i\) and \(j\), as in Eq. (19), a normalized valence-bond basis state for \(N\) (even) spins is of the form

\[
|\phi\rangle = N^{-1/4} |(i_1, j_1)(i_2, j_2)\cdots(i_{N/2}, j_{N/2})\rangle,
\]

(20)

where each site index appears exactly once (i.e., each spin belongs to one singlet). This basis is over-complete in the singlet subspace, and, thus, any singlet state can be expanded in these states, but the expansion coefficients are not unique. The valence-bond basis and computational methods using it are discussed in Refs. [47, 48, 49, 50]. Here we just note that some types of states are more naturally expressed in the valence-bond basis than in the standard basis of individual \(\uparrow\) and \(\downarrow\) spins.

For the valence-bond basis to be (over-)complete, arbitrary lengths of the bonds must be allowed. Restricting the lengths renders the basis incomplete (although one can actually restrict the bonds to only connect sites on two different sublattices). Some types
of states are still completely dominated by short bonds (i.e., the probability of a bond of length \( r \) decreases rapidly with \( r \)). Such short-bond states in two dimensions have no magnetic long-range order (while in three dimensions they can be Néel ordered) and are often called spin liquids or resonating valence-bond (RVB) states. Various types of crystalline order can also form in the bond configurations, leading to periodic modulations in observables such as \( \langle \mathbf{S}_i \cdot \mathbf{S}_j \rangle \) (where \( i, j \) are nearest-neighbor sites). Such ordered states (which break lattice symmetries) are called valence-bond solids (VBSs), or valence-bond crystals. Representative configurations of valence bonds in RVB and VBS states are illustrated in Fig. 2.

2.3. One-dimensional systems

1D systems are rather special and deserve their own introduction. Studies of quantum spin chains date back to Bethe’s exact solution of the \( S = 1/2 \) Heisenberg model, which was published in 1931 [51] and worked out in greater detail some time thereafter [52]. The solution is very complicated (often requiring complex numerical calculations [53, 54]), however, and many properties of the heisenberg chain were only obtained much later and with complementary methods (in particular, renormalization-group treatments of effective low-energy field-theories [55, 56, 57, 58, 59]).

As we have already noted, there can be no magnetic order in a 1D Heisenberg system (but VBS order is allowed, since it breaks a discrete symmetry). The spin correlation function \( C(r) = \langle \mathbf{S}_i \cdot \mathbf{S}_{i+r} \rangle \) of the Heisenberg chain decays with the distance \( r \) as \( (-1)^r/r \) (with a multiplicative logarithmic correction, which we will discuss later). Thus, the ground state is critical (or quasi long-range ordered, on the verge of ordering). Including a frustrated second-nearest-neighbor interaction causes, when sufficiently strong, a quantum phase transition into a VBS state (some times also called the spin-Pierls state), where the spin correlations decay exponentially with distance [60].

While quasi-1D antiferromagnets were actively studied experimentally already in the 1960s and 70s, these efforts were further stimulated by theoretical developments in the 1980s. Haldane conjectured [55], based on a field-theory approach, that the Heisenberg chain has completely different physical properties for integer spin (\( S = 1, 2, \ldots \)) and “half-odd integer” spin (\( S = 1/2, 3/2, \ldots \)). It was known from Bethe’s solution that the \( S = 1/2 \) chain has a gapless excitation spectrum (related to the power-law decaying spin correlations). Haldane suggested the possibility of the \( S = 1 \) chain instead having a ground state with exponentially decaying correlations and a gap to all excitations; a kind of spin liquid state [26]. This was counter to the expectation (based on, e.g., spin wave theory) that increasing \( S \) should increase the tendency to ordering. Haldane’s conjecture stimulated intense research activities, theoretical as well as experimental, on the \( S = 1 \) Heisenberg chain and 1D systems more broadly. There is now completely conclusive evidence from numerical studies that Haldane was right [61, 62, 63]. Experimentally, there are also a number of quasi-one-dimensional \( S = 1/2 \) [64] and \( S = 1 \) [65] (and also larger \( S \) [66]) compounds which show the predicted differences in the excitation spectrum. A rather complete and compelling theory of spin-\( S \) Heisenberg chains has emerged (and includes also the VBS transitions for half-odd integer \( S \)), but even to this
date various aspects of their unusual properties are still being worked out [67]. There are also many other variants of spin chains, which are also attracting a lot of theoretical and experimental attention (e.g., systems including various anisotropies, external fields [68], higher-order interactions [69], couplings to phonons [70, 71], long-range interactions [72, 73], etc.). In Sec. 4 we will use exact diagonalization methods to study the $S = 1/2$ Heisenberg chain, as well as the extended variant with frustrated interactions (and also including long-range interactions). In Sec. 5 we will investigate longer chains using the SSE QMC method. We will also study ladder-systems consisting of several coupled chains [9], which, for an even number of chains, have properties similar to the Haldane state (i.e., exponentially decaying spin correlations and gapped excitations).

### 2.4. Models with quantum phase transitions in two dimensions

The existence of different types of ground states implies that phase transitions can occur in a system at $T = 0$ as some parameter in the hamiltonian is varied (which experimentally can be achieved in quantum magnets, e.g., as a function of pressure or external magnetic field [74]—it should be noted, however, that the possibilities are more limited than in models). As in classical, temperature driven phase transitions, such quantum phase transitions can be continuous or first-order, as illustrated in Fig. 3. Normally a phase transition is associated with an order parameter, which is zero in the disordered phase and non-zero in the ordered phase. e.g., the magnetization $m$ of a ferromagnet or the sublattice magnetization $m_s$ of an antiferromagnet. In Secs. 4 and 5 we will also discuss examples of order-order (Néel-VBS) transitions. Continuous phase transitions are associated with scaling and universality, which we will discuss in more detail in Sec. 3. In these lecture notes we will not discuss topological phase transitions, which are not associated with any local order parameter. The different states across such transitions are distinguishable only through some global topological quantity [76].

In this section we introduce some spin models exhibiting quantum phase transitions in two dimensions. As we have already seen, the 2D $S = 1/2$ Heisenberg model with nearest-neighbor interactions has Néel order at $T = 0$. The question is then how to destroy this order and create some different kind of ground state. Specific studies based on QMC calculations will be presented in Sec. 5. The discussion here serves as an introduction to the kind of models and physical quantities that we will have in mind when
FIGURE 4. Dimerized systems with two different coupling strengths between nearest neighbors; a bilayer (a) with the dimers across the layers and single layers with columnar (b) and staggered (c) dimers. 

discussing computational techniques. Some QMC results will be shown for illustration purposes already in this section, but at this point we do not need to worry about how they were obtained—it suffices to say that the data are numerically exact (to within statistical errors that are in most cases too small to be discerned in the figures). 

There are also 1D analogues of most of the transitions discussed here and some of them will be discussed in Sec. 4.3. Systems in three dimensions can exhibit transitions similar to those in two dimensions, but most computational research is currently focused on 1D and 2D systems, partially because 3D systems are much more challenging. The quantum fluctuations are normally also more prominent in 1D and 2D systems, and some of the most interesting open questions related to real materials are associated with the quasi-2D nature of the systems (although there are also interesting 3D systems [74]).

2.4.1. Dimerized systems

The perhaps simplest way to obtain a non-magnetic ground state of a 2D Heisenberg model is to dimerize the system [75], i.e., to introduce weak and strong antiferromagnetic couplings (bonds), \( J_1 > 0 \) and \( J_2 > J_1 > 0 \), respectively, in a pattern such that each spin belongs exactly to one strong bond. There are several ways to do this, three of which are illustrated in Fig. 4. When \( J_1 = 0 \), these systems consist of \( N/2 \) independent pairs of spins (dimers) with intra-dimer coupling \( J_2 \), and, as we have already discussed, the ground state of each such a pair is a singlet; thus the ground state of the whole system is a singlet-product (valence-bond) state, which clearly has no magnetic order. On the other hand, for \( J_2 = J_1 \) the ground state has Néel order. The question is then how the ground state evolves as a function of the coupling ratio \( g = J_2/J_1 \). One might perhaps think that some amount of Néel order should appear once the dimers are coupled, i.e., for any \( g < \infty \). It turns out, however, that there is actually a phase transition at some critical value \( g_c \) (which depends on the dimer arrangement). While this transition can be analyzed using several analytical and numerical approaches, finite-size scaling of QMC data is currently the only way to obtain unbiased quantitative results (see, e.g., [77] for a discussion of how spin wave theory breaks down close to the phase transition).

Fig. 5 shows some QMC results for the columnar dimer model of Fig. 4(b). The order parameter is the staggered magnetization, with the corresponding operator defined in Eq. (3). This is a vector operator, and, for a finite lattice, its expectation value vanishes due to the spin-rotational symmetry of the hamiltonian. Its square, \( \langle m_z^2 \rangle \), was computed in the simulations. Fig. 5(a) shows results for \( L \times L \) lattices versus the coupling ratio
**FIGURE 5.** QMC results for the squared sublattice magnetization in the two-dimensional Heisenberg model with columnar dimerization. (a) shows results versus the coupling ratio \( g \) for different lattice sizes and (b) shows the size dependence for several values of \( g \). A quantum phase transition where the Néel order vanishes occurs at \( g \approx 1.9 \).

\( g \), and Fig. 5(b) shows data for several values of \( g \) graphed versus \( 1/L \) (which is often the most convenient way of graphing data when examining the convergence to a non-zero value for \( L \to \infty \)). Here it is clear that the behavior changes at \( g \approx 1.9 \); below this coupling the sublattice magnetization extrapolates to a non-zero value in the thermodynamic limit, whereas for larger \( g \) it decays to zero.

In the Néel state, the leading finite-size corrections to \( \langle m_s^2 \rangle \) are \( \propto 1/L \) (a result which can be obtained using spin wave theory \([78, 79]\)). This can be seen quite clearly in Fig. 5(b) for \( g = 1 \) and 1.5. The extrapolation for \( g = 1 \) (see Ref. [50], from which the results for this coupling are taken) gives \( \langle m_s \rangle = 0.3074 \). This is only about 1% higher than the linear spin wave result, \( \langle m_s \rangle = 0.3034 \). As discussed above in Sec. 2.1.1, the success of spin wave theory for this model can be explained *a posteriori* by the fact that the quantum fluctuations are rather weak, reducing the sublattice magnetization only by about 40% from the classical value. For larger \( g \), the agreement between spin wave theory and QMC results quickly becomes much worse. In this kind of dimerized model linear spin wave theory typically over-estimates the critical \( g \) \([80, 81, 77]\), while in other cases, such as a system with uniform couplings with different strengths in the \( x \) and \( y \) lattice directions (coupled chains) linear spin wave theory predicts a critical point at non-zero coupling when, in fact, there is none \([82]\). Various improvements can be made of the spin wave theory (going to higher order in \( 1/S \), incorporating self-consistency, etc. \([6, 40, 41, 80, 77, 83]\)), but a system at or close to a quantum phase transition cannot be captured correctly by these approximative methods.

An important question regarding the quantum phase transition in dimerized systems is its *universality class* (a concept to be discussed further in Sec. 3). A quantum many-body system in \( d \) dimensions can formally be mapped, using path integrals, onto an effective classical system in \( 2+1 \) dimensions (as we will discuss in Sec. 5), and effective continuum field-theories can be constructed for the low-energy behavior, including the quantum phase transition. Many properties have been predicted in this way based on renormalization-group treatments of one such field theory—the nonlinear \( \sigma \)-model in \( 2+1 \) dimensions \([5, 84]\). Based on symmetry arguments alone, one would then expect
the transition to be in the universality class of the 3D classical Heisenberg model. There are, however, subtle issues in the quantum-classical mapping, and QMC simulations are therefore needed to test various predictions. We will see examples of such comparisons between results of simulations and field theories in Sec. 5. While results for the transition in the bilayer (a) [85] and columnar dimer (b) [86] systems in Fig. 4 (and several other cases [87, 88]) are in good agreement with the expectations, recent studies of the staggered dimers (c) show unexpected deviations [89] that are still not understood.

2.4.2. Frustrated systems

The prototypical example of frustration is a system with antiferromagnetic interactions on a triangular lattice. Looking at this problem first within the Ising model, the spins on a single triangle cannot simultaneously be anti-parallel to both their neighbors—there are six configurations with minimum energy, and these all have one “frustrated” bond (two parallel neighbors), as shown in Fig. 6. Being a consequence of the lattice, this is often referred to as geometric frustration. Upon increasing the system size, the ground-state degeneracy grows with the system size, and in the ensemble including all these configurations there is no order of any kind [90, 91]. In the case of the classical XY (planar vector) or Heisenberg (vectors in three dimensions) model, there is, however, order at $T = 0$ (but not at $T > 0$, according to the Mermin-Wagner theorem). The energy is minimized by arranging the spins in a plane at 120° angle with respect to their neighbors on the same triangle, as shown for a single triangle in Fig. 6. This is referred to as a three-sublattice Néel state. There have been many studies of the $S = 1/2$ variant of this model. This was, in fact, the system for which the RVB spin-liquid state was initially proposed [92]. There is now, however, compelling numerical evidence for the three-sublattice Néel order actually surviving the quantum fluctuations [93, 94].

On the square lattice, frustration effects can be investigated by adding interactions beyond nearest neighbors. A well studied case is the $J_1$-$J_2$ model, where $J_1$ and $J_2$ refer, respectively, to the strengths of the nearest- and next-nearest-neighbor interactions. The system is frustrated if both $J_1 > 0, J_2 > 0$ or if $J_1 < 0, J_2 > 0$. Even with Ising spins, this is a highly non-trivial system, with unresolved questions still attracting interest [95, 96, 97]. We will discuss the frustrated Ising system further in Sec. 3.4.

In the case of quantum spins, the $S = 1/2$ $J_1$-$J_2$ Heisenberg model with all antiferromagnetic couplings is one of the prototypical models with a quantum phase transition out of the standard Néel state, in this case as a function of the coupling ratio $g = J_2/J_1$. While many different calculations show rather consistently that the Néel order vanishes...
FIGURE 7. The square-lattice $S = 1/2$ Heisenberg model with only nearest-neighbor interactions $J_1 > 0$ has an antiferromagnetic ground state, in (a) illustrated by open and solid circles for $\langle S_i^z \rangle > 0$ and $\langle S_j^z \rangle < 0$. In (b), next-nearest-neighbor interactions $J_2 > 0$ are shown as dashed lines. When $0.4 < J_2/J_1 < -0.6$ (approximately) the ground state may be a columnar VBS, with $\langle S_i^z \rangle = 0$ on all sites but modulations in the bond correlations $\langle S_i \cdot S_j \rangle$ (where $i,j$ are nearest-neighbors) as shown with thicker lines for the more strongly correlated bonds forming a columnar pattern. For $J_2/J_1 > 0.6$, the ground state has collinear (striped) magnetic order, as shown in (c).

at a critical coupling ratio $g_c \approx 0.4$ [98, 99, 100, 101, 102, 103, 104], the order of the phase transition and the nature of the non-magnetic ground state are still controversial issues. Most studies indicate some type of VBS state, with a columnar one being the prime candidate, but an RVB spin liquid state has also been proposed [105].

For larger $g$, above $g \approx 0.6$, there is again magnetic order. This can be understood in the limit $g \to \infty$, where the system decouples into two separate square-lattice Heisenberg antiferromagnets. 

The three different ground states of the $J_1$-$J_2$ Heisenberg model with $S = 1/2$ spins are illustrated in Fig. 7. Note that a classical version of this model (Ising, XY, or Heisenberg) has a direct (first-order) Néel-collinear $T = 0$ transition exactly at $g = 1/2$ (which can be easily verified just by computing the energies of those states for $g < 1/2$ and $g > 1/2$).

The magnetically disordered state is thus induced by quantum fluctuations, and it has no direct classical analogue. It is not clear whether this kind of state persists for $S = 1$ or higher spins—for some large $S$, perhaps already $S = 1$, it must give way to a first-order Néel-Collinear transition, as in the classical system.

The reason why it has been so difficult to reach a firm conclusion on the nature of the non-magnetic state and the quantum phase transition between it and the Néel state is that large-scale unbiased computational studies of the $S = 1/2 J_1$-$J_2$ model are currently not possible, because of “sign problems” affecting QMC calculations of frustrated systems.

We will discuss this issue further in Sec. 5. There are no other unbiased method that can reach sufficiently large lattices, e.g., exact diagonalization can reach only $N \approx 40$ spins.\(^2\)

---

\(^2\) Recently developed variational methods based on tensor-product states in principle become unbiased in the limit of large tensors [21]. They have been applied to various frustrated spin models [106]. The computational complexity of such calculations is, however, currently too demanding to reach tensors sufficiently large to produce unbiased results in practice.
2.4.3. The J-Q class of models

VBS states of quantum spin systems in two dimensions were predicted theoretically more than two decades ago [107]. The VBS formation is associated with a spontaneously broken translational lattice symmetry. The VBS state and quantum phase transitions into it are therefore quite different from the non-magnetic states and transitions in the “manually” dimerized systems discussed above in Sec. 2.4.1. Although there is a pattern of “strong” and “weak” spin correlations in both cases, the quantum fluctuations in systems with manually and spontaneously broken translational symmetry are different (being much more interesting in VBS states). Until recently, large-scale computational studies of VBS states and Néel-VBS quantum phase transitions had not been carried out starting from microscopic hamiltonians, because of the QMC sign problems affecting frustrated Heisenberg models (which at first sight are the most natural systems in which to explore the physics of non-magnetic states).

VBS states and the Néel-VBS transition have come into renewed focus with a proposal by Senthil et al. [16] that this transition is generically continuous and, thus, violates the “Landau rule”, according to which an order-order transition (between states breaking unrelated symmetries) should be first-order (except at fine-tuned multi-critical points). In the “deconfined” quantum-criticality scenario, the VBS and Néel order parameters are manifestations of spinon confinement and condensation, respectively. Spinons can be thought of as $S = 1/2$ degrees of freedom, but not just corresponding to the bare individual spins on the lattice sites, but more complex collective objects “dressed” by interactions. A non-compact CP$^1$ field theory was proposed to describe such spinons coupled to an emergent gauge field (which corresponds to the properties of the valence-bond background in which the spinons exist) [16]. A central question is then whether this low-energy physics of a continuum field theory really can arise starting from a reasonable microscopic hamiltonian. Answering this question requires large-scale computational studies of models exhibiting Néel-VBS transitions.

Since the QMC sign problem prohibits large-scale studies of the $J_1$-$J_2$ Heisenberg model and other similar frustrated systems, we have to try something else. In the “J-Q” class of models [17, 108, 109], the Néel order is destroyed by an interaction (Q) which is not frustrated, in the standard sense, but still competes with the Heisenberg (J) interaction. To understand these J-Q models, note first that the Heisenberg interaction is, up to a constant, equal to a singlet projector operator: $H_{ij} = -S_{ij} + \frac{1}{4}$, where

$$S_{ij} = \frac{1}{4} - \vec{S}_i \cdot \vec{S}_j. \quad (21)$$

The pair-singlet, Eq. (19), is an eigenstate of this operator with eigenvalue 1, whereas a triplet state is destroyed by it;

$$S_{ij} |\phi^s_{ij}\rangle = |\phi^s_{ij}\rangle, \quad S_{ij} |\phi^{t,m}_{ij}\rangle = 0, \quad (m = 0, \pm 1). \quad (22)$$

Thus, when $S_{ij}$ acts on a singlet-triplet superposition, only the singlet component survives (is “projected out”—note that the property $S^2_{ij} = S_{ij}$ required of a projection operator is satisfied). The standard Heisenberg interaction thus favors the formation of singlets on pairs of nearest-neighbor sites, but, as we discussed in Sec. 2.2, the fluctuations of these singlets among many different pairings of the spins leads to Néel order in
FIGURE 8. Graphical representation of possible arrangements of products of singlet-projector operators $S_{ij}$ in the J-Q model and its generalizations. (a) is the Heisenberg exchange, (b) the four-spin interaction of the original J-Q model, and (c) a six-spin interaction which leads to more robust VBS order. These operators, and their 90°-rotated analogues, are summed over all positions on the square lattice.

The ground state. The idea behind the J-Q models is to project singlets on two or more bonds in a correlated fashion, using products of several $S_{ij}$ operators on a suitable set of different bonds. This favors a higher density of short valence bonds, thereby reducing or completely destroying the antiferromagnetic order.

The original J-Q hamiltonian [17] on the square lattice can be written as

$$H = -J \sum_{\langle ij \rangle} S_{ij} - Q \sum_{\langle ijkl \rangle} S_{ij}S_{kl},$$  

(23)

where both the $J$ and $Q$ terms are illustrated in Fig. 8. The $Q$ interaction involves four spins on a $2 \times 2$ plaquette. An interaction with three singlet projectors in a columnar arrangement is also shown, and operators with even more projectors, or with the projectors arranged on the lattice in different (non-columnar) patterns, can also be considered [109]. With $J > 0$ and $Q > 0$ [and the minus signs in front of the interactions in Eq. (23)], correlated singlets are favored on the lattice units formed by the product of singlet projectors. It is still not clear just from the hamiltonian whether a VBS state is realized for large $Q/J$—since the hamiltonian does not break any symmetries (the interactions in Fig. 8 are summed over all distinct lattice translations and rotations), a VBS state only forms if the hamiltonian also contains in it, implicitly, some effective interactions that favor singlets in some ordered pattern. The J-Q model (23) does exhibit a Néel-VBS transition, at $Q_c/J \approx 22$ [17, 108, 110], and at lower $Q_c$ if more than two singlet projectors are used [108, 109]. We will discuss this in more detail in Sec. 5. Here we just look briefly at examples of correlation functions useful for characterizing the Néel and VBS states and the quantum phase transition between them.

Order parameters. As an alternative to the square of the full spatially averaged sublattice magnetization (3), we can detect the presence or absence of of Néel order using the spin correlation function,

$$C(r_{ij}) = \langle S_i \cdot S_j \rangle,$$  

(24)

at long-distances. Fig. 9(a) shows J-Q model results for the largest separation of the spins, $r_{\text{max}} = (L/2, L/2)$, on periodic $L \times L$ lattices. The results have to be analyzed carefully to determine the transition point, but already this raw data suggest that the Néel order vanishes, i.e., $C(r_{\text{max}}) \to 0$ when $L \to \infty$, for $J/Q < 0.04$.

VBS order can be detected in the dimer (or bond) correlation function, defined as

$$D_{xx}(r_{ij}) = \langle B_x(r_i)B_x(r_j) \rangle,$$  

(25)

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FIGURE 9. QMC results for the long-distance spin (a) and staggered dimer (b) correlation functions (corresponding to the Néel and VBS order, respectively) in the ground state of the J-Q model (23) versus the coupling ratio \( J/Q \). A careful analysis of these correlation functions, as well as other quantities, indicates a single critical point \((J/Q)_c \approx 0.045\) where both the Néel order and VBS order vanish continuously.

where the bond operator is given by

\[
B_x(r_i) = S(r_i) \cdot S(r_i + \hat{x}).
\]  

(26)

Here, instead of using a subscript \( i \) on the spin operator for some arbitrary site labeling, it is more convenient to use the corresponding lattice position vector \( r_i \). Then \( r_i + \hat{x} \) corresponds to the site immediately next to \( r_i \) in the positive \( x \) direction. The subscripts \( xx \) in (25) indicate that the two bond operators \( B_x \) are both oriented in the \( x \) direction, and this correlation function is, by symmetry, equal to \( D_{yy} \) on an \( L \times L \) lattice. One can also consider bond correlations \( D_{xy} \), where the two bond operators are oriented differently. In a VBS state such as the one illustrated in Fig. 7, one expects \( D_{xx}(r) \) to exhibit a columnar pattern of smaller and larger values. A VBS order parameter can then be defined as a suitable difference between these modulated correlations, e.g.,

\[
D^*_x(r) = D_{xx}(r) - \frac{1}{2}[D_{xx}(r - \hat{x}) + D_{xx}(r + \hat{x})].
\]  

(27)

This correlation function is shown in Fig. 9(b) at the largest lattice separation for several different system sizes. Here it is clear that VBS order exists for \( J = 0 \), up to \( J/Q \approx 0.04 \), roughly where the Néel order sets in. As we will discuss in Sec. 5, all calculations so far point to a single critical point, without any intervening third phase or region of coexistence of both Néel and VBS order.

3. CLASSICAL PHASE TRANSITIONS, MONTE CARLO SIMULATIONS, AND FINITE-SIZE SCALING

Many aspects of quantum phase transitions, and how to analyze them based on numerical finite-lattice data, are very similar to classical phase transitions. Here we discuss this common formalism in the simpler context of classical phase transitions, before turning to calculations and data analysis for quantum systems.
In classical statistical mechanics the prototypical example of a system with a continuous phase transition is the 2D Ising ferromagnet, the exact solution [111, 112] of which shows rigorously that such a phase transition exists. This model is defined by

\[ E_{\sigma} = -J \sum_{\langle ij \rangle} \sigma_i \sigma_j - h \sum_{i=1}^{N} \sigma_i, \]  

(28)

which is just the (potential) energy as a function of the spins \( \sigma_i = \pm 1 \). We use \( \sigma \) to collectively denote an entire spin configuration; \( \sigma = (\sigma_1, \ldots, \sigma_N) \). With the interaction bonds \( \langle ij \rangle \) restricted to nearest-neighbors on the simple 2D square lattice, the exact critical temperature for an infinite system is

\[ T_c/J = 2/\ln(1 + \sqrt{2}) \approx 2.269. \]

The order parameter is the magnetization \( \langle m \rangle = \langle \sigma_i \rangle \). It has the asymptotic \( T \to T_c (T < T_c) \)

form \( m \propto |t|^\beta \), where \( t \) is the reduced temperature, \( t = (T - T_c)/T_c \), and the exponent \( \beta = 1/8 \). Such critical exponents and other aspects of scaling behavior at continuous phase transitions will be discussed in this section.

The exact solution of the Ising model is very special, and normally one studies phase transitions in other ways. Critical exponents appear already in simple mean-field theories, but their values are typically not correct. Mean-field theory is nevertheless essential as a starting point, which we here outline for the Ising model. The most important theoretical framework for phase transitions is the renormalization group, which explains how universal (depending only on symmetries and dimensionality, not details of the interactions) non-trivial exponents (i.e., different from the generic mean-field values) can arise (see, e.g., the book by Cardy [15]). To compute critical exponents and other properties in an unbiased way, one normally uses Monte Carlo simulations, where spin configurations are stochastically sampled according to the Boltzmann distribution. In this section the basics of Monte Carlo simulations are outlined (for more details, see, e.g., the books by Landau and Binder [115] and Newman and Barkema [116]). To motivate and illustrate finite-lattice calculations in general, we will also discuss how properties in the thermodynamic limit can be extracted from Monte Carlo data.

Section outline. The standard mean-field treatment of the Ising model is discussed in Sec. 3.1. In Sec. 3.2 the Monte Carlo method is introduced and used to illustrate how phase transitions and symmetry-breaking can occur in practice for large systems (formally in the limit of infinite system size). In Sec. 3.3 we review the key aspects of critical behavior (defining exponents, etc.) in the thermodynamic limit, then discuss the finite-size scaling hypothesis and demonstrate its usage by analyzing Monte Carlo results for the 2D Ising model. First-order transitions are discussed in Sec. 3.4, using a frustrated 2D Ising model as an example of finite-size scaling methods for detecting discontinuities. In Sec. 3.5 the important concept of spin stiffness (which corresponds to an elastic modulus of a solid) is introduced in the context of XY (planar vector) spin models. The scaling properties of the spin stiffness are illustrated with Monte Carlo simulations.
results for the 3D and 2D XY models, the latter of which does not exhibit a normal phase transition into an ordered state, but exhibits a different \textit{Kosterlitz-Thouless} transition into a critical low-temperature phase. In Sec. 3.6 we briefly discuss how the classical criticality concepts are generalized to quantum phase transitions.

### 3.1. Mean-field theory of the Ising model

In mean-field theories the environment of a subsystem of an infinite lattice is replaced by an external field representing the average interactions between the subsystem and the environment. The subsystem can be a single spin or a cluster of several spins. Here we will just consider the simplest case of a single-spin calculation for the Ising model, i.e., the infinite environment of a single spin is replaced by an effective field.

It is convenient to write the hamiltonian (28) in an extended form with general interactions $J_{ij}$ between all the spins (on an arbitrary lattice), and also include an external magnetic field;

$$E_\sigma = -\frac{1}{2} \sum_{i=1}^{N} \sum_{j=1}^{N} J_{ij} \sigma_i \sigma_j - h \sum_{i=1}^{N} \sigma_i.$$  \(29\)

Note the factor $1/2$, which compensates for each interacting pair being included twice in the sum (and $J_{ii} = 0$). We do not impose any restrictions of the signs and magnitudes of $J_{ij}$, but for simplicity we assume that if there is a phase transition, the ordered state is ferromagnetic. To construct and motivate the single-spin approximation, we first group together all the interactions in (29) involving an arbitrary spin $i$;

$$E_i = -\sigma_i \left( \sum_j J_{ij} \sigma_j + h \right).$$  \(30\)

Note that there is no factor $\frac{1}{2}$ in front of the sum here, and we have used $J_{ij} = J_{ji}$.

We now assume that $h > 0$, so that $m = \langle \sigma_i \rangle > 0$. We will investigate spontaneous ordering in the absence of the field, by eventually letting $h \to 0$. Adding and subtracting a constant, we can write the terms within the parentheses in (30) as

$$\sum_j J_{ij} \sigma_j + h = m \sum_j J_{ij} + h + \sum_j J_{ij} (\sigma_j - m).$$  \(31\)

In its most basic formulation, mean-field theory amounts to neglecting the second sum in (31)—the fluctuation term—after which we are left with the easily solvable problem of a single spin in an effective magnetic field of strength $J_s m + h$;

$$E_\sigma = -(J_s m + h) \sigma,$$  \(32\)

where $J_s$ is the sum of the original couplings

$$J_s = \sum_j J_{ij},$$  \(33\)
and we assume a translationally invariant system, so that this sum is independent of $i$. The magnetization $m$ in (32) is at this stage unknown and will be determined through a self-consistency condition; $\langle \sigma \rangle = m$.

Mean-field theory in the present formulation can be justified if the neglected fluctuations are much smaller, on average, than the other terms in (31), i.e., if

$$\delta_m = \frac{\left\langle \sum_j J_{ij} (\sigma_j - m) \right\rangle}{J_s m + h} \ll 1.$$  \hspace{1cm} (34)

Here $\langle \rangle$ denotes the expectation value under the actual probability distribution (the Boltzmann distribution) of the spins, which we cannot compute exactly. In principle the fluctuations can also be calculated within mean-field theory, as an internal consistency check. Even without doing any calculations, we can roughly deduce the conditions under which $\delta_m$ will be small. Clearly, $\delta_m$ is small if there is substantial order, i.e., when $(1 - m) = (1 - \langle \sigma_j \rangle) \ll 1$, because then most $\sigma_j = 1 \approx m$. This is the case if $h$ is large. It is also true for $h = 0$ if the symmetry is spontaneously broken and $m$ is close to 1, i.e., for $T \ll T_c$ if there is a phase transition. Moreover, $\delta_m$ can be small even if $m$ is not very large, if the sum over $j$ involves many non-zero (and relatively large) coupling constants $J_{ij}$—because of cancellations of fluctuations, the sum $\sum_j J_{ij} \sigma_j$ will then typically (in most of the statistically important spin configurations) be close to $m \sum J_{ij}$. In the extreme case of infinite-range uniform interactions, $J_{ij} = J/N$ for all $i, j$, and $N \to \infty$ (where we regard $J$ as a finite constant, e.g., $J = 1$, in order to have a finite energy) all the fluctuations cancel out exactly and $\delta_m = 0$ (and the mean-field theory is then exact). This holds true exactly also for short-range interactions on an infinite-dimensional lattice. Thus, in general, even if $m$ is not large, the fluctuation measure $\delta_m$ can be expected to be small for systems in high dimensions and/or for long-range interactions. These are then the conditions under which mean-field theory can be expected to be quantitatively accurate. Even in cases where it is not quantitatively accurate, mean-field theory can still provide valuable insights qualitatively.

Let us now actually solve the mean-field problem (32), i.e., the single-spin problem (32) under the self-consistency condition $\langle \sigma \rangle = m$. The magnetization is

$$\langle \sigma \rangle = \frac{\sum \sigma e^{\sigma (J_s m + h) / T}}{\sum e^{\sigma (J_s m + h) / T}} = \tanh[(J_s m + h) / T],$$  \hspace{1cm} (35)

and thus the self-consistency condition reads

$$m = \tanh[(J_s m + h) / T].$$  \hspace{1cm} (36)

This equation in general has to be solved numerically, which can be done easily using successive bracketing of the solution. For small $m$ and $h$, we can proceed analytically by expanding to leading order in $m, h$. First, when the external field $h = 0$, $m = 0$ is a solution for all $T$. Looking for other possible solutions, expanding (36) to third order in $x = J_s m + h$, $\tanh(x) = x - x^3 / 3$, we have

$$m^2 = \frac{3 T^2 J_s - T}{J_s^2},$$  \hspace{1cm} (37)
from which we can identify the critical temperature $T_c = J_s$ below which the magnetization can be non-zero. The asymptotic $T \to T_c$ behavior is

$$m = \left(3 \frac{T_c - T}{T_c} \right)^{1/2}, \quad (h = 0, \ T < T_c). \quad (38)$$

We can also obtain the $h \to 0$ field dependence of the magnetization at $T_c$ from (36). Keeping the leading-order terms in $h$ of the third-order expansion at $T = J_s$ we get

$$m = \left(3 \frac{h}{J_s} \right)^{1/3}, \quad (T = T_c). \quad (39)$$

It is instructive to also look at the full numerical solution for $m$. Fig. 10(a) shows some examples of the field dependence of $m$ at different temperatures. For $T > T_c$ the behavior is analytic across $h = 0$, with a singularity described by (39) developing as $T \to T_c$. Below $T_c$ the behavior is discontinuous, which corresponds to a first-order transition versus $h$ between the $m < 0$ and $m > 0$ states. The discontinuity corresponds to a spontaneous magnetization at zero field, the full numerical solution of which is graphed in Fig. 10(b) and the asymptotic $T \to T_c^-$ behavior of which is given by (38). Here the role of $h \to 0^+$ or $h \to 0^-$ in the calculation is to break the degeneracy between the $\pm m$ solutions. The degeneracy for $h = 0$ also corresponds to co-existence of the two ordered states exactly at the first-order transition, as we will discuss more generally in Sec. 3.4.

Translating the result $T_c = J_s$ with $J_s$ given in Eq. (33) gives $T_c = 4J$ for the 2D Ising model with nearest-neighbor interactions of strength $J$. This is well above the correct value, $T_c/J \approx 2.269$. An over-estimation of $T_c$ can be expected on account of the neglected fluctuations (which naturally lower $T_c$).

The exponents $\beta = 1/2$ and $\delta = 1/3$ in (38) and (39) are generically the values of these critical exponents within mean-field theories. Other exponents can also be computed [15]. While power laws are indeed correct generic features of continuous phase transitions, the mean-field values are not correct in general. As already mentioned, for the 2D Ising model, $\beta = 1/8$ from Onsager’s exact solution. In three dimensions its value

FIGURE 10. Mean-field solution of the Ising model. (a) Magnetization versus external field for temperatures above, at, and below $T_c$. The discontinuity for $h \to 0^+$ and $h \to 0^-$ corresponds to the spontaneous magnetization at $h = 0$. (b) Temperature dependence of the spontaneous magnetization.
in the Ising universality class is $\beta \approx 0.33$, as determined using Monte Carlo simulations and series expansion methods (and also, less precisely, using field-theoretical methods and other analytical approaches). The mean-field critical exponents are exact in four and higher dimensions (four being the upper critical dimension for the Ising model—the dimensionality above which the mean-field critical exponents become exact). We will discuss critical behavior in greater detail further below, after developing the technical aspects of Monte Carlo simulations to study criticality in practice.

### 3.2. Monte Carlo simulations of the Ising model

In a Monte Carlo simulation, the goal is to generate a sequence of spin configurations, $\sigma(1), \sigma(2), \ldots, \sigma(K)$, representing a statistically unbiased sample from the Boltzmann distribution, i.e., the probability $P(\sigma)$ of an arbitrary configuration $\sigma$ to be among the sampled ones should be proportional to the Boltzmann weight at temperature $T$;

$$W_\sigma = e^{-E_\sigma / T}, \quad (40)$$

where we work in units such that $k_B = 1$. The actual (properly normalized) Boltzmann probability is $P_\sigma = W_\sigma / Z$, where $Z$ is the partition function

$$Z = \sum_\sigma e^{-E_\sigma / T}. \quad (41)$$

In Monte Carlo simulations we do not need the full partition function, only the un-normalized weights (40). It is also important to note that the sequence $\sigma(1), \ldots, \sigma(K)$ can be correlated (as we will discuss further below), but we only need the probability $P(\sigma) \propto W_\sigma$ and for now do not need to worry about joint probabilities such as $P(\sigma_1, \sigma_2)$.

**The Metropolis algorithm.** The simplest way to generate a valid sequence of configurations is with the Metropolis algorithm [117]. This kind of simulation starts from an arbitrary spin configuration $\sigma(1)$ (e.g., randomly generated). Thereafter, each successive $\sigma(k + 1)$ is obtained stochastically from its predecessor $\sigma(k)$ according to a few simple steps based on flipping randomly selected spins with a probability related to the desired distribution. We only need to store the current configuration and from now on suppress the “time” index $k$. We denote by $\sigma^{-i}$ the configuration obtained when the $i$th spin of $\sigma$ has been flipped; $\sigma^{-i} = (\sigma_1, \ldots, -\sigma_i, \ldots, \sigma_N)$. Normally one defines a Monte Carlo sweep as $N$ such flip attempts, so that $\propto N$ spins are flipped, on average, during this size-normalized unit of simulation time. A Monte Carlo sweep can be carried out according to the following simple algorithm:4

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4 This pseudocode segment resembles a computer language such as Fortran, but with mathematical notation making it easier to read. Such pseudocodes will be used throughout these notes to illustrate implementations of algorithms (showing the essential steps, but without trivial details that may in practice make real working code somewhat longer). Numbers in curly brackets, here \{1\}, will be used to label codes. In most cases the syntax will be rather self-explanatory and does not require any detailed discussions of
do j=1,N
   i = random[1,...,N]
   if (random[0−1]) < W_{σ−i}/W_{σ}) σ_i = −σ_i
endo

This Metropolis algorithm is based on the detailed balance principle, which is a general theorem for a stochastic process (a Markov chain in some arbitrary configuration space) that should generate a probability distribution \( W \). By this we mean, roughly speaking, that the set of sampled configurations should approach the distribution \( W \) as the number of configurations is increased, independently of the initial condition. Denoting by \( P(A \rightarrow B) \) the transition probability of “moving” to configuration \( B \) if the current one is \( A \), the detailed balance principle states that the desired distribution is generated if all the transition probabilities satisfy the condition

\[
\frac{P(A \rightarrow B)}{P(B \rightarrow A)} = \frac{W(B)}{W(A)},
\]

for all pairs of configurations \( A, B \) for which \( P(A \rightarrow B) > 0 \). In addition, the sampling should be ergodic, i.e., any configuration \( C \) with non-zero weight \( W(C) \) must be reachable, in principle, with non-zero probability through a series of moves starting from an arbitrary configuration.

In the Metropolis algorithm, as implemented in code \{1\}, the transition probability consists of two factors;

\[
P(σ \rightarrow σ^{-i}) = P_{\text{select}}(i)P_{\text{accept}}(σ^{-i}),
\]

where \( P_{\text{select}}(i) \) is the probability of randomly selecting spin \( i \), which here always equals \( 1/N \). The full transition probability in (42) can therefore be replaced by the probability \( P_{\text{accept}} \) of actually carrying out (accepting) the spin flip. This probability is not unique. In the Metropolis algorithm it is taken as

\[
P_{\text{accept}}(σ^{-i}) = \min \left[ \frac{W_{σ^{-i}}}{W_{σ}}, 1 \right].
\]

One can easily confirm that this satisfies the detailed balance condition (42). A probability formally has to be \( \leq 1 \), and this is taken care of above with the “minimum of” function. In code \{1\}, comparing the weight ratio with a random number in the range \([0, 1)\) automatically achieves the same result. Note that the ratio of the Boltzmann factors depends only on the spins interacting with the flip-candidate \( σ_i \) (in the simplest case just its nearest neighbors) and can be rapidly evaluated. In more physical terms, a spin flip leading to a lower energy is always accepted, whereas a configuration with higher energy is accepted only with probability

\[P_{\text{accept}}(σ^{-i}) = \exp[(E_σ - E_{σ^{-i}})/T].\]

\[\]
FIGURE 11. Time series for the magnetization generated in Monte Carlo simulations of the Ising model on $L \times L$ lattices with $L = 8$ (top) and $L = 16$ (bottom) at temperature $T/J = 2.2$ ($< T_c \approx 2.269$). In both cases the starting configuration was fully polarized, $m = 1$, and the subsequent points are separated by $N = L^2$ Metropolis spin flip attempts (constituting one Monte Carlo sweep).

The Metropolis algorithm leads to the correct distribution after some transient time, which depends on the initial configuration, the temperature, and the system size. In a simulation one should therefore discard some number of configurations before “measuring” physical observables. After this equilibration, measurements are normally carried out after every or every few sweeps.

In the case of the ferromagnetic Ising model, the most important quantity to calculate is the magnetization. It should be computed using all the spins to take advantage of self-averaging to improve the statistics. Thus, we will normally use

$$m = \frac{1}{N} \sum_{i=1}^{N} \sigma_i.$$  \hspace{1cm} (45)

We will here consider only simulations with the external field $h = 0$. We have already discussed that fact that relevant symmetries of the hamiltonian (here the discrete spin-inversion symmetry) are not broken in simulations of finite systems. For the Ising model, one should therefore compute spin-inversion invariant expectation values such as $\langle m^2 \rangle$ or $\langle |m| \rangle$ in order to detect the phase transition into the ferromagnetic state.

Symmetry-breaking and finite systems. To obtain a qualitative understanding of how symmetry-breaking in the thermodynamic limit is manifested in practice for large lattices, it is useful to first look at an actual Monte Carlo time series for $m$. Examples for two small systems at a temperature below $T_c$ are shown in Fig. 11. Here one can see that the magnetization fluctuates between positive and negative values, and that the typical time taken to reverse the sign of $m$ is longer for the larger system. Plotting the series over a longer time makes this clearer, but note that the time the system spends close to $m = 0$ is visibly much smaller for $L = 16$ than $L = 8$, and this is of course directly related to a typically longer reversal time. It is clear from results such as these that the distribution of $m$ values is peaked at non-zero positive and negative values for $T < T_c$. For $T > T_c$ the
distribution is instead a single peak centered at \( m = 0 \). Examples of such distributions are shown in Fig. 12. In thermodynamics language, these two qualitatively different distributions can be understood as a consequence of the free energy \( F(m) = E(m) - TS(m) \) at low \( T \) being dominated by the internal energy \( E \) (which is low in large-\( |m| \) configurations), and at high \( T \) by the entropy \( S \) (which is large for small \( |m| \)).

A double-peaked magnetization distribution at low temperature suggests that even a finite system can be considered as ordered, although the spin-inversion symmetry is not broken in the simulations. It appears very plausible that the typical reversal time should diverge with \( L \) (and it is not difficult to check this with simulation data for several \( L \) values and a suitably definition of a reversal), and then no reversals would take place for large \( L \), even during very long simulations. The reason for this divergent time scale is that, in order for the magnetization to reverse, a series of local spin flips must necessarily take the system through many configurations with \( m \approx 0 \), which have increasingly high energy for increasing system size (and, thus, lower Boltzmann probability). For a large system the distribution for \( T < T_c \) is therefore only sampled among the subset of configurations with fixed sign of \( m \)—the stochastic process in practice becomes non-ergodic. Broken symmetry and non-ergodic sampling are manifested strictly only for \( N = \infty \), but in practice also for large but finite systems on time scales less than the typical magnetization reversal time. This time scale of course depends on the details of how the spin configurations are thermally sampled—in Monte Carlo simulations and in a real magnets—but diverges as \( N \to \infty \) for any local sampling scheme.

Normally, in Monte Carlo simulations one does not investigate the time series and the full distribution of physical quantities (although sometimes this is useful). By computing \( \langle m^2 \rangle \) or \( \langle |m| \rangle \), we do not have to worry about the time scale of reversals. We normally want to extrapolate finite-\( N \) results to the thermodynamic limit, where \( \langle |m| \rangle = \langle m \rangle = \langle m^2 \rangle^{1/2} \) (but note that \( \langle m^2 \rangle^{1/2} \neq \langle |m| \rangle \) for finite \( N \), because of the finite width of the peaks in the \( m \)-distribution).

**Autocorrelations and statistical errors.** Before calculating expectation values, we have to discuss how to analyze Monte Carlo data statistically. Consecutive configurations generated with the Metropolis algorithm are not statistically independent—only configurations separated by a number of sweeps much larger than the autocorrelation time are statistically independent.
time are statistically independent. The autocorrelation function for a quantity \( Q \) is defined as

\[
A_Q(t) = \frac{\langle Q(i+t)Q(i) \rangle - \langle Q \rangle^2}{\langle Q^2 \rangle - \langle Q \rangle^2},
\]

where \( t \) and \( i \) denote simulation time, normally in units of the Monte Carlo sweeps defined above, and averages are over the reference time \( i \). The normalization is such that \( A_Q(0) = 1 \) and \( A_Q(t \to \infty) = 0 \). The asymptotic decay is exponential, \( A_Q(t) \sim e^{-t/\tau_Q} \), which can be used to define the autocorrelation time \( \tau_Q \). Normally one instead uses the integrated autocorrelation time,

\[
\tau_Q^{\text{int}} = \frac{1}{2} + \sum_{t=1}^{\infty} A_Q(t).
\]

Here we will not discuss autocorrelations at length, but only summarize their underlying role in determining the statistical precision ("error bars") of computed quantities.

The autocorrelation time for \( m \) of the Ising model roughly corresponds to the typical time between magnetization reversals (as in Fig. 11). Other quantities, such as \( m^2 \) and \( |m| \), that are not sensitive to the sign of \( m \), have shorter autocorrelation times (however, often much longer than a single Monte Carlo sweep). A long autocorrelation time does not bias a computed average (i.e., it is not wrong to measure \( Q \) after every Monte Carlo sweep even if \( \tau_Q \gg 1 \)), provided that the total simulation time is much longer than \( \tau_Q \). The autocorrelation time does, however, come into play (explicitly or implicitly) when computing statistical errors.

To calculate the statistical errors, one can subdivide a simulation into a number \( B \) of bins, each containing some number \( M \) of Monte Carlo sweeps. For some quantity \( Q \), averages \( \bar{Q}_b, b = 1, \ldots, B \) are computed over each bin, and the final average \( \bar{Q} \) and error bar \( \sigma_Q \) (one standard deviation of the average of the bin averages) are calculated according to

\[
\bar{Q} = \frac{1}{B} \sum_{b=1}^{B} \bar{Q}_b, \quad \sigma_Q^2 = \frac{1}{B(B-1)} \sum_{b=1}^{B} (\bar{Q}_b - \bar{Q})^2.
\]

The final estimate of the true expectation value \( \langle Q \rangle \) should then be quoted as \( \bar{Q} \pm \sigma_Q \).

The reason for binning the data is that, according to the central limit theorem, the distribution of bin averages is Gaussian for large \( M \) (unlike the distribution of individual measurements, as seen clearly for the magnetization in Fig. 12), and the computed error bar then has a well defined unique meaning (e.g., we know that the probability of \( \langle Q \rangle \) being within one error bar of \( \bar{Q} \) is about 2/3). This is only true if the bin length \( M \) is also much longer than the autocorrelation time, so that the bin averages can be regarded as statistically independent. If that is the case, the error bar should depend only on the total number of sweeps; \( \sigma_Q \propto 1/\sqrt{MB} \), where the factor of proportionality is \( \propto \sqrt{\tau_Q} \) (and of course also depends on the detailed form of the distribution of the individual measured \( Q \) values). It is not necessary to calculate \( \tau_Q \) explicitly. If there is any doubt about the bins being sufficiently long, one can check this by using a rather large number of bins (e.g., in the range 100-1000) and saving all the bin averages on disk. The data can then
be re-binned into longer bins post-simulation, and the convergence of $\sigma_Q$ as a function of the bin length can be tested. Saving the bin averages on disk is always advisable, not only for the purpose of analyzing the error bars, but also in order to make it easy to add more data at some later time (to improve the results, if needed).

An important point that has to be mentioned is that the autocorrelation time of any local updating scheme, e.g., the Metropolis algorithm, diverges for $T \to T_c$ and $N \to \infty$. We previously saw an example of this in the magnetization reversal time of the Ising model, but divergent autocorrelation times also affect any quantity (in particular $m^2$ and $|m|$) that is sensitive to the fluctuations of the magnitude of $m$ (which in practice is the case for most quantities of interest). It is therefore difficult to obtain good results for large systems close to a critical point. In many cases, including the Ising model, this problem can be solved (or almost solved) by using cluster algorithms [118, 119], where clusters of spins (constructed so as to satisfy detailed balance) are flipped collectively (instead of flipping individual spins one-by-one). While we will consider an analogous loop-cluster algorithm for QMC simulations in Sec. 5, we do not have to discuss classical cluster algorithms here; they have been described extensively in the literature (e.g., in [115] and [116]). It should be noted, however, that the Ising results to be discussed below, for systems with up to $1024^2$ spins, were in fact obtained using a cluster algorithm. It would be very difficult to generate (within reasonable time) data of the same quality using the Metropolis scheme for such large systems in the critical region.

### 3.3. Finite-size scaling and critical exponents

Fig. 13(a) shows the temperature dependence of the squared magnetization $\langle m^2 \rangle$ of the 2D Ising model for several lattice sizes. An increasingly sharp feature develops with increasing $L$ in the neighborhood of the known $T_c$, and it appears very plausible that $\langle m^2 \rangle$ vanishes for $N \to \infty$ for $T > T_c$ and remains non-zero for $T < T_c$. In this plot, error bars are not visible because they are much smaller than the line widths. The data were generated on a very fine $T$-grid to produce continuous-looking curves.

The size dependence of $\langle m^2 \rangle$ at three different temperatures is shown in Fig. 13(b). For $T < T_c$, it converges with $L$ to a non-zero value, while for $T > T_c$ it decays to 0 as $L^{-2}$. Exactly at $T_c$ the decay follows a non-trivial power-law; $\langle m^2 \rangle \propto L^{-2\beta/\nu}$, where $\beta = 1/8$ is the same exponent as in the $N = \infty$ magnetization for $T < T_c$, and $\nu$ (the correlation-length exponent, to be discussed below) equals 1 for this model. This behavior is an example of finite-size scaling. A perfect power-law behavior at criticality holds strictly only when $L \to \infty$, while for small systems there can be significant corrections to scaling (which are unusually small in the case of the Ising model discussed here). Normally we do not know the exact value of $T_c$ (which may of course be one reason why a simulation is carried out), and procedures for locating the critical point then have to be devised. There are many ways to do this, all building in some way or another on the fact that the order parameter and related quantities should behave as non-trivial power laws (with known or unknown exponents) for large lattices at a critical point.

Note that the squared magnetization is related to the spin correlation function,

$$C(r_{ij}) = \langle \sigma_i \sigma_j \rangle,$$

\text{(49)}
FIGURE 13. (a) The squared magnetization as a function of temperature for several $L \times L$ Ising systems. (b) The finite-size dependence at, below, and above $T_c$. At $T_c$ the behavior is a power law, $m^2 \sim L^{-1/4}$, which on this log-log scale corresponds to a line with slope $-1/4$. For $T > T_c$ the decay to zero is of the form $L^{-2}$, and this is also the rate of convergence for $T < T_c$.

where $r_{ij}$ is the distance between the spins. If there is long-range order, then $C(r) \to \langle m^2 \rangle$ when $r \to \infty$ in an infinite lattice. For finite systems, the same is true for $C(r_{\text{max}})$ in the limit $L \to \infty$, where $r_{\text{max}}$ is the longest distance on a periodic lattice, e.g., $r_{\text{max}} = \sqrt{2}L$ for an $L \times L$ lattice. If there is no order, then $C(r) \to 0$, according to a power law at $T_c$ and exponentially for $T > T_c$. Below $T_c$, in the infinite system the “connected” correlation function,

$$C^+(r) = C(r) - \langle m \rangle^2,$$

(50)

decays to zero exponentially as $r \to \infty$. The exponential forms of both $C(r)$ and $C^+(r)$ are characterized by a correlation length $\xi$, which diverges as $T \to T_c$ from either side. Note also that $\langle m^2 \rangle$ can be written exactly as a sum over the spin correlations;

$$\langle m^2 \rangle = \frac{1}{N} \sum_r C(r).$$

(51)

The $L^{-2}$ convergence of $\langle m^2 \rangle$ (both below and above $T_c$) is simply related to the finite correlation length.

Here it should be pointed out that the nature of the spin correlations below $T_c$ depends on the symmetry of the order parameter we are dealing with. For a vector order parameter, e.g., in the Heisenberg model (with classical or quantum spins), the correlations of the spin components parallel (longitudinal) to the order parameter of a symmetry-broken system decay exponentially, as in the Ising model. The transverse correlations decay as a power law, however. This is related to the continuous symmetry of the order parameter. In the Ising model the symmetry is discrete—the ordered state breaks the spin-inversion symmetry and the free-energy cost of a local magnetization fluctuation is large (proportional to the boundary of a flipped domain). This leads to the exponentially decaying (connected) correlation function in the ordered phase. In the Heisenberg model, on the other hand, there are gapless spin-wave excitations (which are excitations of the spins in the directions transverse to the ordering direction and are more generally called Goldstone modes). Fluctuations due to these lead to a power-law form of the transverse spin...
correlations. The transverse correlation length is then formally infinite, and in a finite-lattice calculation in which the rotational symmetry is not explicitly broken (so that the computed $\langle m^2 \rangle$ contains contributions from both longitudinal and transverse correlations), the size corrections to $\langle m^2 \rangle$ are $\propto 1/L$. We already discussed this behavior in the ground state of the 2D $S = 1/2$ Heisenberg antiferromagnet in Sec. 2.4, and Fig. 5(b) shows the $1/L$ corrections very clearly.

3.3.1. Scaling and critical exponents

To discuss finite-size scaling close to a critical point in more detail, we first have to review some basic aspects of critical phenomena in the thermodynamic limit. We will only list some of the key results and definitions here; see any standard text on critical phenomena (e.g., the book by Cardy [15]) for more details.

The correlation length is one of the most important concepts underlying the theory of phase transitions and critical phenomena. In an infinite system, as the critical temperature is approached, the correlation length diverges according to a power law:

$$\xi \sim |t|^{-\nu}. \quad (52)$$

The exponent $\nu$ is the same upon approaching $T_c$ from above or below, but the prefactor in (52) is in general different for $t \to 0^+$ and $t \to 0^-$ (however, ratios of these prefactors—called amplitude ratios—are often universal). Within mean-field theory, $\nu = 1/2$. Exactly at $T_c$, although the correlation length is formally infinite, the system is not yet ordered. Instead, for an infinite system the correlation function exactly at $T_c$ has the power-law form,

$$C(r) \sim r^{-(d-2+\eta)}, \quad (53)$$

where $d$ is the dimensionality of the system and $\eta$ is another critical exponent (also called the anomalous dimension, because it can be related to the fractal dimensionality of ordered domains at the critical point), the mean-field value of which is $\eta = 0$. Long-range order sets in only infinitesimally below $T_c$, where the asymptotic long-distance correlation approaches a constant; $C(r \to \infty) = \langle m^2 \rangle$.

We have already discussed the behavior of the order parameter in the symmetry-broken state as $t \to 0^-$:

$$\langle m \rangle \sim |t|^\beta. \quad (54)$$

We will also be interested in the corresponding susceptibility, defined as

$$\chi = \left. \frac{d \langle m \rangle}{dh} \right|_{h \to 0} = N \frac{T}{T} \left( \langle m^2 \rangle - \langle |m| \rangle^2 \right), \quad (55)$$

where $h$ is the strength of a field coupling to the order parameter, e.g., in the case of the Ising ferromagnet a term $-h \sum \sigma_i$ in the hamiltonian. The last expression in (55) shows explicitly how $\chi$ is directly related to the fluctuations of the order parameter. The susceptibility diverges at $T_c$:

$$\chi \sim |t|^{-\gamma}. \quad (56)$$
Thus, upon approaching a critical point, the system becomes infinitely sensitive to a field \( h \) coupling to the order parameter, and exactly at \( T_c \) the linear response form \( \langle m \rangle = \chi h \) ceases to be valid. Instead, at \( T_c \), the order parameter depends on the (weak) field as \( h^{1/\delta} \). According to the result in Eq. (39), the mean-field value of the exponent here is \( \delta = 3 \).

The specific heat is also singular at \( T_c \),

\[
C \sim |t|^{-\alpha},
\]

where \( \alpha \) can be positive or negative. In mean-field theory \( \alpha = 0 \). When \( \alpha < 0 \), there is no divergence, only a cusp singularity at \( T_c \). In some cases, e.g., for the 2D Ising model, \( \alpha = 0 \) but there is still a weak, logarithmic divergence of the specific heat.

The critical exponents \( \nu, \beta, \eta, \) etc., that we have encountered above are not all independent of each other. Relationships between the exponents are explained by the renormalization group theory, which, in the type of order-disorder transitions we are discussed here, shows that there are two more fundamental exponents, in terms of which the physically observable exponents can be written [15]. Exponent relations had been found using other arguments even before the advent of this ultimate theory of phase transitions (in the most complete form by Widom in the mid 1960s), e.g.,

\[
\begin{align*}
\gamma &= \nu(2 - \eta), \\
\nu d &= 2 - \alpha, \\
\alpha + 2\beta + \gamma &= 2.
\end{align*}
\]

Such relations are very useful for checking the consistency of numerical calculations of the exponents. Exponent relations involving the dimensionality \( d \) are called hyper-scaling relations and are less generic than the other relations. They are not applicable, e.g., within mean-field theory, and, therefore, for any system at or above the upper critical dimension.

### 3.3.2. The finite-size scaling hypothesis

The basic assumption underlying finite-size scaling theory [120] is that deviations from the infinite-size critical behavior should occur when the correlation length \( \xi \) (of the infinite system) becomes comparable with the finite system length \( L \). If \( L \gg \xi \), the fact that the system is finite should be irrelevant and the infinite-size behavior applies. On the other hand, if \( L \ll \xi \), then \( L \), not \( \xi \), should be the most relevant length-scale. In order to see how the two length scales come into play, it is useful to express quantities of interest in terms of the correlation length. Consider a quantity \( Q \) which exhibits a power-law divergent behavior at \( T_c \) (reduced temperature \( t \to 0 \)),

\[
Q \sim |t|^{-\kappa},
\]

e.g., the susceptibility (in which case \( \kappa = \gamma \)). We can use Eq. (52) to express \( |t| \) as a function of the correlation length;

\[
|t| \sim \xi^{-1/\nu},
\]
and using this we can write \( Q \) as
\[
Q \sim \xi^{\kappa/\nu}.
\] (61)

This form should apply for \( \xi \ll L \), but when \( \xi \sim L \) the divergence can no longer continue on the finite lattice. The maximum value \( Q_{\text{max}} \) attainable by \( Q \) on the finite lattice should then be obtained by replacing \( \xi \to L \) in Eq. (61), giving
\[
Q_{\text{max}}(L) \sim L^{\kappa/\nu}.
\] (62)

In the same way, from Eq. (60) we can also deduce the scaling of the reduced temperature at which \( \xi \) reaches \( L \), which should also be the temperature at which the maximum value of the quantity \( Q \) is reached;
\[
|t_{\text{max}}(L)| \sim L^{-1/\nu}.
\] (63)

It should be noted that this is just a proportionality, and the number in front of \( L^{-1/\nu} \) depends on the quantity. The shift also applies to non-divergent quantities—any feature which develops singular behavior as \( T \to T_c \) should shift at the rate \( L^{-1/\nu} \). If there is a well-defined maximum or other distinguishable feature in some quantity at \( T = T^*(L) \), then this temperature can be used as a size-dependent critical temperature (and, again, this temperature is not unique but depends on the quantity considered).

The finite-size scaling laws (62) and (63) follow from a more general finite-size scaling hypothesis [120], which, like the scaling theory for infinite systems, was initially proposed based on phenomenological considerations, but was later derived using the renormalization group theory. The hypothesis is that an observable which is singular at \( T_c \) in the thermodynamic limit scales with the system size close to \( T_c \) as a power of \( L \) multiplied by a non-singular function of the ratio \( \xi/L \). Any singular quantity (not necessarily divergent) should thus be of the form
\[
Q(t, L) = L^{\sigma} f(\xi/L),
\] (64)

which, using \( \xi \sim |t|^{-1/\nu} \), we can also write as
\[
Q(t, L) = L^{\sigma} g(tL^{1/\nu}).
\] (65)

This scaling law should hold both above \( (t > 0) \) and below \( (t < 0) \) the critical point. Exactly at \( T_c \), we recover the size-scaling \( Q(0, L) \sim L^{\sigma} \). To relate \( \sigma \) to the standard critical exponents, we can use the fact that, for fixed \( t \) close to 0, as the system grows the behavior for any \( t \neq 0 \) eventually has to be given by Eq. (59); \( Q(t, L \to \infty) \sim |t|^{-\kappa} \) (where \( \kappa \) is negative for a singular non-divergent quantity, e.g., the for the order parameter we have \( \kappa = -\beta \)). To obtain this form, the scaling function \( g(x) \) in (65) must asymptotically behave as \( g(x) \sim x^{-\kappa} \) for \( x \to \infty \). In order for the size-dependence in (65) to cancel out, we therefore conclude that \( \sigma = \kappa/\nu \), i.e.,
\[
Q(t, L) = L^{\kappa/\nu} g(tL^{1/\nu}).
\] (66)

To extract the scaling function \( g(x) \) using numerical data, one can define
\[
y_L = Q(t, L)L^{-\kappa/\nu}, \quad x_L = tL^{1/\nu},
\] (67)
FIGURE 14. Monte Carlo results for the susceptibility (55) of the Ising model on several different $L \times L$ lattices. (a) shows the temperature dependence, with the vertical line indicating $T_c$. Note the vertical log scale. In (b) the data has been scaled using the exact values of the Ising exponents, $\gamma = 7/4$ and $\nu = 1$, and the exact value of $T_c$ in $t = (T - T_c)/T_c$.

and plot $y_L$ versus $x_L$ for different system sizes. If the scaling hypothesis is correct, data for different (large) system sizes should fall onto the same curve, which then is the scaling function (this is referred to as curves collapsing onto each other): $g(x) = y_{L \to \infty}(x)$. Fig. 14 illustrates this using Monte Carlo data for the magnetic susceptibility of the 2D Ising model. The peak location in panel (a) clearly moves toward the known $T_c$ with increasing $L$. After scaling the data according to the above procedures, as shown in panel (b), the curves indeed collapse almost onto each other close to $t = 0$, but further away from the critical point deviations are seen for the smaller systems. These are due to corrections to scaling, which in principle can be described with subleading exponents.

We can apply the scaling form (66) to the correlation length itself, for which $\kappa = \nu$ and the $L$-scaling is independent of model-specific exponents. In cases where the universality class is not known a priori, this is useful for extracting the exponent $\nu$ by curve-collapsing $\xi/L$ data without having to simultaneously adjust another exponent $\kappa$ in (66).

Practical definitions of the correlation length. The correlation length can be defined in various ways, not necessarily just based on the asymptotic decay of the correlation function (which is often difficult to extract reliably). One practical and commonly used correlation length definition is based on the Fourier transform of the correlation function, often called the (static) structure factor,

$$S(q) = \langle \sigma_q \sigma \rangle = \sum_r e^{-iq \cdot r} C(r) = \sum_r \cos(q \cdot r) C(r),$$

(68)

where $\sigma_q$ is the Fourier transform of an individual spin configuration,

$$\sigma_q = \frac{1}{\sqrt{N}} \sum_j \sigma_j e^{-iq \cdot r_j}.$$  

(69)

We denote by $Q$ the wave-vector of the dominant correlations—for a ferromagnet $Q = 0$, for a 2D antiferromagnet $Q = (\pi, \pi)$, etc. To simplify the notation, $q$ will be used for the
deviation from \( Q \). Then \( q_1 = 2\pi / L \) corresponds to one of the wave-vectors closest to \( Q \), e.g., \( Q + (2\pi / L)\hat{x} \), where \( \hat{x} \) is the reciprocal-space unit vector in the \( x \)-direction.

A correlation length \( \xi_a \) can be defined using the structure factors at \( q = 0 \) and \( q_1 \):

\[
\xi_a = \frac{1}{q_1} \sqrt{S(0) / S(q_1)} - 1. \tag{70}
\]

One can show that, for a \( d \)-dimensional lattice, if the correlation function is given by the Ornstein-Zernike form (obtained in mean-field treatments [15]),

\[
C_{OZ}(r) \sim r^{-\frac{d}{2}(d-2)} \exp(-r/\xi), \tag{71}
\]

then \( \xi_a \) is related to the original correlation length \( \xi \) appearing in the exponential decay of this correlation function according to

\[
\xi_a = \xi \sqrt{\frac{(1+d)(3+d)}{8d}}. \tag{72}
\]

Thus, for \( d = 1 \) and 3, \( \xi_a = \xi \), whereas the 2D case is special, with \( \xi_a = \xi (15/16)^{1/2} \) (or, one may say that \( d = 1, 3 \) are special cases, since the factor differs from one also for all \( d > 3 \)). The Ornstein-Zernike form is normally valid in disordered phases for \( r \gg \xi \) [15]. Deviations from this form at short distances imply that (72) is not true exactly, but regardless of the short-distance behavior this relation holds exactly when \( \xi \to \infty \).

In the case of a long-range ordered classical system, \( \xi_a \) normally diverges as \( T \to 0 \), for any \( L \), because there are no fluctuations in the ground state (and, thus, the structure factor vanishes for \( q \neq 0 \)). In order to remove the contributions from the non-decaying part of the correlation function in an ordered system, we can use the connected correlation function (50). Although \( \langle m \rangle^2 \) is not uniquely defined for a finite system, one can, e.g., subtract \( C(r_{\text{max}}) \). As an alternative, we can use a different correlation length definition, based on the structure factor at \( q_1 \) and \( q_2 = 2q_1 = 4\pi / L \):

\[
\xi_b = \frac{1}{q_1} \sqrt{\frac{S(q_1)/S(q_2) - 1}{4 - S(q_1)/S(q_2)}}. \tag{73}
\]

One can show that (72) also holds for this definition if \( C(r) \) is given by the Ornstein-Zernike correlation function, also when a constant corresponding to the long-range order is added for \( T < T_c \) [since this affects only \( S(0) \)].

Fig. 15 shows results for \( \xi_a / L \) and \( \xi_b / L \) for the 2D Ising model. Note that a logarithmic scale is used in the case of the \( T \to 0 \) divergent \( \xi_a / L \), while \( \xi_b / L \) is convergent and graphed on a linear scale. Both quantities exhibit size independence (curves crossing each other) at \( T_c \), but the values there are clearly different. This is because the Ornstein-Zernike form of the correlation function applies asymptotically only for \( T > T_c \), and there is no reason why the two definitions \( \xi_a \) and \( \xi_b \) should agree exactly at \( T_c \) (although their values should be related). Their values are very similar for large systems close to \( T_c \) in the disordered phase. The crossing points (or peak location of \( \xi_b \)) can be used to
extract $T_c$ in systems where it is not known. The temperature axis can also be scaled in the same way as in Fig. 14 to extract the correlation-length exponent.

Note that for the small number of $q$-points needed, $S(q)$ can be efficiently evaluated using the Fourier transform (69) of the spin configurations generated in Monte Carlo simulations. Since the structure factor is real-valued we have

$$S(q) = \left\langle \text{Re}\{\sigma_q\}^2 \rightangle + \left\langle \text{Im}\{\sigma_q\}^2 \rightangle.$$  \hspace{1cm} (74)

Computing the full correlation function $C(r)$ and Fourier-transforming it post-simulation is much more time-consuming.

**Binder ratio and cumulant.** Besides $\xi/L$, there are also other dimensionless quantities that are size-independent at the critical point and useful for extracting $T_c$ independently of the values of the critical exponent. The perhaps most frequently used one is the Binder ratio [121, 122]:

$$R_2 = \frac{\langle m^4 \rangle}{\langle m^2 \rangle^2}. \hspace{1cm} (75)$$

At $T_c$, the power laws cancel out, and the ratio is $L$-independent (and also universal), up to subleading finite-size corrections. Normally, graphing $R$ versus $T$ for different system sizes produces curves that intersect each other close to $T_c$. Locating the points where $R_2$ for pairs of system sizes (e.g., $L$ and $2L$) cross each other, one obtains a size dependent critical point which typically converges faster than the $L^{-1/\nu}$ shift in (63). One can think of this as being a results of the leading corrections canceling in a quantity involving two system sizes, and one is then left with something which approaches $T_c$ according to a faster, higher-order scaling correction. One can also define ratios similar to (75) based on other powers of $m$, e.g., $R_1 = \langle m^2 \rangle / \langle |m| \rangle$. The curve-crossing method for locating $T_c$ can also be applied in the same way with $\xi/L$.

The Binder ratio also has other interesting properties. In the case of a scalar order parameter (e.g., for the Ising model), the Binder *cumulant* is defined as

$$U_2 = \frac{3}{2} \left(1 - \frac{1}{3}R_2 \right). \hspace{1cm} (76)$$

![FIGURE 15. Temperature dependence of the two correlation length definitions, Eqs. (70) and (73), normalized by the size $L$ for $L \times L$ Ising models. The vertical lines indicate $T_c$.](image-url)
In an ordered state, $U_2 \to 1$ for $N \to \infty$, because the magnetization distribution $P(m)$ then approaches two delta-functions at $\pm \langle |m| \rangle$, and, therefore, $R_2 \to 1$. In contrast, in a disordered phase, the fluctuations of $m$ are Gaussian around $m = 0$ (following from the central limit theorem, since fluctuations in regions separated by distance $\gg \xi$ in a large system are uncorrelated). Based on Gaussian integrals, $R_2 \to 3$ and $U_2 \to 0$.

Generalizing the Binder cumulant to an $n$-component order parameter (where $n = 1$ for the Ising magnetization, $n = 2$ for the XY model, etc.), one should keep in mind that $m^2 = m \cdot m$ and $m^4$ in (75) are insensitive to angular fluctuations of the order parameter. Integrating a Gaussian distribution of $|m|$ over the $n$-dimensional space to compute the averages in (75) does, however, introduce $n$-dependent factors. To reproduce the above properties of $U_2$ one therefore has to define it for a general order parameter as

$$U_2 = \frac{n + 2}{2} \left( 1 - \frac{n}{n + 2} R_2 \right).$$

Fig. 16 shows Monte Carlo results for $U_2$ as a function of $T$ for several 2D lattices. The evolution into a step-function at $T_c$ with increasing system size can be seen clearly. In this case all the curves cross each other very close to the known $T_c$, reflecting very small subleading corrections in this model. It is common in other systems that the crossings exhibit some drift, and a careful extrapolation of crossing points have to be carried out (e.g., based on data sets for sizes $L$ and $2L$, or some other aspect ratio).

Finite-size scaling in practice. We briefly discuss how to carry out finite-size data-collapse in practice. The number of parameters involved (i.e., $T_c$ as well as one or two exponents, and possibly also exponents of subleading corrections to be discussed below) is rather small, and normally one has some rough knowledge of their values just from looking at raw data and doing some initial experimentation, e.g., by just locating a non-trivial power-law behavior as in Fig. 13 [and this may often be enough to determine the exponent ratio $\kappa / \nu$ in Eqs. (62) and (66)]. An analysis of the Binder cumulant or $\xi / L$ may already have given $T_c$ to adequate precision, but it may still be useful to check the

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**FIGURE 16.** The Binder cumulant (76) for $L \times L$ Ising models. In (a) one can see the approach to the limiting values $U_2 \to 0$ (for $T > T_c$) and $U_2 \to 1$ (for $T < T_c$) for increasing $L$. In (b) the data close to $T_c$ (vertical line) are graphed on a more detailed scale and for larger $L$ to show the crossings of the curves.
sensitivity of other fits to its value. Thanks to the power of modern computers, as an alternative to using some complicated multi-dimensional optimization procedure, one can write a simple brute-force computer program to search for the best set of parameters on a suitable finite mesh. The goodness of the data collapse produced by a set of parameters can be quantified as the $\chi^2$-value obtained by fitting a single high-order polynomial through all the scaled data points $(x_L, y_L)$, defined in (67), simultaneously for all $L$ for which data are available. Normally there is a large number of data points, for different couplings and system sizes, and, since the scaling function should be well behaved, a polynomial of reasonable order (3th-8th, as a rough guideline) should work well within the window where the data can be collapsed. The size of this window also has to be adjusted until the data collapse well. As an example, for the data in Fig. 14, $x$ in the range $(-0.5, 0.5)$ should be appropriate, although the window also depends on the system sizes included in the analysis and the the error bars (which determine the sensitivity to neglected subleading scaling corrections).

It is not always easy to determine reliable error bars for parameters obtained in this kind of fitting. Beyond the purely statistical errors (which can be determined, e.g., by repeating the $\chi^2$ minimization several times with Gaussian noise, of magnitude equal to the error bars, added to the data), there are also systematical errors due to scaling corrections, which can be difficult to estimate. If $\chi^2$ is statistically reasonable (i.e., close to 1 per degree of freedom), one would normally assume that the neglected corrections have not influenced the parameters beyond the statistical uncertainties.

Fig. 17 shows an example of data collapse. The critical exponents and $T_c$ of the 2D Ising model were determined using data for $L \in \{64 - 512\}$ in the window $|t|L^{1/\nu} < 0.5$ (where $t$ contains the variable $T_c$ adjusted in the procedure). The results, listed in the figure caption, are in excellent agreement with the known $T_c$ and the 2D Ising exponents, and the fit is also statistically very good. Using the same fitting window, a statistically
acceptable fit cannot be obtained if much smaller lattices are included, e.g., including $L = 32$ gives $\chi^2 \approx 2$ (per degree of freedom), which is marginally too large for the number of degrees of freedom of the fit. In this case $T_c = 2.26924 \pm 0.00002$, about 3 error bars above the true value, while the exponent are correct to within two error bars.

Corrections to scaling. As a check, it is often a good idea to include some corrections to the leading scaling forms, and some times it is even necessary to do so in order to obtain good fits. The most commonly used method is to do the data collapse with

$$y_L = Q(t, L)L^{-\kappa/\nu}(1 + aL^{-\omega})^{-1}, \quad x_L = tL^{1/\nu},$$

where $\omega$ is a subleading exponent and $a$ is a constant. Here there is no correction to the $x_L$ scaling (the finite-size shift of the critical point), but such a correction can in principle be included as well [85]. Normally, good fits with corrections can be obtained also when lattice sizes are included that have to be left out if no corrections are used. A larger range of system sizes can partially compensate for the fact that the statistical uncertainties of all parameters increase when more parameters are included. If consistent leading critical exponents are obtained in fits both with and without corrections, then one can be reasonably certain that the results are correct.

3.4. First-order transitions

The scaling properties we discussed in the preceding section apply at continuous phase transitions, where the correlation length diverges. At first-order (discontinuous) transitions, the correlation length remains finite at the transition point and the order parameter, as well as other quantities, exhibit discontinuous jumps. The discontinuities develop in the limit of infinite system size, normally according to power-laws which can also be studied using finite-size scaling techniques. The exponents associated with these powers-laws are typically trivially related to the dimensionality of the system [123, 124, 125]. For instance, the specific heat diverges with the system size as $L^d$ at a first-order transition, instead of $L^{\alpha/\nu}$, with a typically very small (or even negative) $\alpha$, at a continuous transition. The shift of the critical point with the system size scales as $L^{-d}$, instead of the $L^{-1/\nu}$ shift of a continuous critical point.

Although finite-size scaling with exponents equal to $d$ in principle makes it easy to recognize a first-order transition, studies of weak first-order transitions are difficult, because they exhibit large corrections to the leading scaling forms. It may then be difficult, with system sizes accessible in practice, to clearly distinguish slowly developing discontinuities from weaker singular behavior at a continuous transition.

Strongly first-order transitions are also difficult to study, for a completely different reason. A Monte Carlo simulation may get stuck in a meta-stable state, in which case computed quantities do not correspond to correct thermal averages (which, on the other hand, is completely analogous to real systems, for which metastability and hysteresis effects are hall-marks of first-order transitions). It may then even be difficult to accurately locate the transition point. To alleviate such problems, various multi-canonical Monte Carlo methods have been developed in which the configurations are sampled in an ex-
tended ensemble where the temperature of the system is also fluctuating (tempering or parallel tempering [126, 127] methods), or with a distribution different from the Boltzmann probability [128, 129, 130] (to which the measurements are re-weighted), which makes it easier for the system to explore the configuration space.

### 3.4.1. Phase coexistence

One of the most important characteristics of a first-order transition is phase coexistence. In Monte Carlo simulations at the transition point (in practice in a small window which shrinks to a single temperature with increasing system size) this is manifested in the generation of two types of configurations, corresponding to the two distinct phases existing just above and below the transition temperature. This is provided that the full configuration space can be ergodically sampled, which, as discussed above, is not always possible in practice. In small systems, there will also be configurations that cannot be clearly associated with one of the phases—they correspond to the fluctuations (domain walls) required for the system to transition between the two phases. Fig. 18 illustrates this schematically for an Ising order parameter. The transition window in which a three-peak distribution can be observed narrows rapidly with increasing system size, and the peaks develop into delta-functions. When the weight between the peaks becomes very small, it may in practice not be possible to ergodically sample the configurations, as the system gets trapped within the sub-space corresponding to just one of the peaks (in a way exactly analogous to the symmetry-breaking discussed in Sec. 3.2).

Note that the type of order-parameter distribution in Fig. 18 does not apply to the field-driven first-order transition of the Ising model, with magnetization curves illustrated in Fig. 10. In that case there would be just two peaks, with weight transferring between them as the field is tuned through $h = 0$. The phase coexistence at $h = 0$ (and $T < T_c$) corresponds to the two symmetric peaks in the distributions in Fig. 12. The similarity between this case and a paramagnetic-magnetic transition (with a general $n$-component order parameter $m$) can be made clearer by considering the one-dimensional distribution of $|m|$. This distribution has two peaks, one at $|m| = 0$ and one at $|m| > 0$, when an ordered and disordered phase coexist. Note again that in continuous transitions (Fig. 12) the central peak continuously splits into two peaks below $T_c$, in contrast to the two ordering peaks emerging at a non-zero value of $|m|$ in the first-order case.

The Binder ratio, Eq. (77) in the case of a generic $n$-dimensional order parameter depends only on the distribution of $|m|$. It has a very interesting property at a first-order transition. One can easily check, using, e.g., an idealized co-existence distribution of the type $P(|m|) = (1 - p)G(|m|) + pδ(|m| - m^*)$, where $G$ is a normalized “half-Gaussian” (defined for $|m| ≥ 0$ and $0 < m^* ≤ 1$, that $U_2$ exhibits a negative divergence when the ordered-phase weight $p$ is tuned from 0 to 1, and if the width of the Gaussian vanishes (corresponding to infinite system size). While the full $m$ distribution of course contains more information, locating a window of negative Binder cumulant and checking for a divergence of the peak value can be a practical way to analyze a first-order transition.
FIGURE 18. Evolution (schematic) of the magnetization distribution of a finite Ising ferromagnet close to a first-order transition. There is a rapid transfer of weight between the central peak (corresponding to the disordered phase) and the two peaks at non-zero magnetization (the two ferromagnetic states) as the temperature is tuned through the transition region. The peaks become delta-functions in the limit of infinite system size. The transition temperature $T_c$ can be defined as a point with some specific (but essentially arbitrary) feature of the three peaks, e.g., equal weight or equal height of the peaks.

3.4.2. Frustrated Ising model

Here we look at a particular example of a first-order transition, in the frustrated 2D square-lattice Ising model with Hamiltonian

$$E_\sigma = -J_1 \sum_{\langle ij \rangle_1} \sigma_i \sigma_j + J_2 \sum_{\langle ij \rangle_2} \sigma_i \sigma_j,$$

(79)

where both couplings $J_1, J_2 > 0$ (but note the different signs in front of the parameters). The first term (where $\langle ij \rangle_1$ denotes nearest-neighbor spins) is then the standard Ising ferromagnet, whereas the second term (where $\langle ij \rangle_2$ refers next-nearest-neighbor spins, i.e., across the diagonals on $2 \times 2$ plaquettes, as in Fig. 7) is antiferromagnetic and causes frustration. For coupling ratios $g = J_2/J_1 < 1/2$ the ground state of the system is ferromagnetic (fully polarized). In the limit $g \to \infty$ the system reduces to two decoupled antiferromagnets, with four striped (or collinear) ground states, such as the one illustrated in Fig. 7(c). These remain the ground states for all $g > 1/2$. At the point $g = 1/2$ the ground state is massively degenerate. Note that the case of $J_1 < 0$ is equivalent to $J_1 > 0$, through the invariance of $E_\sigma$ under the transformation $\sigma_i \to -\sigma_i$ on one of the checker-board sublattices; thus in this case the ferromagnetic phase is replaced by an antiferromagnetic one, whereas the striped phase is not changed.

The frustrated Ising model has been studied for a long time, but some features of its phase diagram are still debates or were resolved only recently [95, 96, 97]. For $g < 1/2$, the transition is of the standard Ising type, but is difficult to study close to $g = 1/2$, due to large scaling corrections and long Monte Carlo autocorrelation times. Here we will consider only $g > 1/2$, for which there is a first-order transition between a high-temperature paramagnet and a low-temperature striped phase, up to a coupling $g^* \approx 0.8$ [97] after which the transition becomes continuous. We analyze results obtained with the standard single-spin Metropolis algorithm (as cluster Monte Carlo methods do not work in the presence of frustration). Temperatures will be quoted in units of $J_1$. 

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Figure 19. Order parameter distributions in the \((m_x, m_y)\) plane (in the full space \(|m_x| \leq 1, |m_y| \leq 1\)) of the 2D frustrated Ising model of size \(L = 128\) at coupling ratio \(g = 0.55\). Brighter features correspond to higher probability density. The temperatures (indicated in the panels in units of \(J_1\)) are in the first-order transition region, with the phase coexistence (a central peak as well as four peaks corresponding to \(x\)- and \(y\)-oriented stripes) seen most clearly at \(g = 0.772\).

Order-parameter distribution. The ordered phase can have its stripes oriented either along the \(x\) or \(y\)-axis, with the corresponding order parameters

\[
m_x = \frac{1}{N} \sum_{i=1}^{N} \sigma_i (-1)^{x_i}, \quad m_y = \frac{1}{N} \sum_{i=1}^{N} \sigma_i (-1)^{y_i},
\]

where \(x_i\) and \(y_i\) are the (integer) lattice coordinates of site \(i\). Let us first look at the order-parameter distribution \(P(m_x, m_y)\). If the four possible ordered states are sampled equally below \(T_c\), the distribution should be four-fold symmetric, with peaks located on the negative and positive \(x\)- and \(y\)-axis. In the paramagnetic phase there should be a single central peak. Coexistence at a first-order transition should hence be reflected in the presence of five peaks in this case (instead of the 3-peak distribution for the scalar order parameter in Fig. 18). Fig. 19 shows results for an \(L = 128\) system at \(g = 0.55\), for three temperatures in the transition window. A very distinct four-fold symmetry can be seen in all cases. Four symmetric peaks clearly indicate an ordered phase at \(T = 0.771\), but there are strong fluctuations, reflected in weight extending to the center of the distribution. At \(T = 0.773\), the distribution is peaked at the center, but weight also extends far into the ordered regions. Between these temperatures, at \(T = 0.772\), one can discern both the four ordering peaks and a central peak. All these plots show the hall-marks of coexistence; even when there are not five peaks present, there is still significant probability for both ordered and paramagnetic configurations. Away from the narrow transition window, the distribution rapidly turns into one with either distinctly ordered or disordered features.

Binder cumulant. Let us analyze the Binder cumulant, using \(m^2 = m_x^2 + m_y^2\) in Eq. (77). The question now is what order-parameter dimensionality \(n\) to use in this definition. At high temperatures, fluctuations of \(m_x\) and \(m_y\) in different parts of a large system are uncorrelated. This implies a rotationally-symmetric (circular) distribution \(P(m_x, m_y)\). The \(n\)-dependent factors are intended to make \(U_2 \to 0\) for \(T \to \infty\), and in this case we should therefore use \(n = 2\). This is correct also at low temperatures (to guarantee \(U_2 \to 1\)), because in an ordered phase the Binder ratio, Eq. (76), \(R_2 \to 1\) when the
system size diverges, regardless of the order parameter structure. Fig. 20 shows results as a function of temperature at two coupling ratios; \( g = 0.55 \) and 0.70. For \( g = 0.55 \), a negative Binder cumulant can be seen for \( L \geq 8 \), with the negative peaks becoming very narrow and apparently diverging as the system size is increased. At \( g = 0.70 \), the transition is still first-order, but with weaker discontinuities that start to manifest themselves as co-existence and a negative Binder cumulant only around size \( L = 32 \). In both cases, the Monte Carlo simulations were still ergodic, but a large number of updating sweeps (\( \approx 10^8 \) for each case) had to be carried out to obtain smooth curves.

**Discontinuities and finite-size scaling.** In first-order transitions that are not very strong, as in the systems above, it may not be easy to accurately extract the size of discontinuities in physical observables. An example is shown in Fig. 21, where the temperature dependence of the internal energy is graphed for several lattice sizes. While a discontinuity (latent heat) clearly develops with increasing \( L \), it is not easy to determine exactly between which two energies the jump will eventually take place. This requires a careful analysis using larger lattices.

Fig. 21 also shows the maximum value of the specific heat versus the system size for two different coupling ratios. At \( g = 0.51 \), the transition is rather strongly first-order, and a scaling consistent with the expected \( C_{\text{max}} \sim L^2 \) can be observed for the largest lattices. In contrast, at \( g = 0.55 \) the behavior appears to follow a different power law (with an exponent close to 1.2) up to \( L \approx 64 \). For larger lattices the results start to show a somewhat more rapid divergence, however, and eventually, for very large lattices, one would expect the exponent to be equal to \( d = 2 \) also in this case.

### 3.5. Spin stiffness and the Kosterliz-Thouless transition

An important aspect of a system with long-range magnetic order is that it exhibits a non-zero spin stiffness. For a system with continuous vector spins (XY or Heisenberg
FIGURE 21. Left: Internal energy versus temperature for the frustrated Ising model at $g = 0.55$. The discontinuity developing with increasing lattice size corresponds to the latent heat. Right: Finite-size scaling of the peak value of the specific heat at $g = 0.51$ and 0.55. The line shows the expected asymptotic $L^2$ scaling at a first-order transition.

models), the spin stiffness is the analogue of an elastic modulus of a solid. It is also often refereed to as the helicity modulus [131]. To study this concept, we here consider the XY model, with the hamiltonian written as

$$H = -J \sum_{\langle i, j \rangle} \cos(\Theta_i - \Theta_j),$$

(81)

where $\Theta_i \in [0, 2\pi]$ is the angle characterizing spin $i$. For simplicity we assume only nearest-neighbor interactions, but generalizations to arbitrary interactions are straightforward. When deriving an expression for the spin stiffness, we will consider the 2D square lattice (and some times use a 1D chain for simplicity), but later will study also a 3D cubic system. In 2D XY systems, the spin stiffness is an important quantity characterizing the unconventional (topological) Kosterlitz-Thouless transition exhibited by this system, which we also discuss briefly in this section.

3.5.1. Definition of the spin stiffness

Loosely speaking, the spin stiffness $\rho_s$ characterizes the tendency of ordered spins to adapt in response to perturbations imposing modulations of the direction of the order parameter (in contrast to the susceptibility, which measures the tendency of the order parameter to change in response to a field applied in a fixed direction). It is analogous to the shear modulus in continuum mechanics, which characterizes the tendency to shape deformation of an elastic object (while the compressibility corresponds to the tendency to volume change with maintained shape). The definition of the spin stiffness is easiest to understand at $T = 0$, which we consider first, before generalizing to $T > 0$.

The stiffness at $T = 0$. We will first consider a system with open boundaries in the $x$ direction, while periodic boundaries in the $y$ direction can be assumed (and later we
FIGURE 22. A 2D classical XY model with a phase twist in the x-direction imposed by fixing the spins in the boundary column at a relative angle $\Phi$. To minimize the energy the total twist $\Phi$ is distributed evenly, so that the spins in neighboring columns are twisted by $\phi = \Phi / L_x$ relative to each other.

will generalize to periodic boundaries also in the x direction). Fig. 22 illustrates how ferromagnetic XY spins at $T = 0$ adapt in order to minimize the energy when an over-all change in the spin angle $\Phi$ between the left and right boundaries is imposed (e.g., due to strong magnetic fields applied at the boundary columns). To minimize the energy, each column is twisted with respect to the following column by an angle $\phi = \Phi / (L_x - 1)$, where $L_x$ is the length of the system in the x direction (the number of columns). At $T = 0$, we are interested in the energy in the presence of this twist, which for the 2D XY model is simply given by

$$E(\phi) = -J(L_x - 1) L_y \cos(\phi) = E(0) + J(L_x - 1) L_y [1 - \cos(\phi)].$$

(82)

For small $\phi$ we get $E(\phi) - E(0) = (J/2)(L_x - 1)L_y\phi^2$ to leading-order. Motivated by this result, the $T = 0$ spin stiffness is defined as

$$\rho_s = \frac{1}{N} \frac{\partial^2 E(\phi)}{\partial \phi^2},$$

(83)

and we have $\rho_s = J$ for large $N$ [or for any $N$ if we normalize by the number $L_y(L_x - 1)$ of interacting x bonds, over which the energy cost due to the twist is distributed].

Normally it is more convenient to consider a periodic system. To derive an expression for the spin stiffness in this case, a phase twist is first imposed at the boundary. For simplicity we work this out for a 1D chain, with spin angles $\Theta_x, x = 0, \ldots, N - 1$, but the calculation can be trivially generalized to higher dimensionality. The interaction energy for each bond is $E_x = -J \cos(\Theta_{x+1} - \Theta_x)$, except at the boundary, where a twist $\Phi$ corresponds to $E_N = -J \cos(\Theta_0 - \Theta_{N-1} + \Phi)$. The configurations minimizing the energy have $\Theta_{x+1} - \Theta_x = \delta$, where $\delta$ is independent of $x$, which gives the total energy

$$E(\Phi) = -(N - 1) \cos(\delta) - \cos(\Phi - [N - 1] \delta),$$

(84)

which is minimized by $\delta = \phi \equiv \Phi / N$, with $E(\phi) = E(0) + JN[1 - \cos(\phi)]$, and the spin stiffness defined according to (83) is again $\rho_s = J$.

It is useful to consider also another way of twisting the spins in a periodic system, by introducing a twist field $\Phi_x = x\phi$ in the hamiltonian. For a 1D system the energy for each bond in the presence of this field is $-J \cos(\Theta_{x+1} - \Theta_x + \Phi_{x+1} - \Phi_x)$. To treat the boundary correctly, $\Phi_x$ should be considered as a function of a continuous variable $x$.  

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which jumps discontinuously from $N\phi$ to 0 at $x = N$. The phase difference appearing in the XY interaction should then be interpreted as

$$\Phi_{x+1} - \Phi_x = \int_x^{x+1} \Phi_x dx = \int_x^{x+1} \phi dx = \phi,$$

which holds also at the boundary ($x + 1 = N$). Apart from factors, $\phi$ is analogous to a flux threading the ring. The energy in the presence of this twist is minimized for $\Theta_x = \Theta$ independently of $x$, giving $E(\phi) = E(0) + JN[1 - \cos(\phi)]$ as before in the case of the twisted boundary condition. One can consider the twist field as a way of transforming away the twisted boundary condition (rotating to the reference frame of the spins in the presence of the boundary twist), which makes many calculations easier in practice. We will work with the twist field from now on.

**The stiffness at $T > 0$.** Now we consider non-zero temperatures, in which case the spin stiffness is defined as

$$\rho_s = \frac{1}{N} \frac{\partial^2 F(\phi)}{\partial \phi^2},$$

where $F(\phi)$ is the free energy in the presence of a twist field (or, equivalently, a twisted boundary condition), which in turn is related to the partition function according to

$$F(\phi) = -\frac{1}{\beta} \ln[Z(\phi)].$$

For $T \to 0$, Eq. (86) clearly reduces to the ground-state energy definition (83).

Applying a twist field which only depends on $x$, $\Phi(x, y) = x\phi$, in the 2D XY model, we have the hamiltonian

$$H(\phi) = -J \sum_{\langle i,j \rangle_x} \cos(\phi + \Theta_j - \Theta_i) - J \sum_{\langle i,j \rangle_y} \cos(\Theta_j - \Theta_i),$$

where we can assume periodic boundary conditions in both directions (although in principle the $y$ boundary could be open). We can simplify the dependence on $\phi$ by using a standard trigonometric equality,

$$\cos(\phi + \Theta_j - \Theta_i) = \cos(\Theta_j - \Theta_i) \cos(\phi) - \sin(\Theta_j - \Theta_i) \sin(\phi),$$

which we expand to second order in $\phi$:

$$\cos(\phi + \Theta_j - \Theta_i) \to \cos(\Theta_j - \Theta_i)(1 - \phi^2) - \sin(\Theta_j - \Theta_i) \phi + O(\phi^3).$$

The hamiltonian in the presence of a small ($\phi \to 0$) twist can then be written as

$$H(\phi) \to H(0) + \frac{1}{2} \phi^2 H_x - \phi I_x,$$

where $H_x$ is the $x$-bond part of the hamiltonian at $\phi = 0$ and $I_x$ is the total spin “current” in the $x$ lattice direction:

$$I_x = J \sum_{\langle i,j \rangle_x} \sin(\Theta_j - \Theta_i).$$
The partition function in the presence of a small twist can then be written as

\[ Z(\phi) = \int d[\Theta] e^{-\beta H(\phi)} \]

\[ \rightarrow \int d[\Theta] e^{-\beta H(0)} [1 - \frac{1}{2} \beta \phi^2 H_x + \ldots ] [1 + \beta \phi I_x + \frac{1}{2} (\beta \phi I_x)^2 + \ldots ], \]

where the exponentials involving \( H_x \) and \( I_x \) have been Taylor expanded to the orders needed. We can now write this in a form with expectation values over the distribution for \( \phi = 0 \). To second order in \( \phi \):

\[ Z(\phi) \rightarrow Z(0) [1 - \frac{1}{2} \beta \phi^2 \langle H_x \rangle + \beta \phi \langle I_x \rangle + \frac{1}{2} \beta^2 \phi^2 \langle I_x^2 \rangle]. \]

By symmetry \( \langle I_x \rangle = 0 \) and the free energy (87) is given by

\[ F(\phi) = F(0) + \frac{1}{2} \phi^2 \left( \langle H_x \rangle - \beta \langle I_x^2 \rangle \right), \]

and from this we can extract a simple expression for the spin stiffness (86):

\[ \rho_s = \frac{1}{N} \left( \langle H_x \rangle - \beta \langle I_x^2 \rangle \right), \]

which can be evaluated using Monte Carlo simulations. This result is for a twist field in the \( x \) lattice direction. For a \( d \)-dimensional isotropic system, we can average over all the equivalent directions and write the spin stiffness as

\[ \rho_s = \frac{1}{Nd} \left( \langle H \rangle - \beta \sum_{a=1}^{d} \langle I_a^2 \rangle \right), \]

where the index \( a \) corresponds to the current in lattice direction \( a \). For an anisotropic \( d \)-dimensional system, the stiffness in general is different for all lattice directions, i.e., there are \( d \) different stiffness constants, each of them given by a form like (96).

The twist-field definition of the stiffness can be easily generalized to systems with longer-range interactions. The forms (96) and (97) remain valid, with the current \( I_x \) containing contributions from all interactions exactly as in the Hamiltonian.

**Relation to superfluidity and superconductivity.** A very interesting aspect of the spin stiffness of the XY model is that it can be directly related to the superfluid density of a superconductor (or a superfluid such as \( ^4 \text{He} \)) [131, 132]. Like the magnetization of the XY model, the order parameter of a superconductor or a superfluid is \( U(1) \) symmetric (corresponding to the global phase of the wave function), and the twist field we discussed above is directly analogous to a magnetic flux (in the case of a superconductor). A non-zero stiffness corresponds to the Meissner effect exhibited by a superconductor. Monte Carlo simulations of the 2D XY model directly aimed at properties of thin superfluid films are discussed in Refs. [133, 134].
Spin stiffness scaling. The critical scaling properties of the spin (or superfluid) stiffness were first worked out in the context of superfluids [131, 132]. In the infinite $d$-dimensional system with $d > 2$, it was shown that

$$
\rho_s \sim (T_c - T)^{(d-2)\nu},
$$

for $T \to T_c$ from below. Here $\nu$ is the standard correlation length exponent. According to the general finite-size scaling relation (66), the size dependence of the stiffness exactly at $T_c$ is then given by

$$
\rho_s \sim L^{2-d}.
$$

The stiffness is therefore, like the correlation length and the Binder ratio, a useful quantity for locating the critical point without having to adjust any unknown (or not precisely known) exponents. This is illustrated with Monte Carlo results for the 3D XY model in Fig. 23, where $L\rho_s$ is size independent at the critical point.

3.5.2. Kosterlitz-Thouless transition in two dimensions

In the 2D XY model there can be no transition into a phase with long-range magnetic order at $T > 0$, according to the Mermin-Wagner theorem [38] (as discussed in Sec. 2). Remarkably, this system exhibits a different kind of phase transition, where no long-range order develops but the spin correlations change from exponentially decaying to a power-law form [135, 136]. This Kosterlitz-Thouless (KT) transition is topological in nature, being a consequence of proliferation of unbound vortices (which are topological defects) in the spin configurations at temperatures $T > T_{KT}$. For $T \leq T_{KT}$, vortices also exist (at a density which vanishes at $T \to 0$) but they are all bound in vortex-antivortex pairs, which have no net vorticity (and therefore vanish upon course graining of the spins). The power-law form of the spin correlations, $C(r) \sim r^{-\eta}$, applies for all $0 < T \leq T_{KT}$. The exponent $\eta$ is temperature dependent, with the value $\eta = 1/4$ exactly at $T_{KT}$ and $\eta \to 0$ as $T \to 0$, so that true long-range order exists at $T = 0$. 
Although there is no long-range order for $0 < T \leq T_{\text{KT}}$, the spin stiffness is actually non-zero in the KT phase—power-law correlations with a sufficiently small exponent are enough to support an energy cost of a boundary twist. There is no power-law onset of the stiffness, but instead an even more prominent signal of the transition; a discontinuous jump at $T_{\text{KT}}$ from $\rho_s = 0$ to a non-zero value. Renormalization-group calculations for the continuum field theory corresponding to the 2D XY model have given very detailed information about the KT transition. One of the most important results is a rigorous relationship (due to Nelson and Kosterliz [139]) between the transition temperature $T_{\text{KT}}$ and the spin stiffness exactly at this temperature (i.e., the size of the discontinuity);

$$\rho_s(T_{\text{KT}}) = \frac{2T_{\text{KT}}}{\pi}. \quad (100)$$

For finite lattices, the stiffness at $T_{\text{KT}}$ approaches the infinite-size value with a logarithmic size correction [138]:

$$\rho_s(T_{\text{KT}}, L) = \rho_s(T_{\text{KT}}, \infty) \left(1 + \frac{1}{2\ln(L) + c}\right), \quad (101)$$

where $c$ is a system dependent parameter. Thus, one can say that the general finite-size scaling law (99) for $\rho_s$ at a critical point holds, but with a logarithmic correction to the leading-order size-independent form obtaining when $d = 2$.

Fig. 24(a) shows Monte Carlo results for the stiffness of the 2D XY model versus temperature. The jump expected in the thermodynamic limit is approached very slowly as a function of size. This can be related to the fact that the correlation length for $T > T_{\text{KT}}$ does not diverge as a power-law, but according to the exponential form

$$\xi \sim e^{a/(T - T_{\text{KT}})^{1/2}}, \quad (102)$$

where $a$ depends on details of the lattice and the interactions. Therefore, using the same arguments as we in the case of a standard critical point in Sec. 3.3.2, the finite-size shift of $T_{\text{KT}}$ [defined using, e.g., the temperature $T^*(L)$ at which $\rho_s$ drops the most rapidly] is given by the form,

$$T^*(L) - T_{\text{KT}} \sim \frac{1}{\ln^2(L)}, \quad (103)$$

which is much slower than the conventional power-law shift $T^*(L) - T_c \sim L^{1/\nu}$.

Using the standard finite-size scaling hypothesis (64) and replacing $L^\sigma$ (the size dependence of the singular quantity exactly at the critical point) by the logarithmic size correction in (101), we can write a hypothesis for the size and temperature dependence of the spin stiffness at the KT transition as

$$\rho_s(T, L) = \left(1 + \frac{1}{2\ln(L) + c}\right) f\left(e^{a/(T - T_{\text{KT}})^{1/2}} / L\right), \quad (104)$$

which, after taking the logarithm of the argument of the scaling function $f(x)$, can be written in terms of another function $g[\ln(x)]$ as

$$\rho_s(T, L) \left(1 + \frac{1}{2\ln(L) + c}\right)^{-1} = g[\ln(L) - a/(T - T_{\text{KT}})^{1/2}]. \quad (105)$$
Here $T \to T_{\text{KT}}$ corresponds to the argument $z = \ln(L) - a/(T - T_{\text{KT}})^{1/2} \to -\infty$.

The KT transition temperature of the 2D XY model has been extracted in different ways in many studies, e.g., in [133, 137]. Fig. 24(b) shows a test of the scaling form (105), using $T_{\text{KT}} = 0.8933$, as obtained in [137], and with the constants $a$ and $c$ adjusted to obtain the (approximately) best data collapse onto a common scaling function for system sizes $L = 8, 16, \ldots, 128$. The data are indeed well described by this form. This kind of plot confirms three different aspects of the KT transition (which were theoretically deduced at different stages of the history of the KT transition) at the same time; the exponentially divergent correlation length (102) [135, 136], the logarithmic correction (101) [138], and the Nelson-Kosterlitz relation (100) [139].

### 3.6. Quantum Phase Transitions

In the following sections of these notes, we will study several examples of quantum phase transitions, which take place in the ground state of a system as a function of some model parameter [1]. The scaling properties we have discussed above for classical continuous and first-order transitions still apply, with some important extensions and modifications. The nature of a classical thermal phase transition can be traced to singularities in the free energy. At a quantum phase transition, it is instead the ground state energy which exhibits singular behavior, which is manifested also in other quantities. This can be understood as arising from the $T > 0$ free energy of the quantum system, which when $T \to 0$ becomes the ground state energy. That is of course also true classically, but the ground states of classical models normally do not evolve continuously as a function of
the parameters (as we saw in the example of the frustrated Ising model in Sec. 3.4.2) and are therefore strongly first-order. In contrast, as already discussed in Sec. 2.4, quantum systems have non-trivial ground states, with zero-point fluctuations that evolve as the parameters are varied. Continuous phase transitions driven by divergent quantum fluctuations are common.

As we will see in connection with quantum Monte Carlo methods in Sec. 5, a $d$-dimensional quantum system can formally be mapped, using path integrals, onto an equivalent classical statistical-mechanics problem in $d + 1$ dimensions (albeit some times with a non-positive-definite distribution function). The size of the system in the new “imaginary time” dimension is the inverse temperature; $L_\tau = c / T$, where $c$ is a velocity. At $T > 0$, this dimension is finite, while the spatial dimensions can be infinite. The system is then $d$-dimensional with a “thickness” $L_\tau$. The strict $(d + 1)$-dimensionality applies when also $T \to 0$. In that case, a tunable parameter in the hamiltonian can play a role very similar to the temperature in a classical system. Interestingly, changing the temperature in this case is analogous to finite-size scaling in $L_\tau$ [5], which can be used to deduce finite-temperature scaling properties close to quantum-critical points [84].

In some cases, the $d$-dimensional quantum system has the low-energy properties of the same kind of classical system in $d + 1$ dimensions. This is the case, e.g., for the dimerized Heisenberg models discussed in Sec. 2.4. The low-energy physics of these models can be mapped onto a 3D classical Heisenberg model, which is normally done via continuum field theories, such as the nonlinear $\sigma$-model [5, 84] (and, it, should be noted, the dimerization is only a means of tuning the strength of the quantum fluctuations, which in a course-grained uniform effective model is just represented by the coupling constant of a field theory, or the temperature in an effective 3D uniform classical model [140]). One can then presume that the quantum phase transition driven by tuning the dimerization strength should be in the universality class of the temperature-driven transition of the 3D classical Heisenberg model. In other cases, the low-energy mapping may give an effective system that does not correspond to any familiar classical model, but one can still say that the system corresponds to some $(d + 1)$-dimensional effective model.

The dynamic critical exponent. In many cases, such as the dimerized Heisenberg models mentioned above, the time dimension arising in the mapping to $d + 1$ dimensions is equivalent (in an asymptotic sense) to the spatial dimensions. The scaling properties of such a system at a quantum phase transition are then obtained by just replacing $d$ by $d + 1$ in the critical correlation function (53), in hyperscaling relations such as the second Eq. (59), and in the scalin forms (98) and (99) of the spin stiffness. In other cases, the correlations in the new dimension may be fundamentally different from those in the spatial dimensions. The dynamic exponent $z$ relates the power-laws associated with spatial and temporal correlations, e.g., if the spatial correlation length $\xi$ diverges when some parameter $g$ is tuned to its critical value as $|g - g_c|^{-1/\nu}$, then the temporal correlation length diverges as $|g - g_c|^{z/\nu}$, and in classical scaling relations one should replace $d$ by $d + z$ for the quantum critical point.

The dynamic exponent derives its name from the fact that it also governs the dispersion $\omega(q) \sim q^z$ of excitations of wave-number $q$. An important aspect of this is that the finite-size excitation gap is obtained by setting $q \propto 1/L$, giving the gap scal-
ing $\Delta L \sim 1/L^z$. This result can be used directly in numerical calculations, and also has indirect consequences for the scaling properties of quantities that depend on the excitation spectrum (e.g., various susceptibilities). Extracting the dynamic exponent is an important aspect of computational studies of quantum phase transitions. Apart from this, quantum phase transitions, both continuous and first-order ones, can be analyzed with the same finite-size scaling methods as the classical transitions discussed above.

It should be noted that there is a dynamic exponent also in classical systems, but this does not come into play in equilibrium statistical mechanics (because if a system has a kinetic-energy part of the hamiltonian, the associated phase-space probability distribution cancels out in expectation values). The classical dynamic exponent depends on how dynamics is introduced into the system [141]. The Metropolis Monte Carlo algorithm for the Ising model is an example of classical dynamics (called Glauber dynamics), and the long-time decay of the autocorrelations at criticality is governed by an associated dynamic exponent. This dynamics does not, however, determine the equilibrium properties. In quantum mechanics, the dynamics cannot be separated out, but is an integral part of the equilibrium properties, because the full hamiltonian always enters and contains in it the static as well as dynamic properties [142].

4. EXACT DIAGONALIZATION METHODS

By exactly diagonalizing its hamiltonian, complete knowledge of a quantum spin system can be obtained—with the eigenstates available, any static or dynamic quantity can be computed. In principle, all eigenstates can be computed exactly for a finite quantum system, by constructing the hamiltonian matrix and diagonalizing it numerically. In practice, however, such exact diagonalization studies are limited to rather small lattices, a few tens of spins, because of the exponential increase of the basis size with the number of spins ($2^N$ states in the case of $S = 1/2$). Great care therefore has to be taken in drawing conclusions about the thermodynamic limit, which may not even be possible (if the available lattices are too small to accommodate the infinite-size physics). Insights gained from exact diagonalization studies are nevertheless very useful, in their own right and as a complement to other calculations. Exact results for small lattices are also indispensable for testing the correctness of, e.g., quantum Monte Carlo programs. In addition, exact diagonalization methods provide a concrete path for learning many important aspects of quantum mechanics, in particular the symmetry properties of many-body states.

**Block diagonalization.** Given a hamiltonian $H$, the first step of an exact diagonalization calculation is to choose a basis in which it and other operators of interest will be expressed. The working basis for a spin-1/2 system normally consists of the single-spin states $\uparrow_i$ and $\downarrow_i$, $i = 1, \ldots, N$ (with the quantization direction normally taken as $z$). However, because of the $2^N$ growth of the Hilbert space with the number of spins in the system, symmetries should be used whenever possible to first reduce the hamiltonian to a block-diagonal form, as illustrated in Fig. 25. In such a scheme, the spin states are combined and ordered with the aid of applicable symmetry operations. The blocks correspond to states with different conserved quantum numbers related the symmetries, e.g.,...
crystal momentum conservation following from lattice translational symmetry or the conserved $z$-component of the total spin (the magnetization). The blocks can be diagonalized independently of each other, at a much reduced computational cost. In addition to the reduced computational effort, immediate access to the quantum numbers is very useful for classifying excitations.

Some symmetries are relatively easy to take advantage of (e.g., the conservation of the magnetization) whereas others require some more work and lead to more complicated basis states (e.g., momentum states). Some symmetries that could be used in principle are normally not implemented, because the practical complications of the calculation may outweigh the benefits. Total spin conservation is an example of this.

The use of symmetries in exact diagonalization can be discussed using the language of group theory [143]. This formalism is not necessary (and often confusing), however, and here a more practical approach is taken, with no reference to group theory terminology. Group theory is actually very useful when dealing with complex lattices, but the power of its formalism can perhaps be better appreciated after a thorough understanding of symmetry operations and block-diagonalization has been gained through less formal methods for simple lattices. Here we consider 1D chains and 2D simple square lattices.

Outline of this Section. Our discussion will first be framed around the $S = 1/2$ Heisenberg chain as a concrete example in Sec. 4.1. We first discuss its symmetries and introduce a computer representation of the basis states using bits of integers in 4.1.1. In 4.1.2 we use this representation in pseudocodes to construct the full hamiltonian without any symmetries, and then block-diagonalize using magnetization conservation. Momentum states are discussed in 4.1.3, the use of parity (reflection symmetry) in 4.1.4, and spin-inversion symmetry in 4.1.5. Complete diagonalization to obtain finite-temperature thermodynamic properties is illustrated with some results in 4.1.6. To compute a small number of eigenstates for larger chains, the Lanczos method is developed in 4.2. The utility of this technique for several 1D systems is illustrated in Sec 4.3. The ground state and low-energy excitations of the Heisenberg chain are discussed in 4.3.1. The dimerization transition taking place in the presence of a frustrating second-nearest-neighbor interaction is investigated in 4.3.2, and in 4.3.3 an extended model with long-range interactions is considered. Diagonalization of 2D systems is briefly described in 4.4. Momentum states and other square-lattice symmetries are considered. Results pertaining to

FIGURE 25. Schematic illustration of block diagonalization. In the original basis, the hamiltonian has no apparent structure (left). By constructing states labeled by a conserved quantum number, the matrix breaks up into blocks (with all matrix elements zero outside the shaded squares) that can be diagonalized independently of each other (middle). Applying another symmetry (conservation law), the blocks can be further broken up into smaller blocks (right) labeled by two different quantum numbers, etc.
the Néel ground state of this system are discussed.

4.1. Diagonalization of the Heisenberg chain

We will study the $S = 1/2$ antiferromagnetic Heisenberg chain with hamiltonian

$$H = J \sum_{i=0}^{N-1} \mathbf{S}_i \cdot \mathbf{S}_{i+1} = J \sum_{i=0}^{N-1} [S^z_i S^z_{i+1} + \frac{1}{2} (S^+_i S^-_{i+1} + S^-_i S^+_{i+1})],$$

(106)

where, for reasons that will become apparent below, it will here be practical to label the spins $i = 0, \ldots, N - 1$. Periodic boundaries, $S_N = S_0$, will be assumed when we consider momentum states, but before that the boundary condition is arbitrary.

4.1.1. Representations of states and symmetries

Lattice transformations. We use the standard notation $|S^z_0, \ldots, S^z_{N-1}\rangle$ for the basis states, with $S^z_i$ from left to right always corresponding to the spin states on lattice sites numbered 0, 1, 2, ..., irrespective of the ordering of the subscripts $i$. Thus, if we write a state $|S^z_1, S^z_0\rangle$, this is different from $|S^z_0, S^z_1\rangle$ unless $S^z_0 = S^z_1$. The former could refer to the state obtained from the latter when the two spins are switched by a permutation operator $P$, which can be operationally defined so as to affect the site indices; $P|S^z_0, S^z_1\rangle = |S^z_1, S^z_0\rangle$, e.g., $P|\uparrow\downarrow\rangle = |\downarrow\uparrow\rangle$. Generalizing this to $N$-spin lattice transformations, reflections, translations, and rotations (in two and three dimensions) are defined in terms of permutations of the spin indices. As an example, for a periodic chain we define the translation operator as moving the spins one step cyclically to the “right”;

$$T|S^z_0, S^z_1, \ldots, S^z_{N-1}\rangle = |S^z_{N-1}, S^z_0, \ldots, S^z_{N-2}\rangle.$$  

(107)

This corresponds to decreasing the spin index by one (modulus the system size $N$) at each location $i$ in the ket: $S^z_i \rightarrow S^z_{i-1}$. When writing specific states with up and down spins denoted by $\uparrow$ and $\downarrow$, e.g., $|\uparrow\uparrow\downarrow \uparrow\downarrow\ldots\rangle$, indices are normally not needed (and, for, the sake of compactness of the notation, the arrows are also not be separated by commas).

The Heisenberg hamiltonian (106) with periodic boundary conditions is invariant with respect to translations, i.e., it commutes with $T$: $[H, T] = 0$. We can therefore construct momentum states $|\Psi(k)\rangle$, which by definition are eigenstates of the translation operator,

$$T|\Psi(k)\rangle = e^{\imath k} |\Psi(k)\rangle.$$ 

(108)

Here the allowed momenta are $k = 2n\pi/N$, with $n = 0, \ldots, N - 1$, following from the fact that $T^N = 1$. States with different $k$ form their own individually diagonalizable blocks of the hamiltonian. How the momentum states are constructed in practice and used in a computer program will be discussed in Sec. 4.1.3.

The Heisenberg hamiltonian also commutes with the reflection (parity) operator, which we will define in a way generalizing the two-spin permutations already consid-
ered, in terms of the spin index transformation $i \to N - 1 - i$;

$$P|S_0^z, S_1^z, \ldots, S_{N-1}^z\rangle = |S_{N-1}^z, \ldots, S_1^z, S_0^z\rangle.$$  \hspace{1cm} (109)

For an eigenstate of $P, T|\Psi(p)\rangle = p|\Psi(p)\rangle$, where $p = \pm 1$ since $P^2 = 1$. We will use $T$ and $P$ for block-diagonalization, although, as we will discuss in detail in Sec. 4.1.4, they cannot always be used simultaneously because $[T, P] = 0$ only in a sub-space of the Hilbert space. For a system with open boundaries, $T$ is not defined, but $P$ can be used.

**Spin quantum numbers.** Since the Hamiltonian is spin-rotationally invariant, its eigenstates also have to be eigenstates of the square $S^2$ of the total spin, where

$$S = \sum_{i=0}^{N-1} S_i.$$  \hspace{1cm} (110)

For an eigenstate we have $S^2|\psi(S)\rangle = S \cdot S|\psi(S)\rangle = S(S + 1)|\psi(S)\rangle$. Here there is potential for confusion, as the same symbol $S$ is used for both the spin magnitude of the individual spins (i.e., $S_i = S$) and the total spin of a many-body state. The context should always make the meaning clear (and we anyway only consider $S_i = 1/2$ here).

With the total spin conserved, we know that the states can form blocks labeled by the quantum numbers $(S, m_z)$, where $m_z \in \{-S, -S+1, \ldots, S\}$ is the total magnetization in the direction of the quantization axis,

$$m_z = \sum_{i=0}^{N-1} S_i^z.$$  \hspace{1cm} (111)

If we use momentum states, each $k$-block of course also splits into smaller blocks, $(k, S, m_z)$, because $[T, S] = 0$. It is easy to block diagonalize using $m_z$, but implementing the conservation of total $S$ is normally cumbersome (except for a very small number of spins) and therefore rarely used (although $S = 0$ states in the valence-bond basis are some times used [144]). We will use $m_z$ conservation in combination with lattice symmetries. Note that $m_z$ conservation is more general than conservation of the total $S$—even if we introduce some anisotropy in the Hamiltonian by giving a different prefactor to, e.g., the Ising (diagonal) term in the Hamiltonian (106), $m_z$ is still conserved although total $S$ is not. All the techniques discussed in this section can therefore be applied directly also to such anisotropic models.

For the special (and most important) case $m_z = 0$ (for even $N$), we can block-diagonalize using a discrete subset of all the possible rotations in spin-space; the spin-inversion symmetry, i.e., invariance with respect to flipping all the spins. This is defined formally by an operator we call $Z$;

$$Z|S_0^z, S_1^z, \ldots, S_{N-1}^z\rangle = | -S_0^z, -S_1^z, \ldots, -S_{N-1}^z\rangle.$$  \hspace{1cm} (112)

For this operator we again have $Z^2 = 1$ and the eigenvalues $z = \pm 1$. Since $Z$ commutes with both $P$ and $T$, it can be used together with these operators to further block-diagonalize $H$, which we will do in Sec. 4.1.5.
The total-spin operator $S^2$ can be written in a form resembling the Heisenberg model with equal interactions among all the spins:

$$S^2 = \sum_{i=0}^{N-1} \sum_{j=0}^{N-1} S_i \cdot S_j = 2 \sum_{i<j} S_i \cdot S_j + \frac{3}{4}N.$$ \hspace{1cm} (113)

Constructing the matrix form of this operator, which we need for computing the quantum number $S$, is therefore very similar to constructing the hamiltonian matrix.

**Bit representation of spin states.** $S = 1/2$ models are special because $\downarrow$ and $\uparrow$ spins can be represented directly in the computer by the bit values 0 and 1 of an integer. We will take advantage of this here. The bits are conventionally labeled starting from 0, and that is why we also number the spins in that way. To refer to the bits $i = 0, \ldots, 31$ of an integer $s$ (or $i = 0, \ldots, 63$ for a “long” integer), we will use the notation $s[i]$. A basis state $|S_{\uparrow}^z, \ldots, S_{N-1}^z\rangle$ for $N$ spins is thus represented in the computer by an integer $s$ with $s[i] = S_i^z + 1/2$ for $i = 0, \ldots, N-1$ and $s[i] = 0$ for $i > N-1$.

Most computer languages have functions for examining and manipulating bits. The off-diagonal terms of the Heisenberg hamiltonian (106) flip two spins. In pseudocodes we will accomplish this operation using a bit function $\text{flip}(s, i, j)$, which flips $0 \leftrightarrow 1$ bits $i$ and $j$ of the integer $s$ representing the state. How this is implemented in practice depends on the language used. One possibility is to use a bitwise exclusive-or operation with a mask, as illustrated in Fig. 26. In Fortran 90, this operation can be implemented as $\text{ieor}(s, 2^{i+j})$. Later, we will also need functions that accomplish the various symmetry transformations of the states; translation, reflection, and spin-inversion.

With standard 4-byte integers, the bit representation with a single integer works up to $N = 32$. Using long integers, one can extend the scheme up to $N = 64$. The latter is well beyond the maximum size for which exact diagonalization techniques can be used in practice, except for magnetization sectors with $2m_z = n_\uparrow - n_\downarrow$ large enough for the block size $N!/(n_\uparrow!n_\downarrow!)$ to be manageable. Magnetization $m_z = 0$ and other low-$m_z$ sectors are typically of primary interest, however.

Discussing algorithms for constructing the basis set and the hamiltonian matrix, we will start from the simplest method, in which no symmetries at all are employed, and then implement the magnetization conservation. In Secs. 4.1.3-4.1.5 we include
more symmetries. The actual diagonalization of the hamiltonian matrix and the use its eigenstates to calculate physical observables will be deferred to Sec. 4.1.6.

4.1.2. Computer generation of the hamiltonian

If we do not make use of any conservation laws, the hamiltonian consists of a single $2^N \times 2^N$ matrix. We then simply use the numbers $a = 0, 1, \ldots, 2^N - 1$ to label the basis states. The bit-values of these integers correspond directly to the spin states. Determining the diagonal contributions $\langle a | S_i^z S_{i+1}^z | a \rangle = \pm 1/4$ to the hamiltonian matrix just involves examining the corresponding bit pairs $a[i], a[i+1]$ (same or different), whereas an off-diagonal operator $(S_i^+ S_{i+1}^- + S_i^- S_{i+1}^+)/2$ acting on a state $|a\rangle$ with $a[i] \neq a[i+1]$ generates the state $|b\rangle$ where the two bits have been flipped. The matrix element is then $\langle b | H | a \rangle = 1/2$. For a state with $a[i] = a[i+1]$ there is of course no off-diagonal matrix element. The following piece of pseudocode generates the full hamiltonian matrix $H$:

```pseudocode
do a = 0, 2^N - 1
  do i = 0, N - 1
    j = mod(i + 1, N)
    if (a[i] = a[j]) then
      H(a, a) = H(a, a) + 1/4
    else
      H(a, a) = H(a, a) - 1/4
      b = flip(a, i, j); H(a, b) = 1/2
    endif
  enddo
enddo
```

Here $j = \text{mod}(i + 1, N)$ is the “right” nearest neighbor of $i$, with the mod function taking care of the periodic boundary. In Fortran 90, the test for $a[i] = a[j]$ can be implemented with the boolean function btest(a, i) to examine bit $i$ of $a$. Each bond operator in the hamiltonian corresponds to a single off-diagonal matrix element, while the diagonal elements have $N$ different contributions. Matrices corresponding to other operators of interest can of course be constructed in an analogous way, by examining and flipping bits according to whatever combinations of $S_i^z$ and $S_i^\pm$ operators that are involved.

Using fixed-magnetization blocks. Moving up in sophistication, we next implement magnetization conservation. We want to construct the block hamiltonian acting on all states with given $m_z = (n_\uparrow - n_\downarrow)/2$. There are $M = N!/(n_\uparrow! n_\downarrow!)$ such states, and we need a list of them. The order within this list will be used as a label of the states of the block, i.e., we write $|a\rangle$ for the $a$:th state in the list. We then also need a list of integers $\{s_a\}$, the bits of which represent the spin configuration of the $a$:th state. Later, we will have to search the list $s_a$ to find the position label $a$ of a particular given state-integer $s$, and it is therefore practical to make the list ordered: $s_a < s_{a+1}$. We will some times use the notation $|s_a\rangle$ instead of $|a\rangle$ when referring explicitly to the spins, $|s_a\rangle = |s_a[0] - 1/2, \ldots, s_a[N-1] - 1/2\rangle$. The context will make it clear if a label inside
\(|\rangle\) refers to the position in the list of \(M\) states or to the integer containing the spins. To construct the state list, we loop over the integers \(s = 0, \ldots, 2^N - 1\) and check whether the number of set bits (the number \(n\uparrow\) of spins \(\uparrow\) in the state) corresponds to the target sector; \(n\uparrow = m_z + N/2\). After initializing a state counter \(a = 0\), we can use the following pseudocode to generate all the states with given magnetization:

\[
\begin{align*}
\text{do } s &= 0, 2^N - 1 \\
\text{if } (\sum_i s[i] = n\uparrow) \text{ then } a &= a + 1; s_a = s \text{ endif} \\
\text{enddo; } M &= a 
\end{align*}
\]

We now have a basis of size \(M\) stored as the integers \(s_a, a = 1, \ldots, M\).

To construct the Hamiltonian, we loop over the labels \(a = 1, \ldots, M\) and use bit operations as before to act on the corresponding state-integers \(s_a\). When an off-diagonal operation on \(|s_a\rangle\) leads to another state \(|s^*\rangle = |s_b\rangle\), and have to find the position \(b\) of the integer \(s^*\) in the state list. Since this list is ordered, we can do this by a bisectional search in \(\sim \log_2(M)\) steps on average. Such a search proceeds through a series of bracketings, where in each step we can halve the possible range of \(b\) by examining the state at the mid-point of a range \([b_{\text{min}}, b_{\text{max}}]\), with the brackets \(b_{\text{min}} = 1\) and \(b_{\text{max}} = M\) initially. The following subroutine finds the position \(b\) of a state-integer \(s^*\):

\[
\text{subroutine findstate}(s^*, b) \\
b_{\text{min}} = 1; b_{\text{max}} = M \\
\text{do} \\
b = b_{\text{min}} + (b_{\text{max}} - b_{\text{min}})/2 \\
\text{if } (s^* < s_b) \text{ then} \\
\quad b_{\text{max}} = b - 1 \\
\text{elseif } (s^* > s_b) \text{ then} \\
\quad b_{\text{min}} = b + 1 \\
\text{else} \\
\quad \text{exit} \\
\text{endif} \\
\text{enddo}
\]

Division of an integer by 2 should here be regarded in the standard way, i.e., \(i/2\) for odd \(i\) equals \((i - 1)/2\). The exit from the loop occurs when the basis state \(s_b\) equals the target state \(s^*\). One can also use more efficient search procedures, using so-called hash-tables [145]. The bisection is much simpler, however, and sufficiently fast in most cases.

Using subroutine \(\{4\}\), the part of the Hamiltonian originating from operation on a spin pair \((i, j)\) in the state \(|a\rangle\) can be constructed with the following modification of code \(\{2\}\):

\[
\begin{align*}
\text{if } (s_a[i] = s_a[j]) \text{ then} \\
\quad H(a, a) &= H(a, a) + \frac{1}{4} \quad \{5\} \\
\text{else} \\
\quad H(a, a) &= H(a, a) - \frac{1}{4} \\
\quad s^* &= \text{flip}(s_a, i, j); \text{ call findstate}(s^*, b); \\
\quad H(a, b) &= \frac{1}{2} \\
\text{endif}
\end{align*}
\]

If one is just interested in obtaining quick results for some very small lattice, or if the
system is not periodic (an open chain or a system with random couplings, in which case the momentum is not conserved), it may be sufficient to construct the Hamiltonian in this form and proceed to diagonalize it (successively for all the $m_z$-blocks desired). For serious work on translationally invariant (periodic) systems, it is worth implementing additional symmetries to further block-diagonalize the fixed $m_z$ blocks.

4.1.3. Momentum states

We now construct eigenstates of the translation operator $T$ defined Eq. (107). Translating $N$ steps brings the spins back to their original state. Thus, $T^N = 1$, which implies eigenvalues $e^{ik}$, where the set of $N$ non-equivalent momenta can be chosen as,

$$k = m \frac{2\pi}{N}, \quad m = -N/2 + 1, \ldots, N/2,$$

with the lattice constant equal to 1. A momentum state can be constructed using a reference state $|a\rangle$ (a single state in the $z$-component basis) and all its translations;

$$|a(k)\rangle = \frac{1}{\sqrt{N_a}} \sum_{r=0}^{N-1} e^{-ikr} T^r |a\rangle.$$

(115)

It can easily be verified (by a shift of the summation index allowed due to the periodic boundaries) that operating with the translation operator (107) on this state gives $T|a(k)\rangle = e^{ik}|a(k)\rangle$, which is the definition of a momentum state.

To construct the momentum basis for given $k$ (and normally also given $m^z$) we have to find a set of representatives resulting in a complete set of normalizable orthogonal states. Clearly, for two states $|a(k)\rangle$ and $|b(k)\rangle$ to be orthogonal, the corresponding representatives must obey $T^r |a\rangle \neq |b\rangle$ for all $r$. Therefore, among all the states of the set of translated states $|a(r)\rangle = T^r |a\rangle$, $r = 0, \ldots, N - 1$, only one should be used as a representative. With the labels referring to the bit representation, it will be practical to always choose the representative as the one for which the integer $a(r)$ is the smallest (as determined by carrying out all translations).

**Normalization and excluded representatives.** If all the translated states $T^r |a\rangle$ are distinct, the normalization constant in (115) is just $N_a = N$. Some reference states have periodicities less than $N$, however, and this affects the normalization. The periodicity of a state is defined as the smallest integer $R_a$ for which

$$T^{R_a} |a\rangle = |a\rangle, \quad R_a \in \{1, \ldots, N\}.$$

(116)

If $R < N$, then there are multiple copies of the same state in the sum in (115), and the normalization constant must be modified accordingly. We could then also restrict the summation in (115) to $r = 0, \ldots, T_a - 1$, but it is more practical in formal manipulations of the states to keep all $N$ terms regardless of $R_a$.

An important aspect of the momentum basis is that the periodicity of the representative has to be compatible with the momentum in order for (115) to be a viable state. The
compatibility is related to normalizability. The sum of phase factors associated with the representative state $|a\rangle$ in the sum in (115) is

$$F(k, R_a) = \sum_{n=0}^{N/R_a-1} e^{-i\pi n R_a} = \begin{cases} N/R_a, & \text{if } kR_a \text{ is a multiple of } 2\pi, \\ 0, & \text{otherwise}. \end{cases} \quad (117)$$

The normalization constant is then

$$N_a = \langle a(k)|a(k)\rangle = R_a|F(k, R_a)|^2, \quad (118)$$

and, therefore, if $F(k, R_a) = 0$, no state with momentum $k$ can be defined using the reference state $|a\rangle$. Thus, for given $|a\rangle$ the allowed momenta are those for which $kR_a$ is a multiple of $2\pi$, or

$$k = \frac{2\pi}{R_a} m, \quad m = 0, 1, \ldots, R_a - 1. \quad (119)$$

For the allowed momenta, the normalization constant in (115) is

$$N_a = \frac{N^2}{R_a}. \quad (120)$$

When the reference state is not equal to any non-trivial translation of itself, then $R_a = N$ and $N_a = N$. Note again that a given reference state can only appear in a single basis state. It is then clear that the momentum states are orthonormal; $\langle b(k')|a(k)\rangle = \delta_{ab} \delta_{kk'}$. Also note again that reference states that are not compatible with a given momentum will not appear in that block of states. An important aspect of constructing the momentum basis is to check the compatibility of a potential representative state with the momentum.

The Hamiltonian matrix. Next, we construct the Hamiltonian matrix in the momentum basis. The periodic Heisenberg Hamiltonian under consideration here is translationally invariant and consists of $N$ bond operators $S_i \cdot S_{i+1}$. It is convenient to lump all the diagonal terms together and consider the off-diagonal terms separately. To simplify the formalism to follow, we define operators accordingly;

$$H_0 = \sum_{j=1}^{N} S_{j}^z S_{j+1}^z, \quad (121)$$

$$H_j = \frac{1}{2}(S_{j-1}^+ S_j^- + S_j^+ S_{j+1}^-), \quad j = 1, \ldots, N, \quad (122)$$

so that $H = J \sum_{j=0}^{N} H_j$. We now set $J = 1$. We need to find the state resulting when $H$ acts on the momentum state (115). Since $[H, T] = 0$ we can write

$$H|a(k)\rangle = \frac{1}{\sqrt{N_a}} \sum_{r=0}^{N-1} e^{-i\pi r T} H|a\rangle = \frac{1}{\sqrt{N_a}} \sum_{j=0}^{N} \sum_{r=0}^{N-1} e^{-i\pi r T} H_j |a\rangle, \quad (123)$$

and we need to operate with the Hamiltonian operators $H_j$ only on the reference state. For each operation we get a different state, or, in the diagonal ($j = 0$) case, the same
state. In either case we can write \( H_j|a\rangle = h_j(a)|b'_j\rangle \), where \( h_j(a) \) is the matrix element coming from (121) or (122), and we do not, for simplicity of the notation, include any explicit indicator that \( |b'_j\rangle \) also depends on \( |a\rangle \). The prime in \( |b'_j\rangle \) is there to indicate that this new state is not necessarily one of the reference states used to define the basis and, therefore, a momentum state should not be written directly based on it. Provided that \( |b'_j\rangle \) is compatible with the momentum, there must be a reference state \( |b_j\rangle \) which is related to it by some number of translations;

\[
|b_j\rangle = T^{l_j}|b'_j\rangle, \quad (124)
\]

and using this relation we have

\[
H_j|a\rangle = h_j(a)T^{-l_j}|b_j\rangle, \quad l_j \in \{0, 1, \ldots, N - 1\}. \quad (125)
\]

Here the notation is again simplified by not making explicit that \( l_j \) depends on the actual state \( |b_j\rangle \) (and therefore on \( |a\rangle \)). We can now write (123) as

\[
H|a(k)\rangle = \sum_{j=0}^{N} \frac{h_j(a)}{\sqrt{N_a}} \sum_{r=0}^{N-1} e^{-ikrT^{(r-l_j)}}|b_j\rangle, \quad (126)
\]

and by shifting indices in the summation, and also noting that \( |b_j\rangle \) may have a different normalization factor (periodicity) than \( |a\rangle \), we obtain

\[
H|a(k)\rangle = \sum_{j=0}^{N} h_j(a)e^{-ikl_j} \sqrt{\frac{N_{b_j}}{N_a}}|b_j(k)\rangle. \quad (127)
\]

We can now simply extract the matrix elements of the hamiltonian operators \( H_j \):

\[
\langle b_j(k)|H_j|a(k)\rangle = h_j(a)e^{-ikl_j} \sqrt{\frac{N_{b_j}}{N_a}}, \quad (128)
\]

Strictly speaking, for the off-diagonal bond operators \((j > 0)\), this is not the only matrix element, because individually these are not translationally invariant operators. They therefore have matrix elements also between states with different momenta. Here we have in mind summing over all \( j \), after which only the elements diagonal in \( k \) survive.

There can be several terms in \( H \) that contribute to the same matrix element (128), because it is possible (in fact very likely) that \( H_j|a\rangle \propto T^{-l_j}|b_i\rangle \) and \( H_j|a\rangle \propto T^{-l_j}|b_j\rangle \) with \( |b_i\rangle = |b_j\rangle \) (but \( l_i \neq l_j \)). Note also that the matrix element \( h_j(a) \) in (125) is zero for an off-diagonal operator acting on two parallel spins. It should also be kept in mind that \( |b_j(k)\rangle \) may not exist even if \( h_j(a) > 0 \) in in (125), if \( |b_j\rangle \) is incompatible with the momentum. The momentum-state matrix element (128) then does not exist. With these caveats, which we will deal with in the implementations below, (128) specifies all the non-zero matrix elements of the hamiltonian. For the specific Heisenberg model considered here, we can substitute \( h_j(a) \) by the actual values of the diagonal and off-
FIGURE 27. Bit representation of the 8-spin state $|s\rangle = |↑↓↑↑↓↑\rangle$ (top row) along with its translations, $T^r|s\rangle$, with $r = 1, \ldots, R$, where $R = 4$ is the periodicity. The integers $s(r)$ correspond to $r$ successive cyclic permutations (to the left) of the bits. The representative is $s(1) = 51$; the lowest integer in the set.

Table: Bit representation of the 8-spin state $|s\rangle$ and its translations $T^r|s\rangle$

| $r$ | $s(r)$ | $T^r|s\rangle$ |
|-----|--------|---------------|
| 0   | 153    | 1|0|0|1|1|0|0|1 |
| 1   | 51     | 0|0|1|1|0|0|1|1 |
| 2   | 102    | 0|1|1|0|0|1|1 |
| 3   | 204    | 1|1|0|0|1|1|0|0 |
| 4   | 153    | 1|0|0|1|1|0|0|1 |

Matrix elements of other translationally invariant operators (which is what should be used in the momentum basis) can be obtained in exactly the same way if the momentum transferred by the operator is 0 (i.e., a sum of identical local operators $O_i$, $i = 0, \ldots, N - 1$). For an operator $A_q$ which transfers momentum $q \neq 0$ (such as $S_q^z$, the Fourier transform of $S_i^z$) the procedures differ only in that the basis states obtained when operating, $A_q|a(k)\rangle$, have momentum $k + q$ (and in that case the basis sets for both momenta involved have to be stored).

Constructing the momentum-state basis in a program. We will now again use a label $a$ to refer to the position of the state in the basis, and store the corresponding spins in the form of the bits of integers $s_a$, $a = 1, \ldots, M$. A momentum state is defined in terms of its representative $|s_a\rangle$, from which the full state is generated using the translation operator $T$ according to (115). In this case we do not know the basis size $M$ a priori, because the number of compatible representatives depends on the momentum. If all the translations of all states were unique, and there were no states incompatible with the momentum, then the number of states would equal $[N!/(n↑!n↓!)]/N$. However, since many states have periodicities less than $N$ and some are incompatible with $k$ (unless $k = 0$), this is only an approximate basis size. The $k = 0$ momentum block is always the largest, because there are no states incompatible with this momentum. For large systems the fractions of disallowed states and periodicities $< N$ are small, and the above approximation to the basis size is then quite good.

To implement the momentum basis in a computer program, we first need to decide how the representatives are chosen—in principle the representative of a state $|s\rangle$ could be any one of the members of the group of states related to it through translations; $|s(r)\rangle = T^r|s\rangle$, $r = 0, \ldots, N - 1$. Fig. 27 shows an example of all the unique translations of an 8-spin state in the bit representation. We have to construct a list of the representatives.
and should be able to easily identify the representative corresponding to an arbitrary spin state. As already mentioned, when using the bit representation it is natural to pick as the representative the translated state $|s(r)\rangle$ for which the integer $s(r)$ is the smallest.

To generate the basis, we loop over the integers $s = 0, \ldots, 2^N - 1$, and, as in code \{3\}, process only those corresponding to a chosen magnetization. The operations needed to determine whether a state is a valid new representative are carried out by bit operations. When translating $|s\rangle$, if some $s(r) > 0 < s = s(0)$, then the representative of $|s\rangle$ is already in the basis and should not be used again [as in Fig. 27, where the original integer $s = s(0) = 153$ but the translation $s(1) = 51 < s(0)$]. If all $s(r) \geq s(0)$, then the representative of $|s\rangle$ is not yet in the list. However, $|s\rangle$ may still not be allowed, due to its periodicity potentially being incompatible with the momentum, according to (119).

In the pseudocode segments below, the momentum will be represented just by the integer $\in \{-N/2 + 1, \ldots, N/2\}$ multiplying $2\pi/N$ in (114), which we here call $k$. We will define a subroutine which checks whether or not a state-integer $s$ is a new valid representative and, for a valid representative, delivers its periodicity $R$. To carry out the translations of the state $|s\rangle$, we use cyclic permutations of its bits. We assume that this is accomplished with a function `cyclebits$(i, n)$`, which performs a cyclic permutations to the “left” (as exemplified in Fig. 27) of the $n$ first bits of the integer $i$. In Fortran 90, exactly this operation is available with the intrinsic function `ishftc$(s, d, n)$`, which cyclically right-shifts the first $n$ bits of $s$ by $d$ steps. The definition of $T$ according to (107) corresponds to $d = -1$. Our state checking subroutine can be implemented as:

```fortran
subroutine checkstate(s, R)
    R = -1; t = s
    do i = 1, N
        t = cyclebits(t, N)
        if (t < s) then
            return
        elseif (t = s) then
            if (mod(k, N/i) /= 0) return
            R = i; return
        endif
    enddo
enddo
```

Here the integer $R$ is first initialized to $-1$, and this value will be returned (and later used as an indicator of a disallowed representative) unless the input state-integer $s$ is the smallest among all the translated integers and the periodicity is compatible with the momentum (and then $R$ will equal the periodicity upon return). We can now easily construct the list of $M$ basis states:

```fortran
do s = 0, 2^N - 1
    call checkstate(s, R)
    if R \geq 0 then a = a + 1; s_a = s; R_a = R endif
enddo
M = a
```

The list $R_a$ of periodicities will be needed when constructing the hamiltonian.
Constructing the hamiltonian matrix. To generate the hamiltonian, we loop over the representatives $|s_a\rangle$, $a = 1, \ldots, M$. For each of them, we check the bits corresponding to all nearest-neighbor pairs $(i, j)$. The diagonal matrix element can be handled exactly as in code {5}. For the off-diagonal part, we need a few minor modifications. After two spins have been flipped, the resulting state $|s'\rangle$ is typically not a representative, and we need to find it using Eq. (124). The matrix element (130) also requires the number $l_j$ of translations used to bring $|s'\rangle$ to its representative $|s_b\rangle$. We implement these tasks as a subroutine, the contents of which are rather similar to the subroutine checkstate, code {6}, that we used when constructing the basis. Given the state-integer $s$, we translate its bits in all possible ways and store the corresponding smallest integer $r$ (the potential representative) found so far, along with the corresponding number of translations, $l$;

```
subroutine representative(s', r, l)
  r = s'; t = s'; l = 0
  do i = 1, N - 1
    t = cyclebits(t, N)
    if (t < r) then r = t; l = i endif
  enddo
Having found the representative $r$, we need to locate its position $b$ in the list $\{s_a\}$. This is done in the same way as before, with the subroutine findstate, in code {4}. Note, however, that because of the periodicity constraint imposed by the momentum, it is now possible that the potential representative $r$ is actually not present in the list. Therefore findstate is slightly modified, so that $b = -1$ is returned if there is no element $r$ in the list, using the following piece of code after the if...endif statements in code {4};

```
if (b_{min} > b_{max}) then
  b = -1; return
endif
```

For each spin flip, we can now add the contribution to the hamiltonian with the following code replacing the next-to-last statement in code {5};

```
s' = flip(s_a, i, j)
call representative(s', r, l)
call findstate(r, b)
if (b \geq 0) H(a, b) = H(a, b) + \frac{1}{2}(R_a/R_b)^{1/2}\exp(i2\pi kl/N)
```
The hamiltonian is complex, except when $k = 0$ or $N/2$ (actual momentum 0 or $\pi$).

4.1.4. Reflection symmetry and semi-momentum states

We will next consider in addition to the translated states $T^r|a\rangle$ those that are generated by the reflection (parity) operator $P$ defined in (109). The operators $T$ and $P$ do not commute, and so it would at first sight appear that we cannot construct states that are eigenstates of both operators simultaneously. These operators do, however, commute in the $k = 0, \pi$ momentum blocks, as we will show explicitly below. In addition, we will
construct *semi-momentum* states that are also parity eigenstates for any \( k \). An advantage of such states is that they (and the hamiltonian) are real-valued, in contrast to the standard complex momentum states.

*States with parity.* Consider the following extension of the momentum state (115);

\[
|a(k, p)\rangle = \frac{1}{\sqrt{N_a}} \sum_{r=0}^{N-1} e^{-ikr} T^r (1 + pP) |a\rangle,
\]

(131)

where \( p = \pm 1 \). Clearly, this is a state with momentum \( k \) [i.e., it satisfies Eq. (108)], but is it also an eigenstate of \( P \) with parity \( p \)? We can check this by explicit operation with \( P \), using \( P^2 = 1, p^2 = 1 \), and the relationship \( PT = T^{-1}P \):

\[
P |a(k, p)\rangle = \frac{1}{\sqrt{N_a}} \sum_{r=0}^{N-1} e^{-ikr} T^{-r} (P + p) |a\rangle
\]

\[
= p \frac{1}{\sqrt{N_a}} \sum_{r=0}^{N-1} e^{ikr} T^r (1 + pP) |a\rangle.
\]

(132)

This is not exactly of the form (131), unless \( k = 0 \) or \( \pi \), for which \( e^{ikr} = e^{-ikr} \) (i.e., the momentum \( k \) is equivalent to \( -k \), and there is no directionality associated with the state). Thus, in these two special cases, parity and translational invariance can be used simultaneously for block-diagonalization and \( |a(k, p)\rangle \) is indeed a momentum state with parity \( p \) (or, in other words, \([T, P] = 0\) in the sub-spaces with momenta \( k = 0 \) and \( \pi \)).

*Semi-momentum states and parity.* Except for the special cases \( k = 0, \pi \), parity cannot be used to further block diagonalize a momentum block of \( H \). We can, however, use parity in combination with the momentum in a different way, by mixing momentum states with \( \pm k \). We consider the sum and difference of these states;

\[
|a^\sigma(k)\rangle = \frac{1}{\sqrt{N_a}} \sum_{r=0}^{N-1} C_k^\sigma(r) T^r |a\rangle,
\]

(133)

where \( \sigma = \pm 1 \) and we have, for convenience, introduced a function \( C_k^\sigma(r) \);

\[
C_k^\sigma(r) = \begin{cases} 
\cos(kr), & \sigma = +1 \\
\sin(kr), & \sigma = -1
\end{cases}
\]

(134)

We we will here refer to \( k \) in (133) as the *semi-momentum*. Note that \( \sigma \) is not a conserved quantum number, just an indicator for how the momentum states have been combined into semi-momentum states. Strictly speaking, the special values \( k = 0, \pi \) are still conventional (crystal) momenta (and the \( \sigma = -1 \) states do not exist for these \( k \)) and only \( k \) in the range \( 0 < k < \pi \) should be referred to as a semi-momenta. We will here consider all \( 0 \leq k \leq \pi \) on the same footing (where it should be noted that only half of the first Brillouin zone is used, as the other half corresponds to the same states). The
normalization constant is different for the special cases;

\[ N_a = \left( \frac{N}{R_a} \right)^2 \sum_{r=1}^{R_a} [C_k^\sigma (r)]^2 = \frac{N^2 g_k}{2R_a}, \]  

(135)

where we have introduced the factor

\[ g_k = \begin{cases} 
1, & 0 < k < \pi, \\
2, & k = 0, \pi.
\end{cases} \]  

(136)

In practice, \( g_k \) will not matter here, because we are only considering matrix elements of \( H \), which are diagonal in \( k \). The \( g \)-factors therefore cancel out in the ratios of normalization constants appearing in the matrix elements.

Checking overlaps, states with the same \( k \) but different \( \sigma \) are orthogonal,

\[ \langle a^-\sigma (k) | a^\sigma (k) \rangle = \frac{1}{N_a} \sum_{r=1}^{R_a} \sin(kr) \cos(kr) = 0, \]  

(137)

and other requirements for orthonormality, \( \langle a^\tau (k') | a^\sigma (k) \rangle = \delta_{\sigma \tau} \delta_{kk'} \), of the semi-momentum basis can also easily be verified. The advantage of semi-momentum states is that they are real-valued for all \( k \), in contrast to the complex momentum states.

The following equalities—standard trigonometric identities—are useful when manipulating semi-momentum states;

\[ C_k^\pm (-r) = \pm C_k^\pm (r), \]  

(138)

\[ C_k^\pm (r + d) = C_k^\pm (r) C_k^\pm (d) \mp C_k^{\mp} (r) C_k^{-} (d), \]  

(139)

where \( \pm \) stands for \( \sigma = \pm 1 \). Using (139) it is easy to see that the Hamiltonian acting on a semi-momentum state mixes \( \sigma = \pm 1 \) states. With \( H_j |a\rangle = h_j (a) T^{-l_j} |b_j\rangle \) we get

\[ H |a^\pm (k)\rangle = \sum_{j=0}^{N} h_j (a) \sqrt{\frac{R_a}{R_{b_j}}} \left( C_k^+ (l_j) |b_j^+ (k)\rangle \mp C_k^- (l_j) |b_j^- (k)\rangle \right), \]  

(140)

from which the Hamiltonian matrix elements can be extracted and written as

\[ \langle b^\tau (k) | H_j | a^\sigma (k) \rangle = h_j (a) \tau^{(\sigma - \tau)/2} \sqrt{\frac{N_{b_j}}{N_a}} C_k^\sigma \tau (l_j). \]  

(141)

**Incorporating parity.** Since the Hamiltonian (141) is not diagonal in \( \sigma, \tau \), the number of states in a semi-momentum block is twice that in a conventional momentum block, and thus it would appear that there is not much to be gained over complex momentum states by making Hamiltonian real in this way (by making the states real). However, we can also incorporate parity in a semi-momentum state, by defining

\[ |a^\sigma (k, p)\rangle = \frac{1}{\sqrt{N_a}} \sum_{r=0}^{N-1} C_k^\sigma (r) (1 + pP) T^r |a\rangle. \]  

(142)
This is also a semi-momentum state, because the reflected component \( P|a^\sigma(k)\rangle = \sigma|Pa^\sigma(k)\rangle \) is the semi-momentum state obtained by using the reflected representative state \( P|a\rangle \) instead of \( |a\rangle \). It can also easily be verified that (142) is an eigenstate of the parity operator for any semi-momentum: \( P|a^\sigma(k,p)\rangle = p|a^\sigma(k,p)\rangle \) with \( p = \pm 1 \). This is simply due to the fact that in (142) the operator \((1 + pP)\) appears before the translation operators \( T^r \), in contrast to the momentum state (131) where \((1 + pP)\) is written after \( T^r \). The Hamiltonian is thus diagonal in \( p \), and the number of states in each \((k,p)\) block is roughly half of that in the original semi-momentum \( k \) blocks. We are then back to the same block size as with the conventional momentum states, but with states with purely real coefficients.

Orthogonality and normalization of semi-momentum states. Examining the orthogonality of the states (142), we have an apparent problem: When \( T^mP|a\rangle = |a\rangle \) for some \( m \), the states \( |a^+(k,p)\rangle \) and \( |a^-(k,p)\rangle \) are not orthogonal. We can then use Eqs. (138) and (139) to write the parity-conserving semi-momentum state (142) as a linear combination of non-parity semi-momentum states (133);

\[
|a^\pm(k,p)\rangle = \sqrt{\frac{N_a}{N_a^\pm}} \left( (1 \pm pC_k^+(m))|a^\pm(k)\rangle - pC_k^-(m)|a^\mp(k)\rangle \right). \tag{143}
\]

It is then clear that \( \langle a^{-\sigma}(k,p)|a^\sigma(k,p)\rangle \) can be non-zero. To look at this more closely, we first determine the normalization constant \( N_a^\sigma \) in (142). Note that we have attached \( \sigma \) as a superscript to indicate that the normalization constant of the parity-conserving semi-momentum states can, unlike the normalization (135) of the plain semi-momentum states, depend on \( \sigma \) (in addition to the implicit dependence on \( k, p \)). In the case \( T^mP|a\rangle \neq |a\rangle \) for all \( m \), the calculation of \( N_a^\sigma \) is trivial. When \( T^mP|a\rangle = |a\rangle \) for some \( m \) we can use (143) and the orthonormality of the pure semi-momentum states \( |a^\sigma(k)\rangle \), resulting in

\[
N_a^\sigma = \frac{N_a^2g_k}{R_a} \times \begin{cases} 1, & T^mP|a\rangle \neq |a\rangle \ \forall m, \\ 1 + \sigma p \cos(km), & T^mP|a\rangle = |a\rangle. \end{cases} \tag{144}
\]

In the same way, we can calculate the overlap between the \( \sigma = \pm 1 \) states when \( T^mP|a\rangle = |a\rangle \) for some \( m \) and find

\[
\langle a^+(k,p)|a^\mp(k,p)\rangle = -p, \quad (T^mP|a\rangle = |a\rangle \text{ for some } m). \tag{145}
\]

This is of course under the assumption that both the \( \sigma = \pm 1 \) states exist (i.e., \( N_a^\sigma \neq 0 \)). Thus, the \( \sigma = \pm 1 \) states in the case \( T^mP|a\rangle = |a\rangle \) differ at most by a sign, and we should include only one of them in the block of states. For definiteness, we can choose \( |a^+(k,p)\rangle \) if \( N_a^+ \neq 0 \) and \( |a^-(k,p)\rangle \) else. We still have to pay attention to the non-zero overlap (145) in some formal manipulations with the semi-momentum states, as we shall see shortly.

The semi-momentum Hamiltonian. To calculate the Hamiltonian matrix elements, we first note that, instead of (124) for a plain momentum state, now when we act with a Hamiltonian operator \( H_J \) on \( |a\rangle \) we get a state \( |b^J_a\rangle \) which is related to another
representative state \(|b_j\rangle\) by a number of translations and possibly a reflection:

\[
|b_j\rangle = T^{l_j} P^{q_j} |b'_j\rangle, \quad l_j \in \{0, \ldots, N-1\}, \quad q_j \in \{0, 1\}. \quad (146)
\]

The result of \(H\) acting on a semi-momentum state (or a \(k = 0, \pi\) momentum state with parity) can therefore be written in in the form

\[
H|a^\sigma(k, p)\rangle = \sum_{j=0}^{N} h_j(a)(\sigma p)^{q_j} \frac{N^\sigma_{b_j}}{N^\sigma_a} \sum_{r=0}^{N-1} C_k^\sigma (r+l_j)(1+pP)T^r|b_j\rangle. \quad (147)
\]

where the representative \(|b_j\rangle\) is related to \(H_j|a\rangle\)

\[
H_j|a\rangle = h_j(a)P^{q_j}T^{-l_j}|b_j\rangle, \quad l_j \in \{0, \ldots, N-1\}, \quad q_j \in \{0, 1\}. \quad (148)
\]

Using the relation (139) we can write (147) as

\[
H|a^\sigma(k, p)\rangle = \sum_{j=0}^{N} h_j(a)(\sigma p)^{q_j} \sqrt{\frac{N^\sigma_{b_j}}{N^\sigma_a}} \times \\
\left( \cos(kl_j)|b_j^\sigma(k, p)\rangle - \sigma \sqrt{\frac{N_{b_j}^{-\sigma}}{N_{b_j}^\sigma}} \sin(kl_j)|b_j^{-\sigma}(k, p)\rangle \right). \quad (149)
\]

The ratio of the \(\sigma = \pm 1\) normalization constants is 1 if \(T^m P|b_j\rangle \neq |b_j\rangle\) for all \(m\), and otherwise it can be written as

\[
\sqrt{\frac{N_{b_j}^{-\sigma}}{N_{b_j}^\sigma}} = \sqrt{\frac{1 - \sigma p \cos(km)}{1 + \sigma p \cos(km)}} = \frac{|\sin(km)|}{1 + \sigma p \cos(km)}. \quad (150)
\]

Using the overlap (145) we can extract the matrix elements. The ones diagonal in \(\sigma\) are,

\[
\langle b_j^\sigma(k, p) | H_j | a^\sigma(k, p) \rangle = h_j(a)(\sigma p)^{q_j} \sqrt{\frac{N_{b_j}^\sigma}{N^\sigma_a}} \times \\
\left\{ \begin{array}{ll}
\cos(kl_j), & P|b_j\rangle \neq T^m|b_j\rangle, \\
\frac{\cos(kl_j) + \sigma p \cos(kl_j - m)}{1 + \sigma p \cos(km)}, & P|b_j\rangle = T^m|b_j\rangle,
\end{array} \right.
\quad (151)
\]

whereas the off-diagonal ones are

\[
\langle b_j^{-\sigma}(k, p) | H_j | a^\sigma(k, p) \rangle = h_j(a)(\sigma p)^{q_j} \sqrt{\frac{N_{b_j}^{-\sigma}}{N^\sigma_a}} \times \\
\left\{ \begin{array}{ll}
-\sigma \sin(kl_j), & P|b_j\rangle \neq T^m|b_j\rangle, \\
\frac{-\sigma \sin(kl_j) + \sigma p \sin(kl_j - m)}{1 - \sigma p \cos(km)}, & P|b_j\rangle = T^m|b_j\rangle.
\end{array} \right.
\quad (152)
\]
Degeneracies. In the conventional momentum basis, states with \( \pm k \) are degenerate, but are orthogonal states (except when \( k \neq 0, \pi \)). In the semi-momentum basis, \( \pm k \) states are really the same state, differing at most (in the case \( \sigma = -1 \)) by a sign (and that is why the semi-momentum Hilbert space only includes \( 0 \leq k \leq \pi \)). The degeneracy has here been moved to the parity sector, with the \( p \pm 1 \) states being degenerate (but still orthogonal) for \( k \neq 0, \pi \). This can be seen by letting \( (p, \sigma, \tau) \to -(p, \sigma, \tau) \) in the matrix elements (152), which leaves unchanged the set of matrix elements for \( \tau, \sigma = \pm 1 \). Thus, in calculations we need to consider only, e.g., \( p = +1 \) for \( k \neq 0, \pi \) (while for \( k = 0, \pi \) we only have \( \sigma = \tau = 1 \) and the \( p = \pm 1 \) sectors are not degenerate).

Constructing the basis of semi-momentum states. Generalizing the convention we used for momentum states, we now choose as the representative of a state \( |s\rangle \) the state \( |s(r, q)\rangle = T^r P^q |s\rangle \) for which the corresponding integer \( s(r, q) \) is the smallest (where \( r \in \{0, \ldots, N - 1\}, q \in \{0, 1\} \)). States can now be disallowed in a given block not only due to incompatibility with the momentum, but also because of restrictions imposed by the reflection quantum number \( p \) and the non-conserved state label \( \sigma \). We thus have to determine which one, or both, of the \( \sigma = \pm 1 \) semi-momentum states (142) should be included, according to the conditions discussed above. Fig. 28 illustrates all the combination of translations and reflections in the bit representation of a state which is related to its own reflection according to \( |s\rangle = T^m P |s\rangle \), in which case only the \( \sigma = -1 \) or \( \sigma = +1 \) variant of the state should be included in the basis (and the state is compatible with all momenta, because its periodicity \( R = N \)).

When both \( \sigma = \pm 1 \) states are required, we will store two consecutive copies of the same representative, so that we can continue to use the location in the list \( s_d \) as the label of each state within the block. We then also need to store the \( \sigma \) labels for each representative, as well as the number of translations \( m \) by which the reflection of a state is brought back onto itself, which is needed in the normalization and the hamiltonian.

As before, we will use a subroutine \texttt{checkstate} to determine whether or not a state is a new representative to be added to the list. In addition to the periodicity \( R \) of a new representative, this subroutine now also delivers the reflection-translation number \( m \), when applicable. We can use \( m = -1 \) as a flag indicating that there is no \( m \) for which

\[
\begin{array}{c|ccc|ccc}
  r & T^r & T^r P & T^r P \\
  0 & 27 & 0 & 0 & 0 & 1 & 0 & 1 & 1 & 1 & 1 & 0 & 0 & 0 & 0 & 0 & 0 \\
  1 & 54 & 0 & 0 & 1 & 1 & 1 & 0 & 1 & 0 & 1 & 0 & 0 & 0 & 0 & 0 & 1 \\
  2 & 108 & 0 & 1 & 1 & 0 & 1 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 1 & 1 \\
  3 & 216 & 1 & 1 & 0 & 1 & 1 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 1 & 1 \\
  4 & 177 & 1 & 1 & 0 & 1 & 1 & 1 & 0 & 0 & 0 & 0 & 0 & 0 & 0 & 1 & 1 \\
  5 & 99 & 0 & 1 & 1 & 0 & 0 & 1 & 1 & 1 & 0 & 1 & 0 & 0 & 0 & 0 & 1 \\
  6 & 198 & 1 & 1 & 0 & 0 & 0 & 0 & 0 & 1 & 1 & 1 & 0 & 0 & 0 & 0 & 1 \\
  7 & 141 & 1 & 0 & 0 & 0 & 0 & 1 & 1 & 1 & 0 & 1 & 0 & 0 & 0 & 0 & 1 \\
\end{array}
\]

FIGURE 28. A state \( |s\rangle = |\uparrow\uparrow\downarrow\uparrow\downarrow\downarrow\rangle \) (upper left) along with all its transformations by the translation, \( T \), and reflection, \( P \), operator in the bit representation. The corresponding base-10 integer \( s \) and its transformations are also shown—the smallest integer, \( s = 27 \), corresponds to the representative state. For this particular state, \( T^5 P |s\rangle = |s\rangle \), i.e., \( m = 5 \) in the normalization constant (144).
$T^m P|s\rangle = |s\rangle$ (and otherwise $m \geq 0$). The operations on the reflected state are carried out after the instructions in \{6\} have been performed, by first reflecting the spin bits in the state integer $s$ and storing it as $t, t[i] = s[N - 1 - i]$ for $i = 0, \ldots, N - 1$. This reflection is accomplished by a function reflectbits$(s, N)$. Then the resulting integer is translated as before. We thus modify code \{6\} according to:

```fortran
subroutine checkstate(s, R, m)
  ...
  t = reflectbits(s, N); m = -1
  do i = 0, R - 1
    if (t < s) then
      R = -1; return
    elseif (t = s) then
      m = i; return
    endif
    t = cyclebits(t, N)
  enddo
  ...
end subroutine checkstate
```

Here ... represents all operations in code \{6\}. Note that the loop in the code above starts at $i = 0$, and the translations are carried out after comparing $t$ with $s$, since a state can be its own reflection (in which case $m = 0$). Also, since we have already determined the periodicity $R$ of the state, we need to go only up to $i = R - 1$ in this loop. The function reflectbits has to be implemented by hand; a corresponding internal function is normally not available. In Fortran 90 one can use the functions btest$(i, b)$, setbit$(i, b)$, and clrbit$(i, b)$, to examine, set, and clear individual bits $b$ of an integer $i$.

A state which has passed the above checks is still not necessarily a valid representative for both $\sigma = \pm 1$, and the state must therefore be further examined to determine which (one or both) of the $\sigma = \pm 1$ copies of the representative should be added to the basis. The following code segment stores the state information for $\sigma = \pm 1$ states that are to be included in the block based on the criteria discussed above:

```fortran
call checkstate(s, R, m)
  do $\sigma = \pm 1$ (do only $\sigma = +1$ if $k = 0$ or $k = N/2$)  \{12\}
    if ($m \neq -1$) then
      if ($1 + \sigma p \cos(ikm2\pi/N) = 0$) $R = -1$
      if ($\sigma = -1$ and $1 - \sigma p \cos(ikm2\pi/N) \neq 0$) $R = -1$
    endif
    if $R > 0$ then $a = a + 1$; $s_a = s$; $R_a = \sigma R$; $m_a = m$ endif
  enddo
```

Recall that here $m = -1$ means that there is no $m$ such that $T^m P|s\rangle = |s\rangle$, whence both $\sigma = \pm 1$ states should be included. On the other hand, if there is such an $m$, then only one of the states is included—we pick $\sigma = 1$ if the normalization constant (144) for it is non-zero. This is accomplished above by the two consecutive if statements (where $R = -1$ is set for an invalid representative). We have to store $\sigma$, and since the periodicity $R$ is positive and non-zero, we can store it and $\sigma$ jointly as $\sigma R$ in a list $R_a$. A more dense packing of $R, \sigma, m$ into a single integer is of course also possible.
Constructing the semi-momentum hamiltonian. For general semi-momentum, the same representative can appear once or twice in the list \( s_a \), whereas for the special cases \( k = 0, \pi \) there is always only a single (\( \sigma = 1 \)) copy of each representative. We here construct code that can treat both cases, but simpler code can be written for the two special momenta. When building the hamiltonian by looping over the state indices \( a \), with the corresponding representatives \( s_a \), we first check the previous and next items in the list. If the previous item \( s_{a-1} = s_a \), then we skip over this representative, because it will already have been taken care of in the previous pass through the loop. If the next representative, \( s_{a+1} \) equals \( s_a \), then we store as \( n = 2 \) the number of same representatives, and else set \( n = 1 \). This part of the hamiltonian building is accomplished by:

\[
\begin{align*}
\text{do } a &= 1, M \\
\quad \text{if } (a > 1 \text{ and } s_a = s_{a-1}) \text{ then } & \quad \{13\} \\
\quad & \quad \text{skip to next } a \\
\quad \text{elseif } (a < M \text{ and } s_a = s_{a+1}) \text{ then } & \\
\quad & \quad n = 2 \\
\quad \text{else } & \\
\quad & \quad n = 1 \\
\text{endif} \\
\end{align*}
\]

Here the \texttt{skip} command skips to the next iteration of the loop and \ldots stands for the bulk of the loop, the main features of which we discuss next.

Considering first the diagonal matrix elements \( H(a, a) \), for a given \( a \) we now first sum up all the diagonal contributions from the bit configurations \( s_a \) and store the result as \( E_z \). Taking into account the possibility of one or two entries of the representative (\( n = 1 \) or 2), we can use the following loop to assign the diagonal matrix elements accordingly:

\[
\begin{align*}
\text{do } i &= a, a + n - 1 \\
\quad H(a, a) &= H(a, a) + E_z \\
\text{mando} \\
\end{align*}
\]

Turning next to the off-diagonal operations, looping over site pairs \( (i, j) \), if the bits \( s_a[i] \neq s_a[j] \), then the spins are flipped, resulting in a state-integer named \( s' \) as in code \{10\}. We again have to identify the corresponding representative state \( |r⟩ = T^l P^q |s'⟩ \) by carrying out all the symmetry operations to determine the transformation indices \( l \) and \( q \) appearing in the hamiltonian matrix elements (151) and (152). The subroutine accomplishing this is a simple modification of code \{8\}:

\[
\begin{align*}
\text{subroutine representative}(s', r, l, q) \\
\quad \ldots \\
\quad t &= \text{reflectbits}(s', N); \quad q = 0 \\
\text{do } i &= 1, N - 1 \\
\quad t &= \text{cyclebits}(t, N) \\
\quad \text{if } (t < r) \text{ then } r = t; \quad l = i; \quad q = 1 \text{ endif} \\
\text{mando} \\
\end{align*}
\]
Here ... stands for the code after the header in \{8\}. After having called this subroutine, we can search for the representative \( r \) in the list \( \{s_a\} \). This is done in the same way as we did before, with the subroutine \texttt{findstate}(r,b).

Since the hamiltonian is more complicated than in the standard momentum basis, we also define a function \texttt{helement}(a,b,l,q), which returns the matrix element according to (151) or (152). The function takes as input the labels \( a \) and \( b \) (locations in the matrix) of the two representatives and the transformation indices \( l, q \) delivered by \texttt{representative}. The matrix element also of course depends on the momentum \( k \) and the parity quantum number \( p = \pm 1 \), which we do not indicate explicitly here. The function is straightforward to implement based on (151) and (152), and we do not list any code here.

As with the state-integer \( s_a \) in code segment \{13\}, we also here need to take into account that the representative \( s_b \) can appear once or twice (indicated below with \( m = 1 \) or 2) in the list of representatives. In case there are two copies, we do not know whether the subroutine \texttt{findstate} has returned its first or second location. Thus, we again need to examine positions in the list of representatives adjacent to the one delivered. The following code segment does all that and then assigns the matrix elements:

\[
\begin{align*}
& s' = \texttt{flip}(s_a,i,j) \\
& \texttt{call representative}(s',r,l,q) \\
& \texttt{call findstate}(r,b) \\
& \text{if } (b \geq 0) \text{ then} \\
& \quad \text{if } (b > 1 \text{ and } s_b = s_{b-1}) \text{ then} \\
& \quad \quad m = 2; b = b - 1 \\
& \quad \text{elseif } (b < M \text{ and } s_b = s_{b+1}) \text{ then} \\
& \quad \quad m = 2 \\
& \quad \text{else} \\
& \quad \quad m = 1 \\
& \text{endif} \\
& \text{do } j = b, b + m - 1 \\
& \text{do } i = a, a + n - 1 \\
& \quad H(i,j) = H(i,j) + \texttt{helement}(i,j,l,q) \\
& \text{endo} \\
& \text{endo} \\
& \text{endif}
\end{align*}
\]

This piece of codes replaces the much simpler four-line code segment \{10\} for the pure momentum basis. The advantage is that the matrix \( H \) is now real-valued.

### 4.1.5. Spin-inversion symmetry

Spin-inversion symmetry, with the operator \( Z \) defined in (112), can be used to reduce the hamiltonian block size in the magnetization sector \( m^z = 0 \). The fact that the magnetization is conserved implies, however, that there is nothing to be gained by using this symmetry in the \( |m_z| > 0 \) sectors (since such a basis would consist of mixed \( \pm |m_z| \) states, and the blocks of symmetric and anti-symmetric combinations would be of the same size.
as the original blocks). In models that do not conserve \( m_z \), e.g., the transverse-field Ising model, spin inversion can be exploited for all states. We denote by \( z \) the eigenvalue of \( Z \). Since \( Z^2 = 1 \), we again have \( z = \pm 1 \).

Unlike the parity operation considered in the previous section, the spin-inversion operator \( Z \) in (112) commutes with the translation operator, \([T, Z] = 0\). The associated quantum number \( z \) is therefore conserved together with \( k \) in all momentum sectors. In the magnetization sector \( m_z = 0 \), we can therefore always split a momentum block into two smaller ones by using states of the form

\[
|a(k, z)\rangle = \frac{1}{\sqrt{N_a}} \sum_{r=0}^{N-1} e^{-ikr} T^r (1 + zZ) |a\rangle,
\]

(153)

where the normalization constant is easily obtained as

\[
N_a = \frac{2N^2}{R_a} \times \begin{cases} 
1, & T^mZ|a\rangle \neq |a\rangle \forall m, \\
1 + z\cos(km), & T^mZ|a\rangle = |a\rangle.
\end{cases}
\]

(154)

For a hamiltonian operation on \(|\alpha\rangle\) resulting in

\[
H_j|a\rangle = h_j(a) Z^{g_j} T^{-l_j} |b_j\rangle, \quad l_j \in \{0, \ldots, N-1\}, \quad g_j \in \{0, 1\},
\]

(155)

we obtain the matrix element

\[
\langle b_j(k, z)|H_j|a(k, z)\rangle = h_j(a) z^{g_j} e^{-ikl_j} \sqrt{\frac{N_{b_j}}{N_a}},
\]

(156)

which is valid for any \( k \). As always, for \( j = 0 \) this reduces to just \( h_0(a) \).

**Spin-inversion symmetry with semi-momentum states.** We can also consider semi-momentum states incorporating spin-inversion symmetry;

\[
|a^\sigma(k, p, z)\rangle = \frac{1}{\sqrt{N_a}} \sum_{r=0}^{N-1} C_k^\sigma (r)(1 + pP)(1 + zZ) T^r |a\rangle,
\]

(157)

These states are eigenstates of \( Z \) as well as \( P \).

When calculating the normalization and constructing the hamiltonian we now have to consider five different types of reference states, depending on which combinations of symmetry operations transform the reference state into itself:

1) \( T^mP|a\rangle \neq |a\rangle, \quad T^mZ|a\rangle \neq |a\rangle \quad T^mPZ|a\rangle \neq |a\rangle, \)

2) \( T^mP|a\rangle = |a\rangle, \quad T^mZ|a\rangle \neq |a\rangle \quad T^mPZ|a\rangle \neq |a\rangle, \)

3) \( T^mP|a\rangle \neq |a\rangle, \quad T^mZ|a\rangle = |a\rangle \quad T^mPZ|a\rangle \neq |a\rangle, \)

4) \( T^mP|a\rangle \neq |a\rangle, \quad T^mZ|a\rangle \neq |a\rangle \quad T^mPZ|a\rangle = |a\rangle, \)

5) \( T^mP|a\rangle = |a\rangle, \quad T^mZ|a\rangle = |a\rangle \quad \Rightarrow T^{m+n}PZ|a\rangle = |a\rangle. \)

(158)

Here the inequalities should hold for all \( m \) and the equalities for some \( m, n \). The calculations follow the procedures of the previous section, and we just list the results. The
normalization constants are:

\[ N_a^\sigma = \frac{2N^2}{R_a g_k} \begin{cases} 
1, & 1 \\
1 + \sigma p \cos(k m), & 2 \\
1 + \sigma p \cos(k m), & 3 \\
1 + \sigma p \cos(k m), & 4 \\
[1 + \sigma p \cos(k m)] [1 + \sigma \cos(k n)]. & 5 
\end{cases} \] (159)

For cases 2), 4), and 5), only one state out of a pair with \( \sigma = \pm 1 \) should be included in the basis as the two states within such a pair differ merely by a factor. To be definite, we can again pick \( \sigma = 1 \) if that makes \( N_a^\sigma > 0 \) and \( \sigma = -1 \) else. Acting with a hamiltonian operator on a representative now leads to a new representative \(|b_j\rangle\) according to

\[ H_j |a\rangle = h_j(a) P^j Z^j T^{-j} |b_j\rangle. \] (160)

The \( \sigma \)-diagonal matrix elements of \( H \) are

\[ \langle b_j^\sigma(k,p) | H_j | a^\sigma(k,p) \rangle = h_j(a) (\sigma p)^{g_j z^j} \sqrt{N_{b_j}^\sigma} \times \begin{cases} 
\cos(k l_j), & 1), 3), \\
\cos(k l_j) + \sigma p \cos(k[l - m]), & 2), 5), \\
\cos(k l_j) + \sigma p \cos(k[l - m]), & 4), 
\end{cases} \] (161)

whereas the ones off-diagonal in \( \sigma \) are,

\[ \langle b_j^{-\sigma}(k,p) | H_j | a^\sigma(k,p) \rangle = h_j(a) (\sigma p)^{g_j z^j} \sqrt{N_{b_j}^\sigma} \times \begin{cases} 
-\sigma \sin(k l_j), & 1), 3), \\
-\sigma \sin(k l_j) + ps \sin(k[l - m]), & 2), 5), \\
-\sigma \sin(k l_j) + ps \sin(k[l - m]), & 4), 
\end{cases} \] (162)

These expressions may seem rather complicated, but it should be noted that they are completely general for 1D systems, not just for the Heisenberg chain considered here. Once they have been implemented and tested for some model, the code can easily be reused for other systems.

Program implementation; combining all symmetries. Incorporating spin-inversion symmetry in the semi-momentum basis with \( m_z = 0 \), with states \(|a^\sigma(k,p,z)\rangle\) defined in (157), we have to augment the subroutine \texttt{checkstate} in code segment \{11\}. It should return the translation numbers corresponding not only to reflection \( P \) of a representative, \( T^{m_p} P |s\rangle = |s\rangle \), but also translation numbers in symmetry relationships involving \( Z \) and \( PZ; T^{m_z} Z |s\rangle = |s\rangle \) and \( T^{m_p} PZ |s\rangle = |s\rangle \). To accomplish the spin inversion in the bit representation, we define a function \texttt{invertbits}(s,N) which flips all the bits \( s[i] \rightarrow 1 - s[i] \), \( i = 0,N - 1 \). This can be easily accomplished without individual bit-level
TABLE 1. Size of the $k = 0$ state blocks for magnetization $m_z = 0$ and different parity and spin-inversion quantum numbers $(p, z)$.

<table>
<thead>
<tr>
<th>$N$</th>
<th>$(+1,+1)$</th>
<th>$(+1,-1)$</th>
<th>$(-1,+1)$</th>
<th>$(-1,-1)$</th>
</tr>
</thead>
<tbody>
<tr>
<td>8</td>
<td>7</td>
<td>1</td>
<td>0</td>
<td>2</td>
</tr>
<tr>
<td>12</td>
<td>35</td>
<td>15</td>
<td>9</td>
<td>21</td>
</tr>
<tr>
<td>16</td>
<td>257</td>
<td>183</td>
<td>158</td>
<td>212</td>
</tr>
<tr>
<td>20</td>
<td>2518</td>
<td>2234</td>
<td>2136</td>
<td>2364</td>
</tr>
<tr>
<td>24</td>
<td>28968</td>
<td>27854</td>
<td>27482</td>
<td>28416</td>
</tr>
<tr>
<td>28</td>
<td>361270</td>
<td>356876</td>
<td>355458</td>
<td>359256</td>
</tr>
<tr>
<td>32</td>
<td>4707969</td>
<td>4690551</td>
<td>4685150</td>
<td>4700500</td>
</tr>
</tbody>
</table>

operations as $s = 2^{N-1} - s$ (for $N$ less than the number of bits in the integers used; otherwise a different formula has to be used). Codes $\{6\}$ and $\{11\}$ can be easily modified to also check the inverted and reflected-inverted states to determine $m_z$ and $m_{zp}$.

If the modified checkstate returns a potential representative, $R \neq -1$, then again the state has to be examined further to determine whether it satisfies the further criteria for a representative for one or two states $\sigma = \pm 1$. An allowed representative now falls into one of five different classes depending on its symmetry properties as summarized in (158). The class, $c \in \{1, \ldots, 5\}$ can be determined from the translation integers $m_p, m_z, m_{zp}$ delivered by checkstate. After $c$ has been determined, the three translation integers can be reduced to two; the $m$ and $n$ in the normalization constants (159). The class $c$ can be packed along with $m, n$ into a single integer, e.g., as $m + n(N+1) + c(N+1)^2$, which is stored along with the periodicity and $\sigma$ index packed as $\sigma R$ as before.

For the construction of the hamiltonian, the subroutine representative, the main parts of which are described in code segments $\{8\}$ and $\{15\}$, has to be amended further in order to also return an index $g$ corresponding to the number, 0 or 1, of spin-inversions needed, along with $q$ reflections and $l$ translations, to transform the state $|s\rangle$ generated by a spin flip into the corresponding representative; $|r\rangle = T^l P^q Z^g |s\rangle$. The construction of the hamiltonian then proceeds exactly as before, with a modification only of the function helement to include also $g$ as an argument in code segment (5.15). This is straightforward, and there is no need to list any code here.

Examples of state block sizes. This is as far as we will go with applying symmetries for 1D systems. For the Heisenberg model, the total spin is also conserved, but incorporating that symmetry in the basis is much more complicated. In principle it can be done for total spin singlets, using the valence-bond basis [144], but it is rarely done in practice because the resulting hamiltonian is very dense, unlike the sparse matrices obtaining with the symmetries implemented here (the number of non-zero elements is proportional to $M \times N$ for an $M \times M$ matrix, and $N \ll N$). The sparseness will be very useful in the Lanczos calculations discussed in Sec. 4.2.

Table 1 shows the size of the Hilbert space blocks for several different chain sizes $N$ when all the symmetries are used. Here the momentum $k = 0$ and the number of states is shown for all combinations of the parity and spin-inversion quantum numbers ($p = \pm 1, z = \pm 1$). The largest block is always the fully symmetric one. As the system
size grows, the relative variations in the block size diminish, and for large $N$ all sizes are rather well approximated by $N!/[4(\frac{N}{2})^2N]$. For $k = \pi$, the blocks are approximately of the same size as in the table for $k = 0$, and for other momenta, where both $\sigma = 1$ and $-1$ states are allowed, the blocks are roughly twice as large.

### 4.1.6. Expectation values and thermodynamics

We are now ready to diagonalize the hamiltonian and calculate physical observables. Here we will first consider complete diagonalization, meaning that we compute all the eigenvalues and eigenstates of $H$. In principle, the eigenvalues $\lambda_n$, $n = 1, \ldots, M$, of a non-singular $M \times M$ matrix $H$ can be obtained by solving the secular equation,

$$\det[H - \lambda_n I] = 0,$$

(163)

where $\det[]$ denotes the determinant and $I$ is the unit matrix. The eigenvectors $v_n$ can subsequently be obtained by solving the linear system of equations

$$Hv_n = E_n v_n.$$

(164)

However, since the secular equation is very complicated for a large matrix, this method is not used in practice. Most numerical matrix diagonalization methods are based on an iterative search for a unitary transformation $U$ such that

$$U^{-1} H U = E,$$

(165)

where $E$ is the diagonal eigenvalue matrix with $E_{nn} = E_n$ and, for a complex hermitian matrix $H$, the inverse $U^{-1}$ of the unitary matrix $U$ is the transpose of its complex conjugate matrix; $U^{-1} = U^{*T}$. If the matrix is real and symmetric, we have $U^{-1} = U^T$.

The columns of the diagonalizing matrix $U$ contain the eigenvectors of $H$. This can be seen by multiplying (165) with $U$ from the left, giving $HU = UH$. Since $E$ is diagonal, the $n$th column of $U$ on the right side of this equality is multiplied by the $n$th eigenvalue (i.e., the matrix element $E_{nn} = \lambda_n$). In the multiplication by $H$ on the left side, the $n$th column of $U$ gives the $n$th column $(HU)_n$ of $HU$, i.e., $HU_n = E_{nn} U_n$, and thus the eigenvectors $v_n$ of $H$ are identified as $v_n = U_n$.

The expectation value of some observable (operator) $A$ in an eigenstate of $H$ is given by the diagonal element of the corresponding matrix transformed to the energy basis;

$$\langle n | A | n \rangle = [U^{-1} A U]_{nn}.$$

(166)

One would typically be interested in the ground state, $n = 1$ (assuming the eigenvalues to be ordered from the lowest to the highest), of the block in question (especially the actual ground state, in the block with the lowest energy of all). For a thermal average, all eigenvalues and/or eigenstates of all symmetry blocks are needed;

$$\langle A \rangle = \frac{1}{Z} \sum_{j} \sum_{n=1}^{M_j} e^{-\beta E_{jn}} [U_j^{-1} A_j U_j]_{nn}, \quad Z = \sum_{j} \sum_{n=1}^{M_j} e^{-\beta E_{jn}}.$$

(167)
TABLE 2. The eight lowest energies for a 16-site chain with momentum \( k = 0 \) in the blocks with parity and spin-inversion quantum numbers \((p = \pm 1, z = \pm 1)\).

<table>
<thead>
<tr>
<th>(n)</th>
<th>(E(+1,+1))</th>
<th>(S)</th>
<th>(E(+1, -1))</th>
<th>(S)</th>
<th>(E(-1,+1))</th>
<th>(S)</th>
<th>(E(-1,-1))</th>
<th>(S)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>-7.1422964</td>
<td>0</td>
<td>-4.9014133</td>
<td>1</td>
<td>-4.1926153</td>
<td>2</td>
<td>-5.7475957</td>
<td>1</td>
</tr>
<tr>
<td>2</td>
<td>-6.1223153</td>
<td>2</td>
<td>-4.1067293</td>
<td>1</td>
<td>-3.6537528</td>
<td>0</td>
<td>-4.9014133</td>
<td>1</td>
</tr>
<tr>
<td>3</td>
<td>-5.5912905</td>
<td>0</td>
<td>-3.9398439</td>
<td>1</td>
<td>-3.6085498</td>
<td>2</td>
<td>-4.6986358</td>
<td>1</td>
</tr>
<tr>
<td>4</td>
<td>-5.0981578</td>
<td>2</td>
<td>-3.7756347</td>
<td>1</td>
<td>-3.2192241</td>
<td>2</td>
<td>-4.1067293</td>
<td>1</td>
</tr>
<tr>
<td>5</td>
<td>-4.8142442</td>
<td>0</td>
<td>-3.6808576</td>
<td>1</td>
<td>-3.2129324</td>
<td>0</td>
<td>-4.0007340</td>
<td>1</td>
</tr>
<tr>
<td>6</td>
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<td>0</td>
<td>-3.5785191</td>
<td>3</td>
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<td>1</td>
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<td>7</td>
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<td>2</td>
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<td>1</td>
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<td>3</td>
<td>-3.1169772</td>
<td>2</td>
<td>-3.6808576</td>
<td>1</td>
</tr>
</tbody>
</table>

Here \( \beta = T^{-1} \) is the inverse temperature (in units where \( k_B = 1 \)), and the index \( j \) collectively denotes the different quantum numbers, \( m_z, k, p, z \), of the blocks of size \( M_j \).

Matrix diagonalization is a standard linear algebra operation, and sophisticated sub-routines are available in many software libraries. We will therefore not discuss the inner workings of matrix diagonalization procedures here. Almost always, a diagonalization routine delivers the eigenvalues in ascending order in a vector, along with the matrix \( U \) with the corresponding eigenvectors.

The number of operations needed to diagonalize an \( M \times M \) matrix generally scales as \( M^3 \), and the memory required for storage is \( \sim M^2 \) (even for a sparse matrix, as intermediate steps normally do not maintain sparsity). This severely limits the size of the matrices that can be fully diagonalized in practice. Currently \( M \approx 10^4 \) can be handled without too much effort on a workstation, and a few times larger on a supercomputer. Looking at Table 1, it is then clear that one cannot realistically carry out complete diagonalizations for Heisenberg chains larger than \( N \approx 20 \). Calculations aiming at just the ground state, and possibly some number of excited states, can be carried out in other ways for larger systems, using, e.g., the Lanczos method discussed in Sec. 4.2.

Total spin. It is useful to know the total spin \( S \) of the energy eigenstates. Since we do not use the conservation of \( S \) when block-diagonalizing the hamiltonian, we have to calculate \( S^2 \) using the states. As we have already noted, this operator is formally equivalent to a Heisenberg hamiltonian with long-range interactions, as written explicitly in (113). We can use a slightly modified version of the procedures we developed for constructing the hamiltonian to obtain the matrix for \( S^2 \). We then transform it with \( U \), after which we have \( S(S+1) \) in the form of the diagonal matrix elements as in (166).

Let us look at some results obtained for a 16-site chain in the \( k = 0 \) momentum sector. The eight lowest energies are listed for each of the symmetry sectors \((p, z)\) in table 2, along with the calculated spin of the states. The lowest-energy state is a singlet with \((p = 1, z = 1)\). The first excited state of the system is actually not in the \( k = 0 \) sector; it is a \( k = \pi \) triplet with \((p = -1, z = -1)\), at energy \( E = -6.87210668 \).

One interesting feature to note in the table is that, in every symmetry sector, all the states have either even or odd spin. This is due to the spin-inversion symmetry. An \( m_z = 0 \) state with \( N \) sites and total spin \( S \) can be written as a linear combination of states made up of \( N/2 - S \) singlets, \((|\uparrow_i \downarrow_j\rangle - |\downarrow_i \uparrow_j\rangle)/\sqrt{2}\), and \( S \) triplets, \((|\uparrow_i \downarrow_j\rangle + |\downarrow_i \uparrow_j\rangle)/\sqrt{2}\) [48].
Since the singlet is odd under spin inversion but the triplet is even, it follows that a spin \( S \) state of an \( N \)-site system has spin-inversion quantum number \( z = +1 \) if \( N/2 \) and \( S \) are both even or both odd, and has \( z = -1 \) else. Thus, even if we do not calculate \( S \) we have some limited knowledge of it also from \( z \). The low-energy states typically also have low spin, \( S = 0 \) or 1, and these can then be distinguished by \( z \) alone. Note, however, that the lowest state in a given \((p, z)\) sector does not necessarily have the lowest (even or odd) spin, as exemplified in Table 2 by the lowest \((p = -1, z = 1)\) level, which has \( S = 2 \), not \( S = 0 \). However, this is not a low-energy state, as there are 11 states with lower energy in the \( k = 0 \) sector (and numerous additional ones in other momentum sectors). The \( k = 0 \) state with second-lowest energy also has \( S = 2 \), but there are four \( S = 1 \) states below it in other momentum sectors.

**Magnetic susceptibility and specific heat.** As an example of thermodynamics, let us calculate two important properties of the Heisenberg chain; the specific heat \( C \) and the magnetic susceptibility \( \chi \). These can be evaluated according to the formulas,

\[
C = \frac{d\langle H \rangle}{dt} = \frac{1}{T^2} \left( \langle H^2 \rangle - \langle H \rangle^2 \right),
\]

\[
\chi = \frac{d\langle m^2 \rangle}{dh} = \frac{1}{T} \left( \langle m^2 \rangle - \langle m \rangle^2 \right),
\]

which can both be easily derived using (167). In the definition of \( \chi \), \( h \) is an external magnetic field in the \( z \)-direction, which couples through a term \(-hm_z\) added to the hamiltonian. We will here consider the zero-field case only, and thus \( \langle m_z \rangle = 0 \).

Both \( C \) and \( \chi \) are special quantities in the sense that they do not depend on the states, just the energy spectrum. For the Heisenberg model one can calculate these quantities by just considering the \( m_z = 0 \) sector, because of the conservation of the total spin \( S \). A spin \( S \) state is \((2S + 1)\)-fold degenerate with magnetization \( m_z = -S, \ldots, S \). In the case of \( C \), we therefore just have to sum 168 over the \( m_z = 0 \) levels and weight them by this degeneracy factor to obtain the full average over all \( m_z \). In the case of \( \chi_z \), we can proceed in the same way, using also the fact that, in a spin-\( S \) multiplet, the average \( \langle m_z^2 \rangle = \langle S^2 \rangle / 3 \). In (169).

The situation is slightly complicated by the fact that there can be “accidental” degeneracies not related to any apparent symmetry of the hamiltonian. If such degenerate states have different \( S \), then the diagonalization procedure will give some random (depending on exactly how the diagonalization is done) mixed spins for the states. One should then in principle diagonalize \( S^2 \) in the degenerate subspace to obtain the spin eigenvalues. However, in practice these accidental degeneracies are very rare. For small Heisenberg chains they occur only in the \( k = \pi, p = 1, z = 1 \) sector. For \( N = 10 \), there are two such states with \( E = 0.5 \), for \( N = 12 \) and 14 there are no accidental mixed-\( S \) degeneracies, and for \( N = 16 \) there are four mixed levels at energy \( 2J \). With such a small number of degeneracies, and at such high energies, we can just ignore them.

Fig. 29 illustrates how the results for \( C \) and \( \chi \) converge to the thermodynamic limit with increasing system size. Going up to \( N = 16 \), the results are well converged down to \( T/J \approx 0.25 \). It is natural that the convergence is more rapid at high temperatures, because in the limit \( T \to \infty \) the spins are statistically decoupled and the system behaves
FIGURE 29. Temperature dependence of the specific heat and the magnetic susceptibility for chains of length $N = 4, 8, 12, \text{ and } 16$. The thin curves show the leading high-temperature forms.

as a set of independent spins. Thus the susceptibility per spin for $T \to \infty$ has the single-spin Curie form; $\chi \to 1/4T$. For $C$, an analytic high-temperature form can be obtained by calculating the energy exactly for the 2-site system (a single spin has constant energy, thus giving $C = 0$), taking the derivative, and keeping only the leading term of an expansion in $T^{-1}$. This gives $C \to 3/13T^2$ when $T \to \infty$. The high-$T$ limits are also shown in Fig. 29. In the case of $C$, this form describes the behavior well down to $T/J \approx 1.5$, whereas in the case of $\chi$ the agreement is good only at higher temperatures.

When $T \to 0$, both $C$ and $\chi$ approach zero exponentially for a finite system, as a consequence of the finite gap between the ground state and excited states (due to which only the ground state contributes to thermal averages as $T \to 0$) and, in the case of $\chi$, the fact that the ground state of the Heisenberg chain for even $N$ is always a singlet (while for odd $N$ the ground state is an $S = 1/2$ doublet and $\chi$ thus diverges as $T^{-1}$). In the infinite system, $C \to 0$ but $\chi$ approaches a constant non-zero value.

Note that here, as well as in many cases we will encounter later on, when we talk about $T = 0$ properties it is important to consider the order in which the limits $T \to 0$ and $N \to \infty$ are taken. The susceptibility of a chain with finite (even) $N$ always vanishes as $T \to 0$. The temperature at which the exponential drop to zero commences depends on the finite-size spin gap, which in the case of the Heisenberg model is $\propto N^{-1}$ (which we will also discuss further and illustrate with data in Sec. 4.3.1). Thus, for given $N$, one can expect essentially thermodynamic-limit results down to some $T/J \propto N^{-1}$ (which can also be roughly seen in Fig. 29), and if one takes the limit $N \to \infty$ before $T \to 0$, then the susceptibility remains non-zero at $T = 0$.

It is not possible to use exact diagonalization results to extrapolate reliably the properties of the Heisenberg chain to the thermodynamic limit at low temperatures. For the susceptibility, an extrapolation of exact diagonalization results [146], using a particular functional form [147], was already attempted decades ago—the result is known as the Bonner-Fisher susceptibility. This extrapolation used also the known exact value of $\chi$ at $T \to 0$ ($N = \infty$), which was available from the exact Bethe ansatz solution of the ground state. The Bonner-Fisher curve has been commonly used in analyzing experimental results for quasi-one-dimensional antiferromagnets (weakly coupled chains). The most important feature here is the location and shape of the maximum at $T/J \approx 0.6$, which
is already well converged for small systems, as seen in Fig. 29. It has for some time now been possible to use other computational methods to study various properties of the Heisenberg model on very long chains (as we will discuss in Sec. 5.3.1), and thus the low-temperature limit can be accessed more reliably. There are also essentially exact results from field-theoretical studies combined with the Bethe ansatz [7]. These calculations show an anomalous feature; the approach to the $T \to 0$ limit is logarithmic, and therefore very difficult to reproduce with finite-$N$ calculations. The low-$T$ form of the Bonner-Fisher curve is therefore incorrect. Experimentally the most important feature is the broad maximum. Often effects beyond the Heisenberg chain appears at lower temperatures (e.g., due to inter-chain couplings, spin anisotropies, or disorder). The spin susceptibilities of some quasi-one-dimensional materials actually follows $\chi$ of the Heisenberg chain down to temperatures as low as $\approx J/50$, where the logarithmic behavior is prominent [8].

### 4.2. The Lanczos method

As we have seen in the preceding section, a complete diagonalization of the hamiltonian (or individual blocks) becomes prohibitively time consuming for $S = 1/2$ systems with more than $\approx 20$ spins. For higher $S$, the situation is of course even worse. If we restrict ourselves just to the ground state, and possibly a number of low-lying excitations, we can reach systems roughly twice as large, by using a Krylov-space technique, such as the Lanczos method.

#### 4.2.1. The Krylov space

The Krylov space is a sub-space of the full Hilbert space, constructed in such a way that the low-lying states of a hamiltonian $H$ of interest should be well approximated within it. Consider an arbitrary state $|\Psi\rangle$, e.g., a one with randomly generated vector elements $\Psi(i), i = 1, \ldots, M,$ in the $M$-dimensional Hilbert space in which $H$ is defined, and the expansion of this state in terms the hamiltonian eigenstates $|\psi_n\rangle$ (in order of increasing eigenvalues, which we here label $n = 0, 1, \ldots, M - 1$). We operate with a power of the hamiltonian on this chosen state;

$$H^\Lambda |\Psi\rangle = \sum_{n=0}^{M-1} c_n E_n^\Lambda |\psi_n\rangle = c_{\text{max}} E_{\text{max}}^\Lambda |\psi_{\text{max}}\rangle + \sum_{n \neq \text{max}} c_n \frac{E_n}{E_{\text{max}}}^\Lambda |\psi_n\rangle.$$  \hspace{1cm} (170)

If the power $\Lambda$ is large, the state corresponding to the eigenvalue $|E_{\text{max}}|$ with largest magnitude (i.e., max = 0 or max = $M - 1$) will dominate the sum, provided that the expansion coefficient $c_{\text{max}} \neq 0$. Hence, acting many times with the hamiltonian on the state will project out the eigenvector with the eigenvalue $E_{\text{max}}$. If we want to make sure that the ground state $|\psi_0\rangle$ is obtained (exactly for $\Lambda \to \infty$ or approximatively for finite $\Lambda$), we can instead of $H^\Lambda$ use $(H - c)^\Lambda$, where $c$ is a positive number large enough to ensure that $|E_0 - c| > |E_{M-1} - c|$. Here we will in the following assume that such a constant, if required, has already been absorbed into $H$. 

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While Eq. (170) is guaranteed to produce the ground state when \( \Lambda \) is sufficiently large, a more efficient way to construct a state which approaches the ground state as \( \Lambda \to \infty \) is to consider not only \( H^\Lambda |\Psi\rangle \), but the whole subspace of the Hilbert space spanned by the set of states \( H^m |\Psi\rangle, m = 0, \ldots, \Lambda \). These states can be constructed one-by-one by successive operations with \( H \) on the initial state \( |\Psi\rangle \). In this subspace, an optimal linear combination of vectors approximating the ground state (minimizing the energy expectation value) exists, and the way to find it is to diagonalize \( H \) in the generated subspace of \( \Lambda + 1 \) vectors. In addition to projecting out the ground state for relatively small \( \Lambda \) (often many orders of magnitude smaller than the size \( M \) of the full Hilbert space), this approach can also accurately reproduce a number of low-lying excited states.

The subspace of the Hilbert space obtained by acting multiple times with \( H \) on an initial state is called the Krylov space. We are here discussing hamiltonians in quantum mechanics, but Krylov space methods of course apply to eigenvalue problems more generally as well, and are very widely used in many areas of science and engineering.

### 4.2.2. The Lanczos basis and hamiltonian

In the Lanczos method [148], an orthogonal basis is constructed using linear combinations of the Krylov space states such that the hamiltonian written in this basis is tridiagonal. In a standard approach, which will be described below, a basis \( \{ |f_m\rangle \}, m = 0, \ldots, \Lambda \), is first constructed that is orthogonal but not normalized. These states will subsequently be normalized to yield an orthonormal set \( \{ |\phi_m\rangle \} \), in which the hamiltonian takes a particularly simple tridiagonal form. We will also discuss a slightly different approach of directly generating the normalized basis \( \{ |\phi_m\rangle \} \).

**Generation of the Lanczos basis.** The construction of the basis \( \{ |f_m\rangle \} \) starts from an arbitrary normalized state \( |f_0\rangle \), of which it is required only that it is not orthogonal to the ground state of \( H \) (if the goal is to find the ground state). This should be the case for a randomly generated \( |f_0\rangle \), but one can also start from some vector which is known to have a substantial overlap with the ground state. The next state is given by

\[
|f_1\rangle = H|f_0\rangle - a_0|f_0\rangle,
\]

where the constant \( a_0 \) should be determined such that \( |f_1\rangle \) is orthogonal to \( |f_0\rangle \). To this end, we examine the overlap between the two states;

\[
\langle f_1 | f_0 \rangle = \langle f_0 | H | f_0 \rangle - a_0 \langle f_0 | f_0 \rangle = H_{00} - a_0 N_0,
\]

where we have introduced notation for the normalization constants and diagonal matrix elements of the hamiltonian that will also be used for subsequent states;

\[
N_m = \langle f_m | f_m \rangle, \quad H_{mm} = \langle f_m | H | f_m \rangle.
\]

For the overlap (172) to vanish we must choose \( a_0 = H_{00} / N_0 \). The following, \( m = 2 \) state is written in terms of the two preceding ones as

\[
|f_2\rangle = H|f_1\rangle - a_1|f_1\rangle - b_0|f_0\rangle,
\]

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where $a_1$ and $b_0$ can be chosen such that $|f_2\rangle$ is orthogonal to both $|f_0\rangle$ and $|f_1\rangle$. The overlaps are, using $H|f_0\rangle$ and $H|f_1\rangle$ obtained from Eqs. (171) and (174):

$$
\langle f_2|f_1\rangle = H_{11} - a_1 N_1, \quad \langle f_2|f_0\rangle = N_1 - b_0 N_0.
$$  \hfill (175)

The appropriate coefficients are thus $a_1 = H_{11}/N_1$ and $b_0 = N_1/N_0$. For all subsequent iterations, Eq. (174) generalizes to

$$
|f_{m+1}\rangle = H|f_m\rangle - a_m|f_m\rangle - b_{m-1}|f_{m-1}\rangle,
$$  \hfill (176)

and the coefficients rendering this state orthogonal to $|f_m\rangle$ and $|f_{m-1}\rangle$ are generalizations of the expressions we already found for $a_0$, $a_1$, and $b_0$:

$$
a_m = H_{mm}/N_m, \quad b_{m-1} = N_m/N_{m-1}.
$$  \hfill (177)

This can easily be checked by direct computation of the overlaps. It remains to be shown that with these coefficients the state $|f_{m+1}\rangle$ is orthogonal also to all previous states $|f_k\rangle$ with $k < m - 1$. In an inductive proof, we can use the fact that all previously generated states are orthogonal to each other, and obtain,

$$
\langle f_{m+1}|f_{m-k}\rangle = \langle f_m|H|f_{m-k}\rangle - a_m \langle f_m|f_{m-k}\rangle - b_{m-1} \langle f_{m-1}|f_{m-k}\rangle
$$  \hfill (178)

$$
= \langle f_m|f_{m-k+1}\rangle + a_{m-k} \langle f_{m-k}|f_{m-k}\rangle + b_{m-k-1} \langle f_{m-k-1}|f_{m-k}\rangle = 0,
$$

where we also used $H|f_{m-k}\rangle$ in the form given by Eq. (176).

This iterative procedure (176) is continued until $m = \Lambda$, where $\Lambda$ can be determined automatically, on the fly, according to some convergence criterion for computed quantities, as we will discuss below. First we need the hamiltonian matrix elements.

**The hamiltonian in the Lanczos basis.** Having constructed a set of $\Lambda + 1$ Lanczos vectors $|f_m\rangle$, $m = 0, \ldots, \Lambda$, the hamiltonian in this basis can be constructed. We can use the expression for $H$ acting on one of the basis states, obtained from Eq. (176);

$$
H|f_m\rangle = |f_{m+1}\rangle + a_m|f_m\rangle + b_{m-1}|f_{m-1}\rangle,
$$  \hfill (179)

and thus the non-zero matrix elements are

$$
\langle f_{m-1}|H|f_m\rangle = b_{m-1} N_{m-1} = N_m,
$$

$$
\langle f_m|H|f_m\rangle = a_m N_m,
$$

$$
\langle f_{m+1}|H|f_m\rangle = N_{m+1}.
$$  \hfill (180)

The normalized basis states are

$$
|\phi_m\rangle = \frac{1}{\sqrt{N_m}} |f_m\rangle,
$$  \hfill (181)

and $b_m = N_{m+1}/N_m$ according to (177). The non-zero matrix elements are therefore

$$
\langle \phi_{m-1}|H|\phi_m\rangle = \sqrt{b_{m-1}},
$$

$$
\langle \phi_m|H|\phi_m\rangle = a_m,
$$

$$
\langle \phi_{m+1}|H|\phi_m\rangle = \sqrt{b_m}.
$$  \hfill (182)
This is a tridiagonal matrix, which can be diagonalized using special methods that are faster than generic diagonalization algorithms (available in many linear algebra subroutine libraries). The main advantage is not, however, that the matrix can be diagonalized more easily—it is anyway typically not very large and even if a generic diagonalization routine is used the time spent on that part of the calculation is negligible. The advantage is that this matrix can be constructed relatively quickly, especially when the matrix $H$ is sparse and only its non-zero elements have to be considered (stored or generated on the fly when acting with $H$ on a state). Some times, when pushing the method to very large matrices, it is also useful that only three of the large state vectors $|f_m\rangle$ have to be stored in the process. The tridiagonality is also very convenient when calculating spectral functions (dynamic correlations), as discussed, e.g., in Refs. [43, 143].

Note that with the basis states labeled $m = 0, \ldots, \Lambda$, the size of the tridiagonal matrix (the actual basis size) is $\Lambda$, not $\Lambda+1$, because for $m+1 = \Lambda$ in (179) the last diagonal matrix element generated for use in (183) is $a_{\Lambda-1}$. The state $|f_\Lambda\rangle$ does not have to be constructed; only the term $H|f_{\Lambda-1}\rangle$ is needed to calculate the last coefficient $a_{\Lambda-1}$.

Since the cost of diagonalizing the Lanczos hamiltonian matrix is very low, one can do that after each new basis state has been generated, and follow the evolution of the eigenvalues. One can stop the procedure based on some suitable convergence criterion, e.g., the desired eigenvalues changing by less than some tolerance $\varepsilon$ between iterations. The basis size required for convergence of course depends on the system studied, but typically, for quantum spin systems, $\Lambda$ in the range tens to hundreds should suffice.

Alternative formulation with normalized vectors. The generation of the Lanczos basis is normally discussed in terms of the un-normalized states $|f_m\rangle$, as we have done above. However, a direct computer implementation of this procedure occasionally leads to numerical problems, because the normalization constants $N_m$ can become exceedingly large (if the eigenvalues of $H$ are large). It may then be better to work directly with the normalized states $|\phi_m\rangle$ (as an alternative to multiplying $H$ by a suitable factor). In fact, in practice this formulation is even simpler, and the additional computational cost is merely the normalizations at each step (which is normally small compared to other costs).

We start with a normalized state $|\phi_0\rangle$ and generate the second state according to

$$|\phi_1\rangle = \frac{1}{N_1}(H|\phi_0\rangle - a_0|\phi_0\rangle).$$

(183)

Here $N_1$ is a normalization constant, which is determined by direct computation of the scalar product of the constructed state $H|\phi_0\rangle - a_0|\phi_0\rangle$ with itself. Orthogonality with $|\phi_1\rangle$ again requires $a_0 = H_{00}$. For each following state one can easily show that

$$|\phi_{m+1}\rangle = \frac{1}{\sqrt{N_{m+1}}}(H|\phi_m\rangle - a_m|\phi_m\rangle - N_m|\phi_{m-1}\rangle) = \frac{|\gamma_{m+1}\rangle}{N_{m+1}}$$

(184)

is orthogonal to all previous states. Here the definitions of $a_m$ and $N_m$ differ from the previous ones in Eq. (177);

$$a_m = \langle \phi_m | H | \phi_m \rangle,$$

$$N_m = \langle \gamma_m | \gamma_m \rangle.$$
where $|\gamma_m\rangle$ is the generated state before normalization as in Eq. (184) above. The Hamiltonian matrix elements are

$$
\begin{align*}
\langle \phi_{m-1} | H | \phi_m \rangle &= \sqrt{N_m}, \\
\langle \phi_m | H | \phi_m \rangle &= a_m, \\
\langle \phi_{m+1} | H | \phi_m \rangle &= \sqrt{N_{m+1}}.
\end{align*}
$$

In this formulation all the stored numbers are well behaved.

**Degenerate states.** It should be noted that the Lanczos method cannot produce more than one member of a multiplet; out of a degenerate set of states, only a particular linear combination of them will be obtained (which depends on the initial state $|f_0\rangle$). To see the reason for this, we again look at the expansion (170) of a state $H^\Lambda |\Psi\rangle$, in which we assume that there are two degenerate states $|\psi_i\rangle$ and $|\psi_j\rangle$, $E_i = E_j$. In the expansion we can isolate these states from the rest of the terms;

$$
H^\Lambda |\Psi\rangle = E_i^m(c_i |\psi_i\rangle + c_j |\psi_j\rangle) + \sum_{m \neq i,j} c_m E_m^\Lambda |\psi_m\rangle.
$$

For any $\Lambda$, the expansion contains the same linear combination of the states $|\psi_j\rangle$ and $|\psi_j\rangle$. Hence, in the subspace spanned by the set of states $H^m |\Psi\rangle$, $m = 0, \ldots, \Lambda - 1$, there is no freedom for obtaining different linear combinations of the two degenerate states. This of course generalizes also to degenerate multiplets with more than two states.

**Loss and restoration of orthogonality.** When the basis size $\Lambda$ becomes large, the Lanczos procedure typically suffers from numerical instabilities. Round-off errors accumulated in the course of constructing the basis set will eventually introduce some non-orthogonality among the states. Such numerical errors can escalate and lead to successive sudden appearances (within some narrow ranges of $\Lambda$) of several identical eigenvalues (recall that the Lanczos scheme should never produce degenerate states). We will see an example of this further below.

Loss of orthogonality and the appearance of multiple copies of the same states is normally not a problem when the aim is to obtain only the ground state. It can complicate calculations of excited states, however. To remedy this, one can add to the basis construction a step where each new Lanczos vector constructed is explicitly orthogonalized with respect to all previous basis vectors. In the simplest implementation of such a stabilization procedure, all the Lanczos vectors are stored (in primary memory or secondary storage). This can become problematic when dealing with very large basis sets, but the scheme is very simple. Working with the normalized states, each $|\phi_{m+1}\rangle$ constructed according to (184) is orthogonalized with respect to all previous states, according to

$$
|\phi_{m+1}\rangle \rightarrow \frac{|\phi_{m+1}\rangle - q |\phi_i\rangle}{1 - q^2}, \quad q = \langle \phi_i | \phi_{m+1} \rangle,
$$

successively for $i = 0, \ldots, m$. This makes it possible to study a much larger number of excited states (in principle only limited by computer memory).
Instead of converging several excited states in the same run, one can also target excited states one-by-one, starting each time from a vector which is orthogonal to all previous ones. Re-orthogonalization should then also be done with respect to those.

_**Eigenstates and expectation values.**_ Diagonalizing the tridiagonal Lanczos hamiltonian results in eigenvalues $E_n$ and eigenvectors $v_n$, $n = 0, \ldots, \Lambda - 1$. We want these eigenvectors expressed in the original basis $\{|a\rangle\}$, in which we are able to evaluate the matrix elements $\langle b|O|a\rangle$ of operators $O$ of interest. First, the Lanczos basis states are

$$|\phi_m\rangle = \sum_{a=1}^{M} \phi_m(a)|a\rangle, \quad m = 0, \ldots, \Lambda - 1,$$

and we denote the desired eigenvectors of the hamiltonian as

$$|\psi_n\rangle = \sum_{a=1}^{M} \psi_n(a)|a\rangle, \quad n = 0, \ldots, \Lambda - 1. \quad (190)$$

The first few eigenvectors $v_n$ of the tridiagonal matrix accurately represent eigenstates of the hamiltonian (essentially exactly for sufficiently large $\Lambda$) in the Lanczos basis;

$$|\psi_n\rangle = \sum_{m=0}^{\Lambda - 1} v_n(m)|\phi_m\rangle = \sum_{m=0}^{\Lambda - 1} \sum_{a=1}^{M} v_n(m)\phi_m(a)|a\rangle, \quad (191)$$

and thus the wave function coefficients we want to construct are given by

$$\psi_n(a) = \sum_{m=0}^{\Lambda - 1} v_n(m)\phi_m(a), \quad a = 1, \ldots, M. \quad (192)$$

If all the Lanczos vectors have been stored during the basis construction, this transformation can be carried out in a straight-forward manner. If there is enough computer memory available, one should store all Lanczos vectors, but often calculations are pushing the limits of computer capacity, and then only the bare minimum of information can be stored, at the cost of longer calculation times. If we did not store the full set of Lanczos vectors during the basis construction, we need to generate them again, in the same iterative fashion as before, except that we already have the coefficients $a_m$ and $N_m$ available (and $b_m$ if the scheme with un-normalized Lanczos vectors $|f_m\rangle$ is used) and do not need to recompute them. We can transform the states according to (192) on the run with the eigenvectors $v_m$, building up one or several of the eigenstates of the hamiltonian.

Having generated one or several eigenstates, now assumed to be stored in the form of the coefficients $\psi_m(a)$ in (190), an expectation value of some operator $O$ can be obtained by first acting on the state, giving an un-normalized state that we call $|\psi_n^O\rangle$,

$$O|\psi_n\rangle = |\psi_n^O\rangle = \sum_{a=1}^{M} \psi_n(a)O|a\rangle$$

$$= \sum_{a=1}^{M} \sum_{b=1}^{M} \psi_n(a)|b\rangle\langle b|O|a\rangle. \quad (193)$$
\[
\psi_n(a) = \sum_{b=1}^{M} \psi_n(b) \langle b | O | a \rangle.
\]

We then evaluate the scalar product of this state with the original state \( | \psi_n \rangle \):

\[
\langle \psi_n | O | \psi_n \rangle = \langle \psi_n | \psi_n^O \rangle = \sum_{a=1}^{M} \psi_n(a) \psi_n^O(a).
\]  

(194)

This somewhat cumbersome way of writing the expectation value corresponds to a typical computational procedure of first computing the state vector \( \psi_n^O \) and then computing its scalar product with \( \psi_n \).

It is interesting to note that, in Lanczos calculations, we are often dealing with four different bases: First, we have the original “computational” basis of single states of \( \uparrow \) and \( \downarrow \) spins. From these we generate a basis incorporating symmetries, e.g., momentum states (which normally would be the states denoted as \( | a \rangle \) above). We then construct the Lanczos vectors, which are particular linear combinations of those states. Finally, diagonalizing the tridiagonal matrix (which is an effective hamiltonian in the low-energy sector), we obtain the desired energy eigenstates. To do calculations with those states, we effectively do the basis transformations in reverse. The matrix elements in (193) and (194) are in the end carried out in the computational basis of \( \uparrow \) and \( \downarrow \) spins (i.e., with the representative states used, e.g., to build the momentum states). We next discuss all these procedures in practical program implementations.

### 4.2.3. Programming the Lanczos method

While the Lanczos method can be applied to any symmetric (or hermitian) matrix, in the case of a spin hamiltonian there is an added advantage in that the hamiltonian is a sparse matrix. Although the size of the hamiltonian (an individual symmetry-block) can be very large, the number of non-zero matrix elements is much smaller. For a model with short-range interactions on a lattice of \( N \) sites, a hamiltonian block of \( M \) states has on the order of \( NM \) non-zero elements, which for large \( M \) is much smaller than the total number of elements \( M^2 \). Since the most time consuming part of the construction of the Lanczos basis is the repeated operations with the hamiltonian matrix on a state vector, to generate the next basis state according to Eq. (176) or (184), the sparseness allows for enormous time savings. There are similarly significant memory savings advantages as well. The non-zero elements of the hamiltonian should then be stored in a compact form, or generated on the fly as needed (which, when many symmetries are used, typically takes longer than even reading the elements from disk storage). We here implement the Lanczos method with the hamiltonian stored in a compact form in primary memory.

We will generate the Lanczos basis of states \( \{ | \phi_m \rangle \} \) that are normalized at each step, using Eqs. (184) and (183). In a computer program, these states are stored in the form of their vector components \( \phi_m(a) \), in terms of which the Lanczos states are given by Eq. (189). Here \( a = 1, \ldots, M \) labels the states of our working basis, which in the case of maximal use of symmetries would be momentum or semi-momentum states, e.g., \( | a \rangle = | k, p, z, m_z = 0, a \rangle \), but for simplicity we do not write out all the quantum numbers.
The nature of the basis states only come into play when acting with the hamiltonian (or some operator to be measured) on the basis states, and for this we need exactly the matrix elements that have already been discussed for the various symmetry implementations in Secs. 4.1.3, 4.1.4, and 4.1.5. We will write pseudocodes assuming real-valued wavefunction coefficients; the changes needed for complex states are self-evident.

**Lanczos basis construction and eigenstates.** To discuss the general structure of a Lanczos program, we begin by assuming that we have implemented a subroutine \( \text{hoperation}(\phi, \gamma) \) which acts with the hamiltonian on a state vector; \( |\gamma\rangle = H|\phi\rangle \). Later, we will describe the implementation of this subroutine, which is the only part of the Lanczos procedure that requires a specification of the model and the symmetries implemented. We also use a subroutine \( \text{normalize}(\phi, n) \), which first computes \( n = \langle \phi | \phi \rangle \) and then rescales \( \phi \) so that \( \langle \phi | \phi \rangle = 1 \) upon return.

To start the basis generation, we first load the initial state \( |\phi_0\rangle \) with a randomly generated normalized state. We denote the elements \( \phi_0(i), i = 1, \ldots, M \) and \( \langle \phi_0 | \phi_0 \rangle = \sum_i \phi_0(i)^2 = 1 \). The next Lanczos state is generated according to (183);

\[
\text{call hoperation}(\phi_0, \phi_1) \quad a_0 = \langle \phi_0 | \phi_1 \rangle; \quad \phi_1 = \phi_0 - a_0\phi_1 \\
\text{call normalize}(\phi_1, n_1)
\]

If we are not storing all the Lanczos vectors, we can cycle between three vectors, \( \phi_0, \phi_1, \phi_2 \), that contain the information we need at each step when implementing the iterative basis generation according to (184);

\[
\text{do } m = 1, \Lambda - 1 \\
\quad \text{call hoperation}(\phi_1, \phi_2) \quad a_m = \langle \phi_1 | \phi_2 \rangle; \quad \phi_2 = \phi_2 - a_m\phi_1 - n_m\phi_0 \\
\quad \text{call normalize}(\phi_2, n_{m+1}) \\
\quad \phi_0 = \phi_1; \quad \phi_1 = \phi_2 \\
\text{enddo}
\]

If possible, we should store all the states, to avoid having to regenerate them at the later stage when computing expectation values (which doubles the computation time). We have to store all the states if we want to carry out additional re-orthogonalization, to ensure that numerical truncation errors do not eventually degrade the Lanczos basis. In that case we insert an additional loop to orthogonalize with respect to all previously generated states, according to Eq. (188);

\[
\text{do } m = 1, \Lambda - 1 \\
\quad \text{call hoperation}(\phi_m, \phi_{m+1}) \quad a_m = \langle \phi_m | \phi_{m+1} \rangle; \quad \phi_{m+1} = \phi_{m+1} - a_m\phi_m - n_m\phi_{m-1} \\
\quad \text{call normalize}(\phi_{m+1}, n_{m+1}) \\
\quad \text{do } i = 0, m \\
\quad \quad q = \langle \phi_{m+1} | \phi_i \rangle; \quad \phi_{m+1} = (\phi_{m+1} - q\phi_i) / (1 - q^2) \\
\quad \text{enddo} \\
\text{enddo}
\]
After completing either code segment \{18\} or \{19\}, we have the contents of the tridiagonal matrix in Eq. (187) and can proceed to feed this information into a diagonalization routine which delivers the eigenvalues $E_n$ and eigenvectors with elements $v_n(i)$, $n = 0, \ldots, \Lambda - 1$, $i = 1, \ldots, M$. It should be noted again that, with the conventions we have been using, the Lanczos basis size (the size of the matrix) is $\Lambda$; the last state generated in \{18\} and \{19\}, with $m + 1 = \Lambda$, is actually not needed (only the coefficient $a_{\Lambda-1}$ generated in the last iteration is needed).

The computational effort of the diagonalization is very small compared to the time spent on the basis construction. We may therefore as well diagonalize after each new Lanczos vector has been generated. We can then monitor how the energies evolve with the basis size. One can then stop when some convergence criterion is satisfied. One can, e.g., demand that the change in the ground state energy (or the highest excitation of interest) changes between steps $m - 1$ and $m$ by less than some small number $\epsilon$. The Lanczos method is normally capable of converging energies to the numerical precision of the computer (and double precision should always be used). As we will see below, other properties converge slower than the energies.

**Expectation values.** First, let us consider calculations of operator expectation values (besides the energy) in a program where we have not stored all the Lanczos basis vectors. Then, in order to be able to transform the ground state from the Lanczos basis to the original basis, where we can carry out “measurements” on the state, we have to repeat the Lanczos basis construction starting from the same initial state as in the first run. If we had initialized with a random $\phi_0$, as in code \{17\}, and did not save this state, we can re-generate it by initializing the random number generator with the same seeds as in the first construction. In any case, with $\phi_0$ at hand, we can proceed as in code segment \{18\}, with the minor simplification that we now do not have to calculate the coefficients $a_m, nm$, because we already have them. We can use $nm$ for normalizing, instead of calling the subroutine normalize in \{18\}. During the basis re-construction, we transform the states with the ground state vector $v_0$ (or $v_n$ for an excited state $i$) that resulted from the diagonalization of the tridiagonal matrix, building up the ground state in a vector $\psi$:

\begin{verbatim}
call hoperation(\phi_0, \phi_1); \phi_1 = (\phi_1 - a_0 \phi_0) / \sqrt{n_1}
\psi = v_0(0) \phi_0 + v_0(1) \phi_1
\text{do } m = 2, \Lambda - 1
\text{call hoperation}(\phi_1, \phi_2)
\\phi_2 = (\phi_2 - a_{m-1} \phi_1 - n_{m-1} \phi_0) / \sqrt{n_m}
\psi = \psi + v_0(m) \phi_2
\phi_0 = \phi_1; \phi_1 = \phi_2
\text{endo}
\end{verbatim}

The elements of the vector $\psi$ now contain the ground-state wave-function coefficients in whatever basis is used and implemented through hoperation. If we have stored all the Lanczos vectors, we of course do not need this step and can directly transform the states generated in code \{19\}.

It is particularly easy to calculate expectation values of operators that depend only on the $z$-components of the spins and is invariant under all the symmetry operations.
used in the basis. We then just have to weight the quantity calculated using the representative states with the corresponding wave-function coefficient squared (the state’s probability). As an example, the spin-spin correlation function \( \langle S^+_i S^+_i \rangle \) for all distances \( r = 0, \ldots, N/2 \) can be obtained as:

\[
\text{do } a = 1, M \\
\text{do } r = 0, N/2 \\
\text{do } i = 0, N - 1 \\
\quad j = \text{mod}(i + r, N) \\
\quad \text{if } (s_a[i] = s_a[j]) \text{ then} \\
\quad \quad C(r) = C(r) + \psi^2(a) \\
\quad \text{else} \\
\quad \quad C(r) = C(r) - \psi^2(a) \\
\text{endif} \\
\text{enddo} \\
\text{enddo} \\
\text{enddo} \\
C = C/4N
\]

Here \( \psi^2(a) \) is the probability of the state \( |a\rangle \), which when symmetries are incorporated is represented by a number \( s_a \). As before, bit tests can be used to determine the relative spin orientation, now of spins separated by \( r \) lattice spacings. Note that although the final spin correlation \( \langle S^+_i S^+_i \rangle \) only depends on \( r = |i - j| \), we still have to average over all \( i \) above, because we are only using the representative states (which do not by themselves, without acting with the symmetry operators, obey any lattice symmetries). In principle we should also average over both reflections and spin-inversions of the representative, but after the translational averaging has been done the spin correlation function is also invariant with respect to the other two symmetries (which is not the case for all operators). Note that since the Heisenberg Hamiltonian is spin-rotation invariant, the correlation function \( \langle S^+_i S^+_i \rangle \) calculated here equals \( \langle S_i \cdot S_{i+r} \rangle / 3 \).

Calculating expectation values of \( z \) off-diagonal operators that cannot be simply related to \( z \) diagonal ones require explicit operations on the states, and thereafter evaluation of a scalar product, according to Eqs. (193) and (193). This is in principle easy—essentially proceeding as in the construction of the Hamiltonian in the preceding sections—but more time consuming than diagonal operators.

**Compact storage of the hamiltonian.** We now discuss the inner workings of the subroutine \texttt{hoperation}(\( \phi, \gamma \)) that we employed in the Lanczos procedures. It implements the operation

\[
H|\phi\rangle = |\gamma\rangle = \sum_{a=1}^{M} \sum_{b=1}^{M} \phi(a)\langle b|H|a\rangle|b\rangle.
\]

This is of course where all the details of the symmetries employed will enter. The main difference with respect to the construction of the Hamiltonian in complete diagonalization is that we do not want to store \( H \) as a full \( M \times M \) matrix, because we have in mind calculations were the number of basis states \( M \) can be up to many millions (e.g., for...
N = 32 in Table 1). We therefore have to devise a convenient way of only storing its non-zero elements, of which there are of the order $NM$ in (195).

We define the following data structures for the compact hamiltonian storage: For each $a = 1, \ldots, M$ in (195) we store the number $e_a$ of non-zero matrix elements $\langle b | H | a \rangle$. We store the locations $b$ of these non-zero elements as consecutive integers in a vector with elements $B(i)$. In a program, it is convenient to use the elements $i = 1, \ldots, e_1$ for $a = 1$, followed by $i = e_1 + 1, \ldots, e_1 + e_2$ for $a = 2$, etc. The required size of the vector $B$ (the number of non-zero elements) is initially not known, and, depending on the programming language used, may have to be allocated using an estimated size. We can store the corresponding non-zero matrix elements in a vector with floating-point values $H(i)$. We can also take advantage of the fact that the hamiltonian is a symmetric matrix and only store one of its “triangles”.

With the above notation, carrying out the operation (195) is now as simple as:

```
subroutine hoperation(\phi, \gamma)
  \gamma = 0; i = 0
  do a = 1, M
    do j = 1, e_a
      i = i + 1
      \gamma(B(i)) = \gamma(B(i)) + H(i)\phi(a)
      \gamma(a) = \gamma(a) + H(i)\phi(B(i))
    enddo
  enddo
enddo
```

The counter $i$ keeps track of the position of the elements in the data structures $H$ and $B$. Diagonal matrix elements, $B(i) = a$, are double counted in this procedure since the contributions from the upper and lower triangle of the hamiltonian matrix are added. It is therefore assumed that their stored values have been divided by 2.

The matrix elements $H(i)$ and their locations $B(i)$ are generated in a way similar to what we did when constructing the complete hamiltonian, e.g., as in code segments \{13\}–\{16\} in the semi-momentum basis with parity. We also discussed the simple extensions involving spin-inversion symmetry. Now we discuss the modifications needed when we wish to load only the non-zero elements into our compact storage.

In the case of the semi-momentum basis, which we will consider here, the procedures are again somewhat complicated by the fact that the same representative can appear once or twice ($\sigma = \pm 1$) in the state list, and we want to take care of these at the same time in order to avoid repeating tasks unnecessarily. We therefore carry out the loop over the basis states $a$ and determine the number of same representatives $n$ as in code \{13\}. We cannot put the matrix elements directly into the storage vector $H$, because our scheme in code \{22\} requires consecutive storage of all matrix elements for each column. With the way the individual bond operators in the hamiltonian are treated one-by-one in code \{16\}, when $n = 2$ the two columns would be mixed up if we store each matrix element as it is generated. We therefore use temporary storage for the one or two columns being currently processed, and then later copy their contents into the appropriate positions of the final storage. Referring to the first ($a$) and potential second ($a + 1$) column as $c = 1$ and $c = 2$, we store the data temporarily as $B_c(k)$ and $H_c(k)$, $k = 1, \ldots, n_c$, where $n_c$ is...
the number of elements for the column in question. The diagonal elements (divided by 2, as discussed above) are entered first into these temporary storage lists;

\[
\begin{align*}
    n_1 &= 0; \ n_2 = 0 \\
    \text{do } c &= 1, n \\
    B_c(1) &= a + c - 1; \ H_c(1) = E_z / 2; \ n_c = 1
\end{align*}
\]

\textbf{enddo}

For the off-diagonal matrix elements we proceed as in code segment \{16\}. Note, however, that several bond operations may lead to the same state \(|b\rangle\) when acting on a given basis state \(|a\rangle\), but in the compact storage we do not want to store individual contributions to the same matrix element separately. Everything is taken care of by replacing the loops over \(i\) and \(j\) in code \{16\} by this extended version;

\[
\begin{align*}
    \text{do } i &= a, a + n \\
    \text{do } j &= \max(b, i), b + m \\
    c &= i - a + 1 \\
    E &= \text{helement}(i, j, l, q) \\
    \text{do } k &= 1, n_c \\
    &\quad \text{if } B_c(k) = j \text{ exit} \\
    \text{enddo} \\
    &\quad \text{if } (k > n_c) \text{ then } n_c = k; \ B_c(k) = j \text{ endif} \\
    &\quad \text{if } (j = i) \text{ then } H_c(k) = H_c(k) + E / 2 \text{ else } H_c(k) = H_c(k) + E \text{ endif} \\
    \text{enddo} \\
\end{align*}
\]

\textbf{enddo}

Here \(\max(i, b)\) on the second line ensures that only the elements in matrix triangle \(j \geq i\) are constructed. The innermost loop over \(k\) checks whether there is already a stored contribution to the matrix element \(|j\rangle |H|\langle i|\). It has been assumed that when the loop over \(k\) has been completed (without exiting before \(k = n_c\)), then \(k\) takes the value \(n_c + 1\) (which is the case in many computer languages). Then \(k = n_c + 1\) if there is no prior location \(j\) in the list, which means that it should be added to the list (and the size of the list is then \(k\)). We have also assumed that all \(H_c(k)\) are initially set to zero, so that each new contribution \(E\) can be added in the appropriate location of \(H_c\). Note that although we are carrying out off-diagonal operations here, in the basis we are using, such operations can also lead to diagonal matrix elements, in which case we have to divide \(E\) by 2.

After having generated all the matrix elements originating from the current representative (i.e., completed the loop over all nearest-neighbor spin pairs), we copy the contents of the temporary storage vectors into the permanent full storage;

\[
\begin{align*}
    \text{do } c &= 1, 2 \\
    \text{do } i &= 1, n_c \\
    n_H &= n_H + 1; \ B(n_H) = B_c(i); \ H(n_H) = H_c(i)
\end{align*}
\]

\textbf{enddo}

\textbf{enddo}

Here \(n_H\) is a counter for the total number of non-zero matrix elements added so far. After this, the loop over state labels \(a\) is closed, as in code \{13\}. This completes the
FIGURE 30. Example of the convergence as a function of the Lanczos basis size of the energy (left) and the total spin (right) of the four lowest levels of the 24-site Heisenberg chain in the symmetry sector \( k = 0, p = 1, z = 1 \). The spin \( S \) is extracted using the assumption that \( \langle S^2 \rangle = S(S + 1) \).

4.2.4. Convergence of Lanczos calculations

The Lanczos method is essentially exact if a sufficiently large number of Lanczos vectors \( \Lambda \) is used, and typically this number does not even have to be very large; on the order of a few tens to hundreds. The convergence should be checked by carrying out calculations for several \( \Lambda \), until no changes can be detected in the energies and expectation values of interest. The ground state converges the fastest, and energies converge faster than expectation values. As an example, Fig. 30 shows results for an \( N = 24 \) chain. The energy and the total spin of the four lowest levels in the symmetry sector of the ground state are shown versus \( \Lambda \). The spin quantum number is calculated by acting with the squared total spin operator \( S^2 \) on the states, according to Eqs. (193) and (194), and extracting \( S \) assuming \( S^2 = S(S + 1) \) (which is valid only when the
states have converged to eigenstates of the operator. The energies are seen to converge monotonically, whereas this is not necessarily the case for other quantities, as seen clearly for the $S = 2$ states in this case. The details of the convergence of course depend on the initial state from which the Lanczos basis is constructed (which in this case was a random state). In this case all the four levels shown ($E$ as well as $S$) were converged to better than 10 decimal places at $\Lambda = 60$, with the ground state having converged at that level already at $\Lambda = 30$. Going to larger system sizes, the convergence becomes a little slower, but for this particular model there are no difficulties in converging several levels up to the largest system sizes that can feasibly be studied.

One can accelerate the convergence of a Lanczos calculation by starting from a state which is already close to the ground state. Such states may be constructed in a number of ways, e.g., based on some approximate analytical method. But if there are no convergence problems this may not be worth the additional effort. However, if a series of calculations are carried out as a function of some parameter in the hamiltonian, then subsequent calculations can be started from the ground state of the preceding parameter value, which is likely to have a significant overlap with the next ground state. However, it should be noted that if the initial state is a good approximation to the ground state, it will have very small overlaps with the first few excited states, and hence only the ground state is likely to converge rapidly in such a calculation. If excited states are also needed, this problem can be circumvented by starting the next calculations using a linear combination of eigenstates from prior calculations.

**Loss and recovery of orthogonality.** The Lanczos basis vectors should all be completely orthogonal to each other, but numerical truncation errors build up and eventually lead to escalating loss of orthogonality for some $\Lambda$. This manifests itself as artificial degeneracies, with excited states “falling down” onto lower states. An example of this is shown in Fig. 31(a), where the four lowest energies of a 16-site chain in the ground-state symmetry sector are graphed versus $\Lambda$. The higher energies are seen to successively collapse onto the immediately lower energies, with only a few iterations taken for the levels...
to become completely degenerate after signs first appear of the imminent collapse. The basis then includes multiple copies of the same states.

The loss of orthogonality may at first sight seem like a serious problem, but in practice this is no necessarily the case. Calculations for the ground state are not much affected by there being more than one copy of it, as long as one makes sure that it is properly normalized before using it to calculate expectation values, and the first excited states are normally (but not always) well converged before they fall down onto lower states.

Loss of orthogonality often occurs sooner for small systems than large ones. This is related to the fact that there are less low-energy states for small systems, whence the lowest states in the Lanczos basis can become “overcomplete” in this subspace (if the Lanczos basis is larger than the total number of states, it clearly is truly overcomplete and the calculation will not work).

One can easily supplement the Lanczos method by an explicit re-orthogonalization step, as in Eq. (188) and implemented in code {19}. The drawback is that this requires storage of all the Lanczos basis vectors, which may not be possible for large systems. The procedure may also become time consuming if the number of states is large. As shown in Fig. 31(b), re-orthogonalization enables convergence of many more eigenstates (limited only by memory and time constraints).

4.3. 1D states and quantum phase transitions

In this section we discuss several related 1D calculations using the Lanczos method and finite-size scaling methods for up to \(N = 32\) spins. First we investigate the critical ground state of the standard Heisenberg chain, and also analyze some properties of excited states. Then we introduce frustration, studying the dimerization transition in the \(J_1-J_2\) chain. Finally, we add long-range interactions in addition to frustration, in which case the continuous dimerization transition evolves into a first-order transition between a Néel state (which is possible even in a 1D system if the interactions are sufficiently long-ranged) and a dimerized VBS state.

4.3.1. Ground state and excitations of the Heisenberg chain

Although the Heisenberg chain has an exact Bethe ansatz solution [51], the wave function is very complicated and in many cases numerical calculations for finite-size systems have to be used to extract information on physical properties from it [53, 54]. The energy of the ground state [52] and the low-lying excitations [149] can be calculated exactly both for finite chains and in the thermodynamic limit, however. Some other quantities can also be extracted for very large chains [150]. In addition to using the exact solution, many properties of this class of system (i.e., with a wider range of interactions maintaining the symmetries of the system) are known based on an asymptotically exact low-energy field-theory description; the Weiss-Zumino-Witten non-linear \(\sigma\) model with topological coupling (or central charge) \(c = 1\) [56]. Equivalently, the Heisenberg chain also represents a special case (because of its spin-rotational invariance) of the Luttinger-
liquid state, which describes a broad range of interacting 1D spin, fermion, and boson systems [151, 152]. Numerical studies based on exact diagonalization and other unbiased finite-lattice techniques (such as QMC and DMRG) have played an important role in guiding and confirming these theories (see, e.g., [153] and [7]). Computational studies are also required for extracting non-universal (short-distance, higher-energy) properties that are not captured by universal continuum field theories (including quantities directly accessible to experiments, as discussed in, e.g, Refs. [154, 155]). Numerical results for the Heisenberg chain are also very important for enabling rigorous benchmark tests of extrapolation techniques when studying other spin models, for which less is known from analytical calculations.

**Ground state and low-energy excitations.** For a Heisenberg chain of size \( N = 4n \), the ground state has momentum \( k = 0 \), parity \( p = 1 \), and spin-inversion number \( z = 1 \); it is a fully symmetric singlet state. For \( N = 4n + 2 \) the lowest-energy state is in instead in the completely antisymmetric sector; \( k = \pi, p = -1, z = -1, S = 0 \).

Let us look at the most important excitations of the Heisenberg chain. For \( k \) not equal to the ground-state momentum, the lowest-energy state of a finite system is a triplet; hence \( z = -1 \) and \( +1 \) for chains of size \( N = 4n \) and \( 4n + 2 \), respectively. Fig. 32 shows the momentum dependence of the excitation energies \( \omega(k) = E_S(k) - E_0 \) for both the lowest \( z = +1 \) and \( z = -1 \) states of a 32-site chain. The lowest \( z = +1 \) state always has \( S = 1 \), while the lowest \( z = -1 \) state has either \( S = 0 \) or \( S = 2 \). The triplet energies are quite close to the exact \( N = \infty \) triplet dispersion obtained from the Bethe ansatz [149], \( \omega_S(k) = \pi|\sin(k)|/2 \), especially close to \( k = 0 \). Close to \( k = \pi \) the deviations are larger. The \( S = 0, 2 \) excitations are mostly slightly higher in energy, except that the \( S = 2 \) state close to \( k/\pi = 0.85 \) is actually marginally lower than the triplet (likely a small-\( N \) anomaly), and at \( k/\pi = 2/N \) it is much higher.

For large \( N \to \infty \), one would expect the singlet and triplet excitations to become degenerate with dispersion \( \omega_{0,1}(k) = \pi|\sin(k)|/2 \). This can be understood as originating
Singlet and triplet gaps of the Heisenberg chain versus the inverse system size. The inset shows the gaps multiplied by $N$, illustrating the presence of multiplicative logarithmic corrections to the dominant $\sim 1/N$ scaling (dynamic exponent $z = 1$).

from excitations of pairs of spin-1/2 soliton-like degrees of freedom called spinons. These spinons are very weakly interacting and in the infinite chain behave as independent particles, hence forming four degenerate levels (from which $S = 0$ and 1 states can be formed). The dispersion graphed in Fig. 32 is only the lowest edge of a continuum of spinon excitations, which can be calculated in detail using the Bethe ansatz [53] and has also been observed experimentally in quasi-1D antiferromagnets [156].

The very lowest excitation for a $N = 4n$ chain is a triplet at $k = \pi$, with $p = -1, z = -1$, whereas for $4n + 2$ the lowest triplet has $k = 0, p = 1, z = 1$ (i.e., the difference in momentum with respect to the ground state is always $\pi$, and $p, z$ are minus their ground-state values). Fig. 33 shows the finite-size scaling of these lowest triplet and singlet excitation energies (the singlet and triplet finite-size gaps) versus the inverse system size. They are both seen to scale to zero as $1/N$, corresponding to a dynamic exponent $z = 1$. There is a weak correction to this form originating from logarithmic corrections, which can be seen clearly when plotting the gaps multiplied by $N$ (in the inset of Fig. 33). This gap scaling has been predicted in detail based on the continuum field theory approaches [57, 153]. The prefactor of the gap scaling is directly related to the velocity of the spinon excitations, but it is not easy to extract it reliably based on the finite-size data because of the log corrections. The velocity can also be extracted from the linear parts of the dispersion relation close to $k = 0$ and $\pi$, $\omega_S(k)/k$ or $\omega_S(k)/(\pi - k)$, with the $k = 0$ behavior being the easier to analyze, as seen clearly in Fig. 32.

Spin correlation function. The spin correlation function is one of the key characteristics of any quantum spin system. The correlations in the ground state are of primary interest. For a spin-isotropic and translationally invariant chain we can write it as

$$C(r) = \langle S_i \cdot S_{i+r} \rangle = 3\langle S^+_i S^-_{i+r} \rangle,$$

for any reference site $i$. As we discussed in Sec. 2, the Mermin-Wagner theorem rules out antiferromagnetic long-range order in the Heisenberg chain. Instead, in the ground
state $C(r)$ decays as $(-1)^r/r$ for large $r$, up to a multiplicative logarithmic correction (originating from a marginally operator in the field-theoretical description [57, 58, 59]).

Fig. 34 shows ground-state results for $C(r)$ as a function of $r$ for system sizes $N = 16, 24, \text{and } 32$. The results have been multiplied by $r$, so that an $r$-independent behavior should obtain for large $r$ if the asymptotic form is $\sim 1/r$. It is also useful to multiply by $(-1)^r$, in order to cancel out the $\pm$ oscillations of the staggered phases. Note that there are remaining even-odd oscillations in $C(r)(-1)^r$. Such oscillations are quite common for various correlation functions of 1D systems. They diminish with increasing system size but are still strong for the small system studied here. The form of the oscillations is also predicted by the Luttinger liquid theory; they decay as $1/r^2$ [152].

In Fig. 34, clear deviations from the $1/r$ form of $C(r)$ can be seen which are mainly due to the periodic boundary conditions, which enhance the correlations close to $r = N/2$. Open boundaries cause even more severe finite-size effects (in addition to not allowing the use of translational symmetry for block-diagonalization). The logarithmic corrections should play some role as well. As the chain length is increased, the correlation function at fixed $r$ converges to the infinite-$N$ limiting form. However, as can be seen in the figure, the maximum $r$ for which the results are approximately converged is a rather small fraction of $N$. For $N = 32$, one could safely say that $C(r)$ is converged for $N$ only up to $r \approx 4$, which is not enough to say much about the long-distance behavior. An alternative, also shown in the figure, is to investigate the correlation function at fixed $r/N$ as a function of $N$, with $r/N = 1/2$ the most natural choice for checking the long-distance behavior. Even though periodic boundaries enhance the correlations significantly at this point, the functional form of $C(r = N/2)$ is proportional to the infinite-size $C(r)$ [although the overall prefactor of function $C(r = N/2)$ versus $N$ will clearly be different from that in the actual $N \to \infty$ converged $C(r)$]. The leading $1/r$ form appears quite plausible based on the data in Fig. 34, and the remaining enhancement is consistent with the predicted logarithmic corrections. In Sec. 5.3.1 we will further investigate the long-distance correlations based on quantum Monte Carlo calculations for much longer chains.
4.3.2. Frustration-driven quantum phase transition

A very interesting aspect of the $S = 1/2$ Heisenberg chain is that it exhibits several ground-state phase transitions (quantum phase transitions) when a next-nearest-neighbor interaction is added to the hamiltonian:

$$H = J_1 \sum_{i=1}^{N} S_i \cdot S_{i+1} + J_2 \sum_{i=1}^{N} S_i \cdot S_{i+2}. \quad (197)$$

This model goes under the name of the $J_1$-$J_2$ chain, or the Majumdar-Ghosh model (after the authors of the first comprehensive study of the system [60]). Here we will discuss the most important (and most well understood) transition, which occurs when both $J_1$ and $J_2$ are antiferromagnetic (positive). For convenience we define the ratio $g = J_2 / J_1$.

For $g < g_c$, $g_c \approx 0.2411$ [157], the system is in the same phase as the Heisenberg chain with $J_2 = 0$ discussed in the previous section. The ground state is critical with antiferromagnetic correlations decaying as $1/r$ and the finite-size gaps scale as $1/N$ (up to log corrections in both cases). Only the prefactors (e.g., the velocity of the excitations) and the strength of the log corrections change as a function of $g$.

For $g > g_c$ the system is in a completely different kind of state, with exponentially decaying spin correlations and a triplet excitation gap which remains finite in the thermodynamic limit. The ground state has long-range dimer order (a simpler, 1D version of the 2D VBS states discussed in Sec. 2.2). The nearest-neighbor bond strengths (the $J_1$ energy contribution), $\langle B_i \rangle = \langle S_i \cdot S_{i+1} \rangle$, are of the alternating (period two) form $B_i = B + \delta(-1)^i$ in the symmetry-broken infinite-size state (of which there are two degenerate ones, with oscillations out-of-phase relative to each other). Unlike magnetic order, this kind of order is allowed at zero temperature in one dimension because the broken symmetry (the lattice translational symmetry) is discrete. The modulation $\delta$ becomes non-zero at $g = g_c$ and thereafter increases with $g$. The spin correlations are initially staggered (peaked at momentum $k = \pi$ in reciprocal space), but at $g \approx 0.52$ the change to $k = \pi/2$ [158], and for $g > 1$ the peak-value may change continuously with $g$ (a spiral state) [159].

The model with ferromagnetic (negative) $J_1$ is also interesting. There are several transitions between states with different periodicities [160]). Here we will consider exclusively the antiferromagnetic case, in the regime $g < 1$. We will use Lanczos results to investigate the phase transition into the VBS state, and also discuss the properties of this ordered state.

The Majumdar-Ghosh point. Before we discuss the Lanczos results, it is worth noting that the existence of VBS order can be shown exactly at the special point $g = 1/2$ (the Majumdar-Ghosh point), where the ground state is very simple [60]. On a ring with even $N$, the ground state is a two-fold degenerate singlet product, as illustrated in Fig. 35. One can demonstrate this rather easily by just acting on the states with the hamiltonian, to show that they are eigenstates (while the proof of them being the lowest states is more involved [161, 162, 163]). Out of these degenerate states, $|\Psi_A\rangle$ and $|\Psi_B\rangle$, which break the translational symmetry, one can form symmetric and anti-symmetric states, which have momentum $k = 0$ and $\pi$, respectively:

$$|\Psi(0)\rangle = (|\Psi_A\rangle + |\Psi_B\rangle) / \sqrt{2}, \quad |\Psi(\pi)\rangle = (|\Psi_A\rangle - |\Psi_B\rangle) / \sqrt{2}. \quad (198)$$
These are the states obtaining in Lanczos calculations with conserved momentum. Thus, the order parameter $\langle B_i \rangle = E_1$ is featureless when calculated on finite periodic systems. As always in exact finite-lattice calculations, the symmetry-breaking has to be observed by calculating correlation functions (unless we add some perturbation that breaks the symmetry between the two possible ordering patterns, which has its own complications as the limit of vanishing perturbation has to be taken). The ground state away from the point $g = 1/2$ is more complicated than the simple singlet-products in Fig. 35, with fluctuations in the singlet pairings, but with remaining alternating higher and lower density of singlets on the nearest-neighbor bonds. The translationally invariant $k = 0$ and $\pi$ states on finite rings correspond approximately to symmetric and anti-symmetric combinations as in Eq. (198). It is, however, only at $g = 1/2$ that these two states are exactly degenerate for finite $N$. In other cases, the states become degenerate only in the limit $N \to \infty$.

**Dimer order parameter.** We already discussed VBS ground states in Sec. 2.2 and an example of how to characterize this kind of order by dimer (or bond) correlations in Sec. 2.4.3. We now investigate the dimer correlation function introduced in Eq. (25), written explicitly for a 1D system as

$$D(r) = \langle B_i B_{i+r} \rangle = \langle (S_i \cdot S_{i+1}) (S_{i+r} \cdot S_{i+1+r}) \rangle. \quad (199)$$

This function should alternate between “weak” and “strong” values for large $r$ if there is VBS order (even if the symmetry is not broken). The calculation of this correlation function with the Lanczos method can be simplified by taking advantage of the rotational symmetry and compute $\langle (S_i \cdot S_{i+1}) S_{i+r}^z S_{i+1+r}^z \rangle$, which is $1/3$ of $D(r)$. In principle, we could make it even simpler by defining the order parameter solely in terms of the $z$ components, $\langle S_i^z S_{i+1}^z S_{i+r}^z S_{i+1+r}^z \rangle$, which is not just a constant times (199) but still a valid order parameter for a dimerized state. Here we consider the full $D(r)$.

Fig. 36(a) shows results for a 32-site system at three different couplings. At $g = 0$, the standard Heisenberg model, the correlations decay rapidly with $r$ and are very small at $r = N/2$. At the transition point, $g_c \approx 0.2411$ (which we will determine more precisely below), the correlations are clearly stronger. The Luttinger-liquid theory applied to spin chains predicts dimer correlations decaying as $1/r$ for $g \leq g_c$ (with stronger logarithmic corrections when $g < g_c$) [152]. Going to larger coupling ratios, Fig. 36(a) shows correlations clearly indicative of long-range order at $g = 0.4$. To confirm the presence of long-range order, a finite-size scaling analysis has to be carried out. As we did with the spin correlation functions of the Heisenberg chain in the preceding section, it is

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**FIGURE 35.** Illustration of the degenerate ground states at the Majumdar-Ghosh point. The thick arcs illustrate singlets; the product of these singlets on alternating bonds is an exact ground state at $g = 1/2$. 

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then typically best to look at the correlations at the longest distance, \( r = N/2 \), versus the system size. In the case of the dimer correlations, where the “bare” correlation function is non-zero even in the disordered phase, one has to subtract off the constant corresponding to this background. One possibility is to subtract \( \langle B_i \rangle^2 \), which is the value to which \( D(r) \) should converge for large \( r \). Another option, which we will choose here, is to instead use the difference \( D(N/2) - D(N/2 - 1) \). This quantity will be non-zero for \( N \to \infty \) only if \( D(r) \) oscillates, i.e., if there is long-range dimer order. Results for three different system sizes are shown versus the coupling ratio in Fig. 36(b). Here the order parameter appears to be well converged in the range \( 0.4 < g < 0.6 \) confirming that there is indeed long-range order.

**Finite-size extrapolations of spin and dimer correlations.** To extract the infinite-size dimer order parameter quantitatively, it is in most cases necessary to perform an extrapolation, unless the system size is sufficiently large for there to be no remaining size-dependence of significance [which based on Fig. 36(b) is the case close to \( g = 1/2 \)]. Fig. 37 shows results for both dimer and spin correlations versus the inverse system size for representative \( g \) values both inside and outside the VBS phase.

The dimer correlations, shown in the right panel of Fig. 37, are of the \( 1/r \) form for \( g < g_c \). There are logarithmic corrections, which are small exactly at \( g_c \). For \( g > g_c \) inside the VBS phase, the correlations extrapolate to a non-zero value. The asymptotic \( N \to \infty \) convergence should be exponential in this case (as can be demonstrated explicitly at \( g = 1/2 \)) but close to the transition it is in practice not possible to reach system sizes sufficiently large to observe this behavior. Instead, closer to \( g_c \) the behavior appears to be essentially linear in \( 1/N \), as seen in the figure at \( g = 0.4 \). However, at \( g = 0.45 \) one can see that the behavior is actually non-monotonic, with the large-\( N \) behavior consistent with an approach to the infinite-\( N \) value from below, as at \( g = 1/2 \). As seen in the right panel of Fig. 36, for \( g > 0.6 \) the dimer order parameter decreases with \( g \). Larger systems are required for proper extrapolations in this case, and other methods have to be used. It is believed that the system remains dimerized for all \( g > g_c \) [158, 159].

The spin correlations should change from \( 1/r \) to exponentially decaying as the VBS
phase is entered. To detect this change, the correlation function $C(r = N/2)$ is multiplied by $N/2$ in the left panel of Fig. 37. As we saw before, for $g = 0$ there is a log-correction which enhances the long-distance spin correlations. At $g = g_c$ these corrections are much smaller—it is known that the leading log corrections vanish at the dimerization transition [164]—although the behavior is still not purely $1/r$ (which can be expected only in the limit $N \to \infty$). For $g$ inside the VBS phase, $rC(r)$ decays to zero, as expected. Close to $g_c$ one of course has to go to large system sizes—larger than the spin correlation length (which diverges at $g_c$)—in order to observe a pure exponential fall-off.

For the $g$-values used in Fig. 37, the spin correlations are staggered, i.e., peaked at $k = \pi$ in reciprocal space. For $g \approx 0.52$ the peak position changes rapidly to $\pi/2$ [158], and for $g > 1$ the spin structure most likely evolves into a spiral with continuously varying pitch [159] (as is the case for the classical version of the model). The correlations always decay exponentially, however (unlike the classical long-range spiral).

**Determining the dimerization transition point.** The dimerization transition is known to be similar to the Kosterlitz-Thouless transition of the classical 2D XY model. Unlike a conventional phase transition, the order parameter does not follow a power law at $g_c$, but is exponentially small close to $g_c$. It is therefore not possible to extract the infinite-$N$ order parameter close to $g_c$ based on the small systems accessible to Lanczos calculations—due to the cross-over behavior it is even difficult relatively deep inside the VBS phase. It would then appear to be very difficult to determine the location of the dimerization transition. There is, however, a very elegant way to extract the critical coupling in an indirect way, based on excited-state energies [157].

As we saw in Sec. 4.3.1, the lowest excited state of the pure Heisenberg chain ($g = 0$) is a triplet. On the other hand, we also know that in the VBS state the ground-state should be two-fold degenerate, and both these ground states must be singlets. At the exactly solvable point $g = 1/2$, the degeneracy is exact for any $N$, but away from this special point the two states become degenerate only in the infinite-$N$ limits. The approach to degeneracy should be exponential in $N$ (as will be illustrated with data below), whereas

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**FIGURE 37.** Spin (left panel) and bond (right panel) correlations at the longest distance in periodic chains versus the inverse system size. The couplings ratios for the five curves are $g = 0, g_c, 0.40, 0.45, 0.50$, in the order top to bottom (the left panel) and bottom to top (right panel), i.e., the three sets shown with black circles correspond to the VBS phase.
in the case of the triplet excitation of the Heisenberg chain the gap closes as $1/N$, and the lowest singlet also approaches the ground state as $1/N$ (as shown in Fig. 33). This being the case, for fixed $N$ there must be some coupling ratio $g_{\text{cross}}(N)$ at which the singlet and triplet excitations cross each other. This point can be taken as a size-dependent definition of the transition point, and as $N \to \infty$ it should approach the actual critical coupling: $g_{\text{cross}}(N \to \infty) = g_c$.

Fig. 38(a) shows the three energy levels of interest for a 16-site chain, with the corresponding quantum numbers indicated as well (these quantum numbers apply to all system sizes $N = 4n$). Note how the two singlets become degenerate as $g \to 1/2$. As expected, the singlet and triplet excited levels cross; for this system size at $g \approx 0.242$. The crossing points converge very rapidly as a function of $N$, as shown in Fig. 38(b)—the leading corrections are known to be proportional to $1/N^2$ [164]. Based on these results it is possible to determine $g_c$ very precisely. Fitting a high-order polynomial to crossing points for $8 \leq N \leq 32$ gives $g_c = 0.2411674(2)$, where (2) indicates the uncertainty in the last digit based on fluctuations in the extrapolated value when different ranges of system sizes are included in the fits and the order of the polynomial is varied.

Finite-size gaps in the VBS phase. As already mentioned, to accommodate symmetry breaking in the thermodynamic limit, one would expect the singlet-singlet finite-size gap to close exponentially fast with increasing system size inside the VBS phase (with the exception of the Majumdar-Ghosh point, where the degeneracy is exact for any size). The exponential gap scaling is demonstrated in the left panel of Fig. 39 for some representative values of $g$. At $g_c$ the behavior is instead $\propto 1/N$, which is not clearly seen in this lin-log plot, but more clearly in the log-log plot in the right panel. In the right panel triplet gaps are also shown versus $1/N^2$. Here extrapolations to non-zero values inside the VBS phase are apparent. The finite gap essentially corresponds to the energy required to promote a singlet bond into a triplet. Note that only the ground state is exactly solvable
FIGURE 39. Singlet (left panel) and triplet (right panel) gaps as a function of system size for different coupling ratios. Here $g_{\text{cross}}$ is the size-dependent coupling at which the lowest singlet and triplet excitations are degenerate (and thus $\Delta_s = \Delta_t$). The gap at $g_{\text{cross}}$ decays as $1/N$ (as seen more clearly in the right panel). In the left panel a log scale for $\Delta_s$ versus $N$ is used to show the exponentially rapid closing of the gap between the two singlet states that become degenerate for $N \to \infty$ in the ordered VBS phase. To the right, the triplet gap is plotted versus $1/N$ to show its convergence to a non-zero value in the VBS phase.

at the Majumdar-Ghosh point, and the triplet gap is size dependent also here. At $g_c$ the gaps scale as $1/N$—in the figure the behavior at the size-dependent crossing points $g_{\text{cross}}$ is shown and also scales as $1/N$.

4.3.3. Chains with long-range interactions

While long-range spin ordering is not possible in Heisenberg chains with finite-range interactions, long-range interactions make magnetic order possible at $T = 0$. A transition between a Néel state and a quasi-long-range-ordered (QLRO) state (the Heisenberg critical state with spin correlations decaying as $1/r$, discussed in the preceding sections) takes place in a system with distance-dependent couplings of the form $J_r \propto (-1)^{r-1}/r^\alpha$ [72]. The signs here favor antiferromagnetic order, and there is no frustration. When the exponent $\alpha < \alpha_c \approx 2$, the ground state has true long-range Néel order, while for $\alpha > \alpha_c$ the system is in the QLRO phase. The critical value of the long-range interaction parameter $\alpha_c$ depends on details of the couplings, e.g., on the nearest-neighbor coupling $J_1$ when all other $J_r$ are fixed, and the critical exponents at the transition are continuously varying (in contrast to the constant exponents throughout the QLRO phase).

Another interesting example of a system with long-range interactions is the Haldane-Shastry chain [165, 166], with frustrated interactions $J_r = 1/r^2$. It has a critical ground state similar to that of the standard Heisenberg chain, but, in field-theory language, the marginal operator causing the leading log-corrections vanishes [167]. The system is, thus, right at a dimerization transition such as the one discussed above for the $J_1$-$J_2$ chain. It can be noted that $J_2/J_1 = 1/4$ in the Haldane-Shastry model, which is quite close to the critical ratio $g_c = 0.2411$ of the $J_1$-$J_2$ chain. Thus, the interactions beyond distance $r = 2$ in the Haldane-Shastry chain only play a minor role (but, importantly, actually make the system exactly solvable [165]).
FIGURE 40. Ground state phases of the chain with frustration parameter \( g \) and exponent \( \alpha \) in the long-range interaction, reproduced from Ref. [73]. The dashed curves indicate continuous phase transitions, whereas the thick solid curve represents a first-order transition. The thin solid curve for \( g < 0 \) corresponds to the system with unfrustrated interaction studied in [72]. At \( \alpha^{-1} = 0 \) (the \( J_1-J_2 \) chain) the dominant spin correlations in the VBS state change from \( q = \pi \) to \( \pi/2 \) at \( g \approx 0.52 \) [158]. The curve where this change occurs for \( \alpha^{-1} > 0 \) connects to the multi-critical point where all the phase boundaries come together.

In the presence of long-range interactions one can also realize a direct 1D quantum phase transition between a Néel state and a VBS. Here we discuss the system introduced in Ref. [73], which combines unfrustrated long-range interactions and short-range frustration (at separation \( r = 2 \)) according to the Hamiltonian

\[
H = \sum_{r=1}^{N/2} J_r \sum_{i=1}^{N} \mathbf{S}_i \cdot \mathbf{S}_{i+r},
\]

where the distance dependence of the couplings is given by

\[
J_1 = \frac{1}{J_\Sigma}, \quad J_2 = g, \quad J_{r>2} = \frac{1}{J_\Sigma} \left( -1 \right)^{r-1} r^{-\alpha}, \quad J_\Sigma = 1 + \sum_{r=3}^{N/2} \frac{1}{r^{\alpha}}.
\]

Here, with the normalization by \( J_\Sigma \) of all couplings but \( J_2 \), the sum of all non-frustrated interactions \( |J_r| \) equals 1, and the limit \( \alpha \to \infty \) therefore corresponds exactly to the \( J_1-J_2 \) chain with \( g = J_2/J_1 \). This will be useful when comparing the two models and also guarantees a finite energy per spin for \( N \to \infty \), even for \( \alpha < 1 \). Instead of summing \( J_r \) up to \( r = N/2 \), one could also include \( N/2 < r < N \). This should not affect the phase boundaries and critical exponents for \( \alpha > 1 \), however.

We will study the evolution if the dimerization transition occurring as a function of the frustration strength \( g \) as the inverse \( \alpha^{-1} \) of the long-distance interaction exponent is increased from 0. As shown in the semi-quantitative phase diagram in Fig. 40 (constructed on the basis of Lanczos results, as will be discussed below) this continuous transition persists until \( \alpha \approx 2 \), while for smaller \( \alpha \) it evolves into a first-order transition between the Néel state and a state with coexisting VBS order and critical (or possibly long-ranged) spin correlations at wave-number \( q = \pi/2 \). This state is denoted in the phase diagram as VBS+QLRO(\( \pi/2 \)).

**Evolution of the dimerization transition.** We have discussed how to study the dimerization transition based on crossings of excited-state energies for the \( J_1-J_2 \) chain, with
FIGURE 41. Low-energy levels of a 16-spin system at two values of the long-range exponent; $\alpha = 4$ (left) and $\alpha = 1$ (right). The spin $S$ and the momentum $k$ of the states are indicated in the left panel (and note that two levels are shown with $S = 0, k = 0$). The inset in the right panel shows the avoided level crossing of the two $k = 0$ singlets in greater detail. There is a very small gap between these singlets where they come together, but this is much smaller than what can be resolved in this figure.

An illustration in Fig. 38(a). The same physics applies in the presence of the long-range interaction as well, if $\alpha$ is sufficiently large. This is shown for a 16-spin chain at $\alpha = 4$ in the left panel of Fig. 41. The lowest $k = 0$ and $k = \pi$ singlets should become degenerate in the VBS phase for $N \to \infty$ (so that a symmetry-broken dimerized states can be formed). A region of very near degeneracy for $g > 1/2$ can be seen in the figure. The region of approximate degeneracy, which is not easy to demarcate precisely, expands very slowly toward smaller $g$ with increasing $N$. As shown in Fig. 42, the singlet-triplet crossing point converges with the system size [although slower than the $1/N^2$ convergence shown for $J_1-J_2$ the chain in Fig. 38(b)] and can reliably give the QLRO($\pi$)--VBS phase boundary $g_c(\alpha)$ for $\alpha > 2$.

Another interesting feature of the energy levels is that, upon decreasing $\alpha$ below $\approx 2$, the broad maximum in the ground state energy versus $g$ becomes increasingly sharp. As seen in the right panel of Fig. 41, at $\alpha = 1$ it has developed into a sharp tip due to an avoided level crossing with the second singlet at $k = 0$. The real singlet-triplet crossing has moved to the same region. An avoided level crossing between two states with the same quantum numbers, leading to a discontinuity in the derivative of the ground state energy with respect to $g$ for $N \to \infty$, is the hallmark of a first-order transition. It should be noted that it is not just the two lowest singlets that exhibit this kind of avoided level crossing. Other low-energy states as well come together in the neighborhood of the transition, in a complicated cascade of level crossings—all of these should converge to a single point at the first-order transition when $N \to \infty$. The nature of the phases at this transition will be discussed below. First, let us investigate in more detail how the dimerization transition evolves from continuous to first-order.

Fig. 42 shows the size dependence of the level crossing $g_{\text{cross}}$ and the location $g_{\text{peak}}$ of the maximum in the ground state energy. In the standard Majumdar-Ghosh frustrated chain the size correction to the crossing point is $\propto 1/N^2$ [as seen in Fig. 38(b)], which also can be seen for large $\alpha$. For smaller $\alpha$, the corrections instead seem to be $\propto 1/N$, but a cross-over to $1/N^2$ for large $N$ seems likely as long as the transition remains
FIGURE 42. Dependence on the inverse chain length of the singlet-triplet crossing point $g_{\text{cross}}$ and the location $g_{\text{peak}}$ of the ground state energy maximum for different long-range interaction exponents $\alpha$. The two lines show extrapolations of the $g = 2.0$ data to $N = \infty$.

continuous. The peak location moves in the opposite direction. For some $\alpha$ and $N \to \infty$, $g_{\text{cross}}$ and $g_{\text{peak}}$ should coincide. The results indicate that both $g_{\text{cross}}$ and $g_{\text{peak}}$ have dominant $1/N$ corrections at this point. Line fits are shown in Fig. 42 at $\alpha = 2$, were there is still a small gap between the two extrapolated values (and the extrapolations may not be completely accurate, if the linear scaling holds asymptotically only at the special $\alpha$-value for which $g_{\text{cross}} = g_{\text{peak}}$). At $\alpha = 1.7$, where the transition is first-order, they should coincide (and then the asymptotic size correction should be exponential).

First-order Néel–VBS transition. To confirm an avoided level crossing with a discontinuous energy derivative for $\alpha > 1.8$ (approximately), the second derivative of the ground state energy at its maximum is graphed on a lin-log scale in Fig. 43. These results were obtained based on calculations of the energy on a very dense grid of points close to the peak value, to which a polynomial could be reliably fitted. The second derivative extracted from this polynomial grows exponentially with $N$ for $\alpha = 1.5$, showing that the slope of the energy curve indeed changes discontinuously for an infinite chain. In contrast, at $\alpha = 3$ the second derivative decreases for large $N$. For $\alpha = 2$ convergence to a finite value also seems plausible, whereas $\alpha = 1.7$ and 1.8 appear to be close to a separatrix (where the form of the divergence is consistent with a power law) between the two different behaviors. This analysis suggests that the continuous dimerization transition changes smoothly into a first-order transition at ($g_m \approx 0.41, \alpha_m \approx 1.8$). The singlet-triplet crossing moves toward the ground-state energy maximum and coincides with it at the multi-critical point ($g_m, \alpha_m$), beyond which it develops into a first-order singularity.

To analyze the states involved in the first-order transition, we next investigate the standard spin correlation function $C(r)$ and the dimer correlation function $D(r)$ defined in (199). We will also study the Fourier transforms of these correlation functions; the static spin ($s$) and dimer ($d$) structure factors:

$$S_s(q) = C(0) + C(N/2) + 2 \sum_{r=1}^{N/2-1} \cos(qr)C(r),$$

(202)
FIGURE 43. Size dependence of the second derivative of the ground state energy with respect to the frustration parameter $g$ at the point $g_{\text{peak}}$ where the ground state energy takes its maximum value.

$$S_d(q) = D(0) + D(N/2) + 2 \sum_{r=1}^{N/2-1} \cos(qr)D(r).$$

If there is long-range order at some wave-vector $q = Q$, then the corresponding structure factor is proportional to $N$ (for large $N$), and order parameters can therefore be defined as $S_{s,d}(Q)/N$. Here $Q = \pi$ for both the Néel state and the dimerized VBS state. Fig. 44 shows the $g$-dependence of $S_d(\pi)/N$ and $S_s(\pi)/N$ at $\alpha = 1.5$. Discontinuities are seen to develop at $g \approx 0.42$, in agreement with the first-order scenario (and at $g = 1$, not shown here, the discontinuities are much sharper). Both the Néel and VBS order parameters are well converged to non-zero values in their respective phases, and small in the other phase (where they should vanish when $N \to \infty$, but it is not possible to observe this clearly because of the small system sizes). The spin structure factor at $q = \pi/2$ is also shown in the figure. Interestingly, it also becomes large in the VBS phase, while in the Néel phase it should decay to zero with increasing $N$ (which is plausible based on this data, but, again, not possible to see clearly for these small systems). Based on these results one might conclude that the VBS state obtaining at the first-order transition has co-existing long-range magnetic order at $k = \pi/2$ (i.e., a spiral with period four). Considering the small system sizes, this cannot be guaranteed, however. The behavior could also reflect a slow power-law decay of the spin correlations.

Fig. 45 shows the real-space spin and dimer correlations at $\alpha = 1$ for two $g$ values, at either side of the first-order transition. At $g < g_c$, in the Néel state, the spin correlations are staggered and clearly long-ranged (with almost no decay seen as a function of $r$ for $r > 1$). At this coupling there is no structure in $D(r)$, i.e., there is no VBS order. For $g > g_c$ there is clearly VBS order, with $D(r)$ showing the characteristic staggered pattern. Strong period-four spin correlations are also observed, but it is not clear whether these are long-ranged or decay slowly to zero with increasing $r$. The behavior is consistent with an $1/r$ decay, but much longer chains would be needed to extract the behavior reliably. An interesting point to note here is that the dimer correlations oscillate around zero, even though no constant has been subtracted. This is in contrast to the oscillations...
around a positive value in the $J_1$-$J_2$ chain, as seen in Fig. 36. Looking at the definition of the dimer correlation function in Eq. (199), this behavior is another manifestation of period-four spin structure, i.e., at odd separation of the bond operators $S_i + S_{i+1}$, a correlation between a ferromagnetic and an antiferromagnetic bond is measured. While the nature of the magnetic structure in this VBS is not completely clear, the correlation functions seem to indicate a tendency to bond-ordering of the form $|ststst\cdots\rangle$, where $s$ and $t$ represent nearest-neighbor bonds with high singlet and triplet density, respectively. In Fig. 40 the state has been denoted as magnetically quasi-long-range ordered, a VBS–QLRO($\pi/2$) coexistence state, but it could in fact also be a state with coexisting VBS and long-range magnetic order. It should also be noted that this phase could in principle be very sensitive to boundary conditions on small lattices, and it cannot be excluded that the actual magnetic structure is spiral-like, with continuously varying pitch (as in the $J_1$-$J_2$ chain with $g > 1$ [159]).

The VBS+QLRO($\pi/2$) state should have gapless spin excitations, regardless of the spin correlations as long as they are not exponentially decaying. The lowest triplet is at $k = \pi/2$. It is, however, difficult to demonstrate the gaplessness based on data for small systems, because the size dependence of the gaps (and other quantities) for system sizes $N = 4n$ exhibit large even-odd $n$ oscillations (as well as other irregular size effects). In the conventional VBS phase (in the low-right part of the phase diagram in in Fig. 40) the lowest triplet is at $k = \pi$, even when the spin correlations (which are exponentially decaying in this phase) are peaked at $k = \pi/2$. The level crossing between the lowest $k = \pi$ and $k = \pi/2$ triplets can therefore in principle be used to extract the boundary between the VBS and VBS+QLRO($\pi/2$) phases. The size dependence of the crossing point is not smooth, however, and cannot be extrapolated very reliably. The boundary between dominant $k = \pi$ and $k = \pi/2$ spin correlations has also not been extracted accurately. This change in the spin correlations may be associated with a transition to a state with periodicity four [158], although there are no sign of the VBS order changing.

**QLRO–Néel transition.** Let us return to Fig. 41 for another interesting feature of the level spectrum: The lowest singlet excitation for small $g$ has momentum $k = \pi$ for $\alpha = 4$ but $k = 0$ for $\alpha = 1$. The switching of the order of these levels as a function of $\alpha$ for
FIGURE 45. Spin (left) and dimer (right) correlations in a 32-spin chain at $\alpha = 1$ for two values of the frustration parameter. At $g = 0.25$ and $0.45$ the system is in the Néel and VBS–QLRO($\pi/2$) phase, respectively. The first-order transition in this case occurs at $g \approx 0.39$.

$g < g_c$ is associated with the Néel–QLRO($\pi$) transition. The level crossings can be used to extract this phase boundary very accurately up to $g \approx 0.25$ (while for higher $g$ the $N \to \infty$ extrapolations become difficult). While it is not completely clear why the $k = 0$ and $k = \pi$ excited singlets cross each other at this transition, it can again be noted that the QLRO($\pi$) phase the low-energy excitations arise from two deconfined spinons. The lowest (for $N = 4n$) is a triplet at $k = \pi$. The lowest singlet is also at $k = \pi$, with a small finite-size gap to the triplet due to weak spinon-spinon interactions (and Fig. 33 shows and example of how these levels become degenerate as $N \to \infty$). In the Néel phase the spinons are no longer deconfined, and the structure of the low-energy spectrum changes.

As we will discuss below in Sec. 4.4.2, the 2D Néel state on finite lattices has low-energy excitations analogous to quantum-rotor states, which have $S = 1, 2, 3, \ldots$ and become degenerate with the ground state as the system size increases. They have momenta $k = (0,0)$ and $k = (\pi, \pi)$ for even and odd $S$, respectively. One might expect this kind of quantum rotor “tower” in the 1D Néel state as well. Apparently, the lowest $k = \pi$ singlet, which would not be part of such a rotor tower, is then pushed up much higher in energy, and the $k = 0$ singlet takes over as the lowest singlet excitation.

It is useful to compare the level crossing approach with the QMC calculation in Ref. [72] for the unfrustrated model with $J_1 = 1$ and $J_{r>1} = \lambda (-1)^{r-1}/r^\alpha$. A reparametrization of this model for $\lambda = 1$, to the convention used in (201), gives the curve for $g < 0$ shown in Fig. 40. Finite-size scaling of QMC data for the Néel order parameter gave $\alpha_c = 2.225 \pm 0.025$, for $N$ up to 4096 [72]. Extrapolating the $k = 0, \pi$ singlet crossing points (which have size corrections $\propto 1/N^{\beta}$, with $\beta \approx 1.50$ for $N \leq 32$ gives a marginally higher (and probably more reliable) value; $\alpha_c = 2.262 \pm 0.001$. Analyzing the singlet and triplet gaps at the crossings, assuming $\Delta \sim N^{-z}$, gives the dynamic exponent $z = 0.764 \pm 0.005$, in very good agreement with Ref. [72]. For the frustrated model (201) on the QLRO($\pi$)–Néel boundary, the gap scaling in $1/N$ give $z \approx 0.75$ for $g$ up $\approx 0.25$, while for larger $g$ it is not possible to reliably extract $\alpha_c$ and $z$ this way, because of large scaling corrections and the absence of level crossings for increasingly large systems as the multi-critical point ($g_m \approx 0.41, \alpha_m \approx 1.8$) is approached (for reference $\alpha_c = 2.220 \pm 0.005$ for $g = 0$ and $2.170 \pm 0.01$ for $g = 0.2$).
At the multi-critical point \( z \) can be extracted using gaps at the singlet-triplet crossing point and the ground state energy maximum, which gives \( z \approx 0.8 \). It is, thus, possible that the dynamic exponent is constant, \( z \approx 3/4 \), on the whole QLRO(\( \pi \))–Néel boundary, including the multi-critical point where all phase boundaries come together.

**Technical notes.** The model we have studied here is rather challenging for a 1D system, from a technical standpoint, because of the long-range interactions. The combination of long-range interactions and frustration makes it difficult to apply other computational techniques. While efficient QMC techniques can be applied to systems with unfrustrated long-range interactions [168, 72], this is no longer possible in the presence of the frustrating \( J_2 \) term, due to the sign problem (discussed in Sec. 5). The DMRG method [28, 29], on the other hand, can handle frustration but not easily long-range interactions. It would still be interesting to try to apply DMRG, or related techniques based on matrix-product states [21], to this system, in particular to study further the spin structure in the putative VBS-QLRO(\( \pi/2 \)) state.

The Lanczos calculations presented above exploited all the symmetries discussed in Sec. 4.1. The Hamiltonian was stored in compact form on disk, with a separate file for each interaction distance \( r \) in (201). These files were read and used one-by-one in each operation with the Hamiltonian according to code \{22\}. For \( N = 32 \), the number of Lanczos iterations needed was typically 60 \( \sim \) 80 (and less for smaller systems). All basis states were saved in primary memory, and re-orthogonalization to all previous states was carried out after each iteration (as in code \{19\}).

### 4.4. Two-dimensional systems

We now discuss exact diagonalization in two dimensions, using the important case of the Heisenberg model on the simple periodic square lattice as an example. As in one dimension, we can use momentum conservation to block diagonalize the Hamiltonian within the sectors of fixed magnetization. It is a little more complicated to take maximum advantage of lattice symmetries to further split some of the blocks, because of the larger number of symmetry operations commuting with the Hamiltonian. On the positive side, the reward for implementing all the lattice symmetries is that, for high-symmetry momenta, where lattice reflections and/or rotations can be used, the blocks are smaller for given \( N \) than in one dimension. In practice, this may be of little help, since the maximum linear size \( L \) of an \( L \times L \) lattice that can be diagonalized is still pitifully small (\( L = 6 \)). Exact diagonalization studies are still of great value.

In one dimension, we used the reflection (parity) operation \( P \) and constructed momentum eigenstates based on representatives \( |a\rangle \) and their reflections, \( (1 + pP)|a\rangle \) (and later we added also spin-inversion). Although \( P \) does not commute with the translation operator, we found that for the two special momenta \( k = 0, \pi \), both \( k \) and \( p \) are still good quantum numbers. For general \( k, p \) is not a valid quantum number, but with semi-momentum states, mixing \( \pm k \) states, we could use \( P \) to accomplish a real-valued representation. In two dimensions one can in principle also construct a real-valued Hamiltonian for any momentum, but this is much more complicated in practice than in one dimension, and...
in the end it is not worth the effort (since it does not increase the system sizes that can be studied). Here we will therefore work with standard complex momentum states.

**Defining the lattice.** As in one dimension, we use the state notation $|S^z_0, \ldots, S^z_{N-1}\rangle$, which is practical with the bit representation of the spins. For a 1D system the spin indices correspond directly to the chain geometry, but for a 2D lattice we have to establish a labeling convention for the sites. We will consider rectangular lattices with $N = L_x \times L_y$ sites, at coordinates $(x,y)$ with $x = 0, \ldots, L_x - 1$ and $y = 0, \ldots, L_y - 1$. A natural choice is to number the sites $i = 0, \ldots, N - 1$ such that $i = x_i + y_i L_x$, as illustrated for a $4 \times 4$ system in Fig. 46. In a notation not explicitly dependent on the lattice, we can write the Heisenberg Hamiltonian as

$$H = J \sum_{b=1}^{N_b} \mathbf{S}_{i(b)} \cdot \mathbf{S}_{j(b)},$$

(204)

where $[i(b), j(b)]$ are the two nearest-neighbor sites connected by bond $b$. Specifying the lattice then just amounts to creating this list of site pairs.

### 4.4.1. Momentum states in two dimensions

We again define translations of the spin indices, now in both the $x$ and $y$ directions, as illustrated in Fig. 47, with corresponding operators $T_x$ and $T_y$ defined by

$$T_x |S^z_0, \ldots, S^z_{N-1}\rangle = |S^z_{T_x(0)}, \ldots, S^z_{T_x(N-1)}\rangle,$$

$$T_y |S^z_0, \ldots, S^z_{N-1}\rangle = |S^z_{T_y(0)}, \ldots, S^z_{T_y(N-1)}\rangle,$$

(205)

---

5 A real maximally blocked basis exists because the Hamiltonian is real and symmetric, which guarantees real eigenvalues and eigenvectors. If the Hamiltonian is diagonalized numerically as a single block, the eigenvectors in degenerate multiplets will be mixed, and the translation operators will not be diagonal (because their eigenvalues are complex). The degenerate complex momentum states have been mixed in such a way as to make the linear combinations real. Such a superposition can in principle be applied to block-diagonalize the original Hamiltonian matrix into semi-momentum like real-valued blocks.
Ty \uparrow
T_x \rightarrow

FIGURE 47. Symmetries used for the two-dimensional square lattice. $T_x$ and $T_y$ translate the spins by one lattice spacing in the positive $x$ and $y$ directions, respectively. The lattice is periodic; the open circles represent sites of the opposite edges. $P_x$ and $P_y$ are reflections with respect to the $x$- and $y$-axis, and $P_d$ and $P_e$ reflect with respect to the two diagonal axes. For even $L$ (which normally should be used for an antiferromagnet), the $P_x$ and $P_y$ axes pass between lattice sites. The reflections $P_d$ and $P_e$ are about the lines connecting the far corners, which go through the sites on the diagonals (and hence leave the spins on those sites unchanged upon reflection).

where the translated spin indices are

$$T_x(i) = [x_i - 1]L_x + y_iL_x,$$
$$T_y(i) = x_i + ([y_i - 1]L_y)L_x,$$  \hfill (206)

with $[\gamma - 1]L_\gamma$ denoting the modulus of $\gamma - 1$ with respect to $L_\gamma$, i.e., $[-1]L_\gamma = L_\gamma - 1$.

Using these translations, a momentum state based on a representative $|a\rangle$ is defined as

$$|a(k)\rangle = |a(k_x, k_y)\rangle = \frac{1}{\sqrt{N_a}} \sum_{x=0}^{L_x-1} \sum_{y=0}^{L_y-1} e^{-i(k_x x + k_y y)} T_y^y T_x^x |a\rangle,$$  \hfill (207)

where the possible momenta are

$$k_\gamma = \frac{2\pi}{L_\gamma} m_\gamma, \quad m_\gamma = 0, 1, \ldots, L_\gamma - 1, \quad \gamma \in \{x, y\}.$$  \hfill (208)

The normalization constant $N_a$ depends on the translational properties of the representative, i.e., the number of different states $D_a$ obtained among the group of $L_x \times L_y$ translations of the representative $|a\rangle$ (e.g., for a state with the spins in a checker-board pattern $D_a = 2$). A representative is incompatible with the momentum if the sum of phases $F_a$ in (207) over the translations bringing $|a\rangle$ onto itself vanishes. For a compatible state, the normalization constant $N_a = D_a |F_a|^2$. The easiest way to compute this in a program is simply to carry out all the translations and sum up $D_a$ and $F_a$ in the process, instead of using explicit formulas as we did for 1D systems (where one can of course also use the more brute-force approach).

The construction of the hamiltonian matrix proceeds as in the 1D case. We again split the hamiltonian into a diagonal piece $H_0$ and off-diagonal bond operators $H_j$, as in (121) and (122), where now $j = 1, \ldots, N_b$. Acting with these operators on the representative $|a\rangle$, we again may have to translate the resulting state in order to obtain the new representative $|b_j\rangle$ corresponding to $H_j|a\rangle$, i.e.,

$$H_j|a\rangle = h_j(a) T_y^y T_x^x |b_j\rangle.$$  \hfill (209)
Acting on a momentum state, we thus get

$$H_j |a(k_x, k_y)\rangle = h_j(a) \frac{L_x-1 L_y-1}{\sqrt{N_a}} \sum_{x=0}^{L_x-1} \sum_{y=0}^{L_y-1} e^{-i(k_x x + k_y y)} T_y^{(y-l_y)} T_x^{(x-l_x)} |b_j\rangle$$

(210)

$$= h_j(a) \frac{e^{-i(k_x l_x + k_y l_y)}}{\sqrt{N_a}} \sum_{x=0}^{L_x-1} \sum_{y=0}^{L_y-1} e^{-i(k_x x + k_y y)} T_y^{(y-l_y)} T_x^{(x-l_x)} |b_j\rangle,$$

and taking the overlap with $\langle b_j(k_x, k_y) |$ gives the matrix element

$$\langle b(k_x, k_y) | H_j |a(k_x, k_y)\rangle = h_j(a) e^{-i(k_x l_x + k_y l_y)} \frac{N_{b_j}}{N_a},$$

(211)

The normalization constants (or some integer mapping to their possible numerical values) should be stored along with the representatives.

**Incorporating other lattice symmetries.** We here consider a quadratic, $L \times L$, system. There are eight independent rotations and reflections of such a lattice, which can be taken as, e.g., the four $90^\circ$ rotations and a reflection about a horizontal or diagonal axis. It can be easily verified that additional reflections can be written as combinations of rotations and the first reflection. The easiest way to see this is to just consider all rotations and reflections of a $2 \times 2$ array;

$$
\begin{pmatrix}
4 & 3 \\
1 & 2 \\
\end{pmatrix}
\begin{pmatrix}
3 & 2 \\
4 & 1 \\
\end{pmatrix}
\begin{pmatrix}
2 & 1 \\
3 & 4 \\
\end{pmatrix}
\begin{pmatrix}
1 & 4 \\
2 & 3 \\
\end{pmatrix}
\begin{pmatrix}
3 & 4 \\
2 & 1 \\
\end{pmatrix}
\begin{pmatrix}
2 & 3 \\
1 & 4 \\
\end{pmatrix}
\begin{pmatrix}
1 & 2 \\
4 & 3 \\
\end{pmatrix}
\begin{pmatrix}
4 & 1 \\
3 & 2 \\
\end{pmatrix}.
\end{pmatrix}

(212)

Here the first row contains all the rotations of the first array, and in the second row those arrays have been reflected by exchanging the two columns. Any other reflection will just produce an array which is already in the above set of eight. Other permutations of the elements do not correspond to a combination of rotations and reflections (and hence do not correspond to symmetries of hamiltonians we are normally interested in). In practice, we can use any convenient set of reflections, or rotations and reflections, with which the eight unique transformations can be generated.

Using translations as well as a set of other symmetry operators, which we for now collectively add together into a single operator $Q$, with corresponding quantum numbers $\{q\}$, a momentum state can be defined as

$$|a^\sigma(k, \{q\})\rangle = \frac{1}{\sqrt{N_a}} \sum_{r_x=0}^{L_x-1} \sum_{r_y=0}^{L_y-1} e^{-i(k_x r_x + k_y r_y)} T_y^{r_y} T_x^{r_x} Q |a\rangle.$$ 

(213)

As we have seen above, there are more symmetry operations than unique transformations. We can select the ones that are most convenient for given momentum. Instead of using the rotations of the square lattice, we will here use the reflections $P_x$ and $P_y$ about
the $x$ and $y$-axis, respectively. These correspond to symmetries of the hamiltonian for any rectangular ($L_x \times L_y$) lattice (and note that the $180^\circ$ rotation is equivalent to $P_x P_y$). For generic momenta, the reflections do not commute with the translations, and there can be no further blocking using $P_x$ and $P_y$, but there are high-symmetry momenta for which $P_x$ or $P_y$, or both, can be used. For $k_x = \pm k_y$ we will instead use the reflections $P_d$ and $P_e$ about the two perpendicular diagonal axes. All the reflections are defined in Fig. 47. For the most special momenta, $k = (0,0)$ and $(\pi, \pi)$, and $L_x, L_y$ arbitrary, we can also use $P_d$ (or $P_e$) in addition to $P_x, P_y$.

To achieve maximal block-diagonalization for a given momentum $(k_x, k_y)$, the operator $Q$ in (213) can be chosen according to:

$$Q = \begin{cases} 1, & \text{general } k \\ (1 + p_x P_x), & k = (0, k_y), (\pi, k_y) \\ (1 + p_y P_y), & k = (k_x, 0), (k_x, \pi) \\ (1 + p_d P_d), & k_x = +k_y (L_x = L_y) \\ (1 + p_e P_e), & k_x = -k_y (L_x = L_y) \\ (1 + p_y P_y)(1 + p_x P_x), & k = (0, 0), (\pi, \pi), p_x \neq p_y \\ (1 + p_d P_d)(1 + p_p P_p)(1 + p_x P_x), & k = (0, 0), (\pi, \pi), p_x = p_y \end{cases} \tag{214}$$

where again all the reflection quantum numbers take the values $\pm 1$. Note the restriction $p_x = p_y$ in case (g), i.e., there are no such states for $p_x \neq p_y$. In that case, we can of course also use option (f), but using (g) splits the blocks further into two.

To prove that the symmetry operators $Q$ defined above indeed do have the corresponding conserved quantum numbers within the different momentum sectors, the following relationships between the lattice transformations are useful:

$$P_x T_x = T_x^{-1} P_x \quad P_y T_y = T_y^{-1} P_y \\
P_d T_x = T_x P_d \quad P_d T_y = T_y P_d \\
P_e T_x = T_x^{-1} P_e \quad P_e T_y = T_y^{-1} P_e \\
P_d P_x = P_x P_d \quad P_d P_y = P_y P_d \\
P_e P_x = P_y P_e \quad P_e P_y = P_x P_e. \tag{215}$$

The remaining pairs are commuting operators; $[T_x, T_y] = 0$, $[T_x, P_y] = 0$, $[T_y, P_x] = 0$, $[P_x, P_y] = 0$, $[P_e, P_d] = 0$. Let us just check the most complicated case, $(g)$ in (214). The permutations commute with the translations for these special momenta, for the same reasons as discussed for 1D systems in Sec. 4.1.4. We then only have to investigate the properties of $Q$. It is clear that the state is an eigenstate of $P_d$, that being the left-most reflection operator. To check whether it is also an eigenstate of $P_x$ and $P_y$, we use some of the relationships in (215) to verify that $P_x Q = p_x Q$ and $P_y Q = p_y Q$. After a little algebra we get

$$P_x (1 + p_d P_d)(1 + p_y P_y)(1 + p_x P_x) = P_x (1 + p_x p_y p_d P_d)(1 + p_y P_y)(1 + p_x P_x), \tag{216}$$

and an analogous result for $P_y$. Thus, if $p_x = p_y$, the state is indeed, for the special momenta $(0,0)$ and $(\pi, \pi)$, an eigenstate also of $P_x$, $P_y$, and $P_d$. 

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If we want to study all momentum sectors, using the reflection symmetries does not buy us much, because in most of the blocks we have to use the generic option $Q = 1$ in (214)(a). These symmetries are useful when investigating the ground state, which often is in the maximally blockable sector (g). The most useful aspect of the reflections may still be just the fact that they give insights into the symmetry-aspects of the states (e.g., to classify excitations). Apart from the lattice symmetries, spin-inversion symmetry can be used in the $m_z = 0$ sector, exactly as in the 1D case.

We have considered combinations of reflection symmetries but could instead have used a combination of rotations and reflections. This would give us different quantum numbers, but the same size of the blocks. In group theory, there are irreducible representations, with standardized names, corresponding to various lattice symmetries. Instead of working out the applicable symmetries for a new case, one can look up the irreducible representations and their corresponding character tables and employ these to construct states with quantum numbers of a chosen irreducible representation. This is very useful when studying more complicated lattices. Ref. [143] has a discussion of this more formal approach, with examples for chains and square lattices.

We have here used symmetries applicable only to $L_x \times L_y$ lattices with the edges along the $x$- and $y$-axis on the square lattice. With $L_x = L_y = L$, which is the most interesting if we are interested in approaching the infinite 2D lattices, we are then limited to $L = 4$ and $L = 6$ (since odd $L$ introduces undesirable frustration effects for an antiferromagnet). To increase the set of accessible lattice sizes $N$, one can also consider “tilted” lattices, with edges that are not parallel to the square-lattice axes [143].

**Implementation.** To study a $6 \times 6$ lattice we have to use long (8-byte) integers for the bit representation. To save time, it is ten better not to loop over all the $2^N$ possible state-integers and single out the ones corresponding to a given magnetization (as is done, e.g., in code {7}), but to use a scheme which from the outset only constructs states with a given $m_z$ (i.e., given number $n\uparrow$ of $\uparrow$ spins). This can of course also be done for 1D systems. Such a more sophisticated generation of specific $m_z$ states starts with the integer $a = 2^{n\uparrow + 1} - 1$, in which the first $n\uparrow$ bits are 1 and the rest are 0, and then execute:

```plaintext
do
    call checkstate(a, pass)
    if (pass) then $M = M + 1$; store representative information endif
    $c = 0$
do $b = 1, N$
    if ($a[b-1] = 1$) then
        if ($a[b] = 1$) then
            $c = c + 1$
elsee $a[0], \ldots, a[c-1] = 1$; $a[c], \ldots, a[b-1] = 0$; $a[b] = 1$; exit
        endif
    endif
endo
if ($b = N$) exit
endo
```
TABLE 3. Sizes $M(L)$ of the $k = (0,0)$ state blocks for $L \times L$ lattices ($L=4,6$) with magnetization $m_z = 0$ and different reflection and spin-inversion quantum numbers. For $p_x = -p_y$, there is no quantum number $p_d$. For $p_x = -1, p_y = 1$ the block structure is the same as for $p_x = 1, p_y = -1$.

<table>
<thead>
<tr>
<th>$p_x$</th>
<th>$p_y$</th>
<th>$p_d$</th>
<th>$z$</th>
<th>$M(L=4)$</th>
<th>$M(L=6)$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$+$1</td>
<td>$+$1</td>
<td>$+$1</td>
<td>$+$1</td>
<td>107</td>
<td>15,804,956</td>
</tr>
<tr>
<td>$+$1</td>
<td>$+$1</td>
<td>$+$1</td>
<td>$-$1</td>
<td>46</td>
<td>15,761,166</td>
</tr>
<tr>
<td>$+$1</td>
<td>$+$1</td>
<td>$-$1</td>
<td>$+$1</td>
<td>92</td>
<td>15,796,390</td>
</tr>
<tr>
<td>$+$1</td>
<td>$+$1</td>
<td>$-$1</td>
<td>$-$1</td>
<td>38</td>
<td>15,752,772</td>
</tr>
<tr>
<td>$-$1</td>
<td>$-$1</td>
<td>$+$1</td>
<td>$+$1</td>
<td>50</td>
<td>15,749,947</td>
</tr>
<tr>
<td>$-$1</td>
<td>$-$1</td>
<td>$+$1</td>
<td>$-$1</td>
<td>45</td>
<td>15,739,069</td>
</tr>
<tr>
<td>$-$1</td>
<td>$-$1</td>
<td>$-$1</td>
<td>$+$1</td>
<td>42</td>
<td>15,741,544</td>
</tr>
<tr>
<td>$-$1</td>
<td>$-$1</td>
<td>$-$1</td>
<td>$-$1</td>
<td>36</td>
<td>15,730,582</td>
</tr>
<tr>
<td>$+$1</td>
<td>$-$1</td>
<td>$+$1</td>
<td>$+$1</td>
<td>75</td>
<td>31,481,894</td>
</tr>
<tr>
<td>$+$1</td>
<td>$-$1</td>
<td>$-$1</td>
<td>$+$1</td>
<td>108</td>
<td>31,525,574</td>
</tr>
</tbody>
</table>

Here the inner loop is simply searching for the lowest bit position $b - 1$, for which a set (1) bit of $a$ can be moved one step to the left (i.e., to a position $b$ where the bit currently is 0). After such a position has been found, all the previously set bits below this position (the number of which is kept track of with the counter by $c$) are moved to the lowest positions ($0,\ldots,c-1$). This bit evolution corresponds exactly to how the digits of a base-2 odometer advance from right to left. The contents of checkstate depend on what symmetries are used, but it would be very similar to the implementations we discussed for the 1D case, apart from the fact that the translations [defined in (205)] are more complicated than just cyclic bit permutations and have to be implemented by hand.

The normalization constant, needed when constructing the hamiltonian matrix elements, should be delivered by checkstate (if a representative has passed the tests). The simplest way to compute the normalization (instead of using formal expressions as we did in the 1D case) is again just to carry out all the symmetry operations of the representative state and add up the sum $F_a$ of factors (the complex momentum phases as well as the plus or minus signs from the reflection quantum numbers) in (213) for each symmetry operation bringing the representative onto itself. That number, along with the number $D_a$ of non-equivalent transformations of $|a\rangle$, gives the normalization $N_a = D_a|F_a|^2$ (and again $N_a = 0$ if the representative is incompatible with the quantum numbers). We do not discuss further details of how to implement the basis generation and the construction of the hamiltonian matrix, as these tasks are straight-forward generalizations of the one-dimensional implementations discussed in Secs. 4.1.3 and 4.1.4. Spin-inversion symmetry in the $m_z = 0$ sector can be implemented as discussed in Sec. 4.1.5.

Example of block sizes. Table 3 lists block sizes for the square-lattice systems of interest in Lanczos calculations, for one of the special momenta, $k = (0,0)$, where the largest number of symmetries can be exploited. The ground state of the Heisenberg model is in this block (in the sub-block with all other quantum numbers equal to 1). For $L = 6$ even the smallest blocks have more than 15 million states, and the largest blocks...
(for general \( k \)) are about eight times larger. Blocks of this size can still be handled in Lanczos calculations (the smaller one rather easily on a standard workstation).

To test the basis construction, it is useful to check the sum of the sub-block sizes, which should equal the total number of states in the block. For \( L = 4 \) there are 822 states with \( k = (0,0) \), which equals the sum of all the reflection-block sizes in Table 3.

4.4.2. The Néel state and its quantum rotor excitations

We now illustrate 2D Lanczos calculations with results for the Heisenberg model. Although the small lattices accessible with this method are not sufficient for quantitatively accurate extrapolations to the thermodynamic limit, the calculations do illustrate some important aspects of systems with Néel order (beyond what we discussed in the framework of spin-wave theory in Sec. 2.1). We introduce the quantum rotor mapping of the low-energy states of finite systems, and based on these discuss the magnetic susceptibility. We also calculate the sublattice magnetization.

Two-spin model of quantum-rotor states. If there is antiferromagnetic order, the spins on sublattice A are predominantly oriented in the same direction, and the ones on sublattice B are predominantly in the direction opposite to those on A. If the number of spins \( N \) is finite and the symmetry is not broken, the over-all direction, defined, e.g., by the sublattice A spins, is not fixed, however. This situation can be captured by considering the sum of the spins on the individual sublattices A and B [169],

\[
S_A = \sum_{i \in A} S_i, \quad S_B = \sum_{i \in B} S_i,
\]

as two fixed-length spins \( S_A = S_B = N/2 \) (more precisely we would write, \( S_A = S_B = m_s N \) for large \( N \), but the constant is irrelevant), as illustrated in Fig. 48. The two large spins are assumed to be antiferromagnetically coupled to each other in the simplest possible rotationally invariant way, which is through an effective Heisenberg interaction;

\[
H_{AB} = J_{AB} S_A \cdot S_B = \frac{1}{2} (S^2 - S_A^2 - S_B^2),
\]

where \( S = S_A + S_B \) is the total spin. Here \( S_A^2 \) and \( S_B^2 \) are just constants proportional to \( N^2 \), which can be neglected when we discuss excitation energies. However, these constants imply that the coupling constant \( J_{AB} \) should be \( \propto 1/N \), in order for the total ground state energy to be \( \propto N \). The ground state of (218) has total spin \( S = 0 \) and excitations with \( S = 1, 2, \ldots \) at energies \( J_{AB} S(S+1)/2 \) above the ground state.

We define a new \( N \)-independent coupling \( J_{\text{eff}} = NJ_{AB}/2 \) and write the energies as

\[
\Delta_S = J_{\text{eff}} \frac{S(S+1)}{N}.
\]

These excitations are referred to as the tower of quantum rotor states. The states with \( S \ll \sqrt{N} \) become degenerate as \( N \to \infty \), and combinations of them can then be formed which are ground states with fixed direction of the Néel vector (in analogy with the infinite number of momentum states required to localize a particle in quantum mechanics),
FIGURE 48. Effective description of the rotationally invariant Néel vector $\mathbf{m}$, in terms of two large spins, $S_A, S_B$, corresponding to the sum of the spins on the two sublattices. There is an effective antiferromagnetic coupling between these spins, leading to a singlet ground state and a “tower” of quantum rotor excitations of total spin $S = 1, 2, \ldots$ at energies $\Delta S \sim S(S + 1)/N$ above the ground state.

thus allowing for the symmetry breaking that is the starting point for spin-wave theory. In the thermodynamic limit, the direction of the ordering vector is fixed (as the time scale associated with its rotations diverges [169]), and the quantum rotor-states are then in practice not accessed. They are neglected in standard spin-wave calculations (discussed in Sec. 2.1) from the outset because the order is by construction locked to the $z$ direction.

One can still access the rotor energies in spin-wave theory, by considering systems in an external magnetic field, tuned to give a ground state with total magnetization $S_z = S$ [170, 171]. The rotor states are of great significance in finite clusters.

The effective coupling $J_{eff}$ in (219) for a given system can be determined if we can relate it to some physical quantity which depends on the rotor excitations. An obvious choice is the uniform magnetic susceptibility, $\chi = d\langle m_z \rangle/dh$. Calculating it for the two-spin model when $T \rightarrow 0$ gives $\chi = 3/J_{eff}$. For the real Heisenberg model on a finite cluster in dimensions $d \geq 2$, $\chi$ should be dominated by the quantum rotor states when $T \ll 1/L$, because the lowest spin wave energy scales as $\propto 1/L$ (while the quantum rotor states scale as $1/L^d$). Thus, we can write the effective quantum rotor tower for a Heisenberg model with Néel ground state as

$$\Delta S = \frac{S(S + 1)}{3\chi N},$$

where $\chi$ should be evaluated in the limit $N \rightarrow \infty$ (first) and $T \rightarrow 0$. Note that $I = (3/2)N\chi$ here plays the role of a moment of inertia, giving an analogy between (220) and the energy spectrum of a rigid rotor in quantum mechanics.

The relation (220) can also be used as a way to compute the susceptibility of a Heisenberg models numerically; by extracting the lowest energies as a function of $S$ (for small $S$, where the quantum-rotor mapping should apply). More precisely, the small-$S$ energies gives an estimate for $\chi$ as the $N \rightarrow \infty, S \rightarrow 0$ limit of the quantity $\chi(S, N)$:

$$\frac{1}{\chi(S, N)} = \frac{3N^2(E_S - E_0)}{S(S + 1)}.$$

Here $E_S$ denotes the lowest energy for total spin $S$. Note that we have to subtract the ground state energy ($S = 0$) because in the two-spin effective model we only computed the excitation energies $\Delta S$ with respect to the ground state energy (and the latter is not given accurately by the two-spin model). One would expect an $S$-independent behavior.
only for $S \ll L$, as the higher rotor states should be influenced by effects not taken into account in the two-spin model.

In analogy with the low-energy 1D quantum numbers discussed in Secs. 4.1.6 and 4.3.1, on the 2D square lattice (where $N = 4n$ for all even $L$) the quantum rotor states correspond to momentum $(\pi, \pi)$ and $(0,0)$ for odd and even $S$, respectively. For even $S$, the lowest states have reflection quantum numbers $p_x = p_y = p_d = 1$ in (214), while for odd $S$ the appropriate quantum numbers are $p_x = p_y = -1, p_d = 1$. The lowest state for given $S$ can be obtained in the magnetization sector $m_z = S$ (and since $m_z \neq 0$ we cannot use the spin-inversion symmetry here). The $(S=0)$ ground state is in the fully symmetric sector; the momentum is $(0,0)$ and $p_x = p_y = p_d = z = 1$.

Lanczos results for $L = 4$ and $L = 6$ are shown in Fig. 49(a). There are clearly large corrections to (220), as there is a significant decrease in $\chi^{-1}(S,N)$ with $S$ and an increase with $N$. For fixed $N$, the difference between $S = 1$ and $S = N/2$ is roughly 10%. Considering the fact that the limit $S \ll L$ cannot really be studied based on the small lattices, deviations of this order are not surprising. The rotor states have been studied using QMC calculations for much larger lattices [171, 33]. The most precise calculation for small $S$ and large $N$ gives $\chi^{-1}(S,N) \to 22.8$ [33].

To understand the deviations from the rotor picture, one can use the analogy of a slightly non-rigid quantum rotor, which seems natural considering that the two-spin model is defined with fixed-length spins, while clearly in the real system the sublattice spins fluctuate (in a way which can depend on the total spin $S$). It may be possible to take these effects into account by adding higher-order terms $(S_A \cdot S_B)^2$, etc., in the interaction (218). Details of this extended two-spin picture have not been worked out, however.

Transverse susceptibility. For an infinite system at $T = 0$ the spin-rotational symmetry is broken. One can then consider transverse and longitudinal components (with respect to the Néel vector), $\chi_\perp$ and $\chi_\parallel$, of the susceptibility. In the two-spin model it is clear that $\chi_\parallel = 0$ and this is also true in the Heisenberg model. In a large system in which the
symmetry is not broken, the components are rotationally averaged, giving $\chi = (2/3)\chi_\perp$. The transverse susceptibility is relevant in practice in real Néel order quasi-2D systems (such as the high-$T_c$ cuprates) at $T > 0$, because of anisotropies and/or 3D couplings due to which the system can have a finite ordering temperature. The energy scales associated with the ordering are small, however, and the $T \to 0$ value of $\chi_\perp = (3/2)\chi$ of an isolated 2D plane (modeled by the Heisenberg antiferromagnet) gives a good estimate of this quantity in the more complicated system.

**Sublattice magnetization.** The first attempt to compute the sublattice magnetization of the 2D Heisenberg model was made using Lanczos calculations by Oitmaa and Betts in 1978 [172]. At that time, the $6 \times 6$ lattice was beyond reach, and extrapolations based on smaller lattices resulted in a value that is too large. Fig. 49 shows the spin correlation function at all possible separations $r$ on $4 \times 4$ and $6 \times 6$ lattices. There is a slow decay with $r$, and it at least appears plausible that the results are approaching a value $\langle m^2 \rangle > 0$ as $r \to \infty$ (and $N \to \infty$). QMC calculations, results of which were already shown in Fig. 5, give the $r \to \infty$ result indicated with a dashed line in Fig. 49. The existence of long-range order in the 2D Heisenberg model was, in fact, under debate until the first reliable QMC simulations were done in the late 1980s [173].

One can attempt to extrapolate Lanczos results to $N \to \infty$ for, e.g., frustrated systems, were other calculations are very challenging [98, 99]. The results must in general be viewed with caution, however, because it is assumed that the behavior for very small lattices are already exhibiting the ultimate asymptotic behavior. Fitting procedures based on “Betts clusters” [174] of different shapes may then be misleading, even when the size dependence is smooth. There are examples where cross-overs occurs at larger distances [82], due to the presence of some finite length-scale in the system, which has to be exceeded by $L$ before asymptotic behavior can be observed.

5. QUANTUM MONTE CARLO SIMULATIONS AND THE STOCHASTIC SERIES EXPANSION METHOD

Feynman’s path integral formulation of quantum statistical mechanics [175] has played a major role in the development of QMC methods. In the case of spin systems and related lattice models, methods based on the path integral in imaginary time are commonly referred to as world line methods [176]. These techniques were originally based on an approximate discretization of imaginary time—the Suzuki–Trotter decomposition of the Boltzmann operator $\exp(-\beta H)$ [177, 178]. Later, exact algorithms operating directly in the imaginary time continuum were developed [179, 32]. The first practically useful QMC method did not, however, use the path integral. Already in the early 1960s, Handscomb developed an approximation-free method for the Heisenberg ferromagnet based on the power-series expansion of $\exp(-\beta H)$ and exactly computable traces of products of permutation operators (in terms of which the Heisenberg exchange can be written) [180, 181]. Although this scheme was also generalized to the Heisenberg antiferromagnet [182, 183, 184] and some other systems [185], world line methods were in general more efficient and dominated the field for a long time. The power-series approach to QMC calculations was revived with the introduction of a more generally...
applicable exact stochastic series expansion (SSE) formulation [186, 187], in which also the traces are sampled (thus circumventing the previous reliance on permutation operator algebra). In addition to improved computational utility and efficiency, this generalization of Handscomb’s approach also shows clearly how closely the discrete power series is related to the path integral in continuous imaginary time. Modern QMC algorithms based on the two formulations are also in the end rather similar [33, 31]. In particular, in both cases the spin configurations for some models (such as the Heisenberg model) can be sampled using highly efficient loop-cluster updates [188, 189, 179, 190], which are generalizations [191] of the classical Swendsen-Wang [118] cluster algorithm to quantum systems. Further generalizations of the loop concept to “worms” [32] and “directed loops” [190, 33] have an even wider applicability (e.g., in the presence of external fields), and have enabled large-scale studies of a broad range of quantum spin and boson models. Some of these can now be studied at a level of detail approaching the state of the art for classical systems.

In the case of frustrated spin systems, QMC methods are in general hampered by “sign problems”, i.e., a non-positive definite path integral or series expansion [192]. While calculations can still be carried out in principle, and there has been some progress in controlling the sign fluctuations at high temperatures [193], the statistical errors due to the mixed signs become uncontrollable at low temperatures. Apart from discussing the origins of the sign problem for frustrated models, we will here only consider unfrustrated (bipartite) systems. Despite constituting just a subset of the systems which we are in principle interested in, such models still exhibit a wealth of interesting physics and continue to provide important insights in cutting-edge research.

In these notes we will discuss implementations and applications only of the SSE approach, which for spin systems is normally more efficient and technically simpler than the continuous-time path integral. It is still useful to understand the relationships between the two schemes. In Sec. 5.1 we therefore first review path integrals in quantum statistical mechanics and world line QMC methods, in both discretized and continuous imaginary time. We introduce the general series expansion formulation of quantum statistical mechanics in Sec. 5.1.3 and investigate its mathematical relationship to the path integral. An efficient implementation of the SSE method for the $S = 1/2$ Heisenberg model is described in Sec. 5.2. Illustrative calculations and results for several 1D and 2D systems are discussed in 5.3; single chains in 5.3.1, ladder systems in 5.3.2, the uniform 2D lattice in 5.3.3, dimerized systems in 5.3.4 (focusing on the quantum phase transition between the Néel state and a non-magnetic state), and J-Q models in 5.3.5 (with examples of both continuous and first-order Néel–VBS transitions).

5.1. Path integral and series expansion formulations of quantum statistical mechanics

The main technical problem in quantum statistical mechanics is how to deal with the Boltzmann operator $\exp(-\beta H)$. As we saw with exact diagonalization methods in the previous section, a direct construction of the corresponding matrix becomes infeasible for systems with more than a few tens of spins. The path integral method offers a way
to transform the trace of this operator (the partition function) into a form that can be sampled using Monte Carlo methods. An alternative is to start from a power-series expansion of the exponential. Here we introduce both these approaches as a foundation for implementing the SSE method for Heisenberg models in Sec. 5.2.

5.1.1. The imaginary-time path integral

The starting point of the path integral formulation is to write the exponential operator at inverse temperature $\beta$ as a product of $L$ operators with $\Delta \tau = \beta / L$ in the exponent;

$$Z = \text{Tr}\{e^{-\beta H}\} = \text{Tr}\left\{\prod_{l=1}^{L} e^{-\Delta \tau H}\right\}. \quad (222)$$

The trace can be expressed as a sum of diagonal matrix elements in any basis. We can also insert a complete set of states between each of the exponentials. The partition function then takes the form of an $L$-dimensional sum of products of matrix elements;

$$Z = \sum_{\alpha_0} \sum_{\alpha_1} \cdots \sum_{\alpha_{L-1}} \langle \alpha_0 | e^{-\Delta \tau H} | \alpha_{L-1} \rangle \cdots \langle \alpha_2 | e^{-\Delta \tau H} | \alpha_1 \rangle \langle \alpha_1 | e^{-\Delta \tau H} | \alpha_0 \rangle. \quad (223)$$

Formally, the exponential operator is equivalent to the Schrödinger time evolution operator $\exp(-iHt)$ (with $\hbar = 1$) at imaginary (Euclidean) time $t = -i\Delta \tau$. One can therefore consider (223) as a mapping of a $d$-dimensional quantum system to an equivalent system in $d+1$ dimensions, where the new dimension is imaginary time.\(^6\) The state index $l$ corresponds to a discrete set of imaginary time points $\tau_l = l\Delta \tau$, with $0 \leq \tau_l \leq \beta$ and periodic time-boundary conditions ($\alpha_L = \alpha_0$) for the states. Often the discrete times are referred to as “time slices”.

From a technical perspective, the purpose of writing $Z$ in the form (223) is that, while the matrix elements of $\exp(-\beta H)$ are difficult to evaluate, we can use some approximation to evaluate the matrix elements of $\exp(-\Delta \tau H)$ when $\Delta \tau$ is small (the number of time slices is large). We can then compute the weights for the different time-periodic “paths” $\alpha_0 \to \alpha_1 \to \cdots \to \alpha_{L-1} \to \alpha_0$ over which the system can evolve in the chosen basis. In QMC calculations, these paths are importance-sampled according to their weights in (223).

\(^6\) Often the equivalent $(d + 1)$-dimensional system is referred to as an equivalent classical system. This terminology is, however, appropriate only in cases where all the path weights are positive, which we have not yet ascertained. It will be true for some classes of systems only, and those are the ones for which QMC calculations can be performed in practice. In some cases, e.g., the transverse-field Ising model, the effective model is an anisotropic version of the classical model in $d + 1$ dimensions. In most cases, the equivalent classical system is not, however, the same kind of model as the original one with just one more dimension—typically the path integral corresponds to some completely different statistical mechanics problem in $d+1$ dimensions, with no apparent resemblance to the original $d$-dimensional system. In some cases, one can, however, SHOW that the quantum system is equivalent, on large length and time scales, to the same classical system in $d + 1$ dimensions, e.g., the 2D quantum Heisenberg antiferromagnet at low temperatures has the same properties as a 3D classical Heisenberg model [5]. This mapping is normally carried out using a basis of coherent spin states, as discussed, e.g., in the book by Auerbach [4].
To discuss the possible space-time paths and their weights, let us first consider a seemingly extreme method of approximating the exponential operator by just its Taylor expansions to linear order in $\Delta_\tau H$, i.e.,

$$Z \approx \sum_{\{\alpha\}} \langle \alpha_0 | 1 - \Delta_\tau H | \alpha_{L-1} \rangle \cdots \langle \alpha_2 | 1 - \Delta_\tau H | \alpha_1 \rangle \langle \alpha_1 | 1 - \Delta_\tau H | \alpha_0 \rangle,$$

(224)

where $\{\alpha\}$ refers collectively to all the states $|\alpha_0\rangle, \ldots, |\alpha_{L-1}\rangle$. Since each factor now has an error of order $\Delta_\tau^2$ and there are $L = \beta / \Delta_\tau$ factors, the relative error in $Z$ at fixed $\beta$ is of the order $\Delta_\tau$. The leading neglected term in each expanded factor also contains $H^2$, and we might therefore suspect that the discretization error should also scale as $N^2$. However, to find exactly how the error scales with $\beta$ and $N$ requires a more careful analysis than just naively counting the neglected terms at the level of $\langle H^2 \rangle \propto N^2$ and $L = \beta / \Delta_\tau$, because the paths contributing to the error are not the same as those contributing to the approximate $Z$ written as (224). Based on arguments discussed later, to achieve an error which is independent of $\beta$ and $N$ we should expect to use at least of the order of $N\beta$ time slices in the linear approximation (224), i.e., $\Delta_\tau \propto 1/N$.

Far better approximations of the time slice operator can be used to reduce the number of slices (to independent of $N$ and proportional to $\beta$, with an error of $\Delta_\tau^2$ or smaller). However, for the purpose of illustrating many basic aspects of the path integral and QMC methods based on them, it is convenient to first use the linear approximation. In fact, as we will see shortly, it is even possible to take the limit $\Delta_\tau \to 0$ within this approximation and formulate QMC algorithms based on the exact continuous time path integral [179, 32, 33]. We therefore discuss the linear version in some more detail, although in practice it is not recommended to implement an actual program using this scheme with $\Delta_\tau > 0$. After understanding the properties of the paths in the linear approximation it will be easy to understand how to take the limit $\Delta_\tau \to 0$ or use a higher-order discrete approximation, such as the Suzuki-Trotter decomposition discussed in Sec. 5.1.2.

**Boson path integral and world lines.** At this stage it is better to continue the discussion with a particular hamiltonian in mind, in order to have a concrete example of the paths and how one might go about sampling them. Consider first a boson system with only kinetic energy. We here work on the lattice, and the purely kinetic-energy hamiltonian can be written as (with an unimportant prefactor set to 1);

$$H = K = - \sum_{\langle i,j \rangle} K_{ij} = - \sum_{\langle i,j \rangle} (a_i^+ a_j + a_j^+ a_i),$$

(225)

where $a_i^+$ and $a_i$ are, respectively, boson creation and destruction operators on the sites $i = 1, \ldots, N$, and $\langle i,j \rangle$ is a pair of nearest-neighbor sites on an arbitrary lattice (in any number of dimensions, although for ease of visualization we will explicitly consider a 1D chain). For simplicity, we will consider hard-core bosons, for which the

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7 See [194] for a review of boson path integrals in continuous space, and [195] for more recent progress on efficient QMC algorithms based on them.
FIGURE 50. Graphical representation of terms in the discrete path integral for a system of three bosons on a chain with 14 sites. There are 20 time slices, labeled by \( l = 0, \ldots, 19 \), which correspond to the states \( |\alpha_l\rangle \) in Eq. (224). The bosons are represented by circles. They are connected to show more clearly the formation of world lines on the space-time lattice. Note the periodic boundary conditions in the time (vertical) direction, i.e., the first and last (\( l = 0 \)) states are the same. Note also that with the linear time slice operator \((1 - \Delta H)\), there cannot be more than one particle hop (diagonal line segment) within a time slice. With periodic spatial boundary conditions, the identical particles can undergo cyclic permutations (winding), as in the configuration to the right.

Site occupation numbers are restricted to \( n_i = 0, 1 \) (which can be thought of as arising from a very strong on-site repulsion). Hard-core bosons also have a simple relationship to \( S = 1/2 \) quantum spins. Occupied and empty sites correspond directly to \( \uparrow \) and \( \downarrow \) spins, and with no interactions the boson Hamiltonian is equivalent to the XY model, with \( H_{ij} = -S_i^+ S_j^- + S_i^- S_j^+ \). There is no chemical potential in (225)—we use the canonical ensemble with fixed number of bosons \( N_B \) (density \( \rho = N_B/N \) or, in the spin language, magnetization \( m = \rho - 1/2 \)). Later, we will see that it is easy to also incorporate interactions.

We work in the basis of site occupation numbers \( n_i, i = 1, \ldots, N \). Consider a matrix element \( \langle \alpha_{l+1} | 1 + \Delta \tau K_{ij} | \alpha_l \rangle \), which appears in Eq. (224) when we also write each instance of \( H \) as a sum over all the individual hopping terms \( K_{ij} \) (and note the cancellation of the minus signs). There are then two possible relationships between the states \( |\alpha_l\rangle \) and \( |\alpha_{l+1}\rangle \) resulting in non-vanishing matrix elements; either \( |\alpha_{l+1}\rangle = |\alpha_l\rangle \) or \( |\alpha_{l+1}\rangle = K_{ij} |\alpha_l\rangle \). In the latter case, there must initially be a particle at site \( i \), which is moved by \( K_{ij} \) to a previously empty site \( j \), or vice versa. Since these conditions must hold for all consecutive matrix elements, \( l = 0, \ldots, L - 1, 0 \), each particle has to follow a “world line” (a term borrowed from the similar concept of a space-time path in relativity theory), which at each step of the imaginary time propagation either stays at the same spatial position or moves (jumps) by one lattice spacing. Two such world line configurations for a 1D system are illustrated in Fig. 50.

Note again the periodic boundary conditions in the imaginary time direction, which follow from the trace over \( \alpha_0 \) in (224). These boundary conditions apply to the boson states (occupation numbers), but not strictly to the world lines. Since the bosons are identical particles, they can be permuted in the course of their evolution from \( \tau = 0 \) to \( \tau = \beta \) and still fulfill the required time-periodicity. The only possible permutation in a
1D hard-core system with nearest-neighbor hopping is a cyclic permutation involving all the particles on a periodic chain, resulting in a net particle current around the ring. An example of such a “winding” configuration is shown to the right in Fig. 50. Other permutations can take place with soft-core bosons (i.e., with no restriction on the occupation numbers \( n_i \)), and also hard core bosons in two or three dimensions (and even in one dimension if hopping beyond nearest neighbors is included), but only winding results in a net current. The net number of times world lines wrap around the system (the total current divided by the system length) is called the \textit{winding number}. It is a topological quantity, characterizing a non-local property of the configuration. In higher dimensions, there can be winding in each of the spatial directions, with corresponding winding numbers. An interesting aspect of winding in bosons systems is that it (a properly normalized variance of the winding number) corresponds to superfluidity \cite{175, 196}. In spin systems, winding is related to the spin stiffness, or helicity modulus, as we will discuss further below.

We now denote by \( W(\{\alpha\}) \) the weight of a proper world line configuration \( \{\alpha\} \) (i.e., one satisfying the constraints discussed above) in the partition function;

\[
Z = \sum_{\{\alpha\}} W(\{\alpha\}). \tag{226}
\]

The weight has a very simple form in the case considered here: Each kinetic operation \( K_{ij} \) is multiplied by the time step \( \Delta \tau \) in (224), and the matrix elements of the boson operators is always 1 in the hard-core case. The path weight is therefore

\[
W(\{\alpha\}) = \Delta^{n_K} \tag{227}
\]

where \( n_K \) is the total number of kinetic jumps, corresponding to diagonal world line segments in Fig. 50. This weight can be simply generalized to the case of soft-core bosons, where the creation and annihilation operators are associated with factors \( \sqrt{n_i} \) and \( \sqrt{n_i - 1} \). Below we will also consider interactions between the bosons, which lead to a more complicated factor multiplying the kinetic contribution (227).

In a QMC simulation, one makes changes in the world lines in such a way that detailed balance is satisfied with the configurations distributed according to \( W(\{\alpha\}) \). Local updates of world lines are illustrated in Fig. 51. Note that while these local moves are ergodic within a sector of fixed winding number, they cannot change the topological winding number. In higher dimensions, local moves of individual world lines also cannot lead to any of the other permutations that need to be included. Local updates involving two world lines can be used to sample permutations, but are not enough to change the winding numbers associated with periodic boundaries. Algorithms based on simple local updates were historically successful in studies of some systems, but more sophisticated and powerful \textit{loop updates} \cite{31}, and related updating methods in an extended configuration space \cite{32, 33}, have been developed more recently. These are much more efficient in evolving the world lines in an ergodic way (instead of “getting stuck” or spending long times in some restricted part of the contributing configuration space) and can also lead to winding number changes.

In this section we are primarily concerned with the path integral \textit{representation}, and we will not yet address the practical implementation of world line sampling...
FIGURE 51. Local updates of world lines, which change the number of kinetic jumps (diagonal world line segments) by $\pm 2$. In (a) the diagonal segments are directly following each other, whereas in (b) they are separated by one time slice. One can consider moves with arbitrary separation of the jumps.

(which we will do in detail only for the rather similar configurations arising in the series expansion formulation, although we will later in this section return to some more quantitative discussion of world line sampling as well). Next, we consider estimators for expectation values that we may want to compute. This will also give us some deeper insights into the properties of the world lines themselves.

**Expectation values and their world line estimators.** Consider an arbitrary operator $O$ and its thermal expectation value;

$$\langle O \rangle = \frac{1}{Z} \text{Tr} \{ O e^{-\beta H} \}. \quad (228)$$

Proceeding with the numerator as we did for $Z$ in Eq. (223), the exact time-sliced form of the expectation value can be written as

$$\langle O \rangle = \frac{1}{Z} \sum_{\{\alpha\}} \langle \alpha_0 | e^{-\Delta \tau H} | \alpha_{L-1} \rangle \cdots \langle \alpha_2 | e^{-\Delta \tau H} | \alpha_1 \rangle \langle \alpha_1 | e^{-\Delta \tau H} O | \alpha_0 \rangle. \quad (229)$$

We would like to express this expectation value in the form appropriate for Monte Carlo importance sampling;

$$\langle O \rangle = \frac{\sum_{\{\alpha\}} O(\{\alpha\}) W(\{\alpha\})}{\sum_{\{\alpha\}} W(\{\alpha\})}. \quad (230)$$

This is not always possible, however, because the paths contributing to the numerator in (229) may be different from those contributing to $Z$. If $W(\{\alpha\}) = 0$ for a configuration contributing to the expectation value, then no estimator $O(\{\alpha\})$ can be defined in (230). One then has to proceed with the calculation in a different way. We begin by discussing classes of expectation values for which (230) does apply.

**Diagonal operators.** For quantities that are diagonal in the occupation numbers, e.g., a density correlator $\langle n_i n_j \rangle$, the form (230) is trivially valid, because $O(\alpha_0) = O(\alpha_0) | \alpha_0 \rangle$ in (229), where $O(\alpha_0)$ denotes an eigenvalue of $O$. The estimator can then be taken as $O(\{\alpha\}) = O(\alpha_0)$. This form of the estimator for a diagonal operator remains valid regardless of how the time-slice evolution operator $e^{-\Delta \tau H}$ is approximated. Because of the cyclic property of the trace, by which the operator $O$ can be inserted anywhere in the product of time slice operators in (229), one can also average over all time slices and use

$$O(\{\alpha\}) = \frac{1}{L} \sum_{l=0}^{L-1} O(\alpha_l), \quad (231)$$
which normally improves the statistics of a simulation. In practice, to save time without significant loss of statistics, one may only perform a partial summation in (231) over, e.g., every $N$th time slice (normally $L > N$), because eigenvalues $O(\alpha_t)$ and $O(\alpha_{t+1})$ of nearby states differ very little (at most corresponding to one single world line jump). States separated by $\propto N$ slices typically differ significantly and contribute independent statistics, at least to some degree (exactly how much can be statistically quantified in terms of an imaginary-time dependent correlation function $\langle O(\tau)O(0) \rangle$).

The kinetic energy. Expectation values of off-diagonal operators are in general more complicated. The kinetic energy $\langle K \rangle$ is an exception. In addition to being easy to evaluate, it is also a quantity of particular interest for understanding the properties of the path integral. At the level of the linear approximation of $e^{-\Delta \tau H}$ in $Z$, we can approximate $Ke^{-\Delta \tau H}$ in the form (229) with $O = K$ by just $K$, because this is done only at a single time-slice and leads to a relative error of order $\Delta \tau$. This is of the same order as the total error from the linear approximation made at all the other time slices in both (224) and (229). Given a configuration $\{ \alpha \}$ that contributes to $Z$, the estimator for a specific kinetic operator $K_{ij}$ is then

$$K_{ij}(\{ \alpha \}) = \frac{\langle \alpha_1 | K_{ij} | \alpha_0 \rangle}{\langle \alpha_1 | 1 - \Delta \tau K | \alpha_0 \rangle}.$$  \hspace{1cm} (232)

Here the numerator is non-zero only if there is a world line jump between sites $i$ and $j$ at the first time slice. In that case $K_{ij}(\{ \alpha \}) = 1/\Delta \tau = L/\beta$. In all other cases the estimator vanishes. We can again average over all time-slice locations of the operator $K_{ij}$ in (229), which results in

$$\langle K_{ij} \rangle = \frac{\langle n_{ij} \rangle}{\beta},$$  \hspace{1cm} (233)

where $n_{ij}$ denotes the number of kinetic jumps in the world line configuration between sites $i$ and $j$. Thus, the total kinetic energy is given by the average of the total number of kinetic jumps $n_K$; $\langle K \rangle = -\langle n_K \rangle/\beta$.

We could have derived the expression for the kinetic energy in a simpler way, by applying the thermodynamic formula for the internal energy (here just the kinetic energy): $E = \partial \ln(Z)/\partial \beta$, with $Z$ given by Eqs. (226) and (227). The more complicated derivation illustrates explicitly how the form (230) involves matching the configuration spaces of the numerator and denominator. This matching is not possible for generic off-diagonal operators; only for ones that are part of the hamiltonian. Before considering other cases, let us discuss another important aspect of the kinetic energy estimator.

The utility of the expression (233) is not just that it enables us to compute the kinetic energy. It also carries with it a fundamental message about the path integral and the nature of the world lines. It is natural to ask how typical world lines will evolve as we increase the number of time slices. Specifically, how many kinetic jumps can we expect in a typical world line configuration?

At low temperatures the kinetic energy should be almost temperature independent (approaching a constant as $\beta \rightarrow \infty$). It should be proportional to the lattice size $N$. Using Eq. (233), we can therefore deduce the expected size and temperature scaling of the number of kinetic jumps as

$$\langle n_K \rangle = -\langle K \rangle \beta \sim N \beta.$$  \hspace{1cm} (234)
This tells us that there is a typical scale of roughness of the world lines; for large $L$, the typical time separation between jumps is $\sim 1/N$. As mentioned in the beginning of the section, we should expect to use of the order $N\beta$ time slices in a discrete path integral with the linear approximation of the time slice operator to avoid errors growing with $N$ and $\beta$. We now have the explanation for that, because it is clear from (234) that a smaller number of time slices cannot accommodate the number of kinetic jumps necessary for the correct behavior of the kinetic energy (and then indirectly any other quantity).

The relationship between $\langle n_K \rangle$ and $\langle K \rangle$ is also important from another perspective; it indicates that it should be feasible to formulate simulation algorithms directly in the continuum limit—even if we let the number of time slices $L \rightarrow \infty$, the number of kinetic jumps (“events”) in the contributing configurations stays finite. The continuous-time world lines can then be stored in the form of just a single state $|\alpha_0\rangle$ and the $n_K$ events; the times $\tau_i, i = 1, \ldots, n_K$ at which they occur and the direction of each jump. With $\langle n_K \rangle$ scaling as $N\beta$, it should be possible to construct Monte Carlo sampling algorithms with this scaling in system size and temperature of the time and memory requirements.

Examples of world line configurations in continuous time is shown in Fig. 52. The configuration to the left contributes to the partition function, whereas the one to the right does not, because it does not satisfy all time-periodicity constraints. Instead, it contributes to the expectation value of an operator $a_i^+a_j$, the Fourier transform of which is the momentum distribution function $\rho(k) = \langle a_k^+a_k \rangle$. We briefly discuss such off-diagonal quantities next.

**Off-diagonal operators and broken world lines.** If we consider a path $\{\alpha\}$ contributing to $Z$ and proceed to treat the expectation value of a general off-diagonal operator $a_i^+a_j$ in the same way as we did for the kinetic terms in (232), it is clear that we always get zero, unless $i, j$ are nearest neighbors (in which case the operator is part of the kinetic energy). On the other hand, we can construct other paths, which do contribute to $\langle a_i^+a_j \rangle$ but not to $Z$, like the one in Fig. 52(b). If the sites $i, j$ are separated by more than one lattice spacing, the expectation value $\langle a_i^+a_j \rangle$ cannot be written in the standard Monte Carlo sampling form (230), because $W(\{\alpha\}) = 0$ for all $O(\{\alpha\}) \neq 0$. It is still possible to evaluate $\langle a_k^+a_k \rangle$, as well as the corresponding imaginary-time dependent function

![Figure 52](image-url)
\[ \langle a_k^+(\tau) a_k(0) \rangle \] (the single-particle Green’s function) by working in the combined space of the periodic world line configurations contributing to \( Z \) and those with two defects corresponding to the presence of a creation and annihilation operator [32].

The path integral including interactions. It is not difficult to generalize the bosonic path integral discussed above to hamiltonians including potential-energy terms in addition to the kinetic energy \( K \) in (225). Let us denote by \( V \) any interactions that are diagonal in the occupation number basis, e.g., attractive \( (v_{ij} < 0) \) or repulsive \( (v_{ij} > 0) \) terms of the form \( v_{ij} n_i n_j \). With \( H = K + V \), we can decompose \( \exp(-\Delta_\tau H) \) as

\[
e^{-\Delta_\tau H} = e^{-\Delta_\tau K} e^{-\Delta_\tau V} + \mathcal{O}(\Delta_\tau^2),
\]

where the error is due to the commutator \([K, V] \neq 0\). With \( V \) diagonal, we can for each time slice in (223) write

\[
\langle \alpha_{l+1} | e^{-\Delta_\tau H} | \alpha_l \rangle \approx e^{-\Delta_\tau V_l} \langle \alpha_{l+1} | e^{-\Delta_\tau K} | \alpha_l \rangle,
\]

where \( V_l \) denotes the potential energy evaluated at the \( l \)th time slice. The weight (227) of a world line configuration is therefore modified in the presence of interactions as

\[
W(\{ \alpha_l \}) = \Delta_\tau^n \exp \left( -\Delta_\tau \sum_{l=0}^{L-1} V_l \right).
\]

The interaction part of the weight can be easily taken into account in elementary world line moves of the kind shown in Fig. 51, and also in more sophisticated treatments in continuous imaginary time [32].

The continuum limit. We have already discussed the fact that the continuum limit of the path integral can be used directly in QMC algorithms. However, examining the configuration weight, (227) for the purely kinetic hamiltonian or (237) in the presence of interactions, we have an apparent problem; the weight vanishes as \( \Delta_\tau \to 0 \). This must clearly be compensated in some way by the number of configurations increasing, in such a way that the partition function remains finite. In Monte Carlo calculations we do not deal with the partition function directly and only need ratios of configuration weights to compute acceptance probabilities for world-line updates. Looking at such ratios for the simplest kinds of updates, the insertions and removals of opposite kinetic jumps illustrated in Fig. 51, they are also problematic in the continuum limit: Formally the probability for insertions and removals is zero and infinity, respectively. This is not just a problem in the continuum, but also for small \( \Delta_\tau \), where the probability of accepting an insertion would be non-zero but very small.

Proper ways to handle the continuum limit were introduced in the context of “worm” [32] and loop algorithms [179], twenty years after simulations based on discrete path integrals were first introduced [177, 178]. Here we only consider local world line moves for a toy model, as an illustration of how Monte Carlo sampling can in fact rather easily be formulated in the continuum.

Fig. 51 shows two different ways to introduce two events moving a world line to and from a neighboring site. Consider a line segment which is initially straight over \( m \).
FIGURE 53. Local world line moves in continuous time. In (a), two opposite kinetic jumps are inserted at randomly chosen times $\tau_a$ and $\tau_b$ between two existing jumps at $\tau_1$ and $\tau_2$. In (b), two opposite jumps are either removed (←) or the times of the two events are randomly changed to new arbitrary times $\tau'_a$ and $\tau'_b$ between $\tau_1$ and $\tau_2$ (→). The acceptance probabilities are given in Eq. (238).

We also need to consider the removal of two events. In order to simplify the discussion, we here consider just a single boson on two sites. This avoids the complications of having to consider also constraints imposed by world lines on neighboring sites. The world line moves for which we will construct probabilities satisfying detailed balance are illustrated in Fig. 53. The event insertion discussed above is illustrated in (a), where it should be noted that the times $\tau_1$ and $\tau_2$ correspond to two consecutive existing events (and in case there are no events, we take $\tau_1 = 0$ and $\tau_2 = \beta$). We call the times of the two new events $\tau_a$ and $\tau_b$, and these are chosen at random anywhere between $\tau_1$ and $\tau_2$. In the opposite update of removing two events, we consider the total weight of two existing events $\tau_a$ and $\tau_b$ within two surrounding events at $\tau_1$ and $\tau_2$. Then we can again compute the total weight of the existing events, not only at the fixed current times $\tau_a$ and $\tau_b$ but at any times $\tau'_a$ and $\tau'_b$ within $[\tau_1, \tau_2]$. That relative weight is again $(\tau_2 - \tau_1)^2/2$, versus 1 for the single configuration with those two events absent.

Thus, once $\tau_1$ and $\tau_2$ have been identified by inspecting the current configuration (which can be stored in the form of a list of events), the acceptance probabilities for two-event insertion and removal are

$$P_{\text{insert}} = \frac{(\tau_2 - \tau_1)^2/2}{1 + (\tau_2 - \tau_1)^2/2}, \quad P_{\text{remove}} = \frac{1}{1 + (\tau_2 - \tau_1)^2/2}. \quad (238)$$
For an accepted insertion, we generate $\tau_a$ and $\tau_b$ at random, while in a rejected removal we also generate new times $\tau'_a$ and $\tau'_b$ for the existing events.

In a larger system, with more than one boson, the above scheme would have to be modified to take into account the constraints of other world lines when a chosen world line is moved. The times $\tau_1$ and $\tau_2$ should then reflect those constraints. In the presence of interactions, the acceptance probabilities would be modified to take into account the potential-energy factor in (237). In practice, one would not use these local updates, however, as loop and worm updates [32, 179, 33] are not much more complicated to implement, but much more efficient. The purpose of the discussion here was to show how the apparent problems of the continuum limit can be overcome in principle.

### 5.1.2. The Suzuki-Trotter decomposition

While modern world line QMC algorithms for lattice models are normally based on the path integral in the continuum limit, historically time-discretized variants based on a Suzuki-Trotter approximation of the time-slice operator $\exp(-\Delta \tau H)$ were developed first. The discrete approach is still sometimes used and is the most practical option in some cases [197]. We here discuss the main features of Suzuki-Trotter based methods and how they are applied to boson and spin systems.

One example of a Suzuki-Trotter (or split operator) approximation [177, 198] was already written down in Eq. (235). This is just one among many possible decompositions of an exponential of two (or more) non-commuting operators into a product of exponentials [199]. The most commonly used approximants are, for any pair of operators $A$ and $B$ and a small factor $\Delta$:

$$e^{\Delta(A+B)} = \begin{cases} e^{\Delta A} e^{\Delta B} + O(\Delta^2) \\ e^{\Delta B/2} e^{\Delta A} e^{\Delta B/2} + O(\Delta^3). \end{cases}$$

(239)

Here the errors are also proportional to the commutator $[A, B]$. Using a larger number of judiciously chosen exponentials of functions of $A$ and $B$, the remaining error can be further reduced, in principle to an arbitrary high power of $\Delta$ [200]. High-order approximants are often too complicated to work with in practice, however.

If we are interested in the trace of a product of exponentials, as in the path integral, it is easy to see that the two low-order approximants (239) are actually equivalent, because

$$\text{Tr} \left\{ \prod_{l=1}^{L} e^{\Delta B/2} e^{\Delta A} e^{\Delta B/2} \right\} = \text{Tr} \left\{ \prod_{l=1}^{L} e^{\Delta A} e^{\Delta B} \right\},$$

(240)

due to the cyclic property of the trace. Thus, although the world line method is often discussed based on the first line of Eq. (235), where the error from each factor is $\propto \Delta^2$, the error is in effect smaller; $\propto \Delta^3$. However, since the number of exponential factors in the path integral is $L = \beta / \Delta \tau$, the total error made in a product such as (240) for fixed $N$ and $\beta$ is $\propto \Delta^2$. In addition, one can show that for sufficiently large $L$, the error also does not grow with $N$ at fixed $\Delta \tau$. In most cases, one can therefore keep $\Delta \tau$ independent of
We then decompose $H$ with this arrangement, all terms in each of the sums $H_{\tau}$ sense of their evolution from some time $N$, with $\tau \geq 0$ and $N - \tau \geq 0$. Considering all time slices and all site pairs $i, i + 1$, we may be able to evaluate $\exp(-\Delta \tau H_A)$ and $\exp(-\Delta \tau H_B)$ exactly. The prototypical example is a one-dimensional system containing only nearest-neighbor kinetic energy and interactions, which we collectively denote $H_{i,i+1}$ for a site pair $i, i + 1$. We then decompose $H = H_A + H_B$ according to

$$H_A = \sum_{\text{odd } i} H_{i,i+1}, \quad H_B = \sum_{\text{even } i} H_{i,i+1}, \quad (241)$$

With this arrangement, all terms in each of the sums $H_A$ and $H_B$ are mutually commuting; $[H_{i,i+1}, H_{i+2m,i+2m+1}] = 0$ for any $i$ and $m$. We can therefore split $\exp(-\Delta \tau H_A)$ and $\exp(-\Delta \tau H_B)$ into two-body operators without making any further errors, e.g.,

$$\exp(-\Delta \tau H_A) = \exp(-\Delta \tau H_{1,2}) \exp(-\Delta \tau H_{3,4}) \cdots \exp(-\Delta \tau H_{N-1,N}). \quad (242)$$

Using products like this in the time-sliced partition function (223), we can insert complete sets of states between all the exponentials. Since each operator now involves only two sites, the matrix elements reduce to the form

$$\langle \beta | \exp(-\Delta \tau H_{i,j}) | \alpha \rangle = \langle n_{\beta_i}, n_{\beta_j} | \exp(-\Delta \tau H_{i,j}) | n_{\alpha_i}, n_{\alpha_j} \rangle, \quad (243)$$

which for hard-core bosons are the elements of a $4 \times 4$ matrix. Since the bond operator $H_{i,j}$ conserves the number of bosons on the two sites involved, this matrix is block-diagonal, with two single-element blocks (for $n_{\alpha_i} = n_{\alpha_j} = n_{\beta_i} = n_{\beta_j}$) and a $2 \times 2$ matrix with both diagonal and off-diagonal elements connecting the states with a single boson on the bond. It is thus easy to evaluate all the matrix elements, also if interactions are included (as long as they do not extend beyond nearest neighbors; otherwise a more complicated decomposition of $H$ has to be used).

Pictorially, each matrix element (243) corresponds to a four-site plaquette with zero, one, or two world line segments going through it, according to the occupation numbers. Considering all time slices and all site pairs $i, i + 1$ in both $\exp(-\Delta \tau H_A)$ and $\exp(-\Delta \tau H_B)$, these plaquettes form a checkerboard pattern, with two adjacent rows corresponding to one time slice, as illustrated in Fig. 54. For obvious reasons, the hamiltonian decomposition (241) is also often called the checkerboard decomposition.

For a hamiltonian conserving the total number of particles, the constraints on allowed world line configurations are similar to those in the linear approximation of the time-slice operator, with a few important modifications. Kinetic jumps are allowed only on the shaded plaquettes in Fig. 54, but there is no further constraint on the number of jumps within a time slice, unlike in the linear time-slicing approximation illustrated in Fig. 50 (where there can be at most one jump in each time slice). The world line configurations still look very similar. They become equivalent when $\Delta \tau \to 0$, in the physically relevant sense of their evolution from some time $\tau$ to some later $\tau'$, with $\tau' - \tau \gg \Delta \tau$.

In a Monte Carlo simulation, the world lines can be updated using simple moves of the kind illustrated in Fig. 51, with the constraint that the diagonal loop segments
FIGURE 54. A 1D world line configuration based on the checkerboard decomposition with the Suzuki-Trotter approximation. Kinetic jumps of the bosons (or flips of a pair of ↑ and ↓ spins) are allowed only across the shaded squares (plaquettes). A time slice of width \( \Delta \tau \) consists of two consecutive rows of plaquettes. The six isolated plaquettes shown to the right correspond to the non-zero matrix elements, which in the case of a spin model with Heisenberg interactions (for world lines and empty sites corresponding to ↑ and ↓ spins, respectively) are given by Eq. (244).

(kinetic jumps) are allowed only on the shaded plaquettes in Fig. 54. More complicated “loop” and “directed loop” updates, in which large segments of several world lines can be moved simultaneously, are used in modern algorithms [31, 191, 33] (which we will discuss in detail below in the context of the stochastic series expansion method).

Application to the Heisenberg model. It is useful to consider a particular example of the path weights in the Suzuki-Trotter approach. Let us compute the plaquette matrix elements for the antiferromagnetic Heisenberg interaction; \( H_{i,i+1} = S_i \cdot S_{i+1} \). In this case the boson occupation numbers in (243) are replaced by spin states ↑ and ↓. We can consider the world lines forming between the ↑ spins (and note that we could also draw world lines for the ↓ spins in pictures such as Fig. 54; they occupy all sites not covered by ↑ world lines and cross those lines at each diagonal segment). The calculation just involves straight-forward algebra and we just list the results for the six allowed (non-zero) matrix elements:

\[
\begin{align*}
\langle \uparrow^i \downarrow^j | e^{-\Delta \tau H_{ij}} | \uparrow^i \downarrow^j \rangle &= + e^{-\Delta \tau / 4} \\
\langle \uparrow^i \downarrow^j | e^{-\Delta \tau H_{ij}} | \uparrow^i \downarrow^j \rangle &= + e^{\Delta \tau / 4} \cosh(\Delta \tau / 2) \\
\langle \downarrow^i \uparrow^j | e^{-\Delta \tau H_{ij}} | \downarrow^i \uparrow^j \rangle &= + e^{\Delta \tau / 4} \sinh(\Delta \tau / 2)
\end{align*}
\]  

(244)

The weight of a world line configuration is a product of these matrix elements, all of which are pictorially represented in the right part of Fig. 54. Note the minus sign in front of the off-diagonal matrix elements. For an allowed world line configuration, all the signs cancel out due to the periodicity constraint on the world lines. This is true also for world line methods applied to bipartite lattices in higher dimensions, but for frustrated systems there is a “sign problem” because of the presence of both negative and positive weights (as we will discuss further in Sec. 5.1.3). In practice, world line methods and similar QMC approaches are therefore useful primarily for studies of bipartite spin systems and bosons models. For a fermion system, permutation of world lines also lead to sign problems, except in one dimension where only global cyclical permutations (winding) are possible (with associated signs that can be avoided by choosing periodic or anti-periodic boundary conditions [176, 187]).
In higher dimensions one can use the Suzuki-Trotter approach with various decompositions analogous to the one discussed above. Local updates such as those we have discussed for 1D systems can be adapted to higher dimensions as well. It is worth reading Refs. [79, 203], which are two pioneering world line studies that firmly established the basic properties of the 2D $S = 1/2$ Heisenberg antiferromagnet.

Although the number of time slices used in the Suzuki-Trotter approach does not have to depend on the system size (because $\propto N$ events can take place within a single time slice), the computational effort of Monte Carlo sampling the world-line configurations scales in the same way as in the continuous time formulation, as $N\beta$. This scaling is achieved in the exact continuum formulation by only storing and manipulating the times of the events, whereas in the Suzuki-Trotter approach one normally works with the full space-time lattice (although one could also in principle only store the events there). For this reason, the continuous-time methods actually normally run faster on the computer, in addition to not being affected by discretization errors. The only reason to use a discrete path integral would be in problems where it is difficult to take the continuum limit in practice, e.g., in effective models including dissipation, which leads to time-dependent interactions among the world lines [197].

### 5.1.3. The series expansion representation

As an alternative to the discrete time-slicing approach or continuum limit of the path integral, one can also construct a configuration space suitable for Monte Carlo sampling by using the Taylor expansion of the Boltzmann operator;

$$e^{-\beta H} = \sum_{n=0}^{\infty} \frac{(-\beta)^n}{n!} H^n.$$  \hspace{1cm} (245)

This approach was pioneered by Handscomb [180, 181], who developed a method for studying the $S = 1/2$ ferromagnet. The Taylor expansion was later considered more generally as a starting point for exact QMC algorithms for a wide range of models [182, 185, 186, 187]. The power-series expansion of the exponential operator is convergent for a finite lattice at finite $\beta$ (which technically is due to the fact that the spectrum of $H$ is bounded). In effect, as we will see below, the series expansion allows for a discrete representation of the imaginary time continuum, thus avoiding approximations but fully retaining the advantages of a finite enumerable configuration space.

Choosing a basis, the partition function can be written as

$$Z = \sum_{n=0}^{\infty} \frac{(-\beta)^n}{n!} \sum_{\{\alpha\}_n} \langle \alpha_0 | H | \alpha_{n-1} \rangle \cdots \langle \alpha_2 | H | \alpha_1 \rangle \langle \alpha_1 | H | \alpha_0 \rangle,$$  \hspace{1cm} (246)

where the subscript on $\{\alpha\}_n$ indicates that there are $n$ states to sum over. This expression can be compared with the linear-order discrete path integral (224). Taking $H$ to be the bosonic kinetic energy in (225), we can clearly draw world line pictures very similar to those in Fig. 50 to represent the contributing terms. The main difference is that the number of “slices”, the expansion power $n$, is varying and for given $n$ there are
particle jumps at each slice. The new “propagation” dimension we have introduced in this representation is different from imaginary time, but it is clear that the label \( p = 0, \ldots, n - 1 \) of the states is closely related to imaginary time in the path integral. We will discuss the exact relationship further below.

In the case of only kinetic energy, all matrix elements in (246) equal \(-1\) for an allowed configuration, the minus signs cancel and the weight is

\[
W(\{\alpha\}_n) = \frac{\beta^n}{n!}.
\]

(247)

In more general cases, this weight will be modified by the product of matrix elements in (246). As in the path integral, the weights are positive definite (or can be made so with some simple tricks) for boson systems and quantum spins without frustration in the off-diagonal terms. We will here proceed under the assumption of positive definiteness and discuss the precise conditions for this further below.

It is useful to derive a general expression for the total internal energy \( U = \langle H \rangle \). For any hamiltonian, we can write it as

\[
E = \frac{1}{Z} \sum_{n=0}^{\infty} \frac{(-\beta)^n}{n!} \sum_{\{\alpha\}_{n+1}} \langle \alpha_0 | H | \alpha_n \rangle \cdots \langle \alpha_2 | H | \alpha_1 \rangle \langle \alpha_1 | H | \alpha_0 \rangle.
\]

(248)

where the presence of an additional matrix element of \( H \) in each term should be noted; for given \( n \), the summation is over \( n + 1 \) states instead of the \( n \) states in the partition function (246). All configurations that contribute in (248) contribute also to the partition function, but the weights differ. We can match the configurations explicitly by writing

\[
E = -\frac{1}{Z} \sum_{n=1}^{\infty} \frac{(-\beta)^n}{n!} \frac{n}{\beta} \sum_{\{\alpha\}_n} \langle \alpha_0 | H | \alpha_n \rangle \cdots \langle \alpha_2 | H | \alpha_1 \rangle \langle \alpha_1 | H | \alpha_0 \rangle.
\]

(249)

We can extend the sum over \( n \) to include also \( n = 0 \), because this term vanishes. The terms in (246) and (249) then match exactly, and \( n/\beta \) can be identified as the energy estimator. Thus, with the configurations sampled according to their weights in the partition function, the energy is simply given by

\[
E = -\frac{\langle n \rangle}{\beta}.
\]

(250)

Writing \( H \) as a sum, \( H = -\sum_i H_i \), and using this for all instances of \( H \) in (246) and (248), we can derive a similar expression for the expectation value of an individual term \( H_i \). The result for this estimator is the average number of times the operator appears in the operator string in the expansion of the partition function;

\[
\langle H_i \rangle = \frac{\langle n_i \rangle}{\beta}.
\]

(251)

In the case of a kinetic-energy term, this is identical to the path-integral expression (233). This shows that there is a one-to-one correspondence between the paths in the two
formulations, if we by path mean just the order in which kinetic events occur, without reference to the times $\tau_i$ of the events in the discrete or continuous path integral. This correspondence holds quite generally. The difference is that in the series expansion, the potential-energy terms are treated in the same way as the kinetic operators (i.e., they are not re-exponentiated, as they are in the path integral). Their presence corresponds to “non-events”, which do not affect the world lines but are associated with diagonal matrix elements that modify the weight of the paths. In the path integral, there is instead the potential-energy factor in Eq. (237), which depends on the actual time-points of the kinetic events.

Eq. (250) shows that although the sum over expansion orders $n$ in (246) extends up to $n = \infty$, it can in practice be truncated at some order $n_{\text{max}} \propto N\beta$. To further motivate why this will only lead to an exponentially small and completely negligible error, one can make use of the estimator for the specific heat, obtained by taking the temperature derivative of (250);

$$C = \langle n^2 \rangle - \langle n \rangle^2 - \langle n \rangle.$$

When the temperature $T \to 0$, $C$ should vanish, and (252) then shows that the variance of the distribution of $n$ equals $\langle n \rangle$. It is therefore clear that the distribution vanishes exponentially beyond some power $n \propto N\beta$. In practice, QMC (stochastic series expansion, SSE) algorithms based on this representation automatically sample the $n$-distribution according to the relative weights of the different $n$ sectors.

By explicitly truncating the Taylor expansion, one can make the relationship between the series and path integral representations even clearer. Truncating at $n = L$, we can formally construct a fixed-size sampling space by augmenting all powers $H^n$ with $n < L$ by $L - n$ unit operators $I$. Allowing for all possible locations of the unit operators in the product of $L$ operators, we can define a modified operator string $S = S_1, S_2, \ldots, S_L$, in which $S_i \in \{H, I\}$, and do a summation over all these sequences. We then have to compensate the weight by the number $\binom{L}{n}$ of equivalent terms, giving

$$Z = \sum_{S} \frac{(-\beta)^n(L-n)!}{L!} \sum_{\{\alpha\}_L} \sum_{\{S_i\}_L} \langle \alpha_0 | S_L | \alpha_{L-1} \rangle \cdots \langle \alpha_2 | S_2 | \alpha_1 \rangle \langle \alpha_1 | S_1 | \alpha_0 \rangle,$$

where $n$ now refers to the number of elements $S_i = H$ in the operator string (and we no longer need an explicit sum over $n$). If we now consider the boson kinetic energy (225) and take the limit $L \to \infty$, the weight reduces to $\langle \beta/L \rangle^n$, which is the same as the path integral weight (227) for $L$ time slices. In this limit, it is clear that the index $p = 0, \ldots, L - 1$ is related to imaginary time according to $\tau = p\beta/L$. The difference is that the series expansion with the full weight, $\beta^n(L-n)!/L!$, is in practice exact for $L \propto \beta N$, whereas in the path integral we have to take the continuum limit to avoid a discretization error. For finite $L$, the series index $p$ does not correspond exactly to imaginary time, but represents a distribution of imaginary times [187]. Because of the close relationship between the propagation index $p$ and imaginary time, it is appropriate to refer to the $p$ space as the “time” dimension also in the series expansion.

The practical advantage of the series expansion is that it provides an exact but discrete representation of the imaginary time continuum. Algorithms based on it are normally easier to implement than continuous-time world lines and can be more efficient computationally. On the other hand, for a hamiltonian with large potential-energy terms, the
number of operators $\langle n \rangle$ in the expansion can be much larger than the number of kinetic events $\langle n_K \rangle$ in the path integral. The latter approach should then be more efficient, although an exact treatment necessitates manipulation of floating-point time variables. For quantum spin systems, there is in general a favorable balance between kinetic (off-diagonal) and potential (diagonal) energies, and the series expansion is then typically preferable. Next, we proceed to develop a QMC algorithm based on it.

5.2. SSE method for the $S = 1/2$ Heisenberg model

We now discuss in detail how to implement the SSE method for the $S = 1/2$ Heisenberg model. Initially we do not have to specify the lattice and consider the Hamiltonian written as a sum of bond operators,

$$H_b = J_b S_{i(b)} \cdot S_{j(b)},$$

(254)

as in (204), with the lattice encoded as a list of sites $[i(b), j(b)]$ connected by the bonds, $b = 1, \ldots, N_b$. An example of a labeling scheme in 2D was given in Fig. 46.

A positive-definite SSE can be constructed for any bipartite lattice, i.e., when all sites $i(b)$ and $j(b)$ belong to sublattice A and B, respectively, in any number of dimensions. This constitutes a large class of interesting and important systems. Initially we will consider just a single antiferromagnetic coupling constant $J_b = J > 0$, but the modifications needed for non-uniform systems are very simple.

5.2.1. Configuration space

It is useful to subdivide the Heisenberg interaction (254) into its diagonal and off-diagonal parts in the standard basis of diagonal $z$ spin components. We then define operators with two indices, $H_{a,b}$, with $a = 1, 2$ referring to diagonal and off-diagonal, respectively, and $b = 1, \ldots, N_b$ is the bond index as before;

$$H_{1,b} = \frac{1}{4} - S_{i(b)}^z S_{j(b)}^z,$$

(255)

$$H_{2,b} = \frac{1}{2} (S_{i(b)}^+ S_{j(b)}^- + S_{i(b)}^- S_{j(b)}^+).$$

(256)

Here we have also introduced a minus sign and a constant in the diagonal operator, so that the full Hamiltonian can be written as

$$H = -J \sum_{b=1}^{N_b} (H_{1,b} - H_{2,b}) + \frac{J N_b}{4}.$$  

(257)

The reason for including the constant in $H_{1,b}$ is to make the series expansion positive-definite, as we will see shortly. The constant $J N_b / 4$ is irrelevant in the algorithm but we will include it when calculating the energy.
**Series expansion of the partition function.** The general form of the partition function in the SSE approach can be written as \((246)\), but here we will initially not write out all the complete sets of states inserted between the operators. We instead focus on the operators, expanding all instances of \(H\) as sums over all the bond operators in \((257)\). The starting point of the SSE algorithm for the Heisenberg hamiltonian is thus

\[
Z = \sum_{\alpha} \sum_{n=0}^{\infty} (-1)^n \frac{\beta^n}{n!} \sum_{S_n} \left\langle \alpha \left| \prod_{p=0}^{n-1} H_{a(p),b(p)} \right| \alpha \right\rangle.
\]  

(258)

Here \(J\) has been absorbed into \(\beta = J/T\), \(S_n\) refers to the products (strings) of the Heisenberg bond operators \((255)\) and \((256)\) originating from \(H^n\),

\[
S_n = [a(0),b(0)], [a(1),b(1)], \ldots, [a(n-1),b(n-1)],
\]

(259)

and \(n_2\) is the total number of off-diagonal operators, i.e., the number of elements with \(b(i) = 2\) in the string. Note that the label \(p\) referring to the position of an operator in the string here takes the values 0, \(\ldots, n - 1\); this labeling will be more convenient than \(p = 1, \ldots, n\) in the program implementation.

When the operator string acts on the state \(|\alpha\rangle\) = \(|S_i^c, \ldots, S_N^c\rangle\) we get a succession of other basis states, with no branching into superpositions of more than one state. We will refer to these as propagated states \(|\alpha(p)\rangle\).

\[
|\alpha(p)\rangle \propto \prod_{i=0}^{p-1} H_{a(i),b(i)} |\alpha\rangle.
\]

(260)

We have not written down the normalization of these states explicitly but will consider all \(|\alpha(p)\rangle\) as properly normalized. These states of course correspond to those in the summations over complete sets in \((246)\), but in practice the SSE approach is framed around the summation over the operator strings in \((258)\), and we do not always have to write out the states explicitly.

With the constant \(1/4\) in \((255)\), all operations on parallel spins destroy the states;

\[
\begin{align*}
H_{1,b} |\uparrow_{i(b)} \uparrow_{j(b)}\rangle & = 0, & H_{2,b} |\uparrow_{i(b)} \uparrow_{j(b)}\rangle & = 0, \\
H_{1,b} |\downarrow_{i(b)} \downarrow_{j(b)}\rangle & = 0, & H_{2,b} |\downarrow_{i(b)} \downarrow_{j(b)}\rangle & = 0.
\end{align*}
\]

(261)

An operator-state configuration \((\alpha,S_n)\) contributing to \(Z\) thus has to involve only operations on anti-parallel spins, and the propagated states \((260)\) are defined under this assumption. The corresponding matrix elements are

\[
\begin{align*}
\langle \uparrow_{i(b)} \downarrow_{j(b)} | H_{1,b} | \uparrow_{i(b)} \uparrow_{j(b)}\rangle & = \frac{1}{2}, & \langle \downarrow_{i(b)} \uparrow_{j(b)} | H_{2,b} | \uparrow_{i(b)} \uparrow_{j(b)}\rangle & = \frac{1}{2}, \\
\langle \downarrow_{i(b)} \uparrow_{j(b)} | H_{1,b} | \downarrow_{i(b)} \uparrow_{j(b)}\rangle & = \frac{1}{2}, & \langle \uparrow_{i(b)} \downarrow_{j(b)} | H_{2,b} | \downarrow_{i(b)} \uparrow_{j(b)}\rangle & = \frac{1}{2}.
\end{align*}
\]

(262)

The fact that all these are equal will be very useful in the sampling algorithm, and this was the reason for including the constant \(1/4\) in the diagonal operators.

In addition to the local constraints of only operations on anti-parallel spins, for the matrix element of the full operator product in \((258)\) to be non-zero, the propagation also has to satisfy the periodicity \(|\alpha(n)\rangle = |\alpha(0)\rangle\), where \(|\alpha(0)\rangle = |\alpha\rangle\).
Although one can formulate Monte Carlo sampling procedures in the space of operator strings of fluctuating length \( n \) [180, 187, 204], it is normally in practice somewhat easier to work within the fixed-size truncated space discussed in the previous section (and mathematically it is also easier to prove detailed balance in such a space).

In actual simulations, the cut-off \( L \) will be determined automatically by the program, as we will discuss below, such that \( L \) safely exceeds the largest \( n \) ever sampled. Note again that the truncation then does not cause any detectable errors and should not be considered as an approximation. Defining the unit operators used for augmenting strings with \( n < L \) as \( H_0, 0 = I \) and including \( [a(p), b(p)] = [0, 0] \) as an allowed element in the index list (259), the partition function can be written as

\[
Z = \sum_{\alpha} \sum_{S_L} (-1)^{n^2} \frac{\beta^n (L-n)!}{L!} \left\langle \alpha \left| \prod_{p=0}^{L-1} H_{a(p),b(p)} \right| \alpha \right\rangle , \tag{263}
\]

where \( n \) refers to the number of non-[0,0] elements in the fixed-length operator string \( S_L \) (and \( n \) is summed implicitly by the sum over \( S_L \)). Since all matrix elements equal 1/2, the weight of an allowed configuration is given by

\[
W(\alpha, S_L) = \left( \frac{\beta}{2} \right)^n \frac{(L-n)!}{L!}. \tag{264}
\]

Here we have left out the sign factor \((-1)^{n^2}\), which, as we will discuss further below, always is positive for a bipartite (unfrustrated) system. Note also that \( L! \) is an irrelevant normalization factor and can be left out as well.

With \( \uparrow \) and \( \downarrow \) spins represented by the presence and absence of a hard-core boson, respectively, we could draw pictures like Fig. 50 to illustrate the rules for the contributing SSE configurations. However, we now also have diagonal operators that do not change the world lines but modify the path weight. In the path integral approach the diagonal operators are all brought together into a common exponential factor, Eq. (237), but there is no simple way to do this in the series expansion. As mentioned above in Sec. 5.1, the SSE approach should not be used when the diagonal energy dominates, but in the case considered here we have \( \langle H_1, b \rangle \approx \langle H_2, b \rangle \) and the diagonal operators do not dominate. As we will see below, the presence of a significant number of diagonal operators is in fact exploited in the SSE algorithm. When discussing the SSE method, it is useful to draw pictures showing explicitly the diagonal and off-diagonal operators. An example is given in Fig. 55, which includes also illustrations of the data structures that we will later use in a computer implementation of the method. The states will be represented by integers \( \sigma(i) = \pm 1 \) corresponding to \( S_z = \pm 1/2 \). For compact storage of the operator string, it is also useful to combine the indices \([a(p), b(p)]\) into a single list of integers \( s(p) = 2b(p) + a(p) - 1 \). The full propagated states do not have to be stored simultaneously. They can be generated as needed from the single stored state \(|\alpha(0)\rangle\) and

---

\(^8\) A case where it is clearly better to work with sequences of fluctuating length \( n \) is when the SSE approach is combined with a multi-canonical ensemble, in which a range of temperatures is sampled and therefore \( n \) can fluctuate over a very wide range [204].
FIGURE 55. An SSE configuration for an 8-spin chain, with all the propagated states shown. Open and solid bars indicate diagonal $H_{1,i}$ and off-diagonal $H_{2,i}$ operators respectively, while no bar between states corresponds to a “fill-in” unit operator $H_{0,0}$. The $\uparrow$ and $\downarrow$ spins of the state $|\alpha\rangle$ are stored as $\sigma(i) = \pm 1$, and the operator string $S_L$ is encoded using even and odd integers for diagonal and off-diagonal operators, respectively, according to $s(p) = 2b(p) + a(p) - 1$.

FIGURE 56. An example of three off-diagonal operations (indicated by bars) bringing all spins on a triangle back to their original states. Each spin flip is associated with a minus sign, resulting in a negative path weight and a “sign problem” (due to cancellations of configurations with different signs) in QMC simulations of this and other frustrated systems.

Frustrated interactions and the “sign problem”. At first sight, it appears that we have a sign problem—a non-positive definite expansion—because of the factor $(-1)^{n_2}$ in (258). Actually, all the terms are positive for a bipartite lattice. This is because an even number $n_2$ of off-diagonal operators are required in every allowed configuration, in order to satisfy the “time” periodicity $|\alpha(L)\rangle = |\alpha(0)\rangle$. We already discussed this in the context of the world line method, where the off-diagonal matrix elements in (244) are negative, but the continuity of the world lines require an even number of these. This is yet another example of the close relationship between the two approaches.

For frustrated systems, the series expansion is not positive-definite (and neither is the path integral, for exactly the same reason). This can be easily demonstrated for a system of three spins on a triangle. As shown in Fig. 56, an allowed configuration can in this case contain three off-diagonal operators, resulting in an over-all minus sign. This is true for any system in which loops with an odd number of sites can be formed between antiferromagnetically interacting spins—this can be used as the definition of frustration.

Positive-definiteness for a bipartite system can also be proved in a different way, by
carrying out a unitary transformation of the spin operators on one of the sublattices, say \( B \), such that \( S^+_j \rightarrow -S^-_j \) and \( S^-_j \rightarrow -S^+_j \) (and no change in the diagonal operators \( S^z_j \)) for \( j \in B \). This does not affect the spectrum of the model (since the commutation relations among all spin operators remain unchanged), but the sign in front of the off-diagonal terms in the hamiltonian (257) changes to \(+\). The factor \((-1)^{n^2}\) in (258) is then absent.

For a frustrated system, no such transformation can remove all the signs. Note that only the off-diagonal part of the interaction causes a sign problem. One can study systems with frustration in the diagonal part, but then the interaction is no longer spin-isotropic. Just neglecting the sign for a frustrated Heisenberg system corresponds to antiferromagnetic diagonal couplings but ferromagnetic off-diagonal couplings. Here we only consider isotropic interactions and restrict the discussion to unfrustrated systems.

For a bipartite system with periodic boundaries, the absence of sign problems holds strictly only if the lattice lengths \( L_\gamma \) are even in all directions \( \gamma = 1, \ldots, d \). Winding configurations, like the one shown for the path integral in Fig. 50(b), are negative if there is an odd number of windings around an odd number of sites. Clearly, in such cases there is also frustration, due to the boundaries. We will therefore consider only systems of even length (in all periodic lattice directions).

**Linked-vertex storage of the configurations.** We have discussed computer storage of the SSE configuration as a state \(|\alpha(0)\rangle\) and an operator string \( S_L \). By acting sequentially with the operators, it is easy to generate all the propagated state \(|\alpha(p)\rangle\) illustrated in Fig. 55. However, during the Monte Carlo sampling we will need to access operators and some properties of the propagated states also in non-sequential order—given an operator and the spins it acts on, we will need to know which operators act on those spins next (when moving up as well as down in the operator sequence). It would be prohibitively time consuming to propagate a single state back and forth to extract this information, and also it would not be practical to store all the propagated states. We therefore use also another kind of data structure, in which the “connectivity” of the operators is explicit and represented as a network in a compact way. This linked vertex structure is illustrated in Fig. 57, using the same SSE configuration as in Fig. 55. Here, in the pictorial representation, the constant state of each spin between operators have been replaced by straight lines, which in a computer program will correspond to links (pointers). The operators are shown along with only the two spin states before and after each operator has acted. We will call these spin-operator objects vertices and refer to the four spins as the legs of the vertices. One stage of the Monte Carlo sampling procedures will involve making changes to the vertices (while their locations are changed at another stage). The links will allow us to quickly move between the vertices and make a series of changes maintaining all the constraints.

The allowed vertices are dictated by the hamiltonian. In the case of the isotropic Heisenberg system considered here there are four of them, depicted in Fig. 58. They of course correspond to the non-zero matrix elements (262), and, again, the constant \( \frac{1}{4} \) in the diagonal operator is the reason why there is no vertex with all four spins equal. Anisotropic interactions or an external magnetic field would necessitate inclusions of additional vertices [190, 33] (and the algorithm discussed here would then also have to be modified). Although the spin states at the four legs uniquely identify the vertices, we
FIGURE 57. Linked vertex storage of the configuration in Fig. 55. In the graphical representation to the left, constant spin states between operators have been replaced by lines (links) connecting the spins just before and after the operator acts. The links can be stored in a list $X(v)$, where the four elements $v = 4p + l$, $l = 0, 1, 2, 3$, correspond to the legs (with the numbering convention shown in Fig. 58) of the vertex at position $p$ in the sequence $S_L$. For two linked legs $v$ and $v'$, $X(v) = v'$ and $X(v') = v$.

FIGURE 58. Allowed vertices for the isotropic $S = 1/2$ Heisenberg model. The numbering $l = 0, 1, 2, 3$ of the vertex legs corresponds to the position $v = 4p + l$, in linked-list storage illustrated in Fig. 57.

will continue to use also the open and solid bars in pictures, to indicate diagonal and off-diagonal vertices, respectively, for added clarity.

For a given position $p$ in the operator sequence $S_L$, the corresponding list element $s(p)$ tells us the operator type (diagonal or off-diagonal) and the bond $b$ on which it acts (as explained in Fig. 55). As will become clear below, along with this information, we only have to store the connectivity of the vertices, not their spin states. The links allowing us to jump between connected vertex legs are stored as a list $X(v)$, as explained in Fig. 57. For clarity of the illustration, the one dimensional list has here been arranged in four columns, with elements labeled $v = 4p + l$, corresponding to each type of leg, $l = 0, 1, 2, 3$, with the labeling specified in Fig. 58. We will later describe an efficient way to construct this linked list, given the operator sequence. For now, it is sufficient to know that for a given operator at location $p$ in the sequence, the position of its $l$:th leg in the linked vertex list is $v = 4p + l$. This leg is linked to another vertex leg with list address $v' = X(v)$. This kind of structure constitutes a doubly-linked (bi-directional) list, with $X[X(v)] = v$, in which we can move both “up” and “down”. From a position $v$ in the list we can extract the corresponding operator location in $S_L$, $p = v/4$ (its integer part)
FIGURE 59. (a) A pair of off-diagonal operators, indicated by solid-line boxes, which can be replaced by diagonal ones if the spins between the two operators are flipped as well. Such an update cannot be done with the operators enclosed by dashed boxes, because of the illegal spin configuration (a vertex with all four legs in the same spin state) that would result at the operator acting between the two boxed vertices on their left spin. However, as shown in (b), this operator pair can be changed if instead the spins on the opposite sides of the operators are flipped. This also forces a change in the stored state $|\alpha\rangle$.

and leg index $l = \text{mod}[v, 4]$. We can move “sideways” to the other leg on the same vertex by changing the leg label $0 \leftrightarrow 1$ or $2 \leftrightarrow 3$, which in both cases can be expressed as a simple rule of an even$\leftrightarrow$odd change in the list address $v$. These movements will allow us to construct closed loops of changed spins and operators.

5.2.2. Monte Carlo sampling procedures

When Monte Carlo sampling the partition function (258), we have to make changes in the operator sequence $S_L$ as well as in the stored state $|\alpha\rangle$. It is clear from Figs. 55 and 57, however, that updates in these two data structures are not necessarily independent. Allowed changes (i.e., ones maintaining all configuration constraints) in operators at locations $p \in \{p_1, \ldots, p_2\}$ lead to corresponding changes in the propagated states in the range $|\alpha(p_1 + 1)\rangle, \ldots, |\alpha(p_2)\rangle$. Fig. 59 illustrates such an update of two operators, where only the types of the operators are changed, diagonal$\leftrightarrow$off-diagonal (which leads to changes in the states similar to the world line update in Fig. 51). As the changed operator at $p_1$ acts on $|\alpha(p_1)\rangle$, the state $|\alpha(p_1 + 1)\rangle$ and subsequent states also change (by two flipped spins in the example), but when the last updated operator at $p_2$ has acted the modifications are “healed” and the resulting state $|\alpha(p_2 + 1)\rangle$ is the same as before the update. Since the propagated states form a cyclically permutable periodic structure, there is nothing special with the stored state $|\alpha\rangle = |\alpha(0)\rangle$ and it can change as well, depending on the locations of the updated operators.

Because of the time $(p)$ periodicity of $|\alpha(p)\rangle$, the way state changes are forced by operator updates is not unique. In the above example, where we assumed $p_2 > p_1$, we could also have acted first on $|\alpha(p_2)\rangle$ with the new operator at $p_2$. Then the states $|\alpha(p_2 + 1)\rangle, \ldots, |\alpha(L - 1)\rangle$ as well as $|\alpha(0)\rangle, \ldots, |\alpha(p_1)\rangle$ would be affected. There are always two ways of changing the states in this way, but only one may be allowed in
any given case because of the constraints, as illustrated in Fig. 59. In most cases, such updates done at random locations would not at all satisfy the constraints, and one has to specifically look for two (or more) operators that can be updated [205].

As seen in Figs. 55 and 57, the state |α⟩ can some times be updated without any changes to the operator sequence—since there is no operator acting on the spin \( \sigma_1 \), its state is arbitrary. Such free spins are very rare at low temperatures, when the average number of operators on each spin is large.

Focusing on the operator sequence, we will sample the number \( n \) of non-unit operators \([a, b]_p \neq [0, 0]_p\), their positions \( p \in \{0, \ldots, L-1\} \) in \( S_L \), bond indices \( b \in \{1, \ldots, N_b\} \), and types induces \( a \in \{1, 2\} \). The general strategy is to let the expansion order \( n \) change only in exchanges of diagonal operators with the fill-in unit operator; \([0, 0]_p \leftrightarrow [1, b]_p \) (where \( n \rightarrow n \pm 1 \)). Then, keeping all lattice locations \( b \) of the operators fixed, we change the type index, \([1, b]_p \leftrightarrow [2, b]_p \) for some set of operators (as in the example with two operators above, but we will do it in a much more efficient way involving an arbitrary number of operators). With these off-diagonal replacements carried out properly, the sampling is ergodic in combination with the diagonal \( n \rightarrow n \pm 1 \) updates. We next consider the details of these updating procedures. We first assume that the expansion cut-off \( L \) has been properly determined and then discuss how this can be ensured in practice.

**Diagonal updates.** Updates of single diagonal operators, illustrated in Fig. 60, can be carried out sequentially at the locations \( p = 0, \ldots, L-1 \) in \( S_L \). There are no constraints involved in such an update in the direction \([1, b]_p \rightarrow [0, 0]_p \) (removal of a diagonal operator), whereas an insertion of a diagonal operator, \([0, 0]_p \rightarrow [1, b]_p \), is allowed only if the spins on bond \( b \) are antiparallel, \( \sigma(i(b)) \neq \sigma(j(b)) \), in the propagated state \(|\alpha(p)\rangle\) on which the operator acts. We therefore have to generate these states (storing only the one currently needed), which is simply done by flipping the two spins \( \sigma(i(b)) \) and \( \sigma(j(b)) \) each time an off-diagonal operator \([2, b]_p \) is encountered (in which case no single-operator update can be carried out at \( p \)).

The weight ratio to use in the Metropolis acceptance probability is easily obtained from (264). Note, however, that we also need to correct for the inherent imbalance of these update attempts—there is only a single unique way of carrying out the update in the direction \([1, b]_p \rightarrow [0, 0]_p \), whereas when changing \([0, 0]_p \rightarrow [1, b]_p \) the bond \( b \) should be generated at random among the \( N_b \) possible locations [and if, for the chosen \( b \), the spins \( \sigma(i(b)) = \sigma(j(b)) \), the update is immediately rejected and we proceed to the next \( p \)]. Therefore, following the criteria for detailed balance discussed in Sec. 3.2, specifically the transition probability (43) written as a product of selection and acceptance probabilities, we have to include the ratio \( N_b \) of the selection probabilities for updates.
involving a specific bond $b$. We then obtain the following acceptance probabilities:

$$P_{\text{accept}}([0,0] \to [1,b]) = \min \left[ \frac{\beta N_b}{2(L-n)}, 1 \right],$$  \hspace{1cm} (265)

$$P_{\text{accept}}([1,b] \to [0,0]) = \min \left[ \frac{2(L-n+1)}{\beta N_b}, 1 \right],$$  \hspace{1cm} (266)

where $n$ is the number of operators before the update (and after $n \to n+1$ or $n \to n-1$).

To prove detailed balance for the diagonal updates, it may seem necessary to carry them out at random positions $p$, not sequentially. Otherwise, after having carried out an update at some position $p$, there is zero probability of carrying out the reverse update as the next step. However, if we consider the whole sequence (sweep) of updates from $p=0$ to $P=L-1$ and follow this by a reverse updating sequence, starting from $P=L-1$ and ending at $p=0$, then detailed balance holds for the sweeps. In practice, one does not have to switch between the two directions of updating the operator sequence.

**Off-diagonal updates.** Updates involving off-diagonal operators clearly have to involve at least two operators in order to maintain the periodicity constraint on the propagated states. The simplest kind of pair update was already discussed and illustrated in Fig. 59. For a 1D system, such updates are ergodic with open boundary conditions, but with periodic boundaries local updates cannot change the topological winding number (which is clear from Fig 50, where no local deformations can affect cyclic permutations). This may still not be a very serious problem in principle, because for any $T > 0$ only the sector with zero winding number contributes when the system size $N \to \infty$ (in practice for some large $N$, larger for lower $T$). If we are interested in ground state properties, we also can obtain the correct result when $T \to 0$ for fixed $N$, because in this limit an effective winding number defined within a finite range $p_1, \ldots, p_{D_p}$, with $D_p \gg N$, of the propagated states $|\alpha(p)\rangle$ (corresponding to a finite range of imaginary times) can still fluctuate freely, irrespective of the constraint of zero global winding number. Physical quantities are then insensitive to the winding number [206]. To converge to the ground state, much lower $T$ has to be used, however.

In addition to their inability to change the winding number in one dimension, in higher dimensions the kind of operator substitutions shown in Fig. 59 cannot lead to any of the permutations of same spins (permutation of particles in the boson language) that should be included (while in one dimension, cyclical permutations of all spins are the only possible permutations in the kind of system we consider here). To remedy this problem, one can construct other kinds of local updates, e.g., involving two operators acting on the corners of a plaquette of $2 \times 2$ sites on a 2D square lattice [205], which can sample among all such permutations within a given sector of winding numbers. Updates changing the topological winding number has to involve lines of sites wrapping around the boundaries, but for lattices of length larger than $\approx 20$ these targeted updates become very unlikely (i.e., it becomes too difficult to satisfy the constraints when a large number of operators are changed simultaneously) [186, 205].

Fortunately, instead of dealing with updates specifically targeting permutations and winding number sectors, there is a very efficient and simple type of *loop update* that accomplishes all these things automatically. This class of updates was initially intro-
FIGURE 61. A linked-vertex SSE configuration with one loop traced out and shown in both of its “orientations”, along with the corresponding operator-index sequences. All spins covered by the loop are flipped, and operators are changed, diagonal ↔ off-diagonal, each time the loop passes by (with no net change of an operator visited twice). Every vertex leg (spin) belongs uniquely to one loop, and spins not acted upon by any operator (here the one at \( i = 1 \)) can also be regarded as forming their own loops.

duced as a generalization of a cluster algorithm for the Ising model to a model where the flipped clusters take the form of loops; the classical six-vertex model [191]. The effective world line system for the \( S = 1/2 \) Heisenberg model constructed using the discrete Suzuki-Trotter decomposition is exactly equivalent to an anisotropic six-vertex model, and the loop update for it was therefore at the same time a generalization of the classical cluster update to a quantum mechanical system. These ideas were subsequently applied also to continuous-time world lines [179] as well as to the off-diagonal updates in the SSE method [190]. The improvements in performance relative to local updates are enormous (as in the classical case, leading to a much reduced dynamic exponent) and brought simulations of quantum spin systems to an entirely new level. Like classical cluster algorithms, the loop updates are in practice limited to certain classes of models, of which the isotropic Heisenberg systems is one. Generalizations of the loop concept to worms [32] and directed loops [33] (both of which can be regarded as loops that are allowed to self-intersect during their construction, unlike the original loop updates where no self-intersection is allowed) are applicable to a wider range of systems.

For the \( S = 1/2 \) model considered here, there is no reason to even discuss local off-diagonal updates in any greater detail, and we will just focus on how to implement the much more powerful loop updates. In the case of the SSE method, the operator string is again the main focus, and the loop update corresponds to constructing a loop of operators (vertices) connected by the links in the linked-list representation.

**Operator-loop updates.** An example of an operator-loop and how it is flipped is shown in Fig. 61. Here “flipping” refers to the spins along the loop (explicitly those on the vertex legs and implicitly in all propagated states covered by the loop) as well as the operators themselves when we map the changes back to the sequence \( S_L \); a diagonal
vertex with two legs attached to a loop changes to off-diagonal, and vice versa. In the case of all four legs of a vertex belonging to the same loop (and example of which is present in the figure), the operator type does not change.

The key aspect of the loop update is that the weight (264) only depends on the number of operators, which does not change when the operator-loop is flipped. Therefore, according to the detailed balance rules, such a flip can always be accepted. It is also clear that loops can be large, and a loop flip can therefore lead to changes in the global spin permutations. The previously discussed pair substitutions correspond to flipping loops of two operators on the same bond. Any operator substitution specifically constructed to sample permutations can also be formulated as an operator-loop, including ones that change a winding number.

Note that the loops are completely deterministic once the operator locations have been specified and each spin in the full space-time configuration belongs uniquely to one loop. The diagonal update is the de facto mechanism by which the loops are changed, and the purpose of the loop update is just to identify and flip some of the loops. It is then best to construct all the loops and to flip each of them with probability 1/2 (which also maintains detailed balance), instead of constructing loops at random and always flipping them (in which case some loops would be constructed more than once and flipped unnecessarily). Free spins, on which there are no vertices (such as the first spin in Fig. 61), can also be considered as loops, since flipping such a spin implicitly flips a whole line of spins in the full configuration of time-periodic propagated state.

5.2.3. Computer implementation

A convenient definition of a Monte Carlo sweep in the SSE method is a full sequence of diagonal updates, followed by construction (and flip with probability 1/2) of all loops. From the loop illustration in Fig. 61, it is clear why the linked-vertex storage $X(\cdot)$ of the SSE configuration is useful; constructing a loop corresponds to moving in this list using very simple rules (following a link or moving laterally to the adjacent spin on the same vertex). On the other hand, in the diagonal update the original storage illustrated in Fig. 55, using a single spin state $\sigma(\cdot)$ and the operator string $s(\cdot)$, is more convenient. The configuration will always be stored in this way, and the linked vertices will be constructed before each set of loop updates. The loop updates are carried out by moving in the linked list $X(\cdot)$, updating the corresponding operators in $s(\cdot)$ at the same time (as indicated in Fig. 57). The stored spins $\sigma(\cdot)$ can also be affected by the loop updates, and this can be taken care of after all loop flips have been carried out. We now discuss pseudocode implementations of all the main steps involved in a Monte Carlo sweep.

**Diagonal update.** We assume that we currently have a valid configuration stored (which in the beginning of the simulation can be a random spin state and an operator sequence with only $[0,0]$ elements), using $\sigma(i) = \pm 1$ for the individual spins and the compact storage of $[a,b]_p$ as single integers $s(p)$, with even and odd numbers $2b$ and $2b + 1$ corresponding to diagonal and off-diagonal operators, respectively (as illustrated in Fig. 55). A sweep of diagonal updates can then be implemented in the following way;
\begin{verbatim}
  do p = 0 to L - 1
    if (s(p) = 0) then
      b = random[1, ..., N_p]; if \( \sigma(i(b)) = \sigma(j(b)) \) skip to next p
      if (random[0 - 1] < P_{insert}(n)) then s(p) = 2b; n = n + 1 endif
    elseif (mod(s(p), 2) = 0) then
      if (random[0 - 1] < P_{remove}(n)) then s(p) = 0; n = n - 1 endif
    else
      b = s(p)/2; \( \sigma(i(b)) = -\sigma(i(b)); \sigma(j(b)) = -\sigma(j(b)) \)
    endif 
  enddo
\end{verbatim}

Here \( P_{insert}(n) \) and \( P_{remove}(n) \) are the acceptance probabilities (265) and (266) for inserting and removing a diagonal operator when the current expansion order is \( n \). These probabilities can be precalculated or evaluated on the fly. One can also just precalculate the \( n \)-independent constant \( \beta N_p/2 \) and accept a removal if \( \text{random}[0 - 1] \times (L - n) < \beta N_p/2 \) and an insertion if \( \text{random}[0 - 1] \times \beta N_p/2 < L - n + 1 \). This avoids the more time consuming divisions required with the full acceptance probabilities (265) and (266), or storage of a large number of the \( n \)-dependent probabilities.

The sites belonging to bond \( b \) can be stored in a list \([i(b), j(b)]\). With the exception of this list, the SSE sampling algorithm is lattice independent. It is also easy to modify the above code for non-uniform (\( b \)-dependent) Heisenberg coupling strengths. The diagonal update is the only stage at which the interactions appear explicitly, and no modifications at all are required in the loop update.

**Construction of the linked vertex list.** We will construct the linked list by traversing the operators \( s(p) \), starting from \( p = 0 \). Recall that for vertex leg number \( l = 0, 1, 2, 3 \) of an operator at \( p \), the corresponding links are stored as \( X(v) \) with \( v = 4p + l \), as illustrated in Figs. 57 and 58. To construct the list, we will make use of two other data structures. The position \( v \) in \( X() \) of the first vertex leg for spin \( i \) will be stored as \( V_{\text{first}}(i) \), while \( V_{\text{last}}(i) \) will be the position of the last leg on \( i \) encountered so far. Note that the first leg on a given spin is always below the operator (before the operator has acted, \( l = 0, 1 \)), whereas the last leg is above the operator (\( l = 2, 3 \)). Initially all \( V_{\text{first}}(i) \) and \( V_{\text{last}}(i) \) are set to \(-1 \). Later, when encountering an operator \( s(p) \) acting on spin \( i \) and \( V_{\text{last}}(i) \geq 0 \), then \( V_{\text{last}}(i) \) is the previous list element corresponding to an operation on \( i \). We then use the corresponding lower leg (\( l = 0 \) or 1, depending on which of the two spins acted on equals \( i \)) of the new operator and set the links between \( v = 4p + l \) and \( V_{\text{last}}(i) \); \( X(v) = V_{\text{last}}(i) \) and \( X(V_{\text{last}}(i)) = v \). On the other hand, if \( V_{\text{last}}(i) = -1 \), then we have found the first operation on \( i \) and set \( V_{\text{first}}(i) = v \). In both cases, we set the element for the last operation on \( i \) as \( V_{\text{last}}(i) = v + 2 \), where the addition of 2 corresponds to the vertex leg on spin \( i \) after the operation (leg index \( l = 2 \) or 3). For each operator, we have to examine both the spins \( i_1 \) and \( i_2 \) on which it acts. In the following pseudocode, the vertex list positions to be filled for the operator at \( p \) are \( v_0 + l \), with \( v_0 = 4p \), where the leg pairs \( l = 0, 2 \) and \( l = 1, 3 \) correspond to spins \( i_1 \) and \( i_2 \), respectively. We use \( v_1 \) and \( v_2 \) to denote the list elements of the last (previous) operation on the two spins. All links, except those across the boundary \( p = L - 1, 0 \), can be constructed according to;
\begin{verbatim}
do p = 0 to L - 1
    if (s(p) = 0) skip to next p
    v0 = 4p; b = s(p)/2; i1 = i(b); i2 = j(b); v1 = V_{last}(i1); v2 = V_{last}(i2)
    if (v1 ≠ -1) then X(v1) = v0; X(v0) = v1 else V_{first}(i1) = v0 endif
    if (v2 ≠ -1) then X(v2) = v0; X(v0) = v2 else V_{first}(i2) = v0 + 1 endif
    V_{last}(i1) = v0 + 2; V_{last}(i2) = v0 + 3
endo

To connect the links across the time boundary we can use the arrays containing the first
and last list positions;

\begin{verbatim}
do i = 1 to N
    f = V_{first}(i)
    if (f ≠ -1) then l = V_{last}(i); X(f) = l; X(l) = f endif
endo
\end{verbatim}

Since there are also positions in $X()$ containing no links, corresponding to fill-in operators $s(p) = 0$, we should have some way to distinguish these. We can set all $X(v)$ to a
negative number, e.g., $X(v) = -1$, before constructing the links. Then any $v$ for which
$X(v) ≥ 0$ can be used as a starting point to trace a loop.

\textit{Implementation of the operator-loop update.} We want to trace all the loops and flip
each of them with probability $1/2$. In order to make sure that no loop is considered
more than once, we should always start a new loop from a position in $X()$ not previously
visited. We then need some flag to indicate whether a position has been visited or not.
Instead of allocating separate storage for such flags, we can actually use the list $X()$ itself—the links stored in it only have to be used once, and as soon as a link at $v$ has been used we can set $X(v)$ to a negative number, to indicate that this position should not be used again. To start a new loop, we can look for the first position $v_0$ for which
$X(v_0) ≥ 0$. Before traversing the loop, we make the random decision of whether or not to
flip it. If we do not flip, the loop tracing still has to be carried out, in order to flag all the
对应的 $X(v)$ as visited. For a flipped loop, we should also change the operators
in $s(\cdot)$ to reflect the change diagonal↔off-diagonal for each visited vertex. We will take
care of possible flips of free spins in the stored state $|α\rangle$ later. The structure of a code
for tracing and flipping the loops is;

\begin{verbatim}
do v0 = 0 to 4L - 1 step 2
    if (X(v0) < 0) skip to next v0
    v = v0
    if (random[0 - 1] < 1/2) then
        traverse the loop; for all $v$ in loop, set $X(v) = -1$
    else
        traverse the loop; for all $v$ in loop, set $X(v) = -2$
        flip the loop (change operator types $s(p = v/4)$ while loop is traversed)
    endif
endo
\end{verbatim}
\end{verbatim}

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Here \( v_0 \) is a tentative starting point for a new loop, and we actually start a new loop only if \( X(v_0) \) is a non-negative number. Note that we only have to consider even starting points (the step 2 on the first line of \{30\}), because of the structure of the vertex list and the loops, i.e., two consecutive (even, odd) elements in \( X(v) \) always correspond to the same loop. To not visit the same loop more than once, we set \( X(v) = -1 \) for loops that are only visited, and \( X(v) = -2 \) for loops that are also flipped. We will need the distinction between flipped and only visited loops later, when updating the stored state \(|\alpha\rangle\) to reflect the flipped loops.

To trace a loop starting at some \( v_0 \), we can use the leg index \( l_0 = \text{mod}[v_0, 4] \) and move to the adjacent leg \( l'_0 \) on the same vertex. With the leg labeling convention in Fig. 58, the rule for the leg adjacent to a leg \( l \) can be summarized as \( l = (0, 1, 2, 3) \rightarrow l' = (1, 0, 3, 2) \). This rule can be very efficiently implemented using a bit-level operation on \( l \), as \( l' \) corresponds to flipping \((0 \leftrightarrow 1)\) the lowest bit of \( l \). Here we just assume that such functionality is available in the programming language used [e.g., in Fortran 90 one can use the exclusive-or operation of the integer \( l \) with 1; \( l' = \text{ieor}(l, 1) \)] and assign \( l'_0 = \text{flipbit}(l_0, 0) \), where 0 is the bit flipped. We actually do not need to extract the leg index itself, but can do the corresponding move to the adjacent leg just by manipulating its location \( v_0 = 4p_0 + l_0 \) in \( X() \). We can use the same bit flip method to find it; \( v'_0 = \text{flipbit}(v_0, 0) \). To move to the next vertex, we use the link; \( v_1 = X(v'_0) \). This completes one step of the loop tracing procedure. We next proceed in the same way to find the location \( v'_1 \) of to the adjacent leg, and from there we move to the leg \( v_2 \) linked to it. This continues until at some step \( k \) we find \( v_k = v_0 \). We have then completed a full loop. In pseudocode form the traversal of a loop and flipping it can be accomplished by (where no subscripts or primes are needed on the visited list locations \( v \));

\[
\begin{align*}
v &= v_0 \\
do & \ \ \ \ \ \ \ {\{31\}} \\
p &= v/4; \quad s(p) = \text{flipbit}(s(p), 0); \quad X(v) = -2 \\
v &= \text{flipbit}(v, 0); \quad X(v) = -2 \\
v &= X(v); \quad \text{if } (v = v_0) \ \text{exit} \\
enddo
\end{align*}
\]

Here the flip of the operator type diagonal↔off-diagonal is also accomplished using the \text{flipbit} procedure, which corresponds exactly to the changing the operator code \( s(p) \) from \( 2b \) (diagonal operator) to \( 2b + 1 \) (off-diagonal operator) or vice versa. In the case in code \{30\} where the loop is not to be flipped, the only difference is that the two statements involving \( p \) and \( s(p) \) in \{31\} are absent, and we mark all visited locations as \( X(v) = 1 \) instead if \(-2 \) (information which will be used to update the state \(|\alpha\rangle\)).

After all loops have been traced, we also have to update the spins in the stored state \(|\alpha\rangle\). For each site \( i \) which has a flipped loop passing through it, the stored spin \( \sigma(i) \) should be flipped. We should also flip with probability \( 1/2 \) any spin that has no loop connected to it (i.e., spins on which no operator acts). Both these tasks can be taken care of using the information stored in \( V_{\text{first}}(i) \) during the construction of the linked vertex list. If \( v = V_{\text{first}}(i) = -1 \), that implies that there is no operator acting on spin \( i \). Otherwise, we can use the flag \(-1 \) or \(-2 \) now stored in \( X(v) \) to determine whether the loop passing through \( i \) has been flipped [with \( X(v) = -2 \) for flipped loops]. In pseudocode form:
do $i = 1$ to $N$
  $v = V_{\text{first}}(i)$
  if ($v = -1$) then
    if (random[0-1] $< 1/2$) $\sigma(i) = -\sigma(i)$
  else
    if ($X(v) = -2$) $\sigma(i) = -\sigma(i)$
  endif
endo

This completes the loop update, and we can repeat the cycle consisting of: (i) performing a sweep of diagonal updates, (ii) constructing the linked vertex list, and (iii) tracing/flipping all the loops and updating $|\alpha\rangle$ accordingly. These procedures constitute one Monte Carlo sweep in the SSE method.

Equilibration and expansion truncation. We now discuss the expansion cut-off $L$ (when the fixed-length scheme is used). We should make sure that $L$ exceeds the maximum expansion order $n$ that will be sampled, so that the truncation in practice is not an approximation. Since new diagonal operators are inserted through exchanges with the fill-in unit operators, we also want there to be a reasonably large number of these $s(p) = 0$ elements present. But since the memory and CPU time scales with $L$, we should not make $L$ excessively large. One could in principle define an optimal $L$, for which the acceptance rate of the diagonal updates is maximized. In practice, it does not matter much, however, whether $L$ is really optimal, since the off-diagonal updates are anyway more important in determining the autocorrelation times. We will therefore just make sure that some significant fraction of the elements $s(p) = 0$, so that the sampled $n$ never reaches $L$ and that many insertions of operators can be attempted in the diagonal update.

We can aim for $L \approx (1 + a) \langle n \rangle$ with, e.g., $a = 1/3$. Since the relative fluctuations of $n$ are small, $\sim \sqrt{n}$, we can do this by letting $L \to a \times n$ and augmenting $S_L$ with additional $s(p) = 0$ elements after each Monte Carlo sweep when $a \times n > L$ [setting $s(p) = 0$ for $L_{\text{old}} + 1 < p < L_{\text{new}}$ or distributing the zeros randomly. Fig. 62 shows an example of how this procedure works in practice, during equilibration of a $16 \times 16$ system at inverse temperature $\beta = 16$. Initially $L$ was set to $N/2 = 128$. The adjustment of $L$ quickly converges to an acceptable value, and, as shown in the inset, in a subsequent long simulation $n$ never comes close to $L$.

5.2.4. Estimators for physical observables

The autocorrelation time in SSE simulations with loop updates is typically very short; often just a few Monte Carlo sweeps or less [33, 31]. Calculations of physical observables (“measurements”) should therefore normally be performed after every (or every few) Monte Carlo sweep. As discussed in Sec. 3.2, the autocorrelation times do not dictate how often measurements may be done, only with what frequency independent data are generated. Autocorrelations, thus, affect the error bars but have no effect on the correctness of computed expectation values (i.e., it is not wrong to measure correlated

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Evolution of the expansion cut-off $L$ at the initial stage of an equilibration run of a $16 \times 16$ Heisenberg system at $\beta = 16$. The number of operators $n$ in the string after each Monte Carlo sweep is also shown, along with the maximum $n$ reached so far. The final cut-off after 5000 sweeps was $L = 6764$. The inset shows the distribution of $n$ in a subsequent run of $10^7$ MC sweeps.

data, although it may be wasteful if the measurements take a long time to evaluate) provided that the total simulation time is much longer than the autocorrelation time. Data binning should be used to compute error bars reliably, as also discussed in Sec. 3.2. We here discuss several types of observables of interest in SSE calculations. More details and derivations of the expressions can be found in Refs. [186, 187, 207].

Energy and specific heat. We already discussed some observables in Sec. 5.1.3; the internal energy (250) and the related expectation value (251) of an individual operator in the hamiltonian, as well as the specific heat (252). Given that $\langle H \rangle = -\langle n \rangle / \beta$, it may seem surprising that the specific heat $C = (\langle H^2 \rangle - \langle H \rangle^2) / T^2$ is not given just by the fluctuation in $n$, but there is also a term $-\langle n \rangle$. This is because $\langle H^2 \rangle = \langle n(n-1) \rangle / \beta^2$, which can be easily shown using the same procedures leading to $\langle H \rangle = -\langle n \rangle / \beta$ in Sec. 5.1.3. Note that this expression includes the constant $1/4$ subtracted from each bond operator in (256), but that this constant cancels out in the specific heat expression (252). The specific heat in practice becomes difficult to compute reliably (i.e., its statistical error is large) at low temperatures (where it becomes small), because it is the difference of two large numbers ($\sim N^2 \beta^2$).

Diagonal operators. Expectation values of operators diagonal in the $z$-component basis are also easy to evaluate, using averages over the propagated states;

$$\langle O_z \rangle = \frac{1}{L} \sum_{p=0}^{L-1} \langle \alpha(p) | O_z | \alpha(p) \rangle = \frac{1}{L} \sum_{p=0}^{L-1} O_z(p).$$  

(267)
It is not necessary to evaluate this average using all \( p \), as successive propagated states differ by at most two flipped spins. One can instead use a partial summation over \( p \), e.g., \( O_z(0) + O_z(N) + O_z(2N) + \ldots \), to save time. In some cases, however, one may as well just propagate \( O_z(p) \) from \( p = 0 \) to \( p = L - 1 \) by first computing \( O_z(0) \) and then update the value \( O_z(p) \) according to the spin flips occurring when propagating \( |\alpha(p)\rangle \). For example, the fully averaged contribution to the expectation value of the squared staggered magnetization from an SSE configuration can be computed as follows:

\[
m = (1/2) \sum_i \phi_i \sigma(i); \quad m_{s2} = 0
\]

\[
do p = 0 \text{ to } L - 1
\]

\[
\text{if } \left( \text{mod}(s(p), 2) = 1 \right) \text{ then}
\]

\[
b = s(p)/2; \quad \sigma(i(b)) = -\sigma(i(b)); \quad \sigma(j(b)) = -\sigma(j(b))
\]

\[
m = m + 2\phi_i \sigma(i(b))
\]

\[
\text{endif}
\]

\[
\text{if } (s(p) \neq 0) \quad m_{s2} = m_{s2} + m^2
\]

\[
endo
\]

\[
m_{s2} = m_{s2} / (nN^2)
\]

Here \( \phi_i = \pm 1 \) is the staggered phase factor for site \( i \) and \( m \) contains the staggered magnetization, which evolves as the operator string is traversed. The change in \( m \) when two antiparallel spins are flipped can be expressed as \( 2\phi_i \sigma(i(b)) \), because \( \phi_i \sigma(j(b)) = \phi_i \sigma(i(b)) \). The average of \( m \) squared is accumulated as \( m_{s2} \), in this example only using the \( n \) propagated states generated by the original index sequence without the fill-in unit operators [i.e., skipping the steps where the operator \( s(p) = 0 \)]. One can also sum over all \( L \) instances of the states, in which case the result would be divided by \( L \) instead of \( n \) on the last line. As written above, the special case \( n = 0 \) is not treated correctly, but the code is easy to modify by just using \( m_{s2} = m^2/N^2 \) in that case.

The computational effort of the measurement in \{33\} scales as \( \beta N \), i.e., the same as the SSE sampling algorithm. To compute the Fourier transform of the spin correlations (the static structure factor) at some arbitrary momentum [noting that \( m^2 \) corresponds to \( S(\pi) \), with \( \pi \) denoting the staggered wave-vector, e.g., \( \pi = (\pi, \pi) \) in two dimensions], e.g., for use in the correlation-length definition (70) or (73), one can use a similar procedure for the real and imaginary parts of \( m(q) \),

\[
m(q) = \sum_r S_i^z \cos(q \cdot r) + i \sum_r S_i^z \sin(q \cdot r), \quad (268)
\]

with \( \phi \) in \{33\} replaced by the corresponding sine and cosine factors [and explicitly considering the changes from both the flipped spins at \( i(b) \) and \( j(b) \)]. The structure factor is then accumulated according to \( S(q) = S(q) + \text{Re}^2\{m(q)\} + \text{Im}^2\{m(q)\} \).

For computing the full correlation function,

\[
C_z(r_{ij}) = \langle S_i^z S_j^z \rangle, \quad (269)
\]

averaged over all \( i, j \) [or the Fourier transforms for all \( q \), which can be written as \( \langle m(q)m(-q)\rangle \)], it would not be practical to update all these \( \propto N \) different functions after each spin flip as in \{33\}. It is then better to compute all the correlations from scratch in, e.g., every \( N \)-th propagated state. This still leads to a rather expensive scaling \( \propto \beta N^2 \).
of the effort to measure all the correlations—a factor $N$ worse than the sampling. One should then judge whether it is better to compute only a subset of the correlations [e.g., along some lines in the $(x,y)$ plane] or just measure them less frequently, in order for the measurements not to completely dominate the calculation.

**Susceptibilities.** Another important class of observables are generalized susceptibilities, i.e., linear response functions of the form

$$\chi_{AB} = \frac{\partial \langle A(b) \rangle}{\partial b}, \quad (270)$$

where $b$ is the prefactor in a field term $bB$ added to the Hamiltonian and $A$ is the operator whose response to this perturbation we want to compute. For example, we may be interested in the response $\chi_{ij}$ at site $j$ when a magnetic field acts only on site $i$, in which case $A = S_j^z$ and $B = S_i^z$. Such a susceptibility is given by the Kubo formula [186]

$$\chi_{AB} = \int_0^\beta d\tau \langle A(\tau)B(0) \rangle - \beta \langle A \rangle \langle B \rangle, \quad (271)$$

where $A(\tau) = e^{-\tau H} A e^{\tau H}$. If both $A$ and $B$ are diagonal, this Kubo integral can be evaluated in SSE simulations using the generic formula

$$\int_0^\beta d\tau \langle A(\tau)B(0) \rangle = \left\langle \frac{\beta}{n(n+1)} \left( \sum_{p=0}^{n-1} A(p) \right) \left( \sum_{p=0}^{n-1} B(p) \right) \right\rangle + \left\langle \frac{\beta}{n+1} \sum_{p=0}^{n-1} A(p)B(p) \right\rangle. \quad (272)$$

The sums over $A(p)$ and $B(p)$ can be computed using code similar to {33}.

We are often interested in susceptibilities for which $A = B$, e.g., the magnetic response $\chi(q)$ at wave-vector $q$, in which case $A = B = m(q)$, with $m(q)$ the Fourier transform of the spin configuration given by (268). Since the exact value of $\chi(q)$ must be real-valued, one only has to compute the real part of (272). If all local response functions $\chi_{ij} = \chi(r_{ij})$ are computed in a simulation, the Fourier transform $\chi(q)$ can later be evaluated using these. These response functions are directly accessible in NMR experiments; see Ref. [208] for an example.

Note that if $A$ and $B$ commute with the Hamiltonian, then $A(\tau) = A(0) = A$ [and $A(p) = A$ and $B(p) = B$ are independent of $p$ in (272)] and the susceptibility reduces to the classical expression $\chi_{AB} = \beta (\langle AB \rangle - \langle A \rangle \langle B \rangle)$. In practice, for the Heisenberg model the only case where this form applies is the uniform magnetic susceptibility;

$$\chi = \chi(0) = \frac{\beta}{N} \langle M_z^2 \rangle, \quad M_z = \sum_{i=1}^N S_i^z. \quad (273)$$

Here the term $\beta \langle M_z^2 \rangle$ in (271) vanishes, since $H$ does not include a magnetic field.

If the operators $A$ and $B$ in (271) are not diagonal, the susceptibility is more complicated in general. Here we will only discuss an important special case, in which the
estimator actually is very simple. If the operators involved are two terms of the hamiltonian, as defined in Eqs. (255) and (256) and here referred to just as $H_A$ and $H_B$ for any two instances of those (any two diagonal or off-diagonal bond operators), then the susceptibility measurement just involves counting the numbers $N(A)$ and $N(B)$ of those operators in the sampled SSE operator sequences;

$$
\chi_{H_A H_B} = \frac{1}{\beta} \left[ \langle N(A) N(B) \rangle - \delta_{A,B} \langle N(A) \rangle \right].
$$

The most important example of this type is the current susceptibility $\chi_{I_x I_x}$, where the spin current operator $I_x$ (here in the lattice $x$ direction, for definiteness) is defined by

$$
I_x = \sum_{i=1}^{N} \left[ S_i^- (r_i) S_i^+ (r_i + \hat{x}) - S_i^+ (r_i) S_i^- (r_i + \hat{x}) \right],
$$

where we assume, for simplicity, a hamiltonian with only nearest-neighbor interactions. For longer-range interactions, there would be corresponding current terms between the same site pairs as in the hamiltonian. Although (275) is not exactly a sum of the off-diagonal operators (256) used in the SSE sampling, those operators can be written as sums of two parts,

$$
H_{2,b} = H_{b,+} + H_{b,-}, \quad H_{b,+} = S_{j(b)}^+ S_{i(b)}^-, \quad H_{b,-} = S_{j(b)}^- S_{i(b)}^+,
$$

where the site pairs $[i(b), j(b)]$ in each case are assumed to be ordered in such a way that the $+$ and $-$ terms transport one unit of spin in the positive and negative direction $r_j - r_i$ of the bond, respectively. A key point here is that, although the SSE configurations contain the full off-diagonal operators $H_{2,b}$, only the $+$ or $-$ part in (276) contributes in each instance (with the other part destroying the state it acts on). One can therefore use (274) to evaluate local current-current response functions of the form

$$
\Lambda_{b_1 b_2} = \int_{0}^{\beta} d\tau \langle I_{b_1}(\tau) I_{b_2}(0) \rangle,
$$

where $I_b$ here denotes the current operator at bond $b$. Using (274) one obtains

$$
\Lambda_{b_1 b_2} = \langle [N^+(b_1) - N^-(b_1)][N^+(b_2) - N^-(b_2)] \rangle \delta_{b_1 b_2} (N^+(b_1) - N^-(b_1)),
$$

where $N^+(b)$ and $N^-(b)$ denote the number of operators in the SSE operator sequence transporting spin in the positive and negative direction, respectively, across the bond $b$. The meaning of this becomes clear when looking at a graphical representation of an SSE configuration, such as Fig. 55. There one just has to sum the number of off-diagonal events in which an $\uparrow$ spin is moved to the right (the positive direction) minus the number of left-moving events. The term $N^+(b) - N^- (b)$ is zero, unless the winding number in the lattice direction defined by bond $b$ is not zero, which can be seen in the illustration of winding numbers in Fig. (50). It can be noted that the form (278) of the current response is identical in SSE and path integral (world line) methods.
Spin stiffness. We discussed the spin stiffness in the context of the classical XY model in Sec. 3.5. The basic definitions of this quantity at $T = 0$, Eq. (83), and at $T > 0$, Eq. (86), are identical for quantum spin systems (XY or Heisenberg models) as well as bosonic systems more broadly (where the analogous quantity is the superfluid phase stiffness, which is proportional to the superfluid density). Deriving Monte Carlo estimators for SSE or world line methods, one finds that the term $\beta \langle I_x^2 \rangle$ in (97), which is a classical response function, should be replaced by the corresponding quantum mechanical Kubo formula, giving the spin stiffness in the form

$$
\rho_s = \frac{1}{N} \langle H_x \rangle - \frac{1}{N} \beta \int_0^\beta d\tau \langle I_x(\tau)I_x(0) \rangle,
$$

where we assume that the phase twist is imposed in the lattice $x$ direction. The Kubo integral here consists of a sum of bond-current response functions of the form (277), and using the result (278) one finds that the energy term $\langle H_x \rangle$ is exactly canceled by the $\delta$-function terms coming from (278). This leads to a very simple expression for the spin stiffness in terms of the SSE (or world line) configurations:

$$
\rho_s = \frac{1}{\beta N} \langle (N^+_x - N^-_x)^2 \rangle.
$$

(280)

Here $N^+_x$ and $N^-_x$ denote the total number of operators transporting spin in the positive and negative $x$ direction, respectively. For a spatially isotropic system in two or three dimensions, this can of course be averaged over the dimensions, whereas for an anisotropic system the stiffness depends on the direction.

The spin stiffness is often expressed as the fluctuation of the winding number (here in the $x$ direction, with analogous expressions for other directions),

$$
W_x = \frac{1}{L_x} (N^+_x - N^-_x),
$$

(281)
i.e., the size normalized current ($W_x = 0, \pm 1, \pm 2, \ldots$). For a $d$-dimensional system with $N = L^d$ the stiffness is just $\rho_s = \langle W^2_a \rangle / \beta$ for any direction $a$, but for other shapes (e.g., $N = L_x \times L_y$ with $L_x \neq L_y$) the lengths also enter and (280) looks simpler.

In derivations of the spin stiffness, it is normally for simplicity assumed that the spins order in the XY plane (in spin space), whereas in a Heisenberg model the direction of the order parameter in spin space is not restricted to this plane (the symmetry is not broken) in an SSE simulation of a finite system. A simple rotational averaging argument shows that the expression (280) should be multiplied by $3/2$ in order to obtain the correct stiffness of a symmetry-broken state (i.e., when considering the thermodynamic limit). The expression (280) applies also to the quantum XY model [209], where the result should not be multiplied by $3/2$.

A winding number estimator completely analogous to (280) was originally derived in the context of the superfluid phase stiffness of bosons in continuous space [210]. The role of global cyclical permutations of particles in superfluidity dates back to Feynman’s pioneering work on path integrals [175].
5.2.5. Improved estimators

The operator-loop update in the SSE method (as well as loop updates more broadly [31, 191]) is an example of a cluster update. Such non-local updates were first developed for classical Monte Carlo simulations of the Ising model [118]. One aspect of cluster methods is that it is possible to take averages of estimators for physical quantities over all orientations of the clusters, because the configuration weight does not change upon flipping a cluster. This is immediately clear in SSE simulations of $S = 1/2$ Heisenberg models, because the weight (264) only depends on the number of operators $n$ in the sequence, which does not change when a loop is flipped. If the number of clusters (here operator-loops) is $m$, then the total number of equal-weight configurations is $2^m$, and the average over all of these configurations can provide a much less noisy estimator than one depending on just a single configuration. The crucial point here is that, for many important quantities, this average can be computed analytically, and the resulting improved estimator is of a simple form that can be evaluated rapidly in simulations. Here we only discuss the rather simple cases of the static (equal-time) structure factor and the uniform magnetic susceptibility. For improved estimators for some other quantities, see the review article by Evertz [31].

Consider a propagated state $|\alpha(p)\rangle$, e.g., the stored $|\alpha(0)\rangle$. In the linked vertex representation of the SSE configuration, illustrated in Fig. 61, there is a loop passing through each of the spins in this state (with spins without operators acting on them also considered as individual loops). The same loop can go through many spins in $|\alpha(0)\rangle$, and all spins belonging to the same loop form a cluster, in the sense that if the loop is flipped all the spins in the clusters are flipped simultaneously. Note that the loops are objects in space-time, while the clusters discussed here are defined on a cut at fixed time (here propagation index $p$). A cluster can consist of several parts that appear disconnected in space, since such pieces can be connected in the larger space-time volume where the loops exists. An example of clusters on a 2D lattice is shown in Fig. 63.

Since we are dealing with a bipartite lattice, and because the loop structure is such that the spin on a vertical loop segment (referring to pictures such as Fig. 61) changes each time one changes direction when moving along a loop, the spins within a cluster formed at a given state $|\alpha(p)\rangle$ always have a staggered structure. The staggered magnetization $m_s(j)$ of a cluster labeled $j$, with $j = 1, \ldots, C$, where $C$ is the total number of
clusters, is then \( m_s(j) = \pm n_j \), where \( n_j \) is the number of spins in cluster \( j \). For a given configuration, the total staggered magnetization \( M_s = \sum_{j=1}^C m_s(j) \). When averaging the square of this sum over all the different realizations of cluster orientations, the cross terms \( \langle m_s(i)m_s(j) \rangle = 0 \) (for \( i \neq j \)). One is then left with just the \( i = j \) contributions, and the staggered structure factor is simply given by

\[
S(\pi) = \frac{1}{4N} \left\langle \sum_{j=1}^C n_j^2 \right\rangle. \tag{282}
\]

Structure factors at other wave-vectors \( \mathbf{q} \) are only marginally more complicated, demanding in place of the cluster sizes \( n_j \) a summation over each cluster of the phases \( \phi_r \exp(i \mathbf{q} \cdot \mathbf{r}) \), where \( \mathbf{r} \) refers to sites on a given cluster and \( \phi_r = \pm 1 \) is the staggered phase factor, which takes care of the staggered spin structure within the clusters (and the denominator 4 corrects for the fact that the spin values are \( \pm 1/2 \)). One can here also use the fact that the true structure factor must be real-valued for any \( \mathbf{q} \). In principle, equal-time correlation functions such as the structure factor can also be averaged (fully or partially) over the propagation index \( p \), as in Eq. (267). This, however, requires more work for the improved estimator than in code \{33\} for the simple estimator, because it takes some book keeping during the loop update to construct the clusters for several fixed \( p \), and doing so may not always pay off. Without this averaging, however, a simple \( p \)-averaged estimator, such as the one implemented in code \{33\}, may actually give better results at low-temperatures, where the gain due to averaging can be very significant. The case \( \mathbf{q} = 0 \) is special in this regard, because this corresponds to the total squared magnetization, which is a conserved quantity (i.e., independent on the SSE propagation index \( p \)), and no further averaging over \( p \) can then be done to improve the statistics further. The optimal estimator for the uniform susceptibility (273) is therefore

\[
\chi = \frac{\beta}{4N} \left\langle \sum_{j=1}^C \left( \sum_{i=1}^{n_j} \phi_i \right)^2 \right\rangle. \tag{283}
\]

Susceptibilities at other wave-vectors involve the full space-time loop structure, not just the clusters (cut through the loops) formed at a fixed state. For example, the staggered susceptibility is given by the sum of the squares of all the loop sizes [31].

5.2.6. Program verification

QMC programs should always be verified by comparing results for small systems with exact diagonalization data. When correctly implemented, the SSE method should be exact, which means that the deviation of a computed quantity from its true value should be purely statistical, due to the finite number of sampled configurations. We have discussed how to quantify the statistical fluctuations in terms of “error bars” in Sec. 3.2. Deviations beyond the error bars are due either to programming errors or flaws in the random number generator used. While most programming errors would lead to obviously wrong results, there are also possible subtle errors that may only lead to
minute deviations from exact results for small lattices. Such systematical errors may grow with the system size, with potentially grave consequences. The same may be said about random number generators; bias effects due to imperfect random numbers may be very small for small lattices, but can become more significant for larger systems. It is therefore important to make comparisons with exact diagonalization results based on very long SSE runs, to detect possible small deviations. While it is impossible to strictly prove that a program is correct in all respects, agreement to within very small error bars with exact data makes this very likely.

**Test results.** We now discuss some test results for 1D and 2D Heisenberg models. The random number generator used in these calculations (and most other SSE calculations discussed in these lecture notes) was a simple 64-bit linear congruential generator with multiplier 2862933555777941757 [211] and addition of 1013904243.

Fig. 64 shows the susceptibility per spin of a $4 \times 4$ system. For each temperature $10^{10}$ updating sweeps and measurements were carried out, with the data subdivided into 100 bins for computing the statistical errors. For $T = J/10$, this required approximately 100 CPU hours on a mid-range PC workstation. The error bar (defined as one standard deviation of the estimated fluctuation of the average, as discussed in Sec. 3.2) is typically $\approx 10^{-6}$ when the standard estimator for $\chi$ is used, and even smaller with the improved estimator (283). The relative error (the error bar divided by $\chi$) is $\approx 5 \times 10^{-6}$ for $T/J \approx 1$. The deviations of the averages are completely consistent with the size of the error bars. Recall that for correctly computed statistical errors, one should expect about 2/3 of the data points to bracket the true data within one error bar. The gain in precision with the improved estimator can be much larger than in Fig. 64 for larger systems.

While it is essential to confirm the unbiased nature of the SSE calculations on small
lattices, one might still worry about potential problems with the random number generator for larger systems. In one dimension, we can also test SSE calculations against exact Bethe ansatz results for very long chains in the ground state. To approach the ground state in SSE calculations, it is convenient to use inverse temperatures of the form $\beta = 2^m$ and go to sufficiently large $m$ for calculated quantities to become $m$ independent. This approach is illustrated in Fig. 65, which shows the internal energy, $E = \langle H \rangle$, computed using the simple expansion-order estimator (250) for chains of length $N = 1024$ and 4096. Numerically computed Bethe ansatz results for these chain lengths are listed in [212]. The agreement between the calculations is perfect within error bars, with no detectable temperature dependence for the last three or four points for each $N$. This good agreement, to within relative statistical errors as low as $5 \times 10^{-7}$, shows quite convincingly that the calculation is for practical purposes completely unbiased and that ground states of even quite large systems can be studied.

5.3. Applications of SSE to 1D and 2D systems

We next discuss several illustrative results for 1D and 2D Heisenberg models obtained with the SSE method. Results showing clearly the logarithmic corrections to critical behavior in the 1D chain are presented in 5.3.1. The qualitatively different ground states of $n$-chain ladder systems with even and odd $n$ are discussed in 5.3.2. The sublattice magnetization and some other low-energy parameters associated with the Néel state are computed for the standard 2D Heisenberg model in 5.3.3, and in 5.3.4 dimerization is introduced in this model, to drive a quantum phase transition into a plain non-magnetic state. Quantum critical finite-size scaling behaviors of various quantities are analyzed. The more complex case of a Néel–VBS transition is investigated in 5.3.5, using two types of J-Q models that exhibit, respectively, continuous and first-order transitions.
Spin correlations at $T = 0$. In Sec. 4.3.1 we discussed Lanczos results for the spin correlation function of the Heisenberg chain and saw some hints of the expected logarithmic correction to the $\sim 1/r$ critical behavior (Fig. 34). The system sizes accessible with the Lanczos method are not sufficient for studying these scaling corrections quantitatively, however. As we saw above, with the SSE method unbiased studies of the ground state is possible for chains of several thousand spins (with careful checks of the convergence to the $T \to 0$ limit). Fig. 66 shows the spin correlations for $N = 1024$ and 4096 at distances $r = 2^m$, graphed on a log-log scale. To save time, only the correlations at these distances were computed [for a scaling $N \log(N)$ of the time to carry out spatially averaged measurements]. The results for the two system sizes coincide closely for $r$ up to $2^7$, indicating convergence to the infinite size values up to this distance for $N = 1024$ (and therefore up to $r \approx 2^9$ for $N = 4096$, since the convergence behavior should scale approximately linearly with $N$).

The expected form $|C(r)| = A \ln^{1/2}(r/r_0)r^{-1}$ [57, 58, 59] is very well reproduced up to $r = 2^9$ for $N = 4096$. The parameters $A$ and $r_0$ obtained from a fit are listed in the figure caption. If one leaves the exponent $\sigma = 1/2$ of the logarithm as a free parameter to be obtained from the data based on a fit, the exponent indeed comes out close to 0.5, but with a rather large error bar of, roughly, $\pm 0.1$. To really investigate the exponent carefully, one should further increase the chain length (which is possible in principle).

If one did not know about the existence of a log correction and tried to extract the form of the spin correlations on the basis of numerical calculations alone, one might at first sight conclude that the decay is $\propto 1/r^\alpha$ with $\alpha \approx 0.85$, based on the data in Fig. 66. There are, however, small but significant deviations from a pure power-law, which can be detected only if the relative statistical errors are sufficiently small. In the data shown
in the figure, the error bars are too small to discern. While the absolute errors are small, typically $\approx 2 \times 10^{-6}$ for $N = 4096$ (based on approximately $10^6$ Monte Carlo sweeps at $\beta = 2^{14}$ and using an improved estimator), the relative error is much larger, about 0.002 for $r = 2^7$, growing to 0.02 for the longest distance $r = 2^{11}$.

**Low-temperature magnetic susceptibility.** Logarithmic corrections are also important in the Heisenberg chain at $T > 0$. The most prominent example is the uniform magnetic susceptibility, for which a renormalization-group study of the low-energy field-theory predicted the form \[\chi(T) = \frac{1}{2\pi c} + \frac{1}{4\pi c \ln(T_0/T)},\] (284)

where the $T \to 0$ value $\chi(0) = 1/2\pi c$, where $c = J\pi/2$ is the spinon velocity, agrees with the Bethe ansatz solution of the ground state of the Heisenberg chain. The Bethe ansatz can also be extended to $T > 0$. Good agreement with the asymptotic expression (284), with the parameter $T_0 = 7.7$ (adjusted to fit the Bethe ansatz results), was found at low temperatures ($T/J < 0.02$) [7]. Note that the logarithmic correction in (284) implies a very slow convergence to the $T = 0$ limit, and it would therefore be difficult to extrapolate numerical results without knowing about the presence of this kind of correction. A higher-order logarithmic correction to (284) is also known [213], but here we will just consider the leading-order correction.

**Fig. 67** shows SSE results for the susceptibility obtained using chain lengths of the form $N = 2^n$ and temperatures $T/J = 2^{-m}$. Due to the excitation gap in chains with finite $N$ (discussed in Sec. 4.3.1), $\chi$ decays exponentially to zero below a temperature $T \sim 1/N$. The log-lin scale used in the figure makes these finite-size effects very clear and also shows how the results converge rapidly to the thermodynamic limit form once
the finite-size gap becomes smaller than \( T \). The agreement with the form (284), using the value \( T_0 = 7.7 \) determined in Ref. [7], is very good for \( T/J \) below 0.05 (in fact, for some unknown reason, the agreement appears to be even better than in Ref. [7]).

The main utility of the form (284), beyond its role in establishing the correctness of the low-energy field-theory, is that it is valid not only for the simple Heisenberg chain considered here, but for any 1D spin system (at sufficiently low temperatures) which is in the same phase as the Heisenberg chain. This includes the frustrated \( J_1-J_2 \) chain discussed in Sec. 4.3.2, for \( J_2/J_1 \) less than the dimerization point \( (J_2/J_1)_c \approx 0.241 \). Exactly at the dimerization point the leading logarithmic correction should vanish [164] (but there are still other, higher-order corrections), while above the transition point a spin gap opens and \( \chi \to 0 \) even for \( N = \infty \). The asymptotic form (284) should be valid throughout the QLRO\((\pi)\) phase of the chain including longer-range interactions [the model discussed in Sec. 4.3.3], as well as in many other systems. The parameters \( c \) and \( T_0 \) depend on the model parameters and fitting of numerical data provides a way to extract, in particular, the velocity \( c \). As we will see in the next section, ladder systems consisting of an odd number of chains also are in the same critical phase as the single chain, and (284) applies also there.

5.3.2. Ladder systems

A ladder lattice consists of a fixed number \( L_y \) of coupled 1D chains (often referred to as the legs of the ladder) with the chain length \( L_x \) taken to infinity (or, in practice, \( L_x/L_y \) sufficiently large to give results converged to this limit). Strongly-correlated quantum systems in this geometry [9, 214, 215] play an important role as a means of interpolating between one and two dimensions. Many materials exhibit a structure of weakly coupled ladders with small \( L_y \), which motivates studies of \( L_y = 2, 3, \) etc. On a more fundamental level, it is very interesting to see how the special properties of 1D systems evolve with increasing \( L_y \) and approach the 2D limit [216]. Here we will investigate some of the most essential properties of Heisenberg ladders with \( L_y \) up to 6.

The most important aspect of the physics of \( S = 1/2 \) Heisenberg ladders is that they have completely different low-energy properties for even and odd \( L_y \) (the number of sites on each rung of the ladder). For even \( L_y \) there is always a spin gap (which vanishes as \( L_y \to \infty \)), while odd-\( L_y \) systems are gapless and have properties similar to a single 1D chain (below an energy or temperature scale which vanishes as \( L_y \to \infty \)). This difference can be understood roughly based on a simple picture of valence bonds [217], illustrated in Fig. 68. If there is no coupling between the rungs (\( J_x = 0, J_y > 0 \)), the ground state of the 2-leg ladder is just the product of rung singlets—an unique (non-degenerate) state with a gap \( J_y \) to the first excited state (in which one of the singlets is promoted to a triplet). For the 3-leg ladder, on the other hand, the ground state of an individual rung is two-fold degenerate, with \( S_z = \pm 1/2 \). When coupling the rungs (\( J_x > 0 \)), this degeneracy is lifted, but, for any ratio \( J_x/J_y \), there is a remnant of the degeneracy, which can be understood as arising from \( L_x \) interacting \( S = 1/2 \) degrees of freedom. This leads to low-energy properties similar to those of a single \( S = 1/2 \) chain. In the 2-leg ladder, the singlet-triplet excitation gap remains non-zero for any \( J_x/J_y \), and the low-energy excitations...
FIGURE 68. Illustration of the ground states of the 2-leg and 3-leg Heisenberg ladders at inter-rung coupling $J_x = 0$. The vertical bars indicate valence bonds (singlets), and the circles show unpaired $\uparrow$ and $\downarrow$ spins. The ground state of the 2-leg ladder is a unique singlet-product, whereas in the 3-leg case each rung-state is two-fold degenerate, with $S_{\text{rung}} = \pm 1/2$. Each of these states is a symmetric combination of the two states with the unpaired spin on the upper and lower chain. Here a random configuration of the location and $S^z$ of the unpaired spin is shown for each rung.

above this gap form a band of propagating rung triplets [218]. The spin correlations are exponentially decaying with distance, with the correlation length diverging as $L_y \to \infty$.

In the valence bond basis (discussed in Sec. 2.2), the 2-leg ladder with $J_x > 0$ is still dominated by short bonds (one can say that it is an RVB spin liquid, but, due to the constraints of the ladder geometry, it is also appropriate to call it a VBS), whereas the critical state of the 3-leg ladder requires bond probabilities that decay with the bond length as a power-law [217] (and such a state may be called a critical RVB state). The same pictures remain valid also for the ground states of ladders of larger width, but the energy scale associated with ladder behavior vanishes as $L_y \to \infty$, with cross-overs into 2D behavior at higher energies. The relevant energy scale is the spin gap for even $L_y$ and the spin stiffness for odd $L_y$—as we discussed in connection with the KT transition in Sec. 3.5, power-law correlations can be sufficient to sustain an non-zero spin stiffness, and this is the case with the $\sim 1/r$ correlations in the single chain and odd-leg ladders.

Here we will discuss results only for spatially isotropic couplings; $J_x = J_y$. For even $L_y$, the behavior is similar for periodic and open boundary conditions in the $y$ direction (and for $L_y = 2$ periodic boundaries only corresponds to doubling $J_y$), while for odd $L_y$ the gapless nature of the system for all $J_x/J_y$ applies only for open $y$ boundaries. In the “tube” geometry (periodic $y$ boundaries), odd $L_y$ leads to frustration, and the behavior is then much more complex, with several possible ground state phases [219]. In the SSE calculations discussed below, open $y$ boundaries were used in all cases, while the $x$ boundaries were periodic.

Spin correlations. In the preceding section we studied the spin correlations of the single chain and confirmed the presence of a logarithmic correction to the $\propto 1/r$ critical behavior. We now test this behavior for odd $L_y > 1$, and also look at the exponentially decaying correlations for even $L_y$.

With open boundary conditions in the $y$ direction, the correlator $\langle S_i \cdot S_j \rangle$ is not a function just of the separation $r_{ij}$ between the two spins, but depends on the $y$ coordinates of both spins. Here we use the maximal reflection-symmetric (in the $y$-direction) distance between two spins, i.e., for a given spin at $(x,y)$, with $x = 1, \ldots, L_x$ and $y = 1, \ldots, L_y$, the correlation is computed with the spin at $(x + L_x/2, y + L_y - 1)$, with the periodic boundary condition taken into account in the $x$ direction. Averages are then taken over all $(x,y)$. This is just one convenient choice, and it is not important exactly how the separation in the $y$ direction is treated, because the correlations anyway depend only weakly on it when the $x$ separation is large.
Fig. 69 shows the long-distance correlation function versus $x = \frac{L_x}{2}$ for ladders of width $L_y = 1 - 6$. The qualitative difference between even and odd $L_y$ is clear, and in the case of odd $L_x > 1$ the behavior for large $L_x$ is very similar to the $L_x = 1$ system, with deviations from the form $C(x) \propto \frac{1}{x}$ that can be explained by a logarithmic correction (which we will not analyze in more detail here). For even $L_x$, there is an exponential decay for large $x$, but for $L_y = 6$ a different short-distance behavior can already be seen emerging. For very large $L_y$ and $x$ up to $\sim L_y$, one should expect the correlation function to be of the 2D form, for both even and odd $L_y$, i.e., $C(x)$ should approach the value $m_s^2 \approx 0.095$ of the 2D squared staggered magnetization (illustrated with data in Fig. 5).

For $x \gg L_y$, there should be a cross-over into either an exponential fall-off (for even $L_y$) or the $1/x$ form with a logarithmic correction (for odd $L_y$). In Fig. 69 one can see that the correlation at $x = 8$ is almost the same for $L_y = 4 - 6$, but for larger $x$ there is still an increase with $L_y$. To clearly observe 2D behavior followed by cross-overs into either even- or odd-leg ladder asymptotic behavior, much larger $L_y$ would be required.

Susceptibility. The temperature dependence of the susceptibility provides a convenient way to extract the spin gap $\Delta$ for even-$L_y$ ladders, and it is also interesting to investigate the applicability of the expected asymptotic form (284) for odd $L_y$. Fig. 70 shows SSE results for $L_y = 1 - 6$, with $L_x$ sufficiently large for each temperature to converge accurately to the thermodynamic limit. For the lowest temperatures $L_x = 1024$ was used. For fixed $T$, convergence as a function of $L_y$ can be seen for $T/J > 0.2$, where the results for $L_y = 4, 5,$ and 6 are almost the same. The converged curve corresponds to the 2D limit (for which results for large $L \times L$ lattices are shown in Fig. 73). The qualitative difference between ladders of even and odd width is clear at low temperatures, with $\chi$ for even $L_x$ decaying to zero exponentially below an $L_y$ dependent temperature. For odd $L_y$, it can be noted that the $T \to 0$ value decreases with $L_y$. One can argue that the $T \to 0$
susceptibility per rung should be roughly independent of $L_y$ for small $L_y$ [216], and, thus, the susceptibility per spin should scale approximately as $1/L_y$. We will not do any fitting to the asymptotic form (284) here, because it is valid only for very low temperatures for $L_y \geq 3$ (and a higher-order logarithmic correction may be necessary unless extremely low temperatures are used [216]). Note that the $\chi(0) \sim 1/L_y$ form is not inconsistent with the 2D behavior $\chi_{2D}(0) > 0$, because the ladder behavior (for both even and off $L_y$) applies only below some $T^*(L_y)$ which goes to zero when $L_y \to \infty$.

Let us now analyze the $L_y$ dependence of the spin gap $\Delta$ for the even-width ladders. One expects roughly an exponential low-temperature susceptibility, $\chi \sim \exp(-\Delta/T)$, due to the fact that the ground state is a singlet (non-magnetic, with $\langle M_z^2 \rangle = 0$) and the first excited state is a triplet (magnetic, with $\langle M_z^2 \rangle = 2/3$). This behavior is, however, modified by the fact that there is a whole continuum of magnetic states (for $L_x \to \infty$) above $\Delta$. A low-temperature form of the uniform susceptibility of the 2-leg ladder,

$$\chi(T) = \frac{a}{\sqrt{T}} e^{-\Delta/T}, \quad (285)$$

has been obtained by analyzing the limit of weak inter-rung couplings perturbatively [218]. This form can be expected to hold asymptotically for $T \to 0$ for any $J_y/J_z$ and even $L_y$ (for $L_x \to \infty$). The gap can be extracted from numerical $\chi(T)$ data by considering the logarithm of $\sqrt{T} \chi$ in (285), giving

$$-T \ln(\sqrt{T} \chi) = \Delta - T \ln(a). \quad (286)$$

Thus, the intercept of a line fitted to $-T \ln(\sqrt{T} \chi)$ versus $T$ equals the gap $\Delta$. Fig. 71 shows data analyzed in this way. In all cases, a linear behavior obtains at low temperatures, and line fits give the gaps listed in the figure. A field theoretical treatment shows that the gap for a spin-$S$ ladder system should decrease with $L_y$ as $\Delta \sim \exp(-\pi SL_y)$ [220], which is not in good agreement with the results obtained here (and less precise
earlier calculations [216]). Calculations for larger $L_y$ would have to be carried out (which can certainly be done) to investigate this issue in more detail.

It would also be interesting to study the odd-leg ladders for larger $L_y$ and lower temperatures than what has been done until now, to extract the $L_y$ dependence of the parameters $c$ and $T_0$ in (284). An alternative approach is to map the low-energy properties of a ladder onto a single chain with longer-range interactions [221], which arise because the $S = 1/2$ degrees of freedom of the isolated rungs are not completely localized to individual rungs once they are coupled. The localization length increases with $L_y$, and eventually, for $L_y \to \infty$, an effective model with sufficiently long-ranged interactions to produce Néel order (as discussed in Sec. 4.3.3) should obtain. The mapping procedure has not yet been tested for ladders with large $L_y$, however, and one should therefore also study the full ladder systems at lower temperatures and for larger $L_y$.

### 5.3.3. Long-range order in two dimensions

We already discussed the nature of the antiferromagnetically ordered (Néel) ground state of the 2D Heisenberg model in Secs. 2.1 and 4.4.2. Extrapolation of the best currently available QMC results [50] for the squared sublattice magnetization, shown in Fig. 5 gives $m_s = 0.30743(1)$ for the infinite system, where (1) denotes the statistical error in the preceding digit. This result deviates only by about 1% from the linear spin wave result $m_s = 0.3034$. Higher-order spin wave calculations [40, 41, 42] give $m_s \approx 0.3070$. As discussed in Sec. 2.1, this good agreement with the actual value can be traced to the fact that the quantum fluctuations are not that strong, reducing the sublattice magnetization by only about 40% from the classical value. Other important ground state quantities, such as the spin stiffness [222], the transverse magnetic susceptibility, and the spin wave velocity, are in similar good agreement with spin wave theory (but we will...
After discussing some general data fitting issues, we will here compute the energy and the spin stiffness using finite-size extrapolations of $T \rightarrow 0$ converged SSE results (with proper convergence confirmed using checks such as those shown in Fig. 65 for all quantities of interest). In addition to the $L \times L$ lattices normally used, we will also consider rectangular $L_x \times L_y$ lattices with $L_x = 2L_y$. This enables a consistency check of the extrapolations. We also discuss the finite-temperature susceptibility and extrapolate it to zero temperature, using $L \times L$ lattices sufficiently large to completely eliminate finite-effects for the range of temperatures considered. We finally discuss the divergence of the correlation length as $T \rightarrow 0$, which is another important manifestation of the Néel ordered ground state [5].

Extrapolations of the ground state parameters of the 2D Heisenberg model have been presented in a large number of papers previously, e.g., Refs. [223, 224, 205, 225, 226]. For most practical purposes, the precision already achieved is quite sufficient. Since the model is one of the most important prototypical systems in quantum magnetism, it is, however, useful to continue to establish more precise benchmark calculations (which can be useful, e.g., for testing other methods [94]). Some of the results presented below, in particular the ground state energy, represent the most precise calculations to date. In addition to the extrapolated $N = \infty$, $T = 0$ values of the various physical quantities, their finite-size and temperature corrections are also of interest, because they have been predicted in great detail based on field-theoretical methods [227], and numerical test are useful to establish the range of validity of the low-energy theories. The corrections are statistically much noisier and require very long simulations to establish precisely. The discussion here is only intended to give a flavor of what can be done.

**Data fitting issues.** When extrapolating finite-size result to the thermodynamic limit, it is useful to know the expected form of the size corrections based on analytical calculations. Away from a critical point, the size corrections for systems of dimensionality $d > 1$ normally take the form of a polynomial in the inverse system length $1/L$, but some times the leading correction is $\propto 1/L^a$ with $a$ an integer larger than 1. As discussed in Sec. 2.4.1, the leading size correction to the sublattice magnetization of the 2D Heisenberg model can be obtained from the spin wave theory of the Néel state and is $\propto 1/L$. This result is also a more general consequence of the fact that the order parameter is a vector, as discussed in Sec. 3.3. The ground state energy $E/N$ per site has a leading correction $\propto 1/L^3$, and thanks to this high power it is relatively easy to obtain the energy to high precision even based on rather small lattices [205]. The leading correction to the spin stiffness is $\propto 1/L$ [228]. Even if the leading power is not known, one can normally find it empirically based on data, provided that the numerical precision is sufficiently high. When fitting data to a polynomial one may find that the coefficient of the linear term (and possibly higher-order terms) is very small, which then makes it likely that it actually should be exactly zero. It is best to use a fitting program which allows one to specify exactly what powers in $1/L$ should be included.

For a given set of data (here a quantity computed for a set of lattice sizes, but we will use the same technique also for fitting temperature dependent quantities as powers in $T$) one should include high enough powers of $1/L$ for the fit to be statistically sound, i.e., $\chi^2$ per degree of freedom ($\chi^2$/dof) should be close to 1 for a large data set (as discussed
Size dependence of the ground state energy (left) and spin stiffness (right) of 2D Heisenberg lattices with aspect ratio $R = 1$ (where $N = L^2$) and $2$ (where $N = 2L^2$). For $R = 2$, the spin stiffness on a finite lattice is different in the $x$ and $y$ direction. Both values are shown here along with the average. The curves are polynomial fits discussed in the text.

Ground state energy and spin stiffness. The energy per site $E/N$ and the spin stiffness $\rho_s$ were evaluated using the estimators (250) and (280), respectively, with the energy adjusted for the constant $1/4$ subtracted from each bond operator in the SSE simulations. As discussed in Sec. 5.2.4, the value of the spin stiffness extrapolated to the thermodynamic limit should be multiplied by a factor $3/2$, to account for the fact that the spin-rotational symmetry is not broken in the SSE simulations.

Comparing extrapolations for lattices of different shapes is a good way to check for detectable consequences of finite-size corrections beyond those included in the data fits [82, 94]. Fig. 72 shows results for both $N = L \times L$ (aspect ratio $R = 1$) and $N = 2L \times L$ ($R = 2$) lattices, graphed versus $1/\sqrt{N}$ ($= 1/L$ for $R = 1$ and $\propto 1/L$ for $R = 2$). These results were computed using inverse temperatures $\beta$ as high as $32 \times L$ for the largest systems ($L$ up to 40 for $R = 1$ and up to 32 for $R = 2$). For the largest systems several hundred CPU hours were used. The error bars are too small to be visible in the figure. An example of the results produced, for the $32 \times 32$ system $E/N = -0.6695115(8)$ and $\rho_s = 0.12606(5)$, based $5 \times 10^6$ Monte Carlo sweeps. The error bars are of similar magnitude for the other systems as well.
For $R = 1$ using the systems with $L \geq 6$ and a 5th-order polynomial (without the linear and quadratic terms, which are predicted not to be present, as discussed above) gives $E/N = -0.6694421(5)$, while a fit of the same order to the $R = 2$ data with $L \geq 6$ delivers $E/N = -0.6694422(6)$. These results are in perfect statistical agreement with each other, and it is then permissible to use their statistically weighted average, $E/N = -0.6694421(4)$, as a final estimate of the ground state energy. It should be noted that the numerical values depend slightly on what lattices are included in the fit and the order of the polynomial used. Once the fit is statistically sound, these fluctuations should be consistent with the statistical errors.

Turning now to the stiffness, note first that for a finite lattice with aspect ratio $R \neq 1$, there are two stiffness constants, for phase twists imposed in the $x$ and $y$ direction. Both of them are graphed for $R = 2$ in the right panel of Fig. 72, along with the arithmetic average of the two and the $R = 1$ values. It can be seen here that the $R = 1$ stiffness and the average of the $R = 2$ stiffnesses are better behaved for extrapolations than the individual $x$ and $y$ values for $R = 2$, and no fits are therefore included for the latter. Quadratic fits for $R = 1$ and $R = 2$ (using $L \geq 6$ and $L_x \geq 8$ data) gives $\rho_s = 0.12065(4)$ and $\rho_s(\infty) = 0.12070(6)$, respectively. Taking the average of these statistically consistent values, and including the rotational factor $3/2$, gives the final estimate $\rho_s = 0.18100(5)$ for the infinite system.

**Properties at $T > 0$.** Since the Mermin-Wagner theorem rules out magnetic order in the 2D Heisenberg model for any $T > 0$, the correlation length must diverge as $T \rightarrow 0$ for the behavior to be consistent with the ordered ground state. The behavior is similar in the classical 2D Heisenberg model, where the correlation length diverges exponentially. The long-distance correlations and fluctuations of the quantum system can in fact formally be mapped onto a classical system with renormalized couplings [5], and the low-temperature regime in which such a mapping holds is referred to as the renormalized classical regime. Many predictions in this regime have resulted from field-theoretical treatments [5, 227, 84]. Remarkably detailed results have been derived for the temperature dependence of, e.g., the correlation length and the susceptibility. The predicted forms, which are believed to be asymptotically ($T \rightarrow 0$) exact, depend only on the ground state parameters (e.g., the spin stiffness and the spin wave velocity $c$).

Let us first investigate the uniform magnetic susceptibility, which should have the following form for $T \rightarrow 0$ (in an infinite system) [227] :

$$
\chi(T) = \chi(0) \left[ 1 + \frac{T}{2\pi\rho_s} + \left( \frac{T}{2\pi\rho_s} \right)^2 \right].
$$

(287)

The only model dependent parameter here is the ground state spin stiffness, which we determined above. To compare numerical data with this result, we first have to make sure that we can achieve the thermodynamic limit. For fixed temperature, we can study the behavior as a function of the system size, where we here use quadratic lattices with $N = L^2$. The left panel of Fig. 73 shows results for several lattice sizes. For any finite system the susceptibility vanishes as $T \rightarrow 0$ due to the singlet ground state and finite-size gap in the spectrum. As we discussed in Sec. 4.4.2, the lowest excitations are the quantum rotor states, with gaps scaling as $1/N$. The $T/J < 1/L$ susceptibility for a finite
FIGURE 73. Uniform susceptibility of the 2D Heisenberg model computed using several $L \times L$ lattices. The right panel shows low-$T$ results for $L = 128$ and 256 on a more detailed scale (with the two data sets coinciding within statistical errors, indicating convergence to the thermodynamic limit). The solid curve in the right panel is a fit to the $L = 256$ data (using only the $T/J \leq 0.18$ data points) of the form $\chi = a + bT + cT^2$ with the parameters discussed in the text. Error bars are too small to discern but are typically $\approx 2 \times 10^{-5}$ for the low-temperature $L = 256$ results.

system is dominated by these excitations. Once $T > 1/L$ spin waves can also be excited, and one may therefore suspect that a system size of at least $L > 1/T$ will be required for convergence to the thermodynamic limit. Here we will not study the convergence in detail, but just conclude based on the data in Fig. 73 that the $L = 256$ results are safely converged down to $T/J = 0.03$ and can be used to test the form (287).

Tests of analytical predictions can be carried out in different ways. We could here use the value of $\rho_s$ extracted from the finite-size extrapolations of the $T = 0$ data above and check the agreement between the SSE results for $\chi(T)$ and Eq. (73), with only $\chi(0)$ adjusted as a fitting parameter to obtain the best agreement. Another way would be to also adjust $\rho_s$, the best-fit value of which can be compared with the result of the ground state extrapolations. Here we will proceed in a different way, which tests both the $T$ and $T^2$ corrections in (287). Fitting SSE data to a form $\chi = a_0 + a_1 T + a_2 T^2$, we can use the form of the coefficients in (287) to extract corresponding estimates $\rho_s(1)$ and $\rho_s(2)$ for the spin stiffness from the linear and quadratic coefficients $a_1$ and $a_2$. These estimates should agree with each other and with the result of the ground state calculation if the temperatures used in the fit are sufficiently low. At higher temperatures, corrections including higher powers of $T$ will be important and should lead to a bad fit to the quadratic form and disagreements between the different stiffness estimates.

The quadratic fit is good and consistent values of the exponents are obtained if only data for $T/J \leq 0.18$ are included. The fit is shown in the right panel of Fig. 73, and the stiffness constants extracted from the parameters are $\rho_s(1) = 0.180(3)$ from slope and $\rho_s(1) = 0.179(6)$ from the quadratic correction. These values are in good agreement with (but noisier than) the ground state result $\rho_s = 0.18100(5)$. Fixing the form (287) and extracting the best-fit values of $\chi(0)$ and $\rho_s$ gives a less noisy estimate [225], but since we are still dealing with corrections to a $T = 0$ quantity it is not easy to achieve the same precision as we did by extrapolating the finite-size values of $\rho_s$ at $T = 0$. 

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FIGURE 74. The temperature dependent correlation length (graped versus the inverse temperature) of the 2D Heisenberg model computed for different lattice sizes using the spin structure factor definition (70), divided by the factor $\sqrt{15/16}$ coming from Eq. (72). The dashed curve is the expected asymptotic ($L = \infty, T \to 0$) form (289) with the parameters $\rho_s = 0.1810$ and $c = 1.660$ determined previously.

The $T \to 0$ susceptibility extracted from the above fit is $a_0 = \chi(0) = 0.04378(3)$. As we discussed in Sec. 4.4.2, the transverse susceptibility $\chi_\perp$ in the symmetry-broken state at $T = 0$ is this number multiplied by $3/2$, giving $\chi_\perp = 0.06567(4)$. In combination with the spin stiffness, this value gives us access to another ground state parameter; the spin wave velocity. We discussed the spin stiffness constant as an elastic modulus in Sec. 3.5. This elastic-medium approach to the long-wavelength properties of quantum magnets is also referred to as the hydrodynamic description [229]. The transverse susceptibility is there analogous to a mass density, and one can relate the spin wave velocity to the spin stiffness and the susceptibility according to

$$c = \frac{\sqrt{\rho_s}}{\chi_\perp}. \quad (288)$$

Using the values extracted above for $\rho_s$ and $\chi_\perp$ gives the velocity $c = 1.6602(6)$. Recall that linear spin wave theory (Sec. 2.1.1) gives $c = \sqrt{2}$. The difference between these two values is captured very well by spin wave theory including $1/S$ corrections, and the “renormalization factor” of $c$ can also be computed quite precisely using a series expansion around the Ising model [230].

The temperature dependence of the correlation length of the infinite system is also known, including a correction to the leading exponential divergence [5, 227]:

$$\xi(T) = \frac{e}{8 \pi \rho_s} \left( 1 - \frac{4 \pi \rho_s}{T} \right) \exp \left( \frac{2 \pi \rho_s}{T} \right). \quad (289)$$

We have already determined the two parameters involved here, the ground state spin stiffness and velocity, and we will use their values quoted above to directly test the
analytical prediction against SSE results based on the “second moment” structure factor definition (70) of the correlation length.

Since the correlation length is a rapidly divergent quantity, very large lattices are required to converge the calculations to the thermodynamic limit at low temperatures. The lattice length \( L \) should be several times larger than \( \xi \) to achieve complete convergence. Fig. 74 shows results for several lattices sizes up to \( L = 256 \). While larger systems have been studied in the past [225, 226], already these results, which are well converged for \( T/J \geq 0.25 \), show that the form (289) describes the numerical data very well. The small deviations have been discussed in the literature [225, 226] and can be understood as arising from higher-order corrections to the form (289).

5.3.4. Quantum phase transition in a dimerized system

We now study the quantum phase transition in the 2D columnar dimerized Heisenberg model illustrated in Fig. 4(b). The phase transition of the ground state takes place as a function of the coupling ratio \( g = J_2/J_1 > 1 \) and is caused by quantum fluctuations, which here correspond to an increasing density of singlets on the dimers and at some point lead to the loss of the Néel order existing when \( g \approx 1 \). We already examined some SSE results for the sublattice magnetization of this system in Fig. 5, and the behavior indicated a phase transition between a Néel-ordered and a nonmagnetic ground state at coupling ratio \( g \approx 1.9 \). In this section we apply the machinery of finite-size scaling at criticality, which we discussed in the context of classical systems in Sec. 3.3 and generalized to quantum systems in Sec. 3.6. Here we first analyze the critical behavior of several quantities calculated at sufficiently low temperatures to access the ground state critical behavior, and then discuss consequences of the \( T = 0 \) quantum critical point at non-zero temperatures. Before that, let us spend a few words on the implementation of the SSE algorithm for a model with non-uniform couplings.

**SSE method.** An appealing feature of the SSE algorithm for the Heisenberg model is that the coupling strengths only enter in the acceptance probabilities (265) and (266) in the diagonal updates, where the coupling \( J \) is absorbed in the inverse temperature \( \beta = J/T \). For non-constant couplings, we only have to define \( \beta_b = J_b/T \) for each bond \( b \). In the implementation in pseudocode \{27\}, some scheme has to be used to identify the coupling corresponding to a generated bond \( b \) in an insertion attempt. In a system with a large number of different couplings (which could even be random, different for every bond) the couplings (in the form of \( \beta_b \)) should be stored as a list. For systems with just two different couplings, such as the dimerized lattice considered here, it is better to just order the bonds in such a way that the coupling can be quickly determined, e.g., with \( b \leq N/2 \) and \( b > N/2 \) corresponding to \( J_2 \) and \( J_1 \), respectively. Note that for an operator removal, unlike the case of uniform couplings in code \{27\}, we now also need to extract the current bond, \( b = S(p)/2 \), in order to determine the probability.

**Finite-size scaling for \( T \to 0 \).** When analyzing the ground state, we can proceed as in the case of the standard uniform 2D Heisenberg model in the previous section, simulating systems at sufficiently low temperatures to achieve \( T = 0 \) convergence of all the
quantities of interest. This approach is discussed for various dimerized systems in, e.g., Refs. [85, 88] (as well as in many older works). Another approach is to study systems at inverse temperature $\beta = L^z$, where $z$ is the dynamic critical exponent (which we discussed in Sec. 3.6) [231]. This is motivated in the following way, by a generalization of the finite-size scaling hypothesis (64): In a quantum system the scaling function $f(\xi/L)$ should be replaced by a function with two arguments, $f(\xi/L, \xi_T/L_T)$, where the correlation length in the imaginary time dimension depends on the spatial correlation length $\xi$ according to $\xi_T \sim \xi^z$ (which defines the dynamic exponent) and the length of the system in the imaginary time direction is $L_T = c/T \sim \beta$ (where $c$ is a velocity). If we choose $\beta \propto L^z$, then the scaling function can be written as $f(\xi/L, (\xi/L)^z)$, which is a function of the single argument $\xi/L$. Thus, the finite-size scaling procedures can be used exactly as in the classical systems discussed in Sec. (3.3.2). This is the case also if we take the limit $\beta \to \infty$ for each $L$ (in practice finite $\beta$ large enough for convergence to this limit), because then $\xi_T/L_T \to 0$, and there is again only one argument $\xi/L$ left in the scaling function.

There is plenty of evidence already that $z = 1$ in dimerized Heisenberg models, and we will here use systems with $\beta = L$. This allows for studies of larger systems than in the $\beta \to \infty$ limit, although it is not a priori clear which approach is in the end better, since the corrections to the leading finite-size scaling behavior can be different. Here we use $L$ up to $L = 128$. We will also test explicitly that systems with $\beta = L$ exhibit behavior consistent with $z = 1$, by studying quantities which depend on $z$.

We first locate the critical coupling by examining quantities that should be size independent at $g_c$. Fig. 75 shows the $g$ dependence of both the Binder cumulant and the spin stiffness, with the latter multiplied by $L$ to compensate for the expected quantum critical scaling form $\rho_s \sim 1/L$, obtained the classical form (99) with $d \to d + z = 3$.

The Binder cumulant is defined according to (77), with the number of components $n = 3$. Note, however, that (77) is defined with the full scalar product $m^2 = \mathbf{m} \cdot \mathbf{m}$ in (75), whereas with the SSE method we here only compute the $z$ component expectation values $\langle m_z^2 \rangle$ and $\langle m_z^4 \rangle$ (the off-diagonal components being more difficult to evaluate.
FIGURE 76. Size dependent critical coupling for the dimerized Heisenberg model extracted from $(L, 2L)$ crossing points of the Binder cumulant, the spin stiffness constants, and the correlation lengths. The curves show fits to the form $g_c(L) = g_c(\infty) + a/L^\omega$.

One can easily find the geometrical factors relating these by integrating the $z$ component $\cos(\Theta)$ of a classical 3D unit vector over the the angles, giving $\langle m^2 \rangle = 3\langle m_z^2 \rangle$ and $\langle m^4 \rangle = 5\langle m_z^2 \rangle^2$. For locating the critical point, these factors play no role, and we could also use the plain Binder ratio defined as $R_2 = \langle m_z^4 \rangle / \langle m_z^2 \rangle^2$.

Since the dimerized lattice does not have $90^\circ$ rotational symmetry, the stiffness constants in the $x$ and $y$ directions are different. Although the numerical values are indeed quite different, their scaling behaviors close to the critical point is very similar, however [the $x$ stiffness is approximately a factor 2 larger—the dimers are oriented in the $x$ direction as in Fig. 4(b)]. Only the $x$ stiffness is shown in Fig. 75(b).

Curve crossings are indeed seen in Fig. 75 for both $U_2$ and $\rho_y L$, and after some significant drift of the crossing points (e.g., for systems of size $L$ and $2L$) for small $L$, they seem to converge to roughly the same value in both cases. Note that the crossing points for $U_2$ and $\rho_y$ approach $g_c$ from opposite directions, which can be useful for bracketing the critical value [85, 88]. Crossing points can be located numerically by fitting a polynomial of suitable order to some of the data points, repeating the procedure several times with added Gaussian noise to compute error bars. Fig. 76 shows results of such procedures for the Binder ratio, the $x$ and $y$ stiffness constants, as well as the correlation lengths [computed using the definition (70)] in both the $x$ and the $y$ direction. Fits to the data points of the form $g_c(L) = g_c(\infty) + a/L^\omega$ are also shown. This form describes well all the data for $L \geq 10$ (the sizes shown in the figure). All the extrapolated values of $g_c$ fall within the range [1.9094, 1.9096], and if the $L = 10$ data are excluded the range narrows even further. The exponent $\omega$ is in the range $2 \sim 2.5$ for all quantities (being largest for $U_2$). Treating all five values obtained in these extrapolations as independent statistical data gives $g_c = 1.90948(4)$ as a final estimate for the critical point. This is in good agreement with (but with smaller error bar than) a recent estimate $g_c = 1.9096(2)$ obtained using $T \to 0$ data for the same quantities on lattices with $L$ up to 64. The crossing point shift exponents $\omega$ are also in good agreement.
The existence of a limiting value of the location $g_c$ at which the $L\rho_s$ curves cross does not prove that $z = 1$. One also should check that the values of $L\rho_s$ are well behaved, i.e., that a crossing point in the plane $(g, \rho_s L)$ really forms. A plot such as Fig. 76 for $\rho s L$ at the crossing points for lattices of size $L$ and $2L$ confirms that this is the case. Examining the Binder cumulant in a larger window of couplings, one can not find any indications of negative values, which would be a sign of phase coexistence at a first-order transition (as discussed in Sec. 3.4). Thus, the scaling behavior so far supports at continuous quantum critical point with $z = 1$.

We could now proceed to perform data collapse fits in order to find the correlation length exponent $\nu$, as we discussed for classical systems in Sec. 3.3. This has been done for dimerized Heisenberg model, including also scaling corrections [85, 88], and the results are in good agreement with the expected 3D classical Heisenberg universality class. Here we instead just discuss the exponent $\eta$ appearing in the critical correlation function (53), where we should again replace $d$ by $d + z = 3$. The staggered structure factor $S(\pi, \pi)$ is the spatial integral (sum on the 2D lattice) of the correlation function (269), while the Kubo integral (271) for the staggered susceptibility $\chi(\pi, \pi)$ corresponds to a 3D space-time integral (a 2D lattice sum and an imaginary time integral). Performing these integrals with the above critical form of the correlation function, with a cut-off equal to the system size $L$, gives $S(\pi, \pi) \sim L^{d-z-\eta}$ and $\chi(\pi, \pi) \sim L^{d-\eta}$. We will test this behavior of the critical system and extract the exponent $\eta$, using the value of $g_c$ found above. Instead of performing new simulations at this point (which is known only to within a statistical error), one can perform polynomial interpolations within the existing data sets. One can then also easily check the behavior for points slightly off the best estimate of $g_c$ (plus and minus one error bar), to check the sensitivity of the fitted $\eta$ to the location of the critical point. The simplest way to analyze the critical behavior is to fit a straight line to $\ln(S)$ and $\ln(\chi)$ versus $\ln(L)$ (as was done for $S(\pi, \pi)$ of this model in [88]). Some corrections to scaling are always expected, and if the data have small error bars a straight line can only be fitted to large lattices. With the data sets used here, statistically acceptable linear fits are only obtained when using system sizes $L \geq 48$. We will therefore also include subleading corrections and assume the following forms

$$S(\pi, \pi) = aL^{1-\eta} + bL^\omega, \quad \chi(\pi, \pi) = aL^{2-\eta} + bL^\omega,$$  \hfill (290)

where one would expect the subleading exponent $\omega$ to be much smaller (possibly even negative) than the leading exponents. One could in principle perform a combined fit with $\eta$ fixed to be the same for the two quantities (with $\omega$ and the constants different), but here we will fit the data sets independently.

In order to more clearly see the role of the subleading correction, $S(\pi, \pi)$ and $\chi(\pi, \pi)$ are graphed in Fig. 77 with the dominant $L$ and $L^d$ factors divided out. The asymptotic $L \to \infty$ behavior is then in both cases $\sim L^{-\eta}$, where $\eta$ is expected to be small. The currently best estimate for the classical 3D Heisenberg universality class is $\eta = 0.0375(5)$ [113]. The fits to (290) give $\eta = 0.029(2)$ for $S(\pi, \pi)$ and $\eta = 0.020(3)$ for $\chi(\pi, \pi)$. In principle the forms (290) should of course work only exactly at $g_c$, but in practice, for a finite range of system sizes, they fit the data well in some window around the true critical point. As seen in Fig. 77, the interpolated values of the two quantities at $g_c$ plus and minus one error bar deviate visibly from those at the midpoint, but the fits
FIGURE 77. Finite-size scaling of the staggered structure factor (left) and susceptibility (right) in close proximity of the estimated critical coupling ratio $g_c = 1.90948(4)$. The powers of $L$ corresponding to $z = 1$ and $\eta = 0$ have been divided out. The remaining asymptotic size dependence should then be governed by the actual value of $\eta$. The fits to the forms in (290) give $\eta = 0.029(3)$ from $S(\pi, \pi)$ and $\eta = 0.020(4)$ from $\chi(\pi, \pi)$. The fits were based on $L \geq 8$ data, but the resulting curves also go through the $L = 4$ and 6 points. The dashed curves shows the behavior at $g = 1.90948$ without the scaling corrections (i.e., with $\eta$ and $\alpha$ kept at their values obtained in the fit including the corrections).

are statistically acceptable in all three cases. The statistical errors quoted above arise predominantly from the uncertainty in the critical coupling. The subleading exponents in (290) are $\omega = -0.2(2)$ for $S(\pi, \pi)$ and $\omega = 0.6(2)$ for $\chi(\pi, \pi)$.

Fig. 77 also shows the fitted functions with the subleading corrections disregarded (with the other parameters kept at their values obtained in the fit with corrections). Clearly the corrections are quite significant, being completely responsible for the maximums in both curves at $L \approx 10$. One can of course obtain much better fits to the larger lattices without subleading corrections. As mentioned above, with the rather small error bars of the data used here, only systems with $L \geq 48$ can be included in such a fit. Even then, there must be some influence of the neglected corrections. The values of $\eta$ do in fact come out somewhat lower if no corrections are included. With the corrections, all the data ($L \geq 4$) can be included, but to be on the safe side only $L \geq 8$ data were included in the fits quoted here and graphed in the figure. These fits still do pass very closely through the $L = 4$ and 6 data points, which further reinforces the quality of the functional form used.

The $\eta$ values obtained here, and also in Ref. [88], are a bit lower (by a few error bars) than the best available classical 3D Heisenberg value [113] quoted above. Most likely, these small discrepancies are due to further scaling corrections, but it would still be good to push QMC calculations for various dimerized Heisenberg models to even higher precision (using larger lattices, a denser grid of lattices sizes, and also reducing the error bars of the computed quantities) in order to establish the agreement with the classical exponents for sure. This is particularly interesting and important in light of the fact that the staggered dimer model illustrated in Fig.4(c) (and also some other dimerization patterns) seems to show small but statistically significant deviations from the expected 3D Heisenberg exponents [89, 233, 234]. These systems may still not be in...
FIGURE 78. Cross-overs (indicated by thick curves) in the coupling-temperature plane of a generic dimerized 2D Heisenberg model. The quantum-critical point (circle) controls the behavior in the $T \geq 0$ quantum critical (QC) regime, up to some temperature where lattice effects become important (i.e., when $\xi \sim 1$). For couplings away from the critical point there is a low-temperature cross-over into either renormalized classical (RC) or quantum disordered behavior. For a 2D system there is Néel order only exactly at $T = 0$. The thickness of the curves separating the different regimes reflect the fact that the cross-overs are not sharp, but can take place over significant temperature windows.

a different universality class, as originally proposed [89], because the deviations could also originate from anomalously large scaling corrections [233, 234]. Regardless of the underlying reason for the deviations, their origin remains unclear and should be further examined (and such work is in progress [234]).

Quantum critical scaling at $T > 0$. One of the most remarkable and important aspects of quantum criticality is that the properties (the universality class) of a $T = 0$ critical point also can strongly influence a system at $T > 0$, often up to rather high temperatures [84] and also if the system does not have exactly the couplings corresponding to a $T = 0$ critical point (but is near such a point, in a sense which we will make more precise below). Critical fluctuations and scaling behavior can therefore be much more generic features of quantum systems than classical systems [142], where critical behavior often can be observed only very close to the critical point.

In a quantum system, critical behavior can be expected when the temperature is the dominant energy scale. In dimerized 2D Heisenberg models, the energy scales on the Néel and nonmagnetic side of the transition are, respectively, the spin stiffness $\rho_s$ and the singlet-triplet excitation gap $\Delta$. One can therefore expect manifestations of quantum criticality for $T \gg \rho_s$ and $T \gg \Delta$, for $g < g_c$ and $g > g_c$, respectively. In continuum field theories there is no upper-bound temperature for the quantum critical behavior, while for a lattice Hamiltonian there has to be a cross-over to a different behavior above some temperature where the spins become essentially independent due to thermal fluctuations (which cannot be captured by continuum field theories, where independent discrete spins do not exist). The cross-over to the high-temperature limit could be defined, e.g., on the basis of the Curie (independent spin) susceptibility $\chi \to 1/(4T)$ obtaining for $T \gg J$. As we will see below, the quantum critical susceptibility is completely different.

Putting all this together, for a 2D dimerized Heisenberg model one expects a generic “cross-over diagram” [5, 84] of the type shown in Fig. 78. For temperatures below the high-$T$ cross-over, there are three different “regimes”, in which the behavior is controlled
be the corresponding largest energy scale in the problem; $\rho_s, \Delta, \text{ or } T$. In the renormalized classical (RC) regime, the spin stiffness dominates, and the correlation length diverges as $T \to 0$ according to (289), as we saw explicitly for the standard uniform Heisenberg model in Sec. 5.3.3. In the quantum disordered (QD) regime, the correlation length saturates at some finite value as $T \to 0$, with $\xi \to$ constant roughly below $T \approx \Delta$. In the quantum critical (QC) regime, the correlation length should diverge at $g_c$ when $T \to 0$ according to

$$\xi \sim 1/T^z.$$  (291)

This behavior can be understood based on the path integral mapping of the quantum system in $d$ dimensions onto a classical $(d + z)$-dimensional system. There the coupling $g$ of the quantum system corresponds to $T$ of the $(d + z)$-dimensional system. We can then think of temperature scaling in the quantum system as a kind of finite-size scaling in the imaginary time dimension, in which the thickness of the effective system is finite; $L_\tau \propto 1/T$ (i.e., we keep $T$ constant and just change the thickness of the effective system). By definition of the dynamic exponent, at $g_c$ we have

$$\xi \sim \xi_\tau^{1/z}. \quad (292)$$

If the time dimension thickness $L_\tau = \infty$, both the correlation lengths $\xi$ and $\xi_\tau$ diverge at this point, but with finite $L_\tau$ (but infinite spatial size $L$) we should replace $\xi_\tau$ by $L_\tau \propto 1/T$ in scaling formulas (exactly as we do in classical finite-size scaling when $L < \xi$), whence (292) gives (291).

Quantum critical scaling forms have been derived for many quantities based on these ideas worked out quantitatively and in great detail [84] using renormalization group methods and large-$N$ expansions [with $N$ here being the number of components of SU($N$) symmetric spins, with $N = 2$ for the physical $S = 1/2$ spins] within the continuum field theory [the $(2 + 1)$-dimensional nonlinear sigma-model]. The main point is that one can observe scaling, with some corrections (the size of which depends on the quantity), also when $g \neq g_c$, in the QC regime illustrated in Fig. 77. Away from $g_c$, there is a crossover into either RC or QD behavior. Here we only look at two important quantities; in addition to the correlation length we also analyze the uniform susceptibility, which is perhaps the quantity for which QC behavior away from $g_c$ is the most robust. The predicted form is linear in $T$ exactly at $g_c$, with a constant shift away from $g_c$,

$$\chi = \frac{a}{c^2}T + b, \quad (b = 0 \text{ at } g = g_c), \quad (293)$$

where $c$ is the spin wave velocity. The constant $a$ is not known exactly, but its value computed based on the leading terms in an $1/N$ expansion is believed to be close to the actual value [84]. The linear behavior should apply strictly only when $T \to 0$, since there are higher powers of $T$, with unknown prefactors, which are not included in (293).

We begin by investigating the finite-size convergence of the susceptibility at $g_c$ (using $g_c = 1.9095$, within the error bars of the critical point determined above), in order to make sure that we can reliably obtain results reflecting the thermodynamic limit at low temperatures. The left panel in Fig. 79 shows the temperature dependence of $\chi/T$ (which is easier to look at in graphs than $\chi$ itself) for several power-of-two system sizes up to
FIGURE 79. Susceptibility divided by temperature for the dimerized system at the critical point \((g = 1.9095)\) computed using different lattice sizes. The peaks seen in the left panel for small systems is a finite-size feature which moves toward \(T = 0\) as \(1/L\). The solid curve in the right panel is a cubic polynomial fit to the \(L = 512\) data \((T/J_1 \leq 0.20)\), which gives the prefactor \(a/c^2 = 0.0916(1)\) in Eq. (293). For comparison, in the right panel, size-converged results for for \(g = 1.9090\) are also shown (the lowest curve, marked with \(+\)). In this temperature range, quantum critical behavior is seen also at this coupling, which is slightly on the nonmagnetic side of the transition (and hence \(\chi/T \to 0\) eventually as \(T \to 0\)).

\(L = 512\). As expected, the results converge quickly at high temperatures, with finite-size effects entering at approximately \(T \sim 1/L\) (which can be expected on account of the dynamic exponent \(z = 1\)). The right panel of Fig. 79 shows the low-\(T\) results on a more detailed scale, including also results for \(L = 512\). Based on this comparison of results for different sizes, one can conclude that the thermodynamic limit can be studied with \(L \leq 512\) lattices down to \(T/J_1 \leq 0.03\) (and probably even a bit below that).

The right panel of Fig. 79 also shows a fit of \(\chi/T\) to a cubic polynomial at low temperatures. The corrections to the asymptotic linear form (293), is quite prominent, leading to an \(\approx 8\%\) increase in \(\chi/T\) \[which can be considered as effective temperature dependent prefactor \(a/c^2\) in (293)\] from the minimum around \(T/J_1 \approx 0.3\) to the eventual \(T \to 0\) value. In principle the extrapolated \(T = 0\) value \(a/c^2 = 0.0916\) can be used to extract the spin wave velocity, but since the prefactor \(a\) is not known exactly \[84\], this estimate would likely not be very precise. Another use of the result obtained here would be to use it in combination with an accurate estimate of \(c\) obtained in some other way, which would allow a test of the approximate calculation \[84\] of the prefactor. This is beyond the scope of the discussion here, however.

Fig. 80(a) shows the temperature dependence of \(\chi\) at \(g = g_c\) and at two values some distance away on either side of the critical point. There is a broad maximum at \(T \approx J_1\), which corresponds to the cross-over into the eventual high-temperature Curie behavior. Below the maximum, these near-critical systems all exhibit an approximately \(T\)-linear susceptibility, in accord with the form (293) with \(b < 0\) and \(b > 0\) for \(g > g_c\) and \(g < g_c\), respectively. At still lower temperatures, there will be cross-overs into RC or QD behavior, which are not seen clearly here because the temperatures are still too high. In the QD regime \(\chi \to 0\) exponentially for \(T\) below the gap \(\Delta\). The RC form is also linear, like the QC form, but the slope changes in the cross-over region. Such a cross-over can...
FIGURE 80. Temperature dependence of the susceptibility (left) and the correlation length (right) in critical ($g = 1.9095$) and near-critical systems. The solid lines show the asymptotic quantum-critical $T$-linear behavior of $\chi$ (from the $T = 0$ value of the fit in Fig. 79) and the $1/T$-linear behavior of $\chi_x$ [following the form (292) with a small negative constant correction].

actually be seen in the standard 2D Heisenberg model, corresponding to $g = 1$, which may appear to be too far away from the critical point. In the left panel of Fig. 73 one can nevertheless see an approximately linear behavior of $\chi$ in the range $T/J \in [0.3 \sim 0.5]$, before the RC behavior sets in at lower temperatures. The slope of $\chi$ in the narrow window is in very good agreement with that obtained with the known spin wave velocity and the approximately calculated prefactor $a$ in (79), which indicates that this behavior really is a manifestation of quantum critical behavior far from a quantum critical point [84, 225] (although the very good agreement may be to some extent fortuitous, since, as we concluded above, there are significant corrections to the purely linear form exactly at $g_c$, at much lower temperatures than the QC window at $g = 1$).

Fig. 80(b) shows the correlation length in the $x$ direction (which is about 30% larger than the $y$ correlation length at $g_c$) at the same near-critical couplings as in Fig. 80(a). While the behavior is very linear, with a small constant correction to the asymptotic form (292) with $z = 1$, the results for the systems slightly off the critical point deviate significantly from linearity below $T/J_1 \approx 0.3$. Being a divergent quantity for $g \leq g_c$, the correlation length has much larger corrections to the critical form than the non-divergent uniform susceptibility.

5.3.5. Néel–VBS transitions in J-Q models

With the J-Q models introduced in Sec. 2.4.3, one can study quantum phase transitions at which not only the antiferromagnetic long-range order vanishes, but a different symmetry is broken in the nonmagnetic state as a VBS forms. Superficially, a VBS may seem rather similar to the nonmagnetic state of a dimerized Heisenberg model, because in both cases the system exhibits a pattern of strongly and weakly correlated nearest-neighbor pairs. These states are fundamentally very different, however, because in a “manually” dimerized Heisenberg model the hamiltonian itself breaks the translational
symmetry, and the strongly correlated bonds also correspond to the strongly coupled spin pairs. In contrast, in J-Q models, the Hamiltonian obeys all the symmetries of the lattice, and the translational symmetry is spontaneously broken in the VBS state. The quantum fluctuations of this VBS, especially close to the phase transition, are therefore very different (and much more interesting).

On the square lattice, which we will consider here, and with the types of interactions we choose, the VBS can form in four equivalent patterns; hence the broken symmetry is $Z_4$. We will first discuss the standard J-Q model with four-spin couplings [17] (and the term J-Q model will then refer just to that particular case). In this case the Néel–VBS transition appears to be continuous and shows many of the hallmarks of the proposed [16, 235] (and still controversial [236]) deconfined quantum critical point. The VBS state in this case is most likely columnar, but the exact nature of the state is masked by an emergent U(1) symmetry in the VBS state (which in this model is always near-critical, with large fluctuations of the VBS order parameter). This is one of the salient features of the putative deconfined quantum critical point. Despite the many ways in which the behavior of the J-Q model agrees with the theory, there are also aspects of the Néel–VBS transition that remain unexplained—although not necessarily in conflict with the theory, because detailed analytical calculations are very difficult [235, 237] and many properties of the deconfined quantum critical point in SU(2) spin systems are simply not known quantitatively based on the field theory proposed to describe the transition. We will here look at one example of strong corrections (possibly logarithmic) to the quantum critical scaling behavior, which had not been predicted by the theory of deconfined quantum critical points.

We will also discuss some results for a different kind of multi-spin coupling, which leads to a staggered VBS pattern [like the dimer pattern in Fig. 4(c)]. The phase transition in this case is clearly first-order [109]. We will discuss the reasons for the qualitatively different nature of the transitions in the two models.

**SSE implementation for the J-Q model.** We first discuss some implementation issues for studies of the J-Q model with the SSE method. The model was first studied with a projector QMC technique formulated in the valence bond basis (which we will briefly discuss in Sec. 6). It is, however, also easy to implement the SSE method for it [238], which was initially done using “directed loop” updates (a generalized loop algorithm, which we will also summarize Sec. 6). An SSE algorithm for the J-Q model can also be devised which is almost identical to the standard algorithm with operator-loops for the isotropic Heisenberg model. To see this, we can write the J-Q Hamiltonian with four-spin interactions, Eq. (23), in terms of the diagonal and off-diagonal bond operators (255) and (256) used in the SSE algorithm for the Heisenberg model:

$$H = -J \sum_{[b]} (H_{1b} - H_{2b}) - Q \sum_{[bc]} (H_{1b} - H_{2b})(H_{1c} - H_{2c}).$$

(294)

As in the Heisenberg model $[b]$ is a bond with two interacting spins $[i(b), j(b)]$, while $[bc]$ denotes two parallel bonds $[i(b), j(b); k(c), l(c)]$, corresponding to the singlet projector product $S_{ij}S_{kl}$ in (23). We will consider these bonds arranged as in Fig. 8, with the summation in (294) taken over all translations on the lattice and including horizontal
FIGURE 81. Examples of vertices in the J-Q model. Only some of the allowed spin states of the vertex legs (open and solid circles) corresponding to the operators are shown. Open and solid bars indicate diagonal and off-diagonal bond operators, respectively. The J-vertices are identical to those in the Heisenberg model. The Q-terms in Eq. (294) are products of two bond operators, which when expanded out include all combinations of diagonal and off diagonal factors. Allowed loops pass only through one of the operator factors in the case of Q-vertices, as illustrated here with loop segments at all the permissible leg pairs. Flipping loops can lead to any combination of allowed operators and spin states.

as well as vertical bond orientations. The scheme is, however, independent on how the singlet projectors are arranged, and also the generalization to an arbitrary number of bonds in the Q term is trivial. The only constraint is that we have to avoid sign problems, which we do if both $J > 0$ and $Q > 0$ [with the minus signs in (294), which corresponds to energetically favoring singlets on the bonds included in both the J and Q terms]. The absence of sign problems was discussed based on a sublattice rotation in [238], and it can also be demonstrated using the simple operator counting arguments used for bipartite Heisenberg models in Sec. 5.2.

We now have J-vertices with four legs as well as Q-vertices with eight legs, as illustrated in Fig. 81. The Q-vertices can be considered as two J-vertices joined together, in all possible combinations of diagonal and off-diagonal parts arising from the four operators in the Q term of (294). It is then clear that we can proceed in the same way as we did for the Heisenberg model, updating the operator string and a stored state using a combination of diagonal and loop updates. The key is here again that the matrix elements are the same for all J- and Q-vertices (the values being $J/2$ and $Q/4$), which means that a loop update in which the type of vertex (J or Q) is not changed can always be accepted. In the case of the Q-vertices, the loops satisfying this constraint enter and exit at the same operator factor, as illustrated with loop segments in Fig. 81.

Both J and Q diagonal operators are inserted and removed in the diagonal update. The simplest way to insert diagonal operators is to choose completely randomly among all the possible single-bond $[b]$ and double-bond $[bc]$ instances in (294). There are $N$ each of horizontal and vertical bonds $[b]$ and also $N$ each of horizontal and vertical bond pairs $[bc]$, for a total of $4N$ cases to choose from. The spins in the current state have to be antiparallel on the bond or bonds acted on by the chosen operator, and if that is the case the acceptance probability is a simple modification of Eq. (265), with either $\beta_J = J/(2T$) or $\beta_Q = Q/(4T)$ replacing $\beta$, depending on the type of operator inserted. The number of bonds $N_b$ is replaced by the total number of bonds and bond pairs, i.e., $4N$. The same modifications apply to the removal probability (265) as well. If the ratio $Q/J$ is much different from 1, which is the case in the VBS state and at the phase transition, it is better to take this ratio into account already when generating the bonds or bond pairs (but even the trivial random operator generation actually works very well). Note that $Q$ is much larger than $J$ in the parameter regime we are interested in, and it is therefore best to define the temperature in units of $Q$, i.e., setting $Q = 1$ in the program.
FIGURE 82. Binder cumulant of the staggered magnetization of the J-Q model. The left panel shows results up to $J/Q = 1$, where $U_2$ approaches 1, indicating a Néel state. For smaller $J/Q$ the cumulant vanishes with increasing size, as shown in greater detail in the right panel, demonstrating a nonmagnetic state for $J/Q < 0.045$ (below the crossing points, which accumulate at the critical point). The cumulant remains positive for all $J/Q$, showing that there is no phase coexistence, i.e., based on these results the transition is continuous.

The loop update is identical to the one we developed for the Heisenberg model, with the simple extension that the linked vertex list (illustrated in Fig. 57 for the Heisenberg model) now has eight elements for each vertex, instead of only four in the case of the Heisenberg model. Although the J-vertices have only four legs, the most practical way is to allocate eight storage elements for all vertices, using only the first four of them for the J-vertices (and filling the unused ones with a negative number, so that they are never visited in the loop update).

Continuous quantum phase transition in the J-Q model. The initial study of the J-Q model [17] established the existence of a VBS ground state for small $J/Q$ and a Néel–VBS transition compatible with a $z = 1$ quantum critical point. These results were soon thereafter confirmed in SSE studies of the $T > 0$ quantum critical regime [238, 239]. A later world line QMC study claimed evidence for a first-order transition [240], and this scenario was argued for also based on studies of other systems [236]. Recent calculations on very large lattices (up to $L = 256$) do not, however, find any indications of first-order behavior [110]. This is in violation of the “Landau” rule, according to which a direct transition between states breaking unrelated symmetries should be generically discontinuous, but it agrees with the theory of deconfined quantum critical points [16]. There are, however, significant unexpected scaling corrections in some quantities (which may have been mistaken for signs of a first-order transition in [240]). Here we will first discuss some of the evidence for a single continuous Néel–VBS transition.

The Binder cumulant is a useful quantity for analyzing both continuous and first-order phase transitions. As we discussed in Sec. 3.4, with an example in Fig. 20, phase coexistence at a first-order transition leads to a negative divergent peak in the cumulant at the transition point. Fig. 82 shows results for the antiferromagnetic Binder cumulant $U_2$ of the J-Q model, calculated with the SSE method using $\tilde{\beta} = Q/T = L$ (motivated by the previous work showing that $z = 1$). For $J/Q > 0.05$, $U_2$ increases with $L$ and...
tends toward 1, showing that the system has long-range Néel order. For smaller coupling ratios the cumulant decreases to 0, with no indications of a negative peak developing. The crossing points are well behaved and indicate a critical point at $(J/Q)_c \approx 0.045$.

Next, we examine the correlation length, computed using the structure factor definition (70) for both the spin-spin and dimer-dimer correlations. While the wave-vector for the spin (Néel) order parameter is $(\pi, \pi)$, a columnar VBS corresponds to $(\pi, 0)$ or $(0, \pi)$, which then is used as the reference point $q = 0$ in (70). To compute the dimer correlations, we use the definition (25) but only include the diagonal correlations in the bond operator (26) i.e.,

$$B_{xx}(r) = S_z(r)S_z(r + \hat{x}).$$

(295)

Although the finite-size scaling properties of the correlation functions based on the rotational-symmetric and the $z$ component definitions may be different [238], the correlation lengths extracted from these functions should still both exhibit finite-size scaling $\sim L$ (up to scaling corrections) at the critical point.

We can use the scaled correlation lengths $\xi_s/L$ (spin) and $\xi_d/L$ (dimer) to test whether the two order parameters vanish at the same point, or whether there could possibly be two different transitions. Fig. 83 shows the raw SSE data, which again was generated using $\beta = L$. The curves for different system sizes cross each other, although there is some drift, both horizontally and vertically, even for the largest lattices. While the crossing $J/Q$ values for $\xi_s/L$ and $\xi_d/L$ are quite far apart for small lattices, they move toward each other with increasing system size, in a way consistent with a single critical point.

Fig. 84 shows crossing points extracted from $L$ and $2L$ data for several different quantities (those discussed above as well as the spin stiffness). It is interesting to compare this graph with the corresponding one for the dimerized Heisenberg model, Fig. 76. The convergence of the critical point is clearly much slower for the J-Q model. While the behavior for all quantities is almost linear in $1/L$ over some range of system sizes, for the largest systems the behavior starts to change, in a way which suggests an eventual convergence with some higher power of $L$ to a common critical point $(J/Q)_c \approx 0.045$. Recall that for the dimerized Heisenberg model the behavior is $\sim 1/L^\omega$. 

FIGURE 83. Finite-size scaling of the spin (left panel) and dimer (right panel) correlation lengths of the J-Q model. Note that the crossing points for $\xi_s$ and $\xi_d$ move with increasing lattice size in different directions, toward each other.
with $\omega$ in the range $2 \sim 2.5$, and this behavior applies already for rather small lattices ($L \approx 8$). Because of the slow convergence, it is difficult to estimate the critical point of the J-Q model precisely based on these individual crossing point estimates. There is, however, a remarkable feature seen in Fig. 84: The $\xi_s/L$ and $\xi_d/L$ crossing points form curves that look very symmetric, approaching the apparent asymptotic value at the same rate but from different sides. Therefore, the average of these two estimates exhibits almost no size dependence at all (as also shown in the figure). Based on these average values one can therefore obtain a much better estimate of the critical point than what might initially have been expected. The result based on the four largest-$L$ points (which agree completely within statistical errors) is $(Q/J)_c = 0.04498(3)$.

The slow convergence of the crossing points indicates large scaling corrections in the underlying physical quantities. Note that the Binder cumulant crossings have the weakest size dependence of the quantities analyzed in Fig. 84, which is also clear when comparing the raw data in Figs. 82 (cumulant) and 83 (correlation lengths). Since the Binder cumulant involves a ratio of two similar quantities, anomalous scaling corrections in these may partially cancel each other. It is not at all clear why the finite-size corrections in the crossing points for the two correlation lengths cancel each other almost completely. It may indicate some hidden symmetry between the near-critical VBS and Néel states—perhaps some kind of duality inherent to deconfined quantum criticality [which is the case for the U(1) version of the theory [16], but not explicitly in the SU(2) variant, which is the one of relevance for the J-Q model]. Note that it is not just the locations of the crossing points that are symmetric, but in fact the correlation length curves themselves look rather symmetric in Fig. 83. By reflecting one of the graphs about the critical point, $J/Q - (J/Q)_c \rightarrow -[J/Q - (J/Q)_c]$, the spin and dimer correlation lengths for the largest system fall almost on top of each other, i.e., even the scaling functions for these correlation lengths appear to be the same.
Quantum critical susceptibility scaling at $T > 0$. At a conventional quantum critical point, one would expect an asymptotic linear temperature dependence of the uniform magnetic susceptibility, as we saw in the dimerized Heisenberg model in Fig. 79, where corrections in the form of higher powers of $T$ are also visible in the ratio $\chi/T$. In the J-Q model there are much stronger corrections to the expected quantum critical form. As shown in Fig. 85, there appears to be a multiplicative logarithmic correction, i.e., the asymptotic $T \to 0$ critical behavior may be of the form $\chi \sim T \ln(1/T)$. The results are graphed both on linear and logarithmic temperature scales, in order to clearly convey how the behavior differs from that of the dimerized model in Fig. 79. The behavior in the J-Q model does not appear to be compatible with a conventional higher power-law correction, while a logarithm describes the behavior very nicely.

Logarithmic corrections have not been suggested based on the present theory of deconfined quantum criticality. All analytical calculations within the non-compact $\mathbb{CP}^1$ field theory proposed to describe the transition have been performed using large-$N$ expansions of the SU($N$) generalization of the system (the field theory generalized to $\mathbb{CP}^{N-1}$, where $N = 2$ is the physical number of spin components) [16, 16, 237]. It is possible that different features appear for $N = 2$. On the other hand, Monte Carlo calculations of lattice versions of the field theory [241, 242, 236] have not pointed to any anomalous scaling corrections (although one only found first-order transitions [236]). Since these calculations are in disagreement with each other, the situation for the $N = 2$ theory remains unsettled. Logarithmic corrections in the susceptibility and other quantities do appear in related fermionic gauge field theories [243]. As we discussed in Sec. 4.3 and 5.3.1, in field theory language logarithmic corrections can appear as a consequence of marginal operators, but other explanations are also possible. In principle the corrections may also not really be logarithmic—they could also be due to almost marginal (but in the end irrelevant) operators. Corresponding power-law corrections with a very small exponents may not be distinguishable from logs for the system.
FIGURE 86. Illustration of how local fluctuations of valence bond pairs can lead to a gradual change of a columnar state (a) into a plaquette state (d). In (b) and (c) a small number of bond pairs have been flipped. Local horizontal ↔ vertical fluctuations of the maximum number of all bond pairs can take place with the plaquettes (bond superpositions) arranged as in (d) [one of four equivalent translated patterns]. A fraction \( p \in [0, 1] \) of vertical bond pairs on these plaquettes corresponds to a VBS angle \( \phi \in [0, \pi] \), defined using the \( x \) and \( y \) dimer order parameters (296). In these figures, angles \( \phi \in [\pi, 2\pi] \) correspond to shifting the vertical bonds up by one step up.

sizes accessible. While this scenario is possible for some quantities, in the case of the uniform susceptibility that would also be highly unusual, since normally all corrections to quantum critical behavior in 2D antiferromagnets are integer powers of \( T \) [84].

**Emergent U(1) symmetry in the VBS.** A prominent feature of the theory of deconfined quantum criticality is that, in addition to the normal correlation length \( \xi \) (which can be taken as the dimer correlation length \( \xi_d \) in the Néel state and the spin correlation length \( \xi_s \) in the VBS state), there is another length scale \( \Lambda \) in the VBS phase. This is the distance scale on which excited \( S = 1/2 \) spinons (which are defects, a kind of vortices, in the VBS [245]) are confined into \( S = 1 \) “triplons” (i.e., \( \Lambda \) is the size of the spinon bound state). The same length scale should also be associated with an emergent U(1) symmetry, related to a particular kind of quantum fluctuations connecting two different kinds of VBS ordering patterns—the columnar state and the plaquette state. These “angular” VBS fluctuations and the two kinds of VBS states are illustrated in Fig.86. The angular fluctuations occur on length scales less than \( \Lambda \) (i.e., on length scales less than \( \Lambda \), the system is in a superposition of the two different VBS states). The excited spinons can move freely away from each other up to distance \( \Lambda \), but at larger distances they become confined, due to an effective potential mediated by the VBS long-range order (and in this regard one can note interesting connections to confinement/deconfinement in certain gauge theoreis in particle physics [18]). As the critical point is approached, \( \Lambda \) should diverge as a power of the correlation length, \( \Lambda \sim \xi^a \), with the exponent \( a > 1 \).

The existence of an approximate U(1) symmetry in the VBS order parameter can be tested by examining the distribution \( P(D_x, D_y) \) of order parameters \( D_x \) and \( D_y \), corresponding to columnar order with \( x \)- and \( y \)-oriented bonds and defined as

\[
D_x = \sum_{i=1}^{N} S(r_i) \cdot S(r_i + \hat{x})(-1)^{x_i}, \quad D_y = \sum_{i=1}^{N} S(r_i) \cdot S(r_i + \hat{y})(-1)^{y_i}. \tag{296}
\]

For a given Monte Carlo configuration, \( D_x \) and \( D_y \) can be evaluated at fixed imaginary time (propagated state in the SSE), using the improved estimator for the correlation function discussed in Sec. 5.2.5. The improved estimator is spin-rotational invariant, which can best be understood by considering the equivalence between the loop formulation
and the valence bond basis [244, 31, 50] (which we briefly discuss in Sec. 6). In fact, the improved loop estimator for an equal time correlation function is exactly equivalent to an expectation value of the operator in a particular valence bond state, provided that the temperature is sufficiently low for the simulation only to sample the singlet sector.

While the calculation of \( P(D_x, D_y) \) could be done with the SSE method, the results to be discussed below were instead generated with a ground state method in the valence bond basis (which in its most recent formulation [50] actually is very similar to the SSE method, as discussed in Sec. 6). The rotationally invariant estimator is not strictly needed here [109], and one can also just use the \( z \) spin components in (296), as we did in the correlation function Eq. (295). In the valence bond basis it is, however, natural to compute rotation-invariant quantities. Note that \( P(D_x, D_y) \) is not an operator expectation value (i.e., it is not a normal physical observable), just a distribution of individual measurements in a simulation carried out in a particular basis. Its symmetry properties nevertheless reflect the actual symmetries of the VBS order parameter of the system.

We have already looked at a classical analogy of \( P(D_x, D_y) \); the distribution \( P(M_x, M_y) \) of stripe order parameters in the frustrated Ising model, with results shown in Fig. 19. In that case the four-fold symmetry of the order parameter at and below the transition temperature is obvious, while going to very high temperatures (not shown in the figure) the distribution turns into a central peak with full rotational symmetry [i.e., U(1) symmetry] because of the independent Gaussian fluctuations of the short-range order in different parts of the system. In a similar way, in a VBS one should expect \( P(D_x, D_y) \) to be U(1) symmetric in the Néel state (where there is only short-range VBS correlations), and become four-fold (\( Z_4 \)) symmetric inside the VBS state. As seen in Fig. 87 (which shows recent results for larger systems than in [17, 240, 108]), this is not quite the case, however. Inside the VBS phase, at \( J/Q = 0 \), the distribution is ring shaped, with no weight at the center and no signs of a four-fold symmetry. This shows that the

![Figure 87. VBS order parameter distributions \( P(D_x, D_y) \) calculated for \( L = 128 \) systems at \( J/Q = 0 \) (left) and 0.045 (right). The histograms are shown in regions \( |D_x|, |D_y| \leq D_{\text{cut}} \), with \( D_{\text{cut}} \) chosen for the two coupling ratios to cover the region where the weight is significant. Brighter features correspond to higher probability density. The absolute scale is not important here, only the circular shapes of the distributions. The weight is not distributed completely uniformly in these histograms [obeying neither perfect \( Z_4 \) nor U(1) symmetry], due to the inefficiency of the simulations to fluctuate the VBS angle for large systems.](image-url)
magnitude $D$, with $D^2 = D_x^2 + D_y^2$, of the VBS order parameter has formed, $\langle D^2 \rangle > 0$, but its angular symmetry has not yet been established, i.e., while the magnitude of the VBS order parameter has relatively small fluctuations, the fluctuations in the individual components $D_x$ and $D_y$ [which define an angle $\phi$ of the VBS in the $(D_x, D_y)$ plane] are very large. Close to the critical point, at $J/Q = 0.045$ in the figure, the distribution is peaked at the center, as expected, but again there is no $Z_4$ symmetry.

In the continuum field theory, the operator responsible for the VBS formation is dangerously irrelevant [16, 245, 246], which means that although it is a perturbation with $Z_4$ symmetry, which leads to a $Z_4$ symmetry-broken VBS state, this perturbation does not affect the critical point. The theory of deconfined quantum critical points has built-in $U(1)$ symmetry, and this is unchanged in the presence of the VBS operator. The $J$-$Q$ model has no explicit $U(1)$ symmetry—what is built in here is instead the $Z_4$ discrete rotational symmetry of the square lattice, and naively one would expect the VBS forming on this lattice to exhibit $Z_4$ symmetry, exactly as the stripe order of the Ising model discussed in Sec. 3.4. Instead, we see a $U(1)$ symmetric order parameter distribution $P(D_x, D_y)$, which in light of the theory of deconfined quantum critical points should be expected if the system is smaller than the deconfinement length scale $\Lambda$ (which therefore has to be large, $\Lambda > 128$ even at $J/Q = 0$ according to the results in Fig. 87). If we increase the system size, we should eventually, for $L > \Lambda$, observe a four-fold symmetric distribution (which has been seen in some related systems [108, 247, 248]). A columnar state corresponds to peaks at $(0, \pm D)$ and $(\pm D, 0)$.

As discussed above and illustrated in Fig. 86, the emergent $U(1)$ symmetry is a consequence of local fluctuations between columnar and plaquette VBS patterns [245], in such a way that all possible values of $D_x$ and $D_y$ compatible with a magnitude $D$ (weakly fluctuating) are sampled equally (or almost equally—perfect $U(1)$ symmetry only applies at the critical point). There is an analogy in elementary quantum mechanics; a particle in a slightly asymmetric double well. If the barrier between the wells is very high, then the particle is localized in the deeper well, but if the barrier is not very high (relative to the kinetic energy), then the particle fluctuates between both wells and the wave function also has weight inside the barrier. In the VBS, the wells correspond to the columnar and plaquette states, at angles $n\pi/2$ (columnar) and $n\pi/2 + \pi/4$ (plaquette), with $n = 0, 1, 2, 3$, and fluctuations between them correspond to non-zero probability in the continuum of angles between these points. In the deconfinement theory, the columnar and plaquette VBS states are almost degenerate, becoming exactly degenerate at the critical point. Approaching the critical point, the barrier between them also is reduced. In the double-well analogy, the wells are becoming more degenerate and the barrier between them is reduced. There will then be significant fluctuations between columnar and plaquette states in finite systems (up to $L \sim \Lambda$), and this implies $P(D_x, D_y)$ with weight at all the angles and an $U(1)$ to $Z_4$ cross-over of the symmetry when $L > \Lambda$. This applies also to the order parameter computed on subsystems of finite length $L$ in an infinite (or very large) system (which has been studied in classical models with a $q$-fold order parameter which exhibits $U(1)$ to $Z_q$ symmetry cross-over [249, 250]).

Since the $Z_4$ symmetry is not yet clearly manifested in the $J$-$Q$ model up to the largest lattices studied so far, the issue of the nature of the VBS remains open. However, a $J$-$Q$ model in which the $Q$ term consists of products of three parallel singlet projectors, the VBS order is more robust and a $Z_4$ symmetric distribution corresponding to a columnar
state is seen clearly [108]. Simulations of SU($N$) generalizations of the standard J-Q model, with $N = 3$ and 4, also show columnar VBS ground states [108]. Most likely, the standard SU(2) J-Q model also has a columnar state, and there is indeed some weak hints of this in recent (ongoing) long simulations for $L = 64$.

For system sizes $L \gg \Lambda$, the VBS distribution should be four-fold symmetric, and by investigating how the change from U(1) to $Z_4$ symmetry takes place one can in principle determine the exponent $a$ relating the deconfinement length scale $\Lambda$ and the correlation length $\xi$. Classical examples of the cross-over phenomenon, which also inspired the theory [16], were studied in Refs. [249, 251, 250, 252]. A finite-size scaling method for a quantity sensitive to $Z_4$ symmetry can be used to extract the exponent $a$ [252]. In the standard J-Q model, such an analysis cannot yet be performed, because the cross-over into a $Z_4$ symmetric order parameter is not seen clearly enough in the currently accessible lattices. The analysis has, however, been carried out in the J-Q model with columnar six-spin interactions, with the result that $a \approx 1.2$.

**Other critical exponents.** In addition to the dynamic exponent $z = 1$, other critical exponents have been analyzed in several QMC studies [17, 238, 239, 108, 110]. The most noteworthy result is that $\eta_s$ (the exponent of the critical spin-spin correlation function) is anomalously large, $\eta_s \approx 0.3$, which is in qualitative agreement with the theory of deconfined quantum critical points. A “large” $\eta_s$ was argued for [16] based on the $N = \infty$ value $\eta_s = 1$ in the $1/N$ expansion of the $CP^{N-1}$ field theory (i.e., different from the conventional mean-field value $\eta_s = 0$). The $1/N$ corrections are difficult to compute, but in general support an unusually large exponent [237] (although it is not possible to extend the results reliably to $N = 2$). The behavior of the dimer-dimer exponent $\eta_d$ obtained in QMC calculations for SU($N$) systems [108] seems to follow the scaling with $N$ predicted in the theory [246, 253] (where $\eta_d$ diverges with $N$).

In light of the presence of significant scaling corrections, possibly logarithmic, in the susceptibility analyzed in Fig. 85, and also in the spin stiffness [110], an important question is whether the calculated exponents also are affected. This has so far not been apparent, because standard data collapse procedures with reasonable values of the subleading exponents work very well [108]. The best current estimates for $\eta_s$, based on both $T > 0$ [238] and $T = 0$ [108] QMC calculations agree well with each other. On the other hand, the results discussed here and in [110] put the critical point at slightly higher $J/Q$ than previously, and also the correlation length exponent $\nu$ analyzed with a logarithmic correction included [110] is significantly smaller than other estimates [238, 108]. It would therefore be good to repeat the studies of all the exponent, using several different quantities and larger lattices (which is work in progress).

There is also evidence for logarithmic corrections in the J-Q model from studies of its response to impurities. Introducing a vacancy into the system and studying the spatial distribution of the resulting magnetization imbalance, one can normally, at conventional quantum critical points in dimerized models, observe scaling with the system size in this distribution [254, 255]. In the J-Q model, such scaling analysis again requires the introduction of a multiplicative logarithmic correction [255].

**Applications of other methods to the J-Q model.** One interesting and puzzling aspects of the J-Q model is that it seems very difficult to capture its properties correctly
In the J-Q model studied here, three singlet projectors are arranged in a staggered pattern. All distinct orientations (as shown) and translations of the projector products are included.

using analytical many-body techniques, even ones that can rather accurately locate the Néel–VBS transition in the 2D frustrated J₁-J₂ Heisenberg model. A mean-field treatment starting from a columnar dimer state gives a critical J/Q very far from the QMC result [256]. This approach can be improved to better take into account some of the local fluctuations on plaquettes, which improves the value of the critical point but seems to results in a strongly first-order transition [256]. Cluster mean-field calculations converge very poorly with the cluster size [257]. These results point to unusually strong non-local quantum fluctuations [256], which cannot be easily captured with local approaches starting from small clusters or conventional fluctuations around a fixed dimer pattern. The reason for these difficulties to capture the VBS state may be the emergent U(1) symmetry, which makes it difficult to obtain both the correct long-distance behavior (likely columnar order) as well as the strong fluctuations between columnar and plaquette order on shorter length scales.

**First-order transition in a staggered J-Q model.** One way to test the link between emergent U(1) symmetry and a continuous Néel–VBS transition is to construct a model in which the local fluctuations responsible for rotating the VBS angle are suppressed. Intuition for how to accomplish this comes from the Rokhsar-Kivelson (RK) quantum dimer model [258, 259], which can be regarded as an effective model for an extreme nonmagnetic system dominated by short valence bonds (for which the internal singlet structure is also neglected—the bond configurations are regarded as orthogonal states). The RK hamiltonian on the square lattice can be written as \( H_{\text{RK}} = V - K \), where \( V \) is the diagonal (potential-energy) operator, which counts the number of flippable plaquettes [parallel bond pairs, exactly as in Fig. 86(b,c)] and \( K \) is an off-diagonal (kinetic) term which flips such a pair. This model has a critical point at \( k = v \) which separates a plaquette VBS state [similar to the one in Fig. 86(d)] for \( v < k \) and a staggered VBS state [with the bond pattern exactly as in Fig. 4(c)]. While the plaquette state is destroyed continuously by quantum fluctuations as \( v \to k^- \), the staggered state (of which there are four symmetry-related equivalent ones) has no fluctuations, because it has no flippable plaquettes. The transition upon \( v \to k^+ \) is therefore first-order.

This simple picture of the RK model suggests that an actual staggered VBS in a spin model also should have strongly suppressed local fluctuations, and therefore should not be associated with an emergent U(1) symmetry. Due to the suppression of local fluctuations (and therefore also of large-scale fluctuations), the transition between it and the Néel state should be first-order. The picture is not complete, however, because clearly there must be some fluctuations in the staggered VBS state, considering that a reasonable spin hamiltonian will be quite far from a dimer model and the valence
bond configurations making up the ground state should also include some fraction of longer bonds (in the presence of which some local fluctuations are always possible). It is therefore worth testing this picture in simulations of a model whose ground state can be tuned from a Néel antiferromagnet to a staggered VBS. Such a model can easily be constructed within the J-Q framework, by arranging the singlet projectors of the Q term in a staggered fashion, instead of within a 2 × 2 plaquette (or in columns of three or more projectors, as in [108]). It turns out, however, that a Q term consisting of two staggered projectors is not sufficient for destroying the Néel state. With three projectors, arranged as in Fig. 88 (and the coupling of which we call $Q_3$), a staggered VBS is stabilized, however. We here briefly summarize the evidence for a first-order Néel—VBS found in a recent SSE study of the staggered J-Q$_3$ model [109].

Some results are shown in Fig. 89. The Binder cumulant of the Néel order parameter exhibits a negative peak, which grows with increasing system size. This is a strong indication of a first-order transition, as discussed in Sec. 3.4 (and the results can be compared with those of the classical frustrated Ising model in Fig. 20). The data become very noisy for $L > 12$, because of the inefficiency of the simulations in transitioning between the two coexisting states. For $L > 24$ the system can get completely trapped in a metastable state. On the VBS side close to the transition, if the simulation happens to enter a Néel like configuration during the equilibration, it will stay in a metastable Néel state for a very long time (and vice versa on the other side of the transition). One can use this trapping to investigate the transition in other ways. By doing a series of runs at closely spaced values of $Q_3/J$, starting either far inside the VBS or Néel phase, and starting each new run from the last configuration of the previous run, one can follow the ground state into its corresponding metastable state at the “wrong” side of the transition. The energies computed in two such runs, with $Q_3/J$ either increasing or decreasing,
are shown in Fig. 89. The two branches cross each other, which clearly shows the first-order nature of the transition. The level crossing is most likely an avoided one, between two states with the same quantum numbers, but the spacing between the levels should be extremely small here. This spacing should be inversely related to the tunneling time between the two states at the coexistence point (but note that the relationship between simulation time and the real time dynamics of the system is not known).

The crossing point of the two energy branches can be taken as a finite-size definition of the critical coupling. It shifts with the system size approximately as $1/L^3$, which is consistent with a first-order transition with dynamic exponent $z = 1$ [260]. The extrapolated transition point is $Q_3/J = 1.1933(1)$. At the transition, the VBS order parameter is about 75% of its maximum value (in a perfect staggered valence bond state), which is rather large and motivates the classification of the transition as strongly first-order.

6. SURVEY OF RELATED COMPUTATIONAL METHODS

In these lecture notes we have discussed exact diagonalization methods and the SSE QMC method in some detail and also looked at some illustrative calculations with results for $S = 1/2$ models. There are many issues that were left untouched, regarding methods as well as physics, by restricting the discussion to $S = 1/2$ systems, and within that class also to spin-isotropic systems. The exact diagonalization approach can be easily used for any spin model, but the rapidly growing size of the Hilbert space with $S$ imposes even more severe restrictions on the lattices sizes for $S > 1/2$ systems. The world line and SSE QMC methods can be generalized to any spin model, as long as there is no frustration leading to sign problems. In this section we briefly summarize some QMC schemes for more general spin models, as well as some further recent developments for isotropic $S = 1/2$ systems.

There are several other important computational methods that go beyond the scope of these lecture notes, such as series expansions techniques (high-temperature expansions and expansions around some solvable limit of the ground state, e.g., the Ising limit of an anisotropic Heisenberg model) [261, 262] and the DMRG method [28, 29]. While series expansions can in principle be applied to any model, there are issues with extrapolating the series in a controllable manner, and one cannot, in most cases of interest, expect to reach the same level of accuracy as in QMC simulations of sign problem free models (e.g., in studies of quantum phase transitions). Series expansion methods are nevertheless one of the most powerful classes of methods currently available for 2D frustrated quantum spin systems [100], where QMC methods are limited by the sign problem (but note recent progress in controlling the sign problem at high temperatures [193]). More powerful series expansion schemes are also still being actively developed [263], and one can expect progress to continue on this front. The DMRG method is very powerful for 1D systems and can also be used for 2D systems of moderate size [94]. Extensions of the DMRG method and the related matrix-product states [26, 264, 21] to higher-dimensional “tensor-network” states [21, 106, 265] are currently being explored very intensely. This is an exciting line of research, that may eventually lead to viable schemes for unbiased studies of frustrated spins (and even fermionic systems). It appears unlikely, however, that these methods will, for the foreseeable future, be able to probe quantum spin systems...
FIGURE 90. Examples of the four possible paths through vertices in the anisotropic $S = 1/2$ Heisenberg model (which may include also an external magnetic field) and the new vertices generated when moving along the paths in the directed loop algorithm. The vertical lines at the vertices to the left in each box indicate the entrance and exit legs (with either direction of movement being possible). Given an entrance leg, the exit must be chosen such that a new allowed vertex is generated when the leg spins are flipped (the vertices to the right, without the path indicated). In the “bounce” process (d), the exit is at the same leg as the entrance and the vertex is not modified. There are six allowed vertices in the generic anisotropic Heisenberg model—those in which the spin (z component) is conserved. In the “deterministic” loop algorithm discussed in Sec. 5, there are only four vertices and only paths of type (a) are allowed at the level of precision possible with existing QMC methods for unfrustrated systems.

In light of the above advantages and disadvantages of different methods, the author advocates a two-pronged approach in research on quantum spin models and related systems: (i) Explore new and improved schemes (including also QMC schemes) that may be useful to study models beyond the reach of currently available QMC techniques. (ii) Study interesting physics with state-of-the-art QMC methods for sign problem free models. It is far from true that everything that can be done has already been done with sign problem free models (although one can some times hear such claims)! With the QMC methods available now, such as the SSE algorithm described in Sec. 5, and with the power of modern computers (which still is increasing at an amazing pace, e.g., with a steadily increasing numbers of cores per CPU), it is possible to access very interesting physics that was beyond reach only a few years ago. The discussion of the dimerized and J-Q classes of models in Sec. 5.3.5, in particular, has hopefully convinced the reader that there is much to explore in such systems, and beyond (e.g., random systems exhibiting unusual states and quantum phase transitions [266, 267, 268, 278]).

**Directed loop QMC algorithms.** The directed loop algorithm is a generalization of the SSE operator-loop scheme, introduced in [33] (building on a previous less efficient formulation [190]) and applied there to $S = 1/2$ Heisenberg systems with interaction anisotropies (Ising and XY) and external magnetic field. The main difference between loops and directed loops (in both SSE and world line formulations) is that the path through the vertices is not unique in the directed loop scheme. The path (the exit leg, given an entrance leg to a vertex) is chosen according to certain probabilities, constructed to maintain detailed balance in a space with two defects—the open loop ends (in a way similar in spirit to the “worm” algorithm [32]). For $S = 1/2$ systems, there are four types of vertex paths, corresponding to the location of the exit relative to the entrance, as illustrated and discussed in more detail in Fig. 90. The path probabilities, which should be solutions of the corresponding *directed loop equations* (for detailed balance) are in general not unique. For some models, eliminating the “bounce” process in Fig. 90(d) reduces the directed loop scheme to one of the standard loop updates previously developed for $S = 1/2$ (as in the isotropic system discussed in Sec. 5)
and higher-$S$ models [191, 31, 189]. In other cases the directed loops (and worm) algorithms allow for efficient simulations where the standard loop algorithms are not applicable [33, 269, 270, 271]. Here it can also be mentioned that it is sometimes useful in SSE directed loop algorithms to work with operators defined on cells larger than those containing the elementary operators of the Hamiltonian (e.g., the two-spin bond operators of the Heisenberg model) [272]. This allows for more options in choosing how operators are reconfigured when vertices are traversed by the loops, which sometimes can make simulations much more efficient.

**QMC algorithms in the valence-bond basis.** In Sec. 2.2 we briefly discussed the valence bond basis, in which a basis state is a product of two-spin singlets, as in Eq. (20), and in which any total singlet singlet state can be expanded. The basis is overcomplete and non-orthogonal, which implies that such an expansion is not unique. One way to work with the valence bond basis is to construct and optimize variational states, the simplest type of which is an amplitude product states [273]. Such a state is a superposition

$$|\Psi\rangle = \sum_{\alpha} \Psi_{\alpha} |V_{\alpha}\rangle,$$  \hspace{1cm} (297)

where $|V_{\alpha}\rangle$ is a valence bond product state of the form (20) and the expansion coefficients are products of amplitudes $h(r_{\alpha,i})$ corresponding to the “shapes” of the bonds (the bond lengths in the $x$ and $y$ direction in the case of a 2D system), with $i = 1, \ldots, N/2$ referring to the $N/2$ valence bonds in the configuration (bond tiling) labeled by $\alpha$;

$$\Psi_{\alpha} = \prod_{i=1}^{N/2} h(r_{\alpha,i}).$$  \hspace{1cm} (298)

Amplitude product states of this kind can closely reproduce the ground states of many bipartite Heisenberg systems, for which each singlet $(a, b) = (|\uparrow_a\downarrow_b\rangle - |\downarrow_a\uparrow_b\rangle)/\sqrt{2}$ should be defined with site $a$ on sublattice $A$ and site $b$ on sublattice $B$. With all positive expansion coefficients $\Psi_{\alpha}$ in (297), this convention for the singlet sign corresponds to *Marshall’s sign rule* for the ground state wave function of a bipartite system [274], which in the standard $\uparrow, \downarrow$ spin basis can be written as

$$\text{sign}[\Psi_{\alpha}] = \Psi_{\alpha}/|\Psi_{\alpha}| = (-1)^{n_A},$$  \hspace{1cm} (299)

where $n_A$ is the number of $\downarrow$ spins on sublattice $A$.

The properties of these bipartite amplitude product states can be studied by Monte Carlo sampling of the valence bonds [273, 275, 50]. For the 2D Heisenberg model, the state with all the amplitudes $h(x,y)$ variationally optimized is a Néel state with properties in very good agreement with the true ground state [275], e.g., the energy is within 0.1% and the sublattice magnetization within 1% of the values obtained in QMC calculations. For frustrated systems, appropriate sign rules are not known (and may be too complicated to write down in practice).

The idea to use amplitude product states as a starting point for *projector QMC* simulations is quite old [276, 277], but was only recently developed into a generic and efficient tool [47, 49, 50]. The general idea of a projector scheme is the same as
FIGURE 91. Example (from Ref. [50]) of a loop configuration in the valence-bond projector QMC method formulated in a combined space of spins (open and solid circles) and valence bonds (the arcs capping the loops on the left and right boundaries). The bars with four spins are diagonal and off-diagonal vertices with the same meaning and function as in the SSE operator-loop method (illustrated in Fig. 61). Loops form according to the connectivity of the vertices, and also through the valence bonds at the boundaries, and can be flipped without changing the configuration weight. The sum over all possible loop configurations corresponds exactly to the formulation of the projector scheme in the pure valence bond basis (i.e., without using the spins at all) [47]. Using the spins allows for a more efficient sampling of the configurations. Expectation values are evaluated using loop estimators at the mid-point indicated with a vertical dashed line, and this is analogous to the improved estimators (discussed in Sec. 5.2.5) in the finite-temperature versions of the loop algorithm.

we discussed in the context of Krylov space methods in Sec. 4.2: By acting with a high power $H^\Lambda$ of the hamiltonian on an arbitrary state $|\Psi\rangle$, only the component of that state with the largest energy eigenvalue—normally the ground state (and if not this can be arranged by subtracting a constant from $H$)—survives in the limit $\Lambda \to \infty$, as demonstrated by Eq. (170). Expectation values of the form

$$\langle A \rangle = \frac{\langle \Psi| H^\Lambda A H^\Lambda |\Psi\rangle}{\langle \Psi | H^{2\Lambda} |\Psi\rangle},$$

(300)

where $|\Psi\rangle$ is a valence bond state or superposition (e.g., an amplitude product state), can be sampled using Monte Carlo simulations. In the original formulations of this approach [62, 47], $H$ is written as a sum of singlet projector operators (as in the Heisenberg antiferromagnet), or products of singlet operators (as in J-Q models), and strings of such operators are sampled (along with the valence bond configurations of $|\Psi\rangle$). The singlet projectors $S_{ij}$, defined in Eq. (21), lead to simple reconfigurations of bond pairs when acting on valence bond states. When acting on a valence bond, this operator is diagonal with eigenvalue 1,

$$S_{ab}(a,b) = (a,b),$$

(301)

while acting on a pair of different valence bonds leads to a simple reconfiguration of those bonds, with matrix element $1/2$;

$$S_{bc}(a,b)(c,d) = \frac{1}{2}(c,b)(a,d),$$

(302)

This can be shown easily by going back to the basis of $\uparrow$ and $\downarrow$ spins. Note the order of the indices within the singlets, which reflects consistently the convention corresponding to Marshall’s sign rule discussed above; $a, c$ are on sublattice $A$ and $b, d$ on sublattice $B$. These rules allow for a well-defined path integral (or SSE) like propagation of the states, and these propagation paths can be sampled using a Monte Carlo scheme.
In a more recent formulation of the valence bond projector method [50], the $\uparrow$ and $\downarrow$ spin basis is reintroduced, by expressing all singlets in terms of their sums over antiparallel spin pairs. This leads to an algorithm very similar to SSE and world line loop methods. Essentially, the periodic time boundaries used when simulating a system at a fixed temperature are cut open and replaced by two separate boundaries at which the valence bond states live. This is illustrated and discussed in more detail with a simple configuration for a four-site Heisenberg chain in Fig. 91.

The valence bond basis (or its translation to loop methods [244, 50]) has some unique aspects which makes it possible to access observables that are normally out of reach or difficult to calculate. In an extended valence bond basis for the triplet sector, one can study some properties of excited states [47, 49, 278]. One can also generalize the valence bond basis to include one [279] or several [278] unpaired spins, which is useful for studies of, e.g., the magnetization distribution in systems with unequal sublattice occupation of the spins. Simulations in the valence bond basis have also recently found applications in studies of entanglement entropy [23, 280, 281, 282, 283]. One can also extend valence bond projector methods to SU(N) spins [108] (including even non-integer $N$ generalizations [248]) and other related symmetry groups [282].

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