A Novel Fuzzy Approach towards *in silico* B-cell Epitope Identification Inducing Antigen-Specific Immune Response for Vaccine Design

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Introduction

Importance

B-cell epitopes are present on surface of pathogen which is recognized by the human antibody. But, it is found that only certain regions on the antigen surface induce immune response rather than whole antigen.

This makes determination of these specific regions (i.e., ‘B-cell epitopes’) that elicit antigen-specific immune response crucial for immuno-detection & immuno-therapeutic applications, including the development of safe & high efficacy vaccines.

Ahmad, A. et al., “B-cell epitope mapping for the design of vaccines and effective diagnostics.” (2016).
Introduction

Current Challenges

Identifying diagnostically useful epitopes is a difficult, time-consuming, & resource-intensive procedure. Thus, in silico prediction has gained immense attention recently due to its low cost, fast results, & less labor-intensive method compared to NMR spectroscopy & 3D X-ray structural analysis of antibody-antigen complexes.

However, one of the major problems that most established models confront is gathering huge volumes of data. Moreover, most models do not achieve high levels of accuracy. The underlying high complexity & noisy nature of the data further makes the task challenging.

More than 90% of B-cell epitopes in Protein antigens are estimated to be discontinuous/conformational (Barlow et al., Nature) making it challenging to predict them.

Ahmad, A. et al., "B-cell epitope mapping for the design of vaccines and effective diagnostics." (2016).
Previous methods and limitations

- Structure based prediction using antigen structure, propensity scales, geometric attributes
  (-) Gets very complicated for 3D structure
- Mimotope-based method that combines mimotope sequences from phage display experiments with 3D antigen structure. Locates the best alignment sequences and predicts possible epitopic regions by mapping mimotopes back to parent antigen surface.
  (-) Not found effective in actual practice to a great extent.
- Sequence-based (feature matrix formation by scoring amino acids as input antigen chain (Sweredoski et al., Ansari et al., Jespersen et al.)
  (-) These attained low AUC scores & required large quantities of data to attain low positive false rate.
- Various ML & deep learning models have been developed including attention-based LSTM networks, deep ensemble learning, etc. (Kavitha et al., Noumi et al., Sun et al.)
  (-) Training data being highly limited and skewed, most deep learning models may be prone to overfitting.

Proposed Methodology

The current work is the first to propose a ‘Fuzzy’ approach to in silico B-cell epitope prediction. The effectiveness of the proposed approach is demonstrated on severely imbalanced and limited datasets through several experiments. The results show that using the proposed method enhances both accuracy and precision when compared to existing approaches.

Further, the model is tested on the SARS-CoV-1 antigen-antibody PDB complex. The proposed approach outperforms state-of-the-art ML models trained on the same data. Results obtained indicate that applying the proposed method improves the prediction compared to the other approaches.
Model Architecture

Model Architecture

Model Architecture

General Fuzzy Min Max Neural Network

Classifying Layer Neuron (CLN)

\[ b_j(a_h, V_j, W_j) = \min_{i=1,n} \left( \min \left[ \frac{1 - f(a_{hi} - W_{ji} - V_j)}{1 - f(V_{ji} - a_{hi} - y)} \right] \right) \]

Antigen

Feature Vector \( a_h \)

V & W

Model Architecture

General Fuzzy Min Max Neural Network

Classifying Layer Neuron (CLN)

$$b_j(a_h, V_j, W_j) = \min_{i=1,n} \left( \min \left( 1 - f(a_{hi} - W_{ji}, V_j), 1 - f(V_{ji} - a_{hi}, Y_j) \right) \right)$$

Containment Compensation Neuron (CCN)

$$e_j = -1 \times T(b_j(a_h, V, W) - 1)$$

Feature Vector $$a_h$$

Antigen

Model Architecture

Classifying Layer Neuron (CLN)

\[ b_j(a_h, V_j, W_j) = \min_{i=1,n} \left( \min \left( 1 - f(a_{hi} - W_{ji} V_i), 1 - f(V_{ji} - a_{hi} V_i) \right) \right) \]

Containment Compensation Neuron (CCN)

\[ e_j = -1 \times T(b_j(a_h, V, W) - 1) \]

Overlap Compensation Neuron (OCN)

\[ d_{jp} = T(b_j(a_h, V_j, W_j) - 1) \times \left( -1 + \frac{1}{n} \sum_{i=1}^{n} \max \left( \frac{a_{hi} v_{pi}}{w_{pi} w_{ai}}, \frac{v_{pi}}{a_{hi}} \right) \right) \]

Here, \( f(x, y) = \begin{cases} 1 & \text{if } xy > 1 \\ xy & \text{if } 0 \leq xy \leq 1 \\ 0 & \text{if } xy < 0 \end{cases} \)

\[ T(x) = \begin{cases} 1 & \text{if } x \geq 0 \\ 0 & \text{if } x < 0 \end{cases} \]
Model Architecture

General Fuzzy Min Max Neural Network

Hyperbox Containment Compensation

Hyperbox Overlap Compensation

Biologically Inspired Section

Antigen

Feature Vector $a_h$

MAT. U

$e_j = -T(b_j(a_h, V, W) - 1) e_j$

MAT. Z

$T(x) = \begin{cases} 
1 & \text{if } x \geq 0 \\
0 & \text{if } x < 0 
\end{cases}$

MAT. Y

$O_i = O_i$

Feature Vector $a_h$

$O_k = O_k$

$\mathbf{f}(x, y) = \begin{cases} 
1 & \text{if } xy > 1 \\
0 & \text{if } 0 \leq xy \leq 1 \\
0 & \text{if } xy < 0 
\end{cases}$

$\mathbf{f}(x, y) = \begin{cases} 
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\end{cases}$
Model Architecture

General Fuzzy Min Max Neural Network

Hyperbox Containment Compensation

Hyperbox Overlap Compensation

Antigen

Epitope

Antibody

Epitope
Experiments

Dataset

- Epitope candidate amino acid sequence (IgG) and the activity label data from IEDB and UniProt was used. To minimize high-class imbalance, the epitope data was converted to binary classification. For this study, ‘Positive-High,’ ‘Positive-Intermediate,’ and ‘Positive Low’ samples were all regarded as ‘Positive’ samples. Nonetheless, high-class imbalance makes it harder to get positive samples compared to negative ones, making DL models inefficient.

Performance Evaluation Metrics

- In datasets skewed towards a particular class, accuracy alone may be misleading while accessing model performance. Thus, Precision & MCC along with Acc. is considered a better evaluation metric.
- Same dataset, feature selection and pre-processing techniques are used while implementing baselines.
Feature-wise KDE-plots for epitope targets (in Red) and non-epitope (in Blue) antigen regions. A weak negative co-relation can be seen.
Results and Comparison

- Proposed model outperforms ML models by a significant margin & can predict B-cell epitopes with comparable accuracy & higher precision, while being less data intensive.
- As number of training samples varies, the second-best performing model loses consistency, i.e., for n = 200, the Random Forest is the second best-performed model, its performance decreases for n = 300, where Ridge classifier & GBC classifiers are seen performing better. This illustrates that no single model exhibits robust and consistent performance while training on limited data.

Comparison of B-Cell Epitope Prediction Results with ML Classifiers for n = 200, \(imb = 0.75\). Here, learning parameters \(\theta = 0.25, \gamma = 2\) for proposed method

<table>
<thead>
<tr>
<th>Classification Model</th>
<th>Accuracy (%)</th>
<th>Precision (%)</th>
<th>Data Subset</th>
<th>Configuration - I</th>
<th>Recall (%)</th>
<th>F1-Score (%)</th>
<th>MCC (%)</th>
<th>TT (sec) (s)</th>
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<tbody>
<tr>
<td>Ada Boost Classifier</td>
<td>66.25</td>
<td>26.50</td>
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<td>47.37</td>
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</table>

Comparison of B-Cell Epitope Prediction Results with ML Classifiers for n = 300, \(imb = 0.75\). Here, learning parameters \(\theta = 0.20, \gamma = 2\) for proposed method

<table>
<thead>
<tr>
<th>Classification Model</th>
<th>Accuracy (%)</th>
<th>Precision (%)</th>
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<th>TT (sec) (s)</th>
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<td>22.22</td>
<td>34.78</td>
<td>0.3289</td>
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</table>
Results and Comparison

• In contrast to the previous experiment, where the model performance was assessed on a small & severely unbalanced data, here, we analyze the model on SARS-CoV-1 antigen sequence. It should be noted that while the model was trained using IEDB and Uniprot data, the antigen sequence was previously unknown to the model. This translates to real-time circumstances involving the prediction of B-cell epitopes of new antigen sequences. The Proposed model outperforms ML models by a significant margin (≈ 5.77% on Acc.)
Parametric Analysis

- The model's overall training time is extremely short (i.e., 1 to 5 sec), the sample test time is rather long, ranging from 5 to 10 sec. Such a substantial gap is not found in low-dimensional data categorization tasks.
- As hyperbox expansion coefficient ($\theta$) increases, number of hyperboxes created during training increases in an ‘exponential’ manner rather than a ‘linear’ one.
- The model training time climbs steeply until 0.2, after which it reduces ‘exponentially’ on both configurations.

![Graphs showing parametric study results](image-url)

Parametric study results (a) No. of hyperboxes formed vs. expansion coefficient ($\theta$) (b) Training time (sec) vs. expansion coefficient ($\theta$)
Conclusion

Summary

• Prediction of B-cell epitopes with high accuracy prior to laboratory tests can greatly reduce experimental costs while also accelerating the identification process.
• Proposed a fuzzy approach towards B-cell epitope identification.
• Addressed the problem of limited availability of datasets and high-class imbalance seen in them.

Key findings

• Fuzzy classifier-based models are more suited towards problems with limited/highly imbalanced data.
• Prediction through ‘fuzzy models’ is a promising approach towards B-cell epitope identification.

Future work

• Future work may focus on reducing model’s lengthy sample testing time.
• Developing an algorithm for tuning hyperbox expansion coefficient (θ) & fuzziness coefficient (γ).
Thanks

Feel free to contact us at
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