Cluster-Algorithm-Amenable Models of Gauge Fields and Matter

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Typical fermion algorithms require the computation (or sampling) of the fermion determinant. We focus instead on cluster algorithms which do not involve the determinant and involve a more physically relevant sampling of the configuration space. We develop new cluster algorithms and design classes of models for fermions coupled to $\mathbb{Z}_2$ and $U(1)$ gauge fields that are amenable to being simulated by these cluster algorithms in a sign-problem free way. Such simulations should contain rich phase diagrams and are particularly relevant for quantum simulator experiments.

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1. Introduction

Fermions that strongly interact with each other via gauge fields are an essential ingredient for many models in particle physics. For example, the fermions may constitute the matter component, such as leptons in electromagnetic and weak interactions, or quarks in strong interactions, and then the gauge fields may constitute the photon, the \( W^\pm, Z \), and the gluons respectively. Gauge fields are also becoming increasingly important to condensed matter systems, from frustrated magnetism to theories of deconfined quantum criticality. It thus is interesting to develop efficient simulations for interacting theories of gauge fields and matter.

While Quantum Monte Carlo (QMC) methods are robust simulation methods for non-perturbative studies of the aforementioned systems, they are also vulnerable to the sign problem \([1]\). QMC methods work by performing importance sampling of configurations that make up the partition function. Since fermions anti-commute, their sign problem can be straightforwardly understood when the configurations considered are worldlines: whenever fermions exchange positions an odd number of times, the configuration weight acquires another negative sign factor, leading to huge cancellations in the summation and an exponential scaling in the volume for calculations.

Meron cluster methods \([2]\), so named due to the presence of merons (half-instantons) in the first model for which they were developed to simulate (the 2d \( O(3) \) sigma model with \( \theta = \pi \)), can solve sign-problems in four-fermion Hamiltonians for certain parameter regimes, as well as for free fermions with a chemical potential \([3]\). Because these methods sample worldlines, computing the weights scales linearly with the volume of the system, and negative terms in the partition function are taken care of by avoiding merons — this is what distinguishes them from bosonic simulations. Because there are none of the stabilization issues that can arise in determinantal methods, and the weight computations scale favorably compared to determinantal methods, these cluster methods are an attractive choice for simulation when applicable. Correspondingly, exciting opportunities open up when new interesting physical models are found which can be simulated using this method.

Recently, there has been experimental development regarding the physics of confinement and quantum spin liquids using quantum simulation. The models used to capture the physics involve fermions interacting with (Abelian) gauge fields. In these proceedings we introduce new cluster algorithms and develop designer\([4]\) classes of experimentally relevant models which are sign-problem-free for these cluster simulations, enabling a robust elucidation of their phase diagrams. More details may be found in our full manuscript \([5]\). Moreover, the worldline nature of the configurations makes it easy to produce measurement “snapshot” synthetic data to compare with quantum simulators, and such configurations are promising inputs for machine learning algorithms.\([6]\)

2. Models

We build these cluster-algorithm amenable models by beginning with the half-filled \( t-V \) model,

\[
H = \sum_{\langle xy \rangle} \left[ -\frac{t}{2} \left( c_x^\dagger c_y + c_y^\dagger c_x \right) + V \left( n_x - \frac{1}{2} \right) \left( n_y - \frac{1}{2} \right) \right].
\]

Here \( \langle xy \rangle \) are nearest neighbor sites, \( c^\dagger \) and \( c \) are creation and annihilation operators respectively, and the repulsive interaction \( V \) is given in terms of the occupation number \( n = c^\dagger c \). We begin with this model because it is simuable by meron clusters for \( V \geq 2t \) \([3, 7]\).
With this starting point, we now build up cluster-amenable Hamiltonians involving gauge fields. They are organized into $\mathbb{Z}_2$- and $U(1)$- gauge symmetric families, which involve the following terms:

$$H_{N_f}^{(g)} = -\sum_{(xy),f}^{N_f} \left( H_{(xy),f}^{(g)} + H_{(xy),f}^{(g),\text{des}} \right)$$

The label $g \in \{U(1), \mathbb{Z}_2\}$ is the gauge symmetry, with

$$H_{(xy),f}^{\mathbb{Z}_2} = t \left( c_{x,f}^+ s_{xy,f}^1 c_{y,f} + c_{y,f}^+ s_{xy,f}^1 c_{x,f} \right),$$

$$H_{(xy),f}^{U(1)} = t \left( c_{x,f}^+ s_{xy,f}^1 c_{y,f} + c_{y,f}^+ s_{xy,f}^1 c_{x,f} \right).$$

The hopping of spinless fermions between the nearest neighbor sites $xy$ are now governed by the presence of gauge fields, represented by spin-1/2 operators, $s_{xy}^k$, on the bond. Figure 1 illustrates the lattice with the gauge field and matter degrees of freedom. These terms are lower-dimensional versions of quantum electrodynamics (QED). Then $H_{(xy),f}^{(g),\text{des}}$ is a designer term[4] that makes the models particularly amenable to the cluster algorithms, (we consider $V = 2t$, for simplicity)

$$H_{(xy),f}^{\mathbb{Z}_2,\text{des}} = -2t \left( n_{x,f} - \frac{1}{2} \right) \left( n_{y,f} - \frac{1}{2} \right),$$

$$H_{(xy),f}^{U(1),\text{des}} = -t \left( n_{x,f} - \frac{1}{2} \right) \left( n_{y,f} - \frac{1}{2} \right) - t s_{xy,f}^3 \left( n_{y,f} - n_{x,f} \right) + \frac{t}{4}$$

For the $\mathbb{Z}_2$ gauge theory, the Pauli matrix $s^1 = \sigma^1/2$ couples to fermions, and one can see the local $\mathbb{Z}_2$ symmetry from the operator $Q_x$, which commutes with the $H^{\mathbb{Z}_2}$ and is given by $Q_x = (-1)^{\sum_f n_{x,f}} \prod_{f, \hat{a}} s_{x+\hat{a},f}^3 s_{x-\hat{a},f}^3$, where $\hat{a}$ are the unit vectors in a $d$-dimensional square lattice. For the $U(1)$ theory, one can see the $U(1)$ symmetry from the unitary operator $V_{U(1)}$ which commutes with $H^{U(1)}$ and is given by $V_{U(1)} = \prod_x e^{i\theta_x G_x}$, with $G_x = \sum_f \left[ n_{x,f} - \sum_{\hat{a}} s_{x,\hat{a},f}^3 - s_{x,-\hat{a},f}^3 \right] + \left( (-1)^x - 1 \right)/2$. These models are quantum link models [9], which realize the continuous gauge invariance using finite-dimensional quantum degrees of
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Figure 2: Worldlines for the $t$-$V$ model are in (a)-(c), the $\mathbb{Z}_2$ theory in (d)-(f), and the $U(1)$ theory in (g)-(i). Image (a) shows the imaginary time direction and the $(1+1)$-d trotterization, which is the same for all images. Filled circles are sites occupied by fermions, and empty circles are holes. Figures in the second two rows also have link variables because they correspond to gauge theories: the upward triangles correspond to spin $+1/2$ and the downward triangles correspond to spin $-1/2$. While the fermionic worldlines are the same in each column, some configurations that are allowed for the $t$-$V$ model have zero weight for the $\mathbb{Z}_2$ and $U(1)$ theories. These are crossed out, and zero-weight plaquettes are shaded red.

The identification with usual gauge field operators is given by $U_{xy,f} = s^+_{xy,f}$, $U_{xy,f}^\dagger = s^-_{xy,f}$, $E = s^3_{xy,f}$. We note that a straightforward application of the meron idea necessitates the introduction of an equivalent flavor index for gauge links as fermion flavors. The total Gauss law can be expressed through a product ($\mathbb{Z}_2$) or sum ($U(1)$) of the Gauss law of individual flavors degrees of freedom, and the resulting theories have $\mathbb{Z}_2^{\otimes N_f}$ and $U(1)^{\otimes N_f}$ gauge symmetry. However, flavored gauge-interactions can also be turned on in the $U(1)$ model (as explained in our manuscript [5]), $H_{\mathbb{Z}_2}^{U(1)} \to H_{\mathbb{Z}_2}^{U(1)} + J \sum_{(xy)} s^3_{x,y,1} s^3_{x,y,2}$, or through a Hubbard-U interaction for both $\mathbb{Z}_2$ and $U(1)$-symmetric models [7]. These additions would directly cause ordering for either the gauge fields or fermions, with the coupling between them leading to the interesting question of how the other degrees of freedom are affected by this ordering.

3. Algorithm

We first consider the possible worldline configurations for the base models defined in (2) and (3). Here we use the occupation number basis for the fermions and the electric flux (spin-$z$) basis. The partition function in $(1+1)$-d is then

$$Z = \text{Tr} \left( e^{-\beta H} \right) = \sum_{(s,n)} \langle s_1, n_1 | e^{-\epsilon H_e} | s_{2N_1}, n_{2N_1} \rangle \langle s_{2N_1}, n_{2N_1} | e^{-\epsilon H_o} | s_1, n_1 \rangle \\ \times e^{-\epsilon H_o} \ldots e^{-\epsilon H_e} | s_2, n_2 \rangle \langle s_2, n_2 | e^{-\epsilon H_o} | s_1, n_1 \rangle,$$

where $H = H_e + H_o$, and $H_e$ ($H_o$) consists of Hamiltonian terms that correspond to even (odd) links. This is a Trotter approximation, and all terms within $H_e$ and $H_o$ commute with each other. We
thus have a sum of terms that consist of discrete time-slices $1, \cdots, 2N_t$, with defined electric flux and fermion occupation numbers for each of the time-slices. Each of the terms in (5) is a worldline configuration. Figure 2(a)-(c) give examples of such configurations for the $t$-$V$ model as simulated by meron clusters. We note that these configurations are similar to the “snapshots” obtained from experiment in [6], but involve more information from the additional imaginary time direction (one snapshot would be one horizontal line in a configuration from Figure 2), so we are able to obtain imaginary time displaced correlations from these configurations.

In the $Z_2$ case, for each time-slice a fermion may hop to an unoccupied nearest neighbor site of the same flavor. If the fermion hops, then the electric flux on the bond between the nearest neighbor sites that shares the same fermion flavor index must also flip--this is due to the $s^I_{xy}$ operator. Figure 2(d)-(f) gives example configurations for the $N_f = 1$ version of this model. Due to the trace, it is impossible to have odd winding numbers because these would cause the spins in the initial state to not match the spins in the final state. The possible worldline configurations for the $U(1)$ are similar to the $Z_2$ case, but more restrictive. The $s^+_{xy}$ and $s^-_{xy}$ operators allow the hopping for a given flavor of fermion only in a single direction for each bond, which depends on the orientation of the same flavored flux on the bond. Figure 2(g)-(i) illustrates an example configuration and restrictions for the single flavor version of the $U(1)$ model. In (1 + 1)-d it is clear that all allowed configurations must have zero winding number.

The worldline configurations are a tool to obtain cluster configurations by introducing appropriate breakups, which decompose the terms in (5) into further constituents. In considering the allowed worldline configurations given in (2) for the $U(1)$ theory, for example, each of the active plaquettes in each time-slice (shaded in gray) must be one of the plaquettes given in Table 1. The plaquettes in each row share the same weight, computed using $\langle s_{b}, n_{b} | e^{-\epsilon H_b} | s'_{b}, n'_{b} \rangle$, from (5), where $b$ is a nearest neighbor bond, $b = \{x, y\}$. The corresponding breakup cell for each row gives allowable breakups: if all fermion occupations/spins are flipped along any one of the lines, the resulting plaquette also exists in this table. From the table, we see two such breakups are defined, $A$ and $D$. These breakups resemble the breakups from the original fermionic meron cluster algorithms, but also involve the link variables--either as additional lines for the $A$ breakups, or as binding lines extending outward from the horizontal $D$ breakup lines. This is a key difference for the new gauge version of the algorithm. By computing the matrix elements for the plaquettes in each grouping, we find that for the $U(1)$ theory, the corresponding breakup weights $w_A$ and $w_D$ must obey:

$$w_A = 1$$

$$w_D = \exp (\epsilon t) \sinh \epsilon t$$

$$w_A + w_D = \exp (\epsilon t) \cosh \epsilon t,$$

(6)

to satisfy detailed balance. We can then simulate this system similarly to the simulation of the $t$-$V$ model at $V = 2t$ [3], by exploring a configuration space where each configuration is defined by the worldlines and breakups. By assigning breakups to all active plaquettes, we form clusters, and then updates involve flipping all fluxes and fermions within a cluster, which physically corresponds to generating a new worldline configuration. The algorithm begins by putting the system in a reference configuration, defined by the fermionic worldlines only, where the weight is known to be positive. It is straightforward to see that all configurations can be returned to the fermion worldlines.
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Table 1: Plaquettes and breakups for the $U(1)$-symmetric Hamiltonian. The middle vertical cluster lines on the $A$-breakups and gauge binding lines on the $D$-breakups distinguish them from the purely fermionic versions of these breakups.

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of this configuration by cluster flips. For both the $U(1)$ and $Z_2$ theories, the reference configuration has a staggered Fermionic occupation, where fermions are stationary throughout imaginary time. Fluxes and breakups may be initially attached to spacetime plaquettes in any nonzero configuration according to Table 1. A QMC sweep is then:

1. Go through the list of the active plaquettes and update each breakup sequentially.

   (a) If the breakup can be changed for a plaquette, change it with probability dependent on the breakup weights.

   (b) If the breakup is changed, consider the new configuration that would result. If it contains a cluster where flipping the fermion occupation causes the fermions to permute in a way that produces a negative sign, then it is a meron. In that case, change the breakup back to its initial state. Rules for identifying merons generalize [3] and are given in greater detail below.

2. Identify the new clusters from the breakups in the new configuration. For each cluster, flip all fermion occupation states and spins with probability $1/2$.

This describes sampling of the zero-meron sector only, but sectors with other numbers of merons are relevant for off-diagonal observables. [3] While these cluster rules implement the Hamiltonian dynamics, Gauss law constraints are not included.

4. Results

To illustrate the efficacy of the algorithm, we discuss results obtained by simulating the $(1+1)$-d $H_{N_f=1}^{U(1)}$ model in (2), which is related to the massless quantum-link Schwinger model and the PXP model, where quantum scars were first demonstrated experimentally [8]. We simulate the model
Figure 3: Number of configurations versus Gauss law sector index $\sum_x [G_x + 2] \cdot 4^x$ (not all indices correspond to actual sectors) for 50000 equilibrated configurations. Two sectors emerge at large $\beta$: $G_x = 0$ and $G_x = (-1)^x$.

Figure 4: The types of clusters that appear as a function of temperature. None of them are merons because the simulation is in $1 + 1d$.

for different temperatures $\beta = 1/T$, without imposing the Gauss law. A filter may then be applied to study the physics in the desired Gauss law sector. The first non-trivial result is the emergence of two Gauss’ law sectors at low temperatures, as shown in Figure 3, other results are given in our manuscript [5].

For purely fermionic models, as shown in [3], a loop is a meron if the quantity $n_w + n_h/2$ is even, where $n_w$ is the number of temporal windings and $n_h$ is the number of fermionic hops in the loop. For the new gauge-fermion algorithm, we now can bind two or more loops to each other through the gauge fields, and thus we have the following new meron criterion for a cluster

$$\text{Meron if } \begin{cases} n_w + n_h/2 \text{ odd, even # of loops} \\ n_w + n_h/2 \text{ even, odd # of loops} \end{cases}$$

(7)

It can be seen immediately that (7) reduces to the original definition in the case of one loop. In one dimension there should be no merons. Figure 4 plots the $n_h$ and $n_w$ for 100 equilibrated clusters at low temperature, and indeed we see that by the criterion, none of these clusters are merons.

5. Conclusions

We have generalized the construction of the meron algorithm to cases where staggered fermions are coupled to quantum link gauge fields. This construction of the Monte Carlo algorithm is agnostic to the space-time dimension, and paves the way for unbiased studies of large scale gauge-fermionic system with odd or even numbers of fermionic flavours, and includes models not simulable using determinantal Quantum Monte Carlo. It is possible to add different microscopic terms by increasing the allowed ways of bonding the fermions and gauge links. Future extensions could include gauge fields with larger spin representation and non-Abelian gauge fields as well.
References


