First-principles calculation of electronic transport in low-dimensional disordered superconductors

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We present a novel formulation to calculate transport through disordered superconductors connected between two metallic leads. An exact analytical expression for the current is derived and applied to a superconducting sample described by the negative-U Hubbard model. A Monte Carlo algorithm that includes thermal phase and amplitude fluctuations of the superconducting order parameter is employed, and a new efficient algorithm is described. This improved routine allows access to relatively large systems, which we demonstrate by applying it to several cases, including superconductor-normal interfaces and Josephson junctions. Moreover, we can link the phenomenological parameters describing these effects to the underlying microscopic variables. The effects of decoherence and dephasing are shown to be included in the formulation, which allows the unambiguous characterization of the Kosterlitz-Thouless transition in two-dimensional systems and the calculation of the finite resistance due to vortex excitations in quasi-one-dimensional systems. Effects of magnetic fields can be easily included in the formalism, and are demonstrated for the Little-Parks effect in superconducting cylinders. Furthermore, the formalism enables us to map the local super and normal currents, and the accompanying electrical potentials, which we use to pinpoint and visualize the emergence of resistance across the superconductor-insulator transition.

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I. INTRODUCTION

Chief among the remarkable effects observed in superconductors is their eponymous perfect conductivity. Within BCS theory, where superconductivity arises due to pairing between electrons, the effects of temperature \( T \), magnetic field \( B \), and disorder are well understood: as the pairing amplitude is suppressed by these physical parameters, the system becomes normal, and attains a finite resistance. For low-dimensional systems, on the other hand, it has been long understood that phase fluctuations of the pairing amplitude play a major role in the loss of perfect conductance. In two-dimensional systems, for example, it has been demonstrated that as the temperature increases there is a critical temperature \( T_{\text{KT}} \) where vortices and antivortices unbind and proliferate through the system, leading to the loss of global phase coherence and superconductivity, even though the pairing amplitudes remain finite. Indications of such a Berezinsky-Kosterlitz-Thouless (BKT) transition have been observed in Josephson-junction arrays, in superconducting (SC) thin films, and possibly in high-\( T_c \) cuprates.

In recent years there has been a reinvigoration of research into low-dimensional superconductors. This has been motivated by intriguing experimental observations of electronic transport through disordered SC thin films, such as a huge magnetoresistance peak and a “super-insulator” phase, and by the technological progress in producing two-dimensional superconductors in the interface between two oxides and in making ultra-thin cuprate superconductors. Many of these observations have not yet been satisfactorily explained, chiefly because there is no theory that can calculate the current, even numerically, through a disordered superconductor, based on a microscopic model.

The calculation of the resistance within the BCS picture, usually based on the Bogoliubov-de Gennes (BdG) mean-field approach, is straightforward. Blonder, Tinkham, and Klapwijk (BTK) studied the reflectance and transmission at a metal-superconductor junction, and an analogous study was performed at superconductor-metal-superconductor junctions. Similar approaches utilized the Buttiker-Landauer picture for noninteracting Cooper pairs to study scattering through a SC region. (A difficulty with the direct application of the BdG formalism is the nonconservation of charge, which can be overcome by studying a normal ring containing a SC segment.) An alternative approach near to the BCS critical temperature is to use a scaling assumption for the conductivity. The current through diffusive normal metal-superconductor structures has also been calculated using a Keldysh scattering matrix theory. All these approaches neglect phase fluctuations so they cannot be used to study two-dimensional superconductors that exhibit a BKT-like transition at low temperatures.

The resistance of low-dimensional superconductors can also be calculated using phenomenological models. The conductivity of uniform systems can be probed analytically by studying phase slips across the sample within the Ginzburg-Landau approach. Thermally excited phase slips explain both nonlinear conductivity and vortex creep induced resistance, while quantum activated phase slips can drive SC wires insulating. However, phenomenological calculations are neither underpinned by a microscopic model nor include Coulomb repulsion or disorder except for the introduction of a phenomenological normal state resistance.

Here we develop a new formalism to calculate the current and its spatial distribution in a superconductor taking into account phase fluctuations in the presence of disorder, finite \( T \) and \( B \), and Coulomb repulsion. The approach we detail here is based on the Landauer-Buttiker scheme, where one attaches metallic leads to the sample and then calculates its conductance. The lead-superconductor tunneling barriers ensure that the conductance of the system is always finite, even in the SC phase. A previous attempt using the quantum Monte Carlo approach to calculate current in disordered...
systems employed the fluctuation-dissipation theorem via the current-current correlation function.23

The Landauer formula14,15 is a widely adopted method to calculate the current through a mesoscopic sample that contains noninteracting particles. Meir and Wingreen24 have generalized the formula to produce an exact expression for the current through any interacting region attached to noninteracting leads, which has been successfully applied to a wide range of systems. Following this approach, we partition the system into the three parts shown in Fig. 1: the left-hand lead, the central interacting region, here a superconductor, and the right-hand lead. In the leads the natural particle basis set is comprised of electrons, and in the sample the natural basis set is Bogoliubons. To circumvent this mismatch of particle basis sets we reformulate the Meir-Wingreen formula in a Bogoliubon basis set to derive an exact analytical expression for the current flow through a possibly SC region, attached to two metallic leads.

Having derived a general, exact formula, we then model the SC region by a generalized negative-\(U\) Hubbard model based on the lattice shown in Fig. 1. Introducing two local auxiliary fields (which reduce to the local density and gap at zero temperature), we decouple the interacting fermions. While the conductance formula is exact, in order to evaluate correlation functions, while integrating exactly over the fermions, we neglect quantum fluctuations of the auxiliary parameters, and integrate numerically over their thermal fluctuations\(^{25,26}\) using a Monte Carlo method. A significant advantage of the formalism is that it allows us to construct current and potential maps of the system. These enable us to diagnose the microscopic features that increase the resistance of the sample. This paper details the new procedure and presents a number of applications to simple systems, where one can compare with existing theories and shed light on several phenomena. These calculations represent the first \textit{ab initio} studies of these effects, and so allow us to provide a link between the phenomenological parameters often used to describe them and the underlying microscopic variables.

Moreover, we go further and study regimes not accessible by the phenomenological models.

The paper is organized as follows. In Sec. II we first derive an exact analytical expression for the current through a SC region. Using this expression, in Sec. III we describe the approximation and numerical procedure, and outline improvements to the auxiliary field approach that allows us to study large systems. Having developed the new formula for the current and accompanying computational tool, it is imperative to carefully test it against a series of known results. Therefore, in Sec. IV A we study superconductor-normal interfaces in clean systems, and compare with the BTK transmission formulas, while in Sec. IV B we study the temperature-dependent current in a Josephson junction. In Sec. IV C we describe how effects of decoherence and dephasing are manifested in the formalism. We investigate the temperature dependence of resistance in Sec. IV D in which we uncover the temperature dependence of the resistance that characterizes the BKT transition in two dimensions and vortex excitations in quasi-one-dimensional systems. We then, in Sec. IV E, apply an external magnetic field to probe the Little-Parks effect. Finally, we demonstrate how to construct current and potential maps for the system, and use them to study the microscopic behavior at the superconductor-insulator transition in Sec. IV F. The details of the analytical derivation and the numerical procedure are described in the Appendixes.

II. ANALYTICAL DERIVATION

A. Current formula

To calculate the current for interacting particles, we start with the general formula for the current\(^{24}\) through an interacting region, connected between two noninteracting leads

\[
J = \frac{ie}{2\hbar} \sum_\sigma \int d\epsilon \left[ \text{Tr} \left( f_L(\epsilon) G_{L\sigma}^{\left(\uparrow\right)} - f_R(\epsilon) G_{R\sigma}^{\left(\downarrow\right)} \right) (G_{\sigma}^+ - G_{\sigma}^-) \right] + \text{Tr} \left( (\Gamma_L - \Gamma_R) G_{\sigma}^- \right). \tag{1}
\]

Here \( f_i(\epsilon) = \frac{1}{1 + e^{\beta(\epsilon - \mu_i)}} \) with \( \mu_i \in \{L,R\} \) the Fermi distribution of the left (L) and right-hand (R) leads that are held at chemical potentials \( \mu_L \) and reduced temperature \( \beta \equiv 1/k_B T \) (where \( k_B \) is the Boltzmann constant). The imposed potential difference \( eV \equiv \Delta \mu = \mu_L - \mu_R \) between the leads drives the current \( J \) through the system. The integral is over all electronic energies \( \epsilon \), \( \Gamma_0^{ij} \equiv 2\pi \sum_{\kappa \rho} \rho_\kappa(\epsilon) Y_{\kappa a} Y_{\rho b}^{\ast} \) for channels \( a \) in lead \( \chi \), and \( Y_{\kappa a} \) is the tunneling matrix element from channel \( a \) in the lead to site \( i \) in the sample. Finally, \( G_{\sigma ij} = G_{\sigma ij}^+ \) are the retarded, advanced, and lesser Green’s functions (in the site basis) for electrons of spin \( \sigma \) in the sample calculated in the presence of the leads.

Equation (1) is exact and it captures, via the electronic Green’s function \( G_{\sigma ij} \), all the processes that can transfer an electron through the system. When the intermediate regime has SC correlations, some of these processes involve Andreev scattering—absorption of an electron pair by the condensate and a propagation of the remaining hole. To expose these processes, allowing us below to separate the SC and normal channels, it is convenient to transform from the electron basis set \((c_{i\sigma}^\dagger, c_{i\sigma})\) with site index \( i \) into the Bogoliubov basis.
set \( \langle y_{m\sigma} y_{n\sigma} \rangle \), using the Bogoliubov-de Gennes relations 
\[ c_{i\sigma} = \sum_n u_i (n) y_{n\sigma} - \sigma v_i (n) y_{-n-\sigma} \]  
(at present \( u_i \) and \( v_i \) are arbitrary, except for the unitarity condition, but later on they will be determined by the actual Hamiltonian that will be used for the SC region). The Bogoliubov basis will also allow us to identify the low-energy excitations in the system, enabling us to speed up the numerical calculation by concentrating on these states, see Appendix B. Green’s functions transform from the electron basis \( G_{\sigma} \) into the energy basis set of Green’s functions \( \{ G_{\sigma}^\ast, G_{\sigma} \} \) and the family of anomalous Green’s functions \( H_{\sigma}(m,n) = -i \langle y_{m-\sigma} y_{n\sigma} \rangle \), \( H_{\sigma}^\ast(m,n) = i \langle y_{m-\sigma} y_{n\sigma} \rangle \), Green’s functions are diagonal so it does not demand the expensive matrix inversion embodied in Eq. (A2) to find the general equation for the current. Due to its usefulness we also note that an analogous expression can be derived for the normal current when injecting electrons outside of the gap 
\[
J = \frac{e}{h} \sum \int d\epsilon \left[ f_\epsilon (e) (\Gamma^+_{uu} + \Gamma^+_{vv}) - f_{\bar{\epsilon}} (e) (\Gamma^+_{uu} + \Gamma^+_{vv}) \right] \text{Im} G^\ast_{\epsilon} \, ,
\]
where \( \text{Im} \) stands for the imaginary part. Since this term represents the normal current, it has a direct \( Y^2 \) dependence on the tunneling matrix element. Though they offer a considerable computational advantage, these perturbative formulas cannot be used on the border of the superconductor-insulator transition where the superconductor gap breaks down and \( \Delta < Y \). Therefore, unless specified, we use the full expression for the current, Eq. (4), in our numerical calculations.

### B. Current and voltage maps

Equation (6), with a coefficient of \( Y^2 \), describes the normal current that enters and leaves the system as single electrons, whereas Eq. (5) with a coefficient of \( Y^4 \) corresponds to a tunneling supercurrent. However, the normal and supercurrent can interchange inside the sample. To understand the microscopics behind phenomena in the disordered superconductor, it will be advantageous to be able to probe the spatial distribution of the current and its normal or SC nature through the sample. Therefore, here we extend our formalism to map out the flow of current within the sample. To calculate the current distribution map we use the general expression for the current crossing a single bond from site \( i \) to \( j \)
\[
J_{ij} = \frac{2e}{h} \sum \int \frac{d\epsilon}{2\pi} \left[ t_{ij} G^\ast_{\sigma} (j,i) - t_{ji} G^\ast_{\sigma} (i,j) \right] \, .
\]
Transforming again into the diagonalized basis, the local current is
\[
J_{ij} = \frac{2e}{h} \sum \int \frac{d\epsilon}{2\pi} \left[ \text{Tr} \left[ \Lambda_{ij}^\dagger - \Lambda_{ij} \right] G^\ast_{\epsilon} - \left[ \Lambda_{ij}^\dagger - \Lambda_{ij} \right] G^\ast_{\epsilon} \right] + \sigma \left[ \Lambda_{ij}^\dagger - \Lambda_{ij} \right] H^\dagger_{\epsilon} + \sigma \left[ \Lambda_{ij}^\dagger - \Lambda_{ij} \right] H^\dagger_{\epsilon} \, ,
\]
where \( \Lambda_{ij}^\dagger (m,n) = t_{ij} u_i (m) v_j (n) \). As before, the normal \( G^\ast \) and anomalous Green’s functions \( H^\ast \) are calculated in Eq. (A3) in the presence of the leads. Moreover, we note that the current comes in two flavors, the contribution to the current from the normal Green’s function \( G^\ast \) is associated with the normal current and that from the anomalous Green’s function \( H^\ast \) gives the Cooper pair current. In Sec. IV F we verify that this intersite current yields the correct net conservation of charge.
To provide an additional probe into the nature of the superconductor-insulator transition we extend the formalism to map the local chemical potentials across the sample. This should reveal any weak links and the location of the sources of resistance in a sample. To determine the local effective potential at a specific site we add a weak link from that site to a third lead (a “tip”). The tunneling current from the tip into the sample is then calculated, and the chemical potential of the tip adjusted until that current flow is zero. This chemical potential thus corresponds to the effective local chemical potential in that site. To calculate the current flow into the tip we first evaluate the full Green’s functions $G$ in the sample in the presence of a voltage drop between the left and right reservoirs, Eq. (A2), but without the tip. We then use the perturbative formula for the current, Eq. (5), but with one lead representing the left- and right-hand leads, and the other the perturbative tip. This process is repeated for each site in the sample (due to the perturbative nature of the tip, this calculation can be done simultaneously for all sites). In Sec. IV F we demonstrate how maps of the potential can expose weak links in the sample and help diagnose the microscopic mechanisms that give rise to resistance.

III. MODEL AND NUMERICAL PROCEDURE

In the previous section we developed an exact analytical formula for the current through an arbitrary intermediate region, which may include SC correlations. We now use a specific model to describe this SC region: the negative- $U$ Hubbard model, a lattice model that includes on-site attraction and even long-range repulsive interaction (which we will not deal with in this paper). The Hamiltonian is

$$\hat{H}_{\text{Hubbard}} = \sum_{i,\sigma} \epsilon_{i\sigma} c_i^{\dagger} c_i - \sum_{i} U_i c_i^{\dagger} c_i c_i^{\dagger}$$

$$- \sum_{\langle i,j \rangle,\sigma} (t_{ij} c_i^{\dagger} c_j + t_{ij}^{\ast} c_j^{\dagger} c_i),$$

(9)

where $\epsilon_{i\sigma}$ is the on-site energy, $t_{ij}$ the hopping element between adjacent sites $i$ and $j$, and $U_i$ the on-site two-particle attraction, which is taken to be uniform and $i$ independent in this paper. An orbital magnetic field can be incorporated into the phases of the hopping elements $t_{ij}$, while a Zeeman field splits the spin-dependent on-site energies $\epsilon_{i\sigma}$. Unless noted differently, all $t_{ij} \equiv t$ are taken to be equal. In this paper we will only deal with orbital fields. To account for disorder, $\epsilon_i$ will be drawn from a Gaussian distribution with characteristic width $W$. The intersite spacing is $a$. Unlike, for example, the disordered $XY$ model, the negative-$U$ Hubbard model can lead to a BCS transition, a BKT transition, or a percolation transition, and thus this choice is general enough not to limit a priori the underlying physical processes. Importantly, the model includes the fermionic degrees of freedom which may be relevant to some of the experimental observations.

Calculation of correlation functions, for example, Green’s functions that enter the current formula, requires thermal averages. To perform the thermal average we need to decouple the quartic interaction term, so we employ the exact Hubbard-Stratonovich transformation

$$\exp \left[ \int_0^\beta d\tau \sum_i U_i c_i^{\dagger} c_i c_i^{\dagger} c_i \right]$$

$$= \int D\Delta D\tilde{\Delta} \exp \left[ \int_0^\beta d\tau \sum_i \frac{-|\Delta_i(\tau)|^2}{U_i} \right.$$

$$\left. + \Delta_i(\tau) c_i^{\dagger} c_i^{\dagger} + \tilde{\Delta}_i(\tau) c_i c_i \right].$$

(10)

which is basically a Gaussian integration $[D\Delta \equiv \prod_{\tau} d\Delta_\tau(\tau)$, where the product runs over all times and all sites]. Note that the field $\Delta_i(\tau)$ is just an integration variable that decouples the two-body term in the SC channel, and should not be confused with $|U_i||c_i c_i^{\dagger}$]. Similarly, one introduces the integration fields $\rho_\alpha(\tau)$, that couple to the spin density $\langle c_i^{\dagger} c_i \rangle$, and leads to an additional term $-\sum_{\tau,\sigma} \langle U_i | \rho_\sigma - \sigma \tau | c_i^{\dagger} c_i \rangle$ in the action (which, in the mean-field approximation gives rise to the Hartree-Fock contribution).

The Hubbard-Stratonovich decoupling in both the $\Delta$ and $\rho$ channels is not only provides access to both soft degrees of freedom, but also the saddle-point solution gives the standard mean-field results for those fields and furthermore guarantees that the action expanded to Gaussian order corresponds to the random phase approximation. Since our main interest lies in thermal effects, for example, the thermal BKT phase transition or thermal activation of vortices, we now neglect quantum fluctuations ($\tau$ dependence) of the auxiliary field $\Delta$. One can then write the partition function for the Hubbard model as

$$Z = \text{Tr}[e^{-\beta \hat{H}_{\text{Hubbard}}}] = \int \mathcal{D}(\Delta, \rho) \text{Tr}_f \left[ e^{-\beta \hat{H}_{\text{bag}}(\Delta, \rho)} \right].$$

(11)

where the latter trace is over all fermionic degrees of freedom. $\hat{H}_{\text{bag}}(\Delta, \rho)$ is the Bogoliubov-de Gennes Hamiltonian with a given set of $\Delta$ and $\rho$, where these vectors designate the set of values of these parameters on all lattice sites:

$$\hat{H}_{\text{bag}} = \sum_{i,\sigma} (\epsilon_i + \rho_\sigma) c_i^{\dagger} c_i - \sum_{\langle i,j \rangle,\sigma} (t_{ij} c_i^{\dagger} c_j + t_{ij}^{\ast} c_j^{\dagger} c_i)$$

$$+ \sum_i (\Delta_i c_i^{\dagger} c_i^{\dagger} + \tilde{\Delta}_i c_i c_i) + \sum_i |\Delta_i|^2 + \rho_\sigma^2.$$  

(12)

Given the explicit form of the diagonalizable BdG Hamiltonian, we can calculate expectation values and correlation functions,

$$\text{Tr}[\hat{\rho} \hat{O}] = \int \mathcal{D}(\Delta, \rho) e^{-\beta E_0} \sum_{n=1}^N e^{-\beta E_n} \langle n | \hat{O} | n \rangle,$$

(13)

where the sum is taken over all positive eigenvalues (quasi-particle excitations) of the BdG Hamiltonian. In our simulations we calculate the current and so replace $\hat{O}$ with the expression for the current in Eq. (4). Here $E_0$, $E_n$, and $|n\rangle$ are the ground-state energy, excitation energies, and excitation wave functions, respectively, for the BdG Hamiltonian, for the specific configuration of $\Delta$ and $\rho$. It is straightforward to see that in this case, the saddle-point approximation of the partition function gives rise to the mean-field BdG.
eq

equations (and then \( \Delta \) indeed corresponds to \(|U|(|c_{\downarrow}|c_{\uparrow})\)).

The calculation of the full integral, using the (classical) Monte Carlo approach,\(^{34}\) includes also the contributions of thermal fluctuations of the amplitude and phase of the order parameter. In using the Monte Carlo summation, we take a statistical average over different field configurations, which introduces statistical uncertainty into our observables. This uncertainty gives rise to the 1\(\sigma\) error bars on the numerical results in this paper. In Appendix B we detail how we improve on contemporary methods to perform the Monte Carlo calculation in \(O(N^{1.9}M^{1/3})\) time, where \(N\) is the number of sites and \(M\) the order of a Chebyshev expansion.

IV. APPLICATIONS

We have derived a new expression for the current flow through a superconductor, and demonstrated how to calculate the current in mesoscopic systems. Before applying it to understand and predict novel phenomena, it is important to verify it across a variety of exemplar systems, where one can compare against well-established theories. At the same time, as we are using \textit{ab initio} methods we present the first comparison between the underlying microscopic variables and the phenomenological parameters often used to describe these effects. As the main novelty of the approach is the inclusion of thermal fluctuations, we pay particular attention to verifying the formalism in two dimensions, especially looking for signatures of the BKT transition driven by phase fluctuations. In Sec. IV A we probe the current through a clean superconductor, and check that we can recover the BTK results for the contact resistance. A key effect in such systems is Josephson tunneling, so in Sec. IV B we study the temperature dependence of the resistance of a single Josephson junction. Dephasing (by temperature averaging) and decoherence (by electron-electron interactions) are studied in Sec. IV C. In Sec. IV D we examine the temperature dependence of the resistance in the vicinity of the BKT transition and compare to analytical results. In Sec. IV E we introduce finite magnetic field (flux) and probe the Little-Parks effect in the presence of disorder. Finally, in Sec. IV F we demonstrate how plotting maps of the current and potential across the system can illuminate the microscopic processes at the superconductor-insulator transition, finding that the rise in resistance is driven by the emergence of weak links. Throughout we use an attractive interaction of \(U = 1.6t\) to describe the SC region. To avoid the Van Hove singularity at half filling\(^{35}\) we study systems at an average 38.7% filling, except for Sec. IV C, where we focus on wires with a low filling fraction of 20%. All the calculations were carried out in the linear response regime, with a small potential difference of \(eV = 0.02t\). These parameters gave a typical superconducting order parameter at \(T = 0\) of \(\Delta = 0.32t\). Systems were typically two dimensional, so a single lattice site thick, 12 lattice sites wide, and 48 lattice sites long.

A. Clean systems

Well below the BKT transition, where thermal fluctuations of the pair amplitude and phase may be neglected, the resistance through a clean SC region is solely due to the contact resistance at the two interfaces. This resistance has been calculated by Blonder, Tinkham, and Klapwijk.\(^{11}\) By assigning a tunneling strength \(1/Z\) to the barriers, they showed that if the intermediate sample is in the normal state, then the current is purely due to electrons tunneling across the barrier, and the transmission coefficient is given by \(1/Z\).\(^{211}\) On the other hand, if the sample is in the SC state, then the SC gap, \(\Delta\), inhibits electrons from directly tunneling into it. Instead, these electrons Andreev tunnel accompanied by a hole. For a large barrier \(Z \gg 1\) the transmission coefficient becomes \(\Delta^2/4Z^4(\Delta^2 - E^2)\),\(^{11}\) where \(E\) is the electron energy. Electrons with an energy outside of the gap can either tunnel alone with a corresponding normal transmission coefficient \((E + \sqrt{E^2 - \Delta^2})/(2Z^2\sqrt{E^2 - \Delta^2})\), or Andreev tunnel with an accompanying hole, and have a transmission coefficient of \(\Delta^2/4Z^4(E^2 - \Delta^2)\). We first compare the results of our numerical calculations to these BTK formulas, and then demonstrate that for the simple case of a single SC site, the BTK results can be derived analytically from our current formula. Second, we go beyond the low-temperature regime accessible with standard theory by studying the situation at higher temperature where fluctuations become important.

To verify that the model recovers the correct behavior at the tunneling barrier, we focus on the weak coupling limit. In this limit, once a Cooper pair tunnels through the first barrier, it has an equal probability of continuing to either the left or the right lead, and consequently the current through the double barrier will be half that of a single barrier.\(^{27}\) For a long enough system the finite bias and temperature smear any Fabry-Perot type interference. To study the effect of the changing order parameter \(\Delta\), we focus first on a low-temperature system, where \(\Delta\) is indeed equal to the pair correlation \(|U|(|c_{\downarrow}|c_{\uparrow})\), vary the interaction strength \(U\), and monitor the various components of the tunneling current. For a pristine system with \(W = 0\), all of the resistance stems from the two tunneling barriers, and we verified that the current flow was independent of the length of the SC region. A relatively large potential bias of \(eV = 0.1t\) was applied across the leads. This allows us to explore all tunneling processes, either for \(\Delta < eV\) or \(\Delta > eV\) by changing the interaction parameter \(U\) and as a result \(\Delta\). Our results for the conductance are depicted in Fig. 2(a), and \(Z \approx 1.4\). At \(\Delta = 0\) the current is entirely normal. As shown in Fig. 2(a), increasing \(\Delta\) gives rise to a resonance in the Andreev current when \(E = \Delta\). At the same time the normal current falls as fewer electrons can be directly injected outside of the SC gap. As \(\Delta\) increases still further, so that the SC gap exceeds the chemical potential difference, resonant electrons are no longer injected into the divergent density of states at the SC gap, and the Andreev current falls. In agreement with the BTK calculation, at large \(\Delta\) the Andreev current adopts its final value, \(1/4\) of the normal \(\Delta = 0\) conductance, and no normal current flows.

Figure 2(b) depicts our results for higher temperatures, signaling the limitation of the BTK formalism. The numerical results are compared to the prediction of the BTK formalism, where the effects of temperature are taken into account in the mean-field reduction in \(\Delta\) and the broadening of the lead distribution function. The evident reduction of the calculated resistance, compared to the BTK prediction, indicates the importance of thermal fluctuations in determining the resis-
average superconducting order parameter

In the negative-clean superconductors (Hubbard model by an intermediate region consisting of two fluctuation, and the lower (b) at $T = 0$ showing a marked difference between the mean-field BTK formula and the numerical data that include thermal fluctuations. In Sec. IV D we discuss in more detail the temperature dependence of the resistance due to phase fluctuations.

As a last point in this section, we now demonstrate explicitly that in the weak coupling limit $Y \ll 1$, for the special case when the SC region consists of a single site, the BTK results can be derived from our formalism straightforwardly. In the linear response regime where a potential $V$ is put across the sample such that injected electrons are entirely within the SC gap, we find that the normal current is zero and we recover the analytic result for the Andreev current, $J = e^2V\Delta^2/8hZ^2(\Delta^2 - \mu^2)$, where $Z = \sqrt{\mu/\pi V/Y}$, $\mu$ is the chemical potential, and $v$ is the density of states at the Fermi surface. If the sample is normal we find that there is no Andreev current, and the normal current is $J = e^2V/2hZ^2$. These results are what would be expected from the BTK formalism, and coupled with the numerical results they confirm that the formalism properly treats tunneling between the leads and the SC sample.

B. Josephson junction

Another simple example that we wish to explore is a single Josephson junction, which will be modeled in the negative-$U$ Hubbard model by an intermediate region consisting of two clean superconductors ($W = 0$), between which we insert a weak link where the nearest-neighbor hopping element $t'$ is small ($t' \ll t$), see Fig. 3. Studying this system will allow us to probe how a phase difference across a barrier can affect the current flow through it. We first set $t' = 0$ to disconnect the left- and right-hand sides, and numerically evaluate the current through the central region. This is by no means trivial. The current formula, through the anomalous Green’s function, allows an absorption of a pair from the incoming lead into the condensate on one side of the barrier, and an emission of another pair into the outgoing lead. This current, however, will depend on the phase difference between the SC order parameter on the two sides of the barrier. For $t' = 0$, i.e., an infinite barrier, the phases of the left- and right-hand order parameter are uncorrelated, and thus all phase differences are degenerate in energy. Therefore, the current vanishes, but only after averaging over all states, which is done automatically in our numerical procedure. In the other limit, when the hopping matrix elements are the same as the hopping through the rest of the superconductor, $t' = t$, the phase of the superconductor is locked, so we see the standard free SC current flow.

We now model the situation with a moderately sized central barrier. This splits the superconductor in two, but crucially a Josephson supercurrent flows between the two sides, thus allowing the current to flow with no additional resistance. The current $J(T) = J_\text{J}(T)\cos(\phi_L - \phi_R)$ is maintained by the...
phase difference $\phi_L - \phi_R$ between the left- and right-hand superconductors, and the maximum value of the dissipationless current is the critical Josephson current $J_J(T) = (\pi |\Delta|/2eR_n)\tanh(|\Delta|/2k_BT)$, where $R_n$ is the resistance of the central barrier when the system is in the normal state.

To numerically study the thermally driven disruption of the Josephson current, it is vital that this breakdown occurs before the BKT transition occurs, which as we show in Sec. IV D, by itself reduces the current flow through the system. We thus use a small hopping element for the tunneling barrier of $t' = 0.005t$, which has a large $R_n$ and therefore small Josephson current $J_J$. In Fig. 3 we show the current as a function of temperature, in the presence of the weak link. When there is no voltage drop across the Josephson junction, the current $J_M$ that flows through it is given by $J_M = V/R$, where $R$ is the contact resistance to the normal leads. This current is maintained as long as the critical Josephson current $J_J$ is larger than $J_M$. As temperature is increased, thermal fluctuations will weaken the phase lock between the two SC regions, and the critical current is reduced. When $J_J$ is reduced below $J_M$, a finite voltage develops across the Josephson junction. This drives the phase difference across the junction to increase with time, which in turn leads to an oscillating current. This current has a non-zero time average,1 leading to a total resistance $R_J(T) = R/(1 - \sqrt{1 - \lambda^2})$, where $\lambda = J_J/J_M < 1$.1 This time-averaged current is exactly the quantity calculated in our Monte Carlo procedure. Figure 3 depicts a comparison between this simple model and our full numerical calculation, with reasonable agreement.

The critical current can be modified by varying the resistance of the central barrier, $R_n$. The intermediate case has $R_n = 0.075h/e^2$, the stronger coupling with $R_n = 0.06h/e^2$ is obtained by lowering the barrier to $t' = 0.01t$, and the weaker coupling with $R_n = 0.085h/e^2$ by widening the original barrier ($t' = 0.005t$) to four lattice sites. This wider barrier weakens the coupling between the superconductors so the Josephson resistance emerges at a lower temperature. A lower barrier strengthens the coupling thereby raising the temperature required for the emergence of resistance. Both these regimes are consistent with the simple model. We also verified that at very strong coupling where the temperature required for the breakdown of phase coherence becomes of the order of the BKT transition temperature, this simple model for the current flow no longer captures the full physics of the system. This study validates that our formalism can correctly model the presence of the Josephson supercurrent across the weak link introduced into the superconductor, and can therefore be used to model disordered systems that may contain multiple SC grains.

C. Decoherence and dephasing

The issue of decoherence and dephasing plays a significant role in transport at low temperatures. Here we define decoherence as the many-body phenomenon that leads to the loss of coherence via interactions among the electrons or interactions with the environment. On the other hand, dephasing can occur in a noninteracting system, and emerges from the fact that due to the finite temperature, electrons possess a range of energies, of the order of $k_BT$. Electrons of different energies acquire different phases along their respective trajectories, and if these phases differ by $2\pi$ or more when their energy changes by $k_BT$, then interference phenomena will average out to zero.

Decoherence. The effects of decoherence due to electron-electron interactions are more profound in one-dimensional wires in the normal phase. Since the original Hubbard model employed in this calculation [Eq. (9)] is an interacting model, one expects decoherence to arise naturally from the calculation. However, though the original formula for the current is exact, the approximation employed above [Eq. (13)] does not include quantum fluctuations. This means that it neglects the imaginary component of the self-energy which corresponds to damping due to interactions, and the resulting decoherence. To test the effects of such decoherence due to many-body interactions, we introduce into the normal Green’s function for momentum $k$ (as here we study a wire in the normal phase), by hand, the self-energy

$$\lim_{\delta \to 0} \frac{U^2}{2\pi^2} \sum_{p,q} n(\xi_p)[1 - n(\xi_q)][1 - n(\xi_{p+q})] \delta_{p,q} - \xi_{p-q} - \xi_{p+q} = i\delta,$$

which is the lowest-order contribution to the single-particle self-energy, and where $\xi_p$ are the momentum energy eigenstates of the Hamiltonian.

A similar approach37 has been applied to interacting electrons in a continuous one-dimensional system with repulsive contact interactions (the second-order contribution to the self-energy does not depend on the sign of the interaction). In this case, it has been shown, for wires with parabolic dispersion and chemical potential $\mu$, that this damping leads to a change in the distribution function and reduction in the conductivity by a factor of $1 - \pi^2(k_BT/\mu)^2L/12[L + \ell\exp(\mu/k_BT)]$,35 where the wire length $L$ is long enough that the smearing of the Fermi surface due to scattering (which occurs over the relaxation length scale $\ell$) outweighs that due to temperature. For sufficiently long wires $L \gg \ell\exp(\mu/k_BT)$, the reduction in conductivity becomes length independent, $1 - \pi^2(k_BT/\mu)^2/12$.

To compare our work against this theory (which relies on the parabolic dispersion), we focus on a system with a low filling fraction of 20%, near the bottom of the band, and set the disorder to $W = 0.1t$. We employ the perturbative expression for the current, Eq. (6), to give us access to long wires with $L \gg \ell$. In the upper panel of Fig. 4 we show the fall in current, as a function of length, due to the inclusion of self-energy at $k_BT = 0.17\mu$, here $\ell \approx 27t$. The overall change of $-0.5\%$ is small due to the Pauli blocking of scattering processes near the Fermi energy. We see reasonable agreement with the model over a range of length scales. The middle panel of Fig. 4 depicts the change in current with temperature for a system of a fixed length. We highlight the agreement to the expected variation in the fall in conductance with temperature.35 At low temperatures ($k_BT \ll \mu$) the damping is severely Pauli blocked so the characteristic damping length scale exceeds the system length, and the current correction $1 - \pi^2(k_BT/\mu)^2\exp(-\mu/k_BT)L/12\ell$ is exponentially suppressed. As temperature increases, the Fermi liquid $T^2$ behavior starts to dominate the correction to the current. At high, usually unphysical temperatures ($k_BT \gg \mu$), numerics see a smaller current shift than predicted by theory.
as the details of the specific Hubbard band dispersion versus the parabolic dispersion in which the model was developed become important.

In the lower panel of Fig. 4, we examine the effect of a SC phase on decoherence. At low temperature, the presence of the SC gap suppresses many-body scattering processes. However, when the temperature is raised above the BKT phase transition, scattering events are possible, though they have a smaller impact on the current than in the normal phase, due to the still finite local pair correlations. Above the mean-field BCS phase transition the current follows the expected parabolic profile as in the normal phase. Thus we have demonstrated that while quantum fluctuations, as they affect decoherence, can be taken into account in our formalism, their effect on the current, for the range of parameters studied here, is usually small at \( \lesssim 1\% \). We are thus justified in neglecting them in this study.

**Dephasing.** Having observed decoherence in the sample, we now turn to study dephasing due to thermal averaging. To verify that our formalism captures this important phenomenon, we study the length dependence of the conductance in noninteracting systems. We first verified, for a noninteracting clean system, that the net macroscopic current increases noninteracting one-dimensional wire with disorder \( W = 0.2t \) at two different temperatures. The straight (red) trend lines show a linear drop off in conductance with length, and the curved (green) an exponential decay.

**D. Variation of resistance with temperature**

We have now verified that our formalism captures the basic phenomena of contact resistance in Sec. IV A, Josephson coupling in Sec. IV B, and dephasing and decoherence in Sec. IV C. With these key tests complete, we are now
undergoes a BKT transition characterized by the emergence of vortices across the system, leading to the loss of global phase coherence. At a higher (mean-field) temperature, the SC order is completely suppressed and the system loses the SC correlation even locally. To study how this transition is reflected in the current flow, we performed numerical simulations on a two-dimensional SC system at several different temperatures. Simulations were performed for two different levels of disorder, $W = 0.1t$ and $W = 0.2t$, to determine how the transition and current flow are modified by the normal-state resistance, and extrapolated over length to remove the effects of the contact resistance [Fig. 6(b)].

At temperatures below the BKT transition, in Fig. 6(a), the resistance is zero as the system is in the SC state. At temperatures above the BKT transition, vortices and antivortices can easily unbind, though they may be partially pinned by disorder. The finite conductance $G$ of a sample in this case has been shown by Halperin and Nelson\textsuperscript{21} to be given by

$$G = 0.37G_n(\xi_c/\xi)^2,$$  \hspace{1cm} (15)

where $G_n$ is the normal-state conductance, $\xi_c$ is the SC coherence length, and $\xi$ is the SC order correlation length, which diverges at $T_{KT}$. The critical behavior at temperatures near the BKT transition $T \gtrsim T_{KT}$ leads to the conductance

$$G = 0.37G_n b^{-1} \exp[b(T_c - T_{KT})/(T - T_{KT})].$$  \hspace{1cm} (16)

where $b$ is a number of order unity. At temperatures higher than the (renormalized) mean-field critical temperature $T_c$, the conductance is given by the Aslamasov-Larkin theory\textsuperscript{21,40}

$$G = 0.37G_n(T_c - T_{KT})/(T - T_{KT}).$$  \hspace{1cm} (17)

Finally, we can also estimate the crossover between these two regimes by noting that the difference between the Kosterlitz-Thouless and mean-field transition temperatures critical regime is given by\textsuperscript{21}

$$T_c - T_{KT} \approx 0.17e^2T_c/hG_n.$$  \hspace{1cm} (18)

The difference between these two temperatures therefore widens with falling normal-state conductance.

In Fig. 6(a) we depict the variation of resistance with temperature above the BKT transition, showing the two types of dependence on temperature as is expected by theory. At $W = 0.1t$ we have $T_{KT} = 0.0091t$, $T_c = 0.11t$, $G_n = 17.6e^2/h$, and $b = 0.47$, whereas at $W = 0.2t$ we have $T_{KT} = 0.0046t$, $T_c = 0.16t$, $G_n = 22.2e^2/h$, and $b = 0.26$. The rising disorder reduces the normal-state resistance $G_n$, and also enhances the difference between the Kosterlitz-Thouless and mean-field transition temperatures, which agrees with Eq. (18) within 20%. The high-temperature Aslamasov-Larkin expression for the conductance persists well above the the mean-field critical temperature, where the insets show that the SC state has been totally suppressed. Finally, when the temperature is of the same order as the bandwidth, $k_BT \sim t$, the resistance in Fig. 6(a) increases superlinearly as the Fermi distribution becomes smeared across the whole band structure. Thus, we have been able to demonstrate that our formalism can calculate the resistance from a microscopic model, allowing us to derive the phenomenological parameters from the model.

FIG. 6. (Color online) (a) Variation of resistance with temperature for two different values of disorder calculated numerically (points). The red solid line shows the theoretical low-temperature behavior, and the blue dotted line the theoretical high-temperature behavior. The dashed vertical green lines show the BKT $T_{KT}$ and mean-field $T_c$ temperatures. The insets show the variation of the superconducting order parameter $U(c\uparrow c\downarrow)$ renormalized by its $T = 0$ value. (b) Numerical results (black points) and the deduced linear length dependence (red solid line) of the resistance for three different temperatures (i) $T < T_{KT}$, (ii) $T_{KT} < T < T_c$, and (iii) $T > T_c$. The $L \rightarrow 0$ values correspond to the contact resistance, which is eliminated in the values plotted in (a).


E. Little-Parks effect

Varying an applied magnetic field has long been an important experimental probe of the properties of a superconductor. It is therefore imperative to verify that the current formula developed here, coupled with the Hubbard model for the superconductor, is able to accurately model the effects of an applied magnetic field. In the Hubbard model the effects of the magnetic field are incorporated, via the Peierls substitution, into the phases of the hopping elements, $t_{ij} \rightarrow t_{ij} e^{2\pi i \phi_i/\phi_0}$, where $\phi_0 = \hbar c/e$ is the (single-electron) quantum flux, and the phases $\phi_i$ are defined such that their integral over a closed trajectory is equal to the magnetic flux threading the surface spanned by the trajectory.

To check whether this procedure captures the effect of an orbital magnetic field, we apply it to a clean hollow cylindrical superconductor, of radius $r$, such as that shown in Fig. 7, threaded by magnetic flux $\phi$. As demonstrated by Little and Parks, the flux suppresses superconductivity and the transition temperature falls periodically with the flux. This is often probed by measuring the falling conductance of the cylinder near to the SC transition temperature. The energy of electrons in the cylinder increase with trapped flux $\phi$ as $\hbar^2(n + 2\phi/\phi_0)^2/2mr^2$, where the integer $n$ is chosen to minimize the energy. This results in a periodic parabolic variation of the electron energy with flux and thus a parabolic periodic oscillation in the SC transition temperature $\Delta T_c = \hbar^2(n + 2\phi/\phi_0)^2/16mr^2$. Therefore, for a cylinder held just above its SC transition temperature, the change in the transition temperature with increasing flux results in a similar variation of the conductance with flux, $G(\phi) = [G_0 - DG \min_n (n + 2\Phi/\phi_0)]$, leading to a minima in the conductance at every half integer Cooper-pair flux quantum $\phi = (n + 1/2)\phi_0/2$. This has indeed been observed experimentally.

In Fig. 7 we first take a cylinder held just above its mean-field SC transition temperature at $T = 0.1r > T_c \simeq 0.09r$ and numerically evaluate the conductance as a function of the magnetic flux. The reasonable agreement with the above formula (red line, with $G_0 = 23.5e^2/h$ and $DG = 2.8e^2/h$) demonstrates that the formalism correctly picks up the effects of an applied magnetic field. The deviation from the parabolic predictions of mean-field theory at every half flux quantum is due to thermal fluctuations, which are neglected by the standard mean-field approach. In order to explore the consequences of thermal phase fluctuations further, we take a cylinder held well below its mean-field SC transition temperature, but above the Kosterlitz-Thouless transition temperature at $T = 0.03t > T_KT \simeq 0.01t$. According to conventional mean-field theory the system is superconducting whatever the threaded flux, but when phase fluctuations are taken into account, thermally excited phase slips cause the system to have a finite resistance. As a phase fluctuations are enhanced by the flux, we again see a periodically varying conductivity, though here it is better fitted with a sinusoidal curve. The periodically varying conductivity has a significantly lower amplitude than we saw at higher temperature, showing that though the threaded flux can encourage phase slips the effect is less profound than when at the mean-field phase transition. The power of the formalism to explore the interplay between phase fluctuations, flux and disorder will be employed to study recent puzzling observations in future publications.

F. Current distribution maps

One important feature of our formalism is the new capability to map out the flow of both super and normal currents within a sample and the changes in chemical potential which drive that flow. Since we can now study the current flow around impurities in the sample and expose weak links with large potential drop, we should be able to probe phenomena in the disordered superconductor with unprecedented detail and trace their cause back to a microscopic mechanism. While applications of this formalism to the outstanding problems in this field will be described in future publications, in this section we aim to demonstrate the usefulness of the current and potential maps, first by further studying the Josephson junction with a superconductor containing a central normal region, and second by studying the temperature-driven superconductor-insulator transition in disordered systems. However, we will first verify our current mapping formalism by examining the site-by-site current conservation in a clean system. As the only sources and sinks of current are the two metallic leads, a consistent calculation should obey charge conservation for all of the inner sites of the sample. In Fig. 8(a) we show the average fractional error in conservation of current $\sum_{i=1}^{N} |\Delta J_i|/|J_i|N$ on each site as we vary the number of states $K$ included in the calculation out of a possible $N = 77$ states, as prescribed in the penultimate paragraph of Appendix B. We see that if only 5% of states are included, there is a 20% average leakage of
Due to the strong proximity effect, phase locked and predominantly a Josephson current flows for a narrow $U_0$ when the central $K/N$ included in the calculation of the current.

The electrical potential is dropped on the two contact barriers, and remains constant through the superconductor [upper panel in Fig. 9(b)]. The formalism allows us to monitor the current flow through the system to see it change from SC to normal in character as the intermediate normal region is widened, as shown in Fig. 8(b).

For a narrow $U = 0$ central region the two SC regions are phase locked and predominantly a Josephson current flows [lower panel in Fig. 9(a)]. Due to the strong proximity effect, the system is entirely SC with no reduction in conductance. The electrical potential is dropped on the two contact barriers, and remains constant through the superconductor [upper panel in Fig. 9(a)]. (For the present case of two equal contact barriers the potential in the SC is equal to the average of the chemical potential of the two leads). On the other hand, when the central $U = 0$ region is wide, $L_{\text{norm}} \gtrsim 4a$, the two SC regions are too weakly coupled for a Josephson current to flow, and instead a normal current flows between the two SC regions [lower panel in Fig. 9(b)].

This, in turn, introduces a new resistor into the sample and the conductance drops accordingly. Now the potential drop is mostly across the Josephson junction [upper panel in Fig. 9(b)]; the left-hand superconductor adopts, approximately, the potential of the left-hand lead and the right-hand superconductor that of the right-hand lead. Figure 8 depicts the dependence of the fraction of current, out of the total current, that flows through the intermediate region, changing from zero for a short normal region to unity for a long one. This situation is analogous to current flowing between SC grains in a disordered sample, and the analysis can reveal whether they are coherently coupled or decoupled if no normal current flows. Thus the formalism can be an important ingredient in the study of the origin of resistance in a disordered SC system, and will be used in a subsequent publication, to study the anomalous magnetoresistance observed in experiment.

![FIG. 8. (Color online) (a) Average fractional error in conservation of current $\Sigma_{i=1}^{N} |\Delta J_i|/J_i N$ on each site against the fraction of total states $K/N$ included in the calculation of the current. (b) The changing conductance (black line with arrow toward left) of a Josephson junction with width $L_{\text{norm}}$ of central normal region (left axis), and the fraction of the normal current $J_{\text{norm}}/J_{\text{total}}$ (blue dashed line with arrow toward right) flowing through the central region (right axis).](image)

![FIG. 9. (Color online) The upper panels in each figure show the potential difference $V(x)$ across the sample with total potential drop $V$. The lower panels show current maps for (a) short and (b) long normal intermediate regions. Supercurrent is shown by cyan darts and normal current by violet pointers; the arrow length corresponds to current magnitude and orientation to the direction of current flow. Color density corresponds to the order parameter $|\langle c_{\uparrow}c_{\downarrow}\rangle_0|/|\langle c_{\uparrow}c_{\downarrow}\rangle|$ across the sample with total potential drop $V$.](image)
at $T \approx 0.14 T_c$, there are weak-disorder driven fluctuations in the SC order parameter, but an almost uniform supercurrent. The potential drops mainly in the contacts, and in the sample it is equal to the average of the two leads with small random fluctuations. In Fig. 10(d) at $T \approx 2.3 T_c$, the SC order parameter practically vanishes, there is no supercurrent, and, due to the increasing resistance, only a small normal current flows through the sample. The potential, as expected for normal systems, decays linearly across the sample. At intermediate temperatures $T \approx T_c$ the current map [Fig. 10(c)] highlights the interplay of the normal and SC currents. There is a rough correlation between regions of vanishing SC order parameter and normal current [e.g., point (1)], on the one hand, and finite SC order parameter and supercurrent flow [e.g., point (2)], on the other. However, at point (3) a small normal region separates two SC regions, thereby forming an effective Josephson, resulting in a supercurrent flowing through the zero SC order region. By examining the equipotential lines we see that the normal regions, for example, point (1), are acting as weak links; whereas the potential drop over the SC regions is small. Thus the overall resistance of the sample is dominated by such weak links. The current and potential maps allow us to see the superconductor-insulator transition developing, and we plan to investigate in detail the relation of such a percolative picture to the Kosterlitz-Thouless transition, as was recently suggested.

V. DISCUSSION

In this paper we have developed an exact formula to calculate the current through a superconductor connected to two noninteracting metallic leads with an imposed potential difference. The formula was implemented with a negative-$U$ Hubbard model which included both phase and amplitude fluctuations in the SC order parameter. A new Chebyshev expansion method allowed us to solve the model and calculate the current in $O(N^{-1.9} M^{2/3})$ time, granting access to systems of unprecedented size. The formalism also enables the generation of current and potential maps which show exactly where the supercurrent and separately the normal current flow through the system.

The formalism was used to revisit a series of well-established results, allowing us to expose the link between the phenomenological parameters and underlying microscopic variables. This also demonstrated the accuracy of the procedure and its ability to capture various physical processes relevant to superconductivity in disordered systems, and to correctly model the presence of a magnetic field and finite temperature. These tests indicate that the formalism and accompanying numerical solver can robustly calculate the current through a superconductor across a wide range of systems. In the future we plan to report on the application of the formalism to several outstanding questions, such as the magnetoresistance anomaly on crossing the superconductor-insulator transition,\textsuperscript{44} the Little-Parks effect in nanoscale cylinders,\textsuperscript{42} and dissipation-driven phase transitions in SC wires.\textsuperscript{46}
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APPENDIX A: DERIVATION OF THE CURRENT FORMULA

The formula for the current in the Bogoliubov basis set is

\[
J = \frac{i e}{2 \hbar} \sum_{\sigma} \int d\epsilon \text{Tr}[f_{\downarrow}(\epsilon)P^{\downarrow} - f_{\uparrow}(\epsilon)P^{\uparrow}][u_{\sigma}(G_{\sigma}^{\uparrow} - G_{\sigma}^{\downarrow})u_{\sigma}^{\dagger} + v_{\sigma}(G_{\sigma}^{\uparrow} - G_{\sigma}^{\downarrow})v_{\sigma}^{\dagger} - \sigma \epsilon_{\sigma}^{\downarrow}(H_{\sigma}^{\uparrow} - H_{\sigma}^{\downarrow})u_{\sigma}^{\dagger} + \sigma \epsilon_{\sigma}^{\uparrow}(H_{\sigma}^{\downarrow} - H_{\sigma}^{\uparrow})v_{\sigma}^{\dagger}]
\]

\[
+ \text{Tr}[\Gamma^{\downarrow} - \Gamma^{\uparrow}][u_{\sigma}^{\dagger}G_{\sigma}^{\uparrow}u_{\sigma}^{\dagger} - v_{\sigma}G_{\sigma}^{\downarrow}v_{\sigma}^{\dagger} + \sigma \epsilon_{\sigma}^{\downarrow}H_{\sigma}^{\uparrow}u_{\sigma}^{\dagger} - \sigma \epsilon_{\sigma}^{\uparrow}H_{\sigma}^{\downarrow}v_{\sigma}^{\dagger}]].
\]

We need to determine Green’s functions across the sample, which must be calculated in the presence of the leads. However, as the electrons in the metallic leads are noninteracting, we can start from the bare electronic Green’s functions for the superconductor not coupled to the leads \(G_{\sigma}^{\uparrow}(m,n) = \delta_{m,n}/(\epsilon - \epsilon_{m} + i\delta)\) and \(G_{\sigma}^{\downarrow}(m,n) = \delta_{m,n}/(\epsilon + \epsilon_{m} + i\delta)\), which have energy eigenstates \(\epsilon_{m}\) and \(\delta \to 0^{+}\). We then write Dyson’s equation to self-consistently include the leads

\[
\begin{pmatrix}
G_{\sigma}^{\uparrow} \\
H_{\sigma}^{\downarrow}
\end{pmatrix} = \begin{pmatrix}
\tilde{G}_{\sigma}^{\uparrow} \\
0
\end{pmatrix} + V^{2}
\begin{pmatrix}
\tilde{G}_{co}(u_{\sigma}^{\dagger}v_{\sigma}^{\dagger}u_{\sigma}^{\dagger}p_{\sigma}g_{\sigma}^{\dagger}\chi_{p}u_{\sigma}^{\dagger} + p_{\sigma}g_{\sigma}^{\dagger}\chi_{p}v_{\sigma}^{\dagger}u_{\sigma}^{\dagger}) \\
\tilde{G}_{lo}(u_{\sigma}^{\dagger}v_{\sigma}^{\dagger}u_{\sigma}^{\dagger}p_{\sigma}g_{\sigma}^{\dagger}\chi_{p}v_{\sigma}^{\dagger} - p_{\sigma}g_{\sigma}^{\dagger}\chi_{p}u_{\sigma}^{\dagger})
\end{pmatrix}
\begin{pmatrix}
G_{\sigma}^{\uparrow} \\
H_{\sigma}^{\downarrow}
\end{pmatrix}.
\]

Here \(g_{\sigma}^{\dagger}\chi_{p} = 1/(\epsilon - \epsilon_{p} + \mu_{\sigma} + i\delta)\) is the retarded Green’s function of the noninteracting electrons in the leads, with dispersion \(\epsilon_{p}\), and \(\{u_{\sigma}, v_{\sigma}\}\) are the matrices of the eigenstates multiplied by the lead plane wave states \(p\) at the tunneling barriers. To extract the retarded Green’s function and its anomalous counterpart from this matrix equation, one has to perform a matrix inversion. The Dyson equation is for the retarded and advanced Green’s functions, whereas the current formula Eq. (A1) is in terms of the lesser and greater Green’s functions. To transform these into the retarded and advanced Green’s functions we apply the identity \(G_{\sigma} = \tilde{G}_{\sigma}^{+} + \tilde{G}_{\sigma}^{\Sigma}\Sigma_{\sigma}^{+}G_{\sigma}^{\Sigma} + \tilde{G}_{\sigma}^{\Sigma}\Sigma_{\sigma}G_{\sigma}^{\Sigma}\) recursively to find \(G_{\sigma} = (1 + \Sigma^{\Sigma})G_{\Sigma}^{\Sigma} + G_{\sigma}^{\Sigma}\Sigma_{\sigma}G_{\sigma}^{\Sigma}\), where \(\Sigma_{\sigma}\) is the self-energy. This recursion fixes the chemical potential of the superconductor by including tunneling to and from the leads. This will ensure that the net number of electrons is conserved, analogous to some extensions to the BTK formalism.27

However, as the final chemical potential must be independent of the chemical potential of the uncoupled superconductor, the term containing \(\tilde{G}_{\Sigma}\) must be identically zero, leaving \(G_{\sigma}^{\Sigma} = G_{\sigma}^{+} + \Sigma_{\sigma}G_{\sigma}^{\Sigma}\), and its greater Green’s function counterpart \(G_{\sigma} = G_{\sigma}^{+} + \Sigma_{\sigma}G_{\sigma}^{\Sigma}\).

We now extend this identity to include the anomalous Green’s function and recover

\[
\begin{pmatrix}
G_{\sigma}^{\Sigma} \\
H_{\sigma}^{\Sigma}
\end{pmatrix} = V^{2}
\begin{pmatrix}
\tilde{G}_{co}(u_{\sigma}^{\dagger}v_{\sigma}^{\dagger}u_{\sigma}^{\dagger}p_{\sigma}g_{\sigma}^{\dagger}\chi_{p}u_{\sigma}^{\dagger} + p_{\sigma}g_{\sigma}^{\dagger}\chi_{p}v_{\sigma}^{\dagger}u_{\sigma}^{\dagger}) \\
\tilde{G}_{lo}(u_{\sigma}^{\dagger}v_{\sigma}^{\dagger}u_{\sigma}^{\dagger}p_{\sigma}g_{\sigma}^{\dagger}\chi_{p}v_{\sigma}^{\dagger} - p_{\sigma}g_{\sigma}^{\dagger}\chi_{p}u_{\sigma}^{\dagger})
\end{pmatrix}
\begin{pmatrix}
G_{\sigma}^{\Sigma} \\
H_{\sigma}^{\Sigma}
\end{pmatrix}.
\]

We can now take this, the analogous expression for the greater Green’s function, and their anomalous counterparts, and substitute them into Eq. (A1), which will yield Eq. (4).

APPENDIX B: EVALUATION OF THE MONTE CARLO INTEGRALS

To evaluate the correlation functions [e.g., Eq. (13)], we need to sum over all possible spatial configurations of the auxiliary fields \(\rho\) and \(\{\Delta, \bar{\Delta}\}\), with each configuration carrying the weight \(P(\rho, \Delta) = \exp(-\beta E(\rho, \Delta))/Z\). This distribution is sampled using the Metropolis algorithm,34 which at each step proposes a new configuration of either the field \(\rho\) or \(\Delta\) and calculates the resulting change in the total energy. If this change in the energy is negative the step is accepted, whereas if positive it is accepted with probability \(\exp[-\beta (E(\rho_{\text{new}}) - E(\rho_{\text{old}}))]\) and \(\exp[-\beta (E(\Delta_{\text{new}}) - E(\Delta_{\text{old}}))]\) respectively. Since the walk over \(\rho\) is one-dimensional we choose the step size \(\Delta \rho_{\text{new}} - \Delta \rho_{\text{old}}\) to aim for 50% of the steps to be accepted, whereas the walk over \(\Delta, \bar{\Delta}\) covers a two-dimensional space so we choose a step size \(\Delta \bar{\rho}_{\text{new}} - \Delta \bar{\rho}_{\text{old}}\) so that 35.2% of the steps will be accepted.45 To verify the effect of fluctuations in the density field and to check the robustness of our Hubbard-Stratonovich decoupling scheme, in Fig. 11(e) we compare the conductance of the superconductor-insulator transition of Fig. 10 first with density fluctuations and second without density fluctuations but using simply a self-consistent but static mean-field value for the density. As the superconductor-insulator transition is driven by fluctuations in the phase of the SC order parameter, it is expected that density fluctuations will have little affect on the conductance. We see in Fig. 11(e) that density fluctuations reduce the conductance slightly, but overall the transition remains the same. This verifies that our Hubbard-Stratonovich decoupling scheme is robust against the inclusion of density fluctuations. We saw similar conclusions from checking other transitions from Sec. IV.

Central to the Monte Carlo method used to sample the partition function is the requirement to calculate the energy difference between two different configurations of the auxiliary fields, \(\{\rho_{\text{old}}, \Delta_{\text{old}}\}\) and \(\{\rho_{\text{new}}, \Delta_{\text{new}}\}\). For a lattice with \(N\) sites, to calculate the energy of each proposed configuration requires an effort of \(O(N^{3})\), so an entire sweep over the \(N\) sites that make up the fields \(\rho\) and \(\{\Delta, \bar{\Delta}\}\) requires a computational effort of \(O(N^{5})\). However, a recent method developed by Weiße47 calculates just the difference between the energy of the configurations in a computationally efficient manner. For an update to the \(i\)th site, a Chebyshev expansion with the 0 \(\leq m \leq\)
have calculated this site-by-site through a succession of sparse matrix-vector multiplications, each of cost $O(N M)$, so for an entire sweep over the order parameter the computational effort is $O(N^2 M)$. However, here we optimize the program so that the entire sweep can be performed in $O(N^{1.9} M^{2/3})$ time. Rather than follow a site-by-site approach calculated with sparse matrix-vector multiplications, we instead calculate the matrix elements for the entire sweep simultaneously, which necessitates performing matrix-matrix multiplications. Provided the changes in the order parameters are small the local changes are independent of those of surrounding sites and we can then perform the entire sweep from this data set. Spherical averaging further reduces the influence of changes in the surrounding order parameters. Central to the recursion relation for $T_{m}$ is the costly calculation of $x^{a}$, for $1 < n \leq M$. To evaluate this we divide the calculation of the $M$ matrix products into three stages:

1. The lowest-order matrix products, up to $x^{k}$, are sparse. Therefore, for the elements $1 < n \leq k$ the matrix multiplications involve only sparse matrices, each of peak cost $k N$, and the total cost of calculating them is $O(k^2 N)$.

2. The second stage is to successively calculate every $k$th matrix product. Each of these involves multiplying the dense matrix $x^{p}$ by the matrix $x^{k}$, for integer $1 \leq p \leq M/k$, which costs $O(N^{2.38})$ time. With $M/k$ of these products to calculate the total cost is $O(N^{2.38} M/k)$.

3. The third stage is to construct the entire family of $x^{a}$ by interpolating between the matrices $x^{p}$ found in the second stage. This is done by multiplying the dense matrices found in the second stage by the sparse matrices found in the first stage. Furthermore, as we need only the diagonal elements of the final matrix, each separately costs $O(kN)$ and so the total cost is $O(kNM)$.

Having laid out the prescription of how to calculate the matrix elements, we now examine the total cost, $O(k^2 N + N^{2.38} M/k + kNM)$. The choice $k \sim \sqrt{N^{1.38}M}$ will minimize the total cost to $O(N^{1.9} M^{2/3} + N^{1.46} M^{4/3})$, and as typically $N \gg M$ the cost is $\sim O(N^{1.9} M^{2/3})$. This is a significant improvement over the cost $O(N^2 M)$ of the Chebyshev expansion approach, which for the parameters employed in our simulations corresponds to a speedup by a factor of $\sim 30$. Now that the matrix elements behind the Chebyshev expansion have been found they are applied for the entire sweep.

To verify the Monte Carlo procedure in Fig. 11(a) we first check the convergence of the estimate for the current and that its standard error falls as the root of the number of Monte Carlo iterations. In Fig. 11(b) we compare the results of equilibrated Monte Carlo runs at zero temperature from a variety of initial configurations of the order parameter fields $\rho$ and $\Delta$. Evolution under the Metropolis algorithm drives these starting fields into different relaxed configurations, which because the simulations are restricted here to $T = 0$ are unable to be excited out to explore different configurations. These final configurations yield a variety of different current values, with standard deviation of $\sim \pm 2.4\%$ of the final total current. At finite temperature, thermal excitations can drive the system to explore configurations around the ground state with a narrower standard deviation of $\sim \pm 0.6\%$. Having verified the current statistics, in Figs. 11(c) and 11(d) we show the

\[ M \text{ coefficients containing } \langle i | T_{m}(H/s)| i \rangle \text{ must be calculated, where } T_{m} \text{ is defined by the recursion relation } T_{m}(x) = 2xT_{m-1}(x) - T_{m-2}(x), \text{ } T_{0}(x) = 1, \text{ and } T_{1}(x) = x. \text{ A typical expansion contained } M = 1024 \text{ terms. Previous authors}^{47} \]
results of some timing runs that highlight the improvement of the algorithm to $O(N^{1.9}M^{2/3})$ time over the standard approach of calculating all the energy eigenvalues in $O(N^4)$ time and the standard Chebyshev approach that runs in $O(N^2M)$ time. In particular, by varying the system size, we observe that the method of calculating all the eigenvalues is more efficient for systems smaller than $N \sim 10$, but the new Chebyshev approach is superior for large systems. We took advantage of this development to study large systems. In a typical simulation on a single 2.66-GHz Intel Core 2 processor we can accumulate statistics over 10 000 different field configurations for an 800-site system in approximately 10 h.

The Chebyshev expansion method just described represents a zero-order approximation. However, we can extend this method farther and calculate the lowest-order change in the Chebyshev expansion following a shift in the configuration of the fields $\rho$ and $\Delta$ by $\delta$. The resultant shift in the Chebyshev expansion of $T_i$ is found using the recursion relationships $t_i = \frac{1}{2} \delta T_{i-1} + \frac{1}{2} H t_{i-2} - t_{i-2}$ with $t_0 = 0$ and $t_1 = \Delta/s$. This allows the Chebyshev expansion coefficients to be extrapolated over several configuration space sweeps, and the calculation time falls proportionally. Spherical averaging also reduces the influence of changes in the surrounding order parameters.

In practice it was found that up to ten extrapolation steps could be performed, resulting in a code speedup of a factor of 10.

Though the Chebyshev approach can be used to direct the sampling of the system, to calculate expectation values, such as the current, it is necessary to diagonalize the system and determine the field configurations of its states. Formally this requires $O(N^3)$ time. However, since the current is dominated by the quasiparticle states near the Fermi surface, we instead adopt the implicitly restarted Arnoldi method$^{10}$ to calculate only those particular states. We are also helped by the sparsity of the matrix, which allows us to calculate $K$ eigenstates in $O(KN)$ time. It is usually necessary to calculate a certain fraction of the energy states, so $K \propto N$, and the total cost is $O(N^2)$. The eigenfunctions and energies can then be used to calculate the current for a specific realization of $\rho$ and $\Delta$ using the formalism described in Sec. II. It is then necessary to average over successive realizations of $\rho$ and $\Delta$. However, the contribution from successive Monte Carlo calculations might be serially correlated, which would result in an underestimated value for the uncertainty in the predicted value of the current. To correct for this we calculated the correlation time through the truncated autocorrelation function.$^{50}$ We find a typical correlation time of approximately six Monte Carlo steps, which without autocorrelation corrections would correspond to an underestimate in the uncertainty of a factor of $\sim 2.5$.

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If we consider the auxiliary fields in momentum space, a complete summation over momentum contributions from both $\Delta q$ and $\rho q$ would lead to a double counting of the interaction term. However, in the Monte Carlo calculation, we sample just the low-energy fluctuations in each decoupling channel that will give orthogonal contributions to the interaction term and avoid any double counting. In practice it was found that averaging over fluctuations in the $\Delta q$ field was important, and drove, for example, a Kosterlitz-Thouless transition. However, in Appendix B we verify that the average over fluctuations in the $\rho$ field made only a negligible quantitative change to the results.