Active learning of potential-energy surfaces of weakly-bound complexes with regression-tree ensembles

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(Dated: 2021-09-23)

Several pool-based active learning algorithms (AL) were employed to model potential energy surfaces (PESs) with a minimum number of electronic structure calculations. Theoretical and empirical results suggest that superior strategies can be obtained by sampling molecular structures corresponding to large uncertainties in their predictions while at the same time not deviating much from the true distribution of the data. To model PESs in an AL framework we propose to use a regression version of stochastic query by forest, a hybrid method that samples points corresponding to large uncertainties while avoiding collecting too many points from sparse regions of space. The algorithm is implemented with decision trees that come with relatively small computational costs. We empirically show that this algorithm requires around half the data to converge to the same accuracy in comparison to the uncertainty-based query-by-committee algorithm. Moreover, the algorithm is fully automatic and does not require any prior knowledge of the PES. Simulations on a 6D PES of pyrrole(H2O) show that <15 000 configurations are enough to build a PES with a generalization error of 16 cm−1, whereas the final model with around 50 000 configurations has a generalization error of 11 cm−1.

I. INTRODUCTION

The ability of molecules to form weakly-bound complexes, via van der Waals or hydrogen bonds, is fundamental to many physical, chemical, and almost all biological processes, ranging from the phenomenon of resonances in reactive scattering4 and roaming in chemical reaction5 to biochemical processes in aqueous solution.6 Small microsolvated complexes of aromatic molecules or chromophores are important model systems for studying the interactions established by individual water molecules at specific binding sites.7 They offer an appealing way for the quantitative investigations of the effects of hydration on the photoexcitation and ionization dynamics and the mechanisms of chemical reactions in general.8–10

First principles calculations of potential energy surfaces (PESs) for weakly-bound complexes are challenging and computationally expensive.11–13 The presence of small energy differences in weakly-bound PES means that generally high levels of theory need to be employed to produce correct asymptotic behavior.13 Furthermore, the landscape of these PESs is complex because of the loosely bound character of intermolecular interactions. Thus, a larger number of grid points is generally required to sample the complete configuration space. Moreover, due to the importance of dynamical electron correlation (disper-
sion) and its slow basis-set convergence15,16 calculations for the noncovalent long-range parts of the PES are generally more costly than the ones at short-range.

The standard approach for building ab initio-based molecular PESs employs a regression procedure: one generates a grid of distinct molecular geometries at which the potential energy is computed, then a model is fitted to these data by minimizing a loss function, e.g., root-mean-square (RMS) error, under a regularization constraint17–19 in recent years, many machine learning (ML) models,20–24 have been used to fit molecular PESs. The most extensively used models include permutationally invariant polynomials,25–29 neural networks (NNs),30–36 Gaussian processes (GPs),37–43 and other kernel methods.44–47

The quality of PESs strongly depend on the chosen set of molecular geometries sampled. This choice is often based on humans’ intervention: the regions that will likely be intensively accessed by the dynamics simulations are sampled more densely. If one’s assumptions about the distribution of geometries accessed in dynamics’ simulations are mostly correct, the question is what minimal amount of data points is needed to reproduce the PES to a desired accuracy.

Trying to minimize the number of electronic structure calculations and reducing the amount of humans’ intervention in an active learning (AL) paradigm became increasingly popular during the last few years.48–51 AL encompasses a variety of iterative algorithms aimed at minimizing the cost of training data acquisition. In pool-based AL, the expert defines a pool of unlabeled data, e.g., molecular geometries, and a policy algorithm queries the energies of geometries that would most significantly minimize the generalization error of the PES once labeled and included in the training data.52 The vast majority of AL applications to PESs used uncertainty-based sam-

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which queries points corresponding to the largest uncertainties in their predicted energies. For probabilistic models like GPs, the uncertainties can be directly calculated. For the ML models that do not offer a direct way to compute uncertainties, these can be inferred by training an ensemble of models on the currently labeled training set and selecting the points about which the models disagree the most. This algorithm is called query-by-committee (QBC) and it was used for uncertainty-based sampling of PESs.

Uncertainty-based sampling is known to be prone to querying outliers because it does not take into account the statistical distribution of data in the pool and hence partly ignores the user’s a priori knowledge embedded in the pool. Upper bounds on the generalization error in an AL setting suggest that more optimal AL strategies can be built by sampling points corresponding to large uncertainties while not deviating much from the true distribution of data. In applications to PESs, uncertainty-based algorithms tend to select many molecular geometries from regions sparsely sampled by subsequent dynamics calculations. When all sparse regions of the pool can be clearly identified, e.g., as points with high energy, or if prior knowledge about the minima and saddle points exists, this problem can be solved by introducing a weighting function. In a more general setting, one can combine the uncertainty-based query algorithm with a molecular-dynamics sampler starting from various known critical points of the PES.

Another concern in AL is the execution speed and the scaling with the amount of data, as AL is an iterative procedure that includes training and predicting energies of many geometries in each iteration.

Here, we propose a regression-version of the stochastic query by forest (SQBF) selection algorithm and use it to generate a minimal grid for the intermolecular PES of weakly-bound pyrrole(H_{2}O) complex. The algorithm selects molecular geometries with large uncertainties in corresponding energy predictions and at the same time keeps the distribution of selected geometries similar to that in the pool. The proposed algorithm does not suffer from the shortcomings of uncertainty-based methods and leads to better results. The uncertainty is estimated using an ensemble of regression trees that constitute a random forest regressor (RFR), while the data statistics is partially preserved using a random sampling (RS) procedure. Owing to the greedy optimization algorithm employed in regression trees, this method is computationally fast and scales well with the size of data. In general, the RFR model is not a continuous function and hence not well suited for modeling of PESs. However, due to its high computational efficiency, the RFR is an attractive method for sampling purposes in AL, which can be easily combined with other ML models, e.g., NNs, to produce the final model. Our empirical results show that the SQBF selection scheme combined with a NN model for the fitting of PES require only 30% of the initial pool data (around 50,000 grid points) to achieve the convergence with a generalization error of about 16 cm^{-1}. In comparison to popular QBC and RS schemes, the SQBF queries about two and three times less data, respectively, to converge to the same accuracy. The SQBF algorithm is general, not restricted to weakly-bound complexes, and applicable to larger molecular systems.

II. METHODS

A. A formal look into active learning

We denote the joint distribution of the problem as \( \mathcal{P} \), where \( \mathcal{P}(x, E) = p(x) p(E|x) \). In standard supervised ML, one has a dataset of points that are independently and identically distributed according to \( \mathcal{P} \). In an AL paradigm, the dataset is selectively sampled from another data distribution \( Q(x, E) \) that has the same conditional distribution of \( \mathcal{P} \), i.e., \( Q(x, E) = q(x) p(E|x) \).

We define \( P = \{X_1, \ldots, X_l\} \) to be the set of all distinct molecular geometries in the pool and assume that \( P \sim p(x) \). We define \( S^{(0)} = \{(X_1, E_1), \ldots, (X_m, E_m)\} \) to be the initially labeled subset of \( P \), where \( m \ll l \). Molecular geometries in the initially labeled set are denoted by \( S_X^{(0)} := \{X_1, \ldots X_m\} \), so that \( P^{U} = P \setminus S_X^{(0)} \) determines unlabeled geometries in the pool. The steps of a pool-based AL algorithm in a batch-mode are summarized in Algorithm 1. To decide whether or not the performance is satisfactory, one needs to choose a stopping criterion. We used the root mean square (RMS) error on a test dataset. We refer to an algorithm that performs step (a) as a policy algorithm.

A simple example of a policy algorithm is uniform RS from the pool. Such an algorithm is representativeness based, as it samples more data from dense regions. In the present case, RS queries more molecular geometries with energies close to the dissociation limit, which is due to the distribution of data in the pool.

An example of an uncertainty-based algorithm is QBC. In this algorithm one trains an ensemble of efficient and diverse learners on the currently labeled dataset.
Algorithm 2: Basic steps of a query-by-committee algorithm for regression problems. $n$ learners are trained on the labeled datasets and asked to make predictions on the whole unlabeled dataset. The $B$ elements chosen to be labeled are those that maximize the standard deviation of the prediction among 5 learners.

\begin{algorithm}
\begin{enumerate}
  \item Train an ensemble $\{T_i\}^n_{i=1}$ of models on data $S^{(i)}$.
  \item Compute predictions $\hat{E}_i = T_i(X)$ \forall $X$ $\in$ $P^U$, \forall $i$
  \item Compute the community disagreement $q(X) = \text{std}(\hat{E}_i)$ \forall $X$ $\in$ $P^U$
  \item Take $B$ elements from the unlabeled data that have the highest $q(X)$
\end{enumerate}
\end{algorithm}

\begin{algorithm}
\begin{enumerate}
  \item Fix the number of learners $n$ in the ensemble.
  \item \textbf{Result:} $B$ elements from $P^U$
  \item \textbf{Input:} $P^U$, $S^{(i)}$, $B$
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      \item Take $B$ elements from the unlabeled data that have the highest $q(X)$
    \end{enumerate}
\end{algorithm}

Algorithm 3: Stochastic query-by-committee algorithm: Data to query are chosen by sampling according to a probability distribution that gives more weights to data points whose predictions are uncertain. Uncertainty is inferred by a standard query-by-committee algorithm.

Algorithm 3 proceeds as follows:

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Algorithm 3: Stochastic query-by-committee algorithm

Input: $P^U$, $S^{(i)}$, $B$

1. Fix the number of learners $n$ in the ensemble.
2. Train an ensemble $\{T_i\}^n_{i=1}$ of models on data $S^{(i)}$.
3. Compute predictions $\hat{E}_i = T_i(X)$ \forall $X$ $\in$ $P^U$, \forall $i$
4. Compute the community disagreement $q(X) = \text{std}(\hat{E}_i)$ \forall $X$ $\in$ $P^U$
5. Compute the weights: $L(X) = q(X) \cdot q_{\text{min}}$, and the sampling probability $p(X)$ = \frac{L(X)}{\sum_X L(X)}$
6. Take $B$ elements from the unlabeled data with probabilities $p(X)$
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Whether the inclusion of a certain point would improve the performance of the regressor, these learners are asked to predict the energies of this point. If they produce very different values, we conclude that adding this point to the training set will improve the performance and query the user about the corresponding energy. The algorithm is formalized in Algorithm 2. We note that diversity of learners is crucial in this algorithm. If the learners are not diverse, their predictions for a certain geometry would be almost the same and hence one would not be able to infer the uncertainty. Practically, the diversity of learners is introduced through random perturbations to the learning process.

Uncertainty-based algorithms do not take the statistical properties of data into account when querying points, which is a serious drawback when the distribution of molecular geometries in the grid is nonuniform. Therefore, we have trained the models on 5 learners and asked to make predictions. Then, querying grid points is performed through random sampling according to a probability distribution that gives more weights to data points whose predictions are uncertain. Uncertainty is inferred by a standard query-by-committee algorithm. The balance between sampling points from the sparse and high-density regions is controlled by the function $L$, which is linear with respect to the community disagreement. The probability of a point being sampled decreases linearly with the decrease of the point’s uncertainty. One can have more freedom on this balance by considering powers of this function, i.e., $L^\alpha$, where $\alpha \in \mathbb{R}^+$. For $\alpha \in (0, 1)$, the algorithm will sample more points with low uncertainty and conversely less for $\alpha \in (1, \infty)$. We performed a heuristic study of the effect of different powers $\alpha$. At each AL iteration, we ran SQBF algorithm for different values of $\alpha \in \{0.5, 0.75, 1, 1.25, 1.75\}$. We picked the $\alpha$ that led to the largest improvement in generalization error and collected the corresponding queried points. We proceeded to active learning using this batch as part of the pool. The whole procedure was repeated at every AL iteration. We found only minor improvements of the accuracy when using multiple, optimized, values of $\alpha$.

While any ensemble of ML models can be used, we propose to use the trees of an RFR as members of this ensemble. In the next section, we provide a heuristic study of the effect of different values of powers $\alpha$.

Regression trees are a non-parametric way of solving a regression problem. They are based on the intuition that the output value can be inferred by partitioning the input space. In particular, for solving a regression problem with data pairs $\{(X_i, E_i)\}_{i=1}^n$, a Tree-Regressor aims at finding
A powerful approach to mitigate this problem is to consider an ensemble of trees. The key idea is that averaging a set of independent random variables, which have comparable variances, reduces their overall variance. In an ensemble method, a random perturbation is introduced to the learning process in order to produce several different learners from the same training set. Thus, taking the average of the predictions of the ensemble would result in a reduction of variance. Such a random perturbation can be introduced by bootstrapping, which gives rise to bootstrap aggregation (bagging) methods. Here, $B$ different bootstrapped datasets of size $m_b$ are generated. A tree is built on each model. For a new data point, a prediction is made by taking the average of the predictions of all trees: $T_{bag}(x) = \frac{1}{B} \sum_{i=1}^{B} T_i(x)$.

A further random perturbation in tree models can be introduced by considering, at each split, only a randomly drawn subset of all possible features. This gives rise to extremely randomized trees the average time complexity for training is $\Theta(M \cdot K \cdot N \log_2 N)$ [23]. The average inference complexity of RFR is $\Theta(M \log N)$ [22]. Thus, one AL iteration scales as $O(M \cdot K \cdot \tilde{N} \log_2 \tilde{N})$ with the number of so-far-labeled data $N$. The complexity of RFR is asymptotically inferior to that of a NN, which has a training time complexity of $O(N_e N (\sum_{j=1}^K N^j (N^j+1)))$ with the number of epochs $N_e$ needed for the NN to converge and the number of neurons $N^j$ in layer $i$. However, in the data size regime of our application, the computational costs of an RFR are smaller than that of the NN.

III. SIMULATIONS

We applied different policy algorithms and tested their performance for building the prototypical intermolecular PES of the pyrrole(H$_2$O) complex. Due to the highly fluxional nature of the hydrogen bond, the intermolecular motions in pyrrole(H$_2$O) are highly delocalized, rendering the calculation and representation of the PES very challenging. The intramolecular vibrations in the pyrrole and water moieties can be described with a relatively simple, though multi-dimensional, single-minimum PES and thus, for simplicity of calculations, were not considered here. We fixed the structures of pyrrole and water monomers to the experimentally determined values, see supplementary material, and varied the six intermolecular coordinates, shown in Figure 1. These are defined as follows: the relative position of water with respect to pyrrole is described by the three spherical coordinates $R = [0, 2, 1]$ nm, $\theta = [0, \pi]$, $\phi = [0, \pi]$ and the relative orientation of water is defined by the three Euler angles $\alpha = [0, \pi]$, $\beta = [0, \pi]$, $\gamma = [0, \pi]$. The angles $\phi$, $\alpha$, and $\gamma$ were restricted to the ranges $[0, \pi]$ exploiting the $C_{2v}(M)$ symmetry of the complex.

The pool of molecular configurations was generated a priori as the direct product of one-dimensional grids for every degree of freedom and contained 57500 different molecular geometries covering the potential energy up

$J$ distinct and non-overlapping regions $R_1 \ldots R_J$ in the feature space that minimize

$$\sum_{j=1}^{J} \sum_{i \in R_j} (E_i - \bar{E}_{R_j})^2$$

where $\bar{E}_{R_j} = \sum_{i \in R_j} E_i / m_j$ is the average energy of $m_j$ molecular geometries in region $R_j$. [29] This problem is NP-complete. [29] Therefore, only near-optimal solutions are considered by restricting ourselves to hyper-rectangular regions and using recursive binary splitting, a greedy algorithm to obtain a near-optimal segmentation. A prediction for a new molecular geometry is done by assigning the input geometry to one of the regions.

The average time complexity for training is $\Theta(M \cdot K \cdot N \log_2 N)$ [23]. The average inference complexity of RFR is $\Theta(M \log N)$ [22]. Thus, one AL iteration scales as $O(M \cdot K \cdot \tilde{N} \log_2 \tilde{N})$ with the number of so-far-labeled data $N$. The complexity of RFR is asymptotically inferior to that of a NN, which has a training time complexity of $O(N_e N (\sum_{j=1}^K N^j (N^j+1)))$ with the number of epochs $N_e$ needed for the NN to converge and the number of neurons $N^j$ in layer $i$. However, in the data size regime of our application, the computational costs of an RFR are smaller than that of the NN.

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to 5000 cm\(^{-1}\) above dissociation. All coordinates were sampled more densely in the vicinity of the equilibrium geometry. Also, the angular coordinates were sampled more densely for small radial distances \(R \leq 500\) pm with a sparser grid for \(500 < R \leq 1000\) pm. This led to a nonuniform distribution of energies in the pool, shown in Figure 2. Note that a direct-product grid is not essential for the accumulation of the pool of unlabeled geometries and the test dataset. Here, it was used mainly because it allows the coverage of the whole configuration space that is relevant for the subsequent quantum dynamics’ simulations, and hence prevents biases and holes in the pool and test data. While this method is not arbitrarily extendable to systems with more degrees of freedom, other pool accumulation methods\(^{55}\) could be used without modifications to the SQBF approach. The electronic structure calculations employed the density-fitting explicitly-correlated DF-MP2-F12 level of theory\(^{75,76}\) in the frozen-core approximation using aug-cc-pVDZ-F12\(^{71}\) atomic orbital, cc-pVDZ-F12+/OPTR\(^{53}\) resolution of the identity, and aug-cc-pVDZ/MP2 FIT\(^{55}\) density fitting basis sets. The geminal exponent was fixed at 1.0. The electronic structure calculations were carried out using Molpro\(^{53,55}\). A subset of 10\% of the total number of points in the pool was randomly selected as a test set and taken out of the pool (OOP). 5\% of the remaining data was randomly selected as a validation set. We employed two different machine learning models, RFR, and NN, to fit the data. We used exponential functions of interatomic distances, with all distances considered, as molecular descriptors, see Appendix 1.

Furthermore, based on reviewer suggestions we tested the SQBF algorithm and an NN model on the PES of the \(\text{N}_4\) molecule using previously reported electronic structure data\(^{54}\).

### Performance

For pyrrole\((\text{H}_2\text{O})\) we compared the performance of the RS, QBC, and SQBF AL policy algorithms considering the convergence rate and the fitting accuracy. For QBC (algorithm 2) and SQBF (algorithm 3) we used the trees of an RFR as an ensemble of learners. All policy algorithms started from the same fixed amount of \(m = 2458\) labeled samples and queried the same equal number of \(m\) samples at every AL iteration. For every iteration and query algorithm, we used the RFR and NN models to fit the data. The fitting error is defined as the RMS error of the ML models in predicting the energies on the OOP dataset. This dataset was the same for all policy algorithms and followed the joint distribution of the problem \(P\). The accuracy of a model on this dataset is an estimate of the generalization error. Further technical details on the training process of the RFR and NN models and their hyper parameters are provided in Appendix 1.

The fitting errors of the RFR and NN models for different policy algorithms are plotted in Figure 2 as functions of AL iteration. The SQBF strategy with RFR model leads to the fastest convergence of the error. QBC strategy outperforms RS. Similar convergence behavior of different policy algorithms can be observed for the NN model. For our dataset, the fitting error of NN was smaller than that of RFR for all AL iterations and for all strategies by an average factor of 3.3. Table I summarizes these results. The better performance of NNs is partially due to the fact that NNs are easier to train to higher accuracy and can approximate complex functions with a better control on the bias-variance trade-off, which was enabled by using an early stopping criterion on the validation set, see Appendix 1. The AL iterations were terminated when the pool became empty. In practice, the iterations are to be terminated when the derivative of the fitting error with respect to the amount of labeled data is less than a predefined value\(^{53}\) or simply when the fitting error of the model is small enough.

Similarly, for the \(\text{N}_4\) molecule the SQBF algorithm was used to query geometries from the pool of 16421 molecular geometries reported\(^{54}\). The OOP and validation datasets were each generated using 10\% of the uniform-randomly sampled pool data. An initial batch of 240 geometries was uniform-randomly sampled from the pool and the SQBF algorithm queried 240 geometries at each AL iteration.

### Table I. Out-of-the-pool RMS errors (in cm\(^{-1}\)) of the random forest regressor and neural network models, listed as RFR/NN, computed for various fractions of the total pool data collected by the different AL policies

| AL policy  | 20 % | 40 % | 60 % | 80 % | 100 % |
|------------|------|------|------|------|-------|
| RS         | 183  | 117  | 81   | 52   | 39    |
| QBC        | 141  | 74   | 49   | 41   | 38    |
| SQBF       | 88   | 37   | 36   | 38   | 39    |

FIG. 2. The probability density distribution of the energies corresponding to all molecular geometries in the pool. The histogram was calculated for a bin width of 34.5 cm\(^{-1}\) and has a peak at 1600 cm\(^{-1}\), corresponding to the dissociation limit of pyrrole(\(\text{H}_2\text{O}\)).
FIG. 3. RMS error of out-of-the-pool datasets using (up) the random forest regressor and (down) the neural network models for the RS (triangles), QBC (circles), and SQBF (squares) policies. The SQBF has the fastest convergence. A neural network model, trained on 30% of the total amount of datapoints in the pool achieves an RMS error of 16 cm$^{-1}$. The RMS error on the full dataset is 11 cm$^{-1}$. The neural networks trained on data collected by the QBC or RS algorithms show worse performance. The same convergence patterns hold when using a random forest regressor to train on the data instead of a neural network, albeit at overall somewhat slower convergence.

The same molecular descriptor as described above was used to transform the data and an NN model was used for fitting; details on the NN design are provided in Section 1. This procedure was repeated 100 times and the mean and standard deviation of the resulting NN errors on the entire dataset as a function of the number of training examples is reported in Table II. The SQBF results are compared with the ensemble of 100 GPs used to fit the data collected by the latin hypercube sampling algorithm. The GP method shows a better performance for the first few AL iterations. We attribute this to the fact that it is hard to prevent overfitting with a neural network with a very small set of randomly selected training data. However, already at 1200 training points the two models result in comparable accuracy. With 1680 training points, our SQBF/NN approach achieves the same accuracy as GP with 2400 points, which corresponds to a 30% reduction in the size of the training dataset. All the following further investigations are performed for pyrrole(H$_2$O).

TABLE II. RMS mean errors and standard deviations using the available data (in cm$^{-1}$) of NN using the AL SQBF algorithm with a NN to fit the data (present work) and latin hypercube sampling with GPs to fit the data (in cm$^{-1}$), applied to PES data of the N$_4$ molecule [87].

| No. training data | NN (cm$^{-1}$) | GP (cm$^{-1}$) |
|-------------------|---------------|---------------|
| 240               | 36518 ± 2697  | 13300 ± 2770  |
| 480               | 26207 ± 1871  | 10027 ± 1371  |
| 720               | 11192 ± 1200  | 8401 ± 1102   |
| 960               | 8111 ± 668    | 7544 ± 972    |
| 1200              | 6201 ± 462    | 6806 ± 962    |
| 1680              | 4704 ± 612    | -- ± --       |
| 1800              | 4494 ± 633    | 5551 ± 951    |
| 1920              | 4284 ± 658    | -- ± --       |
| 2400              | 3557 ± 675    | 5012 ± 832    |

B. Distribution of queried data

In Figure 4 we plotted the normalized distributions of the samples' electronic energies of pyrrole(H$_2$O) collected by different AL policies at three different iterations corresponding to 20%, 40%, and 60% of the total pool. We compare these with the distribution of energies in the total pool, which has a peak around 1600 cm$^{-1}$, corresponding to the dissociation limit of pyrrole(H$_2$O). The densities were computed using 200 equally-sized bins covering the energy range from 0 to 6874 cm$^{-1}$ and normalized to the bin width of 34.5 cm$^{-1}$. Evidently the probability density of data sampled by the RS policy most closely resembles the pool distribution. On the other hand, it is clear that the QBC algorithm samples more data with higher energies, whereas SQBF keeps a balance between both the RS and QBC tendencies. As the number of the labeled data increases, all probability density distributions become more similar to the distribution in the pool.

It is reasonable to expect that a model built on a dataset sampled by QBC algorithm will tend to have a better performance for the high-energy regions. This is demonstrated in Figure 5 showing the 2D histograms of OOP energies and the absolute errors of the RFR and NN models in predicting these energies, plotted for different policy algorithms. The histograms were computed using 20 and 50 equally-sized bins for the energy and absolute errors, respectively. The size of the training dataset here corresponds to 40% of the pool’s size. We clearly see that RS achieves good accuracy for the points with low energies, QBC works best for the points with high energies, and the SQBF maintains a more regular accuracy across the whole energy spectrum.
FIG. 4. Normalized probability density distributions of the number of data points $N/N_{\text{tot}}$ across the potential energies plotted for the data collected by the RS, QBC, and SQBF policies at different AL iterations corresponding to 20 %, 40 %, and 60 % of the total pool. The bin width of the histograms is 34.5 cm$^{-1}$.

C. Dependency on batch size and size of initially labeled dataset

We repeated the above calculations with a smaller batch size of 122 points instead of initially used 2458, starting from the same initially labeled dataset. The convergence of the RFR fitting error with the number of training data are plotted in [Figure 6] for different policy algorithms. Here, we note that both QBC and SQBF strategies benefit slightly from using a smaller batch size. This is in accordance with previous studies that showed a decreasing performance of QBC with increasing batch size, which is due to collecting many similar samples.

We also studied the effect of changing the size of initially labeled dataset. [Figure 7] shows the RFR fitting errors for different policy algorithms obtained from initial datasets of 100 and 2458 samples with the batch size of 2458. We observe that RS policy algorithm outperforms QBC, and that the accuracy of QBC declines significantly. This suggests that with a fewer number of initially labeled data, an AL strategy should focus on collecting grid points from dense regions of the configuration space rather than sampling points with high uncertainties in their predictions. Notably, the SQBF performance is not affected by the size change.

IV. SUMMARY AND CONCLUSION

The first principles calculations of molecular PESs, especially for molecules with many fluxional degrees of freedom, are computationally expensive. One of the major bottlenecks originates from the need to perform the expensive quantum chemical calculations for tens and hundreds of thousands of different molecular geometries. Algorithms that allow to reduce the number of necess-
We developed an accurate and automated procedure to efficiently sample the molecular geometry grid points leading to a systematic convergence of the accuracy of PES. We employed a regression version of SQBF, a pool-based active learning algorithm to generate a compact grid of molecular geometries and the RFR and NN ML-models to construct the six-dimensional intermolecular PES of the weakly-bound pyrrole(H$_2$O) complex. The proposed method samples grid points with high uncertainties in the corresponding predictions of energies and at the same time preserves the statistical information embedded in the pool. In our benchmark application to the six-dimensional intermolecular PES of pyrrole(H$_2$O) this led to a roughly two times faster convergence with respect to the number of grid points than the commonly used QBC algorithm to represent the PES to an accuracy of about 16 cm$^{-1}$. Furthermore, the PES fitted on the data sampled by SQBF exhibited a more uniform accuracy across the whole energy spectrum in comparison to QBC. We empirically showed that the SQBF method is not very sensitive to a variation of parameters such as the size of initially labeled data and size of the batch.

In addition, the proposed method is computationally cheap and scales well with the size of the labeled data $N$, i.e., as $\Theta(M \cdot K \cdot N \log^2 N)$, where $K, M$ denote the number of random features sampled at each splitting and the number of trees, respectively, $N = 0.632N$. This makes the method attractive for developing universal ML-potentials where large datasets are needed [94,95]. In the case when the accuracy of the RFR is not sufficient for the application of the PES, we showed that the data can be used equally-well by other ML models like NNs. An alternative would be to employ a bagging algorithm [93] with any other ensemble of learners or even with a model that offers a direct computation of uncertainty.
Overall, the presented procedure is general and can be applied to the PESs of any polyatomic molecule. It can also be used to model other physical properties like dipole-moment or polarizability surfaces. The major advantage of the proposed method over more popular QBC approach is the heuristic sampling procedure that preserves the distribution of data in the pool while keeping the uncertainty as the primal selection criterion. We believe that in the future the general approach can be improved even further by a better tuned balance between uncertainty and representativeness.

SUPPLEMENTARY MATERIAL

See supplementary material for the computed potential energies of pyrrole(H2O), and the structures of pyrrole and water monomers.

ACKNOWLEDGMENT

This work has been supported by Deutsches Elektronen-Synchrotron DESY, a member of the Helmholtz Association (HGF), the Data Science in Hamburg HELMHOLTZ Graduate School for the Structure of Matter (DASHH, HIDSS-0002) and by the Deutsche Forschungsgemeinschaft (DFG) through the cluster of excellence “Advanced Imaging of Matter” (AIM, EXC 2056, ID 390715994). We acknowledge the use of the Maxwell computational resources operated at Deutsches Elektronen-Synchrotron DESY, Hamburg, Germany.

DATA AVAILABILITY

The data that supports the findings of this study are available within the article and its supplementary material. The computer code and all relevant data for this work can be obtained from the git repository at https://github.com/CFEL-CMI/Active-Learning-of-PES.

APPENDIX

1. Technical details on the random perturbations in AL algorithms and the training of machine learning models

The regression trees used to implement the QBC and SQBF algorithms were both built using the scikit-learn (sklearn) Python package.\(^\text{68}\) All AL algorithms used here were written based on the Libact Python package.\(^\text{69}\) For both the QBC and SQBF algorithms we used an ensemble of 100 trees. The training during all AL iterations used an exponential function of the inter nuclear distances as a molecular descriptor: 

\[
1 - \exp(-\left(r - r_0\right)) \quad \text{where} \quad r, r_0 \text{ denote the actual distance and equilibrium distance between two atoms, respectively.}
\]

The perturbation of the learning process is controlled through two parameters: (i) a bootstrapping parameter \(\gamma\) that determines the fraction of data sampled by each tree and (ii) the number of features \(\beta\) sampled randomly by each tree. For the batch sizes used in the simulations we experimented with several combinations of these parameters and obtained the best convergence for \(\gamma = 0, \beta = 12\) for simulations on pyrrole(H2O) and \(\gamma = 0, \beta = 4\) for simulations on N2. The same parameters were used for uncertainty estimation in both the QBC and SQBF algorithms. Minimal cost complexity pruning was used to reduce the overfitting of RFR with complexity parameter \(c = 0.01\). Since we require that algorithm 3 queries exactly \(B\) geometries, one may run into a situation where the number of entries with non-zero probabilities of the distribution \(p\) is less than \(B\). In such a case we chose to query the elements with highest uncertainty. The effect of this choice on the simulations conducted in the manuscript is negligible since we only encountered this case once in one of the last AL iterations.

The NN used is a multilayer perceptron and training was implemented using the Python Tensorflow package.\(^\text{90}\) The NN has three hidden layers with 256, 512, and 256 neurons, respectively, and a single neuron output layer. The second and third layers were \(l_2\)-regularized with a regularization parameter of \(10^{-5}\). All hidden layers used “ReLU” as the activation function. The ReLU activation function could be substituted with a smooth approximation such as Softplus to yield a smooth PES. The same aforementioned molecular descriptor was used. The networks were trained for 250 epochs using the Adam optimization algorithm with an initial learning rate of 0.0025 and a decaying learning rate schedule \((lr_{current} = 0.9825 \times lr_{previous})\). An early stopping callback was employed on the validation set that was taken out-of-the-pool with patience of 25. The NN hyper parameters were set to obtain a sufficiently accurate NN, with test error of around 10 cm\(^{-1}\) when using all the training data.

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