MODELING NUCLEAR PROPERTIES
WITH SUPPORT VECTOR MACHINES

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1. INTRODUCTION

Artificial neural networks and other machine-learning strategies can provide a valuable complement to theory-driven models of the systematics of nuclear data. A significant effort to exploit the potential of data-driven methodologies receives strong motivation from the current thrust toward experimental and theoretical exploration of nuclei far from stability. It is made possible by the availability of a growing body of excellent experimental data on nuclear species numbering in the thousands. In outline, statistical models based on supervised learning are developed as follows. Suppose, for example, we wish to predict the atomic mass $M$ of a nuclear species, or nuclide, specifying only its mass number $A$ and atomic number $Z$, or alternatively its proton and neutron numbers $(Z, N)$. A learning machine has an input interface where $(Z, N)$ are fed to the device in coded form and an output interface where an estimate of the mass appears for decoding. In between there is a system or network of interconnected elements that acts to process the incoming information and produce an appropriate output. These processing elements may resemble biological neurons, receiving signals from other units through weighted connections, and displaying nonlinear response to summed input signals. Given a body of training data to be used as examples of the desired mapping, in this case $(Z, N) \rightarrow M$, a suitable
learning algorithm is used to adjust the parameters of the network, e.g., the weights of the connections between the processing elements, so that the learning machine (i) generates responses at the output interface that reproduce, or closely fit, the atomic masses of the training nuclei, and (ii) serves as a reliable predictor of the masses of test nuclei absent from the training set. This second requirement is a strong one – the system should not merely serve as a lookup table for masses of known nuclei; it should also perform well in the much more difficult task of generalization.

The last two decades have seen much activity and considerable progress in the development and application of supervised learning machines of the type described – which are designed to learn by example. The most popular implementation is the multilayer feedforward neural network (or multilayer perceptron), taught by the backpropagation learning algorithm in one or another of its many variations [1-3]. A significant measure of success has been achieved in constructing global models of nuclear properties based on such neural networks, with applications to atomic masses, neutron separation energies, spins and parities of nuclear ground states, stability versus instability, branching ratios for different decay modes, and beta-decay lifetimes. (For reviews, see Ref. [4], and for recent results on atomic-mass prediction, see Ref. [5].)

The support vector machine (SVM) [6-8], a principled and powerful approach to problems in classification and nonlinear regression, came on the scene in the 1990s. It has become a standard tool in statistical modeling, and for many problems it is considered the method of choice. We have begun to explore the promise of SVMs for modeling and prediction of nuclear properties. The first results of this effort are reported here.

Section 2 provides an introduction to support vector machines and the ANOVA decomposition that facilitates their effective implementation. Section 3 summarizes the results obtained for the atomic-mass problem, and compares the predictive performance of the SVM models with that of multilayer backpropagation networks and state-of-the-art “theory-thick” models. Additional results and comparisons for beta-decay half-lives and for ground-state spins and parities are presented in Secs. 4 and 5, respectively. Concluding remarks are made in Sec. 6.

2. SUPPORT VECTOR MACHINE AND ANOVA DECOMPOSITION

The support vector machine (SVM), pioneered by Vapnik [6-8], may be viewed as an approximate realization of the goal of structural risk minimization [9,3]. Let \((x_1, y_1), ..., (x_P, y_P)\) be a set of training data drawn from a function \(y = f(x)\). Here, \(x\) is the input variable, a vector of dimension \(n\), while \(y\) is the output variable, a unique real number for given \(x\). (In the example considered in Sec. 1, \(x\) is a vector formed from the two components \(Z\) and \(N\), while \(y\) is the mass \(M\).) The support vector machine is based on a suitable nonlinear mapping \(x \rightarrow \varphi(x)\) from the input space to a feature space of higher dimension \(m > n\).

Applied to the task of regression, the SVM learning strategy begins by posing an approximation \(\hat{y}\) to the output \(y\) as a linear combination of certain basis functions \(\varphi_i(x)\) in the feature space, with corresponding linear weights connecting the feature
space to the output space. Thus,

\[ \hat{y} = \hat{f}(x, w) = \sum_{j=1}^{m} w_j \varphi_j(x), \quad (1) \]

where \( w \) is an \( m \)-dimensional vector composed of weights \( w_j, j = 1, \ldots, m \). (A bias term \( b \) may be included in Eq. (1) by starting the sum at \( j = 0 \) and introducing \( w_0 \equiv b \) and \( \varphi_0(x) \equiv 1 \).) To determine the image vectors \( \varphi_j(x) \) and their weights \( w_j \), consider an \( \epsilon \)-insensitive loss function defined, for input \( x \), by \( y - \hat{f}(x, w) - \epsilon \) in case the magnitude of the error \( y - \hat{f} \) exceeds a tolerance \( \epsilon \), and taken zero otherwise. The tolerance parameter \( \epsilon \) is at the disposal of the machine’s user. The primal optimization problem then becomes one of minimizing the overall loss (or cost function, or empirical risk), as given by the sum of the individual losses for all the training patterns,

\[ E_\epsilon(w) = \sum_{i=1}^{P} |y_i - \hat{f}(x_i, w)|_\epsilon, \quad (2) \]

subject to the inequality \( \sum_{j=1}^{m} w_j^2 < c_0 \), where \( c_0 \) is a user-determined constant.

Vapnik has shown that an equivalent solution of this constrained optimization problem can be obtained by solving the corresponding dual problem, which may be stated as follows [3].

1. Choose a kernel of the form

\[ K(x, x_i) = \sum_{j=1}^{m} \varphi_j(x) \varphi_j(x_i), \quad (3) \]

symmetrical in its vector arguments and continuous in their components, and qualifying as an inner product in some space, so as to meet the conditions of Mercer’s theorem [10,3].

2. Given the training sample \( \{(x_i, y_i)\}, i = 1, \ldots, P \), assemble the convex functional

\[ Q(\{\alpha_i, \alpha'_i\}) = \sum_{i=1}^{P} y_i (\alpha_i - \alpha'_i) - \epsilon \sum_{i=1}^{P} (\alpha_i + \alpha'_i) - \frac{1}{2} \sum_{i=1}^{P} \sum_{l=1}^{P} (\alpha_i - \alpha'_i)(\alpha_l - \alpha'_l)K(x_i, x_l). \]

\[ \quad (4) \]

3. Maximize \( Q \) subject to the constraints

\[ \sum_{i=1}^{P} (\alpha_i - \alpha'_i) = 0, \quad 0 \leq \alpha_i, \alpha'_i \leq C, \quad (5) \]

where \( C \) is a user-determined constant. The optimal approximating function then takes the forms

\[ \hat{f}_{\text{opt}}(x, w) = w^T w = \sum_{i=1}^{P} (\alpha_i - \alpha'_i)K(x, x_i), \quad (6) \]
where $\mathbf{w}^T$ the transform of the column vector $\mathbf{w}$. The subset of training patterns $i$ for which $\alpha_i - \alpha'_i$ does not vanish then defines the support vectors of the machine, corresponding to the training examples that are the most salient to solution of the problem.

The parameters $\epsilon$ and $C$ provide the user with control over the complexity of the machine, as measured by the so-called VC dimension [11,3], and hence over its performance in generalization. Careful tuning of these parameters is necessary.

Different choices for the inner-product kernel $K(\mathbf{x}, \mathbf{x}_i)$ yield different versions of the support vector machine. The most popular are (i) the polynomial learning machine, corresponding to

$$K(\mathbf{x}, \mathbf{x}_i) = (\mathbf{x}^T \mathbf{x}_i + 1)^p$$

(with user-selected power $p$), (ii) the radial-basis function (RBF) network, corresponding to

$$K(\mathbf{x}, \mathbf{x}_i) = \exp \left(-\gamma ||\mathbf{x} - \mathbf{x}_i||^2 \right)$$

(with user-selected width parameter $\gamma$), and (iii) the two-layer perceptron [1-3], with

$$K(\mathbf{x}, \mathbf{x}_i) = \tanh(\beta_1 \mathbf{x}^T \mathbf{x}_i + \beta_2)$$

(freedom in setting the parameters $\beta_1$ and $\beta_2$ being restricted by Mercer’s theorem).

We are most interested in creating predictive statistical models capable of estimating a real-valued function $f(\mathbf{x})$ from given values for its independent variables comprising $\mathbf{x}$. For that reason, we have outlined the design of SVMs for solving problems of nonlinear regression. However, the support vector machine was originally introduced to solve yes/no classification problems, and applied to problems in which positive and negative cases are either separable by a hyperplane in the input space (trivial), or not (nontrivial). For problems that are not linearly separable in this sense, the input vectors are mapped nonlinearly into a higher-dimensional feature space, in which separation by a hyperplane becomes possible. The principle of structural risk minimization then dictates that an optimal hyperplane be sought in this space, such that the margin of separation between positive and negative cases is minimized. It is known [7,8] from general learning theory that the error rate of a learning machine on test data (i.e., in generalization or prediction) is bounded by the sum of two terms, namely the error rate on the training data and a term involving the VC dimension. For a linearly separable problem treated by a SVM, the first term is zero and the second is minimized. Thus, good generalization is achieved even without building into the model any explicit knowledge about the problem to be solved, beyond the raw training data. This desirable feature is maintained approximately in application of SVMs to nonseparable classification problems and to the generically more difficult problems of regression.

The support vector machine may be broadly viewed as a kind of feedforward neural network, in that the inner-product kernels $K(\mathbf{x}, \mathbf{x}_i)$ provide a layer of hidden units that effect nonlinear processing of the inputs and provide weighted linear outputs, which are summed by an output unit. As seen above, the familiar structures of radial-basis-function networks and perceptrons with one hidden layer can be realized as special cases by suitable choices of kernel, as specified above. But a support vector
machine does more: it also embodies an algorithm that automatically determines the number of hidden units appropriate to the problem at hand, whatever the choice of kernel. This more general scope of the SVM approach stands in contrast to the backpropagation learning algorithm [1-3], which is designed especially for training multilayer perceptrons.

In addition to the benefits already mentioned, the support vector machine offers other significant advantages over the more traditional approaches to supervised learning based on neural networks, which involve dependence on trial and error, rules of thumb, and heuristics. The support vector machine offers a generic way to control model complexity. The curse of dimensionality is overcome by the pivotal strategy of introducing an inner-product kernel conforming to Mercer’s theorem and solving the constrained optimization problem in its dual version, thereby determining the dimension of the feature space as the number of support vectors distilled from the training set. The procedure naturally incorporates regularization. The use of the $\epsilon$-insensitive cost function (2) in the regression application lends robustness to the machine by avoiding certain drawbacks of the least-square estimator employed in the backpropagation learning algorithm (e.g., sensitivity to outliers and to distributions with additive noise having a long tail). Importantly, the SVM is guaranteed to find a global minimum of the error surface. For a more detailed and systematic development of the properties of SVMs, the reader is directed to Haykin's excellent text [3], as well as the authoritative monographs of Vapnik [7,8].

Our investigations of the potential of support vector machines for the design of global statistical models of nuclear properties make use of the RBF kernel (8), as well as a simplified version of what is called ANOVA decomposition [12]. ANalysis Of VAriance (ANOVA) is a scheme for imposing a structure on multi-dimensional kernels that are generated from one-dimensional kernels, in a way that gives better control over the capacity of the machine (as measured by the VC dimension). An ANOVA kernel we have found to be well suited to the regression problem posed by the nuclear (atomic) mass data is rooted in the RBF kernel and has the form

$$K(x, x_i) = \left( \sum_{l=1}^{n} \exp \left[ -\gamma \left( x^{(l)} - x_i^{(l)} \right)^2 \right] \right)^d,$$

where the user-selected parameter $\gamma$ can take any positive value and the power $d$ is usually an integer. We shall call this the ANOVA kernel.

### 3. SVM MODELS OF NUCLEAR MASS SYSTEMATICS

SVM regression models have been trained to predict $(\Delta M)c^2$ in MeV, where $\Delta M$ is the mass excess (or mass defect) defined by the difference $M - A$ between the atomic mass $M$, measured in amu, and the mass number $A$ of the nuclide in question. In our initial study, we focus on a database given by the union $O \oplus N \oplus NB$ of three data sets. The first consists of the set of 1323 “old” (O) experimental mass assignments which the 1981 semi-empirical droplet-model mass formula of Möller and Nix [13] was intended to reproduce. The second is a set of 351 “new” (N) experimental mass assignments for nuclei that lie mostly beyond the edges of the 1981 data (as viewed
in the $N-Z$ plane). In addition to the O and N sets, a set of 158 nuclides with more recently measured masses (the NB set of “even newer” nuclides) is employed in the modeling process. In earlier work [14-16,5], these three data sets have been used to quantify the extrapolation capability (the so-called extrapability) of different global mass models (based either on nuclear theory or neural networks).

The set $O \oplus N \oplus NB$ is divided by a random-sampling procedure into three nonoverlapping subsets, namely a training set (80%), a validation set (10%), and a test set (10%), in the indicated approximate proportions. (In all work reported in this paper, random samplings are drawn from a uniform distribution.) Training, validation, and test sets are each further subdivided into four subsets labeled EE, EO, OE, and OO, composed respectively of nuclides belonging to the four “even-oddness” classes: even-$Z$-even-$N$, even-$Z$-odd-$N$, odd-$Z$-even-$N$, and odd-$Z$-odd-$N$. For convenience, values of the input variables are encoded by a linear transformation that scales and shifts given values of $Z$ and $N$ to lie in the interval [0, 1]. A similar linear transformation decodes the learning machine’s raw output, which lies in the interval $[-1, 1]$, so as to provide an estimate of the corresponding mass excess in MeV.

Effectively, we divide the mass problem into four separate problems, one for each of the four “even-oddness” classes in $Z$ and $N$. In doing so, we are actually incorporating some domain knowledge into the learning strategy. Distinctive quantum-mechanical features of nuclei, abundantly supported by empirical evidence, include quantized angular momenta, magic numbers, shell structure, and pairing energies, all of which stem from the fact that $Z$ and $N$ are integers, even or odd.

A SVM model is developed individually for each of the four nuclear classes EE, EO, OE, and OO. SVM regression (with ANOVA-RBF specification of kernels) is carried out separately for the respective training sets, thereby constructing a predictive model whose reliability is judged by its performance on the examples in the test set. Following established practice, performance of each of the four models on its corresponding validation set have been used to guide the final determination of the adjustable parameters. Ideally, the test set should have no role in choosing these parameters (although in some cases a weak influence is allowed).

As is usual in global models of the atomic-mass table, the quality of a given model is judged by the smallness of the root-mean-square (rms) error $\sigma$ in the mass excess $\Delta M$, averaged over the data set in question (training, validation, or test set for a given class of nuclides). To be competitive, a model should have values of $\sigma$ below 1 MeV. It should be noted however, that only in a few cases has a rigorous test of predictive performance been made for the traditional theoretical models of semi-empirical character. (An important exception is found in the work of Möller, Nix, and collaborators [15,16], who introduce the notion of extrapability, which is equivalent to our generalization.)

Some of the better results obtained in the present exploratory study are displayed in Table 1. The performance of these models, all with RBF parameter $\gamma = 2.5$ and ANOVA degree $d = 8$, is evidently of high quality.

Similar learning experiments can be found among the studies of Ref. [5] based on multilayer perceptrons and modified backpropagation training, although procedural differences preclude direct comparisons of performance. The best model obtained
Table 1

Performance of SVM global models of atomic mass. For all four models, the RBF parameter $\gamma$ is 2.5 and the ANOVA degree is $d = 8$. The other SVM parameters have been defaulted at $C = 0.1$ and $\varepsilon = 0.001$.

| Classes | Learning Set | Validation Set | Test Set |
|---------|---------------|----------------|----------|
|         | # Nuclides $\sigma$(MeV) | # Nuclides $\sigma$(MeV) | # Nuclides $\sigma$(MeV) |
| EE      | 381 0.58     | 48 0.71        | 48 0.99  |
| EO      | 360 0.89     | 45 0.68        | 45 0.62  |
| OE      | 371 0.70     | 46 0.78        | 46 0.88  |
| OO      | 353 0.75     | 44 0.74        | 45 0.97  |

using O as the training set, NB as validation set, and N as test set gave rms error figures on these sets of 0.71 MeV, 2.28 MeV, and 2.16 MeV, respectively. Another strategy yielded better results. The set $O \oplus N$ was first “purified” by removing 20 nuclides with poorly measured masses. A random sample M1 consisting of 1303 of the remaining 1654 examples (some 79%) was used as the training set. The complementary set, M2, played the role of validation set, and the NB set was used for testing the trained model. The best model found in this way produced rms errors on the three sets of 0.44 MeV (M1), 0.44 MeV (M2), and 0.95 MeV (NB). It should be noted that this level of performance on the mass problem was achieved after more than a decade of successive improvements in the choices of architectures, coding schemes, and training algorithms.

In addition to the four class-specific models SVM-EE, SVM-EO, SVM-OE, and SVM-OO reported on in Table 1, we also constructed a single SVM model (denoted SVM-S) using the full O data set as the training sample, without making a distinction between EE, EO, OE, and OO nuclides. In this case, the NB nuclei are used as a validation set, guiding the determination of the RBF and ANOVA parameters. The parameters associated with the SVM-S model are again $\gamma = 2.5$ and $d = 8$, along with $C = 0.1$ and $\varepsilon = 0.001$. This model yields rms errors of 0.70 MeV on the training set O and 0.75 MeV on the validation set NB, with a $\sigma$ value of 1.41 MeV on the N nuclei, regarded as a test set. (These results are erroneously cited in Ref. [5].) A proper averaging over the four nuclidic classes permits a comparison between the SVM-S model and the four models represented in Table 1. The composite performance of the latter models is then reflected in $\sigma$ values of 0.73 MeV, 0.73 MeV, and 0.88 MeV in training, validation, and testing, respectively.

In some cases, meaningful comparisons may be drawn between the performance of statistical mass models based on multilayer perceptrons and support vector machines, and the traditional mass models based on nuclear theory and phenomenology. Starting with the simple liquid-drop model, such traditional theory-thick models have
evolved over seven decades to achieve a high degree of sophistication and precision. For example, the 1992 FRDM model of Möller and Nix [15] gives $\sigma$ values of 0.67 MeV on the O set (when fitted to this set) and 0.74 MeV on the N set (a true measure of predictive performance of the model). The more enhanced FRDM model of Ref. [16], which is fitted to the data set $M_1 \oplus M_2$, yields rms errors of 0.68 MeV ($M_1$), 0.71 MeV ($M_2$), and 0.70 MeV (NB). The HFB2 model of Pearson and collaborators [17] gives respective errors of 0.67 MeV, 0.68 MeV, and 0.73 MeV. (We note that the result of Ref. [17] on the “test set” NB cannot be regarded as a prediction, since the nuclei involved were used in adjusting model parameters.)

With additional refinements, it is not unreasonable to expect that SVM models can equal (and possibly surpass) the levels of robustness and predictive accuracy achieved with theory-thick models and with multilayer perceptron models. However, a conclusive statement must await a thorough SVM study based on the recent AME03 mass evaluation carried out by Audi et al. [18]

### 4. SVM MODELS OF BETA-DECAY HALFLIVES

We now turn to a second problem of regression in the statistical analysis of nuclear properties via support vector machines, namely fitting and prediction of the beta-decay halflives of nuclides ($Z, N$) that decay 100% via the $\beta^-$ mode. The data for this problem have been culled from the on-line repository at the Brookhaven National Nuclear Data Center (http://www.nndc.bnl.gov). The data employed are current to May 2005 and consist of a total of 932 examples. Restricting attention to examples with halflives below $10^6$ s leaves 633 nuclides. When measured in seconds, the experimental values of $T_{1/2}$ range over 26 orders of magnitude, so it is more appropriate to regress $L = \log T_{1/2}$ instead of the halflife itself, and to adopt the rms error $\sigma_L$ of the estimate of $L$ as a figure of merit in learning, validation, and prediction phases of the analysis.

As in the case of the mass problem, separate SVM models are constructed for EE, EO, OE, and OO classes of nuclides. However, we make the simpler RBF choice of kernel, instead of pursuing the more elaborate ANOVA option. (Implementation based on the ANOVA decomposition is much more demanding in terms of computer time.) Each of the four data subsets (EE, EO, OE, OO) is subdivided into training, validation, and test sets in the approximate proportions 80%, 10%, and 10%, respectively.

The results obtained from the SVM regressions are summarized in Tables 2 and 3. Table 2 gives the parameters and performance measures of the models constructed for the full set of data, regardless of measured lifetime. Table 3 displays the corresponding results when nuclides with $T_{1/2} \geq 10^6$ s are removed from the database.

A similar study [19] (see also Ref. [20]) has been carried out with multilayer feedforward neural networks trained by “vanilla” backpropagation, for data available in 1995 (766 examples in total) However, this study did not employ the now-standard protocol in which a validation set is used in making the final model selection. Also, no subdivision into the four even-oddness classes was made. Instead, the full data set (or the restricted set of examples with $T_{1/2} < 10^6$ s) was split into a training set of approximately 75% of the examples and a test set consisting of the remainder.
Table 2

Performance of SVM global models of $\beta$-decay halflives $T_{1/2}$ (including examples having $T > 10^6$ s). For all four models, $C = 1$ and $\varepsilon = 0.001$.

| Classes | Learning Set | Validation Set | Test Set | RBF kernel |
|---------|--------------|----------------|----------|------------|
|         | # Nuclides   | $\sigma_L$     | # Nuclides | $\sigma_L$ | # Nuclides | $\sigma_L$ | $\gamma$ |
| EE      | 137          | 2.88           | 16       | 3.61       | 15         | 1.72       | 5.44     |
| EO      | 198          | 2.75           | 24       | 2.27       | 22         | 2.17       | 7.27     |
| OE      | 187          | 2.37           | 22       | 2.76       | 20         | 2.38       | 9.99     |
| OO      | 236          | 2.62           | 29       | 2.07       | 26         | 2.96       | 9.55     |

Table 3

Performance of SVM global models of $\beta$-decay halflives (with a cutoff at $10^6$ s). For all four models, $C = 1$, $\varepsilon = 0.001$.

| Classes | Learning Set | Validation Set | Test Set | RBF kernel |
|---------|--------------|----------------|----------|------------|
|         | # Nuclides   | $\sigma_L$     | # Nuclides | $\sigma_L$ | # Nuclides | $\sigma_L$ | $\gamma$ |
| EE      | 96           | 1.34           | 11       | 0.52       | 10         | 1.20       | 1.78     |
| EO      | 140          | 0.90           | 17       | 0.69       | 15         | 1.22       | 9.97     |
| OE      | 122          | 1.55           | 14       | 0.63       | 13         | 1.18       | 0.84     |
| OO      | 159          | 1.00           | 19       | 1.28       | 17         | 1.34       | 8.87     |

Comparison of the rms errors shown in Tables 2 and 3 with the corresponding performance figures from the earlier work [19,20] shows an improvement (reduction) in rms error values by about a factor 2, in both learning and prediction, for both the full and restricted data sets. Comparison may also be made with results from traditional nuclear theory (e.g. Refs. [21-23]). Since the cited neural-network models could already attain performance in fitting and prediction comparable to that exhibited by these theory-thick models, we can say with some confidence that the SVM models are capable of a predictive acuity superior to the best of the traditional global models currently in play.
We should also call attention to the greatly improved quality of neural-network models of $\beta$-decay systematics, achieved in very recent studies [24]. Data based on the AME03 evaluation are divided into training, validation, and test sets in the respective proportions 60%, 20%, and 20%, both with and without the restriction to half-lives not greater than $10^6$ s, but without subdivision into even-oddness classes. In the case where the restriction is imposed, the best results found for the error measure $\sigma_L$ are 0.55 (training), 0.61 (validation), and 0.64 (prediction). The corresponding averages for the model represented in Table 3 are 1.43, 0.89, and 1.24, respectively, so further refinement of the SVM models will be needed to match the performance of the best multilayer perceptrons.

5. SVM MODELS OF GROUND-STATE SPINS AND PARITIES

In a third illustration of what is possible, the SVM approach is applied to construct global statistical models of the ground-state spins and parities of nuclei. (In this context, “spin” refers to the total angular momentum quantum number $J$ of the nuclear state.) As in the exercises described in Secs. 3 and 4, we again divide the nuclei under consideration into EE, EO, OE, and OO classes. In the spin problem, this subdivision is of obvious importance, since the law of angular momentum addition in quantum mechanics dictates that the states of EE and OO nuclei can only have integral values of $J$, whereas the spins of EO and OE nuclei must be half-odd-integral. In fact, all EE nuclei are known to have spin/parity $J^\pi = 0^+$. Clearly, we may exclude this class from consideration, since its modeling is a trivial task for any viable learning machine.

The parity property of nuclear states presents the simplest kind of classification problem, with two mutually exclusive outcomes, even or odd. Moreover, because the spin quantum number $J$ is restricted by quantum theory to a finite set of discrete values, global modeling of spin systematics is also most efficiently treated, within the SVM framework, as a problem of classification rather than function approximation or regression. In our study, we consider $J$ values ranging from 0 to $23/2$ in half-odd-integral steps, the integral values being available for OO nuclei and the half-odd integral values, for EO and OE nuclei. This specification of the problem may be construed as introducing some basic domain knowledge into the model-building process.

Data for the spin and parity nuclear ground states have been taken from the on-line Brookhaven database. Based on simple RBF kernels, separate SVM classifier models of these two properties have been developed for each of the three nontrivial even-oddness cases.

Let us first discuss our findings for the parity problem. In treating this problem, the data for each of the cases EO, OE, and OO are divided at random into training, validation, and test sets in the approximate proportions 80%, 10%, and 10%, respectively. Performance is measured in terms of the percentages of correct classifications within these subsets. The primary results are summarized in Table 4. It is apparent that modeling parity is an easy task for SVMs. Judging from available results [25,14], it is also relatively easy for neural networks (although SVM performance is somewhat superior).
Table 4

Performance of SVM global models of ground-state parity. For all four models, $C = 0.1, \varepsilon = 0.01$. Model selection is guided by best performance on the validation set, consistent with a perfect score on the training set.

| Classes | Learning Set | Validation Set | Test Set | RBF kernel |
|---------|---------------|----------------|----------|------------|
|         | # Nuclides    | # Nuclides | # Nuclides | # Nuclides | Score | Score | Score | Score | \( \gamma \) |
| EO      | 474 100%      | 58 93%     | 52 83%    | 9.232      |
| OE      | 466 100%      | 57 89%     | 51 90%    | 9.482      |
| OO      | 434 100%      | 53 87%     | 48 84%    | 9.176      |

Table 5

Performance of SVM global models of ground-state parity. For all four models, $C = 0.1, \varepsilon = 0.01$. In this case, model selection is guided by best performance on the validation set, allowing for minimal nonzero error rate on the training set.

| Classes | Learning Set | Validation Set | Test Set | RBF kernel |
|---------|---------------|----------------|----------|------------|
|         | # Nuclides    | # Nuclides | # Nuclides | # Nuclides | Score | Score | Score | Score | \( \gamma \) |
| EO      | 474 100%      | 58 91%     | 52 83%    | 0.678      |
| OE      | 466 95%       | 57 84%     | 51 92%    | 0.180      |
| OO      | 434 96%       | 53 83%     | 48 86%    | 0.240      |

For the models of Table 4, performance on the training sets is perfect. If we are willing to make a small sacrifice in the quality of reproduction of the input data, slightly better performance on the validation and test sets can be achieved, as seen in Table 5. It is interesting that this second model corresponds to a quite different error minimum under variation of the parameter \( \gamma \). In general, there may be many such minima of similar depth.

We have not yet conducted a full training-validation-test process for the spin problem. Accordingly, we present only preliminary results, which nevertheless are illuminating. In the first experiment to be reported (see Table 6), each of the three spin data sets EE, OO, and OO is divided randomly into two subsets, a training set and a complementary second set. The training set contains approximately 90% of the examples of the given even-oddness class, and the second set, the remaining \( \sim 10\% \).
Table 6

Performance of SVM global models of nuclear ground-state spin. For all three models, $C = 0.1$, $\varepsilon = 0.01$. Model selection is guided by best on performance on the validation set, consistent with a perfect score on the training set.

| Classes | Learning Set | Validation/Test Set | RBF kernel |
|---------|--------------|---------------------|------------|
|         | # Nuclides | Score | # Nuclides | Score | \(\gamma\) |
| EO      | 528        | 100%   | 58         | 81%    | 9.217 |
| OE      | 522        | 100%   | 57         | 68%    | 9.001 |
| OO      | 488        | 100%   | 54         | 43%    | 4.002 |

Table 7

Performance of SVM global models of nuclear ground-state spin. For all three models, $C = 0.1$, $\varepsilon = 0.01$. The parameter \(\gamma\) is fixed at the value determined for Table 6. The test set influences model choice only indirectly.

| Classes | Learning Set | Validation Set | Test Set | RBF kernel |
|---------|--------------|----------------|----------|------------|
|         | # Nuclides | Score | # Nuclides | Score | # Nuclides | Score | \(\gamma\) |
| EO      | 476        | 100%   | 58        | 79%    | 52         | 60%    | 9.217 |
| OE      | 470        | 100%   | 57        | 61%    | 52         | 79%    | 9.001 |
| OO      | 440        | 100%   | 54        | 39%    | 48         | 38%    | 4.002 |

The second set is used to help pin down the RBF parameter \(\gamma\) and thereby plays a role in model selection. Hence it must be interpreted as a validation set. SVM models are constructed for a range of \(\gamma\) values, and the model whose \(\gamma\) value produces the lowest error on the second data set (while scoring 100% on the training set) is selected. There is no real test set in this experiment.

In an alternative experiment, we have implemented a protocol intermediate between the training-validation scheme leading to Table 6, and the full training-validation-test procedure. The data for each of the three even-oddness classes involved are divided into three subsets as follows. The second subset is taken to be identical to the second subset formed in the first experiment. The first subset, used as the training set, consists of 80% of the examples for the class in question, these being chosen at random from the corresponding training set created in the first experiment. The 10% that are not so chosen constitute the third subset, which is regarded as a
test set. Then, using the same parameter $\gamma$ as determined in the first experiment with the aid of the second subset, new SVM models are developed from the examples in the reduced training set. These models are used to generate spin values for both second and third subsets – values which may differ from those given by the models developed in the first experiment (see Table 7). Although it is not legitimate to interpret the third subset as a test set in the purest sense, its influence on model selection is indirect.

From the results shown in Tables 6 and 7, one may plausibly infer that support vector machines can perform very well on the problem of predicting nuclear ground-state spins. While further experiments are needed to affirm this conclusion, it is already of interest to compare our SVM models with other global models of nuclear spin systematics. Global nuclear structure calculations within the macroscopic/microscopic approach [26] reproduce the ground-state spins of odd-$A$ nuclei with an accuracy of 60% (agreement being found in 428 examples out of 713). (In this work, there is no clear distinction between fitting and prediction, or between training, validation, and test sets.) Multilayer feedforward neural networks do somewhat better [25,14]. Averaging over results of three experiments involving nets having a single hidden layer and trained with backpropagation, the performance for odd-$A$ nuclei reaches 62% on what are effectively validation sets, the training sets being reproduced to an accuracy of 93%. In an experiment in which the connection weights of feedforward nets with one hidden layer are determined by a conjugate gradient procedure, performance at the level of 99.5% on the training set and 73.2% on a validation set has been achieved for OE nuclei. The spins of odd-odd nuclei are notoriously difficult to predict. This is reflected in the performance figures of neural-network (perceptron) models on the OO category, which are typically 75% correct on training-set examples and only 15% in validation or testing.

Placed in the context of earlier work, both statistical and phenomenological, the results in Tables 6–7 for the first SVM models of nuclear spin speak for themselves.

6. CONCLUDING REMARKS

We have made initial studies of the potential of support vector machines (SVM) for providing statistical models of nuclear systematics with demonstrable predictive power. Using SVM regression and classification procedures, we have created global models of atomic masses, beta-decay half-lives, and ground-state spins and parities. These models exhibit performance in both data-fitting and prediction that is comparable to that of the best global models from nuclear phenomenology and microscopic theory, as well as the best statistical models based on multilayer feedforward neural networks. Further work to develop the scope, acuity, and reliability of SVM applications to nuclear physics seems to be warranted. In particular, the full body of data in the AME03 atomic-mass evaluation [18] must be brought to bear in construction of SVM models of mass systematics, and the treatment of the spin problem begun here needs to be completed. Fruitful applications to nucleon separation energies, $\alpha$-decay half-lives, branching ratios of nuclear decay, nuclear deformations, neutron cross sections, and other nuclear properties may also be on the horizon.
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