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Phys. Rev. A **105**, 023322 — Published 22 February 2022 DOI: 10.1103/PhysRevA.105.023322

Cold Atom Quantum Simulator for String and Hadron Dynamics in Non-Abelian Lattice Gauge Theory

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(Dated: February 4, 2022)

We propose an analog quantum simulator for simulating real-time dynamics of (1 + 1)-d non-Abelian gauge theory well within the existing capacity of ultracold atom experiments. The scheme calls for the realization of a two-state ultracold fermionic system in a 1-dimensional bipartite lattice, and the observation of subsequent tunneling dynamics. Being based on novel loop string hadron formalism of SU(2) lattice gauge theory, this simulation technique is completely SU(2) invariant and simulates accurate dynamics of physical phenomena such as string breaking and/or pair production. The scheme is scalable and particularly effective in simulating the theory in weak coupling regime, and also a bulk limit of the theory in strong coupling regime up to certain approximations. This paper also presents a numerical benchmark comparison of the exact spectrum and real-time dynamics of lattice gauge theory to that of the atomic Hamiltonian with an experimentally realizable range of parameters.

INTRODUCTION I.

Gauge field theories constitute an exceptionally powerful theoretical framework that describes at least three of the four fundamental interactions of nature. Non-Abelian gauge symmetry lies at the heart of the standard model of particle physics. Quantum chromodynamics (QCD), which is an SU(3) gauge theory, can accurately represent quark-gluon interactions. In 1974, Wilson proposed a regularization of the gauge theory on space-time lattices [1] that exhibits quark confinement in the strong coupling limit. Wilson's lattice gauge theory (LGT) has been used extensively over the past four-five decades because one can perform lattice QCD calculations by Monte Carlo simulations [2]. World's largest super-computing resources are now being employed for the same [3].

Although the lattice QCD numerical scheme is very efficient, there is the infamous 'sign problem' that limits its applicability [4]. For example, it cannot handle systems with finite and non-zero density or calculate realtime dynamics within the Euclidean framework. Following Feynman's visionary idea [5], quantum simulation of lattice QCD offers hope to address these issues. With the recent technological progresses, there has been a surge of interest towards developing quantum algorithms to study gauge theories using both digital and analog approaches [6–13]. However, the progress in quantum simulation of non-Abelian gauge theories lags far behind its Abelian counterparts. The present work outlines a scheme for simulating real-time dynamics of a non-Abelian gauge theory in an analog way. In particular, we demonstrate the simulation of a manifestly gauge-invariant framework, namely loop-string-hadron (LSH) formalism [14] of SU(2) gauge theory. This scheme successfully bypasses the nontrivial task of imposing the non-Abelian gauge invariance (local constraints) additionally.

The concept of analog quantum simulation involves mimicking a quantum system described by a Hamiltonian (simulated Hamiltonian) by a different quantum system described by some other Hamiltonian (simulating Hamiltonian). Systems of ultracold atoms [15] or ions [16]trapped in optical lattices serve as excellent quantum simulators, as the relevant parameters can be precisely measured and controlled. It can be recalled that though the experimental realization of Bose-Einstein condensate and trapped ultracold fermions [17–23] initially inspired studies on the macroscopic quantum coherent phenomena only [24–26], soon it was discovered that ultracold atoms can serve as wonderful testing grounds for other branches of physics as well. The atom-atom scattering length (and thus, the interaction strength) in ultracold gases can be varied across a wide range via Feshbach resonances. The creation of optical lattices [27] by using two counter-propagating coherent laser beams took this tunability a step further: as the size, shape, and dimensionality of the lattice could be easily controlled. In the past, cold atomic systems have successfully emulated a rich variety of systems and addressed problems in disordered systems, spin liquids, superconductivity, nuclear pairing, artificial gauge fields, and topology [28–30]. Advanced cooling and trapping methods have now led to quantum engineering at an unparalleled precision level. This allows for each individual atom to be monitored, and one can have a perfect quantum simulator.

Over the last few years, there has been a continuous pursuit of using cold atom systems for analog quantum simulating both Abelian and non-Abelian lattice gauge theories. Mostly based on Kogut-Susskind (KS) formalism [31-34] as well as Quantum Link Model(QLM) formulation [35–37] and also Abelian Higgs Model [38–40],

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these schemes involved a careful designing of the set-up so that the system remains in the gauge-invariant Hilbert space throughout the dynamics. Quantum simulating non-Abelian gauge theories using Rydberg atom gates [41] was also proposed.

Following the first experimental demonstration of a digital quantum simulation of a lattice Schwinger model [42], a density-dependent quantum gauge field was experimentally demonstrated [43], which is useful for simulating dynamics of \mathbb{Z}_2 gauge theory. The first analog quantum simulation of \mathbb{Z}_2 gauge theory on a two staggered site lattice by cold atom quantum simulator was reported in [44]. The first experiment demonstrating a scalable quantum simulation of continuous gauge theory [45] has also been reported recently.

The major difficulty in any Hamiltonian simulation of gauge theory is to impose the local constraints (Gauss law) and to keep the dynamics confined within the physical Hilbert space that satisfies the constraint. Recently, such a gauge invariance has been experimentally demonstrated for analog simulation of U(1) gauge theory on a sufficiently large lattice [46]. The notion of gauge invariance becomes many-fold complicated for a non-Abelian theory such as SU(2) where there exists more than one mutually non-commuting constraint at each lattice site. That might be a reason why a practically/immediately realizable analog quantum simulation scheme for simulating dynamics even with the simplest non-Abelian, continuous gauge group such as SU(2) in (1+1)-d is absent in the literature till date .

We aim to quantum simulate the same, i.e SU(2) lattice gauge theory on a (1 + 1)-d dimensional lattice described by Kogut-Susskind (KS) Hamiltonian [47]. In a recent study [48], it has been demonstrated that amongst many variants of Hamiltonian formulation of non-Abelian gauge theories [49–54], the loop string hadron (LSH) formalism [14] is the most convenient and computationally least expensive one for (1+1)-d within the scope of classical computation. The reason is, being a manifestly gauge invariant formalism, the LSH Hamiltonian describes the dynamics of only relevant physical degrees of freedom. In a 1-d spatial lattice, that is precisely the dynamics of strings and hadrons. It can be shown [48, 55] that in (1+1)-d, any gauge theory with open boundary condition can be mapped to a theory of only fermions, i.e., equivalent to the XYZ model and hence much simpler to analyze. The novel LSH formalism shares many features of this purely fermionic formalism but can actually be generalized to periodic boundary conditions as well as to higher dimensions [14].

The present paper exploits this versatility of LSH formalism of SU(2) gauge theory. Here, different parameter regimes of SU(2) gauge theory are mapped to different parameter regimes of an atomic Hamiltonian: that of an ionic Hubbard model. We consider the half-filled Hubbard model to be exactly equivalent to the gauge theory Hilbert space containing a strong coupling vacuum (no matter/anti-matter state). We show that the spectrum, obtained with exact diagonalization of both the simulating and simulated systems compared remarkably in the weak coupling regime. We also provide a benchmark comparison of the dynamics of the atomic system directly mapped to the pair production-string breaking dynamics of the low energy sector of SU(2) gauge theory. The numerical analysis employs parameters and experimental set-ups already realized with ultracold atom systems. We demonstrate two key points: (i) the full gauge theory Hamiltonian can be reduced to an approximated LSH Hamiltonian, which, in turn, can be perfectly mimicked by the atomic system to the low energy dynamics in the weak coupling limit of gauge theory, and (ii) for the strong and intermediate coupling regimes, the difference between the full gauge theory Hamiltonian and the approximated Hamiltonian is slightly more prominent : but it can be compensated by tuning the on-site interaction parameter of the Hubbard Hamiltonian. Thus, one can still access the dynamics of strings and hadrons in presence of a background gauge field in the bulk limit of the lattice. Further improvements of this scheme to include dynamical gauge fields in the higher dimensions, and also generalization to SU(3) gauge theory will take us close to quantum simulating the full QCD.

The plan of the paper is as follows: The section II contains the minimal details of the lattice gauge theory Hamiltonian including LSH framework at different coupling regimes that we aim to quantum simulate. The simulating Hamiltonian is discussed in section III, including the atomic system to be used for the quantum simulation scheme, i.e. a fermionic Hubbard model on a bipartite lattice and specification of the parameters of the simulating Hamiltonian to simulate the gauge theory at wide range of coupling regimes. In section IV the proposed experimental set-up is described. Section V contains numerical study and comparison of the spectrum and real time dynamics of both the simulating and simulated systems using the parameters for the proposed experimental scheme and also . Finally, in section VI the results are summarized and also the future prospects have been discussed.

II. THE THEORY TO BE SIMULATED

In this section, we briefly review the theory we would like to simulate (subsection II A), discuss different parameter regimes of interest (subsection II B), propose a mean field ansatz (subsection II C) and apply the ansatz to a gauge invariant formalism for the same theory (subsection II D). Finally, the Hamiltonian we quantum simulate in presented (subsection II E).

A. Kogut-Susskind Hamiltonian

Hamiltonian or canonical formulation of lattice gauge theories was developed by Kogut and Susskind [47] right after Wilson introduced lattice gauge theory originally in Euclidean formalism [1]. In the classical computing era, lattice gauge theory calculations have explored the original Euclidean formulation extensively, but the Hamiltonian framework remained a relatively uncharted territory. However, the interest in the Hamiltonian description of lattice gauge theories is renewed, as it turns out to be the natural framework to work with in the upcoming quantum simulation/computation era. The mostly used formalism in this context is Quantum Link Model representation of gauge theory as it provides a finite dimensional representation of the gauge fields. There is a drawback though : in smaller dimensions, that are accessible by present-day quantum technology, the quantum link model does not have the desired spectrum as obtained with the original Kogut-Susskind Hamiltonian [48, 49]. In this work we consider the original Kogut-Susskind Hamiltonian for the simplest non-Abelian gauge group, i.e. SU(2) and proceed to construct a quantum simulator for the same in (1+1)-d.

The Kogut-Susskind (KS) Hamiltonian describing SU(2) Yang Mills theory coupled to staggered fermions on (1 + 1)-d (1d spatial lattice and continuous time) [47] can be written as:

$$H^{(\rm KS)} = H_E^{(\rm KS)} + H_M^{(\rm KS)} + H_I^{(\rm KS)}.$$
 (1)

Where, $H_E^{(\text{KS})}$ corresponds to electric part of the Hamiltonian given by,

$$H_E^{(\text{KS})} = \frac{g^2 a}{2} \sum_{j=0}^{N-1} \sum_{a=1}^3 E^a(j) E^a(j).$$
(2)

Here,

$$\sum_{a=1}^{3} E^{a}(j)E^{a}(j) = \sum_{a=1}^{3} E^{a}_{L}(j)E^{a}_{L}(j) = \sum_{a=1}^{3} E^{a}_{R}(j)E^{a}_{R}(j)$$

for left and right electric fields $\mathbf{E}_{L/R}$ associated with a link connecting sites j and j + 1.

The staggered fermionic matter ψ , in the fundamental representation of SU(2) consisting of two components $\begin{pmatrix} \psi_1 \\ \psi_2 \end{pmatrix}$ yields a staggered mass term:

$$H_M^{(\rm KS)} = m \sum_{j=0}^N (-1)^j \left[\psi^{\dagger}(j) \cdot \psi(j) \right].$$
 (3)

 $H_I^{(\text{KS})}$ denotes interaction between the fermionic and gauge fields and is given by:

$$H_I^{(\text{KS})} = \frac{1}{2a} \sum_{j=0}^{N-1} \left[\psi^{\dagger}(j) U(j) \psi(j+1) + \text{h.c.} \right].$$
(4)

The gauge link U(j) is a 2 × 2 unitary matrix defined on the link connecting sites j and j + 1. A temporal gauge is chosen to derive the above Hamiltonian which sets the gauge link along the temporal direction equal to unity. The color electric fields $E^a_{L/R}$ are defined at the left L and right R sides of each link and they satisfy the following commutation relations (SU(2) algebra) at each end:

$$\begin{split} [E_{L}^{a}(j), E_{L}^{b}(j')] &= i\epsilon^{abc}\delta_{jj'}E_{L}^{c}(j), \\ [E_{R}^{a}(j), E_{R}^{b}(j')] &= i\epsilon^{abc}\delta_{jj'}E_{R}^{c}(j'), \\ [E_{L}^{a}(j), E_{R}^{b}(j')] &= 0, \end{split}$$
(5)

where ϵ^{abc} is the Levi-Civita symbol. The electric fields and the gauge link satisfy the following quantization conditions at each site,

$$[E_{L}^{a}(j), U(j')] = -\frac{\sigma^{a}}{2} \delta_{jj'} U(j),$$

$$[E_{R}^{a}(j), U(j')] = U(j) \delta_{jj'} \frac{\sigma^{a}}{2},$$
 (6)

where σ^a are the Pauli matrices. The Hamiltonian in (1) is gauge invariant as it commutes with the Gauss' law operator,

$$G^{a}(j) = E_{L}^{a}(j) + E_{R}^{a}(j-1) + \psi^{\dagger}(j)\frac{\sigma^{a}}{2}\psi(j)$$
(7)

at each site j. The physical sector of the Hilbert space corresponds to the space consisting of states annihilated by (7). Solving the non-Abelian Gauss laws at each site j as given in (7) is non trivial and engineering the same in an analog experiment is the most difficult job.

In a very recent work [48], all available formalisms for non-Abelian gauge theory with gauge group SU(2)in (1 + 1)-d has been analyzed and compared in terms of their applicability in Hamiltonian simulation. As concluded in [48], the recently developed LSH formalism [14] enjoys two unique advantages: (i) It is exactly equivalent to the original Kogut-Susskind Hamiltonian (ii) it removes the non-trivial steps (computational costs) required in the original Hamiltonian formulation to contain the dynamics in the gauge invariant sector of LGT Hilbert space. The second advantage becomes particularly important in designing analog/digital quantum simulator [46, 56]. That is why we choose the novel LSH framework to describe gauge theory and map the same to an atomic Hamiltonian. It is already established [48] that the original Kogut-Susskind Hamiltonian, and the LSH Hamiltonian (given in the Appendix A) share identical spectrum and hence generate the same dynamics. At this point we must mention that all the feasible/implemented past proposals involve QLM formulation of lattice gauge theory, that in lower dimension exhibits a complete different spectrum as well as a different Hilbert space than that of the Kogut-Susskind Hamiltonian.

B. The two coupling regimes

It is convenient to scale the Hamiltonian $H^{(KS)}$ given

in (1) as per [57], so as to make it dimensionless:

$$\tilde{H} = \frac{2}{g^2 a} H^{(\text{KS})}$$

$$= \underbrace{\sum_{j} E^2(j)}_{\tilde{H}_E} + \underbrace{\mu_0 \sum_{j} (-1)^j \left[\psi^{\dagger}(j) \cdot \psi(j)\right]}_{\tilde{H}_M} + \underbrace{x_0 \sum_{j} \left[\psi^{\dagger}(j)U(j)\psi(j+1) + \text{h.c.}\right]}_{\tilde{H}_I}.$$
(8)

Here, $x_0 = \frac{1}{g^2 a^2}$ and $\mu_0 = 2\sqrt{x_0} \frac{m}{g}$ are dimensionless coupling constants of the theory. Evolving this \tilde{H} with scaled time (from zero to $\tilde{\tau}$)

$$\tilde{\tau} = \frac{\tau_{\text{gauge}}}{x_0} \tag{9}$$

is due to the unitary operator:

$$\mathcal{U}(\tilde{\tau}) = \exp\left(-i\tilde{H}\tilde{\tau}\right)$$
$$= \exp\left(-i\frac{2}{g^2a}H^{(\mathrm{KS})}g^2a^2\tau_{\mathrm{gauge}}\right)$$
$$= \exp\left(-i2aH^{(\mathrm{KS})}\tau_{\mathrm{gauge}}\right). \tag{10}$$

Here, $2aH^{(KS)}$ is another scaled Hamiltonian with dimensionless parameters given by,

$$2aH^{(KS)} = \frac{1}{x_0} \sum_{x} E^2(x) + 2\frac{m}{g} \frac{1}{\sqrt{x_0}} \sum_{j} (-1)^j \left[\psi^{\dagger}(j) \cdot \psi(j) \right] + \sum_{j} \left[\psi^{\dagger}(j)U(j)\psi(j+1) + \text{h.c.} \right].$$
(11)

The strong coupling limit is defined for $x_0 \rightarrow 0$, where the interaction part of Hamiltonian become less dominant as evident from both (8) and (11). On the other hand, in the weak coupling limit defined at $x_0 \rightarrow \infty$, interaction part of the Hamiltonian becomes the most important term that cannot be treated perturbatively. These scaling rules work equivalently on the LSH Hamiltonian defined in (A18, A19, A20) as the LSH Hamiltonian is exactly equivalent to the original Kogut-Susskind Hamiltonian.

In the strong coupling regime, lattice gauge theory shows the desired physics such as quark confinement and finite mass gap. In this limit, the interaction terms in (4) that involves transitions between different eigenstates of the electric-field operator becomes insignificant, and hence in the Hamiltonian, diagonal terms dominate over the off-diagonal ones in the strong coupling basis ¹. As $x_0 \to 0$, only very small electric flux configurations on the lattice contribute to the low energy sector of the theory. In this regime, lattice Hamiltonian matrices can be analyzed perturbatively with the electric part as the unperturbed Hamiltonian. Order by order perturbation corrections yield a finite dimensional Hilbert space, within a cut-off imposed on the bosonic quantum number corresponding to gauge flux. The computation cost rises exponentially with increasing Hilbert space dimension, that grows with system size as well as cut-off [48]. As a result, calculating Hamiltonian dynamics for an arbitrary large system even with the largest possible computer seems impossible. However, the continuum limit of the LGT lies in the opposite regime, where $x_0 \to \infty$ $(q \to 0, a \to 0)$ together with bulk limit , i.e lattice size $N \to \infty$. In this regime, the dynamics becomes heavily cut-off sensitive. Here all possible electric flux states contribute to the low energy spectrum with major contributions coming from strong coupling basis states with electric flux values to grow larger with $x_0 \to \infty$. The Hamiltonian moves away from diagonal structure as (4) becomes dominants. As a whole, analyzing the weak coupling limit of lattice gauge theory is extremely difficult on a classical computer except some extrapolation technique of strong coupling analysis.

Now, we propose a mean field ansatz for the low energy sector of the gauge theory Hilbert space in the weak coupling regime. In the next section, we propose an analog quantum simulator to simulate the dynamics of gauge theory in this regime. However, this particular novel proposal accurately simulates the dynamics of gauge theory beyond this particular regime as well, within the mean field ansatz. Quantum simulation of the intermediate coupling regime of the full SU(2) gauge theory involves suitable tuning of atomic interactions as described in later part of this work.

C. Weak coupling limit: Mean field ansatz

As stated earlier, we choose the LSH representation of the Kogut-Susskind Hamiltonian (Given in Appendix A) as it provides the most convenient and economic description of the physical degrees of freedom and their dynamics. The LSH basis is characterized by three integer quantum numbers

$$n_l(j) \in (0,\infty) \& n_i(j), n_o(j) \in (0,1).$$
 (12)

for each site j

The strong coupling vacuum of the theory is defined by zero electric flux state. In the LSH formalism, the

¹ For LGT, the natural and most convenient basis is formed out

of eigenstates of the electric-field operator. Tensor product of fermionic occupation number basis and electric field basis constitute the full Hilbert space. This particular basis, being eigenbasis of the diagonal Hamiltonian $(H_E + H_M)$ in the $g \to \infty$ limit, is called the strong-coupling basis of LGT.

same state is given by $n_l = 0$ in (12) at all lattice sites. However, as one approaches the weak coupling regime, the low energy spectrum of the theory contains states that carry large fluxes. In [58], a weak coupling vacuum ansatz was proposed and justified for the (2+1)-d pure SU(2) gauge theory within prepotential framework. In that proposal, each lattice site contains a large but mean value for the local loop quantum numbers. The (1+1)-d version of that ansatz within LSH framework (i.e prepotential + staggered matter) would be equivalent to each site containing more and more gauge fluxes, i.e $n_l \gg 0$, for all sites as one approaches the weak coupling limit $g \to 0$. As discussed before, the incoming flux or boundary flux l_i fixes the bosonic loop quantum numbers n_i 's at each of the lattice site for any configurations of n_i, n_o throughout the lattice as per (A30). Hence, choosing for $l_i \gg 0$ for any finite lattice would result,

$$n_l(j) = l_i \equiv n_l \quad \forall j. \tag{13}$$

D. Approximate Hamiltonian

In this section, we present a particular form of LSH Hamiltonian that we would like to simulate. This is derived from the Hamiltonian given in (A18, A19, A20), that generates the spectrum of the original Kogut Susskind Hamiltonian discussed before.

Electric Hamiltonian: The electric part of the LSH Hamiltonian as given in (A18) can be written as:

$$H_E^{(\text{LSH})} = \frac{g^2 a}{2} \sum_j h_E(j) \tag{14}$$

At each site j, depending upon the fermionic quantum numbers n_i, n_o , the local contribution to electric energy is given by,

$$\frac{n_{i} \ n_{o}}{0} \ \frac{h_{E}}{0} \ \frac{n_{l}}{2} \left(\frac{n_{l}}{2}+1\right)$$

$$0 \ 1 \ \frac{n_{l}+1}{2} \left(\frac{n_{l}+1}{2}+1\right)$$

$$1 \ 0 \ \frac{n_{l}}{2} \left(\frac{n_{l}}{2}+1\right)$$

$$1 \ 1 \ \frac{n_{l}}{2} \left(\frac{n_{l}}{2}+1\right)$$

$$(15)$$

The site index (j) is omitted in the above equation as it is on one particular site. Within the average electric field ansatz, i.e., for $n_l(j) = n_l \Rightarrow h_E(j) = h_E$ for all sites j, resulting,

$$H_E^{(\text{approx})} = \frac{g^2 a}{2} N h_E^0 \tag{16}$$

where, N is total number of staggered sites on the lattice and $h_E^0 = \frac{n_l}{2} \left(\frac{n_l}{2} + 1\right)$. Note that, for $n_l \gg 0$, one can actually consider $h_E^0 = h_E \equiv \frac{n_l^2}{4}$. At any site j, the onsite electric energy $h_E(j)$ differs from h_E^0 if $n_i(j) = 0$, $n_o(j) = 1$, and that difference, that is relevant in strong coupling regime (for $n_l > 0$) is given by:

$$\Delta h_E = \frac{n_l + 1}{2} \left(\frac{n_l + 1}{2} + 1 \right) - h_E^0 = \frac{n_l}{2} + \frac{3}{4}.$$
 (17)

This correction term to $H_E^{(\text{approx})}$ is particularly important for strong as well as intermediate coupling regime, where we consider mean value of gauge flux, that is not very large compared to that considered in weak coupling regime. Within the mean field ansatz the total electric part of the LSH Hamiltonian Hamiltonian is given by:

$$H_E^{(\text{LSH})} = \frac{g^2 a}{2} \left[Nh_E^0 + \sum_{\{j'\}} \left(\frac{n_l}{2} + \frac{3}{4} \right) \right]$$
(18)

where, $\{j'\}$ denotes the sites with fermionic configuration $n_i(j') = 0, n_o(j') = 1$. In the bulk limit of the lattice, the occurrence of j' will be N/4 for N site lattice. Hence, the total mean field electric Hamiltonian in the bulk limit is given by:

$$H_E^{(\text{mLSH})} = \frac{g^2 a}{2} \left[N \frac{n_l}{2} \left(\frac{n_l}{2} + 1 \right) + \frac{N}{4} \left(\frac{n_l}{2} + \frac{3}{4} \right) \right]$$
(19)

Mass Hamiltonian: The mass term (A19), being independent of gauge field configuration remain the same in the mean field ansatz, also for both the strong and weak coupling regime.

$$H_M^{(\text{approx})} = m \sum_j (-1)^j (\hat{n}_i(j) + \hat{n}_o(j))$$
(20)

Interaction Hamiltonian: The matter-gauge field interaction term is the most complicated within LSH framework as detailed in (A20). In the strong coupling limit of the theory, this particular term gives small contribution to the Hamiltonian (see subsection II B) and can be treated perturbatively. However, in the weak coupling regime, this term becomes significant. The purpose of the present approximation scheme is to bring the interaction Hamiltonian into simple form, yet describing matter gauge dynamics in the weak coupling regime.

The approximation scheme that we follow is replacing the local loop quantum numbers $n_l(j)$ by a constant $n_l \gg 0$ at all of the lattice sites. The interaction Hamiltonian given in (A20) can be written as,

$$H_I^{\rm LSH} = \frac{1}{2a} \sum_{j=0}^{N-2} h_I(j, j+1)$$
(21)

where,

$$h_I(j, j+1) = h_I^1(j, j+1) + h_I^2(j, j+1) + h_I^3(j, j+1) + h_I^4(j, j+1)$$
(22)

Each of these terms, can be further decoupled into left (L) and right (R) parts located at site j and site j + 1

respectively,

$$h_{I}^{[s]}(j, j+1) = h_{I}^{[s]}(L)h_{I}^{[s]}(R)$$
, $[s] = 1, 2, 3, 4.$ (23)

Now, considering each term separately, one would obtain the following:

$$h_I^{[1]}(L) = \frac{1}{\sqrt{\hat{n}_l + \hat{n}_o(j)(1 - \hat{n}_i(j)) + 1}} \hat{\chi}_o^+(\lambda^+)^{\hat{n}_i(j)} \sqrt{\hat{n}_l + 2 - \hat{n}_i(j)} = \hat{\chi}_o^+(\lambda^+)^{\hat{n}_i(j)} \hat{C}_1(L)$$
(24)

$$h_{I}^{[2]}(L) = \frac{1}{\sqrt{\hat{n}_{l} + \hat{n}_{o}(j)(1 - \hat{n}_{i}(j)) + 1}} \hat{\chi}_{o}^{-}(\lambda^{-})^{\hat{n}_{i}(j)} \sqrt{\hat{n}_{l} + 2(1 - \hat{n}_{i}(j))} = \hat{\chi}_{o}^{-}(\lambda^{-})^{\hat{n}_{i}(j)} \hat{C}_{2}(L)$$
(25)

$$h_{I}^{[3]}(L) = \frac{1}{\sqrt{\hat{n}_{l} + \hat{n}_{o}(j)(1 - \hat{n}_{i}(j)) + 1}} \hat{\chi}_{i}^{+}(\lambda^{-})^{1 - \hat{n}_{o}(j)} \sqrt{\hat{n}_{l} + 2\hat{n}_{o}(j)} = \hat{\chi}_{i}^{+}(\lambda^{-})^{1 - \hat{n}_{o}(j)} \hat{C}_{3}(L)$$
(26)

$$h_{I}^{[4]}(L) = \frac{1}{\sqrt{\hat{n}_{l} + \hat{n}_{o}(j)(1 - \hat{n}_{i}(j)) + 1}} \hat{\chi}_{i}^{-}(\lambda^{+})^{1 - \hat{n}_{o}(j)} \sqrt{\hat{n}_{l} + 1 + \hat{n}_{o}(j))} = \hat{\chi}_{i}^{-}(\lambda^{+})^{1 - \hat{n}_{o}(j)} \hat{C}_{4}(L)$$
(27)

and

$$h_{I}^{[1]}(R) = \hat{\chi}_{o}^{-}(\lambda^{+})^{1-\hat{n}_{i}(j+1)} \frac{\sqrt{\hat{n}_{l}+1+\hat{n}_{i}(j+1)}}{\sqrt{\hat{n}_{l}+\hat{n}_{i}(j+1)(1-\hat{n}_{o}(j+1))+1}} = \hat{\chi}_{o}^{-}(\lambda^{+})^{1-\hat{n}_{i}(j+1)}\hat{C}_{1}(R)$$
(28)

$$h_{I}^{[2]}(R) = \hat{\chi}_{o}^{+}(\lambda^{-})^{1-\hat{n}_{i}(j+1)} \frac{\sqrt{\hat{n}_{l} + 2\hat{n}_{i}}}{\sqrt{\hat{n}_{l} + \hat{n}_{i}(j+1)(1-\hat{n}_{o}(j+1)) + 1}} = \hat{\chi}_{o}^{+}(\lambda^{-})^{1-\hat{n}_{i}(j+1)}\hat{C}_{2}(R)$$
(29)

$$h_{I}^{[3]}(R) = \hat{\chi}_{i}^{-}(\lambda^{-})^{\hat{n}_{o}(j+1)} \frac{\sqrt{\hat{n}_{l} + 2(1 - \hat{n}_{o}(j+1))}}{\sqrt{\hat{n}_{l} + \hat{n}_{i}(j+1)(1 - \hat{n}_{o}(j+1)) + 1}} = \hat{\chi}_{i}^{-}(\lambda^{-})^{\hat{n}_{o}(j+1)}\hat{C}_{3}(R)$$
(30)

$$h_{I}^{[4]}(R) = \hat{\chi}_{i}^{+}(\lambda^{+})^{\hat{n}_{o}(j+1)} \frac{\sqrt{\hat{n}_{l} + 2 - \hat{n}_{o}(j+1)}}{\sqrt{\hat{n}_{l} + \hat{n}_{i}(j+1)(1 - \hat{n}_{o}(j+1)) + 1}} = \hat{\chi}_{i}^{+}(\lambda^{+})^{\hat{n}_{o}(j+1)}\hat{C}_{4}(R)$$
(31)

The only approximation made in the above set of equations is $n_l(j), n_l(j+1) \rightarrow n_l$, where n_l is the mean

field value. The explicit operator form of the coefficients $\hat{C}_{[s]}(L/R)$'s are the following:

n	i n	n_o	$\hat{C}_1(L)$	$\hat{C}_2(L)$	$\hat{C}_3(L)$	$\hat{C}_4(L)$	$\hat{C}_1(R)$	$\hat{C}_2(R)$	$\hat{C}_3(R)$	$\hat{C}_4(R)$
C		0	1	1	1	$\sqrt{\frac{n_l+1}{n_l+2}}$	1	$\sqrt{\frac{n_l}{n_l+1}}$	$\sqrt{\frac{n_l+2}{n_l+1}}$	$\sqrt{\frac{n_l+2}{n_l+1}}$
C		1	$\sqrt{\frac{n_l+2}{n_l+1}}$	$\sqrt{\frac{n_l+2}{n_l+1}}$	$\sqrt{\frac{n_l+2}{n_l+1}}$	$\sqrt{\frac{n_l+2}{n_l+1}}$	1	$\sqrt{\frac{n_l}{n_l+1}}$	$\sqrt{\frac{n_l}{n_l+1}}$	1
1	. (0	$\sqrt{\frac{n_l+1}{n_l+2}}$	1	1	$\sqrt{\frac{n_l+1}{n_l+2}}$	1	1	1	$\sqrt{\frac{n_l+1}{n_l+2}}$
1		1	$\sqrt{\frac{n_l+1}{n_l+2}}$	1	1	1	$\sqrt{\frac{n_l+2}{n_l+1}}$	$\sqrt{\frac{n_l+2}{n_l+1}}$	$\sqrt{\frac{n_l}{n_l+1}}$	1

(32)

It is clear from the above set of coefficients that in the

limit $n_l \gg 0$, all of the coefficients can be approximated

to be equal to identity operators, that is their leading order contribution. One can expand the coefficients and add corrections order by order. However, for this work, we confine ourselves to the leading order contribution only.

In this regime we also approximate λ^{\pm} as identity operator as per the approximation, $n_l + 1 \approx n_l$. Hence, the approximated interaction Hamiltonian is given by,

$$H_{I}^{(\text{approx})} = \frac{1}{2a} \sum_{j} \left[\chi_{o}^{+}(j)\chi_{o}^{-}(j+1) + \chi_{o}^{-}(j)\chi_{o}^{+}(j+1) + \chi_{i}^{+}(j)\chi_{i}^{-}(j+1) + \chi_{i}^{-}(j)\chi_{i}^{+}(j+1) \right]$$
(33)

E. The simulated Hamiltonian

In summary, we obtain the following approximate Hamiltonian that acting on the LSH states on the 1d spatial lattice with the boundary flux $l_i \gg 0$ would result the exact dynamics of the full gauge theory ²:

$$H_E^{(\text{approx})} = \frac{g^2 a}{2} \sum_j \left[\frac{\hat{n}_l}{2} \frac{\hat{n}_l}{2} \right]$$
(34)

$$H_M^{(\text{approx})} = m \sum_{j} (-1)^j (\hat{n}_i(j) + \hat{n}_o(j))$$
(35)

$$H_{I}^{(\text{approx})} = \frac{1}{2a} \sum_{j} \left[\hat{\chi}_{o}^{+}(j) \hat{\chi}_{o}^{-}(j+1) + \hat{\chi}_{o}^{-}(j) \hat{\chi}_{o}^{+}(j+1) + \hat{\chi}_{i}^{+}(j) \hat{\chi}_{i}^{-}(j+1) + \hat{\chi}_{i}^{-}(j) \hat{\chi}_{i}^{+}(j+1) \right]$$
(36)

We present the details of an atomic quantum simulation scheme to simulate this mean-field Hamiltonian in next section.

III. THE SIMULATING HAMILTONIAN

In this section, we construct an atomic Hamiltonian that can successfully simulate (34 - 36). We describe the relevant quantum system (subsection III A), and its connection with the LSH Hamiltonian is established (subsection III B). That the model is well-suited for both the weak coupling and strong coupling limits of gauge theory is demonstrated in subsection III C.

A. Atomic Hamiltonian : Hubbard Model on a Bipartite Lattice

We consider a Fermi-Hubbard model in a onedimensional lattice. The optical potential is of the form

$$V(x) = -V_L \cos^2(kx) \tag{37}$$

The minimum of each well corresponds to the physical lattice site with site index j. V(x) can also act as the trapping potential. Here $V_L > 0$, and $k = \pi/d$, d being the lattice periodicity. Next, energy offsets of V' and -V' are added to the odd and even sites respectively. So, a bipartite lattice is created, and effectively the lattice spacing is 2d now.

The fermionic atoms can belong to either of its two accessible hyperfine states : we denote them by the symbols $|\uparrow\rangle$ and $|\downarrow\rangle$ respectively. Let $c_{\uparrow}(j)$ be the fermionic annihilation operator for spin index \uparrow and $c_{\downarrow}(j)$ be the fermionic annihilation operator for spin index \downarrow , both for site j. The corresponding number operators are $\mathcal{N}_{j\uparrow}$ and $\mathcal{N}_{j\downarrow}$. The total number of fermions at site j is given by $\mathcal{N}(j) = \mathcal{N}_{\uparrow}(j) + \mathcal{N}_{\downarrow}(j)$.

The lattice depth V_L can be written as $(V_1 - V_0)$. It is to be noted that V_1 and V_0 are theoretical parameters only, and will be used to make a connection between the atomic Hamiltonian and the LSH Hamiltonian. From an experimental perspective, it is only V_L that is of importance.

The Hamiltonian can be written as :

$$H = H_{\text{hopping}} + H_{\text{int}} + H_{V_0} + H_{V'} \tag{38}$$

where, in terms of the tight-binding parameters,

$$H_{\text{hopping}} = -\sum_{j} t_j \left(c^{\dagger}_{\uparrow}(j) c_{\uparrow}(j+1) + c^{\dagger}_{\downarrow}(j) c_{\downarrow}(j+1) \right) + \text{h.c}$$
(39)

$$H_{\rm int} = u \sum_{j} \mathcal{N}_{\uparrow}(j) \mathcal{N}_{\downarrow}(j). \tag{40}$$

$$H_{V_0} = \sum_j V_0 \mathcal{N}(j) \tag{41}$$

and

:

$$H_{V'} = V' \sum_{j = \text{odd}} \mathcal{N}(j) - V' \sum_{j = \text{even}} \mathcal{N}(j) \qquad (42)$$

Here we have dropped a term $H_{V_1} = \sum_j V_1 \mathcal{N}(j)$ from the Hamiltonian, as this provides just a constant energy shift. The details are there in appendix B.

If the hopping -t is a constant throughout the lattice, this model essentially is a 1D Hubbard model with alternating potential, often termed as the "ionic Hubbard model", defined on a bipartite lattice. Here, in addition to a site-independent hopping -t and the on-site interaction u, there is a difference in the energy offset 2V' between sublattice **A** and sublattice **B**. This model was originally proposed to study transitions in organic crystals [59], and later, found application in the studies

 $^{^2}$ In this limit, the Abelian Gauss law constraint (A1) is automatically satisfied as (A15) and (A16) effectively become equal.



FIG. 1. Structure of the two-color lattice

of ferroelectric transitions [60]. In the recent past, this model has been experimentally realized [61] in a system of ultracold atoms. So we consider this to be a very suitable candidate to simulate lattice gauge theories.

At half-filling, the ionic Hubbard model is capable of describing a band-insulator [62]. However, this model has a rich phase diagram, and at higher inter-atomic interaction strengths, can support transitions to different states, including Mott insulator [62], correlated insulator [63, 64], AFM insulator and half-metal [64] phases; and certain combinations of u and V' can even lead to superfluidity [65]. As we will see in the later part of this work, we have to carefully choose our parameters such that the entire dynamics remains confined to a single paramagnetic phase in order to mimic the dynamics of gauge theory.

B. Mapping the parameters

We are now in a position to compare the weak coupling LSH Hamiltonian and the atomic Hamiltonian. For a particular site j, we make the following identification :

$$n_i(j) = \mathcal{N}_{\uparrow}(j); \qquad n_o(j) = \mathcal{N}_{\downarrow}(j)$$
 (43)

$$\chi_i^+(j) = c_{\uparrow}^{\mathsf{T}}(j); \qquad \chi_i^-(j) = c_{\uparrow}(j) \tag{44}$$

$$\chi_{2}^{+}(j) = c_{1}^{\dagger}(j); \qquad \chi_{2}^{-}(j) = c_{1}(j)$$
(45)

$$m = V' \tag{46}$$

Also, the magnitude of V_0 has to be chosen to be mapped to electric part of the gauge theory Hamiltonian for a particular n_l , fixed by the open boundary condition.

The electric term of approximated LSH Hamiltonian is mapped to:

$$H_E^{(\text{approx})} \to \sum_j V_0 \mathcal{N}(j).$$
 (47)

Similarly, the potential V' is fixed by the mapping:

$$H_M^{(\text{approx})} \to V' \sum_{j=\text{odd}} \mathcal{N}(j) - V' \sum_{j=\text{even}} \mathcal{N}(j)$$
 (48)

and the hopping terms are identically related as,

$$H_{I}^{(\text{approx})} \to -t \sum_{j} \left(c_{\uparrow}^{\dagger}(j) c_{\uparrow}(j+1) + c_{\downarrow}^{\dagger}(j) c_{\downarrow}(j+1) \right)$$

+h.c. (49)

Note that, there is no term in the weak coupling LSH Hamiltonian that corresponds to the on site interaction term (B6). So, in the limit $u \to 0$, one would have a complete mapping between atomic system and weak coupling limit of gauge theory.

We would like to point out that although the LSH Hamiltonian contains explicit bosonic modes $n_l(j)$, these are actually non-dynamical in the weak coupling approximation as discussed before, and hence we do not keep actual bosons in the atomic system. Instead, we incorporate the effect of these bosons in the potential itself, in the form of a constant energy shift. This enables us to (i) keep n_l uniform for each site, and (ii) ensure that the bosonic and the fermionic modes are completely decoupled: as there remains no chance of any boson-fermion scattering.

C. Simulating both weak and strong coupling regimes of gauge theory

We now present the novel features of this proposed scheme that enable one to simulate accurate dynamics of the gauge theory beyond the weak coupling limit emphasized so far.

It is important to note that this particular scheme assumes mean field contribution from the loop degrees of freedom, while the two fermionic degrees of freedom describe the dynamics of the theory. The configurations with large incoming flux l_i correspond to low energy states in the weak coupling regime. On the other hand, the strong coupling vacuum is defined to be the state with zero electric flux. This particular scheme calls for choosing a non-zero value of the incoming flux $l_i \geq N$ for a *N*-site staggered lattice, such that all of the fermionic configurations correspond to physical LSH configurations [48]. Physically this amounts to a constant shift in the vacuum energy level as compared to the strong coupling vacuum of the theory. We propose that different sectors of the gauge theory can be simulated by choosing appropriate values of l_i ($l_i >> 0$ for weak coupling theory, $l_i \approx N$ for strong coupling theory).

Moving away from the weak coupling approximation, the electric part of the Hamiltonian becomes dominant. This part (2) measures the electric flux contribution of each individual link. Within the LSH framework, it has some contribution from the local loop quantum number n_l as well as from the local string quantum numbers n_i , n_o at the site from where the link starts as given in (A18). One can also express the total electric contribution as function of local n_l depending on the local fermionic configurations as listed in (15). Thus, the complete on-site contribution to the electric part of the Hamiltonian can divided into two categories :

- (a) for three of the four allowed fermionic configurations at any site (15), the approximated contribution is a function of n_l only (16).
- (b) For the other fermionic configuration, the electric contribution to the Hamiltonian has an additional correction term (17). In our scheme, this correction term (17) is added to (16) for N/4 sites (as the probability of obtaining one such particular arrangement of fermions is 1/4). Therefore, the total correction in the electric Hamiltonian is:

$$\Delta H_E^{(\text{LSH})} = \frac{g^2 a}{2} \frac{N}{4} \left(\frac{n_l}{2} + \frac{3}{4}\right) \tag{50}$$

The on-site interaction in the atomic Hamiltonian, which does not have an equivalent in the approximate LSH Hamiltonian, can be tuned to recover the exact contribution of (50). For a Hubbard model at half-filling, all the four accessible states $|0\rangle$, $|\uparrow\rangle$, $|\downarrow\rangle$ and $|\uparrow\downarrow\rangle$ are equally likely as long as the system remains in the paramagnetic phase. So,

$$N_{|\downarrow\rangle} \approx \frac{N}{4} \& N_{|\uparrow\downarrow\rangle} \approx \frac{N}{4}$$
 (51)

Here $N_{|\downarrow\rangle}$ denotes the number of sites belonging to state $|\downarrow\rangle$, and $N_{|\uparrow\downarrow\rangle}$ is the number of sites with doublons. N is the total number of lattice sites. It is the doublon configurations that contribute to (B6). Hence, one can

utilize the on-site interaction term to recover the exact correction term,

$$u\sum_{j}\mathcal{N}_{\uparrow}(j)\mathcal{N}_{\downarrow}(j) = u\frac{N}{4} \Rightarrow u \equiv \frac{g^{2}a}{2}\left(\frac{n_{l}}{2} + \frac{3}{4}\right)(52)$$

in order to match it with (50).

Hence, for the bulk limit of the lattice (N >> 0), we have the maximum overlap of the (mean field approximated) strong coupling lattice gauge theory to Fermi-Hubbard Hamiltonian as Then

$$H_E^{(\mathrm{mLSH})} \longrightarrow H_{V_0} + H_{\mathrm{int}}$$
 (53)

$$H_M^{(\text{LSH})} \longrightarrow H_{V'}$$
 (54)

$$H_I^{(\text{approx})} \longrightarrow H_{\text{hopping}}$$
 (55)

provided we fix V_0 and u such that the system, staying in the desired phase, mimics the dynamics of gauge theory as shown in Fig. 2.

The correction to the approximate interaction Hamiltonian is negligible in weak coupling regime, and also insignificant in the strong coupling limit. In this work, we do not consider any correction to the interaction term.

We now, explicitly calculate the parameters of the atomic Hamiltonian, that has to be tuned in the experiment to simulate desired gauge theory dynamics in different coupling regime.

1. Weak coupling regime of gauge theory: u/t = 0

In this particular regime we consider the scaled Hamiltonian given in (8). Within the approximation scheme done for the equivalent LSH Hamiltonian for a finite lattice given in (34), we consider the bosonic loop quantum number to take the average value

$$n_l \approx \mathcal{O}(10^p) \Rightarrow \tilde{H}_E^{(\text{approx})} \approx \mathcal{O}(10^{2p}).$$
 (56)

For a comparative mass and interaction contribution of the Hamiltonian, i.e.

$$\tilde{H}_M^{LSH} \approx \mathcal{O}(10^{2p}) \tag{57}$$

$$\& \tilde{H}_I^{(\text{approx})} \approx \mathcal{O}(10^{2(p+p')}), \qquad (58)$$

is obtained for the following scaling of the parameters:

$$\frac{m}{g} \approx \mathcal{O}(10^{p-p'}) \quad \sqrt{x_0} \approx \mathcal{O}(10^{p+p'}) \quad , \forall p' \in \mathbb{Z}_+.$$
(59)

The exact values of the dimensionless parameters of gauge theory can be taken as:

$$n_l = \tilde{n}_l \times 10^p \tag{60}$$

$$\mu_0 = \tilde{\mu}_0 \times 10^{2p} \tag{61}$$

$$x_0 = \tilde{x}_0 \times 10^{2(p+p')}, \quad \forall \text{ integer } p, p' \tag{62}$$

$$= 10^{2p}$$
 for the choice, $p' = 0, \tilde{x}_0 = 1.$ (63)



FIG. 2. Cartoon representation of dynamics of 1d ionic Hubbard model mimicking that of SU(2) lattice gauge theory in one spatial dimension. (a) Initial state: fully filled odd sites and empty even sites mimicking the strong coupling vacuum consisting of no particles ($n_i = 0, n_o = 0$ on even sites) and no antiparticles ($n_i = 1, n_o = 1$ on odd sites). Under Hamiltonian evolution, one atom hops from an odd site to neighboring even site in Hubbard model, that mimics creation of a particle antiparticle pair at two neighboring staggered site of the gauge theory, connected by one unit of flux to form a gauge singlet string configuration. One further hopping as shown in the figure mimics the dynamics in gauge theory as elongation of the string and creation a baryon on one site. In these three states, the total number of particle (antiparticle) for the gauge theory are respectively 0, 1, 2. (b) Ionic Hubbard model dynamics is mimicking string breaking dynamics of gauge theory. Starting from a string of length 3 unit, pair production occurs and the initial string breaks into two smaller strings.

Now, the dynamics of this scaled Hamiltonian \hat{H} in (8), is to be simulated by the simulating Fermi-Hubbard Hamiltonian given in (38) in the time scale $\tilde{\tau}$ as defined in (9), such that

$$\exp(-i\tilde{H}\tilde{\tau}) \longrightarrow \exp(-iH\tau) \tag{64}$$

where, H is the atomic Hamiltonian given in (38) with the parameters:

$$V' = \tilde{\mu}_0 \tag{65}$$

$$V_0 = \frac{1}{4} \left(\tilde{n}_l^2 + 2\tilde{n}_l \right) \tag{66}$$

$$u = 0 \tag{67}$$

$$t = -1. \tag{68}$$

Here, all the parameters are fixed in units of 't'. The only choice that we have made in setting the parameters is p' = 0 in (62). Gauge theory with a nonzero p' can be equivalently simulated by the same atomic system with tuning V' to smaller values $\tilde{\mu}_0 \times 10^{-2p'}$ in an experiment. This will access all mass values of gauge theory in the quantum simulation protocol.

2. Strong coupling regime of gauge theory: |u/t| > 1

We consider the scaled Hamiltonian in (11) in strong coupling regime $x_0 < 1$. As discussed earlier, the bulk limit of the Fermi-Hubbard Hamiltonian in the paramagnetic phase will correspond to the exact mean field electric term (19) and mass term (A19). Although the interaction term is approximated, will not make a major difference in spectrum and/or dynamics as $x_0 \rightarrow 0$ as it is less dominant compared to diagonal terms. Likewise, in the weak coupling regime, we fix the boundary condition l_i to be a fixed integer, but is of $\mathcal{O}(1)$. We map the gauge theory Hamiltonian to Fermi-Hubbard Hamiltonian with parameters

$$V' = 2\frac{1}{\sqrt{x_0}} \cdot \frac{m}{g} \tag{69}$$

$$V_0 = \frac{1}{x_0} \cdot \frac{l_i}{4} \left(l_i + 2 \right) \tag{70}$$

$$u = \frac{1}{x_0} \cdot \frac{1}{4} \left(2l_i + 3 \right) \tag{71}$$

$$t = -1. \tag{72}$$

Here also, all the parameters are fixed in units of 't'. It is clear from the above relations that for a fixed value of l_i , smaller values of x_0 require larger V_0/t and u/t for the atomic system. However, we will have to be careful to remain in the same paramagnetic phase such that our analysis of compensating errors in electric Hamiltonian from the uniform potential are well compensated by the self interaction term. For this purpose, i.e., in order to keep u/tbelow the critical point for paramagnetic-ferromagnetic phase transition one can not really expect to simulate $x_0 \rightarrow 0$ under the present scheme. However, one can simulate $x_0 < 1$ as well as $x_0 = 1$ besides accurately simulating intermediate coupling range $x_0 \approx 10-100$ as will be demonstrated in the numerical analysis.

Likewise the weak coupling case, the simulating and

simulated dynamics are comparable up to a factor

$$\tau_{\text{atomic}} = 2a \times \tau_{\text{gauge}}.$$
 (73)

where, a is small but finite in strong coupling limit.

In the next section, we propose the precise experimental set-up that is close to already performed experiments for Ionic-Hubbard model following the above mentioned scheme, where strong coupling regime of lattice gauge theory dynamics is mapped to ionic Hubbard model with u/t > 1, whereas the weak coupling regime is mapped to the same with $u/t \approx 0$.

IV. EXPERIMENTAL REALIZATION

The experimental scheme calls for the realization of 1-dimensional Fermi-Hubbard model in a bipartite lattice. In the recent past, the ionic Fermi-Hubbard model was experimentally realized in a honeycomb lattice [61]. and its bosonic counterpart was implemented on a bipartite chequerboard lattice [66]. Also, a 1-dimensional Fermi-Hubbard model was implemented in a experiment by Scherg et al. [67]. Both [61] and [67] used a degenerate gas of fermionic 40 K of numbers $\approx 10^5$ and 10^4 respectively. We propose that a combination of these two methods can successfully yield a 1-dimensional Hubbard model with alternating lattice potentials. The set-up, which is very much realizable with current experimental techniques, is described in IVA. The procedures for preparing the initial states and observing the final dynamics are outlined in IVB and IVC respectively. We discuss the possible sources of errors (that can affect the accuracy of the results in this experiment) in IVD.

A. Proposed set-up

The interference pattern of two counter propagating lasers is used to create an optical lattice. The lattice depth is proportional to the intensity of the laser beam and is measured in units of the recoil energy E_R .

In the experiment by Messer et al.[61], first a regular honeycomb lattice was created, and that fixed the hopping parameter t on each bond. Next a staggered energy offset of Δ was independently applied between sites of **A** and **B** sublattices. In our 1-dimensional structure, an equivalent would be to set up the primary lattice with lattice depth V_L : This fixes the hopping parameter t. Then, on top of it, energy offsets V' and -V' can be independently applied on the odd sites and even sites respectively, so that the lattice depth is $(V_0 + V')$ for odd sites, and $(V_0 - V')$ for even sites.

Just like the hopping t, the on-site interaction u, too depends on the lattice depth. However, u can be independently controlled as well, by means of Feshbach resonance. As for the two fermionic states, any two hyperfine states of a particular atom can be employed. In [67], the hyperfine states

$$|\uparrow\rangle = |F = -9/2; m_{\rm F} = -9/2\rangle \\ |\downarrow\rangle = |F = -9/2; m_{\rm F} = -7/2\rangle$$

of ultracold $^{40}{\rm K}$ atoms were used. In [61], in addition to the above, the combination

$$|\uparrow\rangle = |F = -9/2; m_{\rm F} = -9/2\rangle$$
$$|\downarrow\rangle = |F = -9/2; m_{\rm F} = -5/2\rangle$$

was also employed in order to obtain desired range of u.

In [61], the ionic Hubbard model was studied on a honeycomb lattice. In contrast, our model requires the implementation of the ionic Hubbard model in a simple one-dimensional geometry. Regarding the dimensionality of the system, it may be recalled that in the recent past, ultracold atom experiments have successfully confined bosonic and fermionic atoms to one dimension (1D). The basic idea is to tightly confine the particles in two transverse directions, and make them weakly confined in the axial direction. Thus, their motion in the transverse directions are completely frozen. So effectively, these are quasi-1D systems.

For example, in our proposed set up, suppose both V_y and V_z , the potentials in the transverse directions, are kept fixed at a large value (Like, $33E_R$ as in [68], or $42E_R$ as in [69]). V(x), The lattice depth in the axial direction is kept in a range of $5E_R - 12E_R$. We note that in Hubbard model experiments, the potentials are to be deep enough ($V \ge 5E_R$) so that the single-band description of Hubbard model remains valid. On the other hand, V(x) cannot be as deep as the potentials in the transverse direction, so as to restrict the dynamics in 1-dimension only. The hopping parameter t is a function of the lattice depth, and can be estimated using the Wannier functions [70].

In III A, the lattice depth V_L was written as $(V_1 - V_0)$, so different combinations of V_1 and V_0 can result in the same lattice depth. This offers a tremendous advantage in the experimental pursuit, as the same optical lattice can be assumed to be split in different pairs of V_1 and V_0 : allowing one to explore a wide range of V_0 values (that, in turn, enables one to access a wide range of x_0 and/or l_i as per (70)). It is to be noted that both V_1 and V_0 are theoretical parameters in the model that leads to constant shifts in the energy only : bearing no effect on the dynamics of the fermions.

Accordingly, we consider two configurations :

- (i) $V_1 = 6E_R$ and $V_0 = 0.5E_R$ and
- (ii) $V_1 = 6.5E_R$ and $V_0 = 1E_R$.

In both the cases, the resultant uniform lattice depth V_L is $5.5E_R$ for all the sites. This results in a hopping $t = 0.057E_R$. The combinations we have mentioned translate to

(i)
$$V_0 \approx 8.75t$$
 and (ii) $V_0 \approx 17.5t$

respectively. In addition, an offset of V' and -V' is independently applied on the odd and even sites. In our scheme, we choose V' = 1.6t and stick to this value in all our numerical simulations. The on-site interaction u can be controlled by applying a Feshbach field.

To simulate the weak coupling limit, we restrict ourselves to the weakly interacting atomic limit : u/t << 1, and choose u = 0.1t. On the other hand, simulation of the strong coupling limit calls for the realization of the strongly interacting atomic limit : $u/t \gg 1$, and we choose $u \approx 5.5t$. We note that these V'/t and u/t values comfortably fall in the parameter regimes accessed in recent experiments [61, 67, 71].

B. Initial state preparation:

The initial state has to be prepared in a Chargedensity-wave (CDW) configuration where all the odd sites are occupied by the fermionic particles and the even sites are completely empty. This can be done using some sort of filtering sequence in the experiment. For example, in [71, 72], this was achieved by superposing the primary lattice (with wavelength λ) with an additional long lattice (with wavelength 2λ) in the following way:

$$V(x) = -V_L \cos^2(kx) - \tilde{V}(\cos^2(kx/2 + \phi))$$
(74)

with $k = 2\pi/\lambda$.

The lattice depths V_L and \tilde{V} and the relative phase ϕ can be adjusted independently. Here V_L stands for the depth of the original (and short) lattice; and V is the depth of the additional long lattice. This long lattice is utilized during the preparation of the initial CDW state. Initially, the long lattice is made quite deep (like, $20E_B$, as in [71]), and the short lattice is ramped up to that depth at a non-zero relative phase ϕ to create a tilted lattice of double wells. Now it is so arranged that the odd sites host lower energy wells than the even sites, and it is possible to load all atoms in the odd sites only. The tilt offsets are made sufficiently large so that the particles cannot escape from the odd sites and tunnel to the even sites. After loading all the atoms, the longer lattice is switched off, and the short lattice is ramped down to its desired final value (In our case, $5.5E_R$). The offset V' and -V' is added to the odd sites and even sites respectively, to create the bipartite structure. Now tunneling is possible between adjacent sites, and the dynamics begins.

Moreover, (52) is only valid if $N_{|\downarrow>}=N_{|\uparrow\downarrow>}=N/4$ as in 51. Since here the system is initially prepared in a CDW state containing doublons only, this will not hold true in a short initial time-span, and the interactomic interaction u will overcompensate the correction term in the electric Hamiltonian. Thus, the dynamics from the atomic Hamiltonian will have a departure from that of the full Gauge-theory. However, with time, more and more atoms would hop to the adjacent sites and the distribution would get more even across the sites, resulting in $N_{|\downarrow\rangle} \approx N_{|\uparrow\downarrow\rangle}$. So soon enough, the quantum simulation becomes more accurate, as demonstrated in the numerical results in Section V.

C. Observing the dynamics

The observable can be defined as the population imbalance P between the even sites and odd sites, defined as

$$P = \frac{N_e - N_o}{N_e + N_o}.\tag{75}$$

Here N_e is the total number of atoms in the even sites, and N_o is the total number of atoms in the odd sites.

The time evolution of the parameter P is to be studied in order to visualize the particle number dynamics of gauge theory. A site-resolved technique is thus needed to determine the number of atoms on even and odd lattice sites separately. In [71, 72], a band-mapping scheme was successfully employed using the long lattice. Once the desired time evolution in the primary (short) lattice is over, the long lattice is introduced again to create the tilted lattice, and tunneling stops. The phase ϕ is chosen such a way that the odd sites constitute the lower wells in the array of double wells. Now the population distribution across the odd and even sites gets sealed. Next, the depth of the long lattice is ramped to a much higher value : and the atoms in the even sites get transferred to the third Bloch band of the superlattice. Atoms in the odd sites remain in the first band. The density profile in the different bands can be obtained using Time-of-flight (TOF) images and absorption imaging [71].

D. Possible Source of experimental errors

The particle-antiparticle pair creation and string breaking in gauge theory are mimicked by the ionic Hubbard model dynamics, and the relevant observable at any instant is P, the averaged population imbalance between even sites and odd sites. Thus, individual site-resolved occupancy data is not required. If the lattice is sufficiently long and if the averaging is done properly (like, in [67], each data point was averaged over four measurements), it is possible to obtain a very accurate value of P. The lattice potential and the sub-lattice offset, too, can be precisely monitored [61],[66].

The only major source of possible experimental errors can be the imperfection in the initial state preparation. A CDW state can certainly be realized where only the odd sites are occupied, but then those sites can host zero, single or double occupancy as in [71]. To ensure that all the odd sites have doublons, one needs to monitor the number of atoms precisely. As reported in [61], there is always a systematic uncertainty of 10% in the preparation. However, as demonstrated in the next section, the dynamics is very little affected even with this error margin.

V. SIMULATED DYNAMICS AND OBSERVABLES

In this section, we present numerical analysis of our proposal. The comparison between the spectrum of the simulating and simulated systems is presented in V A, and the corresponding dynamics is presented in V B. In V C, we demonstrate that even if there is an initial error in preparing the system in a perfect CDW state, the dynamics is not affected much; and the departure from the expected result remains well within an acceptable window of tolerance.

A. Spectrum comparison:

In weak coupling regime: We aim to quantum simulate gauge theory Hamiltonian, with the values of dimensionless parameters given by:

$$x_0 = 10^{-10}$$
 & $m/g = 1.6 \times 10^{-10}$

acting on the LSH Hilbert space characterized by

$$n_l = 5 \times 10^5$$

at all sites and correspond to to p = 5, p' = 0 in (60-62). The fermionic (string) configurations remain completely dynamical as n_i, n_o can take all possible values at sites 0, 1, 2, ...2N. Following (65-67) we obtain the parameters of the atomic Hamiltonian to be fixed at:

$$V_0 = 8.75t$$
, $V' = 1.6t$, $u = 0.1t$. (76)

Note that, we have chosen a feasible but small value of the parameter u. Smaller and smaller values of u will enable to mimic the dynamics of gauge theory more accurately as we take $p \gg 1$. We perform exact diagonalization for both the Hamiltonians with a small number of sites, that is doable on a PC. Our scheme, being completely scalable, the agreement in spectrum as in Fig. 3 holds true for any size of lattice as per experimental capabilities.

In strong coupling regime: We aim to quantum simulate gauge theory Hamiltonian (11), with the values of dimensionless parameters given by:

$$x_0 = 0.69 \& m/g = 1.6.$$

This Hamiltonian acts on LSH Hilbert space characterized by $l_i = 6$ as in (13), while n_i, n_o can take all possible values at sites 0, 1, 2, ... 2N. Following (69-71), the mimicking atomic system is defined by parameters:

$$V_0 = 17.5t , V' = 1.6t , u = 5.47t.$$
 (77)

Like the weak coupling case, we also perform exact diagonalization for this case to compare and obtain the spectrum as in Fig. 3. Note that, from our analysis we only expect exact match of spectrum in the $N \to \infty$ limit, and that is beyond the scope of exact diagonalization. However, in Fig. 4, we demonstrate that if the lattice size

keeps increasing, the agreement between the simulating spectrum and the original spectrum gradually improves. Thus, it is expected that in the bulk limit, there would be a perfect agreement. Performing numerical calculations for a longer lattice is beyond the scope of exact diagonalization, and thus, the present work. In principle, it can be carried out using state of the art tensor network techniques, and establish a proper benchmark for the scheme in strong coupling regime. However, tensor network can only calculate a particular (low energy) sector of the theory with the desired accuracy, and quantum simulation is expected to outperform the same.

However, even with limited computational resources, we make the following observations:

- It appears from (69-71) that, by increasing V_0/t in the atomic system, one would be able to access smaller and smaller values of the gauge theory parameter x_0 . However, the consequence is that, in order to mimic exact strong coupling dynamics, u/t has to be increased as well.
- With an increasing u/t (even for a fixed value of V_0/t), gaps are introduced in the atomic spectrum as the atomic system experiences a quantum phase transition (see Fig. 6) and enters the Mott Insulator phase [73]. Then the system can no longer mimic dynamics of gauge theory as there is no such quantum phase transition in the gauge theory spectrum. Hence, this quantum simulation scheme is not suitable for $x_0 \rightarrow 0$.
- Instead, if one can arrange the experimental set-up to fix V_0/t at a smaller value, the atomic system simulates the intermediate coupling regime of the full gauge theory reliably. We illustrate such an agreement for $V_0/t = 0.1$ ($x_0 = 120$) in Fig 3.

B. Simulated Dynamics:

One important dynamical phenomenon to observe in real time dynamics in gauge theory is the dynamics of pair production and string breaking as illustrated via cartoon in Fig. 2. We consider preparing the system in a state in which all even sites are completely empty (no particle) and all odd sites are completely filled (no antiparticle). The real time Hamiltonian evolution of the atomic system involves atoms hopping from one site to another, simulating the event of pair creation and particle number dynamics of gauge theory. Within LSH framework, for the no particle-no antiparticle state $|\Psi_0\rangle$ on a 1d lattice of N staggered sites, we define the following quantity to describe particle density,

$$\rho(\tau) = 1 + \frac{1}{N} \langle \Psi_0 | \hat{\mathcal{U}}^{\dagger}(\tau) \hat{\mathcal{O}} \hat{\mathcal{U}}(\tau) | \Psi_0 \rangle$$
 (78)

where, $\hat{\mathcal{O}} = \sum_{j} \left((-1)^{j} (\hat{n}_{i}(j) + \hat{n}_{o}(j)) \right)$ and $\mathcal{U}(\tau_{\text{gauge}})$ is defined in (10).



FIG. 3. Spectrum of the Ionic Hubbard model, Full SU(2) gauge theory (KS or LSH) Hamiltonian (without any approximation) and the Weak coupling approximated LSH calculated by exact diagonalization for 6 site system and scaled to fit between 0 to 1. (a) The spectrum in weak coupling regime of gauge theory with parameters as per (76) for different values of p as discussed in subsection IIIC1. (b) The spectrum obtained for the string coupling analysis, for $V_0 = 17.5, 8.75 \& 0.1$ respectively. The spectrum demonstrates that the intermediate coupling regime is better accessible by strong coupling analysis if smaller values of V_0 becomes experimentally feasible. We propose to quantum simulate strong coupling spectrum within a mean field approximation and at bulk limit, whereas the plots are only for small lattices and hence showing magnified deviation of the mean field spectrum from that of the full gauge theory. The approximated LSH is only valid in weak coupling regime and matches with full gauge theory for $p \gg 1$.

The simulated dynamics in Hubbard model is measured by the observable P, as defined in (75). Its connection with the particle number dynamics of gauge theory can be obtained by looking at the parameter 1 + P. In Fig. 5 we plot the quantities against a scaled time $\tau = \tau_{\rm atomic} = 2a\tau_{\rm gauge}$ following (9).

As done in the spectrum analysis, we consider the

same parameter values for calculating pair-production and string breaking dynamics as well. From the simulated dynamics we can conclude the following :

• The proposed simulation scheme simulates the dynamics of weak coupling gauge theory perfectly and that is evident even from the numerical analysis



FIG. 4. Spectrum of the simulated Hamiltonian is compared with the spectrum of the full gauge theory Hamiltonian. In this comparative study, the incoming flux for the lattice of size N is taken to be $l_i = N$, and the experimental parameters are adjusted accordingly to simulate the gauge theory with the coupling $x_0 = 1$. The deviation of the Simulated eigenvalues range from (a) (0 - 17)% for N = 2, (b) (0 - 14)% for N = 4, and (c), (d) (0 - 12)% for N = 6, 8. This shows that the simulated dynamics becomes more reliable as the lattice size increases. As explained in the text, in the bulk limit of the lattice, the simulated Hamiltonian would match the original gauge theory Hamiltonian as the natural consequence of statistical distribution of the fermions on the lattice . However, classical simulation is not the perfect tool to predict that statistical result (with at least N 100+) via exact diagonalization of the Hamiltonian as the dimension of the Hilbert space is 2^{2N} .

using a small system. Here the particle density dynamics resulting from i) full gauge theory, ii) the approximated LSH theory and iii) the atomic Hamiltonian all agree very well.

- The difference between the actual dynamics due to the original Hamiltonian and the dynamics due to the approximated Hamiltonian is quite pronounced in the intermediate coupling/ strong coupling regimes. However, by adjusting the on-site interaction parameter, it was possible to recover the correction in the electric energy term (50) substantially, and hence the ionic-Hubbard dynamics is now closer to the dynamics of the full gauge theory, when compared to the same with the approximated LSH formulation.
- The discrepancy that still exists in the intermediate/ strong coupling regimes will surely get reduced if one can simulate using a long enough lattice, such that in the statistical limit, one can really recover the correction in electric energy term (50) in full by choosing the atomic self-interaction accordingly. Considering that we used a small lattice (6 site sys-

tem) for our numerical simulation and yet managed to observe a good agreement, it is extremely likely that in an actual experiment (or, tensor network calculation) involving a large number of lattice sites, the error will be insignificant.

It is discussed in Sec. IV, how one can measure the dynamics in an actual experiment. However, the actual time measured in ms during the experiment is related to the scaled times as:

$$\tau_{\rm exp} = \frac{\hbar \tau_{\rm atomic}}{t} \equiv \frac{\tau_{\rm atomic}}{1.5716} \,\,\mathrm{ms} \tag{79}$$

$$\Rightarrow \equiv \frac{2a\tau_{\text{gauge}}}{1.5716} \text{ms.}$$
(80)

Thus, for different values of lattice spacing, the same experiment would simulate real time dynamics of gauge theory happening in different smaller time scale.

C. Effects of possible experimental errors :

As discussed in IV D, the dominant contribution to the experimental error would come from an imperfection in



FIG. 5. Simulated particle density dynamics, corresponding to the cartoon of Fig. 2(a) is plotted against a scaled time τ . The parameters are identical to that used for spectrum analysis in Fig. 3 for (a) weak coupling regime and (b) strong coupling regime. The simulated dynamics is almost exact to that of the full gauge theory for weak coupling limit. The mismatch between full gauge theory dynamics and Hubbard model dynamics in strong coupling regime is expected to get minimized at bulk limit. The approximated LSH is only valid in weak coupling regime and matches with full gauge theory dynamics than that of the approximated Hamiltonian. This is because the tuned self interaction of atomic Hamiltonian takes care of a significant error that exists in the approximated Hamiltonian.

the initial state preparation. To obtain a rough estimate of the same, we study the simulated dynamics for a 6site lattice. Ideally, all the odd sites should be doubly occupied. We choose two other configurations : (i) 2 odd sites doubly occupied, a singly occupied odd site and a singly occupied even site (an error of ~ 17%) and (ii) 2 odd sites doubly occupied and a doubly occupied even site (an error of ~ 33%). As shown in Fig. 7, the error is large in a very short time-span only (< 0.5 in units of the scaled time τ , while the dynamics was studied in the range 0 to 10 in the same unit). If we exclude this region, the percentage error in the first case is well within the margin of 10% for most of the course of evolution, and its peak value lies at around 20%. In the second case, the percentage error lies well within the margin of 20% for most of the course of evolution, and its peak value



FIG. 6. A Quantum phase transition is observed with the ionic Hubbard model at a particular value of u/t, beyond which the spectrum becomes gapped and hence the Hubbard model can no longer mimic the dynamics of gauge theory. This particular plot is obtained with the parameters of Hubbard model given in (77) except varying u/t. Choosing a larger value of V_0/t corresponds to smaller value of x_0 via (70), but following (71) it will always be in the Mott Insulating phase.

lies slightly above 40%. Thus, one may conclude that as long as the error in the initial state preparation is within a reasonable range (like, in [61], the systematic uncertainty is 10%), the dynamics is not much affected by it.

VI. DISCUSSIONS AND FUTURE DIRECTIONS

This paper presents the very first practically implementable quantum simulation proposal for simulating SU(2) lattice gauge theory in (1 + 1)-d, that specifically simulates the spectrum and dynamics of gauge theory in weak coupling regime as well as intermediate coupling regime for a large lattice with good accuracy. Experimental implementation of this particular scheme will demonstrate why quantum simulators can be a very effective tool to study different aspects of gauge theories.

The experimental scheme is remarkably simple. The parameter regimes that we prescribe are very well accessible in current experiments with ultracold atoms. Moreover, the fact that it is only the averaged populationimbalance between the odd and even sites that is to be measured, (and not the single site resolved statistics) makes it easier to implement. The only possible source of experimental error could be that in the initial state preparation where the CDW state might not be a perfect one. We have shown in our simulation that such an error affects the dynamics in a very short initial timewindow only. The atomic dynamics, as observed in a longer time-scale, can very well simulate the gauge theory dynamics : creation of particle-antiparticle pairs and breaking of strings.

The proposal is completely scalable that accesses different regimes of gauge theory (with a varying degree of accuracy) and quantum simulate different symmetry sectors. A suitable scaling scheme presented in this paper enables one to model different regimes of LGT with a single experimental set up, just by tuning the controllable experimental parameters. For example, the weak and strong coupling limits of gauge theory is accessed by taking u/t to 0 and $u < u_c$ respectively in the atomic system, where u_c is a quantum critical point beyond which the atomic system enters into a Mott insulating phase as observed in this particular study with small lattice (see Fig. 6). The only requirement here is that the system requires to remain in the same paramagnetic phase throughout the course of its dynamics, so that in a bulk limit, all the allowed states are equally probable at halffilling.

The scheme we present here has a wider applicability compared to its past counterparts, due to the following reasons : (a) Any Formalism that deals with the purely fermionic degrees of freedom of gauge theory (involving a complete gauge fixing) is dimension-specific, whereas, in LSH framework, the treatment remains valid for all dimensions. Here the coupling of matter to the gauge field remains the same as in 1-d lattice for any higher dimension. Hence, construction of any building block, as done in 1-d, will remain useful for higher dimensional models as well. (b) Replacing gauge fields by fermions as a solution of Gauss law constraint introduces long range fermionic interactions in the Hamiltonian. However, in the present work, with the mean field approximation of the loop degrees of freedom of the gauge theory, the Hamiltonian contains only on-site interactions for the fermions that characterize the combined bosonfermion (ends of string) excitations. Inclusion of the dynamical loop degrees of freedom as in (13), along with Abelian Gauss law constraint (A14) gives the complete and most general description of the theory. Then it becomes exactly equivalent to the purely fermionic formulation [13, 74] in one spatial dimension [48].

Future works will address the issue of going beyond mean field approximation, as well as going beyond 1 spatial dimension. The LSH formalism for gauge theories in higher dimensions should be equally useful in constructing atomic quantum simulators for the same. Specifically, within LSH framework, the matter gauge coupling remains the same as in 1d in any higher dimension, including the feature of non-dynamic loop degrees of freedom at matter sites [14, 56, 58]. Hence we expect the present proposal to remain as a useful building block for higher dimensional quantum simulators as well. Work is in progress in these directions and will be reported elsewhere. The present scheme can also be generalized for gauge group SU(3) upon generalization of LSH formalism for SU(3) gauge theory and that will build a concrete step towards quantum simulating QCD.



FIG. 7. (a) The particle number dynamics of the original theory is compared against the simulated dynamics for different values of the coupling x_0 ; (b) percentage deviation from the dynamics of the original theory for perfectly prepared initial state as well as imperfect initial state preparation is compared. A lattice of 6 sites is being considered. The initial state of the system is chosen to be the strong coupling vacuum, where, the alternate sites (odd sites) are fully filled by fermions and other sites (even sites are empty). We consider imperfect initial state preparation at one of its even site and one odd site is half filled that corresponds to almost 17% error in the initial state preparation and (ii) one fully filled even site and one fully vacant odd site, representing almost 33% error in the initial state preparation.

ACKNOWLEDGEMENT

We would like to thank Zohreh Davoudi and Rudranil Basu for useful discussions and also for careful reading of the manuscript and helpful comments. We thank Tilman Esslinger for useful comments on the source of experimental errors in the present proposal. R.D. would like to acknowledge support from the Department of Science and Technology, Government of India in the form of an Inspire Faculty Award (Grant No. 04/2014/002342). I. R. is supported by the U.S. Department of Energy (DOE), Office of Science, Office of Advanced Scientific Computing Research (ASCR) Quantum Computing Application Teams (QCAT) program, under fieldwork Proposal No. ERKJ347.

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Appendix A: Loop-String-Hadron (LSH) Hamiltonian

LSH formalism of lattice gauge theory is based on prepotential framework, where, the original canonical conjugate variables of the theory, i.e color electric field and link operators are replaced by a set of harmonic oscillator doublets, defined at each end of a link [58, 75–82]. In prepotential framework, the SU(2) gauge group is confined to each lattice site allowing one to have local gauge invariant operators and states at each site. For pure gauge theory, these local gauge invariant operators and states can be interpreted as local snapshots of Wilson loop operators of original gauge theory. One can now construct local loop Hilbert space by action of local loop operators on strong coupling vacuum of the theory (no flux state) defined locally at each site. At this point, we must mention that, mapping the local loop picture to original loop description of gauge theory requires one extra constraint on each link, that states

$$N_L(j) = N_R(j) \tag{A1}$$

where, $N_{L(R)}$ is occupation number of prepotentials/Schwinger bosons at the left(right) end of a link connecting sites j and j + 1. This constraint is actually a consequence of the constraint $E_L^2 = E_R^2$ mentioned in section II.

Inclusion of staggered fermionic matter fields for SU(2)gauge theory at each lattice site, combines smoothly with local loop description obtained in prepotential framework as both the prepotential Schwinger bosons and matter fields transform as fundamental representation of the local SU(2) at that site. In addition to local gauge invariant loop operators, one can now combine matter and prepotentials to construct local string operators, that denotes start of a string from a particle and/or end of a string at an antiparticle. Matter fields combine into local gauge invariant configurations representing hadrons likewise in the original formalism. This complete description is named as LSH formalism as in [14]. We are not going into the details of the full LSH formalism here. Instead, we will focus on the application of LSH formulation to one spatial dimension only, and describe the appropriate framework.

Within LSH framework, the gauge invariant and orthonormal LSH basis is characterized by a set of three integers $n_l(j), n_i(j), n_o(j)$ that satisfies Gauss' law constraint:

$$G^{a}(j)|n_{l}(j), n_{i}(j), n_{o}(j)\rangle = 0, \quad \forall j, a.$$
 (A2)

These three quantum numbers signify loop, incoming string and outgoing string at each site. The allowed values of these integers are given by

$$0 \le n_l(j) \le \infty \tag{A3}$$

$$0 < n_i(j) < 1 \tag{A4}$$

$$0 \le n_O(j) \le 1 \tag{A5}$$

Pictorially, the LSH quantum numbers are illustrated in FIG. 8. It is clear from the range of the quantum numbers, that n_l is bosonic excitation, whereas n_i, n_o are fermionic in nature. However, it is important to note that, unlike fermionic matter field in the original theory, the fermionic operators building the 'local string' Hilbert space are SU(2) invariant bilinears of one bosonic prepotential operator and one fermionic matter field, yielding overall fermionic statistics. Hence, the string states contain the information of both gauge field and matter content.

At this point, we define a set of LSH operators consisting of both diagonal and ladder operators locally at each



FIG. 8. Two staggered sites in the LSH formulation on a 1d spatial lattice. Each site carries three types of operators namely incoming string, outgoing string and flux. The Hilbert space is characterized by the corresponding quantum numbers n_l, n_i, n_o respectively for each and every site of the lattice.

site as following:

$$\hat{n}_l |n_l, n_i, n_o\rangle = n_l |n_l, n_i, n_o\rangle \tag{A6}$$

$$\begin{array}{l}n_i|n_l,n_i,n_o\rangle = n_i|n_l,n_i,n_o\rangle \tag{A7}$$

$$\lambda^{\pm}|n_l, n_i, n_o\rangle = |n_l \pm 1, n_i, n_o\rangle \tag{A9}$$

$$\hat{\chi}_{i}^{+}|n_{l}, n_{i}, n_{o}\rangle = (1 - \delta_{n_{i},1})|n_{l}, n_{i} + 1, n_{o}\rangle$$
 (A10)

$$\chi_i |n_l, n_i, n_o\rangle = (1 - \delta_{n_i,0})|n_l, n_i - 1, n_o\rangle$$
 (A11)

$$\chi_{o} |n_{l}, n_{i}, n_{o}\rangle = (1 - o_{n_{o}, 1})|n_{l}, n_{i}, n_{o} + 1\rangle \quad (A12)$$

$$\chi_o |n_l, n_i, n_o\rangle = (1 - \delta_{n_o, 0})|n_l, n_i, n_o - 1\rangle \quad (A13)$$

In the above set of equations, we have not mentioned explicit site index as these are considered to be defined at a particular site.

One major benefit of using LSH formalism is that, one no longer needs to solve/satisfy SU(2) Gauss' law (7) at each site as the basis states are SU(2) gauge invariant by construction. Note that, for non-Abelian gauge theories imposing Gauss' law is a non-trivial task and that gives rise to a whole range of complications as discussed in [48]. However, the LSH formalism still carries the constraint (A1) that is necessary to glue SU(2) invariant states residing at neighboring sites to yield original non-local gauge invariant Hilbert space of the theory. In terms of LSH operators, this constraint (A1) reads as:

$$\hat{n}_l(j) + \hat{n}_o(j)(1 - \hat{n}_i(j)) = \hat{n}_l(j+1) + \hat{n}_i(j+1)(1 - \hat{n}_o(j+1))$$
(A14)

Comparing each side of (A14) to that of (A1) upon acting on LSH basis states, we get:

$$N_L(j) = n_l(j) + n_o(j)(1 - n_i(j))$$
(A15)

$$N_R(j) = n_l(j+1) + n_i(j+1)(1 - n_o(j+1))(A16)$$

where, $N_L(j)$ and $N_R(j)$ count bosonic occupation numbers at each end of the link connecting site j and j + 1. As mentioned earlier, the bosonic occupation number at each end of a link has contribution coming from fermionic excitation n_i and n_o as well. Pictorially, left and right side of (A14) and/or (A1) is represented by the number of thick solid lines at left and right end of a link connecting sites j and j + 1 in FIG. 8. As in [14, 48], definition of a hadronic state in LSH basis is given by $|n_l = 0, n_i = 1, n_o = 1\rangle$ at one particular site.

Hamiltonian of the theory, exactly equivalent to the original Hamiltonian (1) in terms of LSH operators is given by:

$$H^{(\text{LSH})} = H_E^{(\text{LSH})} + H_M^{(\text{LSH})} + H_I^{(\text{LSH})}$$
 (A17)

where, $H_E^{(\text{LSH})}$ is the electric energy term, $H_M^{(\text{LSH})}$ is the mass term and $H_I^{(\text{LSH})}$ is the matter-gauge interaction term of the Hamiltonian. Explicitly, in terms of LSH operators defined in (A6-A13), each part of the Hamiltonian is as below:

$$H_E^{(\text{LSH})} = \frac{g^2 a}{2} \sum_n \left[\frac{\hat{n}_l(j) + \hat{n}_o(j)(1 - \hat{n}_i(j))}{2}, \\ \times \left(\frac{\hat{n}_l(j) + \hat{n}_o(j)(1 - \hat{n}_i(j))}{2} + 1 \right) \right]$$
(A18)

$$H_M^{(\text{LSH})} = m \sum_n (-1)^j (\hat{n}_i(j) + \hat{n}_o(j)), \qquad (A19)$$

$$H_I^{(\text{LSH})} = \frac{1}{2a} \sum_n \frac{1}{\sqrt{\hat{n}_l(j) + \hat{n}_o(j)(1 - \hat{n}_i(j)) + 1}} \,\, (\text{A20}) \\ \left[S_o^{++}(j) S_i^{+-}(j+1) + S_o^{--}(j) S_i^{-+}(j+1) \right] \\ + S_o^{+-}(j) S_i^{--}(j+1) + S_o^{-+}(j) S_i^{++}(j+1) \right] \\ \times \frac{1}{\sqrt{\hat{n}_l(j+1) + \hat{n}_i(j+1)(1 - \hat{n}_o(j+1)) + 1}}.$$

Here (A20) contains LSH ladder operators in the following combinations (suppressing the explicit site index),

$$S_o^{++} = \hat{\chi}_o^+ (\lambda^+)^{\hat{n}_i} \sqrt{\hat{n}_l + 2 - \hat{n}_i}$$
 (A21)

$$S_o^{--} = \hat{\chi}_o^- (\lambda^-)^{\hat{n}_i} \sqrt{\hat{n}_l} + 2(1 - \hat{n}_i)$$
 (A22)

$$S_o^{+-} = \hat{\chi}_i^+ (\lambda^-)^{1-\hat{n}_o} \sqrt{\hat{n}_l + 2\hat{n}_o}$$
(A23)

$$S_o^{-+} = \hat{\chi}_i^- (\lambda^+)^{1-\hat{n}_o} \sqrt{\hat{n}_l + 1 + \hat{n}_o}$$
(A24)

and

$$S_i^{+-} = \hat{\chi}_o^- (\lambda^+)^{1-\hat{n}_i} \sqrt{\hat{n}_l + 1 + \hat{n}_i}$$
(A25)

$$S_i^{-+} = \hat{\chi}_o^+ (\lambda^-)^{1-\hat{n}_i} \sqrt{\hat{n}_l} + 2\hat{n}_i$$
 (A26)

$$S_i^{--} = \hat{\chi}_i^- (\lambda^-)^{\hat{n}_o} \sqrt{\hat{n}_l + 2(1 - \hat{n}_o)}$$
(A27)

$$S_i^{++} = \hat{\chi}_i^+ (\lambda^+)^{\hat{n}_o} \sqrt{\hat{n}_l} + 2 - \hat{n}_o.$$
 (A28)

The strong coupling $(ga \gg 1, ma = \text{fixed})$ vacuum of the LSH Hamiltonian is given by:

$$n_{l}(j) = 0 \quad \forall j$$

$$n_{i}(j) = 1 , n_{o}(j) = 1 \text{ for } j \text{ odd}$$
(A29)

$$n_{i}(j) = 0 , n_{o}(j) = 0 \text{ for } j \text{ even}$$

It is easy to check that (A29) satisfies Abelian Gauss law (A14). One should also consider a suitable boundary condition for one dimensional spatial lattice as discussed in detail in [48] as:

Open Boundary Condition (OBC):

$$N_R(0) = l_i^{OBC}$$

Periodic Boundary Condition (PBC):
 $N_R(0) = N_L(N-1) \equiv l_i^{PBC}$.

where, N_L , N_R are defined in (A15) and (A16) for the first (0) and last (N-1) site of a N site lattice. l_i can be any positive semi definite integer. Now, one can easily check that, for any gauge invariant state $\prod_{j=0}^{N-1} |n_l(j), n_i(j), n_o(j)\rangle$, the bosonic quantum numbers $n_l(j)$ for all values of j are completely determined by the boundary flux l_i and constraint (A14) imposed on each and every link of the lattice starting from one end as:

$$n_{l}(j) = l_{i} + \sum_{y=0}^{j-1} (n_{o}(y) - n_{i}(y)) -n_{i}(j) (1 - n_{o}(j)).$$
(A30)

For OBC, any physical state in LSH formalism is completely determined by (n_i, n_o) quantum numbers at each side. For PBC, the gauge invariant or LSH Hilbert space is characterized by many copies of the same fermionic (n_i, n_o) configurations with different winding number of closed loops, that plays the exact role as the l_i and fixes the n_l 's throughout the lattice. We exploit this particular feature in the analog quantum simulation proposal outlined in the present work ³. Note that, n_l being determined does not mean that we describe a static gauge field theory; rather, truly relevant or physical gauge degrees of freedom are contained into the (n_i, n_o) excitation of any physical state.

Appendix B: Fermi-Hubbard Model and Tight Binding Parameters

Let $\psi_{\uparrow}(x)$ and $\psi_{\downarrow}(x)$ be the field operators corresponding to the two hyperfine states of the fermionic atom. Now, if the lattice potentials are sufficiently deep, the field operators can be expanded in terms of single-particle Wannier functions, localized to each lattice site:

$$\psi_{\sigma}(x) = \sum_{j} c_{\sigma}(j) \mathcal{W}(x - x_j); \qquad \sigma = \uparrow, \downarrow, \quad (B1)$$

where $c_{\sigma}(j)$ is the fermionic annihilation operator for spin index σ and site j. The corresponding number operators are $\mathcal{N}_{j\sigma} = c_{\sigma}^{\dagger}(j)c_{\sigma}(j)$. The Hamiltonian can be written as

$$H = H_{\text{hopping}} + H_{\text{int}} + H_0 \tag{B2}$$

Here H_{hopping} denotes the hopping of a fermion from one site to another, H_{int} represents the interaction when one up-spin fermion shares the same site with a downspin fermion, and H_0 is energy offset, arising out of the single-particle Hamiltonian.

Let

$$\mathcal{H}_0(x) = \frac{-\hbar^2}{2m} \frac{\partial^2}{\partial x^2} + V(x) \tag{B3}$$

In our construction, $V(x) = -V_L \cos x^2$. So effectively, $V(x) = -V_L$ for each site.

Hopping: The hopping term, which represents the tunneling between sites is given by :

$$H_{\text{hopping}} = -\sum_{j} t_{i,j} \left(c_{\uparrow}^{\dagger}(j) c_{\uparrow}(i) + c_{\downarrow}^{\dagger}(j) c_{\downarrow}(i) \right) .2 \text{ (B4)}$$

Tunneling to next-nearest neighbors is usually suppressed by one order of magnitude, in comparison with the nearest neighbor tunnelling. So we consider hopping between adjacent sites only. The tunneling rate from site j to (j + 1) is given by the matrix element

$$t_{j,(j+1)} = -\int \mathcal{W}(x - x_j)\mathcal{H}_0\mathcal{W}(x - x_{j+1})dx.$$
 (B5)

On-site interaction term: In the low-energy scattering regime, the atoms usually interact via s-wave scattering. The corresponding coupling constant is given by

$$g_0 = \frac{4\pi\hbar^2 a_s}{m}$$

 a_s being the scattering length. The interacting part of the Hamiltonian (between up-spin and down-spin fermions sharing the same site) is given by:

$$H_{\rm int} = u \sum_{j} \mathcal{N}_{\uparrow}(j) \mathcal{N}_{\downarrow}(j). \tag{B6}$$

Here the on-site interaction matrix element is given by

$$u = g_0 \int |\mathcal{W}(x - x_j)|^4 dx. \tag{B7}$$

Energy offset : The energy offset can be expressed as :

$$H_0 = \sum_j \epsilon_j \mathcal{N}(j). \tag{B8}$$

where

$$\epsilon_j = \int \mathcal{W}(x - x_j) \mathcal{H}_0 \mathcal{W}(x - x_j) dx.$$
 (B9)

In our construction, the lattice potential $V(x) = -V_L \cos kx^2$, and in Section III A we split this potential as $V_L = V_1 - V_0$. At this point, We drop the contribution

³ The numerical analysis performed in this work is for OBC in gauge theory as simulating the same in an experiment is easier than that for PBC.

from the kinetic term and also V_1 , because these simply add a constant energy shift throughout the lattice. H_{V_0} , the contribution from V_0 is a constant, too, but we keep this, in order to make a direct correspondence with the reduced LSH Hamiltonian.

$$H_{V_0} = \sum_j V_0 \mathcal{N}(j) \tag{B10}$$

Next, alternating potentials V' and -V' are added to

the odd and even sites respectively. Thus the relevant part of the energy offset becomes:

$$H_0 = H_{V_0} + H_{V'} \tag{B11}$$

with

$$H_{V'} = V' \sum_{j=\text{odd}} \mathcal{N}(j) - V' \sum_{j=\text{even}} \mathcal{N}(j) \qquad (B12)$$