The Na\textsuperscript{+}/H\textsuperscript{+} antiporters catalyze the exchange of Na\textsuperscript{+} for H\textsuperscript{+} across membranes and play a variety of functions such as the regulation of internal pH, cell volume, and sodium level in the cytoplasm (Padan and Schuldiner, 1996). In Escherichia coli, three antiporters (NhaA, NhaB, and ChaA) are known, and their functional characteristics have been well described (Padan and Schuldiner, 1996). In yeast, the Na\textsuperscript{+}/H\textsuperscript{+} antiporters localized in the prevacuolar membrane (NHX1) (Nass et al., 1997; Nass and Rao, 1998) and the plasma membrane (SOD2) (Jia et al., 1992; Hahnenberger et al., 1996) have been reported. In animals, six kinds of Na\textsuperscript{+}/H\textsuperscript{+} antiporters (exchangers) (NHE1-6) have been found (Orlowski and Grinstein, 1997); the mitochondrial localization was demonstrated in NHE6 (Numata et al., 1998). A new unique Na\textsuperscript{+}/H\textsuperscript{+} antiporter (NhaP), which has no homology to NhaA, NhaB, and ChaA, has been found recently in Pseudomonas aeruginosa (Utsugi et al., 1998).

In plants, Na\textsuperscript{+}/H\textsuperscript{+} antiporters, homologous to the vacuolar localized NHX1, have been isolated from Arabidopsis (AtNHX1) (Gaxiola et al., 1999) and Oryza sativa (Fukuda et al., 1999). The enhanced salt tolerance of Arabidopsis has been demonstrated by the overexpression of AtNHX1 (Apse et al., 1999). The SOS1 gene, which is essential for Na\textsuperscript{+} and K\textsuperscript{+} homeostasis, recently was isolated from Arabidopsis (Shi et al., 2000). The expression of SOS1 was significantly enhanced by NaCl stress, especially in the root. SOS1 exhibited very high sequence homology to the NhaP antiporter from P. aeruginosa (Utsugi et al., 1998), but is more distantly related to a cluster of organellar Na\textsuperscript{+}/H\textsuperscript{+} antiporters such as AtNHX1, NHX1, or NHE6 (Shi et al., 2000). From these results, they suggested that the SOS1 is localized in the plasma membranes (Shi et al., 2000).

From the complete nucleotide sequence of a cyanobacterium Synechocystis sp. 6803, it was suggested that the Synechocystis sp. 6803 contains at least five Na\textsuperscript{+}/H\textsuperscript{+} antiporter genes (Kaneko et al., 1996). However, their functional characterization has never been reported. During the homology search, we found that two genes (synnhaP and synnhaP2) are highly homologous to the NhaP from P. aeruginosa. Moreover, it was suggested that the Asp-138 in SynNhaP is conserved in eucaryotic Na\textsuperscript{+}/H\textsuperscript{+} exchangers and the C-terminal hydrophilic tail in SynNhaP is much longer than that of the NhaP. Therefore, we isolated one of two nhaP genes whose deduced amino acid sequence is more homologous to NhaP and SOS1. We constructed the Asp-138 mutants as well as the C-terminal deleted mutant and expressed it in the E. coli mutant. Membrane vesicles prepared from the E. coli mutant transformed with the SynNhaP exhibited the Na\textsuperscript{+}/H\textsuperscript{+} antiporter activities, and their activities were insensitive to amiloride. Moreover, its activity was very high between pH 5 and 9. The replacement of aspartate-138 in SynNhaP with glutamate or tyrosine inactivated the SynNhaP antiporter activity. The deletion of a part of the long C-terminal hydrophilic tail significantly inhibited the antiporter activity. A topological model suggests that aspartate-138 in SynNhaP is conserved in NhaP, SOS1, and AtNHX1 and is involved in the exchange activity. Thus, it appeared that the SynNhaP would provide a model system for the study of structural and functional properties of eucaryotic Na\textsuperscript{+}/H\textsuperscript{+} antiporters.
coli TO114 cells in which nhaA, nhaB, and chaA genes were deleted (Ohyama et al., 1994; Enomoto et al., 1998). SynNhaP complemented the salt-sensitive phenotype of the E. coli mutants and its Asp-138 and C-terminal tail are important for the Na+/H+ antiporter activity.

RESULTS

Isolation of SynNhaP Gene

The homology search revealed that Synechocystis sp. PCC 6803 did not contain the genes homologous to the nhaA, nhaB, and chaA from E. coli. In contrast, two genes (accession nos. D90910 and D90914) homologous to nhaP from P. aeruginosa were found. The former gene encodes a polypeptide of 527 amino acid residues (SynNhaP), whereas the latter one encodes a polypeptide of 540 amino acid residues (SynNhaP2) as shown in Figure 1A. Two deduced proteins showed high homology to NhaP from P. aeruginosa (approximately 33%–34% identity in amino acids), and SOS1 from Arabidopsis (31%–34%) in the corresponding regions, but slightly lower similarities to NhaP and SOS1 from Arabidopsis (31%–34%) in the corresponding regions, but slightly lower similarities to the human NH1 (29%–30%) and the organellar Na+/H+ antiporters such as AtNHX1 and NHE6 (28%–30%). Because the deduced amino acid sequence of SynNhaP showed higher similarities to SOS1 and NhaP than that of SynNhaP2 (Fig. 1A and B), we examined the isolation of synnhaP gene from Synechocystis sp. PCC 6803 by the PCR technique. The nucleotide sequence of the isolated gene completely coincided with that reported (Kaneko et al., 1996). As shown in Figure 1A, the Asp-138 in SynNhaP was coincided with that reported (Kaneko et al., 1996). As a nucleotide sequence of the isolated gene completely coincided with that reported (Kaneko et al., 1996). As a consequence of these studies, the SynNhaP contained a longer hydrophilic C-terminal tail than NhaP from P. aeruginosa.

Expression of Wild-Type and Mutant SynNhaP Antiporters in E. coli

As described in “Materials and Methods,” the isolated synnhaP gene was ligated into the pTrc-His2C plasmid. The resulting plasmid, pSNhaP, encodes the SynNhaP fused in frame to six histidines. The D138E, D138Y, and C-terminal deleted mutants were expressed by using the plasmids, pSNhaPD138E, pSNhaPD138Y, and pSNhaPDAC, all of which retained the His tag. Due to the absence of Na+/H+ antiporter genes (nhaA, nhaB, and chaA) in E. coli host cells (TO114), the host cell itself could not grow in the presence of 0.2 m NaCl at pH 8.

To examine whether the SynNhaP catalyzes the exchange between Na+ and H+ across membranes, the E. coli TO114 cells were transformed with pTrc-His2C, pSNhaP, pSNhaPD138E, pSNhaPD138Y, and pSNhaPDAC. After expression of these genes, the membrane fractions were isolated and subjected to SDS-PAGE and immunoblotting analysis with the antibody raised against the 6×-His tag. The E. coli cells transformed with pSNhaP, pSNhaPD138E, and pSNhaPD138Y exhibited a single cross-reaction band corresponding to approximately 53 kD (Fig. 2). As expected from its shorter C-terminal tail, the pSNhaPDAC mutant migrated faster than other SynNhaP antiporters. The E. coli cells transformed with the vector alone did not show any cross-reaction band. The accumulation levels of wild-type and mutant SynNhaP were similar as shown in Figure 2. These results indicate that both the SynNhaP and its mutants could be expressed and assembled in E. coli membranes.

Complementation of Salt-Sensitive E. coli Cells

Figure 3 shows the growth curves of E. coli TO114 cells transformed with pTrc-His2C, pSNhaP, pSNhaPD138E, pSNhaPD138Y, and pSNhaPDAC. All these E. coli cells showed similar growth rates in LBK medium, in which NaCl in the original L broth was replaced by KCl (87 mM), at pH 7.0 (Fig. 3A). However, the E. coli cells transformed with pTrc-His2C, pSNhaPD138E, and pSNhaPD138Y could not grow during the 24-h incubation in the presence of NaCl at 200 mM or higher concentration in LBK medium (Fig. 3, B and D). In contrast, the E. coli cells transformed with pSNhaP and pSNhaPDAC could grow even in the presence of 200 mM NaCl, although their growth rates were decreased with increasing concentrations of NaCl (Fig. 3, B and D). E. coli cells transformed with pSNhaPDAC showed lower growth rates than the cells transformed with pSNhaP. These results indicate that the His-tagged SynNhaP could function as Na+/H+ antiporter in E. coli cells and the replacement of Asp-138 with Glu or Tyr abolished the complementation ability of SynNhaP. The deletion of hydrophilic C-terminal 56 amino acid residues partially inhibited the complementation ability of SynNhaP.

Similar results were obtained for the SynNhaP, SynNhaPD138E, and SynNhaPD138Y grown in the LBK medium containing LiCl as shown in Figures 3, C and E. However, different effects were observed in SynNhaPDAC. As shown in Figure 3E, SynNhaPDAC could not complement the LiCl-sensitive phenotype of E. coli mutant during the 11-h incubation in the presence of 4 mM LiCl. However, the E. coli cells transformed with pSNhaPDAC started to grow after 24 h of incubation (Fig. 3E).

Essentially similar results were obtained when the growth medium was pH 8.0, although a lower salt concentration must be used due to the increased sensitivity of E. coli mutant to salt at an alkaline pH (Ohyama et al., 1994). These results suggest that SynNhaP could catalyze the exchanges between Na+ and H+ and also between Li+ and H+, that Asp-138 is essential for the antiporter activity, and that the hydrophilic C-terminal 56 amino acid residues are...
important for LiCl tolerance but have minor effects on NaCl tolerance.

**Na\(^+\)/H\(^+\) Antiporter Activity in the Everted Membrane Vesicles**

To examine directly the antiporter activity of wild-type and mutant SynNhaP, the everted membrane vesicles were prepared and their antiporter activities were monitored by measuring the dequenching of acridine orange fluorescence upon addition of NaCl or LiCl. As shown in Figure 4A, the dequenching of fluorescence was observed upon the addition of NaCl in the SynNhaP-expressing cells, but not in the control (pTrc-His2C) cells, indicating that SynNhaP has Na\(^+\)/H\(^+\) antiporter activity. Multiple sequence alignment and generation of phylogenetic tree were performed with ClustalW and TreeView software, respectively.

**Figure 1.** A, Alignment of the deduced amino acid sequences of Na\(^+\)/H\(^+\) antiporters from seven organisms. The sequences were aligned by the program ClustalW. The alignment is based on the N-terminal 550 amino acid residues in the cases of SOS1, human NHE1, and yeast NHX1. The amino acid residues conserved in all sequences are highlighted in black and conservative substitutions are shown in gray. The conserved Asp (Asp-138 in SynNhaP) is shown by an asterisk. Predicted membrane spanning regions were marked above the alignment. B, Phylogenetic analysis of seven Na\(^+\)/H\(^+\) antiporters. Multiple sequence alignment and generation of phylogenetic tree were performed with ClustalW and TreeView software, respectively.

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show that Li+ could be replaced with Na+, whereas K+ or Ca2+ or Mg2+ could not. It is known that amiloride inhibits the activity of Na+/H+ antiporters from animals (Orlowski, 1993), plants (Blumwald et al., 1987), and some bacteria (Pinner et al., 1995). Figure 4D shows the effects of amiloride on the activities of SynNhaP. SynNhaP was insensitive to amiloride both in Na+/H+ and Li+/H+ exchange activities.

Na+/H+ Antiporter Activity of SynNhaP Mutants

Figure 4B shows that the replacement of Asp-138 with Glu-138 or Tyr-138 abolished the Na+/H+ and Li+/H+ exchange activities of SynNhaP, indicating the importance of Asp-138 for the antiporter activity of SynNhaP.

Figure 4B also shows that SynNhaPΔC had very low Na+/H+ and Li+/H+ antiporter activities. This low activity presumably partially complemented the NaCl-sensitive phenotype of the E. coli mutants but not the LiCl-sensitive phenotype.

Effects of pH on the Na+/H+ Antiporter Activity

Next, we examined the effects of pH on the antiporter activity of the SynNhaP. Although the respiration-driven fluorescence quenching of acidine orange was decreased when the reaction mixture was made acidic (pH 5), the fluorescence dequenching upon addition of NaCl occurred to an extent similar to that obtained at pH 7.0. Upon addition of NH4Cl, the fluorescence intensity recovered almost to the original level (data not shown). Thus, it appeared that the SynNhaP has high Na+/H+ antiporter activities at an acidic pH (pH 5) as shown in Figure 5. Essentially similar results were obtained at an alkaline pH of 9, indicating that the SynNhaP had very high Na+/H+ antiporter activities over a wide range of pH between 5 and 9, which is quite different from those of E. coli NhaA. In E. coli NhaA, the antiporter activity could not be observed below pH 7.5, whereas the activity increased with increasing pH (Padan and Schuldiner, 1996). The SynNhaP exhibited similar pH dependences for the Na+/H+ and Li+/H+ exchange activities (Fig. 5).

The SynNhaPD138E and SynNhaPD138Y mutants had no Na+/H+ and Li+/H+ exchange activities at pH values lower than 8.0, but they showed small activities at an alkaline pH as shown in Figure 5.

Topological Model for SynNhaP

From the analysis of the hydropathy plot (Kyte and Doolittle, 1982) and the transmembrane (TM) prediction program (Hofmann and Stoffel, 1992) of the SynNhaP sequence, we predicted that the SynNhaP has 11 putative transmembrane segments (Fig. 6). The amino acid sequence of SynNhaP shows that it has 49 negative and 34 positive charges. Among them, three negative and two positive charges were located in hydrophobic segments, and only the Asp-138 was conserved in SynNhaP, NhaP, SOS1, At-NHX1, OsNHX1, NHX1, and mammalian NHE1. In this model, 28 negative and 23 positive charges were located in the cytoplasmic and 18 negative and nine positive charges were located in the periplasmic face of the membrane. Although the model is consistent with the so-called “positive inside rule” (von Heijne and Gavel, 1988) and the recently proposed one for human NHE1 (Wakabayashi et al., 2000), this is a tentative one and further study is required to obtain the insight on the structure of membrane spanning domains. Nevertheless, this model together with the experimental results presented in the above suggest that Asp-138 is involved in the exchange between Na+ and H+.

DISCUSSION

The data presented above clearly indicate that a putative eucaryotic type antiporter SynNhaP from Synechocystis sp. PCC 6803 had exchange activities between Na+ and H+ as well as Li+ and H+. This conclusion was based on the finding that the antiporter-deficient E. coli TO114 mutant cells became salt-tolerant by transformation with its gene and also by the direct observation of Na+/H+ as well as Li+/H+ antiporter activities in the transformant membrane vesicles. The most striking functional feature of
SynNhaP is that its activity was observed over a wide pH range between pH 5 and 9. In addition, it was shown that the Asp-138 in SynNhaP was essential for the exchange between Na\(^{+}\) or Li\(^{+}\) with H\(^{+}\).

To date, only a few functional residues have been identified in antiporter proteins. Nothing is known about the NhaP antiporter. Although the importance of amino acid residues involving the amiloride-binding domain and glycosylation site have been reported in the mammalian NHE exchanger (Orlowski, 1993; Orlowski and Grinstein, 1997), the importance of Asp in the hydrophobic region has not been reported in eucaryotic Na\(^{+}\)/H\(^{+}\) antiporters. Figure 1A suggests that the Asp-138 in SynNhaP corresponds to the Asp-238 in NHE1, but the mutagenesis study of Asp-238 in NHE1 has not been reported. In NhaA, the importance of Asp-133, -163, and -164 in the hydrophobic TM segments has been demonstrated (Nakamura et al., 1994; Inoue et al., 1995; Nakamura et al., 1995). Consistent with essentially no homology between SynNhaP and NhaA, the SynNhaP contains only one Asp in the hydrophobic TM segments. However, the local homology in the vicinity of Asp-138 in the SynNhaP and Asp-133 in

![Figure 1A](image1.png)

**Figure 1A.** Effects of NaCl and LiCl on the growth rates of various kinds of *E. coli* cells. The control and transformant cells at logarithmic phase in LBK medium were subjected to salt stress by inoculation into fresh LBK medium containing indicated concentrations of NaCl or LiCl. A, Time courses of growth in LBK medium. B, Time courses of growth in LBK containing 0.1 or 0.2 M NaCl. C, Time courses of growth in LBK containing 1 or 4 mM LiCl. D, Growth after 11 or 24 h incubation with various kinds of concentrations of NaCl. E, Growth after 11 or 24 h incubation with LiCl of various concentrations. ●, Control cells; ○, SynNhaP-expressing cells; ■, SynNhaPD138E-expressing cells; □, SynNhaPD138Y-expressing cells; ▲, SynNhaPD133E-expressing cells. Each value shows the average of three independent measurements (SE was within 15%).
NhaA has been suggested. This and the present data imply that Asp-138 might be involved in the exchange of cations, which must be verified experimentally. The present data also indicate that the replacement of Asp-138 with Glu abolished the Na\(\text{I}^+\)/H\(\text{I}^+\) exchange activity. This suggests that the negative charge on Asp-138 is not sufficient for the exchange activity and the geometry in the vicinity of Asp-138 plays an important role.

It has been reported that NhaP from *P. aeruginosa* (Utsugi et al., 1998) has very low activity of exchange between Li\(\text{I}^+\) and H\(\text{I}^+\). Therefore, NhaP from *P. aeruginosa* could not complement the LiCl sensitivity of *E. coli* TO114 mutant cells although NhaA, NhaB, and NhaD could (Utsugi et al., 1998). These properties of NhaP are different from those observed in SynNhaP. The Li\(\text{I}^+\)/H\(\text{I}^+\) antiporter activity of SynNhaP was almost the same as that of Na\(\text{I}^+\)/H\(\text{I}^+\) antiporter (Figs. 4 and 5) and SynNhaP could confer the resistance not only to NaCl but also to LiCl stress (Fig. 3). At the present time, the molecular mechanisms for the different ion specificity between SynNhaP and NhaP remain to be clarified.

The different pH dependence of the antiporter activities between *E. coli* (NhaA) and *Synechocystis* sp. 6803 (SynNhaP) might be interesting to note. It is clearly shown that the point mutations at His-225 of *E. coli* NhaA caused altered responses of the antiporter activity to pH changes. It will be interesting to examine which amino acid residue(s) in the SynNhaP are involved in the constant activity over the wide pH range.

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**Figure 4.** Na\(\text{I}^+\)/H\(\text{I}^+\) antiporter activity measured by the acridine orange fluorescence quenching method. The control *E. coli* cells and *E. coli* cells expressing SynNhaP, SynNhaPD138E, SynNhaPD138Y, and SynNhaPD138C were grown in LBK medium without addition of extra salts. From these cells the everted membrane vesicles were prepared. A, Typical fluorescence traces. At the time indicated by downward arrows, Tris-lactate (final concentration 2 mm) was added to initiate the respiration induced fluorescence quenching. Then at the time indicated by upward arrows, salts (NaCl, LiCl, KCl, CaCl\(\text{II}\), MgCl\(\text{II}\) were added. The final salt concentration was 5 mm. Then NH\(\text{I}_4^+\) (final concentration 25 mm) was added to the assay mixture. The assay mixture was pH 8.0. The dequenching (%) was calculated from the ratio of NaCl-induced dequenching/lactate-induced quenching. B, NaCl- or LiCl-induced dequenching (%). C, KCl-, MgCl\(\text{II}\)-, or CaCl\(\text{II}\)-induced dequenching (%). D, Effects of amiloride on the NaCl- or LiCl-induced dequenching. Each value shows the average of three independent measurements (SE was within 8%).
that of NhaP and *E. coli* antiporters (Fig. 6). In animals, the long C-terminal hydrophilic tails are believed to play a role in the regulation of transport activity (Wakabayashi et al., 1992; Orłowski and Grinstein, 1997). The present results also suggest that the C-terminal hydrophilic tail might interact with the membrane segments and play an important role for the transport activity. Further studies are required to clarify these points.

The recently discovered SOS1 antiporter has been shown to be essential for Na\(^+\) and K\(^+\) homeostasis and its gene expression is up-regulated in response to NaCl stress (Shi et al., 2000). SOS1 was predicted as a 127-kD protein with 12 transmembrane domains in the N-terminal part and a long hydrophilic cytoplasmic tail in the C-terminal part. Since the transmembrane region of SynNhaP and SOS1 has significant sequence similarities, the functional and structural analysis of SynNhaP might also contribute to the understanding of SOS1 antiporter. Construction of the SynNhaP mutants and expression of a large amount of antiporter should help us to study the structural and functional properties of this important protein at the molecular levels.

**MATERIALS AND METHODS**

Isolation of nhaP Gene

The *nha*P gene from *Synechocystis* sp. 6803 was amplified by PCR using the following primers: forward primer, 5'-CACCATGGATA CAGCGGTCAACG-3'; reverse primer, 5'-TCGAAATTCCGGATGTTCGGCCACAT-3'. The forward primer contains the Ncol restriction site. The reverse primer contains the EcoRI restriction site just before the stop codon. The amplified fragment was ligated into the Ncol/EcoRI sites of the pTrc-His2C plasmid. The resulting plasmid, pSNhaP, encodes the SynNhaP fused in frame to six histidines, and was used to transform *Escherichia coli* TO114 cells. The DNA sequence was determined using an ABI310 genetic analyzer (PE-Applied Biosystems, Foster City, CA) and analyzed with the DNASTAR program (Hitachi Software Engineering, Tokyo).

Construction of SynNhaP Mutants

The site-directed mutagenesis of Asp-138 was carried out by using a PCR method (Ito et al., 1991). The 412 to 414 bases, GAT, encoding Asp-138 of SynNhaP were changed to GAA and TAT, which generated the plasmids pSNhaPD138E and pSNhaPD138Y, respectively. The change of nucleotide was confirmed by DNA sequencing. For the construction of C-terminal tail deleted mutant, the pSNhaP plasmid was digested by the restriction enzymes PstI and EcoRI, blunted ended, and ligated. The resulting plasmid, pSNhaPDC, did not contain the last 56 amino acid residues (Leu-472 to Ser-527), but retained the His-tag derived amino acid residues.

Growth of *E. coli* Cells

*E. coli* TO114 cells, in which *nhaA*, *nhaB*, and *chaA* were deleted, were used as the host cells. Cells were grown in LBK medium at 37°C under aerobic conditions. The pH of the growth medium was adjusted with KOH or HCl. The growth of cells was monitored by measuring the A\textsubscript{620} with a Erma AE-22 photoelectric colorimeter.

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**Na⁺/H⁺ Antiporter Activity**

The Na⁺/H⁺ antiporter activity was examined on everted membrane vesicles prepared from the cells grown in LBK as described (Rosen, 1986). *E. coli* cells were harvested by centrifugation at 3,100 g for 10 min at 4°C and then washed with a suspension buffer TCDS, which contained 10 mM Tris-HCl, pH 7.0, 0.14 M choline chloride, 0.5 mM dithiothreitol, and 0.25 M Suc. The pellets were suspended with 10 mL of TCDS buffer and applied to a French Pressure cell (4,000 psi). Then the solution was centrifuged at 12,000 g for 10 min at 4°C. The supernatant was centrifuged at 110,000 g for 60 min at 4°C and suspended in 600 μL of TCDS buffer. The antiporter activity was based upon the establishment of ΔpH (transmembrane pH gradient) by addition of salt to the reaction mixture that contained 10 mM Tris-HCl (titrated with HCl to the indicated pH), 5 mM MgCl₂, 0.14 M choline chloride, 1 mM acridine orange, and membrane vesicles (50 μg of protein) in a volume of 2 mL. The ΔpH was monitored at 25°C with acridine orange as a probe at an extinction wavelength of 492 nm (band width 1.5 nm) and emission wavelength of 525 nm (band width 3.0 nm) of Shimadzu RF-5300PC spectrofluorophotometer (Hibino et al., 1995). At the onset of the experiment, Tris-β-lactate (2 mM) was added and the fluorescence quenching was recorded. Salt (5 mM) was then added and the new steady state of fluorescence obtained (dequenching) after each addition was monitored.

**Immunoblotting and Other Methods**

SDS-PAGE and immunoblotting were carried out as previously described (Lee et al., 1995; Nomura et al., 1995). Protein was determined by the method of Lowry et al. (1951). An antibody raised against 6-His (6×-His tag) was obtained from R & D Systems (Minneapolis).

**Computer Analysis**

The hydropathy profile of the deduced amino acid sequence of the SynNhaP was predicted by the computer-assisted procedure according to the method of Kyte and Doolittle (1982). The possible TM of the SynNhaP sequence...
was deduced by a computer program TopPredII (Hofmann and Stoffel, 1992).

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