

## Transmembrane helices predicted at 95% accuracy

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### Abstract

We describe a neural network system that predicts the locations of transmembrane helices in integral membrane proteins. By using evolutionary information as input to the network system, the method significantly improved on a previously published neural network prediction method that had been based on single sequence information. The input data were derived from multiple alignments for each position in a window of 13 adjacent residues: amino acid frequency, conservation weights, number of insertions and deletions, and position of the window with respect to the ends of the protein chain. Additional input was the amino acid composition and length of the whole protein. A rigorous cross-validation test on 69 proteins with experimentally determined locations of transmembrane segments yielded an overall two-state per-residue accuracy of 95%. About 94% of all segments were predicted correctly. When applied to known globular proteins as a negative control, the network system incorrectly predicted fewer than 5% of globular proteins as having transmembrane helices. The method was applied to all 269 open reading frames from the complete yeast VIII chromosome. For 59 of these, at least two transmembrane helices were predicted. Thus, the prediction is that about one-fourth of all proteins from yeast VIII contain one transmembrane helix, and some 20%, more than one.

**Keywords:** evolutionary information; integral membrane proteins; multiple alignments; neural networks; protein structure prediction; secondary structure; yeast VIII chromosome

Given the rapid advance of large-scale gene-sequencing projects (Oliver et al., 1992; Johnston et al., 1994), most protein sequences of key organisms will be known in about 5 years' time. Experimental structure determination is becoming more of a routine (Lattman, 1994); and the number of proteins with known sequence for which the three-dimensional (3D) structure can be predicted rather accurately by homology modeling is constantly increasing (today more than 25% of all sequences in the SWISS-PROT sequence data base [Bairoch & Boeckmann, 1994] can be modeled with reasonable accuracy by homology [Sander & Schneider, 1994]). Even in such an optimistic scenario, experimental knowledge about membrane proteins is likely to be sparse. However, membrane proteins represent a very important class of protein structures. To what extent can structural aspects for membrane proteins be predicted from sequence information?

*Two types of membrane proteins.* So far, the 3D structures of two types of membrane proteins have been determined. The first type are helical proteins: photosynthetic reaction center (Deisenhofer et al., 1985), bacteriorhodopsin (Henderson et al.,

1990), and the light harvesting complex II (Wang et al., 1993; Kühlbrandt et al., 1994); these proteins consist of typically apolar helices of some 20 residues that traverse the membrane perpendicular to its surface (Fig. 1). The second type is represented by the structure of porin (Weiss & Schulz, 1992; Cowan & Rosenbusch, 1994), a 16-stranded  $\beta$ -barrel.

*Membrane proteins easier to predict than globular ones.* Typical methods for the prediction of transmembrane segments focus on helical transmembrane (HTM) proteins (von Heijne, 1981, 1986; Argos et al., 1982; Eisenberg et al., 1984a; Engelman et al., 1986; von Heijne & Gavel, 1988). It is commonly believed that the prediction of structure is simpler for membrane proteins than for globular ones as the lipid bilayer imposes strong constraints on the degrees of freedom of structure (Taylor et al., 1994).

*Prediction of transmembrane segments.* Methods for prediction of transmembrane helices are usually based on (1) hydrophobicity analyses (Argos et al., 1982; Kyte & Doolittle, 1982; Engelman et al., 1986; Cornette et al., 1987; Degli Esposti et al., 1990); (2) the preponderance of positively charged residues on the cytoplasmic side of the transmembrane segment (interior), established as the "positive inside rule" (von Heijne, 1981, 1986, 1991, 1992; von Heijne & Gavel, 1988; Sipos & von Heijne, 1993); or (3) statistical procedures that perform significantly bet-

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inferior to these methods. However, using information from multiple sequence alignments as input, neural networks have been shown to yield the most accurate prediction of secondary structure for globular proteins (Rost & Sander, 1993a, 1993c, 1994a). Here, we used a similar system of neural networks to predict transmembrane helices based on evolutionary information (Figs. 1, 2). The goal was to predict the location of transmembrane helices (defined as helix caps given in SWISS-PROT [Bairoch & Boeckmann, 1994]) more accurately than alternative methods (Sipos & von Heijne, 1993; Jones et al., 1994; Persson & Argos, 1994; R. Casadio et al., submitted). The neural network system was tested in fivefold cross-validation on 69 proteins with experimentally well-determined transmembrane helices (Materials and methods). Network input was the information derived for successive windows of 13 adjacent residues from a multiple sequence alignment (Fig. 3). Output were two units, one for each state of the central residue (in membrane helix/not in membrane helix; Fig. 2).

## Results and discussion

### *Evolutionary information improves prediction accuracy significantly*

*Better prediction in terms of per-residue and segment-based scores.* Compared to a simple neural network, the per-residue accuracy of the full three-level system using explicitly various aspects of evolutionary information increased by some five percentage points (Table 1). The improvement in prediction accuracy was even more significant in terms of segment-based scores: from some 75% correctly predicted segments to 94%.

*Reliability index of practical use to refine prediction accuracy.* For some 70% of all proteins, 100% of all segments were predicted correctly (data not shown). The reliability of the prediction (reliability index defined in Fig. 4) can help to estimate whether or not a protein is likely to belong to the majority of proteins for which all segments are predicted correctly (Fig. 4). Furthermore, the reliability index was used to control the filtering procedure (Fig. 5).

### *Performance similar to that of the best alternative methods*

Recently, two groups reported significant improvements in predicting transmembrane helices. Jones et al. (1994) use a new method with five output states (HTM-inside/middle/outside and not-HTM inside/outside, where inside/outside refers to inside/outside the cell). Persson and Argos (1994) use four output states (HTM-begin/middle/end and not-HTM) plus multiple alignment information. The system described here resulted in an accuracy in predicting the transmembrane helices similar to these two methods although we used only two output states. An exact comparison of the performance accuracy is made difficult because for both methods neither are per-residue scores published nor are the segment measures used defined (see footnotes to Table 1). Surprisingly, the errors made by the network system are often different from those made by the two statistical methods (Table 2 in comparison to Jones et al., 1994; Persson & Argos, 1994).

### *High reliability in discriminating between proteins with and without transmembrane helices*

Does the prediction method distinguish transmembrane from nontransmembrane proteins? Two questions are of interest. First, did the network system correctly predict all transmembrane proteins used for the cross-validation analysis as transmembrane proteins? And second, were some globular proteins falsely predicted to contain transmembrane segments?

*Transmembrane proteins correctly identified.* Both the network system using single sequences as input and the network using only profiles identified all but two proteins in the test set as transmembrane proteins: melittin (2mlt) and immunoglobulin G-binding protein precursor (iggb\_strsp). Melittin is a special case because the DSSP (Kabsch & Sander, 1983) assignment of secondary structure splits the long helix of the 26-residue molecule into two that were so short that the filtering procedure would miss this protein even on the basis of the known 3D structure. The ultimate network system PHDhtm missed only melittin; all other membrane proteins were correctly identified.

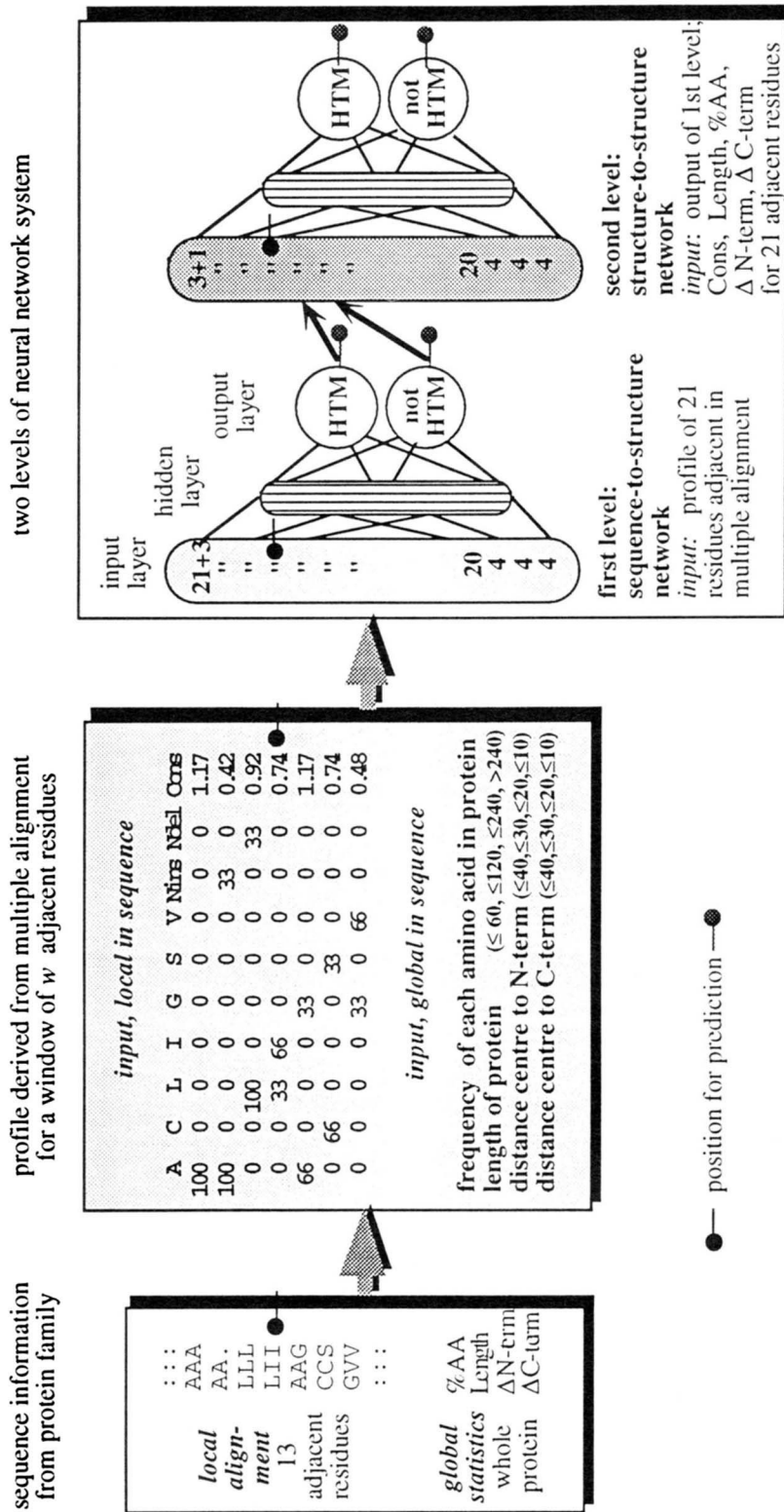
*Fewer than 5% false positives.* To test whether globular proteins were falsely predicted to contain transmembrane helices, we chose a set of 278 unique globular proteins. (No network predicted a transmembrane helix in the  $\beta$ -barrel porin.) PHDhtm mispredicted fewer than 5% of the globular proteins (Table 3). False positives were often globular water-soluble proteins with highly hydrophobic  $\beta$ -strands in the core. An exception was the only globular protein predicted to contain more than three segments: photosynthetic reaction center (4rcr) for which 11 segments with an average length of 21 residues were predicted as transmembrane helices (mandelate race mace [2mnr] was predicted with three long helices). The network using only profiles as input predicted transmembrane helices for less than 2% of the globular proteins.

### *Multilevel system improves significantly over simple neural network*

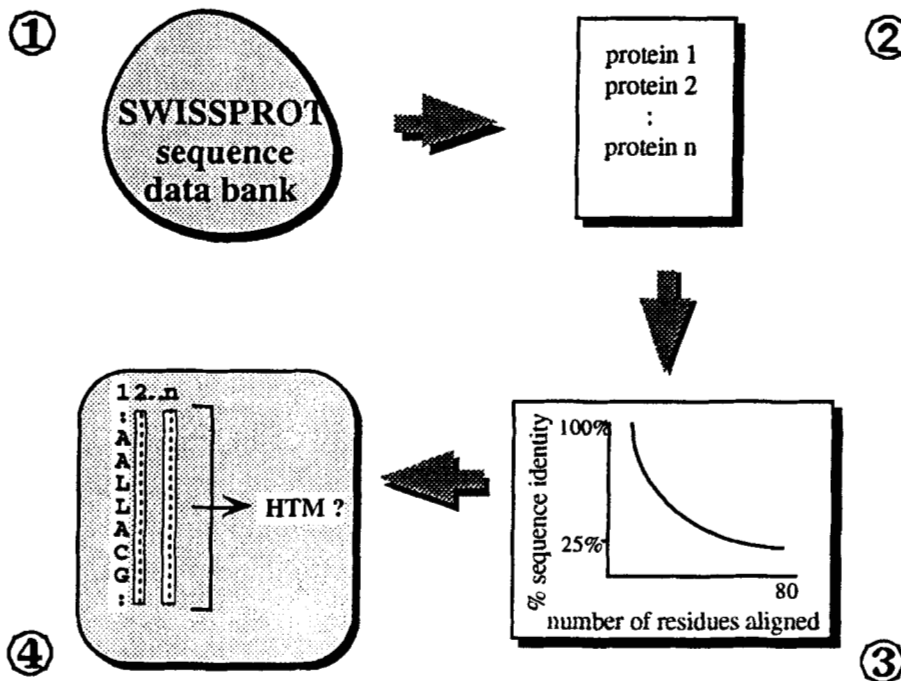
*Alignment information improves performance.* The most significant improvement in prediction accuracy (compared to a simpler neural network prediction) stemmed from including the information contained in multiple alignments. Roughly one half of the improvement attributed to simply using residue substitution frequencies (Table 4), and one half to using additionally more details contained in the alignments (conservation weight, number of insertions and deletions) and information about the whole protein (Table 4).

*Balanced versus unbalanced training.* The balanced training procedure (equally often presenting residues in transmembrane and residues not in transmembrane segments; Materials and methods) tended to overpredict transmembrane helices, whereas an unbalanced training procedure (presentation of examples according to the distribution in the training set; Materials and methods) tended to underpredict transmembrane segments.

*Jury decision finds a compromise between balanced and unbalanced training.* Both balanced and unbalanced training had advantages and disadvantages. Which of the two methods should be used for prediction? A reasonable compromise (effectively between over- and underprediction) was found by the



**Fig. 2.** Two-level system of neural networks for HTM prediction. For each position in the alignment, the amino acid frequencies were compiled, the numbers of insertions and deletions counted, and a conservation weight computed (defined in Rost & Sander, 1993b). Furthermore, "global information" (beyond the window of 13 adjacent residues) about the search sequence was compiled: amino acid composition, length, and the position of the current window with respect to the N- and C-terminal end of the protein. All this information was fed into the neural network input for  $w = 13$  adjacent residues (shown  $w = 7$ ). The input layer was fully connected to a layer with three hidden units, and from there to the two output units coding for the central residues in the window (here "LII") to be in an HTM or not. The output of the first level was fed into a second level of structure-to-structure network, which additionally used the global information and the conservation weight as input. For this network, 15 hidden units were used. The two output units code again for the secondary structure state of the central residues (here "LII"). For first-level input units, local information is coded by  $w \times (21 + 3)$  units, 20 for each amino acid, 1 for a spacer (for allowing windows to extend beyond protein ends, such that the first and last  $w - 1$  residues in a protein can be used as central residue), and 3 for conservation weights, numbers of insertions, and numbers of deletions. Global information is coded by 32 additional units; 20 for the frequency of each amino acid in the protein, 4 for the length of the protein, and 4 for the distance of the central residue to the N- and 4 for the distance to the C-term of the protein. For second-level input units, the local information is coded by  $w \times (3 + 1)$  units, two for each output unit of the first level (HTM, not HTM), one for a spacer, and one for the conservation weight of that residue. Global information is used as in the first-level input.



**Fig. 3.** Generating multiple alignments for the network input. First, for each protein the SWISS-PROT data base of protein sequences (Bairoch & Boeckmann, 1994) was searched for putative homologues with a fast alignment method (FASTA; Pearson & Lipman, 1988; Pearson & Miller, 1992). Second, the list of putative homologues was reexamined with a more sensitive profile-based multiple alignment method (Max-Hom; Sander & Schneider, 1991). Third, a length-dependent cutoff for the sequence identity between the search sequence and the aligned ones was applied to distinguish correct hits for homologues from false positives (for more than 80 residues aligned, the cutoff was chosen 25% + 5%; where the "+5%" reflects a safety margin above the line observed to separate correct and false homologues [Sander & Schneider, 1991]). Fourth, a window of 13 adjacent residues was shifted along the protein sequence. Each such window constituted one training or testing example for the neural network.

**Table 1.** Prediction accuracy cross-validated on helical transmembrane proteins<sup>a</sup>

Set <sup>b</sup>	Method <sup>c</sup>	Overall		Helical transmembrane segments only									
		<i>N</i>	<i>Q</i> <sub>2</sub>	Info	Per-residue score				Segment-based scores				
					%Obs <i>Q</i> <sub>TM</sub>	%Prd <i>Q</i> <sub>TM</sub>	Corr	< <i>L</i> >	%Obs Sov	%Prd Sov	Nseg <sup>d</sup> over	Nseg under	
Set 1	No profiles	69	90	0.45	84	70	0.71	23	90	81	15	47	
	<b>PHDhtm</b>	<b>69</b>	<b>95</b>	<b>0.64</b>	<b>91</b>	<b>84</b>	<b>0.84</b>	<b>23</b>	<b>96</b>	<b>96</b>	<b>5</b>	<b>10</b>	
											6.3%	17%	
											1.9%	3.8%	
Set 2	PHDhtm	37	95		91		0.85	23					
	Edelman (1993)	37	88		90		0.70	26					
Set 3	Jones et al. (1994)	67									15	6	
											4.5%	1.9%	
Set 4	PHDhtm	28									3-2 <sup>e</sup>	3	
											1.6%	2.3%	
											2-3 <sup>e</sup>	3	
	Persson and Argos (1994)	28									1.6%	2.3%	
	Not cross-validated <sup>f</sup>												

<sup>a</sup> *N*, number of proteins used for prediction; *Q*<sub>2</sub>, percentage of correctly predicted residues; Info, information or entropy of prediction (Rost & Sander, 1993b); *Q*<sub>TM</sub>, accuracy of predicting transmembrane helices (HTM); %Obs *Q*<sub>TM</sub>, correctly predicted residues in HTM as percentage of residues observed in HTM; %Prd *Q*<sub>TM</sub>, correctly predicted residues in HTM as percentage of residues predicted as HTM; Corr, Matthews correlation (Matthews, 1975) for residues in HTM; <*L*>, average length of predicted HTM (the observed average is <*L*> = 22); %Obs Sov, segment overlap for HTM computed as percentage of observed segments (Rost et al., 1994); %Prd Sov, segment overlap for HTM computed as percentage of predicted segments (Rost et al., 1994); Nseg over, number of segments predicted but not observed as HTM; Nseg under, number of segments observed but not predicted as HTM. Bold indicates the reference levels.

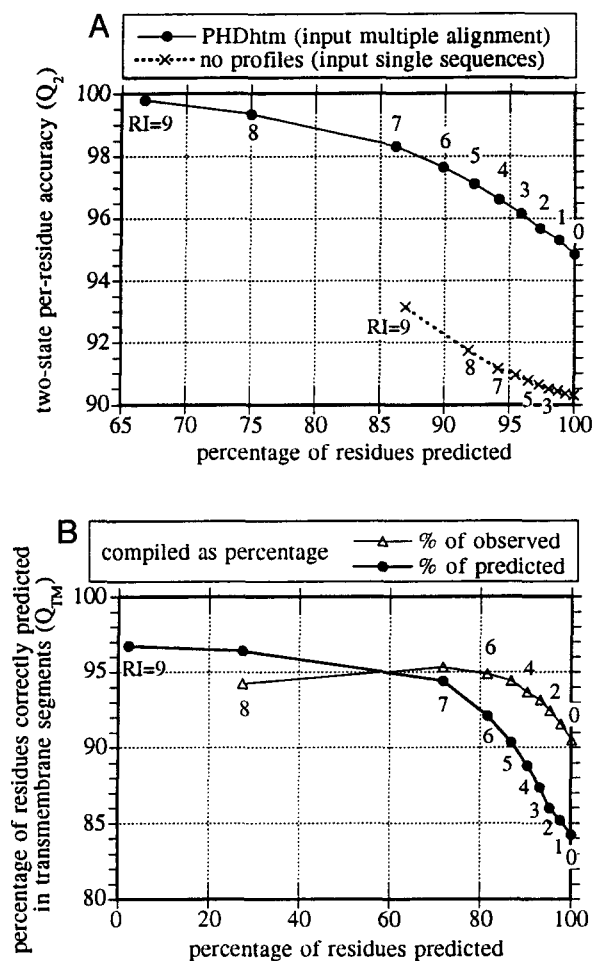
<sup>b</sup> Set 1, set of 69 proteins with experimentally well-determined transmembrane helices (see Materials and methods); set 2, set of 37 transmembrane proteins used by Edelman (1993); set 3, set 1 without glra\_rat and 2mlt; set 4, set of 28 transmembrane proteins used by Persson and Argos (1994).

<sup>c</sup> No profiles, two-level network system using single sequences as input (R. Casadio et al., submitted); PHDhtm, three-level network system + filter using all information from multiple alignments as input (Fig. 2).

<sup>d</sup> Whenever predicted and observed segments overlapped by at least three residues, the segment was counted as correct (Rost et al., 1993, 1994). A similar measure seems to have been used by others. A more reasonable score is the segment overlap Sov (Rost et al., 1994).

<sup>e</sup> Discrepancy in assigning transmembrane helices for atpi\_pea; both methods compared predict five transmembrane helices. In SWISS-PROT only four are annotated; thus, we initially counted our prediction as wrong, whereas Persson and Argos (1994) based their evaluation on the hypothesis that the protein contains five and not four transmembrane helices.

<sup>f</sup> All results except for those in the last row were based on cross-validation tests. Persson and Argos (1994) reported that for their method the results with or without cross-validation analysis are similar and only gave the non-cross-validated results on proteins in their training set.



**Fig. 4.** Reliability of prediction. Reliability index ( $RI$ ) for the prediction was defined as proportional to the difference between the two output units:

$$RI = \text{INTEGER} (10 \times [\text{out}_{\text{HTM}} - \text{out}_{\text{noHTM}}]).$$

The factor 10 scales the reliability index to values 0–9. **A:** Overall two-state per-residue accuracy versus the cumulative percentage of residues with a reliability index  $RI \geq n$ ,  $n = 0, \dots, 9$ . Note that  $RI \geq 0$  is the rightmost point representing 100% of the predicted residues. Results were averaged over the residues in all 69 transmembrane proteins used for the cross-validation test. A network system that used multiple alignments as input was compared to a network system that used single sequence information only. For example, 90% of all residues were predicted with  $RI \geq 6$ . For these, the prediction accuracy for the network using multiple alignment information reached a value of  $Q_2 > 97\%$ . **B:** Percentage of residues correctly predicted in transmembrane helices versus cumulative percentage of residues predicted in transmembrane helices with a reliability index  $RI \geq n$ . Results are given as percentages of the number of residues observed in transmembrane helices (open triangles) and as percentages of the number of residues predicted in transmembrane helices (filled circles). For example, about 70% of all residues predicted in transmembrane segments had a reliability index  $RI \geq 7$ . Ninety-five percent of these were predicted correctly.

jury decision, i.e., the arithmetic average over the output values of balanced and unbalanced networks.

**Second-level elongates helices.** The effect of the second-level (structure-to-structure) network was to elongate or delete short

helical segments. The effect was an increase in the average length of a predicted helical segment from 15 residues for the first level, to 27 residues for the second level (Table 4). In other words, the first-level networks (Fig. 2) yielded an average length for transmembrane segments 5–7 residues shorter than observed; the second-level networks (Fig. 2) resulted in segments up to 13 residues longer than observed. Thus, the second-level networks tended to elongate helices (Table 4).

**Final filtering procedure.** Short loop regions were often missed by the second network, which tended to elongate helices too much (note that the input window is too narrow to learn a maximal length for transmembrane segments). This drawback was compensated by a relatively straightforward filtering procedure (Materials and methods). Filtering improved the prediction accuracy both in terms of per-residue and segment-based measures for prediction accuracy (Table 4).

## Conclusion

**Selection of data set.** The 3D structure is experimentally known for only five (Iprc\_H, Iprc\_L, Iprc\_M, Ibrd, 2mlt) of the 69 protein chains used for the cross-validation analysis. This implies that the results ought to be taken with caution. To increase confidence in the results, we deliberately chose proteins for which there is “reliable” experimental evidence about the locations of the transmembrane regions (list taken from Jones et al., 1994), rather than working with a larger data set including less well-known segments.

**Improved prediction of transmembrane helices.** Using various aspects of evolutionary information improved the overall per-residue accuracy of predicting residues in transmembrane helices by some five percentage points. This improvement could be significant enough to warrant use of the predictions as a starting point for a complete ab initio prediction of 3D structure for transmembrane regions (Baldwin, 1993; Taylor et al., 1994). Our best network system (called PHDhtm) correctly predicted some 94% of all segments and the correct location of some 90% of all residues observed in transmembrane helices. For only 4 of 15 incorrectly predicted (either under-, or overpredicted) segments, the defined reliability index would have led the user to suspect a wrong prediction (Fig. 1).

**Prediction for globular proteins sufficiently accurate.** The two-level network system using only profiles as input mispredicted less than 2% of globular proteins as containing transmembrane helices (Table 3). An unsatisfactory disadvantage of the most accurate network system PHDhtm was that this error rate was clearly higher (<5%). However, for most practical purposes this rate of false positives is sufficiently low. All transmembrane proteins were predicted to contain at least one transmembrane helix, except for melittin, which would not have been recognized as transmembrane helix even from the crystal structure: the strongly bent helix is split into two short helices by the program assigning the secondary structure automatically from 3D structures (DSSP; Kabsch & Sander, 1983).

**Weak point.** A rather inconvenient aspect of the method described here is the necessity to apply a filter procedure (Fig. 5) at the end of the prediction. This disadvantage is one of the details that still has to be improved in a more general tool.

too short helices	
if { $L < 17 \cap RI > 7$ (at either end of helix) }-->	elongate helix by one residue until $L \geq 17$
if { only one helix predicted } if { $L < 17$ }	--> cut helix
if { at least 2 helices predicted } if { $L < 11$ }	--> cut helix
too long helices	
if { $L > 35$ }	--> split helix at position $L/2$ into two helices of length $L/2$
if { $L > n \times 22$ , $n=3,4,\dots$ }	--> split helix into $n$ of length $L/n$

**Fig. 5.** Filtering the prediction. Output of the third level (jury prediction) was filtered to delete too-short and to split too-long predicted transmembrane helices. Splitting of too-long segments was usually done exactly in the middle of the segment by flipping the prediction for one residue from HTM to not-HTM. Two exceptions were: (1) if there was a residue in a three-residue neighborhood of the central residue with a lower reliability index than that of the central one, then splitting was performed at that residue; (2) if the two residues on both sides of the central residue were predicted with an  $RI < 3$ , then up to five residues in total were flipped from the state HTM to not-HTM.

*Possible improvements of the prediction.* There are methods that predict whether or not a loop region is located inside or outside the cell (von Heijne & Gavel, 1988; Nakashima & Nishikawa, 1992; von Heijne, 1992; Sipos & von Heijne, 1993; Jones et al., 1994). Such tools could be used to either complement the network prediction, or directly to train a network to predict transmembrane topology (direction of transmembrane helices with respect to cell).

*$\beta$ -Strand membrane proteins.* How can transmembrane segments for  $\beta$ -barrel proteins such as porin be predicted from sequence? Interestingly, the network system trained on water-soluble globular proteins (PHDsec), predicts the  $\beta$ -strands of the membrane protein porin more accurately than the helices of the photoreaction center, bacteriorhodopsin, or the light harvesting complex. The reason may be that the pore of porin is exposed to solvent and thus resembles globular proteins in some respects. The prediction of  $\beta$ -strands, combined with hydrophobicity scales (Eisenberg et al., 1984b) and/or predictions of solvent accessibility (Rost & Sander, 1994b), has been used to infer which of the porin strands may be in contact with lipids. Unfortunately, however, the structures of very few  $\beta$ -strand membrane proteins are known. Thus, training of neural networks, as well as the application of statistical methods, is premature.

*3D structure prediction.* How can one come closer to the goal of 3D prediction for helical membrane proteins? One way to go from accurate predictions of HTM locations to 3D structure has been indicated by Taylor et al. (1994). Whether or not the network predictions described here, in combination with a prediction of segment orientation relative to the membrane surface, will be useful remains to be shown.

*Keeping up with the flow of genome data.* All results reported here refer to completely automatic usage of PHDhtm. In some cases, prediction accuracy can certainly be improved by expert knowledge, e.g., by fine tuning the alignment. However, fully automatic use permits the analysis of many proteins, e.g., all open reading frames of complete chromosomes. For example, less than an hour of CPU time (on a SUN SPARC10 workstation) was required for the transmembrane helix prediction of all proteins of yeast chromosome VIII (Johnston et al., 1994), given the multiple sequence alignments. For 59 of the 269 proteins at

least two transmembrane helices were predicted (Table 5); for another 27 of the proteins one transmembrane helix was predicted. Given an error rate of 5%, this implies that 20–25% of all yeast VIII proteins were predicted to contain transmembrane helices.

*Availability of the network prediction.* Predictions of transmembrane helices (as well as secondary structure and solvent accessibility for globular proteins) using the method presented here are provided via an automatic electronic mail server. If you send the sequence of your protein, the server will return a multiple sequence alignment and a prediction of the location of transmembrane helices. For further information, send the word *help* to the Internet address *PredictProtein@EMBL-Heidelberg.DE* by electronic mail, or use the World Wide Web (WWW) site <http://www.embl-heidelberg.de/predictprotein/predictprotein.html>.

## Materials and methods

### Database

*Selection of proteins.* We based our analyses on a set of 69 proteins for which experimental information about the location of transmembrane helices is annotated in the SWISS-PROT database (Manoil & Beckwith, 1986; von Heijne & Gavel, 1988; von Heijne, 1992; Sipos & von Heijne, 1993; Jones et al., 1994). This set in particular was chosen to meet three criteria: (1) reliability: the experimental information should be as reliable as possible (Manoil & Beckwith, 1986; von Heijne, 1992); (2) comparability: to enable a comparison to similar methods, the data set should be similar to those used by others; (3) availability: the list (Table 2) was the subset of those proteins used by Jones et al. (1994) that were available in SWISS-PROT when we had started the project (melittin [2mlt] and the glutamic acid receptor [glra\_rat, O'Hara et al., 1993] were added). For the few known 3D structures, the location of the transmembrane regions was taken from DSSP (Kabsch & Sander, 1983). The exact locations of the transmembrane helices are often controversial. To enable a straightforward comparison to future methods and for making our results easily reproducible for others, we decided to always use the definitions found in SWISS-PROT (Bairoch & Boeckmann, 1994).

**Table 2.** Observed and predicted transmembrane helices for 69 proteins<sup>a</sup>

Protein	Observed HTM	Predicted HTM	Protein	Observed HTM	Predicted HTM	Protein	Observed HTM	Predicted HTM
lbrd	23-42	24-43	adt_ricpr	219-239	217-239	glpa_human	92-114	91-114
(bacr_halha)	57-76	55-87	(continued)	280-300	271-298	glpc_human	58-81	57-81
	95-114	92-116		321-341	322-342	glra_rat	539-558	536-557
	121-140	121-143		349-369	348-371		585-603	—
	148-167	145-169		380-400	377-400		614-632	615-636
	191-210	185-211		439-459	444-461		806-826	807-826
	217-236	213-239		466-486	469-485	gmcr_human	321-346	326-351
lprc_H	12-35	12-31	bach_halhm	23-42	24-43	gp1b_human	148-172	147-171
lprc_M	52-76	43-59		57-76	55-87	gpt_crilo	7-32	12-38
	—	63-78		95-114	92-116		58-79	59-83
	111-137	110-130		121-140	121-143		95-114	96-115
	143-166	143-170		148-167	145-169		126-145	127-150
	198-223	198-223		191-210	185-211		165-184	157-181
	260-284	262-292		217-236	213-239		195-211	187-210
lprc_L	33-53	21-38	cb21_pea	62-81	69-75		222-240	224-242
	—	42-58		114-134	115-134		253-269	249-269
	84-111	81-103		182-198	184-196		275-294	277-292
	116-139	115-146	cek2_chick	365-389	371-389		379-397	379-402
	171-198	173-196	cyoa_ecoli	—	12-24	hema_cdvo	35-55	37-58
	226-249	223-255		51-69	44-66	hema_measi	35-55	37-58
2mlt	2-10	—		93-111	90-109	hema_pi4ha	35-59	37-59
	12-25	—	cyob_ecoli	17-35	—	hg2a_human	46-72	50-67
4f2_human	82-104	82-104		58-76	61-77	iggb_strsp	—	18-32
5ht3_mouse	246-272	238-270		102-121	101-131		—	91-103
	278-296	282-301		144-162	146-158		423-443	425-439
	306-324	307-331		195-213	191-212	il2a_human	241-259	235-258
	465-484	457-484		232-250	227-252	il2b_human	241-265	236-267
a1aa_human	54-79	56-79		277-296	286-302	ita5_mouse	356-381	355-383
	92-117	92-116		320-339	315-335	lacy_ecoli	11-33	11-36
	128-150	128-150		348-366	349-368		47-67	46-67
	172-196	173-189		382-401	380-401		75-99	75-98
	210-233	213-235		410-429	415-440		103-125	104-126
	307-331	309-329		457-476	457-470		145-163	148-161
	339-363	—		494-513	498-519		168-187	169-187
a2aa_human	34-59	32-60		588-607	592-608		212-234	219-238
	71-96	69-100	cyoc_ecoli	614-634	612-626		260-281	265-288
	107-129	106-133		32-50	29-50		291-310	294-314
	150-173	151-169		67-85	67-85		315-334	320-337
	193-217	196-221		102-120	101-116		347-366	343-371
	375-399	375-399		143-161	138-162		380-399	377-400
	407-430	405-429		185-203	178-202	lech_human	40-60	40-59
a4_human	700-723	702-722	cyod_ecoli	18-36	20-39	leci_mouse	40-60	40-59
aa1r_canfa	11-33	12-35		46-64	45-64	lep_ecoli	4-22	4-23
	47-69	39-53		81-99	80-101		58-76	63-82
	—	61-74	cyoe_ecoli	10-28	12-24	magl_mouse	517-536	515-534
	81-102	80-110		38-56	44-66	malf_ecoli	17-35	21-35
	124-146	125-144		79-97	90-109		40-58	43-58
	177-201	176-206		108-126	109-127		73-91	71-93
	236-259	235-261		—	142-158		277-295	278-306
	268-292	266-291		—	166-181		319-337	318-339
aa2a_canfa	8-30	10-32		198-216	198-222		371-389	370-390
	44-66	40-71		229-247	228-252		418-436	418-444
	78-100	77-105		269-287	265-287		486-504	486-505
	121-143	122-141	edg1_human	47-71	45-72	motb_ecoli	28-49	30-51
	174-198	174-203		79-107	80-107	mprd_human	186-210	185-211
	235-258	234-260		122-140	116-145	myp0_human	—	14-31
	267-290	266-290		160-185	160-180		154-179	155-183
adt_ricpr	34-54	31-46		202-222	201-227	ngfr_human	251-272	253-272
	68-88	60-87		256-277	254-282			
	93-113	92-115	egfr_human	294-314	288-312			
	148-168	134-148		646-668	648-666			
	—	156-170	fce2_human	22-47	27-47			
	185-205	185-206	glp_pig	63-85	63-84			

(continued)



Table 2. Continued

Protein	Observed HTM	Predicted HTM	Protein	Observed HTM	Predicted HTM	Protein	Observed HTM	Predicted HTM
nep_human	28-50	30-49	ops3_drome	134-152	125-153	opsg_human	219-246	219-245
oppb_salty	10-30	10-29	(continued)	172-196	169-194	(continued)	269-292	269-295
	100-121	96-120		221-248	221-248		301-325	301-325
	138-158	130-162		285-308	285-308	opsr_human	53-77	52-78
	173-190	168-193		317-341	317-340		90-115	90-119
	227-250	228-259	ops4_drome	54-78	53-81		130-149	131-155
	272-293	273-298		91-113	91-115		169-192	168-192
oppc_salty	38-59	39-59		130-149	121-150		219-246	219-245
	102-122	98-126		168-192	166-191		269-292	270-295
	140-160	141-158		217-244	217-244		301-325	301-325
	164-180	166-182		281-304	281-304	pigr_human	621-643	624-643
	216-236	210-225		313-337	313-336			
	-	232-248	opsb_human	34-58	33-59	pt2m_ecoli	25-44	20-42
	268-290	268-289		71-96	71-100		51-69	54-65
ops1_calvi	48-72	47-75		111-130	112-135		135-154	133-156
	85-110	85-110		150-173	149-173		166-184	167-181
	125-144	116-145		200-227	200-227		-	249-262
	164-187	162-187		250-272	251-275		274-291	270-283
	212-239	212-239		282-306	281-306		314-333	312-332
	275-298	275-298	opsd_bovin	37-61	36-62	sece_ecoli	19-36	20-34
	306-330	306-329		74-99	74-104		45-63	42-62
ops2_drome	57-81	55-84		114-133	115-139		93-111	93-123
	94-119	94-118		153-176	152-176	suis_human	13-32	12-33
	134-153	124-153		203-230	203-230	tcb1_rabbit	292-313	285-312
	173-196	171-196		252-276	253-279	trbm_human	516-539	515-536
	221-248	221-248		285-309	285-309	trsr_human	63-88	67-86
	284-307	284-307	opsg_human	53-77	52-78	vmt2_iaann	25-42	27-51
	315-339	315-338		90-115	90-120	vnb_inbbe	19-40	19-42
ops3_drome	58-82	57-85		130-149	131-155			
	95-119	95-119		169-192	168-192			

<sup>a</sup> For the 69 transmembrane proteins used for cross-validation, the following data are listed: (1) the protein name, given by the SWISS-PROT identifier (Bairoch & Boeckmann, 1994); if the 3D structure is known, then the PDB code plus chain identifier is used (Bernstein et al., 1977; Kabsch & Sander, 1983); (2) the positions for the transmembrane helices observed (=SWISS-PROT documentation, or DSSP [Kabsch & Sander, 1983]), counted from the first residue in SWISS-PROT or DSSP; and (3) the cross-validated prediction by the network system PHDhtm. Except for 2mlt and glra\_rat, the list comprises a subset of the proteins used by David Jones (Jones et al., 1994) and Gunnar von Heijne (von Heijne & Gavel, 1988; von Heijne, 1992; Sipos & von Heijne, 1993).

**Generation of multiple alignments.** For each of the initial 69 proteins, a multiple sequence alignment was generated using the program MaxHom (Sander & Schneider, 1991; Fig. 3). All sequences from SWISS-PROT with a sequence identity above a length-dependent cut-off were included in the alignment (Sander & Schneider, 1991), assuming that this is valid not only for globular but also for membrane proteins.

**Cross-validation test.** The set of 69 transmembrane proteins (Table 2) was divided into 52 proteins used for training and 17 used for testing the method. This was repeated five times (five-fold cross-validation), until each protein had been in a test set once. The sets were chosen such that no protein in the multiple alignments used for testing had more than 25% sequence identity to any protein in the multiple alignments of the training set. All results reported are averages over proteins in various test sets.

#### Neural network system

**First level: Sequence-to-structure.** The principles of neural networks for secondary structure prediction (Fariselli et al.,

1993; Rost & Sander, 1993a) and of coding multiple sequence information (Rost & Sander, 1993b, 1994a, 1994b) are described in detail elsewhere. Here, only some basic concepts will be recapitulated and details regarding the application to transmembrane helices will be introduced.

Input to the first-level network consisted of two contributions, (1) one local in sequence, i.e., taken from a window of 13 adjacent residues; and (2) another global in sequence, i.e., compiled from the whole protein (Fig. 2). (1) The local information computed for each residue in the window was the frequency of occurrence of each amino acid at that position in the multiple alignment, the number of insertions and deletions in the alignment for that residue, and a position-specific conservation weight (Fig. 2). (2) As global information, we used the amino acid composition and length of the protein and, furthermore, the distance (number of residues) of the first residue in the window of 13 adjacent residues from the protein begin (N-term), and the distance of the last residue in the window to the protein end (C-term).

Output of the first-level network was two units, one representing examples with the central residue of the window in a

**Table 3.** Prediction accuracy on globular proteins (negative control)<sup>a</sup>

Method	Number of globular proteins used	Number of proteins predicted with HTM	Number of HTM segments longer than 16 residues	% False classifications
No profiles	278	18	7	6.5%
Profiles only	278	5	4	1.8%
PHDhtm	278	12	7	4.3%
Jones et al. (1994)	155	5	—	3.2%
Edelman (1993)	14	3	—	21.4%

<sup>a</sup> Abbreviations for methods as in Table 1 and Table 4. We considered a globular protein to be mispredicted if either at least two transmembrane segments are predicted with more than 10 residues, or at least one with more than 17 residues. Results from Edelman (1993) and Jones et al. (1994) were taken from the literature.

transmembrane helix; the other representing examples with the central residue not in transmembrane helices (Fig. 2).

**Balanced and unbalanced training.** Training was performed with the usual gradient descent (also known as back-propagation [Rumelhart et al., 1986]):

$$\Delta J_{ij}(t+1) = \epsilon \frac{E(t)}{J_{ij}(t)} + \alpha \Delta J_{ij}(t-1),$$

where  $t$  is the algorithmic time step (i.e., change of all connections for one pattern),  $E$  is the error, given by the difference be-

tween actual network output and the desired output (i.e., the value observed for the central residue);  $J_{ij}$  is the connection from unit  $j$  to unit  $i$  on the next layer (input to hidden, hidden to output);  $\epsilon$  is the learning speed, chosen here to be 0.01; and  $\alpha$  the momentum term (permitting uphill moves) chosen here to be 0.2. Two modes were used. First, unbalanced training: at each time step of the error minimization one pattern was chosen at random from the training set, and all connections of the network were changed. Second, balanced training: at each time step of the error minimization (Equation 1), one pattern from the class “transmembrane helix” and one from the class “not transmembrane helix” was used to change all connections.

**Table 4.** Analysis of the performance for each element of the network system<sup>a</sup>

Set	Method <sup>b</sup>	System levels <sup>c</sup>	Overall		Transmembrane helices only					
			$Q_2$	Info	Per-residue score			Segment-based scores		
					%Obs $Q_{TM}$	%Prd $Q_{TM}$	Corr	$\langle L \rangle$	%Obs Sov	%Prd Sov
Set 5	No profiles	2 + filter	90	0.45	84	70	0.71	23	90	81
	Profiles only	2 + filter	94	0.56	86	82	0.80	23	93	90
	<b>PHDhtm</b>	3 + filter	<b>95</b>	<b>0.65</b>	<b>91</b>	<b>84</b>	<b>0.85</b>	<b>23</b>	<b>96</b>	<b>96</b>
Set 1	First unbalanced	1	93	0.52	78	81	0.75	15	84	80
	First balanced	1	91	0.53	91	71	0.76	17	80	72
	First unbalanced–second unbalanced	2	93	0.52	83	80	0.77	22	88	83
	First balanced–second unbalanced	2	93	0.52	83	80	0.77	22	88	83
	First unbalanced–second balanced	2	91	0.55	91	69	0.75	36	71	63
	First balanced–second balanced	2	93	0.58	93	75	0.79	29	80	75
	Jury over four networks	3	91	0.58	94	69	0.75	36	71	63
	<b>PHDhtm</b>	3 + filter	<b>95</b>	<b>0.64</b>	<b>91</b>	<b>84</b>	<b>0.84</b>	<b>23</b>	<b>96</b>	<b>96</b>

<sup>a</sup> See Table 1 for abbreviations of measures. Bold indicates the reference levels for each set.

<sup>b</sup> PHDhtm, three-level network system + filter using all information from multiple alignments as input (Fig. 2); No profiles, two-level network system using single sequences as input (R. Casadio et al., submitted); Profiles only, same as before, but using evolutionary profiles (and no further information derived from the multiple alignment) as input; First unbalanced, first-level network with unbalanced training (see Materials and methods); First balanced, first-level network with balanced training (see Materials and methods); First  $x$ –second  $y$ , a second-level network with  $y$  (balanced or unbalanced) training that uses as input the prediction from a first-level network with  $x$  (balanced or unbalanced) training; Jury over four networks, arithmetic average over the four different second-level networks given above.

<sup>c</sup> Levels of the network system used (Fig. 2): 1, only first level; 2, first and second level; 3, jury average over different second-level networks (see Materials and methods); filter, application of the filtering procedure (Fig. 5). Set 1 contains 69 transmembrane proteins (see Materials and methods). Set 5 is the subset of set 1 without the PDB proteins 2mlt, 1prc (chains H, L, M), and 1brd.

**Table 5.** Prediction of transmembrane helices for yeast chromosome VIII<sup>a</sup>

Identifier	Nres <sup>b</sup>	Nali <sup>b</sup>	Locations of predicted segments			Nhtm <sup>b</sup>	
YHL040c	627	5	75-88 205-216 363-387 568-581	116-127 231-252 404-418	141-157 285-308 429-441	173-190 326-342 458-477	13
YHL047c	637	5	70-83 200-211 358-382 563-576	111-122 226-247 400-413	136-152 280-303 425-436	168-185 321-337 453-473	13
YHR092c	560	21	70-87 215-226 435-459	124-139 247-261 474-492	152-171 369-385 500-518	179-196 400-413	11
YHR096c	592	18	85-101 230-241 450-475	138-154 262-276 489-507	167-186 385-400 515-533	194-212 415-428	11
YHR094c	570	17	64-80 209-220 429-453	118-133 241-255 468-486	146-165 363-379 494-512	173-191 394-407	
YHR026w	213	18	20-37 180-205	56-80	94-122	145-168	5
YHR002w	357	8	37-53 271-281	102-115	141-153	201-227	5
YHL048w	381	4	39-62	70-93	233-252	260-277	4
YHR190w	444	4	272-283	295-310	425-440		3
YHR129c	384	258	137-153	349-360			2
YHR005c	472	153	337-347	377-387			2
YHR183w	489	39	360-371	418-429			2
YHR046c	295	7	103-117	201-216			2
YHR176w	373	6	262-272	338-351			2
YHR039c	644	5	49-66	247-264			2
YHL011c	320	22	73-92				1
YHR028c	818	8	26-44				1
YHR007c	530	7	25-47				1
YHR037w	575	4	209-227				1
YHL016c	735	1	17-33 193-213 358-375 500-516	91-108 256-266 402-421 620-642	137-153 287-311 429-450 651-674	167-186 339-350 458-476	15
YHL035c	1,592	1	33-48 335-357 574-591 1,141-1,158	172-187 378-395 977-998 1,226-1,247	201-217 465-486 1,042-1,058 1,255-1,274	229-239 490-510 1,120-1,137	15
YHL036w	546	1	69-92 211-235 398-413	100-122 261-273 433-445	149-171 298-315 461-477	187-203 345-367 492-519	12
YHR048w	514	1	75-91 197-221 390-407	112-126 229-249 415-438	143-160 308-334 478-498	168-184 343-364	11
YHR050w	549	1	92-106 246-257 434-451	135-156 309-333 518-538	164-181 361-376	199-218 409-423	10
YHR123w	391	2	40-67 267-286	123-156 294-312	177-199 320-342	218-235 350-372	8
YHL003c	411	3	82-100 256-288	133-160 303-319	181-198 353-383	216-238	7
YHL017w	532	2	194-212 331-353	227-243 376-399	260-290 420-438	307-318	7
YHR050w	549	1	92-106 246-257	135-156 309-333	164-181 361-376	199-218 409-423	

<sup>a</sup> As a typical example for the application of the method and as an independent test of the predictive power of the method, we predicted the transmembrane helices for all proteins from the complete yeast chromosome VIII (Johnston et al., 1994). For 59 proteins (of 269), two or more transmembrane helices were predicted. Proteins are labeled by the identifier used in Johnston et al. (1994). Shown are the predictions only for those proteins for which sufficient alignment information was available (P. Bork, C. Ouzounis, & C. Sander, manuscript in prep.) or which were predicted to have more than six transmembrane segments. In some cases, confirmation of the correctness of the prediction comes from detailed sequence analysis (Johnston et al., 1994; P. Bork, C. Ouzounis, & C. Sander, unpubl.): the likely function identified on the basis of sequence similarity to proteins of known function is consistent with the presence of HTM regions. Examples are: YHR026w, an ATPase; YHR048w, a resistance protein, probably works by pumping substances out of the cell through a membrane pore; YHR050w/92c/94c/96c, potential transporters; YHR190w, farnesyltransferase; YHR123w, phosphor transferase; YHR005c, G-protein  $\alpha$  subunit; YHR183w/39c, dehydrogenase.

<sup>b</sup> Nres, length of protein; Nali, number of sequences in the multiple alignment ("1" means that the prediction is based on a single sequence only); Nhtm, predicted number of transmembrane segments.

**Network parameters.** All units were connected to all those on the next layer (input to hidden, hidden to output). Network parameters such as criterion to terminate the training procedure, number of hidden units, training speed ( $\epsilon$  in Equation 1), and momentum term ( $\alpha$  in Equation 1) were chosen arbitrarily based on our experience with secondary structure prediction for globular proteins. In other words, these parameters were not influenced by the test set. Training was stopped when the training set had been learned to an accuracy of 93% for the first- and of 95% for the second-level network. As for the number of hidden units, we started arbitrarily with 3 hidden units for the first level of network and increased the number for the second-level network to 15 because training too often ended in local minima.

**Second level: Structure to structure.** The input to the second-level network consisted—as for the first-level—of a contribution local in sequence and a contribution global in sequence (Fig. 2). (1) For each residue in the input window, the local input were the values of the two output units of the first-level network and the conservation weight. (2) The global input information was the same as for the first-level network. The output of the second-level network—as for the first—consisted of two units for the central residue either being in a transmembrane helix or not.

**Third level: Jury decision.** To find a compromise between networks with balanced and those with unbalanced training, a final jury decision was performed (effectively a compromise between over- and underprediction, Results). The jury decision was a simple arithmetic average over four differently trained networks: all combinations ( $2 \times 2$ ) of first-level network with balanced and unbalanced training, and with balanced or unbalanced training of second-level network. Final prediction was assigned to the unit with maximal output value (“winner takes all”).

**Fourth level: Filtering the prediction.** In contrast to earlier prediction methods (Jones et al., 1992; von Heijne, 1992; Persson & Argos, 1994), which explicitly fix the length of predicted transmembrane segments to typically 17–25 residues, the second-level network occasionally resulted in transmembrane helices that were either too short or too long. This was corrected by a nonoptimized filter that was guided by the experiences of previous work (von Heijne, 1986, 1992; von Heijne & Gavel, 1988; Sipos & von Heijne, 1993; Jones et al., 1994; R. Casadio et al., submitted).

Too long helices were either split in the middle into two shorter helices or were shortened (Fig. 5). Too short helices were either elongated or deleted. All these decisions (split or shorten; elongate or delete) were based both on the strength of the prediction (reliability index, Fig. 2) and on the length of the predicted transmembrane helix (Fig. 5).

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