Data-Driven Dynamical Mean-Field Theory: an error-correction approach to solve the quantum many-body problem using machine learning

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Machine learning opens new avenues for modelling correlated materials. Quantum embedding approaches, such as the dynamical mean-field theory (DMFT), provide corrections to first-principles calculations for strongly correlated materials, which are poorly described at lower levels of theory. Such embedding approaches are computationally demanding on classical computing architectures, and hence remain restricted to small systems, which limits the scope of applicability without exceptional computational resources. Here we outline a data-driven machine learning process for solving the Anderson Impurity Model (AIM) - the central component of DMFT calculations. The key advance is the use of an ensemble error-correction approach to generate fast and accurate solutions of AIM. An example calculation of the Mott transition using DMFT in the single band Hubbard model is given as an example of the technique, and is validated against the most accurate available method. This new approach is called data-driven dynamical mean-field theory (dMFT).

INTRODUCTION

It is evident that many materials which are being used in emerging technologies, such as spintronics, quantum computing and high temperature superconductivity for example, are characterised by interacting \( d \) and \( f \)-band valence electrons situated at the boundary between being localised and delocalised \([1]-[3]\). Computational materials design is hampered by the complexity of these materials (their chemistry and structure) and their significant electron-electron interaction effects. This diminishes both calculation effort and understanding, and hampers technological development.

In recent years significant progress in understanding the basic physics of strong electron correlation effects has been made by the dynamical mean-field theory (DMFT) method \([4]\). Moreover, the harnessing of DMFT to mainstream materials modelling methods, such as density functional theory (DFT) to produce DFT+DMFT theory, has generated material-specific capabilities which have begun to quantitatively address experimental contexts where correlation is a key determinant of material properties \([5]\).

Despite this, significant challenges remain when modelling real materials which feature many interacting electrons in multiple orbitals. At the heart of the LDA+DMFT method is the choice of quantum many-body solver for the Anderson Impurity Model (AIM). DMFT is based on a self-consistent mapping between local properties of a given material onto the AIM, a correlated impurity embedded in an infinite non-interacting bath. The AIM can be solved by high-level many-body methods such as quantum Monte Carlo \([4]\), or its extension, the continuous-time Monte Carlo approach (CTQMC) \([6]-[9]\). The latter provides an exact solution to the full rotationally-invariant AIM Hamiltonian within statistical error bars, but is limited to the imaginary time representation. The evaluation of real-frequency spectral quantities requires the use of ill-defined analytical continuation \([8], [9]\). Cluster extensions of DMFT which use multiple impurity sites suffer from the fermionic sign problem when inter-orbital hybridizations are present. Many body perturbation theory, and for example its implementation via the iterative perturbation theory for DMFT, are a solid starting point for solving quantum many-body problems in the limit of small repulsion \([10]\). They provide good agreement in this limit and can be extended to non-equilibrium and superconductivity.

Other solvers are based on the numerical renormalization group (NRG) \([10]\), which allow real axis calculations and access to Kondo physics, but remain challenging to extend for multi-orbital systems. Another method that provides solver solutions for real frequencies is the exact diagonalisation (ED) approach, where a finite size discretization of the AIM is used, through representation of the infinite bath in terms of a small number of effective bath-sites. In typical implementations, the bath size \( N_b \) is restricted because of the exponential growth of the Hilbert space with the total number of sites \( N_s \) (bath sites and impurity orbitals). Nonetheless, Lanczos-based algorithms allows one to deal with large Hilbert spaces, where the discretization at low temperature \([11],[12]\) is fine enough to compute observables accurately. Some success has been achieved by ED for multi-orbital systems with three or five orbitals \([13],[15]\), but in general the limitation in the bath size limits the scope of applicability for realistically describing transition metal oxides, for example. To achieve of the largest number of bath sites, ED calculations have been extended to handle the single impurity embedding problem, allowing up to \( O(100) \) \([16],[19]\) and \( O(300) \) \([20]\) uncorrelated bath sites. Although the latter approaches generate approximations of the zero temperature Green’s function, the construction of systematically high energy excited states remains challenging. Clearly there is no single applicable AIM solution that is effec-
tive for all contexts and all parameters ranges, and each method has its limitations. The stand-out technique is CTQMC, but it has poor low temperature scaling and is computationally demanding - using it to address f-band materials, for example, demands exceptional computational facilities. This means high-throughput material design applications are beyond the reach of most researchers.

The limitations of current solvers when addressing lanthanide and actinide materials, for example, has stimulated work on alternative solutions to the impurity problem. One conceptually different approach is an ensemble technique that draws upon the solutions of many approximate impurity solvers combined within a machine learning framework. Following the pioneering work of Arsenault [21] and the work of Rigo and Mitchell [22], we use a neural network-based machine learning approach to DMFT and construct a training set of physically reasonable hybridization functions, and determine the impurity Green’s functions corresponding to the training examples. The key feature of the quantum many-body solver presented here is that it uses a data-driven approach which focusses on training for the prediction of errors to calculate the Green’s function rather than attempting to directly predict this quantity. The method we present here is dubbed as "data-driven dynamical mean-field theory" - d³MFT. To demonstrate the robustness of the d³MFT method we give an example calculation of the Mott transition in a half-filled single band DMFT calculation for the Hubbard model on the Bethe lattice, and validate it against a highly accurate CTQMC solution.

The paper is organised as follows: In section I, we describe the machine learning approach for solving the Anderson Impurity Model. In section II, we review the validation and benchmark of the d³MFT approach. In section III, we apply the method to the Mott transition in the Hubbard model. Section IV provides our conclusions to this work. In Appendix A, we review the data augmentation strategy used for our machine learning based approach. In Appendix B, we describe the systematic study of the error scaling. In Appendix C, we provide representative test samples from the quantum database. In Appendix D, we discuss the training and validation loss of the neural network. In Appendix E, we discuss the inference of the trained neural network on representative candidate samples.

I. METHOD: MAPPING THE AIM TO A NEURAL NETWORK

The AIM is a prototypical physical model that explains quantum and thermal fluctuations in low-energy condensed phases of matter. Anderson first used the model and applied it to the Kondo phenomena to explain the increase of resistivity in metals at low temperature [23]. It has also been heavily used in the field of heavy fermion systems [24]. Decades after its initial formulation, it was given a new application in the context of DMFT in an attempt to provide a solution to the Hubbard model [1]. It is often the first port-of-call when going beyond the static mean field approach of DFT to model strongly correlated electronic phenomena in real materials using LDA+DMFT methods [5]. Despite the resurgence of intense research efforts focused on finding solutions to the AIM, they remain elusive. In particular, analytical methods are few, and when successful are inconsistent across the wide range of energy scales present in the model. Challenging parameter regimes are predominantly accessed by numerical methods which follow either the Quantum Monte Carlo (QMC/CTQMC) or Exact Diagonalisation (ED) methodologies. However, despite the conceptual and computational advances for both of these techniques in the past decades there are still major limitations. These include - speed, accuracy at low temperatures, number of correlated orbitals allowed, the fermionic sign problem - to name but a few. As a result, there is considerable interest in developing alternative impurity solvers that go beyond the current paradigm. Even so, a solver that can be (i) reliable across many parameter regimes and (ii) faster than ED or QMC would be a significant achievement, as it would open the field of strongly correlated electron physics to high-throughput techniques and ab-initio molecular dynamics.

The transferability of machine learning and neural network ideas to quantum physics is shown in a collection of seminal works [25–27], which demonstrated that these methods can be used on ab-initio derived databases to replace the computational bottleneck of DFT and Molecular Dynamics (MD). In the last decade, a number of supervised and unsupervised methods have been extended to many more problems in condensed matter physics, and in particular to many-body systems. For example, the pioneering work of Carrasquilla et al. [28] classified the paramagnetic to ferromagnetic phase transition in the two dimensional Ising model using data-driven methods alone, has instigated a number of supervised and unsupervised methods that go beyond the classification of phases of matter such as encoding the quantum many-body state with artificial neural networks [29–31]. Moreover, the QMC method has benefited in terms of speed [32–34] and accuracy [35], where the sign problem has been alleviated - using supervised learning techniques. Another technique that has benefited from data-driven solutions is the procedure of analytically continuing [35] a function of a complex variable to the real axis. However, most relevant to the work presented here paper is the application of machine learning methods to the AIM as in Ref. [21]. Here it was shown that the Green’s function of the AIM could be learned using kernel ridge regression on a database of AIM solutions and their physical parameters alone. Then in Ref. [36] the same authors applied this method to solve the DMFT equations for the single band Hubbard model which worked successfully in different limiting cases of the localised and delocalised limits. More recently, neural networks combined with different
exact solvers have been used to generate larger databases of solutions to the AIM \cite{37,38}.

In contrast to these developments, where the focus has been on building models that relate the physical parameters of the AIM to its Green’s function solution, we propose a new framework where the model relates almost exact solutions of the AIM with their exact solutions. To be specific, we use a neural network to learn the error between a family of solutions that are approximately correct, and what we regard as exact solutions, i.e. from Exact Diagonalisation. By using this approach we find that it retains the full many body features present in the AIM across a wide range of model parameters. Another feature of the work presented here is the automatic determination of a highly versatile and symmetric basis choice for the impurity Green’s functions that is used in the machine learning procedure. We show that the computational resources needed for a highly-accurate impurity solution with the machine learning-driven impurity solver are much reduced, since all it requires for training data is a set of approximate solutions. Ultimately we incorporate the data-driven solvers into a DMFT calculation for the Hubbard model for parameter regimes that span the Mott transition.

Outline of the machine learning protocol

In this section we sketch the overall framework for creating a data-driven machine learning solver for the AIM. To do this coherently we break the problem into three distinct units:

(i) generating a database of solutions for the AIM
(ii) training machine learning algorithms on the database to produce a data-driven solver, and
(iii) testing the efficacy of the data-driven solver.

An intermediate step (between step (i) and step (ii)) discusses some key data augmentations, parametrisations and transformations that enhance the performance of the neural network. This is presented in the Appendix A.

In what follows we address the single-band impurity problem, since there are reliable approximate solutions to it and exact solutions can be produced across a wide range of parameters with relative ease.

i) Database generation

The construction of a high-quality database of training samples is of paramount importance for any data-driven solution to a problem. Specifically, there must be sufficient “good” samples that relate a collection of inputs to a collection of outputs, such that after the training process a new set of input instances will produce the most likely outputs, via the trained neural network.

Before presenting the database construction details, we first need to view the AIM from a data-science perspec-

tive, rather than, a physics one. With this in mind, it is helpful to think of solving the AIM as a black box, and regard it only in terms of its inputs and outputs. In doing so, it will allow for an exact formulation its input samples $X = \{x_1, \ldots, x_{N_S}\}$ and output samples $Y = \{y_1, \ldots, y_{N_S}\}$, where $N_S$ is the number of database samples.

The single-band AIM is completely described by the following set of parameters $\{U, W, \epsilon, \beta, V\}$, where $U$ is the Hubbard $U$ parameter, $W$ is the half-bandwidth of the bath-states, $\epsilon$ is the impurity on-site energy, $\beta$ is the inverse temperature and $V$ characterises the impurity-bath coupling. Furthermore, we can break these contributions down for different sectors of the AIM Hilbert Space, where $\{U, \epsilon\}$ represent the physics of the impurity while $\{W, V\}$ represent the physics of the bath and $\beta$ is a property of the entire system. In practice, these parameters are then passed to an impurity solver and the Green’s function of the impurity, in a specific basis representation, is obtained as the solution of the AIM.

Indeed, in Ref. \cite{21}, the authors used as their input features $x_i = \{U_i, W_i, \epsilon_i, V_i\}$ and $y_i = \{G_{\text{and}}(\tau), G_{\text{and}}(l), G_{\text{and}}(i\omega_n), G_{\text{and}}(\omega)\}$ as the output vector, depending on the choice of basis. The basis choices above are the imaginary-time, Legendre, imaginary-frequency and real-frequency bases, respectively. In this case, each input vector has a dimension of 4 and each output vector has a dimension of $N_b$, where $N_b$ is the number of basis coefficients used to express the functional form of the solution. For example, the number of imaginary-time points is usually $N_\tau > 200$ to ensure subsequent reliable interpolations, while it is known that the number of Legendre coefficients $N_l < 50$ is more compact. \cite{21,39,40}. In other words, this approach addresses the question: given a set of AIM descriptors $x_i = \{U_i, W_i, \epsilon_i, V_i\}$ what is the functional form of its solution, which is basis dependent, i.e. $y_i = \{G_{\text{and}}(\tau), G_{\text{and}}(l), G_{\text{and}}(i\omega_n), G_{\text{and}}(\omega)\}$? This procedure is illustrated in Fig. \ref{IA}. In \cite{37}, the authors employ a similar approach for the input features, but attempt to learn the spectral function $A(\omega)$ instead. In summary, while neglecting the aspects of how the learning is done and what solvers are used to construct the database of solutions, the approaches so far have tried to relate the physical parameters of the Anderson Impurity Model to its solutions, which are defined in the imaginary-time, Legendre, imaginary- or real-frequency bases.

The approach used here is different with regard to the input vectors $x_i$, where instead we use a combination of approximate solutions to the AIM. We illustrate the idea in Fig. \ref{IA} for the $i$th AIM instance, which is still characterised by the set $\{U_i, W_i, \epsilon_i, V_i\}$, but the feature space increases by $N_\lambda N_b - 4$, where $N_b$ is the number of approximate solvers used, while $\lambda$ refers to the different impurity solver types, and $N_b$ is the number of basis coefficients used. For example, if we consider using two different approximate solvers $\{\lambda_1, \lambda_2\}$, then the
In this case, the length of the feature vector is 

\[ \{x_i, y_i\} \]

by trying to learn the relationship between the input parameters \( \{U_i, W_i, \varepsilon_i, V_i\} \) and the length of the output vector is 

\[ \{y_i\} \]

we attempt to learn the error between a set of approximate solutions of the AIM and its exact solution, generated using Exact Diagonalisation. Equivalently, Quantum Monte Carlo can be used as the exact benchmark, but the computational resources required to generate exact solutions using this technique for the single-band Hubbard Model are generally more than those required by ED. We try to learn the error of a set of approximations, rather than using just one, noting that different approximate AIM solvers have their merits in different parts of the AIM parameter space.

For instance, iterative perturbation theory is a well-known and successful diagrammatic method for solving quantum many-body problems in the weak-coupling limit, and can also capture some features of the strongly interacting limit \[11\]. Similarly, the Hubbard-I approach is exact in the weakly hybrised atomic limit, but can qualitatively fail outside of this parameter regimes. These approaches are quite often the first port of call in a many-body theorists arsenal when attacking a quantum many-body problem, due to their simplicity. While the weak-coupling expansion has its merits, for scenarios out at strong-coupling it can be qualitatively wrong. To address this, for example, the Non-Crossing Approximation (NCA) \[42\] is the lowest order strong-coupling perturbative method that sums up all diagrams without crossing hybridisation lines, and like the previous approaches also only requires moderate computational resources. It is also possible to use basis-truncated approximate ED solutions that use less bath sites to represent the Weiss field, which thus provide a more consistent coverage of the parameter AIM space, albeit with larger errors when small numbers of bath fitting parameters are used. Thus, a natural extension for data driven methods lies with the combination of different approximations, which span a wider range of the AIM parameter space, rather than just using one quantum solver alone.

Having established the form of the inputs and their associated outputs, we now discuss the construction of the database. We first deal with inverse temperature, \( \beta \), and unlike the rest of the parameters, the Green’s function is explicitly dependent on it, so each database is constructed at a specific \( \beta \). The next parameters to decide are \( \{U, \varepsilon, D\} \), all of which are randomly distributed between their extremal values. Therefore, for each instance of an AIM, random samples of each are drawn from the uniform distributions \( U \in [U_{\text{min}}, U_{\text{max}}]\), \( W \in [W_{\text{min}}, W_{\text{max}}]\) and \( \varepsilon \in [\varepsilon_{\text{min}}, \varepsilon_{\text{max}}]\).

Determining the hybridisation parameters is slightly more involved, because of how it manifests itself in the impurity solver. The ED solver necessitates a discrete representation of the bath parameters, while this is not true for other impurity solvers. Therefore, for the overall versatility of the machine learning protocol, we need to be able to deal with both discrete and continuous representations of the bath. We first take on the continuous representation, taking the Hilbert transform of specific form of the density of states, \textit{i.e.}

\[ G(z) = \int_{-\infty}^{\infty} \frac{A(\varepsilon)}{z - \varepsilon} d\varepsilon. \]  

(3)

There are two choices in the implementation, the first is a semi-circular DOS given by,

\[ A(\varepsilon) = \frac{2\sqrt{\pi W}}{(\pi W)^2} \Theta(W - |\varepsilon|), \]

(4)

and the second is a constant DOS, given by,

\[ A(\varepsilon) = \frac{\Theta(W^2 - \varepsilon^2)}{2W}, \]

(5)

where \( \Theta \) is the Heaviside step-function. Using either truncates the limits of integration in Eq. \[3\] from \(-W\) to \(+W\), hence \( W \) is known as the half-bandwidth. On the other hand, the discrete representation of the hybridisation function is given by,

\[ \Delta(i\omega_n) = \sum_{i=1}^{N} \frac{V_i^2}{i\omega_n - \varepsilon_i}, \]

(6)

where \( V_i \) and \( \varepsilon_i \) are the bath parameters, and which become an additional parameter in the database construction. Specifically, along with the number of samples in the database and the inverse temperature \( \beta \), the number of bath parameters determine the overall time it takes...
for the construction of the database. After the number of bath sites is chosen, random samples of each are drawn from the uniform distributions $V \in [V_{\min}, V_{\max}]$ and $\epsilon \in [\epsilon_{\min}, \epsilon_{\max}]$. To ensure that the discrete representation of the hybridisation function retains consistent physical characteristic, its bath parameters are all scaled to the chosen values of $D$, such that both $\epsilon_i$ and $V_i$ are normalised by it and $\epsilon_i$ is centered on its weighted arithmetic mean with respect to $V_i^2$. Alternatively, it is possible to create discrete representations of the bath by treating $\epsilon_i$ and $V_i$ as fit parameters in Eq. (6) to a continuous representation generated from the half-bandwidth $W$.

| Sample Index/Basis | Inputs $X$ | Outputs $Y$ |
|-------------------|------------|-------------|
| $S_1$              | $\{G_1^\lambda_1(\tau), G_1^\lambda_2(\tau), \ldots, G_1^\lambda_M(\tau)\}$ | $G_{\text{and}}(\tau)$ |
| Time               |            |             |
| Legendre          | $\{G_1^\lambda_1(l), G_1^\lambda_2(l), \ldots, G_1^\lambda_M(l)\}$ | $G_{\text{and}}(l)$ |
| $S_2$              | $\{G_2^\lambda_1(\tau), G_2^\lambda_2(\tau), \ldots, G_2^\lambda_M(\tau)\}$ | $G_{\text{and}}(\tau)$ |
| Time               |            |             |
| Legendre          | $\{G_2^\lambda_1(l), G_2^\lambda_2(l), \ldots, G_2^\lambda_M(l)\}$ | $G_{\text{and}}(l)$ |
| $\vdots$          |            |             |
| $S_N$              | $\{G_N^\lambda_1(\tau), G_N^\lambda_2(\tau), \ldots, G_N^\lambda_M(\tau)\}$ | $G_{\text{and}}(\tau)$ |
| Legendre          | $\{G_N^\lambda_1(l), G_N^\lambda_2(l), \ldots, G_N^\lambda_M(l)\}$ | $G_{\text{and}}(l)$ |

Figure 2: The database of single-band Anderson Impurity for $M$ approximate ($\lambda_1, \lambda_2, \ldots, \lambda_M$) solutions and exact solutions (from ED) for $S_N$ distinct $\{U_i, W_i, \epsilon_i, V_i\}$ parameter values, where $U$ is the correlation energy, $V$ is the hybridisation, $\epsilon$ is the impurity filling and $W$ is the half-bandwidth in the bath. The database is categorized according to whether a solution is an input $X$ or an output $Y$ for a machine learning model. The Green’s functions are shown to be expressed in the imaginary-time basis and Legendre basis, however they can also be expressed in different representations.

The next step is to generate the database that will be used for the training of the data-driven model. To do this, each instance of $\{U_i, W_i, \epsilon_i, \Delta_i(i\omega_n)\}$ is passed to the set of approximate solvers $\{\lambda_1, \ldots, \lambda_N\}$ as well as one exact solver. In this case, the exact solution is obtained by the ED algorithm using a large number of bath sites, generally between 4 – 6 is enough to ensure a converged solution for the single-site AIM. Fig. 2 illustrates what a typical database looks like, where in this case there are $S_N$ samples that each use $M$ approximate solvers to generate the input data for the model. We also note that we have exclusively addressed the inputs and outputs of the database as being defined in the imaginary-time basis or Legendre polynomial basis, but this can be extended to any other basis representation (e.g. the real-frequency $\omega$ basis) with the appropriate transformation on each entry in the database.

**ii) Training a model solver**

We now outline the details of the multivariate maximally connected neural network regression model that is used for the training against the database we have just constructed. As established above, the set of inputs for the model are $X = \{x_1, \ldots, x_N\}$, where $x_i$ is a set of different approximate solutions of the AIM, while the outputs are $Y = \{y_1, \ldots, y_N\}$, where $y_i$ is an exact solution of the AIM given by ED. For all models trained in this section, they proceed by minimising the cost function,

$$C(X, Y, \alpha) = \frac{1}{N_s} \sum_{i,j}^N \|y_j - g_{\alpha}(x_j)\|^2,$$

with respect to the parameters $\alpha$ to produce a model $g_{\alpha}(x_i) := G^M(x_i)$, where $G^M(x_i)$ is the model Green’s function of the problem. $G^M(x_i)$ is constructed such that the error between it and the true solution $y_i$ is minimised, and therefore $G^M(x_i)$ corrects for the error between the approximate solution $x_i$ and the exact one $y_i$, for all $N_s$ entries in the database. The neural network we use is shown as a schematic in Fig. 3. In the input layer, each neuron evaluates,

$$f(x, w + b) = f \left( \sum_{i,j} w_{ij} G^\lambda_i(k_i) + b_1 \right)$$

with $f(\ldots)$ being the activation function of the input layer neurons (coloured pink), index $i$ is associated to the feature (i.e mesh point) and index $j$ indicates the approximate solver used, $w_{ij}$ are the set of neural weights, and $x_i$ is in general of the format $\lambda_N$ entries, despite the depiction in Figure 8 that suggests that the number of approximate solvers is 2. This procedure then repeats itself as the values propagate forward through the network such that $f(x, w + b)$ of each neuron are used as the inputs for the next layer in the network, until eventually the output layer is reached. As the neural network is being used to solve a regression problem, the output layer applies a linear activation function to its neurons, which doesn’t modify its input data. Therefore, when the output layer is reached the cost function in 7 is evaluated for a "mini-batch" of samples, after which the weights throughout the entire network are updated in accordance with the backpropagation method. This procedure is then repeated until $G^M$ is found with weights $\alpha$ that minimise $C(X, Y, \alpha)$.

It is important to keep in mind the number of parameters in the model $g_{\alpha}(x_i)$ so as to avoid potential overfitting scenarios. For the neural network presented in Fig. 3 the total number of parameters $N_{\alpha}$ can be determined...
Typically a hyperparameter grid search \cite{43} is employed over these parameters for a low value of the epoch number. In doing so, the model is trained multiple times as it scans across different values of its parameters. Ultimately it will produce the final value of the cost function for both the training and validation datasets, where the minimum will inform the final choice of parameters to be used for a comprehensive training of the database.

### iii) Physical applications

After training the neural network, we are left with a data-driven impurity solver for a specific inverse temperature $\beta$. In Fig. 4, we propose a workflow for solving the single-band Hubbard model on a Bethe lattice within the DMFT approximation using a data-driven model that has been trained (in this example) on two approximate solutions. To set up the problem, the local impurity Green’s function takes the form,

$$G(i\omega_n) = \frac{2}{(i\pi W)^2} \int_{-\infty}^{\infty} d\epsilon \frac{\sqrt{W^2 - \epsilon^2}}{i\omega_n - \epsilon} \Theta(W - |\epsilon|),$$  \hspace{1cm} (11)

where $W$ is the half-bandwidth of the system. After setting up the problem the only difference between this scheme and the standard DMFT scheme \cite{4} is that instead of solving the AIM once per DMFT iteration here it is solved twice with the two different approximate solvers ($\lambda_1$ and $\lambda_2$), which are then used as inputs for the trained model $\mathcal{M}$. The trained model then produces the impurity Green’s function $G_{\text{ML}}(\alpha)$ which is subsequently used to calculate the momentum independent self-energy $\Sigma(i\omega_n)$. Additionally, a mixing parameter $\alpha$ is imposed when going from one iteration to the next, i.e. $\Delta(\tau) = \alpha \Delta_{\text{AIM}}(\tau) + [1 - \alpha] \Delta(\tau)$ to ensure that the self-consistency doesn’t precipitously diverge from the previous iteration. By solving this model, it is possible to analyse the Mott transition using the d$^3$MFT data-driven machine learning protocol.

### II. BENCHMARK AND VALIDATION: A NEURAL NETWORK FOR THE AIM

Here we present results that pertain to the d$^3$MFT framework outlined above. We begin with a discussion of the different aspects of the generated databases, and this is followed by detail on the training of an artificial neural network with those generated databases. We conclude by illustrating how the generated data-driven solver is able to capture the Mott transition in the half-filled single-band Hubbard model using the DMFT scheme presented above.
Database of solutions for the Anderson Impurity Model

Using the procedure outlined in the previous section we generate a database of size $N_s = 10,000$ at inverse temperatures of $\beta = \{1, 2, 5, 10, 20, 50\}$ eV$^{-1}$ over the parameter ranges indicated in Table I for discrete sets of bath parameters. The range of temperatures chosen represent the high-temperature and intermediate temperature limits, whereby the features of the Green’s function are smoother, and hence our choice of the range. Each database entry constitutes a random combination of all parameters in Table I. The parameters chosen cover a range of physical features, for example the Hubbard $U$ is uniformly randomly sampled over the range $\{1, \ldots, 10\}$, in addition to $W \in \{1, \ldots, 10\}$ and $\varepsilon \in \{-1, \ldots, 1\}$, then the various combinations of $U, W, \varepsilon$ result in metals or insulators. Take for instance if $\{U, W, \varepsilon\} = \{8, 2, 0\}$ then the result is insulating, and if $\{U, W, \varepsilon\} = \{2, 8, 0\}$ the result is metallic. In Table I we clarify the notation for the Approximate solvers ED-[1,2,3]. ED-1 means solving the AIM with one bath site only, and hence results in a truncated approximation to the exact ED solution of the AIM (which in this case uses 4 bath sites). We justify the inclusion of these ED solvers since they are exponentially faster than the true ED solution and are themselves approximate solutions, similar to IPT or Hubbard-I.

| $U$ (eV)    | $\{1, \ldots, 10\}$ |
|------------|---------------------|
| $N_{\text{bath}}$, $\varepsilon$, $V_i$ | 4 |
| $W$ (eV)    | $\{1, \ldots, 10\}$ |
| $\varepsilon$ | $\{-1, \ldots, 1\}$ |
| $\beta$ (eV$^{-1}$) | $\{1, 2, 5, 10, 20, 50\}$ |
| $N_{\text{samples}}$ | 10,000 |
| $S$         | Hubbard-I, IPT, NCA, ED-[1,2,3] |

Table I: Parameter selection for the database of AIM solutions shown in Fig. 5 where $\{p_i, \ldots, p_f\}$ denotes that a parameter is randomly selected from this interval $[p_i, p_f]$. $U$ is the Hubbard interaction, $N_{\text{bath}}$ stands for the number of bath sites, $W$ is the Half-bandwidth, $\varepsilon$ is the electron on-site energy, $\beta$ is the inverse temperature, $N_{\text{samples}}$ is the number of database entries, and $S$ denotes the total ensemble of approximate quantum solvers used in the ML approach. ED-[1,2,3] denotes the exact diagonalisation solver with respectively 1, 2, 3 bath sites.

Furthermore, in Fig. 5 we show the distribution of all chosen parameters for the 10,000 samples in the database corresponding to $\beta = 20$ eV$^{-1}$. As expected, $\{U, W, \varepsilon\}$ are distributed evenly, $N_{\text{bath}} = 4$ is constant as the number of bath-sites is not changed, and $\{\varepsilon_i, V_i\}$ are chosen by normalising to $W$. While the illustrated database is not the only one that could be considered, it is not a special choice. For all other databases we analysed the distribution of parameters behaves similarly to the $\beta = 20$ eV$^{-1}$ case presented.
In Appendix C, we review the strength and weaknesses of the Hubbard-I, IPT and NCA solvers for representative samples of the database against the corresponding exact diagonalisation results using the solver of Ref. [11] at temperatures $\beta = \{1, 20, 50\}$ eV$^{-1}$ shown in Figure (22). The latter provide valid approximations of the AIM in different limits (Hubbard-I and IPT are good in the weakly hybridised limit, NCA is a good approximation for stronger interactions). In general, the Hubbard-I, IPT and NCA solvers are however in quantitative and qualitative disagreement. For example, the Hubbard-I solver indicates for highly hybridised AIM an insulating solution when in reality the system is metallic. Nevertheless, we emphasise that this behaviour is expected and welcomed, since the end-goal is to systematically correct for the error between the approximate and exact solutions. A systematic analysis of the RMSD error for each of these solvers with respect to the true solution is also reported in Appendix B. This analysis reveals some of the in depth features of the approximate solutions, and highlights potential parameter choices that the network might struggle with.

A Neural Network Impurity Solver

The very first step to training a machine learning model is the hyperparameter grid-search over its independent parameters using tensorflow [15]. Specifically, for our neural network we coarse-grain the number of epochs to 20, set the inverse temperature to $\beta = 1$ eV$^{-1}$ and scan across all combinations of parameters in Table II. Ultimately, 401 different neural networks are trained and the combination of parameters with the minimum cost function $\sim 10^{-6}$ is given by that combination of parameters shown in the "Optima" column of Table II. Additional fixed parameters in the grid search are: evenly spaced time grid, Hubbard-I, IPT and NCA solvers as inputs $x_i$ to the neural network as they all require minimal computational resources in comparison to the ED methods, no data augmentation, and the $T_i$ imaginary-time transformation from Fig. [17] It is noteworthy that either increasing the complexity of the network, i.e. increasing its depth beyond 1, or increasing the learning rate to an order beyond $10^{-2}$ has detrimental effects on the minimisation of the cost function. Practically, it would be computationally prohibitive to perform this grid search for every $\beta$ and their additional free-parameters. In what follows, all networks use the optimal values as specified in Table II and use an 80/20 cross-validation split, i.e. 80% training data and 20% validation data.

With the networks’ hyperparameters chosen, we now outline how the various models $M_i$ are constructed and are used to test the best set of approximate $x_i$ inputs to produce the minimal cost function. These models are differentiated by their input parameters as tabulated in Table III. For example, $M_1$ consists of the IPT, NCA and Hubbard-I solvers as inputs, while $M_3$ combines many more. If each approximate input has $N_b$ mesh points then the feature space for $M_1$ is $3N_b$, while for $M_3$ it is $6N_b$. As a result, the number of weights in the network for $M_3$ will be greater than for $M_1$.

| Hyperparameter | Range | Optima |
|----------------|-------|--------|
| $\eta$         | [0.0001, 0.0002, 0.001, 0.01] | 0.0002 |
| Mini-batch size| {4, 8, 16, 32, 64} | 8 |
| Optimiser      | (Adamax, Nadam) | Nadam |
| Activation functions | {elu, relu, tanh} | tanh |
| Hidden Layers  | {1, 2, 3, 4} | 1 |

Table II: Hyperparameter grid-search over the neural network parameters with a fixed number epochs $= 20$ and $\beta = 1$ eV$^{-1}$

In the Appendix A we propose a collection of data scaling transformations of the input and output data which improve the training of the neural network in the imaginary-time and Legendre bases. Fig. 6 presents the validation loss for these scenarios, for $\beta = 1$ eV$^{-1}$ and model type $M_1$. For the Legendre basis the effect of data transformations is quite significant, as shown in Fig. 6. Here we see that by applying a tanh-type Legendre transformations that the final value of the loss can be improved by at least 2 orders of magnitude, reduced from $10^{-4}$ to $10^{-6}$. We expect the effect of this transformation to be enhanced for larger values of $\beta$ (lower temperatures), where the range of $G_l$ can greatly exceed the value of unity, and therefore necessitates the application of an appropriate data transformation. At higher temperatures (i.e lower $\beta$), the Legendre coefficients are often bounded close to unity, and so the neural network is less sensitive to the untransformed input as compared to lower temperatures.

We show the final value of the validation loss after 30 epochs for the equidistant imaginary-time basis in Fig. 6. For $\beta = 1$ eV$^{-1}$ the converged values are all within $10^{-6}$ of each other, indicating a minimal impact of this transformation scheme at low values of $\beta$. Hereafter, we use the $T_2$ transformation in imaginary-time and $T_{1/2}$ for the Legendre basis. The hypothesis that as $\beta$ increases, i.e. temperature falls, the network becomes harder to train is affirmed, indicating that a deterioration of the error landscape shown in Fig. 18a and Fig. 19a is manifested in the predictive qualities of the network. Notably,
using an adaptive $\tau$-mesh over an equidistant one is more effective in improving the quality of the network at high temperatures, as Fig. 7 illustrated for different values of inverse temperature $\beta$ in the imaginary-time basis with equidistant and adaptive mesh spacings.

To assess this further, in Fig. 9 we show the validation loss function after 40 epochs at different values of $\beta$ for all models $\mathcal{M}_i$ in Table III. The efficacy of the IPT solver is shown here, as all models that do not include it underperform relative to it. In particular $\mathcal{M}_4$ and $\mathcal{M}_6$, which both include the NCA solver (however IPT is absent), are consistently the worst performers. At small values of $\beta$ it is possible to use any of the models and still achieve an accuracy of up to $10^{-5}$, however as $\beta$ increases, the inclusion of the IPT accounts in some cases for an order of magnitude improvement in the loss. The full training and validation loss functions for the additional models studied in Table III are in Appendix D.

In Figure (10) we show the value of the cost function when trained in the Legendre basis for models $\mathcal{M}_1$ and $\mathcal{M}_3$ using the $T_1/2$ data transformation. We note that each value of $\beta$ necessitates a different mesh value $N_b$, as chosen according to the reconstruction criteria detailed in Figure (15) of Appendix A. Moreover, the chosen mesh
value $N_b$ is a critical factor in the complexity of the network, as determined by Eq (9). We observe similar results for the training in the Legendre basis, i.e., that higher temperatures are more amenable to the training procedure and that including more approximate solvers increases accuracy of the final validation loss. Notably, we see that the overall value of the loss function for $\mathcal{M}_3$ and $\mathcal{M}_4$ is lower than that predicted in the imaginary-time basis, regardless of the ensemble of solver types and transformation used. Therefore, we see that by executing suitable basis transformations which are supplemented by a multitude of different approximate solvers then the accuracy of the overall predictive quality of the neural network can be improved.

In Appendix E, we describe the prediction of the optimised network for three representative samples, respectively at small $U \approx 1.88 - 1.99$ eV and larger $U \approx 6.2$ eV in both the Legendre and imaginary-time bases at different temperatures. Of course, we have presented only two possibilities of AIM configurations, where in principle the AIM parameter space is much larger. In general, while it is true the predictive quality in the entire phase-space of $\{U, W, \varepsilon\}$ is poorer for larger values of $\beta$ (c.f. Fig. 9), we have demonstrated that it is still better than what approximate solvers can achieve for some parameter regimes, and even sometimes with effectively error-free solutions in both the imaginary-time and Legendre polynomial bases. Hence, $d^3$MFT provides a systematic improvement over the studied parameters and in many situations.

Moreover, it is noteworthy that the AIM parameter space for real materials is quite restricted, where for example, due to the screened Coulomb interaction, there are few materials with large Hubbard $U$ values, which lie in the Mott limit, with a screened Coulomb interaction $\approx 8 - 10$ eV. The archetypal La$_2$CuO$_4$ is one of the notorious case, where large $U$ effects are obtained by the strong localisation of the atomic $d$-shell. Most charge transfer correlated systems however have Coulomb interactions in the $3 - 6$ eV range, poising $d^3$MFT as a good candidate for high throughput calculations in such cases. This same line of thinking applies for $\{W, \varepsilon\}$ too, and opens up this approach for further dimensional reduction by taking only physically relevant parameter regimes into account.

### III. RESULTS: DATA DRIVEN DYNAMICAL MEAN FIELD THEORY - $d^3$MFT

The motivation behind developing the data-driven impurity solver ($d^3$MFT) is to alleviate DMFT calculations from the intensive computational burden imposed by Ex-
Figure 10: Cost function for the Legendre mesh for different inverse temperatures $\beta$ using as input to the neural network the solvers a [Hubbard-I, IPT, NCA] b [IPT, NCA, ED-1, ED-2, ED-3, Hubbard-I]. IPT is the Iterated Perturbation Theory Solver, NCA is the Non Crossing Approximation solver and ED-[1,2,3] are the truncated ED solvers.

Figure 11: CTQMC and Neural Network solvers used for a DMFT prediction of the quasiparticle weight $Z$ as a function of $U$ at $\beta = 20$ eV$^{-1}$ and $W = 1.0$ eV for the single-band half-filled Hubbard model on a Bethe lattice.

**IV. CONCLUSION**

We have developed an extensible data-driven framework that can readily create error-free neural network impurity solvers capable of capturing the Mott transition using DMFT for the Hubbard model. We call our new method d$^3$MFT. Compared to standard machine learning methods our approach has a relatively small numerical footprint and is straightforward to implement, once the set of impurity solvers are specified. As expected, and in line with Ref. [21], the convergence and overall predictive quality of the training procedure is sensitive to the basis representation of the impurity Green’s function. Using an adaptive imaginary-time $\tau$ mesh or Legendre coefficients under a $\tanh$-type basis transformation to express the Green’s function, supplemented with
Figure 12: The resultant spectral functions as computed with the Neural Network solver for values of $U = 3\, eV$ and $U = 8\, eV$, $\beta = 20\, eV^{-1}$ and $W = 1.0\, eV$ using the Pade method for analytical continuation.

data-augmentation techniques through exploiting physical symmetries, is the most effective route to minimising the overall error of the network's cost function. Nevertheless, using the raw untransformed datasets in either of the two bases can still yield impressive predictive capabilities, often in better agreement with the exact solution over the approximate solvers. The network shows a strong dependence on the temperature at which the database is built for as well as the approximate solvers used. We find that training at higher temperatures using approximate solutions results in consistently reliable results. We anticipate that improved results at higher temperatures could be attained by extending into the intermediate representation \cite{17}, which is currently the most compact representation of the Green's function available. However, we also mention that since the entire protocol is a supervised learning method, it implicitly depends on the quality of the exact solution obtained. Therefore, the quality of the data-driven solver produced at the end is limited by this and will always rely on the precision of the exact model that is used.

A natural extension to the single-band results presented here is to generalise the method for a multi-orbital system. This will enable material-specific calculations to be undertaken rather than model Hamiltonians. However, the extension to multi-orbital physics, especially for realistic materials, must take into account various processes that arise from the Slater-Kanamori representation, such as Hund’s coupling. Furthermore, this aspect means that the simplicity of the perturbative solutions is lost, which will have an effect on correcting their errors in the training process. Additional extensions for this machine learning based approach includes a systematic study of doping, temperature and magnetism for the single-band Hubbard Model. To achieve this, the low-temperature prediction capabilities of the method need to be improved. Notwithstanding these considerations, we expect our proposed framework will be a valuable in stimulating efforts in this direction, and ultimately complement the ongoing research efforts to devise fast and accurate solvers for the AIM so that DMFT calculations can be applied to a much broader class of problems that are out of reach to current methods.

ACKNOWLEDGEMENTS

CW acknowledges insightful and stimulating discussions with Andrew Mitchell. CR thanks AWE for support through its Future Technologies fund. CW is supported by grant EP/R02992X/1 from the UK Engineering and Physical Sciences Research Council (EPSRC). ES is supported by the EPSRC Centre for Doctoral Training in Cross-Disciplinary Approaches to Non-Equilibrium Systems (CANES, EP/L015854/1). IR and FJ acknowledge the support of the UK government department for Business, Energy and Industrial Strategy through the UK national quantum technologies programme (InnovateUK Industrial Strategy Challenge Fund (ISCF) QUANTIFI project). This work was performed using resources provided by the ARCHER UK National Supercomputing Service and the Cambridge Service for Data Driven Discovery (CSD3) operated by the University of Cambridge Research Computing Service (www.csd3.cam.ac.uk), provided by Dell EMC and Intel using Tier-2 funding from the Engineering and Physical Sciences Research Council (capital grant EP/P020259/1), and DiRAC funding from the Science and Technology Facilities Council (www.dirac.ac.uk).

ADDITIONAL INFORMATION

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Appendix A: Data augmentation and data symmetrisation

Once the database of approximate and exact solutions is constructed, and before the data is passed to the machine learning algorithm for training, there are a number transformation operations that allow the database to be augmented through symmetry operations, without having to re-run the impurity solver. In this appendix we explain a number of ways to both extend and transform the database in ways that are amenable to learning a model. For simplicity, we assume a database under consideration is expressed in either the imaginary-time basis on a discrete, evenly spaced grid or the Legendre polynomial basis.

The first symmetry operation makes use of the fact that the Green’s function can be decomposed into its
symmetric and anti-symmetric contributions by decomposing it into the Legendre basis,

\[ G^S(\tau) = \sum_{l>0 \text{ even}} \frac{\sqrt{2l+1}}{\beta} P_l[x(\tau)] G_l \]

and

\[ G^{AS}(\tau) = \sum_{l>0 \text{ odd}} \frac{\sqrt{2l+1}}{\beta} P_l[x(\tau)] G_l, \quad (13) \]

where \( G^S(\tau) \) and \( G^{AS}(\tau) \) are respectively the symmetric and anti-symmetric parts of total Green’s function which give the total Green’s function when summed, i.e.,

\[ G(\tau) = G^S(\tau) + G^{AS}(\tau). \quad (14) \]

In practice, if performed in the \( \tau \) basis this requires an intermediate step of generating the Legendre coefficients, or reading them in from a database which has them stored already. For the size of the databases dealt with in this study (< 40,000) this feature can be added practically at no additional computational cost. Physically, the symmetric part of the Green’s function represents the physics at half-filling while the anti-symmetric component encodes the information away from half-filling. This operation need not only be used for the augmentation of the database, it can similarly be used for partitioning it. Specifically, instead of training a model on both the symmetric and anti-symmetric components simultaneously, two separate models can be trained on the symmetric and anti-symmetric components separately, after which they are recombined to produce the total answer in Eq. (14). The same procedure can be followed in the Legendre basis, where the symmetric part of \( G_l \) is encoded in the even coefficients and the anti-symmetric part in its odd ones. For both bases, this operation allows the database to be augmented by a factor of two and is shown in Fig. 13a.

The second symmetry operation on \( G(\tau) \) that we can consider is particle-hole equivalency, i.e. \( G^c(\tau) = G^h(\beta - \tau) \) where \( G^c \) is the electron Green’s function and \( G^h \) is the hole Green’s function. Therefore, for every entry in the database that is away from half-filling, the corresponding electron (if hole-type) or hole (if electron-type) Green’s function can be generated simply by flipping \( G(\tau) \), and is illustrated in Fig. 13b. If, however, the database is expressed in the Legendre basis instead, this transformation requires that the odd coefficients be multiplied by \(-1\). We note that for both basis representations, this can double the size of the database.

**Data transformation**

In addition to augmenting the database by exploiting symmetry operations, the data must also be transformed into a representation appropriate for how the training data will be manipulated. Here we outline the common ways in which the data are transformed. For lower temperatures the shape of \( G(\tau) \) becomes steeper at its boundaries and flatter further away. If a regularly spaced out \( \tau \)-mesh is used this presents a problem as the behaviour near the boundary may not be taken into account, or in the worst case missed completely. From a data-centric perspective, this would mean that different \( G(\tau) \) functions which describe qualitatively different physics could look very similar to a learning model, since most of the data passed to it will be constant. To remedy this issue it is possible to transform the \( \tau \) domain so that it is sampled more near its boundaries and less near the centre. This can be achieved by transforming the \( \tau \) points from being evenly distributed to being distributed according to a sigmoid-type function \( \sigma \).

Thus, the transformation from a regular equidistant mesh to an adaptive mesh amounts to \( G(\tau) \rightarrow G(\sigma(\tau, \delta)) \), where \( \delta \) is a smearing parameter, and is illustrated in Figure 13: Data augmentation schemes for \( G(\tau) \) where **a** illustrates the extraction of the odd and even parts of \( G(\tau) \) via Legendre transformations and **b** exploiting the particle-hole symmetry of \( G^c(\tau) \rightarrow G^h(\beta - \tau) \) where \( G^c(\tau) \) is the electron Green’s function and \( G^h(\tau) \) is the hole Green’s function.
In this case, if a regular mesh is used then more than 50% of all data-points are zero, due to \( G(\tau) \approx 0 \) between \( 5 < \tau < 15 \). However, if the adaptive mesh is used less than 10% of the points fall in this interval are zero. Indeed, depending on the inverse temperature \( \beta \), the optimal choice of \( \delta \) should ensure that the features of \( G(\tau) \) are retained away from where it plateaus. In practice, the procedure to obtain \( G(\sigma(\tau, \delta)) \) is as follows:

1. Spline the equidistant \( G(\tau_1, \ldots, \tau_N) \) to get its continuous representation.
2. Define the new adaptive grid \( \sigma(\tau, \delta) \) from \([0, \beta] \) on a continuous grid.
3. Evaluate \( G(\sigma(\tau, \delta)) \) on the continuous grid.
4. Sample \( G(\sigma(\tau, \delta)) \) \( N \) times to obtain \( G(\tau_1^{ad}, \ldots, \tau_N^{ad}) \), where \( \tau_i^{ad} \) indicates the \( i \)th \( \tau \) point on the adaptive mesh.

We note that adaptive meshes are not required if the Green’s function is expressed in a polynomial basis. There is still, however, the issue of obtaining spurious Legendre coefficients if a cutoff of \( n_l^{max} \) is not imposed. We illustrate this problem in Fig. 15 for a range of temperatures using 100 Legendre coefficients for each one. For \( n_l > 10^2 \) it is clear that including more coefficients in the expression for \( G(\tau) \) starts to exhibit classical overfitting behaviour. Therefore, when using the Legendre basis a judicious choice of \( n_l^{max} \) must be made, otherwise the model will be trained on data that lacks relevant physical information. As depicted in Fig. 15 when \( \beta \) increases so do they number of Legendre coefficients needed for an accurate expression of the Green’s function. Therefore, depending on the temperature chosen, the size of the feature space changes for the Legendre representation. Whereas in the case of the imaginary-time basis, as \( \beta \) increases the necessity to use an adaptive mesh becomes higher.

Finally, we now turn to the final set of data transformations that must be performed. Scaling the input and output variables so that they are normalised is a standard technique when preparing data for training algorithms such as neural networks. The reason for this is to protect the weights that are learned in the model from becoming too large or biased towards large input values. Specifically, this is essential for when input variables are the Legendre coefficients, as the Legendre basis has no inherent scale for the coefficients. On the other hand, while an inherent scale exists for \( G(\tau) \), i.e. \(-1 \leq G(\tau) \leq 0\) when \( \tau > 0 \), it is also possible to create a family of scaling transformations and test their efficacy throughout the training process. The following scaling transformations work regardless whether the aforementioned symmetrisation or augmentation procedures have been followed.

We first discuss the transformations \( T \) for \( G(\tau) \) as shown in Fig. 16. \( T_0 \) is the unscaled Green’s function and each transformation is a function of \( T_0 \). For \( G(\tau) \) the situation is quite simple, there are only a few transformations that can be done to normalise in between the range \([0, 1] \) or \([-1, 1] \). We note that if \( G^{AS}(\tau) \) is used, i.e. the anti-symmetric part of the Green’s function, it is important to ensure that the scaling operations do not shift the data out of the scaling range, and so an extra constant shift should be applied in these cases to counteract this behaviour. The Legendre basis and its scaling transformations are illustrated in Fig. 16. In this case it is clear that the unscaled data is not normalised. Fortunately, by applying a \( \tanh \) function this can readily be achieved. In the example shown, we see the first anti-symmetric component of the Green’s function, \( G_1 \), is scaled to be much closer to \( G_0 \) and \( G_2 \), the either-side symmetric components. As stated above for the \( \tau \) basis, the dependence of training the model is also assessed as a function of these transformations. Moreover, we emphasise that to recover the physical Green’s function it is necessary to apply the
relevant inverse transformation $T^{-1}$, which are tabulated in Fig. 17.

**Appendix B: error scaling**

As discussed in this work, the error shows a dependence on the chosen parameters $\{U, \varepsilon\}$. Here we investigate the overall features of this dependence by examining it over the entire phase space, restricting the $U$-values of up to 8eV that represent physical realisable scenarios. To this end, we explore the $U - \varepsilon$ phase-space of the error, chosen to be the RMSD, between the approximate solvers and exact one. We select $\varepsilon$ as this combination characterises the physics of the impurity, which is what is being perturbed for the Hubbard-I, IPT and NCA solvers, and hence is where the error should manifest itself. In Fig. 18, we explore this space for the IPT solver for $\beta = 1\, eV^{-1}$ and $\beta = 20\, eV^{-1}$. For $\beta = 1\, eV^{-1}$, illustrated in Fig. 18a-c, we see a clear structure of the error, which is $< 10^{-5}$ for $U < 4\, eV$, and also at near half-filling, clearly highlighted for the 2d-cut in panel b. For large values of $U$, away from half-filling, the solver performance starts to deteriorate. Indeed, this is the expected behaviour of the IPT solver\cite{note}, for these parameter regimes, and we see that there is a high density of points falling near zero in panel c in its error.

On the other hand, for $\beta = 20\, eV^{-1}$ (Fig. 18d-e) a lot of the structure present at $\beta = 1\, eV^{-1}$ disappears. In particular, at low $U$ the error extends to larger RMSD values and the structure in $\varepsilon$ is practically lost. Interestingly, the RMSD error in panel f splits into two regions, one where the error persists to be low while the other is high, and highlights that lowering temperature amplifies the error up to an order of magnitude or greater. While there is still a persistent structure in the error as a function of $U$ at $\beta = 20\, eV^{-1}$, our results suggest that predicting it reliably will be more difficult.

We repeat the same method of analysis for the NCA solver shown in Fig. 19. Again, for $\beta = 1\, eV^{-1}$ there are systematic features to the error where it is lower for larger values of $U$, as shown in Fig. 19a-b. Indeed, this is as expected when using the NCA solver as it is a strong-coupling expansion\cite{note}. Notably, even though the trend favours higher $U$ values, their overall magnitude is larger than for the IPT solver by an order of magnitude, even though the trend is not present there. We also notice that the distribution of the error (panel c) is markedly different, taking a Gaussian-like shape centred around the RMSD value of 0.03. This Gaussian-like behaviour is, however, absent in the $\beta = 20\, eV^{-1}$ case (panel f) where the distributed error reverts back to decaying away from zero. Moreover, in a similar fashion to the IPT case, for $\beta = 20\, eV^{-1}$ a significant portion of structure disappears in the $U - \varepsilon$ dependence of the error, which suggests...
that correcting for it may become increasingly difficult for lower temperatures.

We repeat the same method of analysis for the Hubbard-I solver shown in Fig. 22. Similarly for the IPT and NCA solvers there is a noticeable structure present at $\beta = 1$ eV$^{-1}$, as shown in Figures (20a-c). Interestingly, we see that the error is minimal for $U \approx 5$eV, which is opposite to what was observed for the IPT and NCA solvers. Moreover, at $\beta = 1$ eV$^{-1}$ the distribution of the RMSD error is heavy-tailed and shows a sensitivity away from half filling, similar to the IPT result, for example. Nevertheless, its magnitude is roughly about $10^{-2}$, more in line with the NCA prediction. Therefore, the Hubbard-I solution bears the hallmarks of capturing qualitative features of the exact solution at the intermediate interaction strengths, and places itself as a complementary approximate solution to the IPT and NCA solvers. Finally, we see a striking resemblance to the NCA solver, whereby the error landscape, as seen in Figure (20d-f). Here, we see a striking resemblance to the NCA solver, whereby there is a noticeable structure in the error for large and small values of the Hubbard $U$ parameter.

We also note that the behaviour in the Legendre basis is strikingly similar, although the $U - \varepsilon$ is far more discrete in regions where the solvers perform well. Nevertheless, it is conclusive that a basis-switch is not sufficient to recover the discernible features of its structure at lower temperatures. Moreover, we clarify that the $G(\tau)$ and $G_l$ data analysed is the raw data, without any processing, except for the truncation of the Legendre basis. We use Fig. 21 to determine the $\beta$ dependence of the this cut-off, where the input AIM is a sample from the generated database. In order to maintain an error $< 10^{-5}$ as function of increasing $\beta$ we choose $l_{\text{max}} = 8$ for $\beta = 1$ and follow its diagonal along the matrix in Fig. 21. The over-fitting of $G(\tau)$ with $G_l$ is also evident here, where for example if $\beta = 10$ as $l_{\text{max}}$ is increased there is an eventual increase in the error for $l_{\text{max}} = 100$.

Appendix C: Quantum database samples

In this section we provide representative test examples that are discussed in this manuscript for illustration of the methodology. We first turn our attention to the representative inputs $x_i$ and outputs $y_i$ of the training model (as per Fig. 2) we show the approximate solutions IPT, NCA and Hubbard-I with the corresponding exact solutions in both imaginary-time and Legendre bases for different inverse temperatures $\beta$ (as per Fig. 22). The specific AIM configurations were chosen at random for the database of 10,000 solutions.

For $\beta = 1$ eV$^{-1}$ and the parameter set $\{U = 3.48eV, W = 5.85eV, \varepsilon = -0.61\}$ we see that the IPT approximation is highly accurate in both bases (Fig. 22a-b), even away from half-filling. Moreover, whilst both Hubbard-I and NCA solvers are quantitatively in disagreement with the exact solution, the NCA solver recovers some critical features of the exact solution such as the $G(\tau \rightarrow 0)$ value as well as the overall shape. Evidently, the Legendre basis is significantly more compact than the imaginary-time basis, however the error does not manifest itself so obviously in it, where small changes in the coefficient can give large changes in the overall physical properties.

For $\beta = 20$ eV$^{-1}$ the parameter set $\{U = 6.55eV, W = 6.1eV, \varepsilon = 0.31\}$ is randomly chosen instead, and illustrated in Fig. 22d. The IPT approximation in this case retains a qualitative agreement with the exact answer, however there are obvious deviations that emerge. Furthermore, the NCA and Hubbard-I solvers are in quantitative and qualitative disagreement, and this is illustrated in the second and third Legendre coefficients. For example, the Hubbard-I solver indicates an insulating solution when in reality the system is metallic.

For $\beta = 50$ eV$^{-1}$ the parameter $\{U = 6.2eV, W = 2.69eV, \varepsilon = 0.83\}$ again randomly chosen, and illustrated in Fig. 22f. Here, in contrast with the higher temperature results, both Hubbard-I and NCA solvers are quantitatively and qualitatively in better agreement with the exact solution than the IPT result. Moreover, the IPT result qualitatively estimates an incorrect value for the onsite electron occupation, as estimated by $G(\tau \rightarrow 0)$.

Appendix D: Neural Network loss functions for additional models

Figure 23 illustrates the validation loss for all other models studied in the main text. The overwhelming conclusion that can be made from these results is that the combination of quality approximate solutions results in smaller values of the validation loss when using the imaginary-time basis. As mentioned in the main text, using the IPT approximate solver is a critical component to minimising the overall loss of the neural network.

Appendix E: Neural Network prediction for representative samples

Having optimised the various aspects of the network, and analysed its loss functions, we discuss here how the model is able to make predictions for unseen parameter choices (see Fig. 24). For $\beta = 1$ eV$^{-1}$ and the parameter set $\{U = 1.99eV, W = 3.35eV, \varepsilon = 0.66\}$ we see that the neural network captures the essence of the error in both the imaginary-time and Legendre bases (Fig. 24a-b). This is perhaps most clearly identified at $G(\tau = 0.6)$ in the imaginary-time basis or at second Legendre coefficient in the polynomial basis. For the Legendre basis, we present the data on a log-scale to highlight how the prediction is essentially error-free until the coefficients become $< 10^{-3}$ in log-space, which is practically negligible for the reconstruction in imaginary-time. On the
other hand, for $\beta = 20 \text{ eV}^{-1}$ and the parameter set 
\{\(U = 1.88\text{eV}, W = 4.4\text{eV}, \varepsilon = 0.189\)\} we see that (Fig. 24c-d), while the agreement between the neural network solver is not as refined compared to the $\beta = 1 \text{ eV}^{-1}$ network, it is much better than what the approximate solvers predict. Interestingly, it is only through the odd-components of $G_l$ that the network actually struggles with predicting (up to a critical value of $l$), which suggests that separating these contributions (i.e. the half-filling and away from half-filling) and learning them separately could improve the network. By extending into the lower temperature regime, i.e. $\beta = 50 \text{ eV}^{-1}$, we see that similar features persist in the even/odd decomposition of the coefficients, illustrated for the the parameter set \{\(U = 1.88\text{eV}, W = 4.4\text{eV}, \varepsilon = 0.189\)\} in (Fig. 24e-f). Again, in contrast to the $\beta = 1 \text{ eV}^{-1}$ results, we see that there are slight discrepancies throughout the entire range of Legendre coefficients, rather than just the higher-order ones. Nevertheless, the agreement in the imaginary-time basis indicates that many of the main physically relevant features are captured by the neural network up to $\beta = 50\text{eV}^{-1}$, such as the occupation and metallicity of the solution.
Figure 18: The $U - \varepsilon$ error of the impurity Green’s function $G(\tau)$ calculated with the Iterated Perturbation Theory solver relative to the Exact Diagonalisation solution. For $\beta = 1$ eV$^{-1}$ a 3d-cut of the error b 2d-cut of the error c histogram of the error. For $\beta = 20$ eV$^{-1}$ for d 3d-cut of the error e 2d-cut of the error f histogram of the error.
Figure 19: The $U - \varepsilon$ error of the impurity Green’s function $G(\tau)$ calculated with the Non Crossing Approximation solver relative to the Exact Diagonalisation solution. For $\beta = 1 \text{ eV}^{-1}$ a 3d-cut of the error b 2d-cut of the error c histogram of the error. For $\beta = 20 \text{ eV}^{-1}$ for d 3d-cut of the error e 2d-cut of the error f histogram of the error.
Figure 20: The $U - \varepsilon$ error of the impurity Green's function $G(\tau)$ calculated with the Hubbard-I solver relative to the Exact Diagonalisation solution. For $\beta = 1 \text{ eV}^{-1}$ a 3d-cut of the error b 2d-cut of the error c histogram of the error. For $\beta = 20 \text{ eV}^{-1}$ for d 3d-cut of the error e 2d-cut of the error f histogram of the error.
Figure 21: Reconstructed $G(\tau)$ from $l_{\text{max}}$ Legendre polynomials for the AIM heatmap as a function of inverse temperature $\beta$ and $l_{\text{max}}$. $l > 60$ ensures for $\beta < 100$ a completely accurate representation of $G(\tau)$. 
Figure 22: Representative randomly sampled configurations (from the database of Table I) of the Anderson Impurity Model with a collection of approximate exact solutions for the $\beta = 1 \text{ eV}^{-1}$ a imaginary-time and b Legendre bases. Similarly for $\beta = 20 \text{ eV}^{-1}$ in the c imaginary-time and d Legendre bases. Similarly for $\beta = 50 \text{ eV}^{-1}$ in the e imaginary-time and f Legendre bases.
Figure 23: Cost function for the adaptive $\tau$ mesh for different inverse temperatures $\beta$ using as input to the neural network the solvers a [IPT, NCA, ED-1, ED-2] b [ED-1, ED-2, ED-3, Hubbard-I], c [IPT, ED-1, ED-2], and d [NCA, ED-1, ED-2]. IPT is the Iterated Perturbation Theory Solver, NCA is the Non Crossing Approximation solver and ED-[1,2,3] are the truncated ED solvers.
Figure 24: Representative randomly sampled configurations of the Anderson Impurity Model neural network solution and exact solution for the $\beta = 1 \text{ eV}^{-1}$ trained on the $\mathcal{M}_1$ ensemble of approximate solutions in (a) imaginary-time and (b) Legendre bases. Similarly for $\beta = 20 \text{ eV}^{-1}$ in the (c) imaginary-time and (d) Legendre bases. Finally, for $\beta = 50 \text{ eV}^{-1}$ in the (e) imaginary-time and (f) Legendre bases.