A novel continuous time quantum Monte Carlo solver for dynamical mean field theory in the compact Legendre representation

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Dynamical mean-field theory (DMFT) is one of the most widely-used methods to treat accurately electron correlation effects in ab-initio real material calculations. Many modern large-scale implementations of DMFT in electronic structure codes involve solving a quantum impurity model with a Continuous-Time Quantum Monte Carlo (CT-QMC) solver [1–4]. The main advantage of CT-QMC is that, unlike standard quantum Monte Carlo approaches, it is able to generate the local Green’s functions $G(\tau)$ of the correlated system on an arbitrarily fine imaginary time $\tau$ grid, and is free of any systematic errors. In this work, we extend a hybrid QMC solver proposed by Khatami et al. [5] and Rost et al. [6] to a multi-orbital context. This has the advantage of enabling impurity solver QMC calculations to scale linearly with inverse temperature, $\beta$, and permit its application to $d$ and $f$ band materials. In addition, we present a novel Green’s function processing scheme which generates accurate quasi-continuous imaginary time solutions of the impurity problem which overcome errors inherent to standard QMC approaches. This solver and processing scheme are incorporated into a full DFT+DMFT calculation using the CASTEP DFT code [7]. Benchmark calculations for SrVO$_3$ properties are presented. The computational efficiency of this method is also demonstrated.

I. INTRODUCTION

Density functional theory (DFT), in both its local density (LDA) and generalised gradient (GGA) approximations, is a highly effective method for the calculation of quantitatively accurate ab-initio ground-state properties of a wide range of real materials [8]. However, for some materials DFT’s outstanding predictive capabilities become significantly less reliable. Such materials are found typically amongst the transition metal oxides, as well as in lanthanide and actinide-based compounds. This is problematic because of the potential application of these materials in the fields of quantum computing, data storage and high temperature superconductivity. These materials characteristically have narrow bandwidths ($W \sim 2–3$ eV), so the correlations induced by the Coulomb interaction between valence electrons ($U \sim 4–5$ eV) are strong, implying that i.e. $U/W \gtrsim 1$. When bandwidths are broader the ratio of electron correlation to bandwidth is reduced (i.e. $U/W \lesssim 1$), so the effects of electron correlation are weaker. In the latter case, the typical approximations involved at the DFT level work well, often with a quantitative precision that sustains comparison with experiment.

A single unified framework is sought in which the effects of electron correlation can be addressed for real material calculations over a wide range of temperatures and correlation strengths $U/W$. In recent years, a combination of DFT and dynamical mean-field theory (DMFT) has proved to be an effective way of interpolating between the itinerant and strongly localised limits of the electronic behaviour. The DFT+DMFT approach has been well established, and allows extending the capabilities of DFT-based approaches into calculations where electron correlation effects are significant. DFT+DMFT has evolved into a powerful method for dealing with correlation effects, and is being frequently used for material-specific applications. In addition to DFT+DMFT, the GW+DMFT approach is continuing to mature into an effective way of undertaking material-specific calculations for correlated systems [9].

The critical component of a DFT+DMFT calculation is the so-called “impurity solver”, which represents the quantum many-body physics interactions of the correlated electrons in the material. It is essential that the solver contains an accurate representation of the physics of the interacting electrons, and can efficiently compute solutions in fast and stable way over a wide range of parameters ($U$, $W$ and $T$). A variety of solvers are available for use in DMFT calculations, and they can be selected to match the computational resources available to complete a calculation in a time-efficient way. These solvers can be based on quasi-analytical methods (e.g. Hubbard I, Iterated Perturbation Theory (IPT), Non-Crossing Approximation (NCA) or fluctuation exchange approximation (FLEX)) or numerical methods (e.g. Numerical Renormalisation Group (NRG), Exact Diagonalisation (ED), QMC) [10]. Of all these methods it is the quantum Monte Carlo methods which have proved to be particularly popular. This is because QMC methods are conceptually straightforward and can return stable results (to statistical accuracy) over a wide range of parameter values.

A QMC technique known as continuous-time QMC (CT-QMC) [11,12] has emerged lately as a popular solver
for many DMFT applications. The main advantage of CT-QMC is that it overcomes one of the most significant limitations of conventional QMC methods - namely, the systematic errors which arise due to the Trotter-Suzuki decomposition and the concomitant discretisation of the imaginary time interval. Errors of this kind can generate substantial bias in the results of DMFT calculations. This is because they can significantly shift the converged fixed point of the DMFT iteration [6]. However, a disadvantage of most of the readily available CT-QMC solvers is that the computational effort scales cubically with inverse temperature $\beta = 1/k_B T$, and this severely limits their access to low temperature phases. Moreover, because the order of perturbation of CT-(HYB)/QMC scales with the product of kinetic energy and $\beta$, it is costly to apply to systems far from the localisation-delocalisation transition, especially at low temperatures. Although our method is applicable over a large range of parameters, we concede that in the limit of very large Coulomb repulsion a technique such as CT-HYB may be more effective. For multi-orbital or cluster DMFT applications, the inverse temperature scaling limitation becomes even more significant. A key computational bottleneck in calculations of fully charge self-consistent DFT+DMFT properties of real materials is the requirement to generate accurate impurity model solutions on a fast time-scale with medium-scale computational resources. For such challenging applications QMC methods remain a competitive technique for solving quantum impurity problems, and our intention here is to introduce a complementary QMC technique that has advantage for some applications. Also, when calculating a material equation of state, numerous DFT+DMFT total energy calculations are needed over a range of temperatures and pressures [12]. To perform these calculations to a level comparable with experimental data demands accurate and computationally efficient impurity solvers [13, 14].

In this paper we present a QMC solver which scales linearly in inverse temperature [5, 6] extended to a multi-orbital context, and supplement the solver with a robust Lagrange/Chebyshev-based Green’s function interpolation technique that facilitates controlled extrapolations of the impurity Green’s functions on multiple discretised imaginary time grids to a solution that is quasi-continuous in imaginary time. Because the solver self-energy is very sensitively dependent upon this Green’s function it is imperative that the impurity model solution is an accurate one. In this way we are able to calculate accurate quasi-continuous time quantum impurity model solutions for multi-orbital systems faster than CT-QMC solvers, particularly at temperatures those solvers find challenging to access.

The paper is organised as follows. In Section II we introduce the QMC solver and describe how it is integrated into a DFT+DMFT scheme for real material calculations using the CASTEP plane-wave DFT code [7]. In Section III a new technique designed to generate accurate quasi-CT Green’s functions using fits of multiple low-resolution QMC solutions is presented. Some of the particular issues of fitting Green’s functions from real materials are discussed. In Section IV, to test the technique fully, we undertake a full DFT+DMFT calculation on strontium vanadium oxide SrVO$_3$ by integrating our solver with both the CASTEP DFT code and the quasi-CT Green’s function scheme. Where appropriate, calculations are base-lined against equivalent quasi-CT QMC calculations using the TRIQS solver that is embedded in CASTEP [7, 15]. All energies and temperatures are in eV.

II. QMC IMPURITY SOLVER AND DFT+DMFT CALCULATIONS

1. The QMC algorithm

The purpose of a quantum impurity solver in a DMFT calculation is to adequately capture the physics of spin-dependent electron-electron interactions, using either a numerical model Hamiltonian representation or a quasi-analytic approach [10]. The initial conditions for the model are derived from a material-specific DFT calculation, and then the solver solution is processed and fed back to the DFT code in an iterative cycle. The key to efficient DFT+DMFT calculations of real material properties is finding a fast, accurate and stable solution to the impurity model (without a fermionic sign problem [16]) over a wide range of model parameters ($U, J$ and $W$) and inverse temperatures, $\beta$.

Most early DFT+DMFT calculations employed a Hirsch-Fye (HF) solution of the single impurity Anderson model to represent electron correlation effects [10]. The appeal of this solver is that it is QMC-based method which can be applied over a broad parameter range and band fillings without being compromised by fermionic sign problems. However, this method contains a systematic error inherent in the Trotter decomposition and imaginary time discretisation. Also, its inverse temperature scaling goes as $\beta^3$. More recently HF-QMC has been complemented by the CT-QMC method (and variants thereof) discussed above. But, as noted there, it too scales as $\beta^3$. Another popular impurity solver is based on exact diagonalisation (ED) of the impurity Hamiltonian [17, 18]. The advantage of that method is that it gives an exact Green’s function solution and scales linearly in $\beta$. However, it relies on a discretisation of the Weiss field (bath Green’s function), and the computational load scales exponentially with the number of bath sites which can be limiting for applications beyond application to single site, single band, impurity models.

Khatami et al. [5] and later Rost et al. [6] introduced a novel Hamiltonian-based QMC scheme that combined
the virtues of ED with a standard QMC-type solution, resulting in an impurity solver that scales cubically in the number of discretised bath sites and linearly in inverse temperature. Moreover, it has the same fermion sign performance as HF-QMC. Rost et al. [6] demonstrated the advantages of using this type of solver in DMFT studies of the single impurity model on a Bethe lattice [10]. The solver still exhibits the systematic Trotter errors characteristic of QMC methods, but, as we show below, these can be treated by the application of a novel Green’s function interpolation and extrapolation technique which generates a quasi-CT solution.

The formulation of the QMC algorithm for single band physics is described in detail in [5] and [6]. In what follows we will be extending the method in a multi-band context. For the general case of electrons on a single impurity site which interact both on and between 2M orbital/spin pairs (where 1 < m ≤ 2M), the Hamiltonian is given by

\[ H = H_{LDA}^0 + \sum_{m} U_{mm} n_m \mathbf{\tau}_{m}\mathbf{\sigma}_{m} - \sum_{m\sigma} \mu n_{m\sigma} + \sum_{m < m'} U_{mm'} n_{m\sigma} n_{m'\sigma} + \sum_{m < m'} (U_{mm'} - J_{mm'}) n_{m\sigma} n_{m\sigma'} \]

where \( U_{mm} = U \), \( J_{mm'} = J \), and \( U_{mm'} = U - 2J \) for \( m \neq m' \). Also, \( n_{m\sigma} = c^{\dagger}_{m\sigma} c_{m\sigma} \) where \( c_{m\sigma} \) are the annihilation (creation) operators for electrons in orbital \( m \) with spin \( \sigma \). In this formulation there are Hubbard-like interactions between opposite-spin electrons in the same orbitals and neighbouring orbitals, and Ising-like Hund’s coupling between same-spin electrons on neighbouring orbitals. The full Slater-Kanamori Hamiltonian for multi-orbital interactions is rotationally invariant in spin-space [19] and also includes “spin-flip” and “pair-hopping” terms, which are likewise determined by \( \approx J_{mm'} \). These interactions cannot be straightforwardly factorised into products of quadratic operators \( n_{m\sigma} \), which is essential in any auxiliary field QMC-based implementation. Various schemes have been proposed to include these additional terms into QMC formulations [19][24], but this is not wholly straightforward to do without introducing additional sign problems, especially at low temperatures. To obviate such sign problems we do not include spin-flip and pair-hopping terms in this work, and this is a current limitation of this method. However, future work will address this deficiency. ED and CT-QMC approaches can handle the full Hamiltonian, and this must be borne in mind when comparing results generated by different multi-orbital impurity solvers.

Our solver uses a multi-orbital generalisation of the standard determinantal QMC procedure. For a multi-orbital interaction matrix \( \sum_{m_{m'n_{m'n_{m}}}U_{mm'n_{m'n_{m}}} \rangle m \mathbf{\tau}_{m}\mathbf{\sigma}_{m} \) the usual Hubbard-Stratonovich transformation can be applied, i.e.

\[ e^{-U_{mm'} \Delta \tau (n_{m-1/2}n_{m'1/2})} = \frac{1}{2} \sum_{nmn'm'} e^{\lambda_{nmn'm'}} (n_{m-n_{m'}}) \]

where \( \lambda_{nmn'm'} = \text{acosh}[e^{\frac{1}{2} \Delta (U_{mm'})}] \) and \( U_{mm'} \) is one of \( U \), \( (U - 2J) \) or \( (U - 2J) \), depending on which orbital/spin pair is being considered. There is an auxiliary spin field \( s \) for each inter and intra orbital interaction.

Following conventional QMC procedure, at each imaginary time slice each of the \( M(2M - 1) \) auxiliary spin fields is flipped sequentially and then tested for acceptance or rejection using the ratio \( R = R_m R_{m'} \) where

\[ R_m = 1 + \langle (1 - G_m) \gamma_m \rangle \]
\[ R_{m'} = 1 + \langle (1 - G_{m'}) \gamma_{m'} \rangle \]

with \( \gamma_m = e^{2\lambda_{mm'n_{m'n_{m'}}} - 1} \) and \( \gamma_{m'} = e^{2\lambda_{mm'n_{m'n_{m'}}} - 1} \). If the spin flip is accepted the Green’s function is updated according to the standard BSS [25] prescription

\[ G_{j,k,m} = G_{j,k,m} - (\delta_{j,1} - G_{j,1,m}) \gamma_m G_{j,k,m}/R_m \]
\[ G_{j,k,m'} = G_{j,k,m'} - (\delta_{j,1} - G_{j,1,m'}) \gamma_{m'} G_{j,k,m'}/R_{m'} \]

If the spin flip is rejected the Green’s functions remain the same. The condition for half-filling in this Hamiltonian is given by \( \mu = (U + (M-1)(U - 2J) + (M-1)(U - 3J))/2 \).

In the exact diagonalisation method [17] the infinite lattice surrounding the impurity site is approximated by a discretised bath of finite size \( N_h \), and the identical procedure is followed in this QMC solver. The first step is to parameterise the Weiss field \( \Omega \) for each orbital and spin in terms of a finite number of bath parameters by approximating the Weiss Green’s function in terms of a non-interacting Anderson impurity model as follows:

\[ G_{\text{And},m}(i\omega_n) = i\omega_n + \mu - \epsilon^{\text{imp}}_m - \sum_{k=1}^{N_h} V_{m,k} V_{m,k}^* / i\omega_n - \epsilon_{m,k} \]

where \( k \) is the index for the bath level, \( m \) is the index for each orbital/spin. Essentially, this entails minimising the difference between the Weiss field and equation [6] using a cost function, typically

\[ \chi^2[\epsilon_k, V_k] = \sum_{n=0}^{n_{\infty}} A_n |G_{\text{And}}(i\omega_n : \{\epsilon_k, V_k\}) - \Omega(i\omega_n)|^2 \]

In practice it has been found advisable to weight the cost function towards smaller imaginary frequencies by using pre-factor \( A_n \approx 1/i\omega_n^2 \). This avoids squandering fitting cost on the asymptotic regions of the Green’s function, and has been found particularly apposite when attempting accurate fits to real material Green’s functions. A
robust multi-dimensional fitting routine was used to determine the set of $\epsilon_k$ and $V_k$ for each orbital, which are subsequently used to parameterise the QMC solver. To avoid falling into local minima, multiple fits over a wide range of initial conditions are performed and the best-fit is then selected.

Once a set of $\epsilon_k, V_k$ has been found to fit the Anderson impurity Green’s function the QMC part of the calculation proceeds as a standard Monte Carlo simulation for a multi-orbital context, where the imaginary time interval $0 \leq \tau \leq \beta$ is discretised into $L$ time-slices ($\beta = L \Delta \tau$). A sufficient number of Monte Carlo steps are run on a parallelised processor configuration, and the imaginary time impurity Green’s functions are calculated for each orbital and spin using the procedure of White et al. [26], and Loh and Gubernatis [27].

### 2. DFT+DMFT implementation with CASTEP

In our implementation of DFT+DMFT the plane-wave DFT code CASTEP was used [7]. It is a widely available DFT code that can be readily used for calculating ab-initio material properties. Previous work with CASTEP has demonstrated the integration of a Hubbard I solver [15] and an ED-based solver [18] for materials-specific calculations, and what follows builds on that. The multi-orbital solver described above was integrated with CASTEP to better represent the electron correlation effects. Many of the issues associated with integrating an impurity solver with a DFT code are generic and a variety of methods have been used. Here we give a description of our CASTEP implementation of DFT+DMFT, and this can be compared to other approaches.

A CASTEP calculation generates an Linear Combination of Atomic Orbital (LCAO) basis set using either norm-conserving or ultra-soft pseudopotentials. These two cases can be dealt with on an equal footing by defining an “overlap matrix”, $S$. The basis set functions generated from norm-conserving pseudopotentials are orthogonal by construction, so the overlap matrix is the identity matrix. In the case of ultra-soft pseudopotentials these states are overlapping with a matrix:

$$\langle \chi_{m'} R | S | \chi_m R \rangle = \delta_{m',m} \delta R R' \ .$$

This implies that the Kohn-Sham (KS) equations transform from a standard eigenvalue problem into a generalised one

$$H_k^{KS} | \Psi_{k,\nu} \rangle = E_{k,\nu} S | \Psi_{k,\nu} \rangle \ .$$

where $\Psi_{k,\nu}$ are the KS eigenstates.

### 3. Projectors

The standard DMFT technique is defined in terms of completely localised electronic states (e.g. as in Hubbard model). On the other hand, in a wide class of ab-initio codes, including CASTEP, the electrons are described in terms of completely delocalised plane wave states. Consequently, a key feature of the DFT+DMFT calculation is the selection of a correlated sub-space of orbitals, and the means to bridge the DFT Bloch space basis and the basis of the correlated sub-space. To do this we define the orthonormal projectors $P_{L,\nu}(k)$, where

$$P_{L,\nu}(k) = \langle \chi_L | S | \Psi_{k,\nu} \rangle \ .$$

We now have two distinct spaces - i) the KS Bloch space indexed by $k$ and $\nu$, and ii) the localised (correlated) subspace indexed by $L$.

To go from $\chi_L$ to $\Psi_{k,\nu}$ i.e. “up-folding”, we use

$$| a_{k,\nu} \rangle = \sum_L P_{L,\nu}^{\ast}(k) | b_L \rangle \ .$$

Conversely, to go from $\Psi_{k,\nu}$ to $\chi_L$ i.e. “down-folding”, we use

$$| b_L \rangle = \sum_{k,\nu} P_{L,\nu}(k) | a_{k,\nu} \rangle \ .$$

where $a_{k,\nu}$ is a vector defined in Bloch space and $b_L$ is a vector defined in the space of correlated orbitals.

The projector matrix satisfies the following condition:

$$\sum_{k,\nu} P_{L,\nu}(k) P_{L',\nu'}^{\ast}(k) = \delta_{L,L'} \ .$$

In this scheme an up-folding operation followed by a down-folding operation is equivalent to an identity operation. Both the up-folding and down-folding operations are each used once in each DMFT iteration cycle, as we now discuss.

### 4. The DMFT cycle

To begin the DMFT iteration for SrVO$_3$ we use CASTEP to calculate the Bloch Green’s function

$$G_{\nu,\nu'}^B(k, i\omega_n) = \left\{ \left( i\omega_n + \mu - \epsilon_{k,\nu} \right) \delta_{\nu,\nu'} - \Sigma_{\nu,\nu'}^B(k, i\omega_n) \right\}^{-1} \ .$$

which is the Fourier transform $F.T.\{ [r | \hat{G} | r] \}$, where

$$\hat{G}(r, i\omega_n) = (i\omega_n + \mu + \frac{1}{2} \nabla^2 - \nu_{KS}(r) - \Sigma^B(r, i\omega_n))^{-1} \ .$$

(An adjustment to the chemical potential $\mu$ is necessary at this point in the cycle to ensure correct level of electron occupancy - see Appendix A.) We now consider a
correlated atom at location $R$. The basis functions in the correlated sub-space are denoted by $m$. The local Green’s function for the correlated sites is now obtained from the Bloch Green’s function by down-folding and then averaging over the Brillouin zone, as follows:

$$G_{m,m'}^{loc}(i\omega_n) = \frac{1}{N_k} \sum_{\nu,\nu',k} P_{\nu,\nu',m}(k)G_{\nu,\nu'}^B(k,i\omega_n)P_{\nu,\nu',m'}(k).$$

(15)

We now make the identification of $G^{loc}$ with $G^{imp}$ - so in the DMFT impurity Dyson equation we calculate the Weiss field

$$[G_0(i\omega_n)]^{-1}_{m,m'} = \Sigma_{m,m'}^{imp}(i\omega_n) + [G^{imp}(i\omega_n)]^{-1}_{m,m'}.$$  

(16)

On the first iteration we make a guess for the self-energy (typically $\Sigma_{\nu,m}^{imp} = 0$). We then parameterise the Weiss field $[G_0^{-1}]_{m,m'}$ using equation (15) for $G^{1nd}_{\nu,m}$.  

It is now possible to run the multi-orbital QMC solver (described above) to obtain a set of impurity Green’s functions ($G_{QMC}(\tau)$). Following the interpolation and extrapolation procedure (described in Section III), the Fourier Transformed Green’s functions are used to calculate a revised impurity self-energy, i.e.

$$\Sigma_{m,m'}^{imp} = [G_0(i\omega_n)]^{-1}_{m,m'} - [G_{QMC}(i\omega_n)]^{-1}_{m,m'}.$$  

(17)

At this point it is necessary to make allowance for the “double counting” term, $V_{m,m'}^{DC}$ (see Appendix B), and then up-fold the impurity self-energy back to Bloch space, i.e.

$$\Sigma_{\nu,\nu'}^{B}(k,i\omega_n) = \sum_{m,m'} P_{\nu,m}(k)(\Sigma_{m,m'}^{imp}(i\omega_n) - V_{m,m'}^{DC})P_{\nu,m'}(k).$$  

(18)

The self-energy is purely local when expressed in the set of correlated orbitals, but it acquires momentum dependence when up-folded to the Bloch basis set. The up-folded self-energy is returned to equation (13), and the iteration sequence is continued until an acceptable level of convergence is reached in the self-energy (or chemical potential).

System properties can also be calculated during the DMFT iteration cycle. Of particular value are the Bloch-level occupation matrix:

$$N_e = \frac{1}{N_k} \sum_{\nu,k} \sum_n G_{\nu,\nu'}^B(k,i\omega_n)e^{i\omega_n \rho},$$  

(19)

the spectral density

$$A_{\nu,\nu'}(k,\omega) = -\frac{1}{\pi} \text{Im} G_{\nu,\nu'}^B(k,\omega),$$  

(20)

and the density of states (DOS)

$$D(\omega) = \frac{1}{N_k} \sum_{\nu,k} A_{\nu,\nu'}(k,\omega).$$  

(21)

To calculate the spectral density and DOS, the analytic continuation from imaginary ($i\omega_n$) to real ($\omega$) frequency is made using the Padé approximation [28].

As well as our implementation here, which uses the BSS method to update the Green’s function, a version of the multi-orbital solver code has also been implemented which uses the Nukala et al. [29] fast updating method. The results are identical to BSS, but the Nukala method will be better suited to multi-orbital cluster DMFT applications.

III. A NOVEL QUASI-CONTINUOUS IMAGINARY TIME SOLUTION

The conventional QMC method calculates imaginary time Green’s functions $G(\tau)$ on a grid of ($L$) regularly spaced $\tau$ points over the interval $0 \leq \tau < \beta (= L\Delta\tau)$. In DMFT calculations this Green’s function is Fourier transformed and then used to calculate the self-energy of the impurity problem. However, any attempt to calculate the frequency-dependent self-energy $\Sigma(\omega)$ using the raw QMC Green’s functions immediately generates two problems, both related to the fact that the Green’s function is calculated on a discrete imaginary time grid. Firstly, the Trotter decomposition implies that the resulting self-energy will be in error - to first order $\sim U\Delta\tau^2$. Secondly, attempting to Fourier transform the Green’s function $G_{QMC}(\tau)$ as it stands will introduce aliasing errors in the transformed function $G_{QMC}(i\omega_n)$. Aliasing arises from the fact that the Fourier transform of the imaginary time Green’s function is a periodic function in imaginary frequency with poles at multiples of the Nyquist frequency $(2\pi L/\beta)$. This leads to weight being folded back into the transform from frequencies above the Nyquist frequency, and generates incorrect asymptotic behaviour of $G_{QMC}(i\omega_n)$.

In this section we present a novel multi-scale Green’s function processing technique that simultaneously addresses these two issues. The method interpolates a set of comparatively coarse time-scale QMC impurity Green’s functions on to a fine-scale $\tau$ grid (to eliminate aliasing problems), and extrapolates them to generate a quasi-continuous imaginary time solution for the multi-orbital impurity problem (to eliminate systematic Trotter errors). This method generates Green’s functions and self-energies with the correct behaviour across all imaginary frequencies that can be seamlessly integrated into the DMFT iteration cycle.

Splining procedures (with and without quasi-CT extrapolation) have been used previously to process imaginary time Green’s functions so they can be used in DMFT calculations [3]. In real material calculations the imaginary time Green’s function can substantially change its magnitude over the space of few imaginary time intervals, $\Delta\tau$. For strong electron interactions and low
temperature, this behaviour can be particularly acute. Because of this care must be taken when interpolating and extrapolating Green’s functions so as not to inadvertently incorporate spurious processing pathologies into calculated Green’s functions. Moreover, QMC methods generate noisy data which can cause additional challenges for splining schemes, so here we describe a new Green’s function splining and extrapolation scheme which is robust and can fit noisy data before generating a quasi-continuous time solution to the DMFT Hamiltonian. As we will show subsequently, it is straightforward to integrate this scheme into a full material-specific DFT+DMFT calculation.

1. Polynomial basis method

One way to reduce the systematic error introduced by the Trotter decomposition is to simply reduce $\Delta \tau$ by increasing the number of imaginary time steps $L$. However, the price for an increasingly accurate representation of $G(\tau)$ is a significantly more computationally intensive QMC calculation. For example, the ubiquitous Hirsch-Fye solver scales as $\sim L^3$.

The objective of our new Green’s function processing protocol is to demonstrate that by using a novel polynomial basis interpolation for $G_{QMC}(\tau)$, and an extrapolation scheme, we can perform QMC calculations on multiple, relatively coarse, imaginary time grids to generate a quasi-continuous imaginary time solution.

We expand $G(\tau)$ in an arbitrary orthogonal polynomial basis $P_i^{(k)}[x(\tau)]$ (e.g. Legendre, Chebyshev, etc) where $i$ is the polynomial order, $k$ is the polynomial species and $x(\tau) = \frac{\tau}{\beta} - 1$ is the transformation from $[-1, +1]$ to $[0, \beta]$. The expansion is:

$$G^{(k)}(\tau) = \sum_{i \geq 0} P_i^{(k)}[x(\tau)] G_i^{(k)}. \quad (22)$$

To isolate the basis coefficients we apply the orthogonality constraints obeyed by the polynomials, i.e.,

$$\int_0^\beta d\tau G^{(k)}(\tau) P_{i'}^{(k)}[x(\tau)] W(\tau) = 0, \quad (23)$$

The general orthogonality condition obeyed by the family of polynomials $P_i^{(k)}[x(\tau)]$ is

$$\int_0^\beta d\tau P_i^{(k)}[x(\tau)] P_{i'}^{(k)}[x(\tau)] W(\tau) = \tilde{W}(i) \delta_{i,i'}, \quad (24)$$

and so the basis coefficients can be calculated as

$$G_i^{(k)} = \frac{1}{\tilde{W}(\tau)} \int_0^\beta d\tau G^{(k)}(\tau) P_i^{(k)}[x(\tau)] W(\tau). \quad (25)$$

Here we restrict our analysis only to the Legendre polynomials, where $W(\tau) = 1$ and $\tilde{W}(i) = \frac{1}{2i+1}$.

Calculating $G_i$, allows us to express $G(\tau)$ on an arbitrarily fine imaginary time grid.

It is imperative to first obtain a reliable representation of $G(\tau)$ in the Legendre basis. To achieve this, it is possible to formulate a controlled fitting procedure that uses the Legendre basis coefficients $g_i$ as parameters to be adjusted to the raw QMC data.

2. Green’s function fitting procedure

Expressing $G(\tau)$ in the Legendre basis allows a fitting procedure to be formulated, which amounts to the minimisation of the function

$$\min_{\{g_i\}} [G_{QMC}(\tau) - G_{FIT}^{(g_i)}(\tau)], \quad (26)$$

where the fitted (model) function $G_{FIT}^{(g_i)}(\tau)$ is parametrised by the basis coefficients $g_i$, i.e.

$$G_{FIT}^{(g_i)}(\tau) = \sum_{l \geq 0} N_l \sqrt{2l+1} P_l(x(\tau)) g_i. \quad (27)$$

It is straightforward to find $G_{FIT}^{(G_i)}(\tau)$ using the conjugate gradient method, and since $N_l$ (the number of Legendre coefficients) is generally quite modest. In our case because $N_l \approx 20$, this procedure is exceptionally computationally efficient. Moreover, by shifting the paradigm of dealing with a statistical problem, i.e. Monte Carlo, to that of an optimisation one, allows the advantage of including a-priori information. This strategy becomes particularly attractive when dealing with unrefined QMC data on exceedingly coarse imaginary time grids. In this respect, the Legendre basis proves to be an extremely useful tool for two outstanding reasons; the first is its simple relationship to the moments of the $G_{QMC}(i\omega_n)$, and the second is due to the convergence properties of $g_i$ in the Kernel Polynomial Method (KPM) [30]. The final significant constraint that can be imposed on the parameters $g_i$ in an a-priori fashion is the convexity of $G(\tau)$. We now discuss how incorporating this additional information on $G_{FIT}(\tau)$ can reliably improve its accuracy.

In many-body calculations it is essential that the Green’s function solutions have the correct high frequency tail. Often the high frequency tail, which has a $1/i\omega_n$ behaviour, is fitted onto the low frequency result. However, in the Legendre basis there is an exact relationship between the moments of the Green’s function and the Legendre coefficients [31]. By introducing a set of Lagrange parameter penalty terms into equation [25] information on the tail can be included into the fit, thereby eliminating the need for an ad-hoc fit of the tail. The term added to equation [20] to ensure the correct
high frequency tail asymptotic behaviour is

\[
\lambda_1 \left( c_1 + \sum_{l \geq 0, \text{even}} \frac{2\sqrt{2l+1}}{\beta} g_l \right) + \lambda_2 \left( c_2 - \sum_{l \geq 0, \text{odd}} \frac{2\sqrt{2l+1}}{\beta^2} g_l(l+1) \right) \tag{28}
\]

where \( \lambda_1 \) is the Lagrange parameter controlling the first moment \( c_1 \), and \( \lambda_2 \) is the Lagrange parameter controlling the second moment \( c_2 \). The inclusion of the two Lagrange penalty terms relies on knowing the values for \( c_1 \) and \( c_2 \) for \( G(\omega_n) \) before the fit is performed. Fortunately, for \( c_1 \) it is known that in the high frequency limit of \( G_{\text{QMC}}(\omega_n) \), it behaves as \( 1/(\omega_n) \), and therefore \( c_1 = 1 \). The equivalent considerations for \( c_2 \) are somewhat more involved, but it can be shown that

\[
c_2 = \mu - \varepsilon + \Sigma'(\infty), \tag{29}
\]

where \( \mu \) is the chemical potential, \( \varepsilon \) is the impurity level and \( \Sigma'(\infty) \) is the high frequency real asymptotic self-energy of an isolated impurity. \( \Sigma'(\infty) \) is attainable by solving the Anderson Impurity Model with a finite set of bath orbitals; for simplicity, this can be achieved using an ED-solver [13], or Hubbard I solver [10].

It is an unavoidable fact that the truncation of a function in any polynomial basis that is, in principle, infinite can lead to convergence issues. In the case where there are non-differentiable points or singularities it is especially problematic and can lead to the well established Gibbs oscillations. The severity of the oscillations near these ill-defined points can be damped by the introduction of a kernel \( k_l \) in equation (27) such that \( g_l \rightarrow k_l g_l \). The process of truncating this series and modifying the basis coefficients amounts convolving \( G(\tau) \) with a kernel \( k_l \). Since \( G(\tau) \) is continuously differentiable (except at its boundaries) it is possible to pick a kernel that will guarantee this behaviour, and in the process filter out any spurious noise. In this work, we concern ourselves only with the kernels of Dirichlet and Jackson.

3. Green’s function extrapolation for \( \Delta \tau^2 \) scaling

Representing \( G(\tau) \) in a polynomial basis is the first stage in the two-step quasi-continuous method by generating an accurate parameterisation of the raw QMC Green’s functions on an imaginary time grid of arbitrary resolution. The second step is the systematic removal of the Trotter error by engineering a well defined extrapolation procedure of the Legendre basis coefficients on different Green’s functions \( G_\lambda(\tau) \), where \( \lambda \) is the imaginary time grid index, related to \( \Delta \tau_\lambda = \beta/N_\lambda \), with \( N_\lambda \) being the number of imaginary time points.

The procedure begins with defining a measure of error on each grid and time point, i.e. \( F^\lambda(\tau_i) \) such that,

\[
F^\lambda(\tau_i) = \sum_{l=1}^{N_\lambda} [g^\lambda_l - T_l] P_l[x(\tau_i)] = \alpha(\tau_i) \Delta \tau^2, \tag{30}
\]

where \( T_l \) are Legendre basis coefficients of quasi-continuous Green’s function \( G_{\text{QMC}}(\tau_i) \), absent of the systematic Trotter error. The \( \alpha(\tau_i) \) are the scaling coefficients of each imaginary time point.

The motivation for defining such an object is that we would like to find the set of coefficients \( \{g^\lambda_i\} \) such that \( F_{\text{QMC}}(\tau_i) = 0 \). To find this set of coefficients we define the following minimisation problem:

\[
\min_{T_l, \alpha(\tau_i)} [F^\lambda(\tau_i) - \alpha(\tau_i) \Delta \tau^2], \tag{31}
\]

over the completely continuous target parameters \( T_l \) and scaling coefficients \( \alpha(\tau_i) \). It is possible to simplify this procedure by introducing the matrix \( A_\lambda(\tau_i) \) such that,

\[
A_\lambda(\tau_i) = \sum_{l=1}^{N_\lambda} \frac{[g^\lambda_l - T_l]}{\Delta \tau^2} = \alpha(\tau_i), \tag{32}
\]

and noticing that across each grid \( \lambda \) that \( A_\lambda(\tau_i) \) remains unchanged. As a result, it is possible to map the minimisation problem of equation (31) to that of a simpler one,

\[
\min_{T_l} \left| \frac{A_\lambda(\tau_i)}{A_{\lambda-1}(\tau_i)} - 1 \right|, \tag{33}
\]

that is simply a minimisation with respect to the target parameters \( T_l \) and not the scaling coefficients \( \alpha(\tau_i) \).

Of critical importance in generating a controlled estimate of the high frequency tails of \( G_{\text{QMC}}(\omega_n) \), and thus also \( \Sigma_{\text{QMC}}(\omega_n) \), is the implementation of the additional constraint on the second moment of \( G_{\text{QMC}}(\omega_n) \), as discussed earlier. In principle, it is only sensible to add this constraint here, at the level of the Green’s function without any systematic error, since \( G_{\text{QMC}}(\omega_n) \) truly represents a physically relevant object while the \( G_\lambda(\omega_n) \) are inherently non-physical.

IV. APPLICATION TO SRVO\textsubscript{3}

1. DFT+DMFT for SrVO\textsubscript{3}

In this section we integrate the three main components of our DFT+DMFT scheme: the CASTEP DFT
code, the multi-orbital QMC-based solver, and the quasi-continuous imaginary time Green’s function processing protocol.

We apply this scheme to a full iterated DFT+DMFT calculation of SrVO$_3$ properties, and, where relevant, show comparison calculations made using the TRIQS CT-HYB solver.

The lack of any structural or magnetic phase transition behaviour over a broad temperature range makes the metallic transition metal oxide SrVO$_3$ an ideal test material against which to benchmark our CASTEP first-principles DFT+DMFT technique. There have been previous experimental and theoretical investigations of this material (using a variety of flavours of DFT+DMFT) against which results using our DFT+DMFT scheme can be compared [32, 35]. These studies highlight the necessity to explicitly account for electron correlation when calculating material properties, beyond those given by basic one-particle LDA. SrVO$_3$ has a perovskite structure with completely occupied oxygen 2$p$ bands, and partially occupied vanadium 3$d$ bands.

2. Calculation of SrVO$_3$ Green’s functions and self-energies

A basic electronic structure calculation for SrVO$_3$ was carried out in CASTEP. SrVO$_3$ has a perovskite unit cell (space group of crystal = 221: Pm-3m, -P 4 2 3) with lattice parameters $a = b = c = 3.8421\text{Å}$ [35], giving a unit cell volume $V = 56.72\text{Å}^3$. We have used a $20 \times 20 \times 20$ Monkhorst-Pack $k$-point grid, with 550 irreducible $k$-points. The pseudopotentials for all three elements Sr, V and O were taken from the C17 CASTEP set, and the plane-wave basis cut-off was automatically determined to be $653.07\text{eV}$. The calculations were performed at a temperature $T = 0.1\text{eV}$ ($\beta = 10\text{eV}^{-1}$).

Figure 1 shows the LDA band structure and DoS calculation for SrVO$_3$ obtained from CASTEP. The DoS for $O(2p)$ and $V(3d)$ orbitals are shown, along with the total DoS. There is an isolated set of three partially occupied bands around the Fermi level, which originate mainly from the triply degenerate vanadium $t_{2g}$ orbitals, $(d_{xy}, d_{xz}, d_{yz})$. The contribution from $V(e_g)$ and $O(2p)$ orbitals is minimal in the vicinity of $\epsilon_f$.

Particularising our QMC solver to the three vanadium $t_{2g} - d$ orbitals, $M = 3$, gives $\mu = 5U/2 - 5J$, and $M(2M - 1) = 15$ auxiliary fields are required for the multi-orbital quantum Monte Carlo simulation.

Previous studies with ED-like solvers have used $< 5$ bath levels to parameterise the Weiss field [35, 36] at inverse temperatures of $\beta \approx 10$. In our calculation we set the number of bath levels $N_b = 5$. The QMC solver can in fact deal with a large number of bath orbitals, and test calculations with $N_b$ as great as 9 showed no significant change to the accuracy of the impurity Green’s function at this temperature. It should be noted that the computational load of the QMC simulation scales as $\propto N_b^3 \beta$ [5]. Furthermore the fitting error of the parametrised Weiss field can be systematically controlled and in this calculation was of the order of $10^{-5}$.

The results of the impurity QMC Green’s function are shown in Figure 2a for the first iteration of the DMFT calculation for SrVO$_3$ across the interval $0 \leq \tau \leq \beta$. Two QMC Green’s functions are shown for one of the $t_{2g} - d$ orbitals for imaginary time discretisations $L = 50$ and $L = 80$, along with fits to these functions using the method described in Section III. The parameters chosen for this calculation are $U = 4$, $J = 0$ and $\beta = 10$, and match those used in other DFT+DMFT studies of SrVO$_3$. That said techniques such as constrained-RPA have been successfully used to give ab-initio estimates of model parameters for this kind of application [37]. The effect of the Trotter error is evident, and in Figure 2b a detailed portion of the Green’s function is shown, along with the Green’s function that results from our integrated fitting and extrapolation procedure. In this example a set of grids $L = 50, 60, 70, 80$ were used. However identical
results are also seen when using either $L = 50, 60, 70$ or $L = 60, 70, 80$, for instance. Additionally, a minimum of $N_l = 22$ Legendre polynomials are needed to correctly express $G(\tau)$ across the interval, but especially so near the boundaries where its derivative is larger. For comparison, the equivalent TRIQS CT-QMC result is also shown, and the agreement between the two results can be readily observed. This demonstrates that the QMC solver combined with our interpolation-extrapolation method is capable of generating quasi-continuous time solutions to the impurity problem.

To test the method further, an additional calculation was performed for $U = 8, J = 0.65$ and $\beta = 10$. Figure 3a shows the Green’s functions and their fits for the same time discretisations as before, illustrating the enhanced Trotter error for this larger $U$ value. In Figure 3b a comparison with the extrapolated Green’s function with a QMC calculation for $L = 200$ is shown. It is expected that the quasi-continuous solution will be very close to the very fine-scale (but computationally demanding) QMC calculation, and this is indeed the case.

The next step is the calculation of the self-energy, which involves Fourier transforming the Green’s function from $\tau$ to $i\omega_n$ i.e.

$$\Sigma(i\omega_n) = G_0^{-1}(i\omega_n) - G_QMC^{-1}(i\omega_n) \quad (34)$$

where $G_QMC^{-1}$ is the inverse of the calculated quasi-continuous time Green’s function. Figure 4 shows the self-energy result for the first DMFT iteration for a set of different $L$ values at $U = 8, J = 0.65, \beta = 10$. The extrapolated result, for both the real (Figure 4a) and imaginary (Figure 4b) parts, is a close match to the fine-scale QMC calculation. Clearly, discretisation of the imaginary time interval has a significant effect of self-energies.
FIG. 4. The DMFT self-energy at $U = 8$, $J = 0.65$ and $T = 0.1$ in the limit of $\Delta \tau \to 0$ for the first DMFT iteration. The excellent agreement of the extrapolated data in both real and imaginary parts with the $L = 200$ result is a consequence of the systematic nesting of the coarse-grid set of discrete BSS-QMC results, their fitting and subsequent extrapolation.

In Figure 5 the self-energy calculations are now taken to full DMFT convergence. It is not computationally feasible to perform a fully converged self-consistent calculation using an imaginary time discretisation of $L = 200$. As justified in Figure 4, by extrapolating a finite set of coarse discrete time grids it is, however, possible to replicate the continuous result with this method. Therefore, it is possible to converge the calculation using a subset of successively coarse grids that achieve the same accuracy as that of the more expensive fine grid calculation. We see that after three iterations the DMFT converges, as SrVO$_3$ is relatively weakly correlated.

The QMC solver and its associated quasi-continuous time protocol can be straightforwardly applied to ever-larger values of electron correlation. In Figure 6 calculations are shown for $U = 12$, $J = 0.65$, $\beta = 10$. For this large $U$ value the Trotter error on the Green’s function calculation is non-negligible. For one of the orbitals, Figure 6a shows the raw QMC Green’s function (with interpolations), along with an extrapolation to quasi-continuous time. Figure 6b shows that at this significant level of correlation the Trotter error is dramatically enhanced in contrast to smaller values of $U$. As a result, it is necessary to extrapolate with a finer set of discrete grids, i.e. $L = 80, 90, 100$, while also increasing the size of the Legendre basis to $N_l = 28$ polynomials. In doing so, it is possible to recover the correct form of the extrapolation scaling and therefore remove the non-trivial Trotter error induced by such strong interactions.

We conclude this section with some observations on the comparative computational efficiency of the TRIQS CT-QMC solver and the quasi-CT BSS-QMC solver method described above, using $U = 4$, $J = 0$ at $\beta = 10$.

Figure 7a compares the BSS-QMC calculation method run for 20 core hours to the 100 core hour CT-QMC(TRIQS) result. A factor five times speed-up gen-
FIG. 6. Upper panel: (6a) BSS-QMC impurity Green’s function in the strongly interacting limit of $U = 12$, $J = 0.65$ and $T = 0.1$. Lower Panel: (6b) The relative separation of the successive grids is indicative of the severe systematic Trotter error present in BSS-QMC calculations for large values of $U$. Using the quasi-continuous orthogonal polynomial method, by extrapolating respectively the $L = 80, 90, 100$ grids with $N_l = 28$ Legendre polynomials, illustrates how it can be remedied by comparing with the approximately continuous $L = 200$ data.

FIG. 7. Upper panel: (7a) Comparison of CT-QMC and BSS-QMC impurity Green’s functions over the full imaginary time range. Lower Panel: (7b) Detail of the Green’s function showing the comparison of a 50 core hour calculation with the CT-QMC result.

V. CONCLUSIONS

In recent years the DFT+DMFT framework has emerged as a powerful and effective procedure for undertaking *ab-initio* calculations of the properties of real materials when electron correlation effects are a significant influence. The method builds on the well-established capabilities of DFT, whilst seeking to enhance the way the electron correlation problem is addressed by utilising a quantum many-body physics approach.

A big step towards demonstrating the full potential of the DFT+DMFT scheme would be to attempt fully charge self-consistent calculations for many more mate-
FIG. 8. A calculation comparable with TRIQS can be achieved an order of magnitude faster using the BSS-QMC solver.

materials. However, to do this requires computationally efficient and accurate quantum impurity solvers that work over a very broad range of parameters. The work presented in this paper presents one possible implementation of a QMC-based solver that addresses this need. Additionally, applications such as the calculation of material equations of state, where many repeated DFT+DMFT calculations are needed, also require very fast and accurate QMC solvers. The solver we have presented here has the advantage of scaling linearly in inverse temperature, and moreover provides a quasi-continuous imaginary time solution. We have integrated the solver with the popular DFT code CASTEP, thus opening the way to fully charge self-consistent calculations on real materials. Though the validation of this approach has been done here against SrVO$_3$ it will be of particular value when used in the more challenging context of modelling the properties of f-band materials, and particularly for calculating materials’ equations of state where the availability of an accurate and computationally efficient solver will be essential.

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APPENDICES

Appendix A: Updating the chemical potential

During each cycle in the DMFT iteration it is necessary to adjust the chemical potential in the Bloch Green’s function (equation (13)) in order to maintain the overall correct number of electrons. Essentially, at each iteration a value of $\mu$ is sought that ensures the constancy of the overall electron occupancy, equation (19). A search algorithm employing the Brent method [35] is used to fix $\mu$.

Appendix B: Double-counting correction

CASTEP calculations already include a partial representation of electron correlation effects through the DFT exchange-correlation potential. In order not to count the contribution due to electron correlation twice the first step is to subtract the effect of the DFT potential through the “double-counting” approximation.

Double counting is not unambiguously resolved in DFT+DMFT, and there is no incontestable way to perform this correction. In CASTEP there is a choice of three double counting corrections: i) Fully localised limit (FLL); ii) Around mean-field limit, and iii) Held’s correction [10] [39] [40]. A selection can be made according to the modelling context. Taking each in turn:

i) FLL: In this approximation it is assumed that an orbital occupation is either 0 or 1 i.e. empty or full. Denoting $N_\sigma = \sum_m n_{m\sigma}$, $N_{tot} = \sum_\sigma N_\sigma$ and taking an average value for $U$ and $J$ as follows

$$U_{\text{avg}} = U = \frac{1}{(2l + 1)^2} \sum_{m,m'} U_{mm'}$$

and

$$U_{\text{avg}} - J_{\text{avg}} = U - J = \frac{1}{2(l + 1)} \sum_{m,m'} (U_{mm'} - J_{mm'})$$

the double counting is found to be

$$E_{\text{DC}} = \frac{1}{2} UN_{tot}(N_{tot} - 1) - \frac{1}{2} \sum_\sigma N_\sigma(N_\sigma - 1)$$

Differentiating with respect to $N_\sigma$ gives

$$V_{\sigma}^{\text{DC}} = U \left( N_{tot} - \frac{1}{2} \right) - J \left( N_\sigma - \frac{1}{2} \right)$$

This approximation is better suited to insulating systems.

ii) AMF: Here it is assumed that the average occupation of an orbital $(n_{m\sigma})$ is independent of $m$, so that

$$n_{m\sigma} = n_\sigma = \frac{N_\sigma}{2l + 1}$$
where $N_{\sigma}$ is the total occupation of the impurity site (with spin $\sigma$) and $l$ orbitals. The double-counting energy is given by

$$E_{DC} = U n_{\uparrow} n_{\downarrow} + \frac{2l}{2l+1} U - J \left( n_{\uparrow}^2 + n_{\downarrow}^2 \right)$$

and the potential by

$$V_{DC} = U \left( N_{\text{tot}} - \frac{n_{\sigma}}{2l+1} \right) - J \left( n_{\sigma} - \frac{n_{\sigma}}{2l+1} \right).$$

This approximation is better suited to metallic systems.

iii) Held’s correction: In this approximation an average Coulomb repulsion $U_{av}$, is introduced as follows

$$U_{av} = \frac{U + (l-1)(U-2J) + (l-1)(U-3J)}{2l-1}$$

where $l$ is the degeneracy of the shell. The double counting and associated potential as given as

$$E_{DC} = \frac{U_{av} N_{\text{tot}} (N_{\text{tot}} - 1)}{2}$$

and

$$V_{\sigma}^{DC} = U_{av} \left( N_{\text{tot}} - \frac{1}{2} \right).$$

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[1] A. N. Rubtsov, V. V. Savkin and A. I. Lichtenstein, Continuous-time quantum Monte Carlo method for fermions. Physical Review B 72, 035122 (2005).
[2] P. Werner, A. Comanac, L. de Medici, M. Troyer and A. J. Millis, Continuous-Time Solver for Quantum Impurity Models. Physical Review Letters 97, 076405 (2006).
[3] P. Werner and A. J. Millis, Hybridization expansion impurity solver: General formulation and application to Kondo lattice and two-orbital models. Physical Review B 74, 155107 (2006).
[4] E. Gull, A. J. Millis, A. I. Lichtenstein, A. N. Rubtsov, M. Troyer and P. Werner, Continuous-time Monte Carlo methods for quantum impurity models. Reviews of Modern Physics 83, 349 (2011).
[5] E. Khatami, C. R. Lee, Z. J. Bai, R. T. Scalettar and M. Jarrell, Cluster solver for dynamical mean-field theory with linear scaling in inverse temperature. Physical Review E 81, 056703 (2010).
[6] D. Rost, F. Assaad and N. Blumer, Quasi-continuous-time impurity solver for the dynamic mean field theory with linear scaling in the inverse temperature. Physical Review E 87, 053305 (2013).
[7] S. J. Clark, M. D. Segall, C. J. Pickard, P. J. Hasnai, M. J. Probert, K. Refson and M. C. Payne, First principles methods using CASTEP”, Zeitschrift fuer Kristallographie 220(5-6), 567-570 (2005).
[8] Materials Modelling using Density Functional Theory: Properties and Predictions. F. Giustino. Oxford University Press, Oxford, 2014.
[9] J. M. Tomczak, P. Liu, A. Toschi, G. Kresse and K. Held, Merging GW with DMFT and non-local correlations beyond. European Physics Journal (Special Topics) 226, 2565–2590(2017).
[10] A. Georges, G. Kotliar, W. Krauth and M. J. Rozenberg, Dynamical mean-field theory of strongly correlated fermion systems and the limit of infinite dimensions. Reviews of Modern Physics 68, 13 (1996).
[11] M. Iazzi and M. Troyer, Efficient continuous-time quantum Monte Carlo algorithm for fermionic lattice models. Physical Review B 91, 241118(R) (2015).
[12] S. Mehta, G. D. Price and D. Allé, Ab-initio thermodynamics and phase diagram of solid magnesium: A comparison of the LDA and GGA. The Journal of Chemical Physics 125, 194507 (2006).
[13] E. R. Ylvisaker, DFT and DMFT: Implementations and Applications to the Study of Correlated Materials. PhD Thesis, University of California Davis (2008).
[14] A. K. McMahan, Combined local-density and dynamical mean field theory calculations for the compressed lanthanides Ce, Pr and Nd. Physical Review B 72, 115125 (2005).
[15] E. Plekhanov, P. Hasnip, V. Sacksteder, M. Probert, S. J. Clark, K. Refson and C. Weber, Many-body renormalisation of forces in $f$-materials. Physical Review B 98, 075129 (2018).
[16] R. R. dos Santos, Introduction to Monte Carlo simulations for fermionic systems. Brazilian Journal of Physics 33, 36 (2003).
[17] M. Caffarel and W. Krauth, Exact diagonalisation approach to correlated fermions in infinite dimensions: Mott transition and superconductivity. Physical Review Letters 72, 1545 (1994).
[18] C. Weber, A. Americi, M. Capone and P. B. Littlewood, Augmented hybrid exact-diagonalization solver for dynamical mean field theory. Physical Review B 86, 115136 (2012).
[19] S. Sakai, A. Ritta, K. Held and H. Aoki, Quantum Monte Carlo study for multiorbital systems with preserved spin and orbital rotational symmetries. Physical Review B 74, 155102 (2006).
[20] S. Sakai, R. Arita, and H. Aoki, Numerical algorithm for the double-orbital Hubbard model:Hund-coupled pairing symmetry in the doped case. Physical Review B 70, 172504 (2004).
[21] J.E. Han, Spin-triplet s-wave local pairing induced by Hund’s rule coupling. Physical Review B 70, 054513 (2004).
[22] Y. Motome and M. Imada, A Quantum Monte Carlo Method and Its Applications to Multi-Orbital Hubbard Models. Journal of the Physical Society of Japan 66, 1872 (1997).
[23] Y. Motome and M. Imada, Numerical Study for the Ground State of Multi-Orbital Hubbard Models. Journal of the Physical Society of Japan 67, 3199 (1998).
[24] K. Held and D. Vollhardt, Microscopic conditions favoring itinerant ferromagnetism: Hund’s rule coupling and orbital degeneracy. European Physics Journal B 5, 473 (1998).
[25] R. Blankenbecler, D. J. Scalapino and R. L. Sugar, Monte Carlo calculations of coupled boson-fermion systems I. Physical Review D 24, 2278 (1981).
[26] S. R. White, D. J. Scalapino, R. L. Sugar, E. Y. Loh, J. E. Gubernatis, and R. T. Scalettar, Numerical study of...
the two-dimensional Hubbard model. Physical Review B 40, 506 (1989).
[27] E. Y. Loh and J. E. Gubernatis, Stable numerical simulations of models of interacting electrons in condensed matter physics. Chapter 4 in Electronic Phase Transitions ed. W. Hanke and Yu.V. Kopaev. Elsevier Science Publications (1992).
[28] H. J. Vidberg and J. W. Serene, Solving the Eliashberg equations by means of N-point Padé approximants. Journal of Low Temperature Physics 29, 179 (1977).
[29] P. K. V. V. Nukala, T. A. Maier, M. S. Summers, G. Alvarez, and T. C. Shulthess, Fast update algorithm for the quantum Monte Carlo simulation of the Hubbard model. Physical Review B 80, 195111 (2009).
[30] A. Weiße, G. Wellein, A. Alvermann and H. Fehske, The kernel polynomial method. Reviews of Modern Physics 78, 275 (2006).
[31] L. Boehnke, H. Hafermann, M. Ferrero, F. Lechermann, and O. Parcollet, Orthogonal polynomial representation of imaginary time Green’s functions. Physical Review B 84, 075145 (2011).
[32] V. I. Anisimov, D. E. Kondakov, A. V. Kozhevnikov, I. A. Nekrasov, Z. V. Pchelkina, J. W. Allen, S.-K. Mo, H.-D. Kim, P. Metcalf, S. Suga, A. Sekiyama, G. Keller, I. Leonov, X. Ren, and D. Vollhardt, Full orbital calculation scheme for materials with strongly correlated electrons. Physical Review B 71, 125119 (2005).
[33] I. A. Nekrasov, G. Keller, D. E. Kondakov, A. V. Kozhevnikov, T. Pruschke, K. Held, D. Vollhardt and V. I. Anisimov, Comparative study of correlation effects in CaVO$_3$ and SrVO$_3$. Physical Review B 72, 155106 (2005).
[34] F. Lechermann, A. Georges, A. Poteryaev, S. Biermann, M. Posternak, A. Yamasaki, and O. K. Anderson, Dynamical mean-field theory using Wannier functions: A flexible route to electronic structure calculations of strongly correlated materials. Physical Review B 74, 125120 (2006).
[35] B. Amadon, F. Lechermann, A. Georges, F. Jollet, T. O. Wehling, and A. I. Lichtenstein, Plane-wave based electronic structure calculations for correlated materials using dynamical mean-field theory and projected local orbitals. Physical Review B 77, 205112 (2008).
[36] A. Liebsch and H. Ishida, Temperature and bath size in exact diagonalisation dynamical mean-field theory. Journal of Physics, Condensed Matter 24, 053201 (2012).
[37] F. Aryasetiawan, M. Imada, A. Georges, G. Kotliar, S. Bierman and A. I. Lichtenstein, Frequency-dependent local interactions and low-energy effective models from electronic structure calculations. Physical Review B 70, 195104 (2004).
[38] W. H. Press, B. P. Flannery, S. A. Teukolsky and W. T. Vetterling, Numerical Recipes: The Art of Scientific Computing. Cambridge University Press, 2 edition, (1992).
[39] K. Held, Electronic structure calculations using dynamical mean field theory. Advances in Physics 56, 829 (2007).
[40] E. Pavarini, Mott transition: DFT+U vs DFT+DMF. Autumn School on Correlated Electrons: The Physics of Correlated Insulators, Metals, and Superconductors. Forschungszentrum Jülich (2017).