




# 1 solid\_dmft: gray-boxing ab initio DFT+DMFT utilizing 2 TRIQS

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DOI: [10.xxxxxx/draft](https://doi.org/10.xxxxxx/draft)

## Software

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Submitted: 01 January 1970

Published: unpublished

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## 8 Summary

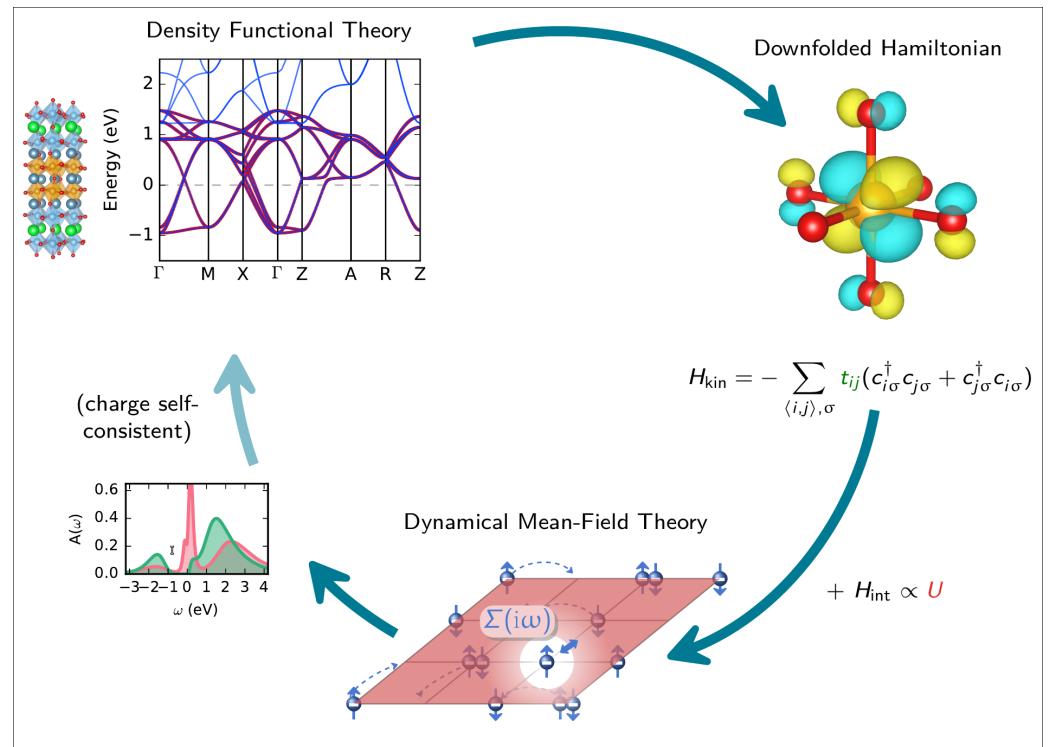
9 Although density functional theory (DFT) plus dynamical mean-field theory (DMFT) has  
10 proven successful in describing correlated electron systems for over two decades, it has only  
11 been very recently that ready-to-use software packages began to become available, with most  
12 scientific research carried out by self-written codes developed and used in research groups.  
13 Given the complexity of the method, there is also the question of whether a black-box approach  
14 is beneficial to the community or whether users should be able to implement the formalism  
15 themselves.

16 The goal of solid\_dmft is to find a middle ground, i.e. a gray-box tool as a ready-to-use  
17 implementation. This means that even if the code provides all functions needed for many  
18 standard DMFT calculations, the code is highly modular, based on open-source and community-  
19 developed software and therefore easily adjustable for specific applications and needs. Hence,  
20 the targeted user group of this software project is researchers that aim to perform DMFT  
21 calculations on top of their DFT simulations to describe the physics of strongly correlated  
22 electron systems, without the need of elaborate coding, but rather using a standardized input  
23 file to control the calculation. On the other hand, they can easily go beyond the implemented  
24 behavior by modifying the corresponding modules in the code.

25 The package is an MPI-parallelized scientific simulation code written in Python 3, allowing  
26 to perform ab initio DFT+DMFT calculations. solid\_dmft utilizes the publicly available  
27 TRIQS software library ([Parcollet et al., 2015](#)), which handles most numerical operations.  
28 The philosophy of the package is to increase reproducibility of DFT+DMFT calculations,  
29 provide clearer convergence metrics, and allow to run calculations for a large variety of systems  
30 without adapting the code manually, i.e. on this level similar to widely available DFT simulation  
31 packages.

## 32 Design Principles

33 The idea is to provide the full functionality of a full a DFT+DMFT calculation by merging the  
34 state-of-the-art implementations provided by the TRIQS library and its applications. This allows  
35 to easily run ab initio calculations for strongly correlated materials, as well as implement and test  
36 new features of TRIQS or benchmark new solvers against existing ones. solid\_dmft manages  
37 the calls of the necessary routines to run the DFT calculations, to create the downfolded  
38 Hamiltonian, solve the resulting Hubbard-like Hamiltonian via DMFT, postprocess the data to  
39 calculate physically meaningful observables, and allow for charge-corrected feedback via charge  
40 self-consistency. The full DFT+DMFT cycle is presented in [Figure 1](#).



**Figure 1:** Full CSC DFT+DMFT cycle. Starting from a DFT calculation (top left), a downfolded Hamiltonian and projector functions are created via optimized projections on a local basis set (top right). By adding a specified interaction Hamiltonian  $H_{\text{int}}$ , a full interacting electron problem is created, to be solved via the DMFT equations (bottom) in TRIQS. After convergence in DMFT is reached, physical observables are calculated (bottom left). For fully CSC calculations a charge density correction is added to DFT and the cycle is restarted.

41 The code is designed to run on top of a DFT calculation or model system providing a low-energy  
42 (downfolded) description of the periodic solid system. The DFT calculations can be performed  
43 with any code that is compatible with TRIQS/DFTTools (Aichhorn et al., 2016). The input for  
44 the DMFT calculation can be either provided directly as a Hamiltonian in reciprocal  $\mathbf{k}$ -space  
45 in a localized basis set, or in terms of the overlap between the localized basis set and the  
46 Kohn-Sham wavefunctions (so-called projector functions), and their respective eigenvalues.  
47 The DFT output is converted by TRIQS/DFTTools into an HDF archive in a standardized  
48 structure to be read or called by solid\_dmft. The code is designed to be modular in the same  
49 philosophy as the TRIQS software package, relying on TRIQS functionalities to perform basic  
50 operations. Therefore, we split each part of the simulation into separate stand-alone functions,  
51 to limit statefulness to a minimum and allow for an easy extension to include new features.  
52 The modularity of the program also allows to run, for example, the DMFT loop only via a call  
53 of a single function with well-defined input and output, i.e. without running solid\_dmft as  
54 a monolithic code. This ensures that the code can be used in other projects. A abstracted  
55 solver class implements the various impurity solvers available in triqs: cthyb, HubbardI,  
56 ForkTPS, ctint, and ctseg. Even though these solvers operate differently, solid\_dmft allows to  
57 seamlessly switch impurity solvers, with a simple input flag and adjusting the solver paramters.  
58 A fully charge self-consistent (CSC) interface is implemented for Quantum ESPRESSO and the  
59 Vienna ab-initio simulation package (VASP). solid\_dmft allows also to perform inhomogenous  
60 DMFT calculations, i.e. the treatment of multiple correlated and uncorrelated shells (impurity  
61 problems) while converging the full lattice self-energy. After self-consistency is reached, either  
62 via full CSC or just within the DMFT cycle, postprocessing scripts are available to perform  
63 analytic continuation of imaginary Green's functions, and to calculate spectral functions.

As of now, `solid_dmft` has been successfully used in various peer-reviewed research studies (Beck & Georges, 2022; Hampel et al., 2019, 2020, 2021; Merkel & Ederer, 2021; Zhang et al., 2022), and provides stable releases matching the releases of the TRIQS library. We provide a full documentation including several tutorials and a reference manual. Examples and benchmark calculations can be found in the tutorials section of the documentation. Furthermore, we utilize an extensive CI workflow on GitHub to test every pull request and commit.

## Statement of need

The number of ready-to-use DFT+DMFT codes is small, and all codes have been developed rather recently. Some of these operate in a black-box way, such as, for example, EDMFT, Amulet (Poteryaev & others, n.d.) and the DMFT implementation included in Abinit (Romero et al., 2020). Other software packages like DFTwDMFT (Singh et al., 2021) and DCORE (Shinaoka et al., 2021) follow a very similar strategy as `solid_dmft` but have not implemented different impurity solvers as of now. `solid_dmft` provides a flagship implementation of the TRIQS functionality to perform DFT+DMFT calculation and is easily extended to beyond-DMFT formalisms. This is beneficial both to developers of TRIQS applications, by enabling them to benchmark their applications in a well-tested framework, and to users, who benefit from the most recent features of TRIQS. Via the gray-box approach the software provides a robust and flexible implementation of the DFT+DMFT method, controlled via a single input file. It is developed in the spirit of a community code and supports external contributions that advance the capabilities of the program.

## Acknowledgements

This research was supported by ETH Zurich and the NCCR MARVEL, a National Centre of Competence in Research, funded by the Swiss National Science Foundation (grant number 182892). The Flatiron Institute is a division of the Simons Foundation.

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