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Symmetry breaking and localization in a random Schwinger model with commensuration

A. A. Akhtar,1 Rahul M. Nandkishore,2 and S. L. Sondhi3

1Department of Physics, University of California at San Diego, La Jolla, CA 92093, USA
2Department of Physics and Center for Theory of Quantum Matter, University of Colorado at Boulder, Boulder CO 80309, USA
3Department of Physics, Princeton University, Princeton, NJ 08544, USA

We numerically investigate a lattice regularized version of quantum electrodynamics in one spatial dimension (Schwinger model). We work at a density where lattice commensuration effects are important, and preclude analytic solution of the problem by bosonization. We therefore numerically investigate the interplay of confinement, lattice commensuration, and disorder, in the form of a random chemical potential. We begin by pointing out that the ground state at commensurate filling spontaneously breaks the translational symmetry of the lattice. This feature is absent in the conventional lattice regularization, which breaks the relevant symmetry explicitly, but is present in an alternative (symmetric) regularization that we introduce. Remarkably, the spontaneous symmetry breaking survives the addition of a random chemical potential (which explicitly breaks the relevant symmetry) in apparent contradiction of the Imry-Ma theorem, which forbids symmetry breaking in one dimension with this kind of disorder. We identify the long range interaction as the key ingredient enabling the system to evade Imry-Ma constraints. We examine spatially resolved energy level statistics for the disordered system, which suggest that the low energy Hilbert space exhibits ergodicity breaking, with level statistics that fail to follow random matrix theory. A careful examination of the structure of the first excited state reveals that disorder induced localization is responsible for the deviations from random matrix theory, and further reveals that the elementary excitations are charge neutral, and therefore not long range interacting.

Quantum electrodynamics in one spatial dimension (also known as the ‘Schwinger model’) has long been an important platform for investigating confinement1–4. More recently, this model has also proven important in connection with many body localization (MBL), a phenomenon whereby disordered quantum systems can exhibit ergodicity breaking, failing to come to equilibrium even at infinite times5–12. In particular, while MBL traditionally occurs in systems with purely short range interactions, and indeed there are arguments to the effect that long range interactions are incompatible with MBL13,14, Ref.15 argued that the interplay of confinement and disorder would cause the disordered Schwinger model to exhibit MBL, despite the long range interaction. Ref.16 further pointed out that the ground state of this model realized an elusive and long sought after phase known as the Mott glass17,18

The discussion of the interplay of disorder and confinement in Refs.15 and 16 was formulated in the continuum. MBL in the continuum is a fairly delicate phenomenon19–21, and a lattice formulation of the argument is greatly desired. A lattice formulation is moreover essential if the scenario of Ref.15 is to be subjected to numerical tests. At the same time, a reformulation of the problem on a periodic lattice brings in the possibility of commensuration effects whereby electrons can scatter freely by reciprocal lattice vectors - and these are entirely absent in the continuum analysis. The interplay of confinement, disorder, and commensuration in lattice regularized versions of the Schwinger model is thus an important open problem.

In this work, we investigate the interplay of confinement, disorder, and lattice commensuration effects in a disordered, lattice regularized Schwinger model, using numerical exact diagonalization (ED) and density matrix renormalization group (DMRG) techniques. The presence of strong lattice commensuration precludes analytic solution of the problem by bosonization, as in Ref. 15, and necessitates a numerical approach. We discover that the ground state of the lattice Schwinger model at commensurate filling spontaneously breaks translation symmetry. This feature is absent in the conventional Kogut-Susskind22 lattice regularization of the Schwinger model, which breaks translation symmetry explicitly. It is however crucial in an alternative (fully symmetric) regularization that we introduce. We demonstrate that the broken symmetry phase survives the addition of a random chemical potential, which itself explicitly breaks lattice translation symmetry (‘random field disorder’ in the statistical mechanics jargon). This result flies in the face of the Imry-Ma theorem23, which states that symmetry breaking in one spatial dimension is unstable to random field disorder. We identify the long range interaction as the key ingredient enabling the system to evade Imry-Ma constraints. We then examine the energy resolved level statistics and hence show that the low energy Hilbert space exhibits ergodicity breaking. While ergodicity breaking could come from either integrability or localization, a careful DMRG based examination of the structure of the first excited state indeed reveals disorder induced localization, as was predicted (in the absence of commensuration) by Ref. 15. The DMRG analysis also reveals that the elementary excitations are charge neutral on average, and therefore lack long range interactions.
This paper is structured as follows. In Sec.I we discuss the regularization of the Schwinger model on a lattice. In Sec.II we discuss spontaneous symmetry breaking first in the clean and then the disordered Schwinger model. In Sec.III we discuss level statistics and ergodicity breaking in the low energy Hilbert space. Finally, in Sec.IV we discuss localization in the low lying states. We conclude with a discussion of our results.

I. LATTICE REGULARIZATION OF THE SCHWINGER MODEL

The Schwinger model is a 1+1 dimensional model for quantum electrodynamics, describing the interaction of a Dirac field $\Psi$ (charge $e$) with an electric field $E$ that is conjugate to a vector potential $A^\mu$. The continuum action (including a random chemical potential) takes the form

$$\mathcal{L} = \bar{\Psi}(i\gamma^\mu \partial_\mu - e\gamma^0 A_\mu + \rho(x))\Psi - \frac{1}{4}F_{\mu\nu}F^{\mu\nu},$$

where the chemical potential depends only on spatial position. In the temporal gauge ($A_0 = 0$) the continuum Hamiltonian takes the form:

$$H = \int dx \left( \bar{\Psi}(-i\gamma^0 \partial_0 + e\gamma^1 A_1 + \rho(x))\Psi + \frac{1}{2}E^2 \right),$$

where $E = -\partial_0 A_1$ is the electric field conjugate to the vector potential. The electric field is fixed by Gauss’s law $\partial_j E = e\Sigma^0 \Psi$ up to a constant of integration, corresponding to a uniform background field. We set the background field to zero in all that follows.

We wish to regularize this model on a lattice and work at half filling (one electron for every two lattice sites), so as to maximize lattice commensuration effects. The Kogut-Susskind staggered potential formulation is the conventional way to do this. A brief summary of this approach can be found in Ref.24. In essence one introduces fermion fields $\phi$ which sit on lattice sites, and gauge fields that sit on links. The conjugate variable to the gauge field is a coordinate $L_n$, which is related to the electric field by $eL_n = E_n$, where $E_n$ is the electric field between sites $n$ and $n+1$. If the gauge field is taken to be compact, then Gauss’s law takes the form

$$L_n - L_{n-1} = \phi_n^+ \phi_n - \frac{1}{2}[1 - (-1)^n].$$

In this conventional Kogut-Susskind regularization, the ground-state expectation value $\langle \phi_n^+ \phi_n \rangle = 0$ (1) for $n$ even (odd). The regularization thus explicitly breaks the translation symmetry of the lattice, giving rise to a staggered potential with a two site unit cell. In essence, the regularization places electrons on odd sites, and positrons on even sites, leading to charge neutrality on average. Underlying this regularization is a picture of a two component plasma, where electrons and positrons are simultaneously present with equal density. This model may then be recast as a spin Hamiltonian by performing a Jordan-Wigner transformation and applying the Gauss’s law constraint. In this manner one obtains a Hamiltonian of the form

$$H = xH_{\text{hop}} + \lambda H_{\text{int}} + \frac{1}{2}\sum_{n=0}^{N-1} V(n)\sigma_n^z$$

where $V(n)$ comes from the random chemical potential and is chosen from the uniform distribution $(-\theta, \theta)$, with $n \in 0...N-1$, where $N$ is the system size. Meanwhile the specific forms of $H_{\text{hop}}$ and $H_{\text{int}}$ depend on the choice of boundary conditions and are tabulated in Table I. The parameters $x, \lambda, \theta$ characterize the hopping, interaction, and disorder strengths of Hamiltonian, respectively.

<table>
<thead>
<tr>
<th>$H_{\text{hop}}$</th>
<th>Open Boundary Conditions (OBC)</th>
<th>Periodic Boundary Conditions (PBC)</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\sum_{n=0}^{N-2} (\sigma_n^+ \sigma_{n+1}^- + \sigma_n^- \sigma_{n+1}^+)$</td>
<td>$\sum_{n=0}^{N-1} (\sigma_n^+ \sigma_{n+1}^- + \sigma_n^- \sigma_{n+1}^+)$</td>
<td></td>
</tr>
<tr>
<td>$H_{\text{int}}$</td>
<td>$\sum_{n=0}^{N-2} (E_n)^2$</td>
<td>$-\sum_{i&lt;j} \rho(i)\rho(j) \sin \left( \frac{\pi}{L} (i-j) \right)$</td>
</tr>
<tr>
<td>Conventional</td>
<td>$E_n = \frac{1}{2} \sum_{k=0}^{N-1} \text{sgn}(n + \frac{1}{2} - k)\sigma_k^z + \frac{1}{4}((-1)^n + 1)$</td>
<td>$\rho(n) = \frac{1}{2} \sigma_n^z$</td>
</tr>
<tr>
<td>Symmetric</td>
<td>$E_n = \frac{1}{2} \sum_{k=0}^{N-1} \text{sgn}(n + \frac{1}{2} - k)\sigma_k^z$</td>
<td>$\rho(n) = \frac{1}{2} \sigma_n^z$</td>
</tr>
</tbody>
</table>

TABLE I. We detail the Hamiltonians for open and periodic boundary conditions respectively. The Hamiltonians are obtained by starting from a lattice regularized open boundary model, performing a Jordan-Wigner transformation, and applying the Gauss’s law constraint. The row marked ‘conventional’ indicates the conventional Kogut-Susskind staggered potential regularization with unit cell of length two. This regularization is based on a picture of the Schwinger model as a two component plasma of electrons and positrons. The row labeled ‘symmetric’ indicates results for the alternative ‘symmetric’ regularization that we introduced, with unit cell of length one. This regularization is based on a picture of a one component plasma of electrons moving on a neutralizing ‘jellium’ background, which is more natural from a condensed matter perspective. Following standard notation, $\sigma_n^\alpha$ is the $\alpha$-Pauli matrix for the Hilbert space on site $n$, and $\sigma_n^z \equiv \frac{1}{2}(\sigma_n^+ \pm i\sigma_n^-)$ are the raising (lowering) operators on site $n$. 

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$$H = xH_{\text{hop}} + \lambda H_{\text{int}} + \frac{1}{2}\sum_{n=0}^{N-1} V(n)\sigma_n^z$$

where $V(n)$ comes from the random chemical potential and is chosen from the uniform distribution $(-\theta, \theta)$, with $n \in 0...N-1$, where $N$ is the system size. Meanwhile the specific forms of $H_{\text{hop}}$ and $H_{\text{int}}$ depend on the choice of boundary conditions and are tabulated in Table I. The parameters $x, \lambda, \theta$ characterize the hopping, interaction, and disorder strengths of Hamiltonian, respectively.
It is illuminating instead to consider an alternative ‘symmetric’ regularization with a one site unit cell, motivated by an alternative picture of the Schwinger model as a one component plasma, where electrons move on a neutralizing ‘jellium’ background. This symmetric model is obtained by removing the staggered potential and modifying the Gauss’s law to

\[ L_n - L_{n-1} = \phi_n^\dagger \phi_n \]  \tag{5}

With this regularization \( \langle \phi_n^\dagger \phi_n \rangle = 0 \forall n \), and the unit cell consists of a single lattice site. Moving to a spin model as before, one obtains again a Hamiltonian of the form (4), where however the ‘interaction’ Hamiltonian takes the distinct form tabulated in Table I. The two distinct choices of Gauss’s law (3, 5) correspond to two distinct lattice regularizations of the same continuum model - however, they have (apparently) very different properties. In particular, the conventional regularization has a unit cell of period two, while the ‘symmetric’ regularization has a unit cell of period one. Nevertheless, as we shall see, they have the same long distance physics, since the model with ‘symmetric’ regularization spontaneously breaks symmetry so that the states have a unit cell of period two.

The Hamiltonian given by Eq.4 together with Table I constitutes the basic model we study in this work. To simplify our analysis, we work throughout in the total spin-zero sector i.e. \( \langle \sum_n \sigma_n^z \rangle \mid j \rangle = 0 \) for the basis states \( \{ \mid j \rangle \} \). This is equivalent to working in a system with zero net charge.

II. SYMMETRY BREAKING IN DISORDERED SCHWINGER MODEL

To develop some intuition for why the low energy physics may be insensitive to the choice of regularization, recall that the classical one component plasma develops long range crystalline order at any temperature\(^{25,26}\), while the two component plasma does not\(^{27}\). One may thus conjecture that the quantum regularization based on a model of a one component plasma may spontaneously break symmetry, exhibiting crystalline order with a two site unit cell. This would then lead to results analogous to those obtained with the conventional regularization, which employs a Hamiltonian with a two site unit cell.

To test this scenario, we first employ exact diagonalization on systems with \( N = 12 \) spins. The Hamiltonian is a sparse, \( n \times n \) matrix with \( n = \binom{N}{N/2} \), constructed in the local \( \sigma^z \) basis, and the low energy eigenstates are found using Lanczos’ algorithm for the disordered systems with \( N \geq 16 \).

Having obtained the low energy eigenstates in this way, we look at the Fourier transform of the density density correlation functions \( \langle \sigma_n^x \sigma_{n+1}^x \rangle \) in the ground and first five excited states. The results are shown in Fig.1. Note that while the results depend on choice of boundary conditions, there is little dependence on the choice of regularization. In particular, regardless of regularization there is a peak at momentum \( k \approx \pi \) indicating crystalline order with a two site unit cell. This supports the basic scenario of spontaneous symmetry breaking in the ‘symmetric’ regularization that we have introduced. The (considerable) additional structure, particularly in the Fourier transform of the excited states, we believe to be a finite size effect.

Cleaner results may be obtained by enlarging the system size. To do this we employ DMRG techniques (following Ref. 24). This allows us to treat chains of up to \( N = 100 \) spins, although we are limited to systems with open boundary conditions, and are not able to easily access excited states. The results obtained in this manner are plotted in Fig. 2. Note that expanding system size leads to significantly cleaner results. Note also the very clear peak at momentum \( k = \pi \), indicating density structure with a two site unit cell, in both the conventional and symmetric lattice regularizations. This further supports our basic thesis that both the conventional and symmetric lattice regularizations encode the same physics, since the symmetric regularization spontaneously breaks symmetry until the system is no more symmetric than with conventional regularization. Why in symmetric regularization the peak is so much sharper in the first excited state than in the ground state is however a mystery.

We now comment on a remarkable feature hidden in Fig.1, 2. Namely, the random scalar potential \( (V(n) \) in the lattice regularization) explicitly breaks translation symmetry. That is, it constitutes ‘random field’ disorder, in the language of the Imry-Ma theorem\(^{23}\). According to this theorem, spontaneous symmetry breaking is supposed to be unstable to random field disorder in one spatial dimension. How then does the system with the ‘symmetric’ regularization manage to develop a two site unit cell, even in the presence of disorder?

To understand how the disordered Schwinger model evade Imry-Ma constraints, we must realize that the ‘spontaneous symmetry breaking’ of the Schwinger model (in its symmetric regularization) corresponds to charge density wave formation with period two (in effect, Wigner crystallization). A domain wall in this charge density wave is a soliton where the system switches from having charge preferentially on odd numbered sites to having charge preferentially on even numbered sites, or vice versa. This kind of ‘phase slip’ necessarily binds a charge. If the phase slip is accomplished by leaving two successive sites unoccupied, then it binds a positive charge; if it is accomplished by placing electrons on two successive sites, it binds a negative charge. Furthermore, charges in the Schwinger model experience a long range interactions. Thus, creating a domain of size \( L \) involves creating two domain walls at separation \( L \), and this carries an energy cost \( \sim L \). For large \( L \), this overwhelms the \( \sim L^{d/2} \) energy gain due to locally aligning with disorder, thus evading the Imry-Ma theorem\(^{28}\). The system can locally
FIG. 1. Fourier transform of the density density correlation function \( \langle \sigma_z^x \sigma_z^x + r \rangle \) in the ground state \( E_0 \) and first five excited states, as obtained using exact diagonalization on a system of twelve spins. The plots in the left hand column are with the conventional regularization (CR), based on a picture of a two component plasma, and with a two site unit cell. The plots in the right hand column are for the alternative ‘symmetric’ regularization (SR), based on a picture of a one component plasma, and with a one site unit cell. The first two rows show data obtained with open boundary conditions (OBC), and the last two rows show periodic boundary conditions (PBC). The labels \( x, \lambda, \theta \) denote the strength of the kinetic, interaction and disorder terms, respectively. Remarkably, the choice of lattice regularization (and the presence or absence of disorder) makes little difference to the density correlation function, which has structure around \( k = \pi \) indicating that the ground state has a two site unit cell.
FIG. 2. Fourier-transformed density-density correlations in the ground and first excited state, obtained using DMRG for a chain with $N = 100$ spins and open boundary conditions. The first column corresponds to the conventional regularization, the second column to symmetric regularization. Bond dimension $D$ is indicated on the plots, as is interaction strength $\lambda$ and disorder strength $\theta$. The disorder strength $\theta = 0.0, 0.5, 0.1$ going top down. The peak at $\pi$ indicates density structure with a two site unit cell, regardless of choice of regularization or addition of disorder.

deform, making small domains that align with disorder in regions where the disorder is small, but the domain walls are always bound together by the long range (confining) interaction, and thus long range order survives. The robustness of long range order to disorder is quite apparent in our numerical results.

As a parenthetical remark, we note that in the symmetric regularization, sites either carry charge $-e/2$ if an electron is present or $+e/2$ if an electron is absent. A domain wall consisting of two consecutive occupied/unoccupied sites thus carries a net half integer charge. This half integer charge is similar in origin to the celebrated Su-Schrieffer-Heeger model (Ref. 29), and may be understood as the polarization charge corresponding to the shift of an electron by half a unit cell. It implies that the domain wall solitons in the symmetric regularization may be viewed as fractionally charged quarks. Of course, these ‘quarks’ are confined into ‘mesons’ (dipoles) by the long range interaction.

III. ERGODICITY BREAKING IN LOW ENERGY HILBERT SPACE

We now search for ergodicity breaking in the low energy Hilbert space of the disordered Schwinger model, using as our principal diagnostic the level statistics ratio $\langle \tilde{r} \rangle$. The level statistics ratio $\langle \tilde{r} \rangle$ is a function of the spectrum
FIG. 3. Energy resolved level statistics ratios for the clean and disordered Schwinger model, obtained using numerical exact diagonalization on $N = 16$ spins, with kinetic scale $x = 1.0$ and interaction scale $\lambda = 0.125$. In the first plot, we consider the clean system in the symmetric regularization with PBC, and further resolve by the symmetry sector. The parity eigenvalue and translation phase angle are indicated in the plot. Even in this case, $\langle r \rangle$ meanders near the Poisson value, suggesting integrability. This is supported by the next figure. In the second plot, we prepared 100 disorder realizations in OBC, with $\theta = 0.05$. Note that at low energies, the mean level statistics ratio approaches the Poisson value 0.38.

We note first that in the absence of disorder $\theta = 0$, the level statistics are quite far from ergodic. We speculate this may be because of proximity to an integrable point (the clean continuum Schwinger model). Upon turning on disorder, the high energy sector of the Hilbert space starts to show ergodic level statistics. However, the level statistics ratio in the low lying Hilbert space exhibits clear signs of ergodicity breaking, with a level statistics ratio close to the Poisson value. Given the strong finite size effects, unfortunately, it is indeterminate whether this is significant, necessitating DRMG.

How should the ergodicity breaking in the low energy Hilbert space be understood? It is tempting to attribute this to localization, but the calculations of Ref.15, with our parameter values, suggest a localization length of order twenty sites, which is not expected to be accessible in a sixteen site simulation. Furthermore, there is a more prosaic explanation - namely a residual integrability of the model. (The lattice Schwinger model is not integrable as far as we are aware, but the clean continuum Schwinger model is integrable, and ‘proximity’ to this integrable point may contaminate our results given the limitations of finite size). To test for this possibility, we have tracked the energies of low lying excited states as a function of interaction strength in Fig. 4. We find clear signatures of level crossings in the low lying energy spectrum of the clean system, indicating that the low energy part of Hilbert space is behaving in an integrable manner at least in the finite sizes that we can access. Level statistics alone therefore cannot discriminate between localization and integrability as potential explanations of the non-ergodic level statistics in the low energy Hilbert space.

### IV. LOCALIZATION IN THE LOW ENERGY HILBERT SPACE

We have observed that level statistics are indicative of ergodicity breaking in the low energy Hilbert space of the disordered lattice Schwinger model, but cannot discriminate between localization and a residual integrability in this finite size system. We therefore directly examine the eigenstates. Since the localization lengths are estimated to be quite long $^{15}$ ($\xi \approx 20$ for the parameter values under consideration$^{31}$). We employ DMRG, which allows us to treat systems with $N = 100$ spins. A detailed explanation of the algorithm is found in Ref. 32. Diagonalizing the network at each iterative step is done using Lanczos’ algorithm. The bond dimension of the Schwinger Hamiltonian MPO is $D = 5$. The ground state is represented by an MPS with $D = 20$. Excited states are calculated as in Ref.24, by first finding the ground state and then adding a penalty term to the original Hamiltonian.

To probe localization, we examine the density imbalance between the ground $|\psi_0\rangle$ and first excited $|\psi_1\rangle$ states. Specifically, we extract

$$\tau(n) = \langle \psi_1 | \sigma_n^z | \psi_1 \rangle - \langle \psi_0 | \sigma_n^z | \psi_0 \rangle \quad n \in \{0...N-1\} \quad (6)$$

The results obtained are plotted in Fig.5 for the conventional regularization, and in Fig.6 for the symmetric
FIG. 4. Eigenenergies of first twenty states as obtained by Lanczos algorithm on a system of sixteen spins, in OBC. We only consider states with the same parity eigenvalue $P = 1$. Note the clear level crossings in the clean system, indicative of residual integrability.

regularization. In both cases, we clearly observe that the excited state contains a localized excitation, the localization length of which grows smaller as the disorder strength is increased. Furthermore, for all parameter values that were considered, the localization length was longer than would be accessible using exact diagonalization, necessitating DMRG. Finally, it is apparent from the figures that the elementary excitation is charge neutral, consistent with general expectations for the Schwinger model that excitations should be dipoles, which in one dimension lack long range interactions. We remained throughout in the regime where kinetic energy was the largest scale in the problem, and disorder the smallest, which is the regime where the analysis of Ref. 15 is expected to apply.

Finally, it is interesting to explore how the localization length scales with disorder strength, and to compare to the analytical predictions in Ref. 15. The localization length can be read off from plots of the form Fig. 5, 6, and the error bars estimated by sampling multiple disorder realizations. However, there is a slight complication. For some disorder realizations, the ‘density imbalance’ turns out to be bimodal, corresponding to existence of a long range resonance in the system (see e.g. Fig. 7a). In such cases, we take the localization length to be the mean of the decay lengths of the two peaks. In other cases, there are two peaks that overlap and cannot be cleanly separated (see e.g. Fig. 7b). Such cases are excluded from our estimate of the localization length. The localization length obtained in this manner is plotted as a function of disorder strength in Fig. 8 for the two choices of regularization. Ref. 15 predicted that the localization length $\xi$ should scale with disorder strength $\theta$ as $\xi \sim \theta^{-2/3}$. A fit to this prediction is in good agreement with the data.

V. DISCUSSION AND CONCLUSIONS

We have explored the interplay of disorder, confinement and lattice commensuration in a lattice regularized version of the Schwinger model. We have pointed out that there are two distinct choices of lattice regularization which at first glance are very different. One is based on a picture of a two component plasma of electrons and positrons, and has a two site unit cell; while the other is based on a picture of electrons moving on a uniform positively charged background, and has a one site unit cell. Nonetheless, both regularizations have the same physics, because the ‘symmetric’ regularization exhibits spontaneous symmetry breaking, such that the eigenstates of the Hamiltonian have a two site unit cell even though the Hamiltonian itself has only a one site unit cell. This spontaneous symmetry breaking survives the addition of disorder, with the Imry-Ma theorem being evaded due to the long range interaction.

We have used exact diagonalization to examine the energy resolved level statistics ratio of the model, and have data suggesting ergodicity breaking in the low energy sector of Hilbert space. However, in the system sizes to which exact diagonalization is limited, we cannot determine whether this is due to localization, due to proximity to an integrable point or simply due to finite size effects. Nevertheless, a DMRG investigation on systems with $N = 100$ spins confirms that the first excited state contains a localized excitation, the localization length for which decreases with increasing disorder strength. This is in outline consistent with the picture of localization in the disordered Schwinger model advanced in Ref. 15. The analytical analysis also predicted a scaling relation for localization length $\xi$ as a function of disorder strength $\theta$ - as $\xi \sim \theta^{-2/3}$. Fig. 8 shows a fit of the data to this form and finds good agreement between the data and the prediction.

There remain a number of interesting open questions for future work. In this investigation, we restricted ourselves to a regime where kinetic energy is the largest energy scale in the problem, and disorder the weakest. This is the regime where the analytic treatment of Ref. 15 is well controlled, and outside this regime there is no analytic understanding that we are aware of. However, dynamics reminiscent of MBL have also been numerically observed in a very different parameter regime when the interaction strength is the largest energy scale in the problem. A systematic exploration of the Schwinger model in all its parameter regimes would be an interesting problem for future work, and may serve as a useful
FIG. 5. Spatial profile of density imbalance (Eq: 6) between ground state and first excited state for a system of $N = 100$ spins, conventional regularization, and open boundary conditions treated using DMRG. Here $x = 1.0$, $\lambda = 0.2$ and $\theta = 0, 0.01, 0.05, 0.1$ indicate the strength of the kinetic, interaction, and disorder terms respectively. Note that the results are clearly indicative of the first excited state (of a disordered system) containing a localized excitation, the localization length of which decreases as disorder strength is increased. Note also that the localization length is too large to be accessible with exact diagonalization (consistent with analytic predictions\textsuperscript{15}) thus necessitating DMRG. Note also that the elementary excitations shown here are charge neutral (the density difference is equally likely to be positive or negative), consistent with general expectations.

guide to the development of an analytical understanding.

We note also that while we were able to confirm localization in the first excited state, in this model localization is expected to persist\textsuperscript{15} even in states with a non-zero density of excitations. This expectation follows because (as is apparent from our numerics), the elementary excitations of the problem are charge neutral, and thus should lack long range interactions. Unfortunately the numerical techniques available to us were not able to tackle states with a finite density of excitations in system sizes large enough to see localization (while remaining in the parameter regime where an analytical understanding is available). Numerical advances will likely be necessary to be able to explore this highly excited regime, and this would also be a worthwhile problem for future work.

Finally, while we have focused on the simplest model exhibiting confinement - the Schwinger model - confinement is a more general phenomenon. The interplay of disorder, confinement and lattice scale effects in more complicated (higher dimensional) settings would also be an interesting topic for future work.

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FIG. 6. Same as Fig. 5 but with the symmetric regularization
FIG. 7. For certain disorder realizations, the density imbalance in the first excited state is bilocalized (a). In this case we take the localization length to be the mean of the decay lengths for the two peaks. In other cases, there are two overlapping peaks (b) which cannot be cleanly separated. We exclude such configurations from our estimate of localization length.

FIG. 8. Localization length $\xi$ (extracted by eye from Fig.5 and Fig.6), plotted against disorder strength. The dashed lines are a best fit to the scaling prediction $\xi \sim \theta^{-2/3}$. The error bars come from sample to sample fluctuations and are estimated by averaging over at least 10 disorder realizations.