A Framework for Simulating Gauge Theories with Dipolar Spin Systems

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Gauge theories appear broadly in physics, ranging from the Standard Model of particle physics to long-wavelength descriptions of topological systems in condensed matter. However, systems with sign problems are largely inaccessible to classical computations and also beyond the current limitations of digital quantum hardware. In this work, we develop an analog approach to simulating gauge theories with an experimental setup that employs dipolar spins (molecules or Rydberg atoms).

We consider molecules fixed in space and interacting through dipole-dipole interactions, avoiding the need for itinerant degrees of freedom. Each molecule represents either a site or gauge degree of freedom, and Gauss’ law is preserved by a direct and programmatic tuning of positions and internal state energies. This approach can be regarded as a form of analog systems programming and charts a path forward for near-term quantum simulation. As a first step, we numerically validate this scheme in a small system study of $U(1)$ quantum link models in $(1+1)$-d with link spin $S = 1/2$ and $S = 1$ and illustrate how dynamical phenomena such as string inversion and string breaking could be observed in near-term experiments. Our work brings together methods from atomic and molecular physics, condensed matter physics, high energy physics, and quantum information science for the study of nonperturbative processes in gauge theories.

Gauge theories are fundamental to descriptions of a wide range of phenomena in many areas of physics. Euclidean lattice field theory [1] has been developed into a general quantitative tool for the study of nonperturbative phenomena of gauge theories using Monte Carlo simulations on classical computers. In particular, the use of lattice gauge theory (LGT) to study QCD (Quantum Chromodynamics) has yielded profound insights into its nonperturbative dynamics and hence our understanding of particle and nuclear physics [2–8]. However, problems involving nonequilibrium dynamics or systems described by complex actions suffer from the sign problem and are not easily amenable to Euclidean LGT simulations.

The possibility of using quantum computations or simulations for these problems has led to renewed interest in Hamiltonian formulations [9]. Quantum link models (QLMs) [10–12], an alternative formulation of LGTs, are well-suited to studies of real-time dynamics on both classical and quantum devices. They use quantum spins in finite integer/half-integer representations of $S$ to replace the infinite gauge degrees of freedom.

Digital quantum computers, typically based on qubits and quantum circuits, are still in the NISQ (noisy intermediate scale quantum) era and are limited by both qubit number and gate depth due to noise and decoherence [13]. Still, for small systems [14] and with the aid of variational techniques [15], digital approaches to the quantum simulation of gauge theory dynamics have shown recent success. Alternatively, analog quantum simulators — special purpose quantum systems designed to implement the real-time evolution of model Hamiltonians — may be more suitable for near-term investigations of LGT dynamics on moderate to large systems [16–18]. While some problems are emulated naturally on physical platforms, such as realizing the Hubbard model with cold atoms in optical lattices [19, 20] or realizing Heisenberg spin models with arrays of polar molecules or Rydberg atoms [21–

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Figure 1. Emulating quantum link models (QLMs) with arrays of dipolar molecules. (a) Mapping between the rotational levels of molecules in an array and the sites and links of the QLM for spin $S = 1/2$. The designation of particular molecules as sites or links ($S_x, L_x, S_{x+1}, L_{x+1}$ for a given unit cell) is enforced through local laser control of level-dependent light shifts. (b) Low-lying molecular rotational levels $|N, m_N\rangle$ and their redefinition in terms of states $|a\rangle$, $|b\rangle$, $|c\rangle$, and $|d\rangle$. (c) The hopping of “fermions” between sites and the associated spin operations on the links are realized by a second-order dipolar exchange of rotational excitations.
elements of a continuous gauge group. In the quantum link model version of LGTs, the link variables are instead represented by non-commuting, finite-dimensional operators that are analogous to quantum spin operators, a feature that makes QLMs more directly accessible to quantum simulation. Here, we consider QLMs of a $U(1)$ LGT in $1+1$ dimensions in the Hamiltonian formulation with staggered fermions \cite{[9]}, which take the form

$$H_{QLM} = -w \sum_x \left[ \psi_x^\dagger U_{x,x+1} \psi_{x+1} + \psi_{x+1}^\dagger U_{x,x+1}^\dagger \psi_x \right] + m \sum_x (-1)^x \tilde{\psi}_x^\dagger \tilde{\psi}_x + \frac{g^2}{2} \sum_x \tilde{E}_{x,x+1}^2,$$

where $x$ labels the spatial lattice sites, $\psi_x$ is the fermion operator with the staggered mass $m(-1)^x$, $w > 0$ is the hopping parameter, $U_{x,x+1}$ is the link variable, $\tilde{E}_{x,x+1}$ is the electric flux for the $(1)$ gauge field on the link between $x$ and $x+1$, and $g$ is the gauge coupling \cite{[9],[12]}. In this paper we focus on QLMs with $S = 1/2$ and $S = 1$ representations for the link variables, but this may be generalized to larger $S$. Note that the underlying LGT is recovered in the $S \to \infty$ limit \cite{[28]}. The physical Hilbert space of the QLM is constrained by the gauge symmetry through Gauss’ law and the gauge transformations are generated by the Gauss’ law operator $\tilde{G}_x = \psi_x^\dagger \psi_x - E_{x,x+1} + E_{x-1,x} + \frac{1}{2}((-1)^x - 1)$, where the last term stems from using staggered fermions.

Dipolar molecules.— A dipolar molecule is effectively a quantum rotor with angular momentum and projection eigenstates $|N_\alpha,m_{N_\alpha}\rangle$. We restrict our consideration to electronic, vibrational, and hyperfine ground states in the absence of large dc electric fields. Furthermore, we only consider rotational angular momentum states with $N_\alpha \in \{0,1\}$, hereafter using the notation $|a\rangle \equiv |0,0\rangle$, $|b\rangle \equiv |1,-1\rangle$, $|c\rangle \equiv |1,0\rangle$ and $|d\rangle \equiv |1,1\rangle$ for the states we consider (see Fig. 1). The Hamiltonian for a system of fixed dipolar molecules (DMH) is

$$H_{DMH} = \sum_{i,\alpha} \left( 2\hbar B_{rot} N_{\alpha}(N_{\alpha} + 1) + \epsilon_{i,\alpha} \right) b_{i,\alpha}^\dagger b_{i,\alpha} + \frac{1}{2} \sum_{i,j} \sum_{\alpha,\beta,\gamma,\eta} V_{i,j}^{\alpha,\beta;\gamma,\eta} b_{i,\alpha}^\dagger b_{j,\gamma}^\dagger b_{j,\beta} b_{i,\alpha}. \quad (2)$$

where $b^\dagger$ and $b$ are hard core bosonic creation and annihilation operators, $B_{rot}$ is the molecule’s rotational constant, $i,j$ label molecular positions, and $\alpha, \beta, \gamma, \eta$ label angular momentum states. The $\epsilon_{i,\alpha}$ are the additional position- and/or state-dependent contributions to the single-molecule energies, which include both spatially uniform but state-dependent terms arising from weak electric fields or nuclear-rotational coupling \cite{[23]}, as well as position-dependent and state-dependent energy shifts, which can be engineered through local differential ac Stark shifts owing to the anisotropic polarizability of molecules \cite{[38]}. In particular, local control over molecular
light shifts could be naturally incorporated in molecule arrays with individual laser addressing [39]. The second line of Eq. 2 represents the dipole-dipole interaction $V_{i,j}^{\alpha,\beta;\gamma,\eta}$, where $\alpha, \beta; \gamma, \eta$ label pairs of initial and final angular momentum states. The form of the interaction terms are naturally restricted by the dipole selection rules $\Delta N = \pm 1$ and $\Delta m_N = 0, \pm 1$, but importantly their dipolar nature allows for the populations of the various molecular levels to be non-conserved (i.e., dipolar interactions allow for the interconversion of rotational and orbital angular momentum [40]). Experimentally, one can use laser power and polarization to tune the relative $\epsilon_{i,\alpha}$ terms, and we consider control of the $V_{i,j}^{\alpha,\beta;\gamma,\eta}$ magnitudes through control of inter-molecular distances [41].

**Effective Hamiltonian.**—We now generate a map between the parameters of the QLM and the physical parameters of the DMH. In our mapping, every site and link in the QLM maps to a different individual molecule; the Hilbert spaces are mapped as in Table I. In this construction, the rotational levels on the “site” molecules are used to represent the “fermions” (which are hard core bosons in one dimension) while rotational levels on the “link” molecules represent the link gauge fields. For a QLM with staggered fermions, each unit cell corresponds to two sites (odd $x$ and even $x$) and two links, with the link between sites $S_x$ and $S_{x+1}$ labeled as $L_x$.

Letting $H_0$ be the one-body terms in the first line of Eq. 2, we tune the energies $\epsilon_{i,\alpha}$ so that Gauss’ law is satisfied in the QLM and so that all of the dipolar configurations satisfying Gauss’ law are nearly degenerate in $H_0$ while all of the other configurations are separated from the Gauss’ law configurations by an energy scale $\Delta \gg V$. While we keep $m, g^2 \ll V \ll \Delta$ to preserve this condition, this still allows for the tuning of $m$ and $g^2/2$ on scales comparable to the hopping $w$. If the molecular system is prepared in an initial state that satisfies Gauss’ law, energy constraints will ensure that the time-evolved state will remain in the physical Hilbert space.

The hopping term $-w \sum_{x} |\psi_x^a U_{x,x+1} \psi_{x+1}^a + h.c.$ of the QLM involves two sites and one link, and implies changes in the states of three molecules, $S_x$, $L_x$, and $S_{x+1}$. Because the DMH (Eq. 2) contains only two-body interactions, matching to the QLM hopping term proceeds by constructing the quasi-degenerate effective Hamiltonian to second order [42] and includes a combination of exchange terms $V_{i,j}$ and state-dependent energies $\epsilon_{i,\alpha}$. In the following two sections we describe the details of this mapping for $S = 1/2$ and $S = 1$.

**Realization and tests of $S = 1/2$ QLM.**—Here we describe details specific to the mapping for the case $S = 1/2$ and we numerically validate this mapping. In this mapping, the molecular state $|c\rangle$ is kept energetically decoupled from gauge-invariant initial states, which can be achieved by means of a small dc electric field. Furthermore, local light shifts can be used to decouple the molecular state $|d\rangle$ from dynamics at the site positions. On the links, while the molecular state $|a\rangle$ does not map directly to anything in the QLM Hilbert space, it is utilized to help mediate the second-order process needed to describe the fermion hopping interaction in the QLM.

A three-body hopping process is illustrated in Fig. 1 for the example: $|a\rangle_{S_x} |d\rangle_{L_x} |b\rangle_{S_{x+1}} \rightarrow |b\rangle_{S_x} |b\rangle_{L_x} |a\rangle_{S_{x+1}}$. In the dipole system this is a second-order process, which can proceed via $|a\rangle_{S_x} |d\rangle_{L_x} |b\rangle_{S_{x+1}} \rightarrow \overset{\text{virtual}}{|b\rangle_{S_x} |a\rangle_{L_x} |b\rangle_{S_{x+1}},}$ where $|b\rangle_{S_x} |a\rangle_{L_x} |b\rangle_{S_{x+1}}$ is an intermediate state outside the physical Hilbert space with an energy difference $\Delta \gg V$.

![Figure 2](image-url) Real-time evolution of densities of sites and links in the dipolar molecular system to simulate the $S = 1/2$ QLM. The dynamics for initialized strings of right-pointing electric fields are shown for the cases of small mass, (a) $m = 0.1w$, and large mass, (b) $m = 2.0w$. Time is in units of the inverse hopping, $w^{-1}$. For small mass (a), the electric field of the string undergoes large-scale oscillations. For large mass (b), the string stays roughly fixed, with only small fluctuations of the charge densities and link spins. To note for both (a) and (b), the outermost sites and links are fixed because of the open boundary conditions. (c) Electric fluxes summed over all dynamical links for $m = 0.1w$ (blue) and $m = 2.0w$ (orange). (d) Top: Fidelity of the dipolar molecular wavefunction versus the QLM wavefunction. Bottom: The effective gauge invariance parameter $G \equiv \sum_x |\langle G_x \rangle|/L [28]$ at the two mass values shown in (a) and (b).
The second-order hopping process has a term given by
\[ -w = \frac{1}{2} \sum_{b,a,d} \sum_{L_x,S_x,+1} \left[ \frac{1}{\Delta \epsilon_{1,x}} + \frac{1}{\Delta \epsilon_{2,x}} \right], \tag{3} \]
with \( \Delta \epsilon_{1,x} = \epsilon_{L_x,a} + \epsilon_{L_x,d} - \epsilon_{L_x,b} - \epsilon_{L_x,a} \) and \( \Delta \epsilon_{2,x} = \epsilon_{L_x,b} + \epsilon_{S_{x+1},a} - \epsilon_{L_x,a} - \epsilon_{S_{x+1},b} \). This yields one equation for every \( x \). While the QLM has a two-site unit cell that repeats, the microscopic parameters of the underlying DMH can be varied slightly between \( x \) and \( x + 2 \), e.g., to allow for the mitigation of undesired processes resulting from the long-ranged dipolar interactions. Given a target \( w \) value, we can then simultaneously solve for all these equations up to corrections of order greater than \( \mathcal{O}(V^2/\Delta) \), which generates relationships between the various \( V \) and \( \epsilon \) values. These parameters are chosen so as to preserve Gauss’ law, and furthermore so that they are physically reasonable. Finally, in addition to the kinetic terms that appear in the second-order effective model, second-order self-interaction terms appear as well. These small diagonal terms can be fully compensated by a slight renormalization of the DMH energy terms \( \epsilon_{i,a} \) (see Supplemental Tables S-3 and S-4 [41]).

We numerically confirm the mapping for \( S = 1/2 \) by simulating the QLM on 3 unit cells with open boundary conditions, and comparing to the full simulation of the DMH. Figure 2a-c shows the DMH dynamics of the sites and links, for an initial product state configuration with staggered site occupations and polarized electric fields. Note that the electric flux energy is always a constant in the \( S = 1/2 \) QLM, and therefore \( g^2 \) is irrelevant [41]. For a small mass, \( m = 0.1w \), the dynamics of the DMH reveal a string inversion of the electric fluxes (Fig. 2a and 2c). For a large mass, \( m = 2.0w \) (more than three times the reported critical mass \( m_c = 0.655w \) [43]), the system shows little dynamics, as there is almost a static flux string with small fluctuations (Fig. 2b and 2c).

We additionally compute the wave-function fidelity and violation of Gauss’ law, shown in Fig. 2d. The fidelity is defined as \(|\langle \psi_{QLM} | \psi_{DMH} \rangle|^2\), where \( \psi_{DMH} \) is the wave-function from the DMH and \( \psi_{QLM} \) is the wave-function mapped from the QLM into the DMH space. The fidelity of our scheme is greater than 0.998 out to \( t = 20w^{-1} \), and Gauss’ law is preserved over this time range at the \( 10^{-7} \) level. For the molecule NaRb (body-frame dipole moment of 3.3 D) and a minimum separation of 0.5 \( \mu m \), these calculations relate to a hopping rate of \( w/h = 83 \) Hz (with \( h \) Planck’s constant). This robust energy scale should be compatible with long molecule trapping times [44] and coherence times [26, 27]. In addition, even larger values of the hopping can be achieved by reducing the scale of imposed energy penalties \( \Delta \) [41], albeit with fidelities lower than those shown in Fig. 2d.

Realization and tests of \( S = 1 \) QLM — The \( S = 1 \) case is realized similarly to \( S = 1/2 \), however more internal levels are used to represent the larger number of link spin values. The link states are represented by the various \( N = 1 \) rotational sub-levels as in Table I. For the sites, the \( |b \rangle \) and \( |d \rangle \) levels are decoupled from the dynamics by large local light shifts. Unlike for \( S = 1/2 \), in the \( S = 1 \) case second-order self-interaction processes, such as \( |c \rangle_s |b \rangle_{L_x} |a \rangle_{S_{x+1}} \xrightarrow{\text{virtual}} |c \rangle_s |a \rangle_{L_x} |c \rangle_{S_{x+1}} \), cannot be entirely removed through coordination of the \( \epsilon_{i,a} \) terms. This comes from the fact that they are not simply renormalized one-body terms. For example, the above process generates an \( \mathcal{O}(V^2/\Delta) \) term of the form \( |b \rangle_{L_x} b_s |a \rangle_{S_{x+1}} a_s |b_s a_s \) that is not in the QLM. Nevertheless, these additional terms are diagonal in the molecular \( \{|N, m_N\} \) basis and still preserve Gauss’ law, but cannot be removed because the simultaneous constraints for \( w \) and setting these “extra” terms to zero results in an over-determined system of equations. To overcome this, we introduce a new length scale.

For every \( x \), we set the distances between molecules representing \( S_x \) and \( L_x \) to be small and denote the characteristic scale of the dipole-dipole interaction between them as \( V_{\text{short}} \). Meanwhile, the distances between molecules representing \( L_x \) and \( S_{x+1} \) are chosen to be larger with a characteristic scale for the dipole-dipole in-

Figure 3. Real-time evolution of fermions and links in the dipolar molecular system to simulate the \( S = 1 \) QLM with \( g^2 = \sqrt{2}w \) starting from a string of right-pointing electric fields at (a) \( m = 0.25 \times \sqrt{2}w \) and (b) \( m = 2.0 \times \sqrt{2}w \). Time is in units of \( (\sqrt{2}w)^{-1} \). With a small mass \( m \), the string breaks on the hopping timescale, resulting in two approximate mesons on the edges and approximate vacuum in between. With a large mass \( m \), the string approximately remains, with small fluctuations in densities. In (a) and (b), the densities of the two sites and two links on the edges are fixed due to the open boundary condition. (c) Electric fluxes summed over all dynamical links for both \( m = 0.25 \times \sqrt{2}w \) (blue) and \( m = 2.0 \times \sqrt{2}w \) (orange). Solid and dashed lines relate to the DMH and QLM dynamics, respectively. (d) Top: Fidelity of the dipolar molecular wavefunction versus the QLM wavefunction. Bottom: The effective gauge invariance parameter \( G \equiv \sum_x |\langle \tilde{G}_x \rangle|/L \) [28] at two values of masses in (a) and (b).
teraction between them of $V_{\text{long}}$, and $V_{\text{long}} \ll V_{\text{short}}$. The hopping parameter $w$ is $\mathcal{O}(V_{\text{short}}V_{\text{long}}/\Delta)$. The second-order self-interaction term for $S_z$ and $L_z$ is $\mathcal{O}(V_{\text{short}}^2/\Delta)$ and that for $L_z$ and $S_{z+1}$ is $\mathcal{O}(V_{\text{long}}^2/\Delta)$, which can be safely neglected. Therefore, we only need to consider the equations between $S_z$ and $L_z$, which decreases the total number of equations and leaves them under-determined. A non-unique solution to these equations can then be found [41], and we can obtain experimental parameters such that these second-order terms are made small or can be removed from the effective Hamiltonian. To note, for larger $V_{\text{short}}/V_{\text{long}}$ ratios, higher-order terms eventually limit this minimization of the extra terms [41].

To benchmark our scheme for $S = 1$, we perform exact diagonalization on three unit cells with open boundary conditions. The initial configuration is chosen as shown in Fig. 3, relating to a flux string connecting static edge. To verify that our scheme reflects the physical string may break into approximate vacuum in the middle region, resulting in the production of two mesons on the edge. To verify that our scheme reflects the physical dynamics properly, we investigate both a small mass scenario (Fig. 3a, $\{m = 0.25 \times \sqrt{2}w, g^2 = \sqrt{2}w\}$) and a large mass scenario (Fig. 3b, $\{m = 2.0 \times \sqrt{2}w, g^2 = \sqrt{2}w\}$), which should result in string breaking and a stabilized string, respectively. Figure 3 shows that the dynamics of the dipolar molecular system (solid lines) reflects this behavior. We find good agreement with the expected dynamics of the target QLM (dashed line). Specifically, for large mass, the string stays approximately in its initial configuration up to small fluctuations, while the string of the small mass case breaks on the hopping timescale. These results are consistent with the estimated critical length $L_c = 4m/g^2 + 3$ [28].

Gauss’ law is preserved to high accuracy (see Fig. 3d), however the fidelity of the DMH “simulator” drops from 1.0 to roughly 0.5 over 10-20 time units (in terms of $(\sqrt{2}w)^{-1}$). This infidelity comes almost entirely from the additional gauge-invariant self-interaction terms that do not appear in the QLM. These terms slightly modify the frequencies of oscillations in the DMH and QLM dynamics, and thus have a large influence on the fidelity at long times, but otherwise do not alter the expected QLM phenomenology (see Fig. 3c).

While one approach to improving this fidelity is simply adding the additional gauge-invariant terms to the target QLM (as is done in Ref. [28]), we can suppress the additional gauge invariant terms and improve the fidelity by adjusting the “long-short” distance ratios. In this work we use the value of $V_{\text{short}}/V_{\text{long}} = 1.5^{3} \approx 3.4$ (see Fig. 3), which represents a non-optimal but more experimentally realistic compromise [41]; the parameters used in Fig. 3 already relate to hopping energies of $\sqrt{2}w/h = 3.2$ Hz for NaRb (3.3 D) and an assumed minimum spacing of 0.5 μm. See the supplemental material for the fidelity increase which can be achieved as a function of the “long-short” distance ratio (Fig. S2 and Fig. S3 [41]).

Conclusions and outlook.— In this paper, we propose a new approach for simulating quantum link models based on the restricted internal-state dynamics of fixed dipolar spins. Our numerical tests of simple $U(1)$ LGTs in $1 + 1$ dimensions show that this approach enables the experimental exploration of important dynamical phenomena such as string inversion and string breaking. Further directions are suggested by the present work, including extensions to realizations of higher $S$ values, non-Abelian LGTs, quantum mechanical $\theta$ angles [47], and QLM dynamics in higher dimensions [48]. Moreover, while our scheme based on dipolar spin interactions is of relevance to a range of systems, such as cold molecules and Rydberg atom arrays, the potential reach could be greatly broadened by generalizing this framework to generic spin systems, as realized by most physical quantum information platforms.

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[1] K. G. Wilson, Phys. Rev. D 10, 2445 (1974).
[2] C. DeTar and S. Gottlieb, Physics Today 57, 45 (2004).
[3] A. S. Kronfeld, Ann. Rev. Nucl. Part. Sci. 62, 265 (2012).
[4] A. X. El-Khadra, in Proceedings, Theoretical Advanced Study Institute in Elementary Particle Physics: Journeys Through the Precision Frontier: Amplitudes for Colliders (TASI 2014): Boulder, Colorado, June 2-27, 2014, WSP (WSP, 2015) pp. 183–233.
[5] H.-T. Ding, F. Karsch, and S. Mukherjee, Int. J. Mod. Phys. E 24, 1530007 (2015).
[6] C. Ratti, Reports on Progress in Physics 81, 084301 (2018).
[7] C. Lehner et al. (USQCD), Eur. Phys. J. A 55, 195 (2019).
[8] S. Aoki et al. (Flavour Lattice Averaging Group), (2019), arXiv:1902.01819 [hep-lat].
[9] J. Kogut and L. Susskind, Phys. Rev. D 11, 395 (1975).
[10] D. Horn, Phys. Lett. B 100, 149 (1981).
[11] P. Orland and D. Rohrlich, Nucl. Phys. B 338, 647 (1990).
[12] S. Chandrasekharan and U.-J. Wiese, Nucl. Phys. B 534, 516 (2016).
Supplementary Material for “A Framework for Simulating Gauge Theories with Dipolar Spin Systems”

Quantum link models

We start with the Hamiltonian for (1 + 1)-d U(1) LGT with staggered fermions in the temporal gauge,

\[ H_{\text{LGT}} = -w \sum_x \left[ \bar{\psi}_x U_{x,x+1} \psi_{x+1} + \bar{U}_{x,x+1}^\dagger \psi_x \right] + m \sum_x (-1)^x \psi_x^\dagger \psi_x + \frac{g^2}{2} \sum_x E_{x,x+1}^2. \]  

(S1)

The link variables \( U_{x,x+1} = \exp(ia g A_{x,x+1}) \) take continuous values in the group U(1), where \( A_{x,x+1} \) is the spatial component of the U(1) gauge field and \( a \) is the lattice spacing. The electric flux \( E_{x,x+1} = -\frac{1}{a g} \partial_A A_{x,x+1} \) is proportional to the canonical momentum of \( A_{x,x+1} \) and can take any integer values. Commutation relations for quantum operators on a link are \([U_{x,x+1}, U_{x,x+1}^\dagger] = 0\), \([E_{x,x+1}, U_{x,x+1}] = U_{x,x+1}\), and \([E_{x,x+1}, U_{x,x+1}^\dagger] = -U_{x,x+1}^\dagger\). In the QLM version of this LGT, the first commutation relation is modified to \([U_{x,x+1}, U_{x,x+1}^\dagger] = 2E_{x,x+1}\). In analogy with quantum angular momentum operators, we can write \( U_{x,x+1} = S_{x,x+1}^+ U_{x,x+1}^\dagger = S_{x,x+1}^- U_{x,x+1}^\dagger = S_{x,x+1}^3 \), so that each link is in a spin-S representation with \( S = 0, 1/2, 1, \cdots \). The Hamiltonians of the QLM and the LGT look exactly the same, but the link variables and the sizes of the Hilbert spaces are different. If the number of sites on the lattice is finite, then the QLM Hilbert space is finite whereas the LGT Hilbert space is infinite. The physical Hilbert space of the QLM is defined through the Gauss’ law \( \hat{G}_x \mid \text{phys} \rangle = 0 \), with the Gauss’ law operator defined in the main text and \( \mid \text{phys} \rangle \) is any state in the physical Hilbert space.

Molecular dipole-dipole interaction

The molecular dipole-dipole interaction [24] is

\[ V = \frac{1}{2} \sum_{ij} \sum_{\alpha,\beta,\gamma,\eta} V_{ij}^{\alpha,\beta,\gamma,\eta} \hat{b}^\dagger_{i,\gamma} \hat{b}^\dagger_{j,\eta} \hat{b}_{j,\beta} \hat{b}_{i,\alpha} = \frac{1}{2} \sum_{ij} \hat{V}_{ij} \]  

(S2)

where

\[ \hat{V}_{ij} = \frac{1}{4\pi \epsilon_0} \frac{\hat{d}_i \cdot \hat{d}_j - 3 \left( \hat{d}_i \cdot \hat{r}_{ij} \right) \left( \hat{d}_j \cdot \hat{r}_{ij} \right)}{r_{ij}^3} = -\frac{\sqrt{6}}{4\pi \epsilon_0 r_{ij}^3} \sum_{p=-2}^{2} (-1)^p T^2_p(C) T^2_p \left( \hat{d}_i, \hat{d}_j \right) \]

(S3)

is the interaction between two molecules at positions \( \hat{r}_i \) and \( \hat{r}_j \), \( \hat{r}_{ij} = \hat{r}_j - \hat{r}_i \) is the vector connecting these two molecules, \( \hat{r}_{ij} = \hat{r}_{ij}/r_{ij} \) is the directional vector, \( \hat{d}_i \) is the dipole operator of the molecule \( i \), and \( \hat{d}_j \) is that of the molecule \( j \). The functions \( T^2_p(C) \) are proportional to spherical harmonics \( Y_{2,-p} \)

\[ T^2_0(C) = \frac{(3 \cos^2 \theta_{ij} - 1)}{2} = 2 \sqrt{\frac{\pi}{5}} Y_{2,0} (\theta_{ij}, \phi_{ij}), \]  

(S4)

\[ T^2_{\pm 1}(C) = \pm \sqrt{\frac{3}{2}} e^{\pm i \phi_{ij}} \sin \theta_{ij} \cos \theta_{ij} = 2 \sqrt{\frac{\pi}{5}} Y_{2,\pm 1} (\theta_{ij}, \phi_{ij}), \]  

(S5)

\[ T^2_{\pm 2}(C) = \sqrt{\frac{3}{8}} e^{\pm 2i \phi_{ij}} \sin^2 \theta_{ij} \cos \theta_{ij} = 2 \sqrt{\frac{\pi}{5}} Y_{2,\pm 2} (\theta_{ij}, \phi_{ij}), \]  

(S6)

where the polar and azimuthal angles \( \theta_{ij} \) and \( \phi_{ij} \) are measured with respect to the quantization axis (which we assume to be defined by the direction of a strong uniform magnetic field in experiments with small or zero dc electric field). The quantization axis does not specify a second direction so the azimuthal angles \( \phi_{ij} \) are defined only up to an overall offset. \( T^2_p \left( \hat{d}_i, \hat{d}_j \right) \) are rank-2 tensor operators

\[ T^2_0 \left( \hat{d}_i, \hat{d}_j \right) = \frac{2}{\sqrt{6}} \left[ \hat{d}_i^\dagger \hat{d}_j + \hat{d}_i^\dagger \hat{d}_j^\dagger + \hat{d}_i \hat{d}_j^\dagger \right], \]  

(S7)
\[
T_{\pm 1}^2 \left( \hat{d}_i, \hat{d}_j \right) = \frac{\hat{d}_0 \hat{d}_{\pm 1} + \hat{d}_{\pm 1} \hat{d}_0}{\sqrt{2}}, \tag{S8}
\]

\[
T_{\pm 2}^2 \left( \hat{d}_i, \hat{d}_j \right) = \hat{d}_{\pm 1} \hat{d}_{\pm 1}, \tag{S9}
\]

where \( \hat{d}_\pm = \hat{d}_x \pm i \hat{d}_y, \hat{d}_0 = \hat{d}_z, \hat{d}_{\pm 1} = \mp (\hat{d}_x \pm i \hat{d}_y)/\sqrt{2}. \hat{d}_{+1} = -\hat{d}_+ / \sqrt{2}, \hat{d}_{-1} = \hat{d}_- / \sqrt{2}. \) It is worth noting the minus sign in the relation \( \hat{d}_{+1} = -\hat{d}_{-1}. \)

The matrix elements of \( \hat{d}_q \) for a given molecule are

\[
\left\langle N', m'_N | \hat{d}_q | N, m_N \right\rangle = d (\mp 1)^{m'_N} \sqrt{(2N' + 1)(2N + 1)} \left( \begin{array}{ccc} N' & 1 & N \\ -m'_N & q & m_N \end{array} \right) \left( \begin{array}{ccc} N' & 1 & N \\ 0 & 0 & 0 \end{array} \right), \tag{S10}
\]

where the parentheses are Wigner 3-j symbols and \( d \) is the electric dipole moment of the molecule. Dipole selection rules \( \Delta N = N' - N = \pm 1 \) and \( \Delta m_N = m'_N - m_N = 0, \pm 1 \) are required to have a nonzero matrix element explicitly by the second Wigner 3-j symbol.

We only consider \( N = 0 \) and \( N = 1 \) states. States with \( N \geq 2 \) are naturally off-resonant from the initialized configurations. We introduce in the main text the notation for these four states \( |a\rangle \equiv |0,0\rangle, |b\rangle \equiv |1,-1\rangle, |c\rangle \equiv |1,0\rangle \) and \( |d\rangle \equiv |1,1\rangle \) for each molecule. Without external electric fields, magnetic fields, or laser fields, and ignoring internal nuclear structures (i.e., only considering the rotational kinetic energy), states \( |b\rangle, |c\rangle, \) and \( |d\rangle \) of a single molecule are degenerate and their energy is greater than the energy of \( |a\rangle \) by \( 2h B_{\text{rot}} \), where \( B_{\text{rot}} \) is the rotational constant.

Single-molecule nonvanishing dipole matrix elements within the four-state subspace stated above are

\[
\left\langle a | \hat{d}_{+1} | b \right\rangle = \left\langle 0, 0 | \hat{d}_{+1} | 1, -1 \right\rangle = -\frac{1}{\sqrt{3}} d, \tag{S11}
\]

\[
\left\langle a | \hat{d}_0 | c \right\rangle = \left\langle 0, 0 | \hat{d}_0 | 1, 0 \right\rangle = \frac{1}{\sqrt{3}} d, \tag{S12}
\]

\[
\left\langle a | \hat{d}_{-1} | d \right\rangle = \left\langle 0, 0 | \hat{d}_{-1} | 1, +1 \right\rangle = -\frac{1}{\sqrt{3}} d, \tag{S13}
\]

and their complex conjugates.

Non-zero matrix elements of the dipole-dipole interaction between a pair of molecules in any combinations of single-molecule states can be easily calculated by multiplying single-molecule matrix elements. Non-zero matrix elements are

\[
V^{a,b,c,d}_{i,j} = \left( V_{i,j}^{a,b,c,d} \right)^* \equiv \left( V_{i,j}^{a,c,b,d} \right)^* = -\frac{1}{4\pi e_0 r_{i,j}^3} \sin^2 \theta_{i,j} \frac{1}{2} d^2, \tag{S14}
\]

\[
V^{a,a,c,c}_{i,j} = \left( V_{i,j}^{a,a,c,c} \right)^* = \left( V_{i,j}^{c,c,a,a} \right)^* = -\frac{1}{4\pi e_0 r_{i,j}^3} \sin \theta_{i,j} \cos \theta_{i,j} \frac{1}{\sqrt{2}} d^2, \tag{S15}
\]

\[
V^{a,a,d,d}_{i,j} = \left( V_{i,j}^{a,a,d,d} \right)^* = \left( V_{i,j}^{d,d,a,a} \right)^* = -\frac{1}{4\pi e_0 r_{i,j}^3} \sin^2 \theta_{i,j} \frac{1}{\sqrt{2}} d^2, \tag{S16}
\]

\[
V^{a,a,b,c}_{i,j} = \left( V_{i,j}^{a,a,b,c} \right)^* = \left( V_{i,j}^{b,c,a,a} \right)^* = -\frac{1}{4\pi e_0 r_{i,j}^3} \sin \theta_{i,j} \cos \theta_{i,j} \frac{1}{\sqrt{2}} d^2, \tag{S17}
\]

\[
V^{a,a,b,d}_{i,j} = \left( V_{i,j}^{a,a,b,d} \right)^* = \left( V_{i,j}^{b,d,a,a} \right)^* = -\frac{1}{4\pi e_0 r_{i,j}^3} \frac{3 \cos^2 \theta_{i,j} - 1}{2} \frac{1}{3} d^2, \tag{S18}
\]

\[
V^{a,a,c,d}_{i,j} = \left( V_{i,j}^{a,a,c,d} \right)^* = \left( V_{i,j}^{c,d,a,a} \right)^* = \frac{1}{4\pi e_0 r_{i,j}^3} \frac{3 \cos^2 \theta_{i,j} - 1}{2} \frac{1}{3} d^2, \tag{S19}
\]
\[ V_{i,j}^{a,b;\ast} = V_{i,j}^{b,a;\ast} = \frac{1}{4\pi\epsilon_0 r_{i,j}^3} \left( 3\cos^2\theta_{i,j} - 1 \right) \frac{1}{3} d^2, \]  
(S20)

\[ V_{i,j}^{a,c;\ast,a} = V_{i,j}^{c,a;\ast} = -\frac{1}{4\pi\epsilon_0 r_{i,j}^3} \left( 3\cos^2\theta_{i,j} - 1 \right) \frac{1}{3} d^2, \]  
(S21)

\[ V_{i,j}^{a,d;\ast,a} = V_{i,j}^{d,a;\ast} = \frac{1}{4\pi\epsilon_0 r_{i,j}^3} \left( 3\cos^2\theta_{i,j} - 1 \right) \frac{1}{3} d^2, \]  
(S22)

\[ V_{i,j}^{a,b,c;\ast} = V_{i,j}^{b,c,a;\ast} = \left( V_{i,j}^{c,a,b;\ast} \right)^* = \left( V_{i,j}^{b,c,a;\ast} \right)^* = -\frac{1}{4\pi\epsilon_0 r_{i,j}^3} e^{-i\phi_{i,j}} \sin\theta_{i,j} \cos\theta_{i,j} \frac{1}{\sqrt{2}} d^2, \]  
(S23)

\[ V_{i,j}^{a,b,d;\ast,a} = V_{i,j}^{b,d,a;\ast} = \left( V_{i,j}^{d,a,b;\ast} \right)^* = \left( V_{i,j}^{a,b,d;\ast} \right)^* = \frac{1}{4\pi\epsilon_0 r_{i,j}^3} e^{-2i\phi_{i,j}} \sin^2\theta_{i,j} \frac{1}{\sqrt{2}} d^2, \]  
(S24)

\[ V_{i,j}^{a,c,d;\ast,a} = V_{i,j}^{c,d,a;\ast} = \left( V_{i,j}^{d,a,c;\ast} \right)^* = \left( V_{i,j}^{a,c,d;\ast} \right)^* = \frac{1}{4\pi\epsilon_0 r_{i,j}^3} e^{-i\phi_{i,j}} \sin\theta_{i,j} \cos\theta_{i,j} \frac{1}{\sqrt{2}} d^2. \]  
(S25)

Molecular internal state energies and their experimental control

The dipole-dipole interactions described in the previous section provide the fundamental mechanism by which dynamics can proceed and by which the densities of various internal states can evolve in the considered system of molecular “spins” fixed in place. In the mapping to the QLM, this will provide a mechanism for “fermions” (or “charges”) hopping, as represented by spin excitations (i.e., hard core bosons) being exchanged between different fixed molecules.

Equally important to our proposed framework is the ability to restrict these dipolar exchange processes in a controlled way. Specifically, by imposing energetic constraints on the various internal-state configurations of the molecules, we can effectively impose gauge invariance or enforce Gauss’ law, by only allowing processes that correlate the hopping of fermions between “sites” with the modification of the spin that resides on the intervening “link.” In this approach, such energetic constraints are imposed directly on the molecules through the single-particle terms \( H_0 \) of the DMH. These energy terms \( \epsilon_{i,a} \) of the DMH depend in general on both on the molecule position (labeled by the index \( i \)) and the internal rotational level (denoted by \( a \)). First off, for typical experiments on ultracold molecules operating at large magnetic fields (near the field values used for magnetoassociation of the atomic constituents), rotational-level dependent energy terms arise due to the weak coupling between molecular rotation and the hyperfine (nuclear) degrees of freedom [23]. These naturally arising shifts to the various rotational levels serve to break the degeneracy of the \( N = 1 \) rotational manifold at the scale of \( \sim 10 – 100 \) kHz, even in zero electric field. In addition to this, the energies of the rotational levels can be modified globally through the addition of weak dc electric fields or off-resonant (and polarized) microwave fields. These can be used, e.g., for the purpose of shifting particular rotational sub-levels of the \( N = 1 \) manifold by a large amount so as to decouple it from near-resonant dipole-driven dynamics.

Finally, and most central to the proposed approach, spatially-resolved control of the internal state energies can be engineered by direct optical addressing, using level-dependent ac Stark shifts to tune the internal state energies. Such an ability to locally address individual molecules arises naturally in implementations based on micro-trapped arrays [39], but could also be achieved by projecting tailored laser patterns onto lattice-trapped samples.

In the proposed scheme, the positions of all molecules are fixed, and the total number of molecules as well as the total number of molecules in the \( N = 0 \) rotational ground state, or level \( |a\rangle \), are conserved. As such, the full tuning of all relevant configurations of molecules can be accomplished through local and level-dependent control of the differential (with respect to \( |a\rangle \)) ac Stark shift of the utilized \( N = 1 \) sublevels. In general, molecules play host to a strongly anisotropic and rotational level-dependent ac polarizability [38]. By control of the local laser intensity and polarization (with respect to the quantization axis, here assumed to be defined by a quantization magnetic field), a large differential tuning of these energies is available for almost any laser wavelength.

For a complete and general control, we consider addressing the array of molecules with a control laser that is tuned near to a narrow optical transition from the molecular ground state to a relatively long-lived electronic excited
state. For commonly used alkalis such as NaRb or KRb, this could for example relate to transitions of the form \( |X^1\Sigma, \nu = 0, N = 1, m_N \rangle \rightarrow |b^1\Sigma_{\alpha}, \nu = 0, N = 0, m_N = 0 \rangle \), characterized by kHz-level linewidths [49, 50]. In particular, for GHz-scale detunings from such a transition, local control of laser intensity and polarization would provide a complete control over all relevant differential rotational level-dependent energies \( \epsilon_{i,\alpha} \) of the DMH, owing to dipole selection rules. Shifts at the necessary scales (even up to order MHz) can be accommodated with modest optical powers in scenarios based on local projection of tightly-focused lasers. For alternative realizations based on arrays of Rydberg atoms, we note that the control of internal state-dependent energies via local state-dependent ac Stark shifts has already been demonstrated [36].

The local detection of the various molecular internal states could be accomplished, e.g., by mapping them on to different atomic levels (in a reversal of STIRAP process) followed by imaging of the atoms, or alternatively by extensions of direct molecular detection methods [51]. Similar capabilities will be equally critical to the development of molecules as qubits or qudit architectures for applications in quantum information science.

### Quasi-degenerate effective Hamiltonians

Before the construction of QLM Hamiltonians, we first introduce the method of quasi-degenerate effective Hamiltonians [42]. This method is a perturbative way of calculating an effective Hamiltonian that will yield similar dynamics as the original Hamiltonian \( H = H_0 + V \) in a subspace \( \alpha \) which we are interested in. Eigenstates \( |m, \alpha \rangle \in \alpha \) of \( H_0 \) are given by \( H_0 |m, \alpha \rangle = E_{m\alpha} |m, \alpha \rangle \). \( E_{m\alpha} \) for different \( m \)’s are nearly degenerate and small variations are allowed. Eigenvalues of eigenstates of \( H_0 \) outside the subspace \( \alpha \) are separated from \( E_{m\alpha} \). To second order, the matrix elements of the effective Hamiltonian for \( \alpha \) are

\[
\langle m, \alpha | H^\alpha_{\text{eff}} | n, \alpha \rangle = E_{m\alpha} \delta_{m,n} + \langle m, \alpha | V | n, \alpha \rangle + \frac{1}{2} \sum_{l,\gamma \neq \alpha} \langle m, \alpha | V | l, \gamma \rangle \langle l, \gamma | V | n, \alpha \rangle \left[ \frac{1}{E_{m\alpha} - E_{l\gamma}} + \frac{1}{E_{n\alpha} - E_{l\gamma}} \right] + \cdots ,
\]

where Greek letters label subspaces and Roman letters label states.

In our case of realizing QLMs, \( \alpha \) is chosen as the subspace of DMHs that maps to the physical Hilbert space of QLMs. There are no two-body interactions in the QLM Hamiltonian so there should be no such terms in the effective Hamiltonian, either. We describe here how to suppress these terms in the effective Hamiltonian. The dipole-dipole interaction decays as a power law of \( r^{-3} \) but it exists even for two molecules far away from each other. We choose parameters in Table S-1, as will be explained below, such that only our wanted DMH states are nearly degenerate and \( \langle i, \alpha | V | j, \alpha \rangle \) always vanishes. In addition, for the two molecules extremely far away from each other such that their dipole-dipole interaction is much smaller than \( w, m, \) and \( g^2 \), then their dipole-dipole interaction can be neglected and the requirement of energy separations can be loosened in this case.

### Construction of Fermions from Hard Core Bosons

We will map the fermion operators in QLMs to spin operators through a Jordan-Wigner transformation, which will be further related to the hard core boson operators in the DMH.

According to the mapping between fermion site states and dipolar molecule states in Table I in the main text for both spin 1/2 and spin 1, the occupied fermion site is always mapped to \( |a \rangle \) while the unoccupied fermion site is mapped to either \( |b \rangle \) or \( |c \rangle \). In our setup, each “fermion” site in the DMH is either occupied or unoccupied. Namely, \( b_{x,a}^\dagger b_{x,a} + b_{x,a}^\dagger b_{x,a} = 1 \), where \( \downarrow = b \) or \( c \). In this situation, each molecule is a two-level system and analogous to spin-1/2.

Since the internal state excitations of dipolar molecules can be described as hard core bosons, in one spatial dimension a Jordan-Wigner transformation can map the hardcore boson states to the fermion states while preserving the locality of local operators. The Jordan-Wigner transformation takes fermion operators to spin-1/2 operators, \( \psi_x^\dagger \prod_{\beta=3}^{x=1} e^{i \tau_x \psi_x^\dagger} \psi_x = S^+_x \), \( \psi_x \prod_{\beta=1}^{x=3} e^{-i \tau_x \psi_x^\dagger} \psi_x = S^-_x \). As a result, \( \psi_x^\dagger \psi_x = S^3_x + 1/2 \) and \( \psi_x^\dagger \psi_{x+1} = S^+_x S^-_{x+1} \). The spin-1/2 operators mentioned above, \( S^+_x, S^-_x, S^3_x \), have no relation with the spin operators on quantum links. Similarly, they are not to be confused with the actual site identifiers of the form \( S_x \) and \( S_{x+1} \) as introduced in the main text.

Finally, we have relations between spin operators and hard core bosonic operators in the DMH, which are \( S^+_x = b_{x,a}^\dagger b_{x,a} - b_{x,a} b_{x,a}^\dagger \), \( S^-_x = b_{x,a}^\dagger b_{x,a} + b_{x,a} b_{x,a}^\dagger \), \( S^3_x = (b_{x,a}^\dagger b_{x,a} - b_{x,a} b_{x,a}^\dagger)^2 / 2 \). They give rise to \( \psi_x^\dagger \psi_x = S^3_x + 1/2 = (b_{x,a}^\dagger b_{x,a} - b_{x,a} b_{x,a}^\dagger + 1)/2 \) and \( \psi_x^\dagger \psi_{x+1} = b_{x,a}^\dagger b_{x,a} b_{x,a}^\dagger b_{x+1,a} + 1/2 \).
Construction of the $S = 1/2$ QLM Hamiltonian

For the $S = 1/2$ QLM, the link operators can be mapped to hard core bosonic operators as follows: $U_{x,x+1} = S^+_{x,x+1} = b^\dagger_{(x,x+1),d} b_{(x,x+1),d}$, $U^\dagger_{x,x+1} = S^-_{x,x+1} = b^\dagger_{(x,x+1),b} b_{(x,x+1),b}$ and $E_{x,x+1} = S^3_{x,x+1} = (b^\dagger_{(x,x+1),d} b_{(x,x+1),d} - b^\dagger_{(x,x+1),b} b_{(x,x+1),b})/2$. With the mappings given above, the $S = 1/2$ QLM Hamiltonian can be written in the dipolar molecular operators

$$H = -w \sum_x \left[ b_{x,d}^\dagger b_{x,b} b_{(x,x+1),d} b_{(x,x+1),b} b_{x+1,a} + h.c. \right] + m \sum_x (-1)^x \frac{1}{2} \left( b_{x,a}^\dagger b_{x,a} - b_{x,b}^\dagger b_{x,b} + 1 \right) + \frac{g^2}{2} \sum_x \frac{1}{4} \tag{S27}$$

where the last term is a constant and can be discarded.

The hopping process $|a\rangle_{S_x} |d\rangle_{L_x} |b\rangle_{S_{x+1}} \xrightarrow{\text{virtual}} |b\rangle_{S_x} |a\rangle_{L_x} |b\rangle_{S_{x+1}} \xrightarrow{\text{virtual}} |b\rangle_{S_x} |b\rangle_{L_x} |a\rangle_{S_{x+1}}$ described in the main text (where $|b\rangle_{S_x} |a\rangle_{L_x} |b\rangle_{S_{x+1}}$ is a virtual intermediary) is illustrated schematically in Fig. S1.

![Figure S1. Schematic hopping on $S_x$, $L_x$, $S_{x+1}$ through a virtual intermediate molecular configuration. The QLM states and the molecular levels are shown together for the initial and final configurations, respectively. The key hopping process of the QLM is achieved through a second-order dipolar exchange process, where the correlation between matter hopping and the changes to the link spin (Gauss’ law) is imposed through energetic constraints.](image)

We choose $\epsilon_{i,\alpha}$ introduced in Eq. 2 in the main text as is shown in Table S-1 for the $n$-th unit cell. $\Delta_{1,n}$, $\Delta_{2,n}$, $\delta_{1,n}$, $\delta_{2,n}$ for any $n$ are at the order of $\Delta$ mentioned in the main text and will be specified in numerical simulations.

For a molecule in the $N = 0$ ($|\alpha\rangle$) state and another molecule in an $N = 1$ state, there are self-interactions to second order, a virtual process in which a state first hops to an intermediate state and then hops back to the initial state. This type of self-interaction always exists in principle for such pairs of molecules, no matter how far they are from each other in space, although the self-interaction decays as $r^{-6}$. If the distance between the two molecules is far away enough such that the self-interaction is much smaller than relevant energy scales in the QLM, $w$, $m$, and $g^2$, then those self-interactions can be neglected. For the $S = 1/2$ QLM, we use a molecular chain with roughly uniform spacing between adjacent molecules (within the order of magnitude), which will be specified later in the numerical details. The second-order self-interaction between next-nearest molecules (such as $S_1$ and $S_2$) is about $w/2^6 = w/64 \ll w$ because of the $r^{-6}$ scaling. Therefore, we can safely neglect self-interactions between next-nearest molecules and those between two molecules that are even farther apart. We only consider self-interactions between the nearest molecules (such as $S_1$ and $L_1$) which are at the order of $w$. 

For our purpose, $\Sigma$ is the coefficient. From the dipole selection rule, $\Sigma$

Therefore, in calculating the denominator in Eq. S29,

TABLE S-1. Energy conditions for the $n$-th unit cell in the $S = 1/2$ QLM, $n = 0, 1, 2, \cdots$. The other molecule states not listed are made off-resonant. $B$ is the molecule’s rotational constant and $h$ is Planck’s constant.

Self-interactions between the molecule $i$ with internal state $\alpha$ and the molecule $j$ with internal state $\beta$ can be denoted as

$$\Sigma_{i,\alpha;j,\beta} \hat{n}_{i,\alpha} \hat{n}_{j,\beta}, \quad (S28)$$

where

$$\Sigma_{i,\alpha;j,\beta} = \sum_{\gamma\neq\alpha, \eta\neq\beta} V_{i,j}^{\alpha,\beta;\gamma,\eta} V_{i,j}^{\gamma,\eta;\alpha,\beta} \frac{1}{\epsilon_{i,\alpha} + \epsilon_{j,\beta} - \epsilon_{i,\gamma} - \epsilon_{j,\eta}} = \sum_{\gamma\neq\alpha, \eta\neq\beta} \left| V_{i,j}^{\alpha,\beta;\gamma,\eta} \right|^2 \frac{1}{\epsilon_{i,\alpha} + \epsilon_{j,\beta} - \epsilon_{i,\gamma} - \epsilon_{j,\eta}} \quad (S29)$$

is the coefficient. From the dipole selection rule, $\Sigma_{i,\alpha;j,\beta} = 0$ if both $\alpha$ and $\beta$ are from $N = 1$, or both are from $N = 0$. For our purpose, $\Sigma_{i,\alpha;j,\beta}$ only needs to be precise at the order of $O(V^2/\Delta)$ and any higher orders can be neglected. Therefore, in calculating the denominator in Eq. S29, $\epsilon_{i,\alpha} + \epsilon_{j,\beta} - \epsilon_{i,\gamma} - \epsilon_{j,\eta}$, corrections at the order of $m, g^2, \Sigma \ll \Delta$ can be neglected, since these corrections when propagated to Eq. S29 are at most at the order of $O(V^3/\Delta^2)$. Note that we are interested in regions of parameter space where $m, g^2$ and $w$ are comparable, and also, $\Sigma \sim O(w)$.

We explain in detail how to suppress the nearest self-interaction for $S = 1/2$. The nearest self-interactions for $S = 1/2$ are products of two number operators, as is shown in Eq. S29. These products of number operators, in the physical Hilbert space, can be rewritten into one-body terms which are just single number operators using the Gauss’ law. Single number operators are one-body potentials and we can compensate them by adding laser light potentials of the opposite values.

The Gauss’ law on the site $S_{2n+2}$, written in terms of molecule number operators, is

$$\hat{n}_{S_{2n+2},a} - \hat{n}_{L_{2n+2},d} + \hat{n}_{L_{2n+1},d} = 0, \quad (S30)$$

where we have used constraints on each molecular position

$$\hat{n}_{S_{2n+2},a} + \hat{n}_{S_{2n+2},b} = 1, \quad (S31)$$

$$\hat{n}_{L_{2n+1},b} + \hat{n}_{L_{2n+1},d} = 1, \quad (S32)$$

$$\hat{n}_{L_{2n+2},b} + \hat{n}_{L_{2n+2},d} = 1. \quad (S33)$$
We can simplify the two-body interaction terms to one-body potentials by using these constraints on number operators. From the Gauss’ law,

\[ \hat{n}_{S_{2n+2},a} - \hat{n}_{L_{2n+2},d} = -\hat{n}_{L_{2n+1},d}, \]  

we square it

\[ \hat{n}^2_{S_{2n+2},a} + \hat{n}^2_{L_{2n+2},d} - 2\hat{n}_{S_{2n+2},a} \hat{n}_{L_{2n+2},d} = \hat{n}^2_{L_{2n+1},d}. \]  

Because \( \hat{n}^2_{i,\alpha} = \hat{n}_{i,\alpha} \) for hard core bosonic states,

\[ \hat{n}_{S_{2n+2},a} + \hat{n}_{L_{2n+2},d} - 2\hat{n}_{S_{2n+2},a} \hat{n}_{L_{2n+2},d} = \hat{n}_{L_{2n+1},d}, \]  

and we arrive at

\[ \hat{n}_{S_{2n+2},a} \hat{n}_{L_{2n+2},d} = \hat{n}_{S_{2n+2},a}. \]  

Similarly,

\[ \hat{n}_{S_{2n+2},a} \hat{n}_{L_{2n+1},d} = 0, \]  

\[ \hat{n}_{S_{2n+2},a} \hat{n}_{L_{2n+2},b} = 0, \]  

\[ \hat{n}_{S_{2n+2},a} \hat{n}_{L_{2n+1},b} = \hat{n}_{S_{2n+2},a}. \]

The Gauss’ law on the site \( S_{2n+1} \), written in terms of molecule number operators, is

\[ \hat{n}_{S_{2n+1},a} - \hat{n}_{L_{2n+1},d} + \hat{n}_{L_{2n},d} = 1, \]  

where we have used constraints on each molecular position

\[ \hat{n}_{S_{2n+1},a} + \hat{n}_{S_{2n+1},b} = 1, \]  

\[ \hat{n}_{L_{2n+1},b} + \hat{n}_{L_{2n+1},d} = 1, \]  

\[ \hat{n}_{L_{2n},b} + \hat{n}_{L_{2n},d} = 1. \]

We can derive that

\[ \hat{n}_{S_{2n+1},a} \hat{n}_{L_{2n+1},d} = \hat{n}_{L_{2n+1},d}, \]  

\[ \hat{n}_{S_{2n+1},a} \hat{n}_{L_{2n},d} = \hat{n}_{L_{2n},d}, \]  

\[ \hat{n}_{S_{2n+1},a} \hat{n}_{L_{2n+1},b} = \hat{n}_{S_{2n+1},a} - \hat{n}_{L_{2n+1},d}, \]  

\[ \hat{n}_{S_{2n+1},a} \hat{n}_{L_{2n},b} = \hat{n}_{S_{2n+1},a} - \hat{n}_{L_{2n},d}. \]

The left-hand sides of Equations (S37) to (S40) and (S45) to (S48) are the only possible combinations of two number operators with nonzero second-order self-interactions. The right-hand sides of Equations (S37) to (S40) and (S45) to (S48) show that the effects of self-interactions are equivalent to one-body terms. These one-body terms from self-interactions are not in the original QLM Hamiltonian, so we want to compensate for them by using laser lights to introduce one-body terms with opposite values.
To know what values should be used to compensate for the self-interactions, we need to calculate \( \Sigma_{i,i;j,i} \). With the general form in Eq. S29, we can plug values of \( V_{i,j}^{\alpha,\beta;\gamma,\eta} \) and \( \epsilon_{i,i} \) to obtain all of the nearest self-interaction coefficients as follows

\[
\Sigma_{L_{2n+1},b;S_{2n+1},a} = \frac{1}{V_{L_{2n+1},S_{2n+1}}^{b,a;b,a}} \frac{1}{\Delta_{1,n} - \delta_{1,n}} \left( \frac{3 \cos^2 \theta_{L_{2n+1},S_{2n+1}} - 1}{2} \right) \left( \frac{1}{3} d^2 \right)^2 \left( \frac{1}{\Delta_{1,n} - \delta_{1,n}} \right)
\]

\( \text{(S49)} \)

\[
\Sigma_{S_{2n+2},d;L_{2n+2},d} = \frac{1}{V_{S_{2n+2},L_{2n+2}}^{d,a;d,a}} \frac{1}{\Delta_{1,n} - \delta_{1,n}} \left( \frac{3 \cos^2 \theta_{L_{2n+2},S_{2n+2}} - 1}{2} \right) \left( \frac{1}{3} d^2 \right)^2 \left( \frac{1}{\Delta_{1,n} - \delta_{1,n}} \right)
\]

\( \text{(S50)} \)

\[
\Sigma_{S_{2n+1},a;L_{2n+1},d} = \frac{1}{V_{S_{2n+1},L_{2n+1}}^{a,d,a,d}} \frac{1}{\Delta_{1,n} - \delta_{1,n}} \left( \frac{3 \cos^2 \theta_{L_{2n+1},S_{2n+1}} - 1}{2} \right) \left( \frac{1}{3} d^2 \right)^2 \left( \frac{1}{\Delta_{1,n} - \delta_{1,n}} \right)
\]

\( \text{(S51)} \)

\[
\Sigma_{L_{2n+2},d;S_{2n+3},a} = \frac{1}{V_{L_{2n+2},S_{2n+3}}^{d,a;d,a}} \frac{1}{\Delta_{1,n} - \delta_{1,n}} \left( \frac{3 \cos^2 \theta_{L_{2n+2},S_{2n+3}} - 1}{2} \right) \left( \frac{1}{3} d^2 \right)^2 \left( \frac{1}{\Delta_{1,n} - \delta_{1,n}} \right)
\]

\( \text{(S52)} \)

\[
\Sigma_{S_{2n+1},a;L_{2n+1},b} = \frac{1}{V_{S_{2n+1},L_{2n+1}}^{a,b,a,b}} \frac{1}{\Delta_{1,n} + \delta_{2,n} - 2\delta_{1,n}} \left( \frac{3 \cos^2 \theta_{L_{2n+2},S_{2n+1}} - 1}{2} \right) \left( \frac{1}{3} d^2 \right)^2 \left( \frac{1}{\Delta_{1,n} + \delta_{2,n} - 2\delta_{1,n}} \right)
\]

\( \text{(S53)} \)

\[
\Sigma_{L_{2n+2},b;S_{2n+3},a} = \frac{1}{V_{L_{2n+2},S_{2n+3}}^{b,a;b,a}} \frac{1}{\Delta_{2,n} - \delta_{2,n}} \left( \frac{3 \cos^2 \theta_{L_{2n+2},S_{2n+3}} - 1}{2} \right) \left( \frac{1}{3} d^2 \right)^2 \left( \frac{1}{\Delta_{2,n} - \delta_{2,n}} \right)
\]

\( \text{(S54)} \)

### Construction of the \( S = 1 \) QLM Hamiltonian

The \( S = 1 \) quantum link model is

\[
H = -w \sum_x \left[ b_{x,a}^\dagger b_{x,a} S^+_{x,x+1} b_{x+1,a}^\dagger b_{x+1,a} + b_{x+1,a}^\dagger b_{x+1,a} S^-_{x,x+1} b_{x,a}^\dagger b_{x,a} \right] + m \sum_x (-1)^x \frac{1}{2} \left( b_{x,a}^\dagger b_{x,a} - b_{x,a}^\dagger b_{x,a} + 1 \right) + \frac{g^2}{2} \sum_x \left( S^3_{x,x+1} \right)^2
\]

\( \text{(S55)} \)
where the $S = 1$ spin operators are

$$S^+_{x,x+1} = \sqrt{2} \left( b^\dagger_{(x,x+1),a} b_{(x,x+1),a} + b^\dagger_{(x,x+1),b} b_{(x,x+1),b} \right),$$  \hspace{1cm} (S56)$$

$$S^-_{x,x+1} = \sqrt{2} \left( b^\dagger_{(x,x+1),b} b_{(x,x+1),c} + b^\dagger_{(x,x+1),c} b_{(x,x+1),b} \right),$$  \hspace{1cm} (S57)$$

$$S^3_{x,x+1} = b^\dagger_{(x,x+1),c} b_{(x,x+1),c} - b^\dagger_{(x,x+1),d} b_{(x,x+1),d}.$$  \hspace{1cm} (S58)$$

The energy conditions are listed in Table S-2. Similar to $S = 1/2$, $\Delta_{1,n}$, $\Delta_{2,n}$, $\delta_{1,n}$, $\delta_{2,n}$ for any $n$ are at the order of $\Delta$ mentioned in the main text and will be specified in numerical simulations.

| Molecule State Energy | Value |
|-----------------------|-------|
| $\epsilon S_{2n+1,a}$ | $-m - \sum_{S_{2n+1,a}} L_{2n+1,b}$ |
| $\epsilon S_{2n+1,c}$ | $2hB + \delta_{1,n}$ |
| $\epsilon L_{2n+1,a}$ | $0$ |
| $\epsilon L_{2n+1,c}$ | $2hB + \Delta_{1,n} - (\delta_{2,n} - \delta_{1,n}) + g^2/2$ |
| $\epsilon L_{2n+1,0}$ | $2hB + \Delta_{1,n}$ |
| $\epsilon L_{2n+1,-1}$ | $2hB + \Delta_{1,n} + (\delta_{2,n} - \delta_{1,n}) + g^2/2$ |
| $\epsilon S_{2n+2,a}$ | $m$ |
| $\epsilon S_{2n+2,c}$ | $2hB + \delta_{2,n}$ |
| $\epsilon L_{2n+2,a}$ | $\sum_{S_{2n+2,a}} L_{2n+2,b}$ |
| $\epsilon L_{2n+2,c}$ | $2hB + \Delta_{2,n} - (\delta_{1,n+1} - \delta_{2,n}) + g^2/2 + \sum_{S_{2n+2,a}} L_{2n+2,d} - \sum_{S_{2n+2,a}} L_{2n+2,b}$ |
| $\epsilon L_{2n+2,0}$ | $2hB + \Delta_{2,n}$ |
| $\epsilon L_{2n+2,-1}$ | $2hB + \Delta_{2,n} + (\delta_{1,n+1} - \delta_{2,n}) + g^2/2 - \sum_{S_{2n+3,a}} L_{2n+3,d} + \sum_{S_{2n+3,a}} L_{2n+3,b}$ |

TABLE S-2. Energy conditions for the $n$-th unit cell the $S = 1$ QLM, $n = 0, 1, 2, \ldots$. The other molecule states not listed are made off-resonant. $B$ is the molecule’s rotational constant and $h$ is Planck’s constant.

The nearest-neighbor self-interactions with two number operators from the second-order effective Hamiltonian are the total sum of the following terms

$$\sum_{S_{2n+1,a}; L_{2n+1,c}, \hat{n}_{S_{2n+1,a}} \hat{n}_{L_{2n+1,c}}} + \sum_{S_{2n+1,a}; L_{2n+1,b}, \hat{n}_{S_{2n+1,a}} \hat{n}_{L_{2n+1,b}}} + \sum_{S_{2n+1,a}; L_{2n+1,d}, \hat{n}_{S_{2n+1,a}} \hat{n}_{L_{2n+1,d}}} + \sum_{S_{2n+1,a}; L_{2n+1,c}, \hat{n}_{S_{2n+1,a}} \hat{n}_{L_{2n+1,c}}} + \sum_{S_{2n+1,a}; L_{2n+1,b}, \hat{n}_{S_{2n+1,a}} \hat{n}_{L_{2n+1,b}}} + \sum_{S_{2n+1,a}; L_{2n+1,d}, \hat{n}_{S_{2n+1,a}} \hat{n}_{L_{2n+1,d}}}$$

$$= \left( \sum_{S_{2n+1,a}; L_{2n+1,c}, \hat{n}_{S_{2n+1,a}} \hat{n}_{L_{2n+1,c}}} - 2\sum_{S_{2n+1,a}; L_{2n+1,b}, \hat{n}_{S_{2n+1,a}} \hat{n}_{L_{2n+1,b}}} + \sum_{S_{2n+1,a}; L_{2n+1,d}, \hat{n}_{S_{2n+1,a}} \hat{n}_{L_{2n+1,d}}} \right) \hat{n}_{S_{2n+1,a}} \hat{n}_{L_{2n+1,c}}$$

$$+ \left( \sum_{S_{2n+1,a}; L_{2n+1,c}, \hat{n}_{S_{2n+1,a}} \hat{n}_{L_{2n+1,c}}} - 2\sum_{S_{2n+1,a}; L_{2n+1,b}, \hat{n}_{S_{2n+1,a}} \hat{n}_{L_{2n+1,b}}} + \sum_{S_{2n+1,a}; L_{2n+1,d}, \hat{n}_{S_{2n+1,a}} \hat{n}_{L_{2n+1,d}}} \right) \hat{n}_{S_{2n+1,a}} \hat{n}_{L_{2n+1,b}}$$

$$+ \left( \sum_{S_{2n+1,a}; L_{2n+1,c}, \hat{n}_{S_{2n+1,a}} \hat{n}_{L_{2n+1,c}}} - 2\sum_{S_{2n+1,a}; L_{2n+1,b}, \hat{n}_{S_{2n+1,a}} \hat{n}_{L_{2n+1,b}}} + \sum_{S_{2n+1,a}; L_{2n+1,d}, \hat{n}_{S_{2n+1,a}} \hat{n}_{L_{2n+1,d}}} \right) \hat{n}_{S_{2n+1,a}} \hat{n}_{L_{2n+1,d}}$$

$$= \left( \sum_{S_{2n+1,a}; L_{2n+1,c}, \hat{n}_{S_{2n+1,a}} \hat{n}_{L_{2n+1,c}}} - 2\sum_{S_{2n+1,a}; L_{2n+1,b}, \hat{n}_{S_{2n+1,a}} \hat{n}_{L_{2n+1,b}}} + \sum_{S_{2n+1,a}; L_{2n+1,d}, \hat{n}_{S_{2n+1,a}} \hat{n}_{L_{2n+1,d}}} \right) \hat{n}_{S_{2n+1,a}} \hat{n}_{L_{2n+1,c}}$$

$$+ \left( \sum_{S_{2n+1,a}; L_{2n+1,c}, \hat{n}_{S_{2n+1,a}} \hat{n}_{L_{2n+1,c}}} - 2\sum_{S_{2n+1,a}; L_{2n+1,b}, \hat{n}_{S_{2n+1,a}} \hat{n}_{L_{2n+1,b}}} + \sum_{S_{2n+1,a}; L_{2n+1,d}, \hat{n}_{S_{2n+1,a}} \hat{n}_{L_{2n+1,d}}} \right) \hat{n}_{S_{2n+1,a}} \hat{n}_{L_{2n+1,b}}$$

$$+ \left( \sum_{S_{2n+1,a}; L_{2n+1,c}, \hat{n}_{S_{2n+1,a}} \hat{n}_{L_{2n+1,c}}} - 2\sum_{S_{2n+1,a}; L_{2n+1,b}, \hat{n}_{S_{2n+1,a}} \hat{n}_{L_{2n+1,b}}} + \sum_{S_{2n+1,a}; L_{2n+1,d}, \hat{n}_{S_{2n+1,a}} \hat{n}_{L_{2n+1,d}}} \right) \hat{n}_{S_{2n+1,a}} \hat{n}_{L_{2n+1,d}}$$

$$= \left( \sum_{S_{2n+1,a}; L_{2n+1,c}, \hat{n}_{S_{2n+1,a}} \hat{n}_{L_{2n+1,c}}} - 2\sum_{S_{2n+1,a}; L_{2n+1,b}, \hat{n}_{S_{2n+1,a}} \hat{n}_{L_{2n+1,b}}} + \sum_{S_{2n+1,a}; L_{2n+1,d}, \hat{n}_{S_{2n+1,a}} \hat{n}_{L_{2n+1,d}}} \right) \hat{n}_{S_{2n+1,a}} \hat{n}_{L_{2n+1,c}}$$

$$+ \left( \sum_{S_{2n+1,a}; L_{2n+1,c}, \hat{n}_{S_{2n+1,a}} \hat{n}_{L_{2n+1,c}}} - 2\sum_{S_{2n+1,a}; L_{2n+1,b}, \hat{n}_{S_{2n+1,a}} \hat{n}_{L_{2n+1,b}}} + \sum_{S_{2n+1,a}; L_{2n+1,d}, \hat{n}_{S_{2n+1,a}} \hat{n}_{L_{2n+1,d}}} \right) \hat{n}_{S_{2n+1,a}} \hat{n}_{L_{2n+1,b}}$$

$$+ \left( \sum_{S_{2n+1,a}; L_{2n+1,c}, \hat{n}_{S_{2n+1,a}} \hat{n}_{L_{2n+1,c}}} - 2\sum_{S_{2n+1,a}; L_{2n+1,b}, \hat{n}_{S_{2n+1,a}} \hat{n}_{L_{2n+1,b}}} + \sum_{S_{2n+1,a}; L_{2n+1,d}, \hat{n}_{S_{2n+1,a}} \hat{n}_{L_{2n+1,d}}} \right) \hat{n}_{S_{2n+1,a}} \hat{n}_{L_{2n+1,d}}$$
\[
\sum S_{2n+2,aL2n+2,c}c \hat{n}s_{2n+2,a} \hat{n}s_{2n+2,c} + \sum S_{2n+2,aL2n+2,b}b \hat{s}s_{2n+2,b} \hat{s}s_{2n+2,b} + \sum S_{2n+2,aL2n+2,d}d \hat{s}s_{2n+2,d} \hat{s}s_{2n+2,d} \\
= (\sum S_{2n+2,aL2n+2,c}c - 2\sum S_{2n+2,aL2n+2,b}b + \sum S_{2n+2,aL2n+2,d}d) \hat{s}s_{2n+2,a} \hat{s}s_{2n+2,c} \\
+ (\sum S_{2n+2,aL2n+2,d}d - \sum S_{2n+2,aL2n+2,b}b) (-\hat{n}s_{2n+2,c} + \hat{n}s_{2n+1,c}) + \sum S_{2n+2,aL2n+2,d}d \hat{s}s_{2n+2,a},
\]

where the Gauss’ law and other molecular number constraints are already used.

We want the coefficients in front of the two-number operators to vanish, so there are four independent equations for each unit cell generated from this, shown as follow

\[
\sum S_{2n+1,aL2n+1,c}c - 2\sum S_{2n+1,aL2n+1,b}b + \sum S_{2n+1,aL2n+1,d}d = 0, 
\]

(63)

\[
\sum S_{2n+1,aL2n,c}c - 2\sum S_{2n+1,aL2n,b}b + \sum S_{2n+1,aL2n,d}d = 0, 
\]

(64)

\[
\sum S_{2n+2,aL2n+1,c}c - 2\sum S_{2n+2,aL2n+1,b}b + \sum S_{2n+2,aL2n+1,d}d = 0, 
\]

(65)

\[
\sum S_{2n+2,aL2n+2,c}c - 2\sum S_{2n+2,aL2n+2,b}b + \sum S_{2n+2,aL2n+2,d}d = 0. 
\]

(66)

Equations (63) to (66) are over-determined as equations for distances \(r\)'s, energy parameters \(\delta\)'s and \(\Delta\)'s, and the angle \(\theta\)'s. As is explained in the main text, we have introduced nonequal inter-molecular separations such that the self-interactions for molecular pairs \(L_x\) and \(S_x\) are negligible. Therefore, Equations (64) and (65) are no longer needed. The remaining equations are under-determined. Equations (63) and (66) with \(\Sigma\)'s plugged in are

\[
\frac{1}{\Delta_{1,n} - \delta_{2,n}} \left| - \frac{1}{4\pi\epsilon_0 r_{S_{2n+1},L_{2n+1}}} (3\cos^2 \theta_{S_{2n+1},L_{2n+1}} - 1) \frac{1}{3} d^2 \right|^2 
\]

\[
- 2 \frac{1}{\Delta_{1,n} - \delta_{1,n}} \left| - \frac{1}{4\pi\epsilon_0 r_{S_{2n+1},L_{2n+1}}} e^{-i\phi_{S_{2n+1},L_{2n+1}}} \sin \theta_{S_{2n+1},L_{2n+1}} \cos \theta_{S_{2n+1},L_{2n+1}} \frac{1}{\sqrt{2}} d^2 \right|^2 
\]

(67)

\[
+ \frac{1}{\Delta_{1,n} + \delta_{2,n} - 2\delta_{1,n}} \left| \frac{1}{4\pi\epsilon_0 r_{S_{2n+1},L_{2n+1}}} e^{-i\phi_{S_{2n+1},L_{2n+1}}} \sin \theta_{S_{2n+1},L_{2n+1}} \cos \theta_{S_{2n+1},L_{2n+1}} \frac{1}{\sqrt{2}} d^2 \right|^2 = 0, 
\]

\[
\frac{1}{\Delta_{2,n} - \delta_{1,n+1}} \left| - \frac{1}{4\pi\epsilon_0 r_{S_{2n},L_{2n}}} (3\cos^2 \theta_{S_{2n},L_{2n}} - 1) \frac{1}{3} d^2 \right|^2 
\]

\[
- 2 \frac{1}{\Delta_{2,n} - \delta_{2,n}} \left| - \frac{1}{4\pi\epsilon_0 r_{S_{2n},L_{2n}}} e^{-i\phi_{S_{2n},L_{2n}}} \sin \theta_{S_{2n},L_{2n}} \cos \theta_{S_{2n},L_{2n}} \frac{1}{\sqrt{2}} d^2 \right|^2 
\]

(68)

\[
+ \frac{1}{\Delta_{2,n} + \delta_{1,n+1} - 2\delta_{2,n}} \left| \frac{1}{4\pi\epsilon_0 r_{S_{2n},L_{2n}}} e^{-i\phi_{S_{2n},L_{2n}}} \sin \theta_{S_{2n},L_{2n}} \cos \theta_{S_{2n},L_{2n}} \frac{1}{\sqrt{2}} d^2 \right|^2 = 0, 
\]

After a bit algebra from the equations, we obtain the constraints

\[
\Delta_{1,n} = \frac{1}{2} (3\delta_{1,n} - \delta_{2,n}), 
\]

(69)

\[
\Delta_{2,n} = \frac{1}{2} (3\delta_{2,n} - \delta_{1,n+1}). 
\]

(70)

From further calculation, there is constraints on angles

\[
\frac{1}{\sqrt{2}} \left| \sin \theta_{S_{2n+1},L_{2n+1}} \cos \theta_{S_{2n+1},L_{2n+1}} \right| = \frac{1}{3} \left| \frac{3\cos^2 \theta_{S_{2n+1},L_{2n+1}} - 1}{3} \right|, 
\]

(71)

\[
\frac{1}{\sqrt{2}} \left| \sin \theta_{S_{2n+2},L_{2n+2}} \cos \theta_{S_{2n+2},L_{2n+2}} \right| = \frac{1}{3} \left| \frac{3\cos^2 \theta_{S_{2n+2},L_{2n+2}} - 1}{3} \right|, 
\]

(72)
which gives

$$\cos^2 \theta_{S_{2n+1},L_{2n+1}} = 0.0220216 \text{ or } 0.917372, \quad (S73)$$

$$\cos^2 \theta_{S_{2n+2},L_{2n+2}} = 0.0220216 \text{ or } 0.917372. \quad (S74)$$

In addition to the solutions above, we also need to impose the condition that tunneling amplitudes $-\sqrt{2}w$ at all positions should be equal, which yields some constraints on the distances $r$'s and angles $\theta$'s. In principle, the chain can be zig-zag. However, in our scheme, we specifically set the chain of molecules to be a straight line. Namely, all of the angles $\theta$'s are the same.

### Details on Numerical Methods and Simulation

We have implemented the Exact Diagonalization (ED) Method to simulate the time evolution of QLMs and the DMH for three unit cells. This section provides the details for the numerical algorithm and the choices of parameters in the simulation.

#### Construction of the Hilbert Spaces

For QLMs, each unit cell has two sites and links $S_{2n+1}, L_{2n+1}, S_{2n+2}, L_{2n+2}$ where each site has 2 degrees of freedom and each link has $2S+1$ degrees of freedom for spin-$S$. Therefore, for $N$ unit cells ($n = 0, 1, \cdots, N-1$), the Hilbert space should have the dimension $(4S+2)^2N$. For the DMH, each unit cell has 4 molecules and each molecule has 4 degrees of freedom. It follows that for $N$ unit cells, the dimension of the Hilbert space is $2^{8N}$. It is clear that the DMH Hilbert space dimension is the bottleneck of the ED method when $N$ is large.

To implement the ED method, we need to reduce the dimensionality of both the DMH and QLM Hilbert spaces while preserving the accuracy of the simulation. To achieve that, we utilize symmetries and quantum numbers in both the DMH and the QLM Hamiltonian. Because of the dipole selection rules, the number of molecules at the state $a$ is exactly conserved and thus a good quantum number. In addition, we can make certain states off-resonant by tuning the laser light. For $S = 1/2$, states $c, d$ on fermion sites and the state $c$ on link sites are off-resonant while for $S = 1$, states $b, d$ on fermion sites are off-resonant. Therefore, those states will be excluded when we construct the DMH Hilbert space. Since the last link $L_{2N}$ never changes its state in QLMs due to open boundary conditions, we fix the state of the molecule which represent $L_{2N}$ to its initial state throughout the simulation for DMH. For the QLM Hamiltonian, the total number of fermions is conserved and we can use this fact to reduce the dimensionality of the QLM Hilbert space. Similar to the DMH, the state of the last link $L_{2N}$ is fixed.

According to the above construction of DMH and QLM Hamiltonian, the DMH Hilbert space is larger than the QLM Hilbert space. The QLM wavefunction can be embedded into the DMH Hilbert space while the DMH wavefunction needs to be truncated when it is projected onto the QLM Hilbert space, where the mapping is provided in Table I in the main text. In this paper, fidelity is computed with the embedded QLM wavefunction in the DMH Hilbert space and DMH observables are computed with the DMH wavefunction which is projected onto the QLM Hilbert space. Since our constructed QLM Hilbert space, onto which the DMH wavefunction is projected, is larger than the physical QLM Hilbert space, we will be able to see slight Gauss’ law violation in DMH simulations. In experiments, to measure the Gauss’ law, one can post-select the measurement outcomes of the DMH from our constructed QLM Hilbert space. The Gauss’ law plots presented in the main text are computed in the same way.

#### Choices of Distances and Energy Conditions

The assumed minimum molecule spacing in experiments is 0.5 $\mu$m. In this section, all of the energies are divided by Planck’s constant $\hbar$ and are thus in units of Hz. With the following specified inter-molecular distances and energy conditions, hopping parameters are $w = 82.7$ Hz for $S = 1/2$ and $w = 3.17$ Hz for $S = 1$. We note that these choices have yielded decent fidelity overlap with the QLM dynamics (excluding the influence of the “extra” gauge invariant terms in the $S = 1$ case), and larger hopping energies can be achieved if sources of dephasing, decoherence, or parameter control disorder serve as practical limitations.
TABLE S-3. Energy conditions used in simulations for the n-th unit cell in the $S = 1/2$ QLM, $n = 0, 1, 2, \cdots$. The other molecule states not listed are made off-resonant. $B$ is the molecule’s rotational constant and $h$ is Planck’s constant.

| Molecule State | Energy |
|----------------|--------|
| $\epsilon_{S_{2n+1},a}$ | $-m - \sum_{k=1}^{S_{2n+1},a} k_{2n+1},b - \sum_{k=2}^{S_{2n+3},a} k_{2n+2},d$ |
| $\epsilon_{S_{2n+1},b}$ | $2hB + \delta_{1,n}$ |
| $\epsilon_{L_{2n+1},a}$ | $0$ |
| $\epsilon_{L_{2n+1},d}$ | $-\sum_{k=1}^{S_{2n+1},a} k_{2n+1},d - \sum_{k=2}^{S_{2n+3},a} k_{2n+1},b + \sum_{k=2}^{S_{2n+3},a} k_{2n+2},d$ |
| $\epsilon_{L_{2n+1},b}$ | $2hB + (\delta_{1,n}/2 + \delta_{1,n}/2) + g^2/2$ |
| $\epsilon_{S_{2n+2},a}$ | $m - \sum_{k=1}^{S_{2n+1},b} k_{2n+2},a - \sum_{k=2}^{S_{2n+3},a} k_{2n+2},d$ |
| $\epsilon_{S_{2n+2},b}$ | $2hB + \delta_{2,n}$ |
| $\epsilon_{L_{2n+2},a}$ | $0$ |
| $\epsilon_{L_{2n+2},d}$ | $2hB + (3\delta_{2,n}/2 - \delta_{1,n+1}/2) + g^2/2$ |
| $\epsilon_{S_{2n+2},b}$ | $2hB + (\delta_{1,n+1}/2 + \delta_{2,n}/2) + g^2/2$ |

Since the energy conditions of $\Delta_{1,n}$ and $\Delta_{2,n}$ for $S = 1/2$ are flexible, for a particular realization of energy conditions for $S = 1/2$, Eq. S69 and Eq. S70 for $S = 1$ are also applicable and can be substituted into Table S-1, which gives rise to Table S-3. For $S = 1$, Eq. S69 and Eq. S70 are substituted into Table S-2 to obtain Table S-4.

For both $S = 1/2$ and $S = 1$, we construct two arithmetic sequences $\delta_{1,n}$, $\delta_{2,n}$ by $\delta_{1,n} = \delta_{1,0} - D_1$ and $\delta_{1,n+1} = \delta_{1,n} - D_2$ to avoid accidental energy degeneracy between two states in different unit cells and thus suppress the first-order interactions in the effective Hamiltonian. We define $V_0 = \frac{1}{4\pi \alpha_0 r^3_{S,1,L_1}} d^2$, where $d$ is the electric dipole moment of one molecule. In particular, we choose $\delta_{1,0} = 12.5 V_0$, $D_1 = 10 V_0$, $D_2 = 70 V_0$, $B = 1000 V_0$.

For $S = 1/2$, in the main text,

$$-w = \frac{1}{2} \sum_{b,a,d} \sum_{k=1}^{S_{2n+1},a} \left[ \frac{1}{\Delta \epsilon_{1,x}} + \frac{1}{\Delta \epsilon_{2,x}} \right],$$

with $\Delta \epsilon_{1,x} = \epsilon_{S_{2n+1},a} + \epsilon_{L_{2n+1},d} - \epsilon_{S_{2n+2},b} - \epsilon_{L_{2n+2},a}$ and $\Delta \epsilon_{2,x} = \epsilon_{S_{2n+1},a} + \epsilon_{L_{2n+1},d} - \epsilon_{S_{2n+2},b} - \epsilon_{L_{2n+2},a}$. $\Delta \epsilon_{1,x}$ and $\Delta \epsilon_{2,x}$ have been specified by energy conditions. The right-hand sides of Eq. S75 at different $x$’s are required to be the same because the hopping parameter $w$ does not depend on $x$. In order to achieve that, we need to alter the dipole-dipole interactions by tuning the inter-molecular distances. Similar tuning needs to be done for $S = 1$ as well for the same reason. In particular, for both $S = 1/2$ and $S = 1$, we set the relative distance ratios as $r_{S_{2n+1},L_{2n+1}} = r_{S_{1,L_1}}$, $r_{S_{2n+2},L_{2n+2}} = \gamma r_{S_{1,L_1}}$, $r_{S_{2n+3},L_{2n+2}} = \gamma r_{S_{1,L_1}}$, $r_{L_{2n+2},a} = \beta \gamma r_{S_{1,L_1}}$ for every $n$, where $\beta = (D_1/D_2)^{1/6} \approx 0.723$ is fixed by energy conditions and $\gamma$ is defined as a variable long-short distance ratio greater than or equal to one, mentioned in the main text. $r_{S_{2n+2},L_{2n+2}}$ are the smallest inter-molecular distances, which we assume can be set to 0.5 $\mu$m in an envisioned experiment which corresponds to $r_{S_{1,L_1}} = 0.692$ $\mu$m. The $\theta$ angles for all positions are set to be the same with all the molecules on a line. For $S = 1/2$, we choose $\cos \theta = 0$. For $S = 1$, from Eq. S73 and Eq. S74, we choose $\cos \theta = 0.1485$. The $\phi$ angles are all the same with a line of molecules and are set to 0 for both $S = 1/2$ and $S = 1$. They are furthermore irrelevant for one-dimensional models with only local hopping terms. The body-frame electric dipole moment of the ground state NaRb molecule is $d = 3.3$ D (Debye) [52]. In the considered experimental setup, $V_0 = 4.96$ kHz when $r_{S_{1,L_1}} = 0.692$ $\mu$m.

In the following, we compute the hopping parameter $w$ with the above energy conditions and with experimentally reasonable parameters. For $S = 1/2$, we set $\gamma = 1$. Since $w$ is the same for every position, we can just compute its value at one position

$$w = \left( \frac{\delta_{1,n} - \delta_{2,n}}{2} \right)^{-1} \sum_{b,a,d} \sum_{k=1}^{S_{2n+1},a} \left( \frac{1}{\delta_{1,n} - \delta_{2,n}} V_0^2 \sin^2 \theta \beta \cos^2 \theta - \frac{1}{6} \right).$$
Molecule State Energy | Value
---|---
$\epsilon_{S_{2n+1}^a}$ | $-m - \Sigma S_{2n+1}^a L_{2n+1}^b$
$\epsilon_{S_{2n+1}^c}$ | $2hB + \delta_{1,n}$

$\epsilon_{L_{2n+1}^a}$ | $0$
$\epsilon_{L_{2n+1}^c}$ | $2hB + (5\delta_{1,n}/2 - 3\delta_{2,n}/2) + g^2/2$
$\epsilon_{L_{2n+1}^b}$ | $\Sigma S_{2n+1}^a L_{2n+1}^b - \Sigma S_{2n+2}^a L_{2n+2}^b$
$\epsilon_{L_{2n+1}^d}$ | $2hB + (3\delta_{1,n}/2 - \delta_{2,n}/2)$

$\epsilon_{S_{2n+2}^a}$ | $m$
$\epsilon_{S_{2n+2}^c}$ | $2hB + \delta_{2,n}$

$\epsilon_{S_{2n+2}^a}$ | $\Sigma S_{2n+2}^a L_{2n+2}^b$
$\epsilon_{S_{2n+2}^c}$ | $2hB + (5\delta_{2,n}/2 - 3\delta_{1,n+1}/2) + g^2/2 + \Sigma S_{2n+2}^a L_{2n+2}^b - \Sigma S_{2n+2}^a L_{2n+2}^b$
$\epsilon_{S_{2n+2}^b}$ | $2hB + (3\delta_{2,n}/2 - \delta_{1,n+1}/2)$
$\epsilon_{S_{2n+2}^d}$ | $2hB + (\delta_{2,n}/2 + \delta_{1,n}/2 + g^2/2 - \Sigma S_{2n+2}^a L_{2n+2}^b + \Sigma S_{2n+2}^a L_{2n+2}^b$

TABLE S-4. Energy conditions used in simulations for the n-th unit cell the $S = 1$ QLM, $n = 0, 1, 2, \cdots$. The other molecule states not listed are made off-resonant. $B$ is the molecule’s rotational constant and $h$ is Planck’s constant.

where as described we have chosen $\delta_{1,n} - \delta_{2,n} = D_1$. From our chosen parameters, it follows that $w = V_0/60 = 82.7$ Hz. For $S = 1$, we calculate

$$-\sqrt{2}w = \left(\frac{3(\delta_{1,n} - \delta_{2,n})}{2}\right)^{-1} \left(\frac{1}{4\pi \epsilon_0 r_1^2 d^2} \frac{3 \cos^2 \theta - 1}{3} \right) \left(\frac{1}{4\pi \epsilon_0 r_1^2 d^2 \delta_{1,n}} \frac{1}{\sqrt{2}} \sin \theta \cos \theta \right)$$

(S77)

We choose the long-short distance ratio $\gamma = 1.5$ in the main text. From our chosen parameters, it follows that $\sqrt{2}w = 3.17$ Hz. We note that this rather small energy scale may be practically challenged by both dephasing and parameter control disorder, and larger values can be achieved by relaxing some of the assumed energy constraint conditions.

**Effects of long-short distances for $S = 1$**

For $S = 1$, we have investigated the effect of long-short distance ratio $\gamma$ in fidelity. Fig. S2 demonstrates the average fidelity up to time $t = 20(\sqrt{2}w)^{-1}$ at various long-short distance ratios for $m = 0.25 \times \sqrt{2}w$ and $m = 2.0 \times \sqrt{2}w$. It is shown that the fidelity is low for both small and large $\gamma$, while the optimal value is achieved at an intermediate ratio between 2 and 3. For small long-short distance ratios, certain second-order self-interactions at the order $O(\sqrt{2}w/\Delta)$ are not sufficiently suppressed, which results in low fidelity. For large long-short distance ratios, higher-order self-interaction terms (such as fourth-order self-interactions at the order $O(V_{\text{short}}^4/\Delta^3)$ for the pair of molecules $S_x$ and $L_x$ separated by a short distance) may become comparable to the hopping parameter $w \sim O(V_{\text{short}} V_{\text{long}}/\Delta)$ and are thus not negligible. Therefore, the optimal choice of $\gamma$ for fidelity should balance the trade-off between suppressing second-order self-interaction terms at the order $O(V_{\text{long}}^2/\Delta)$ and avoiding high-order self-interactions, which is attained in the intermediate ratio. The effect of long-short distance ratios is further presented in (a) and (b) in Fig. S3 for real time fidelity changes.
Figure S2. Average fidelity of the calculated DHM dynamics based on the mapping to the $S = 1$ QLM, plotted versus the “long-short” distance ratio used to mitigate the influence of “extra” gauge-invariant terms. The plotted curves are for mass values of $m = 0.25 \times \sqrt{2}w$ and $m = 2.0 \times \sqrt{2}w$. These represent the fidelity when starting from the initial product state as in Fig. 3 of the main text, comparing the DMH-evolved state to the evolution under the ideal QLM, with averaging over the time period $0 \leq t \leq 20$ (with time in units of $(\sqrt{2}w)^{-1}$). An initial rise in fidelity is seen for moderate ratios as the “extra” gauge invariant terms are suppressed, but it decreases for larger ratios as higher-order terms become important.

The long-short distance ratios that provide the highest fidelity (as compared to the target QLM) may be less ideal from a practical perspective, as they result in lower hopping parameters $w$ for a fixed minimum separation of molecules. However, as discussed in the text, the reduction of the fidelity due to the “extra” gauge invariant terms in the $S = 1$ case does not necessarily preclude the DMH dynamics from displaying the physical processes of interest. In our case, at a sub-optimal long-short distance ratio, even though the fidelity may not be very high, the observed phenomenology is not significantly altered by the extra gauge invariant terms. In Fig. S3 (c) and (d), we compare the sum of electric fluxes, an indicator of string breaking, over various long-short distance ratios. Although different long-short distance ratios have very different fidelities, they result in similar string breaking phenomena except for $\gamma = 1.0$. Qualitatively, at all of the ratios except for $\gamma = 1.0$, our results at $m = 0.25 \times \sqrt{2}w$ reveal string breaking while the results at $m = 2.0 \times \sqrt{2}w$ do not. This provides nice evidence that we may be able to choose sub-optimal (in terms of fidelity with ideal QLM) but experimentally favorable parameters to understand the physics of interest.
Figure S3. Effect of additional gauge-invariant terms generated by the dipolar molecular system for simulation of the $S = 1$ QLM. This figure considers the influence of these terms on the fidelity (when comparing the DMH simulations to the ideal QLM) and the dynamics of key physical observables. (a,b) Fidelity versus time for (a) $m = 0.25 \times \sqrt{2} w$ and (b) $m = 2.0 \times \sqrt{2} w$ with various long-short distance ratios. (c,d) The sum of electric fields for (c) $m = 0.25 \times \sqrt{2} w$ and (d) $m = 2.0 \times \sqrt{2} w$ with various long-short distance ratios. For all panels, time is in units of $(\sqrt{2} w)^{-1}$. While the fidelity (as compared to the ideal QLM) is strongly dependent on the ratio value, the physical observations of string-breaking in (c) and a fixed string with small fluctuations in (d) are robust over a very large range of ratio values.