Pure density functional for strong correlations and the thermodynamic limit from machine learning

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We use density-matrix renormalization group, applied to a one-dimensional model of continuum Hamiltonians, to accurately solve chains of hydrogen atoms of various separations and numbers of atoms. We train and test a machine-learned approximation to $F[n]$, the universal part of the electronic density functional, to within quantum chemical accuracy. Our calculation (a) bypasses the standard Kohn-Sham approach, avoiding the need to find orbitals, (b) includes the strong correlation of highly-stretched bonds without any specific difficulty (unlike all standard DFT approximations) and (c) is so accurate that it can be used to find the energy in the thermodynamic limit to quantum chemical accuracy.

I. INTRODUCTION

Although widely used in solid-state physics, chemistry, and materials science,\textsuperscript{1} Kohn-Sham density functional theory (KS-DFT) with standard approximations fails for strong correlation\textsuperscript{2,3}. The prototype is the H$_2$ molecule. When stretched, the electrons localize on each site while remaining in a singlet state, but this is not captured by such calculations.\textsuperscript{4} These kinds of difficulties have led to the popularity of many beyond-DFT schemes, ranging from the simple addition\textsuperscript{5} of a Hubbard $U$ to the use of dynamical mean field theory\textsuperscript{6} as well as many others.

But even KS-DFT is too slow for many large calculations, such as those using classical MD or continuum mechanics.\textsuperscript{7} The original DFT, first suggested in the Thomas-Fermi approximation\textsuperscript{8,9} and later justified by the Hohenberg-Kohn theorem\textsuperscript{10}, uses only pure functionals of the total density, $n(r)$. This ‘orbital-free’ version has the potential to be much faster than even the most efficient KS implementations, because the KS equations need not be solved\textsuperscript{11}. Several recent attempts have constructed machine learning (ML) kinetic energy functionals specifically to bypass this step\textsuperscript{12,15}. These are designed to be used in conjunction with standard KS approximations to speed up such calculations, but not to improve their accuracy.

Meanwhile, beyond the world of DFT, density matrix renormalization group (DMRG) has become a standard tool for finding extremely accurate solutions to strongly correlated lattice problems.\textsuperscript{16,19} In recent years, a one-dimensional analog of ab-initio Hamiltonians has been developed, using typically about 20 grid points per atom and interactions involving many grid points, with the express purpose of rapidly exploring both conceptual and practical issues in DFT.\textsuperscript{20,21,22} A particular advantage is that, since 2000 grid points is routinely accessible, this includes up to 100 atoms, and extrapolations to the thermodynamic limit are much easier than in three dimensions. Applications include a demonstration of the behavior of the KS gap in a Mott-Hubbard insulator\textsuperscript{20} and a proof of convergence of the KS equations with the exact functional, regardless of the starting point or strength of correlation\textsuperscript{21}.

![FIG. 1. (Color online) Electronic energy of infinite chain from model learned from extrapolated chain densities and energies. The accurate value was calculated with infinite DMRG (see text).
In the present work, we combine all these methodologies to demonstrate several important features. We perform DMRG calculations on a variety of one-dimensional hydrogen atom chains, with from 2 to 20 atoms, and whose interatomic spacing $R$ varies from 1 to 10 Bohr radii, and use these to train a ML model of $F[n]$, the ‘universal’ part of the density functional identified by Hohenberg-Kohn. This simultaneously includes both the non-interacting kinetic energy sought in orbital-free DFT and the exchange-correlation energy that is approximated in KS calculations. We demonstrate that, with reasonable amounts of training, we can self-consistently calculate densities and energies for these chains at new values of $R$, outside the training set, with quantum chemical accuracy. This includes highly stretched systems which are strongly correlated, and where all popular DFT approximations fail. We furthermore extrapolate the DMRG densities from the center of finite chains to the infinite chain limit, i.e., a 1d solid. We train a new ML model and find we can solve self-consistently the solid

\begin{figure}[h]
\centering
\includegraphics[width=0.5\textwidth]{figure1.png}
\caption{(Color online) Electronic energy of infinite chain from model learned from extrapolated chain densities and energies. The accurate value was calculated with infinite DMRG (see text).}
\end{figure}
problem at the same level of accuracy. Fig. 1 shows the convergence of our ML method for a typical separation of the infinite chain with respect to the number of training points. The horizontal lines show two independent DMRG estimates of the energy.

II. BACKGROUND

A. DFT

The Hohenberg-Kohn theorem \cite{Hohenberg1964} establishes that the ground-state energy and density of a many-body problem may be found by minimizing a density functional:

$$E = \min_n \left\{ F[n] + \int d^3r \ n(r) v(r) \right\},$$

where \(n(r)\) is the single-particle density, normalized to \(N\) particles, and \(v(r)\) is the one-body potential. The functional \(F\) can be defined via a constrained search as \cite{Perdew1986}

$$F[n] = \min_{\Psi \rightarrow n} \langle \Psi | \hat{T} + \hat{V}_{ee} | \Psi \rangle$$

where \(\hat{T}\) is the kinetic energy operator and \(\hat{V}_{ee}\) is the electron-electron repulsion operator, while \(\Psi\) is a normalized antisymmetric (for fermions) wavefunction. While this showed that the old Thomas-Fermi theory \cite{Thomas1927, Fermi1927, Becke1988} was an approximation to an exact formulation, few modern approximations exist. An analogous failure occurs for semilocal approximations to \(T_s[n]\) when bonds are stretched in orbital-free DFT. Fig. 2 illustrates the failure of semilocal XC, by comparing the blue restricted LDA curve with the black DMRG curve. There is a huge error in the stretched limit.

B. DMRG benchmark data

It is difficult to overemphasize the utility of benchmark quantum chemical calculations for the development of DFT. The DFT revolution in quantum chemistry was made possible by the existence of the well-tested G2 data set for small molecules, and databases in quantum chemistry have proliferated ever since. On the other hand, calculations of ‘quantum chemical’ accuracy, i.e., errors below 1 kcal/mol, are much more difficult and rarer for solids. A recent heroic effort \cite{&Coulomb1997} was made for benzene, a molecular crystal.

For the present study, we need to consider chains of up to 20 H atoms, with many different values of the interatomic spacing ranging from about 1 to 10 Bohr. Extracting this large amount of data at the required level of accuracy from a quantum chemical code would be extremely demanding, if not impossible, given the strong correlation effects when the bonds are stretched.

Recently, DMRG has been applied to a one-dimensional analog of real-space Coulomb-interacting Hamiltonians, for precisely the purpose of performing demanding, highly accurate benchmark calculations of strongly correlated systems. In particular, the interac-
The density functional is represented as a nonlinear regression method with an L2 regularization [34]. The density functional ridge regression (KRR). It is a nonlinear regression technique constructed by interpolation from accurate examples. To avoid searching in directions for which there is little information, the focus has been on the KS kinetic energy, \( T \), which has previously been shown to be sufficient to converge rapidly with these parameters, and illustrates that the corresponding functional derivative is highly inaccurate. This is unfortunate, as the practical usefulness of an accurate model for \( T \) is in finding the density via solution of the Euler equation (for the KS system):

\[
\frac{\delta T}{\delta n(x)} = -v_t(x),
\]

where \( v_t(x) \) is the KS potential. This difficulty has been surmounted in a sequence of increasingly sophisticated methods [13]–[15], each of which constrains the density search to only the manifold of densities spanned by the data, avoiding searching in directions for which there is insufficient data to evaluate \( T \) accurately. With such techniques, it has been possible to demonstrate an ML \( T \) functional that correctly mimics the KS solution even as a bond stretches [13], something impossible for any local or semilocal approximation to \( T \). The value of this is to cut down the computational cost of large, repetitive KS calculations, but one still uses some standard XC approximation. Thus a machine-learned functional for \( T \) can, at best, reproduce the incorrect LDA curve of Fig. 2.

### C. Machine learning of the KS kinetic energy functional

ML is a set of algorithms developed to find hidden insights in data. It is widely used especially when the pattern behind complicated data is difficult to deduce explicitly. Successful applications include computer vision [29], cybersecurity [30], ancient abstract strategy games [31], etc.

Recently, in chemistry and materials science, machine-learning has become a popular tool for analyzing properties of molecules and materials, and finding specific functions from large data sets [32, 33]. But it has also been applied to the problem of finding density functionals, constructed from interpolation from accurate examples. To date, the focus has been on the KS kinetic energy, \( T \), partially because of the ready availability of data (every cycle of every one of the 30,000 KS-DFT calculations each year [14] produces an accurate example of \( T \)) and because of the enormous potential for speeding up routine DFT calculations.

The ML algorithm we used for modeling \( T \) is kernel ridge regression (KRR). It is a nonlinear regression method with an L2 regularization [23]. The density functional is represented as

\[
T_{\text{ML}}^{\alpha}[n] = \sum_{i=1}^{N_T} \alpha_i k[n, n_i],
\]

where \( N_T \) is the number of training data, \( n_i(x) \) are the training data and \( k[n, n_i] \) is a kernel, some measure of the “similarity” between densities. Throughout this work, we use a Gaussian kernel,

\[
k[n, n'] = \exp(-\|n - n'\|^2/2\sigma^2),
\]

where

\[
\|n - n'\| = \int dx (n(x) - n'(x))^2.
\]

Such a kernel is standard in KRR, and has yielded excellent results in previous studies of \( T \) [14]. The weights \( \alpha = (\alpha_1, \ldots, \alpha_{N_T}) \) are found by optimizing the cost function

\[
C(\alpha) = \sum_{i=1}^{N_T} (T_{\text{ML}}^{\alpha}[n_i] - T[\delta_t])^2 + \lambda \alpha^T K\alpha
\]

where \( K \) is the kernel matrix, \( K_{ij} = k[n_i, n_j] \). The regularization strength \( \lambda \) and length scale \( \sigma \) are hyperparameters which are found via cross validation. A crucial principle in kernel ridge regression is the separation of the training data from the test data. A test set is constructed entirely independently from the training set. The cross-validation to find the hyperparameters occurs using only training data. The resulting approximate functional is tested only on the test data.

While highly accurate results for \( T \) can be found with relatively little data [12], it was immediately realized that the corresponding functional derivative is highly inaccurate. This is unfortunate, as the practical usefulness of an accurate model for \( T \) is in finding the density via solution of the Euler equation (for the KS system):

\[
\frac{\delta T}{\delta n(x)} = -v_t(x),
\]

where \( v_t(x) \) is the KS potential. This difficulty has been surmounted in a sequence of increasingly sophisticated methods [13]–[15], each of which constrains the density search to only the manifold of densities spanned by the data, avoiding searching in directions for which there is insufficient data to evaluate \( T \) accurately. With such techniques, it has been possible to demonstrate an ML \( T \) functional that correctly mimics the KS solution even as a bond stretches [13], something impossible for any local or semilocal approximation to \( T \). The value of this is to cut down the computational cost of large, repetitive KS calculations, but one still uses some standard XC approximation. Thus a machine-learned functional for \( T \) can, at best, reproduce the incorrect LDA curve of Fig. 2.

### III. METHOD

In all applications in this work, we generate a large data set of highly accurate results generated using DMRG. We use a real-space grid with spacing 0.04, which has previously been shown to be sufficient to converge the results [23]. We calculate the energies and densities of chains of even numbers of atoms, from 2 to 20, with interatomic separations between 1 and 10 Bohr. Higher accuracy is achieved when every atom is centered on a grid point, discretizing the set of allowed separations. The specific separations used are listed in the supplemental information.
Then a subset of these data are left out as test set. The training set, with \( N_T \) values of \( R \), are collected from the remaining data. These are chosen to be as close to equally spaced as practical. The test set is excluded from the data where the training set is sampled from, to avoid contamination via the cross-validation process.

### A. Machine-learned functional for a given molecule

![Fig. 3](image-url) The green curves are ML with \( N_T = 5 \) on both the exact (dashed) and ML-optimized (solid) densities. The red solid curve is the ML with \( N_T=20 \) on ML-optimized (solid) densities. Black dashed curve is the exact DMRG curve, matching nearly exactly the \( N_T=20 \) on ML line.

We continue to use the \( H_2 \) molecule to illustrate our method. Contrary to previous work, we apply KRR algorithms to ML the interacting functional \( F[n] \) itself, by training on highly accurate DMRG energies and densities at various values of \( R \). In Table I we list the errors for \( H_2 \) as a function of \( N_T \), both on the exact density and on an optimally constrained density found by the methods of Ref. [35].

To illustrate the procedure, in Fig. 3 we show the energies with only 5 training points, \( R = 1.00, 3.20, 5.48, 7.76, 10.00 \), yielding the smooth, green dashed curve, when evaluated on the exact densities. The curve (almost) exactly matches at the training points, but is noticeably inaccurate inbetween. But note that, in contrast to all previous studies, we are fitting the full \( F[n] \), not just \( T_0[n] \), so that, e.g., our inaccurate curve dissociates \( H_2 \) correctly, while no standard DFT calculation, with a standard XC approximation, can.

The problem is actually much greater than even the smooth dashed green curve would suggest. In practice, we not only need the energy functional, but also its derivative, at least in the vicinity of a solution density. This is because we use the functional to find the density itself, via the Euler equation

\[
\frac{\delta F}{\delta n(x)} = -n(x). \tag{10}
\]

In fact, the derivatives of ML functionals such as that of Eq (6) are highly inaccurate and cannot be used to find the minimizing density. Methods have been developed to constrain the search to the manifold of training data via non-linear gradient denoising (NLGD) [37]. For our \( H_2 \) with \( N_T = 5 \), these lead to the (even worse) solid green curve of Fig. 3. The optimal density is shown in Fig. 4. We clearly see that (a) the accuracy is not high enough and (b) the error is dominated by the error in the densities. (This is called a density-driven error [38] in a DFT calculation.)

However, when we increase to 20 data points, the ML curve (red solid) is indistinguishable from the exact one, and the error at equilibrium is only 0.007 kcal/mol, and shrinks with increasing \( R \). This calculation applies all the principles discussed in Ref. [13] but is now applying them to the many-body problem, not just the KS problem.

Even in the stretched limit, where the system is strongly correlated, there is no loss of accuracy. Note that we are not just fitting the binding curve, as we are reproducing the many-body density at every value of \( R \), starting from data at a limited number of values. In Fig. 4 we plot the optimally-constrained densities at \( R = 4.0 \) (outside all training sets) for \( N_T = 5 \) and \( N_T = 20 \), compared with the exact density.

Here, ML has entirely bypassed the difficulty of solving the many-fermion problem. The machine learns the characteristics of the solution without ever solving the differential equation. Moreover, the HK theorem is a statement of the minimal information needed to characterize the ground-state of the system. In some ways, this ML approach is the purest embodiment of the HK theorem.

### B. Finding a data-driven optimal basis for longer chains

The cost of optimal gradient descent methods, evaluated on a spatial grid, grows very rapidly with the num-
number of grid points, and rapidly becomes unfeasible as the number of H atoms grows. Thus a simpler representation of the density is required. To overcome those difficulties, we introduce a basis set. Inspired by the localized atomic bases used in most quantum chemical codes, we developed a data-driven basis set using Hirshfeld partitioning \cite{37} and principal component analysis (PCA).

To partition a molecular density via the Hirshfeld scheme, begin with the protomolecule of overlapped atomic densities at the nuclear positions of the real molecule. If \( n_i^0(x) = n_i^0(x-(i-1)R) \) is an isolated atomic density at the \( i \)-th nuclear center, spaced \( R \) apart, then

\[
n^0(x) = \sum_{i=1}^{N} n_i^0(x) \tag{11}
\]

is the density of the protomolecule, where \( R \) is the interatomic spacing. We define a weight

\[
w_i(x) = n_i^0(x)/n^0(x), \tag{12}
\]

associated with each atom, and then define the density of each Hirshfeld atom within the real molecule as

\[
n_i(x) = w_i(x)n(x), \tag{13}
\]

where \( n(x) \) is the exact molecular density. The ground state density of a single hydrogen atom \( n_0^0(x) \) is reported in Ref. \cite{23} Fig. 5 shows partition densities \( n_i(x) \) of atoms in one \( \text{H}_8 \).

Next, for a specific chain length \( N \), we consider a range of interatomic separations \( R \), and consider the collection of every atomic density within the chain for every value of \( R \) in a training set, each centered on the origin, as shown in Fig. 6. These individual atomic partition densities reflect the diverse behaviors caused by the interaction between the hydrogen atoms inside the chains. A principal component analysis is applied to these densities, and the eigenvalues are ordered in decreasing magnitude to find a subspace with the maximum variance. Each atomic density can be accurately represented by the base density \( f_0(x) \) (red in Fig. 6) and 7 principal components (Fig. 7),

\[
n_i(R,x) = f_0(x) + \sum_{p=1}^{7} c_{i,p}(R)f_p(x). \tag{14}
\]

Thus the total density of \( \text{H}_N \) with separations \( R \) is \( \sum_i n_i(R,x) \), and is described by just \( 7N \) coefficients. Note that \( f_0(x) \) is very close to an isolated atom density, but we use the average to center our data for the PCA analysis. Our representation greatly reduces the number of variables in the density representation for a given chain length, and saves a significant amount of computational cost when solving for the ground state density of the system. This new basis set is completely data-driven and physically meaningful.

We next repeated these calculations for a sequence of chains of increasing length. In each case, we train \( F_{\text{ML}}[n] \) on a limited training set, and then compare on a test set.
TABLE I. ML performance on different chains $H_N$. $N_T$ is the size of training set. Regularization strength $\lambda$ and kernel length scale $\sigma$ are the model hyperparameters selected by cross validation [14]. The functional-driven error $\Delta E_F/N$ [36] is tested on the entire test set to show the overall accuracy. The total error $\Delta E/N$ is tested on the equilibrium test set to emphasize accuracy around equilibrium position. $E_{R=9.8/N}$ shows that ML can get very accurate dissociation limit. All errors are given in kcal/mol.

| $N$ | $N_T$ | $\lambda$ | $\sigma$ | $|\Delta E_F|/N$ | max $|\Delta E_F|/N$ | $\Delta E/|N$ | max $|\Delta E|/N$ | $E_{ML}^{R=9.8/N}$ | $E_{DMRG}^{R=9.8/N}$ |
|-----|-------|-----------|----------|--------------|-----------------|-------------|-----------------|-----------------|-----------------|
| 2   | 5     | $1.0 \times 10^{-8}$ | 1000     | 2.54         | 7.02            | 9.74        | 20.3            | -421.291        | -425.797        |
| 2   | 20    | $4.6 \times 10^{-10}$ | 2.15     | 0.00121      | 0.00802         | 0.005       | 0.013           | -425.785        | -425.797        |
| 2   | 50    | $1.0 \times 10^{-12}$ | 0.70     | 0.00003      | 0.00034         | 0.050       | 0.304           | -425.798        | -425.797        |
| 4   | 50    | $2.2 \times 10^{-11}$ | 46.4     | 0.0021       | 0.016           | 0.005       | 0.017           | -428.617        | -428.620        |
| 8   | 50    | $1.0 \times 10^{-4}$  | 2.15     | 0.011        | 0.31            | 0.28        | 1.68            | -430.011        | -430.032        |
| 12  | 50    | $1.0 \times 10^{-12}$ | 0.46     | 0.0031       | 0.010           | 0.24        | 0.88            | -430.502        | -430.503        |
| 16  | 50    | $2.2 \times 10^{-11}$ | 0.46     | 0.0042       | 0.012           | 0.08        | 0.41            | -430.738        | -430.738        |
| 20  | 50    | $2.2 \times 10^{-11}$ | 0.46     | 0.0042       | 0.014           | 0.26        | 0.88            | -430.880        | -430.880        |
| $\infty$ | 50 | $1.0 \times 10^{-8}$  | 0.46     | 0.012        | 0.050           | 0.073       | 0.27            | -431.447        | -431.444        |

C. Extrapolation to the thermodynamic limit

Our ultimate goal is to use ML to find the energy of the infinite chain to within chemical accuracy, for all interatomic separations. To do this, we first build a set of infinite chain energies and densities. For each value of $R$, we extrapolate both the density and energy of our finite chains as a function of $N$. This then gives us a set of data for the infinite chain that we can both train and test on and gave rise to Fig. 1.

In an entirely separate calculation, we also performed DMRG directly for the infinite chain, using the method of McCulloch [35] for a four atom unit cell [39]. The system is initialized by solving the equivalent finite size system with box edges at $R/2$. As a part of the iDMRG algorithm [35], a single unit cell is then inserted into the center of the finite system and 15 sweeps are performed over the inserted unit cell. The sequence is repeated—after adding another unit cell—until convergence. We compare these energies with the extrapolated values, finding agreement to within 1 kcal/mol for all values of $R$. This agreement validates our extrapolation procedure. We find that, with 50 data points, the ML result, on the optimized density, also agrees to within 1 kcal/mol. Thus, armed with the 50-data-point machine learned functional, one can self-consistently find the density and energy of the infinite chain to quantum chemical accuracy.

Our final figure simply demonstrates that the error for the infinite chain (and for all the ML calculations) is almost entirely due to the error in the optimized density. The functional-driven error [30] is the energy error made on the exact density:

$$\Delta E_F = E_{ML}[n] - E[n] = F_{ML}[n] - F[n].$$

We see that, at any level of training, $\Delta E_F$ is an order of magnitude smaller than the final energy error on the optimized density. Thus the error is density-driven but, nonetheless, can be forced down to quantum chemical limits with enough data.

![Fig. 8](image-url) (Color online) Learning curves for several 1d H chains. (a) ML using the total density. (b) ML using the bulk partition densities (see text).
FIG. 9. (Color online) Electronic energy per atom in the thermodynamic limit, both via DMRG chains (extrapolated to infinity) and using machine learning with 50 data points per chain.

FIG. 10. (Color online) For a given training set with \( N_T \) training points, the functional driven error, \( \Delta F_F \) per atom is shown in red (lower curve). The upper curve is the total energy error per atom evaluated self-consistently.

IV. DISCUSSION

We have shown that it is in principle possible to construct, via machine learning, the entire interacting functional of Hohenberg and Kohn, \( F[n] \), so accurately that optimized densities and energies evaluated on them are within quantum chemical accuracy. We have done this using the 1d simulation of continuum Hamiltonians established over the last several years, and using DMRG as an efficient solver. We apply the ML methods previously developed for approximating the non-interacting kinetic energy. Here, because we have precise energies for the interacting system to train on, we are able to construct the interacting functional, including all exchange and correlation effects. Our ML functional has no difficulties when bonds are stretched so that correlations become strong. We have even managed to apply this methodology to chains extrapolated to the thermodynamic limit, producing chemically accurate results for solids. This level of accuracy is far beyond that of any existing DFT calculation of a solid.

We conclude with a long list of the many things we have not done. Most importantly, all calculations have been in one dimension, because of the relative ease of setting them up, the efficiency of DMRG, and the rapidity of approach to the bulk limit. We do not know how much additional cost is involved in three dimensional calculations. In fact, other recent work [40] shows that 3d molecular calculations do work for the orbital-free problem, but that both a good choice of basis set is needed, as well as building a density-potential map directly, avoiding the need to minimize the ML density functional. Thus, the extension to 3d appears entirely practical.

A second limitation is the rather large amount of data employed. When we need an accurate calculation at every 0.2 atomic units, how much have we achieved when we interpolate between points? But this extreme amount is needed only because we must produce a self-consistent density from the ML functional, and one that is so accurate as to introduce only a 1 kcal/mol density-driven error. This is a level of accuracy that could only be dreamed of with human orbital-free approximations.

There are two ways in which this requirement might be very dramatically reduced: (a) machine-learning the HK map \( n[v] \) directly, as mentioned above (but here for the interacting density as a functional of the potential) and (b) returning to a KS scheme. The latter would yield highly accurate densities almost always, and a ML \( E_{\text{xc}}[n] \) would almost certainly not produce a significant density-driven error. Of course, the price paid is the cost of solving the KS equations, but that price is acceptable for the overwhelming applications of DFT at present.

A third important limitation is that, throughout this paper, we have used only uniformly spaced chains of atoms. In practice, of far greater interest would be the case of varying separations, even if all atoms (or molecules) are the same kind, as in an MD simulation of liquid water. We believe the density basis created here should prove very useful in constructing a more general ML functional that would apply to a much larger variety of situations.

In short, there are many issues that must be addressed before this functional could be used in practice. But in theory, machine-learning, combined with the Hohenberg-Kohn theorem, can produce quantum chemical accuracy for strongly correlated solids.

V. ACKNOWLEDGEMENTS

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VI. SUPPLEMENTARY MATERIAL

A. Description of Data

The density matrix renormalization group (DMRG) [16,19] has become the gold standard for calculations in one dimension. The ansatz made for the wavefunction is that of a matrix product state (MPS). This ansatz allows for a site-by-site determination of the wavefunction by concentrating on a small number (in our implementation, two) lattice sites at a time. Once the wavefunction is updated on those two sites, the next two sites are treated. The entire system is swept back and forth until convergence which usually occurs very quickly in one dimension.

To evaluate the Hydrogen chains in this work, an extended Hubbard model [20,21,23],

\[ \mathcal{H} = \sum_{j,\sigma} \left( -\frac{1}{2a^2} (c_{j,\sigma}^{\dagger} c_{j+1,\sigma} + \text{h.c.}) - \mu n_j \sigma \right) + \sum_{j} v_{ij} n_j + \frac{1}{2} \sum_{ij} v_{ij} c_{i}^{\dagger} c_{j} (n_j - \delta_{ij}), \]  

(16)

(17)

can be constructed to recover the continuum limit in the limit of many sites. The prefactor on the kinetic energy terms is chosen to match the finite difference approximation for the kinetic energy with grid spacing \( a \). An external potential is applied in the variable \( v_{ij} \) while \( \mu = -\frac{1}{\sigma} \) for chemical potential \( \mu \). Also, an electron-electron term, \( v_{ij} \), is represented by an exponential function [23]. This exponential mimics the soft-Coulomb interaction in 3d but spherically averaged [23]. The similarity between these functions gives the similar behaviors of the 1d atoms and their 3d counterparts when the symmetry is high.

Systems are calculated with open boundary conditions (“box” boundary conditions). The limit where the box boundary is far from the nearest atomic center is always taken, so no finite size effects appear.

A complication is apparent in 1d that does not appear in 3d. There is no angular momentum in 1d. Thus, not all neutral atoms bind their electrons. One can see this in a reduced example as follows: Consider a delta function interaction in 1d of the form \( -\delta(x - R/2) - \delta(x + R/2) \) [11]. When \( R = 0 \), there is only one solution. At any finite \( R \), the number of electrons that will bind increases from two. The same effect occurs for the exponential interaction, though it is not as easy to see.

This implies that a lower cutoff in the exponentially interaction hydrogen chains will impose a lower limit on suitable chain length. We are interested in systems that do bind all electrons, so a systems below a critical \( R \) are ignored. Table I lists the range of interatomic distances used for each chain. For each Hydrogen chain data generated by DMRG, first sample 80 data from the entire test set range in Table II equi-distantly. This test set is inaccessible in the training process. The rest of data in the training set range in Table II are used as grand training set, where the \( N_T \) training data are uniformly sampled to train the model. The equilibrium test set range is a subset of entire test set range, emphasizing the performance around equilibrium positions. The upper bound is around twice the equilibrium position given by DMRG result.

| \( R \) | \( x \) | \( y \) | \( z \) |
|-------|-------|-------|-------|
| 0     | 1.0   | 1.2   | 8.0   |
| 1.0   | 1.2   | 1.6   | 8.0   |
| 1.4   | 1.6   | 1.8   | 8.0   |
| 8.0   | 1.2   | 1.6   | 8.0   |
| 16.0  | 1.8   | 1.8   | 8.0   |
| 20.0  | 1.8   | 1.8   | 8.0   |

TABLE II. Hydrogen chain data. \( N \) is the number of Hydrogen atoms in the chain. \( R \) is the atomic distance between atoms. The number of DMRG data in each range is in parenthesis.

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