

PROTEIN SUBCELLULAR LOCALIZATION PREDICTION WITH WOLF PSORT

PAUL HORTON^A, KEUN-JOON PARK^{A,C}, TAKESHI OBAYASHI^{B,D,E}, AND KENTA NAKAI^C

^a*Computational Biology Research Center, AIST, Tokyo, Japan*
E-mail: horton-p@aist.go.jp

^c*Human Genome Center, Institute of Medical Science, University of Tokyo*
E-mail: park-kj@hgc.jp, knakai@ims.u-tokyo.ac.jp

^b*Graduate School of Bioscience and Biotechnology, Tokyo Institute of Technology*
E-mail: toobayas@bio.titech.ac.jp

^d*Graduate School of Pharmaceutical Sciences, Chiba University*

^e*Core Research for Evolutional Science and Technology*

We present a new program for predicting protein subcellular localization from amino acid sequence. WoLF PSORT is a major update to the PSORTII program, based on new sequence data and incorporating new features with a feature selection procedure. Following SWISS-PROT, we divided eukaryotes into three groups: fungi, plant, and animal. For the 2113 fungi proteins divided into 14 categories; we found that, combined with BLAST, WoLF PSORT yields a cross-validated accuracy of 83%, eliminating about 1/3 of the errors made when using BLAST alone. For 12771 animal proteins a combined accuracy of 95.6% is obtained, eliminating 1/4 of BLAST alone errors, and for 2333 plant proteins the accuracy can be improved to 86% from 84%.

1. Introduction

Protein localization is a central issue in understanding cells. More than 20 papers⁶ have been published in major international journals describing programs for predicting localization from amino acid sequence in eukaryotic cells. Some of these works such as PSORT,⁹ PSORTII,⁷ the SignalP^{11,10} family of programs and others⁴ use sorting signal information for prediction. However the majority of prediction programs developed use amino acid content in some form. These methods exploit the longstanding observation¹² that amino acid content correlates strongly with localization site.

In this paper we present WoLF PSORT, a major extension to PSORTII which combines (mostly signal based) features from PSORT and iPSORT² with amino acid content. Prediction accuracy is increased by using feature selection while retaining

the simple k nearest neighbor classifier used by PSORTII. The dataset constructed and web service are available at wolfsort.org

2. Methods

2.1. Dataset

We prepared the dataset primarily from Swiss-Prot³ Release 45.0 annotation, ignoring entries with weakening qualifiers such as “by similarity”. In addition several hundred *Arabidopsis* entries were added from the Gene Ontology¹ web site (up to the 2004/12/4 release). Entries with evidence codes {TAS, IDA, IMP} were included, with revisions by hand in a few cases.

2.1.1. Site Definition

The distribution of localization sites in our database is shown in Table 1. The sites reflect common usage found in localization labeled Swiss-Prot entries. Table 2 shows the corresponding Gene Ontology numbers for the Gene Ontology derived sequences.

Table 1. The distribution of localization sites for each category of organisms.

| localization | animal | plant | fungi | localization | animal | plant | fungi |
|------------------------|--------|-------|-------|--------------|--------------|-------------|-------------|
| <i>nuclear</i> | 2682 | 433 | 667 | cyto_nucl | 245 | 9 | 91 |
| <i>plasma membrane</i> | 3195 | 160 | 220 | cyto_mito | 18 | 3 | 8 |
| <i>extracellular</i> | 3130 | 113 | 140 | cyto_pero | 10 | 0 | 2 |
| <i>cytosol</i> | 1555 | 452 | 383 | cyts_plas | 5 | 0 | 0 |
| <i>mitochondria</i> | 938 | 200 | 389 | cyto_plas | 4 | 0 | 0 |
| <i>chloroplast</i> | N/A | 744 | N/A | cyto_golg | 4 | 0 | 0 |
| <i>E.R.</i> | 425 | 65 | 66 | E.R._mito | 18 | 0 | 4 |
| <i>peroxisome</i> | 217 | 47 | 77 | E.R._golg | 9 | 0 | 0 |
| <i>lysosome</i> | 132 | N/A | N/A | extr_plas | 19 | 0 | 0 |
| <i>golgi body</i> | 100 | 25 | 38 | mito_pero | 15 | 0 | 0 |
| <i>vacuole</i> | 16 | 72 | 23 | mito_nucl | 2 | 0 | 0 |
| <i>cytoskeletal</i> | 32 | 10 | 5 | sum | 12771 | 2333 | 2113 |

Italics indicate the abbreviated name. Localization names joining two abbreviated names with “_”, such as “cyto_nucl”, indicate dual localization.

2.2. WoLF PSORT system

WoLF is a feature selection program, (the name “WoLF” is loosely inspired by the words “Learning”, and “Weighted Features”). WoLF PSORT is the combination of WoLF with a version of PSORTII slightly extended for this purpose. The extended version outputs amino acid features and some iPSORT² features as well as the PSORT features. An overview of the system is shown in Figure 1.

Table 2. The correspondence between the localization sites used in our study and the GO numbers for the entries derived from GO annotation.

| description | GO numbers | depth | WoLF PSORT site |
|-----------------------|------------|-------|-----------------|
| cytoskeleton | GO:0005856 | 2 | cyts |
| cytosol | GO:0005829 | 0 | cyto |
| endoplasmic reticulum | GO:0005783 | 0 | E.R. |
| extracellular | GO:0005576 | 0 | extr |
| cell wall | GO:0005618 | 0 | extr |
| Golgi apparatus | GO:0005794 | 1 | golgi |
| mitochondrion | GO:0005739 | 0 | mito |
| nucleus | GO:0005634 | 0 | nucl |
| plasma membrane | GO:0005886 | 0 | plas |
| peroxisome | GO:0005777 | 2 | pero |
| vacuolar membrane | GO:0005774 | 2 | vacu |
| chloroplast | GO:0009507 | 0 | chlo |
| thylakoid lumen | GO:0009543 | 0 | chlo |

Depth indicates the number of levels of “part_of” descendents included with the GO number. For example GO:0005856 and all of its “part_of” children and grandchildren were included in cyts. Note that “extr” and “chlo” are the union of two lines from this table.

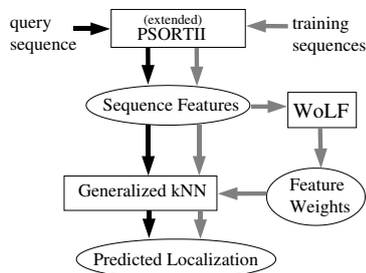


Figure 1. A schematic of the WoLF PSORT system is shown. Rectangles represent programs or procedures and ovals represent computed quantities. The black arrows denote information derived from the query sequence while gray arrows denote information from the training sequences.

2.3. Classification

2.3.1. Candidate Features

We used four kinds of features: PSORT⁹ features, iPSORT² features, amino acid content, and sequence length. Each feature is a deterministic mapping from amino acid sequence to the reals. Since the numerical range of the raw features are not homogeneous, we normalized each feature to its percentile value in the training data. Values observed in test data may not appear in the training data – in which case our programs use linear interpolation (extrapolation) to obtain a pseudo-percentile value.

2.3.2. Classification Algorithm

We adopted a weighted version of the k NN (k -Nearest Neighbors)⁵ algorithm for classification. As in standard k NN, our method classifies based on the k nearest instances in the dataset. However we slightly extended the distance calculation. In our variation two weights, w_{1i} and w_{2i} , are associated with each feature i . More formally, let F_{ji} and F_{ki} denote the values of feature i in protein instance j and k respectively. The distance $d_W(j, k)$ between j and k is defined as:

$$d_W(j, k) = \sum_i w_{1i} |F_{ji} - F_{ki}| + \sqrt{\sum_i w_{2i} (F_{ji} - F_{ki})^2} \quad (1)$$

combining elements of Euclidean and Manhattan (city block) distances.

2.3.3. Extensions for Dual Localization Prediction

The dataset contains some dually localized proteins. We gave partial credit for partially correct predictions as shown Table 3.

Table 3. Example Utility Values.

| label | prediction | utility | label | prediction | utility |
|-----------|------------|---------|-----------|------------|---------|
| nucl | nucl | 1 | nucl_cyto | nucl_mito | 0.333 |
| nucl_cyto | nucl_cyto | 1 | nucl | cyto | 0 |
| nucl_cyto | nucl | 0.5 | mito | nucl_cyto | 0 |
| nucl | nucl_cyto | 0.5 | mito | nucl | 0 |

Examples of the utilities used in this study are shown. “label” denotes the localization site according to the dataset annotation. Note that we chose to use utilities that are symmetric with regard to the label and prediction.

2.3.4. Feature Selection and Weighting

We developed a C++ program, WoLF, for selecting the weights in Equation 1. Given a set of candidate features, WoLF selects non-negative integer weights for each feature (a weight of zero is equivalent to exclusion of the feature). WoLF uses a greedy, neighborhood search algorithm to find a locally optimal set of weights. The program uses the jackknife (leave one out cross-validation) utility on the training data to evaluate weight vectors. In the case of ties the simpler weight vector is chosen.

2.3.5. Reducing Over-reliance on Sequence Similarity

When training WoLF PSORT we employed a *taboo list* which disallows the use of highly similar sequences (with identical localization sites) as a neighbor when classifying a given instance. We determined the threshold by inspection of the

correlation between best hit eValue and co-localization. The thresholds used were -33, -63, -33.4 \log_2 eValue for fungi, plant, and animal respectively.

2.3.6. *Evaluation of WoLF PSORT Accuracy*

5-fold cross-validation was used to estimate the accuracy. For each partition, feature and k value selection were performed using a jackknife test on the training partition.

3. Results

3.1. *Effect of Feature Weighting*

The results of various feature weight vectors, including those selected by WoLF PSORT are shown in Table 4. For those tables the value of k was optimized separately for the taboo and no taboo list cases. The WoLF PSORT weight vectors (one per partition) however were only trained using the taboo list. The confusion matrices for the WoLF PSORT cross-validation for yeast (which had the highest fraction of dual localization annotations) is shown in Table 6.

In addition to cross-validation studies, we also ran the WoLF feature weighting procedure on the complete datasets. The selected features (trained with the taboo list) are shown in table 5.

3.2. *WoLF PSORT Combined with BLAST*

We calculated the utility of combining WoLF PSORT with BLAST in a trivial way; namely using the WoLF PSORT prediction for queries whose best BLAST hit eValue exceeds a given threshold, while predicting the localization of the best BLAST hit otherwise. Ties for the best BLAST hit (especially with eValue=0) were fairly common, in which case we voted amongst the best hits (breaking ties using the overall proportion of each localization in the given dataset). In the rare cases in which no BLAST hit was obtained, the majority classifier was used in lieu of BLAST. The results of this hybrid predictor for the three datasets are shown in Figure 2.

3.3. *WoLF PSORT Server*

The WoLF PSORT server is freely available at wolfpsort.org. Detailed information about the features of the query sequence and its k nearest (by Equation 1) neighbors are given. These tables give the user a chance to examine the *evidence* behind the prediction. For example Figure 3 shows a partial screen shot of the detailed page (one click away from the summary page) when the protein AEP_YARLI is used as a query. From the first row in the displayed table one can see that the variable *gvh*, the signal peptide detecting weight matrix score of Gunnar von Heijne,¹³ has a very high value (93 percentile), consistent with the prediction of

Table 4. Cross-validated utility with various feature weight vectors.

| Fungi Dataset | | | | |
|--------------------|-----------|-------|------------|------------------|
| weight vector type | #weights | taboo | k | % utility |
| psortEuclid | 31 | no | 9.2(2.2) | 64.7(2.8) |
| psortEuclid | | yes | 15.6(6.0) | 61.3(3.1) |
| allEuclid | 56 | no | 4.6(3.5) | 73.9(2.2) |
| allEuclid | | yes | 26.8(11.3) | 69.3(2.9) |
| allWeights | 112 | no | 3.2(1.8) | 74.3(1.7) |
| allWeights | | yes | 31.2(18.0) | 69.4(2.7) |
| WoLF PSORT | 22.2(4.3) | no | 18.2(4.3) | 72.6(0.7) |
| WoLF PSORT | | yes | 18.2(4.3) | 70.7(1.1) |
| Plant Dataset | | | | |
| weight type | #weights | taboo | k | % utility |
| psortEuclid | 31 | no | 4.8(1.9) | 66.5(2.0) |
| psortEuclid | | yes | 41.2(24.3) | 53.6(2.0) |
| allEuclid | 56 | no | 1(0) | 85.3(1.2) |
| allEuclid | | yes | 19(3.8) | 60.0(3.8) |
| allWeights | 112 | no | 1(0) | 85.2(1.2) |
| allWeights | | yes | 20.6(3.2) | 60.0(3.7) |
| WoLF PSORT | 21.2(1.3) | no | 3(2.1) | 76.7(2.4) |
| WoLF PSORT | | yes | 13.8(7.0) | 65.1(2.6) |
| Animal Dataset | | | | |
| weight type | #weights | taboo | k | % utility |
| psortEuclid | 31 | no | 1(0) | 79.7(0.5) |
| psortEuclid | | yes | 36.0(9.3) | 72.2(0.8) |
| allEuclid | 56 | no | 1(0) | 92.3(0.5) |
| allEuclid | | yes | 34.6(6.5) | 77.8(0.6) |
| allWeights | 112 | no | 1(0) | 93.1(0.6) |
| allWeights | | yes | 30.2(7.2) | 79.0(0.7) |
| WoLF PSORT | 25.8(4.2) | no | 39.4(7.2) | 83.2(1.2) |
| WoLF PSORT | | yes | 39.4(7.2) | 79.7(1.0) |

Utility is given as percent of the maximum possible. The number of (non-zero) weights is omitted when it is the same as the row above. Numerical entries represent averages over 5-fold cross-validation with standard deviations given in parenthesis. The “psortEuclid” weight vector has weight 1 for the quadratic term of each PSORT feature, “allEuclid” has weight 1 for the quadratic term of all features, “allWeights” has weight 1 for all possible terms, and WoLF PSORT is the weight vector selected by WoLF PSORT.

extracellular. One can also see that the value of the *mit* feature is very different between the query and two of its neighbors (PEPF_ASPFU and PEPA_ASPOR). *mit* is a variable designed to discriminate between mitochondrial and non-mitochondrial proteins⁹ so this does not seem to weaken the evidence in this case.

4. Discussion

4.1. Interpretable Results

WoLF PSORT alone achieves accuracy estimates (with sequence similarity reduction used for training but not evaluation) of 73%, 77%, 83%, on the yeast, plant

Table 5. Selected Features.

| feature type | feature name | fungi | plant | animal |
|---|---------------------------|----------------|----------------|------------------|
| iPSORT | net charge(1, 25) | 1 | 0 | 1 ² |
| | max negative charge(1,20) | 0 | 1 | 0 |
| | max hydropathy(1,30) | 2 | 1 | 1,1 ² |
| PSORT | gvh | 1 | 0 | 1 ² |
| signal peptide | psg | 0 | 2 | 0 |
| PSORT | mip | 0 | 2 | 1 ² |
| targeting signal | mit | 1 | 2 | 1 |
| PSORT membrane protein related | alm | 1 | 0 | 3 ² |
| | m1a | 1 ² | 1 | 0 |
| | m1b | 1 | 3 | 0 |
| | m3a | 0 | 1 | 0 |
| | m3b | 1 | 1 | 0 |
| | mNt | 1 | 0 | 0 |
| | tms | 2 | 1 ² | 2 |
| PSORT sorting motif | erl | 1 ² | 1 ² | 0 |
| | leu | 1 | 0 | 1 |
| | nuc | 1 | 1 | 1 ² |
| | pox | 1 ² | 1 ² | 0 |
| | tyr | 2 | 1 | 1 ² |
| | yqr | 1 | 0 | 0 |
| | vac | 0 | 0 | 1 |
| PSORT non-sorting motif | dna | 1 ² | 1 ² | 1,4 ² |
| | rib | 2 ² | 1 ² | 1 ² |
| | myr | 1 ² | 0 | 1,1 ² |
| | rnp | 1 | 1 | 1 ² |
| | act | 0 | 0 | 3 ² |
| miscellaneous | length | 0 | 1 | 1 ² |

The selected features are shown with their weights. Features with 0 weight for all datasets were omitted. “1²” and “2²” indicate the quadratic term rather was selected with a weight of 1 or 2 respectively. The amino acid content features (by one letter code) selected for the three datasets were “ARNDEQGIKMFVW”, “ACQHILSV”, and “CIKS” respectively. In each of those cases the weight was 1, the weight type was always linear for the fungi and plant dataset and always quadratic for the animal dataset. “sorting” motifs refer to motifs such as the E.R. retention signal “erl” with a direct causal relationship to localization. Descriptions of the variables can be found on the WoLF PSORT server. For the PSORT variables useful documentation can also be found on the PSORT help page www.psort.nibb.ac.jp

and animal datasets, with a small number of features and the trivially simple k nearest neighbors classifier. We do not claim that this will meet the accuracy of sophisticated classifiers such as the popular support vector machine. However the template based nature of the k NN classifier makes individual classifications easier to interpret. The web server provides a tabular display to facilitate this process.

4.2. Evaluation in the presence of similar sequences

In this study we included many similar sequences. This makes achieving a high accuracy easy. However by comparing with BLAST we were able to show that our method can be effective even when sequence similarity is low. For example the

Table 6. Confusion Matrix for the Fungi Dataset.

| site | nucl | cyto | cyto | cyto | mito | mito | plas | extr | cyto | pero | E.R. | golg | vacu | cyts | sum |
|-----------|------------|-----------|------------|----------|------------|----------|------------|------------|----------|-----------|----------|----------|----------|----------|------|
| | nucl | nucl | pero | pero | nucl | pero | | | | | | | | | |
| nucl | 556 | 33 | 37 | 0 | 27 | 2 | 6 | 0 | 0 | 6 | 0 | 0 | 0 | 0 | 667 |
| cyto_nucl | 49 | 10 | 24 | 0 | 6 | 0 | 0 | 0 | 0 | 2 | 0 | 0 | 0 | 0 | 91 |
| cyto | 88 | 16 | 232 | 3 | 30 | 0 | 1 | 5 | 0 | 7 | 1 | 0 | 0 | 0 | 383 |
| cyto_mito | 1 | 0 | 0 | 0 | 7 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 8 |
| mito | 38 | 3 | 22 | 0 | 299 | 3 | 9 | 6 | 0 | 8 | 1 | 0 | 0 | 0 | 389 |
| mito_nucl | 1 | 0 | 1 | 0 | 2 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 4 |
| plas | 6 | 0 | 3 | 0 | 2 | 0 | 205 | 2 | 0 | 1 | 1 | 0 | 0 | 0 | 220 |
| extr | 2 | 2 | 4 | 0 | 2 | 0 | 0 | 129 | 0 | 0 | 1 | 0 | 0 | 0 | 140 |
| cyto_pero | 1 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 2 |
| pero | 16 | 3 | 23 | 0 | 5 | 0 | 6 | 2 | 1 | 20 | 1 | 0 | 0 | 0 | 77 |
| E.R. | 4 | 2 | 5 | 0 | 2 | 0 | 30 | 11 | 0 | 4 | 7 | 1 | 0 | 0 | 66 |
| golg | 10 | 1 | 1 | 0 | 1 | 0 | 10 | 8 | 0 | 0 | 2 | 5 | 0 | 0 | 38 |
| vacu | 2 | 0 | 2 | 0 | 2 | 0 | 10 | 6 | 0 | 0 | 1 | 0 | 0 | 0 | 23 |
| cyts | 4 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 5 |
| sum | 778 | 70 | 356 | 3 | 385 | 5 | 277 | 169 | 1 | 48 | 15 | 6 | 0 | 0 | 2113 |

The rows represent the dataset labels. The columns represent predictions made by WoLF PSORT

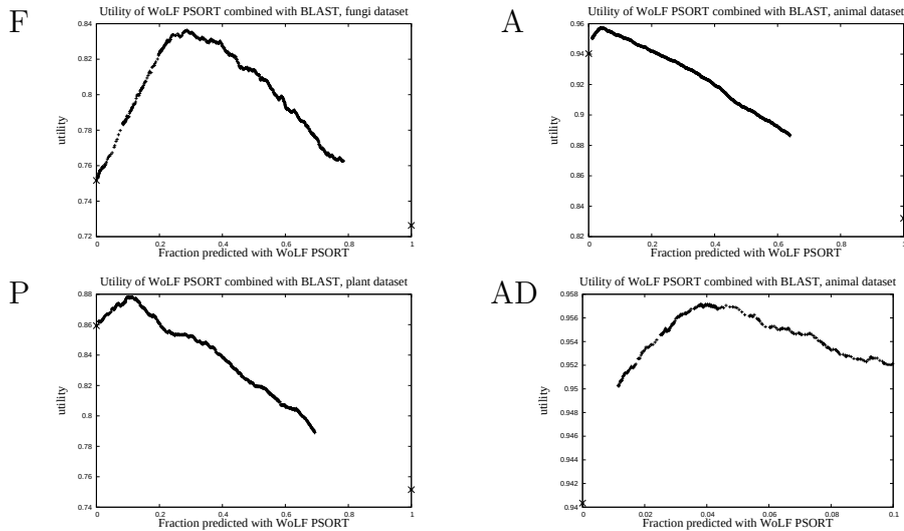


Figure 2. The utility obtained by combining WoLF PSORT and BLAST with a simple eValue threshold is shown for the fungi (F), plant (P), and animal (A) datasets. A detailed view of the animal data (AD) is shown as well. The point at $x = 1$ represents using WoLF PSORT alone. The lack of points between 0.8 and 1 for the fungi data is due to the fact that approximately 20% (30% for plant, 36% for animal) of the proteins have hits with an eValue of “0”, a similar (but smaller) gap occurs at the other end due to proteins with no blast hits.

number of errors produced by using BLAST alone on the animal dataset can be reduced by about a quarter (accuracy of $\approx 95.6\%$ vs $\approx 94.0\%$, yielding 179 errors) by using WoLF PSORT for proteins without good BLAST hits.

Normalized F

| id | site | iPSORT | | PSORT Featu | | | | | | | | | |
|------------|-------|---------------------------------------|--|---------------------|---------------------|---------------------|---------------------|---------------------|---------------------|---------------------|---------------------|---------------------|---------------------|
| | | -1 25 | MxHy1 30 | alm | dna | erl | gvh | leu | mla | mlb | m3b | mNt | mit |
| AEP_YARLI | extr? | 55 | 92 | 21 | 46 | 50 | 93 | 47 | 50 | 49 | 48 | 49 | 58 |
| AEP_YARLI | extr | 55 | 92 | 21 | 46 | 50 | 93 | 47 | 50 | 49 | 48 | 49 | 58 |
| PEPF_ASPFU | extr | 55 | 96 | 17 | 46 | 50 | 96 | 47 | 50 | 49 | 48 | 49 | 83 |
| CAR1_CANPA | extr | 55 | 93 | 20 | 46 | 50 | 96 | 47 | 50 | 49 | 48 | 49 | 55 |
| PEPA_ASPOR | extr | 55 | 93 | 21 | 46 | 50 | 93 | 47 | 50 | 49 | 48 | 49 | 94 |

Figure 3. A screen shot of the server showing the feature table displayed for query and nearest neighbors is shown. The query id, predicted localization, and features are showed aligned with the id, localization, and features of its nearest neighbors. Color is used to attract attention to large differences between the query and nearest neighbors. In the example shown the query is “AEP_YARLI”, which also appears on the next line as the nearest neighbor in the dataset.

4.3. Predicting Dual Localization

WoLF PSORT was designed with dual localization in mind. The only dual localization category for which SWISS-PROT currently contains a significant number of entries is dual localization to the nucleus and cytosol. This is an important category of proteins, for example some transcription factors are regulated by conditional localization to the nucleus⁸.

The prediction results seen in the confusion matrix (Table 6) are mixed. Unfortunately most of the 91 proteins labeled “cyto_nucl” are misclassified as either nuclear or cytosol. On the other hand, perhaps this mistake should not be looked at too harshly – as a “half-right” prediction of a dually localized protein is the *best possible* prediction for existing prediction methods which do not consider multiple localization at all. It seems likely that the current annotation in SWISS-PROT is conservative relative to multiple localization. For example in a recent large scale experiment using GFP fusion proteins to determine localization in yeast, approximately 20% out of a total of over 4000 measured proteins were found to dually localize to the nucleus and cytoplasm.

5. Conclusion

WoLF PSORT achieves a dramatic improvement in prediction accuracy over PSORTII while maintaining the simple, easily understood classifier which has been one reason for PSORTII’s widespread use. Indeed by applying a feature selection algorithm, WoLF PSORT is actually *simpler* than PSORTII in the sense that it uses fewer features for classification. WoLF PSORT is also one of the most serious attempts to date to incorporate dually localized proteins into a prediction scheme for eukaryotic cells.

6. Acknowledgement

C.J.Collier designed the initial server task handler. H. Harada helps maintain the current server. Dr. H. Ohta gave valuable advice in preparing the entries extracted from GO. This work was partly supported by MEXT.

References

1. M. Ashburner, C. A. Ball, J. A. Blake, D. Botstein, H. Butler, J. M. Cherry, A. P. Davis, K. Dolinski, S. S. Dwight, J. T. Eppig, M. A. Harris, D. P. Hill, L. Issel-Tarver, A. Kasarskis, S. Lewis, J. C. Matese, J. E. Richardson, M. Ringwald, G. M. Rubin, and G. Sherlock. Gene ontology: tool for the unification of biology. *Nature Genetics*, 25(1):25–29, 2000.
2. Hideo Bannai, Yoshinori Tamada, Osamu Maruyama, Kenta Nakai, and Satoru Miyano. Extensive feature detection of N-terminal protein sorting signals. *Bioinformatics*, 18:298–305, 2002.
3. B. Boeckmann, A. Bairoch, R. Apweiler, M. Blatter, A. Estreicher, E. Gasteiger, M. J. Martin, K. Michoud, C. O’Donovan, I. Phan, S. Pilbout, and M. Schneider. The SWISS-PROT protein knowledgebase and its supplement TrEMBL in 2003. *Nucleic Acids Research*, 31:365–370, 2003.
4. M.G. Claros and P. Vincens. Computational method to predict mitochondrially imported proteins and their targeting sequences. *European Journal of Biochemistry*, 241:779–786, 1996.
5. Richard O. Duda and Peter E. Hart. *Pattern Classification and Scene Analysis*. John Wiley & Sons, New York, 1973.
6. Paul Horton, Yuri Mukai, and Kenta Nakai. Protein localization prediction. In Limsoon Wong, editor, *The Practical Bioinformatician*, pages 193–215. World Scientific, 5 Toh Tuck Link, Singapore 596224, 2004.
7. Paul Horton and Kenta Nakai. Better prediction of protein cellular localization sites with the k nearest neighbors classifier. In *Proceeding of the Fifth International Conference on Intelligent Systems for Molecular Biology*, pages 147–152, Menlo Park, 1997. AAAI Press.
8. José M. Mingot, Eduardo A. Espeso, Eliecer Díez, and Miguel A. Peñalva. Ambient pH signaling regulates nuclear localization of the *aspergillus nidulans* PacC transcription factor. *Molecular and Cellular Biology*, 21(5):1688–1699, March 2001.
9. Kenta Nakai and Minoru Kanehisa. A knowledge base for predicting protein localization sites in eukaryotic cells. *Genomics*, 14:897–911, 1992.
10. Henrik Nielsen. Machine learning approaches for the prediction of signal peptides and other protein sorting signals. *Protein Engineering*, 12(1):3–9, 1999.
11. Henrik Nielsen, Jacob Engelbrecht, Søren Brunak, and Gunnar von Heijne. Identification of prokaryotic and eukaryotic signal peptides and prediction of their cleavage sites. *Protein Engineering*, 10:1–6, 1997.
12. K. Nishikawa and T. Ooi. Correlation of the amino acid composition of a protein to its structural and biological characters. *J. Biochem.*, 91(5):1821–1824, 1982.
13. Gunnar von Heijne. A new method for predicting signal sequence cleavage sites. *Nucleic Acids Research*, 14:4683–4690, 1986.