SPAAN: a software program for prediction of adhesins and adhesin-like proteins using neural networks

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Received on February 24, 2004; revised on August 20, 2004; accepted on August 31, 2004
Advance Access publication September 16, 2004

ABSTRACT

Motivation: The adhesion of microbial pathogens to host cells is mediated by adhesins. Experimental methods used for characterizing adhesins are time-consuming and demand large resources. The availability of specialized software can rapidly aid experimenters in simplifying this problem. We have employed 105 compositional properties and artificial neural networks to develop SPAAN, which predicts the probability of a protein being an adhesin ($P_{ad}$).

Results: SPAAN had optimal sensitivity of 89% and specificity of 100% on a defined test set and could identify 97.4% of known adhesins at high $P_{ad}$ value from a wide range of bacteria. Furthermore, SPAAN facilitated improved annotation of several proteins as adhesins. Novel adhesins were identified in 17 pathogenic organisms causing diseases in humans and plants. In the severe acute respiratory syndrome (SARS) associated human corona virus, the spike glycoprotein and nsp5 (nsp2, nsp5, nsp6 and nsp7) were identified as having adhesin-like characteristics. These results offer new lead for rapid experimental testing.

Availability: SPAAN is freely available through ftp://203.195.151.45
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INTRODUCTION

Microbial pathogens encode adhesins that mediate their adherence to host cell surface receptors, membranes or the extracellular matrix for successful colonization. Investigations into this primary event of host–pathogen interaction have revealed a wide array of adhesins in a variety of pathogenic microbes (Finlay and Falkow, 1997). New approaches to vaccine development focus on targeting adhesins to abrogate the colonization process (Wizemann et al., 1999). However, the specific roles of particular adhesins in several pathogens remain to be elucidated.

One of the best-understood mechanisms of bacterial adherence is the attachment mediated by pili or fimbriae. The well-studied adhesins in this category are FimH and PapG adhesins of *Escherichia coli* (Hahn et al., 2002) and the Type IV pilu adhesins in *Pseudomonas aeruginosa*, Neisseria, Moraxella, enteropathogenic *E.coli* and *Vibrio cholerae* (Strom and Lory, 1993). Several adhesins from other commonly known bacterial pathogens include MrkD protein of *Klebsiella pneumoniae* (Gerlach et al., 1989), Hia of *Haemophilus influenzae* (Barenkamp and St Geme, 1996) and many others (for further details see http://www.igib.res.in/data/seepath/spaan_data.html).

Several vaccine formulations either currently approved or being evaluated use adhesins as immunizing agents. Examples include filamentous hemagglutinin and pertactin proteins against *Bordetella pertussis* (Halperin et al., 2003), FimH against pathogenic *E.coli* (Langermann et al., 2000), PsaA against pneumococcal disease (Rapola et al., 2003), outer membrane vesicle preparations including BabA adhesin against *Helicobacter pylori* infections (Prinz et al., 2003) and a synthetic peptide anti-adhesin vaccine against *P.aeruginosa* infections (Cachia and Hodges, 2003).

Experimental identification of adhesins is an arduous task. Computational methods such as homology search can aid in identification, but this procedure suffers from limitations when the homologues are not characterized. Sequence analysis based on compositional properties provides relief to this problem. Amino acid composition is a fundamental attribute of a protein and it has significant correlation to its location, function, folding type, shape and *in vivo* stability (Nakashima and Nishikawa, 1994; Nandi et al., 2003). Recently, compositional properties have been applied to the problems as diverse as the prediction of functional roles (Hobohm and Sander, 1995), protein secondary structures (Rost and Sander, 1993), secretory proteins...
and apicoplast-targeted proteins in *Plasmodium falciparum* (Schneider, 1999; Zuegge et al., 2001).

We report a non-homology method using 105 compositional properties combined with artificial neural networks (ANNs) to identify adhesins and adhesin-like proteins in species belonging to a wide phylogenetic spectrum.

**SYSTEMS AND METHODS**

The five attributes

**Amino acid frequencies** Amino acid frequency \( f_i = (\text{counts of } i\text{-th amino acid in the sequence})/l \), where \( i = 1, \ldots, 20 \) and \( l \) is the length of the protein.

**Multiplet frequencies** Multiplets are defined as homopolymeric stretches \((X)_n\), where \( X \) is the amino acid and \( n \) (integer) \( \geq 2 \) (Brendel et al., 1992). After identifying all the multiplets, the frequencies of the amino acids in the multiplets were computed as follows:

\[
f_i(m) = (\text{counts of } i\text{-th amino acid occurring as multiplet})/l.
\]

**Dipeptide frequencies** The frequency of a dipeptide \((i, j)\) \( f_{ij} = (\text{counts of } ij\text{-th dipeptide})/(\text{total dipeptide counts}) \), where \( i, j = 1–20 \).

The theoretical number of possible dipeptides is 400. The recommended ratio for the number of input vectors to the number of weight connections is \( \sim 2 \) to avoid overfitting (Andrea and Kalayeh, 1991). Therefore, we used top 20 dipeptides (when arranged in the ascending order of the \( P \)-values assessed using \( t \)-test) whose frequencies in the adhesin dataset were significantly different from that in the non-adhesin dataset (single-letter code): NG, RE, TN, NT, GT, TT, DE, ER, RR, RK, RI, AT, TS, IV, SG, GS, TG, GN, VI and HR.

**Charge composition** The frequency of charged amino acids (R, K, E and D considering the ionization properties of the side chains at pH 7.2) is given by \( f_c = (\text{counts of charged amino acids})/l \). Furthermore, information on the characteristics of the distribution of the charged amino acids in a given protein sequence was obtained by computing the moments of the positions of the occurrences of the charged amino acids.

The general expression to compute moments of a given order; say ‘\( r \)’ is

\[
M_r = r\text{-th order moment of the positions of charged amino acids} = \sum \frac{(X_i - X_m)^r}{N},
\]

where, \( X_m \) is the mean of all positions of charged amino acids, \( X_m = \sum_{i=1}^{N} X_i/N \); \( X_i \) is the position of \( i\text{-th} \) charged amino acid; and \( N \) is the number of charged amino acids in the sequence.

The frequency of charged amino acids, the length of the protein and the moments of order from of 2 to 19 were used to train the ANN constituting a total of 20 inputs. Moments of order \( \geq 19 \) were not useful in further enhancing the performance.

**Hydrophobic composition** The amino acids were classified into five groups based on their hydrophobicity scores: \((-8 \text{ for } K, E, D \text{ and } R), (-4 \text{ for } S, T, N \text{ and } Q), (-2 \text{ for } P \text{ and } H), (+1 \text{ for } A, G, Y, C \text{ and } W) \) and \(+2 \text{ for } L, V, I, F \text{ and } M \) (Brendel et al., 1992).

The inputs fed into the neural network for each group are as follows:

1. \( f_i = (\text{counts of } i\text{-th group})/(\text{total counts in the protein}), \) where \( i = 1–5 \).
2. \( m_{jj} = j\text{-th order moment of positions of amino acids in } i\text{-th group}, \) where \( j = 2–5 \).

A total of 25 inputs representing the hydrophobic composition of a protein were fed to the neural network.

Taken together, a total of 105 compositional properties in the five modules were used to predict the adhesin-like characteristics of a given protein sequence.

**Database construction**

**Adhesins** Protein sequences were retrieved from [http://www.ncbi.nlm.nih.gov](http://www.ncbi.nlm.nih.gov) using the keyword ‘adhesin’. Furthermore, proteins containing the following keywords were removed from the primary retrieval: ‘transport’, ‘pyrophosphatase’, ‘peroxidase’, ‘myosin’, ‘chaperone’, ‘hydrolase’, ‘gene product’, ‘accessory’, ‘regulatory’, ‘patent’, ‘permease’, ‘hypothetical’, ‘keratin’, ‘agrobacterium’, ‘intimin’, ‘ORF’, ‘ATP binding’, ‘tRNA’, ‘deiminase’, ‘metallopeptinase’, ‘cofactor’, ‘amylase’, ‘methylase’, ‘unknown’, ‘ribosomal’, ‘alternative start’, ‘submitter believes’ and ‘phospholipase’.

The remaining sequences in the adhesin dataset were manually curated to generate a set of well-annotated proteins many of which have been verified experimentally.

**Non-adhesins** The rationale we used here was to collect sequences of enzymes and other proteins that function within the cell. They probably have remote possibility of functioning as adhesins and would differ in compositional characteristics (Nakashima and Nishikawa, 1994). The keywords used were ‘dehydratase’, ‘dehydrogenase’, ‘ribosomal protein’, ‘kinase’, ‘polymerase’, ‘acyl-CoA synthase’, ‘decarboxylase’, and ‘hydrolase’. Since effective implementation of the algorithm requires that the sizes of the two databases to be similar, we selected sequences from *Methanococcus jannaschii*, *E.coli* and *Saccharomyces cerevisiae* as representatives of the three primary kingdoms of life: Archaea, Eubacteria and Eukarya. This selection offers a diverse set for obtaining a broad range of limits for the detection of non-adhesins. In the subsequent step, ‘hypothetical’,
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**Fig. 1.** The neural network architecture of SPAAN. A given protein sequence was processed through five modules, A, C, D, H and M, to quantify the five types of compositional attributes. A, Amino acid frequencies; C, Charge composition; D, Dipeptide frequencies; H, Hydrophobic composition; and M, Multiplet frequencies. The sequence shown is part of the FimH precursor (gi 5524634) of *E.coli*. The direction of arrows show data flow.

'transport', 'unknown' and 'membrane' protein sequences were removed.

**Eliminating redundant entries** We used CLUSTALW (Thompson *et al.*, 1994) to analyze sequence similarities between the sequences in pairwise comparisons. Only one sequence entry was retained among pairs that had a CLUSTALW score of 100. Partial sequence entries were also removed. The total number of adhesins was 469 and the total number of non-adhesin proteins from *E.coli* was 282, *M.jannaschii* was 162 and *S.cerevisiae* was 259 which summed to 703 entries.

**Neural network**

The feed forward error back propagation neural network algorithm was used. The program was downloaded from the website http://www.cs.colostate.edu/~anderson a gift from Charles W. Anderson (Department of Computer Science, Colorado State University, Fort Collins, CO, e-mail: anderson@cs.colostate.edu)

**ALGORITHM**

**Neural network architecture**

The neural network used here has a multilayer feed forward topology. It consists of an input layer, a hidden layer and an output layer. This is a ‘fully-connected’ neural network where each neuron of one layer is connected to each neuron of the next layer (Fig. 1). The weight connections are denoted by $w_{ij}$. The state $I_i$ of each neuron in the input layer is assigned directly from the input data, whereas the states of hidden layer neurons are computed from the states of input layer neurons using the sigmoid function,

$$h_j = 1 / \left[1 + \exp(- (w_{j0} + \sum w_{ij} I_i))\right],$$

where $w_{j0}$ is the bias weight. The back propagation algorithm was used to minimize the differences between the computed output and the target value. The target value for adhesins was set as ‘1’ and for non-adhesins it was set as ‘0’.

In the initial optimization experiments, a training set and a validate set were used. The training set had 367 adhesins and
580 non-adhesins. The validate set had 102 adhesins and 123 non-adhesins. A total of 10,000 cycles (epochs) of training iterations were performed. Subsequently, the best epoch with minimum error on the validate set was identified and the corresponding weight matrix was used for the prediction.

Five networks were prepared, one for each attribute (Fig. 1). The number of neurons in the input layer was equal to the number of input data points for each attribute. The optimal number of neurons in the hidden layer was determined through experimentation for minimizing the error at the best epoch for each network individually. An upper limit for the total number of weight connections was set to half of the total number of input vectors to avoid overfitting as suggested previously (Andrea and Kalayeh, 1991). The final number of neurons in the hidden layer for each module was Amino acid frequencies: 30, Multiplets frequencies: 28, Dipeptide frequencies: 28, Charge composition: 30 and Hydrophobic composition: 30. During predictions, the network is fed with new data from the sequences that were not part of the training set.

**Probability of being an adhesin, the \( P_{ad} \) value**

Query proteins were processed modularly through the networks trained for each attribute. Thus, five probability outputs for each sequence were obtained. Final prediction was computed using the following expression, which is a weighted linear sum of the probabilities from five modules:

\[
P_{ad} = \frac{(P_A \times fc_A + P_C \times fc_C + P_D \times fc_D + P_H \times fc_H + P_M \times fc_M)}{(fc_A + fc_C + fc_D + fc_H + fc_M)}
\]

where \( fc_i \) is the fraction of correlation of \( i \)-th module of the trained neural network, where \( i = A, C, D, H \) or \( M \). The fractions of correlation \( fc_i \) represent the fractions of total entries that were predicted correctly of \( P_{ad, \text{adhesin}} > 0.5 \) and \( P_{ad, \text{non-adhesin}} < 0.5 \) by the trained network on the validate set (Charles W. Anderson, http://www.cs.colostate.edu/~anderson). \( fca = 0.84, fcC = 0.71, fcD = 0.84, fcH = 0.79, fcM = 0.83 \).

**Matthew’s correlation coefficient for assessing the performance of SPAAN**

The Matthew’s correlation coefficient (Mcc) (Matthews, 1975) is defined as follows:

\[
\text{Mcc} = \frac{(TP \times TN) - (FP \times FN)}{(TN + FN)(TN + FP)(TP + FN)(TP + FP)},
\]

where TP stands for true positives, TN the true negatives, FP the false positives and FN the false negatives.

Here TPs are adhesins and TNs are non-adhesins. Adhesins with \( P_{ad} \) value above a chosen threshold are TPs, whereas known non-adhesins with \( P_{ad} \) value below the chosen threshold are TNs. The sensitivity, \( Sn \), is given by \( TP / (TP + FN) \) and specificity, \( Sp \), is given by \( TP / (TP + FP) \).

**SPECIFICATIONS**

Computer programs used to compute individual compositional attributes were written in C and executed on a PC with operating system Red Hat Linux version 7.3 or 8.0.

**Sequence inputs**

SPAAN accepts input sequence files in the FASTA format. Multiple sequences can be present in one file. Protein sequences with ambiguous amino acids and/or of length <50 amino acids were filtered out. Amino acids must use the single-letter code according to the IUPAC-IUB nomenclature system.

**RESULTS AND DISCUSSION**

**Sensitivity, specificity and correlation coefficient**

In designing SPAAN, we developed a non-homology, compositional property based method to predict adhesins and adhesin-like proteins solely from the sequence data. To assess the performance of SPAAN, we prepared a test set of 37 well-annotated adhesins and 37 non-adhesins that were not part of the training set. The results are shown in Figure 2. It is apparent that SPAAN could identify 89% of known adhesins with 100% specificity when examined at \( P_{ad} \geq 0.51 \). At \( P_{ad} \geq 0.51 \), the Mcc (Matthews, 1975) was observed to be highest (0.94). We observed that the combination of five modules provided the best results.

Assessment of the performance in individual modules showed that they performed poorly when compared with the combination of modules. The performances of individual modules were as follows: Charge composition (\( P_c = 0.55 \), Mcc = 0.658, Sn = 0.756 and Sp = 0.848), Dipeptide frequencies (\( P_d = 0.54 \), Mcc = 0.84, Sn = 0.86 and Sp = 0.94), Hydrophobic composition (\( P_h = 0.61 \), Mcc = 0.63, Sn = 0.54 and Sp = 0.9) and Multiplet frequencies (\( P_M = 0.58 \), Mcc = 0.77, Sn = 0.81 and Sp = 0.9). Performance of the Amino acid frequencies’ module could not be assessed unambiguously because the Mcc was nearly flat over a broad range. These observations suggest that it would be fruitful to include multiple modules for obtaining high-quality predictions and are consistent with the experience of Hobohm and Sander (1995).

**SPAAN predicts experimentally characterized adhesins with high \( P_{ad} \) value**

Considering the small size of test set, we examined the general applicability of SPAAN by analyzing several well-characterized adhesins from a wide range of pathogens causing a variety of diseases. The results on 194 adhesins with binding specificity to a wide range of host receptors are displayed in Table 1 (for further details see http://www.igib.res.in/data/seepath/spaan_data.html). It is apparent that except two FimH proteins of *E.coli*, pertactin of *B.pertussis*, protein F of *Streptococcus pyogenes* and PspC
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Fig. 2. Performance of SPAAN. A theoretical polynomial curve of second-order (dashed line) was fitted to the observed curve (smooth line) with a Karl–Pearson correlation coefficient $R^2 = 0.9799$. The maximum point of the theoretical curve was chosen as a reference (vertical dotted line) to identify the maximum Mcc on the observed curve (shown by an arrow). Note that the Mcc does not drop down to the x-axis because the highest $P_{ad}$ value attained using adhesins was 0.939 in comparison with the theoretical attainable limit of 1.0.

Table 1. Prediction of well-characterized adhesins from various bacterial pathogens using SPAAN

| Species        | Disease caused | Adhesin$^a$ | Host ligand                                         | $P_{ad}$ value$^b$ (range) | Reference          |
|----------------|----------------|------------|----------------------------------------------------|---------------------------|--------------------|
| **E.coli**     | Diarrhoea      | PapG (27)  | α-α-gal(1–4) β-β-Gal-containing receptors         | 0.84–0.76                 | Bock et al. (1985) |
|                |                | SfaS (5)   | alpha-sialyl-beta-2,3-b-galactose                  | 0.94–0.94                 | Moch et al. (1987) |
|                |                | FimH (63)  | β-mannosides                                       | 0.96–0.23$^c$            | Hahn et al. (2002) |
|                |                | Intimin (12)| Tyrosine-phosphorylated form of host cell receptor Hf90 | 0.95–0.78                 | Rosenshine et al. (1996) |
| **Non-typeable**| Influenza      | PrsG (5)   | Gal(alpha1–4)Gal                                   | 0.86–0.85                 | Johnson et al. (1997) |
| **H.influenzae** | Whooping cough | HMW1, HMW2 | Human epithelial cells                            | 0.97                      | St Gerne (1996)    |
| **B.pertussis** |                | Hia (8)    | Human conjunctival cells                           | 0.93–0.90                 |                    |
| **Yersinia enterocolitica** | Enterocolitis | Pertactin  | Integrins                                          | 0.43                      | Brennan and Shahin (1996) |
| **Enterococcus faecalis** | Empyema in patients | YadA (5)  | $β_1$ integrins                                    | 0.88–0.79                 | Schulze-Koops et al. (1993) |
|                | with liver disease | EfaA      | Unknown                                            | 0.83                      | Finlay and Falkow (1997) |
| **H.pylori**   | Peptic ulcers  | BabA (17)  | Difucosylated Lewis blood group antigen            | 0.87–0.68                 | Prinz et al. (2003) |

$^a$The number of sequences analyzed from different strains and homologs from related species are shown in parentheses.

$^b$Rounded off to the second decimal.

$^c$Out of 63 FimH proteins, 54 were from E.coli, 6 from Shigella flexneri, 2 from Salmonella enterica and 1 from Salmonella typhimurium. Except two FimH proteins, the rest had $P_{ad} \geq 0.51$. The two exceptions (gi numbers: 5524636 and 1778448) were from E.coli. The gi: 5524636 protein is annotated as a FimH precursor but is much shorter (129 amino acids) than other members of the family. The gi: 1778448 protein is a S.typhimurium homolog in E.coli.

of Streptococcus pneumoniae, the rest 189 adhesins had $P_{ad} \geq 0.51$ indicating an overall sensitivity of 97.4%. These results demonstrate the general applicability of SPAAN. **SPAAN is a non-homology method based on sequence properties**

To examine the non-homology character of SPAAN, we prepared a dataset of 130 adhesins that did not have any protein pairs with CLUSTALW score of 100. Equal number of non-adhesins was selected with the same criterion. A histogram plot of adhesins and non-adhesins in the various ranges of $P_{ad}$ values is displayed in Figure 3a. It is evident that SPAAN is capable of segregating the adhesins and non-adhesins into two distinct cohesive groups. Most of the adhesins (96%) have $P_{ad} \geq 0.51$ whereas all the non-adhesins (100%) have $P_{ad} < 0.51$. 

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Fig. 3. SPAAN is a non-homology-based software program. A total of 130 adhesins and 130 non-adhesins were analyzed to assess whether the predictive power of SPAAN could be influenced by sequence relationships. (a) Histogram plots of the number of proteins in the various \( P_{ad} \) value ranges are shown. Shaded bars represent adhesins and open bars represent non-adhesins. Note the ability of SPAAN to segregate adhesins and non-adhesins into two distinct cohesive groups. (b) Pairwise sequence relationships among the adhesins were determined using CLUSTALW and plotted on the \( x \)-axis. Higher CLUSTALW scores indicate similar pairs. The corresponding differences in \( P_{ad} \) values in the same protein pair was plotted on the \( y \)-axis. Each point in the diagram represents a pair. Arrow points to protein pairs of the FimH family with high \( \Delta P_{ad} \) values in spite of high similarity. Since one of the FimH proteins (gi: 5524636) had very low \( P_{ad} \) value, all pairs with this false negative protein show high \( \Delta P_{ad} \) values. The protein (gi: 5524636) is of much shorter length compared with other members of the same family. (c) Plot for non-adhesins. Data were plotted in the four-quadrant format to enhance clarity.
Table 2. Analysis of predictions carried out using SPAAN on genome scans of a few selected pathogenic organisms

| Organism                                | Disease caused   | Total no. of proteins analyzed | No. of these supported by complementary evidence CDD/BLASTP/PubMed | No. of these supported by complementary evidence BETAWRAP | No. of adhesin-like proteins | No. of false positives |
|-----------------------------------------|------------------|--------------------------------|---------------------------------------------------------------------|----------------------------------------------------------|-----------------------------|------------------------|
| E. coli O157:H7                         | Diarrhoea        | 50                             | 37<sup>b</sup>                                                      | 33                                                       | 12                          | 1                      |
| H. pylori                               | Peptic ulcers    | 50                             | 25<sup>c</sup>                                                      | 36                                                       | 24                          | 1                      |
| Listeria monocytogenes                 | Listeriosis      | 50                             | 23<sup>d</sup>                                                      | 39                                                       | 24                          | 3                      |
| S. pneumoniae                          | Bacterial pneumonia | 40                              | 13<sup>e</sup>                                                      | 12                                                       | 23                          | 4                      |
| Mycobacterium tuberculosis H37Rv       | Tuberculosis     | 50                             | —                                                                  | 32                                                       | 50<sup>f</sup>              | —                      |
| SARS-associated corona virus           | SARS             | 5                              | —                                                                  | —                                                        | 5<sup>g</sup>              | —                      |

*The top-scoring similar sequences with $E < 0.001$ were only considered for assessing sequence-based relationships. The low complexity filter was ‘off’.

<sup>a</sup>Includes Fimbrial adhesins (nine proteins), AidA-I, gamma intimin, hemagglutinin, translocated intimin receptor, putative tail fiber protein and putative major tail protein.

<sup>b</sup>Includes putative vacuolating cytotoxin (VacA) (autotransporter adhesin-like), outer membrane protein (adhesin; 2 proteins), outer membrane protein (porin; 3 proteins), flagellin A and outer membrane proteins (13 proteins).

<sup>c</sup>Includes internalin A (mediates adhesion and invasion), other internalins (seven proteins), pepGlycan-linked protein (similar to autotransporter adhesin, eight proteins), autolysin (amidase, presumably involved in adhesion, three proteins), flagellar hook protein may be involved in adhesion, cell surface protein (adhesin by BLASTP analysis).

<sup>d</sup>Includes PsaA (pneumococcal surface adhesin A), pspA similar to ChpA, ChpD, ChpE, Chf1 and Chp (two proteins) all similar to ChpA, two hypothetical proteins with low-level similarity to ChpA, autolysin.

<sup>e</sup>PPE, PE_PGRS (35 proteins), PPE (12 proteins), two hypothetical proteins with no similarity to either PE or PE_PGRS or PPE proteins.

<sup>f</sup>These proteins were the spike glycoprotein with antigenic properties, and nsp2, nsp5, nsp6 and nsp7.

We computed the pairwise sequence similarities using CLUSTALW (Thompson et al., 1994) and compared with the differences between the $P_{ad}$ values in the pair (denoted by $\Delta P_{ad}$). The relationships for adhesins and non-adhesins are shown in Figures 3b and c, respectively. In both adhesins and non-adhesins, $\Delta P_{ad}$ values are uniformly low and appear nearly independent of sequence relationship. Furthermore, among the protein pairs with score $>2$, 82% of adhesin pairs and 86% of non-adhesin pairs had $\Delta P_{ad} < 0.2$. These data reinforce the non-homology character of SPAAN.

**Application of SPAAN to whole genomes**

The results of the genome scan for selected pathogens of humans and plants are displayed in Table 2 (for detailed data description, see online Supplementary Table at http://www.igib.res.in/data/seepath/spaan_data.html). We used a stringent criterion of $P_{ad} > 0.7$ on the basis of the results shown in Figure 3a to reduce the detection of false positives. Subsequently, we restricted our analysis to a maximum of 50 top-scoring proteins. This serves as a good starting point to examine the performance of SPAAN and to identify top-scoring novel adhesins with high confidence. The experimentally characterized adhesins from a wide range of pathogens form the top list in genome scans. Several of the predicted adhesins are supported by complementary evidence from Conserved Domain Database search (RPS-BLASTP), BLASTP and the beta helix predictor BETAWRAP (Marchler-Bauer et al., 2002; Altschul et al., 1990; Bradley et al., 2001). About 30–78% of these predicted adhesins also contain beta helix motif. The beta helix motif was found to be associated with several adhesins, toxins, virulence factors and surface proteins (Bradley et al., 2001).

In addition, SPAAN guided the improved annotation of a number of adhesins by suggesting re-examination of these proteins using the most commonly used software listed above. It is also evident that the well-known adhesins in these organisms top the list of predictions using SPAAN (Table 2). Several proteins with high $P_{ad}$ values were identified using SPAAN for which either limited or no complementary evidence exist. We have classified these proteins as ‘adhesin-like’. Interestingly, several mycobacterial proteins, namely, 35 PE_PGRS proteins and 12 PPE proteins were identified with high $P_{ad}$ value. SPAAN could identify these putative mycobacterial adhesins even though our training dataset was devoid of mycobacterial proteins. Indeed, experimental analysis has demonstrated that some of these proteins could mediate host–pathogen interactions (Brennan et al., 2001). These results demonstrate that SPAAN could overcome taxonomic limits and can be used for general purpose.

Although SPAAN was primarily trained on bacterial adhesins, we examined its ability to predict putative adhesins from eukaryotic systems. The criteria was relaxed by using the base threshold value of $P_{ad} \geq 0.51$ to scan the genome of SARS-associated human coronavirus. The spike glycoprotein, nsp2, nsp5, nsp6 and nsp7 were identified as adhesins. Spike glycoprotein has been implicated to play a role in viral entry and pathogenesis (Gallagher and Buchmeier, 2001). The role of nsp proteins in viral pathogenesis is not clear. Since SARS is an important public health problem, these results could rapidly aid experiments that characterize host–pathogen interactions.
A few false positives do appear in the list. A judicious approach for experimental characterization could be developed by considering the total number of proteins to be analyzed, prioritizing proteins with other complementary evidence while keeping the number of false positives as low as possible. In summary, SPAAN could serve as an useful guide to perform experimental characterization of proteins for adhesin-like properties.

ACKNOWLEDGEMENTS

We thank Prof. G. Padmanaban, Prof. Samir K. Brahmachari, Dr R. Sonti and A. Maharana for their useful discussions. S.R. thanks Council of Scientific and Industrial Research for a grant under the New Millennium Indian Technology Leadership Initiative (NMITLI) program.

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