Numerical renormalization group calculation of near-gap peaks in spectral functions of the Anderson model with superconducting leads

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Received 8 March 2008, in final form 30 April 2008
Published 4 June 2008
Online at stacks.iop.org/JPhysCM/20/275213

Abstract

We use the numerical renormalization group method (NRG) to investigate a single-impurity Anderson model with a coupling of the impurity to a superconducting host. Analysis of the energy flow shows that, contrary to previous belief, NRG iterations can be performed up to a large number of sites, corresponding to energy differences far below the superconducting gap \( \Delta \). This allows us to calculate the impurity spectral function \( A(\omega) \) very accurately for frequencies \( |\omega| \sim \Delta \), and to resolve, in a certain parameter regime, sharp peaks in \( A(\omega) \) close to the gap edge.

(Some figures in this article are in colour only in the electronic version)

1. Introduction

The vast progress in nanofabrication during the last decades made it possible to study basic physical effects in a very controlled manner. One example of such highly controllable devices are quantum dots [1], which are used, amongst various applications, for a detailed and very controlled study of the Kondo effect [2–5], which is one of the prime examples of many-body phenomena. Below a critical temperature (Kondo temperature \( T_K \)) a local moment, provided by the spin of an electron occupying the quantum dot, gets screened by reservoir electrons within an energy window \( T_K \) around the Fermi energy.

These seminal experimental works on the Kondo effect, together with the possibility to engineer reservoir properties, raise the following intriguing question: What interesting effects may arise if the local moment in the quantum dot is coupled to superconducting leads while parameters like the Kondo temperature or the superconducting gap can be adjusted arbitrarily? In a superconductor as described by Bardeen, Cooper and Schrieffer (BCS) [6], electrons with opposite spin and momentum form Cooper pairs, thereby expelling the lead density of states around the Fermi energy. An energy gap occurs. Obviously, when combining a Kondo quantum dot with superconducting reservoirs both effects compete: screening of the local moment by electrons around the Fermi energy against pair formation of the latter.

This competition has attracted a lot of interest and various methods were used to analyze the properties of the system in the above-mentioned limits, see for example the references in [7] or [8–11]. In the early 1990s Satori et al [12] proposed to apply the numerical renormalization group method (NRG) [13, 14] to the problem. Since then NRG was used to calculate the ground state and subgap bound state properties like position and degeneracy [12, 15, 16] as well as their spectral weight [17, 18]. Also dynamic quantities like the spectral function [18] or the Josephson current [19, 20] can be calculated with NRG, as was done recently.

The first main goal of the present paper is to gain insight into the way NRG works when applied to a system of one
local level coupled to a superconducting lead. NRG is a well established method for solving strongly correlated impurity problems. Actually, it was invented to solve the Kondo problem [13] and since then has been generalized to various schemes involving localized states coupled to fermionic [21] or bosonic [22, 23] baths. For a review, see [14]. The key idea of NRG is to discretize the conduction band logarithmically. This leads to a chain Hamiltonian with exponentially decreasing couplings, the so-called Wilson chain. It can be solved iteratively by enlarging the system site by site: due to the decreasing couplings every new site can be treated as a perturbation of the old system, thus increasing the resolution with every step.

It is, however, not at all obvious whether NRG still works for superconducting leads at energy resolutions well below the gap. This is because the pairing energy $\Delta$ is the same at all energy scales, thus remains a constant on-site contribution also in the above-mentioned chain structure. At iterations where the coupling of the Wilson chain reaches $\Delta$, it is not obvious whether added sites still can be understood as a perturbation in the iterative process or not.

In order to address this problem, we analyze in detail the flow of the eigenenergies during the NRG procedure. We show that NRG is indeed capable to resolve the continuum close to superconducting leads. They report dramatic enhancement to an enhanced linear conductance, as observed in a recent experiment [24] with carbon nanotube quantum dots coupled to superconducting leads. They report dramatic enhancement of the linear conductance (only) in the regime $\Delta \ll T_k$.

We find similar behavior of the spectral function in the non-interacting case where an analytical solution exists. We analyze this solution and compare our NRG results against it. We find excellent agreement especially at energies close to the gap, as expected from our study of the energy spectrum.

The paper is organized as follows. In section 2 we introduce the superconducting-lead Anderson model (SC-AM) and the NRG method. In section 3 the flow of the energy spectrum of the SC-AM is analyzed. NRG is shown to work at resolutions well below the energy scale of the superconducting gap. Section 4 discusses the calculation of the spectral function. In the last part we conclude and summarize our results.

2. The model and the method

In this section we introduce our model for the quantum dot coupled to a left and right superconducting reservoir, as well as the method we use, the numerical renormalization group method [13, 21]. We perform a Bogoliubov as well as a particle–hole transformation. We derive the formulas in a general form. For convenience, the discussions in the later sections will be restricted to a real order parameter of the superconductors, equivalent to only one lead. In the non-interacting limit the system can be understood in terms of a simple single-particle picture. The latter can be solved exactly and will serve as a guideline for gaining a deeper understanding of the problem.

2.1. Superconducting-lead Anderson model (SC-AM)

To describe a quantum state coupled to two superconducting reservoirs, we consider the standard Anderson model (AM) for a local level coupled to two metallic, non-interacting reservoirs [25], and add a BCS-type term $H_\Delta$, describing pair formation in the leads. This SC-Anderson model, to be called SC-AM, is then described by

$$H = H_{\text{dot}} + H_{\text{hyb}} + H_{\text{lead}} + H_\Delta,$$

with

$$H_{\text{dot}} = \sum_{\sigma} \varepsilon_d n_{\sigma d} + U n_{\sigma d} n_{\sigma \downarrow},$$

$$H_{\text{hyb}} = \sum_{l=L,R} \sum_{k,\sigma} V_l (c_{l\sigma}^\dagger d_{k\sigma} + d_{k\sigma}^\dagger c_{l\sigma}),$$

$$H_{\text{lead}} = \sum_{l=L,R} \sum_{k,\sigma} \Delta_k (c_{l\sigma}^\dagger c_{l-k\sigma} + c_{l-k\sigma}^\dagger c_{l\sigma}).$$

Electrons on the dot with spin $\sigma = \{\uparrow, \downarrow\}$ are created by $d_{k\sigma}$ and interact via the Coulomb repulsion $U$ with each other, $n_{\sigma d} = d_{\sigma d}^\dagger d_{\sigma d}$ being the charge operator for spin $\sigma$. In the hybridization term $H_{\text{hyb}}$, the coupling strength $V_l$ between dot and lead states is assumed to be real and independent of the wavevector $k$. $c_{l\sigma}$ creates an electron in lead $l = L, R$, respectively. $H_{\text{lead}}$ describes the conduction band of metallic leads. We assume an isotropic and linearized dispersion. The density of states is then constant, $\rho_l = 1/2D$, where the band ranges from $-D$ to $D$. In the following $D = 1$ will serve as energy unit. The pairing of electrons with opposite spin and momentum (Cooper pairs) is described by $H_\Delta$. $\Delta_k$ is the magnitude (later on often called the gap), $\phi_l$ the phase of the order parameter of the superconductor. For simplicity, we assume left-right symmetry, i.e. $\Delta_l = \Delta, \phi_l = \pm \phi/2$ and $V_l = V$, where $l = L, R$, respectively. Then the unitary rotation

$$
\begin{pmatrix}
  c_{l\sigma} \\
  c_{l\sigma}^\dagger
\end{pmatrix} = \frac{1}{\sqrt{2}} \begin{pmatrix}
  e^{-i\phi/4} & e^{i\phi/4} \\
  -i e^{-i\phi/4} & i e^{i\phi/4}
\end{pmatrix}
\begin{pmatrix}
  c_{l\sigma} \\
  c_{l\sigma}^\dagger
\end{pmatrix}
$$

of the reservoir operators yields a real Hamiltonian.

It is well known [26, 27] that the Hamiltonian of a bulk superconductor (equations (4) and (5)) can be diagonalized by a Bogoliubov transformation. Note, though, that the SC-AM

$$H_{\text{SC-AM}} = \sum_{k,\sigma} \epsilon_k (c_{l\sigma}^\dagger c_{l\sigma} + c_{l\sigma}^\dagger c_{l\sigma}),$$

in which $\epsilon_k$ is the bulk band structure of the superconductor, $c_{l\sigma}$ creates a pair in lead $l$ with spin $\sigma$. $H_{\text{SC-AM}}$ is the SC-Anderson Hamiltonian for a local level coupled to two metallic superconducting reservoirs at zero temperature. As in the Anderson model (with normal states)

$$H = \sum_{k,\sigma} \epsilon_k (c_{l\sigma}^\dagger c_{l\sigma} + c_{l\sigma}^\dagger c_{l\sigma})$$

and lead states is assumed to be real and independent of the wavevector $k$. $\epsilon_k$ is the magnitude of the order parameter of the superconductor. For simplicity, we assume left-right symmetry, i.e. $\Delta_l = \Delta, \phi_l = \pm \phi/2$ and $V_l = V$, where $l = L, R$, respectively. Then the unitary rotation

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  -i e^{-i\phi/4} & i e^{i\phi/4}
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  c_{l\sigma}^\dagger
\end{pmatrix}
$$

of the reservoir operators yields a real Hamiltonian.

It is well known [26, 27] that the Hamiltonian of a bulk superconductor (equations (4) and (5)) can be diagonalized by a Bogoliubov transformation. Note, though, that the SC-AM
cannot be understood as an AM with a superconducting-lead density of states \( \rho_\Delta = |e_k|/\sqrt{\varepsilon_k^2 + \Delta^2} \). This is because the Bogoliubov transformation explicitly depends on \( k \). The hybridization term \( \frac{1}{4} \epsilon_{\omega k} d_\sigma + \frac{1}{4} d_\sigma \epsilon_{\omega k} \) would transform to a complicated object that cannot be simplified by rotating the \( d \) operators of the local dot space.

2.2. The numerical renormalization group method (NRG)

In the 1970s, Wilson came up with a scheme for solving the Kondo problem nonperturbatively: the numerical renormalization group (NRG) \([13]\). Since then it was generalized to various schemes, describing localized electronic states coupled to fermionic \([21]\) or bosonic \([22, 23]\) baths. The NRG allows thermodynamic and dynamic properties of such strongly correlated systems to be calculated at zero as well as at finite temperature. We first discuss the method for the AM (\( \Delta = 0 \)), then for \( \Delta \neq 0 \). For brevity we apply all NRG transformations to the full SC-AM already in the discussion of the AM.

The key idea of NRG is to discretize the conduction band of the reservoir logarithmically. The Hamiltonian can then be transformed to a chain Hamiltonian. In this representation, equations (3)–(5) are mapped onto

\[
H_{\text{hyb}} = \frac{2\pi}{\nu} \sum_{\sigma} \left[ \cos \frac{\phi}{4} f_{\text{e}l} d_\sigma + \text{h.c.} \right] - \sin \frac{\phi}{4} \left( f_{\text{e}l} d_\sigma + \text{h.c.} \right)
\]

\[
H_{\text{lead}} = \frac{1}{\sqrt{\Lambda}} \left( 1 + \frac{1}{\Lambda} \right) \sum_{l=0,\omega,\sigma} \sum_{n=0}^{\infty} \Lambda^{-n/2} \xi_n \left( f_{\text{lead}} \pi f_{\text{lead}} + \text{h.c.} \right)
\]

\[
H_\Delta = -\Lambda \sum_n \left[ \left( f_{\text{e}n+1} f_{\text{e}n+1} + \text{h.c.} \right) - \left( f_{\text{e}n+1} f_{\text{e}n+1} + \text{h.c.} \right) \right].
\]

Electrons on site \( n \) in lead \( l = e, o \) are created by \( f_{\text{e}n} \). The dot level only couples to the zeroth site of the so-called Wilson chain \( H_{\text{lead}} \), where the hybridization is given by \( \Gamma = p_0 2 \nu \) (the factor 2 stems from the two leads). \( \Lambda > 1 \) is the discretization parameter of the conduction band. \( \xi_n = (1 - \Lambda^{-n-1}) (1 - \Lambda^{-2n-1})^{-1/2} (1 - \Lambda^{-2n-3})^{-1/2} \approx 1 \) for large \( n \). The hopping matrix elements between successive sites of \( H_{\text{lead}} \) fall off exponentially with \( \Lambda^{-n/2} \). The resulting energy scale separation ensures that the AM can be solved iteratively. The recursion relation reads

\[
H_0 = 1/\sqrt{\Lambda} \left[ H_{\text{dot}} + H_{\text{hyb}} - \Delta \sum_{l=0,\omega,\sigma} \sum_{\nu} S_l \left( f_{\text{e}l} f_{\text{e}l+1} + \text{h.c.} \right) \right]
\]

\[
H_{N+1} = \sqrt{\Lambda} H_N
\]

\[
+ \frac{1}{2} \left( 1 + \Lambda^{-1} \right) \sum_{l=0,\omega,\sigma} \sum_{\nu} \xi_n \left[ f_{\text{lead}} f_{\text{lead}} + \text{h.c.} \right] - \Delta \Lambda^{N/2} \sum_{l=0,\omega,\sigma} \sum_{\nu} S_l \left[ f_{\text{lead}} f_{\text{lead}} + \text{h.c.} \right].
\]

Here \( S_l = \pm 1 \) for \( l = e, o \), respectively. The initial Hamiltonian of the system is related to the NRG Hamiltonian by \( H = \lim_{N \to \infty} \Lambda^{-(N-1)/2} H_N \). This relation is exact in the limit \( \Lambda \to 1 \) and \( N \to \infty \).

Sites are added successively and at each step the enlarged system is diagonalized. Each added site then acts as a perturbation of order \( \Lambda^{-1/2} \) on the previous part of the chain. Consequently, the typical energy resolution \( \delta_{\text{AM}} \) of the AM at iteration \( n \) is given by \( \delta_{\text{AM}} \propto \Lambda^{-n/2} \). Thus, by choosing the length \( N \) of the chain large enough (so that \( \Lambda^{-N/2} \) is much smaller than all other energies in the problem), all relevant energy scales can be resolved and treated properly. When adding a site to the system, the dimension of the Hilbert space gets multiplied by the dimension \( d \) of the state space of that site, yielding \( d = 4 \) for a single fermionic lead (empty, singly occupied (either up or down), doubly occupied). Therefore the dimension of the Hilbert space increases exponentially with the length of the chain. Wilson proposed a truncation scheme according to which only the lowest \( N_{\text{kept}} \) eigenstates are kept at each iteration, thereby ensuring that the dimension of the truncated Hilbert space stays manageable. Recently, it was shown that by keeping track of the discarded states a complete, but approximate, basis of states can be constructed \([28, 29]\). This can be used to calculate dynamic properties like the spectral function \( A(\omega) \) (see equation (20) below) which rigorously satisfy relevant sum rules \([30, 31]\), like \( \int d\omega A(\omega) = 1 \).

Applying the NRG mapping also to the pairing term of the SC-AM (as already done above), an on-site contribution appears, see equation (8), constant in magnitude for each site. In the limit \( \Lambda^{-n/2} \gg \Delta \), this additional term hardly affects the properties of the system. But, when \( \Lambda^{-n/2} \sim \Delta \), it is not obvious whether the added sites still act as a perturbation in the iterative process (9) or not, that is whether the energy scale separation still works or not. In section 3 we will show that the separation of energy scales does work also at resolutions much smaller than the gap.

2.3. Bogoliubov and particle–hole transformations

Satori \textit{et al} \([12]\) have shown that a computationally more convenient representation of the Hamiltonian can be obtained by performing a Bogoliubov–Valatin transformation \([b_{\text{in}, \sigma} = 1/\sqrt{2} \phi f_{\text{in}, \sigma} + f_{\text{in}, \sigma}, b_{\text{in}, \sigma}^+ = b_{\text{in}, \sigma}^+ \phi \sigma \] as well as a particle–hole transformation \([\tilde{c}_{\text{in}, \sigma} = b_{\text{in}, \sigma}^+ - \sigma \tilde{c}_{\text{in}, \sigma}, \tilde{c}_{\text{in}, \sigma}^+ = b_{\text{in}, \sigma}^+ \sigma \], \( n = -1 \) represents the dot, thus \( d_\sigma = \tilde{c}_{-1, \sigma} \). Applying these transformations to (9), the Hamiltonian reads

\[
H_\text{dot} = \frac{U}{2} (1 - \tilde{n}_d + 2 \tilde{n}_d \tilde{n}_d) - \left( \tilde{\epsilon}_d + \frac{U}{4} \right) (\tilde{d}_d^+ \tilde{d}_d + \tilde{d}_d \tilde{d}_d)
\]

\[
H_{\text{hyb}} = \frac{2\pi}{\nu} \sum_{\sigma} \left[ \cos \frac{\phi}{4} (\tilde{c}_{\text{in}, \sigma}^+ d_\sigma + \text{h.c.} \right] - \sin \frac{\phi}{4} (\tilde{c}_{\text{in}, \sigma}^+ d_\sigma + \text{h.c.})
\]

\[
H_{\text{lead}} = \frac{1}{\sqrt{\Lambda}} \left( 1 + \frac{1}{\Lambda} \right) \sum_{\nu} \sum_{n=0}^{\infty} \Lambda^{-n/2} \phi \left( \tilde{c}_{\text{lead}, \sigma}^+ \tilde{c}_{\text{lead}, \sigma} + \text{h.c.} \right)
\]
of accurately treating superconducting leads. In section 3 the
energy spectrum, thus confirming that NRG is capable
of the Hilbert space. One can use the resulting exact solution as
large number of iterations exactly—that is without truncating
understanding of reasons why NRG does work for the SC-AM.

resulting single-particle picture will serve as a tool to gain a
deeper understanding of reasons why NRG does work for the
SC-AM.

2.4. Single-particle picture

Some properties of the system show up already in the non-
interacting case, \( U = 0 \). The Hamiltonian is then of quadratic
form and we only have to solve a single-particle problem. The
NRG Hamiltonian ((10)-(13)) can be diagonalized up to a
large number of iterations exactly—that is without truncating
the Hilbert space. One can use the resulting exact solution as
benchmark for the NRG result. We obtain very good agreement
in the energy spectrum, thus confirming that NRG is capable
of accurately treating superconducting leads. In section 3 the
single-particle picture will also serve as a tool to gain a deeper
understanding of reasons why NRG does work for the SC-AM.

Without lack of generality we restrict the discussion to
the symmetric case, \( \varepsilon_d = 0 \). Then the NRG Hamiltonian
only contains quadratic terms of the form \( a_i^\dagger a_i \), with \( a_i \)
some fermionic operator. For every iteration \( N \) the single-
particle Hamiltonian can be diagonalized by some unitary
transformation \( T \) to \( H_N = \sum_{j=1}^N \varepsilon_j a_j^\dagger a_j \). Here \( a_j = T_j^\dagger a_i \)
and the eigenstates \( |n_j\rangle = a_j^\dagger |\text{vac}\rangle \) satisfy \( a_j^\dagger a_j |n_j\rangle = n_j |n_j\rangle \).
The many-body eigenstates and the energy spectrum follow from
the Schrödinger equation

\[
H |m\rangle = E_m |m\rangle, \quad E_m = \left( \sum_{n=0}^{N_{\text{e}}} \varepsilon_n \right) - E_0, \tag{14}
\]

with the many-body eigenstates \( |m\rangle = |n_1 \ldots n_N\rangle \) and
eigenenergies \( E_m \) which are calculated w.r.t. the ground state
energy \( E_0 \). In the ground state \( |0\rangle \) all single-particle levels
with energy below the Fermi energy \( \varepsilon_F = 0 \) are occupied, thus \( E_0 = \sum_{\varepsilon_n < \varepsilon_F} \varepsilon_n \).
Expectation values of local operators are evaluated easily,
\( \langle 0| a_{j\downarrow}^\dagger a_{j\uparrow} |0\rangle = \sum_{\varepsilon_n < 0} U_{1\downarrow}^\dagger |0\rangle \langle 0| a_{j\uparrow} a_{j\downarrow}^\dagger |0\rangle U_{1\downarrow} = \sum_{\varepsilon_n < 0} |U_{1\downarrow}|^2. \)

The construction of the many-body spectrum from the
single-particle energy levels using (14) is illustrated in figure 1
for the SC-AM. Figure 1(a) shows a sketch of a typical single-
particle spectrum. The single-particle level spectrum consists of
a continuum above and below the gap, \( \varepsilon_d > \Delta \), and
the corresponding energies \( E_m \) are part of the continuum
energies. A concrete example of the many-body as

\[
\tilde{H}_\Delta = -\sum_{n=0,\sigma}^{\infty} (-1)^n \Delta (\tilde{n}_{\sigma n} - \tilde{n}_{\sigma n}). \tag{13}
\]

Operators in the new basis will always be denoted by a
tilde, e.g. \( \tilde{n}_{d\sigma} = d_j^\dagger d_\sigma \) or \( \tilde{n}_{\text{in}} = \tilde{c}_\text{in}^\dagger \tilde{c}_\text{in} \). The \( \tilde{Q} \)
nonconserving property of \( H_\Delta \) has been transferred to \( \tilde{H}_\Delta \).
This has two useful effects: (i) for the symmetric model not
only the \( z \)-component of the total spin (per iteration), \( \tilde{S}_z = \sum_{n=1}^{N} (\tilde{n}_{\sigma n} = \tilde{n}_{\sigma n}) \), but also the particle number \( \tilde{Q}_N = \sum_{n=1}^{N} (\tilde{n}_{\sigma n} - 1) \) is a conserved quantum number (note that
this definition yields \( \tilde{Q} = 0 \) for the Fermi sea together with
a singly occupied dot at \( \Gamma = 0 \)). Therefore the dimensions of
the matrices to be diagonalized at each iteration (and therefore
the numerical effort) is reduced significantly. (ii) Additionally,
in the non-interacting symmetric case \( (U, \varepsilon_d = 0) \) the
Hamiltonian takes a very simple quadratic form. We will focus
on its exact solution in the next section. In section 3 the
resulting single-particle picture will serve as a tool to gain a
deeper understanding of reasons why NRG does work for the
SC-AM.

For simplicity, we use \( \phi = 0 \) in the following. Then, the
odd channel decouples and the problem reduces to an effective
one-channel problem. The resulting model is equivalent to that
describing an impurity embedded in a bulk superconductor.

Figure 1. Single-particle and many-body eigenenergies for the SC-AM \((U = \varepsilon_d = 0)\). (a) Schematic sketch of a typical single-particle level spectrum of the SC-AM. The continuum is represented by a discrete set of closely-spaced levels. The construction of the lowest lying many-body eigenstates in terms of single-particle states according to (14) is illustrated. The corresponding energies \( E_m \) (w.r.t. the ground state energy) and quantum numbers of the many-body states are also given. Two dots stand for a doubly occupied level. (b) Single-particle energies \( \varepsilon_j (\varepsilon_j^+ > 0; \text{stars}, \varepsilon_j^- < 0; \text{crosses}) \), as well as the corresponding many-body eigenenergies \( E_m \) (lines) for \( \Delta = 10^{-4} \) and \( \Gamma/\Delta = 0.3 \) at a late
NRG iteration \((\delta_\ell \ll \Delta)\). The many-body eigenenergies depicted in (a) are specified on the right-hand side. The single-particle continuum energies go like \( \varepsilon_j^+ - \Delta \propto \lambda^{2j} \), see section 3.2. Due to many-particle excitations, this dense profile repeats as substructures in the many-body spectrum at \( m\Delta + \epsilon_r \ell_0 \) \((m = 1, 2, \ldots, \ell = 0, 1, 2)\).
where $n$ fall off exponentially with $(f))$ Cooper pair formation dominates the low-energy properties and the ground state is a doublet. Favorable to break Cooper pairs and lower the energy by $\Delta_1$, electrons and the Kondo singlet (ground state) forms. ((b), (c), (e), (f)): $\Delta_1 > \Delta_1$. In this section we analyze the many-body energy spectrum generated during the iterative NRG procedure. As already mentioned in the last section, the spectrum consists of a continuum above and subgap bound states below the gap $\Delta$. The competition between the Kondo effect and Cooper pair formation is reflected in the ground state properties of the system. A detailed analysis of the structure of the continuum with the help of the single-particle picture reveals that, interestingly, energy scale separation is even more efficient at energy scales smaller than the gap (compared to the AM). The energy resolution level spacing of the Wilson chain drops exponentially with every added site (see discussion above). The energy resolution $\delta_{10}$, respectively, are indicated by dashed or solid arrows. (a), (d) $\Delta = 0$, AM. Since $\delta_{\text{AM}} \propto 1/n^2$, the eigenenergies (a) fall off exponentially with $n$ and the energy flow diagram (d) converges. At energy scale $T_K$ the localized spin gets screened by the conduction electrons and the Kondo singlet (ground state) forms. For comparison, the result for the AM is indicated (brown (grey), too). For $\Delta > T_K$ it is energetically favorable to break Cooper pairs and lower the energy by $T_K$ by screening the local spin, thus the ground state is a singlet. For $\Delta > T_K$ Cooper pair formation dominates the low-energy properties and the ground state is a doublet.

![Figure 2](image)

**Figure 2.** Energy spectra ((a)–(c)) and the corresponding energy flow diagrams ((d)–(f)) for the lowest 280 eigenenergies of the even NRG iterations of a SC-AM. $U = 0.2$, $\Gamma = 0.01$, $T_K = 1.25 \times 10^{-5}$, $\Delta = 2.5$ and $N_{\text{kept}} = 512$ in all plots. The iteration numbers $N_T$ or $N_{\text{AM}}$, where $\delta_{\text{AM}} \approx T_K$ or $\Delta$, respectively, are indicated by dashed or solid arrows. (a), (d) $\Delta = 0$, AM. Since $\delta_{\text{AM}} \propto 1/n^2$, the eigenenergies (a) fall off exponentially with $n$ and the energy flow diagram (d) converges. At energy scale $T_K$ the localized spin gets screened by the conduction electrons and the Kondo singlet (ground state) forms. Consequently, the flow diagram energies grow with $\Delta \Lambda^{n/2}$ due to rescaling. For comparison, the result for the AM is indicated (brown (grey), too). For $\Delta > T_K$ it is energetically favorable to break Cooper pairs and lower the energy by $T_K$ by screening the local spin, thus the ground state is a singlet. For $\Delta > T_K$ Cooper pair formation dominates the low-energy properties and the ground state is a doublet.

3. Energy spectrum

In this section we analyze the many-body energy spectrum generated during the iterative NRG procedure. As already mentioned in the last section, the spectrum consists of a continuum above and subgap bound states below the gap $\Delta$. The resulting many-body eigenenergies are shown in the vertical axis of the many-body eigenstates constructed in figure 1(a). For the AM the effective energy scale separation is even more efficient at energy scales smaller than the gap (compared to the AM).

![Figure 2](image)

**Figure 2.** Energy spectra ((a)–(c)) and the corresponding energy flow diagrams ((d)–(f)) for the lowest 280 eigenenergies of the even NRG iterations of a SC-AM. $U = 0.2$, $\Gamma = 0.01$, $T_K = 1.25 \times 10^{-5}$, $\Delta = 2.5$ and $N_{\text{kept}} = 512$ in all plots. The iteration numbers $N_T$ or $N_{\text{AM}}$, where $\delta_{\text{AM}} \approx T_K$ or $\Delta$, respectively, are indicated by dashed or solid arrows. (a), (d) $\Delta = 0$, AM. Since $\delta_{\text{AM}} \propto 1/n^2$, the eigenenergies (a) fall off exponentially with $n$ and the energy flow diagram (d) converges. At energy scale $T_K$ the localized spin gets screened by the conduction electrons and the Kondo singlet (ground state) forms. Consequently, the flow diagram energies grow with $\Delta \Lambda^{n/2}$ due to rescaling. For comparison, the result for the AM is indicated (brown (grey), too). For $\Delta > T_K$ it is energetically favorable to break Cooper pairs and lower the energy by $T_K$ by screening the local spin, thus the ground state is a singlet. For $\Delta > T_K$ Cooper pair formation dominates the low-energy properties and the ground state is a doublet.

In contrast to the exponential decaying couplings of the Wilson chain (7), the on-site contribution $\Delta$ of the pairing term (8) is constant in magnitude for each site. Consequently, for $\delta_n < \Delta$, the BCS contribution $\Delta$ is relevant perturbation and determines the physics of the system. Typical energy spectra (or energy flow diagrams, respectively) for finite $\Delta$ are shown in figures 2(b), (c) or 2(e), (f)). At energy scale $\Delta$, the exponential reduction of the eigenenergies crosses over to a saturation towards $\Delta$. The characteristic gap as well as the subgap Andreev bound states form.

The energy spectra and the corresponding energy flow diagrams are specified. (a), (d) $\Delta = 0$, AM. Since $\delta_{\text{AM}} \propto 1/n^2$, the eigenenergies (a) fall off exponentially with $n$ and the energy flow diagram (d) converges. At energy scale $T_K$ the localized spin gets screened by the conduction electrons and the Kondo singlet (ground state) forms. Consequently, the flow diagram energies grow with $\Delta \Lambda^{n/2}$ due to rescaling. For comparison, the result for the AM is indicated (brown (grey), too). For $\Delta > T_K$ it is energetically favorable to break Cooper pairs and lower the energy by $T_K$ by screening the local spin, thus the ground state is a singlet. For $\Delta > T_K$ Cooper pair formation dominates the low-energy properties and the ground state is a doublet.

3.1. Competition between Kondo effect and Cooper pair formation

The system possesses two energy scales determining the characteristics of the system, $T_K$ and $\Delta$. The resulting competition between Kondo effect and Cooper pair formation is reflected in the ground state properties of the system: for $T_K \ll \Delta$, $H_\Delta$ is the dominant term of the Hamiltonian. The system lowers its energy by $\Delta$ by the formation of Cooper pairs leading to effective spin zero bosons (singlets) in the reservoir. The lead density of states gets depleted, a gap from $-\Delta$ to $\Delta$ forms. Therefore no electrons are available to screen the localized spin. Consequently, the ground state is a spin doublet with $S_z = \pm 1/2$. In case when the Kondo effect is dominant ($T_K \gg \Delta$), it is energetically favorable to break Cooper pairs so that the localized spin on the quantum dot gets
screened by the non-paired electrons near the Fermi energy. The energy is lowered by $T_K$ and as ground state the typical Kondo singlet forms ($S_z = 0$). The influences of the different scales is also apparent in the NRG energy flow diagrams, see figures 2(e), (f). For example, for $T_K \gg \Delta$, the effect of $H_\Delta$ sets in at iterations after the Kondo fixed point is reached (i.e. at lower energies). A phase diagram for the singlet and doublet sets in at iterations after the Kondo fixed point is reached (i.e. at $n > N_\Delta$, since here they scale as $\varepsilon_j \propto \Lambda^{2-j-n}$. The line sloping upwards at the left of figures (b) and (c) is due to the fact that at every iteration a degeneracy is split and hence an extra eigenenergy is generated.

3.2. Analysis of the continuum

The key feature of NRG is the energy scale separation: the couplings between successive sites of the Wilson chain describing a normal lead fall off exponentially, therefore each added site can be treated as a perturbation of the previous system. However, when generalizing the AM to superconducting leads, a constant on-site energy $\Delta$ is added at each site (see (9)). In order to understand why NRG works even at resolutions well below $\Delta$, we now take a closer look at the structure of the continuum produced during the iterative NRG procedure.

We therefore analyze the (positive) continuum of the single-particle problem. Figure 3 shows an example of a single-particle spectrum for three different scalings of the vertical axis: In (a) the (unscaled) spectrum is plotted versus the iteration number. The NRG eigenenergies decrease with iteration number $n$ and tend towards $\Delta$. However, the decrease of the continuum eigenenergies depends on whether $n$ is smaller or larger than $N_\Delta$, the iteration number for which

$$\varepsilon_{\Delta} \approx \Delta.$$ We find the following asymptotic behavior:

\[
\begin{align*}
\varepsilon_j &\propto \Lambda^{j-n/2} & &\text{for } n < N_\Delta, \\
\varepsilon_j' \propto \varepsilon_j - \Delta &\propto \Lambda^{2-j-n} & &\text{for } n > N_\Delta.
\end{align*}
\]

Primed energies will henceforth always be understood to be measured relative to $\Delta$. Both relations of (15) are illustrated in figures 3(b) and (c), by plotting the eigenenergies $\varepsilon_j$ and $\varepsilon_j'$ in units of $\Lambda^{-n/2}$ and $\Lambda^{-n}$, respectively.

These results can already be understood by further reducing the problem to only the superconducting reservoir ($H_l + H_\Delta$): for fixed iteration $n$, the $j$-dependence of (15) reflects the standard logarithmic discretization of the continuous conduction band, according to which the single-particle energies of the Wilson chain grow in powers of $\Lambda$, i.e. $\varepsilon_k \propto \Lambda^{j}$ [21]. Inserting this into the single-particle dispersion relation of BCS quasiparticles, the effective discretization of the isolated superconducting reservoir is obtained, as sketched in figure 4. The limits yield

\[
\xi_k = \sqrt{\varepsilon_k + \Delta^2}; \quad \frac{\varepsilon_k^2}{2\Delta} \rightarrow \Lambda^{2j} \quad \text{for } \varepsilon_k \ll \Delta
\]

\[
\xi_k' = \frac{\varepsilon_k}{\lambda_n + \Delta} \quad \text{for } \varepsilon_k \gg \Delta
\]

The $n$-dependence of (15) follows heuristically from considering the coupling of two neighboring sites $n-1$ and $n$, with hopping matrix element $t_n \sim \delta_n^\Delta$ and on-site energy $\pm \Delta$ (see (12) and (13), respectively). Since $t_n \sim \Lambda^{-n/2}$, the eigenstates $\lambda_{\pm}$ of this two-state problem show the asymptotic behavior

\[
\lambda_{n,\pm} = \pm \sqrt{\Delta^2 + t_n^2}; \quad \begin{align*}
\lambda_{n,\pm} &\sim t_n \propto \Lambda^{-n/2} & &\text{for } n < N_\Delta, \\
\lambda_{n,\pm} &\propto \frac{t_n^2}{2\Delta} \propto \Lambda^{-n} & &\text{for } n > N_\Delta.
\end{align*}
\]

Note that the $\Lambda^{-n/2}$ versus $\Lambda^{-n}$ scaling of the two limits of (15) implies that the energy scale separation is actually more efficient in the second limit, when the gap dominates the spectral features. This establishes one of the central results of the present paper: The energy scale separation, being the heart of the NRG approach, is not impaired but rather enhanced (w.r.t. $\Delta$) by the presence of the energy gap in the
superconducting leads. Increasing the chain length leads to an exponential enhancement of the resolution at the continuum edge at $\Delta$.

The many-body spectrum is constructed according to equation (14). For $n < N_\Delta$, as in the AM, the mean level spacing (at fixed iteration) does not depend on energy, as can be seen in figures 2(d)–(f). For $n > N_\Delta$ the gap forms, and every many-body state with energy $E_m < 0$ can only stem from adding one electron (or hole) to the Fermi sea. $\epsilon_0$ is the energy of the single-particle subgap level. Therefore in this regime $E_{r:m} = \epsilon_{m}$. Due to many-particle excitations, this dense profile repeats as substructures in the many-body zero. The spectral function is defined as

$$\sigma(\omega) = \left. \frac{1}{\pi} \text{Im} G^R(\omega) \right|_{\mu = 0}.$$

(18)

where $G^R_0(\omega)$ is the Fourier-transformed of the retarded Green’s function $G^R_0(t) \equiv -i \theta(t) \langle [d_{\mu}(t), d_{\mu}^\dagger(0)] \rangle$. Motivated by the structure of the eigenspectrum we distinguish two contributions to the spectral function,

$$\sigma(\omega) = \sum_{|E_m| < \Delta} w_m \delta(\omega - E_m) + A_\sigma(\omega) \theta(|\omega| - \Delta).$$

(19)

$w_m$ denotes the weight of excitations to the Andreev bound states, which contribute as $\delta$-peaks within the gap between $-\Delta$ and $\Delta$. $A_\sigma(\omega)$ represents the continuum contribution, with $|\omega| > \Delta$. In the following we present results for the continuum part of the spectral function. We focus on the behavior of the continuum contribution of the spectral function close to the gap.

4. Spectral function

In this section we study basic properties of the spectral function that are special for the SC-AM. We therefore begin by analyzing the analytic solution of the non-interacting case and find not only a continuum for energies $|\omega| > \Delta$ together with subgap resonances (as expected from the energy spectrum), but also a sharp peak at the gap edge for $\Delta < \Gamma$. The agreement between NRG results and the analytic solution is excellent, especially when resolving that sharp feature, again confirming that NRG is valid also at resolutions well below the effective temperature zero, i.e. at temperature much smaller than any other energy scale of the problem. Only matrix elements connecting the ground state(s) with excited states then contribute.

As NRG parameters we choose $\Lambda = 1.8$ and use $z$-averaging [33] (where for fixed $\Delta$ data is averaged for different discretizations) with an interval spacing of $\delta z = 0.05$ to improve the results. We keep $N_{\text{kept}} = 1024$ at the first 6 to 10 iterations, and use $N_{\text{kept}} = 512$ for the rest of the iterations. We test NRG against the analytic solution at $U = 0$ and find excellent agreement, implying that the broadening procedure as well as the choice of $N_{\text{kept}}$ are adequate for the present problem.

4.1. NRG

NRG calculations of the spectral function are based on the Lehmann representation:

$$A_\sigma(\omega) = \frac{1}{Z} \sum_{mn} \left( e^{-E_n/\hbar k T} + e^{-E_m/\hbar k T} \right) \left| \langle m | d_{\mu}^\dagger | n \rangle \right|^2 \times \delta(\omega - (E_m - E_n)).$$

(20)

Here $Z = \sum_m e^{-E_m/\hbar k T}$ is the partition function at temperature $T$, and $|m\rangle$ denotes an exact eigenstate of the Hamiltonian with eigenenergy $E_m$. The matrix elements of these operators as well as the eigenenergies can be calculated with NRG at all energy scales. The $\delta$-peaks of the continuum contribution are broadened as described by [31, 32]. As expected from the findings about the spectra, broadening w.r.t. $\Delta$ (i.e. using $\omega' = |\omega| - \Delta$ in [31]) leads to good results, see below. We use the full density matrix NRG [30, 31] at effective temperature zero, i.e. at temperature much smaller than any other energy scale of the problem. Only matrix elements connecting the ground state(s) with excited states then contribute.

4.2. Spectral function for $U = 0$

For analyzing the basic properties of $A(\omega)$, we first review the non-interacting problem. There, the local (retarded) Green’s function is known exactly [18]. The continuum contribution to $G^0$ then reads

$$G^0_{\omega}(\omega) = \frac{1}{D(\omega)} (\omega + i\epsilon) + i \Gamma \rho_{\Delta},$$

(21)

with $D(\omega) = (\omega^2 - \Gamma^2 - \epsilon^2) + i 2 \Gamma \omega \rho_{\Delta}$, and $\rho_{\Delta}$ the density of states of a bulk superconductor. At temperature $T = 0$, the latter has the limits

$$\rho_{\Delta}(\omega) = \begin{cases} \frac{\Delta}{2\omega^2} \theta(\omega), & \text{for } \omega' < \Delta, \\ 1, & \text{for } \Delta < |\omega|. \end{cases}$$

(22)

For $\Delta = 0$, equation (21) simplifies to the well known formula for the AM, $G_{\omega-\omega'}^0(\omega) = (\omega - i\epsilon - i\Gamma)^{-1}$. For the spectral function we obtain from equations (21) and (18),

$$A(\omega) = \frac{(\omega + i\epsilon)^2 + \Gamma^2}{(\omega^2 - \epsilon^2 - \Gamma^2)^2 + (2\Gamma^2 \omega \rho_{\Delta})^2} \Gamma \rho_{\Delta}/\pi.$$

(23)

This function is shown for various parameter combinations in figure 5. The common features are (i) the atomic resonance of
width (half width half maximum) \( \sim \Gamma \) centered at \( \varepsilon_d \), reflecting the level broadening due to the level-lead coupling and (ii) a gap from \(-\Delta\) to \(\Delta\), with (iii) bound states at some energy \( \pm \omega_b \) inside the gap. The energy and weight of the subgap contribution is indicated by the thin lines of the red arrows in the insets of figures 5(a), (b). Here they are calculated with NRG, but can be also obtained analytically, see equation (7) of \[18\]. Note that for finite \( \Gamma \) bound states exist also for \( \varepsilon_d \gg \Delta \), see figure 5(b). They asymptotically approach the gap edge for \( |\varepsilon_d| \to \infty \), in accordance with equation (7) of \[18\].

Additionally, the continuum part of the spectral function may feature near-gap sharp peaks, point of interest in the following discussion. The behavior near the gap edge can be approximated (using equation (22) and writing \( s = \text{sign}(\omega) \)) by

\[
A(\omega) \approx \left\{ \begin{array}{ll}
\frac{(s\Delta + \varepsilon_d)^2 + \Gamma^2}{(\Delta^2 - \varepsilon_d^2 + 2\Gamma^2\Delta^2 + 2\Gamma^2\Delta^3/|\omega|)\Gamma\rho_{\Delta}/\pi} & \text{for } |\omega| \gg \Delta \\
\frac{(s\Delta + \varepsilon_d)^2 + (\Gamma^2 + 2\Gamma^2\Delta^3/|\omega|)\Gamma\rho_{\Delta}/\pi}{(s\Delta + \varepsilon_d)^2 + (\Gamma^2 + 2\Gamma^2\Delta^2)\Gamma\rho_{\Delta}/\pi} & \text{for } \omega' \ll \omega, \omega' \ll \Delta
\end{array} \right.
\]

(24)

(25)

The limits given in (25) are indicated by the thin lines in figures 5(c), (d): \( A(\omega) \) increases as \( \sqrt{|\omega|} \) when \( \omega' \) is increased from 0 (dashed lines), decreasing again for \( \omega' > \omega'_{\text{c}} \approx \frac{2\Gamma^2\Delta}{(\Delta^2 - \varepsilon_d^2 + 4\Gamma^2\Delta^2)} \) (which is the zero of derivative of equation (24)). If \( \omega'_{\text{c}} \ll \Delta, \Gamma \), this leads to a very sharp near-gap peak which decreases as \( \rho_{\Delta} \) (dotted lines). Then the near-gap spectral function is greatly enhanced compared to the AM (where the symmetric case yields \( A(0)|\pi\Gamma = 1 \)).

The solution of the AM (dash–dotted lines) describes the high-energy limit of the SC-AM:

\[
A(\omega) \approx \frac{\Gamma/\pi}{(\omega - \varepsilon_d)^2 + \Gamma^2} = -\frac{1}{\pi} \text{Im} \frac{G_0^0}{(\omega - \varepsilon_d)^2 + \Gamma^2} \quad \text{for } \Delta < |\omega|.
\]

(26)

The emergence of the near-gap peak is depicted in figure 6(a) for the symmetric model, where the gap \( \Delta \) is varied over four orders of magnitude, starting from \( \Delta = 0 \). For \( \Delta = 10^{-4} \), we also show numerical NRG results (fat solid line). Their agreement with the analytical results is excellent.

The height of the near-gap peak does not depend on \( \varepsilon_d \), see figure 6(b), where \( \varepsilon_d \) is increased up to 8\( \Gamma \) for fixed \( \Delta = 10^{-4} \).

4.3. Spectral function for finite \( U \)

Figure 7 shows spectral functions of the SC-AM for finite \( U \) and \(-U < \varepsilon_d < 0 \). The two atomic resonances of width \( \Gamma \) are now separated by the Coulomb repulsion, thereby they are centered near \( \varepsilon_d \) and \( \varepsilon_d + U \). The Coulomb repulsion also drives the Kondo effect, yielding a sharp resonance of width \( T_K \) pinned at the Fermi energy. This resonance is cut by a gap reaching from \(-\Delta\) to \(\Delta\), reflecting the influence of the superconducting lead. Depending on the ratio \( T_K/\Delta \), the Kondo resonance can be cut completely (\( T_K/\Delta \ll 1 \)) or emerge clearly (\( T_K/\Delta \gg 1 \)).

Additionally, a similar near-gap feature as found for the non-interacting case may emerge. For \( T_K/\Delta \gg 1 \), a sharp
resonance forms at the gap edge, highly exceeding the Kondo
resonance, see figure 7 for the symmetric case (where the
height of the Kondo resonance is given by $1/\pi \Gamma$) as well as the
antisymmetric case. The asymptotic behavior of the near-gap
peak is in agreement with that given in equations (25) and (26).
In particular, for $\omega' \to 0$, the continuum edge decreases as $\sqrt{\omega'}$.

**5. Conclusion**

The NRG is a well established method for a variety of
quantum impurity models. It is usually applicable in the whole
parameter range and allows to calculate physical quantities
for a wide range of temperatures and frequencies. In this
paper we showed that in the presence of a superconducting
reservoir, NRG provides information for resolutions far below
the energy scale of the gap $\Delta$. Moreover, Wilsonian energy
scale separation, being the heart of the success of the NRG
approach, is not impaired but rather enhanced by the presence
of the energy gap of the superconducting leads. This allows
sharp features of spectral functions at the continuum gap edge
to be resolved. Our calculations of the impurity spectral
function cover the whole region from $\Delta \gg T_K$ to $\Delta \ll T_K$. In
the latter case, we find a sharp peak at the continuum gap edge,
vastly exceeding the Kondo resonance contribution. We expect
this to result in an enhanced linear conductance, as recently
reported for experiments with carbon nanotube quantum dots coupled to superconducting leads [24].

The ability of the NRG to resolve spectral functions at energy resolutions well below the gap should be useful for other problems as well. As discussed in detail in [34], the problem of an impurity in a superconducting host can be mapped (under certain conditions) to a model in which the superconductor couples to a normal metal, with a modified density of states. For the problem studied in this paper we have performed such a mapping; the resulting Hamiltonian is given in equation (8). In this case, the oscillating on-site energies, $(-1)^n \Delta$, generate a hard gap of width $2\Delta$.

This connection allows us, in principle, to calculate the dynamic quantities for impurity models with arbitrary gapped bath spectral function (but for this one still has to develop an algorithm which produces directly the chain parameters from the hybridization function). Such calculations might also help to improve the resolution of the NRG in DMFT calculations for the Hubbard model where the standard implementation of the NRG does not describe the shape of the Hubbard bands properly (for dynamic DMRG calculations for this problem, see [35]).

Acknowledgments

We acknowledge helpful discussions with G Zarán and A Tóth. This research was supported by the DFG through SFB 484 (RB), SFB-TR12 and De-730/3-2 (AW) and SFB 631 (TH, AW). Financial support of the German Excellence Initiative via the ‘Nanosystems Initiative Munich (NIM)’ is gratefully acknowledged.

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