ProInflam: a webserver for the prediction of proinflammatory antigenicity of peptides and proteins

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Abstract

Background: Proinflammatory immune response involves a complex series of molecular events leading to inflammatory reaction at a site, which enables host to combat plurality of infectious agents. It can be initiated by specific stimuli such as viral, bacterial, parasitic or allergenic antigens, or by non-specific stimuli such as LPS. On counter with such antigens, the complex interaction of antigen presenting cells, T cells and inflammatory mediators like IL1α, IL1β, TNFα, IL12, IL18 and IL23 lead to proinflammatory immune response and further clearance of infection. In this study, we have tried to establish a relation between amino acid sequence of antigen and induction of proinflammatory response.

Results: A total of 729 experimentally-validated proinflammatory and 171 non-proinflammatory epitopes were obtained from IEDB database. The A, F, I, L and V amino acids and AF, FA, FF, PF, IV, IN dipeptides were observed as preferred residues in proinflammatory epitopes. Using the compositional and motif-based features of proinflammatory and non-proinflammatory epitopes, we have developed machine learning-based models for prediction of proinflammatory response of peptides. The hybrid of motifs and dipeptide-based features displayed best performance with MCC = 0.58 and an accuracy of 87.6 %.

Conclusion: The amino acid sequence-based features of peptides were used to develop a machine learning-based prediction tool for the prediction of proinflammatory epitopes. This is a unique tool for the computational identification of proinflammatory peptide antigen/candidates and provides leads for experimental validations. The prediction model and tools for epitope mapping and similarity search are provided as a comprehensive web server which is freely available at http://metagenomics.iiserb.ac.in/proinflam/ and http://metabiosys.iiserb.ac.in/proinflam/.

Keywords: Proinflammatory, Antigens, Prediction, Vaccine, Machine-learning

Background

The role of peptides as therapeutic agents has gained considerable importance recently, and more than 7000 natural peptides have been reported which play a pivotal role in human physiology and have different applications such as, vaccines, and other immunotherapeutics [1]. However, in addition to the desired action, these peptides may show undesirable immuno-activity, for example B cell or T cell activation and other proinflammatory events [2–4]. Similarly in nature, different infectious agents also harbor immunomodulatory properties present in their proteins, which help them in initiation and progression of the disease [5, 6]. Several examples of proinflammatory reactions are known where the pathogens get advantage of inflammation caused by their antigen. A well-known example is the proinflammatory response induced by the peptide Hp(2–20) of Helicobacter pylori which induces proinflammatory activities such as, recruiting and activating various immune cells like neutrophils and monocytes, upregulation of integrins (Mac-1) and activation of the oxygen radical producing NADPH-oxidase. This leads to destruction of host mucosal tissue along with
to the best of authors knowledge, there are no compu-
amino acid sequence and its proinflammatory behavior.
peptide sequences underscore the correlation between
inflammatory behavior, which worsen the disease [11].
ever, examples like C-peptide have an undesirable pro-
therapeutics. There are examples of therapeutic peptides
may be a desirable or undesirable property of peptide
The induction of proinflammatory immune response
Results and discussion

The above evidences of proinflammatory property of
peptide sequences underscore the correlation between
amino acid sequence and its proinflammatory behavior.
To the best of authors knowledge, there are no compu-
tional studies reported till date where any sequence-
based signature or feature has been investigated which
could be responsible for proinflammatory behavior of a
peptide. Although, several studies have focused on the prediction of different kind of immune epitopes, such as
B cell epitopes [12–14], T cell epitopes [15–17], MHC
binders [18], IL4-inducing peptides [19], IFN-gamma
inducing MHC binders [20] and allergenicity [21, 22],
there is no study known where the sequence-based features have been examined to determine the proinflammatory nature of peptides. In this work, we have analyzed
amino acid sequence of experimentally validated proinflammatory epitopes (PiEs) in contrast to non-proinflammatory epitopes (NPiEs) and developed a machine
learning-based classification method incorporating the
sequence-based features, to predict the proinflammatory
nature of peptides and proteins.

Results and discussion

The induction of proinflammatory immune response may be a desirable or undesirable property of peptide therapeutics. There are examples of therapeutic peptides
where inflammation is a desirable property [3, 23]. How-
ever, examples like C-peptide have an undesirable pro-
inflammatory behavior, which worsen the disease [11].
The aim of this study is to develop an in silico method
for predicting PiEs. In this study, we have analyzed the
sequence-based properties which may contribute to its
proinflammatory nature. Although in the past, several
studies have been carried out on allergenic proteins/pep-
tides [21, 22], toxic peptides [24], MHC binders [18], CTL
epitopes [17], and B cell epitopes [12]; this study focus on
investigating the basic property of peptide antigens to ini-
tiate proinflammatory cascade, which involves recruiting
several immune cells, activation of complement proteins
and communication via different immune mediators,
which are also known as cytokines. The cytokines, such as
IL1α, IL1β, TNFα, IL12, IL18 and IL23, are considered
as proinflammatory cytokines [25], which are established
mediators measured during a proinflammatory reaction
assay. In this study, the experimentally validated epitopes
which are assayed positive for these cytokines were con-
sidered as PiEs. The epitopes which gave negative assay
were considered as NPiEs (Fig. 1). The compositional and motif-based analysis were carried out on the main
dataset, however, the prediction models were developed
using both main and alternate dataset, as discussed in
methods section.

Compositional analysis

In order to examine the abundance of amino acids in
PiEs as compared to NPiEs, amino acid composition was
computed for both the epitope classes. The composi-
tional analysis revealed that the average composition of
Ala, Phe, Iso, Leu and Val amino acids is higher in PiEs
as compared to NPiEs (Fig. 2; Additional file 1: Table S1),
whereas, the amino acids Cys, Gly, Pro and Thr are less
abundant in PiEs. It suggests a preferential abundance of
some amino acids in PiEs. Similarly, the dipeptide com-
position was examined in both the classes and several
dipeptides were observed to be significantly abundant
(Welch’s t test, p < 0.05) in PiEs. The composition of 91
dipeptides were found to differ significantly in PiEs as
compared to NPiEs, of which the dipeptides AE, DA, GF,
IN, KA, KD, RK, RM, TL and YA are the top ten dipe-
ptides which showed the most significant differences in
composition. Of these 91 significantly different dipep-
tides, AE, FA, FF, PF, IV, IN were also observed to be
the most abundant in PiEs (Additional file 2: Table S2).
The composition-based analysis revealed that in PiEs,
most of the preferred residues (Ala, Phe, Iso, Leu and
Val) are aliphatic or hydrophobic amino acids. Similarly,
the significant dipeptides, discussed in the above section,
mostly included pairs of aliphatic or hydrophobic amino
acids in different local orders.

Motif analysis

The proinflammatory immune response requires the activation of T cells by presentation of peptide bound
to MHC molecule. Several key amino acids of the peptide bind at certain positions in the binding core of MHC which suggests that peptide may contain specific motif required for binding. In order to investigate if there are particular sequential motifs present in PiEs, the motifs were searched in the training datasets using MERCI software as described in the Methods section. After applying different algorithms available in MERCI software, such as none, Koolman–Rohm and Betts–Russell, several motifs were discovered which were found to be exclusively present in PiEs and NPiEs (Table 1; Additional file 3: Table S3). The overall coverage of a motif represents the total number of epitopes harboring that particular motif and same epitopes might also be covered by other motifs. Out of 583 PiEs present in positive dataset, the Betts–Russell algorithm, Koolman–Rohm algorithm and none algorithm could identify 256, 192 and 179 unique PiEs using 11, 10 and 10 motifs, respectively. Similarly, the above algorithms were used to identify motifs from NPiEs (Additional file 3: Table S3).

The “hydrophobic hydrophobic K hydrophobic hydrophobic” and “hydrophobic aliphatic polar N” were found to be the most frequent motifs and covered 54 and 44 unique proinflammatory epitopes, respectively. Among motifs given by Koolman–Rohm algorithm, “basic A aliphatic” was the most recurring motif covering 41 unique proinflammatory epitopes. Interestingly, the 10 motifs obtained in MERCI analysis with none algorithm were same as the significant dipeptides obtained in the compositional analysis. MERCI motifs, which were discovered exclusively in proinflammatory epitopes against non-proinflammatory epitopes, showed similar conservation of residues as observed in compositional analysis, for example, the motifs having highest coverage as shown in Table 2, are abundant in hydrophobic and aliphatic residues at various positions. Similar observations were also
reported earlier where MBP(85–105) peptide binds to MHC allele DRB1 *1501 with a nonaromatic, hydrophobic anchor (L, V, or I) at position i and by a bulky hydrophobic residue (F or Y) at position i + 3 as primary anchor which may contribute to its immunodominance [26].

**Machine learning-based classification**

The preliminary analysis unveiled that the PiEs differ from NPiEs in amino acid sequence-based features, and thus, sequence-based features can be exploited for their classification into PiEs and NPiEs epitopes using machine learning-based classification. Classification models for different features were developed using 6 different machine learning techniques (SVMlight and RandomForest, BayesNet, NaiveBayes, IBk and J48). However, the performance of RandomForest, BayesNet, NaiveBayes, IBk and J48 models was observed to be lower as compared to SVM-based models (Additional file 4: Table S4). Therefore, SVM-based models were implemented and discussed in the manuscript. Similarly, the performance of models on alternate dataset is mentioned in Additional file 5: Table S5.

**Amino acid composition-based models**

The two classes of epitopes differed in amino acid composition as mentioned in the compositional analysis

| Algorithm for motif discovery | \( N_p \) | \( N_n \) |
|------------------------------|---------|---------|
| Betts–Russell                | 256     | 29      |
| Koolman–Rohm                 | 192     | 15      |
| None                         | 179     | 9       |

For example, Betts–Russell algorithm-based proinflammatory and non-proinflammatory motifs could identify 256 proinflammatory as well as 29 non-proinflammatory unique epitopes, respectively.

| Proinflammatory MERCI motifs | Overall coverage |
|------------------------------|------------------|
| Hydrophobic hydrophobic K hydrophobic hydrophobic          | 54               |
| Hydrophobic aliphatic polar N                                  | 48               |
| K hydrophobic aliphatic polar                                  | 46               |
| Hydrophobic hydrophobic K small hydrophobic                   | 45               |
| Aliphatic R hydrophobic hydrophobic                            | 44               |
| Positive tiny L                                                | 43               |
| Polar tiny hydrophobic aromatic hydrophobic                    | 43               |
| K hydrophobic L                                                | 42               |
| Hydrophobic positive tiny hydrophobic polar                    | 42               |
| Hydrophobic hydrophobic aliphatic polar small aliphatic        | 41               |
| Hydrophobic N aromatic hydrophobic                             | 41               |
Therefore, the amino acid composition was utilized to classify the two classes by developing machine learning models. After optimization of parameters, the best performing SVM-based model was selected with rbf kernel \((t = 2)\), gamma parameter \((g) = 0.005\), trade off factor \((c) = 80\) and a cost factor \((j) = 1\). The model performed with an overall accuracy of 72.9 % and MCC was measured as 0.36. The threshold independent parameter area under curve (AUC) was found to be 0.77 (Table 3; Fig. 3).

**Dipeptide composition-based models**

The models developed on AAC-based feature with vector size of 20, could not perform well both on threshold dependent as well as threshold independent parameters. Further, seeking better performance, dipeptide composition (DPC) was utilized, as input feature. The DPC-based SVM models yielded an overall accuracy of 81.5 % and MCC = 0.45 and AUC = 0.8. The optimized parameters for this model included rbf kernel \((t = 2)\), \(g = 0.001\), \(c = 10\) and \(j = 1\) (Table 3; Fig. 3).

**Physicochemical properties-based models**

While exploring different sequence-based features for better classification of PiEs from NPiEs, the physicochemical property-based models (explained in Methods) could only provide an accuracy of 79 % with MCC = 0.2, which was much lower as compared to AAC and DPC-based models (Table 3; Fig. 3) and hence, were not included in the tool.

**Hybrid model**

Among three sequence-based features mentioned above, the DPC-based model displayed the best performance. In order to further explore the possibility of better performance, the motif information was incorporated into the DPC-based models. A hybrid model of DPC and MERCI motifs were developed using the same methodology which was used to develop the DPC models and validated by tenfold cross-validation. Furthermore, three different models were developed employing three different algorithms of MERCI as described in method section. The overall accuracy of DPC-motif hybrid model with none algorithm (DPCHyb_NONE) was found to be 83 %, whereas, the DPC-motif hybrid model with Koolman–Rohm (DPCHyb_KOOL) gave an accuracy of 84 %. The DPC-motif hybrid with Betts–Russell algorithm yielded an overall accuracy of 87.6 %, MCC = 0.58 along with an AUC of 0.88 and was selected. The model was developed using rbf kernel with \(g = 0.001\), \(c = 8\), \(j = 3\) (Table 3; Fig. 3).

**Performance on validation dataset**

Although, the tenfold cross validation technique is well accepted practice in machine learning methods, there are chances of over-fitting. In order to examine the possibility that the observed performance of the final model could be due to over-optimization, the model was tested on validation dataset. The DPCHyb_BETTS model displayed an accuracy of 83.3 % with MCC = 0.43 and AUC of 0.77 on validation dataset (Table 3).

The DPCHyb_BETTS model could achieve good accuracy and MCC on DPC-motif hybrid inputs. The strategy of giving weightage worked well and enhanced the performance of model in terms of accuracy, MCC and AUC. Since the prediction model performed well on unseen validation dataset, it attests that the performance of the model is not due to over optimization.

The performance of models developed on alternate dataset is mentioned in the Additional file 5: Table S5. In brief, among models developed on alternate dataset, RandomForest performed best using AAC as input feature.

### Table 3 Performance of different classification models developed using support vector machine as machine learning technique

| Feature       | Thre | Sen  | Spec | Acc  | MCC  | AUC  | Parameters |
|---------------|------|------|------|------|------|------|------------|
| **Performance on training data** |      |      |      |      |      |      |            |
| AAC           | 0.6  | 73.58| 70.07| 72.92| 0.36 | 0.77 | t2 g:0.005 c:80 j:1 |
| DPC           | 0.4  | 86.11| 62.04| 81.53| 0.45 | 0.8  | t2 g:0.001 c:10 j:1 |
| PHY           | 0.7  | 91.25| 24.82| 78.61| 0.20 | 0.57 | t2 g:0.001 c:50 j:4 |
| DPCHyb_NONE   | 0.4  | 87.82| 62.04| 82.92| 0.48 | 0.8  | t2 g:0.001 c:20 j:1 |
| DPCHyb_KOOL   | 0.4  | 89.54| 60.58| 84.03| 0.49 | 0.85 | t2 g:0.001 c:4 j:2 |
| DPCHyb_BETTS  | 0.3  | 93.65| 62.04| 87.64| 0.58 | 0.88 | t2 g:0.001 c:8 j:3 |
| **Performance on validation data** | 0.3  | 91.1 | 50   | 83.33| 0.43 | 0.71 |            |

The hybrid model prepared using Dipeptide composition based features and MERCI displayed the best performance with an accuracy of 87.6 %. The same model showed an accuracy of 83.3 % on validation dataset.
The accuracy of the AAC-based RandomForest model was 67.2% with MCC = 0.34.

The DPCHyb_BETTS model (developed on main dataset) was incorporated in the webserver to allow the users to analyze and get predictions for their queries. It could serve as a computational substitute to the costly and time consuming experiments, as mentioned in case of HP(2–20) of H. pylori, gG2p20 of HSV-2, prion peptide PrP(106–126) and C-peptide of proinsulin. Using this web-based tool, users can sort down number of candidates responsible for proinflammatory nature of an antigen, which can further be validated by experimental studies.

Webserver and tools
The predictive modeling yielded classification tools with good accuracy for predicting the proinflammatory property of a peptide/protein and were used to develop a webserver by incorporating prediction model along with different other analysis tools. To show the translational application of "Proinflam" web server, a highly relevant example of the C-peptide from proinsulin protein is included in different modules (Peptide prediction, Protein Scan, Epitopes mapping and Similarity Search) of the web server. The C-peptide is a byproduct of proinsulin, located at position from 57 to 87 amino acids in the proinsulin protein, and was predicted as a proinflammatory epitope from the proinsulin protein using the web server. This peptide is responsible for the proinflammatory events in kidney and vasculature leading to diabetic nephropathy and atherosclerosis, respectively [11]. This example demonstrates the application and ability of the server to predict the proinflammatory epitopes in clinically relevant real proteins.

Peptide prediction
This module is designed for submission of single or batch of peptide/protein sequences in FASTA format with a length ranging from 4 to 30 amino acids. This tool runs the queries through the prediction pipeline with DPCHyb_BETTS model and classifies the queries into PiEs or NPiEs. The threshold option is provided to select the stringency of positive prediction.

As an additional function, virtual screening and designing option has been provided in the result table which allows the resubmission of the selected peptide. The virtual screening and designing involves substitution of amino acids at each position of peptide with the other 19 natural amino acids, which are further predicted in terms of PiEs or NPiEs and the results are displayed as a table. It allows the user to predict the proinflammatory nature of different variants of a given query.

Protein scan
Apart from peptide prediction module, which was meant for small length peptides, this module can be used for identification of antigenic regions in a protein, which can induce proinflammatory response in a host. The provision of window length allows user to select desired length of peptide for prediction. Similar to the peptide prediction module, this module runs query through prediction pipeline and virtual screening and provides the results as a table.

Epitope mapping
The prediction models achieved good accuracy in this study. However, user might want to investigate if there are previous reports of experimentally validated epitopes mapping to the query sequence. Therefore, 'Epitope mapping' module is developed to assist the user for mapping experimentally validated PiEs on the query sequence. Using this tool, the user can map the query sequence with PiEs and can also link to related assays in IEDB database.

Similarity search
In contrast to the epitope mapping module where exact match with experimental data is carried out, the ‘Similarity Search’ module performs Smith-Waterman search of query sequence in the database of experimentally validated PiEs. The top hits are shown with alignment and links to related assays in IEDB database.
Conclusion
The tendency of an antigen to initiate proinflammatory cascade, such as recruiting neutrophils, monocytes, and activate complement proteins, is of great importance in immunology and peptide therapeutics. Therefore, we have investigated sequence-based signatures which could be responsible for the proinflammatory nature of a peptide and developed a machine learning-based prediction tool for the prediction of proinflammatory epitopes. The computational identification of proinflammatory antigenic candidates before going for expensive and time-consuming experiments would be of great help to the scientific community. The developed computational tools are available freely for academic usage as a web server.

Methods
Dataset
The availability of clean experimental data is very crucial for any predictive modeling. Therefore, the Immune Epitope Database (IEDB) [27] for immune assays carried out for different peptide antigen was used to retrieve a clean dataset of 729 epitopes which were reported positive in assays in which any one of the proinflammatory cytokine [IL1α, IL1β, TNFα, IL12, IL18 and IL23] [25] was measured. Since these 729 epitopes were showing proinflammatory response in assays, these epitopes were considered as proinflammatory epitopes (PiEs) in this study. All those peptides which were assayed negative for proinflammatory cytokines were considered as non-proinflammatory epitopes (NPiEs). The total retrieved dataset contained 729 PiEs and 171 NPiEs ranging in length between 4 and 30 amino acids and was termed as ‘Main Dataset’. From the main dataset, 80 % of the data was assigned as training data and 20 % of the data was picked up randomly and kept as validation dataset (Fig. 1). The final training dataset contained 583 PiEs (positive data) and 137 NPiEs (negative data). The validation dataset contained 146 PiEs and 34 NPiEs.

Since the NPiEs (negative dataset) in main dataset was lesser in numbers as compare to PiEs (positive dataset), an additional dataset was constructed where the number of negative examples were kept equal to the number of positive examples. For this, 558 randomly picked non-T cell epitopes (NTCEs) data of 4–31 amino acid length were added to NPiEs of main dataset. The finalized alternate dataset contained 729 PiEs and 729 NPiEs (171 NPiEs + 558 NTCEs). The models were trained and tested in the same manner as done for the main dataset.

The main dataset was used for compositional and motif-based analysis, whereas, both main dataset and alternate dataset were used for the development of machine learning-based models. The models developed using the main dataset were incorporated in web server as prediction tool.

Input features for machine learning

Composition-based features
Amino acid composition (AAC) is the percentages of each amino acid in the given length of amino acid sequence. AAC has widely been applied in different peptide and protein composition based classification method [28, 29]. Since there are 20 amino acids, each peptide/protein is represented by 20 types of compositions or a vector size of 20.

\[
AAC(i) = \frac{\text{Total number of amino acid (i)}}{\text{Total number of all possible amino acids}} \times 100
\]

where, AAC(i) is the amino acid composition of the amino acid (i) and amino acid (i) is one of the 20 amino acids.

Di-peptide composition Similar to the AAC, dipeptide composition (DPC) has also been extensively applied in sequence-based classifications, particularly in the immune epitope prediction algorithms [30, 31]. DPC differs from AAC in having pair of amino acids and thus also provides information on local arrangement. The percentage of every possible pair (dipeptide) of amino acids was calculated. The following equation has been used for this calculation:

\[
DPC(i) = \frac{\text{Total number of dipeptides (i)}}{\text{Total number of all possible dipeptides}} \times 100
\]

where, DPC(i) is the dipeptide frequency of dipeptide (i) and the dipeptide (i) is one out of 400 dipeptides.

Physicochemical properties
In earlier studies, different physiochemical properties of amino acids have been used in several classification methods for predicting immune epitopes [24, 32] and these features are also implemented in this study for developing the prediction models. 10 different physicochemical properties were computed, namely amphipathicity, hydrophobicity, pl value, bulky side chain, hydrophilicity, net-hydrogen, steric hindrance, charge, hydropathy, molecular weight [32, 33].

Motif-based features
Identification of functional motifs in amino acid sequences has widely been exploited in functional annotation of protein/peptide sequences. Several authors have discovered immunologically relevant motifs in immunoinformatics studies [19, 34]. In this analysis, motifs specific to PiEs were identified using MERCI software (http://dtai.cs.kuleuven.be/ml/systems/merci) [35]. MERCI software is a tool to identify exclusive motifs present in positive data by comparing it with negative data. The exclusive motif discovery using MERCI was
carried out using two steps. In the first step, PiEs were taken as positive and NPiEs were taken as negative data, which yielded exclusive motifs present in PiEs. In the second step, in order to get the exclusive motifs present in NPiEs, the datasets were reversed, i.e. NPiEs as positive data and PiEs as negative data.

While discovering the motifs with MERCI software, three algorithms were adopted: (a) none (b) Koolman–Rohm, and (c) Betts–Russell. The length of motif was set as maximum of 9 amino acids because the size of binding core in both MHC I and II is 9 amino acids [36, 37]. The gap length was set to default (1).

**Hybrid feature**
The hybrid of compositional features and motif-based feature has already been used in various prediction tools by different authors [19, 20]. In this study, a hybrid of DPC and MERCI-based motifs was used to improve the classification performance. As described above, two sets of motifs were identified as exclusively found in PiEs and NPiEs, respectively. In order to make a hybrid feature, the presence of proinflammatory and non-proinflammatory motifs was searched in the peptide. If the peptide is positive for proinflammatory motif, the weight of +1 was assigned to the DPC based SVM score. Similarly, if the peptide is positive for non-proinflammatory motif, a weight of −1 was assigned to the SVM score.

**Machine-learning-based prediction models**

**Support vector machine**
In this study, support vector machine is used as the machine learning algorithm implemented using SVM-light package available at http://svmlight.joachims.org/. SVM-based models are trained with a learning algorithm where it draws an optimal hyperplane in a multi-dimensional feature space that creating a boundary dividing the datasets in two classes. Among different machine learning techniques, SVM performs well because it is effective in controlling the classifier’s capacity and associated potential for overfitting [38]. In particular, SVM has been hugely implemented in various immune epitopes prediction tools [31, 39], protein structure prediction [40] and genomic data [41] because of its ability to handle noise and large dataset [42].

**RandomForest**
In this study, RandomForest (RF) has been implemented using randomForest package in R (http://cran.r-project.org/). RF has been widely used for the binary as well as multiclass classification using nucleotide or amino acid compositions as feature inputs [43]. RF classification model, at optimized parameters with lowest out-of-bag (OOB) error, was selected for the classification purpose. The overall performance of the selected model was evaluated in terms of sensitivity, specificity, accuracy and MCC from the confusion matrix.

**WEKA-based techniques**
In addition to the SVM and RF, BayesNet, NaiveBayes, IBk (kNN) and J48 machine learning algorithms were also evaluated through WEKA package. These techniques have already been implemented in immune epitope prediction studies in earlier studies [31, 44]. Similar to SVM and RF, the performances were evaluated using sensitivity, specificity, accuracy, MCC and AUC.

**Evaluating performance of models**
Evaluation and comparison of learning methods is essential part of predictive modeling. Cross-validation technique is among most practiced techniques which involve dividing the data into two segments; the first part is used to train the model and the other holdout or test data is used to test the model. Tenfold cross validation technique has been adopted where at a given instance, nine segments were used in training the model and the rest one segment was used to test the model. The process is repeated ten times such that each segment can be tested. The performance of model was calculated by including results from all the ten predictions taken together. The performance of models can be measured by both threshold dependent as well as threshold independent parameters. For threshold independent parameter, AUC was measured which was calculated by PERF software. For threshold dependent parameters, the parameters like sensitivity (Sen), specificity (Spec), accuracy (Acc) and Matthews’s correlation coefficient (MCC) were calculated. The following equations were used for the calculation of these parameters:

\[
\text{Accuracy} = \frac{TP + TN}{TP + FN + FP + TN}
\]

\[
\text{Sensitivity} = \frac{TP}{TP + FN}
\]

\[
\text{Specificity} = \frac{TN}{TN + FP}
\]

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

where, TP = True Positive, FP = False Positive, FN = False Negative, TN = True Negative.
Additional files

**Additional file 1: Table S1.** Amino acid composition distribution between proinflammatory and non-proinflammatory data.

**Additional file 2: Table S2.** Dipeptide composition distribution between proinflammatory and non-proinflammatory data.

**Additional file 3: Table S3.** MERCI motifs extracted form proinflammatory and non-proinflammatory data.

**Additional file 4: Table S4.** Performance of models developed on main dataset, with RandomForest, BayesNet, NaiveBayes, IBk and J48.

**Additional file 5: Table S5.** Performance of different machine learning models developed on alternate dataset.

**Abbreviations**

AAC: amino acid composition; Acc: accuracy; AUC: area under curve; DPC: dipeptide composition; IEDB: immune epitope database; MCC: Matthews correlation coefficient; MHC: major histocompatibility complex; NPIEs: non-proinflammatory epitopes; NTCEs: non T Cell epitopes; OOB: out-of-bag; PHY: physicochemical properties; PIES: proinflammatory epitopes; RF: RandomForest; Sen: sensitivity; Spec: specificity; SVM: support vector machine.

**Authors’ contributions**

SG and MKM developed web server. SG and VKS conceived the work, AKS and MKM developed RandomForest models and revised the manuscript critically. SG and MKM developed web server. SG and VKS conceived the work, participated in the design of the study, and drafted the manuscript. All authors read and approved the final manuscript.

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**Availability of data and materials**

We have provided prediction models and other tools freely in the public domain at http://metagenomics.iiserb.ac.in/proinflam/ and http://metabiosis.iiserb.ac.in/proinflam/. In order to ensure reproducibility of our study and method, we have also provided the training datasets as well as validation datasets on the webserver.

**Competing interests**

The authors declare that they have no competing interests.

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