Nonequilibrium-induced metal-superconductor quantum phase transition in graphene

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We study the effects of dissipation and time-independent nonequilibrium drive on an open superconducting graphene. In particular, we investigate how dissipation and nonequilibrium effects modify the semi-metal-BCS quantum phase transition that occurs at half-filling in equilibrium graphene with attractive interactions. Our system consists of a graphene sheet sandwiched by two semi-infinite three-dimensional Fermi liquid reservoirs, which act both as a particle pump/sink and a source of decoherence. A steady-state charge current is established in the system by equilibrating the two reservoirs at different, but constant, chemical potentials. The graphene sheet is described using the attractive Hubbard model in which the interaction is decoupled in the s-wave channel. The nonequilibrium BCS superconductivity in graphene is formulated using the Keldysh path integral formalism, and we obtain generalized gap and number density equations valid for both zero and finite voltages. The behaviour of the gap is discussed as a function of both attractive interaction strength and electron densities for various graphene-reservoir couplings and voltages. We discuss how tracing out the dissipative environment (with or without voltage) leads to decoherence of Cooper pairs in the graphene sheet, hence to a general suppression of the gap order parameter at all densities. For weak enough attractive interactions we show that the gap vanishes even for electron densities away from half-filling, and illustrate the possibility of a dissipation-induced metal-superconductor quantum phase transition. We find that the application of small voltages does not alter the essential features of the gap as compared to the case when the system is subject to dissipation alone (i.e. zero voltage). The possibility of tuning the system through the metal-superconductor quantum critical point using voltage is presented.

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I. INTRODUCTION

The landmark experimental realization of an isolated graphite monolayer, or graphene$^{1,2}$, has sparked intense theoretical and experimental interest in the material over the last few years$^{3}$. A source of interest in the study of graphene is the unique properties of its charge carriers. At low energies, these charge carriers mimic relativistic particles, and are most naturally described by the (2+1)-dimensional Dirac equation with an effective speed of light, $c \sim v_F \sim 10^6 \text{ms}^{-1}$. The fact that graphene is an excellent condensed-matter analogue of (2+1)-dimensional quantum electrodynamics (QED) has been known to theorists for over 20 years$^{4,5,6}$. However, it was not until the spectacular experimental realization of isolated graphene that experimentalists began observing signatures of the QED-like spectrum in their laboratories. Consequences of graphene’s unique electronic properties have been revealed in the context of anomalous integer quantum-Hall effect$^{7,8}$ and minimum quantum conductivity in the limit of vanishing carrier concentrations$^{9}$. In addition to its importance in fundamental physics, graphene is expected to make a significant impact in the world of nano-scale electronics. Research efforts in developing graphene-based electronics have been fueled by a strong anticipation that it may supplant the silicon-based technology which is nearing its limits$^{9}$. Graphene is a promising material for future nanoelectronics because of its exceptional carrier mobility which remains robustly high for a large range of temperatures, electric-field-induced concentrations$^{1,2,7,8}$, and chemical doping$^{9}$. Indeed, recent experiments have explored the possibilities of in-plane graphene heterostructures by engineering arbitrary spatial density variation using local gates$^{10,11,12}$. The application of local gate techniques to graphene marks an important first step on the road towards graphene-based electronics.

From a theoretical point of view realizing graphene nanoelectronics requires a theoretical understanding of open nonequilibrium graphene. Naturally, graphene in nanocircuits is subject to decoherence effects due to its coupling to external leads via tunnel junctions. Furthermore, a nonequilibrium treatment of graphene becomes necessary when a charge current is driven through it. To this date, effects of dissipation and nonequilibrium drive on graphene electronic properties have not been addressed. The focus of this paper is to show a theoretical framework in which these effects can be studied and illustrate how they give rise to striking influences on the equilibrium properties of graphene.

This work considers dissipation and nonequilibrium effects on superconducting graphene. Besides the possibility of superconductivity in graphene by proximity effect$^{13}$, some works suggested the potential of achieving plasmon-mediated singlet superconductivity in graphene$^{14,15}$. Several groups have investigated the equilibrium mean-field theory of superconductivity in graphene using the attractive Hubbard model on the honeycomb lattice. Uchoa and Castro-Neto$^{14}$ studied spin singlet superconductivity in graphene at various fillings by considering both the usual s-wave pairing as well as pairing with $p + ip$ orbital symmetry permit-
ted by graphene’s honeycomb lattice structure. Zhao and Paramekanti examined the possibility of s-wave superconductivity on the honeycomb lattice. Both works show that (in the absence of p-wave pairing) half-filled graphene displays a semimetal-superconductor quantum critical point at a finite critical attractive interaction strength $u_c$. Away from half-filling, the system exhibits Cooper instability at any finite $u$ and thus undergoes the usual BCS-BEC crossover as $u$ is increased. The difficulty in achieving superconductivity at half-filling is a result of the vanishing density of states at the Dirac point and the absence of electron screening.

In this work, the superconducting graphene sheet is subjected to dissipation and nonequilibrium drive by coupling it to two semi-infinite particle reservoirs via tunnel junctions. The geometry of the system is shown in Fig.1. While the two reservoirs are independently held in thermal and chemical equilibrium at all times, an out-of-plane steady-state current through graphene is established by equilibrating the reservoirs at two different, but constant, chemical potentials. The leads act as infinite reservoirs and are assumed to be held at a common temperature $T$ at all times. Nonequilibrium theory of BCS superconductivity is formulated using the Keldysh path integral formalism, and the resulting nonequilibrium mean-field equations are used to investigate the gap behaviour at and near half-filling for various attractive interaction strengths. The gap is plotted in the parameter space of filling $n$ and the interaction strength $u$ (see Fig.5), and our results can be directly compared to the gap phase diagram in Fig.2 of the work by Zhao and Paramekanti.

Our main results are now qualitatively summarized. We find that the gap is generally suppressed in the presence of leads. As the paper will discuss in detail, the key to understanding our findings is to notice that the dissipation of electrons into the leads act as a pair-breaking mechanism for the Cooper pairs in the central graphene sheet. This mechanism, and hence the suppression, is present at both zero and finite voltages and for all electron densities. As a consequence, the Fermi liquid ground state of the system remains stable against Cooper pairing up to some density-dependent finite attractive interaction strength $u_c(n)$ at all densities. With respect to the gap phase diagram, dissipation gives rise to a finite region around half-filling in which the gap vanishes (see Fig.5). From these results, we infer that dissipation induces a metal-superconductor quantum phase transition at all fillings, for which the tuning parameter is the attractive interaction strength $u$. The qualitative behaviour of the gap is not appreciably different in the zero and finite voltage cases as long as the voltage is small, i.e. $V \ll \Gamma$, where $\Gamma$ denotes the average tunneling rate of electrons between graphene and the two leads. Finite voltage modifications, however, result due to voltage-induced changes in the graphene electron density.

The paper is organized as follows. In Sec. II, we introduce the Hamiltonian which models our heterostructure. The mean-field treatment of the model is formulated on the Keldysh contour in Sec. III. In Sec. III B, the nonequilibrium gap and number density equations will be derived. The results are presented in Sec. IV. The effects of dissipation in the absence of voltage is discussed in Sec. IVA while the finite voltage effects are included in Sec. IVB. We conclude in Sec. V.

### II. MODEL

The lead-graphene-lead heterostructure considered in this work is shown in Fig.1. Graphene is located on the $z = 0$ plane, and each of its sites is labeled using two coordinates $x_i = (x_i, y_i, z_i \equiv 0)$. The semi-infinite metallic leads extend from both sides of the graphene sheet for $z > 0$ and $z < 0$. We assume the leads are separated from graphene by thin insulating barriers, and the tunneling of electrons through each of the barriers can be described by phenomenological tunneling parameters. Full translational symmetry is present along the planes parallel to the $xy$-plane for $z \neq 0$ while only the discrete translational symmetry of the graphene lattice is present at $z = 0$. The leads are assumed to be in thermal equilibrium with their continuum of states occupied according to the Fermi-Dirac distribution, $f_\alpha(\omega) = \left[ 1 + \exp \left( \beta (\omega - \mu_\alpha) \right) \right]^{-1}$, where $\alpha = L$ (left), $R$ (right) labels the leads. An electric potential bias is set up in the out-of-plane direction by tuning the chemical potentials of the leads to different values.

![FIG. 1: A schematic of the system considered. Chemical potential mismatch in the two leads will lead to a charge current parallel to the $z$-axis.](image)

The Hamiltonian consists of three parts,

$$H = H_{\text{sys}} + H_{\text{res}} + H_{\text{sys-res}}.$$  \hspace{1cm} (1)

The central graphene sheet is modeled using the attractive Hubbard model on the honeycomb lattice. The kinetic term is a tight-binding description for the $\pi$-orbitals of carbon that includes nearest- and next-nearest-neighbour hopping processes. The on-site interaction strength is parametrized by $U$. The Hamiltonian
for the layer is

\[
H_{sys} = -t \sum_{(i,j),\sigma} (c_{i,\sigma}^\dagger c_{j,\sigma} + h.c.) - t' \sum_{(i,j)} (c_{i,\sigma}^\dagger c_{j,\sigma} + h.c.) - U \sum_i e_{i,1}^\dagger e_{i,1} e_{i,1}^\dagger e_{i,1} .
\]  

\[ (i,j) = \quad (2) \]

c\textsubscript{i,\sigma} (c\textsubscript{i,\sigma}) creates (annihilates) electrons on site \( r\textsubscript{i} \) of the graphene honeycomb lattice with spin \( \sigma \) (\( \sigma = \uparrow, \downarrow \)). \( U \) is assumed positive due to attractive interaction, and \( t \) and \( t' \) are the nearest- and next-nearest-neighbour hopping parameters, respectively. Specific values for \( t \) and \( t' \) have been estimated by comparing a tight-binding description to first-principle calculations. Following their estimates, we take \( t = 2.7 \text{eV} \) and fix \( t'/t = 0.04 \).

The honeycomb lattice can be described in terms of two inter-penetrating triangular sublattices, \( A \) and \( B \) (see Fig.2). Each unit cell is composed of two atoms, one each of type \( A \) and type \( B \). Primitive translation vectors, \( e\textsubscript{1} \) and \( e\textsubscript{2} \), are

\[
e\textsubscript{1} = (\sqrt{3}, 0) \quad e\textsubscript{2} = (-\sqrt{3}/2, 3/2) \quad e\textsubscript{3} = e\textsubscript{1} + e\textsubscript{2} ,
\]

where they are expressed in units of \( a \), the distance between two nearest carbon atoms. Any \( A \) atom is connected to its nearest neighbours on the \( B \) lattice by three vectors

\[
d\textsubscript{1} = (0, 1) \\
d\textsubscript{2} = (-\sqrt{3}/2, -1/2) \\
d\textsubscript{3} = (\sqrt{3}/2, -1/2).
\]

In momentum space, the kinetic term reads

\[
H^K_{sys} = \frac{1}{N_\Delta} \sum_{k,\sigma} \left( a_{k,\sigma}^\dagger b_{k,\sigma}^\dagger \right) \left( \lambda_k g_k^\dagger \right) \left( a_{k,\sigma} b_{k,\sigma} \right) ,
\]

where

\[
\lambda_k = -t' \sum_{i=1}^3 e^{i \mathbf{k} \cdot \mathbf{d}_i} + c.c. \\
g_k = -t \sum_{i=1}^3 e^{i \mathbf{k} \cdot \mathbf{d}_i} .
\]

Components of the pseudospinor, \( a_{k,\sigma}^\dagger \) and \( b_{k,\sigma}^\dagger \), describe quasiparticles that belong to sublattice \( A \) and \( B \), respectively. Here, \( N_\Delta \) denotes the number of lattice sites in a triangular sublattice. \( N = 2N_\Delta \) will denote the total number of sites on the honeycomb lattice.

Coupling between leads and the graphene sheet is modeled using the following Hamiltonian,

\[
H_{sys-res} = \int \frac{dk_z}{2\pi} \sum_{\alpha=L,R} \sum_{i,\sigma} \zeta_\alpha \left( A_{i,\sigma,\alpha,k_z}^\dagger c_{i,\sigma} + h.c. \right) .
\]

\[ (2) \]

\( \zeta_\alpha \) is a phenomenological tunneling matrix that describes the tunneling of an electron between site \( i \) on the graphene sheet and an adjacent site on lead \( \alpha \) (see Fig.3). We only consider lead-graphene tunneling processes in which \( (x, y) \) coordinates of the electron in the initial and final states are the same. This assumption simplifies various computational steps without altering the qualitative features of the final results. \( C_{i,\sigma,\alpha,k_z}^\dagger \) creates an electron in lead \( \alpha \) at coordinates \( (x_i, y_i) \) with spin \( \sigma \) and longitudinal momentum \( k_z \). We assume here that the tunneling parameters are independent of frequency and momentum but maintain their lead dependence in order to describe possible asymmetries in the lead-layer couplings. In momentum space, the tunneling Hamiltonian Eq.8 becomes

\[
H_{sys-res} = \sum_{\alpha} \zeta_\alpha \int \frac{dk_z}{2\pi} \frac{1}{N_\Delta} \sum_{k,\sigma} \left( A_{k,\sigma,\alpha,k_z}^\dagger a_{k,\sigma} + B_{k,\sigma,\alpha,k_z}^\dagger b_{k,\sigma} + h.c. \right) .
\]

\[ (9) \]

The inplane momentum, \( k \), is the component of momentum parallel to the graphene plane and the out-of-plane momentum, \( k_z \), is its component normal to the plane. \( A_{k,\sigma,\alpha}^\dagger \) (\( B_{k,\sigma,\alpha}^\dagger \)) corresponds to an electron mode...
propagating in "sublattice $A(B)$" in lead $\alpha$ with spin $\sigma$ and wavevector $\mathbf{k}$. Although the full inplane translational symmetry of the leads implies that $\mathbf{k}$ can take on any value in $\mathbb{R}^2$ the tunneling assumption (see Fig. 3) tells us that the only modes that tunnel are those with $\mathbf{k}$ values that are the allowed modes of the triangular sublattices in the graphene sheet. All other inconsequential modes can eventually be integrated out in the path integral sense and will merely contribute a multiplicative factor in front of the partition function. Therefore, we will not consider these modes further.

Both leads are assumed to be Fermi liquids

$$H_{\text{res}} = \sum_{\mathbf{k},\sigma} \frac{1}{N_\Delta} \int d\mathbf{z} \sum_{\mathbf{k},\mathbf{z},\sigma} \epsilon_{\mathbf{k},\mathbf{z}} \left( A_{\mathbf{k},\mathbf{z},\sigma,\mathbf{z},\sigma} A_{\mathbf{k},\mathbf{z},\sigma,\mathbf{z},\sigma} + B_{\mathbf{k},\mathbf{z},\sigma,\mathbf{z},\sigma} B_{\mathbf{k},\mathbf{z},\sigma,\mathbf{z},\sigma} \right),$$

(10)

with a separable dispersion

$$\epsilon_{\mathbf{k},\mathbf{z}} = \epsilon_{\mathbf{k}} + \epsilon_{\mathbf{z}} = \frac{|\mathbf{k}|^2}{2m_e} + \frac{k^2_z}{2m_e}.$$ 

Besides their role as a particle pump/sink, the leads play an important role as a heat sink. An important assumption we make is that any heat generated in the interacting region due to the application of a transverse electric field is efficiently dissipated into the leads so as to prevent build up of heat in the region. This is a well-justified assumption because the leads are assumed to be infinite and the interacting region has a thin profile.

In equilibrium ($\mu_{\text{res}} = \mu_R = \mu_L$), the central system is expected to reach chemical equilibrium with the reservoirs in the long-time limit so that $\mu_{\text{sys}} = \mu_{\text{res}}$. In the out-of-equilibrium case, the system is coupled to two reservoirs that are not in chemical equilibrium. Therefore, although the electron distribution in the interacting system reaches a static form in the long-time limit, it is in no way expected to have an equilibrium form due to constant influx (outflux) of particles from (into) the leads.

III. KELDYSH PATH INTEGRAL FORMULATION

In this section, we formulate a theory of nonequilibrium BCS superconductivity in graphene using the Keldysh functional integral formalism. The theory is first expressed in terms of a Keldysh partition function using coherent states of fields defined on the time-loop Keldysh contour, $\mathcal{C}$. Following a Hubbard-Stratonovic decoupling of the quartic interaction term in the pair channel a BCS theory for superconducting graphene is obtained by assuming a static, homogeneous gap, integrating out both leads and graphene electrons, and extremizing the effective action with respect to the gap. The resulting mean-field equations, which are a nonequilibrium generalization of the corresponding equilibrium equations, are analyzed in the remainder of the paper.

The starting Keldysh generating functional reads

$$Z^K = \int \mathcal{D}\{a, \bar{a}, b, \bar{b}, A, \bar{A}, B, \bar{B}\} e^{iS^K},$$

(12)

where

$$S^K = S^K_{\text{sys}} + S^K_{\text{res}} + S^K_{\text{sys-res}}.$$ 

(13)

If we introduce 4-component spinors defined in Nambu-sublattice space for both graphene electrons and leads electrons

$$\phi_{\mathbf{k},\mathbf{z},\sigma}(t) = \begin{pmatrix} A_{\mathbf{k},\mathbf{z},\sigma,\uparrow}(t) \\ -\bar{A}_{\mathbf{k},\mathbf{z},\sigma,\uparrow}(t) \\ B_{\mathbf{k},\mathbf{z},\sigma,\downarrow}(t) \\ -\bar{B}_{\mathbf{k},\mathbf{z},\sigma,\downarrow}(t) \end{pmatrix},$$

(15)

the actions in Eq.[13] become

$$S^K_{\text{sys}} = \int_{\mathcal{C}} dt \frac{1}{N_\Delta} \sum_{\mathbf{k}} \tilde{\phi}_{\mathbf{k}}(t) \times \left[i\partial_t - \lambda_{k} \tau_z^N - g_{k} \tau_z^N \tau_+^N - g_{k}^* \tau_z^N \tau_+^N \right] \phi_{\mathbf{k}}(t) + U \int dt \sum_{t} \left[ a_{\mathbf{i},1}(t) \bar{a}_{\mathbf{i},1}(t) a_{\mathbf{i},1}(t) a_{\mathbf{i},1}(t) + b_{\mathbf{i},1}(t) \bar{b}_{\mathbf{i},1}(t) b_{\mathbf{i},1}(t) b_{\mathbf{i},1}(t) \right],$$

(16)

$$S^K_{\text{res}} = \int_{\mathcal{C}} dt \int \frac{dz}{2\pi} \sum_{\mathbf{k}} \sum_{\mathbf{z}} z_{\mathbf{k}} \Delta_{\mathbf{k}} \phi_{\mathbf{k},\mathbf{z},\sigma}(t) \times \left(i\partial_t - \epsilon_{\mathbf{k},\mathbf{z}} \tau_z^N \right) \Phi_{\mathbf{k},\mathbf{z},\sigma}(t),$$

(17)

and

$$S^K_{\text{sys}-\text{res}} = \int_{\mathcal{C}} dt \int \frac{dz}{2\pi} \sum_{\mathbf{k}} \sum_{\alpha} \sum_{\mathbf{z}} \sum_{\mathbf{z}} \tilde{\Phi}_{\mathbf{k},\mathbf{z},\sigma,\alpha}(t) \times \left[ \Phi_{\mathbf{k},\mathbf{z},\sigma,\alpha}(t) \phi_{\mathbf{k}}(t) + \tilde{\phi}_{\mathbf{k}}(t) \Phi_{\mathbf{k},\mathbf{z},\sigma,\alpha}(t) \right].$$

(18)

$\tau_\pm^\nu$ are $2\times2$ matrices given by

$$\tau_\pm^\nu = \frac{1}{2} \left( \tau_x^\nu \pm i \tau_y^\nu \right),$$

(19)

where $\tau_x^\nu, \tau_y^\nu, \tau_z^\nu$ are Pauli matrices. Superscript $\nu$ indicates the space in which the matrices act: $A$ ($N$) denotes sublattice (Nambu) space. The quartic interaction term in Eq.[13] is decoupled using Hubbard-Stratonovic fields $\Delta^N(t)$ and $\Delta^P(t)$. In the BCS mean-field approximation, where this field is assumed static and homogeneous (i.e. $\Delta^N(t) = \Delta^P(t) \equiv \Delta$), the resulting action of the system reads

$$S^K_{\text{sys}} = \int_{\mathcal{C}} \frac{1}{N_\Delta} \sum_{\mathbf{k}} \tilde{\phi}_{\mathbf{k}}(t) \times \left[i\partial_t - \lambda_{k} \tau_z^N - g_{k} \tau_z^N \tau_+^N - g_{k}^* \tau_z^N \tau_+^N \right] + U \Delta \tau_+^N + U \Delta^* \tau_+^N \left[ \phi_{\mathbf{k}}(t) - 2U |\Delta|^2 \right].$$

(20)
The self-consistency condition for the gap is
\[ \Delta = (a_{i,\downarrow} a_{i,\uparrow}) (t) = (b_{i,\downarrow} b_{i,\uparrow}) (t). \]

The time-loop contour integral is carried out by first splitting every field into two components, labeled "+" and "−", which reside on the forward and the backward parts of the time contour, respectively. The continuous action then becomes
\[ S^K = \int_{-\infty}^{\infty} dt \left[ \mathcal{L}_+(t) - \mathcal{L}_-(t) \right], \]
where \( \mathcal{L}_\pm(t) \) is the Lagrangian corresponding to the action defined in Eq. (13) written in terms of \( + \) (−) fields. When time-ordered products of Heisenberg fields in the theory are constructed on the Keldysh contour we obtain four Green functions
\[
\begin{align*}
    iG^T(t, t') &= \langle \Upsilon_+(t) \Upsilon_+(t') \rangle \\
    iG^F(t, t') &= \langle \Upsilon_-(t) \Upsilon_-(t') \rangle \\
    iG^< (t, t') &= \langle \Upsilon_+(t) \Upsilon_-(t') \rangle \\
    iG^> (t, t') &= \langle \Upsilon_-(t) \Upsilon_+(t') \rangle.
\end{align*}
\]
Because these Green functions are not linearly independent, a linear transformation of the fields from the Kadanoff-Baym basis \((+,−)\) to the Keldysh basis \((cl,q)\) is commonly performed. For bosons, the barred fields are related to the unbarred fields simply by complex conjugation and thus the transformation is identical for both,
\[
\begin{pmatrix}
    \Upsilon_{cl} \\
    \Upsilon_q
\end{pmatrix} = \frac{1}{\sqrt{2}} \begin{pmatrix}
    1 & 1 \\
    1 & -1
\end{pmatrix}
\begin{pmatrix}
    \Upsilon_{+} \\
    \Upsilon_{−}
\end{pmatrix}.
\]

For fermions, unbarred fields transformed in the same manner as Eq. (23). For barred fields, we choose a different transformation,
\[
\begin{pmatrix}
    \tilde{\Upsilon}_1 \\
    \tilde{\Upsilon}_2
\end{pmatrix} = \frac{1}{\sqrt{2}} \begin{pmatrix}
    1 & -1 \\
    1 & 1
\end{pmatrix}
\begin{pmatrix}
    \Upsilon_{+} \\
    \Upsilon_{−}
\end{pmatrix}.
\]

In order to express the Keldysh action, Eq. (22) in the Keldysh basis it is now appropriate to define 8-component spinors for graphene electrons and leads electrons defined in the Nambu-sublattice-Keldysh space. Since we are interested in steady-state properties of the system, it is useful to first Fourier transform the fields into frequency space. We define the 8-component spinors as
\[
\psi_k \equiv \begin{pmatrix}
    a_{k,\downarrow}^1 \\
    a_{k,\downarrow}^2 \\
    b_{k,\downarrow}^1 \\
    b_{k,\downarrow}^2 \\
    a_{k,\uparrow}^1 \\
    a_{k,\uparrow}^2 \\
    b_{k,\uparrow}^1 \\
    b_{k,\uparrow}^2
\end{pmatrix}, \quad \Psi_{k,\alpha} \equiv \begin{pmatrix}
    A_{1,\alpha, k, \downarrow, \downarrow} \\
    A_{2,\alpha, k, \downarrow, \uparrow} \\
    B_{1,\alpha, k, \downarrow, \downarrow} \\
    B_{2,\alpha, k, \downarrow, \uparrow} \\
    A_{1,\alpha, k, \uparrow, \downarrow} \\
    A_{2,\alpha, k, \uparrow, \uparrow} \\
    B_{1,\alpha, k, \uparrow, \downarrow} \\
    B_{2,\alpha, k, \uparrow, \uparrow}
\end{pmatrix},
\]
where \( k \equiv (\mathbf{k}, \omega) \) is the energy-momentum 3-vector. The action (Eq. (22) then becomes
\[
S^K_{\text{sys}} = \int_k \tilde{\psi}_k \left\{ (g^{R}_0(k) \tau^N - g^{A}_0(-k) \tau^N) \tau^K \\
    (g^{A}_0(k) \tau^N - g^{R}_0(-k) \tau^N) \tau^K \\
    + U \left[ \Delta_q \tau^N + \Delta^* \tau^N + (\Delta_\alpha \tau^N + \Delta^* \tau^N) \tau^K \right] \right\} \psi_k \]
\[ - 2U \left[ \Delta_q \Delta_q + \Delta^* \Delta^* \right], \]

\[
S^K_{\text{sys-res}} = \int_k \int \frac{dk}{2\pi} \sum_\alpha \Psi_{k,\alpha} \left[ \frac{\alpha}{\sqrt{2}} \tilde{\psi}_k + \tilde{\psi}_k \Psi_{k,\alpha} \right].
\]
Here, \( f_k = \frac{1}{N} \sum \frac{d\omega}{2\pi} \), and \( \tau_{\uparrow,\downarrow} \) are 2×2 matrices defined by
\[
\tau_{\uparrow,\downarrow} = \begin{pmatrix}
    1 & 0 \\
    0 & 1
\end{pmatrix}.
\]

Superscript \( K \) on various \( \tau \) matrices indicate that they act in Keldysh space. \( g^{R,A,K}_0(k) \) denote inverse, advanced and Keldysh Green functions for non-interacting electrons in the graphene sheet while \( \tilde{g}^{R,A,K}_0(k) \) are the corresponding Green functions for lead \( \alpha \). For the graphene sheet, they are given by
\[
g^{R}_0(k) = \omega - \lambda_k + i\delta = g^{A*}_0(k) \]
\[ g^{K}_0(k) = 2i\delta K(\omega). \]

Here, \( K(\omega) \equiv 1 + 2n_F(\omega) \) where \( n_F(\omega) \) is the usual Fermi-Dirac distribution function. \( \delta \) is an infinitesimal regularization parameter. For the non-interacting case, \( g^{K}_0 \) merely serves as a regularization for the Keldysh functional integral. Because a finite self-energy term is anticipated from the coupling of graphene electrons to the leads, \( \tilde{g}^{R,A,K}_0 \), are those corresponding to free fermions.

A. Integrating out the leads

We now integrate out the leads degrees of freedom in order to obtain an effective theory only in terms of fields defined on the graphene sheet. The inverse retarded, advanced and Keldysh Green functions for the leads, \( \tilde{g}^{R,A,K}_0 \), are those corresponding to free fermions.
and because the leads are always in thermal and chemical equilibrium the Keldysh Green function is strictly related to the retarded and advanced Green functions via the fluctuation-dissipation theorem (FDT). They are given by

\[
\tilde{g}^{R}(k) = \omega - \epsilon_{k} + i \delta = g_{\alpha}^{A}(k) \tag{32}
\]

\[
\tilde{g}^{K}(k) = 2i \delta \tanh \left( \frac{\omega - \mu_{\alpha}}{2T} \right). \tag{33}
\]

Upon integrating over the leads, the resulting self-energy action becomes

\[
S_{\Sigma} = \int_{k} \bar{\psi}_{k} \left\{ -\Sigma^{R}(k) \tau_{z}^{N} \tau_{1}^{K} - \Sigma^{A}(k) \tau_{z}^{N} \tau_{1}^{K} \right. \]

\[
\left. -\Sigma^{K}(k) \tau_{1}^{N} \tau_{+}^{K} - \Sigma^{K}(k) \tau_{1}^{N} \tau_{-}^{K} \right\} \psi_{k}, \tag{34}
\]

where

\[
\Sigma^{R}(k) = \sum_{\alpha} \int \frac{dk_{z}}{2\pi} \frac{\epsilon_{\alpha}^{2}}{\omega - \epsilon_{k} + i \delta}
\]

\[
= -i \sum_{\alpha} \pi \rho \epsilon_{\alpha}^{2} = -i \Gamma \tag{35}
\]

\[
= \Sigma^{A}(k),
\]

and

\[
\Sigma^{K}(k) = -2\pi i \sum_{\alpha} \int \frac{dk_{z}}{2\pi} \frac{\epsilon_{\alpha}^{2}}{\omega - \mu_{\alpha}}
\]

\[
\times \delta(\omega - \epsilon_{k} - \epsilon_{k'}),
\]

\[
= -2i \sum_{\alpha} \Gamma_{\alpha} \tanh \left( \frac{\omega - \mu_{\alpha}}{2T} \right). \tag{36}
\]

Here, \( \Gamma_{\alpha} = \pi \rho \epsilon_{\alpha}^{2} \) measures the effective coupling strength between the layer and leads, and \( \Gamma = \Gamma_{L} + \Gamma_{R} \). \( \rho \) is the lead density of states to tunnel into the layer assumed to be constant. The frequency-independent damping coefficient, \( \Gamma \), and the vanishing real energy shift that result from our assumptions indicate that the bath is treated as an Ohmic environment. Combining the actions Eqs.\ref{37},\ref{39}, we obtain the dressed inverse Green functions for electrons in the graphene sheet

\[
g^{R}(k) = \omega - \lambda_{k} + i \Gamma = g^{A}(k), \tag{37}
\]

\[
g^{K}(k) = 2i \sum_{\alpha} \Gamma_{\alpha} \tanh \left( \frac{\omega - \mu_{\alpha}}{2T} \right). \tag{38}
\]

The negative imaginary part of \( \Sigma^{R}(k) \) leads to an irreversible damping in the time-dependent Green function \( G^{R}(k,t) \). The damping term formally describes decoherence suffered by a propagating electron wave due to incoherent escape and injection of electrons into and from the leads.

At this point, it is convenient to shift the energy scale so that all energies are measured with respect to \( \mu = (\mu_{L} + \mu_{R})/2 \). This is equivalent to the following mapping

\[
\omega \rightarrow \omega - \mu
\]

\[
\lambda_{k} \rightarrow \lambda_{k} - \mu
\]

\[
\mu_{\alpha} \rightarrow V_{\alpha}/2,
\]

where \( V_{L,R} = \pm V \) and \( V = \mu_{L} - \mu_{R} \). We assume \( V > 0 \).

Following this choice the inverse retarded Green function, Eq.\ref{37} remains invariant while Eq.\ref{38} becomes

\[
g^{K}(k) = 2i \sum_{\alpha} \Gamma_{\alpha} \tanh \left( \frac{\omega - V_{\alpha}/2}{2T} \right). \tag{39}
\]

Using the dressed inverse Green functions defined in Eqs.\ref{37},\ref{39}, the effective action for the graphene sheet is

\[
S_{\text{sys}}^{\text{eff}} = \int_{k} \bar{\psi}_{k} \tilde{g}^{-1}_{k} \psi_{k} - 2U \left[ \Delta_{cl}^{*} \Delta_{q} + \Delta_{q}^{*} \Delta_{cl} \right], \tag{40}
\]

where the inverse Green function matrix \( \tilde{g}^{-1}_{k} \) is now given by

\[
\tilde{g}^{-1}_{k} = \begin{pmatrix}
\begin{array}{cccc}
g^{R}(k) & \Delta_{q} & -g_{k}^{*} & 0 \\
\Delta_{q}^{*} & -g^{R}(-k) & 0 & g_{k}^{*} \\
-g_{k} & 0 & g^{R}(k) & \Delta_{q} \\
0 & \Delta_{cl} & 0 & -g^{R}(-k) \\
\Delta_{cl}^{*} & g^{K}(k) & 0 & 0 \\
0 & \Delta_{cl} & 0 & \Delta_{cl} \\
0 & 0 & \Delta_{cl} & g^{K}(k) \\
0 & 0 & \Delta_{cl}^{*} & g^{K}(K)
\end{array}
\end{pmatrix}
\right). \tag{41}
\]

\]

B. The Mean Field Equations

In closed equilibrium, solutions to the mean-field gap and number equations on the honeycomb lattice have shown that while graphene exhibits a BCS-BEC crossover behaviour away from the Dirac point for increasing attractive interaction strength, \( u \), superconductivity in graphene at half-filling requires a finite attrac-
tive interaction. In this section, we derive the main results of our work which are the mean-field gap and number equations in the presence of leads and voltage. Solving these equations will allow us to study the effects of dissipation and nonequilibrium current on the gap as a function of attractive interaction strength $u$ and filling $n$ and compare the results to the equilibrium calculations. We begin by obtaining an effective theory for the $s$-wave order parameter alone by integrating out the graphene electrons. From Eq. (40) we obtain

$$iS_{eff}^{K}(\Delta, \Delta^*) = Tr \ln[-i\gamma^{-1}] - 2iU(\Delta^*_q + \Delta_q + c.c.). \quad (42)$$

1. The gap equation

The self-consistent equation for the gap can be obtained from the following classical saddle-point equation

$$\frac{2\Delta}{U} = \int_k \frac{4\Delta \sum_{\alpha} \Gamma_{\alpha} \tanh \left( \frac{\Delta^*_k + \Delta_q + c.c.}{2} \right) ((\omega + E_k)^2 + \Gamma^2)(\omega - E_k)^2 + \Gamma^2} {[(\omega - E_k)^2 + \Gamma^2][\omega - E_+(k)]^2 + \Gamma^2}[\omega + E_+(k) + \Gamma^2][\omega + E_+(k)^2 + \Gamma^2]. \quad (45)$$

The spectra of the two bands are given by

$$E_{\pm}(k) = \sqrt{\xi_{\pm}^2 + \Delta^2}, \quad \xi_{\pm}(k) = \lambda_k \pm |g_k|, \quad (46)$$

and $$E_k = \sqrt{\lambda_k^2 + |g_k|^2 + \Delta^2}. \quad (47)$$

Scaling all energies by bandwidth $\tilde{t}$ and evaluating the $\omega$-integral we obtain

$$\frac{1}{u} = \frac{1}{2\pi N} \sum_k \left[ F_+(\xi_+(k)) + F_-(\xi_-(k)) \right], \quad (48)$$

where

$$F_v(x) \equiv \frac{1}{x} \left[ \tan^{-1}\left( \frac{v + x}{\gamma} \right) - \tan^{-1}\left( \frac{v - x}{\gamma} \right) \right],$$

and

$$\xi_{\pm}(k) = \frac{E_{\pm}(k)}{u}, \quad u = \frac{U}{t}, \quad \gamma_{\alpha} = \frac{\Gamma_{\alpha}}{t}, \quad v = \frac{V}{t}.$$ 

$\gamma = \gamma_L + \gamma_R$ denotes the sum of lead-graphene tunneling rates scaled by $t$. Eq. (47) is the BCS gap equation in the presence of leads ($\gamma$) and voltage ($v$) and is the nonequilibrium generalization of Eq. 2 in ref. (16). Indeed when one takes the limit as $\gamma \to 0$ and $v \to 0$ in Eq. (47) the equilibrium gap equation is recovered.

At low-energies, excitations in graphene at or near half-filling are concentrated near two inequivalent Fermi points at the corners of the hexagonal Brillouin zone. In the vicinity of these points, we have

$$\lambda_k \approx 3t' - \mu \equiv m \quad |g_{\pm k} + k| \approx v_F |k|, \quad (48)$$

where $v_F = 3t/2$ is the Fermi velocity and $\pm K = (\pm 4\pi/3\sqrt{3}, 0)$ are the locations of the inequivalent Fermi points. Within this approximation, the quasiparticle dispersions, $\xi_{\pm}(k)$, become

$$\xi_{\pm}(k) \approx m \pm \epsilon \quad \Xi_{\pm}(k) \approx \sqrt{\xi_{\pm}^2 + \Delta^2}, \quad (49)$$

where $\epsilon = v_F |k|$. Noting that the area per lattice site is $A/N = 3\sqrt{3}/4$ the conversion from $k$-summation to $\epsilon$-integral is given by

$$\frac{1}{N} \sum_k = \frac{3\sqrt{3}}{4\pi v_F} \int_0^{D} \epsilon d\epsilon, \quad (50)$$

The energy cut-off, set by conserving the total number of states in the Brillouin zone, is $D = \sqrt{3}\pi \approx 2.33$ in units of $t$. In the continuum limit, the gap equation then becomes

$$\frac{1}{u} = \frac{3\sqrt{3}}{8\pi^2 v_F} \int_0^{\epsilon_F} \epsilon d\epsilon \left[ F_+(\Xi_+(k)) + F_-(\Xi_-(k)) \right]. \quad (51)$$

2. The number density equation

In equilibrium, the number density is computed using a thermodynamic relation $\partial F_{MF}/\partial \mu = -N_c$. Out of equilibrium, the relation does not hold and the particle density, $n$, must be extracted from one of the four
Green functions, $G^<$ using Kadanoff-Baym Green functions, $\gamma$

\[ n = \frac{-i}{4} \sum \int_k G^<_{\sigma\Lambda}(k). \] (52)

$\sigma$ labels the electron spin and $\Lambda \in \{A,B\}$ labels the
sublattice in which it propagates. In terms of Keldysh Green functions,

\[ n = \frac{-i}{4} \sum \int_k [G^R_{\sigma\Lambda}(K) - G^R_{\sigma\Lambda}(K) + G^A_{\sigma\Lambda}(K)], \] (53)

where $G^{R,A,K}(k)$ are the retarded, advanced and Keldysh
Green functions for the graphene electrons. These Green functions can be obtained by inverting the matrix, $\gamma^{-1}(k)$, in Eqs.\[11\] We find that the form of the Green
functions is independent of spin and sublattice, and the resulting number equation reads

\[ F(\omega, v) = \sum_{\sigma} \gamma_{\sigma} sgn(\omega - v_\sigma) = \frac{\gamma_L}{\gamma} sgn \left( \omega - \frac{v}{2} \right) + \frac{\gamma_R}{\gamma} sgn \left( \omega + \frac{v}{2} \right). \] (55)

An exact evaluation of the $\omega$-integral in Eq.\[54\] is difficult. However, it can be done in the limit where the applied bias is assumed small compared to the bandwidth and the damping coefficient, i.e. $v \ll \min\{1, \gamma\}$. Computing the integral up to quadratic order in $v$ the number density yields

\[ n = \frac{3\sqrt{3}}{4\pi v_F} \int_0^{\beta} \frac{de}{\pi} \]

\[ c_0(10\gamma^2 + \Xi_+^2 + \Xi_-^2) + (\gamma^2 + \Xi_+^2)(\gamma^2 + \Xi_-^2)[2c_2 + c_4(2\gamma^2 + \Xi_+^2 + \Xi_-^2) + c_6(10\gamma^2 + 6\gamma^2\Xi_+^2 + \Xi_-^2 + 6\gamma^2\Xi_+^2 + \Xi_-^2)]
\]

\[ \frac{1}{2} (\gamma^2 + \Xi_+^2)(\gamma^2 + \Xi_-^2)(16\gamma^4 + \Xi_+^2 (8\gamma^2 + \Xi_+^2 - \Xi_-^2) + \Xi_-^2 (8\gamma^2 + \Xi_+^2 - \Xi_-^2)) \]

\[ - \frac{\pi}{2} \left\{ \tan^{-1}\left(\frac{\Xi_+^2}{\Xi_-^2} \right) \right\} \left[ c_1(\Xi_+^2 - \Xi_-^2 - 4\gamma^2) + c_3(\Xi_+^2 + \gamma^2\Xi_+^2 + 4\gamma^4 - \Xi_-^2(\Xi_+^2 - 3\gamma^2)) + c_5[\Xi_+^2 + 6\gamma^2\Xi_+^2 + 9\gamma^4\Xi_+^2 - \Xi_-^2(\gamma^4 - 6\gamma^2\Xi_+^2 + \Xi_-^2) - 4\gamma^6] \right] \right) \]

\[ + \gamma \ln \left( \frac{\gamma^2 + \Xi_+^2}{\gamma^2 + \Xi_-^2} \right) \left\{ 2c_1 + c_3(\Xi_+^2 + \Xi_-^2 + 2\gamma^2) + c_5(\Xi_+^2 \Xi_-^2 - \gamma^2\Xi_+^2 - \gamma^2\Xi_-^2 - 3\gamma^4) \right\} \]

\[ + (2x - 1) \left( \frac{2\gamma c_0}{\pi(\gamma^2 + \Xi_+^2)^2(\gamma^2 + \Xi_-^2)^2} v + \frac{\gamma c_1}{2\pi(\gamma^2 + \Xi_+^2)^2(\gamma^2 + \Xi_-^2)^2} v^2, \right) \]

where $x = \gamma_L/\gamma$ and $\Xi_\pm$ are given by Eq.\[49\] The coefficients $c_0, \ldots, c_6$ are dependent on $\Xi_\pm, \xi_\pm$ and $\gamma$, and are defined as

\[ c_6 = 1, \]

\[ c_5 = \xi_+ + \xi_-, \]

\[ c_4 = 3\gamma^2 - \frac{\Xi_+^2 + \Xi_-^2}{2}, \]

\[ c_3 = 2(\xi_+(\gamma^2 - \Xi_+^2) + \xi_-(\gamma^2 - \Xi_-^2)), \]

\[ c_2 = 3\gamma^4 + (\Xi_+^2 - \Xi_-^2)^2 + \gamma^2(\Xi_+^2 + \Xi_-^2) - \frac{\Xi_+^4}{2} - \frac{\Xi_-^4}{2}, \]

\[ c_1 = \xi_-(\gamma^4 + 2\gamma^2\Xi_+^2 + \Xi_+^4) + \xi_+(\gamma^4 + 2\gamma^2\Xi_-^2 + \Xi_-^4), \]

\[ c_0 = \frac{1}{2}(\Xi_+^2 + \gamma^2)(\Xi_+^2 + \gamma^2)(\Xi_-^2 + \Xi_+^2 + \gamma^2). \]

It can be easily verified that in the limit of $\gamma \to 0$ and
$v \to 0$, Eq.\[56\] reduces to the equilibrium number equation
(c.f. Eq.\[3\] in ref.\[16\]). The mean-field equations Eqs.\[50,55\]
are the central results of this work. These equations will be analysed in the remainder of the paper.

IV. RESULTS

Our main focus will be on obtaining and analyzing gap phase diagrams in the parameter space of interaction strength \( (u) \) and number density \( (n) \) for various leads-graphene couplings \( (\gamma_L, \gamma_R) \) and external biases \( (v) \). A previous work on closed equilibrium graphene\(^{43}\) revealed that, at half-filling, the superconducting instability of the semi-metallic phase requires a critical attractive interaction strength \( u_c \) and, thus, the gap vanishes up to \( u_c \). Away from half-filling, the metallic phase is immediately unstable to superconductivity for arbitrarily weak attractive interaction strength. As a result, the gap remains finite for any finite \( u \) and the system displays a typical BCS-BEC crossover behaviour. In this section we quantitatively discuss the effects of dissipation and nonequilibrium current on the gap phase diagram by numerically solving the generalized mean-field equations, Eqs. 51, 56. The following sections will show that a dramatic modification to the phase diagram is observed by the mere coupling of graphene to its environment, even in the absence of nonequilibrium current. We find that the effects of external biases in addition to dissipation does not substantially alter the qualitative features of the phase diagram from the case in which the system is subject to dissipation alone. However, as the following sections will discuss, the application of an external bias leads to shifts in the metallic region surrounding half-filling which result from voltage-induced changes in the graphene electron density. The results presented here are applicable to the case of small biases \( (v < \min\{1, \gamma\}) \); effects of large biases are not considered here.

A. Finite lead-layer coupling \( \gamma \neq 0 \), zero voltage

\( (v = 0) \)

First, we begin with the case in which the lead-graphene-lead heterostructure is in thermodynamic equilibrium. In particular, this is the situation where \( \mu_L = \mu_R = \mu_{res} \), and in the long-time limit \( \mu_{sys} = \mu_{res} \) is maintained. Here, electron tunneling processes between the central graphene system and the leads is providing a mechanism for decoherence for the particles in the system \( (\gamma \neq 0) \), but an external bias that explicitly breaks time-reversal symmetry of the heterostructure is absent \( (v = 0) \). Consider the case where the central graphene sheet is in a superconducting phase. Because of its coupling to the leads one can envisage a situation in which an electron that constitutes a Cooper pair escapes into the leads. Because the leads are assumed to be infinite the electron that has escaped the system is completely lost in the leads and as a consequence looses its coherence with its former partner. Although a different electron may enter the system from a lead within a time-scale of \( \tau_{inv} \sim 1/T \), the electron will not necessarily pair with the widowed electron since it completely lacks coherence to do so. Because dissipation effectively acts as a pair-breaking mechanism we expect a suppression of the gap throughout the entire region of the phase diagram. Fig.5 plots the gap phase diagrams for various leads-graphene coupling strengths \( (\gamma) \). Fig.5(a) correspond to the closed equilibrium case which has been obtained previously\(^{43} \). Fig.5(b), (c) display the behaviour of the gap as \( \gamma \) is increased. It is apparent from these plots that the suppressed region in the gap (dark blue region) grows as \( \gamma \) is strengthened. Regions of large gap values corresponding to the region with large \( u \) also displays an overall suppression in the gap as \( \gamma \) is increased. The qualitative features of the diagrams are consistent with the expectation described above. Let us now discuss the results more quantitatively.

1. Half-filling \( (n = 1) \)

For the closed equilibrium case at half-filling \( (\gamma = v = 0 \text{ and } n = 1) \) the semimetal-superconductor transition is possible mainly because the divergent nature of the integral on the right hand side of Eq. 51 is cured by particle-hole symmetry. When the integral is convergent, it is clear that a solution to the gap equation does not exist for small \( u \) where \( u^{-1} \) becomes larger than the integral. The value of the critical interaction parameter at which the transition occurs can be easily quantified. At half-filling the number equation, Eq. 55 is satisfied by \( m = 3t' - \mu = 0 \), and thus at the critical point \( (n = 1, \Delta = 0, \text{ and } m = 0) \) the gap equation reads

\[
\frac{1}{u_c} = \frac{3\sqrt{3}}{4\pi v_F} \int_0^D d\epsilon = \frac{1}{2.33}.
\]

For any \( u < u_c \) the equations cannot be solved with any real \( \Delta \) and the system enters the semimetallic phase. In the presence of dissipation \( (\gamma > 0) \) the number equation is still solved by \( m = 0 \) at half-filling, and the gap equation at the critical point yields

\[
\frac{1}{u_c} = \frac{3\sqrt{3}}{2\pi^2 v_F^2} \int_0^D d\epsilon \tan^{-1} \left( \frac{\epsilon}{\gamma} \right) = \frac{3\sqrt{3}D}{2\pi^2 v_F^2} \left[ \tan^{-1} \left( \frac{\gamma D}{2} \right) - \frac{\gamma D}{2} \ln \left( 1 + \frac{\gamma D^2}{2} \right) \right],
\]

where the reduced coupling strength is given by \( \gamma_D = \gamma/D \). The integral on the right hand side of Eq. 59 is convergent and, thus, tells us that the semimetal-superconductor transition exists in the presence of dissipation at half-filling. The behaviour of \( u_c \), as a function of \( \gamma_D \) is plotted in Fig. The value of \( u_c \) increases as \( \gamma_D \) is increased. This is consistent with the above considerations from which we expect that a larger interaction parameter is necessary to achieve pairing since leads-induced decoherence generally suppresses superconductivity. The phenomenon can also be observed in Fig. where the apex of the blue region shifts.
right for larger $\gamma$. The plots show that at $\gamma = 0$ $u_c$ converges to the closed equilibrium value of $u_c \sim 2.33$ as predicted by previous calculations.

What is notable in Fig. 5 is the expansion of the blue region, where the gap is small, as $\gamma$ is increased. The question is whether or not the typical BCS-BEC crossover behaviour observed in the closed equilibrium case is a correct physical picture away from half-filling for finite $\gamma$. The external baths acting as a pair-breaking mechanism makes the issue subtle. The pair-breaking perturbation in a superconductor with magnetic impurities has been shown\cite{24,25} to strongly suppress the transition temperature of the superconductor. Therefore, when such perturbation is strong enough the gap may vanish completely and give rise to a metal-superconductor quantum phase transition at finite doping. The question of whether or not the gap vanishes away from half-filling depends on the convergence of the integral in the gap equation. At $\nu = 0$, the generalized gap equation becomes

$$\frac{1}{u} \propto \int_0^D e^\nu \left[ \frac{1}{\Xi} \tan^{-1} \left( \frac{\Xi_+}{\gamma} \right) + (+ \rightarrow -) \right].$$ (61)

We see that for any $m$ (i.e. regardless of being at half-filling or not), the integral is convergent because for any small $\Xi_\pm$, which is the source of divergence, the arctan factor nullifies the divergence. This implies a finite $u_c$ at both half-filling and away from half-filling. Consequently, the system should undergo a superconductor-to-metal phase transition as the interaction parameter is lowered. Notice that the analysis above infers that the system will eventually enter the metallic phase as $u$ is decreased for any density.

In the closed equilibrium case away from half-filling, $m \neq 0$ and the critical point condition becomes

$$\frac{1}{u_c} = \frac{3\sqrt{3}}{4\pi v_F} \int_0^D e^\nu \left[ \frac{1}{|m+\epsilon|} + \frac{1}{|m-\epsilon|} \right] = \infty. \quad (60)$$

The divergence of the integral results in a solution with $\Delta > 0$ for any small $u > 0$. This gives $u_c = 0$ implying that Cooper instability occurs for any finite $u$ away from half-filling. Let us now investigate how this is modified when $\gamma$ is finite.
equation is solutionless and represents a (semi)metallic phase. Clearly, as \( \gamma \) is increased, the metallic region expands. We find that the superconducting (white) and metallic (black) regions are separated by a second-order phase transition.

The fact that the central graphene sheet becomes metallic away from half-filling certainly defies expectations based on the intrinsic properties of graphene. For closed graphene the semimetal-superconductor quantum criticality emerges at half-filling because of the absence of electron-electron screening that results from the vanishing density of states at the Dirac point. As stated earlier in this section, the physics of the metallic region can be understood using the phenomenon of leads-induced decoherence, and, thus, is not described by intrinsic properties of the central graphene sheet.

**B. Effect of voltage, \( v \neq 0 \)**

So far, we have discussed the effect of leads-induced dissipation on the gap phase diagram in the absence of voltage. We now consider the effects of driving an out-of-plane charge current through the superconducting graphene sheet. Here, we are limited to the regime of small voltages, specifically \( v \ll \min\{1, \gamma\} \). As mentioned before, we assume \( v \propto \mu_L - \mu_R > 0 \) and allow the relative strengths of the two couplings to the leads, \( \gamma_L \) and \( \gamma_R \), to vary. In the absence of current \( (v = 0) \), the gap equations depend only on the sum of these couplings \( \gamma = \gamma_L + \gamma_R \). But Eq.56 shows that in the presence of current \( (v \neq 0) \) the number density now depends on these couplings independently and depending on the relative strengths of these couplings the dominant correction term may change sign. The main qualitative modifications to the gap phase diagram in the presence of finite voltage reflects the influence of this correction term.

In the small voltage regime and for \( \gamma < 1 \), the dominant correction term gives a correction of order \( \gamma v \ll 1 \) to the number density, which is of order unity. Because the modifications to the gap phase diagram due to voltage is expected to be small the effect is more clearly seen by plotting the difference in gap values at finite and zero voltage. This is shown in Fig.4. The gap difference is plotted here for \( 2v = \gamma = 0.2 \) in the vicinity of the apex region. In Fig.4(a) \( \gamma_L/\gamma_R = 4 \) while in Fig.4(b) \( \gamma_L/\gamma_R = 0.25 \). The plots reveal that the metallic region (blue region in Fig.5) shifts vertically away from half-filling. The figure shows that for \( \gamma_L/\gamma_R = 4 \) the apex shifts up while for \( \gamma_L/\gamma_R = 0.25 \) it shifts down. Given \( \mu = (\mu_L + \mu_R)/2 \) and \( v > 0 \) the lowest order voltage correction in Eq.56 tells us that the number density increases or decreases depending on the asymmetry of the lead couplings. If \( \gamma_L > \gamma_R \), \( n \) increases, while if \( \gamma_L < \gamma_R \), \( n \) decreases. The gap equation yields the largest value of \( u_c \) given \( \gamma \) and \( v \) when \( m = 0 \). Thus, the above observation tells us that, for \( \gamma_L > \gamma_R \), \( m = 0 \) is achieved not at half-filling as in the equilibrium case but at \( n > 1 \). This shifts the apex upward. The opposite occurs for \( \gamma_L < \gamma_R \).

![FIG. 7](image-url) A cartoon plot showing the effect of voltage on the boundary of the metallic region. The dashed lines in both plots denote the boundary at \( v = 0 \). The shaded area is the metallic region after a steady-state bias is applied. In both plots, the applied voltage is \( v = 0.1 \). However, \( \gamma_L > \gamma_R \) in (a) while \( \gamma_L < \gamma_R \) in (b). Essentially, the voltage-induced modification is to shift the metallic region to higher values in density or to lower values depending on the polarity of the voltage and the lead-coupling asymmetry.

The nonequilibrium gap equation is convergent for all \( \mu \), thus, a metallic phase is once again expected at all densities.

**V. CONCLUSION**

In conclusion, we have theoretically studied the effects of dissipation and nonequilibrium drive on the properties of superconducting graphene. An external steady-state current was perpendicularly driven through the graphene sheet by attaching it to two leads which were equilibrated at two constant, but different, chemical potentials. The mean-field BCS theory of superconductivity on graphene was extended to the nonequilibrium situation by formulating the theory on the Keldysh contour. After obtaining nonequilibrium gap and number density equations.

![FIG. 8](image-url) A plot of \( u_c \) vs. \( v \) for a fixed \( \mu \). The plot line separates the metallic and superconducting phases of our system. Adjusting \( \mu \) will tune the location of \( u^*_c \) on the x-axis but the general shape of the curve is not modified.
we studied the BCS gap as a function of attractive interaction strength \( u \) and electron density \( n \) for various lead-graphene coupling strengths \( \gamma \) and voltages \( v \). We have shown that dissipation results in a suppression of the BCS gap at both zero and finite voltages. We argued that the coupling of the graphene sheet to external baths acts as a pair-breaking mechanism because it causes an electron that constitutes a Cooper pair to escape into the leads. Once an electron leaves the scattering region, it looses coherence with its time-reversed partner and the destruction of the Cooper pair entails.

A quantitative understanding of why the gap is significantly suppressed by dissipation can be gained by observing how dissipation affects the gap equation. Recall that the BCS gap equation for an ordinary superconductor in closed equilibrium is given by

\[
\Delta = uTN(0)\sum_n \frac{\Delta}{\sqrt{\omega_n^2 + \Delta^2}} \tag{62}
\]

\( N(0) \) is the density of states at the Fermi energy, and \( u > 0 \) is the attractive interaction strength. A general result for these ordinary superconductors is that the gap equation (Eq.62), and hence the gap, is unaffected by time-reversal-invariant perturbations. Take, for example, the influence of non-magnetic impurities on the superconducting state. The gap equation obtained after invoking disorder-averaging and the Born approximation reads

\[
\Delta = uT\bar{N}(0)\frac{\bar{\Delta}}{\sqrt{\omega_n^2 + \bar{\Delta}^2}} \tag{63}
\]

where \( \bar{\omega} \) and \( \bar{\Delta} \) are frequency and order parameter renormalized by the perturbation and \( \bar{N}(0) \) is the density of states in the presence of the perturbation. The essential point is that \( \bar{\omega} \) and \( \bar{\Delta} \) are related to their unrenormalized counterparts by a common factor \( \eta = \eta(\omega_n, \Delta) \), i.e.,

\[
\bar{\omega} = \eta \omega, \quad \bar{\Delta} = \eta \Delta.
\]

Because this factor \( \eta \) cancels out in Eq.63 the gap equation remains invariant and leads to the result that the gap is unaffected by non-magnetic impurities.

Imagine now that a pure ordinary superconductor is coupled to an external bath in equilibrium. The Nambu-Gorkov equations can be straightforwardly derived for this case,

\[
(i\omega_n + isgn(\omega_n)\Gamma - \xi_k) G + \Delta F = 1 \tag{64}
\]

\[
(i\omega_n + isgn(\omega_n)\Gamma + \xi_k) F + \Delta G = 0. \tag{65}
\]

where the ordinary and anomalous Green functions are given by

\[
G(k, \omega_n) = -\int_0^\beta d\tau \left( T_c c_{k,\uparrow}(\tau)e^\dagger_{k,\uparrow}(0) \right) e^{i\omega_n\tau}
\]

\[
F(k, \omega_n) = -\int_0^\beta d\tau \left( T_c c_{k,\uparrow}(\tau)c_{-k,\downarrow}(0) \right) e^{i\omega_n\tau}.
\]

We immediately see from Eqs.64,65 that \( \omega \) and \( \Delta \) scale asymmetrically, namely,

\[
\bar{\omega} = \eta \omega, \quad \bar{\Delta} = \Delta, \quad \eta = 1 + \frac{\Gamma}{|\omega_n|}. \tag{66}
\]

Here, \( \Gamma \) is the rate at which electrons decay into the bath. The asymmetry in the renormalization of \( \omega \) and \( \Delta \) (Eq.66) greatly affects the gap equation, Eq.63 and shows how dissipation can affect the gap significantly. This is consistent with the qualitative argument given above.

The emergence of the metal-superconductor quantum phase transition in the graphene subsystem at both zero and finite voltages gives rise to the possibility of inducing the phase transition using external bias. While fixing the average chemical potential \( \mu \) to some value, \( v \) can be changed by adjusting the interaction strength, \( u \), of the system is at \( u = u^* \). Suppose \( v < v(u^*) \) the system will be metallic. However, when \( v \) is increased and passes \( v = v(u^*) \), the system will become superconducting. \( u^* \) can be tuned by adjusting the average chemical potential \( \mu \). It is clear from Eq.66 that when the average chemical potential \( \mu \) is fixed, the electron density can change as a function of voltage.

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