Finite-temperature density matrix embedding theory

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We describe a formulation of the density matrix embedding theory at finite temperature. We present a generalization of the ground-state bath orbital construction that embeds a mean-field finite-temperature density matrix up to a given order in the Hamiltonian, or the Hamiltonian up to a given order in the density matrix. We assess the performance of the finite-temperature density matrix embedding on the one-dimensional Hubbard model both at half-filling and away from it, and the two-dimensional Hubbard model at half-filling, comparing to exact data where available, as well as results from finite-temperature density matrix renormalization group, dynamical mean-field theory, and dynamical cluster approximations. The accuracy of finite-temperature density matrix embedding appears comparable to that of the ground-state theory, with, at most, a modest increase in bath size, and competitive with that of cluster dynamical mean-field theory.

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I. INTRODUCTION

The numerical simulation of strongly correlated electrons is key to understanding the quantum phases that derive from electron interactions, ranging from the Mott transition [1–4] to high-temperature superconductivity [5–7]. Consequently, many numerical methods have been developed for this task. In the setting of quantum lattice models, quantum embedding methods [8], such as dynamical mean-field theory (DMFT) [9–13] and density matrix embedding theory (DMET) [14–20], have proven useful in obtaining insights into complicated quantum phase diagrams. These methods are based on an approximate mapping from the full interacting quantum lattice to a simpler self-consistent quantum impurity problem, consisting of a few sites of the original lattice coupled to an explicit or implicit bath. In this way, they avoid treating an interacting quantum many-body problem in the thermodynamic limit.

The current work is concerned with the extension of DMET to finite temperatures. DMET so far has mainly been applied in its ground-state formulation (GS-DMET), where it has achieved some success, particularly in applications to quantum phases where the order is associated with large unit cells [17,18,21]. The ability to treat large unit cells at relatively low cost compared to other quantum embedding methods is due to the computational formulation of DMET, which is based on modeling the ground-state impurity density matrix, a time-independent quantity accessible to a wide variety of efficient quantum many-body methods. Our formulation of finite-temperature DMET (FT-DMET) is based on the simple structure of GS-DMET, but includes the possibility to generalize the bath so as to better capture the finite-temperature impurity density matrix. Bath generalizations have previously been used to extend GS-DMET to the calculation of spectral functions and other dynamical quantities [22,23]. Analogously to GS-DMET, since one only needs to compute time-independent observables, finite-temperature DMET can be paired with the wide variety of quantum impurity solvers which can provide the finite-temperature density matrix.

We describe the theory of FT-DMET in Sec. II. In Sec. III we carry out numerical calculations on the one-dimensional (1D) and two-dimensional (2D) Hubbard models, using exact diagonalization (ED) and the finite-temperature density matrix renormalization group (FT-DMRG) [24] as quantum impurity solvers. We benchmark our results against those from the Bethe Ansatz in one dimension, and DMFT and the dynamical cluster approximation (DCA) in two dimensions, and also explore the quantum impurity derived N´eel transition in the 2D Hubbard model. We finish with brief conclusions about prospects for the method in Sec. IV.

II. THEORY

A. Ground-state DMET

In this work, we exclusively discuss DMET in lattice models (rather than for ab initio simulations [15,16,20,25]). As an example of a lattice Hamiltonian, and one that we will use in numerical simulations, we define the Hubbard model [26,27]

$$\hat{H} = -t \sum_{\langle i,j \rangle, \sigma} \hat{a}_{i \sigma}^\dagger \hat{a}_{j \sigma} - \mu \sum_{i, \sigma} \hat{a}_{i \sigma}^\dagger \hat{a}_{i \sigma} + U \sum_{i} \hat{n}_{i \uparrow} \hat{n}_{i \downarrow},$$  \hspace{1cm} (1)

where $\hat{a}_{i \sigma}^\dagger$ creates an electron with spin $\sigma$ on site $i$ and $\hat{a}_{i \sigma}$ annihilates it; $\hat{n}_{i \sigma} = \hat{a}_{i \sigma}^\dagger \hat{a}_{i \sigma}$; $t$ is the nearest-neighbour
(denoted $\langle i, j \rangle$) hopping amplitude, here set to 1; $\mu$ is a chemical potential; and $U$ is the on-site repulsion.

The general idea behind a quantum embedding method such as DMET is to approximately solve the interacting problem in the large lattice by dividing the lattice into small fragments or impurities [8]. (Here we will assume the impurities are nonoverlapping.) The main question is how to treat the coupling and entanglement between the impurities. In DMET, other fragments around a given impurity are modeled by a set of bath orbitals. The bath orbitals are constructed to exactly reproduce the entanglement between the impurity and environment when the full lattice is treated at a mean-field level (the so-called “low-level” theory). The impurity together with its bath orbitals then constitutes a small embedded quantum impurity problem, which can be solved with a “high-level” many-body method. The low-level lattice wave function and the high-level embedded impurity wave function are made approximately consistent, by enforcing self-consistency of the single-particle density matrices of the impurities and of the lattice. This constraint is implemented by introducing a static correlation potential on the impurity sites into the low-level theory. The correlation potential introduced in DMET is analogous to the DMFT self-energy. A detailed discussion of the correlation potential including the comparison to other approaches such as density functional theory (DFT) can be found in Refs. [8,15].

To set the stage for the finite-temperature theory, in the following we briefly recapitulate some details of the above steps in the GS-DMET formulation. In particular, we discuss how to extract the bath orbitals, how to construct the embedding hamiltonian, and how to carry out the self-consistency steps in the GS-DMET formulation. In particular, we discuss following we briefly recapitulate some details of the above approaches such as DMET is to approximately solve the interacting bath formulation, where both ED and FT-DMRG solvers are used. This second choice is because the cost of interacting bath formulation in the 1D Hubbard model where an acting bath formulation in the 2D Hubbard model, we use the non-interacting bath formulation, where both ED and FT-DMRG solvers are used. This second choice is because the cost of treating nonlocal interactions in FT-DMRG is relatively high (and we make the same choice with ED solvers to keep the results strictly comparable).

1. DMET bath construction

We first assume that we have a mean-field (“low-level”) wave function $|\Psi\rangle$ defined on the full lattice of $L$ sites; $|\Phi\rangle$ is an eigenstate of a quadratic lattice Hamiltonian $\hat{h}$. Then, given an impurity $x$ defined over $L_x$ sites, DMET relies on the lemma that $|\Phi\rangle$ can be rewritten in the form

$$|\Phi\rangle = |\Phi_{\text{emb}}\rangle|\Phi_{\text{core}}\rangle,$$

(2)

where $|\Phi_{\text{emb}}\rangle$ lives in a Hilbert space of the $L_x$ sites of the impurity and an additional $L_x$ bath orbitals defined in its environment; $|\Phi_{\text{core}}\rangle$ is a product state defined in the complement of the impurity and environment Hilbert spaces. This basic result states that the entanglement between the impurity and its environment contained in $|\Phi\rangle$ is entirely captured within the small impurity-bath Hilbert space.

The single-particle density matrix $D^{\Phi}$ obtained from $|\Phi\rangle$ contains all information on the correlations in $|\Phi\rangle$. Thus the bath orbitals can be defined from this density matrix. We consider the impurity-environment block $D_{\text{imp-env}}^{\Phi}$ ($D_{ij}$ for $i \in x$, $j \notin x$) of dimension $L^x \times (L - L^x)$. Then taking the thin SVD

$$D_{\text{imp-env}}^{\Phi} = U\Lambda B^\dagger$$

(3)

do the columns of $B$ specify the bath orbitals in the lattice basis. The bath space is thus a function of the density matrix, denoted $B(D)$.

2. Embedding Hamiltonian

After obtaining the bath orbitals, we construct the embedded Hamiltonian of the quantum impurity problem. In GS-DMET, there are two ways to do so, the interacting bath formulation and the noninteracting bath formulation. The conceptually simplest approach is the interacting bath formulation. In this case, we project the interacting lattice Hamiltonian $\hat{H}$ into the space of the impurity plus bath orbitals, defined by the projector $\hat{P}$, i.e., the embedded Hamiltonian is $\hat{H}_{\text{emb}} = \hat{P}\hat{H}\hat{P}$. $\hat{H}_{\text{emb}}$ in general contains nonlocal interactions involving the bath orbitals, as they are nonlocal orbitals in the environment. From the embedded Hamiltonian, we compute the high-level ground-state impurity wave function,

$$\hat{H}_{\text{emb}} |\Psi\rangle = E |\Psi\rangle.$$  

(4)

If $\hat{H}$ were itself the quadratic lattice Hamiltonian $\hat{h}$, then then $|\Psi\rangle = |\Phi\rangle$ and

$$\hat{P}\hat{h}\hat{P}|\Phi\rangle = E|\Phi\rangle.$$  

(5)

Another way to write Eq. (5) for a mean-field state is

$$[\hat{P}\hat{h}\hat{P}, PD^{\Phi} P] = 0,$$

(6)

where $\hat{h}$ denotes the single-particle Hamiltonian matrix and $P$ is the single-particle projector into the impurity and bath orbitals. These conditions imply that the lattice Hamiltonian and the embedded Hamiltonian $\hat{H}_{\text{emb}}$ share the same ground state at the mean-field level, which is the basic approximation in GS-DMET.

In the alternative noninteracting bath formulation, interactions on the bath are approximated by a quadratic correlation potential (discussed below). This formulation retains the same exact embedding property as the interacting bath formulation for a quadratic Hamiltonian. In practice, both formulations give similar results in the Hubbard model [19,28], and the choice between the two depends on the available impurity solvers; the interacting bath formulation generates nonlocal two-particle interactions in the bath that not all numerical implementations can handle. In this work, we use the interacting bath formulation in the 1D Hubbard model where an ED solver is used. In the 2D Hubbard model, we use the non-interacting bath formulation, where both ED and FT-DMRG solvers are used. This second choice is because the cost of treating nonlocal interactions in FT-DMRG is relatively high (and we make the same choice with ED solvers to keep the results strictly comparable).

3. Self-consistency

To maintain self-consistency between the ground state of the lattice mean-field $|\Phi\rangle$, and that of the interacting embedded Hamiltonian $|\Psi\rangle$, we introduce a quadratic correlation potential $\hat{u}$ into $\hat{h}$, i.e.,

$$\hat{h} \rightarrow \hat{h} + \hat{u},$$

(7)
where \( \hat{u} \) is constrained to act on sites in the impurities, i.e., \( \hat{u} = \sum_x \hat{u}_x^i \). To study magnetic order, we choose the form

\[
\hat{u}_x^i = \sum_{j \in x, \sigma \in \{\uparrow, \downarrow\}} u_{ij}^\sigma a_{i\sigma} a_{j\sigma}^\dagger.
\]  

(8)

The coefficients \( u_{ij}^\sigma \) are adjusted to match the density matrices on the impurity that are evaluated from the low-level wave function (\( \Phi \)) and from the high-level embedded wave function (\( |\Psi\rangle \)). In this work, we only match the single-particle density elements of the impurity (impurity only matching [16]). Note also that we will only be considering translationally invariant systems, and thus \( \hat{u} \) is the same for all impurities.

### B. Ground-state expectation values

Ground-state expectation values are evaluated from the density matrices of each high-level impurity wave functions (\( |\Psi\rangle \)). Since there are multiple impurities (in a translationally invariant system, these are constrained to be identical) an expectation value is typically assembled from the multiple impurity wave functions using a democratic partitioning [16]. For example, given two sites \( i, j \), where \( i \) is part of impurity \( x \) and \( j \) is part of impurity \( y \), the expectation value of \( a_x^i a_j^\dagger \) is

\[
\langle a_x^i a_j^\dagger \rangle = \frac{1}{2} [\langle |\Psi\rangle a_x^i a_j^\dagger |\Psi\rangle + \langle |\Psi\rangle a_x^i a_j^\dagger |\Psi\rangle].
\]  

(9)

Note that the pure bath components of the high-level wave functions, e.g., \( \langle |\Psi\rangle a_x^i a_j^\dagger |\Psi\rangle \) for \( i, j \notin x \) are not used in defining the DMET expectation values. Instead, the democratic partitioning is arranged such that an individual impurity embedding contributes the correct amount to a global expectation value so long as the impurity wave function produces correct expectation values for operators that act on the impurity alone, or the impurity and bath together.

### C. Finite-temperature DMET

Our formulation of FT-DMET follows the same rubric as the ground-state theory: a low-level (mean-field-like) finite-temperature density matrix is defined for the lattice; this is used to obtain a set of bath orbitals to define the impurity problem; a high-level finite-temperature density matrix is calculated for the embedded impurity; and self-consistency is carried out between the two via a correlation potential. The primary difference lies in the properties of the bath, which we focus on below, as well as in the appearance of quantities such as the entropy, which are formally defined from many-particle expectation values.

#### 1. Finite-temperature bath construction

In GS-DMET, the bath orbital construction is designed to be exact if all interactions are treated at the mean-field level, giving rise to the commuting condition for the projected single-particle density matrix and projected Hamiltonian in Eq. (6). In general, we can look for a finite-temperature bath construction that preserves a similar property. As pointed out in Sec. II B, the DMET embedding is still exact for single-particle expectation values if the embedded projected single-particle density matrix produces the correct expectation values in the impurity and impurity-bath sectors, due to the use of the democratic partitioning. We aim to satisfy this slightly relaxed condition.

The finite-temperature single-particle density matrix of a quadratic Hamiltonian \( \hat{h} \) is given by the Fermi-Dirac function

\[
D(\beta) = \frac{1}{1 + e^{(\beta \epsilon - \mu) / k_B T}}.
\]

(10)

where \( \beta = 1 / k_B T \) (\( k_B \) is the Boltzmann constant, \( T \) is the temperature). In the following, we fix \( k_B = 1 \), thus \( \beta = 1 / T \). If we could find an embedding directly analogous to the ground-state construction, we would obtain a projector \( P \), such that the embedded density matrix \( PDP \) is the Fermi-Dirac function of the embedded quadratic Hamiltonian, i.e., \( PhP \), i.e.,

\[
PDP = \frac{1}{1 + e^{(\beta PhP - \mu) / k_B}}.
\]

(11)

However, unlike in the ground-state theory, the nonlinearity of the exponential function means that Eq. (11) can only be satisfied exactly if \( P \) projects back into the full lattice basis. Thus a bath orbital construction at finite temperature is necessarily always approximate, even for quadratic Hamiltonians.

Nonetheless, one can choose the bath orbitals to reduce the error between the left-hand side (l.h.s.) and right-hand side (r.h.s.) in Eq. (11). First, we require that the equality is satisfied only for the impurity only, and impurity-environment, blocks of \( D \), following the relaxed requirements of the democratic partitioning. Second, we require the equality to be satisfied only up to a finite order \( n \) in \( h \), i.e.,

\[
[PDP]_{ij} = \left[ \frac{1}{1 + e^{(\beta PhP - \mu) / k_B}} \right]_{ij} + O(h^n) \quad i \in x, j \notin x.
\]

(12)

Then there is a simple algebraic construction of the bath space as (see the Appendix for a proof)

\[
\{B(h^k) \oplus B(h^k) \oplus B(h^k) \ldots B(h^k)\},
\]

(13)

where \( B(h^k) \) is the bath space derived from \( h^k \), \( k = 1, \ldots, n \). Note that each order of \( h \) adds \( L_a \) bath orbitals to the total impurity plus bath space.

We can alternatively choose the bath to preserve the inverse relationship between the density matrix and Hamiltonian

\[
[PhP]_{ij} = \text{inverseFD}(PDP) + O(D^n) \quad \text{not } i, j \notin x,
\]

(14)

where “inverseFD” is the inverse Fermi-Dirac function, and the bath space is then given as

\[
\{B(D) \oplus B(D^2) \oplus B(D^2) \ldots B(D^n)\}.
\]

(15)

The attraction of this construction is that the lowest order corresponds to the standard GS-DMET bath construction.

The above-generalized bath constructions allow for the introduction of an unlimited number of bath sites (so long as the total number of sites in the embedded problem is less than the lattice size). Increasing the size of the embedded problem by increasing the number of bath orbitals (hopefully) increases the accuracy of the embedding, but it also increases the computational cost. However, an alternative way to increase accuracy is simply to increase the number of impurity sites. Which strategy is better is problem-dependent, and we will assess both in our numerical experiments.
2. Thermal observables

The thermal expectation value of an observable $\hat{O}$ is defined as

$$\langle \hat{O}(\beta) \rangle = \text{Tr}[\hat{\rho}(\beta)\hat{O}].$$

(16)

Once $\hat{\rho}(\beta)$ is obtained from the high-level impurity calculation, for observables based on low-rank (e.g., one- and two-) particle reduced density matrices, we evaluate Eq. (16) using the democratic partitioning formula for expectation values in Sec. II B.

We will also, however, be interested in the entropy per site, which is a many-particle expectation value. Rather than computing this directly as an expectation value, we will obtain it by using the thermodynamic relation $\frac{dS}{dE} = \beta$, and

$$S(\beta_0) = S(0) + \int_{E(0)}^{E(\beta_0)} \beta(E)dE,$$

(17)

where $\beta_0$ is the desired inverse temperature, and $S(0) = \ln 4 \approx 1.386$.

III. RESULTS

A. Computational details

We benchmarked the performance of FT-DMET in the 1D and 2D Hubbard models as a function of $U$ and $\beta$. For the 1D Hubbard model, we compared our FT-DMET results to exact solutions from the thermal Bethe Ansatz [29]. For the 2D Hubbard model, the FT-DMET results were compared to DCA and DMFT results [30–35]. We used large DMET mean-field lattices with periodic boundary conditions (240 sites in one dimension, $24 \times 24$ sites in two dimensions). We used exact diagonalization (ED) and finite-temperature DMRG (FT-DMRG) as impurity solvers. There are two common ways to carry out finite-temperature DMRG calculations: the purification method [24] and the minimally entangled typical thermal states (METTS) method [36]. In this work, we used the purification method implemented with the ITENSOR package [37] as the FT-DMRG impurity solver, as well as to provide the finite lattice reference data in Fig. 5. In the FT-DMRG solver, the sites were ordered with the impurity sites coming first, followed by the bath sites (an orthonormal basis for the set of bath sites of different orders was constructed via singular value decomposition, and ordered in decreasing size of the singular values) and the ancillae arranged in between each physical site. In the 1D Hubbard model, we used ED exclusively and the interacting bath formulation of DMET, while in the 2D Hubbard model, we used ED for the four impurity, four bath calculations, and FT-DMRG for the four impurity, eight bath calculations, both within the noninteracting bath formulation. FT-DMRG was carried out using fourth-order Runge-Kutta time evolution. To denote different calculations with different numbers of impurity and bath orbitals, we use the notation $InBm$, where $n$ denote the number of impurity sites and $m$ the number of bath orbitals.

B. 1D Hubbard model

The 1D Hubbard model is an ideal test system for FT-DMET as its thermal properties can be exactly computed via the thermal Bethe ansatz. We thus use it to assess various choices within the FT-DMET formalism outlined above.

We first compare the relative performance of the two proposed bath constructions, generated via the Hamiltonian in Eq. (13) or via the density matrix in Eq. (15). In Fig. 1 we show the error in the energy per site (measured from the thermal Bethe ansatz) for $U = 2, 4$ and half-filling for these two choices. (The behavior for other $U$ is similar). Using four bath sites, the absolute error in the energy is comparable to that of the ground-state calculation (which uses two bath sites) over the entire temperature range. Although the Hamiltonian construction was motivated by the high-temperature expansion of the density matrix, the density matrix construction appears to perform well at both low and high temperatures. Consequently, we use the density matrix derived bath in the subsequent calculations.

We next examine the effectiveness of the density matrix bath construction in removing the the finite-size error of the impurity. As a first test, in Fig. 2 we compare the energy...
error obtained with FT-DMET and I2B2 to a pure ED calculation with four impurity sites (I4) and periodic (PBC) or antiperiodic (APBC) boundary conditions, at various $U$ and $\beta$. For weak ($U = 2$) to moderate ($U = 4$) coupling, FT-DMET shows a significant improvement over a finite system calculation with the same number of sites, reducing the error by a factor of $\sim 2$ for weak $\beta$, thus demonstrating the effectiveness of the bath. The maximum FT-DMET energy error is 8.1, 6.6, 3.1% for $U = 2, 4, 8$. At very strong couplings, the error of the finite system ED with PBC approaches that of FT-DMET. This is because both the finite-size error and the effectiveness of the DMET bath decrease as one approaches the atomic limit.

As a second test, in Fig. 3 we compare increasing the number of impurity sites versus increasing the number of bath orbitals generated in Eq. (15) for various $U$ and $\beta$. Although complex behavior is seen as a function of $\beta$, we roughly see two patterns. For certain impurity sizes, (e.g., I4) it can be slightly more accurate to use a larger impurity with an equal number of bath sites, than a smaller impurity with a larger number of bath sites. (For example, at $U = 8$, one can find a range of $\beta$ where I4B4 gives a smaller error than I2B6.) However, there are also some impurity sizes which perform very badly; for example I3B3 gives a very large error because the (short-range) antiferromagnetic correlations do not properly tile between adjacent impurities when the impurities are of odd size. Thus, due to these size effects convergence with impurity size is highly nonmonotonic, but increasing the bath size [by including more terms in Eq. (15)] is less prone to strong odd-size effects. The ability to improve the quantum impurity by simply increasing the number of bath sites, is expected to be particularly relevant in higher-dimensional lattices such as the 2D Hubbard model, where ordinarily to obtain a sequence of clusters with related shapes it is necessary to increase the impurity size by large steps. Nonetheless, convergence with bath size is also not strictly monotonic, as also illustrated in Fig. 4, where we see the error in the I2B4 entropy can sometimes be less than that of I2B6 for certain ranges of $\beta$. For the largest embedded problem I2B6, the maximum error in the entropy is $4 \times 10^{-3}$, $2 \times 10^{-2}$ for $U = 4, 8$.

The preceding calculations were all carried out at half-filling. Thus, in Fig. 5 we show FT-DMET calculations on the 1D Hubbard model away from half-filling at $U = 4$. We chose to simulate a finite Hubbard chain of 16-sites with PBC to readily generate numerically exact reference data using FT-DMRG (using a maximum bond dimension of 2000, and an imaginary time step of $\tau = 0.025$). The agreement between the FT-DMRG energy per site and that obtained from the thermal Bethe ansatz can be seen at half-filling, corresponding to a chemical potential $\mu = 2$. We see excellent agreement between FT-DMET and FT-DMRG results across the full range of chemical potentials, and different $\beta$, suggesting that FT-DMET works equally well for doped system as well as for undoped systems.

\[ \beta = 0.2 \]
\[ \beta = 0.3 \]
\[ \beta = 0.4 \]

FIG. 5. Energy per site (units of $t$) of a 16-site Hubbard chain with periodic boundary conditions at $U = 4$ as a function of the chemical potential $\mu$ at various $\beta$ values. The difference between the DMRG and DMET (I2B6) energies per site is 0.01–1.4%. Solid lines: DMRG energies; dashed lines: DMET energies; pentagons: Bethe Ansatz.
C. 2D Hubbard model

The 2D Hubbard model is an essential model of correlation physics in materials. We first discuss the accuracy of FT-DMET for the energy of the 2D Hubbard model at half-filling, shown in Fig. 6. The FT-DMET calculations are performed on a $2 \times 2$ impurity, with four bath orbitals ($I4B4$) (green diamond markers) and eight bath orbitals ($I4B8$) (red triangular markers). The results are compared to DCA calculations with clusters of size 34 (orange circle markers), 72 (blue square markers) [35], and $2 \times 2$ (light blue hexagon markers) (computed for this work). The DCA results with the size 72 cluster can be considered here to represent the thermodynamic (computed for this work). The DCA results with the size 72 cluster can be considered here to represent the thermodynamic limit. The DCA ($2 \times 2$) data provide an opportunity to assess the relative contribution of the FT-DMET embedding to the finite-size error; in particular, one can compare the difference between FT-DMET and DCA(72) to the difference between DCA ($2 \times 2$) and DCA(72). Overall, we see that the FT-DMET energies with eight bath orbitals are in good agreement with the DCA(72) energies across the different $U$ values, and that the accuracy is slightly better on average than that of DCA ($2 \times 2$). The maximum error in the $I4B8$ impurity compared to thermodynamic limit extrapolations of the DCA energy [35] is found at $U = 4$ and is in the range of 1–2%, comparable to errors observed in ground-state DMET at this cluster size (e.g., the error in GS-DMET at $U = 4$ and $U = 8$ is 0.3% and 1.8%, respectively). In the $\beta = 8$ case, the FT-DMET calculations with two different bath sizes give very similar results; at low temperature, the bath space constructed by the FT procedure is similar to that of the ground state, and the higher-order bath sites do not contribute very relevant degrees of freedom. Thus even the smaller bath achieves good accuracy in the low-temperature FT-DMET calculations.

A central phenomenon in magnetism is the finite-temperature Néel transition. In the thermodynamic limit, the 2D Hubbard model does not exhibit a true Néel transition, but shows a crossover [38]. However, in finite quantum impurity calculations, the crossover appears as a phase transition at a nonzero Néel temperature. Figure 7(a) shows the antiferromagnetic moment $m$ calculated as $m = \frac{1}{2 \pi} \sum_{i} |n_{i \uparrow} - n_{i \downarrow}|$ as a function of temperature $T$ for various $U$ values. As a guide to the eye, we fit the data to a mean-field magnetization function $m = a \tanh (bm/T)$, where $a$ and $b$ are parameters that depend on $U$. The FT-DMET calculations are performed with a $2 \times 2$ impurity and eight bath orbitals, using a finite-temperature DMRG solver with maximal bond dimension $M = 600$ and time step $\tau = 0.1$. With this, the error in $m$ from the solver is estimated to be less than $10^{-3}$, $m$ drops sharply to zero as $T$ is increased signaling a Néel transition. The Néel temperature $T_N$ is taken as the point of intersection of the mean-field fitted line with the $x$ axis; assuming this form of the curve, the uncertainty in $T_N$ is invisible on the scale of the plot. The plot of $T_N$ versus $U$ is shown in Fig. 7(b), showing that the maximal $T_N$ occurs at $U = 6$. Similar $T_N$ calculations on the 2D Hubbard model with single site DMFT [32] and DCA [31,33,34] are also shown in Fig. 7(b) for reference. Note that the difference in the DMFT results [32] and single-site DCA (formally equivalent to DMFT) [33,34] likely arise from the different solvers used. The behavior of $T_N$ in our $2 \times 2$ FT-DMET calculations is quite similar to that of the 4-site DCA cluster. In particular, we see in DCA that the $T_N$ values obtained from calculations with a single-site cluster ($N_c = 1$) are higher than the $T_N$ values obtained from calculations with a 4-site cluster ($N_c = 4$).

An alternative visualization of the Néel transition in FT-DMET is shown in Fig. 8. The FT-DMET calculations here were performed with a $2 \times 2$ impurity and four bath orbitals using an ED solver. Though less quantitatively accurate than the eight bath orbital simulations, these FT-DMET calculations still capture the qualitative behavior of the Néel
transition. Focusing on the dark blue region of the phase diagram, one can estimate the maximal $T_N$ to occur near $U \approx 9$, an increase over the maximal Néel temperature using the eight bath orbital impurity model. This increase in the maximal $T_N$ appears similar to that which happens when moving from a 4-site cluster to 1-site cluster in DCA in Fig. 7.

IV. CONCLUSION

To summarize, we introduced a finite-temperature formulation of the density matrix embedding theory (FT-DMET). This temperature formulation inherits most of the basic structure of the ground-state density matrix embedding theory, but modifies the bath construction so as to approximately reproduce the mean-field finite-temperature density matrix. From numerical assessments on the 1D and 2D Hubbard model, we conclude that the accuracy of FT-DMET is comparable to that of its ground-state counterpart, with at most a modest increase in size of the embedded problem. From the limited comparisons, it also appears to be competitive in accuracy with cluster dynamical mean-field theory for the same sized cluster. Similarly to ground-state DMET, we expect FT-DMET to be broadly applicable to a wide range of model and \textit{ab initio} problems of correlated electrons at finite temperature [18,25].

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APPENDIX: PROOF OF THE FINITE-TEMPERATURE BATH FORMULA

Let $M$ be an arbitrary $N \times N$ full rank square matrix, and $Q_k$ be the $Q$ derived from the QR decomposition of the first $n$ columns of $M^k$, i.e., $M^k[:, : n] = Q_k R_k$, with $k = 0, 1, \ldots, K$. Let $S (|S| < N)$ be a space spanned by $\{Q_0, Q_1, \ldots, Q_K\}$, and $P$ be the projector onto $S$. The following equality holds:

$$P^l M^l P[:, : n] = (P^l M P)^2[:, : n], \quad l \leq K + 1. \quad (A1)$$

We prove the statement by mathematical induction. First write $M$ in the following form

$$M = \begin{bmatrix} A & B \\ C & D \end{bmatrix}, \quad (A2)$$

where $A$ and $B$ are the first $n$ rows of $M$, $A$, and $C$ are the first $n$ columns of $M$. The projector has the form

$$P = \begin{bmatrix} I & 0 \\ 0 & V \end{bmatrix}, \quad (A3)$$

where $I$ is an $n \times n$ matrix, and $V$ is an $(N-n) \times (K-1)n$ matrix with $(K-1)n < (N-n)$. The columns of $V$ are derived from the QR decomposition of $M^k[:, : n]$, $k = 1, \ldots, K$ and then orthogonalized. We can write $V$ in the form

$$V = [V_1 \ V_2 \ \cdots \ V_K], \quad (A4)$$

where $V_k$ is from the QR decomposition of $M^k[:, : n]$. $P^l M P$ has the form

$$P^l M P = \begin{bmatrix} A & B V \\ V^\dagger C & V^\dagger D V \end{bmatrix}. \quad (A5)$$

The mathematical induction consists of two parts.

(i) We start with $l = 2$. The first $n$ columns of $P^l M^2 P$ and $(P^l M P)^2$ are

$$P^l M^2 P[:, : n] = \begin{bmatrix} A^2 + BC \\ V^\dagger C A + V^\dagger D C \end{bmatrix}, \quad (A6)$$

$$(P^l M P)^2[:, : n] = \begin{bmatrix} A^2 + B V V^\dagger C \\ V^\dagger C A + V^\dagger D V V^\dagger C \end{bmatrix}. \quad (A7)$$

The two are equal when

$$V V^\dagger C = V V^\dagger (V R) = VIR = VR = C, \quad (A8)$$

which is true since $V$ is the $Q_1$ from the QR decomposition of $C$. (Note that $V V^\dagger = I$, but $V V^\dagger \neq I$.) Therefore, Eq. (A1) holds for $l = 2$ when $K \geq 1$.

(ii) Now let us inspect Eq. (A1) for the $l$th order, assuming that Eq. (A1) holds for the $(l-1)$th order, i.e., $P^l M^{l-1} P = (P^l M P)^{l-1}$. Let

$$M^{l-1} = \begin{bmatrix} W & X \\ Y & Z \end{bmatrix}, \quad (A9)$$

and $M^l = M M^{l-1}$ has the form

$$M^l = \begin{bmatrix} A W + B Y & A X + B Z \\ C W + D Y & C X + D Z \end{bmatrix}. \quad (A10)$$

and

$$P^l M^{l-1} P = (P^l M P)^{l-1} = \begin{bmatrix} W & X V \\ V Y & V Z V \end{bmatrix}. \quad (A11)$$
One can prove that $CW$ and $C$ share the same $Q$ space from the QR decomposition: let $C = QR$, then $CW = QRW$, where $R$ and $W$ are square matrices; we then perform another QR decomposition of $RW$, $RW = UR$, where $U$ is a unitary matrix, then $CW = QR$ with $Q = QU$. Therefore, $Q$ and $Q$ span the same space.

The first $n$ columns of $(P^T M' P)$ and $(P^T M P)^j$ are

$$P^T M' P[:, n] = \left[ \begin{array}{c} AW + BY \\ V^T CW + V' DY \end{array} \right].$$

Since $V$ contains $V_{l-1}$, which is derived from the QR decomposition of $Y$, we have $VV^T Y = Y$ as in Eq. (A7).

Combining (i) and (ii) we then see that Eq. (A1) holds for the $l$th order with $K \geq l - 1$ for $\forall l$.