

**METHODOLOGY ARTICLE**

**Open Access**

# Outer membrane proteins can be simply identified using secondary structure element alignment

Ren-Xiang Yan<sup>1,2</sup>, Zhen Chen<sup>1,2</sup> and Ziding Zhang<sup>1,2\*</sup>

## Abstract

**Background:** Outer membrane proteins (OMPs) are frequently found in the outer membranes of gram-negative bacteria, mitochondria and chloroplasts and have been found to play diverse functional roles. Computational discrimination of OMPs from globular proteins and other types of membrane proteins is helpful to accelerate new genome annotation and drug discovery.

**Results:** Based on the observation that almost all OMPs consist of antiparallel  $\beta$ -strands in a barrel shape and that their secondary structure arrangements differ from those of other types of proteins, we propose a simple method called SSEA-OMP to identify OMPs using secondary structure element alignment. Through intensive benchmark experiments, the proposed SSEA-OMP method is better than some well-established OMP detection methods.

**Conclusions:** The major advantage of SSEA-OMP is its good prediction performance considering its simplicity. The web server implements the method is freely accessible at <http://protein.cau.edu.cn/SSEA-OMP/index.html>.

## Background

Outer membrane proteins (OMPs), an important class of proteins, are found in gram-negative bacteria, mitochondria and chloroplasts. Computational discrimination of OMPs from globular proteins and other types of membrane proteins is helpful to accelerate new genome annotation and drug discovery. A variety of OMP identification methods have been elegantly developed [1-27] and some web servers have also been freely accessible to the research community [2,7,13,16,21,27]. However, OMPs are difficult to be discriminated from other types of proteins, and the existing methods are not entirely satisfactory, mainly because the membrane-spanning regions of OMPs are shorter and these regions usually have higher variations in properties when compared with  $\alpha$ -helical membrane proteins [28]. Therefore, the development of new OMP identification methods with improved performance is needed. Meanwhile, it is also hoped that new OMP identification methods will be

helpful to accelerate the exploration of the sequence-structure protein landscape in OMPs.

The existing OMP predictors can be categorized through different ways. According to the adopted algorithms, the predictors can be divided into simple statistical theory- and machine learning-based methods. The highlight of simple statistical methods is that the biological meanings of the established statistical models are comprehensible. Representative statistical theory-based OMP predictors include DD [15], WED [17], WED\_HFS [17], BOMP [2] and TMB-Hunt [29]. The advantage of machine learning algorithms is that they can easily incorporate different information/features into an OMP discrimination system. Some state-of-the-art machine learning algorithms (*e.g.*, Neural Network (NN) and Support Vector Machines (SVM)) have been employed to construct OMP discrimination systems [5,6]. Using amino acid composition as input, Gromiha and Suwa (2006) developed a NN-based method (NN\_AAC) to identify OMPs [5]. Later, they also proposed an SVM-based OMP identification method (SVM\_AAC\_DPC) by adding di-peptide composition information as input, which was reported to have improved performance [6]. Although machine learning algorithms can often result

\* Correspondence: [zidingzhang@cau.edu.cn](mailto:zidingzhang@cau.edu.cn)

<sup>1</sup>State Key Laboratory of Agrobiotechnology, College of Biological Sciences, China Agricultural University, Beijing 100193, PR China  
Full list of author information is available at the end of the article

in excellent performance, some of them are often criticized and labeled as “black box” methods, due to a lack of biological interpretation.

According to the type of input data, the previous predictors can be roughly divided into global amino acid composition- and local sequence features-based methods. For instance, DD, WED and TMB-Hunt mainly used the global amino acid composition as input to identify OMPs. As a typical local sequence features-based method, BOMP employs C-terminal pattern recognition and a sliding window analysis of amino acid composition in alternating positions to identify OMPs. In 2005, Gromiha also proposed a local sequence features-based method, which used frequently occurring motifs to predict OMPs [11]. Moreover, the established OMP predictors can be grouped in terms of the prediction output type. Many OMP predictors such as DD, WED, WED\_HFS, NN\_AAC, TMB-Hunt and BOMP were limited to distinguish OMPs and non-OMPs, while a few OMP predictors (*e.g.*, PRED-TMBB [20,21], TMBETA-NET [23], TBBPred [22] and PROFtmb [30]) were also able to detect transmembrane  $\beta$ -strand regions in query sequences. The existing predictors could also be classified according to whether they use sequence evolutionary information or not. PRED-TMBB, DD, NN\_AAC and SVM\_AAC\_DPC are typical methods that do not use sequence evolutionary information, while profile-based methods such as PROFtmb, HHomp are heavily relied on the use of evolutionary information. PROFtmb is a profile-based Hidden Markov Model (HMM) for the prediction of transmembrane  $\beta$ -strand regions and discrimination between OMPs and non-OMPs. To predict whether a query sequence is an OMP, HHomp builds a profile HMM for a query sequence and compares it with an OMP database by pairwise HMM comparison (*i.e.*, HMM-HMM matching).

Based on the observation that an OMP usually contains 8-24 antiparallel  $\beta$ -strands that form a barrel shape, we propose a simple method to identify OMPs using secondary structure element alignment (SSEA). Predicted secondary structure has long been known to be informative in protein structure prediction and classification. SSEA was initially proposed by Przytycka *et al.* and used as a protein taxonomy method [31]. Subsequently, SSEA has been employed as an important component for fold recognition methods such as MANIFOLD [32], GenTHREADER [33,34] and DescFold [35,36]. SSEA was also used to target new folds [37]. In our previous work, the alignment score resulting from SSEA was used as a descriptor to detect TIM-barrel proteins [38]. Additionally, Fontana and co-workers developed a web server that implements the SSEA algorithm, which is freely accessible at <http://protein.cribi.unipi.it/ssea/>[39].

It should also be clearly pointed out that the information of predicted secondary structure has been incorporated into some OMP predictors [7,8,19]. For instance, Liu and co-workers (2003) [8] found that the transmembrane  $\beta$ -strand regions in OMPs have different amino acid composition compared with the  $\beta$ -strands of globular proteins. Therefore, they proposed a predictor based on the composition of selected amino acids (*i.e.*, Gly, Val, Ile, Lys, Cys and Asn) in predicted  $\beta$ -strands to identify OMPs.

In this study, we explore a new application of SSEA by proposing an SSEA-based OMP detection method (SSEA-OMP). The proposed method was intensively tested against well-established OMP discrimination methods and found to be very competitive, suggesting that it can serve as a useful tool to assist in proteome-wide OMP annotation.

## Results and discussion

### Overall performance of SSEA-OMP

Based on the GS-dataset [15], the overall performance of SSEA-OMP was assessed by a Leave-One-Out (LOO) test. It is worth mentioning that some protein pairs in the GS-dataset may share more than 25% sequence identity. More detailed information of GS-dataset can refer to Methods section. To avoid performance overestimation caused by redundant proteins, we employed a stringent sequence-filtering method in each step of the LOO test, which can be divided into two procedures. In the first procedure, significant sequence similarity between the test protein and the library can be removed. Briefly, only the remaining sequences sharing a sequence identity (global alignment mode inferred from the algorithm of Needleman and Wunsch [40]) less than 25% and a BLAST *e*-value greater than 0.01 with the test protein were kept in the sequence library. Even with the above filtering procedure, the sequence profiles of the remaining sequences in the library, which were required by PSIPRED [41], may still contain some sequence members sharing significant homology with the test protein. Therefore, in the second filtering procedure we further filtered the remaining sequences by using the PSI-BLAST [42]*e*-value. Briefly, all the sequence profiles of the remaining sequences in the library were jumpPSI-BLASTed against the test protein. Only the remaining sequences with a PSI-BLAST *e*-value greater than 0.01 with the test protein were further kept. Based on the above two procedures, we can guarantee that the sequence similarity between the test protein and the filtered library should be very low.

As shown in Table 1, SSEA-OMP resulted in an excellent performance. Four measurements, *i.e.*, Accuracy (*Ac*), Sensitivity (*Sn*), Specificity (*Sp*) and Matthew correlation coefficient (*MCC*), were jointly used to assess

**Table 1 Performance of different OMP discrimination methods based on the GS-dataset**

Method	MCC	Ac (%)	Sn (%)	Sp (%)
DD <sup>a,b</sup>	0.541	82.4	78.8	83.3
NN_AAC <sup>a,b</sup>	0.716	91.0	79.3	93.8
SVM_AAC_DPC <sup>a,b</sup>	0.816	93.9	90.9	94.7
SSEA-OMP <sup>c</sup>	0.772	90.9	72.9	98.1
SSEA-OMP <sup>d</sup>	0.906	96.2	91.5	98.1

<sup>a</sup>DD, NN\_AAC and SVM\_AAC\_DPC were developed in Suwa's group[5,6,15].

<sup>b</sup>The corresponding results are directly cited from [5,6,15].

<sup>c</sup>Based on the stringent sequence-filtering method. Briefly, only the remaining sequences sharing a sequence identity less than 25%, a BLAST e-value greater than 0.01 and a PSI-BLAST e-value greater than 0.01 with the test protein were kept in the sequence library.

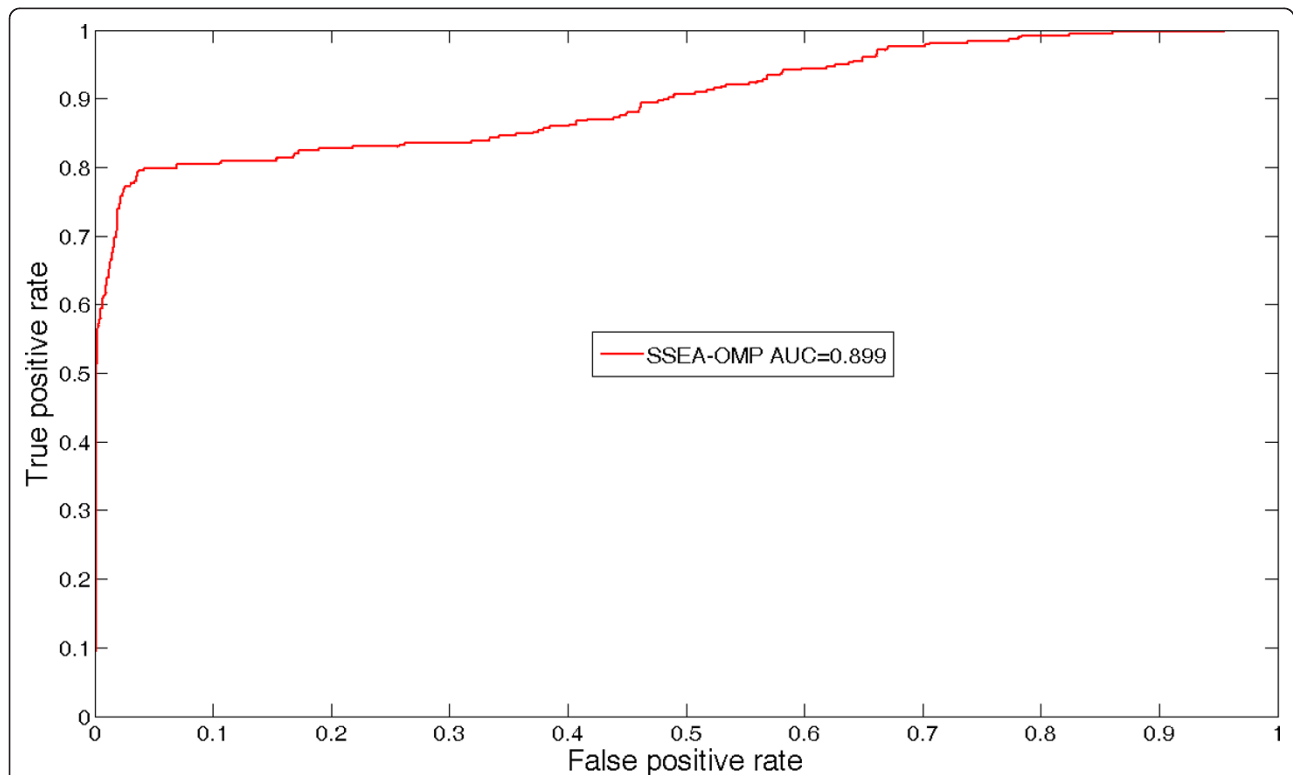
<sup>d</sup>Only the first sequence-filtering procedure was employed. Briefly, only the remaining sequences sharing a sequence identity less than 25% and a BLAST e-value greater than 0.01 with the test protein were kept in the sequence library. It should be emphasized that the performance of SSEA-OMP based on the first sequence-filtering procedure could be overestimated. We list the SSEA-OMP performance based on the first sequence-filtering procedure to allow a generally fair comparison between SSEA-OMP and the other three methods, since the performance of the other three methods were characterized by simple sequence identity-based filtering procedure [5,6,15].

the performance of different methods. In general, *MCC* is a more suitable parameter than *Ac* for assessing the two-class prediction when the numbers of samples in the two classes are not equal. The value of *MCC* was therefore considered the main measurement by which to

assess the performance of SSEA-OMP in this work. For instance, the proposed SSEA-OMP can distinguish between OMPs and non-OMPs with an *MCC* value of 0.772 (*Ac* = 90.9%, *Sn* = 72.9% and *Sp* = 98.1%). The overall performance of SSEA-OMP was further measured by an ROC curve. As can be seen in Figure 1, SSEA-OMP yields an AUC score (*i.e.*, the area under the ROC curve) of 0.899. The performance of SSEA-OMP at low false positive rates is also impressive. For instance, SSEA-OMP was able to correctly identify 62.9% OMPs at a less than 1% false positive rate (Figure 1). For comparison, the performance using only the first sequence-filtering procedure is also shown in Table 1. When only the first procedure was adopted, the *MCC* value of the proposed SSEA-OMP can be dramatically increased to 0.909 (*Ac* = 96.2%, *Sn* = 91.5% and *Sp* = 98.1%), implying the results are strongly affected by the sequence-filtering method.

#### Comparison with existing OMP discrimination methods

In this work, SSEA-OMP was intensively tested against some existing predictors. To make a fair and comprehensive comparison, we tested SSEA-OMP using two established benchmark datasets (the GS-dataset and the R-dataset [7]). Because the performance of some existing methods using these two datasets has been reported in



**Figure 1 ROC curve of SSEA-OMP assessed using the GS-dataset.** The overall performance of SSEA-OMP was assessed by the Leave-One-Out (LOO) test. In each step of the LOO test, only the remaining sequences sharing a sequence identity less than 25%, a BLAST e-value greater than 0.01 and a PSI-BLAST e-value greater than 0.01 were kept and used as the training dataset.

the literature, the corresponding results were used to make a direct comparison with our SSEA-OMP.

Developed in Suwa's group, the DD, NN\_ACC and SVM\_ACC\_DPC methods have been benchmarked in the GS-dataset, which allows us to compare SSEA-OMP with them directly (Table 1). It should be emphasized that the performance of DD, NN\_ACC and SVM\_ACC\_DPC was evaluated in the GS-dataset without any stringent sequence-filtering procedure [5,6,15]. For instance, SVM\_AAC\_DPC [6] only used CD-HIT [43] to ensure that the sequence identity between any two sequences in GS-dataset could be less than 40%. As pointed out in the previous section, the performance of SSEA-OMP is strongly affected by the sequence-filtering method. Even with the same benchmarking dataset, the comparison of different methods should ideally be conducted under the same sequence-filtering method. To have a comparatively fair assessment, we benchmarked SSEA-OMP against these three methods based on the performance of SSEA-OMP when only the first sequence-filtering procedure was used. As shown in Table 1, SSEA-OMP performs better than the DD, NN\_ACC and SVM\_ACC\_DPC methods.

SSEA-OMP was also benchmarked against HHomp based on the R-dataset. As reported by Remmert *et al.* (2009), HHomp used 486 OMP sequence clusters [ftp://ftp.tuebingen.mpg.de/pub/protevo/HHomp/db/HHompDB\\_1.0.hhm](ftp://ftp.tuebingen.mpg.de/pub/protevo/HHomp/db/HHompDB_1.0.hhm), which were derived from 23 structurally solved OMPs, as the reference dataset. Furthermore, the performance of HHomp was evaluated on the R-dataset. To allow an impartial comparison between SSEA-OMP and HHomp, we compiled 486 consensus sequences from the 486 sequence clusters and the non-OMPs in the GS dataset into a library. Thus, the comparison of SSEA-OMP and HHomp was based on the reference datasets with the same OMPs. To strictly test the R-dataset, the stringent sequence-filtering method used in the LOO test of the GS-dataset was also employed. After the sequence filter in each step of the benchmark experiment, we ensured that sequences kept in the library should share a sequence identity less than 25%, a BLAST *e*-value greater than 0.01 and a PSI-BLAST *e*-value greater than 0.01 with the test sequence in the R-dataset. In the original paper of HHomp [7], the performance of BOMP <http://www.bioinfo.no/tools/bomp>, TMB-Hunt [http://bmbpcu36.leeds.ac.uk/~andy/betaBarrel/AACompPred/aaTMB\\_Hunt.cgi](http://bmbpcu36.leeds.ac.uk/~andy/betaBarrel/AACompPred/aaTMB_Hunt.cgi) and PROFtmb [30] based on the R-dataset was also assessed by directly submitting the sequences in the R-dataset to the corresponding OMP prediction servers, which also facilitates us to compare SSEA-OMP with these peer OMP predictors. Because the performance at low false positive rates is more important for real-world application, here we pay more attention on comparing

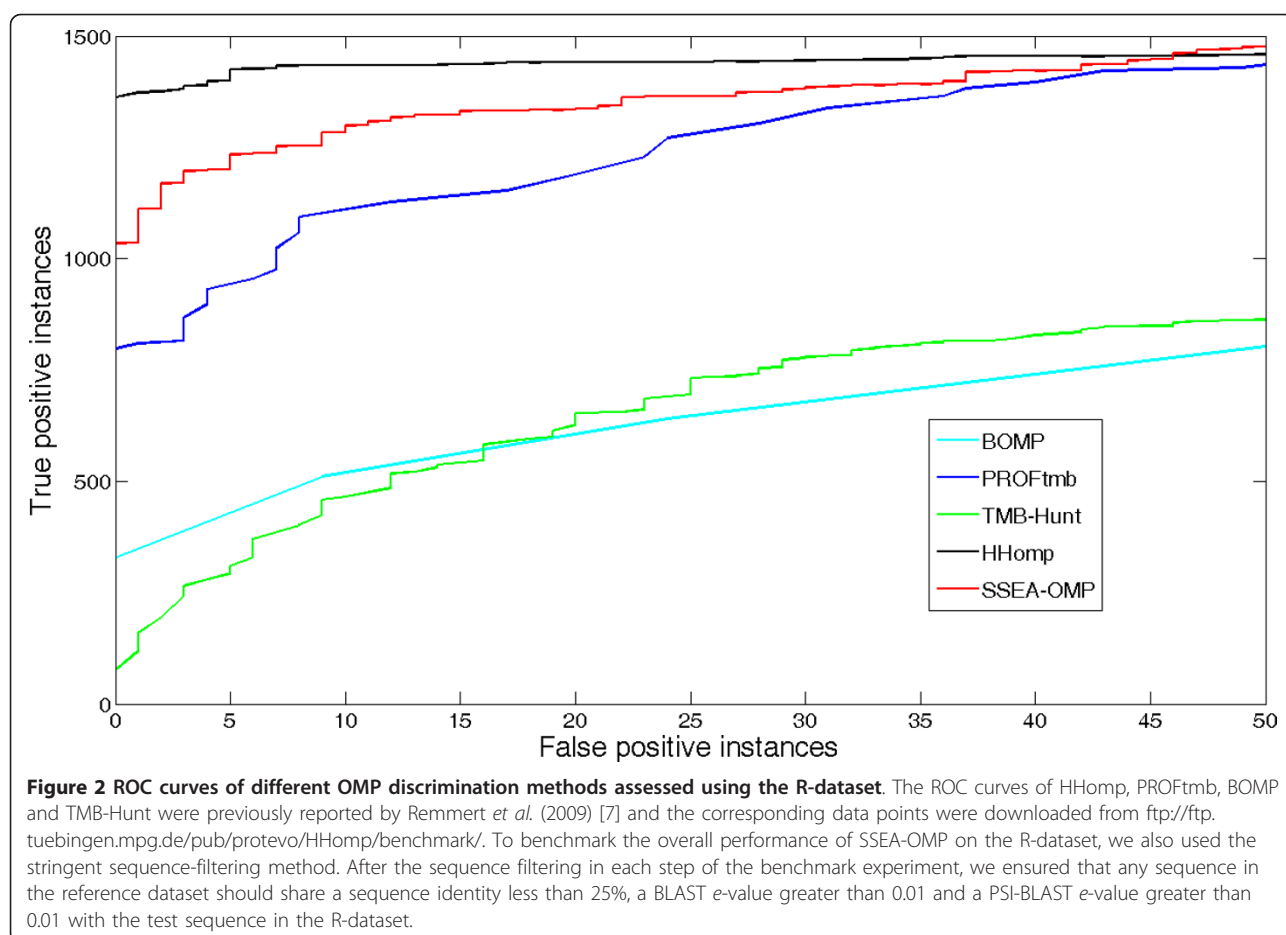
difference methods' performance at low false positive rates. For instance, SSEA-OMP correctly recognizes 1036 OMPs before including the first false positive, whereas HHomp, BOMP, TMB-Hunt and PROFtmb can detect 1363, 329, 76 and 798 OMPs, respectively (Figure 2). At a less than 1% false positive rate (*i.e.*, 50 false positive instances) control, SSEA-OMP can correctly recognize 1476 OMPs, which is slightly better than the performance of HHomp and PROFtmb (1458 and 1429, respectively) and significantly higher than the correctly identified numbers of TMB-Hunt and BOMP (862 and 641, respectively) (Figure 2).

Based on the above benchmark experiments, SSEA-OMP has showed a fully comparable performance to some existing OMP predictors. Although many efforts were taken to make sure that the above benchmark experiments were intensive and strict, it should be pointed that we are still not able to guarantee a fully unbiased assessment. We take the performance comparison in the R-dataset as an example to discuss the potential biases. To obtain the performance of BOMP, PROFtmb and TMB-Hunt, Remmert *et al.* (2009) submitted the R-dataset directly to the corresponding web servers. On the one hand, some proteins in the R-dataset are likely to have been used to training the BOMP, TMB-hunt and PROFtmb servers. Therefore, the performance of these three methods might have been overestimated. On the other hand, the training/reference datasets in these three OMP predictors are not fully identical, although all of them are based on known OMPs with experimentally solved 3D structures. Generally, a training dataset covering a more complete sequence/structure space of known OMPs should result in a more favorable performance. Thus, the comparison bias among these three predictors has also been inevitably caused by the corresponding training datasets. With more and more OMP predictors available to the community, critical assessment of different predictors based on some standard training and test datasets is increasingly important. We hope such datasets will be available in the near future. Thus, different OMP predictors can be more reliably benchmarked. Meanwhile, initiatives in establishing some real-time comparison methods by following similar strategies in assessing different protein structure prediction methods (*e.g.*, Live-Bench [44] and EVA [45]) should also be helpful to advance the method development of OMP identification.

#### Benchmark experiment on $\beta$ -class globular proteins

Since all- $\beta$  globular proteins and OMPs may have high similarity in secondary structure arrangements, one may argue that all- $\beta$  proteins should have higher probability to be predicted as OMPs and this could be a limitation of SSEA-OMP. To investigate whether all- $\beta$  proteins





tends to be predicted as OMPs in our SSEA-OMP, the following benchmark experiment was carried out. We relied on the SCOP database (version 1.75) to compile a large-scale benchmarking dataset. The SCOP\_1.75\_40% dataset with a total of 10567 proteins was first downloaded from the ASTRAL website <http://astral.berkeley.edu/>, in which the sequence identity among the proteins is equal to or less than 40%. Only the sequences in the four major globular protein classes (*i.e.*, all- $\alpha$ , all- $\beta$ ,  $\alpha/\beta$  and  $\alpha+\beta$  proteins) were kept. For each fold in all- $\alpha$ ,  $\alpha/\beta$  and  $\alpha+\beta$  proteins, only one representative sequence was further kept. Thus, we compiled a dataset called SCOP\_1.75\_3000, which contains 2197 all- $\beta$  proteins and 803 non-all- $\beta$  proteins. Using the GS-dataset as the library, the SCOP\_1.75\_3000 dataset was processed by SSEA-OMP. In each step of the benchmark experiment, a BLAST sequence-filtering method was applied to ensure that only the sequences sharing a BLAST *e*-value greater than 0.01 with the test protein were kept in the library. Of the 2197  $\beta$ -class proteins, 63 proteins were predicted as OMPs ( $S_p = 97.1\%$ ). Concerning the prediction of 803 non-all- $\beta$  proteins, 16 of them were predicted as OMPs ( $S_p = 98.0\%$ ). Therefore, SSEA-OMP

does not show a tendency to generate significantly higher false positive rate in predicting all- $\beta$  proteins as OMPs, implying the secondary structure topology difference between OMPs and all- $\beta$  proteins (*e.g.*, the number of  $\beta$ -strand elements, the order of secondary structure elements and the length of each element) can be sensitively captured by SSEA.

#### The web server

To aid the research community, a web server that implements the SSEA-OMP method was constructed and is freely accessible at <http://protein.cau.edu.cn/SSEA-OMP/index.html>. The GS-dataset is used as the library for our web server. For a query sequence, the web server returns the top hit to OMPs and the corresponding SSEA similarity and prediction scores. Moreover, the alignment between secondary structural elements of the query sequence and the top hit is provided. It was estimated that a prediction score  $\geq 0.021$  yields a false positive rate of  $\leq 1\%$  (*i.e.*, 99% confidence level). Currently, a four-CPU Dell Linux machine with 16 GB of main memory hosts the web server. The multi-thread technique was employed and the

computational time required for processing a query sequence of 600 amino acids is usually not more than two minutes.

It should be pointed out that SSEA-OMP's performance is related to the selected library to some extent, which has been clearly reflected in the aforementioned two benchmark experiments. Similar to some other OMP predictors such as HHomp [7], one major limitation of SSEA-OMP is that it can only identify OMPs sharing similar secondary structure topology with known OMPs in the library. Experienced users may prefer to download our in-house SSEA algorithm <http://protein.cau.edu.cn/pdbs/SSEA.tar.gz> and use a library compiled by themselves for some proteome-wide OMP identification tasks.

#### Proteome-wide OMP identification in *Escherichia coli*

To provide a practical application of SSEA-OMP, we conducted a proteome-wide OMP identification in *E. coli*. The whole proteome of *E. coli*, which contains 4126 protein sequences, was downloaded from the NCBI website [ftp://ftp.ncbi.nih.gov/genomes/Bacteria/Escherichia\\_coli\\_K\\_12\\_substr\\_DH10B\\_uid58979/](ftp://ftp.ncbi.nih.gov/genomes/Bacteria/Escherichia_coli_K_12_substr_DH10B_uid58979/). All the *E. coli* protein sequences were directly submitted to the SSEA-OMP web server and 167 proteins were predicted to be OMPs with a false positive rate control of 1% (Additional file 1).

To assess the performance of SSEA-OMP, we collected known *E. coli* OMPs from public databases. In fact, there are 74 proteins annotated as OMPs in the downloaded *E. coli* proteome dataset. 94 *E. coli* proteins in the PSORTdb database [46] are characterized as OMPs through experimental study or computational prediction, and 69 *E. coli* proteins are also annotated as OMPs in the OMPdb database [47]. We extracted all OMP annotations from these three datasets, and compiled a total number of 120 proteins into a known *E. coli* OMP dataset. Of the 167 predicted OMPs, 80 proteins have been included in the known *E. coli* OMP dataset. Therefore, these 80 predicted OMPs should be regarded as true positives with high confidence. Since it is estimated that OMPs consist of 2-3% of the complete proteomes [48], the current OMP prediction apparently resulted in a certain number of false positives. When further searched the PSORTdb database, 55 hits are clearly annotated as non-OMPs in terms of subcellular localization information, suggesting that they are very likely to be false positives. The remaining 32 proteins, whose subcellular localizations are annotated as "unknown" or "this protein may have multiple localization sites" in the PSORTdb database, may be the potential OMPs that have not been previously discovered.

Considering the highly imbalanced numbers of OMPs and non-OMPs in a proteome, it is not surprising that

our SSEA-OMP resulted in a certain number of false positives even at a false positive rate control of 1%. In order to reduce the false positives, we may resort to other bioinformatics tools. For example, false positive predictions could be further reduced employing a signal peptide predictor (e.g., SignalP server [49]) according to the fact that most OMPs have a signal peptide [20,46]. Alternatively, we may choose the threshold value at a higher confidence level, but the identified true positives will be reduced accordingly. It should also be mentioned that 39 known *E. coli* OMPs were not successfully identified by SSEA-OMP, which may be ascribed to the fact that some of these 39 proteins share dissimilar secondary structure topology with known OMPs in the SSEA-OMP library. To maximize the performance of SSEA-OMP, a regularly-updated library which covers all sequence/structure space of known OMPs is highly desired.

#### Conclusions

Taking together these findings, we have clearly demonstrated that OMPs can be simply identified using SSEA-OMP. First, the success of SSEA-OMP should be ascribed to the facts that known OMPs have similar secondary structure topologies and the overall similarity of secondary structure topology between two OMPs can be sensitively detected by SSEA. Although predicted secondary structure has been incorporated into several existing OMP predictors [7,8,19], it should be emphasized that our SSEA-OMP utilizes the information of secondary structure in a different way. Second, the high accuracy of SSEA-OMP also implies that PSIPRED has reached a reasonably high degree of accuracy in predicting the secondary structure of OMPs, even though PSIPRED was not optimized for OMPs. Concerning future development, two aspects should be taken into account. First, optimization of the SSEA scoring scheme may help to improve the current version of SSEA-OMP. Second, SSEA can be used as a key feature in the construction of a new OMP discrimination method. It is hoped that the integration of SSEA with some other well-recognized features can result in a more powerful OMP discrimination system with the assistance of statistical or machine learning methods.

#### Methods

##### Datasets

We relied mainly on Gromiha and Suwa's dataset [15], which we refer to as the GS-dataset, to construct SSEA-OMP. The GS-dataset consists of 377 OMPs, 674 globular proteins, and 268 inner membrane proteins (IMPs). OMP discrimination can be assigned as a binary classification problem. The 377 OMPs were considered positive instances, and 942 non-OMPs (i.e., the 674 globular

proteins and 268 IMPs) were considered negative instances.

To critically benchmark SSEA-OMP against some of the existing methods, the R-dataset [7], which contains 2164 OMPs from the TransportDB database [50] and 5000 non-OMPs randomly selected from the SCOP database (version 1.69) [51], was downloaded from <ftp://ftp.tuebingen.mpg.de/pub/protevo/HHomp/benchmark/>.

The NCBI non-redundant (NR) sequence database (November 2008 version) was downloaded from <ftp://ftp.ncbi.nlm.nih.gov/blast/>. The NR database was further clustered at 90% sequence identity (global alignment mode) by using CD-hit [43], and the resulting NR90 database was used to implement the PSI-BLAST search [42].

### Construction of SSEA-OMP

To develop the SSEA-OMP method, our in-house SSEA algorithm was used. Performing an SSEA for two sequences typically consists of three steps. First, the two sequences were PSI-BLASTed against the NR90 database for three iterations to generate the corresponding PSSM profiles. The *e*-value cutoff for including sequences in the PSSM profiles was set at 0.001. The obtained PSSM profiles were further employed as input to PSIPRED [41] to predict the secondary structures of the two sequences. Second, the predicted secondary structural string for each sequence was converted into secondary structure elements such that “H” represents a helix element, “E” denotes a strand element, and “C” stands for a coil element. Thus, the predicted secondary structural string was shortened and the length of each element was retained for the scoring of SSEA. For example, the secondary structure string HHHHCCCCCEEEHHHH would be shortened to HCEH, and the lengths of all elements (*i.e.*, 4, 5, 5 and 4) would be stored. Third, the two shortened strings were aligned using a modified dynamic programming algorithm with a scoring scheme adopted from Przytycka *et al.* (1999) [31]. The detailed alignment score between two elements with lengths  $L_i$  and  $L_j$  is defined as:

$$Alignment\ Score = \begin{cases} \min(L_i, L_j) & (a) \\ 0.5 \times \min(L_i, L_j) & (b) \\ 0 & (c) \end{cases} \quad (1)$$

(a) This term is applied when match between two identical elements.

(b) This term is applied when match between helix/strand and coil.

(c) This term is applied when match between helix and strand.

where  $\min(L_i, L_j)$  stands for the minimal length between  $L_i$  and  $L_j$ . Additionally, the gap cost in SSEA is set to 0. Since the alignment score between two elements is equal to or less than  $\min(L_i, L_j)$ , the total alignment score between two sequences is equal to or less than the minimal sequence length of these two sequences. To obtain a normalized SSEA similarity score (*SSEA\_Score*), the total alignment score is further divided by the average length of these two sequences. Therefore, the *SSEA\_Score* is always in the range of 0 to 1. Generally, the closer the SSEA similarity score is to 1, the more significant is the secondary structure-based similarity between two sequences. In the original SSEA algorithm [31] both helix and strand elements can be split for alignment with coil elements, and coil elements can be split into either two or three smaller coil elements. However, Fontana *et al.* (2005) [39] argued that the partition of secondary structure elements can not obtain better alignment between two proteins. Therefore, we do not split any secondary structure element in our SSEA algorithm.

With the established SSEA algorithm, SSEA-OMP can be easily implemented by selecting a suitable sequence library (*i.e.*, reference dataset). In this work, we used the GS-dataset as the sequence library. For a query sequence, SSEA-OMP calculated the pair-wise SSEA similarity scores between the query sequence and all proteins in the GS-dataset. The top hit to OMPs and the corresponding SSEA similarity score (*i.e.*,  $SSEA\_Score_{max,OMP}$ ) were recorded. Meanwhile, the maximal SSEA similarity score between non-OMPs in the library and the query protein (*i.e.*,  $SSEA\_Score_{max,non-OMP}$ ) was also calculated. For a query sequence, a prediction score (*i.e.*, *Pred\_Score*) was defined as:

$$Pred\_Score = SSEA\_Score_{max,OMP} - SSEA\_Score_{max,non-OMP} \quad (2)$$

The query protein was predicted to be an OMP if  $Pred\_Score > 0$ ; otherwise, it was predicted to be a non-OMP.

### Performance assessment of SSEA-OMP

The performance of SSEA-OMP was assessed using the LOO test. For each step of the assessment, a protein in the GS-dataset was selected as a test protein, and the remaining proteins were regarded as a sequence library. The test protein was scanned against the sequence library using SSEA, and the prediction result was recorded. When the test was performed over all proteins in the GS-dataset, the overall performance of SSEA-OMP was evaluated with respect to four parameters: *Ac*, *Sn*, *Sp* and *MCC*. These parameters are defined below:

$$Ac = \frac{tp + tn}{tp + fn + tn + fp} \quad (3)$$

$$Sn = \frac{tp}{tp + fn} \quad (4)$$

$$Sp = \frac{tn}{tn + fp} \quad (5)$$

$$MCC = \frac{tp \times tn \times fp \times fn}{\sqrt{(tp + fp) \times (tp + fn) \times (tn + fn) \times (tn + fp)}} \quad (6)$$

where  $tp$ ,  $fp$ ,  $fn$  and  $tn$  denote true positives, false positives, false negatives and true negatives, respectively. The performance of an OMP discrimination method can also be systematically measured by a receiver operating characteristic (ROC) curve. Using a strategy similar to the one reported by Yan *et al.* (2008) [17], we defined a threshold parameter,  $\alpha$ , such that the test protein was predicted to be an OMP if  $Pred\_Score > \alpha$ . The ROC curve of SSEA-OMP plots true-positive instances as a function of false-positive instances for all possible values of  $\alpha$ . Additionally, the ROC curve can be further quantified by the AUC score, which represents the area under the ROC curve.

## Availability and requirements

**Project Name:** SSEA-OMP

**Project home page:** <http://protein.cau.edu.cn/SSEA-OMP/index.html>

**Operating system:** Online service is web based; local version of the software is platform independent.

**Programming language:** Perl and Java.

**Other requirements:** None.

**License:** Free.

**Any restrictions to use by non-academics:** None.

## Additional material

**Additional file 1:** is a text file showing the predicted 167 *E. coli* OMPs.

## Acknowledgements

We thank the anonymous referees whose constructive and insightful comments were very helpful in improving the quality of this work. We are also grateful to Dr. Johannes Söding and Mr. Michael Remmert at Ludwig-Maximilians-University (Germany), Drs. Michael M. Gromiha and Makiko Suwa at the National Institute of Advanced Industrial Science and Technology (Japan) for sharing their datasets. This work was supported by grants from the National Key Basic Research Project of China (2009CB918802) and the State High Technology Development Program (2008AA02Z307).

## Author details

<sup>1</sup>State Key Laboratory of Agrobiotechnology, College of Biological Sciences, China Agricultural University, Beijing 100193, PR China. <sup>2</sup>Bioinformatics Center, College of Biological Sciences, China Agricultural University, Beijing 100193, PR China.

## Authors' contributions

RXY wrote the codes, developed the web server, and drafted the manuscript. ZC participated in the method assessment. ZZ directed the research and revised the manuscript. All authors read and approved the final manuscript.

Received: 2 August 2010 Accepted: 17 March 2011

Published: 17 March 2011

## References

1. Gromiha MM, Ahmad S, Suwa M: Application of residue distribution along the sequence for discriminating outer membrane proteins. *Comput Biol Chem* 2005, **29**(2):135-142.
2. Berven FS, Flikka K, Jensen HB, Eidhammer I: BOMP: a program to predict integral beta-barrel outer membrane proteins encoded within genomes of Gram-negative bacteria. *Nucleic Acids Res* 2004, **32** Web Server: W394-399.
3. Garrow AG, Westhead DR: A consensus algorithm to screen genomes for novel families of transmembrane beta barrel proteins. *Proteins* 2007, **69**(1):8-18.
4. Wu Z, Feng E, Wang Y, Chen L: Discrimination of outer membrane proteins by a new measure of information discrepancy. *Protein Pept Lett* 2007, **14**(1):37-44.
5. Gromiha MM, Suwa M: Discrimination of outer membrane proteins using machine learning algorithms. *Proteins* 2006, **63**(4):1031-1037.
6. Park KJ, Gromiha MM, Horton P, Suwa M: Discrimination of outer membrane proteins using support vector machines. *Bioinformatics* 2005, **21**(23):4223-4229.
7. Remmert M, Linke D, Lupas AN, Soding J: HHomp-prediction and classification of outer membrane proteins. *Nucleic Acids Res* 2009, **37** Web Server: W446-451.
8. Liu Q, Zhu Y, Wang B, Li Y: Identification of beta-barrel membrane proteins based on amino acid composition properties and predicted secondary structure. *Comput Biol Chem* 2003, **27**(3):355-361.
9. Gromiha MM, Majumdar R, Ponnuswamy PK: Identification of membrane spanning beta strands in bacterial porins. *Protein Eng* 1997, **10**(5):497-500.
10. Lin H: The modified Mahalanobis Discriminant for predicting outer membrane proteins by using Chou's pseudo amino acid composition. *J Theor Biol* 2008, **252**(2):350-356.
11. Gromiha MM: Motifs in outer membrane protein sequences: applications for discrimination. *Biophys Chem* 2005, **117**(1):65-71.
12. Gnanasekaran TV, Peri S, Arockiasamy A, Krishnaswamy S: Profiles from structure based sequence alignment of porins can identify beta stranded integral membrane proteins. *Bioinformatics* 2000, **16**(9):839-842.
13. Bigelow H, Rost B: PROFtmb: a web server for predicting bacterial transmembrane beta barrel proteins. *Nucleic Acids Res* 2006, **34** Web Server: W186-188.
14. Martelli PL, Fariselli P, Krogh A, Casadio R: A sequence-profile-based HMM for predicting and discriminating beta barrel membrane proteins. *Bioinformatics* 2002, **18**(Suppl 1):S46-53.
15. Gromiha MM, Suwa M: A simple statistical method for discriminating outer membrane proteins with better accuracy. *Bioinformatics* 2005, **21**(7):961-968.
16. Garrow AG, Agnew A, Westhead DR: TMB-Hunt: a web server to screen sequence sets for transmembrane beta-barrel proteins. *Nucleic Acids Res* 2005, **33** Web Server: W188-192.
17. Yan C, Hu J, Wang Y: Discrimination of outer membrane proteins with improved performance. *BMC Bioinformatics* 2008, **9**:47.
18. Wimley WC: Toward genomic identification of beta-barrel membrane proteins: composition and architecture of known structures. *Protein Sci* 2002, **11**(2):301-312.
19. Zhai Y, Saier MH Jr: The beta-barrel finder (BBF) program, allowing identification of outer membrane beta-barrel proteins encoded within prokaryotic genomes. *Protein Sci* 2002, **11**(9):2196-2207.



20. Bagos PG, Liakopoulos TD, Spyropoulos IC, Hamodrakas SJ: **A Hidden Markov Model method, capable of predicting and discriminating beta-barrel outer membrane proteins.** *BMC Bioinformatics* 2004, **5**:29.
21. Bagos PG, Liakopoulos TD, Spyropoulos IC, Hamodrakas SJ: **PRED-TMBB: a web server for predicting the topology of beta-barrel outer membrane proteins.** *Nucleic Acids Res* 2004, **32** Web Server: W400-404.
22. Natt NK, Kaur H, Raghava GP: **Prediction of transmembrane regions of beta-barrel proteins using ANN- and SVM-based methods.** *Proteins* 2004, **56**(1):11-18.
23. Gromiha MM, Ahmad S, Suwa M: **TMBETA-NET: discrimination and prediction of membrane spanning beta-strands in outer membrane proteins.** *Nucleic Acids Res* 2005, **33** Web Server: W164-167.
24. Freeman TC Jr, Wimley WC: **A highly accurate statistical approach for the prediction of transmembrane beta-barrels.** *Bioinformatics* 2006, **22**(16):1965-1974.
25. Yan C, Hu J, Wang Y: **Discrimination of outer membrane proteins using a K-nearest neighbor method.** *Amino Acids* 2008, **35**(1):65-73.
26. Deng Y, Liu Q, Li YX: **Scoring hidden Markov models to discriminate beta-barrel membrane proteins.** *Comput Biol Chem* 2004, **28**(3):189-194.
27. Bagos PG, Liakopoulos TD, Hamodrakas SJ: **Evaluation of methods for predicting the topology of beta-barrel outer membrane proteins and a consensus prediction method.** *BMC Bioinformatics* 2005, **6**:7.
28. Koebnik R, Locher KP, Van Gelder P: **Structure and function of bacterial outer membrane proteins: barrels in a nutshell.** *Mol Microbiol* 2000, **37**(2):239-253.
29. Garrow AG, Agnew A, Westhead DR: **TMB-Hunt: an amino acid composition based method to screen proteomes for beta-barrel transmembrane proteins.** *BMC Bioinformatics* 2005, **6**:56.
30. Bigelow HR, Petrey DS, Liu J, Przybylski D, Rost B: **Predicting transmembrane beta-barrels in proteomes.** *Nucleic Acids Res* 2004, **32**(8):2566-2577.
31. Przytycka T, Aurora R, Rose GD: **A protein taxonomy based on secondary structure.** *Nat Struct Biol* 1999, **6**(7):672-682.
32. Bindewald E, Cestaro A, Hesser J, Heiler M, Tosatto SC: **MANIFOLD: protein fold recognition based on secondary structure, sequence similarity and enzyme classification.** *Protein Eng* 2003, **16**(11):785-789.
33. Lobley A, Sadowski M, Jones DT: **pGenTHREADER and pDomTHREADER: new methods for improved protein fold recognition and superfamily discrimination.** *Bioinformatics* 2009, **25**(14):1761-1767.
34. McGuffin LJ, Jones DT: **Improvement of the GenTHREADER method for genomic fold recognition.** *Bioinformatics* 2003, **19**(7):874-881.
35. Zhang Z, Kochhar S, Grigorov MG: **Descriptor-based protein remote homology identification.** *Protein Sci* 2005, **14**(2):431-444.
36. Yan RX, Si JN, Wang C, Zhang Z: **DescFold: a web server for protein fold recognition.** *BMC Bioinformatics* 2009, **10**:416.
37. McGuffin LJ, Jones DT: **Targeting novel folds for structural genomics.** *Proteins* 2002, **48**(1):44-52.
38. Si JN, Yan RX, Wang C, Zhang Z, Su XD: **TIM-Finder: A new method for identifying TIM-barrel proteins.** *BMC Struct Biol* 2009, **9**(1):73.
39. Fontana P, Bindewald E, Toppo S, Velasco R, Valle G, Tosatto SC: **The SSEA server for protein secondary structure alignment.** *Bioinformatics* 2005, **21**(3):393-395.
40. Needleman SB, Wunsch CD: **A general method applicable to the search for similarities in the amino acid sequence of two proteins.** *J Mol Biol* 1970, **48**(3):443-453.
41. Jones DT: **Protein secondary structure prediction based on position-specific scoring matrices.** *J Mol Biol* 1999, **292**(2):195-202.
42. Altschul SF, Madden TL, Schaffer AA, Zhang J, Zhang Z, Miller W, Lipman DJ: **Gapped BLAST and PSI-BLAST: a new generation of protein database search programs.** *Nucleic Acids Res* 1997, **25**(17):3389-3402.
43. Li W, Godzik A: **Cd-hit: a fast program for clustering and comparing large sets of protein or nucleotide sequences.** *Bioinformatics* 2006, **22**(13):1658-1659.
44. Rychlewski L, Fischer D: **LiveBench-8: the large-scale, continuous assessment of automated protein structure prediction.** *Protein Sci* 2005, **14**(1):240-245.
45. Eyrich VA, Marti-Renom MA, Przybylski D, Madhusudhan MS, Fiser A, Pazos F, Valencia A, Sali A, Rost B: **EVA: continuous automatic evaluation of protein structure prediction servers.** *Bioinformatics* 2001, **17**(12):1242-1243.
46. Rey S, Acab M, Gardy JL, Laird MR, deFays K, Lambert C, Brinkman FS: **PSORTdb: a protein subcellular localization database for bacteria.** *Nucleic Acids Res* 2005, **33** Database: D164-168.
47. Tsirigos KD, Bagos PG, Hamodrakas SJ: **OMPdb: a database of [beta]-barrel outer membrane proteins from Gram-negative bacteria.** *Nucleic Acids Res* .
48. Wimley WC: **The versatile beta-barrel membrane protein.** *Curr Opin Struct Biol* 2003, **13**(4):404-411.
49. Bendtsen JD, Nielsen H, von Heijne G, Brunak S: **Improved prediction of signal peptides: SignalP 3.0.** *J Mol Biol* 2004, **340**(4):783-795.
50. Ren Q, Kang KH, Paulsen IT: **TransportDB: a relational database of cellular membrane transport systems.** *Nucleic Acids Res* 2004, **32** Database: D284-288.
51. Andreeva A, Howorth D, Brenner SE, Hubbard TJ, Chothia C, Murzin AG: **SCOP database in 2004: refinements integrate structure and sequence family data.** *Nucleic Acids Res* 2004, **32** Database: D226-229.

doi:10.1186/1471-2105-12-76

**Cite this article as:** Yan et al.: Outer membrane proteins can be simply identified using secondary structure element alignment. *BMC Bioinformatics* 2011 **12**:76.

**Submit your next manuscript to BioMed Central and take full advantage of:**

- Convenient online submission
- Thorough peer review
- No space constraints or color figure charges
- Immediate publication on acceptance
- Inclusion in PubMed, CAS, Scopus and Google Scholar
- Research which is freely available for redistribution

Submit your manuscript at  
www.biomedcentral.com/submit

