

# 118. Protein Fold Class Prediction Using Neural Networks Reconsidered

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## 1 Problem description

Predicting the three-dimensional structure of a protein from its amino acid sequence is an important and challenging task in bioinformatics. In the following, we describe an application of feed-forward neural networks (NNs) for the protein fold classification task, introduce a tailored regularization method for sparse data, and compare the results to support vector machines (SVMs [9]). We consider the problem of assigning a primary sequence to one of 42 fold classes. The data are taken from the *database for expected fold-classes* (DEF [7]). Each sequence is represented by its dipeptide-frequency without any additional associated physicochemical properties, i.e., the input dimension of the problem is 400. The data set is split into a set  $D_{\text{APE}}$  for training and a disjoint test set  $D_{\text{TPE}}$  as in [4, 6], consisting of 143 and 125 patterns, respectively.

## 2 Method

A standard NN was employed for classification [2]. We did not reduce the input dimension (as done for the NNs in [4] by PCA), used 1-of-42 output encoding, and set *ad hoc* the number of hidden neurons to 10. This means, the network has by far more weights than training patterns are available. However, the generalization of NNs is surprisingly insensitive to excess capacity, see [1] and [3] for theoretical and empirical studies. *Early stopping* in conjunction with a suitable learning algorithm starting from small initial weights can “be used to stop training large nets when they have learned models similar to those learned by smaller nets of optimal size” [1]. Here, an improved version of the Rprop learning algorithm was used for training [5, 8]. We split  $D_{\text{APE}}$  randomly but ensuring an as equal distribution of the classes as possible into  $D_{\text{train}}$  and  $D_{\text{validate}}$  with 97 and 50 patterns, respectively. However, as we have only 147 training patterns for discriminating 42 classes, it is problematic to restrict the complete training process to a subset of  $D_{\text{APE}}$ , i.e., early stopping can not be applied in the standard way. Thus, we used a two-stage training process: First, the network is trained on  $D_{\text{APE}}$  until the classification error on  $D_{\text{APE}}$  vanished. Then training continued on  $D_{\text{train}}$  until generalization loss is observed on  $D_{\text{validate}}$ . The weight configuration yielding the smallest mean-squared-error on  $D_{\text{validate}}$  is taken as the final solution of the training process.

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### 3 Results

In [4], 11 different classification methods were applied to the described data sets including different discriminant analysis methods, nearest neighbor classification, projection pursuit regression, and NNs. SVMs with different kernels were used in [6] for comparison and gave the best results. We trained our network starting from 10 small random weight initializations for 100 iterations. The apparent prediction error (APE, the classification error on  $D_{\text{APE}}$ ) was 0 in all trials. The average classification error on the test set (TPE, test prediction error) was 22.96 % and the variance  $\sigma^2 = 7.18 \cdot 10^{-5}$ . Most trials including the one with the smallest mean-squared error on training and validation set (i.e., the one that would be selected) resulted in a network with a TPE of 22.4 %. In our (preliminary) experiments,

	SVM RBF	SVM POLY1	SVM POLY2	SVM POLY3	NN
APE	0	0	4.2	1.4	0
TPE	23.2	28.8	32	32.8	<b>22.4</b>

Table 1: Results using the neural network (NN) compared to SVMs with different kernels [6], a radial basis kernel (RBF) and polynomial kernels of degrees 1, 2, and 3 (POLY1, POLY2, POLY3).

NNs combined with an appropriate regularization scheme showed better classification results than reported for SVMs and other classification methods [4, 6], see Tab. 1. Note that it is sufficient to train a single NN for a few generations, but that 42 SVMs are needed for the *one-against-all* multiple-class discrimination approach in [6]. The proposed method to deal with sparse data is not limited to protein fold classification, but may be applicable to a wide range of bioinformatics tasks.

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