## Machine Learning Classification can Significantly Reduce the Cost of Calculating the Hamiltonian Matrix in CI Calculations

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## Abstract

Hamiltonian matrices in electronic and nuclear contexts are highly compute-intensive to calculate, mainly due to the cost for the potential matrix. Typically these matrices contain many off-diagonal elements that are orders of magnitude smaller than diagonal elements. We illustrate that here for vibrational H-matrices for H<sub>2</sub>O, C<sub>2</sub>H<sub>3</sub> (vinyl) and C<sub>2</sub>H<sub>5</sub>NO<sub>2</sub> (glycine) using full-dimensional *ab initio*-based potential surfaces. We then show that many of the these small elements can be replaced by zero with small errors of the resulting full set of eigenvalues, depending on the threshold value for this replacement. As a result of this empirical evidence, we investigate three machine learning approaches to predict the zero elements. This is shown to be successful for these H-matrices after training on a small set of calculated elements. For one vinyl and glycine H-matrices, of order 15 552 and 8 828, respectively, training on a percent or so of elements is sufficient to obtain all eigenvalues with a mean absolute error of roughly 2 cm<sup>-1</sup>. The most general quantum approach to obtain accurate energies and wavefunctions of bound states of electrons (in electronic structure theory) and nuclei (vibrational dynamics) is "configuration interaction" (CI). This approach typically leads to a large Hamiltonian matrix, the cost of which to obtain scales exponentially or factorially with the size of the underlying basis. This bottleneck has stimulated numerous approaches to reduce the size of the basis, which of determines the size of the H-matrix. For approaches in electronic structure, see for example refs. 1–4. Here we focus on vibrational dynamics, which has some prominent differences with electron motion; perhaps the biggest one is the need to obtain hundreds to thousands of eigenvalues from nuclear CI compared to a much smaller number in electronic CI.

To begin, recall that in the widely-used direct-product (Hartree) basis for f-modes the H-matrix is given by

$$\langle n'_1, n'_2, ..., n'_f | H | n_1, n_2, ..., n_f \rangle,$$
 (1)

where the quantum numbers are the usual ones for this basis. For example, these could be harmonic-oscillator functions in normal coordinates of a potential minimum. More sophisticated 1-mode bases are used currently, but the details of these are not relevant. The interested reader can consult relevant chapters in a recent book on computational approaches to vibrational dynamics.<sup>5</sup> In applications, the values of these quantum numbers for each 1mode basis is restricted. In addition, the sum of quantum numbers of all modes is typically bounded. This is probably the most common method to restrict the basis and goes back 40-50 years.<sup>6,7</sup> More fine-grained restriction methods are used in the code MULTIMODE, <sup>8–10</sup> where the excitation space is restricted, as in electronic structure methods, to "singles", "doubles", "triples", "quadruples", etc. Then separate excitation bounds are applied to each level of excitation. Also, when applicable, point-group symmetry is also used to block-diagonalize the *H*-matrix.

The above (and perhaps other) approaches all reduce the basis compared to an unrestricted basis. And this does result in a major reduction in the size of the H-matrix. We might refer to this as a basis-restricted H-matrix. It is possible to go further and directly eliminate elements of the matrix. This has been done, albeit rarely, in the vibrational dynamics literature. One example uses a method suggested by Handy and Carter<sup>11</sup> and used for vibrational energies of C<sub>2</sub>H<sub>4</sub> and CH<sub>3</sub>NO<sub>2</sub>.<sup>12</sup> The method is based on second-order perturbation theory to eliminate elements and borrows from similar ideas in electronic structure theory. A direct approach has been given in ref. 13 as a means to reduce the storage of the H-matrix, illustrated for C<sub>2</sub>H<sub>4</sub>. This is simply to neglect Hamiltonian matrix elements below an absolute value of about 0.00002 cm<sup>-1</sup>. Doing so resulted in a major reduction is the number of elements stored in memory.

While it may be possible to exploit the presence of very small matrix elements in memory storage and even reduce the size of the matrix, it still requires the calculation of the *H*-matrix. Clearly, it would be advantageous if the determination of very small elements could be done without calculating them. Here we report several machine learning classification approaches to achieve this. This is illustrated for  $H_2O$ ,  $C_2H_3$  (vinyl), and  $C_2H_5NO_2$  (glycine).

To proceed, we give histograms of five H-matrix elements for these molecules obtained, with one exception, using MULTIMODE, which uses the exact Watson Hamiltonian with a direct product basis and the *n*-mode representation<sup>14,15</sup> of full-dimensional *ab initio* potential energy surfaces. Potential matrix elements were calculated using optimized numerical quadrature. In one case, 4-mode vinyl, the *H*-matrix was obtained using a direct-product of numerical basis functions in the normal coordinates of the saddle-point separating the two equivalent minima in vinyl. Details of the steps used in these calculations have been given previously for  $H_2O$ ,<sup>8</sup> vinyl,<sup>16,17</sup> and glycine<sup>18</sup> and the interested reader if referred to those references for details.

First, consider the *H*-matrix for  $H_2O$  using a high-level potential.<sup>19</sup> Restriction of the basis was done using an excitation space up to triples. Point group symmetry was also used. The resulting *H*-matrix consists of two blocks ( $A_1$  and  $B_2$ ) of orders 161 and 125, respectively. (Even with these small matrices, the eigenvalues for the fundamentals and the

bend overtone are within less than  $1.0 \text{ cm}^{-1}$  of "exact" results.) A histogram of the matrix elements for the  $A_1$  block is given in Figure 1. As seen many elements are small; 1 192 of the 13 041 elements are less than  $10^{-10}$  hartree (0.00002 cm<sup>-1</sup>) in magnitude and so can be classified as negligible. Of course many more elements are also small, e.g., 7 310 have magnitude less than  $10^{-5}$  hartree (2 cm<sup>-1</sup>). Whether these can classified as negligible, i.e., assigned the value zero, or not requires quantitative investigation and that is done after histograms for vinyl and glycine are given.



Figure 1: Histogram of H-matrix elements in cm<sup>-1</sup> for the  $A_1$  symmetry block of for H<sub>2</sub>O. See text for more details.

Next consider the histogram of H-matrix elements from MULTIMODE for 5-mode vinyl, shown in Figure 2, using an accurate *ab initio*-based potential.<sup>20</sup> Note we used an option in MULTIMODE to consider a subset of modes. In this case 5 modes and a quintuples excitation space. As seen the vast majority of elements have a magnitude of order 0.01 cm<sup>-1</sup>.

The final histogram we show is for 10-mode glycine in Fig. 3, where it appears that the vast majority of elements are either very small or exactly zero. The "exact-zeros" occur only



Figure 2: Histogram of *H*-matrix elements in hartree for 5-mode vinyl on a semi-log scale



Figure 3: Histogram of H-matrix elements for the A'' symmetry block for 10-mode glycine on a semi-log scale

for 10-mode glycine among the five *H*-matrices considered here. We explain this below.

To proceed we determine the quantitative relationship between the magnitude of matrix elements that are be set to zero, based on a threshold, and the resulting mean absolute error (MAE) of all the eigenvalues for each *H*-matrix. (The number is the same as the order of the matrix and this is given for convenience in the second column of Table 1.) This relationship is shown in Figure 4 for five *H*-matrices. The two additional ones are for 3-mode and 4-mode vinyl. The latter one is the exception mention above that was not obtained with MULTIMODE. It was obtained used saddle-point normal modes, following earlier work.<sup>17</sup> As seen, and as expected, the fraction of elements set to zero and the MAE of all eigenvalues increases with increasing threshold. For the threshold of  $10^{-5}$  hartree (2.2 cm<sup>-1</sup>) the MAE is at most 1 cm<sup>-1</sup>. Also at that threshold the percentage of zeroed elements increases from about 25 to 90+ percent. For glycine with a percentage just below 100 %, the histogram includes numerous elements that are set to zero in MULTMODE without calculating them. Glycine is the only example where this occurs and as noted earlier we provide the explanation below, where we examine the n-mode representation of the potential.

We now describe the machine learning classification (MLC) approaches we use to predict zero matrix elements. This is a standard 0/1 classification problem: in the training set, elements with a magnitude greater than the threshold are labelled 1, and elements with smaller magnitudes are labelled 0. The ML model then will predict 0 or 1 for an unknown matrix element; if it predicts 0, the matrix element can be set to zero so the computation can be skipped, and otherwise if the prediction is 1, the actual value matrix element will be calculated.

For demonstration purposes, we use a threshold of  $1 \times 10^{-5}$  hartree, 2.2 cm<sup>-1</sup>, as this achieves good accuracy in the eigenvalues and considerable savings in the computations. Using this threshold, we label *H*-matrix elements as 0 or 1 to create a binary dataset that can be used in machine learning classification. The features ( $\mathbf{X}$ ) for this classification problem are the quantum numbers of the 1-mode basis in *bra* and *ket*,  $\langle n'_1, n'_2, ..., n'_{nmode} |$  and



Figure 4: Fraction of matrix elements classified as zero as a function of the threshold for the 5 *H*-matrices studied in this work (upper); mean-absolute-error (MAE) of all eigenvalues as a function of the threshold (lower). The dashed line shows the threshold we use for the ML classification,  $1 \times 10^{-5}$  hartree (2.2 cm<sup>-1</sup>).

 $|n_1, n_2, ..., n_{nmode}\rangle$ . We investigate three widely used classification algorithms on the water *H*-matrix: random forest (RF), support vector machine (SVM), and multilayer perceptron (MLP). All ML models were trained using scikit-learn package.<sup>21</sup>

First consider the smallest H-matrix (of order 161) for water, with distribution of elements shown in Fig. 1. There are 13 041 matrix elements. These are randomly split into 40% training and 60% testing. For MLP and SVM models, the feature X is standardized to zero mean and unit variance by using the StandardScaler from scikit-learn, while for the RF model, we did not scale the features because it is invariant to feature scaling. The RF uses 20 estimators (also known as "decision trees") with maximum depth of 10. The MLP model uses two hidden layers with 15 neurons in each, and 'relu' as the activation function. Early stopping is used to mitigate overfitting when the validation score (which is based on 20% of the training data, set aside as the validation set) does not improve for 10 consecutive training epochs. The SVM model uses C-Support Vector with radial basis function (RBF) kernel.

For 5-mode and 3-mode vinyl the *H*-matrices are of 1 152 and 1 960, respectively, so the numbers of matrix elements are 664 128 and 1 921 780. The SVM algorithm implemented in scikit-learn becomes too expensive, so only RF and MLP classifiers are considered. In these two cases, only 10% of the data set is used as the training data. This is produces sufficient data to train on. And, more importantly since the goal of the ML approach is to accurately predict negligible matrix elements and thus avoiding calculating them, we would like to keep the training set as small as possible. For these matrices the MLP classifier uses 20 neurons in each hidden layer and The RF classifier use 50 estimators with maximum tree depth of 20.

Moving on to much larger H-matrices, consider 4-mode vinyl using a saddle-point normalmode basis and the B<sub>2</sub>-symmetry block of the 10-mode glycine. The respective H-matrices are order 15 552 and 8 828. Thus there are tens or hundreds of millions of matrix elements, and for vinyl each matrix element requires 4D quadrature. Note, for glycine, although we included 10 normal modes in the basis, we used 4-mode representation of the potential so the highest dimension for the numerical quadrature of potential matrix elements is still 4D. For glycine, as explained in detail below, because the number of modes is larger than than 4, many matrix elements are known to be exactly zero and so not calculated. In these two cases, only 1% of the matrix elements are used to train the ML models. The MLP models for these two examples have three hidden layers of 20 neurons, while the RF models use 50 estimators with maximum depth of 25. For 4-mode vinyl that uses the saddle-point normalmode basis, the *H*-matrix is of order 15 552 and only the MLP classifier is used, since the random forest also becomes too expensive.

Now we present the training and testing accuracy of the ML models on the five Hmatrices in Table 1. It can be seen that all ML models give accurate predictions for the labels, with an accuracy close to or greater than 90%. The random forest models slightly overfit the training set, indicated by the difference in the training and testing accuracy.

Table 1: Training/testing accuracy of the ML predicted labels of the matrix elements for the five H-matrices.

H-matrix	Order	SVM	$\operatorname{RF}$	MLP
H <sub>2</sub> O	161	$0.891 \ / \ 0.885$	$0.963 \ / \ 0.933$	0.889 / 0.882
5-mode vinyl	1152	-	$0.994 \ / \ 0.970$	$0.966 \ / \ 0.963$
3-mode	1960	-	$0.984 \ / \ 0.93 \ \text{vinyl} \ 2$	$0.915 \ / \ 0.914$
10-mode glycine	8 828	-	$0.999 \ / \ 0.992$	$0.994 \ / \ 0.993$
4-mode SP vinyl	15  552	-	$0.975 \ / \ 0.879$	$0.872 \ / \ 0.872$

Next we reconstruct the *H*-matrices using the predicted labels from various ML models. If the predicted label for a matrix element is 1, we keep the exact value of that element, and if the predicted label is 0, we simply set that matrix element to zero. Then we diagonalize the reconstructed *H*-matrices and compare the resulting eigenvalues with the exact eigenvalues of the exact *H*-matrices. In Table 2, we present the mean-absolute-error (MAE) of all the eigenvalues. We can see that the MAE of the eigenvalues are about  $1-2 \text{ cm}^{-1}$ , except for the RF model on glycine. This is probably due to the very unbalanced 0/1 label in this *H*- matrix. When using a threshold of  $10^{-5}$  hartree, 98.9% of the matrix elements are labelled 0, so a small amount of wrong predictions may cause relatively large error. Nevertheless, this level of accuracy in the eigenvalues is satisfactory compared to the the errors caused by the underlying electronic structure theory, fitting precision, etc.

Table 2: Mean-absolute-error (in  $cm^{-1}$ ) in the eigenvalues of the approximate *H*-matrices based on the predicted labels from different ML models on the matrix elements.

<i>H</i> -matrix	SVM	RF	MLP
H <sub>2</sub> O	1.12	0.67	1.42
5-mode vinyl	-	1.31	0.84
3-mode vinyl	-	0.94	1.37
10-mode glycine	-	7.37	1.62
4-mode SP vinyl	-	2.23	2.17

Finally, we present the timings for training these ML models and make predictions on the matrix elements. The goal of pruning the matrix elements is to save computer times for "zeros", so the training and prediction of the ML models must at least be more efficient than actually computing those matrix elements. We only test the timings on the largest H-matrix, i.e., from 4-mode saddle point vinyl, as the matrix elements are expensive to compute in this example. The ML approach may not have the advantage if the matrix size is small to medium, since computing all matrix elements in a small or medium H-matrix is still fast, while the overheads of training ML models make this approach slower than simply computing all the elements. For this 4-mode vinyl H-matrix, training the RF model takes only 48 seconds and predicting the off-diagonal elements take 389 seconds, and the training and prediction of the MLP model takes 151 and 36 seconds, respectively. As a comparison, computing the H-matrix exactly takes hours, so the overhead of the ML model is small. The advantage of the ML approach will become more significant as the size of the H-matrix increases.

In summary, the ML classification approaches are quantitatively accurate for the Hmatrices examined here by training on just 1% of the elements with a major reduction on the number of predicted non-negligible elements to be calculated. For the largest examples, 4-mode vinyl with the saddle point basis and 10-mode glycine the reductions are around 90%. Also the results here show that the percentage reduction increases with the number of modes.

Figure 5 shows a flow chart of our proposed approach that would be implemented for a calculation of the H-matrix. As usual the starting point is the restricted basis for the calculation of the H-matrix.



Figure 5: Flow chart of the machine-learning approach to prune the H-matrix elements. Note: \* Could train committee ML models; \*\* Optionally could try other thresholds.

Finally, to provide insight into the large number of very small matrix elements consider the *n*-mode representation of the potential,<sup>8,10,14</sup> which some readers may know is used in MULTIMODE. In brief it represents the potential by

$$V(Q_1, Q_2, \cdots, Q_f) = \sum_i V_i^{(1)}(Q_i) + \sum_{i,j} V_{ij}^{(2)}(Q_i, Q_j) + \sum_{i,j,k} V_{ijk}^{(3)}(Q_i, Q_j, Q_k) + \sum_{i,j,k,l} V_{ijkl}^{(4)}(Q_i, Q_j, Q_k, Q_l) + \cdots,$$
(2)

where  $V_i^{(1)}(Q_i)$  is the one-mode potential, i.e., the 1D cut through the full-dimensional PES in each mode, one-by-one,  $V_{ij}^{(2)}(Q_i, Q_j)$  is the intrinsic 2-mode potential among all pairs of modes, etc. Here, intrinsic means that any *n*-mode term is zero if any of the arguments is zero. This representation is exact if there are f terms. In MULTIMODE this level goes up to a maximum of  $V_{ijklmn}^{6}$  terms. For H<sub>2</sub>O, C<sub>2</sub>H<sub>3</sub> with 3, 4, and 5 modes this expansion was taken to the level of being exact. For the VCI calculation using saddle point normal modes the full 4-mode potential was used without expansion as above. The 10-mode glycine example the 4-mode representation of the potential was used,<sup>18</sup> as many applications have indicated that this level produces eigenvalues that converged to with a few cm<sup>-1</sup>. For the present work the accuracy of the representation is not relevant. However, the 10-mode glycine example is insightful. A general potential matrix is given by

$$\langle n'_1, n'_2, ..., n'_{10} | V | n_1, n_2, ..., n_{10} \rangle,$$
 (3)

Using a 4-mode representation of V, it should be clear that many of these elements are exactly zero. And indeed MULTIMODE takes note of these and sets them to zero. Referring to the *n*-mode representation, it's clear that only by going to  $V_{ijklmn...}^{(10)}$  would these analytical zeros disappear. If the *n*-mode representation is not used then the potential for 24-mode glycine (the actual number of vibrational modes) would be expressed as usual by  $V(Q_1, ..., Q_{24})$ and the idea of doing general matrix would be truly infeasible and nonsensical. Clearly, at a minimum the excitation space would have to be severely limited. However, even with the evaluating the matrix elements by even the most efficient quadrature methods would be prohibitive in 24 dimensions.<sup>22</sup> This fact notwithstanding, it does appear that the ML approach here will be powerful means to avoid calculating very small matrix elements. And, based on the empirical evidence given here and the discussion above, the number of elements increases rapidly with the number of degrees of freedom. Thus, the ML method proposed will become even more "needed" and effective as a means to deal with this rapid increase.

We note that non-direct product bases are also used in vibrational dynamics. These generally come from partitioning the full Hamiltonian into two sub-Hamiltonian in nonoverlaping modes plus an interaction part. Diagonalization of the sub-Hamiltonians yields a new basis in the corresponding modes, which can then be used in new direct-product representation. Matrix elements in this new basis are then done and a final H-matrix is obtained. The goal here is to reduce the order of this matrix, however, a reasonable expectation is that there also many small elements in this matrix. Major examples of this approach are in refs. 23–25 It does appear that the machine learning classification approach described here for a direct-product basis should be directly applicable to non-direct product bases. That is work for the future.

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