Green’s functions from real-time bold-line Monte Carlo

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We present two methods for computing two-time correlation functions or Green’s functions from real time bold-line continuous time quantum Monte Carlo. One method is a formally exact general-ized auxiliary lead formalism by which spectral properties may be obtained from single-time observables. The other involves the evaluation of diagrams contributing to two-time observables directly on the Keldysh contour. Additionally, we provide a detailed description of the bold-line Monte Carlo method. Our methods are general and numerically exact, and able to reliably resolve high-energy features such as band edges. We compare the spectral functions obtained from real time methods to analytically continued spectral functions obtained from imaginary time Monte Carlo, thus probing the limits of analytic continuation.

I. INTRODUCTION

Strongly correlated electron materials exhibit fascinating collective behavior which has long challenged our understanding. A few notable examples include Mott metal–insulator transitions in transition metal oxides, colossal magnetoresistance in perovskite manganites, quantum criticality in heavy fermion systems, and high temperature superconductivity in copper oxides and pnictides. Of central experimental interest are dynamical properties, both of excited states near the Fermi energy and at highly excited energies. However, the theoretical description of correlated electron materials has proven difficult: because of the absence of a small parameter, perturbation theory is in general unreliable. Traditional materials science techniques, among them the density functional theory (DFT) and the GW approximation, do not capture strong correlation effects, while standard numerically exact lattice methods are limited to small lattices, high symmetry points or one dimensional systems.

An alternative to directly determining quantities in a correlated quantum lattice model is provided by the dynamical mean field theory (DMFT) In the approximation that correlations are local, the physics of the lattice model can be mapped onto a numerically tractable impurity model: a finite interacting system coupled self-consistently to a noninteracting effective bath. Extensions of DMFT systematically relax the approximation of locality.

In addition to their importance for dynamical mean field theory, quantum impurity models are extremely important in their own right, since they appear directly in a variety of problems including nanoscience (where they are used in describing quantum transport) and catalysis and surface science (where they can be used to model the adsorption of molecules on surfaces). The general solution of such impurity models remains a formidable challenge. Exact solutions are only available in specific limits. A variety of more general semi-analytical methods have been used successfully. Numerical approaches—all of which are general in principle, but have different advantages and limitations in practice—include the numerical renormalization group, exact diagonalization, configuration interaction, hierarchical equations of motion, and quantum Monte Carlo.

Continuous time Monte Carlo (CTQMC) algorithms are numerically exact and very general in the sense that their computational complexity is independent of the spectral resolution and band shape, a property which is particularly crucial in the context of DMFT. Most algorithms are formulated on the Matsubara axis, and therefore access to single- and two-particle response functions as measured in experiments requires an analytic continuation to the real axis. This analytic continuation problem is ill-posed: small fluctuations in the Matsubara data, due to Monte Carlo noise, cause large changes in the output real frequency data and render the direct inversion of the continuation kernel unreliable for practical purposes. Instead, analytical continuation algorithms perform the inversion under additional assumptions, e.g. that the function be described by a small number of zeros and poles on the complex plane so that it deviate as little as possible from a ‘default model’ function while being consistent with the input data within some predetermined error bounds or that it be as smooth as possible.

While analytical continuation methods produce spectral functions that are consistent with Matsubara data, the bias that they introduce on the real axis is hard to quantify and, as a consequence, controlled error estimates and confidence intervals on the real axis are not available even for numerically exact Matsubara data. Real time methods, on the other hand, embody a controlled way to study the dynamical properties of quantum impurity models. Monte Carlo impurity solvers have also been formulated on the Keldysh contour such that analytical continuation is unnecessary (although a combination of real time and Matsubara techniques may be beneficial). In the weakly interacting limit, response functions at short times immediately after a quantum quench have been obtained in this way. Using the...
newest generation of real time impurity solvers, bold-line CTQMC single time quantities have been obtained at substantially longer times and at large interaction strength, so that, in combination with reduced dynamics techniques, the long time steady state behavior in the Kondo regime was reached after a quantum quench.

Here we present two ways of obtaining spectral functions from real time CT-QMC in the strong correlation limit. These methods were recently used to access nonequilibrium spectral properties; we emphasize, however, that the methods are equally applicable to nonequilibrium situations. The first method obtains real frequency spectral functions from two-time correlation functions which are then Fourier transformed. The second is based on an auxiliary current formulation, and is more efficient when one is interested in obtaining spectral steady state or equilibrium properties. This second method is compatible with the reduced dynamics technique of Ref. 54 and with any numerical solver applicable to transport. It simulates a direct measurement of the real frequency spectral functions from two-time correlators, and we explore the limitations of analytical continuation.

The coupling densities defined as $\Gamma_{ij} (\omega) = 2\pi \sum_{k \in \ell} t_{ik}^a t_{jk}^d \delta (\omega - \varepsilon_k)$, fully define the properties of bath and hybridizations for the purpose of this paper. We will assume $\Gamma_{ij} (\omega) \equiv \delta_{ij} \Gamma_\ell$ to be diagonal in dot orbital space but allow it to differ for each lead, such that $\Gamma_\ell (\omega) \equiv 2\pi \sum_{k \in \ell} |t_{ik}|^2 \delta (\omega - \varepsilon_k)$. General coupling density matrices will be discussed briefly in Sec. IV.

Information about thermodynamic parameters, e.g. the lead chemical potential or the lead temperature, is encapsulated in the initial conditions of the system. For example, fermionic leads at an inverse temperature $\beta_\ell$ and a chemical potential $\mu_\ell$ are described by an initial Fermi Dirac distribution $f_\ell (\omega) \equiv \frac{1}{1 + e^{\omega/(\beta_\ell - \mu_\ell)}}$.

II. MODEL

We consider a general quantum impurity model described by

$$ H = H_D + H_B + V. \tag{1} $$

$H_D$, the dot Hamiltonian, can be any (generally interacting) Hamiltonian of the form

$$ H_D = \sum_{i=1}^{d} \varepsilon_i d_i^\dagger d_i + \sum_{ijkl} U_{ijkl} d_i^\dagger d_j^\dagger d_k d_l + \cdots, \tag{2} $$

with $d_i^{(d)}$ denoting dot operators. $d$, the total number of degrees of freedom on the dot, is assumed to be small. $\varepsilon_i$ describes on-site energy levels, the interaction $U_{ijkl}$ represents the strength of four-operator (two-body) terms, and the ellipsis all potential higher order interactions.

The bath or lead Hamiltonian is:

$$ H_B = \sum_{\ell} H_{B\ell}, \tag{3} $$

$$ H_{B\ell} = \sum_{k \in \ell} \varepsilon_k a_k^\dagger a_k, \tag{4} $$

$H_B$ is written here as a sum of $N_B$ bath terms $H_{B\ell}$ which may be characterized by different dispersions $\varepsilon_k$ and different thermodynamic parameters (such as temperature and chemical potential) entering through the initial conditions. In a molecular electronics scenario with left and right leads, $N_B = 2$ and the index $\ell$ takes the corresponding values $L$ and $R$. $H_B$ has infinite degrees of freedom described by lead operators $a_k^{(d)}$ and a lead dispersion $\varepsilon_k$ but is noninteracting.

The third term is the hybridization Hamiltonian $V$, which describes population transfer between the dot and leads. Here we assume it to have the bilinear form

$$ V = \sum_{\ell} V_\ell, \tag{5} $$

$$ V_\ell = \sum_{i=1}^{d} \sum_{k \in \ell} (t_{ik} a_k^\dagger d_i + t_{ik}^* d_i^\dagger a_k) \tag{6} $$

characterized by a hopping $t_{ik}$ from dot to lead. The coupling densities $\Gamma_{ij} (\omega)$, defined as $\Gamma_{ij} (\omega) = 2\pi \sum_{k \in \ell} t_{ik}^a t_{jk}^d \delta (\omega - \varepsilon_k)$, fully define the properties of bath and hybridizations for the purpose of this paper. We will assume $\Gamma_{ij} (\omega) \equiv \delta_{ij} \Gamma_\ell$ to be diagonal in dot orbital space but allow it to differ for each lead, such that $\Gamma_\ell (\omega) \equiv 2\pi \sum_{k \in \ell} |t_{ik}|^2 \delta (\omega - \varepsilon_k)$. General coupling density matrices will be discussed briefly in Sec. IV.

III. AUXILIARY CURRENT METHOD

In their seminal 1992 paper, Meir and Wingreen showed that if initial correlations can be neglected, the steady-state current $I_\ell$ out of lead $\ell$ can be written as

$$ I_\ell = \frac{ie}{\hbar} \int d\omega \times \text{Tr} \left\{ \Gamma_\ell (\omega) \left[ f_\ell (\omega) [G^> (\omega) - G^< (\omega)] + G^< (\omega) \right] \right\}. \tag{7} $$

Here, $e$ and $h$ are the electron charge and Planck's constant (both are set to one from here onward); $f_\ell$ is the initial occupation of lead $\ell$; and $G^>$, $G^<$ and $G^<$ are the
As previous authors have pointed out, the spectral function $\Gamma_A(\omega)$ and $\Gamma_L(\omega)$ are independent of energy and the orbital structure of $\ell$ $(\ell \in A)$ is diagonal at all frequencies such that it can be diagonalized by coupling the system to an auxiliary lead. Following Refs. 55 and 56, we use the conservation of current $\sum \mathcal{I}_l + \mathcal{I}_A = 0$ along with Eq. (7) and (8) to construct an expression for $\mathcal{I}_A$ in which $G^\pm$ no longer appears:

$$I_A = I_A - \frac{\eta}{\sum \lambda_\ell + \eta} \left( \sum \mathcal{I}_l + \mathcal{I}_A \right)$$

$$= \pi \int d\omega \text{Tr} \{ \Gamma(\omega) A(\omega) \} \times \left\{ \frac{\eta}{\sum \lambda_\ell + \eta} \left[ f_A \left( \sum \lambda_\ell \right) - \lambda_\ell f_\ell \right] \right\}.$$ 

In the limit where $\eta \rightarrow 0$, system properties such as the Green’s functions are unaffected by the presence and properties of the auxiliary lead. Therefore, the derivative of the current $I_A$ with respect to the chemical potential in the auxiliary lead, $\frac{dI_A}{d\mu_A}$, only contains contributions from $f_A$:

$$\lim_{\eta \rightarrow 0} \frac{d}{d\mu_A} I_A = \lim_{\eta \rightarrow 0} \pi \int d\omega \text{Tr} \{ \Gamma(\omega) A(\omega) \} \times \left\{ \frac{\eta}{\sum \lambda_\ell + \eta} \left[ \frac{df_A}{d\mu_A} \left( \sum \lambda_\ell \right) \right] \right\}.$$ 

With our assumption that $\Gamma$ and $A$ are diagonal, the trace of $\Gamma$ over the impurity degrees of freedom is a product of over their elements (up to a factor of $d$, which we will ignore). If lead $A$ is maintained at low enough temperature, we can also set $\lim_{\beta \rightarrow \infty} \frac{df_A(\omega)}{d\mu_A} = -\delta(\omega - \mu_A)$, which allows us to perform the integration and obtain

$$A(\mu_A) = -\frac{1}{\pi \eta} \Gamma^{-1}(\mu_A) \sum \lambda_\ell \frac{dI_A}{d\mu_A}$$

$$= -\sum \Gamma_\ell(\mu_A) + \Gamma_A(\mu_A) \frac{1}{\pi} \Gamma^{-1}_A(\mu_A) \frac{dI_A}{d\mu_A}.$$ 

This result is identical to the one of Refs. 55 and 56 except in that $\Gamma$ is allowed to have an arbitrary energy dependence. The approach outlined here is, in general, limited to systems with only diagonal dot correlation and hybridization functions.

Eq. (16) and (17) allow sampling the spectral function by measuring the current flow between the dot and a weakly coupled auxiliary lead at a variety of chemical potentials. This could also in principle be done in an experimental setting by varying the gate voltage of an auxiliary lead, and this is perhaps the greatest advantage of this variant of the method. However, the setup is only useful in cases where $\Gamma(\omega)$ is not suppressed. Near regions where $\Gamma(\omega)$ is small, $\frac{d\mathcal{I}_A}{d\mu_A}$ must be very small in order for $A$ not to diverge, and the numerical computation or measurement of the derivative must be performed to a very high accuracy. For this reason, the presence of noise makes the numerical computation of this derivative impractical within a Monte Carlo simulation, at least for cases where one is interested in behavior in or near a gap in the lead’s spectrum.
IV. DOUBLE PROBE SCHEME

In order to bypass the limitation outlined above and the need to compute derivatives, we propose an alternative scheme. The current in the auxiliary lead is (see (7))

\[ I_A = \frac{ie}{2\hbar} \int d\omega \sum_i \left\{ \Gamma_A(\omega) \left[ f_A(\omega) [G^r(\omega) - G^a(\omega)] + G^< (\omega) \right] \right\}. \]

(18)

In the limit \( \Gamma_A \rightarrow 0 \) the partial derivative of this equation with respect to \( \mu_A \) immediately yields Eq. (17) without the factor \( \frac{\sum_i \Gamma_i}{\sum_i \Gamma_i + \Gamma_A} \approx 1 - \frac{\ddot{r}_A}{\sum_i \Gamma_i} = 1 - \frac{\eta}{\sum_i \Gamma_i} \), the higher order terms of which constitute corrections in powers of \( \frac{\eta}{\sum_i \Gamma_i} \). Our form ignores the secondary effects of the auxiliary lead on the currents through other leads, and only becomes accurate in the limit of an auxiliary lead that is weakly coupled to the system (\( \eta \) small), a criterion that can be verified \textit{a posteriori} by varying \( \eta \). Additionally, no assumptions regarding the relationship or proportionality between \( \Gamma_A \) and \( \{ \Gamma_i \} \) need to be made. This freedom allows the experimentally unrealistic but numerically convenient choice of introducing two auxiliary leads, one empty and one full, set up in such a way that they are coupled to the system at only a single frequency \( \omega' \):

\[ f_A' (\omega) = \begin{cases} 0: & i = 0, \\ 1: & i = 1, \end{cases} \]

(19)

\[ \Gamma_A' (\omega) = \eta \delta (\omega - \omega'). \]

(20)

Using these definitions, we define two auxiliary currents

\[ I_A' (\omega') = \frac{i}{2} \eta \{ G^< (\omega') \}, \]

(21)

\[ I_A^\prime (\omega') = \frac{i}{2} \eta \{ [G^r(\omega') - G^a (\omega')] + G^< (\omega) \} \]

(22)

\[ = \eta \frac{\pi}{2} A (\omega') + I_A (\omega'). \]

(23)

The assumption made here, as in Ref. [55] and [56] and the preceding section, is that \( \eta \) is small enough that the properties of the auxiliary lead have a negligible effect on the physical properties of the full system. We note in passing that this same idea also allows for the use of a single wide-bandwidth auxiliary lead in the case where the coupling densities are \textit{not} proportional.

Two independent calculations, one for an empty lead and a second one for a full lead, provide a setup (see right side of Fig. 1) with which the dot spectral function can be obtained. Restoring physical constants for a moment,

\[ A (\omega) = \lim_{\eta \rightarrow 0} \frac{2\hbar}{e i \eta} \left[ I_A^\prime (\omega) - I_A' (\omega) \right]. \]

(24)

This analytically exact result is not restricted to any particular way of solving the impurity model, and is therefore usable within any formalism where one has access to currents. Apart from quantum Monte Carlo this includes hierarchical equation of motion methods. Intuitively, the introduction of two leads of opposite populations simultaneously should correct current conservation to some degree (since the coupling densities are no longer proportional, an exact statement is hard to make), but they can also be connected one at a time at the price of small loss in accuracy, as long as \( \eta \) is taken to be small enough.

Equations Eq. (21) and Eq. (22) also provide direct access to the lesser Green’s function \( G^< (\omega) \), providing the full information about single particle correlations even in nonequilibrium situations. Finally, since \( \Gamma_A(\omega) \) can be chosen to have any dot orbital matrix structure, it is straightforward to find a choice that allows extracting individual elements of \( A_{ij} (\omega) \).

Since no numerical derivative needs to be taken, the double probe scheme performs better within a Monte Carlo simulation, in addition to providing more information (such as \( G^< \)). Any practical implementation will need to approximate the delta function in Eq. (20) by a smooth numerical approximation. In this work, we approximate \( \Gamma_{A}^\prime \) with Gaussians with an amplitude and width small enough that the results are independent of them within the numerical precision bounds and frequency resolution we need (a good rule of thumb is to start the amplitude and width at least an order of magnitude or two below any other energy scale in the problem; one then verifies convergence in both parameters of the Gaussian by systematically varying them). The picture which emerges is physically intuitive: we probe the system using a pair of virtual leads, one of which is empty of electrons while the other is full. The leads are coupled very weakly and only within a narrow frequency range. By measuring the rate at which electrons are injected into one lead and extracted from the other, one obtains information about the average density of single-electron excitations within the frequency range probed.

The auxiliary current method by construction contains two-time information, but strictly speaking this only corresponds to the two-time correlation function in a certain adiabatic limit (where the relaxation of the auxiliary current to its final value is much faster than that of the system’s dynamical timescales). However, the method has a straightforward extension to the handling of two-time correlation functions; this can be seen by considering eq. (15) in Ref. [55] which connects the current through any lead \( \ell \) to the two-time Green’s function \( G^r (t_1, t_2) \) by way of a generalized coupling density \( \Gamma_\ell (\omega, t_1, t_2) \). By considering this expression, it is easy to see how a generalized, time-dependent auxiliary coupling density might be devised, which fully and rigorously probes the full two-time correlation functions. This is beyond the scope of the present work.
V. BOLD-LINE MONTE CARLO

The auxiliary current formalism is only useful in combination with a method for computing currents through interacting quantum dots. This requires the solution of a correlated quantum many-body problem, and therefore a numerical method. Several approaches exist. In this section we present a detailed description of the real-time bold-line continuous-time hybridization expansion quantum Monte Carlo algorithm introduced in Ref. 50. We focus in particular on the observables required for obtaining two-time correlation functions but, for the sake of simplicity, limit ourselves to the single-orbital Anderson impurity model.

The method is based on a stochastic summation of all diagrams containing partially summed (‘bold’ rather than ‘bare’) propagator lines and vertex functions. So far, these partially summed propagators have come from the non-crossing or one-crossing approximations (NCA or OCA\(^{52}\)) but the method is more general: In a first, quasi-analytic step, an underlying diagrammatic approximation selecting some (but not all) diagrams is chosen, and propagators within that approximation are obtained analytically or via the solution of a set of coupled integral equations. In a second step, all corrections to the propagators are summed up using a stochastic Monte Carlo procedure, so that the resulting sum contains all diagrams and therefore becomes numerically exact. The precise choice of the underlying approximation determines the speed of convergence to the exact result, the statistical uncertainty, and the feasibility of the method for any given system, but has no effect on the final answer if convergence is attained. The imaginary time version of bold-line CTQMC has been introduced in Ref. 51. The formulation on the Keldysh contour and the advantages of bold-line CTQMC over bare CTQMC in the non-equilibrium context have been discussed and benchmarked in Ref. 50 and an extension to OCA and to a non-equilibrium context have been discussed and benchmarked in Ref. 51.

We consider the single-orbital Anderson impurity model, where we set \(d = 2\) in Eq. (2) such that there may be up to two electrons having opposite spin indices \(\uparrow\) and \(\downarrow\) on the dot simultaneously. The dot Hamiltonian becomes

\[
H_D = \sum_{\sigma = \uparrow, \downarrow} \varepsilon_\sigma d_\sigma^\dagger d_\sigma + U d_\sigma^\dagger d_\sigma^{\dagger} d_\sigma^{\dagger} d_\sigma, \tag{25}
\]

the lead Hamiltonian

\[
H_B = \sum_{\sigma = \uparrow, \downarrow} \sum_{k \in k_\sigma} \varepsilon_k a_k^\dagger a_k, \tag{26}
\]

and the dot-lead hybridization

\[
V_I = \sum_{\sigma = \uparrow, \downarrow} \sum_{k \in k_\sigma} \left( t_k a_k^\dagger d_\sigma + t_k^* d_\sigma^\dagger a_k \right), \tag{27}
\]

We first derive a version of the bare real time hybridization expansion by introducing a perturbation theory written in terms of many-body atomic state propagators. The expectation value of an operator \(A\) at time \(t\) in the interaction picture with respect to \(H_0 \equiv H - V\) is

\[
\langle A(t) \rangle = \text{Tr} \left\{ \rho U^{\dagger}(t) A_I(t) U(t) \right\}. \tag{28}
\]

Here \(\rho = \rho_D \otimes \rho_B\) is the initial density matrix. We assume that it can be factorized into dot and lead parts (this assumption can be relaxed by adding an imaginary branch to the contour). \(U(t) = e^{iH_0 t} e^{-iH t}\) is the interaction picture propagator, and interaction picture operators are denoted by a subscript \(I\) and given by \(A_I(t) = e^{iH_0 t} A e^{-iH_0 t}\). We then expand \(U\) and \(U^{\dagger}\) in the form\(^{29}\)

\[
U(t) = \sum_{n=0}^{\infty} (-i)^n \int_0^t dt_1 \int_0^{t_1} dt_2 \ldots \int_0^{t_{n-1}} dt_n \times V_I(t_1) \ldots V_I(t_n) , \tag{29}
\]

thus obtaining an infinite series of terms. It is convenient to imagine that interactions coming from \(U^{\dagger}\) (it exist on a forward branch while those coming \(U\) (it are on a backward branch; the union of these two contours forms the real part of the Keldysh contour. Notably, since in our choice of model and expansion \(H_0\) is interacting, Wick’s theorem does not hold and one cannot at this point apply standard diagrammatic tools.

The interaction form of the Hybridization term \(V_I(t)\) is

\[
V_{II}^I(t) = \sum_{\sigma \in \uparrow, \downarrow} \sum_{k \in k_\sigma} \left\{ t_{\sigma k} e^{i(\varepsilon_{\sigma} + U d_\sigma^\dagger d_\sigma - \varepsilon_{\sigma k}) t} d_\sigma^\dagger d_{\sigma k} + t_{\sigma k}^* e^{-i(\varepsilon_{\sigma} + U d_\sigma^\dagger d_\sigma - \varepsilon_{\sigma k})} d_{\sigma k}^\dagger d_\sigma \right\} , \tag{30}
\]

with \(\bar{\sigma}\) denoting the spin opposite of \(\sigma\). All terms of \(V_I^I\) change the state of the dot by adding or removing one electron from it, while time evolution with \(H_0\) leaves the dot state invariant. It is therefore convenient to group certain sets of contributions together in dressed \((G)\) and bare \((G^{(0)})\) propagators

\[
G_{\alpha \beta}(t) \equiv \langle \alpha | \text{Tr}_B \left\{ \rho e^{-iHt} \right\} | \beta \rangle , \tag{31}
\]

\[
G^{(0)}_{\alpha \beta}(t) \equiv \langle \alpha | \text{Tr}_B \left\{ \rho e^{-iH_0 t} \right\} | \beta \rangle , \tag{32}
\]

between the many-body states \(\alpha\) and \(\beta\) on the dot, with the leads traced out. Both atomic state propagators and correlation functions are often referred to as Green’s functions (in addition, the retarded and advanced Green’s functions are often referred to as propagators), but the objects defined here differ from correlation functions in several important ways—most notably, they have an exponentially larger dimensionality, since they contain matrix elements between pairs of many-body states \(\alpha, \beta\) rather than single-particle level indices. Diagrammatically, a propagator over a segment of one branch of the
Using Eq. 30 and Eq. 33 this can be written as

\[
G_{\alpha\alpha}(t-t') = G_{\alpha\alpha}^{(0)}(t-t') + \sum_{\beta} \int_{0}^{t} dt_{1} \int_{0}^{t_{1}} dt_{2} 
× G_{\alpha\beta}^{(0)}(t-t_{1}) G_{\beta\beta}^{(0)}(t_{1}-t_{2}) G_{\beta\alpha}^{(0)}(t_{2}-t')
× \Delta_{\alpha\alpha}^{\beta}(t_{1}-t_{2}) + \ldots,
\]

where

\[
\Delta_{\alpha\alpha}^{\beta}(t_{1}-t_{2}) \equiv \langle \alpha| d_{\beta}^{\dagger} \beta \rangle \langle \beta| d_{\alpha} \rangle
\times \sum_{k \in \ell} | t_{k} |^{2} \text{Tr}_{B} \left\{ a_{l_{1} k}^{\dagger} a_{l_{2} k} \right\}
× \langle \alpha| d_{\beta}^{\dagger} | \beta \rangle \langle \beta| d_{\alpha} \rangle
\times \sum_{k \in \ell} | t_{k} |^{2} \text{Tr}_{B} \left\{ a_{l_{1} k}^{\dagger} a_{l_{2} k} \right\}
\]
defines the hybridization function (note that spin indices have been suppressed for brevity). The hybridization function \(\Delta_{\alpha\alpha}(t_{1}, t_{2}) = \sum_{k \in \ell} | t_{k} |^{2} \text{Tr}_{B} \left\{ a_{k}^{\dagger} a_{k} \right\}\) for each lead \(\ell\) can be expressed in terms of the coupling densities \(\Gamma_{\ell}\) of that lead and its initial occupation probability, both of which are obtained from the lattice Green’s function within DMFT. It is \(\Delta_{\ell}^{\alpha\alpha}\) if \(t_{1}\) appears before \(t_{2}\) on the Keldysh contour, and \(\Delta_{\ell}^{-}\) otherwise, with

\[
\Delta_{\ell}^{\alpha\alpha}(t_{1}, t_{2}) = -2i \int_{-\infty}^{\infty} \frac{d\omega}{2\pi} e^{-i\omega(t_{1}-t_{2})} \Gamma_{\ell}(\omega) f(\omega - \mu_{\ell}),
\]

\[
\Delta_{\ell}^{-}(t_{1}, t_{2}) = 2i \int_{-\infty}^{\infty} \frac{d\omega}{2\pi} e^{-i\omega(t_{1}-t_{2})} \Gamma_{\ell}(\omega) \left[ 1 - f(\omega - \mu_{\ell}) \right].
\]

This second order term contains all the elements occurring in the expansion, namely the bare propagators and hybridization lines. A graphical representation is shown in Fig. 2 in the upper left panel, atomic state propagators for a given state are represented by a pair of lines; the upper one representing spin up, the lower one spin down. Dotted lines represent empty states and solid lines represent occupied states, such that the full set of \(2^{N}\) states is represented by \(N\) lines. Additionally, thick (or bold) lines represent full propagators while thin lines stand for bare propagators. Full propagators are built out of all possible combinations of bare propagators and interaction vertices (shown to the upper right), with all interaction times integrated over. Every hybridization line connects two interactions, and each such line comes with the minus sign of Eq. 33. The second order contribution explicitly written above includes both the 2nd and 3rd term in the expansion of \(G_{\alpha\alpha}\) shown in the bottom part of Fig. 2 where an additional 8th order diagram is also shown.
Figure 3. The matrix elements of the NCA self energy (two terms on the left with a single hybridization line) and of the OCA self energy (all four terms) in diagrammatic form.

\[
\begin{align*}
\Sigma_{00} &= \begin{tikzpicture} [baseline=-0.5ex]
\draw (0,0) -- (0.5,0) -- (0.5,0.5) -- (0,0.5) -- cycle;
\draw (0,0) -- (0.5,0.5);
\draw (0,0.5) -- (0.5,0.5);
\end{tikzpicture} + \begin{tikzpicture} [baseline=-0.5ex]
\draw (0,0) -- (0.5,0) -- (0.5,0.5) -- (0,0.5) -- cycle;
\draw (0,0) -- (0.5,0.5);
\draw (0,0.5) -- (0.5,0.5);
\end{tikzpicture} + \begin{tikzpicture} [baseline=-0.5ex]
\draw (0,0) -- (0.5,0) -- (0.5,0.5) -- (0,0.5) -- cycle;
\draw (0,0) -- (0.5,0.5);
\draw (0,0.5) -- (0.5,0.5);
\end{tikzpicture} + \begin{tikzpicture} [baseline=-0.5ex]
\draw (0,0) -- (0.5,0) -- (0.5,0.5) -- (0,0.5) -- cycle;
\draw (0,0) -- (0.5,0.5);
\draw (0,0.5) -- (0.5,0.5);
\end{tikzpicture}
\end{align*}
\]  
\[
\begin{align*}
\Sigma_{11} &= \begin{tikzpicture} [baseline=-0.5ex]
\draw (0,0) -- (0.5,0) -- (0.5,0.5) -- (0,0.5) -- cycle;
\draw (0,0) -- (0.5,0.5);
\draw (0,0.5) -- (0.5,0.5);
\end{tikzpicture} + \begin{tikzpicture} [baseline=-0.5ex]
\draw (0,0) -- (0.5,0) -- (0.5,0.5) -- (0,0.5) -- cycle;
\draw (0,0) -- (0.5,0.5);
\draw (0,0.5) -- (0.5,0.5);
\end{tikzpicture} + \begin{tikzpicture} [baseline=-0.5ex]
\draw (0,0) -- (0.5,0) -- (0.5,0.5) -- (0,0.5) -- cycle;
\draw (0,0) -- (0.5,0.5);
\draw (0,0.5) -- (0.5,0.5);
\end{tikzpicture} + \begin{tikzpicture} [baseline=-0.5ex]
\draw (0,0) -- (0.5,0) -- (0.5,0.5) -- (0,0.5) -- cycle;
\draw (0,0) -- (0.5,0.5);
\draw (0,0.5) -- (0.5,0.5);
\end{tikzpicture}
\end{align*}
\]  
\[
\begin{align*}
\Sigma_{22} &= \begin{tikzpicture} [baseline=-0.5ex]
\draw (0,0) -- (0.5,0) -- (0.5,0.5) -- (0,0.5) -- cycle;
\draw (0,0) -- (0.5,0.5);
\draw (0,0.5) -- (0.5,0.5);
\end{tikzpicture} + \begin{tikzpicture} [baseline=-0.5ex]
\draw (0,0) -- (0.5,0) -- (0.5,0.5) -- (0,0.5) -- cycle;
\draw (0,0) -- (0.5,0.5);
\draw (0,0.5) -- (0.5,0.5);
\end{tikzpicture} + \begin{tikzpicture} [baseline=-0.5ex]
\draw (0,0) -- (0.5,0) -- (0.5,0.5) -- (0,0.5) -- cycle;
\draw (0,0) -- (0.5,0.5);
\draw (0,0.5) -- (0.5,0.5);
\end{tikzpicture} + \begin{tikzpicture} [baseline=-0.5ex]
\draw (0,0) -- (0.5,0) -- (0.5,0.5) -- (0,0.5) -- cycle;
\draw (0,0) -- (0.5,0.5);
\draw (0,0.5) -- (0.5,0.5);
\end{tikzpicture}
\end{align*}
\]  
\[
\begin{align*}
\Sigma_{33} &= \begin{tikzpicture} [baseline=-0.5ex]
\draw (0,0) -- (0.5,0) -- (0.5,0.5) -- (0,0.5) -- cycle;
\draw (0,0) -- (0.5,0.5);
\draw (0,0.5) -- (0.5,0.5);
\end{tikzpicture} + \begin{tikzpicture} [baseline=-0.5ex]
\draw (0,0) -- (0.5,0) -- (0.5,0.5) -- (0,0.5) -- cycle;
\draw (0,0) -- (0.5,0.5);
\draw (0,0.5) -- (0.5,0.5);
\end{tikzpicture} + \begin{tikzpicture} [baseline=-0.5ex]
\draw (0,0) -- (0.5,0) -- (0.5,0.5) -- (0,0.5) -- cycle;
\draw (0,0) -- (0.5,0.5);
\draw (0,0.5) -- (0.5,0.5);
\end{tikzpicture} + \begin{tikzpicture} [baseline=-0.5ex]
\draw (0,0) -- (0.5,0) -- (0.5,0.5) -- (0,0.5) -- cycle;
\draw (0,0) -- (0.5,0.5);
\draw (0,0.5) -- (0.5,0.5);
\end{tikzpicture}
\end{align*}
\]

The essence of the CTQMC method is that one can evaluate the sum of all diagrams by sampling them stochastically using the Metropolis algorithm. In particular, the ‘continuous time’ descriptor implies that, unlike in previous algorithms, systematic Trotter time-discretization errors are absent. In the hybridization expansion continuous-time algorithm, the stochastic sampling of diagrams is done by insertion of pairs of dot operators and hybridization lines: Namely, in each Monte Carlo step, a hybridization line and two dot operators are created, destroyed, or moved. This set of updates can be shown to respect ergodicity, and moves can be weighed in such a way that detailed balance is also maintained. Since the leads are noninteracting both initially and un-
are four such sets, one for each value of the initial condition (the right index) at a pair of propagation times, beginning of the contour and terminating at some final time correlation function, with a special hybridization line (grey) coupling to the final time. Lower panel: two of the NCA vertex equations which take into account diagrams near the beginning of the contour. The gray parts of the lines are not part of the diagrams, but instead illustrate their location on the Keldysh contour.

Figure 4. Top left panel: a diagram on the two-branch Keldysh contour. Note the branch crossing hybridization line, which is not contained in single-branch propagators. Top right panel: a diagram contributing to the current or a two-time correlation function, with a special hybridization line (grey) coupling to the final time. Lower panel: two of the NCA vertex equations which take into account diagrams near the beginning of the contour. The gray parts of the lines are not part of the diagrams, but instead illustrate their location on the Keldysh contour.

ple shows a contribution to the probability that the dot is in the $\alpha = |\downarrow\rangle$ state at the final time at the right tip of the contour, while beginning in state $|0\rangle$. Inter-branch hybridization lines are a crucial part of the expansion, as they allow changes to the state at the tip.

Generalizing the Monte Carlo process to the double contour poses no difficulties greater than those entailed by the extra bookkeeping (due to the fact that vertices on the two branches contribute opposite signs, while time on the two branches flows in opposite directions). However, the dressed propagators contain no information about inter-branch diagrams. Just as one can use the bold line technique to renormalize the propagators, it is possible and highly advantageous to obtain partial summations of contour-crossing diagrams into vertex corrections. An example of a set of self-consistent equations defining a vertex correction is shown in the lower panel of Fig. 3. This particular vertex function, which we call the $K$ vertex, sums both non-crossing intra-branch and inter-branch diagrams. It gives all such contributions starting from some initial condition (marked by the left index, 0 here) at the beginning of the contour and terminating at some final condition (the right index) at a pair of propagation times, one of which is on either branch. The rest of the contour (shown in gray) can then be filled by any bold diagram not containing contributions already summed within the vertex.

The two equations shown are part of a set of four coupled integral equations which must be solved self-consistently, as they also depend on $K_{02}$ and $K_{03}$. The final two equations have an analogous structure. There are four such sets, one for each value of the initial condition (see also Ref. [50] and Ref. [52] which used a notation that compacts all $16 K$-vertex equations into one equation and also includes the OCA contribution). Since the 16 elements of the vertex are two time objects, they are stored in discretized form in large 2D matrices. Currently a uniform discretization is used and the memory requirements for the partial summation part of the method is a bottleneck of the method. All integrals within the NCA vertices and all but a few within the OCA take the form of a convolution (for a Hamiltonian with no explicit time dependence), and can be performed efficiently in Fourier space. The remaining integrals are amenable to parallelization on clusters, and current implementations scale linearly to over a thousand nodes on high-performance clusters.

Two important examples of observables which will be presented in the results below are the electronic current into the different leads and the two-time correlation function. Both observables are given by diagrams which have a special line at the end of the Keldysh contour, shown as the light gray line on the upper right corner of Fig. 4. This line is created by an operator $d^\dagger_\sigma$ at the final measured time. To measure the current with spin $\sigma$ out of certain leads, the hybridization function for the special line should be replaced by the one defined by Eq. (37) with the sum over $k$ restricted to values within the chosen subset of leads (for instance, to measure the “left” current one only sums over $k$ values in the left lead). To measure two-time correlation functions, the value of the hybridization line becomes unity (as it describes free operators $d^\dagger(t)$ and $d(t')$ which do not come with a corresponding lead operator and coupling), and the contribution is binned with respect to the location of the operator at the other end of the line. With combinations of such lines one can construct any single particle correlation function ($G^<, G^>$ etc.).

As mentioned earlier, in practice all possible contractions of hybridization lines from a diagram are taken into account simultaneously: this means that the final expression for a contribution to (say) the current from any given diagram is identical to the one introduced in Eq. (62-66) of Ref. [52] but with terms already accounted for by the partial resummation excluded from the evaluation of the determinant of Eq. (59) of that reference.

In using bold-line CTQMC within the context of reduced dynamics for dot observables [51] the somewhat more complex $\Phi$ observables were implemented which measure the current through one spin for a preselected occupation of the other spin. It should be clear from the description above how the generalization proceeds: one accepts only current diagrams where the opposite spin meets the desired criterion. In fact, since both possible criteria are needed for the reduced dynamics, all contributions to the single-spin current can be sampled in each Monte Carlo trajectory, as long as during the measurement only the appropriate observable for each configuration is updated. A reduced dynamics treatment which also allows for the computation of non-dot observables
such as the current would require a further addition of observables to the simulation.\cite{footnote}

VI. RESULTS

We now show results for the spectral function of the Anderson impurity model, obtained using bold-line CTQMC and the auxiliary current method and a direct measurement of the time-dependent Green’s function. To constrain the parameter space we restrict ourselves to the particle-hole symmetric case $\epsilon_i = -\frac{\xi}{2}$ and to a single flat band with a soft cutoff $\Gamma(\omega) = \frac{\Gamma}{(1+e^{\frac{|\omega|-\mu}{\beta}})(1+e^{\frac{|\omega|-\mu}{\beta}})}$. We choose $\Gamma \nu = 10$ and hold the lead chemical potential at 0, while varying dot interaction $U$, the lead temperature $\beta$, and the band cutoff energy $\Omega_c$. All results are computed with the real time bold-line CTQMC expansion \cite{aoki2012quantum,aoki2014current} built around the one-crossing approximation (OCA) \cite{schrieffer2011theory} by measuring the current to an auxiliary reservoir defined by

$$\Gamma_A(\omega_A,\omega) = \frac{\eta \beta_A}{\sqrt{\pi}} e^{-[\beta_A(\omega-\omega_A)]^2},$$

(41)

with $\Gamma \beta_A = 10$ and $\eta = 10^{-3} \Gamma$. The dot is initially decoupled from the physical and auxiliary leads, and the coupling is turned on at time zero, after which the system time-evolves according to the full Hamiltonian, Eq. (1).

Little is known about the properties of the spectral function as measured from the auxiliary current, Eq. (21), as a function of time; what we have shown rigorously is only that at steady state or equilibrium it must approach the spectral function known from other methods. At intermediate timescales this quantity contains information about quench dynamics (since the dot begins in an out-of-equilibrium state where it is decoupled from the bath), but the interpretation of this data must be performed with care, since the physical dynamics are mixed to some degree with those of the auxiliary lead. We therefore begin the discussion by exploring it in detail for the three examples shown in Fig. 5. The $A(\omega)$ as measured by our virtual probes is plotted as a function of time. Note that a constant-$t$ cut across the surface at long times forms the steady state spectral function, which can be read from the profile of the plots. With the initial ($t = 0$) condition we have chosen, the auxiliary current $I_A(t = 0)$ is always zero. In all three cases, the band cutoff energy of the leads $\Omega_c$ is set to 10$\Gamma$; the band can therefore be considered essentially flat over the range of frequencies displayed. The long time limit corresponds to the equilibrium situation.

On the left, we display a noninteracting case. Within hybridization expansion CTQMC, the noninteracting case is a stringent test for the algorithm, as it expands about the atomic limit making this exactly solvable limit a difficult case for our approach. First, the final profile has a Lorentzian shape, as expected from a noninteracting dot coupled to a flat lead. Second, the observed relaxation timescale appears to be related to $\Gamma$ rather than the auxiliary parameters $\eta$ and $\beta_A$. This is typical for charge-related properties in the system, and suggests that we are in fact measuring physical system properties and not properties related to our choice of auxiliary lead. This is expected, since the auxiliary lead is coupled to the system only weakly, so that probing the dot only involves its linear response characteristics.

The behavior at both short and long timescales can alternatively be visualized by taking cuts through this data at planes of fixed $\omega$, as shown in the top panel of Fig. 6. Here, we see that relaxation to equilibrium occurs somewhat more slowly for energies near the Fermi level. At short timescales the data contains information about the nonequilibrium evolution of the system in a quench situation, but this is mixed to some degree with the relaxation properties of the auxiliary lead. Exploring the implications of this in detail is beyond the scope of the current work.

In the middle panel of Fig. 5 a strong interaction has been turned on, while the temperature is kept rather high. At long times, excitations spread throughout the band, but not far beyond $\Omega_c$. In addition, noise and oscillations in both time and frequency appear, indicating that convergence to the steady state is much slower. However, as before, the time evolution of the spectral function is mostly converged after a time on the order of $1/\Gamma$ (see also the corresponding middle panel of Fig. 6).

On the right side of Fig. 5 the temperature is lowered while the interaction is reduced. The formula $k_B T_K = U \sqrt{\frac{\nu}{2\pi} e^{-\frac{\pi U}{2\beta}}} \pi e$ is commonly used to estimate the Kondo temperature \cite{hewson1997kondo} and gives $\Gamma \beta_K \approx 4.7$, yet even at $\Gamma \beta_K \approx 3$ the predicted spectrum clearly begins to display the characteristics of Kondo physics, and one can observe the formation of a central peak at the chemical potential and an indication of two side bands. Interestingly, the time scale over which the general profile develops does not appear to be significantly modified, but long-time oscillations which have not fully attenuated by our final simulation time appear. This suggests that obtaining numerical convergence for this particular parameter set requires longer propagation times than required in the strongly interacting regime. On the other hand, it also suggests that obtaining a basic estimate of the spectral function only requires comparatively short propagation times (see also the corresponding lower panel of Fig. 6).

As explained in Section V, within bold-CTQMC two-time correlation functions can also be obtained directly for a given $t$ and $t'$. The result of one such evaluation for $G'_{\omega}(t,t')$ is displayed in Fig. 7. At equilibrium $G'_{\omega}$ must become a function of the difference $t-t'$ between its two time parameters, but for any finite $t$ it exists only for $t'<t$. We expect that for large enough $t$, the correlation function measured as a function of $t-t'$ should converge to the equilibrium value; Fig. 7 illustrates that while this clearly is not the case for $\Gamma t < 2$, convergence occurs rather quickly. For the parameters shown, at $\Gamma t \gtrsim 9$...
Figure 5. The dependence of the probed Spectral function $A(\omega) \equiv -\frac{1}{\pi} \Im \{G_R(\omega)\}$ on time and frequency, as obtained from the auxiliary current method for an initially decoupled quantum dot at $\Omega_c = 10\Gamma$. At $U = 0$ and $\beta\Gamma = 1$ (left), a simple Lorentzian shape develops. At $U = 6\Gamma$ and $\beta\Gamma = 1$ (center), the interaction distorts and widens the spectrum. At $U = 6\Gamma$ and $\beta\Gamma = 3$ (right), the spectral profile typical to the Kondo problem develops, exhibiting a central Kondo peak between two lower Hubbard peaks.

Figure 6. Several cuts through the data of Fig. 5 are shown at fixed $\omega$, making it easier to observe both the convergence and the behavior at short times.

4 one obtains reliable estimates of two-time correlation functions in equilibrium for any $t - t' < t$.

Once the equilibrium correlation function $G^r(t-t')$ is obtained in the time domain, a simple Fourier transform takes us to the frequency domain, where the spectral function may be obtained using Eq. 8. In practice, since $t - t'$ is limited in range by the maximum $t$ reachable, one must also converge the result of the Fourier transform in $t$ at all frequencies. This proves to be difficult, and we will show that when one is interested in frequency domain properties the auxiliary current method offers more accurate results. However, for certain nonequilibrium problems, it is often the time-domain function itself that proves to be of interest.

The dependence of the real part of the retarded equilibrium Green’s function on the physical parameters of the system is explored in Fig. 8. The top panel shows that while the effect of weak interaction is relatively mild, strong interaction induces qualitative changes in the structure of the correlations. The central panel shows that these effects are mitigated but not destroyed by higher temperatures. Finally, the bottom panel shows how a reduction in the bandwidth of the bath results in the introduction of high-frequency oscillations into the correlation function.

To validate our results, Fig. 9 compares the spectral function, as obtained with real time bold-line CTQMC from both the auxiliary current method and the direct Fourier transformation of the time-domain correlation function, to the exact spectrum in the noninteracting case, shown as the gray shaded region terminated by a dashed black line. We note that this is a particularly difficult limit for bold-line CTQMC based on the OCA, as OCA performs poorly for small values of $U$; to decrease the amount of computer time needed, we have limited the maximum order of bold diagrams to 6, despite the fact that higher order contributions are contributing on the order of a few percent. As a basis for assessing the
Figure 8. The real part of the retarded two-time correlation function \( G^r(t-t') \) as a function of the time difference \( t' - t \), for a final time \( t = 10 / \Gamma \). The dependency on different parameters is illustrated by taking different values of \( U \) (top panel), \( \beta \) (middle panel) and \( \Omega_C \) (lower panel) with the other parameters fixed.

Fig. 9 clearly shows that none of the numerical methods perfectly reconstruct the exact spectral function. The real time auxiliary current method (blue line with error bars denoting statistical, but not finite time or order truncation errors) is the only one to perform consistently well at all parameters and frequencies shown. However, some systematic errors remain, in particular at the band edges and near the chemical potential. The Fourier-transformed real time correlation function (green circles) is not as accurate, and in particular displays a high-frequency tail caused by numerical noise (the auxiliary lead formulation circumvents this by the finite width of the auxiliary lead’s coupling density). However, this always induces an averaging over a finite frequency window.

The MaxEnt result computed with a flat model (solid red line) in Fig. 9 appears to behave relatively well only at low temperature and large bandwidth, as seen in the bottom right panel. As we raise the temperature—going from the bottom right to the bottom left panel—MaxEnt fails, in particular at frequencies near the resonance. The peak is at odds with the bias towards flatness present in our default model, and causes a noticeable deviation from the correct result as we go from bottom right to top right panel. While the MaxEnt results are incompatible with the real time results, the back-continuation of the real-time results to the imaginary axis (where imaginary time Monte Carlo results are numerically exact) agrees perfectly with the imaginary time results, and the back-continuation of the continued MaxEnt results also agrees perfectly with the imaginary time results. The fact that two completely incompatible real-frequency spectra can have the same Matsubara frequency representation (within Monte Carlo errors on the imaginary axis) is well known and caused by the large number of very small eigenvalues of the analytic continuation kernel. Continuing further to the top left panel, when both temperature is low and bandwidth is high, MaxEnt results are unreliable both at high and at low frequencies (while, again, imaginary frequency Green’s functions agree within errors). As seen in the inset, no signature of high energy features, e.g. the band edge, is discernible. This also remains true at lower temperatures.

A few caveats regarding the comparison with MaxEnt are in order. First, analytic continuation is an ill-posed problem, and the analytically continued result depends strongly on the algorithm and parameters used. In the case of MaxEnt, it also depends on the model. As the algorithm is based on a penalty for deviations from the default model, a correct guess for the noninteracting system would have lead to the exact result. We therefore stress here that additional information (e.g. from low-order perturbation theory or high-temperature simulations) may yield substantially better results by taking advantage of additional knowledge; it may however also
be a source of bias. In the context of real time method, the ill-posed continuation problem can be avoided and the sources of ambiguity introduced when continuing results to the real axis are absent from the start.

We now turn to the exploration of the results of the direct measurement of the spectral function and of the auxiliary current method in combination with bold-line CTQMC in interacting systems. In Fig. 10 we increase the interaction strength to \( U = 2\Gamma \). The two real time methods are in far better agreement in this case, but deviations at high frequency are visible. Once again, only the real time auxiliary current correctly captures the existence of a band edge, and the MaxEnt method is unable to capture the results both for low band width and high temperature.

The nonzero interaction makes convergence with the order of bold diagrams much easier to obtain, as the quality of the OCA approximation improves and the number of hybridization events decreases. Still, these weakly interacting parameters remain beyond the scope where NCA or OCA are accurate. In fact, even a perturbative weak coupling treatment would be sufficient in this regime. The main result of this plot is therefore that it highlights the ability of OCA based bold-line CTQMC to treat interacting problems substantially beyond the range where OCA itself—without Monte Carlo—is accurate.

Next, we continue to a strongly interacting case, Fig. 11 with \( U = 6\Gamma \). Here \( \frac{U}{\Gamma} \) is large enough that NCA can be a useful approximation, and we include it for comparison. As before, the two real time methods give consistent results at small to intermediate frequencies. At large bandwidth and low temperatures (lower right), all Monte Carlo methods, including imaginary time CTQMC, yield similar results except at high frequencies, while the NCA underestimates the height of the central resonance. However, when either the bandwidth or the temperature is varied, a dramatic change occurs: lowering the bandwidth (upper right) results in the development of a high and sharp band edge, which the MaxEnt method misses completely. In this respect, MaxEnt actually performs even worse than NCA (though results are clearly much better at low frequencies). At higher temperature (lower left of Fig. 11), both MaxEnt and NCA underestimate the central resonance, resulting in a dip where a peak should be and in a qualitatively different spectral shape at low frequencies. When both temperature and bandwidth are varied simultaneously, NCA continues to suffer from the same problem, while MaxEnt suffers from both simultaneously, resulting in a qualitatively incorrect spectral function at all frequencies.

We note that previous authors have found that at least for some parameters, a perturbative expansion including select diagrams beyond the OCA results in substantially better approximations [8]. Using these higher order expansions as a starting point within bold-line CTQMC is expected to improve its convergence properties further, making it potentially easier to obtain reliable, unbiased data independent of the underlying approximation.
VII. SUMMARY AND CONCLUSIONS

To conclude, we have proposed and implemented two methods of obtaining Green’s functions from real-time bold-line CTQMC. The first one directly evaluates two-time correlation functions, while the second one is based on a newly proposed double-probe scheme for the computation of an auxiliary current. The auxiliary current method is general enough to be used with other numerical methods, and is especially helpful when two-time correlation functions are impossible or more expensive to acquire, as is the case in formalisms relying on reduced dynamics. We compared these two methods to each other, to analytically continued data and to data obtained within the non-crossing approximation. In general the real time methods give more accurate results and the auxiliary current method in particular even resolves high-energy features such as hard band edges. Numerically exact convergence accounting for all sources of errors remains technically difficult to achieve. The MaxEnt method (in the particular variation we chose to test) does not reconstruct high-frequency features. At low temperature it performs very well at low frequencies, but at higher temperatures it fails. We demonstrate that for certain choices of model parameters, which are neither special nor extreme, both MaxEnt and NCA produce qualitatively misleading results.

Real time methods remain substantially more expensive than imaginary time methods in terms of the computer time needed to accomplish similar tasks, and will in all probability remain so. As DMFT continues to evolve towards being a general-purpose tool for material science, it is plausible to assume that better scaling methods capable of treating larger impurities with more orbitals per site will become standard, even for real-time dynamics. While in this paper we have considered only expansions about the NCA and OCA for a single-site Anderson impurity model, the ideas and formalism are general: they can also be used for other formal resummations, or for expansions about more complicated models such as those arising in cluster extensions of dynamical mean field theory. Even now, real time methods offer (aside from the ability to address nonequilibrium situations, which was discussed elsewhere\cite{53} an alternative and controlled route to access spectral quantities at high frequency that is not available with other tools, and will find its use in particular in real-time DMFT.

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