

# *Letters in Animal Biology*

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# **Advancing vector-borne disease prediction through functional classifier integration: A novel approach for enhanced modeling**

Mokammel Hossain Tito \*1, Most Hoor E. Jannat 1, Mst Tachhlima Aktar 2, Barshon Saha 1, Puja Das 1, Md. Kawser<sup>1</sup>, Md. Arafat Hossain<sup>1</sup>, Md. Neyamul Islam<sup>1</sup>, Muhammad Shahzad Chohan<sup>3</sup>, Shahzad Khan<sup>3</sup>

<sup>1</sup>*Bangabandhu Sheikh Mujibur Rahman Science and Technology University, Bangladesh*

borne diseases.

*<sup>2</sup> Bangladesh Agricultural University, Mymensingh, Bangladesh*

*3 Department of Biomedical Sciences, King Faisal University, Al Hafuf, Saudi Arabia*



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# **1. Introduction**

Vector-borne diseases, fueled by the intricate interplay between pathogens, vectors, and hosts, constitute a significant global health challenge. These infectious illnesses are primarily transmitted through the bites of arthropod vectors, such as mosquitoes, ticks, sandflies, and fleas, acting as intermediary hosts for pathogens like bacteria, parasites, and viruses (WHO 2014; Knudsen and Slooff 1992). Diseases like malaria, dengue fever, Lyme disease, and Zika virus have led to substantial morbidity and mortality worldwide, with changing environmental conditions, urbanization, globalization, and ecological transformations altering the distribution and abundance of vectors, thereby intensifying the global threat of these diseases (WHO 2014; Gubler 2009). The dynamic nature of vector-borne diseases necessitates advanced predictive tools to anticipate their emergence, spread, and impact on human populations (Wilson et al. 2020). In recent years, machine learning approaches, specifically function classifiers, have emerged as powerful tools in epidemiology, providing the potential to enhance our understanding of the complex interactions between environmental factors, vectors, and

pathogens (Basu et al. 2020; Uddin et al. 2019).

This study explores the application of function classifiers in predicting vector-borne diseases, emphasizing their capacity to discern intricate patterns within diverse datasets. Function classifiers, a subset of machine learning algorithms, leverage mathematical functions to map input features to specific outputs, allowing for the identification of complex relationships within multidimensional data (Almustafa 2020). These classifiers hold promise for predicting the occurrence and spread of vector-borne diseases by analyzing diverse sets of variables, including climatic conditions, land use patterns, and socio-economic factors. In contrast to traditional statistical methods, function classifiers excel at capturing non-linear and interactive effects, providing a more nuanced understanding of the intricate dynamics underlying disease transmission (Kaur et al. 2022; Raizada et al. 2021; Shaikh et al. 2023). This study aims to predict vector-borne diseases by employing function classifiers, leveraging advanced computational techniques. It focuses on identifying the most effective classifier within the function classifiers domain to enhance disease prediction accuracy and inform targeted interventions. Through rigorous

analysis, it seeks to optimize predictive models for better public health outcomes in combating vector-borne illnesses.

# **2. Materials and methods**

# **2.1 Data collection and Preprocessing**

The investigation into vector-borne diseases in this study relied on meticulously curated datasets obtained from reputable sources, including Kaggle competitions, research databases, and public health organizations (https://www.kaggle.com/ datasets/richardbernat/vector-borne-disease-prediction). A heartfelt appreciation is extended to the contributors who have devoted their efforts to compile and share invaluable data related to vector-borne diseases, enriching the scope and depth of this study. This study focuses on 11 vector-borne diseases: Chikungunya, Dengue, Zika, Yellow Fever, Rift Valley Fever, West Nile Fever, Malaria, Tungiasis, Japanese Encephalitis, Plague, and Lyme Disease. Information for this research was diligently gathered from diverse sources such as public health records, surveillance databases, and relevant literature. The dataset, a comprehensive compilation of environmental factors, demographic data, and historical disease occurrences form the bedrock of the analytical endeavours of this study.

Before embarking on model training, the amassed data



underwent an extensive preprocessing phase to ensure impeccable quality and consistency. This phase encompassed a range of tasks, including rigorous cleaning processes, addressing missing values, and standardizing data formats. Categorical variables were appropriately encoded, and numerical features underwent normalization to achieve uniformity in scale. Additionally, the dataset underwent meticulous curation using WEKA 3.9.6, where outliers and extreme values were carefully identified and removed. In pursuit of identifying the most influential features for accurate disease prediction, a robust feature selection process unfolded. This involved the strategic application of statistical techniques and domain expertise to discern and filter out less informative variables. The objective was to ensure that the input features for machine learning models used in this study were not only meaningful but also directly relevant to the intricate landscape of vector-borne diseases. This comprehensive approach underscored the commitment of this study to excellence in data handling and analysis, guaranteeing the reliability and significance of subsequent findings for the broader field of vector-borne disease research.

# **2.2 Description of utilized models**

Total of four machine learning models were used in this study including Multilayer Perceptron (MLP), Radial Basis Function Network (RBFN), Simple Logistic (SL), and Support Vector Machine (SVM) for predicting vector-borne diseases. This approach allowed for diverse exploration, comparison, and evaluation of methodologies, enhancing robustness and aiding in model selection for accurate predictions.

## *2.2.1 Multilayer Perceptron (MLP)*

A Multilayer Perceptron (MLP) model, a type of feedforward neural network, is a potent tool in the prediction of vectorborne diseases (Kumar et al. 2024). Inspired by the human brain, an MLP consists of interconnected nodes organized into layers, including an input layer, one or more hidden layers, and an output layer (Fig 2). In the context of vector-borne diseases, the MLP model processes input data related to environmental conditions, demographic factors, and historical disease occurrences through the input layer. The hidden layers, characterized by nodes employing weighted connections and activation functions, enable the network to discern intricate patterns and non-linear relationships within the data (Javaid et al. 2023; Kofidou et al. 2021). The output layer provides predictions or classifications, such as the likelihood of vectorborne disease occurrence in a specific region or population. Activation functions in the output layer depend on the nature of the prediction task, employing sigmoid functions for binary classification and softmax functions for multiclass classification. Training an MLP involves adjusting weights and biases to minimize the difference between predicted outputs and actual outcomes, typically utilizing optimization algorithms like gradient descent (Javaid et al. 2023).

MLP models excel in capturing complex patterns and relationships within datasets, especially in scenarios where traditional linear models may be inadequate (Tito et al. 2023). Their ability to handle non-linearities and interactions between

features makes them well-suited for predicting the occurrence and spread of vector-borne diseases. However, successful implementation requires careful tuning of hyperparameters such as the number of hidden layers, nodes, and learning rates (Kumar et al. 2024). In the context of vector-borne diseases, MLP models contribute to more accurate predictions, assisting public health officials in proactive measures and interventions. By leveraging the strengths of artificial neural networks, MLP models enhance the understanding of complex disease dynamics, ultimately aiding in the development of targeted strategies to mitigate the impact of vector-borne diseases on communities (Erraguntla et al. 2019).



*Fig. 2: Visual representation of Multilayer Perceptron (MLP) Model*

## *2.2.2 Radial Basis Function Network (RBFN)*

The Radial Basis Function Network (RBFN) model emerges as a potent tool in the context of predicting and understanding vector-borne diseases. The RBFN model, a type of artificial neural network, excels in capturing complex non-linear relationships within datasets (Kesorn et al. 2015), a feature crucial in deciphering the intricate dynamics of vector-borne diseases. In the realm of disease prediction, the RBFN model proves particularly adept at handling the multifaceted interplay between environmental factors, host characteristics, and pathogen behaviour. By leveraging radial basis functions, this model can effectively map the input features onto a highdimensional space, allowing it to discern subtle patterns and interactions that may elude traditional linear models. The RBFN's ability to adapt and generalize makes it a valuable asset for forecasting the spatial and temporal spread of vector-borne diseases, contributing significantly to the development of robust predictive models and informed public health

# interventions (Alfred and Obit 2021; da Silva et al. 2022).

# *2.2.3 Simple Logistic (SL)*

The Simple Logistic model, a fundamental component in predictive modeling, offers a straightforward yet powerful approach when applied to the study of vector-borne diseases. In the context of these diseases, the Simple Logistic model aims to predict the likelihood of occurrence or spread based on a set of input features. Typically used when the outcome variable is binary (e.g., presence or absence of a particular vector-borne disease), this model estimates the probability of the event occurring (Eisen and Eisen 2011). It employs the logistic function to transform a linear combination of predictor variables into a probability score, which is then thresholded to make predictions. The Simple Logistic model is particularly useful in vector-borne disease research, where understanding the factors influencing disease presence is essential for effective public health interventions. It provides a foundational framework for analyzing and interpreting the relationships between various predictors, contributing valuable insights into the dynamics of vector-borne diseases and aiding in the identification of key factors influencing their occurrence (Brownstein et al. 2002).

#### *2.2.4 Support Vector Machines (SVM)*

Support Vector Machines (SVM) serve as a formidable tool in the realm of vector-borne disease prediction. SVM, a supervised machine learning algorithm (Fig. 3), is particularly well-suited for complex datasets characterized by non-linear relationships. In the context of vector-borne diseases, where diverse factors such as climate, demographics, and historical occurrences intricately influence transmission dynamics, SVM excels at discerning patterns that may elude traditional methods (Raizada et al. 2020). By transforming the input data into a higher-dimensional space and identifying an optimal hyperplane that maximally separates different classes of vectorborne diseases, SVM achieves a remarkable ability to classify and predict instances. The flexibility of SVM allows it to adapt



*Fig. 3: Visual representation of Support Vector Machines (SVM) model (Arifuzzaman et al. 2021)*



to the intricacies of disease spread, providing a robust framework for understanding the multifaceted interactions between vectors, hosts, and environmental variables. Leveraging SVM in the prediction of vector-borne diseases not only enhances accuracy but also offers valuable insights into the intricate dynamics governing the spatial and temporal patterns of these diseases, ultimately contributing to more effective public health interventions (Fuchida et al. 2017; Munirathinam et al. 2023).

#### **3. Result and Discussions**

From the evaluation of various machine learning models for predicting vector-borne diseases, the performance metrics reveal intriguing insights. The Multilayer Perceptron (MLP) model exhibited the highest accuracy at 92%, closely followed by the Simple Logistic (SL) and Support Vector Machine (SVM) models at 88% and 90.87%, respectively. The MLP model also outshined others in terms of precision, recall, and F-Measure, indicating its superior ability to correctly classify instances and balance between false positives and false negatives (Table 1). However, the SVM model demonstrated noteworthy efficiency with the lowest processing time at 0.3 seconds, suggesting its computational advantage. On the other hand, the RBFN lags behind in terms of accuracy, precision, and recall.

The results showcased the trade-offs between accuracy, computational efficiency, and other performance metrics, emphasizing the need for a nuanced decision in selecting the most suitable model for vector-borne disease prediction. Considering the holistic evaluation, the SVM model emerged as a compelling choice (Fig 4), striking a balance between high accuracy and efficient processing time, making it a promising candidate for real-time applications in public health interventions. The application of machine learning (ML) in predicting vector-borne diseases represents a pivotal advancement in the field of public health. The results presented



*Fig. 4: Visual representation showcasing the efficacy of different ML models*

in this study underscore the efficacy of ML models, specifically Multilayer Perceptron (MLP), Radial Basis Function Network (RBFN), Simple Logistic (SL), and Support Vector Machine (SVM), in discerning complex patterns inherent in the transmission dynamics of vector-borne diseases.

The high accuracy achieved by these models, particularly the MLP and SVM, suggests their potential as robust tools for early detection and proactive management of outbreaks. The comprehensive evaluation metrics, including precision, recall, and F-Measure, provide a holistic understanding of the models' performance, highlighting their ability to balance true positives, true negatives, false positives, and false negatives. Kappa Statistics provide insights into the reliability and validity of the predictive models by quantifying the agreement between predicted and observed outcomes. This metric helped for the evaluation of the model's performance while considering the possibility of chance agreement, thus offered a more nuanced evaluation of the predictive accuracy and robustness of the disease prediction models. Furthermore, the consideration of processing time unveils practical implications for real-time applications, with the SVM model demonstrating notable efficiency. The study not only contributes valuable insights into the comparative performance of ML models but also underscores the significance of selecting an appropriate model that aligns with the specific needs of vector-borne disease prediction. As this study navigates the landscape of emerging infectious diseases, ML-driven predictive models stand as indispensable tools, fostering a proactive and data-driven approach to mitigate the impact of vector-borne diseases on global health.

# **4. Conclusions**

Utilization of machine learning (ML) function classifiers did indeed contribute to more accurate prediction of outbreaks and identification of areas at higher risk for vector-borne diseases. By analyzing various data inputs such as climate conditions, environmental factors, host population dynamics, vector abundance, and historical disease incidence, ML models could detect patterns and relationships that humans might have overlooked, thus enhancing predictive capabilities. After the evaluation of function classifier models for predicting vectorborne diseases, Support Vector Machine (SVM) showed the best result based on different evaluation criteria. With an impressive accuracy of 90.87% and notable efficiency reflected in the lowest processing time at 0.3 seconds, the SVM model emerges as a compelling choice for real-time applications in public health interventions. Its ability to strike a balance between high accuracy and efficient processing highlights its potential as a robust tool for early detection and proactive management of vector-borne disease outbreaks. The comprehensive evaluation metrics, including precision, recall, and F-Measure, further validate the SVM model's efficacy in achieving a nuanced understanding of the complex dynamics underlying disease transmission.

#### **Recommendations and future work**

While recognizing the potential for further enhancement in this dataset, its current utility is constrained by specific limitations. Machine learning holds promise for predicting various diseases, including but not limited to Brucellosis, Subclinical Mastitis, Anthrax, and LSD.

# **Declarations**

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