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Sum rules and bath parametrization for quantum cluster theories

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We analyze cellular dynamical mean-field theory (CDMFT) and the dynamical cluster approximation (DCA). We derive exact sum-rules for the hybridization functions and give examples for dynamical mean-field theory, CDMFT, and DCA. For impurity solvers based on a Hamiltonian, these sum rules can be used to monitor convergence of the bath-parametrization. We further discuss how the symmetry of the cluster naturally leads to a decomposition of the bath Green matrix into irreducible components, which can be parametrized independently, and give an explicit recipe for finding the optimal bath parametrization. As a benchmark we revisit the one-dimensional Hubbard model. We carefully analyze the evolution of the density as a function of chemical potential and find that, close to the Mott transition, convergence with cluster size is unexpectedly slow. Going from one to two dimensions we find that fitting the bath becomes in general significantly more difficult, requiring a large number of bath sites. For such large baths our symmetry-adapted approach should prove crucial for finding a reliable bath-parametrization.

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I. INTRODUCTION

Strongly correlated materials are characterized by the interplay of kinetic energy and sizable short-range electronic repulsion, which cannot be described with single-particle approaches or standard perturbative theories. The dynamical mean field theory^{1,2} (DMFT) has proven extremely powerful, in particular for the description of the correlation-driven Mott transition. Yet DMFT is strictly local. A number of recent developments are aimed at overcoming this limitation and are greatly contributing to the understanding of the physics of strongly correlated systems.³⁻⁹ In quantum cluster theories⁷ the k dependence is introduced by considering a small number of sites, instead of the single correlated site of DMFT, and embedding them in a bath, i.e., a dynamical mean-field host determined self-consistently. This is a good approximation when the self-energy is reasonably localized, which can be systematically improved by considering larger and larger clusters.¹⁰

Cluster extensions of DMFT are not unique. Here we consider the two main flavors, cellular dynamical mean-field theory⁵ (CDMFT) and the dynamical cluster approximation³ (DCA). Finite temperature Monte Carlo provides an efficient cluster solver, however, to obtain spectra on the realfrequency axis, data for imaginary time must be analytically continued, usually applying maximum entropy. If the spectrum has structures on a small energy scale, this approach can lead to problems. It is then natural to explore Hamiltonian-based solvers, like the Lanczos method, which directly give result on the real-frequency axis.

Lanczos is extensively applied to DMFT and results are reliable and very accurate. The critical step in such calculations is the fitting of the bath degrees-of-freedom. In cluster methods this step gets more involved. This is one of the reasons why it found few applications for cluster methods: To the best of our knowledge, it has so far only been used for CDMFT calculations of very small clusters. Our aim is therefore to give a systematic formulation, exploiting symmetries to arrive at an optimal parametrization of the bath. In addition we derive exact sum rules for the bath Green's functions. As an application, we investigate the one-dimensional Hubbard model with CDMFT in detail,^{11–13} paying particular attention to the convergence with the size of both bath and cluster.

The paper is organized as follows: To fix the notation, in Sec. II, we give a unified formulation of CDMFT and DCA in a formalism using a Hamiltonian solver. In Sec. III we derive general sum rules for the hybridizations and give examples for DMFT, CDMFT, and DCA. In Sec. IV we discuss how the symmetry of the cluster naturally leads to a decomposition of the bath Green matrix into irreducible components, which can be parametrized independently. We give an explicit recipe for finding the optimal bath-parametrization and discuss how this approach relates to the technique of cluster replica. In Sec. V we analyze how the cluster approaches work for the Hubbard chain and carefully reexamine the evolution of the density as a function of chemical potential. In Sec. VI we give our conclusions and an outlook.

II. METHOD AND NOTATION

We consider the Hubbard model

$$H = -\sum_{ij\sigma} t_{ij} c_{i\sigma}^{\dagger} c_{j\sigma} + U \sum_{i} n_{i\uparrow} n_{i\downarrow}.$$
(1)

To fix the notation we briefly sketch the self-consistency loop for cellular DMFT and the dynamical cluster approximation using, e.g., exact diagonalization as impurity solver. Let N_c be the number of cluster sites, N_b the number of bath sites. For simplicity we suppress spin indices.

Given an $N_c \times N_c$ bath Green matrix \mathcal{G}^{-1} ,

(1) Fit parameters of an Anderson model with N_b bath sites

$$\boldsymbol{\mathcal{G}}_{And}^{-1}(\boldsymbol{\omega}) \approx \boldsymbol{\omega} + \boldsymbol{\mu} - \mathbf{H}_c - \boldsymbol{\Gamma}[\boldsymbol{\omega} - \mathbf{E}]^{-1} \boldsymbol{\Gamma}^{\dagger}$$
(2)

to \mathcal{G}^{-1} , where Γ is the $N_c \times N_b$ -dimensional hybridization matrix, and **E** the $N_b \times N_b$ -dimensional bath-matrix. **H**_c is specified below,

(2) Solve the $N_c + N_b$ -site Anderson model H_{And} (specified below) to obtain the $N_c \times N_c$ cluster Green matrix \mathbf{G}_c ,

(3) Get the cluster self-energy matrix

$$\boldsymbol{\Sigma}_{c}(\boldsymbol{\omega}) = \boldsymbol{\mathcal{G}}^{-1}(\boldsymbol{\omega}) - \mathbf{G}_{c}^{-1}(\boldsymbol{\omega}), \qquad (3)$$

(4) Calculate the local Green matrix for the cluster by integrating over the reduced Brillouin-zone of the cluster

$$\mathbf{G}(\boldsymbol{\omega}) = \int d\widetilde{\mathbf{k}} [\boldsymbol{\omega} + \boldsymbol{\mu} - \mathbf{H}(\widetilde{\mathbf{k}}) - \boldsymbol{\Sigma}_{c}(\boldsymbol{\omega})]^{-1}, \qquad (4)$$

where $\mathbf{H}(\tilde{\mathbf{k}})$ is the single-electron part of the of the Hubbard Hamiltonian (1) in the reduced Brillouin-zone of the cluster,

(5) Determine the new bath Green matrix (self-consistency condition)

$$\mathcal{G}^{-1}(\omega) = \Sigma_c(\omega) + \mathbf{G}^{-1}(\omega).$$
 (5)

These steps are iterated to self-consistency.

The Anderson model to be solved in step 2 is given by

$$H_{\text{And}} = H_{\text{clu}} + \sum_{lm,\sigma} E_{lm,\sigma} a_{l\sigma}^{\dagger} a_{m\sigma} + \sum_{li,\sigma} \Gamma_{il} (a_{l\sigma}^{\dagger} c_{i\sigma} + \text{H.c.}), \quad (6)$$

where the operator $a_{l\sigma}^{\dagger}$ creates an electron of spin σ on bathsite *l*. The cluster Hamiltonian H_{clu} is obtained from the original Hamiltonian (1) by transforming to the reciprocal space of the superlattice of clusters, and projecting to the cluster. Writing the single-electron part of $H(\tilde{\mathbf{k}})$ as the matrix $\mathbf{H}(\tilde{\mathbf{k}})$, the single-electron part of H_{clu} is given by

$$\mathbf{H}_{c} = \int d\tilde{\mathbf{k}} \mathbf{H}(\tilde{\mathbf{k}}). \tag{7}$$

The interaction terms are simply those of Eq. (1), restricted to the cluster.

The Hamiltonian $H(\mathbf{k})$ in the reciprocal space of the superlattice $\{\tilde{\mathbf{r}}\}\$ of clusters can be obtained by changing to the basis of operators

$$\tilde{c}_{\mathbf{R}_{i},\sigma}^{\text{CDMFT}}(\tilde{\mathbf{k}}) = \sum_{\tilde{\mathbf{r}}} e^{-i\tilde{\mathbf{k}}\tilde{\mathbf{r}}} c_{\tilde{\mathbf{r}}+\mathbf{R}_{i},\sigma}.$$
(8)

The resulting quantum cluster approximation is CDMFT. Alternatively, we can start from the operators in the reciprocal space of the *lattice* to obtain

$$\tilde{c}_{\mathbf{R}_{i},\sigma}^{\mathrm{DCA}}(\tilde{\mathbf{k}}) = \sum_{\tilde{\mathbf{r}}} e^{-i\tilde{\mathbf{k}}(\tilde{\mathbf{r}}+\mathbf{R}_{i})} c_{\tilde{\mathbf{r}}+\mathbf{R}_{i},\sigma}.$$
(9)

Now we obtain the DCA. The choice of the operators in the two approaches differs just by local phase factors. In CDMFT this gauge¹⁴ is chosen such that phases appear only in matrix elements involving different clusters. Thus all matrix elements on the cluster are the same as in the original Hamiltonian. The price for retaining the original matrix elements on the cluster is a breaking of the translation symmetry of the original lattice. DCA opts instead to retain this

symmetry by distributing the phase change uniformly over the cluster sites. The price for retaining translation invariance is that the matrix elements in the cluster Hamiltonian differ from those in the original Hamiltonian (coarse graining). In both cases, CDMFT and DCA, the eigenvalues of $\mathbf{H}(\mathbf{\tilde{k}})$ are identical to the eigenvalues of the noninteracting part of *H*

III. HYBRIDIZATION SUM-RULES

While the most general parametrization for the bath is given by expression (2),^{4,11} we can always diagonalize the hopping matrix **E** among the bath sites to obtain

$$\boldsymbol{\mathcal{G}}_{\text{And}}^{-1}(\{\boldsymbol{\varepsilon}_{l}, \mathbf{V}_{l}\}; \boldsymbol{\omega}) = \boldsymbol{\omega} + \boldsymbol{\mu} - \mathbf{H}_{c} - \sum_{l} \frac{\mathbf{V}_{l} \mathbf{V}_{l}^{\dagger}}{\boldsymbol{\omega} - \boldsymbol{\varepsilon}_{l}}.$$
 (10)

The hybridization matrix is then given by the tensor product of the vectors \mathbf{V}_{l} , where

$$V_{l,i} = \sum_{m} \Gamma_{i,m} \phi_{l,m} \tag{11}$$

and ϕ_l are the eigenvectors of **E** with eigenvalues ε_l .

To obtain sum-rules for the hybridizations, we write the inverse of the bath Green matrix as

$$\boldsymbol{\mathcal{G}}^{-1}(\boldsymbol{\omega}) = \boldsymbol{\Sigma}_{c}(\boldsymbol{\omega}) + \left\{ \int d\tilde{\mathbf{k}} [\boldsymbol{\omega} + \boldsymbol{\mu} - \mathbf{H}(\tilde{\mathbf{k}}) - \boldsymbol{\Sigma}_{c}(\boldsymbol{\omega})]^{-1} \right\}^{-1}.$$

Considering the limit $\omega \rightarrow \infty$, expanding to order $1/\omega^2$, using Eq. (7), and comparing to Eq. (10) we find

$$\sum_{l} \mathbf{V}_{l} \mathbf{V}_{l}^{\dagger} = \int d\tilde{\mathbf{k}} \mathbf{H}^{2}(\tilde{\mathbf{k}}) - \left[\int d\tilde{\mathbf{k}} \mathbf{H}(\tilde{\mathbf{k}}) \right]^{2}.$$
 (12)

To illustrate this hybridization sum-rule we consider a representative set of examples.

A. Single site

We consider a *d*-dimensional lattice with hoppings t_n to the z_n *n*th-nearest neighbors. For $N_c=1$ we have $\mathbf{H}(\mathbf{k}) = \varepsilon_{\mathbf{k}}$. Thus we find for the hybridizations

$$\sum_{l} V_l^2 = \frac{1}{(2\pi)^d} \int d\mathbf{k} \varepsilon_{\mathbf{k}}^2 = \sum_{n} z_n t_n^2,$$
(13)

where the integral is just the second moment of the density of states, so that the last equation follows as in the recursion method.¹⁵ For a Bethe lattice of connectivity z with hopping matrix element t/\sqrt{z} the sum-rule reduces to $\sum_l V_l^2 = t^2$.

B. CDMFT

We start by considering a linear chain with nearestneighbor hopping t and a three-site cluster $N_c=3$. In the CDMFT gauge we have

$$\mathbf{H}(\tilde{k}) = -t \begin{pmatrix} 0 & 1 & e^{-3i\tilde{k}} \\ 1 & 0 & 1 \\ e^{3i\tilde{k}} & 1 & 0 \end{pmatrix}$$
(14)

so that \mathbf{H}_c is the original single-electron Hamiltonian restricted to the cluster:



FIG. 1. CDMFT sum-rules for a one-dimensional three-site cluster and nearest and next-nearest-neighbor hoppings t and t", respectively: (a) $\Sigma_l |V_{l,1}|^2 = t^2 + t''^2$, (b) $\Sigma_l \overline{V}_{l,1} V_{l,2} = tt''$, and (c) $\Sigma_l |V_{l,2}|^2 = 2t''^2$. The hybridizations are given by the two-step hopping processes that are lost when cutting the cluster out of the original lattice.

$$\mathbf{H}_{c} = \frac{3}{2\pi} \int_{-\pi/3}^{\pi/3} d\tilde{k} \mathbf{H}(\tilde{k}) = -t \begin{pmatrix} 0 & 1 & 0 \\ 1 & 0 & 1 \\ 0 & 1 & 0 \end{pmatrix}.$$
 (15)

The sum-rule (12) then is

$$\left(\sum_{l} V_{l,i} \overline{V}_{l,j}\right) = \begin{pmatrix} t^2 & 0 & 0\\ 0 & 0 & 0\\ 0 & 0 & t^2 \end{pmatrix},$$
 (16)

i.e., only the sites on the surface of the cluster couple to the bath.

The general CDMFT hybridization sum-rule (12) can be easily visualized. The integral over the Brillouin-zone of the cluster projects the single-electron part of the full Hamiltonian to the cluster [see Eq. (7)]. The matrix elements of \mathbf{H}_c^2 are thus the two-step hoppings that are possible on the cluster. Likewise the integral over the Hamiltonian squared gives the second moments, only that here the intermediate site is not restricted to the cluster. Thus the sum-rule matrix is given by the second-order paths between cluster sites that proceed via a site outside the cluster. This is illustrated in Fig. 1. As a special case, for a single site we obtain the second equality in Eq. (13).

The vanishing of a matrix element in the sum-rule only implies that the corresponding matrix element of the bath Green matrix decays faster than $1/\omega$ for large ω . For a diagonal element, however, all terms in $\sum_{l} V_{l,i} \bar{V}_{l,i}$ are positive. Thus a vanishing sum means that all terms must be zero. Hence the sum-rule implies that cluster sites that are so far in the interior that they cannot be reached by hopping from outside the cluster do not couple to bath and that all matrix elements of the bath Green function involving such a site *i* are given by $\mathcal{G}_{ij}^{-1}(\omega) = \omega + \mu - (\mathbf{H}_c)_{ij}$ for all ω . In that sense the bath hybridizes only to the surface of the cluster and we see that the hybridization strength to these sites does not decrease for increasing cluster size N_c .

C. DCA

We start again by considering the three-site cluster. In the DCA gauge we write

$$\mathbf{H}(\tilde{k}) = -t \begin{pmatrix} 0 & e^{i\tilde{k}} & e^{-i\tilde{k}} \\ e^{-i\tilde{k}} & 0 & e^{i\tilde{k}} \\ e^{i\tilde{k}} & e^{-i\tilde{k}} & 0 \end{pmatrix}.$$
 (17)

Now \mathbf{H}_{c} has translation symmetry, but the hopping matrix element is rescaled by $\sin(\pi/N_{c})/(\pi/N_{c})$:

$$\mathbf{H}_{c} = \frac{3}{2\pi} \int_{-\pi/3}^{\pi/3} d\tilde{k} \mathbf{H}(\tilde{k}) = -\frac{3\sqrt{3}}{2\pi} t \begin{pmatrix} 0 & 1 & 1\\ 1 & 0 & 1\\ 1 & 1 & 0 \end{pmatrix}.$$
 (18)

Since all matrices in Eq. (12) are periodic, it is convenient to transform to k space. With $V_{l,K}=\sum_i V_{l,i}e^{iKr_i}/\sqrt{N_c}$ and the coarse-graining factor $\tau=3\sqrt{3}/2\pi$ we find

$$\sum_{l} |V_{l,K=0}|^2 = (2 + \tau - 4\tau^2)t^2.$$
$$\sum_{l} |V_{l,K=\pm 2\pi/3}|^2 = (2 - \tau/2 - \tau^2)t^2.$$

The hybridization sum-rule (12) is then, likewise, diagonal in the cluster-momenta **K**

$$\sum_{l} |V_{l,\mathbf{K}}|^2 = \int d\tilde{\mathbf{k}} \varepsilon_{\mathbf{K}+\tilde{\mathbf{k}}}^2 - \left(\int d\tilde{\mathbf{k}} \varepsilon_{\mathbf{K}+\tilde{\mathbf{k}}}\right)^2, \quad (19)$$

while all terms $V_{l,\mathbf{K}}\overline{V}_{l,\mathbf{K}'}$ mixing different cluster momenta vanish. As a special case, for a single site the above sum-rule is just the first equality in Eq. (13). Expanding $\varepsilon_{\mathbf{K}+\mathbf{k}}$ around **K**, we find that for a *d*-dimensional system $\Sigma_l |V_{l,\mathbf{K}}|^2$ decreases with cluster size as $1/N_c^{2/d}$, while all cluster-sites couple with the same strength to the bath.¹⁶

D. Discussion

From the sum-rules we recover⁷ that the individual hybridizations in CDMFT are independent of cluster size, while for DCA they decrease with cluster size as $N_c^{-2/d}$. Interestingly this means that for a *d*-dimensional system in CDMFT the overall coupling to the bath scales as $N_c^{(d-1)/d}$, while in DCA it scales as $N_c^{(d-2)/d}$. For nonlocal properties a DCA calculation is therefore expected to converge faster with cluster size.¹⁷ For a calculation where we represent the bath by discrete degrees of freedom this decrease in hybridization strength does, however, not help very much as we still need bath sites to fit the hybridizations, even if they are small. With increasing DCA cluster size we thus have to parametrize N_c baths, one for each **K**. In CDMFT the situation is more fortunate, as the sum-rules imply that many hybridizations surface sites to the bath.

The lack of translational invariance in CDMFT has two important practical implications. First, the full Green matrix has to be calculated, instead of just its diagonal. Second, when calculating local quantities, like the density per site, in CDMFT we have a choice of inequivalent sites, or we could consider the average over all sites. In a gapped system the best choice is the innermost site,¹⁸ however, in such a situation it might be better to do a straight Lanczos calculation with N_c+N_b cluster sites, instead of using N_b bath sites.¹⁹

IV. SYMMETRIES

In the absence of spontaneous symmetry breaking the symmetries of the cluster (point symmetries in CDMFT and additionally translation symmetry in DCA) are reflected in the Green matrix. In a symmetry broken state with longrange order, such as an antiferromagnet or a charge-density wave, the symmetry of the Green matrix is accordingly lowered. To exploit this symmetry we introduce vectors on the cluster that transform according to its irreducible representations. We write these vectors as $\mathbf{w}_{I,\nu}$ where I is the irreducible representation and $\nu = 1 \dots N_I$ counts linear independent vectors transforming according to I. On an N_c -site cluster we can choose N_c such vectors that are orthonormal. Defining the matrix $\mathbf{W} = (\mathbf{w}_{I\nu})$ of these vectors, we can block diagonalize the bath Green matrix: $\mathbf{W}^{\dagger} \boldsymbol{\mathcal{G}}^{-1} \mathbf{W}$ has blocks of dimension N_I corresponding to the different irreducible representations I. Since $\mathbf{W}^{\dagger} \mathcal{G}^{-1} \mathbf{W}$ is block diagonal for all ω , it follows from Eq. (10), that W must also block diagonalize the individual hybridization matrices $\mathbf{V}_{l}\mathbf{V}_{l}^{\dagger}$. Therefore the hybridization vectors must transform according to an irreducible representation; they can be written as $\mathbf{V}_{l} = \sum_{\nu} V_{l,l,\nu} \mathbf{w}_{l,\nu}$ for some irreducible representation I. If the V_l also had components $\mathbf{w}_{J,\nu}$ of a different irreducible representation $J \neq I$ this would produce a hybridization matrix that could not be block diagonalized.

We thus find that the bath sites can be arranged into sets corresponding to the different irreducible representations. For fitting a block of the symmetrized bath Green matrix we need then only consider bath sites of the respective irreducible representation. If the block is one dimensional we can choose the corresponding hybridization real. An early example is the bonding-antibonding transformation introduced in Ref. 20.

A. CDMFT

As an example we consider a linear cluster of 3 sites as shown in Fig. 2. The symmetry is C_2 (see Table I). Transforming to the basis vectors $\mathbf{w}_{A,1} = (|1\rangle + |3\rangle)/\sqrt{2}$ and $\mathbf{w}_{A,2}$ = $|2\rangle$ of symmetry A (see Table I) and $\mathbf{w}_B = (|1\rangle - |3\rangle)/\sqrt{2}$, we find the transformed bath Green matrix

$$\mathbf{W}^{\dagger} \boldsymbol{\mathcal{G}}^{-1} \mathbf{W} = \begin{pmatrix} \boldsymbol{\mathcal{G}}_{11}^{-1} + \boldsymbol{\mathcal{G}}_{13}^{-1} & \sqrt{2} \boldsymbol{\mathcal{G}}_{12}^{-1} & 0\\ \sqrt{2} \boldsymbol{\mathcal{G}}_{21}^{-1} & \boldsymbol{\mathcal{G}}_{22}^{-1} & 0\\ 0 & 0 & \boldsymbol{\mathcal{G}}_{11}^{-1} - \boldsymbol{\mathcal{G}}_{13}^{-1} \end{pmatrix}.$$

A bath site of irreducible representation A contributes to the first block and has the same hybridization $V_{A,1}$ to the outer cluster-sites plus an independent hybridization parameter $V_{A,2}$ to the central site. A bath site of irreducible representation B contributes to the second block. For such a bath site



FIG. 2. Hybridization of bath-sites of symmetry A and B to a three-site cluster. As defined in Table I, A is the unit representation, so a bath site of type A has the same hybridization V to all cluster sites that are equivalent by symmetry. B is the antisymmetric representation, so the hybridization of a bath site of type B to cluster sites that are related by mirror symmetry have the opposite sign. Consequently the hybridization to the central site of a linear cluster with an odd number of sites vanishes in the B representation.

the hybridization to cluster sites that are related by mirror symmetry have opposite signs. Consequently, the hybridization to the central site vanishes.

The situation is slightly more complicated when the symmetry group has irreducible representations of dimension higher than one. The simplest example is the 2×2 cluster with C_{4v} symmetry. With $\mathbf{w}_{A_1} = (|1\rangle + |2\rangle + |3\rangle + |4\rangle)/2$, $\mathbf{w}_{B_2} = (|1\rangle - |2\rangle + |3\rangle - |4\rangle)/2$, and the pair $\mathbf{w}_{E,1} = (|1\rangle - |2\rangle - |3\rangle + |4\rangle)/2$, $\mathbf{w}_{E,2} = (|1\rangle + |2\rangle - |3\rangle - |4\rangle)/2$ we find that $\mathbf{W}^{\dagger} \mathcal{G}^{-1} \mathbf{W}$ is diagonal with diagonal elements

$$(\mathbf{W}^{\dagger} \boldsymbol{\mathcal{G}}^{-1} \mathbf{W})_{11} = \boldsymbol{\mathcal{G}}_{11}^{-1} + 2\boldsymbol{\mathcal{G}}_{12}^{-1} + \boldsymbol{\mathcal{G}}_{13}^{-1}$$
$$(\mathbf{W}^{\dagger} \boldsymbol{\mathcal{G}}^{-1} \mathbf{W})_{22} = \boldsymbol{\mathcal{G}}_{11}^{-1} - 2\boldsymbol{\mathcal{G}}_{12}^{-1} + \boldsymbol{\mathcal{G}}_{13}^{-1}$$
$$(\mathbf{W}^{\dagger} \boldsymbol{\mathcal{G}}^{-1} \mathbf{W})_{33} = \boldsymbol{\mathcal{G}}_{11}^{-1} - \boldsymbol{\mathcal{G}}_{13}^{-1}$$
$$(\mathbf{W}^{\dagger} \boldsymbol{\mathcal{G}}^{-1} \mathbf{W})_{44} = \boldsymbol{\mathcal{G}}_{11}^{-1} - \boldsymbol{\mathcal{G}}_{13}^{-1}.$$

A bath site of symmetry A_1 has the same hybridization to all cluster sites while for a bath-site of symmetry B_2 the hybridizations have alternating signs: $\mathbf{V}_l^{\dagger} = \overline{V}_l(1, -1, 1, -1)$. To realize the two-dimensional representation E we need two bath

TABLE I. Character tables of the point groups C_2 , C_{3v} , and C_{4v} .

	C_2	$E \sigma_v$	_	~	_		~1		
	A	1 1		C_{4v}	E	$2C_4$	C_{4}^{2}	$2\sigma_v$	$2\sigma_d$
	В	1 -1		A_1	1	1	1	1	1
		I		A_2	1	1	1	-1	-1
C_3	v E	$E 2C_3$	$3\sigma_v$	B_1	1	-1	1	1	-1
A_1	1	l 1	1	B_2	1	-1	1	-1	1
A_2	2 1	l 1	-1	E	2	0	-2	0	0
E	4	2 -1	0						



FIG. 3. Hybridization of bath sites of symmetry A_1 , B_2 , and E to a 2×2 cluster. For a given irreducible representation the absolute value of the hybridization to all cluster-sites is the same, while the signs are indicated in the figure. Nontrivial hybridizations corresponding to irreducible representations A_2 or B_1 only appear for larger clusters.

sites l_1 and l_2 with degenerate energies $\varepsilon_{l_1} = \varepsilon_{l_2} = \varepsilon_l$ and hybridizations: $\mathbf{V}_{l_1}^{\dagger} = \overline{V}_l(1, -1, -1, 1)$ and $\mathbf{V}_{l_2}^{\dagger} = \overline{V}_l(1, 1, -1, -1)$. This is illustrated in Fig. 3.

B. DCA

As an example for DCA we consider a three-site cluster with periodic boundary conditions. The symmetry group is C_{3v} (translations and inversion). Hence we introduce the basis vector $\mathbf{w}_{A_1} = (|1\rangle + |2\rangle + |3\rangle)/\sqrt{3}$, corresponding to k=0, while the vectors formed by $\sin(2\pi/3)$ and $\cos(2\pi/3)$ give the *E* representation: $\mathbf{w}_{E,1} = (|1\rangle - |2\rangle)/\sqrt{2}$ and $\mathbf{w}_{E,2} = (|1\rangle + |2\rangle - 2|3\rangle)/\sqrt{6}$

$$\mathbf{W}^{\dagger} \boldsymbol{\mathcal{G}}^{-1} \mathbf{W} = \begin{pmatrix} \boldsymbol{\mathcal{G}}_{11}^{-1} + 2\boldsymbol{\mathcal{G}}_{12}^{-1} & 0 & 0 \\ 0 & \boldsymbol{\mathcal{G}}_{11}^{-1} - \boldsymbol{\mathcal{G}}_{12}^{-1} & 0 \\ 0 & 0 & \boldsymbol{\mathcal{G}}_{11}^{-1} - \boldsymbol{\mathcal{G}}_{12}^{-1} \end{pmatrix}.$$

In general bath sites corresponding to the gamma point have the same hybridization to all cluster sites, while those corresponding to $k=\pi$ have alternating hybridizations. For all other *k* points we need two degenerate bath sites, with hybridizations $V_{l_1,\mu} = V_l \sin(k\mu)$ and $V_{l_2,\mu} = V_l \cos(k\mu)$ to cluster site μ .

C. Cluster replica

Instead of implementing the symmetry of the Green matrix as described above, one might construct the bath out of replica of the N_c -site clusters.¹³ For a two-site cluster this means that bath sites come in pairs, with on-site energy $\tilde{\varepsilon}$, hopping $-\tilde{t}$ between the bath-sites, and hybridization \tilde{V}_{11} and \tilde{V}_{12} to the cluster as illustrated in Fig. 4. Diagonalizing such a bath pair, we obtain one bath site of symmetry A with on-site energy $\varepsilon_A = \tilde{\varepsilon} - \tilde{t}$ and hybridization $V_A = (\tilde{V}_{11} + \tilde{V}_{12})/\sqrt{2}$ and one bath site of symmetry B with $\varepsilon_B = \tilde{\varepsilon} + t$ and $V_B = (\tilde{V}_{11} - \tilde{V}_{12})/\sqrt{2}$.

To generalize this approach, let \tilde{H}_b be the Hamiltonian for an N_c -site cluster with general on-site energies and hoppings that respect the symmetry of the Green matrix. Furthermore let bath site *n* hybridize to cluster site *i* with \tilde{V}_{ni} . These hybridizations are chosen symmetric under simultaneous



FIG. 4. Replica of a two-site cluster in the bath. Diagonalizing the Hamiltonian for the two bath sites leads to a bath-site of type A (even representation) with energy $\varepsilon_A = \tilde{\varepsilon} - \tilde{t}$ and hybridization V_A $= (\tilde{V}_{11} + \tilde{V}_{12})/\sqrt{2}$ and a site of type B with $\varepsilon_B = \tilde{\varepsilon} + t$ and $V_B = (\tilde{V}_{11} - \tilde{V}_{12})/\sqrt{2}$.

symmetry transformations of the original cluster and the bath replica. Diagonalizing \tilde{H}_b we obtain N_c eigenstates φ_l with energy ε_l . These can be considered as bath sites that hybridize to cluster site *i* with $V_{l,i}=\sum_n \varphi_{l,n} \tilde{V}_{n,i}$.

Being the eigenstates of the Hamiltonian \tilde{H}_{h} , the φ_{l} transform according to the irreducible representations of the symmetry of the Green matrix.²¹ Therefore they can be written as linear combination $\varphi_l = \sum_{\nu} \alpha_{l;l,\nu} \mathbf{w}_{l,\nu}$ for some irreducible representation I. From this we can conclude that a cluster replica gives rise to N_I bath sites of symmetry I. By working with cluster replica we thus sacrifice the freedom of choosing the irreducible representations for the bath individually. Moreover, it is not straightforward to find a proper parametrization. \tilde{H}_b must be chosen such that all accidental degeneracies can be lifted. For symmetries with higher-dimensional irreducible representations there will be, however, corresponding essential degeneracies. Moreover, working with cluster replica, we cannot fit the individual blocks of the bath Green matrix with the minimal set of symmetry-adapted parameters, but have to solve the optimization problem for the full bath Green matrix and all parameters. Thus, using cluster replica is less flexible than using individual irreducible representations and it leads to a more complicated fitting procedure, in particular when considering large baths.

V. HUBBARD CHAIN

We now discuss the one-dimensional Hubbard model, for which exact results are available from the Bethe ansatz²² and which has been studied with CDMFT using Lanczos^{11–13} and quantum Monte Carlo,²³ as well as with the variational cluster approximation.²⁴ Here going from single-site DMFT to a cluster description makes a qualitative difference; for a paramagnetic single-site calculation antiferromagnetism is completely suppressed, while on a cluster we will have short-ranged antiferromagnetic correlations, even if we impose a paramagnetic bath.

This inclusion of antiferromagnetic correlations might well be the cause for the spectacular difference between the single-site and two-site CDMFT calculations of the density as a function chemical potential reported in Ref. 12. For illustration, in Fig. 5, we compare the density as a function



FIG. 5. (Color online) Density as a function of chemical potential for a Hubbard model on the Bethe lattice with half band width D and U=2D in single-site DMFT with paramagnetic (para) and antiferromagnetic (AF) bath (blue squares and red circles, respectively). The AF curve comprises a left and a right branch: the left one has been obtained by fixing the magnetization from 0.01 to 0.82 (from left to right) and determining the chemical potential. The right branch has been obtained by just decreasing the chemical potential from the half-filling value ($\mu = U/2$) down to the point where convergence is no longer found ($\mu = 0.6$). The magnetization in the right branch is about 0.85. In the inset we compare the total energy of the AF and para phases as a function of doping. The AF phase is the stable one up to about $\delta = 0.2$. We have added $\delta U/2 - E_{AF}(0)$ to $E(\delta)$ to allow better comparison with Fig. 1 of Ref. 26 and Fig. 8 of Ref. 27.

of the chemical potential for a single-site calculation with paramagnetic and antiferromagnetic bath.²⁵ They represent two limiting cases, the former being always metallic, the latter yielding a gap, which is overestimated as we are using an antiferromagnetic bath to mimic the short-ranged correlations present in the Hubbard chain.

A. Bath Green matrix

For a one-dimensional lattice the CDMFT bath Green matrix simplifies drastically: When removing the N_c -site cluster from the lattice, we are left with two disconnected pieces. Thus a vanishing sum-rule (12) in one dimension means that the hybridization matrix vanishes for all frequencies. For the irreducible representations of the bath this means that bath sites of symmetry A and B come in degenerate pairs: $\varepsilon_{A_l} = \varepsilon_{B_l}$ and $V_{A_l} = V_{B_l}$. Thus for the Hubbard chain with nearestneighbor hopping only the outer-most cluster sites hybridize with the bath and the bath parametrizations are identical by symmetry.¹³ The evolution of the bath Green matrix element \mathcal{G}_{11}^{-1} with cluster size is shown in Fig. 6. We find that the bath Green matrix elements hardly depend on cluster size and even for a single-site calculation the bath is already similar to that for a large cluster.

In contrast, in the DCA we get a nonvanishing hybridization for each *K* point of the cluster. This is shown in Fig. 7. While the hybridization strength per *K* point decreases with cluster size, we still have to parametrize all of them, possibly except for K=0 and π , which almost vanish already for moderate cluster sizes.



FIG. 6. (Color online) Plot of the CDMFT bath Green function $\mathcal{G}_{11}^{-1}(\omega) - (\omega + \mu)$ on the real axis for linear clusters with nearestneighbor hopping *t*. Increasing the size of the cluster the hybridization hardly changes. The plots show calculations for chains with U=6t and $\mu=0.5t$.

B. Fitting the bath Green matrix

The most critical step in calculations with a finite bath is the determination of the parameters for the impurity Hamiltonian (6). This is usually done by fitting the bath Green function on the imaginary axis; a fictitious temperature $1/\beta$ is introduced and the sum of the squared difference between \mathcal{G} and its parametrized version [Eq. (10)] over the Matsubara frequencies up to some cutoff is minimized.² This procedure is fairly robust if the number of bath sites is sufficiently large. For cluster calculations the number of effective bath sites per fitted bath Green function can, however, be quite small. In such a situation details of the fitting procedure are important and are accordingly discussed in the literature.^{11–13,28–30}

To fit the Anderson parameters V_l and ε_l we use the distance function

$$\sum_{<\omega_n<\omega_{\rm cut}}\frac{\|\mathbf{W}^{\dagger}\{\boldsymbol{\mathcal{G}}^{-1}(i\omega_n)-\boldsymbol{\mathcal{G}}_{\rm And}^{-1}[(\varepsilon_l,\mathbf{V}_l);i\omega_n]\}\mathbf{W}\|_{I}}{\omega_n^N},$$

where $\|\cdot\|_{l}$ is the 1-norm for the block of irreducible representation I and N determines how strongly large Matsubara frequencies ω_n are weighted. The distance function will only be finite in the limit $\omega_{\rm cut} \rightarrow \infty$ if the sum-rules are fulfilled exactly, or if $N \ge 2$. Thus for N < 2 increasing the cutoff emphasizes the sum-rules. The same is true for decreasing β . If the distance function is dominated by the large frequency asymptotics, the optimization mainly focuses on just the sum-rules. This means that in practice the Anderson parameters can become under determined. Since the fictitious temperature is only used for fitting, while the calculations are actually for T=0 there is a strong dependence of the physical quantities on the Anderson parameters. Hence for small β the self-consistent results strongly differ for different initial Anderson parameters. This is illustrated in Fig. 8. Interestingly, the situation is opposite to calculation at finite temperature, where at higher temperature the physical quantities become less dependent on the details of the fitting.³¹

0<



FIG. 7. (Color online) Plot of the DCA bath Green functions $\mathcal{G}_{K}^{-1}(\omega) - (\omega + \mu - \int d\tilde{k}\varepsilon_{K+\tilde{k}})$ on the real axis for linear clusters with nearest-neighbor hopping *t*. Increasing cluster size ($N_c=2$, 4, 6, and 8 from top to bottom) the number of independent functions to fit increases. Note that K=0 and $K=\pi$ are associated with single bath-sites, while all the other *K* points need to be described by pairs of bath sites. From the figures it is clear how the total spectral weight decreases with increasing N_c . The plots show calculations for chains with U=6t and $\mu=0.5t$.

To avoid an under determination of the Anderson parameters, it is important to ensure that the features of the bath Green matrix close to the real axis are properly weighted in the distance function. This was already pointed out in Ref. 12. We find that a good compromise between fitting the large- and small-frequency behavior is given by $\beta = 256/t$, N=1, and $\omega_{cut} = 200t$, which, if not explicitly specified otherwise, is used in the calculations reported in this work.

C. Convergence with number of bath-sites

To check how many bath sites we need to reach a satisfactory fit of the bath Green matrix, we consider a two-site cluster for increasing N_b . As example we show in Fig. 9 the density for μ =0. For CDMFT we find that we need at least eight bath sites to obtain a converged density. Also for DCA we obtain convergence for N_b =8. Both these results translate to four bath sites per nonvanishing element of the bath Green matrix. It is interesting to note that DCA converges to a density above the Bethe-ansatz result. This could be an artifact of the two-site cluster, for which the coarse-grained hopping is larger than *t*, because for periodic boundary conditions the hopping on the cluster and across the boundary add up. Averaging over different choices of boundary conditions



FIG. 8. (Color online) Density for a two-site cluster with U = 4t and $\mu = 0$ in CDMFT (left panel) and DCA (right panel) calculations with 12 bath sites using different fictitious temperatures $1/\beta$ but constant cutoff $\omega_{cut} \approx 200t$ and N=0 and 1, starting from different Anderson parameters. For low fictitious temperature results are fairly independent of the starting points, while for larger temperature the Anderson parameters are essentially under determined by the distance function and consequently the results of the supposedly self-consistent calculation strongly depend on the initial values.



FIG. 9. (Color online) Density as a function of the number of bath-sites N_b for a two-site cluster for U=4t and $\mu=0$. To give a measure of the reproducibility of the results, we plot the densities of several converged CDMFT and DCA runs (multiple symbols at the same N_b).

or going to a larger cluster might improve the situation.³²

Checking the hybridization sum-rule for the diagonal elements of the bath Green matrix, we find that the density is already converged while the sum of the hybridizations $\Sigma_l |V_l|^2$ is only at about 70% of its exact value. As shown in Fig. 10 we need to go to even larger baths to properly fulfill the sum-rule. We tried also to fit the bath Green matrix imposing the sum-rule, i.e., fixing one Anderson parameter per irreducible representation. We found, however, that for small N_b this does not give particularly good results, as it weights the large frequency behavior of the bath Green matrix too strongly, while for large N_b it is not necessary.

Going to larger clusters, we expect that we will need more bath sites for a converged calculation. A notable exception is CDMFT for the linear chain with nearest-neighbor hopping only. As discussed above, in this case there are two identical baths coupling to one surface site each. Since the bath Green function that these baths have to fit is fairly independent of cluster size (see Fig. 6) we expect that the number of bath sites needed for convergence is independent of cluster size.



FIG. 10. (Color online) Fraction of hybridization sum-rule for a two-site cluster as a function of N_b . As more and more bath sites are added, the exact hybridization sum-rule is approached. Shown is the percentage of the sum-rule for irreducible representation A in CD-MFT and K=0 in DCA. For irreducible representation B and $K = \pi$ we find very similar behavior.



FIG. 11. (Color online) Density for linear CDMFT clusters of increasing size N_c with U=4t at $\mu=0$ (a) and $\mu=t$ (b). All cluster calculations are for $N_b=8$. Circles denote the average density per cluster site. (Green) Open squares are the individual cluster-site occupations. The size of the squares indicates how close the site is to the center of the cluster. To assess the reproducibility of the calculation we show for each even value of N_c the result of at least two to seven runs with different choice of the starting set of Anderson parameters. The dotted line represents the exact Bethe-ansatz result.

D. Convergence with number of cluster sites

We now analyze the convergence of CDMFT with the number of cluster sites N_c . As before we focus on the density on the cluster n, which is shown in Fig. 11 for U=4t and chemical potential $\mu=0$ and $\mu=t$. Considering the series of odd or even N_c separately, we see that with increasing cluster size the average density systematically approaches the exact result for the infinite chain. Interestingly, going from a cluster with an even number of sites N_c to N_c+1 the average density hardly changes. For both chemical potentials already the smallest cluster gives a significant improvement over a single-site calculation. For $\mu=0$ the exact density is basically obtained for $N_c=2$. For $\mu=t$ convergence to the infinite-chain result is only reached at $N_c=6$.

In Fig. 12 we show how the density versus chemicalpotential curve for CDMFT calculations of increasing cluster size approach the exact result for the infinite Hubbard chain. We find that the closer we come to the metal-insulator transition the harder it gets to reach the infinite-size limit. This does not come as a complete surprise, as the self-energy is



FIG. 12. (Color online) Density *n* as a function of chemical potential μ at U=4t for linear CDMFT clusters of increasing size and $N_b=8$ compared to the exact result for an infinite Hubbard chain (Bethe Ansatz). The crosses give the results from Fig. 4.7 of Ref. 13. The inset shows how the convergence of the density with increasing cluster size to the Bethe result becomes progressively slower close to the Mott transition. $n_{BA}-n_{CDMFT}$ jumps when the self-consistent solution changes sector ($N_{\uparrow}, N_{\downarrow}$). Close to these sector changes the results slightly depend on the initial conditions, i.e., there is a hysteresis between calculations increasing or decreasing μ . This is shown as multiple symbols for a given chemical potential. The upper plot shows the average density for two-, four-, and six-site clusters, the lower plot the density on the two central sites.

expected to become strongly k dependent at the Mott transition.^{33,34}

We note that our results for $N_c=2$ agree with Fig. 4.7 of Ref. 13. They are not compatible with Fig. 4.1 of Ref. 13 and Fig. 2 of Ref. 12. We have checked that our calculation is properly converged by starting from a number of different initial points, always converging to essentially the same density. To achieve this, for chemical potentials $\mu \ge 1.3t$ we increased β from 256/t to 512/t, in line with the trend shown in Fig. 8. We can, however, get significant variations in the density by putting restrictions on the bath parametrization. Using, e.g., only six bath sites, the CDMFT result happens to be closer to the one for the infinite system, in the vicinity of the Mott transition.¹³ This is shown in Fig. 13. For small baths the calculated density is very sensitive to N_h and can be either larger or smaller than the density for the infinite chain. For such small baths results will therefore critically depend on the fitting. We can, e.g., artificially "improve" the result by forcing a pair of bath energies to zero. Other restrictions



FIG. 13. (Color online) Average density as a function of the number of bath-sites N_b for two- and four-site clusters with U=4t, $\mu=1.3t$, and $\beta=512/t$. As for smaller chemical potential the density is converged for $N_b=8$ (cf. Fig. 9). We observe that for larger μ unconverged calculations with small baths, e.g., $N_b=6$, can give densities closer to the one of the Hubbard chain.

on the bath parameters instead move the densities further away from the Bethe curve. In all these cases we find that the restricted bath-parametrization results in a significantly deteriorated fit of the bath Green matrix. That is, the sensitivity of these calculations to technical details merely shows the effects of an inadequate fitting of the bath Green matrix. By increasing N_b , the bath-parametrization improves and the calculated density converges, as shown in Fig. 13. Nevertheless, differently from the behavior at smaller chemical potential (cf. Fig. 9), the converged value is substantially smaller than the Bethe ansatz one. As this does not improve much with increasing N_c , we can conclude that the clusters are still too small to accurately capture the behavior of the infinite system close to the Mott transition.

VI. CONCLUSIONS

The central problem of dynamical mean-field calculations with a solver that uses a small number of sites is to find a good parametrization of the bath. To address this problem for dynamical cluster approximations we have presented a systematic formulation for the bath degrees of freedom. We have found sum-rules which allow to identify what hybridizations vanish and hence need not be parametrized at all. In addition the sum-rules can be used to check convergence for small baths. For the nonvanishing hybridization functions, we have introduced a bath-parametrization based on the irreducible representations of the cluster Green matrix. In this approach the fitting of the bath sites is broken into independent fits of irreducible blocks of the Green matrix. This leads to a significant simplification of the fitting procedure which is particularly important when dealing with large baths. The symmetry-based approach should also benefit the variational cluster approximation,²⁴ where the determination of the parameters requires a Lanczos calculation in each optimization step.

As an application we have revisited the Hubbard chain. While this one-dimensional problem is the worst case scenario for DMFT, which is exact in infinite dimensions, it is technically the easiest case for CDMFT, because it requires only a minimal bath which is essentially independent of cluster size. This allowed us to study the results of CDMFT using linear clusters of increasing size, extending previous work that was limited to two- and three-site clusters.^{12,13} Analyzing the density as a function of chemical potential, we find that results significantly improve already going from a single-site DMFT to a two-site cluster and become systematically better for larger and larger cluster sizes. Close to the Mott transition the convergence with N_c critically slows down, implying that the *k* dependence of the self-energy gets more and more important.

The fortuitous independence of the CDMFT bath on the cluster size for the Hubbard chain is lost in higher dimensions. Already for a cluster as small as 2×2 we have to fit three functions, one of which belongs to a doubly degenerate E representation. In that case $N_b=8$ translates to only two effective bath sites per bath Green function and the fit in general becomes very poor, as illustrated in Fig. 14, except in special cases like large U at half-filling. For comparison, in our one-dimensional calculations we need at least four bath sites per bath Green function for a converged bath, implying that it is extremely hard to converge a 2×2 CDMFT calculation with a Lanczos solver.

The same is true for DCA, since the number of baths increases with cluster size, independently of the cluster or the nature of the hopping. Nevertheless, it might help somewhat that these baths appear to be easier to fit as suggested by Fig. 7. Still, for zero-temperature cluster calculations it seems necessary to move to impurity solvers that can handle large baths, e.g., DMRG.³⁵ For these calculations with large baths the efficient parametrization of bath and fitting of irreducible blocks will become even more important.



FIG. 14. (Color online) Fitting the bath Green function for a 2 \times 2 cluster in CDMFT with U=12t, $\mu=0$, $N_b=8$. The imaginary part of the block-diagonalized bath Green matrix (shown are only the three inequivalent blocks labeled by the irreducible representation) and its corresponding fit are plotted for two values of the exponent *N* (cf. Sec. V B). The quality of the fit is clearly very poor in both cases (for N=1 compare full red/gray line with full black one and for N=0 compare dashed green/gray line with dashed black one).

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