Quantum simulation of Anderson and Kondo lattices with superconducting qubits

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(Rceived 11 December 2007; revised manuscript received 6 January 2008; published 24 January 2008)

We introduce a mapping between a variety of superconducting circuits and a family of Hamiltonians describing localized magnetic impurities interacting with conduction bands. This includes the Anderson model, the single impurity one- and two-channel Kondo problem, as well as the one-dimensional Kondo lattice. We compare the requirements for performing quantum simulations using the proposed circuits to those of universal quantum computation with superconducting qubits, singling out the specific challenges that will have to be addressed.

DOI: 10.1103/PhysRevB.77.024522

PACS number(s): 03.67.Lx, 74.50.+r, 75.30.Mb, 85.35.Gv

I. INTRODUCTION

Quantum simulation1 consists of tuning the dynamics of a flexible quantum mechanical system to simulate the properties of another physical system or of a quantum mechanical model whose solution is unknown. While a universal quantum computer can efficiently simulate the dynamics of any quantum system,2,3 designing a quantum simulator for a specific model should simplify the experimental requirements. In particular, quantum simulators require neither high fidelity thresholds nor error correction, and thus fewer qubits are needed to get interesting results which are out of reach for classical numerical computations. The successful simulation of the Bose-Hubbard model using cold atoms in optical lattices4,5 is a paradigmatic example.

In the last years, there have been tremendous experimental achievements in the context of superconducting qubits, improving both quantum control and coherence times.6–15 While the aim is the implementation of a scalable universal quantum computer,14–16 this technology can be used in the quantum simulation of many-body physics with Josephson-Junction arrays.17–19

With this motivation, we first design a circuit to simulate the Anderson model for an impurity ion coupled to conduction electrons.20 This is done through an exact mapping from the effective models describing the superconducting circuits to the fermionic Hamiltonian for the impurity problem. In a certain parameter regime of our circuit, this leads to the Kondo Hamiltonian modeling the nontrivial physics of itinerant electrons that interact with a localized impurity having nonzero magnetic moment.21 This is one of the central models for strongly correlated electrons with implications going beyond condensed matter physics. Furthermore, our constructions are versatile enough to simulate other relevant Hamiltonians. With an appropriate choice of the geometry, these include the Kondo lattice22 and the multichannel Kondo model.23 The latter exhibits the simplest example of non-Fermi liquid behavior, which is a strong deviation from the standard model of condensed matter systems introduced by Landau.

We will also discuss the experimental challenges for implementing such circuits. In particular, while coherence and interaction times of current superconducting charge qubits seem good enough to simulate the long-time dynamical and static properties of these lattices, further experimental work is needed. Finally, we propose a set of measurements to gather information about transport, correlation, and energy spectra and comment on possible implications of the present work.

II. CIRCUIT DESIGN

A. Elementary components

Let us start by describing the proposed architecture. We consider two or more lines of low-capacitance superconducting islands coupled by Josephson junctions [Fig. 1(a)]. While not shown here, each island is itself part of a circuit like that of a charge qubit, with an external lead which capacitively induces an offset potential on the island, and possibly a Josephson coupling to a superconducting reservoir of Cooper pairs.24

Our quantum simulation protocols are based on two building blocks. First, as we will show below, each island will have a very low capacitance and be constrained to have at most one Cooper pair, which can be treated as an impenetrable boson. Second, selected pairs of islands will be coupled capacitively [Fig. 1(b)]. The coupling and voltages of these islands will be adjusted to create a suitable energy landscape that favors having a single excess Cooper pair in either the upper or lower island, states which we associate with an effective pseudospin. The appropriate choice of energies has the following form:

\[ E(n_{0\uparrow}, n_{0\downarrow}) = Un_{0\uparrow}n_{0\downarrow} - \epsilon(n_{0\uparrow} + n_{0\downarrow}). \] (1)

Here, \( n_{0\uparrow} \) and \( n_{0\downarrow} \) represent the excess of Cooper pairs on the upper and lower island in the impurity [Figs. 1(a) and 1(b)]. The constants \( U \) and \( \epsilon \) denote, respectively, the repulsive interaction energy coming from the capacitive coupling between islands and an energy offset of these islands with respect to the rest of the circuit.

As mentioned before, when we embed such an element in a circuit, the energy offset \( \epsilon \) enforces the impurity to host at most a single excess Cooper pair. In Fig. 1(c) we have de-
lected to an array of ordinary islands, with total population
in the impurity and the rest of the circuit. By external voltages,
perconducting phase, respectively.

Here, \(n_i\) represents a magnetic impurity. The Josephson energy for tunneling be-

Two occupation states for the impurity representing a pseu-
doing several levels of the circuit containing a single
impurity, with occupation numbers \(n_{01}\) and \(n_{02}\), con-
ected to an array of ordinary islands, with total population
\(N_r\). A possible configuration is shown in Fig. 1(a). If we neglect the kinetic energy of hopping pairs, we find that
the states where the impurity is occupied by a single pair, \(n_{01} = 1\) or \(n_{02} = 1\), have lower energy than states where this pair is
in the rest of the circuit. Multiply occupied impurities, \(n_{01} + n_{02} > 1\), have an even larger energy, \(U\), so that we can ne-
glect these states.

**B. Quantum circuit model**

In Secs. II D–II G, we will combine the previous elements
in circuits that simulate specific Hamiltonians. In order to
perform this task we must first introduce the mathematical
description of such circuits and the approximations under
which they will operate. We start considering the standard
description of such circuits and the approximations under

Finally, it is interesting to note that the total number of
pairs in the ground state can be tuned with very small volt-
tages that take islands away from degeneracy and which play
the role of a local chemical potential,

\[
\mu_{ir} = \sum_j U_{ij} \left( n_{ij}^{(s)} - \frac{1}{2} \right).
\]

**C. Hard-core boson model**

Summarizing the previous approximations, we have an
array of superconducting islands coupled both capacitively
and via Josephson tunneling. At the same time we have en-
sured that each superconducting island has a 0 or 1 excess
Cooper pair, all other states being energetically unfavorable.
In other words, we have imposed the hard-core bosons con-
ditions on the Cooper pairs: they become like impenetrable
bosonic particles such that no two pairs with the same effec-
tive “spin”\(^3\) can coexist on the same site.

In the limit we can describe any configuration of the cir-
cuit using the occupation numbers \(n_{ij}\). Due to the hard-core
condition, the dynamics is well-approximated by the projec-
tion of the circuit Hamiltonian (2) onto the relevant energy
subspace, \(n_{ij} \in \{0,1\}\). To perform this projection we will need to express the phase terms in the basis of occupation
numbers,

\[
\sum_{ij} J_{ij} \cos(\phi_{ij} - \phi_{ja}) = \sum_{ij} \frac{1}{2} J_{ij} \sum_{n_{ja} = 0} n_{ja} + 1, n_{ja} - 1 \langle n_{ij}, n_{ja} \rangle 
+ \text{H.c.}
\]

Note that this equation has a very simple interpretation: Coo-

\[
\text{FIG. 1. (a) Quantum circuit simulating the single Anderson or}
\text{Kondo impurity model. The big square boxes are superconducting}
\text{islands, connected by crossed squares which are Josephson junc-
\text{tions. A selected pair of islands is capacitively coupled and repre-
\text{sents a magnetic impurity. The Josephson energy for tunneling be-
\text{tween junctions are } J \text{ for the main lattice and } J \text{ around the impurity.}
\text{(b) Two occupation states for the impurity representing a pseudo-
\text{spin state. (c) Energy levels of the impurity and a neighboring site.}
\text{n}_{01}, n_{02} \text{ and } N_r \text{ represent the number of pairs of the two islands}
\text{in the impurity and the rest of the circuit. By external voltages,}
\text{ } \epsilon, \text{ and the capacitive coupling, } U \sim 1/C, \text{ it becomes favorable}
\text{to have a single pair in the impurity.}
\]
per pairs are transferred one by one between any two islands connected by the hopping matrix $\overline{J}_{ij}$.

The projection is implicitly performed by means of a more concise mathematical representation that includes an exclusion principle for particles on the same site. We introduce hard-core bosonic operators, $a_{i\alpha}^\dagger$ and $a_{i\alpha}$, which, respectively, create and annihilate a Cooper pair on a given site. They act on the basis elements as follows:

$$a_{i\alpha}|n_{i\alpha}=1\rangle = |n_{i\alpha}=0\rangle, \quad a_{i\alpha}(0)=0, \quad a_{i\alpha}^\dagger(0) = |1\rangle, \quad a_{i\alpha}^\dagger(1)=0$$

have the usual bosonic commutation relations

$$[a_{i\alpha}, a_{j\sigma}^\dagger] = \delta_{ij}\delta_{\sigma\sigma'}, \quad [a_{i\alpha}, a_{j\sigma}] = 0, \quad \delta_{ij}=0$$

and enforce the hard-core condition simply because $a_{i\alpha}^\dagger a_{i\alpha}=0$. Using these operators we can express the number of particles $n_{i\alpha}=a_{i\alpha}^\dagger a_{i\alpha}$ and the projected phase operator,

$$2\cos(\phi_{i\alpha} - \phi_{j\alpha}) = a_{i\alpha}^\dagger a_{j\sigma} + a_{j\sigma}^\dagger a_{i\alpha}$$

which adopts the form of a hopping term.

\section*{D. Single impurity Anderson model}

We will apply all the approximations and techniques introduced before to study a particular circuit, which consists of two impurity islands coupled to two 1D Josephson junction arrays [Fig. 1(a)]. The low energy dynamics of this circuit is described by the bosonic Hamiltonian,

$$H_A = \sum_{\langle i,j \rangle = 1} (a_{i\alpha}^\dagger a_{j\alpha} + H.c.) - i \sum_{\sigma} (a_{0\alpha}^\dagger a_{1\alpha} + H.c.) + E(n_{01}, n_{01}) + \sum_{i\alpha} \mu_{i\alpha} n_{i\alpha}.$$  

Out of all terms in the hopping matrix $\overline{J}_{ij}$, we have left only the Josephson energies between sites in the conduction array, $\overline{J}_{ij}=J$ for $i>0$, and the coupling between the impurity site and this band, $\overline{J}_{01}=\overline{J}_{10}=t$ [see Fig. 1(a)]. Finally, as mentioned in Sec. II C, we have included an effective chemical potential which depends on deviations from the degeneracy point of the local potentials acting on each island.

We can map this Hamiltonian to a fermionic model using the Jordan-Wigner transformation,\cite{PhysRevB.63.235305,PhysRevB.81.075318}

$$a_{i\alpha} = c_{i\alpha}^\dagger (-1)^{\sum_{j<i} \delta_{\alpha\sigma}} c_{j\sigma}^\dagger,$$

$$a_{i\alpha} = c_{i\alpha}^\dagger (-1)^{\sum_{j<i} \delta_{\alpha\sigma}} c_{j\sigma}^\dagger + N_{i\alpha},$$

$$n_{i\alpha} = c_{i\alpha}^\dagger c_{i\alpha} = a_{i\alpha}^\dagger a_{i\alpha}.$$  

The new operators satisfy the usual fermionic anticommutation relations $\{c_{i\alpha}, c_{j\sigma}^\dagger\} = \delta_{ij}\delta_{\sigma\sigma'}$ and $N_{i\alpha} = \sum_{\sigma} n_{i\alpha}$. With this transformation, our effective Hamiltonian (9) converts identically into a fermionic model.

\section*{E. Kondo model}

A special limit of the Anderson model is obtained when we impose a low capacitance on the impurity, $U \rightarrow \infty$, and make the energy displacement large compared to the coupling between the impurity and its neighbors, $t \ll \epsilon$. Using standard second order perturbation theory, the result is the single impurity Kondo Hamiltonian,\cite{PhysRev.144.433}

$$H_K = \sum_{\langle i,j \rangle = 1} c_{i\alpha}^\dagger c_{j\alpha} + J_K \tilde{S}_0 \cdot \tilde{s}_1.$$  

Here $\tilde{S}_0$ and $\tilde{s}_1$ are the pseudospins of the impurity and of the first lattice site,

$$\tilde{S}_0 = \sum_{\sigma,\sigma'} \frac{1}{2} \tilde{\sigma}_{\sigma\sigma'} c_{0\sigma}^\dagger c_{0\sigma'},$$

$$\tilde{s}_1 = \sum_{\sigma,\sigma'} \frac{1}{2} \tilde{\sigma}_{\sigma\sigma'} c_{1\sigma}^\dagger c_{1\sigma'},$$

respectively, and $J_K = t^2/\epsilon$ is the antiferromagnetic coupling between the Kondo impurity and the free fermions. We want to stress that by tuning the parameters $J$, $t$, and $\epsilon$, we can realize a crossover from a low energy regime below the Kondo temperature $T_K$ to a high energy limit.

\section*{F. Two-channel Kondo model}

The previous setup and treatments can be reused to simulate a two-channel Kondo system.\cite{PhysRevB.77.024522} As shown in Fig. 2(a), the impurity is in the middle of a one-dimensional lattice, coupled to two different conduction bands. Following similar steps, the new effective model becomes

$$H_K = - \sum_{\langle i,j \rangle = 1} c_{i\alpha}^\dagger c_{j\alpha} + J_K \sum_{\alpha=1,2} \tilde{S}_0 \cdot \tilde{s}_{1\alpha},$$  

where $\alpha \in \{1,2\}$ denotes the two fermionic channels of our problem. Note that in this Hamiltonian we have removed the
the shape of the bosonic Hamiltonian. We will, however, perform an incomplete Jordan-Wigner transformation which acts only on the conduction arrays, \(a_{i\sigma}\), leaving the \(b_{k\sigma}\) operators untouched,

\[
a_{i\uparrow} = c_i (1 - \Sigma_{\alpha<\uparrow} n_{i\alpha}^{(a)}),
\]

\[
a_{i\downarrow} = c_i (1 - \Sigma_{\alpha<\downarrow} n_{i\alpha}^{(a)} + n_{i\alpha}^{(b)}),
\]

where \(N_{\alpha}^{(a)} = \Sigma n_{i\alpha}^{(a)}\). With this unitary map, and working out the second order perturbation theory, one arrives at the standard Kondo lattice model,

\[
H_{KL} = -J \sum_{(i,j),\sigma} c_i^\dagger c_j + \sum_{k,\alpha,\alpha'} \frac{J_{KL}}{2} \sum_{\alpha,\alpha'} c_{i\alpha}^\dagger \sigma_{\alpha\alpha'} c_{j\alpha'}.
\]  

Here, \(\hat{S}_k\) denotes the spin of the \(k\)th impurity which is coupled to the site \(i_k\) and is defined using bosonic operators \(\hat{S}_{i_{\alpha}}^{\uparrow} = \Sigma_{\alpha<\uparrow} \sigma_{\alpha\alpha'} c_{i\alpha}^\dagger c_{j\alpha'}\). This general notation allows for having less impurities than a conduction electron, as well as them being placed on arbitrary positions. The coupling is once more antiferromagnetic \(J_K = -t^2/\epsilon\) and just as tunable as in the case of a single impurity.

### III. ENERGY SCALES AND DECOHERENCE

Let us consider what are the experimental challenges for implementing these mappings. We begin with the energy scales of our problem. On the one hand we have free fermions, which hop with a tunneling amplitude \(J\) and a Fermi energy \(\epsilon_F \sim 2J\). In current experiments with charge qubits \(J\) can be roughly of the order of a few gigahertz. On the other hand we have the Kondo coupling \(J_K = t^2/\epsilon\), which will be in general smaller. For this coupling to be relevant, the temperature at which the experiment is performed should lay below the Kondo temperature, \(k_B T_K = \epsilon_F \exp(1/J K \rho(k))\), where \(\rho(k)\) is the density of states at the Fermi energy. Since in our case the density of states at half filling \((\mu=0)\) is roughly \(\rho(k) \approx \pi J\), using \(\rho J\), we have \(k_B T_K \sim 2J \exp(-\epsilon J / 2\pi)\). This means that, conservatively, for \(\epsilon J = 8 - 10\), the Kondo temperature \(T_K \sim 40 - 75\) mK would be larger by 20 mK, the refrigerator temperature of current experiments.

Another important challenge is decoherence, which in our setup arises mainly from charge fluctuations. As is the case of scalable quantum computation, the characterization of decoherence in multiqubit setups is still an open problem that deserves further investigation. As a guide, we consider the time scale for the decoherence of a single qubit is \(T_2 \sim 0.5\) μs, which is achieved for dispersive readouts. This time scale gives a frequency of 2 MHz which is well below \(J/h \sim 10\) GHz and two orders of magnitude away of the lowest frequency \(T_2/h = 10^7\) Hz that we find here. Furthermore, associated to this decoherence rate we can establish an effective temperature of 0.1 mK which is also well below the Kondo temperature.

When compared to other systems, such as cold atoms in optical lattices, the superconducting circuit approach looks
to be enjoying real advantages. First, the superconducting qubits are not constrained to periodic or quasiperiodic structures. Second and most important, the introduced setup has the potential quality of a good quantum simulator, which is measured by the number of times a particle can hop before the wave function is affected by decoherence. For cold atoms in the lowest band of an optical lattice,\textsuperscript{5} coherence times of 500 ms are to be compared with a hopping rate of 1 kHz between lattice sites, giving a ratio of 500 or better. A similar ratio with superconducting circuits, where the hopping is around 10 GHz, means the decoherence time can be 0.05 μs, an order of magnitude faster than current experiments.\textsuperscript{8,28} On the other hand, superconducting circuits have the disadvantage of being one-time experiments because the geometry of the couplings cannot be modified in real time. In addition, it may be complicated to calibrate all islands to reach the sweet spot. However, if we realize that the couplings can be initially switched to zero, this task is as difficult as tuning the individual superconducting qubits in a scalable quantum computer.

IV. MEASUREMENT

While we have demonstrated that certain quantum circuits can be used to simulate interesting fermionic models, the equivalence between circuit and the Hamiltonian involves the use of Jordan-Wigner transformations (8). Therefore not all measurements on the quantum circuit will give directly properties of the respective fermionic system. One can envision three kinds of measurements in these systems which do not have this problem. The first ones are global transport measurements that mimic past experiments with Josephson junction arrays.\textsuperscript{17} Since they are based on total particle numbers, they are the same for fermionic and bosonic models.

Next we have spectroscopic measurements and measurement of energy gaps. The coupling between the impurity and the conduction band leads to a binding energy of order $\Delta = k_B T_k$ given above. This energy can be measured by trying to polarize the impurity with an oscillating voltage applied to the island. We expect this to have an effect either when the potential is of order $\Delta$ or when it oscillates with a frequency of order $\Delta/\hbar$.

The third kind of experiments corresponds to measuring the individual qubits and extracting information about certain correlation functions. In the case of Fig. 1(a), the formation of a singlet with the conduction band electrons gives rise to a strong spin correlation in the vicinity of the impurity. The correlation length is of order $\xi = \exp[1/|J_K p|]$ sites and can be seen in the connected correlator as a function of the site $m$,

$$C_m = \langle S_0 S_m \rangle - \langle S_0 \rangle \langle S_m \rangle \sim \exp(- m/\xi).$$

This is known as the Kondo cloud and has not been directly detected by condensed matter setups. In our case, $C_m$ can be computed from the statistics of the population difference between islands, $s_i = (n_i - n_{i-1})/2$, which is best measured using dispersive probes.\textsuperscript{28,29}

V. CONCLUSIONS

Summing up, in this paper we have presented a mapping between some superconducting circuits to a family of relevant Hamiltonians describing magnetic impurities in conduction bands. While there are still implementation issues to be solved, they seem less demanding than those of a fully scalable quantum computer implementation with the same technology. We thus expect that our work will motivate further developments, both on the experimental and on the theoretical side, searching better multiqubit coherence times and other mappings which are based on flux or hybrid qubits.

ACKNOWLEDGMENTS

We thank Vitaly Golovach, Stefan Kehrein, and Jan von Delft for their comments. E.S. acknowledges financial support from DFG SFB 631, EU EuroSQIP projects, and the German Excellence Initiative via the “Nanosystems Initiative Munich (NIM).” M.A.M.D. and J.J.G.R. acknowledge financial support from Projects No. FIS2006-04885 (Spanish M.E.C.) and No. CAM-UCM/910758. J.J.G.R. acknowledges support from the Ramon y Cajal Program of the M.E.C.

31 That is, belonging to the same junction array.
32 Note that this procedure, though equivalent in the results, differs in the method from Ref. 30, where perturbation theory is carried on in momentum space.