Optimal ramp shapes for the fermionic Hubbard model in infinite dimensions

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We use non-equilibrium dynamical mean field theory and a real-time diagrammatic impurity solver to study the heating associated with time-dependent changes of the interaction in a fermionic Hubbard model. Optimal ramp shapes $U(t)$ which minimize the excitation energy are determined for a noninteracting initial state and an infinitesimal change of the interaction strength. For ramp times of a few inverse hoppings, these optimal $U(t)$ are strongly oscillating with a frequency determined by the bandwidth. We show that the scaled versions of the optimized ramps yield substantially lower temperatures than linear ramps even for final interaction values comparable to the bandwidth. The relaxation of the system after the ramp and its dependence on the ramp shape are also addressed.

I. INTRODUCTION

Correlated electron systems play a central role in modern condensed matter physics. While most theoretical studies have focussed on the equilibrium properties of these systems and many interesting phenomena remain to be understood, there is a growing effort to also address the non-equilibrium dynamics. These investigations are motivated by ongoing experimental progress in this field, in particular measurements of the relaxation dynamics in correlated materials by means of pump-probe spectroscopy and experiments on cold-atom systems. The latter provide an experimental realization of theoretical models such as the fermionic one-band Hubbard model and allow to explore the time-evolution after rapid changes in model parameters.

A computationally tractable method to simulate the time evolution of a correlated lattice model after a quench or external perturbation is nonequilibrium dynamical mean field theory. This approach neglects spatial correlations, and local correlation functions and their dynamics are obtained from an appropriately defined quantum impurity model. The repeated numerical solution of this impurity model on the real-time (Keldysh) contour is the most challenging aspect of nonequilibrium DMFT calculations, but several perturbative and exact impurity solvers have recently been developed to tackle this problem. Nonequilibrium dynamical mean field theory has been applied to interaction quenches in the Hubbard model as well as in the Falicov-Kimball model and to study the time evolution in the presence of external fields. Interesting phenomena have been observed in these theoretical investigations, such as an apparent dynamical phase transition in the relaxation dynamics after an interaction quench.

A change in the interaction parameter affects the total energy and therefore the temperature of the system after equilibration, and nonadiabatic heating effects can be substantial. The dynamical phase transition after a sudden interaction quench in the $T = 0$, noninteracting Hubbard model for example, occurs at an energy which translates into a temperature of about 0.2 times the bandwidth, which is far higher than the temperatures for which a metal-insulator transition is found in equilibrium. While it is an interesting observation that an apparently sharp dynamical transition occurs in such a highly excited system, one would like to avoid the heating effect in other contexts. In cold-atom systems, where much effort is devoted to realize a fermionic Hubbard model in the low-temperature regime, interactions and the optical lattice must be switched on in such a way that the heating is minimized. If there are constraints on the ramping time, it can become a subtle question whether an equilibrium state in a desired low-temperature phase can be reached at all by means of a ramp which starts from another, more easily preparable state of the system. And since in most cases one will be interested in preparing an equilibrium state of the interacting system, a second relevant question concerns the time-scale on which the system relaxes and its dependence on the ramp shape. Another example is the recently proposed conversion of repulsive interactions into attractive ones by means of external periodic electric fields (also essentially an interaction quench), which raises the interesting possibility of AC-field induced superconductivity. Also in this context the heating and the thermalization time associated with the ramp protocol are important issues.

The purpose of this theoretical study is to determine the optimal ramping procedure between two different parameter regimes of the Hubbard model, where “optimal” means the passage in which the system is least excited. In particular when the ramping time is restricted to only a few times the inverse hopping, one can expect a strong dependence of the excitation energy and of the relaxation dynamics on the ramping protocol. However, an unbiased optimization over the infinite space of ramp shapes is not possible, because even the computation of the excitation energy for a single ramp protocol requires considerable numerical effort. For “small ramps”, on the other hand, in which the parameter is changed by only a small amount, one can resort to perturbation theory in the ramp amplitude to disentangle the influence of the ramp shape and many-body effects on the excitation energy. This allows us to perform an efficient optimization over a wider space of ramp shapes. In the present work we demonstrate that ramp shapes which are optimized for such infinitesimal parameter changes yield a considerably lower excitation energy than the generic linear ramp or other simple ramping protocols, even when applied to ramps with an amplitude comparable to the...
We compute the real-time evolution of the impurity model within the DMFT selfconsistent loop using a continuous-time Quantum Monte Carlo (CT-QMC) method which is based on an expansion of the partition function in powers of the interaction term. This method is free of systematic errors but restricted to relatively short times due to a dynamical sign problem. Nevertheless, it has proven useful for interaction quench calculations in the Hubbard model and for the simulation of transport through quantum dots. Here, we briefly recapitulate the weak-coupling formalism to show that it can account for time-dependent interactions, simply by replacing weight factors associated with individual vertices by time-dependent factors. The remaining part of the DMFT self-consistency, i.e., the computation of lattice observables and lattice Green functions from the local Green function of the impurity system, is unchanged with respect to the interaction quench setup. A detailed description of these equations can be found in Ref.\textsuperscript{13}.

We start by writing the partition function $Z = Tr[e^{-\beta H}]$ as

$$Z = Tr\left[e^{-\beta H_1}Te^{-\int_0^\tau dt H_2}\right]$$

where $\int_c$ denotes the integral along the Keldysh-contour $0 \rightarrow -i\beta \rightarrow t_{\text{max}} \rightarrow 0 \rightarrow -i\beta$. In the simulations, we choose the length of the contour $t_{\text{max}}$ somewhat larger than the ramp time $\tau$. The operators $H_1$ and $H_2$ correspond to the hopping and interaction part of the impurity Hamiltonian. Specifically, we use

$$H_2 = H_U - k(t)/t_{\text{max}},$$

with

$$H_U = U(t)(n_\uparrow n_\downarrow -(n_\uparrow + n_\downarrow)/2).$$

The nonzero real function $k(t)$ was introduced to enable an auxiliary field decomposition of the interaction term. Expansion of the contour-ordered exponential in powers of $H_U - k(t)/t_{\text{max}}$ and the application of the decoupling formula\textsuperscript{22}

$$1 - t_{\text{max}}H_U/k(t) = \frac{1}{2} \sum_{s=-1,1} e^{\gamma(t)s(n_\uparrow - n_\downarrow)},$$

$$\cosh(\gamma(t)) = 1 + \frac{t_{\text{max}}U(t)}{2k(t)}$$

at every interaction vertex results in an expression of the partition function as a sum over all possible collections of Ising spin configurations on the contour with weight

$$\omega\left\{(t_1, s_1), (t_2, s_2), \ldots, (t_n, s_n)\right\} = (-i^{n_-})(i^{n_+})(k(t)dt/(2t_{\text{max}}))^{n_-+n_+} \prod_{\sigma} \det N^{-1}_{\sigma}.$$  

Here, $n_+$ and $n_-$ are the number of spins on the forward and backward branch of the contour, and we have used the fact that in a noninteracting initial state there are no spins on the...
imaginary-time branch. The matrices $N^{-1}_{\sigma}$ are determined by
the location of the interaction vertices on the contour $C$ and are
given by
\[
N^{-1}_{\sigma} = e^{S_{\sigma}} - (iG_{0,\sigma})(e^{S_{\sigma}} - I),
\]
with $G_{0,\sigma}$ the bath Green’s function and $e^{S_{\sigma}} = \text{diag}(e^{\gamma(t_1)\sigma}, \ldots, e^{\gamma(t_n)\sigma})$. In the actual calculations, we
choose $\gamma(t) = \gamma$ time-independent, so that the time dependence of the interaction manifests itself only in a time dependence of the parameter $k(t) = \frac{1}{2}t_{\text{max}}U(t)/(\cosh(\gamma) - 1)$.

The sampling procedure consists of generating all spin configurations on the contour $C$ through random insertions and removals of spins. During the sampling, we measure the quantity
\[
X_{\sigma}(s_1, s_2) = \left\langle i \sum_{i,j=1}^{n} \delta_C(s_1, t_i)[(e^{S_{\sigma}} - 1)N_{\sigma}]_{i,j} \delta_C(s_2, t_j) \right\rangle_{MC}.
\]
The impurity Green’s function is then obtained as
\[
G_{\sigma}(t_1, t_2) = G_{0,\sigma}(t_1, t_2) + \int_{C} ds_1 \int_{C} ds_2 G_{0,\sigma}(t_1, s_1)X_{\sigma}(s_1, s_2)G_{0,\sigma}(s_2, t_2).
\]

III. PERTURBATIVE ANALYSIS FOR SMALL RAMPS

A. General considerations

In this section we briefly recall the perturbative analysis of Ref. 24 for an infinitesimal interaction ramp. To be specific, we consider a ramp of the form
\[
U(t) = U r(t/\tau),
\]
where $r(x)$ is the ramp-shape function which satisfies $r(x) = 0$ for $x \leq 0$ and $r(x) = 1$ for $x \geq 1$. The excitation energy per lattice site is defined as
\[
\Delta E(\tau) = E(\tau) - E_0(\tau),
\]
where $E_0(\tau)$ is the groundstate energy per lattice site of the model with interaction $U$, and $E(\tau) = \langle H(t) \rangle$ is the energy expectation value of the system with time-dependent interaction. After expanding in powers of $U$, the second-order result for the excitation energy can be decomposed into contributions from a ramp spectrum $F(x)$, which depends only on the ramp shape but not on the properties of the system, and from an excitation density $R(\omega)$, which depends on the system but not on the ramp shape.

\[
\Delta E(\tau) = U^2 \mathcal{E}(\tau) + O(U^3),
\]
\[
\mathcal{E}(\tau) = \int_{0}^{\infty} \frac{d\omega}{\omega} R(\omega) F(\omega \tau),
\]
\[
F(x) = \left[ \int_{0}^{1} ds r'(s) e^{i x s} \right]^2.
\]

In this expression, the excitation density is defined by the Lehmann representation,
\[
R(\omega) = \frac{1}{L} \sum_{n \neq 0} \langle \phi_n | W | \phi_0 \rangle^2 \delta(\omega - E_n + E_0),
\]
where $W = (n \uparrow - \frac{1}{2})(n \downarrow - \frac{1}{2})$ is the operator which couples to the time-dependent parameter $U(t)$ in the Hamiltonian, $L$ is number of lattice sites, and $|\phi_n\rangle$ and $E_n$ are the eigenfunctions and eigenvalues of the Hamiltonian. The function $R(\omega)$ can be evaluated easily for the Hubbard model with $U = 0$, leading to 24
\[
R(\omega) = \int_{-\omega}^{\omega} d\epsilon \rho(\epsilon) \int_{0}^{\mu+\omega} d\mu \rho(\mu - \omega - \epsilon) \int_{0}^{\mu} d\nu \rho(\nu) \rho(\mu - \nu).
\]

In Fig. 2 we plot the function $R(\omega)/\omega$ for the semi-elliptical density of states together with ramp spectra for different ramp shapes. At $U = 0$, the only relevant energy scale is the hopping $V (= 1)$: $R(\omega)/\omega$ has a peak near $\omega \approx 3$ and vanishes for $\omega > 8 \approx 2 \cdot$ bandwidth. According to Eq. (14), a small overlap of the excitation density and ramp spectrum results...
in a small excitation energy. If the ramp spectrum $F(x)$ falls off rapidly at large $x$, the main weight of $F(\omega \tau)$ and the main contribution to the integral in Eq. (14) comes from frequencies $\omega \lesssim 1/\tau$. Using the asymptotic form $R(\omega) \propto \omega^3$ for small $\omega$ one can prove that the system approaches the adiabatic limit $\tau \to \infty$ with a power law behavior $\Delta E(\tau) \propto 1/\tau^3$, provided that the high-frequency tail of $F(x)$ falls off faster than $1/x^3$.

For the linear ramp $r(x) = x$, for which $F(x)$ falls off as $1/x^2$, one finds

$$\Delta E(\tau) \propto 1/\tau^2.$$  

In the present paper we are interested in the excitation energies for ramp times which are so short that the asymptotic power law behavior $\Delta E(\tau) \propto 1/\tau^n$ does not yet hold, and we attempt to minimize $\Delta E(\tau)$ in Eq. (14) with respect to the ramp shape $r(x)$. The reduction of the excitation energy can be understood as a consequence of a suppression of the ramp spectrum $F(\omega \tau)$ in the frequency range around the maximum of $R(\omega)/\omega$, which is achieved by suitably designing the ramp $r(x)$. Clearly, the optimal ramp shape will strongly depend on the ramp time $\tau$. For example, for the ramp $r(x) = x + 0.2 \sin(2\pi x)$ the function $F(\omega \tau)$ is suppressed around $\omega = 3$ for $\tau \approx 1.25$, while for $\tau = 2.5$ the same ramp shape is apparently not favorable (Fig. 2).

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**B. Minimization in a tent basis**

To find the optimal ramp-protocol $r(x)$ which minimizes the excitation energy $\mathcal{E}(\tau)$ given some constraints on $r(x)$, we discretize the interval $[0, 1]$ into an equidistant mesh of $N + 2$ points $x_k = k\Delta x (k = 0, \ldots, N + 1; \Delta x = 1/(N + 1))$, and consider ramp functions which linearly interpolate between the values $r(x_k) = x_k + c_k$. The numbers $c_k$ measure the deviations from the linear ramp $r(x) = x$, so $c_0 = c_{N+1} = 0$. Equivalently, this means that the function $r(x)$ is expanded in a basis of tent-shaped functions $\phi_k(x)$,

$$r(x) = x + \sum_{k=1}^{N} c_k \phi_k(x),$$  

with

$$\phi_k(x) = \begin{cases} \frac{x - x_k - 1}{x_k - x_{k-1}} & \text{if } x \in [x_{k-1}, x_k], \\ \frac{x - x_k + 1}{x_{k+1} - x_k} & \text{if } x \in [x_k, x_{k+1}], \\ 0 & \text{otherwise}. \end{cases}$$

The first derivative of the ramp function $r(x)$ is given by

$$r'(x) = 1 + \sum_{k=1}^{N} c_k \phi'_k(x)$$  

with

$$\phi'_k(x) = \begin{cases} \frac{1}{\Delta x} & \text{if } x \in [x_{k-1}, x_k], \\ \frac{1}{\Delta x} & \text{if } x \in [x_k, x_{k+1}], \\ 0 & \text{otherwise}. \end{cases}$$

The calculation of $F(\omega \tau)$ requires the evaluation of the integral

$$\int_0^1 r'(s)e^{i\omega \tau s}ds = \frac{1}{i\omega \tau} (e^{i\omega \tau} - 1) + \frac{2(1 - \cos(\omega \tau \Delta x))}{i\omega \tau \Delta x \sum_{k=1}^{N} c_k e^{i\omega \tau x_k}}.$$  

Multiplying the right hand side of Eq. (22) with its complex conjugate and remembering that $F(x)$ is real gives

$$\mathcal{E}(\tau) = \int_0^\infty d\omega \frac{R(\omega)}{\omega} F(\omega \tau) = \text{const} + c^T f + c^T M c,$$  

where the coefficients $c_k$ are written as an $N$-component vector $c$, and the $N$-component vector $f$ and the $N \times N$ matrix $M$ are given by

$$f_k = \int_0^\infty d\omega \frac{R(\omega)}{\omega^2} (1 - \cos(\omega \tau x_k)) \frac{1}{|\omega|^2 \Delta x} \times [(\cos(\omega \tau (1 - x_k)) - \cos(\omega \tau x_k)),$$

$$M_{kk'} = \int_0^\infty d\omega \frac{R(\omega)}{\omega^2} (1 - \cos(\omega \tau x_k))^2 \frac{1}{|\omega|^2 \Delta x^2 \cos(\omega \tau (x_k - x_{k'})).}$$

The minimization of this quadratic problem can be performed using standard techniques.

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**C. Unconstrained optimization and symmetries**

In order to get a qualitative understanding of the optimal ramp shapes we first note the following symmetry properties of $M$ and $f$:

1. $M$ is symmetric, i.e. $M_{i,j} = M_{j,i}$,
2. $M_{i,j}$ is constant on each subdiagonal $|i - j| = k$,
3. $f$ is antisymmetric, i.e. $f_i = -f_{N+1-i}$.

The optimal, unconstrained ramp shape can be found by solving $\partial \mathcal{E}(\tau; \{c_i\})/\partial c_i = 0$ and therefore, using the symmetry of $M$

$$2Mc + f = 0.$$  

It follows that the solution $c$ is antisymmetric, i.e. $c_i = -c_{N+1-i}$. In fact, setting $N = 2k$, the first $i = 1, \ldots, k$ components of Eq. (26) are

$$c_k = -\sum_{j=1}^{k} (M_{i,j} c_j + M_{i,2k+1-j} c_{2k+1-j}).$$  

Using the symmetry $f_i = -f_{2k+1-i}$, we get

$$0 = \sum_{j=1}^{k} (M_{i,j} + M_{2k+1-i,j}) c_j + (M_{i,2k+1-j} + M_{2k+1-i,2k+1-j}) c_{2k+1-j}.$$
The symmetry properties of $M$ thus imply

$$0 = \sum_{j=1}^{k} (M_{i,j} + M_{i,2k+1-j}) (c_j + c_{2k-1-j}),$$

which can only be satisfied for all $i = 1, \ldots, N$ if $c_j = -c_{2k-j+1}$.

IV. RESULTS

A. Small ramp amplitude

We first consider unconstrained paths and solve Eq. (26) for different values of $\tau$. As illustrated in Fig. 3, the optimal paths oscillate around the linear ramp, and the number of oscillations increases with increasing $\tau$. Thus, the somewhat unexpected result is that paths which minimize the excitation energy, at least according to the perturbative formula (14), may involve excursions to positive and negative interaction values which are much larger in absolute value than the final interaction $U$.

Note that the unconstrained optimization as described here becomes numerically unstable for large $N$. This is because one can always add to the ramp a highly oscillating component $\delta r(x)$ whose ramp spectrum $\delta F(x)$ lies almost completely outside the support of the excitation density $R(\omega)$, and thus does not influence the excitation energy. In other words, there are directions in the parameter space along which the excitation energy hardly changes. The corresponding eigenvalues of the matrix $M$ are almost zero, and the linear equation (26) becomes ill conditioned for large $N$.

However, optimal ramps with large amplitude oscillations and sign changes may be difficult to realize in experiments. We therefore also compute optimal ramps with the constraint $0 \leq r(x) \leq 1$, using the reflective Newton method implemented in MATLAB. The resulting paths for different $\tau$ are shown in Fig. 4. It is evident from these results that the optimal ramp shapes are characterized by a roughly constant oscillation frequency $\omega_0 = 2\pi n_{\text{osc}}/\tau \approx 9 - 10$. To gain some insight into the origin of these oscillations we plot in Fig. 5 the ramp spectra $F(\omega\tau)$ for the ramps shown in Fig. 4. The inset shows a close-up view of the optimal ramp spectra in the frequency range where $R(\omega)$ is large. We see that $F(\omega\tau)$ is optimized in such a way that the overlap with $R(\omega)$ is minimal. The main panel shows the ramp spectra in the range $0 \leq \omega \leq 14$. A large peak in $F(\omega\tau)$ is evident at $\omega \approx 9.5$ ($\tau = 1.25$), $11$ ($\tau = 2.25$), $9.75$ ($\tau = 3.25$) and $9$ ($\tau = 4.25$), i.e., just above the largest value of $\omega$ for which $R(\omega) > 0$.

We can understand from Eq. (13) that a sharp peak in $F(\omega\tau)$ at $\omega \approx \omega_0$ corresponds to an oscillating $r'(s) \sim \cos(\omega_0 \tau s)$. So, the frequency which appears in the optimal ramp shapes is determined by the support of the function $R(\omega)$, which itself is defined by the density of states $\rho(\omega)$. In the case of a symmetric $\rho(\omega)$ considered here, the support of $R(\omega)$ is twice the bandwidth (Eq. (14)). The optimal ramp shapes have therefore an oscillating component with an oscillation frequency roughly given by $\omega_0 \approx 2 \cdot \text{bandwidth}$.

B. Monte Carlo results for larger $U$

In this subsection, we use Monte Carlo simulations to compute the heating effect for ramps with amplitudes beyond the perturbative regime, and compare the heating produced by linear ramps to the heating produced by ramps which were optimized for infinitesimal ramp amplitudes.

First, we would like to confirm the validity of the perturbative analysis for ramps to small interaction. In Fig. 6 we plot the excitation energy computed by means of nonequilibrium DMFT for different ramps to $U = 1$. The red line with crosses, and the green line with stars show the result of Eq. (14) for linear ramps and ramps of the form $r(x) =$
larger values of \(m\) yield low excitation energies also for shapes in Fig. 4 (Fig. 2 and explain the nonmonotonic behavior of the excitation energy for an interacting ground state energy estimated to be \(E_0 = 0.6195\).

This value of \(E_0\) gives the best agreement between analytical and DMFT results, and is consistent within error bars with the ground state energy \(E_{0}^{\text{DMFT}}(U = 1) = 0.618(2)\) estimated from equilibrium DMFT calculations. Hence, for ramps to \(U = 1\) the formula (14) gives accurate excitation energies. The ramp spectra \(F(\omega \tau)\) for several values of \(\tau\) are plotted in Fig. 5 and explain the nonmonotonic behavior of the excitation energy in the case of \(r(x) = x + 0.2 \sin(2\pi x)\).

A nontrivial question is whether the ramp shapes optimized using Eq. (14) yield low excitation energies also for larger values of \(U\). In the following we will consider ramps to \(U = 3\). This value is close to the critical interaction strength \(U_c \approx 3.2\) for instantaneous quenches, where the system was found to thermalize within a time of less than two inverse hoppings. To quantify the heating effect we translate the excitation energy into an effective temperature (inset of Fig. 7), which is defined as the temperature for which an equilibrium system with interaction \(U\) has total energy \(E(\tau)\).

The effective temperature for linear ramps to \(U = 3\) is plotted as a function of ramp time in the main panel of Fig. 7 (black line with squares). We see that the temperature after the quench drops rapidly with \(\tau\) until about \(\tau \approx 1.5\) and then more slowly for longer ramp times. The times \(\tau\) which are accessible with Monte Carlo are not sufficient to determine the asymptotic behavior of the excitation energy \((\tau \rightarrow \infty)\), but it is expected to be \(\Delta E(\tau) \sim 1/\tau^2\) based on the perturbative analysis. Figure 8 shows the effective temperatures for \(\tau = 1.25\) obtained from the optimized unconstrained ramps for different numbers \(N\) of basis functions (see top panels of Fig. 5). The lowest temperature for \(\tau = 1.25\), \(N = 5\) and \(\tau = 2.25\), \(N = 7\), are indicated by the red and green crosses in Fig. 7. For short ramp times \((\tau = 1.25)\), the effective temperature can be reduced by about a factor of 2 (compared to linear ramps) if an oscillating ramp shape is used. For \(\tau = 2.25\), the reduction is only about 10%. The constrained optimized ramps yield comparable reductions in the effective temperature. In Fig. 7 the blue line with circles shows the effective temperature for \(N = 25\) basis functions and the constraint \(0 \leq r(x) \leq 1\).

C. Accuracy of the perturbative results

The results shown in Fig. 7 demonstrate that the perturbative analysis of Section III may be used to compute ramp shapes with considerably lower excitation energy than linear ramps. On the other hand, it is not yet clear how quantitatively accurate the estimated energies are in the case of ramps...
to intermediate or strong $U$. If the difference between the true and the estimated excitation energy is large and the error depends sensitively on the ramp shape, then the true optimal ramp might look very different from the shape obtained by our procedure. To test the reliability of our estimated excitation energy and its dependence on the ramp shape we consider the simple one-parameter families $r_1(x) = x + a \sin(2\pi x)$, and $r_2(x) = x + a \sin(4\pi x)$ with $\tau = 1.25$ and $U = 3$, and compare the optimal values for the parameter $a$ obtained from the perturbative analysis to the Monte Carlo results. The oscillation frequency of $r_1(x)$ is not compatible with the ramp time, while the oscillation frequency of $r_2(x)$ is identical to that found in the constrained optimized infinitesimal ramp (top left panel of Fig. 4) and thus results in lower excitation energies.

In Fig. 5 the blue line with squares plots the excitation energy obtained from Eq. (14) as a function of the shape parameter $a$. The red line with circles shows QMC results for the total energy which are shifted in such a way that the minima of the curves coincide. The necessary shifts $E_0 = -0.279$ (for $r_1(x)$), and $E_0 = -0.277$ (for $r_2(x)$) are still compatible to the ground state energy $E_0^{DMFT} = -0.284(3)$ obtained from a $T \to 0$ extrapolation of equilibrium DMFT energies for the $U = 3$ model, but the perturbative formula appears to underestimate the excitation energy of the system by $\delta E_0 \approx 0.005 - 0.007$. Still, the variation of the excitation energy as a function of the ramp-parameter $a$ is quite accurately reproduced. In particular, the perturbative analysis yields the correct values for the optimal parameters ($a_{\text{opt}} = 0.175$ for $r_1(x)$ and $a_{\text{opt}} = 0.87$ for $r_2(x)$), and thus the correct optimal ramp shape.

For interactions $U \gtrsim 3.5$, the analytical prediction for the excitation energy is no longer in quantitative agreement with the Monte Carlo results. In the lower panel of Fig. 5 we show Monte Carlo data for $U = 4$ and 5, with an arbitrary offset (chosen in such a way that the minima are at around the same energy). We see that as the interaction is increased, the minimum in the excitation energy shifts to smaller values of $a$, while the curvature at the minimum increases. This implies that the minimization of Eq. (14) yields ramp shapes which do not minimize the excitation energy, and which in the large-$U$ limit may be qualitatively wrong.

D. Relaxation after the ramp

We would finally like to address the relaxation dynamics after the ramp. If the goal of an experiment is to prepare an equilibrium state at given interaction strength, then it is not only the ramp time, but also the thermalization time, which determines the relevant time scale of this process. The relaxation dynamics after an (sudden) interaction quench has been studied in Refs. 12,13,14. These calculations demonstrated the existence of a “dynamical phase transition” at some critical interaction strength $U_{\text{quench}}^c \approx 3.2$, which separates two qualitatively different relaxation regimes. After a quench to $U_{\text{quench}}^c$, the system thermalizes within a few inverse hoppings, whereas away from this particular interaction value thermal-
The critical interaction can be identified for example by plotting the time evolution of the quantity\textsuperscript{12}
\begin{equation}
\Delta n(t) = n(\epsilon_k = 0, t) - n(\epsilon_k = 0, t),
\end{equation}
which is the size of the discontinuity of the momentum distribution function $n(\epsilon_k, t) = \langle c_{k,\sigma}^\dagger(t)c_{k,\sigma}(t) \rangle$ at the Fermi energy. After a quench to $U < U_c^{\text{quench}}$, the system initially settles into a nonthermal quasistationary state characterized by a nonzero $\Delta n(t)$. For $U > U_c^{\text{quench}}$, $\Delta n(t)$ exhibits collapse-and-revival oscillations. At $U \approx U_c^{\text{quench}}$, $\Delta n(t)$ rapidly vanishes, with no sign of trapping in an intermediate nonthermal state.

A very similar relaxation dynamics can be observed after an interaction ramp with relatively short ramp time $\tau$. In Fig. 10 we plot $\Delta n(t)$ for different values of the interaction. The top panel shows results for linear ramps with ramp time $\tau = 1.25$, and the lower panel for ramps of the form $r(x) = x + 0.87 \sin(4\pi x)$ with ramp time $\tau = 1.25$. The critical interaction strengths (just before the onset of the collapse-and-revival oscillations) are $U_c^{\text{linear}} \approx 3.75$ and $U_c^{\text{oscillating}} \approx 4.25$. Apparently, the critical interaction strength for given ramp time $\tau$ depends on the ramp shape and this degree of freedom can be used to design protocols for which the desired interaction strength corresponds to a dynamical phase transition point.

The green curves with triangles and the insets of Fig. 10 demonstrate that also in the ramp case, the critical interaction strength is associated with fast thermalization. At the critical point, $\Delta n(t)$ vanishes within a few inverse hoppings, both for linear and oscillating ramps. In the insets we show a comparison of $n(\epsilon, t = 3.125)$ with an equilibrium distribution function for the interacting system (temperature $T = 0.547$ for the linear ramp and $T = 0.551$ for the oscillating ramp). The temperatures of the thermalized systems have been computed by comparing the total energy after the ramp to the temperature-dependent total energy obtained from equilibrium DMFT simulations. The fast thermalization at the dynamical transition and the ramp shape dependence of the critical interaction strength imply that it is possible to prepare thermal equilibrium states over a range of interaction values by suitably designing the ramp protocol.

\section{V. Conclusion}

Using nonequilibrium DMFT, we investigated the excitation energy of the one-band Hubbard model after ramping up the interaction within a time $\tau$ of the order of a few inverse hoppings. Based on the perturbative analysis of Ref. \textsuperscript{24} we determined optimal ramp protocols within a general set of piecewise linear ramp functions. This analysis indicates that the optimal ramp shape is oscillating around the linear ramp $U(t)/U(\tau) = t/\tau$, with a period which is determined by the support of the excitation density $R(\omega)$. For small values of $U$, the latter depends mainly on the density of states. Applying these optimized ramp shapes to larger interactions ($U = 3$ for a model with bandwidth $4$) yields considerable reductions in the effective temperature, compared to linear ramps. For short ramp times $\tau \lesssim 1.5$, both unconstrained paths and paths constrained to the interaction range $[0, U]$ result in effective temperatures which are 30\% to 50\% lower than those obtained with linear ramps.

Up to intermediate interaction values, the optimal ramps determined from the perturbative approach coincide well with those determined from the QMC calculations. Our analysis thus suggest that one may use the perturbative formula to determine optimal ramps when an optimization using QMC simulations is not possible. This is true in particular for slow ramps with ramp times up to 100 inverse hoppings, which are used for the preparation of states in cold atom systems. Guided by the optimal ramp shape at short times (oscillations...
superimposed on a linear ramp), one can, e.g., try to improve on the linear ramp by optimizing the amplitude \( a \) in a ramp of the form \( U(t)/U(\tau) = t/\tau + a \sin(\omega t) \). Using the “optimal” frequency \( \omega \) from Fig. 9, for example, the perturbative approach predicts that an optimization of \( a \) results in a decrease of the excitation energy of more than 30% with respect to the linear ramp, even in the limit of large \( \tau \), where the absolute value of the excitation energy becomes small (\( \sim 1/\tau^2 \)).

The precise shape (oscillation frequency) of the optimal ramps found in this study is specific to the \( U = 0 \) initial state. For ramps within the insulating phase, the ramp spectrum will be determined by the interaction energy \( U \) rather than the bandwidth, and the optimal ramp shapes will be different. However, in view of our results a perturbative analysis should still provide a numerically efficient way to compute ramp shapes which result in low excitation energies.

In addition to minimizing the excitation energy, we have briefly addressed the question of thermalization after the ramp. We have demonstrated that the thermalization after a ramp strongly depends on the final interaction and the ramping parameters. Our results indicate the existence of a dynamical transition (associated with rapid thermalization), similar to the behavior after an interaction quench. The ramp shape dependence of this dynamical transition point may be exploited to design ramp protocols which yield a thermal equilibrium state within a time of a few inverse hoppings after the switch-on of the interaction.

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