Tensor field theory with applications to quantum computing

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We discuss the successes and limitations of statistical sampling for a sequence of models studied in the context of lattice QCD and emphasize the need for new methods to deal with finite-density and real-time evolution. We show that these lattice models can be reformulated using tensorial methods where the field integrations in the path-integral formalism are replaced by discrete sums. These formulations involve various types of duality and provide exact coarse-graining formulas which can be combined with truncations to obtain practical implementations of the Wilson renormalization group program. Tensor reformulations are naturally discrete and provide manageable transfer matrices. Combining truncations with the time continuum limit, we derive Hamiltonians suitable to perform quantum simulation experiments, for instance using cold atoms, or to be programmed on existing quantum computers. We review recent progress concerning the tensor field theory treatment of non-compact scalar models, supersymmetric models, economical four-dimensional algorithms, noise-robust enforcement of Gauss’s law, symmetry preserving truncations and topological considerations.

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I. INTRODUCTION

Quantum field theory models on space or space-time lattices play an important role in our understanding of strongly interacting particles, nuclei, superconductivity and phase transitions. In high-energy and nuclear physics, lattice quantum chromodynamics (QCD) provides an ab-initio theory of strong interactions. In the QCD context, the lattice is a non-perturbative ultraviolet regularization which preserves local gauge invariance. As the lattice is not a physical feature, we need to approach the continuum limit where the lattice spacing is small compared to the physical length scales involved in the problem.

In the context of condensed matter and solid state physics, a physical lattice spacing of the order of a nanometer can be present, however, large correlation lengths appear near critical points and universal behavior independent of the microscopic details can be observed. It is also possible to create actual optical lattices in laboratories by trapping cold atoms in counter-propagating laser beams, and tune the interaction in order to quantum simulate lattice models with interactions similar to Hubbard models (Bloch et al., 2008). This can be called an analog computing method or a quantum simulation experiment. Again, it is possible to tune the parameters to reach universal behaviors related to quantum phase transitions with large correlation lengths.

More generally, we are getting better control in the manipulation of small quantum systems evolving in small Hilbert spaces and the idea of using physical quantum systems to study theoretical quantum models (Feynman, 1982) has generated many exciting developments (Georgescu et al., 2014). As bits—that can be either on or off—can be thought of as the basic building blocks of classical computers, one can envision qubits that can each be used as a two-dimensional Hilbert space as the building blocks of a quantum computer. If we want to use the $2^N$ dimensional Hilbert space provided by $N$ qubits to represent the Hilbert space of a quantum field theory problem we need to apply discretizations and truncations. Discretization of space can be achieved by the lattice approximation, while the discretization of continuous field integration can be done by using character expansion, as will be discussed extensively in this review, or by other methods (Alexandru et al., 2019a; Hackett et al., 2019; Jordan et al., 2014; Klco and Savage, 2019; Lamm et al., 2019).

General arguments (Lloyd, 1996) show that for local interactions, a quantum computer will reduce the computational effort, for problems like the real-time evolution, to a polynomial in the size of the system rather than an exponential for a classical computer.

From a purely theoretical point of view, studying models with a large number of strongly correlated degrees of freedom is very challenging. In order to deal with this situation, L. Kadanoff (Kadanoff, 1966) suggested to consider the average field or spin in cells of variable sizes often called “blocks”. The procedure is often called “blockspinning” and it played a crucial role in the development of the renormalization group (RG) ideas (Wilson and Kogut, 1974). Sometimes great theoretical intuitions can take a long time to be practically realized. Despite its visual appeal, the blockspinning procedure is not easy to implement numerically. It typically involves approximations that are difficult to improve.

Successful applications of the RG idea were made possible without requiring numerical implementations of the original blocking idea. A well-known example is the discovery of asymptotic freedom (Gross and Wilczek, 1973; Politzer, 1973) which initially relied on a one-loop calculation of the Callan-Symanzik beta function. Typically, the interplay between small and large energy scales is more easily seen in the momentum representation. However, if blocking in configuration space could be practically achieved, the computational cost grows like the logarithm of the volume. Below, we briefly explain the practical issues that have prevented the use of blocking for quantitative purposes and how new tensorial methods can be used to make progress in this direction.

A simple way of blocking consists in introducing 1 in the partition function in the following generic form:

$$\prod_{B} \int d\Phi_B \delta(\Phi_B - \sum_{x \in B} \phi_x) = 1,$$

(1)

where the blocks $B$ form a partition of the original lattice. For instance, on a three-dimensional cubic lattice, the blocks can be chosen as cubes with a linear size of two lattice spacings and contain eight sites. The $\phi_x$ are
the original lattice fields and \( \Phi_{B} \) the block fields which inherit new effective interactions after one performs the integration over the original fields. Conceptually, this sounds easy, however in practice it appears to be more complicated than the original problem. As an example, one can try to write a simple algorithm for the two-dimensional Ising model on a square lattice by replacing four spins in a \( 2 \times 2 \) square block by a single variable and write an effective energy function (or at least some effective measure) for the new block variables. The procedure becomes more intricate as we proceed, and finding the effective energy function is nontrivial (Liu et al., 2013).

For this reason, approximate procedures were developed such as the Migdal-Kadanoff approximation (Migdal, 1975), the approximate recursion formula (Wilson and Kogut, 1974) or other hierarchical approximations (Meurice, 2007), where no new interactions are generated by the blocking process. However, in these examples, the lack of reference to an exact procedure to handle the original model with localized interactions makes the systematic improvement of these approximations difficult. Similar issues appear in non-perturbative functional methods based on the momentum space representation (Berges et al., 2002), where the local potential approximation allows high-accuracy estimates of the critical exponents (Bervillier et al., 2007), but its improvement with methods such as the derivative expansion remains difficult (Bervillier, 2013). For Ising models, it is possible to deal with the proliferation of couplings generated by the blocking process by starting with the most general set of interactions (Kadanoff, 1975; Kadanoff and Houghton, 1975; Niemeijer and van Leeuwen, 1976). They introduce the identity in terms of probabilities \( P(\{\sigma'\}, \{\sigma\}) \) such that

\[
\sum_{\{\sigma'\}} P(\{\sigma'\}, \{\sigma\}) = 1, \tag{2}
\]

where \( \{\sigma'\} \) are new Ising spins associated with blocks. As we will discuss later, this special setup allows us to write formal expressions for the effective couplings as double partition functions and write RG equations. However, from a computational point of view the locality of the interactions is lost and additional assumptions are needed to proceed.

In contrast, reformulations of the partition function of classical spin models as the trace of a product of local tensors provides a new type of blocking procedure in configuration space called Tensor RG (TRG) (Gu et al., 2010; Gu and Wen, 2009; Levin and Nave, 2007; Nishino and Okunishi, 1996; Xie et al., 2012; Xie et al., 2009). The TRG is based on exact blocking formulas with numerically controllable truncations. The basic reason is that the TRG blocking separates neatly the degrees of freedom inside the block (which are integrated over), from those kept to communicate with the neighboring blocks (Meurice, 2013).

At early stages of the TRG development, Singular Value Decomposition (SVD) methods were used extensively. This is reviewed in Ref. (Efrati et al., 2014). Some of the SVD procedures can be simplified by using character expansions (Liu et al., 2013) when applied to most models studied in the context of lattice gauge theory (Bazavov et al., 2019, 2015; Butt et al., 2019; Denblyker et al., 2014; Kadoh et al., 2018, 2019, 2020; Kuramashi and Yoshimura, 2019; Liu et al., 2013; Nakamura et al., 2019; Shimizu and Kuramashi, 2014a, 2018; Takeda and Yoshimura, 2015; Unmuth-Yockey, 2019; Yoshimura et al., 2018; Yu et al., 2014; Zou et al., 2014). Tensorial methods are also used in the context of quantum gravity (Asaduzzaman et al., 2019; Dittrich et al., 2016; Perez, 2013).

In general, tensorial methods represent a new approach of lattice field theory that we will call Tensor Field Theory (TrFT). The “r” between T and FT stands for the last letter of the word tensor, or for the often used trace symbol Tr, and is used to avoid confusion with Topological Field Theory. TrFT can be used for purposes more general than the blocking procedure. In particular TrFT provides a very convenient, discrete, framework to perform quantum computations or simulations.

In this review article, we introduce TrFT for lattice models studied in the context of lattice gauge theory and report progress made for blocking and quantum computing, two “competing” methods that attempt to reduce the computing time logarithmically. The models targeted are introduced in Sec. II. We advocate a roadmap starting with the Ising model and culminating with QCD, which we call the “Kogut sequence” (Kogut, 1979, 1983). This sequence is sometimes called a “ladder” and has been followed successfully in situations where importance sampling methods such as the Metropolis algorithm are effective. Lattice QCD has become a very reliable precision tool to study the static properties of hadrons. As we are writing this article, we are roughly in the middle of the sequence. However, we hope that recent progress with four-dimensional theories (Kadok and Nakayama, 2019) could be combined with the methods that we describe to deal with gauge fields, fermions, and non-Abelian (non-commuting) symmetries in order to attempt calculations directly related to lattice QCD in the coming years.

In Sec. III, we discuss situations where importance sampling cannot be used and where quantum computations or simulations could provide alternate ways to perform computations. This includes unitary, real-time evolution and other situations where a sign problem is encountered. One important long-term goal with potential impact on the interpretation of high-energy collider data is doing ab-initio real-time calculations relevant to fragmentation processes and parton distribution functions. In other words, starting with lattice QCD, we would like to be able to perform calculations that would ultimately replace the use of event generators such as Pythia (Sjos-
Real-time evolution requires a quantum Hamiltonian. We provide a first look at the transfer matrix which connects smoothly the “classical” Lagrangian approach to the Hilbert space used in the Hamiltonian formalism. We discuss various types of dualities (geometrical and topological) that are often used together and taken for each other.

For the models in the Kogut sequence, the bosonic field variables and the symmetry groups are compact. General mathematical theorems, notably the Pontryagin duality (Pontryagin, 1939) and the Peter-Weyl theorem (Peter and Weyl, 1927), guarantee that functions over compact groups can be expanded in terms of discrete sums of representations. This is called the “character expansion” and was exploited to calculate strong coupling expansions (Balian et al., 1975) or introduce new variables on geometrically dual lattice elements (Savit, 1980).

The discreteness of the character expansion provides a natural starting point for building approximate formulations of lattice models suitable for quantum computing or quantum simulation experiments. The Ising model is an elementary example where the Hilbert space of the transfer matrix can be implemented with a set of qubits, the basic components of actual quantum computers which exist in a linear superposition of two states, |0⟩ and |1⟩, rather than being just “on” or “off” like the bits of a classical computer. For models with continuous fields, character expansions allow us to perform the “hard integrals” analytically without the need of approximate numerical discretizations which break the continuous symmetries. Demonstrating the power of the character expansion is one of the main goals of this review. In Sec. IV, we clarify the use of “classical” and “quantum” in various contexts.

Sec. V introduces the tensor reformulation for the Ising model. SVD, truncation and the TRG method are discussed in Sec. VI. Spin models with an O(2) symmetry or with discrete subgroups are discussed in Sec. VII. In Sec. VIII, we derive expressions for local tensors in the complex-field case. For the complex-field case.

In Sec. XI we present tensor formulations for models with fermionic degrees of freedom. In general fermions fit in well with the tensor (and discrete) approach thanks to the nilpotency of the Grassmann variables. In the section, various models that contain fermions such as pure fermions, gauged fermions, and fermions combined with scalars are discussed.

In Sec. XII we re-discuss the transfer matrix using the tensor formalism. Possible quantum simulations and computations inspired by TrFT are discussed in Sec. XIII. Recent TrFT developments regarding symmetries, topological solutions and quantum gravity are discussed in Sec. XIV.

II. LATTICE FIELD THEORY

A. The “Kogut sequence”: from Ising to QCD

In the early 70’s, QCD appeared as a strong candidate for a theory of strong interactions involving quarks and gluons. However the perturbative methods that provided satisfactory ways to handle the electroweak interactions of leptons failed to explain confinement, mass gaps and chiral symmetry breaking. A non-perturbative definition of QCD was needed. In 1974, K. Wilson proposed (Wilson, 1974) a lattice formulation of QCD where the SU(3) local symmetry is exact. As this four-dimensional model is fairly difficult to handle numerically, a certain number of research groups started considering simpler lattice models in lower dimensions and then increased symmetry and dimensionality. This led to a sequence of models, sometimes called the “Kogut ladder” which appears in J. Kogut’s review articles (Kogut, 1979, 1983), and later with small modifications in textbooks by A. Polyakov (Polyakov, 1987) and C. Itzykson with J. M. Drouffe (Itzykson and Drouffe, 1991).

The sequence is approximately the following:

1. $D = 2$ Ising model
2. $D = 3$ Ising model and its gauge dual
3. $D = 2$ O(2) spin and Abelian Higgs models
4. $D = 2$ fermions and the Schwinger model
5. $D = 3$ and 4 $U(1)$ gauge theory
6. $D = 3$ and 4 Non-abelian gauge theories
7. $D = 4$ lattice fermions
8. $D = 4$ QCD

This sequence should not be understood in a rigid way as if each step is necessary for the next step. For instance steps 3. to 5. could be interchanged and the problems involving fermions have very specific features that are not
easily compared to those involving only bosonic fields. The message that we want to convey is that there is an approximate roadmap that has proven to be very effective for the “classical” approach of lattice field theory in order to deal with static problems using importance sampling (Monte Carlo) methods. We advocate to follow a similar path to develop the quantum versions of these models and deal with real-time evolution and other problems not accessible with classical methods. The difference between quantum and classical is explained more precisely in Sec. IV.A. In addition, a similar path is being followed to develop numerical coarse-graining.

B. Classical lattice models and path integral

In this subsection we introduce lattice versions of classical field theory models. At this point, we would like to point out that while we will provide definitions of the fields used, notations, and acronyms/initials, more details on basic quantum field theory and lattice field theory can be found in textbooks and review articles e.g. (Itzykson and Drouffe, 1991; Kogut, 1979; Montvay and Münster, 1994; Peskin and Schroeder, 1995). We use a Euclidean time and treat space and time on the same footing. The metric is simply a Kronecker delta in D dimensions. We then discretize space and time. We use a D-dimensional (hyper) cubic Euclidean space-time lattice. The sites are denoted \( x = (x_1, x_2, \ldots, x_D) \), with \( x_D = \tau \), the Euclidean time direction. In lattice units, the space-time sites are labelled with integers. In the following, the lattice units are implicit. The links between two nearest neighbor lattice sites \( x \) and \( x + \mathbf{\mu} \) are labelled by \((x, \mathbf{\mu})\) and the plaquettes, the smallest squares on a square or (hyper) cubic lattice, delimited by four sites \( x, x + \mathbf{\mu}, x + \mathbf{\mu} + \mathbf{v} \) and \( x + \mathbf{v} \) are labelled by \((x, \mathbf{\mu}, \mathbf{v})\). By convention, we start with the lowest index when introducing a conventional circulation at the boundary of the plaquette. The total number of sites is denoted \( V \). Unless specified periodic boundary conditions are assumed. They preserve a discrete translational symmetry. If we take the time continuum limit, we obtain a quantum Hamiltonian formulation in \( D - 1 \) spatial dimensions.

In the continuum, the Lagrangian density for \( N \) real scalar fields with a \( O(N) \) global symmetry reads

\[
L^{(O(N))}_{\text{Euclidean}} = \frac{1}{2} \partial_\mu \tilde{\phi} \cdot \partial_\nu \tilde{\phi} \delta^{\mu\nu} + \lambda (\tilde{\phi} \cdot \tilde{\phi} - v^2)^2, \tag{3}
\]

with \( \tilde{\phi} \) a \( N \)-dimensional vector. Here \( \lambda \) is a coupling constant, whose size determines fluctuations of the \( |\tilde{\phi}| \) field around some value, \( v \). The potential has degenerate minima on a \( N \)-1 dimensional hyper-sphere \( S_{N-1} \) and a local maximum at \( \tilde{\phi} = 0 \). For \( N = 2 \), the low energy part of the potential has a shape reminiscent of the bottom of a wine bottle. The degenerate minima form a circle at the very bottom. We can study the small fluctuations about a given minimum on the circle. Note that the choice of a minimum breaks the \( O(2) \) symmetry. There are “soft” fluctuations along the circle that restore the symmetry and “hard” fluctuations in the radial direction.

We can extend this analysis for arbitrary \( N \). We have one massive mode (fluctuations in the symmetry breaking direction) with mass \( 2\sqrt{2} \lambda v \) and \( N - 1 \) massless modes (“Nambu-Goldstone” (NG) modes) which is the number of broken generators.

We can write the Euclidean action for the NG modes on a \( D \)-dimensional lattice with isotropic lattice spacing \( a \) as

\[
S_{\text{NLSM}} = \frac{1}{2} \sum_{x}^{D} D^{D-2}(\tilde{\phi}_{x+\mathbf{\mu}} - \tilde{\phi}_{x}) \cdot (\tilde{\phi}_{x+\mathbf{\mu}} - \tilde{\phi}_{x}). \tag{4}
\]

This model is called the nonlinear sigma model (NLSM). The constraint \( \tilde{\phi}_{x} \cdot \tilde{\phi}_{x} = v^2 \) (which enforces the “nonlinear” part of its name) can be expressed by introducing unit vectors: \( \tilde{\phi}_{x} = v \tilde{\sigma}_{x} \) such that

\[
\tilde{\sigma}_{x} \cdot \tilde{\sigma}_{x} = 1. \tag{5}
\]

Redefining \( a^{D-2}v^2 = \beta \), we get the simple action

\[
S_{\text{NLSM}} = \beta \sum_{x, \mu} (1 - \tilde{\sigma}_{x+\mathbf{\mu}} \cdot \tilde{\sigma}_{x}). \tag{6}
\]

These models are often called spin models as well. The first term in the action \( \beta \sum_{x, \mu} 1 \) is a constant that is often dropped. However, for large \( \beta \), the configurations with almost constant \( \sigma_{x} \) dominate the partition function and since under these circumstances \( \tilde{\sigma}_{x+\mathbf{\mu}} \cdot \tilde{\sigma}_{x} \approx 1 \) it is useful to subtract the constant in order to just keep the small fluctuations.

The case \( N = 1 \) is the well-known Ising model with \( \sigma_{x} = \pm 1 \). For \( N = 2 \), the terminology “planar model” or “classical XY model” is common and if we use the circle parameterization

\[
\sigma_{x}^{(1)} = \cos(\varphi_{x}), \text{ and } \sigma_{x}^{(2)} = \sin(\varphi_{x}), \tag{7}
\]

then

\[
\tilde{\sigma}_{x+\mathbf{\mu}} \cdot \tilde{\sigma}_{x} = \cos(\varphi_{x+\mathbf{\mu}} - \varphi_{x}). \tag{8}
\]

There is another class of models which breaks the \( O(2) \) symmetry in Eq. (8) into a discrete \( Z_{q} \) symmetry, i.e. the possible angles are restricted to those of the \( q^{th} \) roots of unity in the complex plane. They are called the \( q \)-state clock models. These models have the same action from Eq. (6) with the identification of the angles as being discrete,

\[
\varphi_{x} = \frac{2\pi n_{x}}{q} \tag{9}
\]

with \( n_{x} = 0, 1, 2, \ldots, q - 1 \). With this identification, the \( O(2) \) model emerges as the \( q \to \infty \) limit of the \( q \)-state
clock models, and the Ising model is simply the $q = 2$ model.

For $N = 3$, the symmetry becomes non-Abelian and the model is sometimes called the “classical Heisenberg model”. In the large-$N$ limit, the model becomes solvable if we take the limit in such a way that $N/\beta(N) = \lambda$ remains constant (Coleman et al., 1974).

It is instructive to rewrite the O(2) model using the complex form

$$\Phi_x = e^{i\varphi_x}. \quad (10)$$

Dropping the constant term, the O(2) action reads

$$S_{O(2)} = \frac{\beta}{2} \sum_{x, \mu} (\Phi_{x+\mu} - \Phi_x) \cdot (\Phi_{x+\mu} - \Phi_x)^*$$

$$\to -\frac{\beta}{2} \sum_{x, \mu} (\Phi_x^* \Phi_{x+\mu} + \text{h.c.})$$

$$= -\frac{\beta}{2} \sum_{x, \mu} \cos(\varphi_{x+\mu} - \varphi_x). \quad (11)$$

The O(2) model has a global symmetry

$$\varphi_x \to \varphi_x + \alpha. \quad (12)$$

With the complex notation, this transformation becomes

$$\Phi_x \to e^{i\alpha} \Phi_x. \quad (13)$$

We would like to promote this symmetry to a local one

$$\Phi_x \to e^{i\alpha_x} \Phi_x, \quad (14)$$

i.e. one that is site dependent. This can be achieved by inserting a phase $U_{x, \mu}$ between $\Phi_x^*$ and $\Phi_{x+\mu}$ which transforms like

$$U_{x, \mu} \to e^{i\alpha_x} U_{x, \mu} e^{-i\alpha_{x+\mu}}. \quad (15)$$

The procedure can be extended for arbitrary N-dimensional complex vectors $\Phi_x$ with a local transformation involving a $U(N)$ matrix $V_x$:

$$\Phi_x \to V_x \Phi_x. \quad (16)$$

In addition, we can also introduce $U(N)$ matrices $U_{x, \mu}$ transforming like

$$U_{x, \mu} \to V_x U_{x, \mu} V_{x+\mu}^\dagger. \quad (17)$$

The action

$$S_{U(N)} = -\frac{\beta}{2} \sum_{x, \mu} (\Phi_x^\dagger U_{x, \mu} \Phi_{x+\mu} + \text{h.c.}) \quad (18)$$

has a local $U(N)$ invariance which we call gauge invariance. If we consider two successive links in positive directions, then the local transformation at the middle site cancels and

$$U_{x, \mu} U_{x+\mu, \nu} \to V_x U_{x, \mu} U_{x+\mu, \nu} V_{x+\mu, \nu}^\dagger. \quad (19)$$

If the second link goes in the negative direction, we use the Hermitian conjugate and a similar property holds

$$U_{x, \mu} U_{x+\mu, \nu}^\dagger \to V_x U_{x, \mu} U_{x+\mu, \nu}^\dagger V_{x+\mu, \nu}^\dagger. \quad (20)$$

We can pursue this process for an arbitrary path connecting $x$ to some $x_{\text{final}}$. The transformation on the right will be $V_{x_{\text{final}}}$. If we close the path and take the trace, we obtain a gauge-invariant quantity. We call these traces of products of gauge matrices over closed loops “Wilson loops” (Wilson, 1974). In the case where the loop goes around the imaginary time direction, we often call it a “Polyakov loop” (Polyakov, 1978).

On a square, cubic or hypercubic lattice, the smallest path that gives a non-trivial Wilson loop is a square. We call this square a plaquette. Claude Itzykson coined this terminology after Ken Wilson’s seminar in Orsay in 1973. The corresponding matrix is

$$U_{\text{plaquette}} = U_{x, \mu\nu} = U_{x, \mu} U_{x+\mu, \nu} U_{x+\mu, \nu}^\dagger U_{x, \mu}^\dagger \quad (21)$$

The simplest gauge-invariant lattice model has an action, called Wilson’s action:

$$S_{\text{Wilson}} = \beta_{\text{pl}} \sum_{\langle x, \mu\nu \rangle} \left[ 1 - \frac{1}{2N} \text{Tr} \left[ U_{x, \mu\nu} + \text{h.c.} \right] \right]. \quad (22)$$

where $\sum_{\langle x, \mu\nu \rangle}$ indicates a sum over all plaquettes. Here each $U_{x, \mu}$ is related to the vector potential, or gauge field, in the continuum theory through $U_{x, \mu} = e^{iA_{x, \mu}}$. In the Abelian case ($N = 1$), the matrix reduces to a phase

$$U_{x, \mu} = e^{iA_{x, \mu}}, \quad (23)$$

and there is no need to take the trace.

Another generalization of the $N = 1$ expression of the complex phase given in Eq. (10), consists of replacing $\Phi_x$ by a $SU(N)$ matrix $U_x$. This is called the principal chiral model,

$$S_{\text{PCM}} = -\frac{\beta}{2} \sum_{x, \mu, \nu} \left[ \text{Tr} \left[ U_{x, \mu}^\dagger U_{x, \mu} \right] + \text{h.c.} \right]. \quad (24)$$

This model has a global rotational symmetry under the $U(N)$ group, such that $U_{x, \mu} \to U_{x, \mu}' = V U_{x, \mu} V^\dagger$, just as in Eq. (17) in the case of a uniform $V$ in all of space-time.

We can also consider the $N = 1$ case for Eq. (3) when the group is no longer compact, i.e. $\phi_x$ can take on the values of any real, or complex, number. The action on the lattice is then,

$$S_{\text{scalar}} = \sum_x \left( \frac{1}{2} \sum_{\mu, \nu = 1}^D |\phi_{x+\mu} - \phi_{x\nu}|^2 + \lambda (|\phi_{x\nu}|^2 - \nu^2)^2 \right) \quad (25)$$

or equivalently,

$$S_{\text{scalar}} = \sum_x \left\{ \frac{1}{2} \sum_{\mu, \nu = 1}^D |\phi_{x+\mu} - \phi_{x\nu}|^2 - \frac{\mu_0^2}{2} |\phi_{x\nu}|^2 + \frac{\lambda'}{4} |\phi_{x\nu}|^4 \right\} \quad (26)$$
with the substitution $\mu^2/2 = 2\nu^2$ and $\chi' = 1$, and an overall constant is ignored.

Besides scalar fields, we will also consider fermionic fields (or Grassmann fields on the lattice). In the case of free fermions, we can write down a lattice action using a straightforward discretization due to Wilson,

$$S_{WD} = \sum_x \bar{\psi}_x (D\psi)_x$$  \hspace{1cm} (27)

where the Wilson-Dirac operator is defined by

$$D_{xx'} = (am + rD) \delta_{xx'} + \frac{1}{2} \sum_{\mu=1}^D \{ (r - \gamma_\mu) \delta_{xx',\mu} + (r + \gamma_\mu) \delta_{xx',\mu} \}.$$  \hspace{1cm} (28)

where $\gamma_\mu$ are the gamma matrices in $D$ dimensions, and $r$ is the “Wilson parameter” to control species doubling. $\psi_x$ and $\bar{\psi}_x$ are multi-component Grassmann variables, this means they follow different rules compared to commuting variables. Grassmann variables anti-commute, as opposed to commute for typical scalar variables. There is another formulation of lattice fermions, where the different components of the fermion fields are located at different lattice sites, called staggered fermions. This comes about from a transformation which mixes the fermion components and space-time components (Kogut and Susskind, 1975). The action for free fermion fields is given by,

$$S_F = \frac{1}{2} \sum_{x=1}^N \sum_{\mu=1}^D \eta_{x,\mu} [\bar{\psi}_x U_{x,\mu} \psi_{x+\mu} - \bar{\psi}_{x+\mu} U_{x,\mu}^\dagger \psi_x]$$  \hspace{1cm} (29)

where

$$\eta_{x,\mu} = (-1)^{\sum_{\nu < \mu} x_\nu}$$  \hspace{1cm} (30)

with $x_\nu$ the coordinate in the $\nu$th direction. One can include gauge fields in a gauge-invariant manner by inserting $U_{x,\mu}$ like,

$$S_F = \frac{1}{2} \sum_{x=1}^N \sum_{\mu=1}^D \eta_{x,\mu} [\bar{\psi}_x U_{x,\mu} \psi_{x+\mu} - \bar{\psi}_{x+\mu} U_{x,\mu}^\dagger \psi_x].$$  \hspace{1cm} (31)

These models with appear in the rest of the review and will be very briefly reintroduced in each section.

C. Physical applications

The sequence of models described in Sec. II.A is designed to handle lattice QCD which is so far our best definition of the theory describing strongly interacting particles observed in a large number of experiments. Some of the models discussed in Sec. II.B are also studied in condensed matter. For instance, the $O(2)$ model with a chemical potential can be seen as an effective theory for the Bose-Hubbard model (Sachdev, 2001). $U(1)$ gauge theories with either scalar of fermion fields are studied in the context of superconductivity (Herbut, 2007). Tight-binding approximations for solids leads to interesting lattice models, for instance a sheet of graphene can be described with fermions on a hexagonal lattice having a dispersion relation similar to the one for a massless Dirac fermion at half-filling. (Castro Neto et al., 2009).

In the continuum limit, models compatible with relativistic invariance, local gauge invariance and renormalizability have a very small number of free parameters provided that the interactions among the fields and their first derivatives are kept local. The standard model of electroweak and strong interactions has only 18 free parameters if we ignore the QCD vacuum angle and the additional parameters related to the masses and mixing of the neutrinos. It is not possible to tweak the theory each time a new experiment is completed. This makes the standard model a very predictive theory. In the lattice formulation, the local interactions involving derivatives in the continuum are replaced by interactions involving fields located on neighboring sites, links and plaquettes and can be considered quasi-local.

D. Computational methods beyond perturbation theory

Besides the RG methods mentioned in the Introduction, lattice models have been studied with a variety of analytical and numerical methods. For example, expansions at small and large coupling for spin and gauge models (Itzykson and Drouffe, 1991; Kogut, 1979, 1983; Parisi, 1998; Polyakov, 1987) have been investigated extensively. As power series, they can be used to check numerical calculations in their respective limits, however, it seems hard to capture the non-perturbative effects such as the generation of a dynamical mass gap in the continuum limit. After Wilson’s original proposal (Wilson, 1974) a very suggestive Hamiltonian picture was developed by Kogut and Susskind (Kogut and Susskind, 1975), however it became clear that the size of the Hilbert space would make numerical calculations impractical.

In many practical situations, the path integral formulation uses a real Euclidean action $S_E$. This allows importance sampling. A typical way to proceed is to start with a random field configuration and then apply some random changes on this configuration. If the new configuration has a lower action, it is accepted. If the new configuration has an action larger by $\delta S_E$, then it is accepted with a probability $\exp(-\delta S_E)$. The fact that the fields are only connected to a few neighboring fields makes the calculation of $\delta S_E$ easy and the exploration of the important configurations controllable.

Mathematician are often amazed that it is possible to obtain very reliable results with this method (Villani,
Several decades after its inception, numerical lattice gauge theory has become an area of research where corrections on the order of a few percent are considered important. For instance, in heavy flavor physics the determination of the mixing angle $|V_{ub}|$ is quoted as $(3.72\pm0.16)10^{-3}$ (Bailey et al., 2015). This result was obtained using ensembles of lattices of various lattice spacings and sizes, some of them as large as $64^3 \times 192$. For a review of averages of numerical results with estimated errors see, for instance, the Flavour Lattice Averaging Group (FLAG) summary (Aoki et al., 2020).

### III. QUANTUM COMPUTING

#### A. Situations where importance sampling fails

The numerical successes of lattice gauge theory can be linked to the fact that when the action of the Euclidean-time path integral $S_E$ is real, importance sampling works remarkably well for the selection of configurations with a Boltzmann distribution $\exp(-S_E)$. However, if we return to real time or introduce a chemical potential that makes the action complex, this powerful tool becomes ineffective.

Real-time evolution requires a Hamiltonian acting on a Hilbert space. This can be done by noticing that the Euclidean path-integral can be recast as the trace of a transfer matrix (Fradkin and Susskind, 1978; Kogut, 1979; Luscher, 1990; Wilson and Kogut, 1974). With generic notations for a lattice model with $N_r$ sites in the Euclidean time direction

$$Z = \int \mathcal{D}\Phi \ e^{-S[\Phi]} = \text{Tr}(\mathcal{T}^{N_r}).$$

(32)

If the lattice spacing $a_r$ in the Euclidean time direction is small compared to the physical time scales involved, we have

$$\mathcal{T} \propto e^{-a_r H}.$$  

(33)

For $N_r$ large enough, the use of Euclidean time provides a projection in the low energy sector of the Hilbert space. This property remains effective if we insert operators that create and destroy states with non-trivial quantum numbers.

Analysis of the exponential decays of correlations as the Euclidean time is increased is a standard tool to extract masses and form factors for momenta that are small compared to the lattice cutoff. On the other hand, real-time evolution and a large Hilbert space involving states with large momenta are needed to describe aspects of hadron fragmentation in real space-time and deep-inelastic scattering (Bauer et al., 2019; Lamm et al., 2020; Mueller et al., 2020). The real-time evolution operator does not provide a positive measure nor a projection onto a small Hilbert space. Another situation where sampling methods are challenged is the construction of interpolating operators for nuclei as the number of Wick contractions grows rapidly with the number of light quarks involved (Detmold and Orginos, 2013).

One could consider the possibility of abandoning the Lagrangian formulation and directly considering the Kogut-Susskind Hamiltonian (Kogut and Susskind, 1975) for QCD and/or the standard many-body formalism in condensed matter and nuclear physics (Fetter and Walecka, 2003) where the Hilbert space is generated by creation operators on a Fock-space vacuum. For the sake of argument, let us consider the simple case of $N$ conjugate pairs of fermionic creation and annihilation operators. They generate a Hilbert space of dimensions $2^N$. The exponential growth of this number with $N$ rapidly restricts our ability to store the matrix elements of operators using classical computers. Computations involving spatial lattices with $64^3$ sites as made possible in the Lagrangian formulations would be completely out of the question if we had to setup a corresponding quantum Hilbert space with existing classical computers.

#### B. Computing with qubits or cold atoms?

The building blocks of an ordinary (classical) computer memory are bits taking the values 0 or 1. For instance, in some designs of Dynamic Random Access Memory (DRAM) devices, this is achieved by using small capacitors that are either on or off. The typical units of capacitance are femtofarads with voltages of the order of 1V. These capacitors store a few thousand electrons and can be charged or discharged in times of order $10^{-15}$ sec. It is interesting to consider the limit of miniaturization where electrons, atoms or photons could be manipulated individually and where the peculiarities of the quantum behavior become prominent (Lloyd, 1996).

Following the physical examples of the electron spin or the photon polarization, one could envision some ideal and generic quantum system, where the on/off concept for bits is replaced by the linear superposition of two states. This basic unit of quantum computing is often called a qubit (Schumacher, 1995) and can be represented as

$$|\text{qubit}\rangle = \alpha_0 |0\rangle + \alpha_1 |1\rangle,$$

(34)

with two complex numbers $\alpha_0$ and $\alpha_1$ such that $|\alpha_0|^2 + |\alpha_1|^2 = 1$. A set of $N$ qubits spans a Hilbert space of
dimension $2^N$. If we use a classical computer to apply a dense unitary matrix representing the real time evolution on an arbitrary state, we need to perform on the order of $2^{2N}$ arithmetic operations. On the other hand, if the Hamiltonian is such that any qubit is only connected to a restricted number of other qubits which is fixed by the dimensionality of space and the internal symmetries, then it is possible to design a method that performs the evolution with a number of quantum manipulations which only scales polynomially with $N$ (Lloyd, 1996). This will be discussed in Sec. III.F. Nowadays, commercial quantum computers provide sets of qubits and the possibility of preparing, evolving and measuring quantum states.

More generally, the idea that quantum devices could be used to perform computations for quantum problems involving many degrees of freedom is very appealing (Feynman, 1982). Physical systems involving cold atoms (see (Bloch et al., 2008) for a review of the early developments) or trapped ions (see (Leibfried et al., 2016) for a recent example) can be used to mimic the behavior of simplified many-body models such as various types of spin chains or Hubbard models.

C. From Euclidean transfer matrices to Hilbert spaces

For the lattice models introduced in Sec. II.B, the Euclidean time was treated on equal footing with the $D - 1$ spatial dimensions. In order to discuss real-time evolution, we first need to single-out the time direction. Evolution then occurs along this direction according to a transfer matrix. The key ingredient then is to connect the Lagrangian formulation to the Hamiltonian formalism and to identify the Hilbert space from the transfer matrix introduced in Eq. (32). The general idea is to organize the partition function sums or integrals into operations performed on successive time slices. This can be accomplished in different ways. A general procedure to construct the transfer matrix of lattice models in configuration space is discussed in (Luscher, 1990). A “dual” method based on character expansions which are at the heart of TrFT are discussed in Sec. XII. We will illustrate these two possibilities with examples.

This simplest possible example is the one-dimensional Ising model,

$$ S_{\text{Ising}} = \beta \sum_{\tau} \left( 1 - \sigma_{\tau+1} \sigma_{\tau} \right) $$

with partition function

$$ Z = \sum_{\{\sigma\}} e^{-S}. $$

This is a product of exponentials which each share one spin variable with the next factor. Then we can write the partition function as

$$ Z = \text{Tr}[T^{N_{\tau}}] $$

with

$$ T_{\alpha \alpha'} = \exp \left\{ -\frac{\beta}{2} \left( \sigma_{\tau+1}^{(\alpha)} - \sigma_{\tau}^{(\alpha')} \right)^2 \right\} $$

and $\sigma^{(\alpha)} = 1, -1$ for $\alpha = 0, 1$, respectively. Along the diagonal of the transfer matrix we see only unity. On the off-diagonal, a spin flip comes with weight $e^{-2\beta}$. Then to leading order in the temporal lattice spacing, $T \simeq 1 - a_{\tau} \hat{H} + \ldots$ which allows us to identify

$$ \hat{H} = -h_x \hat{\sigma}^x, $$

with $h_x \equiv e^{-2\beta}/a_{\tau}$ and $\hat{\sigma}^x$ the $x$-Pauli matrix. For this case, to extract a Hamiltonian from the original Lagrangian formulation, we required that the coupling, $\beta$, goes to infinity to match the temporal lattice spacing, $e^{-2\beta} \propto a_{\tau}$.

In the above we found the Hamiltonian and Hilbert space in configuration space; however, one can use a “dual” method as well by expanding the original Boltzmann weights. Consider the action for the Ising model in Eq. (35) with the constant dropped,

$$ S_{\text{Ising}} = -\beta \sum_{\tau} \sigma_{\tau+1} \sigma_{\tau}. $$

The Boltzmann weight can be expanded in the form,

$$ T_{\alpha \alpha'} = e^{\beta \sigma_{\tau+1}^{(\alpha)} \sigma_{\tau}^{(\alpha')}} $$

$$ = \cosh(\beta) \sum_{n=0}^{1} \left( \sigma_{\tau+1}^{(\alpha)} \sigma_{\tau}^{(\alpha')} \right)^n \tanh^n(\beta). $$

which is simply the Euler identity for imaginary angles. Ignoring the factor out front as it does not affect the Hamiltonian, in this form we can easily do the summation over the values of $\alpha$ at all lattice sites. The transfer matrix becomes a diagonal matrix with matrix elements labeled by the integers $n$, from each Boltzmann factor,

$$ T_{nn'} = \begin{pmatrix} 1 & 0 \\ 0 & \tanh(n \beta) \end{pmatrix}. $$

In the literature, the integers $n$ are often called dual variables or characters indices. The terminology is discussed more systematically in Sec. III.D. To recover a Hamiltonian, the transfer matrix must have the form $1 - a_{\tau} \hat{H} + \ldots$ for small times. This is found by taking $\beta \rightarrow \infty$, and recalling $\tanh \beta = 1 - 2e^{-2\beta} + \ldots$ for large $\beta$. Then the Hamiltonian in these new variables takes the form,

$$ \hat{H} = h_x (1 - \hat{\sigma}^z) $$
with $h_z \equiv e^{-2\beta}/a_z$, and $\sigma_z^x$ the $z$-Pauli matrix. These are two distinct procedures which give Hamiltonians in the time continuum limit; one, in the original configuration variables, and the second in the dual variables.

The transfer matrix of the Ising model in higher dimensions can be constructed in a similar manner (Kaufman, 1949). In higher dimensions there are now two types of interactions. If we consider a particular time slice, we can first collect all the time links connected to the next time slice, each with a representation given in Eq. (35).

In addition, we have all the spatial links in the time slice with nearest-neighbor interactions. We can represent any spin configuration in a time slice as an element of a tensor product of eigenstates of the Pauli matrix $\sigma_z^x$:

$$\{\text{configurations}\} = \bigotimes_x |\pm 1\rangle_x. \tag{46}$$

We can introduce operators $\sigma_z^x$ or $\sigma_z^x$ acting on this Hilbert space that can be identified with a set of qubits. Following (Kaufman, 1949), we can collect the two types of interactions in two matrices. Using the relation

$$e^{\delta \sigma_z^x} = (2 \sinh(2\beta))^{1/2} (\exp(\beta \sigma_z^x))^{\alpha \alpha'}, \tag{47}$$

where $\bar{\beta}$ is the dual inverse temperature introduced by Kramers and Wannier (Kramers and Wannier, 1941) which satisfies the relation $\tanh(\bar{\beta}) = e^{-2\beta}$, the contribution of time links can be summarized with the matrix connecting the configuration of the two time slices

$$\mathcal{V}_1 = (2/\sinh(2\bar{\beta}))^N \delta^{D-1/2} e^{\beta \sum_x \sigma_z^x}. \tag{48}$$

On the other hand, the spatial links can be recast in the diagonal matrix

$$\mathcal{V}_2 = e^{\beta \sum_x j \sigma_z^x \sigma_z^{x+i}}. \tag{49}$$

We can now write the transfer matrix as

$$\mathcal{T} = \mathcal{V}_2^{1/2} \mathcal{V}_1 \mathcal{V}_2^{1/2}, \tag{50}$$

where the matrix indices label the spatial configurations. Geometrically, the Hilbert space is located on the time slices. Alternatively, we can work in a dual representation where the $\sigma_z^x$ are diagonal

$$\tilde{\mathcal{T}} = \mathcal{V}_1^{1/2} \mathcal{V}_2 \mathcal{V}_1^{1/2}, \tag{51}$$

where the matrix indices label sets of group characters. Geometrically, the Hilbert space is located between the time slices. Graphical illustrations of this situation will be provided in Sec. XII. The construction generalizes easily for finite Abelian groups and in a non-trivial way for continuous and compact Abelian groups. The advantage of using the second (dual) representation is that it remains discrete and as we will see in Sec. XIV.A, it preserves the symmetry when truncations are applied (Meurice, 2019, 2020a).

### D. Topological and geometrical dualities

In Sec. III.C we used the concept of duality in two occasions. The first was the relation between $\beta$ and $\bar{\beta}$ which interchanges their low and large values regimes. The second occasion was the discussion of the two ways to represent the transfer matrix. In addition, in Secs. VII.C and IX.C, we will use the geometrical duality reviewed in (Savitt, 1980). Duality is a general concept used in many branches of mathematics. According to Atiyah (Atiyah, 2007) duality gives “two different points of view of looking at the same object”. In the following we clarify the various usages of the concepts in the rest of this review.

An important notion of duality is the so-called “Pontryagin duality” (Pontryagin, 1939) coming in the study of topological groups. It relates an Abelian group and its characters (for instance, Fourier modes). It states that if the former is compact, the later is discrete and vice-versa. The simplest situation is a finite group which is compact and discrete. In the case of finite cyclic Abelian groups the characters form a finite group which is isomorphic to the group itself (Serre, 1973). A simple example is $\mathbb{Z}_q$ the additive group of integers modulo $q$. If $x$ denotes an element of $\mathbb{Z}_q$, the characters have the form

$$\chi_k(x) = \exp \left( \frac{2\pi i}{q} kx \right), \tag{52}$$

and clearly satisfy the character property

$$\chi_k(x + x') = \chi_k(x) \chi_k(x'). \tag{53}$$

The product of two characters is another character

$$\chi_k(x) \chi_{k'}(x) = \chi_{k+k'}(x), \tag{54}$$

and one sees that they also form a $\mathbb{Z}_q$ group. They obey the orthogonality relations

$$\frac{1}{q} \sum_{x=0}^{q-1} \chi_k(x) \chi_{k'}(x) = \delta_{k,k'}, \tag{55}$$

and

$$\frac{1}{q} \sum_{k=0}^{q-1} \chi_k(x) \chi_{k'}(x') = \delta_{x,x'}. \tag{56}$$

A simple example of a continuous (nondiscrete) and compact group is $U(1)$, the multiplicative group of complex numbers with modulus one. The group properties can be reformulated in an additive manner by introducing the phases $z = e^{i\varphi}$ which are added modulo $2\pi$. Topologically it is a circle which is a compact manifold. Its characters are discrete and labelled by the integers. They are the usual Fourier modes $e^{i m \varphi}$. The orthogonality relations appear in an asymmetric way

$$\int_{-\pi}^{\pi} \frac{d\varphi}{2\pi} e^{i m \varphi} (e^{i n \varphi})^* = \delta_{m,n}. \tag{57}$$
\[ \sum_{n=-\infty}^{\infty} e^{i\varphi} (e^{i\varphi'})^* = 2\pi \sum_{m=-\infty}^{\infty} \delta(\varphi - \varphi' + 2\pi m) \] (58)

In practice, a deep understanding of the mathematical statements made above is not necessary and we just need to remember a few character expansions. For the Ising models we have functions over the multiplicative group \( \sigma = \pm 1 \), and we can recall the expansion from Sec. III.C as a character expansion

\[ \langle \sigma' | \| \sigma \rangle = \exp(\beta \sigma' \sigma) \]
\[ = \frac{1}{2} \sum_{n=0}^{1} \lambda_n(\beta)(\sigma' \sigma)^n \]
\[ = \cosh(\beta) + \sigma' \sigma \sinh(\beta). \] (60)

A similar representation can be obtained for the \( \mathbb{Z}_q \) spin models. If we replace the discrete angle variables \( \frac{2\pi}{q} x \) by continuous ones \( \varphi \), we obtain a “matrix” with continuous elements which can be calculated using the standard Fourier transform

\[ \langle \varphi' | \| \varphi \rangle = \exp(\beta \cos(\varphi' - \varphi)) \]
\[ = \sum_{n=-\infty}^{\infty} I_n(\beta) \exp(in(\varphi' - \varphi)), \] (62)

where \( I_n(\beta) \) is the modified Bessel function of order \( n \). We will show in Sec. VII.B that finite versions of this expansion hold for the finite \( \mathbb{Z}_q \) subgroups.

Generalizations of Pontryagin duality to compact non-Abelian groups appear in the Peter-Weyl theorem (Peter and Weyl, 1927). As an example, this translates into expansions in spherical harmonics for problems involving the \( O(3) \) symmetry.

From the point of view of quantum computing, we see that using compact fields guarantees that we can replace the continuous integrals by discrete sums. Our strategy is to associate the indices of these sums with quantum states. One important aspect of TrFT is that when the fields are compact, we don’t need to discretize the integrals using numerical approximations. Instead, we can use characters expansions as Eq. (62) where some integrals have been done exactly and result in Bessel functions that we can input with any desired accuracy. In other words, the difficult part of the classical path-integral approach can be done efficiently with classical methods. After that, the original integrals reduce to orthogonality relations and can be performed exactly.

Another notion of duality is of a geometrical nature. It is related to the Levi-Civita tensor \( \epsilon^{i_1 \cdots i_d} \). Its meaning is dimension-dependent and relates objects of dimension \( d \) to objects of dimension \( D - d \). For instance, in \( D = 4 \), a dual field-strength tensor with two indices is obtained by contracting the original field-strength tensor with the Levi-Civita tensor. This duality transformation interchanges the electric and magnetic fields and reduces to the identity when repeated twice. It also relates sites to four-dimensional hypercubes, and plaquettes to plaquettes. In \( D = 3 \) it relates the field-strength tensor to a divergenceless pseudovector \( \epsilon^{ijk} \partial_j A_k \), links to plaquettes and sites to cubes.

The various notions of duality are often used simultaneously. A common example is the “dual formulation” of the Ising model. As we will explain in Sec. V.B, the new set of indices from Eq. (60) which is a consequence of Pontryagin duality, leads to a representation in terms of paths. We can then try to represent these paths as the boundaries of surfaces, which brings the geometrical duality and new “dual variables” together (Savit, 1980).

There are also occasions where the phrase “dual variables” has become associated with generic integer fields, regardless of their origin or their relation to the concepts of duality mentioned above (Bruckmann et al., 2016, 2015; Gattringer et al., 2018b, 2015a; Marchis and Gattringer, 2018). These integer fields can arise from a Taylor series of the Boltzmann weight, e.g. in the case of the Ising model,

\[ e^{\beta \sigma \sigma'} = \sum_{q=0}^{\infty} \frac{\beta^q}{q!}(\sigma \sigma')^q. \] (63)

This expansion associates a natural number with the links of the lattice, similar to the character expansion from before. Also just as before, this expansion allows one to perform the path-integral sums over the \( \sigma \) fields, leaving one with a theory of constrained, positive, integer fields on the links of the lattice. However, instead of two values, i.e. \( n = 0, 1 \), these integers can take on an infinite number of values. So while this expansion accomplishes similar feats, it can be seen as less economical parameterization of the model. Of course, the character expansion from before can be found within the Taylor expansion by summing the even and odd integers, respectively, leaving one with two terms (0 and 1).

E. Real-time evolution with qubits

We are now ready to discuss the possibility of using quantum devices to represent states of the Hilbert space emerging from the transfer matrix construction and to design methods to apply unitary transformations corresponding to the real time evolution. For the sake of definiteness, we assume that we have at our disposal a set of qubits. For the spin Ising model, the construction of the transfer matrix leads to a Hilbert space with an obvious qubit structure given by Eq. (46). The method to take the time continuum limit and identify a Hamiltonian using Eq. (32) is well-known (Fradkin and Susskind, 1978; Kogut, 1979); we deform the original transfer matrix by
increasing $\beta$ in the time direction (which makes the dual value $\tilde{\beta}$ small) and decreasing $\beta$ in the spatial directions. The arguments of the exponentials in $\mathcal{V}_1$ and $\mathcal{V}_2$ become infinitesimal and provide the two non-commuting pieces of the “quantum Hamiltonian”. The role played by $\mathcal{V}_1$ is quite special in the intuitive picture that we are trying to convey: it only acts on “single qubits” without connecting them together. Consequently, working in the representation where $\sigma_\mathbf{x}$ is diagonal is a good starting point. Next, we can “turn on” the terms in $\mathcal{V}_2$. At lowest order in the time lattice spacing they only connect qubits which are nearest neighbours. We will show in Sec. III.F that this type of situation permits quantum computation in a time scaling polynomially with the size of the system (Lloyd, 1996).

Character expansions and TrFT provide the natural tools to perform similar constructions for the models presented in Sec. II.B. A first step consists in isolating “building blocks” which are localized in space and have a very simple real-time evolution. The models have interactions associated with links and plaquettes. As a first approximation we set the interactions on spatial links and space-space plaquettes to zero.

For spin models, this results in a collection of $N_s^{D-1}$ isolated one-dimensional spin models. These isolated models are the building blocks. They are solvable and it is easy to calculate their evolution at real time. For gauge models, we have in addition, electric degrees of freedom which can be associated with the spatial links of a given time slice of the lattice. They are required to satisfy a constraint called “Gauss’s law”, however, when this condition is satisfied, the real-time evolution in the isolation limit described above is straightforward as we will explain in Sec. XII.B. The Hilbert space of the isolated building blocks depend on the model considered. For the Ising spin models, a single qubit is all we need. For models with continuous symmetries, the exact treatment requires an infinite Hilbert space, however, small size truncations provide good approximations and preserve the symmetries of the models (Meurice, 2019, 2020a). This is a very attractive feature of TrFT which is discussed in Sec. XIV.A. Having setup a finite Hilbert space with isolated building blocks, our next step is to restore the interactions associated with the spatial links and the space-space plaquettes. This will be done for a variety of models in Sec. XII. Independently of the model-specific aspect of this procedure, it is clear that each building block is only connected to a limited numbers of other building blocks. For instance, for a spin model, there are two connections for each spatial direction. This quasilocality is crucial to implement real-time evolution with a quantum computer.

F. Lloyd-Trotter product formula

An important motivation for using a quantum computer is to calculate the real-time evolution for systems with many degrees of freedom having quasilocal interactions in the sense discussed in Sec. III.E. Ideally the time to perform computations should scale polynomially with the size of the system rather than exponentially. A general argument leading to these conclusions has been put forward by S. Lloyd in (Lloyd, 1996) where he states that: “Feynman’s 1982 conjecture, that quantum computers can be programmed to simulate any local quantum system, is shown to be correct”. The proof is based on the basic idea behind the Trotter product formula (Reed and Simon, 1980), namely that for two non-commuting operators $\hat{A}$ and $\hat{B}$ and sufficiently small $\epsilon$

$$e^{i\epsilon(\hat{A}+\hat{B})} \simeq e^{i\epsilon\hat{A}}e^{i\epsilon\hat{B}} + \mathcal{O}(\epsilon^2).$$

In the standard construction of the path integral in quantum mechanics, it is applied to the kinetic and potential energy, but it can also be applied to all the quasilocal parts of the Hamiltonian.

The argument goes as follows (Lloyd, 1996). Consider a system composed of $N$ variables with Hamiltonian

$$\hat{H} = \sum_{j=1}^{\ell} \hat{H}_j$$

where each $\hat{H}_j$ acts on a space that involves at most $k_{max}$ of the variables. It is assumed that $\ell$ increases linearly with $N$ but that $k_{max}$ is fixed by the dimension and the symmetries and independent of $N$. The individual $\hat{H}_j$ can be represented as finite matrices in their local subspace. Under these assumptions, it is shown (Lloyd, 1996) that the error associated with the approximation

$$e^{i\hat{H}t} \simeq (e^{i\hat{H}_1t/n} \ldots e^{i\hat{H}_\ell t/n})^n + \ldots$$

can be controlled by taking $n$ large enough. In addition, once the accuracy goal is determined, the computing time scales linearly in $N$ and $\ell$.

In order to fix the ideas, for most of the available quantum computers, the $\hat{H}_j$ act on one or two qubits and can be represented by $2 \times 2$ or $4 \times 4$ matrices in this restricted space. A simple quantum circuit for the quantum Ising model used in Refs. (Gustafson et al., 2019a,b) is displayed in Fig. 1. The basic elements are rotations generated by $\hat{\sigma}_z^j$ or $\hat{\sigma}_x^j$ and acting on the $j$-th qubit only and CNOT gates acting on a pair of qubits and flipping the target qubit when the control qubit is in the $|1\rangle$ state. The circuit can be repeated in the space (vertical) and time (horizontal) directions and conveys the linear scalings stated above.

It is instructive to compare the computational resources to perform the unitary rotation $\hat{\sigma}_z^j$ which flips
the \( j \)-th qubit and does not act on the rest of the system. With a quantum computer, one expects that this flip keeps the coherence of the qubit with the rest of the system, present in the original state, but the cost is ideally independent on the size of the system. On the other hand, the same operation on an arbitrary state performed with a classical computer involves a matrix with \( 2^N \) non-zero and non-diagonal matrix elements and the resources necessary increase exponentially with the size of the system.

**G. Dealing with noise in the NISQ era**

The discussion in of Sec. III.F discuss algorithmic aspects in an idealized situation where computer errors or noise can be neglected. The quantum computing technology is still in its early development. The current Noisy Intermediate Scale Quantum (NISQ) computer hardware platforms can only accommodate small depth circuits and and sources of errors need to be understood in detail.

Various types of noise affect the single-qubit gates in a way that can be parametrized \( (Nielsen \text{ and Chuang, 2000}) \) in terms of the density matrix \( \hat{\rho} \):

\[
\mathcal{E}(\hat{\rho}; p_x, p_y, p_z) = (1 - p)\hat{\rho} + p_x\hat{\sigma}^y\hat{\rho}\hat{\sigma}^y + p_y\hat{\sigma}^y\hat{\rho}\hat{\sigma}^y + p_z\hat{\sigma}^z\hat{\rho}\hat{\sigma}^z. \tag{67}
\]

The values \( p_x, p_y, \) and \( p_z \) correspond to the probabilities of a \( \sigma^x, \sigma^y, \) or \( \sigma^z \) error, respectively, occurring and \( p = p_x + p_y + p_z \). The error channel for two qubit gates is given by \( \mathcal{E}(2) = \mathcal{E} \otimes \mathcal{E} \). In practice, if a classical simulation of the qubit evolution is performed, each unitary evolution operation needs to be followed by applying one of the four possibilities \( (1, \hat{\sigma}^x, \hat{\sigma}^y, \hat{\sigma}^z) \) with respective probabilities \( 1 - p, p_x, p_y, \) and \( p_z \) on the qubits involved.

In addition, readout errors (misidentifying a \( |1\rangle \) for a \( |0\rangle \) or vice-versa) should be taken into account for current superconducting qubit quantum computers. Given estimates of the probabilities for these errors, it is possible to correct the actual measurements by a multiplicative factor \( (Kandala et al., 2018) \).

A common NISQ strategy for error mitigation is to increase the source of error in a controllable way and then extrapolate to the limit where the error is not present. Examples with superconducting qubits are given in Refs. \( (Gustafson et al., 2019a,b; Kandala et al., 2019; Klco et al., 2018; Temme et al., 2017) \). A simple way to increase the error is to insert two successive CNOT gates. Their exact multiplication is the identity, however for a NISQ devices it increases the chances of errors.

In this context, the choice of Trotter step \( \delta t \) is crucial because the number of steps is limited by the loss of coherence and the noise. A concrete discussion can be found in Ref. \( (Gustafson et al., 2019a) \). If we pick a small \( \delta t \) with a good control on the \( (\delta t)^2 \) error, we may not be able to reach a time scale relevant for what we want to learn. However, it appears that by picking a \( \delta t \) significantly larger, the rigorous bound is not sharp and the empirical bound is much tighter as shown in Fig. 2

where we plot the operator norm of the Trotter error

\[
\Delta_2 U = e^{-ih_T \frac{J}{2} T} e^{ih_T \frac{J}{2} T} e^{-iJ \hat{H}_N} \delta t \approx \frac{h_T J}{2} [\hat{H}_T, \hat{H}_N](\delta t)^2. \tag{68}
\]

The nonlinear aspects of the error are quite interesting and may involve resonances \( (Gustafson et al., 2019a) \).

**Figure 2** \( ||\Delta_2 U|| \) versus \( \delta t \) for the quantum Ising model with \( h_T = 1 \) and \( J = 0.02 \). The vertical lines are at \( \pi/2, 3\pi/4 \). For details see Ref. \( (Meurice, 2020b) \).

**IV. QUANTUM VERSUS CLASSICAL**

In this section, we discuss the meaning of “classical” and “quantum” for models, phase transitions and tensor networks.

**A. Models**

In textbooks on quantum mechanics, it is a common procedure to start with the Hamiltonian formulation and derive a path-integral representation which can extended to field theory. The path-integral formalism allows for formulations that are manifestly gauge-invariant.
and treats space and time on equal footing. For these reasons, it can be argued that the fundamental definition of relativistic models should be done in terms of the action and the measure of integration over all the possible configurations without reference to a Hamiltonian in the first place. Examples of such actions are given in Sec. II.B.

It is very common to call models formulated with the path-integral “classical” while the corresponding formulation using a Hamiltonian acting on a Hilbert space is called “quantum”. However, except for possible discretization artifacts, the two formulations describe the same quantum behavior. In the path-integral formulation for bosonic fields, the action is calculated in terms of c-numbers as in the classical formulation but the sum over all the configurations provides a quantum description which cannot be reduced to the classical solutions of the equations of motion. In other words, the path-integral is an alternate method of quantization which is very convenient in Euclidean time.

Starting with a classical action, a Hamiltonian can be constructed from the action by using the transfer matrix formalism. This was briefly demonstrated in Sec. III.C, and will be discussed in detail in Sec. XII for the models introduced in Sec. II.B. A discussion with detailed references on the connection between statistical mechanics in $D$ dimensions and quantum Hamiltonians in $D - 1$ dimensions, can be found in the classic work of Wilson and Kogut (Wilson and Kogut, 1974).

### B. Phase transitions

The actions for spin and gauge models introduced in Sec. II.B contain parameters $\beta$ generically associated with links or $\beta_{pl}$ for pure gauge theories, that are often called the “inverse classical temperature” and can be associated with “classical phase transitions”. For instance the $D = 2$ classical Ising model has a spontaneous magnetization when $\beta > \beta_c = (1/2) \ln(1 + \sqrt{2})$.

In contrast, given a Hamiltonian $H$, we can define a thermal quantum partition function with temperature $T_{qu}$ in the usual way

$$Z(T_{qu}) = \text{Tr}[e^{-\hat{H}/T_{qu}}],$$  (69)

where $T_{qu}$ has in general a different meaning than $1/\beta$ in the classical formulation. With the lattice formulation at Euclidean time as a starting point, we have the identification

$$1/T_{qu} = N_\tau a_\tau,$$  (70)

and the non-zero temperature is associated with the finite extend of the temporal dimension. A typical situation of interest is to start in an ordered phase at $T_{qu} = 0$, corresponding to the infinite Euclidean time limit, and induce a finite-temperature phase transition into a disordered phase by taking a sufficiently small temporal extent. Sometimes, the transition can be understood in terms of the classical phase diagram in $D - 1$ dimensions. A more detailed discussion can be found in Cardy’s monograph (Cardy, 1996).

### C. Tensor networks

In Secs. V to XI we will introduce “classical tensors” in order to reformulate classical models as defined in Sec. IV.A. The partition function of these models can be visualized as an assembly obtained by “wiring” (tracing) together objects carrying multiple “legs” (tensor indices) and attached to the sites, links or plaquettes of a Euclidean space-time lattice. This type of classical constructions has been inspired (Gu and Wen, 2009; Levin and Nave, 2007) by various quantum tensor networks (Fannes et al., 1992; Perez-Garcia et al., 2006; Shi et al., 2006; Verstraete and Cirac, 2004; Verstraete et al., 2008; Vidal, 2007, 2003, 2004), to mention a few references, developed in various contexts often related to the density matrix renormalization group (DMRG) method (White, 1992). There is abundant literature on the subject reviewed for instance in (Cirac and Verstraete, 2009; Montangero, 2018; Orus, 2014; Ran et al., 2020; Schollwock, 2005, 2011; Silvi et al., 2017; Verstraete et al., 2008).

One important idea is the representation of quantum states by matrix product states (MPS) which appear in several references mentioned above. As an example, for a one-dimensional quantum chain problem with $N_s$ sites, an arbitrary element of the Hilbert space can be written as

$$|\psi\rangle = \sum_{i_1, \ldots, i_{N_s}} c_{i_1, \ldots, i_{N_s}} |i_1, \ldots, i_{N_s}\rangle,$$  (71)

where

$$|i_1, \ldots, i_{N_s}\rangle = |i_1\rangle \otimes \cdots \otimes |i_{N_s}\rangle,$$  (72)

and each of the indices runs over a local Hilbert space of dimension $d_H$ attached to a site. The dimension of the Hilbert space is $d_H^{N_s}$. It represents an exponential growth with the size of the system that rapidly becomes computationally unmanageable. For a MPS state, one assumes the form

$$c_{i_1, \ldots, i_{N_s}} = \text{Tr}[A_{i_1} \ldots A_{i_{N_s}}],$$  (73)

where the $A_{i_j}$ are $d_B \times d_B$ matrices for each value of $i_j$, $d_B$ is called the bond dimension. A graphical representation of such a state is shown in Fig. 3 for open boundary conditions. The filled circles represent the matrices, the vertical lines represent open indices with $d_H$ values and the horizontal lines traced indices with $d_B$ values. The size of the MPS sub-Hilbert space is only growing like
similar shapes will appear as “times slices” of the classical construction. In the case where the indices take an infinite number of values and the sums are truncated in a way that the number of indices kept in the time direction (similar to $d_H$), is the same as the number in the space directions (similar to $d_B$), we will use the isotropic notation $D_{\text{cut}}$. On the other hand, we will discuss the anisotropic situation where the time direction is singled out to define the transfer matrix and the Hamiltonian in Sec. XII.

MPS have been used for 1+1 dimensional models including the Schwinger model (Bañuls et al., 2013; Buyens et al., 2014, 2016), SU(2) gauge theory (Bañuls et al., 2017b; Kühn et al., 2015) and the O(3) models (Bruckmann et al., 2019). Tensor network techniques for lattice gauge theories are also discussed in (Haegeman et al., 2015; Pichler et al., 2016; Rico et al., 2014; Silvi et al., 2016, 2019; Tagliacozzo et al., 2014; Zohar et al., 2015b) and reviewed in (Bañuls et al., 2018). For related work in 2+1 dimensions see (Felser et al., 2019; Robaina et al., 2020; Tagliacozzo and Vidal, 2011).

V. TENSOR METHODS EXPLAINED WITH THE ISING MODEL

A. Tensor formulation

In this section we will construct a tensor formulation for the Ising model in $D$ dimensions. The partition function for the Ising model is

$$Z_{\text{Ising}} = \prod_x \sum_{\sigma_x = \pm 1} e^{\beta \sum_{\mu \neq \mu} \sigma_x \mu \bar{\sigma}_x}.$$  (74)

For each link $(x, \mu)$ we use the expansion

$$e^{\beta \sigma_x \mu \bar{\sigma}_x} =$$

$$\cosh(\beta) \sum_{n_{x,\mu} = 0, 1} [\sigma_x + \mu \sqrt{\tanh(\beta)} \sigma_x \sqrt{\tanh(\beta)}]^{n_{x,\mu}},$$ (75)

This attaches an index $n_{x,\mu}$ at each link $(x, \mu)$. It is then possible to pull together the various factors of $(\sqrt{\tanh(\beta)} \sigma_x)^{n_{x,\mu}}$ from links coming from a single site $x$, and sum over $\sigma_x$,

$$\sum_{\sigma_x = \pm 1} \prod_{\mu = 1}^D (\sqrt{\tanh(\beta)} \sigma_x)^{n_{x,\mu} + n_{x,\mu}}$$

$$= (\sqrt{\tanh(\beta)}) \sum_{\mu} n_{x,\mu} + 2 \delta(\text{mod}[n_{x,\mu}, 2]) \quad (76)$$

using

$$\sum_{\sigma = \pm 1} \sigma^n = 2 \delta(\text{mod}[n, 2]).$$ (77)

The expression $\delta(\text{mod}[n, 2])$ is 1 when $n$ is even (0 modulo 2) and 0 otherwise. We can rewrite the partition function as the trace of a tensor product,

$$Z = (2 \cosh(\beta))^{V_D} \text{Tr} \left[ \prod_x T^{(x)}_{n_{x-1,1},n_{x,1},\ldots,n_{x,D}} \right].$$ (78)

$\text{Tr}$ means contractions (sums over 0 and 1) over the link indices (the $n_{x,\mu}$s). The local tensor $T^{(x)}$ has $2D$ indices. The explicit form is

$$T^{(x)}_{n_{x-1,1},n_{x,1},\ldots,n_{x-D,D},n_{x,D}} =$$

$$(\sqrt{\tanh(\beta)})^{n_{x,\mu} + n_{x,\mu}} \times 2 \delta(\text{mod}[n_{x,\mu} + n_{x,\mu}, 2]) \quad (79)$$

with the definitions

$$n_{x,\mu} \equiv \sum_{\mu = 1}^D n_{x,\mu}$$ (80)

$$n_{x,\mu} \equiv \sum_{\mu = 1}^D n_{x,\mu}.$$ (81)

The notions of “in” and “out” refer to position of the link with respect to the positive directions. The basic tensors and their assembly in two and three dimensions are illustrated in Figs. 5-8.

Any kind of boundary condition can be accommodated by adapting the method of integration to the link configuration at the boundary. This will be discussed in subsection V.C.

The Kronecker delta in Eq. (79) implies the discrete conservation law

$$\sum_{\mu} (n_{x,\mu} - n_{x,\mu}) = 0 \text{ mod } 2,$$ (81)
which we also call the “tensor selection rule”. It implies that only an even number of \( n_s \) are allowed to take on the value one. For instance, for \( D = 2 \), there are in principle 16 tensor values, however, only eight are nonzero, one with all four indices as zero (zeroth order in \( \tanh(\beta) \)), six with two zeros and two ones (linear in \( \tanh(\beta) \)), and one with four ones (quadratic in \( \tanh(\beta) \)). Note that if a symmetry breaking term like a magnetic field coupling to the total spin is introduced, then all the 16 tensors elements will be nonzero generally.

There are also tensor formulations which use the singular value decomposition on each nearest-neighbor Boltzmann factor to factorize the spins. The nearest neighbor interactions can be represented as a matrix, whose indices are the spin variables themselves,

\[
e^{\beta \sigma_{x+\mu} \sigma_x} = \begin{pmatrix} e^{\frac{\beta}{2}} & e^{-\frac{\beta}{2}} \\ e^{-\frac{\beta}{2}} & e^{\frac{\beta}{2}} \end{pmatrix}
\]

One can then of course perform the singular value decomposition to get,

\[
e^{\beta \sigma_{x+\mu} \sigma_x} = \sum_{\alpha, \beta} U_{\sigma_{x+\mu} \alpha} \lambda_{\alpha \beta} U^T_{\beta \sigma_x},
\]

and in the case of the Ising model the singular value decomposition has the same left and right unitary matrices.

This factorizes the spins and allows the creation of the matrix,

\[
W_{\sigma_x \alpha} \equiv U_{\sigma_x \alpha} \sqrt{\lambda_{\alpha}}.
\]  

The main local tensor is then the contraction of all \( W \) matrices which have a common spin,

\[
T_{i_1 \cdots \alpha} = \sum_{\sigma_x} W_{\sigma_x i_1} W_{\sigma_x j} \cdots W_{\sigma_x N}.
\]

In fact, the singular value decomposition in this case can be completely related to the expansion in Eq. (75). Consider the character expansion from Sec. V.A for the nearest neighbor interaction,

\[
e^{\beta \sigma_{x+\mu} \sigma_x} = \frac{1}{2} \sum_{n_{x,\mu}=0}^{1} e^{n_{x+\mu} \sigma_{x} C_{n_{x,\mu}(\beta)} \sigma_{x}},
\]

with matrices,

\[
\sigma_{x,\mu}^{n_{x,\mu}} = \begin{pmatrix} \frac{1}{2} & \frac{1}{\sqrt{2}} \\ \frac{1}{\sqrt{2}} & \frac{1}{2} \end{pmatrix}
\]

whose columns are the normalized eigenvectors of the Pauli-\( x \) matrix, and \( C_0 = 2 \cosh \beta \) and \( C_1 = 2 \sinh \beta \). The elements of this matrix are from the four values \( n_{x,\mu} \) and \( \sigma_x \) can take, normalized to be unitary. Since the variables are compact, operators of them have a discrete spectrum, and the character expansion is exactly the spectrum decomposition for this matrix. For any matrix, \( M \), by definition the singular value decomposition is found from the eigenvalues and eigenvectors of \( MM^\dagger \)
and \(M^1M\). In this case (we drop any specific spacetime lattice indices for these steps since they are completely general),

\[
\sum_{\sigma_j} e^{\beta\sigma_i} e^{\beta\sigma_j} e^{\beta\sigma_k} = \sum_{\sigma_j} \sum_{n,m} e^{\sigma_j^n} C_{n} e^{\sigma_j^m} C_m e^{\sigma_k^m} = \sum_{n,m} \sigma_j^n C_{n} \sigma_j^m C_m \delta_{n,m} = \sum_{n} \sigma_j^n C_{n} \sigma_j^m.
\]

Then the singular values are given by \(\lambda_1 = 2|\cosh \beta|\) and \(\lambda_2 = 2|\sinh \beta|\), as one would expect from Eq. (83) and Eq. (86).

**B. Remarks on the forms of duality**

The tensor representation can be used to reproduce the set of closed paths appearing in the expansion in powers of \(\tanh(\beta)\) (Itzykson and Drouffe, 1991; Parisi, 1998) for the Ising model. The links associated with the set of indices \(n_{x,\mu} = 1\) form a graph (a set of sites connected by links). The selection rule means that each site is attached to an even number of nonzero links. These graphs are closed paths with specific connectivity which can in principle be enumerated order by order in their length using geometric constructions and combinatoric techniques.

One can try to construct these closed paths by assembling the most elementary contributions, namely closed loops on a single plaquette. The way they can be assembled depends on the dimension. For instance for \(D = 2\), we can decide that when two loops around two plaquettes share a link, this link is “erased” from the path. Alternatively, one can introduce “dual variables” originally conceived by Kramers and Wannier (Kramers and Wannier, 1941) which are spins on the dual lattice located at the centers of the plaquettes. Each dual spin is then associated with a closed plaquette loop around it on the original lattice.

Furthermore, the Pointryagin dual variables \(n_{x,\mu}\) can be expressed as \((1 - \tilde{\sigma} \tilde{\sigma}')/2\) with \(\tilde{\sigma}\) and \(\tilde{\sigma}'\) the two dual spin variables connected by a dual link crossing the original link. A picture showing the dual variables in their different locations can be seen in Fig. 9. This illustrates that the notions of duality are often combined in a way that may be seen as confusing at first sight. Note that the dual domains with a given dual spin have boundaries that can be interpreted as the closed paths of the of the original model. The questions of completeness and multiplicity need to be addressed with specific boundary conditions.

Similarly in higher dimensions it is possible to introduce dual spins with interactions involving \(2(D-1)\) spins. For \(D = 3\), this leads to a gauge theory with plaquette interactions (Wegner, 1971). It is also possible to introduce dual variables for Ising models with arbitrary all-to-all spin interactions (Meurice, 1994). Duality questions related to Gauss’s law will be discussed in Sec. XII.C.

**C. Boundary conditions**

In Eq. (78), the trace is a sum over all the link indices. We need to specify the boundary conditions. Periodic boundary conditions (PBC) allow us to keep a discrete translational invariance: the tensors themselves are translation invariant and assembled in the same way at every site. Open boundary conditions (OBC) can also be implemented by introducing new tensors that can be placed at the boundary. Their construction is similar to the tensors in the bulk. The only difference is that the “outside links” which would be attached at sites on the boundary have an index set to zero. With the normalization introduced in Eq. (75) the indices carrying a zero index carry a unit weight. This construction can be understood as decoupling the system from a larger environment by setting \(\beta\) on the links connecting to this environment to zero because \(\tanh(0) = 1\) and \(\tanh(0) = 0\). This is illustrated in Fig. 10.

**D. Exact blocking**

An important feature of the tensor representation presented in Sec. V.A is that it allows an exact local blocking procedure in contrast to what can be done in configuration space. We divide the original lattice into cells having a linear size of two lattice spacings (“blocks”) in such a way that the boundaries are half-way between lattice sites. In other words, the boundaries are normal to links.
and cut them in the middle. As an example for $D = 2$, the blocks are square enclosing four sites and their four sides cross eight links. This cell partition separates the link degrees of freedom into two disjoint categories: those completely inside the block, which can be integrated over, and those shared by neighboring blocks and kept to communicate between the blocks. This is a generic feature of the method (Meurice, 2013) which motivated a systematic study of lattice models (Liu et al., 2013). Note that when translation invariance is present, all the blocks are identical and we only need to do one calculation.

We now describe explicitly this exact blocking for $D = 2$ using generic notations inspired by those of Refs. (Meurice, 2013; Xie et al., 2012). The extension to higher dimensions on hypercubic lattices is straightforward. We contract the four tensors located at the four sites inside the block along the four indices located strictly inside the block as well. The remaining eight indices associated with the eight links piercing the block are left as new degrees of freedom. By taking the tensor product between the two indices in each of the four directions coming out of the block, we obtain a new rank-4 tensor $T'_{XX'YY'}$. In the case of the Ising model, each index now takes four values. This provides a simple isotropic formula:

$$T'_{X(x_1,x_2)X'(x'_1,x'_2)Y(y_1,y_2)Y'(y'_1,y'_2)} = \sum_{x_U,x_D,x_R,x_L} T_{x_U,x_D,x_R,x_L} T_{x_U,x_D,x_R,x_L} T_{x_U,x_D,x_R,x_L} T_{x_U,x_D,x_R,x_L},$$

where $X(x_1,x_2)$ is a notation for the product states i.e., if we regroup the indices with the same parity together, $X(00) = 1$, $X(11) = 2$, $X(10) = 3$, $X(01) = 4$. This contraction and redefinition, relative to the block, is illustrated in Fig. 11.

After this blocking, the partition function can again be written as

$$Z = \text{Tr} \prod_{2x} T''_{X''X''Y''Y''},$$

where $2x$ denotes the sites of the coarser lattice with twice the lattice spacing of the original lattice. This coarse-graining provides an exact representation of the original partition function. However, the number of states associated with each index is the square of the number of states in the original tensor. If this exact procedure had to be carried numerically, this rapid growth would quickly run into practical limitations. Truncations are thus necessary. It is important to appreciate that truncations are the only approximations that will be needed. We now proceed to discuss truncations.

VI. TENSOR RENORMALIZATION GROUP

A. Block-spinning through SVD

Once partition functions and physical quantities are expressed as tensor networks, one needs to contract them to obtain numerical values. However, contracting exactly requires an extraordinary amount of computational resources. Then, in this subsection we work through the original idea of how one can perform tensor contractions approximately, and how truncations appear in this approximation. To do that a coarse-graining algorithm

Figure 11 Graphical representation of the block (dotted square) and $T'_{XX'YY'}$, as in (Meurice, 2013).
for tensor networks is introduced. The original idea of such an approach was proposed by Levin and Nave in (Levin and Nave, 2007), where tensor networks are simply coarse-grained by using the SVD (see Sec. A.3). This method is similar in spirit to the real-space renormalization group approach, and, in this sense, it is called the tensor renormalization group.

In the standard renormalization group procedure (Wilson and Kogut, 1974), the blocking process is supplemented by a sorting of the resulting information according to their degree of relevance. As far as universal properties characterizing the continuum limit, it is acceptable to discard the information which only reflects the microscopic details of a specific lattice formulation. In the context of the tensor formulation discussed above, it means that, possibly after a certain number of exact contractions, we need to restrict the number of states associated with the tensor indices to a fixed number $D_{\text{cut}}$. We are then mapping a tensor with $D_{\text{cut}}^2$ entries into another tensor of the same shape and the question of fixed points becomes meaningful. An important goal of the renormalization procedure is to identify fixed points. Note that the updated tensor remains a local object which supersedes the notion of action or Hamiltonian.

Here we assume that a partition function is expressed as a two dimensional tensor network with bond dimension $D_{\text{cut}}$:

$$Z = \text{Tr} \prod_x T^{(x)}_{x'xyy'}, \quad (90)$$

and that the periodic boundary conditions are imposed in all directions.

First, the tensor $T$ can be regarded as a $D_{\text{cut}}^2 \times D_{\text{cut}}^2$ matrix, and can be approximately decomposed using the SVD in two ways:

$$T_{x'xyy'} = M_{(x'y)(xy')}^{[13]} \approx \sum_{m=1}^{D_{\text{cut}}} S_{(xy)m}^{[1]} \lambda_m^{[13]} S_{m(xy')}^{[3]}, \quad (91)$$

$$T_{y'x'yx} = M_{(y'x)(yx')}^{[24]} \approx \sum_{m=1}^{D_{\text{cut}}} S_{(yx'm)}^{[2]} \lambda_m^{[24]} S_{m(yx')}^{[4]}, \quad (92)$$

where $\lambda^{[13]}$ and $\lambda^{[24]}$ are the singular values (assumed to be the descending order: $\lambda_1 \geq \lambda_2 \geq \cdots \geq \lambda_{D_{\text{cut}}}^{[13]} \geq 0$), and $S^{[1]}$, $S^{[2]}$, $S^{[3]}$, $S^{[4]}$ are unitary matrices. Here the degree of the approximation is set to be $D_{\text{cut}}$ for simplicity. Of course one can freely choose this parameter, and it becomes the bond dimension of the coarse-grained tensors.

Using the decomposed tensors $S^{[i]}$ ($i = 1, 2, 3, 4$), a coarse-grained tensor is defined by

$$T^{\text{new}}_{x'xyy'} = \sqrt{\lambda_x^{[13]} \lambda_y^{[24]} \lambda_{x'}^{[13]} \lambda_{y'}^{[24]}} \times \sum_{a,b,c,d=1}^{D_{\text{cut}}} S_{x(cdy)}^{[3]} S_{y(das)}^{[4]} S_{(ab)x'}^{[1]} S_{(bc)y'}^{[2]}, \quad (93)$$

The number of tensors on the network is now reduced by 1/2, and the bond dimension of $T^{\text{new}}$ is a free parameter and here set to be the same as that of $T$, $D_{\text{cut}}$. Repeating this procedure, one can make an effective tensor network that consists of a few tensors, and then one can take the contraction of the tensor indices. A graphical explanation of the TRG is given in Fig. 12.

After a coarse-graining step, the network is rotated by 45 degrees, and then how one defines the new unit vectors is one’s choice. One possible way is to define them by $1^* = 1 + 2$ and $2^* = 1 - 2$, where $1^*$ and $2^*$ are the unit vector along the $1$- ($2$-)direction of the original lattice and that on the coarse-grained lattice, respectively (see Fig. 12). Using this definition, the orientation of the network is recovered after every two coarse-graining steps; i.e., $1^{**} = 1^* + 2^* = 2 \cdot 1$ and $2^{**} = 1^* - 2^* = 2 \cdot 2$.

In this procedure the relevant degrees of freedom are decided by the SVD during the decomposition of the $T$ tensor into an intermediate sum-over-states. In the next section we will see there are improved methods to pick out the relevant states during truncation.

### B. Optimized truncations

The previous section discussed the first work at a renormalization group procedure using a tensor formulation. Here we will discuss refinements which occurred later. These refinements incorporate an environment tensor from which relevant states are determined and kept.

We begin with assuming that one has completed contraction of a single block. Each tensor index of the blocked tensor now possesses $D_{\text{cut}}^2$ states. The next step is to find a way to decide which of the $D_{\text{cut}}^2$ states (or possibly a linear combination of them) corresponding to the tensor product associated with each index should be kept. Ideally, we would like to address this question in terms of the environment of a single tensor. We can write

$$\text{Tr} \prod_x T^{(x)} = \sum_{XX'YY'} E_{XX'YY'} T_{XX'YY'}^{(0)}, \quad (95)$$

where $E_{XX'YY'}$ is obtained by tracing all the indices except for four pairs of indices associated to a single block.

![Figure 12](image_url)
that we have located at the origin. This is illustrated in Fig. 13. Obviously, the environment tensor $E_{XX'YY'}$ is very close to what are ultimately trying to calculate and so it is not immediately available. The only purpose of using $E_{XX'YY'}$ is to rank order the states of the tensor product and its exact form does not appear in later calculations. Consequently, the exact form may not be too important and as a first step, we can try approximations. A very simple approximation is to ignore the details of the environment and use (Meurice, 2013)

$$E_{XX'YY'}^{\text{app}} = C \delta_{XX'} \delta_{YY'},$$

for some positive constant $C$. We can then optimize the truncation by maximizing the approximate partition function expressed in terms of the trace of a matrix $G$ such that

$$\text{Tr } G = (1/C) \sum_{XX'YY'} E_{XX'YY'}^{\text{app}} T_{XX'YY'}^{(0)},$$

which can be achieved with

$$G_{XX'} = \sum_{YY'} T_{XX'YY'}^{(0)}.$$  

By looking at the expression in terms of the original tensors one realizes that $G_{XX'}$ is in fact the square of another matrix (Meurice, 2013) and if the eigenvalues of this matrix are real than all the eigenvalues of $G$ are positive and we can optimize the truncations by selecting the states corresponding to the largest eigenvalues of $G$.

A more refined approximation is to assume that the environment is a “mirror image” of the tensor itself (Xie et al., 2012):

$$E_{XX'YY'}^{\text{app}} = C' T_{XX'YY'}^{(0)*},$$

The trace of $G$ can then be identified with the tensor norm:

$$\text{Tr } G = \sum_{XX'YY'} T_{XX'YY'}^{(0)} T_{XX'YY'}^{(0)*} = ||T^{(0)}||^2,$$

which is clearly a sum of positive terms. This can be accomplished with the Hermitian matrix

$$G_{XX'} = \sum_{XX'YY'} T_{XX'YY'}^{(0)} T_{XX'YY'}^{(0)*}. \quad (101)$$

The problem is then reduced to selecting the states that provide the best approximation of Tr$G$ which is obvious when all the eigenvalues are positive.

The procedure that we just described is isotropic. It is however possible to coarse grain one direction at a time (Xie et al., 2012) in order to reduce the size of the summed expressions. For instance, the summation over the tensor product indices $Y$ and $Y'$ in Eq. (101) could be first replaced by summations over single indices,

$$M_{X(x_1,x_2)X'(x'_1,x'_2)yy'} = \sum_a T_{x_1x_2}^{a} T_{x'_1x'_2}^{a} T^{(0)}_{XX'YY'}. \quad (102)$$

The tensor $M$ can then be used in Eq. (101) in place of $T^{(0)}$ as the “blocked” tensor to find the most relevant states for the $X$ indices. This provides a coarse-graining in the first direction. It is then necessary to coarse-grain in the second direction using sums over single indices in the first direction. As our discussion is focused on $D = 2$, these two steps constitute a coarse-graining that doubles the lattice spacing in all directions. In (Xie et al., 2012), these calculations were conducted using higher-order generalizations of the SVD method called the higher-order tensor renormalization group (HOTRG).

A better description of the environment can be reached by following a local truncation procedure as described above for a sufficiently large but finite number of times at which point it is assumed that there is no environment and Eq. (96) can be used. In other words, by working with a finite lattice, the procedure is terminated by approximating the partition function as a trace of the last coarse-grained expression for the tensor. It is then possible to move “backward” (Xie et al., 2009; Zhao et al., 2010) and reconstitute the approximate environment of a single tensor coarse-grained one less time. Explicit expressions based on the HOTRG construction can be found in (Xie et al., 2012). This can be pursued recursively until we reach the first coarse graining level illustrated in Fig. 13.

The analogy with the backward propagation used in machine learning has been exploited to design new algorithms recently (Chen et al., 2020). More generally, algorithmic improvement in the TRG context is a subject of active investigation (Bal et al., 2017; Fishman et al., 2018; Morita and Kawashima, 2020)

C. Higher dimensional algorithms

While the Levin–Nave type TRG can be applied to two dimensional systems, higher dimensional systems are

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure13.png}
\caption{A blocked tensor and its environment.}
\end{figure}
deal with using other algorithms. One such algorithm is the higher-order TRG (HOTRG) (Xie et al., 2012) mentioned in the previous section. Using the HOTRG, in principal, any dimensional tensor network can be coarse-grained. When a $D$ dimensional tensor network is build by tensors with the bond dimension $D_{\text{cut}}$, the computational complexity of the HOTRG is $O(D_{\text{cut}}^{4D-1})$, and the memory complexity is $O(D_{\text{cut}}^{2D})$.

Recently, cheaper algorithms have been invented. The anisotropic tensor renormalization group (ATRG) (Adachi et al., 2019), whose graphical description is given in Fig. 14, achieved the time and the memory complexities $O(D_{\text{cut}}^{2D+1})$ and $O(D_{\text{cut}}^{D+1})$ that are significant reductions from the HOTRG. The ATRG introduces an approximation of an approximation, and indeed, when the bond dimensions are the same, the ATRG is less accurate compared to the HOTRG. However, thanks to the cheaper complexity, the ATRG leads to better accuracy with fixed CPU time. Using the ATRG, four dimensional systems, where the HOTRG is much more expensive, have begun investigation (Akiyama et al., 2020a, 2019, 2020b).

![Figure 14 Graphical description of the ATRG. (a) Decompose $T$ into $(A$ and $B)$ and $(C$ and $D)$ with the SVD. (b) Swap the bonds of $B$ and $C$. The swapping can be done also by the SVD: $\sum B_{x,y} C_{x,y'} = (BC)_{x,y'} C'_{x,y} = \sum B'_{x,y} S_{x,y} C_{x,y}$. (c) Contract $A', B', C', D$ while truncating the dimensions of horizontal bonds.](image)

Another approach is coarse-graining on a triad tensor network representation (Kadoh and Nakayama, 2019). Although we do not describe the details, some comparisons to the HOTRG and the ATRG are done in the paper.

**D. Niemeijer-Van Leeuwen equation**

In Sec. V.D we have constructed a coarse-grained tensor that can be used to give an exact expression for the partition function. Despite the fact that we integrated over microscopic degrees of freedom, the number of tensor indices needed for this exact representation grows exponentially with the size of the blocks. In Secs. VI.A and VI.B, we introduced truncations where a fixed number of indices $D_{\text{cut}}$ was kept at each step. This procedure discards some information but allows us to compare the tensors before and after the coarse-graining. Typically, the tensors tend to grow exponentially with the number of coarse graining steps and it is important to renormalize their absolute size or to only consider their ratios. After such a renormalization takes place, we obtain a RG transformation. The fixed points of this transformations are the central objects of the RG approach and it is useful to compare the TRG equations with standard RG equations due to Niemeijer and van Leeuwen (NvL)(Niemeijer and van Leeuwen, 1976).

NvL were aware of the difficulty of controlling the new couplings generated by the blocking procedure, so they started immediately with the most general Ising interactions in a finite volume $V$. If no conditions are imposed, there are as many couplings as Ising configurations so this is not suitable for numerical purposes. They then introduced 1 in the partition function as in Eq. (1), in order to define new Ising spins $\{\sigma'\}$ in a volume $V' = V/b^D$ with a new lattice spacing rescaled by the linear size of the blocks $b$. They were able to give a formal expression for the new couplings in terms of the original ones as $K'(K)$. Strictly speaking there are less couplings after the coarse-graining because they considered the most general case involving the products of spins in arbitrary domains, but they assumed that only a certain number of quasi-local couplings were important and had the same form before and after the coarse-graining. In addition they assumed that the dependence of the free energy density $f(K)$ on these couplings is the same after the coarse-graining. This led to the NvL equation

$$f(K) = g(K) + b^{-D} f(K').$$

The function $g = G/V$, comes from

$$G = \sum \ln \left( \sum_{\{\sigma'\}} P(\{\sigma'\}, \{\sigma\}) \exp \left( H(\{\sigma\}) \right) \right),$$

and is defined by the condition

$$\sum_{\{\sigma'\}} H'(\{\sigma'\}) = 0,$$

where $H$ and $H'$ are the Hamiltonians before and after coarse graining.

Even though computing the new couplings and the functions may be very difficult in practice (the new couplings are double partition functions), NvL succeeded to obtain a formal relation which can be iterated and linearized near a fixed point. This allows us to identify the relevant directions and it is often taken as the starting point for the introduction of the RG method in textbooks (Cardy, 1996).

For the TRG, we can factor out the increasing size of the tensors, for instance, by imposing the normalization condition

$$T_{0000} = 1,$$
at each step. In other words we divide all the tensors by one unnormalized tensor element. We now have the two steps (coarse-graining and renormalization) that define a RG transformation. We can write the exact identity

\[
\ln(\text{Tr} \prod_{\text{sites}} T^{(\text{sites})}_{xx'yy'})/V = (1/4) \ln(T'_{0000}) + (1/4) \ln(\text{Tr} \prod_{\text{sites}'} T^{(\text{sites}')}_{XX'Y'Y'})./V'.
\]

\(T'_{0000}\) is the unnormalized tensor element that we constructed in Sec. V.D. \(T^{(\text{sites})}_{XX'Y'Y'}\), is the renormalized tensor meaning the unnormalized tensor divided by \(T_{0000}\). Bearing in mind that \(b^{-D} = 1/4\) we see the analogy with the NvL equation (103). \((1/4) \ln(T'_{0000})\) plays the role of \(g(K)\). Note that \(\text{Tr} \prod_{\text{sites}}\) has a different meaning in both sides of the equation. However, if we assume that the coupling dependence of the densities are the same before and after as in NvL and we obtain a RG equation. In both cases, neglecting couplings can be justified by the fact the the RG transformation has only a small number of important directions in the space of couplings. This will be illustrated with a simple example in the next subsection. More details can be found in Ref. (Meurice, 2020b).

E. A simple example of TRG fixed point

In the following we discuss the two-state truncation for the Ising model. In other words, we keep the same number of states for each index as for the initial tensor. With the indices taking two values, the rank four tensor has in principle 16 independent entries, however because of the Ising selection rule the sum of the indices must be even and so eight of the tensor values are zero. In addition, if we preserve the symmetry under the rotation of the square lattice by \(\pi/2\), this imposes that

\[
T_{1010} = T_{0110} = T_{1001} = T_{0101} = t_1,
\]

and

\[
T_{1100} = T_{0011} = t_2.
\]

In addition, we define

\[
T_{1111} = t_3.
\]

For the initial tensor, we have

\[
t_1 = t_2 = \tanh(\beta) \quad \text{and} \quad t_3 = t_1^2.
\]

The property \(t_1 = t_2\) is not preserved by the blocking procedure which can be expressed as a mapping of the three dimensional parameter space \((t_1, t_2, t_3)\) into itself that we denote \(t'_1(t_1, t_2, t_3)\).

As a numerical example, we use the method of Eq. (96) and the normalization from Eq. (106) which is discussed after Eq. (10) in (Meurice, 2013). The results for \(t_1\) as a function of the initial \(\beta\) for 6 iterations are shown in Fig. 15. We see that for values of \(\beta\) low enough, \(T_{1010}\) goes to zero at a faster rate as the the number of iterations increases. On the other hand, for values of \(\beta\) large enough, \(T_{1010}\) goes to 1. As the number of iterations increases, the transition becomes sharper and sharper and singles out a critical value \(\beta_c = 0.394867858...\) where the curves for successive iterations intersect. This also singles out a fixed point value for \(t_1\) near 0.4. The graphs for \(t_2\) and \(t_3\) have similar features. Near \(\beta_c\), the departure from the fixed point value is approximately linear in \(\beta - \beta_c\) with a slope of the form \(\lambda_1^\ell\), where \(\ell\) is the number of iterations. It is possible to rescale \(\beta - \beta_c\) by a factor \(\lambda_1\) at each iteration to obtain a “data collapse” shown at the bottom of Fig. 15. Numerically (Meurice, 2013), \(t_1^* = 0.42229, t_2^* = 0.28637\) and \(t_3^* = 0.27466\) and \(\lambda_1 = 2.00931069\) which provides a critical exponent \(\nu = \log b/\log \lambda_1 \approx 0.993\) which is surprisingly close to the exact value 1 given that the truncation is quite drastic. Very similar results are obtained in a dual version of the map in (Aoki et al., 2009).

Figure 15 \(t_1 = T_{1010}\), versus \(\beta\) (top) and versus \((\beta - \beta_c)\lambda_1^\ell\) (bottom) for \(\ell = 1, \ldots, 6\) iterations of the two-state approximation. The dotted line is at the critical value. As the iteration number increases, the color smoothly changes from blue to green.

One would think that by adding a few more states, we could get even better results, however this is not the case (Efrati et al., 2014). One of the reasons is explained in
the next section.

F. Corner double line structure on tensor network

In this subsection we discuss a fixed point of the TRG (Levin and Nave, 2007) that is called the corner double line (CDL) tensor (Gu and Wen, 2009). Here let us consider a toy model of short-range correlations that is expressed as a tensor network spanned by tensors with the following form:

\[ T_{ijkl}^{\text{CDL}} = \delta_{i_1,j_2} \delta_{j_1,i_2} \delta_{k_1,j_2} \delta_{l_1,k_2}, \]

(112)

where each tensor index has two components: e.g. \( i = (i_1, i_2) \) (see Fig. 16). We call \( T_{ijkl}^{\text{CDL}} \) the CDL tensor, and it describes interactions on plaquettes as seen in Fig. 17. Now, let us consider a TRG step for this tensor network.

Then, by contracting the decomposed components, the coarse-grained tensor is given by

\[ (T_{ijkl}^{\text{CDL}})_{ijkl}' = \sum_{a,b,c,d=1}^{D_{\text{cut}}} \delta_{i_1,c_1} \delta_{d_2,i_2} \delta_{j_1,d_2} \delta_{a_1,j_2} \delta_{b_1,a_1} \delta_{b_2,k_2} \]

\[ \cdot \delta_{l_1,b_2} \delta_{c_1,l_2} \delta_{a_2,b_1} \delta_{c_2,d_1} \delta_{d_1,a_2} \]

\[ \propto T_{ijkl}^{\text{CDL}}. \]

(114)

Each step is graphically displayed with the assignments of indices in Figs. 18–19. Surprisingly the CDL tensor has turned out to be a fixed point of the TRG up to a constant factor. This is not a physical but an artificial fixed point, and unfortunately this fact means that the TRG leaves short-range correlations on coarse-grained tensors. This is because the SVD is the best approximation of a tensor but is not the best for a network.

To avoid such an unexpected fixed point, one needs to consider more global blocking procedures. One ideal way is to insert unknown tensors (called “(dis)entangler”) on a network and variationally tune them to remove CDL structures. This method is called the tensor network renormalization (TNR) (Evenbly and Vidal, 2015) and leads to more precise computations. In principal this can be extended to three or higher dimensions although the computational complexities would be extremely demanding. In two dimensions, more low-cost methods are invented so far (Evenbly, 2018; Hauru et al., 2018; Yang et al., 2017) although we do not mention in detail here. The common concept in such approaches is to consider global cost functions and to remove the CDL structures on a network.
VII. TENSORS FOR SPIN MODELS WITH AN ABELIAN SYMMETRY

In this section, we discuss tensor formulations of generalizations of the $O(2)$ nonlinear sigma model introduced in Sec. II.B where the spin variables form a two-dimensional unit vector varying continuously over a circle. We will show later that the results extend easily to the clock models with a discrete $\mathbb{Z}_q$ symmetry.

A. $O(2)$ nonlinear sigma model

The partition function for the $O(2)$ model reads

$$Z_{O(2)} = \prod_x \int_{-\pi}^\pi \frac{d\varphi_x}{2\pi} e^{-S_{O(2)}},$$

with

$$S_{O(2)} = -\beta \sum_{x,\mu} \cos(\varphi_{x+\mu} - \varphi_x).$$

We use the Fourier expansion to expand the Boltzmann weights,

$$e^{\beta \cos(\varphi_{x+\mu} - \varphi_x)} = \sum_{n,\mu=1}^{\infty} e^{i n \mu \varphi_{x+\mu}} I_n(\beta) e^{-i n \mu \varphi_x}.$$

This expansion factorizes the $\varphi$ fields. We then integrate over all the $\varphi$ fields using the orthogonality relations of the Fourier modes, e.g.,

$$\int_{-\pi}^\pi \prod_{\mu=1}^D \sum_n e^{i (n_x - n_{x\mu} - n_{x,\mu}) \varphi_x} = \delta_{n_{\text{in}}, n_{\text{out}}, 0},$$

with $n_{\text{in}}$ and $n_{\text{out}}$ defined in the same way as in Sec. V. We can rewrite the partition function as the trace of a tensor product:

$$Z = \text{Tr} \prod_x T_{n_{x-1},n_x,1,\ldots,n_x,D}^{(x)}.$$

The local tensor $T^{(x)}$ has $2D$ indices. The explicit form is

$$T_{n_{x-1},n_x,1,\ldots,n_x,D}^{(x)} = \sqrt{\prod_{\mu=1}^D I_{n_{x\mu}} I_{n_{x-D\mu}}} \delta_{n_{\text{out}}, n_{\text{in}}},$$

where

$$\sum_{\mu=1}^D (n_{x\mu} - n_{x-D\mu}) = 0, \quad \sum_{\mu=1}^D (n_{x\mu} - n_{x-D\mu}) = 0, \quad \sum_{\mu=1}^D (n_{x\mu} - n_{x-D\mu}) = 0,$$

and

$$\sum_{\mu=1}^D (n_{x\mu} - n_{x-D\mu}) = 0, \quad \sum_{\mu=1}^D (n_{x\mu} - n_{x-D\mu}) = 0, \quad \sum_{\mu=1}^D (n_{x\mu} - n_{x-D\mu}) = 0,$$

which is a discrete version of Noether current conservation if we interpret the $n_{x\mu}$ with $j = 1, \ldots, D-1$ as spatial current densities and $n_{x,D}$ as a charge density. At finite $\beta$, the relative size of the higher order Bessel functions compared to the zeroth decay rapidly with their order, and it is justified to introduce a truncation. If any of the indices of a tensor element are larger in magnitude than a certain value $n_{\max}$, we approximate the tensor element by zero. The compatibility of this type of truncation with the symmetries of the model is discussed in Sec. XIV.A.

The tensor formulation here can be seen from another perspective, i.e., by using the SVD on each nearest neighbor factor, and then doing the $\varphi$ integrals. The Fourier expansion of the nearest neighbor interaction given by Eq. (117) can be understood as the spectral decomposition of the Boltzmann weight. Here the “matrix” $e^{i\varphi}$ is unitary, and is parameterized by the “indices” $\varphi$ and $n$. To find the singular values, we multiply by the Hermitian conjugate and diagonalize (just as in the Ising case),

$$\begin{align*}
\int_0^{2\pi} \frac{d\varphi_2}{2\pi} e^{\beta \cos(\varphi_1 - \varphi_2)} e^{\beta \cos(\varphi_1 - \varphi_2)} & = \int_{\varphi_1}^{2\pi} \frac{d\varphi_2}{2\pi} \sum_{n,m} e^{i\varphi_1} I_n e^{-i\varphi_2} I_m e^{-i\varphi_2} \\
& = \sum_{n,m} e^{i\varphi_1} I_n I_m e^{-i\varphi_2} \delta_{m,n} \\
& = \sum_{n} e^{i\varphi_1} I_n^2 e^{-i\varphi_2}
\end{align*}$$

which gives the expected result, that the singular values are the absolute value of the modified Bessel functions, $\lambda_n(\beta) = |I_n(\beta)|$. We see from these examples that performing the character (or Fourier) expansion of the nearest neighbor interaction during the tensor formulation is similar to the SVD of that same factor, and in fact when the coupling is positive, they are equivalent.

For convenience when dealing with the group $U(1)$ (or $O(2)$ for that matter) we factorize all the $I_n(\beta)$ factors which dominate the small $\beta$ regime and define the ratios

$$t_n(\beta) = \frac{I_n(\beta)}{I_0(\beta)} = \begin{cases} \frac{1 - \frac{\beta^2}{2n!} + O(1/\beta^2)}{I_0(\beta)}, & \beta \to \infty \\ \frac{\beta^n}{2n!} + O(\beta^{n+2}), & \beta \to 0 \end{cases}$$

This factorization helps elucidate the role of the boundary conditions in the tensor formulation. As discussed in Sec. V.C, with open boundary conditions, one sets the boundary tensor indices to 0, which in the above definition reduces the weights on those links to value 1.
B. \textit{q}-state clock models

The results of this section hold for the $Z_q$ restrictions. The infinite sums are replaced by finite sums with $q$ values. The modified Bessel functions are replaced by their discrete counterparts:

$$ I_n(\beta) \to I_n^{(q)}(\beta) \equiv (1/q) \sum_{t=0}^{q-1} e^{\beta \cos(\frac{2 \pi}{q} t)} e^{-in\frac{2 \pi}{q} t}, \quad (124) $$

which in the large $q$ limit turns into the usual integral formula. In the Ising case ($q = 2$), we have

$$ I_0(\beta) \to \cosh(\beta), \quad \text{and} \quad I_1(\beta) \to \sinh(\beta). \quad (125) $$

The selection rules in Eq. (121) remain valid modulo $q$.

For recent numerical TRG-inspired work on clock models and discussion of the second transition see (Chen \textit{et al.}, 2017; Li \textit{et al.}, 2020).

C. Dual reformulations with unconstrained variables

In Secs. III.D and V.B, we mentioned the possibility of expressing the closed paths of the expansion in powers of $\tanh(\beta)$ of the Ising model using dual variables. These ideas can be generalized for a large class of models with Abelian symmetries (Banks \textit{et al.}, 1977; Einhorn and Savit, 1978, 1979; Kogut, 1979; Savit, 1977, 1980). In this subsection, we discuss the case of spin models, with interactions on links. Models with interactions on plaquettes and higher dimensional simplices will be discussed in Sec. IX.C.

Consistent with the rest of the article, in this subsection we use a Euclidean metric with all lower indices as well as implicit summations of repeated indices in order to make a stronger connection with the covariant formulation of Maxwell’s equations, unless specified otherwise. The discrete form of Noether current conservation, given in Eq. (121), which also holds modulo-$q$ for $q$-state clock models can be written in a compact way as

$$ \nabla_{\mu} n_{\mu} = 0, \quad (126) $$

where $\nabla_{\mu}$ is a discrete derivative

$$ \nabla_{\mu} f_x = f_{x-\mu} - f_x. \quad (127) $$

Since more indices will appear we keep the reference to the site $x$ implicit. Following the example of Maxwell’s equations written in a relativistically covariant way, we can express a conserved current as the gradient of an antisymmetric tensor of order 2:

$$ n_{\nu} = \nabla_{\mu} C_{\mu\nu}. \quad (128) $$

This holds in arbitrary dimensions $D$.

Because of the divergenceless condition Eq. (126), $n_{\mu}$ has $D - 1$ degrees of freedom per link. On the other hand, $C_{\mu\nu}$ has $D(D - 1)/2$ degrees of freedom which is $(D - 1)(D - 2)/2$ more than $D - 1$. The redundancy which appears for $D \geq 3$ can be made more obvious by introducing a dual tensor with $D - 2$ indices (Savit, 1977, 1980) and also $D(D - 1)/2$ components:

$$ C_{\mu\nu} = \frac{1}{(D - 2)!} f_{\mu\nu\mu_1...\mu_{D-2}} C_{\mu_1...\mu_{D-2}}. \quad (129) $$

The $\tilde{C}$ field is precisely the dual field which lives on the dual lattice. If we plug this dual form into Eq. (128), we see that $\tilde{C}_{\mu_1...\mu_{D-2}}$ can be shifted by antisymmetrized derivatives of lower rank tensors. For $D = 3$, $\tilde{C}_\mu$ has 3 components and is defined up to a gradient so we end up with the desired 2 independent degrees of freedom. For $D \geq 4$, the redundancy becomes nested: we need to count the redundancy of the redundancy etc. For instance for $D = 4$, $\tilde{C}_{\mu\nu}$ has 6 components. The shift by the gradient of a 4-vector naively subtracts 4 degrees of freedom, but this 4-vector can itself be shifted by a gradient without affecting the initial shift and we end up with $6 - 4 + 1 = 3$ degrees of freedom.

D. Chemical potential, complex temperature, and importance sampling

Since the tensor renormalization group—and tensor network methods generally—do not rely on sampling from probability measures, situations where sampling methods would falter or fail due to the loss of real, positive definite weights never arise. Instead, only linear algebra is needed in the form of tensor contractions. This allows the method to address the “sign problem”, which can occur during the inclusion of a chemical potential, and a complex coupling. In (Denbleyker \textit{et al.}, 2014) this was addressed in the case of a complex temperature. The authors studied the zeros in the complex temperature plane, \textit{i.e.} Fisher zeros, and found the tensor renormalization group is able to out perform the re-weighting method using in classical Monte Carlo studies that involve imaginary parts of the action.

In Refs. (Yang \textit{et al.}, 2016; Zou \textit{et al.}, 2014) a purely real chemical potential, $\mu$, is added to the action of the two-dimensional $O(2)$ nonlinear sigma model in the form,

$$ S_{\mu} = -\beta \sum_{x,\mu} \cos(\theta_{x+\mu} - \theta_x - i\mu). \quad (130) $$

In this form, the action has a complex sign problem. The authors studied the phase diagram of the model in the $\beta$-$\mu$ plane both in the discrete time, and continuous time limits. Reference (Banerjee and Chandrasekharan, 2010) studied this action as well using a sampling method known as the “worm algorithm” using a Fourier expanded
form of the Boltzmann weight which eliminates the sign problem completely.

In fact, the worm algorithm is intimately related to the tensor formulation. When the tensor elements which appear in a tensor formulation of a model are positive definite, that model admits a formulation of the worm algorithm where tensor elements are weighted against each other in an “accept-reject” Metropolis style algorithm. In this way, a lattice configuration is populated by tensor indices at their respective locations (sites, links, etc.). These indices correspond to tensor elements, and hence weights. These weights are used to update the tensor indices throughout the lattice using the standard Metropolis update. Moreover, the tensor interpretation—along with the exact blocking procedure (Liu et al., 2013)—allows one to use the worm algorithm on exactly coarse-grained lattice models (when the coarse-grained tensor elements are again positive-definite) and therefore use the renormalization group exactly with Monte Carlo studies. This procedure of “tensor sampling” is an interesting direction and deserves more attention.

VIII. Tensors for Spin Models with Non-Abelian Symmetries

A. \(O(3)\) Nonlinear Sigma Model

Consider the action for the \(O(3)\) nonlinear sigma model in \(D\) dimensions,

\[
S = -\beta \sum_{x=1}^{N} \sum_{\mu=1}^{D} \sum_{a=1}^{3} \sigma_{x+\mu}^{(a)} \sigma_{x}^{(a)}
\]

\[
= -\beta \sum_{x=1}^{N} \sum_{\mu=1}^{D} \cos \gamma_{x+\mu, x},
\]

where \(\gamma\) is the angle in the plane created by the two vectors.

As discussed before, the important insight in constructing the tensors here is to find a formulation which introduces discrete variables (the tensor indices) and factorizes the partition function in a such a way that the original field variables can be integrated out. We will discuss two different possible ways to do this here. One will use the Fourier expansion, and express the model in terms of its irreducible representations, while the second uses the Taylor expansion to create discrete variables (the natural numbers) which become the tensor indices.

The first tensor construction is based on the global symmetry group of the model. This construction has been explored and used successfully in Refs. (Bruckmann et al., 2019; Liu et al., 2013; Unmuth-Yockey et al., 2015) for classical tensor network calculations and matrix product-states calculations in the Hamiltonian formulation. Here we give the classical tensor formulation. Since each term in the action is a dot product between vectors of length-one, we can expand on basis functions for the sphere, i.e. the spherical harmonics. First consider the partition function,

\[
Z = \int d\Omega e^{-S} = \prod_{x} \int d\Omega_{x} \prod_{x, \mu} e^{\beta \cos \gamma_{x+\mu, x}},
\]

where \(d\Omega\) is the normalized measure on \(S_{2}\), i.e. \(d\Omega = -d(\cos \theta) d\phi / 4\pi\) since each Boltzmann factor is a function of the cosine of the angle between the vectors we can expand using Legendre polynomials straight-forwardly.

\[
e^{\beta \cos \gamma_{x+\mu, x}} = \sum_{l=0}^{\infty} \frac{2l + 1}{4\pi} A_{l}(\beta) P_{l}(\cos \gamma_{x+\mu, x}).
\]

This step is advantageous since it gives the \(A\) coefficients only \(l\) dependence. The \(A\) can be solved for by inverting the above,

\[
A_{l}(\beta) = 4\pi l! \frac{\partial}{\partial \beta} j_{l}(-i\beta),
\]

where \(j_{n}(z)\) are the spherical Bessel function. The Legendre Polynomials can then be rewritten in terms of spherical harmonics using the addition theorem for spherical harmonics,

\[
P_{l}(\cos \gamma_{x+\mu, x}) = \frac{4\pi}{2l + 1} \sum_{m=-l}^{l} Y_{lm}^{*}(\theta_{x+\mu}, \phi_{x+\mu}) Y_{lm}(\theta_{x}, \phi_{x}).
\]

This step separates the dependence on the coupled \(x+\mu\) and \(x\) sites and allows the factors to be treated individually. With the \(\theta\) and \(\phi\) dependence decoupled between neighboring sites we can perform the angular integration for the field at each site. In \(D\) dimensions there are \(2D\) nearest-neighbors for each site, giving an integral of the form,

\[
\int d\Omega_{x} \prod_{\mu=1}^{D} Y_{(lm)_{\lambda\mu}}(\theta_{x}, \phi_{x}) Y_{(lm)_{-\lambda\mu}}^{*}(\theta_{x}, \phi_{x}).
\]

This integral can be evaluated with the use of the Clebsch-Gordan series,

\[
Y_{l_{1}m_{1}}(\theta, \phi) Y_{l_{2}m_{2}}^{*}(\theta, \phi) = \sum_{l=|l_{2}-l_{1}|}^{L} \sum_{M=-L}^{L} C_{l_{1}m_{1}l_{2}m_{2}}^{LM} C_{l_{1}00m_{2}}^{L0} \times \sqrt{\frac{(2l_{1} + 1)(2l_{2} + 1)}{4\pi(2L + 1)}} Y_{LM}(\theta, \phi),
\]

along with the orthogonality of the spherical harmonics.
We now restrict to $D = 2$ and continue explicitly. The integral in Eq. (136) takes the form,

$$\int d\Omega_x Y_{(lm)_{x,1}}^* Y_{(lm)_{x,2}}^* Y_{(lm)_{x,3}}^* Y_{(lm)_{x,4}}^* (\theta_x, \phi_x).$$  (138)

If we make the change of notation for $(lm)_{x,1}$ with $l_1 m_1$, $(lm)_{x,2}$ with $l_2 m_2$ etc..., we find for Eq. (138),

$$\int d\Omega_x Y_{l_1 m_1} Y_{l_2 m_2} Y_{l_3 m_3} Y_{l_4 m_4} (\theta_x, \phi_x)$$  

$$= \sqrt{(2l_1 + 1)(2l_2 + 1)(2l_3 + 1)(2l_4 + 1)} \times$$

$$\sum_{L,M} \frac{1}{(4\pi)^2 (2L + 1)} C_{lm_1 l_2 m_2}^{LM} C_{lm_3 l_4 m_4}^{LM} C_{l_1 0 l_2 0}^{LO} C_{l_3 0 l_4 0}^{LO}.$$

The Clebsch-Gordan coefficients constrain the surrounding $l$'s around a site to satisfy the triangle inequalities, and enforce a conservation law between the $m$'s. This constraint must be imposed at every site. Let us define the constraint from Eq. (139) as $C_x$, along with a composite index defined formally as $L \equiv (l, m)$ which has dimension $(l_{\text{max}} + 1)^2$. This index contains all the states from $l = 0$ up to some $l_{\text{max}}$ given by $\sum_{l=0}^{l_{\text{max}}}(2l + 1) = (l_{\text{max}} + 1)^2$. We can write down the local tensor as,

$$T_{l_{x-1},l_{x},L_{x-2},L_{x-2}}^{(x)} = \sqrt{A_{l_{x-1},1} A_{l_{x},1} A_{l_{x-2},1} A_{l_{x-2},1}} C_x.$$  (140)

The constraint $C$ in Eq. (139) is somewhat complicated, but the physical content is that it simply demands the four $l$'s around a site to satisfy the triangle inequalities according to the typical addition of angular momenta, while enforcing a conservation law on the $O(2)$ subgroup $m$. A pleasing feature of this formulation is that the weights, $A_x$, only depend on $l$.

The second tensor formulation for the $O(3)$ nonlinear sigma model uses the Taylor expansion of the Boltzmann weight to recast the model in terms of discrete fields. This formulation follows directly from (Bruckmann et al., 2016, 2015; Wolff, 2010).

Starting with the partition function,

$$Z = \int \mathcal{D}\Omega \ e^{-S}$$

$$= \prod_{x=1}^{N} \frac{1}{4\pi} \int \sin \theta_x d\theta_x d\phi_x \prod_{x=1}^{N} \prod_{\mu=1}^{2} \prod_{a=1}^{3} e^{\beta \sigma_x^{(a)} \sigma_x^{(a)} + \mu}.$$  (141)

We now expand the Boltzmann weight in a Taylor series,

$$e^{\beta \sigma_x^{(a)} \sigma_x^{(a)} + \mu} = \sum_{n_{x,\mu}^{(a)}=0}^{\infty} \frac{\beta n_{x,\mu}^{(a)}}{n_{x,\mu}^{(a)}!} \left( \sigma_x^{(a)} \sigma_x^{(a)} + \mu \right)^{n_{x,\mu}^{(a)}}$$  (142)

associating three natural numbers, $n_{x,\mu}^{(a)}$, with each link. Reordering and collecting the same spin-field at a site we can write the partition function as,

$$Z = \sum_{\{n\}} \left( \prod_{x} \prod_{\mu} \prod_{a} n_{x,\mu}^{(a)} \right) \times$$

$$\left( \prod_{x} \prod_{\mu} \prod_{a} \frac{1}{4\pi} \int (\sigma_x^{(a)})^n_x \sin x d\theta_x d\phi_x \right).$$  (143)

The first factors in parenthesis are the new weights associated with a configuration of $n$s. The integrals inside the second factor in parenthesis must be evaluated for each site. They are all identical, so we focus on a single site and perform the integration. For one site the integral we must evaluate looks like

$$\frac{1}{4\pi} \int (\sigma_x^{(1)})^n_x \sin x d\theta_x d\phi_x.$$  (144)

Using the explicit expressions for $\sigma^{(1)}$, $\sigma^{(2)}$, and $\sigma^{(3)}$ in terms of $\phi$ and $\theta$, we can perform the $\phi$ and $\theta$ integrals separately. We find for $\theta$,

$$\Theta_x = \frac{1}{2} \int_{0}^{\pi} \sin \theta_x \sum_{\mu=1}^{2} (n_{x,\mu}^{(b)} + n_{x,\mu}^{(b)}) + 1 \times$$

$$(\cos \theta_x) \sum_{\mu=1}^{2} (n_{x,\mu}^{(b)} + n_{x,\mu}^{(b)}) d\theta_x.$$  (145)

and for $\phi$,

$$\Phi_x = \frac{1}{2\pi} \int_{0}^{2\pi} \sin \phi_x \sum_{\mu=1}^{2} (n_{x,\mu}^{(b)} + n_{x,\mu}^{(b)}) \times$$

$$(\cos \phi_x) \sum_{\mu=1}^{2} (n_{x,\mu}^{(b)} + n_{x,\mu}^{(b)}) d\phi_x.$$  (146)

Eq. (145) can be computed exactly and gives,

$$\Theta_x = \frac{1}{2} \int_{0}^{\pi} \sin \theta_x \sum_{\mu=1}^{2} (n_{x,\mu}^{(b)} + n_{x,\mu}^{(b)}) + 1 \times$$

$$(\cos \theta_x) \sum_{\mu=1}^{2} (n_{x,\mu}^{(b)} + n_{x,\mu}^{(b)}) d\theta_x$$

$$= \frac{1}{2} \delta_{\text{mod}2} \sum_{\mu} (n_{x,\mu}^{(b)} + n_{x,\mu}^{(b)}) \times$$

$$B \left[ \frac{1}{2} \left( 1 + \sum_{\mu} (n_{x,\mu}^{(b)} + n_{x,\mu}^{(b)}) \right) \right],$$  (147)

where $B(p, q)$ is the beta function, and $\delta_{\text{mod}2}$ is the Kronecker delta but the indices need only be equal modulo.
2. Similarly Eq. (146) can be computed as well giving,

\[
\Phi_x = \frac{1}{2\pi} \int_0^{2\pi} (\sin \phi_x)^\sum_{\mu}(n_\mu^{(2)}+n_{\bar{\mu}}^{(2)}) d\phi_x \\
= \frac{1}{\pi} \sum_{\mu}(n_\mu^{(1)}+n_\mu^{(1)}) \times B \left( \frac{1}{2} \left(1 + \sum_{\mu}(n_\mu^{(1)}+n_{\bar{\mu}}^{(1)}) \right) \right) , \\
\frac{1}{2} \left(1 + \sum_{\mu}(n_{\bar{\mu}}^{(1)}+n_{\bar{\mu}}^{(1)}) \right) .
\] (148)

These are two constraints which need to be imposed at each site of the lattice. With these constraints and weights from the Taylor series expansion we can now define a tensor at every lattice site.

First, define a collective index given by \( N_{x,\mu} \equiv n_\mu^{(1)} \otimes n_{\bar{\mu}}^{(2)} \otimes n_\mu^{(3)} \), as well as a weight associated with a link,

\[
w_{x,\mu} = \frac{\beta \frac{1}{2} \sum_{\mu} n_\mu^{(1)}}{\sqrt{\sum_{\mu} \left(n_\mu^{(2)}+n_{\bar{\mu}}^{(2)}\right)}}. \tag{149}
\]

Now the tensor at site \( x \) in \( D \) dimensions is given by,

\[
T_{N_{x,\mu}=1,N_{x,\mu}=1,...,N_{x,\mu}=D} = \prod_{\mu=1}^{D} w_{x,\mu,\mu} w_{x,\mu} \Theta_x \Phi_x.
\] (150)

This tensor has the nice property that—assuming \( \beta > 0 \)—the tensor elements are positive. This follows from the positivity of \( \beta \) and that the \( n_\mu \) are non-negative. The constraints coming from the \( \theta \) and \( \phi \) integrals are positive as well, since the \( \beta \) functions are positive for positive arguments. This allows this formulation to be used in sampling methods, which it has been (see (Bruckmann et al., 2016; Wolff, 2010)). However, there are more indices necessary in this description, which increases the cost numerically in a tensor renormalization group algorithm.

B. SU(2) principal chiral model

The SU(2) principal chiral model consists of a SU(2) matrix associated with each site on the lattice that interacts with its nearest neighbors. The action on a \( D \)-dimensional square lattices with periodic boundary conditions is given by,

\[
S = -\frac{\beta}{2} \sum_{x=1}^{N} \sum_{\mu=1}^{D} \text{Tr}[U_x U_{x+\mu}^\dagger]. \tag{151}
\]

The partition function for the model is given as the Haar integration over each of the fields on the lattice,

\[
Z = \int \mathcal{D}U e^{-S} = \prod_x \int dU_x e^{\frac{\beta}{2} \sum_{\mu=1}^{D} \text{Tr}[U_x U_{x+\mu}^\dagger]}. \tag{152}
\]

The partition function only depends on the trace of group elements which means the characters of the group can be expanded on. We expand the Boltzmann weight,

\[
e^{\frac{\beta}{2} \sum_{\mu=1}^{D} \text{Tr}[U_x U_{x+\mu}^\dagger]} = \sum_{r=0}^{\infty} F_{r,\mu}(\beta) \chi^r(\mu)(U_x U_{x+\mu}^\dagger) \tag{153}
\]

where the sum runs over all half-integer irreducible representations of the group. The expansion coefficients can be solved for by inverting Eq. (153) using the orthogonality of the characters. The characters are traces over matrix representations of the group. This allows the group elements to be split and factorized, \( \chi^r(U_x U_{x+\mu}^\dagger) = \sum \chi^r_a(U_x) \chi_{a+\mu}^r(U_{x+\mu}) \), and subsequently integrated over. Collecting all the \( D \) matrices associated with the same site we find an integral of the form,

\[
\int dU_x \prod_{\mu=1}^{D} D_x^r \chi^r(\mu)(U_x). \tag{154}
\]

where the matrix indices have been suppressed. We can perform this integral with the help of the Clebsch-Gordan series,

\[
D_{m_1n_1}^r(U) D_{m_2n_2}^s(U) = \sum_{r_1+r_2=r} \sum_{M=-R}^{R} \sum_{N=-R}^{R} C_{r_1m_1r_2m_2}^{RM} C_{r_1n_1r_2n_2}^{RN} D_{m_3n_3}^{r_1} D_{m_4n_4}^{r_2} \tag{155}
\]

along with the orthogonality of the \( D \) matrices\(^1\).

We now restrict to the case of \( D = 2 \). Eq. (154) takes the form,

\[
\int dU_x D_{m_1n_1}^{r_1} D_{m_2n_2}^{r_2} D_{m_3n_3}^{r_3} D_{m_4n_4}^{r_4} (U_x) \tag{156}
\]

and using the steps mentioned above we find,

\[
\int dU_x D_{m_1n_1}^{r_1} D_{m_2n_2}^{r_2} D_{m_3n_3}^{r_3} D_{m_4n_4}^{r_4} (U_x) = \sum_{R,M,N} \sum_{R,M,N} \sum_{r_1+r_2+r_3+r_4=r} \sum_{m_3=m_4} \sum_{n_3=n_4} C_{r_1m_1r_2m_2}^{RM} C_{r_1n_1r_2n_2}^{RN} \tag{157}
\]

\(^1\) The orthogonality is with the complex conjugate, not Hermitian conjugate:
where \( d_r = 2r + 1 \) is the dimension of the representation. This is the constraint associated with a site. Similar to the \( O(3) \) nonlinear sigma model, this constraint constrains the surrounding representation numbers on the links around a site through the triangle inequalities. If we define the constraint from Eq. (157) as \( C_r \), and define a composite index formally as \( X_{r,1} \equiv \{ r_{x,1}, m, n \} \), where \( m \) and \( n \) are the matrix indices naturally associated with an \( r \) on a link. Then we can define a local tensor at each site by,

\[
T_{X_{r,1},X_{r,2},Y_{r,2}} = \sqrt{F_{r_{x,1}}F_{r_{x,2}}F_{r_{x-1,1}}F_{r_{x-2,2}}(\beta)} C_r.
\]

By contracting this tensor with itself recursively one rebuilds the original partition function.

A possible alternative way to formulate a local tensor is to use the same discrete variables used in (Gattringer et al., 2018a). This formulation follows along the same lines as the second tensor formulation for the \( O(3) \) nonlinear sigma model by expanding the Boltzmann weight in a Taylor series. However, in this reference this formulation was used in sampling methods. We do not attempt to give the tensor formulation using these variables, but the required steps seem straightforward, and mimic the steps in the second formulation of the \( O(3) \) tensor.

### C. Truncations and asymptotic freedom

An important question is how do the previous tensor formulations, and specifically the expansions before-hand which lead up to the tensor definitions, affect universality. Looking at Eq. (135) for the \( O(3) \) nonlinear sigma model it is clear that this expansion does not affect the global \( O(3) \) invariance of the model, since one expands on the dot product between nearest neighbor vectors. This interaction is \( O(3) \) invariant, so long as each spin is rotated by the same rotation matrix, and so a polynomial in this interaction remains \( O(3) \) invariant. Similarly a truncation in the \( l \) variable to a finite \( l_{\text{max}} \) leaves the expansion \( O(3) \) invariant for the same reason. Because of this, after truncation but before integration, the model consists of a local nearest neighbor interaction which is \( O(3) \) invariant and in the same number of dimensions we started in indicating, naively, that this truncated model lies in the same universality class as the original \( O(3) \) nonlinear sigma model. Likewise, the expansion in Eq. (142) is also globally \( O(3) \) invariant. So any truncation on the \( n \) variables leaves the expansion only dependent on spins which interact with their neighbors in an \( O(3) \) invariant fashion.

The \( O(3) \) nonlinear sigma model in two dimensions is known to be asymptotically free (Hasenfratz et al., 1990). On a two-dimensional lattice, the continuum limit is approached by taking the nearest-neighbor coupling, \( \beta \), infinitely large. In this limit one expects the mass gap to obey the continuum perturbative result which predicts (Hasenfratz et al., 1990),

\[
am = \frac{8}{\epsilon} a \Lambda_{\text{MS}} = 128\pi\beta \exp(-2\pi\beta).
\]

An initial study of the asymptotic scaling of the mass gap in this model was done in (Unmuth-Yockey et al., 2015) using the tensor renormalization group. There they compared tensor renormalization group calculations of the mass gap with results from Monte Carlo simulations. They found a slow convergence to the expected result as a function of \( l_{\text{max}} \). A more thorough study in the Hamiltonian limit was done in (Bruckmann et al., 2019) using matrix product states at different truncations and different volumes. While they found relatively good convergence to the asymptotic result for \( l_{\text{max}} > 2 \) and in the large volume limit, which support this idea of universality, the lower \( l_{\text{max}} \) values did not converge as well.

For the action in Eq. (151) one has the freedom to rotate all group elements by the same global matrix like \( U_x \rightarrow U_x' = V U_x V^\dagger \). This leaves the action invariant, as well as the measure. The expansion in Eq. (153) retains this freedom since the thing that is expanded on is the trace of group elements which is the type of interaction which allows for this freedom. Furthermore, a truncation on the sum of representations in Eq. (153) does nothing to this symmetry. Again, a truncation then naively leaves the model with the same nearest neighbor interaction with the same symmetries in the same number of dimensions and one expects this truncated model to lie in the same universality class as the original principal chiral model.

### IX. Tensors for lattice gauge theories

In this section we will discuss gauge theories with Abelian symmetries. The gauge Ising model is the simplest model that we can consider. However, as for the spin models, we will start with the continuous case and then obtain the models with discrete symmetries such as the Ising model and the gauge clock models using the substitutions described in Sec. VII.B.

#### A. Pure gauge \( U(1) \)

The partition function for the pure gauge \( U(1) \) model introduced in Sec. II.B reads

\[
Z_{\text{PG}} = \prod_{x,\mu} \int_{-\pi}^{\pi} \frac{dA_{x,\mu}}{2\pi} e^{-S_{\text{Wilson}}},
\]

where \( d_r = 2r + 1 \) is the dimension of the representation. This is the constraint associated with a site. Similar to the \( O(3) \) nonlinear sigma model, this constraint constrains the surrounding representation numbers on the links around a site through the triangle inequalities. If we define the constraint from Eq. (157) as \( C_r \), and define a composite index formally as \( X_{r,1} \equiv \{ r_{x,1}, m, n \} \), where \( m \) and \( n \) are the matrix indices naturally associated with an \( r \) on a link. Then we can define a local tensor at each site by,

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\]

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\]
with the action
\[
S_{\text{Wilson}} = -\beta_{\text{pl}} \sum_{x, \mu < \nu} \cos(A_{x, \mu} + A_{x+\hat{\mu}, \nu} - A_{x+\hat{\nu}, \mu} - A_{x, \nu}).
\] (161)

It possess a local symmetry
\[
A'_{x, \mu} = A_{x, \mu} - (\alpha_{x+\hat{\mu}} - \alpha_x).
\] (162)

Using the Fourier expansion
\[
e^{\beta_{\text{pl}} \cos(A_{x, \mu} + A_{x+\hat{\mu}, \nu} - A_{x+\hat{\nu}, \mu} - A_{x, \nu})} =
\sum_{m_{x, \mu, \nu} = -\infty}^{+\infty} e^{im_{x, \mu, \nu}(A_{x, \mu} + A_{x+\hat{\mu}, \nu} - A_{x+\hat{\nu}, \mu} - A_{x, \nu})} I_{m_{x, \mu, \nu} (\beta_{\text{pl}})},
\]
(163)
to factorize the gauge fields, and integrating over \(A_{x, \mu}\) using the orthgonality of the \(U(1)\) elements, we obtain the selection rule
\[
\sum_{\nu > \mu} [m_{x, \mu, \nu} - m_{x, \nu, \mu}] + \sum_{\nu < \mu} [-m_{x, \mu, \nu} + m_{x, \nu, \mu}] = 0.
\] (164)

In \(D\) dimensions, there are \(2(D-1)\) plaquettes attached to each link. This selection rule constrains the \(m\) values associated with those plaquettes. It is convenient to introduce a tensor that is associated with the plaquettes of the lattice. It has four indices which can be naturally associated with the four links bounding a plaquette. Since each plaquette has a single \(m\) value associated to it, the four tensor legs attached to a given plaquette \((x, \mu)\) must carry the same index, \(m\). Following the terminology of Ref. (Liu et al., 2013), we introduce a “\(B\)-tensor” for each plaquette
\[
B^{(x, \mu)}_{m_1, m_2, m_3, m_4} = \begin{cases} 
  t_{m_1} (\beta_{\text{pl}}), & \text{if all } m_i \text{ are the same} \\
  0, & \text{otherwise.} 
\end{cases} 
\] (165)

where \(t_m\) is defined in Sec. VII.A. The \(B\)-tensors are assembled (traced) together with “\(A\)-tensors” attached to links with \(2(D-1)\) legs orthogonal to the link \((x, \mu)\)
\[
A^{(x, \mu)}_{m_1, \ldots, m_{2(D-1)}} = \delta_{m_{m, \text{out}}},
\] (166)
where \(\delta_{m_{m, \text{out}}}\) is a short notation for Eq. (164). Notice that in contrast to Ref. (Liu et al., 2013), the weight of the plaquettes is carried by the \(B\)-tensor. The partition function with PBC can now be written as
\[
Z = (I_0(\beta_{\text{pl}}))^{V(D-1)/2} \times \text{Tr} \prod_{l} A^{(l)}_{m_1, \ldots, m_{2(D-1)}} \prod_{\text{pl.}} B^{(\text{pl.})}_{m_1, m_2, m_3, m_4},
\] (167)
where the trace means index contraction following the geometric procedure described above. The tensor assembly is illustrated in Fig. 20 for \(D = 2\) and in Fig. 21 for \(D = 3\).

**B. Discrete Maxwell equations**

In the previous section we again found a selection rule which could be recast as a constraint on the \(m\) values which surround a link. Here we will show that Eq. (164) represents a discrete version of Maxwell’s equations: \(\partial_{\mu} F^{\mu \nu} = 0\). For this purpose, we define the “electric integers”
\[
e_{x,j} \equiv m_{x,j D},
\] (168)
with \(j = 1, \ldots, D-1\), which are associated with temporal plaquettes and which can be interpreted as electric fields. Eq. (164) for \(\mu = D\) reads
\[
\sum_{j=1}^{D-1} (e_{x,j} - e_{x-j,j}) = 0.
\] (169)

This is a discrete form of Gauss’s law, \(\nabla \cdot \mathbf{E} = 0\), in the source-free model.
For $D \geq 3$, we can introduce magnetic fields in a dimension dependent way. For $D = 3$, we define
\[ b_x \equiv m_{x,12}. \]  
Eq. 164 for $\mu = 1, 2$ are
\[ e_{x,1} - e_{x-\tau,1} = -(b_x - b_{x-\tau}), \]
\[ e_{x,2} - e_{x-\tau,2} = (b_x - b_{x-\tau}). \]  
These are a discrete version of the $D = 3$ Euclidean Maxwell’s equations
\[ \partial_1 B = \partial_2 E_2, \]
\[ \partial_2 B = -\partial_1 E_1, \]  
with $B = F^{12}$. However, there is no discrete equation corresponding to the Maxwell equation for the dual field strength tensor
\[ \partial_\mu \epsilon^{\mu\nu\sigma} F_{\nu\sigma} = 0. \]  
An example of a legal configuration violating the discrete version of Eq. (173), also written $\partial_\tau B = -\nabla \times E$, can be constructed.

For $D = 4$, we define
\[ b_{x,j} \equiv \epsilon_{jkl} m_{x,kl}, \]  
and obtain a discrete version of
\[ \partial_\mu \mathbf{E} = -\nabla \times \mathbf{B}, \]  
with the Euclidean magnetic field
\[ F^{jk} = +\epsilon^{jk} B^j. \]  
Note that the sign in Eq. (175) is different in Euclidean and Minkowskian spaces. Again there is no discrete version of the homogeneous equations for the dual field strength $\partial_\tau \mathbf{B} = -\nabla \times \mathbf{E}$ and $\nabla \cdot \mathbf{B} = 0$.

C. Abelian gauge duality

The dual construction of Sec. V.B for spin models can be extended for models with plaquettes and higher-dimensional simplex interactions (Savit, 1977). First, if we define $m_{\mu\nu} = -m_{\nu\mu}$ when $\mu > \nu$, the discrete Maxwell’s Eqs. (164) take the obvious form
\[ \nabla_\nu m_{\mu\nu} = 0, \]  
As explained in Sec. VII.C this expression is divergenceless and represent $D - 1$ conditions. We can introduce a dual tensor with $D - 3$ indices (Savit, 1977, 1980)
\[ m_{\mu\nu} = \frac{1}{(D-3)!} \epsilon_{\mu\nu\rho_1...\rho_{D-3}} \nabla_{\rho} \hat{C}_{\rho_1...\rho_{D-3}}, \]  
which provides an automatic solution of Eq. (177). After using the $D - 1$ conditions of Eq. (177) we are left with $(D-1)(D-2)/2$ independent components for $m_{\mu\nu}$. For $D = 3$, there is no redundancy and we have one degree of freedom. For $D = 4$, $\hat{C}_\mu$ is defined up to a gradient and we recover the 3 degrees of freedom.

D. The compact Abelian Higgs model

The compact Abelian Higgs model (CAHM) is a gauged version of the $O(2)$ model where the global symmetry under a $\varphi$ shift becomes local
\[ \varphi'_x = \varphi_x + \alpha_x. \]  
Its partition function is
\[ Z_{CAHM} = \prod_{x,\mu} \int^{\pi}_{-\pi} \frac{d\varphi_x}{2\pi} \prod_{x,\mu} \int^{\pi}_{-\pi} \frac{dA_{x,\mu}}{2\pi} e^{-S_{Wilson} - S_{U(1)}}, \]  
with
\[ S_{U(1)} = -\beta \sum_{x,\mu} \cos(\varphi_x + \mu \varphi_x + A_{x,\mu}) \]  
and $S_{Wilson}$ as in Eq. (161). Using the same $U(1)$ Fourier expansions as before, the $A$-field integration can be carried out. The integration over $A_{x,\mu}$ yields the selection rule
\[ \sum_{\nu > \mu} [m_{x,\mu\nu} - m_{x,\nu\mu}] + \sum_{\nu < \mu} [-m_{x,\nu\mu} + m_{x,\nu\nu}] + n_{x,\mu} = 0. \]  
which simply inserts the $n_{x,\mu}$ in Eq. (164) and corresponds to the Maxwell equations with charges and currents
\[ \partial_\mu F^{\mu\nu} = J^\nu. \]  
Equation (164) means that the link indices $n_{x,\mu}$ can be seen as determined by unrestricted plaquette indices $m_{x,\mu\nu}$. We write this dependence as $n_{x,\mu}(\{m\})$ as a shorthand for Eq. (182).

Note that for $n_{x,\mu}(\{m\})$, the discrete current conservation Eq. (121) is automatically satisfied (Meurice, 2019), and as long as the gauge fields are present, there is no need to enforce Eq. (121) independently. This is a discrete version of the fact that Maxwell’s equations with charges and currents (182) imply $\partial_\mu J^\mu = 0$.

With the introduction of the matter fields, we need to update the definition of the $A$-tensors. We now have quantum numbers on the links, $n_{x,\mu}$, which are completely fixed by the plaquette quantum numbers, and they bring a weight $t_{n_{x,\mu}}(\beta_\mu)$. This translates into
\[ A^{(x,\mu)}_{m_1...m_{D-1}} = t_{n_{x,\mu}}(\{m\})(\beta_\mu). \]  
since Eq. (182) gives $n_{x,\mu}$ in terms of the surrounding $m$s. The partition function with PBC can now be written as
\[ Z_{CAHM} = \left( I_0(\beta_{pl}) \right)^{V(D-1)/2} (I_0(\beta_\mu))^V \right) \prod_{x,\mu} A^{(x,\mu)}_{m_1...m_{D-1}} \prod_{x,\mu} P^{(x,\mu)}_{m_1m_2m_3m_4}. \]
SU(2) gauge theory

SU(2) gauge theory in D dimensions is governed by an action of the form,

$$S_{\text{Wilson}} = -\frac{\beta_D}{2} \sum_{x=1}^{N} \sum_{\mu<\nu=1}^{D} \text{ReTr}[U_{x,\mu}U_{x+\hat{\mu},\nu}U_{x+\hat{\nu},\mu}^{\dagger}].$$

(186)

In order to construct a local tensor we will proceed as before and use the character expansion, since the action only depends on the trace of group elements. The partition function for this model can be written as the Haar integration over the group elements on the links of the lattice,

$$Z = \int DU e^{-S_{\text{Wilson}}}$$

$$= \int DU_{x,\mu} \prod_{x} \prod_{\mu<\nu} \left[ e^{\beta_D \text{ReTr}[U_{x,\mu}U_{x+\hat{\mu},\nu}U_{x+\hat{\nu},\mu}^{\dagger}]} \right]$$

$$= \int DU_{x,\mu} \prod_{x} \prod_{\mu<\nu} \left[ e^{\beta_D \text{ReTr}[U_{x,\mu\nu}]} \right].$$

(187)

where $U_{x,\mu\nu}$ is the product of gauge fields around a plaquette. While this model is trivial in $D = 2$ for this model at the time of writing this; however, there have been tensor studies of other gauge models (Kuramashi and Yoshimura, 2019; Unmuth-Yockey, 2019; Zohar et al., 2015a, 2016).

To proceed we expand the Boltzmann weight,

$$e^{\frac{\beta_D}{2} \text{ReTr}[U_{x,\mu\nu}]} = \sum_{r,\nu,\mu=0}^{\infty} F_{r,\nu,\mu}(\beta_D) \chi_{r,\mu\nu}^{x}(U_{x,\mu\nu}).$$

(188)

This expansion associates an $r$ with each plaquette on the lattice. The characters can be written as the trace of the product of matrix representations of the group,

$$\chi_{r,\mu\nu}^{x}(U_{x,\mu\nu}) = \sum_{a,b,c,d}^{D} D_{ab}^{r,\mu\nu}(U_{x,\mu\nu}) D_{bc}^{r,\mu\nu}(U_{x+\hat{\mu},\nu}) \times$$

$$D_{cd}^{r,\mu\nu}(U_{x+\hat{\nu},\mu}) D_{da}^{r,\mu\nu}(U_{x,\nu}).$$

(189)

By factorizing the group elements in this way, we can perform the link integration link-by-link, reformulating the model in terms of the discrete representations, and the matrix indices. In $D$ dimensions, there are $2(D-1)$ plaquettes associated with each link. The integral over the group element associated with link $(x,\mu)$ then has the form

$$\int dU_{x,\mu} \prod_{\nu<\mu} D_{x,\mu\nu}^{r,\nu} D_{x-\hat{\mu},\mu}^{r,\nu} \prod_{\nu>\mu} D_{x,\mu\nu}^{r,\nu} D_{x-\hat{\nu},\mu}^{r,\nu},$$

(190)

where the matrix indices have been suppressed, and the $D$-matrices are all the same $U_{\mu}$ rotation matrix or its Hermitian conjugate. This integral is in general quite complicated but is simplified by using the Clebsch-Gordan series to systematically reduce Eq. (190) to an integral over only two $D$-matrices. The Clebsch-Gordan series is given by,

$$D_{m_{1},n_{1}}^{R} D_{m_{2},n_{2}}^{R} = \sum_{R_{1},R_{2}}^{R} \sum_{M=-R}^{R} \sum_{N=-R}^{R} C_{RM}^{R_{1}R_{2}} C_{R_{1}R_{2}N}^{R} D_{MN}^{R},$$

(191)

Using this we can collect the daggered, and non-daggered $D$-matrices in Eq. (190), and simplify them in pairs. The $D$-matrices are orthogonal according to,

$$\int dU \; D_{mn}^{R}(U) D_{m'n'}^{R}(U) = d_{r}^{-1} \delta_{r,\nu'} \delta_{mm'} \delta_{nn'}$$

(192)

with $d_{r} = 2r + 1$ the dimension of the representation.

The final expression is tedious to write down, but there is nothing subtle about it. Here we will write the final expression for $D = 3$ for a link in the $\mu = 2$ direction, and subsequently write the local tensors for $D = 3$ as well,

$$\int dU_{x,2} D_{m_{1},n_{1}}^{r_{2},23} D_{m_{2},n_{2}}^{r_{3},23 \dagger} D_{m_{3},n_{3}}^{r_{4},12} D_{m_{4},n_{4}}^{r_{5}-1,12},$$

$$= \sum_{R,M,N}^{R} d_{R}^{-1} C_{RM}^{r_{2}r_{3}} C_{R_{2}r_{1},-1,12}^{R_{1}} C_{r_{2}r_{3},m_{1},n_{1}}^{R_{1}} C_{r_{4}r_{5},-1,12}^{n_{4}}$$

$$C_{r_{4},12,m_{2},n_{2}}^{n_{3}} C_{r_{5},12,n_{3}}^{R_{2}} C_{r_{5},12,n_{3}}^{R_{2}},$$

(193)

Then for each link in the lattice there is a constraint of this form. Let us define this constraint as $C(x,\mu)$ for a general link at site $x$ in direction $\mu$. If we define a composite index formally as $R_{x,\mu\nu} = \{r_{x,\mu\nu}, m_{1}, n_{1}\}$ we can define a tensor associated with the links of the lattice whose indices are associated with the shared plaquettes as,

$$A_{R_{x,\mu\nu}}^{(x,\mu)} R_{x-\hat{\mu},\nu} R_{x,\mu\nu} R_{x-\hat{\nu},\mu} = C_{R_{x,\mu\nu}}^{(x,\mu)}$$

(194)

for $\nu \neq \rho \neq \mu$. The $A$ tensor is defined as the constraint on a link. An illustration of the three-dimensional tensor can be found in Fig. 22.

This is not the whole story though, since this tensor is not enough to reproduce the partition function of the original model. The weight factors, $F_{r}(\beta)$, still need to be accounted for. To include the weight factors we define an additional tensor associated with the plaquettes of the lattice; however, there is a slightly subtle aspect with this tensor. That is the circulation of the $D$-matrix indices in Eq. (189) around the plaquette. These indices—which are now a part of the $A$ tensor—are still required to be contracted in the pattern found in Eq. (189). To enforce
Figure 22 An illustration of the three-dimensional tensor $A$ for $SU(2)$ gauge theory in three dimensions.

In this circulation we assign Kronecker deltas to the new tensor in such a way that the contraction pattern of the matrix indices in Eq. (189) is reproduced. To be clear, consider Eq. (189) again, rewritten,

$$r_{x;}(U_{x;}) = D_{r_{x;}}^{ab}(U_{x;})D_{r_{x;}}^{cd}(U_{x;})D_{r_{x;}}^{ef}(U_{x;})D_{r_{x;}}^{gh}(U_{x;})D_{r_{x;}}^{ha}(195)$$

with an implied sum over repeated indices here. These Kroneker deltas will be moved onto the new plaquette tensor,$$B_{(r_1ab)(r_2cd)(r_3ef)(r_4gh)} = \delta_{bc} \delta_{de} \delta_{fg} \delta_{ha} \times \begin{cases} F_{pl}^{\beta} & \text{if all } rs \text{ are the same} \\ 0 & \text{otherwise.} \end{cases} (196)$$

Each index of the $B$ tensor is associated with one of the four links which border the plaquette. This makes this tensor identical for all dimensions. An illustration of this tensor can be seen in Fig. 23. By contracting this $B$ tensor on the plaquettes with the $A$ tensors on the links the full partition function is constructed exactly. The contraction pattern between these indices can be seen in Fig. 21.

F. The non-Abelian Higgs model

The lattice $SU(2)$ gauge-Higgs model in $D$ dimensions consists of three main parts. There is the pure Yang-Mills lattice action, a gauge-matter interaction term, and a matter potential term. For the pure Yang-Mills term we use the standard Wilson action,

$$S_{\text{Wilson}} = -\frac{\beta_{\text{pl.}}}{2} \sum_x \sum_{\mu<\nu} \text{ReTr}[U_{x;\mu}U_{x;\nu}U_{x;\mu}^{\dagger}U_{x;\nu}]$$

where one takes a product of the gauge fields associated with the links around an elementary square (plaquette) for each square of the lattice. For the gauge-matter coupling term we have,

$$S_{U(2)} = -\frac{K}{2} \sum_{x=1}^{N} \sum_{\mu=1}^{D} \Phi_x^{\dagger} U_{x;\mu} \Phi_x.$$ (198)

The $\Phi$ field can be re-expressed in terms of a $2 \times 2$ matrix (Montvay and Münster, 1994) and the gauge-matter term becomes,

$$S_{U(2)} = -\frac{K}{2} \sum_x \sum_{\mu} \text{ReTr} \left[ \phi_x^{\dagger} U_{x;\mu} \phi_x \right],$$ (199)

where $\phi$ is now a $2 \times 2$ matrix. Since $\phi_x^{\dagger} \phi_x = \rho_x^2 \mathbb{1}$, $\phi_x$ can be written as $\phi_x = \rho_x \alpha_x$ with $\rho_x \in \mathbb{R}$, $\rho_x \geq 0$, and $\alpha_x \in SU(2)$. This expresses $\phi_x$ in terms of the Higgs ($\rho_x$) and Goldstone ($\alpha_x$) modes, respectively. This allows the gauge-matter term to be again re-written as,

$$S_{U(2)} = -\frac{K}{2} \sum_x \sum_{\mu} \rho_x \rho_x \text{ReTr} \left[ \alpha_x^{\dagger} U_{x;\mu} \alpha_x \right].$$ (200)

Finally, the Higgs potential,

$$V = \sum_x (|\Phi_x|^2 + \lambda(|\Phi_x|^2 - 1)^2)$$ (201)

only couples same-site fields and therefore—in terms of the matrix $\phi_x$—only involves the Higgs mode,

$$V = \sum_x \rho_x^2 + \lambda(\rho_x^2 - 1)^2.$$ (202)
The partition function for this model is then,

\[ Z = \int DU \, D\rho \, D\alpha \, e^{-S_{\text{Wilson}} - S_U(2)} - V \]  

(203)

where the integration over \( U \) and \( \alpha \) is the \( SU(2) \) Haar measure, and the integration measure over \( \rho \) is given by \( \rho^3 \, d\rho \) over \( [0, \infty) \).

We only consider the limit in which \( \lambda \to \infty \), \( \rho_x \to 1 \), which is when the Higgs mode becomes infinitely massive. In addition we perform a change of variables on the gauge which is when the Higgs mode becomes infinitely massive. Similar steps to the previous Sec. IX.E and constructions in the Hamiltonian formulation have been considered (Kühn et al., 2015). In fact the expansion for the Yang-Mills term is identical to Eqs. (188) and (189). The expansion over only two matrices using the singular value decomposition, exactly. This contraction pattern is shown in Fig. 20. As mentioned before, the tensor associated with the plaquettes from the pure Yang-Mills term is the same as Eq. (196), and is the same regardless of dimension for the \( SU(2) \) gauge-Higgs model as well. With the \( A \) and \( B \) tensors mentioned here, one can contract them in the appropriate pattern to construct the partition function exactly. This contraction pattern is shown in Fig. 20.

In fact, in \( D = 2 \) it is possible to go one step further and define a single tensor which can be contracted with itself to construct the partition function. Since the \( A \) tensor only has two indices, it can be factored into a product of two matrices using the singular value decomposition,

\[ A_{RR'} = \sum \alpha, \beta U_{\alpha \beta} \lambda_{\alpha \beta} V_{\beta R'}^\dagger \]

(211)

As mentioned before, the tensor associated with the plaquettes from the pure Yang-Mills term is the same as Eq. (196), and is the same regardless of dimension for the \( SU(2) \) gauge-Higgs model as well. With the \( A \) and \( B \) tensors mentioned here, one can contract them in the appropriate pattern to construct the partition function exactly. This contraction pattern is shown in Fig. 20.

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the same and real,

\[ A_{RR'} = \sum_{\gamma} \left( \sum_{\alpha} U_{R\alpha} \sqrt{\lambda_{\alpha\gamma}} \right) \left( \sum_{\beta} \sqrt{\lambda_{\gamma\beta}} U^{T}_{\beta R'} \right) = \sum_{\gamma} L_{R\gamma} L^{T}_{\gamma R'}. \quad (213) \]

Using the \( L \) matrix and the \( B \) tensor we can make a local tensor associated with a plaquette by summing over all four \( R \) indices in the \( B \) tensor,

\[ T_{\alpha \beta \gamma \delta} = \sum_{R_1R_2R_3R_4} B_{R_1R_2R_3R_4} L_{R_1\alpha} L_{R_2\beta} L_{R_3\gamma} L_{R_4\delta}. \quad (214) \]

In this form it is clear that contracting this tensor with itself appropriately one reconstructs the \( A \) matrix at each link which is already manifestly contracted with the adjacent \( B \) tensors, and hence recursively contracting this \( T \) tensor with itself constructs the partition function.

Within this tensor reformulation it is also possible to define the Polyakov loop straightforwardly. For \( SU(2) \) in the fundamental representation the Polyakov loop at site \( x^* \) is given by

\[ P_{x^*} = \text{Tr} \left[ \prod_{n=0}^{N_x-1} D^{\frac{1}{2}} \langle U_{x^*+n^*}\rangle \right], \quad (215) \]

where \( \tau \) indicates a direction chosen as time. The expectation value of this operator is,

\[ \langle P \rangle = \frac{1}{Z} \int \mathcal{D}U \ P \ e^{-\mathcal{S}}. \quad (216) \]

One can recast this average in terms of local tensors by performing the same steps as before. The only difference in this case is that for a particular spatial site, \( x^* \), all the temporal links have an additional \( D \)-matrix associated with them, altering the integral found in, say, Eq. (209) by the inclusion of a fourth \( D \)-matrix whose representation is \( 1 \). However, one proceeds as before using Eq. (191) to make the integral manageable. The integral on the temporal links of the Polyakov loop have the form,

\[ \sum_{k} \int dU_{x,2} D_{kk}^{r^*} D_{m_1 n_1}^{r^*} D_{m_2 n_2}^{r^*} D_{ij}^{\frac{1}{2}} = \sum_{k,R,M,N} C_{R}^{RM} C_{R}^{MN} C_{R}^{MN} \]

\[ \int dU_{x,2} D_{MN}^{r^*} D_{m_1 n_1}^{r^*} D_{ij}^{\frac{1}{2}} = \sum_{k,R,M,N} d_{r^*}^{-1} C_{R}^{RM} C_{R}^{MN} C_{R}^{MN} \]

If we define this constraint as \( \tilde{C}_{R_{r^*12},R_{r^*12}12} \) then we can write down the tensor on the Polyakov loop links,

\[ \tilde{A}_{R_{r^*12},R_{r^*12}12} = \sum_{r^*2} F_{r^*2} \tilde{C}_{R_{r^*12},R_{r^*12}12} \quad (218) \]

This tensor has two more indices than the typical \( A \) tensor. This is because of the additional \( D \)-matrix from the Polyakov loop insertion. These additional matrix indices are contracted with each other and traced over as in the definition in Eq. (215).

With the tensors from Eqs. (214) and (218) it is possible to using coarse-graining schemes to approximate the free energy, and compute expectation values. Using the higher-order tensor renormalization group for the case of \( D = 2 \), in (Bazavov et al., 2019) derivatives of the free energy were computed along with the Polyakov loop, and Polyakov loop correlator. Of the derivatives of the free energy, one of primary interest is the average of the gauge-matter interaction, and its fluctuations,

\[ \langle L_\phi \rangle = \frac{1}{V} \frac{\partial \ln Z}{\partial \kappa} \quad \chi_{\phi} = \frac{1}{V} \frac{\partial^2 \ln Z}{\partial \kappa^2}. \quad (219) \]

These were computed while taking the continuum limit. The continuum limit in this model is controlled by the Yang-Mills coupling and the system volume, since the Yang-Mills coupling is dimensionful in \( D = 2 \). By fixing the ratio, \( \beta/V = c \), with \( c \) a constant, and increasing the system volume one approaches the fixed-volume continuum limit. An interesting result from this study was evidence for a cross-over transition between a confining—pure Yang-Mills—regime, and a Higgs-regime. This can be seen from the expectation value of the squared fluctuations of the gauge-matter interaction in Fig. 24. In this figure one can see a gradual convergence as the continuum limit is approached, and the presence of a peak around \( \kappa \approx 1.4 \) separating the two regimes.

This is further supported by the behavior of the Polyakov loop correlation function on either side of the peak value. Figs. 25 and 26 show examples of the potential between static charges—\( \mathcal{V} \)—in the \( \kappa < 1.4 \) and \( \kappa > 1.4 \) regimes, respectively. The potential is found from the logarithm of the correlator. In Fig. 25 there is a linear confining potential which persists for long distances as the continuum limit is approached. In the Higgs-like regime seen in Fig. 26 there is a linear potential for short distances, however after a certain distance the potential flattens and the forces between charges is zero (string breaking).

**X. Tensors for Models with (Non-Compact) Scalars**

For tensor networks with discrete indices, scalar fields have to be discretized in a proper manner. There are sev-
In this section we discuss the cases of Lagrangian path integrals. Specifically, we consider TRG studies of the real Yang-Mills theory, and a Higgs regime where string-breaking occurs. Here $\beta/V = 0.01$ was held fixed. The colored symbols are computed using the HOTRG, while the hollow black markers are from Monte Carlo data as a check. The maximum representation used in the HOTRG calculation was $r = 1$, and the final number of states kept was 50.

There is a peak around $\kappa \approx 1.4$ which seems to indicate a cross-over between a confining regime characteristic of a pure Yang-Mills theory, and a Higgs regime where string-breaking occurs. Here $\beta/V = 0.01$ was held fixed. The maximum representation used in the calculation was $r = 1$, and the final number of states kept in the calculation was 50.

Several ways to apply simple discretization rules to scalar fields to make transfer matrices (Chung, 1999; Iblisdir et al., 2007; Lay and Rudnick, 2002; Nishiyama, 2001a,b). In this section we discuss the cases of Lagrangian path integrals. Specifically, we consider TRG studies of the real $\phi^4$ theory, and the complex $\phi^4$ theory in two dimensions.

This section is organized as follows. We first present the definition of the real $\phi^4$ model in Sec. X.A. In Sec. X.B, a tensor network representation of the real $\phi^4$ model is made via the Gaussian quadrature rule. In Sec. X.C, the definition of the complex $\phi^4$ model and its tensor network representation are given.

Figure 24 The susceptibility in the gauge-matter interaction. For this value of $V$ we see string breaking at small systems $r = 0$, as a function of distance, taking the continuum limit. Here we find the phenomenon of string breaking which persists even as we take the continuum limit after some distance. Here $\beta/V = 0.01$ was held fixed. The maximum representation used in the calculation was $r = 1$, and the final number of states kept in the calculation was 50.

Figure 25 The potential, $V$, between two static charges, for $\kappa = 0.5$, as a function of distance, taking the continuum limit. For this value of $\kappa$ we see string breaking at small systems $r = 0$, but as the large lattice volume limit is taken we find a linear potential across long distances. Here $\beta/V = 0.01$ was held fixed. The maximum representation used in the calculation was $r = 1$, and the final number of states kept in the calculation was 50.

Figure 26 The potential, $V$, between two static charges, for $\kappa = 2$, as a function of distance, taking the continuum limit. Here we find the phenomenon of string breaking which persists even as we take the continuum limit after some distance. Here $\beta/V = 0.01$ was held fixed. The maximum representation used in the calculation was $r = 1$, and the final number of states kept in the calculation was 50.

### A. Real $\phi^4$ theory

The Euclidean action of the real $\phi^4$ theory in two dimensions is

$$S_{\text{cont.}} = \int d^2 x \left\{ \frac{1}{2} \left( \partial_\mu \phi \right)^2 + \frac{\mu^2_0}{2} \phi^2 + \frac{\lambda}{4} \phi^4 \right\}, \tag{220}$$

where $\mu_0$ and $\lambda$ are the bare mass and the bare coupling, respectively. $\phi$ is an one-component real scalar field. This model possesses the spontaneous breaking of the $Z_2$ symmetry, where the expectation value of the field $\langle \phi \rangle$ is an order parameter.

From here on, we treat the model on a square lattice with periodic boundary conditions. Using the lattice spacing $a$, the dimensionless parameters are defined by

$$\mu^2 = a^2 \mu^2_0, \quad \lambda = a^2 \lambda. \tag{221}$$

Then the lattice action is given by

$$S_{\text{scalar}} = \sum_x \left\{ \frac{1}{2} \sum_{\nu=1}^2 \left( \phi_{x+\mu} - \phi_x \right)^2 + \frac{\mu^2_0}{2} \phi_x^2 + \frac{\lambda}{4} \phi_x^4 \right\}, \tag{222}$$

where $x$ is the lattice coordinate and $\hat{\nu}$ denotes the unit vector along the $\hat{\nu}$-direction.

In two-dimensional scalar theories, one has to take care of the divergence of the one-loop self energy. In this section the following renormalization condition for the squared-mass,

$$\mu^2 = \mu^2_0 + 3 \lambda A (\mu^2), \tag{223}$$
is used to define the renormalized squared-mass $\mu$. $A(\mu^2)$ denotes the one-loop self energy on the lattice

$$A(\mu^2) = \frac{1}{V} \sum_{k_1,k_2=1}^{N} \frac{1}{\mu^2 + 4 \sin^2(\pi k_1/N) + 4 \sin^2(\pi k_2/N)}$$

with the lattice volume $V = N \times N$. To provide numerical results the nonlinear equation in Eq. (223) is solved to translate the bare squared-mass into the renormalized one. Note that the coupling constant is free of the renormalization; this is a common property in two dimensional scalar theories.

### B. Tensors from Gaussian quadratures

In this subsection a tensor network representation of the real $\phi^4$ theory is derived by using the Gaussian quadrature rule. This method is given and used in tensor network studies for Lagrangian path integrals in (Kadoh et al., 2018, 2019, 2020). For the original tensor network study of the real $\phi^4$ model, consult (Shimizu, 2012). The new tensor network representation discussed in this section is more accurate than the original one by Shimizu although here we just draw the attention of the reader to the original work and do not conduct any quantitative comparison.

The partition function on the lattice is defined by

$$Z = \left( \prod_x \int_{-\infty}^{\infty} \right) e^{-S_{\text{scalar}} - S_h},$$

where

$$S_h = \sum_x -h\phi_x.$$

For later use we introduce the external field $h$ here. An important step to build a tensor network representation is generating discrete degrees of freedom that are the candidates for tensor indices. Then, in the following, we mainly discuss how to extract the discrete degrees of freedom from the continuous and non-compact scalar fields.

Since the action has only the nearest neighbor interactions, the Boltzmann weight can be rewritten as a product of local factors

$$e^{-S_{\text{scalar}} - S_h} = \prod_x \prod_{\nu=1}^2 f(\phi_x, \phi_{x+\nu})$$

where the local factor is given by

$$f(\phi_1, \phi_2) = \exp \left\{ -\frac{1}{2} (\phi_1 - \phi_2)^2 - \frac{\mu^2}{8} (\phi_1^2 + \phi_2^2) - \frac{\lambda}{16} (\phi_1^4 + \phi_2^4) + \frac{h}{8} (\phi_1 + \phi_2) \right\}.$$

To derive the discrete formula, we briefly summarize the Gaussian quadrature rule for a weighted integral of a single variable function. We consider to discretize the (well-defined) target integral of a function $g$

$$I = \int_{-\infty}^{\infty} dx \, W(x) \, g(x),$$

where $W$ is a weight function. A successful way to discretize this type of integral is the Gaussian quadrature method. The quadrature rule gives a simple replacement of the integral with a discrete summation

$$I \approx \sum_{i=1}^{K} w_i \, g(y_i),$$

where $y_i$ and $w_i$ are the $i$-th root of the order $K$ orthonormal polynomial, and corresponding weight, respectively. Comprehensive definitions for the Gaussian quadrature rule (including the definition of weights) are given in (Abramowitz and Stegun, 1965). The species of the orthonormal polynomial is one’s choice and corresponds to the form of the weight function $W$. Typical choices are the Legendre polynomials, and the Hermite polynomials that correspond to $W(y) = 1$ and $W(y) = e^{-y^2}$, respectively. If we consider that the mass term in the action plays the role of the weight function, it seems to be natural to use the Hermite polynomials. Indeed, this choice has been used in Refs. (Kadoh et al., 2018, 2019, 2020), and we exclusively use the Hermite polynomials for the Gaussian quadrature rule in this sectionootnote{As mentioned in the main text, the choice does not matter to the numerical accuracy as long as the degree of the orthonormal polynomial is sufficiently large.}.

When $g$ is a polynomial function of degree $2K - 1$ or less, the Gaussian quadrature reproduces the exact value. Even if not, if $g$ is well approximated by a polynomial function of degree $2K - 1$ or less, the Gaussian quadrature would be accurate. We apply this quadrature rule to each integral of the scalar field in the path integral.

By applying the Gauss–Hermite quadrature to the partition function, a discrete form is introduced as

$$Z(K) = \prod_{\{\alpha\}} \prod_x w_{\alpha_x} e^{v_{\alpha_x}} \prod_{\nu=1}^{2} f(\phi_{\alpha_x}, \phi_{\alpha_{x+\nu}}).$$


where \( \sum_{\{\alpha\}} \) denotes \( \prod_x \sum_{\alpha_x=1}^{K} \). The discrete form depends on the order of the Hermite polynomial \( K \), and this parameter would be set to be large for accurate results. In practice \( K \geq 64 \) could be regarded as sufficiently large (Kadoh et al., 2018, 2019, 2020), though, in a later section on numerical results from a TRG analysis of the real \( \phi^4 \) theory, \( K = 256 \) is taken.

Note that the method is applied numerically and that so far there is no analog of character expansions and orthogonality relations used in the previous sections. In Sec. XII.D we show that for the Gauss-Hermite quadratures, it is possible to interpret the construction in terms of a truncated version of the harmonic oscillator algebra of creation and annihilation operators.

In Eq. (231) the local Boltzmann factors can be regarded as \( K \times K \) matrices, and one can perform the SVD for them:

\[
f(y_{\alpha x}, y_{\beta x}) = \sum_{i_{x,v}=1}^{K} U_{\alpha x\beta x} \lambda_{i_{x,v}} V_{i_{x,v}\beta x}^\dagger , \tag{232}
\]

where \( \{\lambda\} \) is the singular values that are assumed to be in descending order \( \lambda_1 \geq \lambda_2 \geq \cdots \lambda_K \geq 0 \), and \( U \) and \( V \) are unitary matrices. Now a tensor network representation of \( Z(K) \) is defined by

\[
Z(K) = \left( \sum_{\{X,T\}} \prod_x T(K)_{X_{x-1}X_xT_xT_{x-1}} \right) \tag{233}
\]

where \( \sum_{\{X,T\}} \) denotes \( \prod_x \sum_{X_x=1}^{K} \sum_{T_x=1}^{K} \), and we have made the replacement \( i_{x,1} \rightarrow X_x \) and \( i_{x,2} \rightarrow T_x \). The tensor (at any site) is defined by

\[
T(K)_{ijkl} = \sqrt{\lambda_i \lambda_j \lambda_k \lambda_l} \sum_{\alpha=1}^{K} w_{\alpha} e^{y_{\alpha}^2} U_{\alpha \alpha} U_{\alpha \beta} V_{\beta \gamma}^\dagger V_{\gamma \alpha}^\dagger . \tag{234}
\]

At this stage the bond dimension of the tensors is \( K \). To take a balance of computational cost and numerical accuracy, one may initially truncate the bond dimension to a certain value \( D_{\text{cut}} \leq K \). In (Kadoh et al., 2019), \( D_{\text{cut}} \leq 64 \) is taken for actual computations and the sufficiency of this choice is numerically shown.

Physical quantities can also be expressed as tensor networks. A key point is to respectively treat the denominator and the numerator in the right-hand side of

\[
\langle \phi \rangle = \frac{Z_1}{Z} , \tag{235}
\]

where

\[
Z_1 = \left( \prod_{x} \int_{-\infty}^{\infty} d\phi_x \right) \phi_x e^{-S_h - S_{\text{scalar}}} \tag{236}
\]

with \( \bar{x} \neq x \). The presence of \( \phi_x \) does not affect the tensor construction procedure, it merely alters what integral is being approximated, so Gaussian quadrature rule and the SVD of the local factors works as before. However, the resultant tensor network representation contains an "impurity tensor" owing to \( \phi_x \). To perform a coarse-graining of a tensor network that contains impurities, one needs a little ingenuity; the details are shown in (Kadoh et al., 2019). Finally, one can calculate \( Z \) and \( Z_1 \) separately, and using them the value of \( \langle \phi \rangle \) is obtained using Eq. (235).

Here the numerical results for the real \( \phi^4 \) theory are shown. The target quantity in this section is the critical coupling constant and its continuum limit value; in (Kadoh et al., 2019), the continuum limit extrapolation proceeds as: 1) Take the thermodynamic and zero-external field limits to get a susceptibility \( \chi \) for given (bare) mass and (bare) coupling constant, 2) Find the critical mass where \( \chi \rightarrow \infty \), 3) Extract renormalized critical squared-mass from Eq. (223), 4) Compute the dimensionless critical coupling constant \( \lambda^2/\mu_0^2 \), 5) Vary \( \lambda \) from 0.1 to 0.005 and repeat above procedure to take the continuum limit, 6) Take a linear extrapolation to find the critical coupling constant at \( \lambda = 0 \). As mentioned in the previous sections, the parameters for the tensor network analysis are set to \( K = 256 \) and \( D \leq 64 \). The legitimacy of this choice is confirmed numerically in (Kadoh et al., 2019).

Figures 27 and 28 show the results of the thermodynamic limit and zero-external field limit, respectively. In both cases, (bare) parameters and the bond dimension of the tensor are \( \mu_0^2 = -0.1006174 \), \( \lambda = 0.05 \), and \( D_{\text{cut}} = 32 \) as an example. In Fig. 27 the ratio \( \langle \phi \rangle / h \) behaves as a constant in the extremely large space-time volume where \( L \geq 10^6 \), so that one can consider that the system reaches the thermodynamic limit for \( L \geq 10^6 \). In Fig. 28 the ratio also behaves as a constant for \( h \leq 10^{-11} \), so that \( \langle \phi \rangle \approx \chi h \) holds. Then the susceptibility \( \chi \) can be obtained from the relation.

From the susceptibilities for several masses, the critical mass where \( \chi \rightarrow \infty \) is determined. In (Kadoh et al., 2019),

\[\text{Note that a fast decay of the singular values guarantees the accuracy of such an approximation. Although the decay rate would be weak near the criticality, a notable accuracy of the critical coupling constant is achieved in (Kadoh et al., 2019). This is reviewed in the later section.}\]

\[\text{4 Because of the translation invariance, a subscript that denotes the coordinate in Eq. (235) is omitted.}\]

\[\text{5 Note that, if we do not omit showing the lattice spacing \( \alpha \), \( \alpha \lambda \rightarrow 0 \) means the continuum limit.}\]

\[\text{6 Actually the ratio shows a quadratic behavior for \( h \leq 10^{-11} \), so it is proper to take the susceptibility using more suitable fitting function. In (Kadoh et al., 2019) the susceptibility is defined in such a way.}\]
In (Kadoh et al., 2019) the following fitting formula is used:

$$\chi^{-1/1.75} = A \left| \mu_{0, c} - \mu_0^2 \right|^{\gamma/1.75}. \quad (237)$$

Figure 29 shows the susceptibility with the fit result with fixed $\gamma = \gamma_{\text{sing}}$. The critical bare squared-mass $\mu_{0, c}$ is obtained as $\mu_0^2$-intercept of the line. The parameters are set to $\lambda = 0.05$ and $D_{\text{cut}} = 32$.

Taking above procedures for several values of $\lambda$ one can obtain the dimensionless critical coupling constant $\lambda/\mu_0^2$ as a function of $\lambda$, and the remaining procedure is to take the continuum extrapolation. In (Kadoh et al., 2019), the $\lambda = 0$ value of the dimensionless critical coupling is calculated by a linear extrapolation with a reasonable chi-squared value $\approx 0.026$. The result is

$$\lim_{\lambda \to 0} \frac{\lambda}{\mu_0^2(\lambda)} = 10.913(56). \quad (238)$$

The error is mainly due to a fluctuation in the large $D_{\text{cut}}$-region. In the paper, it is shown that the $D_{\text{cut}}$-dependence is the main source of the error.

Figure 30 shows a comparison among recent Monte Carlo works in (Bosetti et al., 2015; Bronzin et al., 2019; Schaich and Loinaz, 2009; Wozar and Wipf, 2012) and the TRG work in (Kadoh et al., 2019). The TRG result shows notable accuracy and has achieved the smallest $\lambda$-value that is essentially important for the continuum extrapolation.

In (Kadoh et al., 2019) the legitimacy of fixing $\gamma$ to the exact value is supported by reasonable reduced chi-squared values for fittings.
C. Complex $\phi^4$ theory

The continuum action of the complex $\phi^4$ theory with finite chemical potential $\mu$ is given by

$$S_{\text{cont.}} = \int d^2 x \left\{ \sum_{\nu=1}^{2} \left| \partial_{\nu} \varphi \right|^2 + \left( m^2 - \mu^2 \right) |\varphi|^2 \right\} + \mu \left( \varphi^* \partial_2 \varphi - \varphi \partial_2 \varphi^* \right) + g |\varphi|^4,$$

where $\varphi$, $m$, $g$ are the complex scalar field, the mass, and the coupling constant, respectively. This is a model of relativistic Bose gas with finite density and a typical model that suffers from the complex action problem (sign problem).

As in the case of the real $\phi^4$ theory, we introduce the lattice action that takes the form

$$S = \sum_{x} \left[ \left( 4 + m^2 \right) |\varphi_x|^2 + g |\varphi_x|^4 - \sum_{\nu=1}^{2} \left( \epsilon^{\mu \nu \alpha \beta} \varphi_x^{\mu} \varphi_{x+\nu}^{\alpha} + \epsilon^{\mu \nu \beta \alpha} \varphi_x^{\mu} \varphi_{x+\nu}^{\beta} \right) \right],$$

where the periodic boundary conditions are assumed and all parameters are dimensionless under the lattice units where $a = 1$. The partition function is defined by

$$Z = \int D\varphi e^{-S}.$$

(241)

Since the complex field $\varphi_x$ has two real components: $\varphi_x = (1/\sqrt{2}) \left( \varphi_x^{(1)} + i\varphi_x^{(2)} \right)$ with $\varphi_x^{(1)}, \varphi_x^{(2)} \in \mathbb{R}$, the measure in Eq. (241) $D\varphi$ means $\prod_x d\varphi_x^{(1)} d\varphi_x^{(2)}$. Hereafter we consider how to make a tensor network representation of Eq. (241). The same technologies used in the case of the real scalar field can be used in the complex case for the two real components which comprise the complex field. We follow the same procedure given in (Kadoh et al., 2020) while emphasizing the differences owing to the multi-component variables.

Since the action includes only the nearest neighbor interactions, the Boltzmann factor $e^{-S}$ can be expressed as a product of local factors:

$$e^{-S} = \prod_x f_1(\varphi_x, \varphi_{x+1}) f_2(\varphi_x, \varphi_{x+2}),$$

(242)

where

$$f_\nu(\varphi_1, \varphi_2) = \exp \left\{ - \left( 1 + \frac{m^2}{4} \right) \left( |\varphi_1|^2 + |\varphi_2|^2 \right) - \frac{g}{4} \left( |\varphi_1|^4 + |\varphi_2|^4 \right) + \epsilon^{\mu \nu \alpha \beta} \varphi_1^{\mu} \varphi_2^{\alpha} + \epsilon^{\mu \nu \beta \alpha} \varphi_1^{\mu} \varphi_2^{\beta} \right\},$$

(243)

The scalar fields are discretized by the Gauss–Hermite quadrature. An important point here is to discretize the two components of the complex fields independently:

$$Z(K) = \sum_{\{\alpha, \beta\}} \prod_x w_{\alpha_x} w_{\beta_x} \exp \left( y_{\alpha_x}^2 + y_{\beta_x}^2 \right) \prod_{\nu=1}^{2} f_\nu \left( \frac{y_{\alpha_x} + iy_{\beta_x}}{\sqrt{2}}, \frac{y_{\alpha_{x+\mu}} + iy_{\beta_{x+\mu}}}{\sqrt{2}} \right),$$

(244)

where $y_i$ and $w_i$ are the $i$-th root of the $K$-Hermite polynomial and corresponding weight, and $\sum_{\{\alpha, \beta\}}$ denotes $\prod_{\alpha, \beta} \sum_{\alpha_n=1}^{K} \sum_{\beta_n=1}^{K}$. Then the local factor now can be decomposed using the singular value decomposition:

$$f_\nu \left( \frac{y_{\alpha} + iy_{\beta}}{\sqrt{2}}, \frac{y_{\gamma} + iy_{\delta}}{\sqrt{2}} \right) = \sum_{i=1}^{K^2} U_{\nu}^{[\alpha \gamma]} \sqrt{\lambda^{[\nu]}_i} V_{\nu}^{[\nu \delta]} i, \ldots,$$

(245)

where $\{\lambda^{[\nu]}_i\}$ and $U^{[\nu]}_i$, $V^{[\nu]}_i$ are the singular values and unitary matrices, respectively. The tensor network representation of $Z(K)$ is given by

$$Z(K) = \sum_{\{X,T\}} \prod_x T(K)_x x_{x-1} x_{x} t_{x},$$

(246)

where a tensor at any site is given by

$$T(K)_{ijkl} = \sqrt{\lambda^{[1]}_{ij} \lambda^{[2]}_{kl} \lambda^{[1]}_{ij} \lambda^{[2]}_{kl}} \sum_{\alpha, \beta=1}^{K} w_{\alpha \beta} \exp \left( x_{\alpha}^2 + x_{\beta}^2 \right) \cdot U^{[1]}_{\{\alpha \gamma\}} U^{[2]}_{\{\alpha \beta\}} V^{[1]}_{i,ij} V^{[2]}_{i,ij},$$

(247)

the $\sum_{\{X,T\}}$ denotes $\sum_{x} \sum_{t_{x-1}=1}^{K^2}$, and we have used the same notation as in Sec. X.B.

Here the TRG results for complex $\phi^4$ theory are shown. The truncation order of the coarse-graining steps is called $D_{\text{cut}}$, and this is set to 64 in this whole subsection. Note that, as in the case of the real $\phi^4$ theory, the bond dimension of tensors is initially truncated. The bond dimension of the initial tensors are also $D_{\text{cut}} = 64$ in this subsection. The order of the Hermite polynomial $K$ is also set to 64. The sufficiency of these choices is discussed and validated with numerical results in (Kadoh et al., 2020).

Figure 31 shows the average value of the phase factor $e^{i\theta}$ for $m^2 = 0.01$ and $g = 1$. This quantity is defined as the ratio

$$\langle e^{i\theta} \rangle_{pq} = \frac{Z_{pq}}{Z_{pq}},$$

(248)

where $Z_{pq}$ is the phase quenched version of the partition function. The average phase factor shows the difficulty to conduct the Monte Carlo simulation, and then, from the figure, one can find that the severe sign problem occurs.
when the chemical potential $\mu$ and the volume $V$ are large.

Figures 32 and 33 show the particle number density and the expectation value of the squared-norm of $\varphi$, respectively. The number density is calculated by the numerical differentiation of the free energy:

$$\langle n \rangle = \frac{1}{V} \frac{\partial \ln Z}{\partial \mu}, \quad (249)$$

and $\langle |\varphi|^2 \rangle$ is calculated by using the impurity tensor method \(^8\). In the finite density systems, thermodynamic quantities do not react to the chemical potential below an onset in the non-zero $\mu$ region. This is called the Silver Blaze phenomenon. Using the TRG, the phenomenon is clearly observed even in the large $\mu$- and $V$-region, where the sign problem is severe in Monte Carlo simulations.

In this section we reviewed progress using the TRG on the real $\phi^4$ theory (Kadoh et al., 2019) and the complex $\phi^4$ theory (Kadoh et al., 2020). The former work supports the accuracy of the TRG approach, and the latter shows a sustainability of the TRG in the severe sign problem region. In both works, the Gaussian quadrature rule is made use of to discretize the scalar fields.

D. Open questions/Possible directions

For more precise works near criticality, it is natural to use improved coarse-graining algorithms such as the (loop-)TNR (Evenbly and Vidal, 2015; Yang et al., 2017), the graph-independent local truncations (GILT) (Hauru et al., 2018), and the full environment truncation (Evenbly, 2018). The common concept is to properly deal with the corner double line (CDL) structure on tensor networks (see (Gu and Wen, 2009) and Sec. VI.F). Recently the GILT has been applied to the 2D real $\phi^4$ model and a precise value of the critical coupling constant has been reported (Delcamp and Tilloy, 2020). The result is comparable to that of (Kadoh et al., 2019), recent Monte Carlo works, and other computational schemes. When using such a deterministic approach, the systematic error should be properly understood, and the definition of the error is important for more concrete discussions.

There could be a better scheme for generating tensor networks for scalar bosons although the tensor network representation via the Gaussian quadrature rule seems work quite well. One big concern is that the Gaussian quadrature rule effectively puts a cut-off for the scalar fields, so that one cannot deal with models whose local Boltzmann factor $f(\phi_1, \phi_2)$ has a long (or maybe infinite) tail in the $\phi_1$-$\phi_2$ space; e.g. massless free scalar bosons could not be suitably treated by the Gaussian quadrature rule, which requires a fast damping of the local Boltzmann factor. The authors of (Delcamp and Tilloy, 2020) generate discrete degrees of freedom using the Taylor series expansion instead of the Gaussian quadrature rules,

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\(^{8}\) Note that any local operator can be expressed by tensor networks including impurities. Although the number density can also be expressed directly by a tensor network, in (Kadoh et al., 2020) it was calculated via the differentiation to check the numerical stability.
XI. MODELS WITH FERMIONS

In this section we discuss tensor network representations, and coarse-graining algorithms for fermion systems. An important point is that in fermion tensor networks additional Grassmann variables are generated stemming from the original field variables. Since there are not Grassmann valued data types on computers, one needs some special treatments for the Grassmann variables on tensor networks. The coarse-graining procedure for fermion tensor networks was given in (Gu, 2013; Gu et al., 2010), and applications for relativistic fermion systems are given in (Shimizu and Kuramashi, 2014b; Takeda and Yoshimura, 2015).

A. Tensor representation for free Wilson fermions

In this section we construct a tensor network representation of the two-dimensional Wilson–Dirac fermion system. Interactions are not discussed here, but local interaction terms could be easily introduced as in (Kadoh et al., 2018; Shimizu and Kuramashi, 2014b; Yoshimura et al., 2018), where \( U(1) \), four fermion, and Yukawa-type interactions are discussed, respectively. The Lagrangian density of the target system is given by

\[
\mathcal{L}_x = \bar{\psi}_x (D\psi)_x
\]

where the Wilson–Dirac operator is defined by

\[
D_{x,x'} = (m + 2) \delta_{x,x'} - \frac{1}{2} \sum_{\mu=1}^2 \{(1 + \gamma_{1\mu}) \delta_{x,x'+\mu} + (1 - \gamma_{1\mu}) \delta_{x,x'-\mu}\}. \tag{251}\]

\( \psi \) and \( m \) are a two-component spinor field: \( \psi_x = (\psi_x^{(1)}, \psi_x^{(2)})^T \), and the mass, respectively. The Grassmann variables satisfy anti-commutation relations. We assume periodic boundary conditions in all directions in what follows in this section.

The partition function of the system is given by

\[
Z_F = \int D\psi D\bar{\psi} e^{-\sum \mathcal{L}_x}.
\]

Under this representation of gamma matrices:

\[
\gamma_1 = \sigma_1 = \begin{pmatrix} 0 & 1 \\ 1 & 0 \end{pmatrix}, \quad \gamma_2 = \sigma_3 = \begin{pmatrix} 1 & 0 \\ 0 & -1 \end{pmatrix}, \tag{253}
\]

the hopping factors are expanded as

\[
e^{-\sum_x \bar{\psi}_x (D\psi)_x} = \prod_x e^{-(m+2)\bar{\psi}_x \psi_x} \cdot \sum_{X_{x,1}} \left( \begin{array}{c} \bar{\chi}^{(1)}_{x+1} \chi^{(1)}_x \end{array} \right)^{X_{x,1}} \cdot \sum_{X_{x,2}} \left( \begin{array}{c} \bar{\chi}^{(2)}_{x+1} \chi^{(2)}_x \end{array} \right)^{X_{x,2}} \tag{254}\]

where \( \chi \) and \( \bar{\chi} \) are linear combinations of \( \psi \) and \( \bar{\psi} \):

\[
\psi_x = \begin{pmatrix} \psi_x^{(1)} \psi_x^{(2)} \end{pmatrix}, \quad \bar{\psi}_x = \begin{pmatrix} \bar{\psi}_x^{(1)} + \bar{\psi}_x^{(2)} \bar{\psi}_x^{(1)} - \bar{\psi}_x^{(2)} \end{pmatrix}, \quad \chi_x = 1/\sqrt{2} \left( \psi_x^{(1)} + \psi_x^{(2)}, \psi_x^{(1)} - \psi_x^{(2)} \right). \]

Each expansion is a binomial because of the nilpotency of Grassmann variables, and, at this point, discrete indices have arisen at each link.

Next, we integrate out the old degrees of freedom. An important point here is to break the hopping factors into Grassmann even structures to freely shuffle them between one another. To do that the following identities are useful:

\[
\begin{aligned}
\left( \psi^{(1)}_{x+\mu} \psi^{(1)}_x \right) \\
\left( \psi^{(2)}_{x+\mu} \psi^{(2)}_x \right)
\end{aligned}
\]

\[
\begin{aligned}
\left( \bar{\psi}^{(1)}_{x+\mu} \bar{\psi}^{(1)}_x \right) \\
\left( \bar{\psi}^{(2)}_{x+\mu} \bar{\psi}^{(2)}_x \right)
\end{aligned}
\]
the same coordinate are gathered up without involving sign factors. Then the tensor network representation of the partition function is defined by

$$Z_F = \sum_{\{X,T\}} \prod_x T_F x_{x-1} x_x T_{x+1} G_x x_{x-1} x_x T_{x+1},$$

where

$$T_{F,kijl} = \int dA(1) d\tilde{A}(1) dA(2) d\tilde{A}(2) e^{-(m+2)\Lambda},$$

$$\mathcal{A} = (A^{(1)}, A^{(2)})^T, \quad \tilde{A} = (\tilde{A}^{(1)}, \tilde{A}^{(2)}),$$

with

$$G_{x,kijl} = d\xi^{(1)}_{x^*+1} d\xi^{(2)}_{x*+2} d\xi^{(1)}_{x^*+1} d\xi^{(2)}_{x*+2} \cdot \left( \eta^{(1)}_{x^*+1} \right)^{i_1} \left( \eta^{(2)}_{x^*+1} \right)^{i_2} \cdot \left( \tilde{\eta}^{(1)}_{x^*+1} \right)^{j_1} \left( \tilde{\eta}^{(2)}_{x^*+2} \right)^{j_2},$$

with the new binary index \(m_i = (i_1 + i_2 + j_1 + j_2) \mod 2 = (k_1 + k_2 + l_1 + l_2) \mod 2 \). This decomposition is analogous to the decomposition that takes place in the original TRG. On the right-hand-side of Eq. (261), each factor is Grassmann-even thanks to the inclusion of the new Grassmann variables and the definition of the new binary index. This is similar to the construction of the fermion tensor network (see Eqs. (261)–(264)). \(x^*\) denotes the new coordinate on the coarse-grained square lattice, and the unit vectors on the coarse-grained lattice is defined by \(1^* = 1 + 2\) and \(2^* = 1 - 2\). (See also VLA.)

We have another way of decomposing \(G\)

$$G_{x,kijl} = (-1)^{i_1} d\gamma^{(2)}_{x^*+1} d\gamma^{(1)}_{x^*+2} \left( \tilde\gamma^{(2)}_{x^*+1} \right)^{i_1} \left( \tilde\gamma^{(1)}_{x^*+2} \right)^{i_2},$$

with the new binary index \(m_i = (i_1 + i_2 + j_1 + j_2) \mod 2 = (j_1 + j_2 + k_1 + k_2) \mod 2 \).

Using the above two ways of decomposing \(G\), we can now integrate out the old Grassmann variables, yielding a phase factor,

$$\int \Theta_{x+2} \Theta_{x+1} \Theta_{x} \Theta_{x+1} \Theta_{x+2} \Theta_{x+1} = (-1)^{f_{x+1} x_{x+2} \Theta_{x+1}} \Theta_{x+2} \Theta_{x+1} = (-1)^{f_{x+1} x_{x+2} \Theta_{x+1}} \Theta_{x+2} \Theta_{x+1}.$$

where

$$\Theta_{x+2} = X_{x+2}(X_{x+1} + X_{x+2}) + T_{x+1}(T_{x+1} + T_{x+2})$$

$$+ T_{x+2}(T_{x+1} + T_{x+2})$$

$$+ (X_{x+1} + X_{x+2} + X_{x+2} + X_{x+2})$$

$$\cdot (T_{x+1} + T_{x+2} + T_{x+1} + T_{x+2}).$$

Note that the details of the phase factor depend on the ordering of the \(\Theta_s\) in Eq. (267) and is not unique. Finally, the effect of the coarse-graining of the Grassmann part is interpreted in terms of a (non-Grassmann) phase factor and constraints:

$$(-1)^{T_{x+1} + T_{x+2} + X_{x+2} T_{x+1} + X_{x+2} T_{x+1}}$$

$$\cdot \delta(X_{x+2} + X_{x+2} + T_{x+1} + T_{x+2}) \mod 2, X_{x+2}$$

$$\cdot \delta(T_{x+1} + T_{x+1} + X_{x+2} + X_{x+2}) \mod 2, T_{x+2}$$

$$\cdot \delta(X_{x} + X_{x} + T_{x} + T_{x}) \mod 2, X_{x+1}$$

$$\cdot \delta(T_{x} + T_{x} + X_{x} + X_{x}) \mod 2, T_{x+1}.$$
where the indices labeled with “f” are the new binary indices introduced above. The phase and the Kronecker deltas are to be incorporated into the bosonic tensors, and the coarse-grained Grassmann part $G^*$ that consists of the new Grassmann variables is defined by

$$G^*_{x,kijl} = d_l^{ij} d_{x}^{ij} d_{x+1}^{ij} (\bar{\eta}_{x+1}, \eta_{x})^{ij} (\bar{\xi}_{x+2}, \xi_{x})^{ji}. \quad (270)$$

The above procedure is iteratively executed along with the normal coarse-graining steps for the bosonic tensors.

C. 2D Schwinger model with Wilson fermion

In (Shimizu and Kuramashi, 2014a), the critical behavior at $\theta = \pi$ of the two dimensional Schwinger model is studied with Wilson fermions. By studying the Fisher zeros, the authors of (Shimizu and Kuramashi, 2014a) have confirmed that there is a critical point around $\kappa = 0.2415$ and that the phase transition belongs to the 2D Ising universality class, where $\kappa$ is the inverse of the Wilson fermion mass $m$: $1/\kappa = 2 (m + 2)$. In addition, they have done the Lee–Yang zero analysis to seriously study the phase structure. Assuming that the gauge part of the lattice action is given by the usual Wilson action along with a topological term,

$$S_G = -\frac{1}{g^2} \sum_x \cos \left( A_{x+1} + A_{x+1,2} - A_{x+2,1} - A_{x,2} \right)$$

$$- i\theta Q, \quad (271)$$

where $g^2 = 1/\beta_{pl}$ is the gauge coupling, and $Q$ is the topological charge, a scaling behavior of the partition function zeros in the complex $\theta$-plane is studied with fixed $\text{Re} \theta = \pi$. At the critical mass $\kappa_c$, the position of a partition function zero $\theta_0 (L)$ would obey

$$\text{Im} \theta_0 (L) - \text{Im} \theta_0 (\infty) \propto L^{-(2\nu-\beta)/\nu} \quad (272)$$

with the critical exponents $\nu$ and $\beta$. If a first-order phase transition occurs (conjectured that being at $\kappa < \kappa_c$), it is expected that $\text{Im} \theta_0 (L) \propto L^{-2}$ with vanishing $\text{Im} \theta_0 (\infty)$. On the other hand, if there are not phase transitions (conjectured being at $\kappa > \kappa_c$), it is expected that $\text{Im} \theta_0 (\infty) \neq 0$.

Figure 34 shows the scaling behaviors of $\text{Im} \theta_0 (L)$, and the fitting results are summarized in table I. The bond dimension of tensors are fixed to $D_{\text{cut}} = 160$. These results clearly show that a) for $\kappa < \kappa_c$, $\text{Im} \theta_0 (\infty)$ vanishes, and the exponent $y$ is close to 2, b) for $\kappa > \kappa_c$, $\text{Im} (\infty)$ has a non-zero value, c) for $\kappa = 0.2415 \approx \kappa_c$, $y = 1.869 (10)$ is consistent with $y = 1.875$ that is the same as in the 2D Ising universality class.

Summarizing above, on the line $\theta = \pi$, there are no phase transition at $\kappa > \kappa_c$, there is the second order phase transition belonging to the 2D Ising universality class at $\kappa = \kappa_c \approx 0.2415$, and there are first order phase transitions at $\kappa < \kappa_c$. This is exactly the expected result.

<table>
<thead>
<tr>
<th>$\kappa$</th>
<th>$y$</th>
<th>$\text{Im} \theta_0 (\infty)$</th>
<th>fit range</th>
<th>$\chi^2/\text{DOF}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.2400</td>
<td>2.000(12)</td>
<td>0.000034(59)</td>
<td>$L \in [32\sqrt{2}, 256]$</td>
<td>0.65</td>
</tr>
<tr>
<td>0.2415</td>
<td>1.869(10)</td>
<td>$-0.000016(64)$</td>
<td>$L \in [32\sqrt{2}, 256]$</td>
<td>0.41</td>
</tr>
<tr>
<td>0.2430</td>
<td>1.850(15)</td>
<td>0.00442(12)</td>
<td>$L \in [32\sqrt{2}, 256]$</td>
<td>0.78</td>
</tr>
</tbody>
</table>

There is a further study of the Berezinskii–Kosterlitz–Thouless transition in the same model by the same authors (Shimizu and Kuramashi, 2018), but here we just draw the readers attentions to the paper.

Figure 34 Adapted from (Shimizu and Kuramashi, 2014a). Lee–Yang zeros for $\kappa = 0.24$ (top), 0.2415 (middle), and 0.243, respectively. $1/g^2 = 10$. Solid lines are the fit results with a function $\text{Im} \theta_0 (L) = \text{Im} \theta_0 (\infty) + a L^{-y}$ via three parameters $\text{Im} \theta_0 (\infty)$, $a$, and $y$. 

<table>
<thead>
<tr>
<th>$\kappa$</th>
<th>$y$</th>
<th>$\text{Im} \theta_0 (\infty)$</th>
<th>fit range</th>
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<tr>
<td>0.2415</td>
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<td>$L \in [32\sqrt{2}, 256]$</td>
<td>0.41</td>
</tr>
<tr>
<td>0.2430</td>
<td>1.850(15)</td>
<td>0.00442(12)</td>
<td>$L \in [32\sqrt{2}, 256]$</td>
<td>0.78</td>
</tr>
</tbody>
</table>
D. 2D Thirring model

In (Takeda and Yoshimura, 2015), the two dimensional Thirring model is studied with finite density. This is a first application of the Grassmann TRG to a system with finite chemical potential $\mu$. Figure 35 shows the fermion number density

$$n = \frac{1}{V} \frac{\partial \ln Z}{\partial \mu}.$$  (273)

Even though the numerical differentiation would cause a loss of significance, a Silver Blaze phenomenon is clearly observed, and, in the large $\mu$-region, the Grassmann TRG results reach the saturation density 1 regardless of parameters. This result supports the efficiency of the Grassmann TRG at the finite density region.

E. 3D free fermions

For three dimensions or higher, the Grassmann parts of tensors can be coarse-grained in a similar way to the higher-order TRG (Sakai et al., 2017), and expectation values of physical quantities can be calculated by using impurity tensor techniques (Yoshimura et al., 2018). Figures 36-38 show the free energy density, the chiral condensate $\langle \bar{\psi} \psi \rangle$, and the correlator $C(x_1, x_2) = \langle \bar{\psi}^{(1)}(x_1) \psi^{(2)}(x_2) \rangle$, respectively.

In Fig. 36, the convergence in the number of states, $D_{\text{cut}}$ is extremely rapid, and one cannot see the difference between the different Grassmann HOTRG results and the exact values in this resolution. Additionally, no mass-dependence is observed. By contrast, in Fig. 37, there is clearly a mass-dependence in the accuracy of the Grassmann HOTRG. Converging to the exact result at smaller mass values appears difficult. This could be due to the choice of unitary matrices used in coarse-graining steps. The authors state that the shorter correlation length of the system prevents them from accurately measuring the correlations at large separation.

F. 2D $\mathcal{N} = 1$ Wess–Zumino model

The interacting two dimensional $\mathcal{N} = 1$ Wess–Zumino model displays a vanishing partition function (Witten,
Numerical treatment of Majorana fermions on discrete space-

time lattice is given, and an integer partition function for the free \( N = 1 \) Wess–Zumino model is numerically shown.\(^9\)

The Euclidean continuum action is given by

\[
S_{\text{cont.}} = \int d^2 x \left\{ \frac{1}{2} (\partial_\mu \phi)^2 + \frac{1}{2} W' (\phi)^2 \right. \\
+ \left. \frac{1}{2} \bar{\psi} (\gamma_\mu \partial_\mu + W'' (\phi)) \psi \right\},
\]

(274)

where \( \phi \) and \( \psi \) are a one-component real scalar field and a two-component Majorana spinor field, respectively.\(^11\)

For a review on lattice supersymmetry see (Catterall et al., 2009). \( W'(\phi) \) is an arbitrary function of \( \phi \) and called the superpotential, which is the source of the Yukawa- and \( \phi^2 \)-interactions.

The spinor field \( \psi \) satisfies the Majorana condition

\[
\bar{\psi} = -\psi^T C^{-1}
\]

(275)

with the charge conjugation matrix \( C \):

\[
C^T = -C, \quad C^\dagger = C^{-1}, \quad C^{-1} \gamma_\mu C = -\gamma_\mu^T.
\]

(276)

The continuum action (274) is invariant under the supersymmetry transformation

\[
\delta \phi = \bar{\epsilon} \psi, \quad \delta \psi = (\gamma_\mu \partial_\mu - W'' (\phi)) \epsilon,
\]

(277)

(278)

where \( \epsilon \) is a two-component Grassmann number and \( \bar{\epsilon} \) is defined as in Eq. (275).

Using the symmetric difference operator \( \partial^S_\mu = (\partial_\mu + \partial^*_\mu)/2 \) with the forward difference \( \partial \) and the backward difference \( \partial^* \), the lattice action is given by

\[
S = \sum_x \left\{ \frac{1}{2} (\partial^S_\mu \phi_x)^2 + \frac{1}{2} \left( W' (\phi_x) - \frac{r}{2} \partial_\mu \partial^*_\mu \phi_x \right)^2 \\
+ \frac{1}{2} \bar{\psi}_x (D \psi)_x \right\}.
\]

(279)

It is perturbatively proven that the broken supersymmetry on the lattice is restored in the continuum limit for this construction of the action (Golterman and Petcher, 1989). The Dirac operator on the lattice is defined by

\[
D_{xx'} = \left( \gamma_\mu \partial^S_\mu - \frac{r}{2} \partial_\mu \partial^*_\mu \right)_{xx'} + W'' (\phi_x) \delta_{xx'},
\]

(280)

where \( r \) is a nonzero real parameter that is called the Wilson parameter.

The partition function is given by

\[
Z = \int \mathcal{D} \phi \mathcal{D} \psi e^{-S}
\]

(281)

with the path integral measures \( \int \mathcal{D} \phi = \prod_x \int_{-\infty}^{\infty} d\phi_x/\sqrt{2\pi}, \int \mathcal{D} \psi = \prod_x \int d\psi_x^{(1)} d\psi_x^{(2)} \).

First we construct a tensor network representation of the fermion part in Eq. (281)

\[
Z_F = \int \mathcal{D} \psi e^{-\frac{1}{2} \sum_x \bar{\psi}_x (D \psi)_x}.
\]

(282)

Even in the Majorana case, the basic construction procedure for the Dirac case works as discussed in Sec. XI.A. and the resulting tensor network representation is given by

\[
Z_F = \sum_{\{X_i, T_i\}} \prod_x T_F (\phi_x)_{X_{x-1}X_xT_xT_{x-2}} \mathcal{G}_{x,x_{x-1}X_xT_xT_{x-2}},
\]

(283)

where \( \sum_{\{X_i, T_i\}} \) denotes \( \prod_x \sum_{X_{x-1},X_x,T_x,T_{x-2}=0}^1 \). Here the Grassmann part of the tensor is defined by

\[
\mathcal{G}_{x,ijkl} = \delta_{i\xi_x}^a \delta_{j\xi_{x+1}}^b \delta_{k\xi_x}^c \delta_{l\xi_{x+2}}^d \delta_{i\xi_x}^a \delta_{j\xi_{x+1}}^b \delta_{k\xi_x}^c \delta_{l\xi_{x+2}}^d \cdot (\bar{\xi}_{x+1} \xi_x)^{11} (\bar{\xi}_{x+2} \xi_x)^{12} (\bar{\xi}_{x+2} \xi_x)^{21} (\bar{\xi}_{x+1} \xi_x)^{22}.
\]

(284)

\(^9\) The partition function with the periodic boundary conditions is equivalent to the trace of the fermion number operator \((-1)^F\).

\(^10\) The free case also posses the sign problem although the partition function does not vanish.

\(^11\) Numerical treatment of Majorana fermions on discrete space-time lattice is discussed in e.g. (Wolff, 2008).

\(^12\) The normalization constant is necessary to reproduce the integer partition function.
The definition of $T_F$ is
\[
T_F(\phi)_{kijl} = \int d\Psi d\bar{\Phi} e^{-W''(\phi) + 2r} \Psi \Phi
\]

\[
\cdot \bar{\Psi}^{l_1} \bar{\Psi}^{j_1} \bar{\Phi}^{k_1} \bar{\Phi}^{j_2} \Psi^{i_1} \Psi^{j_1} \Phi^{i_2} \Phi^{j_2} \cdot (\sqrt{\frac{1 + r}{2}})^{i_1+j_1+k_1+l_1} \cdot (\sqrt{\frac{1 - r}{2}})^{i_2+j_2+k_2+l_2}
\]

(285)

with single-component Grassmann variables $\Psi, \bar{\Phi}, \bar{\Psi} = (\Phi + \Psi)/\sqrt{2}, \Phi = (\Phi - \Psi)/\sqrt{2}$. Note that the elements of the tensor depend on the scalar field. This is due to the presence of the Yukawa interaction.

Next, a tensor network representation of the boson part
\[
Z_B = \int D\phi e^{-S_B},
\]

(286)

where
\[
S_B = \sum_x \left\{ \frac{1}{2} (\partial_\mu \phi_x)^2 + \frac{1}{2} \left( W'(\phi_x) - \frac{r}{2} \partial_\mu \partial_\nu \phi_x \right)^2 \right\},
\]

(287)

is considered. A main difficulty here is that $S_B$ has next-nearest-neighbor interaction terms due to the square of the Wilson term.

A simple way to make a nearest-neighbor form is to introduce auxiliary fields. By using two auxiliary fields $G$ and $H$, one can obtain a nearest-neighbor action
\[
\tilde{S}_B = \frac{1}{2} \sum_x \left\{ (\partial_\mu \phi_x)^2 + (W'(\phi_x))^2 + G_x^2 + H_x^2 \right. \\
- (r W'(\phi_x) + \alpha G_x + \beta H_x) (\phi_{x+1} + \phi_{x-1} - 2\phi_x) \\
- (r W'(\phi_x) + \alpha G_x - \beta H_x) (\phi_{x+2} + \phi_{x-2} - 2\phi_x) \right\}
\]

(288)

with $\alpha = \sqrt{(1 - 2r^2)/2}$ and $\beta = 1/\sqrt{2}$. Note that $\alpha$ is real for $|r| \leq 1/\sqrt{2}$ and pure imaginary for $|r| > 1/\sqrt{2}$. Using the nearest-neighbor form of the action, the partition function is written by
\[
Z_B = \int D\phi DGDHE^{-\tilde{S}_B},
\]

(289)

where the measures of $G_x$ and $H_x$ are given in the same way as that of $\phi_x$. Now the scalar part has become a model of a three-component scalar field with nearest-neighbor interactions\textsuperscript{13}, and a tensor network representation for this type of model can be constructed in the same way as the complex $\phi^4$ theory discussed in Sec. X.C.

Using ingredients given above, the total partition function is written as
\[
Z = \int D\phi DGDH e^{-\tilde{S}_B} Z_F,
\]

(290)

where the tensor network representation of the fermion part $Z_F$ is a functional of $\phi$. Even in the presence of $Z_F$, the Gaussian quadrature rule can be used for the discretization of the scalar fields, and a tensor network representation of Eq. (290) can be derived straightforwardly. Note that we do not specify the details of the superpotential, and the above construction can be applied to the $\mathcal{N} = 1$ Wess–Zumino model with any choice of the superpotential.

Here we show the (Grassmann) TRG result (Kadoh et al., 2018) for the partition function of the free Wess–Zumino model, whose superpotential is defined by
\[
W(\phi) = \frac{1}{2} m \phi^2
\]

(291)

with the mass parameter $m$. In order not to break the supersymmetry, periodic boundary conditions are assumed in all directions for both fermions and bosons. The Wilson parameter $r$ is set to $1/\sqrt{2}$ to have $G$ decoupled (see footnote 13). The order of the Hermite polynomial used in Gaussian quadrature rule is set to 64. For a reasoning behind this choice, see the original paper (Kadoh et al., 2018).

In Fig. 39, the partition function (called the Witten index in this model) of the free $\mathcal{N} = 1$ Wess–Zumino model is shown for several lattice volumes: $V = 2 \times 2, 8 \times 8, \text{and} 32 \times 32$. In the free case, the partition function can be analytically obtained, and the exact solution is $Z = \text{sign} \{m (m + 4r)\}$. Thus the exact solution is 1 for the $m > 0$ region shown in the figure. The TRG results tend to converge to the exact value 1 with increasing $D_{\text{cut}}$, the number of singular values that are picked up in coarse-graining steps. The significantly less accurate results in the small $m$ region is due to the lack of the fast damping factor in the local Boltzmann factor that is required for the Gaussian quadrature rule to be effective, but such a bad behavior is a special case for the non-interacting model. When one deals with the interacting Wess–Zumino model that has $\phi^n$-interaction terms, they guarantee the presence of fast damping.

In this subsection a tensor network representation of the $\mathcal{N} = 1$ Wess–Zumino model is discussed, and the numerical result in the non-interacting case (Kadoh et al., 2018) is shown. The integer partition function in the model is reproduced by using the TRG method. Even though the given result is for the non-interacting case, the construction procedure for the tensor network representation is immediately applicable to any interacting case.

The interacting $\mathcal{N} = 1$ Wess–Zumino model would be the next target. The formulation can be immediately ap-

\textsuperscript{13} Actually, by setting the Wilson parameter to be $r = \pm 1/\sqrt{2}$, $G$ is decoupled from the other fields. This choice of the parameter is useful to reduce the computational complexity for tensor calculations and adopted in this section.
Figure 39 Adapted from (Kadoh et al., 2018). The partition function as a function of \( m \) on \( V = 2 \times 2 \) (top), \( 8 \times 8 \) (center), \( 32 \times 32 \) (bottom) lattices.

G. 2D Schwinger model with staggered fermion

While the Wilson fermion case needs special treatment for the Grassmann variables, the staggered one has an excellent feature; its tensor network representation does not have Grassmann variables. i.e. one can represent this staggered fermion system as a purely bosonic tensor network. In this subsection a tensor network representation of the Schwinger model with staggered fermions is discussed.

The one-flavor staggered action for the massless Schwinger model on a two-dimensional lattice has the action,

\[
S = S_F + S_g
\]

with

\[
S_F = \frac{1}{2} \sum_{x=1}^{N} \sum_{\mu=1}^{2} \eta_{x,\mu} \left[ \bar{\psi}_x U_{x,\mu} \psi_{x+\hat{\mu}} - \bar{\psi}_{x+\hat{\mu}} U_{x,\mu}^{\dagger} \psi_x \right]
\]

and

\[
S_g = -\beta \sum_{x=1}^{N} \text{Re} \left[ U_{x,12} \right]
\]

is the usual Wilson action. Here \( \eta_{x,\mu} \) is the staggered phase which for \( \eta_{x,1} = 1 \) and \( \eta_{x,2} = (-1)^{x_1} \) with \( x_1 \) the 1-component of \( x \). The partition function for this model is then given by

\[
Z = \int D\bar{\psi} D\psi e^{-S} = \int D\bar{\psi} D\psi e^{-\beta \sum_{x=1}^{N} \text{Re} \left[ U_{x,12} \right]} Z_F(U)
\]

with

\[
\int D\bar{\psi} D\psi = \prod_x \int_{-\pi}^{\pi} dA_{x,\mu} / 2\pi, \quad \int D\bar{\psi} D\psi = \prod_x \int d\bar{\psi}_x d\psi_x, \quad \text{and} \quad Z_F \text{ represents the part of the partition function that depends on the fermion fields.}
Following (Gattringer et al., 2015b) to formulate the model in terms of discrete degrees of freedom, we first integrate out the fermions and generate an effective action depending only on the gauge fields. As a first step we redefine the link variables such that the staggered fermion phases $\eta_{x,\mu}$ can be absorbed into modified link variables $\hat{U}_{x,\mu} \rightarrow \eta_{x,\mu} U_{x,\mu}$. Under this transformation the gauge action picks up an overall negative sign but the measure is invariant. The Boltzmann factors associated with each bilinear fermion term can be Taylor expanded yielding a partition function

$$Z_F = \int D\tilde{\psi} D\psi \times \prod_x \prod_{\mu} \sum_{k_{x,\mu} = 0}^{1} \left( -\frac{1}{2} \bar{\psi}_x U_{x,\mu} \psi_{x+\hat{\mu}} \right)^{k_{x,\mu}} \times \sum_{\tilde{k}_{x,\mu} = 0}^{1} \left( \frac{1}{2} \bar{\psi}_{x+\hat{\mu}} U_{x,\mu}^\dagger \psi_x \right)^{\tilde{k}_{x,\mu}}.$$  \hspace{1cm} (296)

Notice that higher order terms in the expansion of the Boltzmann factors vanish because of the Grassmann nature of the fermions. There are several ways to generate a non-zero contribution to $Z_F$. In each case, the Grassmann integration at each site must be saturated. To saturate the Grassmann integrations, exactly one forward and one backward hopping term must be associated with each site. On the one hand, there may be a single forward and backward hop along the same link. This saturates the integration, and is referred to as a dimer. On the other hand, there may be a forward and backward hop on two different links at a site. This indicates the passage of fermionic current through the site, and again saturates the integration measure there. Furthermore because of gauge invariance any non-dimer contribution to $Z_F$ must correspond to a closed loop.

For a loop $\ell$ with length $L(\ell)$ one finds a contribution with absolute value

$$\left( \frac{1}{2} \right)^{L(\ell)} \prod_{x,\mu \in \ell} (U_{x,\mu})^{k_{x,\mu}} (U_{x,\mu}^\dagger)^{\tilde{k}_{x,\mu}}.$$ \hspace{1cm} (297)

where on a given link only a single $k$ or $\tilde{k}$ is nonzero. In addition each loop carries a certain $Z_2$ phase which depends on the length of the loop and its winding along the temporal direction given by

$$-(-1)^{\frac{1}{2}L(\ell)}(-1)^W(\ell).$$ \hspace{1cm} (298)

Here, the overall negative sign is the usual one for closed fermion loops while the second factor keeps track of the number of forward hops which is exactly half the total length of the loop for a closed loop. Finally the factor $(-1)^W(\ell)$ of the loop will be determined by the number of windings of the loop along the temporal direction assuming anti-periodic boundary conditions for the fermions.

Using dimers and loops as basic constituents for non-zero contributions to the fermionic partition function we can write

$$Z_F = \left( \frac{1}{2} \right)^V \sum_{\{\ell,d\}} (-1)^{N_\ell + \frac{1}{2} \sum_{\ell} L(\ell) + \sum_{\ell} W(\ell)} \times \prod_{\ell} \left[ \prod_{x,\mu \in \ell} (U_{x,\mu})^{k_{x,\mu}} (U_{x,\mu}^\dagger)^{\tilde{k}_{x,\mu}} \right],$$ \hspace{1cm} (299)

where $\sum_{\{\ell,d\}}$ indicates a sum over all valid loop and dimer configurations, and $N_\ell$ is the number of loops. We construct a local tensor which reproduces the nonzero configurations of this partition function.

Let us ignore the overall sign for now and just deal with the magnitude. We allow two types of indices per link to capture separately the incoming and outgoing fermion lines making the fermion site tensor a rank eight object. To write down a tensor, first, let us fix the coordinates so that right (1-direction) and up (2-direction) are positive. We construct a local tensor which reproduces the nonzero contributions to the fermionic partition function we can write

$$T_{k_{-1,1}k_{-1,1}k_{1,1}k_{x,2}k_{x,2}k_{x-2,2}k_{x-2,2}} = \begin{cases} 1 & \text{if } k_{x-1,1} + k_{x-2,2} + \tilde{k}_{x,1} + \tilde{k}_{x,2} = 1 \\ & \text{and } k_{x,1} + k_{x,2} + \tilde{k}_{x-1,1} + \tilde{k}_{x-2,2} = 1 \\ 0 & \text{otherwise} \end{cases}.$$ \hspace{1cm} (300)

where each $(k_i, \tilde{k}_i) = 0, 1$. The pairs of indices are ordered (left, right, up, down). A graphical representation of this tensor is shown in Fig. 40 (a).

By repeatedly contracting this site tensor with copies of itself over the lattice it can be seen that we generate the full set of closed loops and dimers for the model at zero gauge coupling excluding the overall factor of minus one for each closed fermion loop. The absolute value of the partition function at zero gauge coupling is then,

$$Z_F^{\beta=\infty} = \sum_{\{k,\tilde{k}\}} \prod_{x} T_{k_{x-1,1}k_{x-1,1}k_{x,1}k_{x,2}k_{x-2,2}k_{x-2,2}}.$$ \hspace{1cm} (301)

Here, $\sum_{\{k,\tilde{k}\}}$ is an unrestricted sum over zero and one for all $k, \tilde{k}$ values on the entire lattice, since the $T$ tensor now destroys invalid configurations.

The fermion partition function does not include any contribution or interaction with the gauge fields. To proceed further we will employ a character expansion of the
Boltzmann factors associated with the gauge action.

\[ e^{-\beta \cos [A_{x+1,2} - A_{x+2,1} - A_{x,2}]} \]

\[ \sum_{m_{x,12} = -\infty}^{m_{x,12} = \infty} I_{m_{x,12}} (-\beta) e^{im_{x,12}[A_{x,1} + A_{x+1,2} - A_{x+2,1} - A_{x,2}]} \]

Each plaquette is now labeled by an integer \( m_{x,12} \) (which we shorten to \( m_x \) since there are only temporal plaquettes in two dimensions). Note that \( I_{m_x} (-\beta) = (-1)^{m_x} I_{m_x} (\beta) \). In two dimensions each link is shared by two plaquettes. For a link in the \( \mu = 1 \) direction the two plaquettes give factors of \( e^{im_x A_{x,1}} \) and \( e^{-im_{x-2} A_{x,1}} \). In the \( \mu = 2 \) direction, \( e^{-im_x A_{x,2}} \) and \( e^{im_{x-1} A_{x,2}} \). In addition, the link carries a factor of \( e^{-ik_x \mu A_{x,\mu}} \) or \( e^{-i\bar{k}_x \mu A_{x,\mu}} \) coming from \( Z_F \). Thus, in total, links carry two \( m \) indices inherited from their neighboring plaquettes together with a \( k \) and a \( \bar{k} \) index associated with the fermionic hopping terms. The integral over a link variable is given by

\[ \int_{-\pi}^{\pi} \frac{dA_{x,\mu}}{2\pi} e^{i(k_{x,\mu} - \bar{k}_{x,\mu}) A_{x,\mu}} \prod_{\nu>\mu} e^{i(m_{x,\nu} - m_{x,\nu}) A_{x,\mu}} \times \prod_{\nu<\mu} e^{i(m_{x,\nu} - m_{x,\nu}) A_{x,\mu}} = \prod_{x,\mu} \delta^{(x,\mu)}(m_{x,\nu} - m_{x,\nu}) + \sum_{\nu<\mu} (m_{x,\nu} - m_{x,\nu}) + k_{x,\mu} - \bar{k}_{x,\mu}, 0^+ \]

This allows us to write the partition function as a sum over \( m \) and \( k, \bar{k} \) variables,

\[ Z = \sum_{\{m\}} \sum_{\{k, \bar{k}\}} \left( \prod_{x} I_{m_x} (\beta) \right) \times \left( \prod_{x} T_{x,k_{x,\mu}, \bar{k}_{x,\mu}} \right) \times (-1)^{N_F + N_P + \frac{1}{2} \sum_{\ell} L(\ell) + \sum_{\ell} W(\ell)} \prod_{x,\mu} \delta^{(x,\mu)}(m_{x,\nu} - m_{x,\nu}) + \sum_{\nu<\mu} (m_{x,\nu} - m_{x,\nu}) + k_{x,\mu} - \bar{k}_{x,\mu}, 0^+ \]

where \( N_F = \sum m_x \). At this point we have included all minus signs for completeness. It was proven in (Gattringer et al., 2015b) that every valid contribution to the partition function is positive in the case of periodic boundary conditions, and so from here on we ignore the factor of \( (-1)^{N_L + N_P + \frac{1}{2} \sum_{\ell} L(\ell) + \sum_{\ell} W(\ell)} \).

Now, associated with each link is a constraint between the \( k \) and \( \bar{k} \) fields on the link and the adjacent \( m \) fields on the plaquettes given by Eq. (303). This is a natural object to use to form a tensor. We define a tensor on each link by,

\[ A^{(x,\mu)}_{m_x m_{x-1}} k_{x,\mu} k_{x+1,\mu} k_{x+2,\mu} \]

\[ \delta^{(x,\mu)}_{m_{x-1}}(m_{x,\nu} - m_{x,\nu}) + \sum_{\nu<\mu} (m_{x,\nu} - m_{x,\nu}) + k_{x,\mu} - \bar{k}_{x,\mu}, 0^+ \]

Here the \( k^i, \bar{k}^i \) indices are associated with the two ends of a link. These indices are diagonal as indicated by the Kronecker deltas. A diagram showing the relative position of the fermion and plaquette indices is shown in Fig. 40 (b).

Finally, we construct a tensor associated with the plaquettes of the lattice. This is the same tensor used before in previous tensor formulations of Abelian gauge theories. Since there is only a single \( m \) associated with each plaquette, a tensor definition must only depend on that single \( m \). A plaquette tensor, \( B \), can be defined as,

\[ B_{m_1 m_2 m_3 m_4} = \begin{cases} I_{m_x} (\beta) & \text{if } m_1 = m_2 = m_3 = m_4 \\ m_x & \text{otherwise.} \end{cases} \]
A graphical representation for the $B$ tensor is shown in Fig. 40 (c). The contraction over these three ($T$, $A$, and $B$) unique tensor types can be represented as the tensor network shown in Fig. 41. Since the fermionic $k$ indices always come in $k^i, \bar{k}^i$ pairs, we can form a product state of those two indices to reduce the complexity of the notation. Define $K_{x, \mu}^{a} = k_x^i \otimes \bar{k}_x^i$, then,

$$T(x) \rightarrow T'(x)$$

$$= T(x)_{(k_{x-1,1} \otimes k_{x-1,1}) (k_{x,1} \otimes k_{x,1}) (k_{x,2} \otimes k_{x,2}) (k_{x-2,2} \otimes k_{x-2,2})}$$

$$= T(K_{x-1,1} K_{x,1} K_{x,2} K_{x-2,2})$$

and

$$A(x, \mu) \rightarrow A'(x, \mu)$$

$$= A_{m_2 m_{x-\bar{\theta}} K_{x, \mu}}^{(x, \mu)}(k_{x, \mu} \otimes k_{x, \mu}^2)$$

(307)

(308)

The new enlarged $K$ indices take values from 0 to 3, enumerating the four possible states each link can have: un-occupied, incoming, outgoing, and dimer. The $A$ tensors are still diagonal in the new $K$ indices.

Figure 41 Contraction pattern of basic tensors.

It is possible to include a topological term in the original action with the addition of,

$$S_\Theta = \frac{i \Theta}{2\pi} \sum_x \text{Im}[U_{x,12}].$$

(309)

Taking the staggered phase into account, and expanding the Wilson plaquette term and this term simultaneously,

$$e^{-\beta \text{Re}[U_{x,12}]} + \frac{\Theta}{2} \sum_x \text{Im}[U_{x,12}] =$$

$$\sum_{m_x = -\infty}^{\infty} C_{m_x}(\beta, \Theta) e^{i m_x (A_{x,1} + A_{x+1,2} - A_{x+2,1} - A_{x,2})},$$

(310)

(311)

Figure 42 The average plaquette as a function of the $\Theta$ parameter on a $4 \times 4$ lattice. Here $N_{\text{gauge}} = 5$ indicates a truncation on the $m$ numbers such that $m$ runs from $-2$ to $2$. That is to say that $D_{\text{cut}} = 5$ on the $B$ tensor initially.

The previous steps in formulating a tensor network can be followed straightforwardly. One can solve for the Cs numerically or analytically (Gattringer et al., 2015b). It amounts to the replacement,

$$I_{m_x}(\beta) \rightarrow I_{m_x}(2\sqrt{\eta}) \left( \frac{\eta}{\eta + 2 - \Theta/4\pi} \right)^{m_x/2}$$

(312)

in the definition of the $B$ tensor, with $\eta = \beta/2 - \Theta/4\pi$ and $\bar{\eta} = \beta/2 + \Theta/4\pi$.

Using these tensors, one can perform numerical calculations using a coarse graining scheme. In (Butt et al., 2019) the authors used the higher-order tensor renormalization group to calculate the free energy for the massless Schwinger model with and without the presence of a topological term. From the free energy they calculated the average plaquette, and the topological charge both as a function of the gauge coupling and the $\Theta$ parameter. The average plaquette and topological charge are given by,

$$\langle U_p \rangle = \frac{1}{V} \frac{\partial \ln Z}{\partial \beta}$$

(313)

and

$$\langle Q \rangle = \frac{1}{V} \frac{\partial \ln Z}{\partial \Theta},$$

(314)

respectively. These were compared with Monte Carlo calculations from (Gosch et al., 2017) when possible. A figure from (Butt et al., 2019) showing the average plaquette as a function of the the $\Theta$ parameter is shown in Fig. 42 for a $4 \times 4$ lattice. In Fig. 43 we see a comparison between the tensor calculation and Monte Carlo for a fixed volume on a $4 \times 4$ lattice of the topological charge. The authors reported difficulty at larger volumes, perhaps owing to how the coarse-graining scheme handles
which states are kept before knowing the boundary conditions on the lattice.

Investigations of the Schwinger model with staggered fermions in the Hamiltonian limit have also been conducted, with and without a topological term. In (Baâuls et al., 2013) the mass spectrum of the model is studied using the density matrix renormalization group with matrix product states. In (Baâuls et al., 2017a) the phase diagram of the Schwinger model with two flavors of fermion in the presence of a chemical potential was studied. The authors investigate the isospin as a function of the chemical potential, and map the phase diagram in the chemical potential-mass plane. A more recent study of the Schwinger model with the inclusion of a Θ term was done in (Funcke et al., 2020) again using matrix product states. The authors looked at different thermodynamic quantities as a function of the Θ parameter, as well as the spectrum of the model. They also considered the continuum and chiral limits of the model where the Θ parameter becomes irrelevant.

H. Open questions/Possible directions

One of the big goals in lattice gauge theory is the successful simulation of four dimensional QCD at finite density. However, the computational time of the Grassmann HOTRG in four dimensions is extremely demanding: $\propto D_{cut}^{15}$ with the bond dimension $D_{cut}$. To achieve the goal, further improvements of the algorithm would be required, such as Monte Carlo approximation/sampling of the tensors, and effective truncation of bonds.

From an application point of view, non-trivial models in three dimensions would be within the range. The three dimensional Thirring model that has a non-trivial phase structure and $2+1$ dimensional domain-wall fermion systems would be interesting targets.

As described in Sec. XI.E, the choice of the unitary matrices used in (Yoshimura et al., 2018) during coarse-graining is not optimal. This leaves open the possibility to improve their results by tuning the unitary matrices.

In fermion systems, the spectra of tensors tend to have milder hierarchies than purely bosonic ones. Indeed, in the works on the Schwinger model (Shimizu and Kuras-mashi, 2014a,b, 2018), the bond dimension of tensors is taken to be 160. This is very large, so that one cannot easily reproduce their results on standard, say desktop or laptop, computers. Then, even in two dimensions, serious calculations require improvements of the algorithms. In two dimensions, there are several improved schemes for bosonic tensor networks such as (loop-)TNR (Evenbly and Vidal, 2015; Yang et al., 2017), graph-independent local truncations (Hauru et al., 2018), and full environment truncation (Evenbly, 2018). Then Grassmann versions of them would all be possible directions.

XII. TRANSFER MATRIX AND HAMILTONIAN

We now move on from the topic of reformulating the partition function in terms of tensors, to arranging their contractions so as to deduce a transfer matrix which can be used to rewrite the partition function as in Eq. (32). Once the partition function has been written entirely in terms of local tensor contractions, it is possible to organize these index contractions into time layers. The natural choice is to use the indices attached to time links and/or space-time plaquettes to be the indices of the transfer matrices. Geometrically, the Hilbert space is located in between two time slices while the transfer matrix is centered on a time slice and connects two copies of the Hilbert space. To the best of our knowledge, interchanging the role of these two types of layers is only possible by going back to configuration space.

In the rest of this section, we target models with continuous Abelian symmetries (the O(2) spin model and U(1) gauge theory) and describe their transfer matrices from a tensor perspective. However, it is not difficult to extend the discussion to other models.

A. Spin models

For spin models (Zou et al., 2014), the transfer matrix can be constructed by taking all the tensors on a time slice and tracing over the spatial indices. This is illustrated for $D = 2$ and $D = 3$ in Fig. 44.

For the O(2) model, the Hilbert space $\mathcal{H}$ is the product of integer indices attached to time links between two time
slices.

\[ \mathcal{S} = |\{n\} \rangle = \bigotimes_{x,j} |n_{x,j}\rangle. \]  

(315)

For \( D = 2 \) with \( N_s \) sites and periodic boundary conditions, the matrix elements of the transfer matrix \( T \) have the explicit form

\[ \langle \{n'\} | T | \{n\} \rangle = \sum_{\vec{n}_1 \ldots \vec{n}_{N_s}} T^{(1,\tau)}_{\vec{n}, \vec{n}_1 \ldots \vec{n}_1, \vec{n}_2} \cdot \ldots \cdot T^{(N_s,\tau)}_{\vec{n}, \vec{n}_1 \ldots \vec{n}_N, \vec{n}_N', \vec{n}_N} \]  

(316)

with the individual tensors provided in Sec. VII. The transfer matrix can be coarse-grained in the spatial dimension (Yang et al., 2016; Zou et al., 2014) as illustrated in Fig. 45. This method was used to perform numerical calculations in Refs. (Unmuth-Yockey et al., 2018; Yang et al., 2016; Zhang et al., 2018; Zou et al., 2014).

![Graphical representation of the coarse-graining truncation of the transfer matrix](image)

Figure 45 Graphical representation of the coarse-graining truncation of the transfer matrix described in the text.

The symmetries of the model are completely encoded in Kronecker deltas appearing in the definition of the tensor (Meurice, 2019). This corresponds to a divergenceless condition and with either periodic or open boundary conditions, the charges carried by the indices cannot flow out in the spatial directions. For the \( O(2) \) model, the sum of the time indices going in the time slice equals the sum of the indices going out. This conserved quantity can be identified as the charge of the initial and final states and the transfer matrix commutes with the charge operator which counts the sum of the in or out indices. As we will explain in Sec. XIV.A, setting some matrix elements to zero if some of the local indices exceed some value, \( n_{\text{max}} \), in absolute value will not affect this property.

The transfer matrix can be used to define a Hamiltonian by taking an anisotropic limit where \( \beta \) becomes large on time links and small on space links (Fradkin and Susskind, 1978; Kogut, 1979). We define \( V = 1/(\beta \tau a_\tau) \), \( \mu = \mu/a_\tau \) and \( J = \beta s/a_\tau \). The Hamiltonian is defined by

\[ \mathcal{T} = 1 - a_\tau \hat{\mathcal{H}} + \mathcal{O}(a_\tau^2). \]  

(317)

It will inherit the symmetry properties of the transfer matrix. It explicit form is

\[ \hat{\mathcal{H}} = \sum_x \left[ \frac{\hat{V}}{2} \hat{L}_x^2 - \hat{\mu} \hat{L}_x - \frac{J}{2} \sum_{x,\mu} (\hat{U}_{x+\mu}^\dagger \hat{U}_x^\dagger + \text{h.c.}) \right] \]  

(318)

with the operator \( \hat{L} |n\rangle = n |n\rangle \), and the operator \( \hat{U} = e^{i\varphi} \) which corresponds to the insertion of \( e^{i\varphi} \) in the path integral and raises the charge

\[ \hat{U} |n\rangle = |n + 1\rangle, \]  

(319)

while its Hermitian conjugate lowers it

\[ (\hat{U}^\dagger) |n\rangle = |n - 1\rangle. \]  

(320)

This implies the commutation relations

\[ [\hat{L}, \hat{U}] = \hat{U}, \quad [\hat{L}, \hat{U}^\dagger] = -\hat{U}^\dagger, \]  

(321)

and

\[ [\hat{U}, \hat{U}^\dagger] = 0. \]  

(322)

B. Gauge models

The Hilbert space for the compact Abelian Higgs model, or its pure gauge \( U(1) \) limit, \( \mathcal{S}_G \) can be constructed with the indices associated with space-time plaquettes

\[ \mathcal{S}_G = |\{e\} \rangle = \bigotimes_{x,j} |e_{x,j}\rangle. \]  

(323)

The electric layer is a diagonal matrix \( \mathcal{T}_E \) with matrix elements

\[ \langle \{e'\} | \mathcal{T}_E | \{e\} \rangle = \delta_{\{e\},\{e'\}} T_E(\{e\}). \]  

(324)
where $T_E(\{e\})$ are traced products of $A$ tensors on time links with $B$ tensors on space-time plaquettes

$$T_E(\{e\}) = \text{Tr} \prod_{\text{time } l} A^{(l)}_{m_1, \ldots, m_2(D-1)} \prod_{\text{sp.-time } pl} B^{(pl)}(e).$$

(325)

The $A$ tensor of the compact Abelian Higgs model is given in Eq. (184). It enforces Gauss’s law in the pure gauge limit. The electric layer is illustrated in Fig. 46.

Similarly, we define the magnetic matrix elements $\langle\{e\}|T_M|\{e'\} \rangle$ with the indices $e$ and $e'$ carried by the time legs of the $A$-tensors located on time links

$$\langle\{e'\}|T_M|\{e\} \rangle = \text{Tr} \prod_{\text{space } l} A^{(l)}_{m_1, \ldots, m_2(D-1)}(e, e') \prod_{\text{sp.-sp. } pl} B^{(pl)}.$$  

(326)

The traces are taken over the spatial legs of the tensors, while the time legs are left open and carry the the indices $e$ and $e'$. The magnetic layer is illustrated in Fig. 47.

We define the transfer matrix $\mathbb{T}$ as

$$\mathbb{T} \equiv (e^{-\beta_{\text{pl}} I_0(\beta_{\text{pl}})})^{(V/N_r)D(D-1)/2} (e^{-\beta_l I_0(\beta_l)})^{(V/N_r)D} \times T_E^{1/2} T_M^{1/2},$$

(327)

with $N_r$ the number of sites in the temporal direction.

---

**Figure 46** Electric layer of the transfer matrix for $D = 3$ between two time slices (top) and “from above” (bottom).

**Figure 47** Magnetic layer of the transfer matrix for $D = 3$ in a time slice (top) and “from above” (bottom).

Proceeding as for the spin model (Fradkin and Susskind, 1978; Kogut, 1979), we define

$$\beta_{\tau_{\text{pl}}} = \frac{1}{a_r g^2_{\text{pl}}}, \quad \text{and} \quad \beta_{\tau_l} = \frac{1}{a_r g^2_l},$$

(328)

for the couplings related to the time direction and

$$\beta_{s_{\text{pl}}} = a_r J_{\text{pl}}, \quad \text{and} \quad \beta_{s_{\text{l}}} = a_r h_l,$$

(329)

for the spatial couplings. We then obtain (Meurice, 2020a; Unmuth-Yockey, 2017)

$$\hat{\mathcal{H}} = \frac{1}{2} g^2_{\text{pl}} \sum_{x,j} (\hat{\epsilon}_{x,j})^2$$

$$+ \frac{1}{2} g^2_l \sum_{x} (\sum_{j} (\hat{\epsilon}_{x,j} - \hat{\epsilon}_{x-\hat{j},j}))^2$$

$$- h_l \sum_{x,j} (\hat{U}_{x,j} + h.c.)$$

$$- J_{\text{pl}} \sum_{x,j<k} (\hat{U}_{x,j} \hat{U}^{\dagger}_{x+\hat{j},k} \hat{U}^{\dagger}_{x,k} + h.c.).$$

(330)

Notice that for the compact Abelian Higgs model the matter fields can absorb non-zero values in Eq. (169) (Gauss’s law). However, in the limit where the link couplings are set to zero, we recover the pure gauge $U(1)$ model where the $A$-tensors for time links enforce Gauss’s law.

So far everything we have done has been manifestly gauge-invariant because the tensors resulted from a complete integration over the gauge fields. The partition function remains unchanged if we use a temporal gauge
If we gauge away the gauge fields on a time link instead of integrating over them, we lose the Gauss’s law enforcement associated with that time link. However, the discrete Maxwell equations of Sec. IX.B imply that if Gauss’s law is satisfied on one electric layer, then it is also satisfied on all the other layers. With open boundary conditions in time, Gauss’s law is trivially satisfied on the first and last layers. With periodic boundary conditions, we cannot gauge away the Polyakov loops and we need to keep the integration over the temporal links for one layer. This is sufficient to enforce Gauss’s law in that layer and consequently everywhere.

If we prepare an initial state which satisfies Gauss’s law, the exact time evolution will preserve this property. However, in the noisy intermediate-scale quantum (NISQ) era, various types of errors can introduce Gauss’s law violations. For this reason, it has been argued (Bender and Zohar, 2020; Kaplan and Stryker, 2018; Meurice, 2020a; Unmuth-Yockey, 2019, 2020) that it would be desirable to find a parametrization of the Hilbert space where Gauss’s law is automatically satisfied. One possibility discussed next in Sec. XII.C is to use the unconstrained variables introduced in Sec. IX.C. A simple solution (Meurice, 2020a) for the Hilbert space $\mathcal{H}_G$ introduced in Eq. (323), is to write the $\epsilon_{x,i}$ as the discrete divergence of antisymmetric tensors. For $D = 3$, we only need one field instead of two and we obtain an optimal representation similar to what was proposed in Refs. (Kaplan and Stryker, 2018; Unmuth-Yockey, 2019).

\[
\begin{align*}
\epsilon_{x,1} &= -c_x + c_{x-2} \\
\epsilon_{x,2} &= +c_x - c_{x-1} \\
\epsilon_{x,3} &= = c_{x-3} + c_{x-2} - c_{x-1,2} \\
\end{align*}
\]

For $D = 4$, we can write the electric field as the curl of a three component vector (Meurice, 2020a). As this new vector is defined up to a gradient we can attempt to use this freedom to remove say the first component. This would provide an expression of the form

\[
\begin{align*}
\epsilon_{x,1} &= -c_{x,3} + c_{x-2,3} + c_{x,2} - c_{x-1,3} \\
\epsilon_{x,2} &= +c_{x,3} - c_{x-1,3} \\
\epsilon_{x,3} &= -c_{x,2} + c_{x-1,2}.
\end{align*}
\]

However, the global implementation depends on the boundary conditions (Meurice, 2020a). A more recent discussion of Gauss law for PBC and OBC can be found in Ref. (Bender and Zohar, 2020). In summary, it is possible to enforce Gauss’s law with no unphysical degrees of freedom that would waste computational resources. This can be done in any dimension and is better understood using the dual formulation discussed in the coming section.

C. Duality revisited and Gauss law

The passage to the unconstrained variables discussed in the Lagrangian formalism Sec. IX.C solve Gauss’s law in $D = 3$ and remove any gauge freedom from the model. In the continuous-time limit, when the transfer matrix is close to the identity and one can identify a Hamiltonian, there is no residual gauge freedom, and in fact the model is recast as a spin model.

In $D = 4$, the unconstrained variables which solve the divergence-less constraint in Abelian models are left with a redundancy themselves. That is, there is a local operation which leaves the new Hamiltonian unchanged, and so the question of physical states remains. This arises from,

\[
m_{x,\mu} = \epsilon_{\mu \nu \rho a} \Delta_\rho C_{x^\nu, a}
\]

which introduces a new “gauge field” on the links of the dual lattice. The field strength tensor for this gauge field possess a similar redundancy to that of the original field, i.e. $C_{x,\mu} \to C_{x,\mu}’ = C_{x,\mu} + \Delta_\mu \phi_x$ leaves the quantum Hamiltonian unchanged. In the electric basis (the $L^z$ basis) this symmetry is manifested in an operator which raises and lowers all angular momentum numbers around a site by one, $G_x = \Pi_{\mu=1} U_{x,\mu} U_{x-\mu, \mu}$. This operator commutes with the Hamiltonian. This identifies physical states as those which do not differ from others by arbitrary applications of $G_x$.

D. Algebraic aspects of the Hamiltonian formulation

As practical implementations require a finite number of states, we need to discuss the effect of a truncation on the algebra defined by Eqs. (321) and (322). By truncation we mean that there exists some $n_{\max}$ for which

\[
\hat{U} |n_{\max}\rangle = 0, \text{and } (\hat{U}^\dagger) |−n_{\max}\rangle = 0.
\]

It is clear that these modifications contradict Eq. (322) because if $\hat{U}$ and $\hat{U}^\dagger$ commute, we can apply $\hat{U}^\dagger$ on the first equation (334) and obtain that $\hat{U} |n_{\max} − 1\rangle$ is also zero and so on. If we consider the commutation relations with the restriction (334), we see that the only changes are

\[
\langle n_{\max} | [\hat{U}, \hat{U}^\dagger] | n_{\max} \rangle = 1,
\]

\[
\langle −n_{\max} | [\hat{U}, \hat{U}^\dagger] | −n_{\max} \rangle = −1,
\]

instead of 0. The important point is that the truncation does not affect the basic expression of the symmetry in Eq. (322). It only affects matrix elements involving the $\hat{U}$ operators but not in a way that contradicts charge conservation. For a related discussion of the algebra for the O(3) model see Ref. (Bruckmann et al., 2019).

Other deformations of the original Hamiltonian algebra defined by Eqs. (321) and (322) appear in the quantum link formulation of lattice gauge theories (Brower et al., 1999). In this approach, one picks a representation of the SU(2) algebra and replace $\hat{U}$ by the raising
operator $S^\dagger$. Eq. (334) is then satisfied if the dimension of the representation is $2n_{\max} + 1$ but Eq. (322) becomes

$$[\hat{S}^+, \hat{S}^-] = 2\hat{S}^z. $$ (336)

Finally, we would like to comment about algebraic aspects of the Gaussian quadratures discussed in Sec. X.B. This numerical integration method averages over a finite number of sampling points which are the zeros of a Hermite polynomial of sufficiently large order $n_{\max} + 1$. This can be related to a truncation of the standard harmonic oscillator algebra in the following way. If we use the standard raising and lowering operators on energy eigenstates $|n\rangle$ to calculate $(x| \hat{x} |n\rangle$, we recover the Hermite polynomial recursion formula. These relations still hold for the zeros of $H_{n_{\max}+1}$ until we reach level $n_{\max}$. Iterating one more time provides a relation equivalent to

$$\hat{a}^\dagger |n_{\max}\rangle = 0. $$ (337)

The modified commutation relation become

$$[\hat{a}, \hat{a}^\dagger] = 1 - (n_{\max} + 1) |n_{\max}\rangle \langle n_{\max}|. $$ (338)

A better algebraic understanding of the results of Sec. X.B would certainly be of great interest.

**XIII. QUANTUM COMPUTATIONS AND SIMULATIONS**

The idea of using actual quantum devices to mimic or study theoretical quantum models has been a subject of intense activity in recent years. In particular the use of optical lattices (Bloch et al., 2008) to quantum simulate lattice gauge theories has been developed extensively and reviewed in Refs. (Tagliacozzo et al., 2013; Wiese, 2013; Zohar et al., 2015b). For recent developments combining condensed matter and gauge theories ideas see (Clark et al., 2018; Kasper et al., 2017; Li et al., 2016; Schweizer et al., 2019). Trapped ions (Leibfried et al., 2003) have provided new opportunities to approach lattice gauge theory models (Davoudi et al., 2020a). Rydberg atoms offer a versatile platform to probe many-body dynamics (Bernien et al., 2017) and out-of-equilibrium properties of spin models (Keesling et al., 2019) and gauge theories (Celi et al., 2019). The Schwinger model is often the first target to develop new approaches (Davoudi et al., 2020a; Kasper et al., 2017; Kharzeev and Kikuchi, 2020; Klco et al., 2018; Magnifico et al., 2019; Martinez et al., 2016; Surace et al., 2020). For recent work on non-Abelian models see (Dasgupta and Raychowdhury, 2020; Davoudi et al., 2020b; Raychowdhury and Stryker, 2020a,b; Silvi et al., 2019). In the following, we only cover selected aspects. For a recent review of lattice gauge theories within quantum technologies see (Bañuls et al., 2020).

**A. The Ising model**

As stressed in this review, the Ising model is the quintessential model to use to check computational tools and proposed algorithms. This is because the model possesses non-trivial features, but has been solved exactly. It is not surprising then that this is one of the first models to be tried and tested in different forms of quantum computation. There have been various investigations into how to simulate the model on a quantum computer, and what interesting observables to measure (e.g. phase shift, and thermodynamics) (Gustafson et al., 2019a; Lidar and Biham, 1997; Mostame and Schützhold, 2008; You et al., 2013). In terms of actual experimental (or machines) there are several main approaches: superconducting qubit machines, trapped-ion machines, cold atoms trapped in optical lattices, and Rydberg atom simulators.

On the superconducting qubit front, Refs. (Heiblum et al., 2017; Zhang et al., 2017) made initial simulations, and in some cases at relatively large system sizes. Their calculations of various spin observables matched the corresponding quantities in exact diagonalization well. In (Cervera-Lierta, 2018) a simulation of the Ising model using a few spins was also carried out, and a comparison between theory and computation was made for the average magnetization. In Refs. (Gustafson et al., 2019a; Smith et al., 2019) The Ising model was simulated using a few qubits on IBM’s machines, and in (Gustafson et al., 2019a), with the addition of removing noise through error reduction techniques such as Richardson extrapolation. In addition, in (Chen et al., 2019) a quantum-classical approach was taken which uses a variational algorithm to compute the ground-state wave function; the so-called Variational Quantum Computing, where the quantum computer performs an energy minimization given a trial wave function whose parameters are tuned by a classical computer using the quantum output.

Using trapped ions, there are a couple of extensive investigations from student’s work (Friedenauer, 2010; Korotkov, 2013). These are pioneering investigations into the trapped-ion platform. Refs. (Edwards et al., 2010; Islam et al., 2011; Kim et al., 2010, 2011) look at the phase structure of the model by simulating a few spins. They calculate the phase diagram of the model using the probability of the state to be in a ferromagnetic state, or moments of the magnetization. There has also been an investigation into measuring the Rényi entropy using digital quantum gates. In (Linke et al., 2018) the authors consider a two-site antiferromagnetic Ising model, and using the SWAP gate measure the parity of two copies of the system (Johri et al., 2017). Ref. (Islam et al., 2015) used bosonic many-body states and interfered the copies to extract sub-system parities. They then calculate the Rényi entropy from sub-system parities.
Finally, another promising approach is to use highly excited (Rydberg) states of atoms, which allows for strong atom-atom interactions across relatively large distances. In (Kim et al., 2017) the group uses a chain of 19 Rubidium atoms whose interactions they control through tuning lattice parameters to simulate the model. In (Guardado-Sanchez et al., 2018) out-of-equilibrium dynamics are explored through a quench using an array of Lithium atoms, again placed in a Rydberg state. Reference (Simon et al., 2011) uses trapped Rubidium atoms to simulate the anti-ferromagnetic Ising model. They are able to identify a phase transition between para- and anti-ferromagnetic phases, and observe magnetic domains using a site-resolved atomic microscope and noise correlations measurements. In (Bernien et al., 2017) the authors demonstrate a reconfigurable, programmable array of atoms, and use it to simulate an Ising-like model on 51 qubits. Using this array and atoms they observe a phase transition between symmetric and ordered phases. For a review of this entire topic see (Schauss, 2018).

B. O(2) and O(3) model

For the O(2) nonlinear sigma model, (Zou et al., 2014) uses a mapping between the O(2) model, and the Bose-Hubbard model. They relate the two phase diagrams in the hopping-chemical potential plane, and give the explicit mapping between variables between the two models. A similar approach can be seen as the limiting behavior of the Abelian Higgs model in Refs. (Bazavov et al., 2015; Zhang et al., 2018) when the gauge coupling is taken to zero. In (Unmuth-Yockey et al., 2017) the authors describe a method to measure the second-order Rényi entropy for the O(2) model with a chemical potential in the limit where it appears as the Bose-Hubbard model. They do this in the case of an ultra-cold atomic species trapped in an optical lattice at half-filling. They also consider the experimental cost to extract the central charge from measurements of the Rényi entropy.

There are a couple of results in progress towards quantum simulation of the O(3) nonlinear sigma model, but none on the principal chiral model at the time of writing. In (Schützhold and Mostame, 2005), a proposal for an analogue quantum simulator for the O(3) nonlinear sigma model in two dimensions is discussed. The set-up involves an idealized circuit of superconducting and insulating spheres and wires. The $\sigma$ field is identified with the position of an electron living on the surface of an insulating sphere. The nearest neighbor potential is discrete in space and is identified with the difference in positions between adjacent electrons. This is mapped to the spatial gradient of the $\sigma$ field. These two identifications are used to match couplings between the circuit model and the original nonlinear sigma model. Possible experimental parameters are discussed as well as an analysis of noise contributions to the simulation.

The reference (Alexandru et al., 2019b) discusses an approach to quantum simulating the O(3) nonlinear sigma model using “digital” quantum computers implementing qubits. The original Hamiltonian is re-expressed in the angular momentum basis. In this basis (and discussed above) a truncation is made which preserves the O(3) symmetry of the model, but reduces the local state space to four states. This is a natural truncation to the $l_{\text{max}} = 1$ state which possesses a singlet and triplet state coming from $l = 0$ and $l = 1$ states, respectively. This is precisely what one finds in the addition of angular momentum between the product of two spin-$\frac{1}{2}$ states. In this truncation and representation the model is cast in terms of two-qubit operators. Finally the authors use a Suzuki-Trotter decomposition to write the Hamiltonian evolution in short, discrete steps. Each step is mapped to a quantum circuit over qubits. The authors simulate the Hamiltonian evolution on a classical computer and discuss results. They also perform runs on a quantum computer; however, at the time, they find “mostly noise.”

C. Quantum simulation of Abelian Higgs model with cold atoms

There are a few concrete proposals on how to simulate the Abelian Higgs model using cold atoms trapped in an optical lattice. These methods either make use of the similarity between multi-species Bose-Hubbard models and the Abelian Higgs model, or create an effective model only in terms of gauge degrees of freedom and construct the local Hilbert space of the model directly as a physical dimension and include operators for the new dimension.

In (Bazavov et al., 2015) a two-species Bose-Hubbard model is proposed to simulate the Abelian Higgs model in the limit of infinite Higgs mass in 1+1 dimensions. The authors use the Fourier expansion for the Abelian fields and rewrite the model in terms of discrete variables. Then, the matter degrees of freedom are integrated out creating an effective theory only in terms of the discrete gauge-field degrees of freedom. Finally, the authors make use the mapping between Schwinger Bosons and $SU(2)$ angular momentum operators to write the model in terms of Bosonic creation/annihilation operators and number operators whose form matches that of a two-species Bose-Hubbard model. The authors compare the energy spectra between the original model and the Bose-Hubbard system and find good agreement.

In (González-Cuadra et al., 2017) another multi-species Bose-Hubbard model is proposed to simulate the Abelian Higgs model in 2+1 dimensions. The authors use a six-species Bose-Hubbard Hamiltonian, and again find a mapping between their electric field and paral-
A symmetry is defined as a field transformation
\[ \phi_\ell \rightarrow \phi'_\ell = \phi_\ell + \delta \phi_\ell \] (341)
such that the action and the integration measure are preserved. These invariances imply that
\[ \langle f(\Phi) \rangle = \langle f(\Phi + \delta \Phi) \rangle. \] (342)

If the action is not exactly invariant, \( \exp(\delta S) \) gets inserted in the expectation value on the right-hand side of the equation.

The O(2) model discussed in Sec. VII.A is invariant under the global shift
\[ \varphi' \equiv \varphi + \alpha. \] (343)

Assuming that the function \( f \) is 2\( \pi \)-periodic in its \( M \) variables, we expand in Fourier modes and after using Eq. (342), we obtain
\[ \langle e^{i(n_1 \varphi_{r_1} + \cdots + n_M \varphi_{r_M})} \rangle = 0. \] (344)

This global selection rule can be explained (Meurice, 2019) in terms of the selection rule of the microscopic tensors at each site given in Eq. (121). It is a divergenceless condition and it can be interpreted as a discrete version of Noether’s theorem. If we enclose a site \( x \) in a small size (compared to the lattice spacing) \( D \)-dimensional cube, the sum of indices corresponding to positive directions \( (n_{r,\text{out}}) \) is the same as the sum of indices corresponding to negative directions \( (n_{r,\text{in}}) \). For instance, in two dimensions the sum of the left and bottom indices equals the sum of the right and top indices. By assembling such elementary objects (tracing over indices corresponding to their interface) we can construct an arbitrary domain. Each tracing automatically cancels an “in” index with an

XIV. ADDITIONAL ASPECTS

A. Symmetries and truncations

As explained in the Introduction, the implementation of field theory calculations with quantum computers requires discretizations and truncations of the problems considered. As symmetries play a crucial role in most of these calculations, we need to understand the effects of discretization on the realization of the original symmetries. The effects of the discretization of space-time are well understood and the remaining discrete symmetries—discrete translations and rotations—are used consistently by lattice practitioners. On the other hand, the fate of internal continuous symmetries in reformulations involving discrete character expansions and truncations is a more complicated question. We report here recent progress on this question that have a great deal of generality and apply to global, local, continuous and discrete symmetries (Meurice, 2019, 2020a).

We consider generic symmetries for a generic lattice model with action \( S(\Phi) \) where \( \Phi \) denotes a field configuration of fields \( \phi_\ell \) attached to locations \( \ell \) which can be sites, links or higher dimensional objects. The partition function reads
\[ Z = \int \mathcal{D}\Phi e^{-S(\Phi)}, \] (339)
with \( \mathcal{D}\Phi \) the measure of integration over the fields. We define expectation values of a function of the fields \( f \) as
\[ \langle f(\Phi) \rangle = \frac{1}{Z} \int \mathcal{D}\Phi f(\Phi)e^{-S(\Phi)}. \] (340)

A symmetry is defined as a field transformation
\[ \phi_\ell \rightarrow \phi'_\ell = \phi_\ell + \delta \phi_\ell \] (341)
such that the action and the integration measure are preserved. These invariances imply that
\[ \langle f(\Phi) \rangle = \langle f(\Phi + \delta \Phi) \rangle. \] (342)

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\[ \langle e^{i(n_1 \varphi_{r_1} + \cdots + n_M \varphi_{r_M})} \rangle = 0. \] (344)

This global selection rule can be explained (Meurice, 2019) in terms of the selection rule of the microscopic tensors at each site given in Eq. (121). It is a divergenceless condition and it can be interpreted as a discrete version of Noether’s theorem. If we enclose a site \( x \) in a small size (compared to the lattice spacing) \( D \)-dimensional cube, the sum of indices corresponding to positive directions \( (n_{r,\text{out}}) \) is the same as the sum of indices corresponding to negative directions \( (n_{r,\text{in}}) \). For instance, in two dimensions the sum of the left and bottom indices equals the sum of the right and top indices. By assembling such elementary objects (tracing over indices corresponding to their interface) we can construct an arbitrary domain. Each tracing automatically cancels an “in” index with an
“out” index and consequently, at the boundary of the domain, the sum of the “in” indices remains the same as the sum of the “out” indices. This discrete version of Gauss’s theorem is illustrated in $D = 2$ in Fig. 49. We can pursue this process until we reach the boundary. For PBC, the “in” and “out” cancel and for OBC, all the indices at the boundary are zero. In both cases, the system is “isolated” in the sense that no flux escapes to or comes from the environment. If we now remove one site out of the entire domain, the point-wise conservation inside the rest of the domain implies that the indices connecting to the missing site satisfy the divergenceless condition independently. This has a very simple interpretation in terms of global symmetry of the model (Meurice, 2020a). We can use the symmetry to fix the value of $\phi$ at the missing site instead of integrating over the possible values because as we just explained, the divergenceless condition resulting from this integration is redundant.

$$T_{nn'} = \delta_{nn'} \sqrt{I_n(\beta)I_{n'}(\beta)}$$ (345)

Figure 49 Example of flux cancelations in $D = 2$. The total flux in and out the upside-down L-shaped domain is $+1$.

We can now understand the global selection rule of Eq. (344). The insertion of various $e^{inQx}$ is equivalent to inserting an “impure” tensor which differs from the “pure” tensor by the Kronecker symbol replacement $\delta_{n,x,\text{out},n,x,\text{in}} \rightarrow \delta_{n,x,\text{out},n,x,\text{in}+nQ}$. Proceeding as before for PBC or OBC, this implies that the sum of the charges should be zero.

To summarize, the global selection rule is a consequence of the selection rule at each site which is the Kronecker delta in the expression of the tensors. It is independent of the particular values taken by the tensors (like Bessel functions). So if we set some of the tensor elements to zero as we do in a truncation, this does not affect the global selection rule and truncation are compatible with symmetries (Meurice, 2019).

The reasoning can be extended to local symmetries (Meurice, 2020a). For the CAHM, the divergenceless condition for the $n_{x,\mu}$ is redundant with the selection rule coming from the integration over the gauge fields as expressed in Eq. (182). This means that we can eliminate the $\phi$ field with the unitary gauge. It was shown (Meurice, 2019) that in the pure gauge limit the set of equations (164) are not independent. If we pick a site, we can construct an in-out partition for the legs attached to links coming out of this site. The sum of “in” indices is the same as the sum of the “out” indices, and if we assemble them on the boundary of a $D$-dimensional cube, one of the divergenceless conditions follows from the other $2D - 1$ conditions. This is illustrated for $D = 2$ in Fig. 50 where three of the delta functions on the $A$-tensors attached to the links imply the fourth one.

![Figure 50](image.png) Illustration that one divergenceless condition is redundant for $D = 2$.

The redundancy argument extends to discrete $\mathbb{Z}_q$ subgroups of $U(1)$ where the divergenceless condition is expressed modulo $q$ and the infinite set of Bessel functions are replaced by the $q$ discrete ones. We conclude that Noether’s theorem can be expressed in the tensor formulation context as: for each symmetry, there is a corresponding tensor redundancy. This applies to global, local, continuous and discrete Abelian symmetries (Meurice, 2020a).

**B. Topological considerations**

In classical field theory, the boundary conditions play an important role in the investigation of topological solutions. As a simple example, if an angle variable $\varphi$ satisfies Laplace equation in $D = 1$, then PBC allow the existence of solutions with any winding number. On the other hand, for arbitrary Dirichlet boundary conditions, the concept of winding number is not applicable because the one-dimensional interval does not have the topology of a circle.

In the $D = 1$ $O(2)$ model, we observe features which are reminiscent of this observation. For PBC, we can assemble tensors with any index $n$,
and
\[ Z_{\text{PBC}} = \text{Tr}[T^{N_x}] = \sum_{n=-\infty}^{\infty} I_n(\beta)^{N_x}. \]  
(346)

On the other hand for OBC, we have a zero index at the ends and
\[ Z_{\text{OBC}} = I_0(\beta)^{N_x} - 1. \]  
(347)

It is often believed that the contributions for various \( n \) in \( Z_{\text{PBC}} \) correspond to the topological sectors of the classical equations of motion which become Laplace’s equation in the continuum limit (so for large \( \beta \)). This is not correct because the \( I_n(\beta) \) differ from \( I_0 \) by corrections of \(-n^2/(2\beta)\) as shown in Eq. (123), while in the semi-classical solution, one expects suppression of the form \( \exp(-\beta n^2 C) \) for some calculable constant \( C \). However, the two types of behaviors are swapped after the Poisson summation
\[ \sum_{\ell=-\infty}^{\infty} e^{-\frac{B}{2} \ell^2} = \sqrt{\frac{2\pi}{B}} \sum_{n=-\infty}^{\infty} e^{-\frac{(2\pi)^2 n^2}{B^2}}, \]  
(348)

A detailed analysis of the classical solutions (Meurice, 2020a) shows that \( B = \beta(2\pi)^2/N_x \) and that summation over the winding numbers using Poisson summation and the calculation of the quadratic fluctuations reproduce precisely the leading behavior of Eq. (346) in the large \( \beta \) limit. Similar observations were made in Ref. (Akerlund and de Forcrand, 2015) for a version of the O(2) model where the fluctuations are limited.

Similar results were obtained for the \( D = 2 \) pure gage \( U(1) \) model. In these calculations, the possibility of fixing the values of variables that lead to redundant selection rules to arbitrary values as discussed in Sec. XIV.A removes the zero modes from the quadratic fluctuation calculations. Note also that it is possible to construct models where the large \( \beta \) approximations are exact. The questions of topological configurations and duality are discussed for Abelian gauge models of this type in various dimensions in Refs. (Banks et al., 1977; Gattringer et al., 2018a; Savit, 1977; Sulejmanpasic and Gattringer, 2019).

C. Quantum gravity

Tensor networks have also found a use in the study of quantum gravity. One of the directions where tensor networks have appeared is in the “spin-foam” formulation of quantum gravity (Perez, 2013). In (Dittrich et al., 2016), a tensor network formulation is developed along with a coarse-graining scheme where the intention is to use it on a spin-foam partition function, although it is applicable in other situations. The algorithm is sketched out, and some numerical results for the two- and three-dimensional Ising model are presented. In (Asaduzzaman et al., 2019) the authors start with the partition function for two-dimensional gravity where the gauge symmetry has been extended to unify the tetrad and spin-connection variables into a single connection. They present a tensor formulation and study the zeros of the partition function (Fisher’s zeros) in the complex-coupling plane.

XV. CONCLUSIONS

In summary, TrFT provides new ways to approach models studied by lattice gauge theorists. For models with compact field variables, character expansions and orthogonality relations provide ways to do the difficult integrals exactly and replace them by discrete sums. For continuous field variables, the sums are infinite and need to be truncated for practical implementations. These truncations preserve global and local symmetries.

We showed that by combining tensor blocking and truncations, we can obtain coarse-grained versions of the original model where the new “effective” tensors are assembled in the same way as the original tensors while taking different values. The TRG flows in the space of tensors replace the RG flows in the space of effective interactions. The effective tensors remain local in the coarse-grained system of coordinates. When the Euclidean action is real, TRG calculations can be compared with accurate results obtained using importance sampling in the original Lagrangian formulation. Tensor sampling could also be conducted by generalizations of the worm algorithm. TRG calculations can be extended to the case of complex actions and evade sign problems.

It is important to realize that if a reasonable control of the truncations can be reached for values of \( D_{\text{cut}} \) that are achievable with current computers, the computation cost scales logarithmically with the volume of the system which is exceptionally efficient. The scaling of the cost with the dimension can be seen as an obstacle, however recent progress in four dimensions provide an optimistic outlook.

The continuum limits of lattice models can be constructed in the vicinity of RG fixed points. Universal quantities such as the critical exponents can be extracted by linearizing the RG transformation near the fixed point. The naive approach for this program has to be amended due to the existence of unphysical fixed points. This requires a detailed understanding of the UV and IR entanglement.

TrFT allows for the construction of transfer matrices and smoothly connects the classical Lagrangian approach at Euclidean time to the Hamiltonian approach. The discreteness of the reformulation combined with truncations provides approximate Hamiltonians which are suitable for quantum simulation experiments and quantum computations. TrFT is a natural tool to design quantum cir-
cuits. In the NISQ era, benchmarking is crucial to assess the progresses in this direction. Hybrid formalisms combining real and imaginary time can be accommodated easily by TrFT and may play an important role in the near future.

We think that the TrFT program is making good progress towards the long term goal of performing QCD calculations. We expect that it will play an important role in developing practical methods to approach nuclear matter, jet physics and fragmentation. Active collaborations between the lattice gauge theory and condensed matter, jet physics and fragmentation. Active collaboration, for useful discussions and comments. This work was supported in part by the U.S. Department of Energy (DOE) under Award Numbers DE-SC0010113, and DE-SC0019139.

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Appendix A: Review of mathematical results

1. Character expansions

One of the most important relations we use in this review is the change of basis called the character expansion. In the cases considered here, this relates a compact variable (discrete and bounded, or continuous and bounded) to their Pontryagin dual. The relevant relation can be written,

\[ f(x_a) = \sum_{k_a} \lambda_{k_a} \chi_{k_a}^* (x_a). \]  

(A1)

Here \( x_a \) is the compact variable under consideration (it can be a matrix, or a spin, for example), and \( f \) must be a “class function”, a function that only depends on the trace of the compact variables. \( k_a \) is the dual variable, which takes on the values of the irreducible representations of the group that \( x_a \) belongs. Finally, \( \chi_{k_a}^* \) are the characters of the group, which are complete and orthogonal. In practice we use the following,

\[ e^{\beta \sigma} = \sum_{n=0}^{1} \lambda_n(\beta) \sigma^n, \]  

(A2)

\[ e^{\beta \cos \theta} = \sum_{n=-\infty}^{\infty} I_n(\beta) e^{im\theta}, \]  

(A3)

\[ e^{\beta \text{Tr}[U]} = \sum_{r=0}^{\infty} F_r(\beta) \chi^r(U), \]  

(A4)

for the groups \( Z_2, \ U(1), \) and \( SU(2), \) where \( \lambda_n, I_n, \) and \( F_r \) are the expansion coefficients, and \( \sigma^n, \ e^{im\theta}, \) and \( \chi^r(U) \) are the characters of the respective groups.

2. Orthogonality & completeness

As mentioned above, the expansions in Eq. (A2) are examples of the completeness and orthogonality of the characters. In each of those cases we have,

\[ \frac{1}{2} \sum_{\sigma} \sigma^n \sigma^m = \delta_{n,m}^{(2)}, \]  

(A5)

\[ \int_{-\pi}^{\pi} \frac{d\theta}{2\pi} e^{i\theta} e^{-i\theta} = \delta_{n,m}, \]  

(A6)

\[ \int dU \chi^r(U) \chi^r(U) = (2r + 1)^{-1} \delta_{r,r'}, \]  

(A7)

where \( \delta^{(2)} \) is a Kronecker delta with the equivalency taken modulo two. We also use the orthogonality of the matrix representations of group elements under the Haar measure,

\[ \int dUD^r_{mn}(U)D^{r'}_{m'n'}(U) = (2r + 1)^{-1} \delta_{r,r'} \delta_{m,m'} \delta_{n,n'} \]  

(A8)

with * as complex conjugation without transposition. The \( D \)-matrices are related to the characters through the trace, \( \chi^r(U) = \text{Tr}[D^r(U)] \).

On the other hand these characters obey a completeness relation. This is given by the sum over the representations, rather than the original group variables,

\[ \sum_{n=0}^{\infty} \sigma^n \sigma^m = \delta_{\sigma,\sigma'}, \]  

(A9)

\[ \sum_{n=-\infty}^{\infty} e^{in(\theta-\theta')} = \delta(\theta-\theta'), \]  

(A10)

\[ \sum_{r=0}^{\infty} (2r + 1) \text{Tr}[D^r(U)D^r(U')] = \delta(U,U'). \]  

(A11)

\[ \sum_{r=0}^{\infty} (2r + 1) \chi^r(UU'^{-1}) = \delta(U,U'). \]  

(A12)
3. Singular value decomposition

The singular value decomposition (SVD) plays a central role in many different parts of this review, and when dealing with compact variables it is completely determined by the character expansion. Here we give bare working knowledge of how it will be used. One of the basic ideas is to regard Boltzmann factors $f(x_\alpha, x_\beta)$ which appear in the partition function, as $K$-sic ideas is to regard Boltzmann factors $f$ working knowledge of how it will be used. One of the bare algorithms is to regard Boltzmann factors $f$ which can be expressed in terms of the characters discussed in Sec. III.D. For the $Z_q$ group, we have $K = q$,

$$U_{x_\alpha,j} = \exp \left( \frac{2\pi}{q} j x_\alpha \right), \quad (A15)$$

and

$$V_{x_\beta,j} = \exp \left( \frac{2\pi}{q} j x_\beta \right). \quad (A16)$$

Another case which occurs often is to use the SVD to split a higher-dimensional array in two. Consider a generic tensor $A_{ijkl}$ with dimensions $(d,d,d,d)$. Suppose one wanted to somehow factorize this tensor into two smaller tensors, one with indices $i$, $j$, and the other with indices $k$, $l$. Then one can do the following,

$$A_{ijkl} \rightarrow A_{(i\otimes j)(k\otimes l)} \rightarrow A_{ij} M \sum_{M,M'=1}^d U_{M\alpha}^\dagger V_{M'\beta}^\dagger, \quad (A17)$$

$$= \sum_{M,M'=1}^d U_{M\alpha}^\dagger V_{M'\beta}^\dagger, \quad (A18)$$

$$= \sum_{M=1}^d (U_{IM} \sqrt{\lambda_M}) (V_{M'J} \sqrt{\lambda_{M'}}) \quad (A19)$$

$$= \sum_{M=1}^d B_{ijM} C_{Mkl} \quad (A20)$$

which is the split we were looking for. This allows any tensor to be split exactly into smaller tensors, with the sum over intermediate states as the price.

REFERENCES


Bäniüs, Mari Carmen, Krzysztof Cichy, J. Ignacio Cirac, Karl


Chen, Jing, Hai-Jun Liao, Hai-Dong Xie, Xing-Jie Han, Rui-Zhen Huang, Song Cheng, Zhong-Chao Wei, Zhi-Yuan Xie, and Tao Xiang (2017), “Phase Transition of the q-


Korenblit, Simcha (2013), “Quantum simulations of theising model with trapped ions:devil’s staircase and arbitrary lattice proposal.”


Meurice, Yannick (2020b), Quantum field theory: a quantum


Schweizer, Christian, Fabian Grusdt, Moritz Berghuber, Luca Barbiero, Eugene Demler, Nathan Goldman, Im-

Serre, JP (1973), A Course in Arithmetic, Graduate Texts in Mathematics (Springer).


Unrmut-Yockey, Judah Francis (2017), Duality methods and the tensor renormalization group: applications to quantum simulation (PhD (Doctor of Philosophy) thesis, University of Iowa).


Villani, C (2012), Théorème vivant, Littérature Francaise (Grasset).


Wiese, Uwe-Jens (2013), “Ultracold Quantum Gases and Lattice Systems: Quantum Simulation of Lattice Gauge...