Real-time evolution of Anderson impurity models via tensor network influence functionals

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In this work, we present and analyze two tensor network-based influence functional approaches for simulating the real-time dynamics of quantum impurity models such as the Anderson model. Via comparison with recent numerically exact simulations, we show that such methods accurately capture the long-time nonequilibrium quench dynamics. The two parameters that must be controlled in these tensor network influence functional approaches are a time discretization (Trotter) error and a bond dimension (tensor network truncation) error. We show that the actual numerical uncertainties are controlled by an intricate interplay of these two approximations, which we demonstrate in different regimes. Our work opens the door to using these tensor network influence functional methods as general impurity solvers.

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I. INTRODUCTION

The numerical simulation of quantum many-body systems is, in principle, an exponentially difficult problem due to the growth of Hilbert space dimension with system size. While the advent of tensor network techniques has made the equilibrium problem more tractable, the application of tensor network methods to dynamics has been limited by the growth of entanglement during time evolution, which can cause exponentially growing resource requirements to accurately describe the dynamics. Given the complex nature of this problem, developing new numerical methods and furthering the understanding of dynamical phenomena is best done within the context of simple, yet nontrivial models.

Quantum impurity models offer one such possibility, with the additional benefit that they are of practical physical importance, having led to an understanding of phenomena ranging from the Kondo effect in solids [1] to the survival of macroscopic quantum coherence effects in condensed phases [2,3]. In the past few decades, impurity models have gained additional significance as a key component of computational embedding frameworks such as dynamical mean-field theory [4] and density matrix embedding theory [5,6]. Here, one must compute the real-time dynamics of the impurity to obtain an approximation to the local dynamics of the full problem via a self-consistently refined effective bath. Thus the efficacy of such embedding methods is constrained by the flexibility, speed, and accuracy with which the dynamics of an impurity coupled to a bath can be simulated.

Numerical approaches for solving impurity problems must contend with issues around the treatment of a large (infinite) number of bath degrees of freedom, while simultaneously needing to ameliorate the dynamical sign problem for impurity dynamics. The issue of treating continuous baths poses an impediment to methods such as diagonalization-based techniques [7–10] and is partially solved by the advent of tensor network approaches [11–16]. The treatment of the sign problem saddles dynamical Monte Carlo methods [17–23] with an exponentially scaling numerical cost. Although this scaling can be tamed by inchworm diagrammatic expansions [24–27], these approaches still incur statistical errors. Methods based on the Feynman-Vernon influence functional (IF), such as iterative path integral methods [28–30] and the hierarchical equations of motion (HEOM) approach [31,32], have been used with great success in spin-boson-type problems, but are far less explored in problems with fermionic baths such as the Anderson model. [33] Generally speaking, treating the problem of impurity dynamics using the IF provides a natural formalism to consider continuous baths without invoking stochastic sampling which introduces dynamical sign problems.

Of the above approaches, discretized path integral-based IF methods, e.g., the quasi-adiabatic path integral method (QUAPI) [28,29,34,35], are promising, as they reduce the problem to one of managing Trotter errors and memory truncation. However, models such as the Anderson model are more complex than the spin-boson model, in part because the system part of the system-bath coupling cannot be written in a simple diagonal form. Early attempts to generalize these approaches to the dynamics of Anderson-like models provided a window to obtain exact nonequilibrium dynamics in some parameter regimes, but suffered from memory length issues that limited their range of applicability [36,37]. In the intervening years, significant progress has been made marrying tensor network methods and IF methods, particularly for spin-boson-like models. Specifically, one can view
the Trotterized dynamics as a tensor network and employ tensor contraction over the environment degrees of freedom for a fixed propagation time, instead of contracting along the temporal direction [38–44]. The resulting object can be viewed as a matrix product state representation of the exact discretized influence functional (MPS-IF), which exists on a temporal lattice. Such an approach has been successfully applied to harmonic baths [41,42,45–49] as well as finite baths of bosons, fermions, or spins [41,50].

Recently, Abanin and co-workers have focused on the behavior of MPS-IF approaches for fermionic models, and have usefully detailed how entanglement properties of the influence functional temporally evolve [43,51,52]. In this paper, we follow Abanin et al., considering the explicit nonequilibrium dynamics of the Anderson impurity model directly in the continuous fermionic bath limit. We use a more general and flexible formulation of the system-bath coupling to allow for the treatment of different bath densities of state. In addition, we take advantage of the Gaussian nature of the bath to construct the MPS-IF in two ways, either by leveraging the Gaussian form of the IF when the bath is noninteracting or by propagating it forward iteratively in a similar spirit to the QUAPI method [29]. The two methods as presented here have different asymptotic costs with similar errors in the dynamics, which is suggestive of a hybrid scheme being useful for simulations to long times. Using this approach, we present a numerical solution of the Anderson impurity model including a comparative analysis and optimization of the errors arising from the required time discretization and bond dimension truncation. We thus obtain approaches to the impurity problem where the convergence is not determined by the standard bath size and sign issues, and which can produce a description of the true non-Markovian evolution of the impurity with polynomially scaling numerical effort and error control.

This paper is organized as follows: In Sec. II, we briefly outline the approach to real-time dynamics using discretized influence functionals and we give two schemes for constructing a matrix product state representation for the IF. In Sec. III, we compare our approach for the nonequilibrium dynamics of the Anderson model to recent exact calculations on the model. This comparison points to some important specific details of our approaches with respect to convergence, which are then discussed in detail. In Sec. IV, we conclude and discuss outstanding questions for future study. Details of the derivations are contained in the Supplemental Material [53].

II. INFLUENCE FUNCTIONALS AND THEIR REPRESENTATIONS

We consider the quench dynamics of the single impurity Anderson model,

\[
\hat{H} = \sum_{k,\sigma} E_{k,\sigma} c_{k,\sigma}^\dagger c_{k,\sigma} + \sum_{k,\sigma} (V_{k,\sigma} c_{k,\sigma}^\dagger \hat{d}_{\sigma} + \text{H.c.}) + U\hat{n}_1\hat{n}_\perp + \sum_{\sigma} \epsilon_{\sigma} \hat{n}_{\sigma}. \tag{1}
\]

Here, \(\epsilon_{\sigma}\) is the on-site energy for electrons with spin \(\sigma = \{\uparrow, \downarrow\}\) residing on the impurity, \(U\) is the Coulomb repulsion for two electrons that reside on the impurity, \(E_{k,\sigma}\) is the conduction (bath) electron energy with momentum \(k\), and \(V_{k,\sigma}\) characterizes the strength of the coupling between the impurity and bath electrons. For the remainder of this paper, we will refer to the terms in the first and second lines of Eq. (1) as \(\hat{H}_0\) and \(\hat{H}_1\), respectively; the bath-only terms within \(\hat{H}_0\) will also be denoted by \(\hat{H}_B\).

The dynamics we consider starts from an initially nonequilibrium state, in which the impurity is decoupled from the bath, \(\hat{\rho}_{\text{init}} = \hat{\rho}(0)\hat{\rho}_B\) [54]. The ensuing evolution of the impurity is approximated by a second-order Trotter decomposition as

\[
\dot{\rho}(N\Delta t) = \text{Tr}_B \left[ \left( e^{-\hat{H}_0} \hat{\rho}(0) e^{-\hat{H}_1} \right)^N \right] \left( \hat{\rho}(0)\hat{\rho}_B \right),
\]

where, for compactness, we have defined the superoperators \(e^{-i\hat{L}_0\Delta t} \equiv e^{-i\hat{H}_0\Delta t} \hat{A} e^{i\hat{H}_0\Delta t} \). The trace over the bath can be performed in the basis of coherent states when the statistics of the bath is Gaussian, e.g., \(\hat{\rho}_B \propto \exp(-\beta \hat{H}_B)\) (see Supplemental Material [53]). What remains then is the impurity dynamics captured by trajectories over coherent states on the forward (backward) contour, \(|\eta_n\rangle\langle\eta_n|\). Each of these trajectories is weighted by the influence functional \(I_N\) containing properties of the bath as well as the impurity-bath coupling, leading to the representation of the Trotterized dynamics as

\[
\langle \eta_N^* | \dot{\rho}(N\Delta t) | \eta_N \rangle = \int \left( \prod_n \mathcal{D}\eta_n \mathcal{D}\eta_n^* \right) I_N \{\eta_n\}, \{\eta_n^*\} \times \langle \eta_0^* | \hat{\rho}(t_N) | \eta_0 \rangle e^{\int_0^{t_N} \mathcal{S}_{\text{imp}}(\eta_\tau,\eta_\tau^*)(\tau)} \tag{2}
\]

Since \(\hat{H}_0\) is quadratic by construction, the influence functional takes the general Gaussian form,

\[
I_N[\eta, \eta^*] = \exp \left[ \begin{array}{c} \eta_1 \\ \vdots \\ \eta_N \end{array} \right] \cdot G \cdot \left[ \begin{array}{c} \eta_1 \\ \vdots \\ \eta_N \end{array} \right] \tag{3}
\]

where \(G\) is a matrix describing the temporal correlations in the impurity’s trajectories. For the single-impurity case that we consider here, the trajectory at time step \(N\) is specified using four states. In particular, for each branch on the Keldysh contour (forward and backward), we must keep track of the impurity’s state before and after it is acted on by \(\exp(\pm i\hat{H}_1\Delta t)\) at time \(N\Delta t\). Thus, the IF at the \(N\)th time step requires \(G\) to be a \(4N \times 4N\) matrix. The explicit expressions of Eqs. (2) and (3) are rather involved and are presented fully in [55].

To represent and compute quantities associated with \(I_N[\eta, \eta^*]\), we can treat the Grassmann variables \(\eta_n\) and \(\eta_n^*\) as fermionic operators [51,56]. This turns the \(I_N[\eta, \eta^*]\) into a generalized Gaussian state \(|\eta_N\rangle\),

\[
|\eta_N\rangle \propto \exp \left[ \frac{1}{2} \sum_{i,j} \hat{c}_i^\dagger G_{i,j} \hat{c}_j \right] |0_1 \ldots 0_{4N}\rangle. \tag{4}
\]

Having constructed \(G\), the brute of the numerical effort now lies in representing \(|\eta_N\rangle\). This can be accomplished using matrix product states (MPSs), which circumvent memory resource requirements growing exponentially with \(N\). Moreover, it has been recently demonstrated that for impurity models of the type considered here, the maximum entanglement entropy in \(|\eta_N\rangle\) typically saturates to an area-law
behavior as $N \to \infty$ [43,51]. This suggests that $|\Omega_N\rangle$, and therefore the impurity dynamics, can be efficiently simulated using an MPS with low bond dimension, at least for some classes of impurity models. The efficiency hinges crucially on the approach used to construct an MPS approximation of $|\Omega_N\rangle$ (MPS-IF). We now outline two methods for doing so: one which directly constructs each site tensor of the MPS-IF to produce an optimal low-rank MPS approximation, and another which reuses information from previous time steps.

### A. Direct construction

A many-body MPS can be constructed from its site tensors by considering the overlaps of Schmidt matrices for two different bipartitions of $|\Omega_N\rangle$ [57]. Here, one can leverage the fact that $|\Omega_N\rangle$ is a Bardeen-Cooper-Schrieffer (BCS) state [51] and therefore can be transformed into a Hartree-Fock state, for which an efficient method exists to construct its MPS representation directly [58]. The transformation into a Hartree-Fock state proceeds by finding the Bogoliubov quasiparticles, which can be done by diagonalizing the matrix $G$ in Eq. (4) at a cost of $O(N^3)$ for the $N$th time step, and performing a particle-hole transformation. This allows for a direct construction of the Schmidt decomposition of any bipartition of $|\Omega_N\rangle$ between sites $(\ell, \ell + 1)$ as

$$|\Omega_N\rangle \propto \prod_{i=1}^{2N} \left[ \sqrt{v_i^{(\ell)} \phi_{iL}^{(\ell)}} + \sqrt{1 - v_i^{(\ell)} \phi_{iR}^{(\ell)}} \right] \langle \Omega^{(\ell)}_L \rangle \otimes \langle \Omega^{(\ell)}_R \rangle. \tag{5}$$

The $\phi_{i}^{(\ell),L/R}$ is a set of orthonormal single-particle operators acting on the left/right partition, and $v_i^{(\ell)}$ are their associated eigenvalues [59]. Note that the left and right Schmidt states are guaranteed to be Hartree-Fock states. We can directly pick out the $D$ most relevant Schmidt states without needing to directly construct them. While we can do so at the single-particle level (i.e., approximating all but $\log_2 D$ values $v_i^{(\ell)}$ with 0 or 1), we will keep the $D$ Schmidt states of highest weight.

Having picked out the relevant Schmidt states, the site tensor $A_{\ell}^{(\ell)\sigma}$ at the $\ell$th site is specified by the overlaps,

$$A_{\alpha_L}^{(\ell)\sigma} = (\sigma_L)_{\alpha_L}. \tag{6}$$

where $\alpha_L$ labels the left Schmidt states of the bipartition at the $\ell$th bond. Via Wick’s theorem, the elements of the site tensor are straightforwardly found from determinants of overlaps of the single-particle states, e.g., $\langle \Omega_L^{(\ell)} \phi_{iL}^{(\ell-1)} \phi_{iR}^{(\ell)} \Omega_R^{(\ell)} \rangle$. Note that the matrix of overlaps only needs to be computed once per site tensor, as the required determinants can be formed from its submatrices. The cost of constructing all $O(N)$ single-particle operators $\phi_{i}^{(\ell),L/R}$ and the overlap matrix is $O(N^3)$. Furthermore, the evaluation of determinants for each site matrix can be sped up by the fact that certain orbitals are always occupied in all the considered Schmidt states. The determinant calculation can be broken up into the product of the determinant of the occupied block with the determinant of its Schur complement. In all, a single site tensor can be constructed at a cost of $O(N^3 + D^2N^2 \log_2 D)$ [58], meaning that the cost of constructing the full MPS-IF at time step $N$ is $O(N^4 + D^2N^3 \log_2 D)$. Note that the above procedure constructs each tensor independently, which leads to trivial parallelization, but the time may also be reduced if the truncated tensors are constructed in the basis of preceding tensors.

### B. Iterative construction

In addition to using the generalized Gaussian nature of $|\Omega_N\rangle$ to directly construct an MPS-IF, we can take an approach in the same spirit as similar methods for bosonic impurity problems, i.e., the time-evolving matrix product operator (TEMPO) reformulation of the QUAPI approach [45,46,49,50]. In TEMPO, the MPS-IF can be propagated from the $(N-1)$th to $N$th time step by contracting it with a layer of matrix product operators (MPOs), which contain information on how the impurity state at time step $N$ correlates with the impurity state at all previous times. To work in the same spirit, let us start from Eq. (4) and decompose the quadratic exponent into the form

$$|\Omega_N\rangle = \prod_j \exp \left[ \sum_{i=1}^{j-1} \Delta G_{ij} \hat{c}_i^{\dagger} \hat{c}_j \right] |\Omega_{N-1}\rangle, \tag{7}$$

where $\Delta G$ denotes the change in $G$ between time steps $N - 1$ and $N$ [53]. Let us examine one grouping of operators at fixed $j$ and define, for convenience, $g_i \equiv \Delta G_{ij} \equiv |g_i|e^{i\phi}$,

$$\exp \left[ \sum_{i=1}^{j-1} g_i \hat{c}^{\dagger}_i \hat{c}_j \right] = 1 + \sum_{i=1}^{j-1} g_i \hat{c}^{\dagger}_i \hat{c}_j. \tag{8}$$

This operator explicitly contains long-range couplings. While there are more sophisticated ways to represent such an object as an MPO [60], we will instead use a straightforward approach of decomposing it as a sequence of nearest-neighbor unitary transformations $V_{i-1,i}$. These are defined such that they transform a linear combination of two fermions into a single effective fermion,

$$g_{i-1} \hat{c}^{\dagger}_{i-1} + g_i \hat{c}^{\dagger}_i = V_{i-1,i} (e^{i\phi} \sqrt{|g_{i-1}|^2 + |g_i|^2} \hat{c}^{\dagger}_{i-1}) V^{\dagger}_{i-1,i},$$

from which one finds

$$V_{i-1,i} = \exp \left[ i \left( \frac{g_{i-1} - g_i}{2} \right) \hat{c}^{\dagger}_{i-1} \hat{c}_i - \text{H.c.} \right] \exp \left[ i \left( \frac{|g_{i-1}|}{|g_i|} \right) \left( \hat{c}^{\dagger}_{i-1} \hat{c}_i - \text{H.c.} \right) \right]. \tag{9}$$

Combining these transformations gives a representation of Eq. (8) for a single $j$. Constructing the MPS-IF at the $N$th time step from the $(N - 1)$th time step requires the $O(N)$ applications of Eq. (8), each of which involves $O(N)$ two-site gates. To keep the bond dimension manageable, truncations via singular-value decompositions (SVDs) are performed after
are results with a time step of the order of $10^{-3}$ over the time range $\Gamma t \in [0, 5]$. The deviations are generally larger than violations of the trace condition, $\text{Tr} \, \rho(t) = 1$ [61]. Similar magnitudes of error are present from the iteratively constructed MPS-IF, which we show in [55]. Notably, the dynamics from both construction methods do not perfectly coincide due to the use of SVDs to compress the iteratively constructed MPS-IF. For larger bond dimensions, this problem is exacerbated by the fact that the Schmidt values of $|\psi_i\rangle$ can fall below the precision of 64-bit floating point numbers so that the accuracy of the associated Schmidt vectors found by SVD cannot be guaranteed. This, however, poses no issue for the direct construction method, for which the Schmidt values are found from products of the $\sqrt{\Gamma}$ and $\sqrt{\Gamma - \nu_i}$ in Eq. (5).

The above considerations suggest that truncation error in the MPS construction must be carefully examined at the bond dimensions we use and must be considered in addition to Trotter error. We thus undertake a closer examination of these errors below.

A. Convergence analysis in the noninteracting limit

We begin our discussion by considering the dynamics for the case $U = 0$, $\epsilon_\sigma \neq 0$ since this admits exact numerical solutions [62]. In Fig. 2, we show the deviation of the double occupancy $\langle \hat{n}_i \hat{n}_j \rangle$, as computed by the direct and iterative methods, from the exact values over a range of on-site energies $\epsilon_\sigma$. The most striking feature in these plots is a discontinuous rate of growth in the error $\delta \langle \hat{n}_i \hat{n}_j \rangle = \langle \hat{n}_i \hat{n}_j \rangle - \langle \hat{n}_i \hat{n}_j \rangle_{\text{ref}}$, most prominently exhibited in simulations using MPS-IF with smaller bond dimensions and in the cases where the direct construction approach is used; see Figs. 2(a)–2(c). Specifically, this behavior appears in Fig. 2(b) at $\Gamma t \approx 1.2$. In contrast, the nearly discontinuous behavior appears to be smoothed over in the iterative construction [Figs. 2(d)–2(f)], although the deviations in the double occupancy generally follow the same trends as in the directly constructed cases. We observe $t^*$ to increase modestly with increasing bond dimension. For instance, at $D = 256$, $\Gamma t^* \approx 1.6$. Since the deviations of the iteratively constructed method and direct method are qualitatively similar, below we focus on the behavior of the direct method. The error incurred by the MPS approximation can be isolated by examining the $\delta = 0$ case, where there is no Trotter error. These results are shown in the insets of Figs. 2(b) and 2(e). It can be seen that the error increases dramatically with propagation time, growing by five decades over 20 time steps for $D = 128$ before saturating. The error can be suppressed by increasing the bond dimension. We generally see that doing so decreases the error algebraically with $D$ (Fig. 3). Note that this holds at both intermediate ($\Gamma t \sim 1$) and long ($\Gamma t \gtrsim 2$) times for which the impurity dynamics is close to its steady-state behavior [see bottom-most curve in the inset of Fig. 2(c)].

At the same time, over the time range for which the dynamics are converged with respect to the available bond dimensions ($\Gamma t \lesssim 0.6$), the Trotter error scales as $(\Delta t)^2$ [55]. Thus, ensuring that the Trotter error is smaller than a tolerance $\delta$ implies that $\Delta t \propto \delta^{1/2}$ and $N \propto \delta^{-1/2}$. If the cost to reach a truncation error $\delta$ also scales asymptotically as a power law in

\[ \Delta(\omega) = \Gamma \sqrt{W^2 - \omega^2}/\pi, \]  

where $W = 10\Gamma$ and $\omega \in [-W, W]$. The bath is initially equilibrated at a temperature $\Gamma/\beta = 2$ and is decoupled from an unoccupied impurity $\hat{\rho}(0) = |0\rangle \langle 0|$. For $U = -2\epsilon_\sigma = 2.5\Gamma$, we compare our results, generated using the direct and iterative constructions without memory length truncations, to those of the MPS time-dependent variational principle calculations of Kohn and Santoro [16].

In Fig. 1, we show the quench dynamics of the populations, $P_{\langle \psi | \hat{\rho} | \psi \rangle}$, from the directly constructed MPS-IF with a maximum bond dimension $D = 64$. The time step used is $\Gamma \Delta t = 0.05$. As seen in the inset, the absolute difference between our results and those of Kohn and Santoro is of the order of $10^{-3}$ over the time range $\Gamma t \in [0, 5]$. The deviations are generally larger than violations of the trace condition, $\text{Tr} \, \hat{\rho}(t) = 1$. Similar magnitudes of error are present from the iteratively constructed MPS-IF, which we show in [55]. Notably, the dynamics from both construction methods do not perfectly coincide due to the use of SVDs to compress the iteratively constructed MPS-IF. For larger bond dimensions, this problem is exacerbated by the fact that the Schmidt values of $|\psi_i\rangle$ can fall below the precision of 64-bit floating point numbers so that the accuracy of the associated Schmidt vectors found by SVD cannot be guaranteed. This, however, poses no issue for the direct construction method, for which the Schmidt values are found from products of the $\sqrt{\Gamma}$ and $\sqrt{\Gamma - \nu_i}$ in Eq. (5).

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FIG. 2. Difference between the double occupancy computed from the MPS-IF with $\Gamma \Delta t = 0.05$ and its exact value from the solution of a differential equation [55]. The bath is defined by Eq. (10) in the $U = 0$ limit at temperature $\Gamma \beta = 2$, with an initially unoccupied impurity. The MPS-IF is constructed (a)–(c) directly and (d)–(f) iteratively, and $D = (a),(d)$ 64, (b),(e) 128, and (c),(f) 256. Inset of (a) and (e): The absolute difference for $\epsilon = 0$ across different bond dimensions. Inset of (c): The exact double occupancies used as reference in the main panels. The red dashed line is the steady-state value for $\epsilon = 0$.

1/δ, then the total cost to simulate the noninteracting impurity dynamics with the MPS-IF to the specified error tolerance will scale polynomially with 1/δ.

B. Convergence analysis in the interacting limit

Given that the influence functional is independent of the details of the impurity Hamiltonian $\hat{H}_1$, vestiges of the irregularities in the error from the noninteracting dynamics should also appear for the interacting dynamics. This fact is useful since it implies that by examining the noninteracting dynamics, we can anticipate points in time to focus convergence efforts for the interacting problem, e.g., close to the $t^*$ identified in the previous section. However, the information gleaned from the noninteracting dynamics does not tell the full story. For example, Fig. 2 shows that the double occupancy appears to be adequately converged with respect to $D$ around $\Gamma t \approx 0.7$. Yet in the full problem with $U = -2\epsilon_\sigma = 2.5\pi \Gamma$ as shown in Fig. 4, we see that the error can behave rather differently as a function of both $D$ and $\Delta t$. First, for $D \gtrsim 256$, we see that the deviations are most sensitive to the time-step size, though there are small residual truncation errors independent of the Trotter error. We surmise that the errors shown in Fig. 4(a) are mostly due to Trotter error since $\delta \langle \hat{n}_\uparrow \hat{n}_\downarrow \rangle$ are identical for $D = 256$ and 512. At the same time, the error for $D = 64$ is mostly unaffected by decreasing $\Delta t$, indicating that truncation errors are dominant. We conclude that decreasing the time step size does not necessarily decrease the overall error given by the MPS-IF at fixed bond dimension. As noted in Ref. [51], the half-cut von Neumann entanglement entropy of the MPS-IF vanishes as $\Delta t \rightarrow 0$. However, this vanishing of the entanglement entropy, which is due to the scaling of the Schmidt values with $\Delta t$ [52,55], does not imply that the accuracy of the MPS-IF approximation with fixed $D$ improves as $\Delta t \rightarrow 0$. Our lack of rigorous understanding of the overall error is highlighted by the growth of deviations with decreasing $\Delta t$ for $D = 128$, seen in Figs. 4(c)–4(e). We observe a similar error behavior from the iterative construction, shown in the dashed lines in Fig. 4(c).

Finally, we can make a similar comparison of the intermediate- to long-time dynamics as in Fig. 3 for the full $U \neq 0$ problem. Restricting ourselves to the direct construction of the MPS-IF, we show in Fig. 5 the errors in the double occupancy across bond dimensions, holding $\Gamma \Delta t = 0.05$. As

FIG. 3. Absolute deviations from the exact double occupancy for $U = 0$, $\epsilon = 0$ for various bond dimensions $D$ at fixed times approaching the steady state of the dynamics. The bath is initially at temperature $\Gamma \beta = 2$ and the impurity is initially unoccupied. The MPS-IF is constructed directly with a time step of $\Gamma \Delta t = 0.05$. 

FIG. 4. Absolute deviations from the exact double occupancy for $U = 0$, $\epsilon = 0$ for various bond dimensions $D$ at fixed times approaching the steady state of the dynamics. The bath is initially at temperature $\Gamma \beta = 2$ and the impurity is initially unoccupied. The MPS-IF is constructed directly with a time step of $\Gamma \Delta t = 0.05$. 

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in the noninteracting case, the errors at long times $\Gamma t \gtrsim 2$ suggest a power-law decay with $D$ with a similar exponent. In contrast, errors at the intermediate time $\Gamma t = 1$ do not exhibit such a decay, taking into account that the $D = 128$ dynamics (Fig. 4) suffers from larger-than-expected errors. Similar to Fig. 4(a), it is likely that the errors at $\Gamma t = 1$ for $D \neq 128$ are dominated by Trotter errors. Achieving convergence with respect to $D$ and $\Delta t$ may require first decreasing $\Delta t$ by a factor of $\sim 3$; cf. Figs. 4(a) and 4(c). While these results are suggestive of the long-time ($\Gamma t \gtrsim 2$) dynamics being convergeable with polynomial effort for a specified error tolerance, larger bond dimensions will be required to reach this asymptotic convergence regime at intermediate times.

IV. DISCUSSION

In this paper, we have presented two representations of fermionic bath influence functionals with matrix product states to simulate the real-time dynamics of the single-impurity Anderson model. We have found that we can obtain good agreement with other real-time propagation methods with modest numerical effort. We have shown that both construction approaches yield similar errors in the resulting impurity dynamics. The two ways of constructing the IF have costs that scale differently with the number of propagation time steps, meaning that a hybrid approach may be a good strategy for dealing with long-time propagation. Computational costs of both methods as presented here can be further attenuated by introducing additional approximations, which we did not consider here as these would complicate the convergence analysis. We note that the MPS-IF can treat arbitrary bath densities of state through the specification of its hybridization function [55], and obtain systematically convergeable simulations of non-Markovian dynamics. We stress here that unlike other dynamics methods, once we have constructed and saved the MPS-IF, calculations with different impurity Hamiltonians $\hat{H}_1$ and different initial impurity states $\hat{\rho}(0)$ can be performed trivially. This allows for the treatment of, e.g., time-dependent forms of $\hat{H}_1$ so that problems with external driving can be treated with no additional cost. Furthermore, viewing the IF as a “process tensor” of an open quantum dynamics [63] means that we can easily extract arbitrary multitime impurity correlation functions within the same formalism.

We have also shown that the outstanding sources of error, i.e., Trotter error and truncation error of the MPS-IF, can likely be controlled with only polynomially growing resource requirements, but the two errors do not necessarily go hand in hand. In particular, our construction of the IF as a process tensor allows us to isolate the MPS truncation error, and gives us a guide to where convergence efforts should be focused. As presented, our approach should be readily generalizable to other impurity problems, and with some effort can be adapted as an impurity solver for dynamical mean-field theory. These considerations, as well as modifications to improve computational efficiency, will be taken up in future work.

As it stands, we currently do not have a complete understanding of the major determinants of errors stemming from truncating the MPS-IF. We anticipate that insights in this direction will help make the direct construction of the MPS-IF more efficient. Further explorations along these lines are forthcoming.

Recently, we became aware of a similar work by Thoenniss et al. [64].

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[53] See Supplemental Material at http://link.aps.org/supplemental/10.1103/PhysRevB.107.125103 for additional data, along with details of the derivation of the IF and the computation using the MPS-IF.
[54] In cases where all terms of the impurity-bath coupling can be simultaneously diagonalized, $\Delta G$ is zero everywhere except for the rows/columns containing correlations between the “newest” impurity states $\hat{c}_{4n-3}^\dagger \ldots \hat{c}_{4n}^\dagger$ and the previous states.
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Such violations are due solely to the approximation of $|k\rangle$ as a finite bond dimension MPS, since the IF represents unitary Trotterized dynamics.

Unlike for other methods, the dynamics of the noninteracting case are just as difficult to compute via the IF approach as they are for $U \neq 0$, since the presence or absence of $U$ does not alter the IF itself.

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