VALLEY-FORECAST: FORECASTING COCCIDIOIDOMYCOSIS INCIDENCE VIA ENHANCED LSTM MODELS TRAINED ON COMPREHENSIVE METEOROLOGICAL DATA

RESEARCH ARTICLE

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ABSTRACT

Coccidioidomycosis (CM), commonly known as Valley Fever, is a fungal infection caused by Coccidioides species that poses a significant public health challenge, particularly in the semi-arid regions of the Americas, with notable prevalence in California and Arizona. Previous epidemiological studies have established a correlation between CM incidence and regional weather patterns, indicating that climatic factors influence the fungus's life cycle and subsequent disease transmission. This study hypothesizes that Long Short-Term Memory (LSTM) and extended Long Short-Term Memory (xLSTM) models, known for their ability to capture long-term dependencies in time-series data, can outperform traditional statistical methods in predicting CM outbreak cases. Our research analyzed daily meteorological features from 2001 to 2022 across 48 counties in California, covering diverse microclimates and CM incidence. The study evaluated 846 LSTM models and is currently assessing 176 xLSTM models with various fine-tuning metrics. To ensure the reliability of our results, these advanced neural network architectures are cross analyzed with Baseline Regression and Multi-Layer Perceptron models, providing a comprehensive comparative framework. Preliminary results indicate that LSTM-type architectures outperform traditional methods, demonstrating an increased predictive accuracy of up to 87%. This improvement in predictive capability suggests a strong correlation between temporal microclimatic variations and regional CM incidences. The increased predictive power of these models has significant public health implications, potentially informing strategies for CM outbreak prevention and control. These findings contribute to the ongoing efforts to address CM, offering a new approach to understanding and potentially mitigating the impact of the disease in affected regions.

Keywords Coccidioidomycosis · LSTM · xLSTM · Epidemiological forecasting · Meteorological data · Machine learning · Time-series analysis · Public health · California · Climate factors

1 Introduction

Coccidioidomycosis (CM), or Valley Fever, is a fungal disease that presents a growing challenge to public health systems in the Americas, particularly in arid and semi-arid regions [7]. The causative agents, Coccidioides immitis and C. posadasii, are soil-dwelling fungi that release spores into the air when the soil is disturbed, leading to respiratory infections in humans and animals [1, 7]. The disease's impact extends beyond individual health outcomes, affecting communities, healthcare systems, and local economies in endemic areas [8].

The spectrum of CM manifestations is broad and often unpredictable. While many infected individuals remain asymptomatic, others experience symptoms from mild flu-like illness to severe pneumonia [7]. In rare but serious cases, the infection can disseminate beyond the lungs, leading to meningitis, osteomyelitis, or cutaneous lesions. These severe

forms of CM can result in long-term disability or even death, particularly among immunocompromised individuals, pregnant women, and certain ethnic groups who are at higher risk for disseminated disease [5, 7].

Current strategies to combat severe CM cases exist but are limited in effectiveness and must be better established [7]. Early diagnosis relies on a combination of clinical suspicion, radiological findings, and serological tests, which can be challenging in non-endemic areas where awareness is low [7]. Treatment typically involves antifungal medications such as Fluconazole or Itraconazole for symptomatic cases, with more aggressive therapies reserved for severe or disseminated disease [7]. However, these treatments are often prolonged, costly, and associated with significant side effects [7]. Prevention efforts include dust control measures, public education campaigns, and targeted interventions for high-risk populations to reduce exposure to fungal spores. While these approaches have been implemented, particularly in prison systems, Coccidioidomycosis incidence rates continue to rise as seen in Figure 1. Ongoing research is focused on developing more effective risk-reduction strategies, especially for vulnerable groups in endemic areas [11].

Coccidioidomycosis Cases by County and Year (2001-2022)

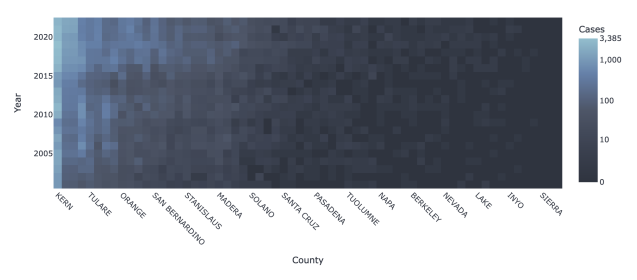


Figure 1: Heatmap distribution of Coccidioidomycosis (CM) cases across California counties from 2001 to 2022. The color intensity represents the number of cases on a logarithmic scale, ranging from 0 (darkest) to 3,385 (lightest). Counties are sorted by total case count, with Kern County showing consistently high incidence. The visualization reveals a general upward trend in CM cases over time, with notable increases in endemic areas and sporadic outbreaks in less affected counties.

The economic impact of CM is substantial and growing. In California alone, the lifetime cost burden associated with CM is estimated to exceed \$700 million, covering direct medical expenses, lost productivity, and ongoing care for chronic cases. With incidence rates showing a consistent upward trend, particularly in California and Arizona, there is an urgent need for more effective predictive models and preventive strategies [8].

Our research addresses this need by developing advanced machine-learning models based on comprehensive meteorological data to forecast CM incidence cases. We focus on California, analyzing daily weather features from 2001 to 2022 across 48 counties, representing a diverse range of microclimates and CM incidences. By leveraging the power of Long Short-Term Memory (LSTM) networks and their extended variant (xLSTM), we aim to capture the complex, long-term dependencies between environmental factors and disease occurrence [14].

The choice of LSTM and xLSTM architectures is motivated by their proven effectiveness in handling time-series data and their ability to retain information over extended periods [18]. These characteristics are particularly relevant to our study, given the seasonal nature of CM and the potential lag between environmental conditions and disease manifestation [2, 3, 6, 9, 10, 17]. Our approach builds upon previous studies that have established correlations between climatic variables and CM rates using regression models, particularly noting peak incidences during the fall season following wet and then dry seasons [3, 6, 17].

By developing more accurate predictive models, our research aims to provide public health officials with a powerful tool for anticipating CM outbreaks and allocating resources more effectively. The potential applications of this work extend

beyond California, offering a framework that could be adapted to other endemic regions facing similar challenges with fungal diseases influenced by environmental factors.

The remainder of this paper is structured as follows: Section 2 details our comprehensive data preparation and processing methodologies, providing a foundation for our analytical approach. Section 3 presents an in-depth exploration of the models employed in this study, elucidating their theoretical underpinnings, practical implementations, and individual performance outcomes. In Section 4, we synthesize and compare the results across all models, offering a comprehensive view of their relative efficacies in predicting CM incidence. Finally, Section 5 concludes with a discussion of our findings and their implications for public health strategies and proposes promising avenues for future research in this domain.

2 Methodologies

2.1 Hardware Specifications

All computational tasks were performed on a single, high-performance workstation to maintain consistency and enable efficient processing of large-scale meteorological and epidemiological data. The system specifications were: Processor: AMD Ryzen 9 5950X (16 cores, 32 threads, Memory: 128 GB DDR4 RAM, Graphics Processing Unit: NVIDIA GeForce RTX 4090 with 24 GB VRAM.

2.2 Data Sourcing

Our study utilized two primary data sources to construct a comprehensive dataset for analysis. The epidemiological data on Coccidioidomycosis (CM) cases were obtained from the California Department of Public Health (CDPH). This dataset contained a temporal range from 2001 to 2022, a geographic scope of 64 distinct locations within California, and four key variables (county name, year, number of cases, and incidence rate). See Figure 2 to the right for the counties mapped to the cumulative incidences used in this study. For the meteorological component, we focused on the 48 counties from the CDPH dataset that exhibited the most significant CM case numbers annually. This selection criterion helped to guarantee a lower chance of counties being used with inaccurate case numbers that would lead to erroneous results. The weather data was sourced using the OpenWeather API's historical data service (https://openweathermap.org/api). This dataset included an hourly temporal resolution, a temporal range of 2001-2002, and a geographical scope of the 48 counties selected from the epidemiological dataset. The OpenWeather API provided a rich set of meteorological variables, including but not limited to temperature, humidity, precipitation, wind speed, and atmospheric pressure. Figure 3 below provides a visualization of these weather variables spanning over the year's average to represent the trends inside the meteorological data. The high temporal resolution of this data allowed for a detailed analysis of weather patterns concerning CM incidence.

2.3 Data Preparation

The preparation of our dataset involved several steps to guarantee data quality, relevance, and suitability for our machine-learning models. The epidemiological and meteorological datasets were integrated based on temporal and geographical alignment. We excluded the incidence rate as an outcome variable for the epidemiological data refinement due to reliability concerns flagged by the CDPH. We also filtered down the number of geographical locations from 64 initial locations to 48 locations, as shown in Figure 2. This was due to the 16 locations needing to meet the minimum threshold for having at



Figure 2: Distribution of Coccidioidomycosis (CM) Cases in California Counties from 2001-2022. This choropleth map illustrates the cumulative number of CM cases across California counties over a 22-year period. The color gradient ranges from light (fewer cases) to dark red (more cases), revealing significant regional variations in CM incidence. Southern and central California counties show notably higher case numbers, particularly in the San Joaquin Valley and Los Angeles area, while northern and coastal regions generally report fewer cases.

least 60 percent incidences for the timespan. This choice was to maintain the data integrity and thin out locations with possibly less accurate reporting.

We temporally aligned the dataset for the meteorological data processing to fit the same timeframe (2001-2022) for the epidemiological dataset. From there, the original resolution of the dataset needed to be lowered to better match our needs. So, we aggregated the sequence length from 8760 (hourly for a year) to 365 (daily averages for a year). This decision ensured that the model was manageable and fit the small training set. Additionally, we opted to feature encode the categorical weather descriptors (e.g., cloudy, sunny). This method made it possible for us to preserve the features for numerical analysis [19]. Finally, a MinMax normalization was applied to all weather features to standardize the scale of different meteorological variables, improving model performance and stability. LSTMs have been found to perform much better with accuracy when normalization is applied [19,20].

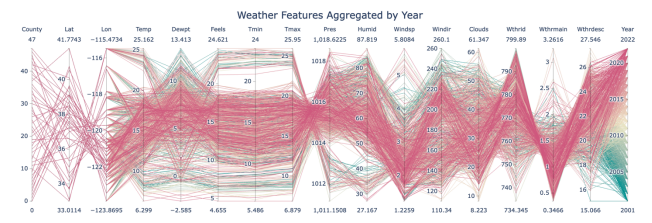


Figure 3: Aggregated Weather Features for California Counties, 2001-2022. This parallel coordinates plot displays various meteorological parameters across years and counties. Each line represents a county-year combination, with different colors indicating each year. The plot shows the relationships and variations among features such as latitude, longitude, temperature, dewpoint, precipitation, humidity, wind speed, and cloud cover. This visualization helps to illustrate the complex interplay of weather conditions that may influence Coccidioidomycosis (CM) incidence over time and space.

Data Structuring

The final dataset was shaped to accommodate the requirements of LSTM models. The final input shape was (1056, 365, 19), i.e. (number of sequences, days per year, features per day). The final output shape consequently is (1056), corresponding to the CM case counts for each yearly sequence.

Data Augmentation

To strengthen model robustness and mitigate overfitting, we implemented a data augmentation strategy following established methods [21, 22]. The dataset was randomly split into training (844 sequences), validation (158 sequences), and test (54 sequences) sets. The training set was then augmented 20-fold, a decision made through empirical observation of model performance on augmented datasets spanning from 2-fold to 100-fold increases. Augmentation was achieved by adding Gaussian noise to the original data. The noise variance was scaled logarithmically from 1e-2 to 1e-5, creating 20 distinct noise levels. The augmentation process can be described by the following equations:

$$N_a = 20$$
 number of augmentations (1)

$$\sigma_i = 0.01 \cdot 10^{-i/19}, \qquad i = 0, 1, \dots, N_a - 1$$
 (2)

$$\epsilon_i \sim \mathcal{N}(0, \sigma_i^2)$$
 noise distribution (3)

$$x_i^a = x + \epsilon_i,$$
 $i = 0, 1, \dots, N_a - 1$ (4)

$$\epsilon_i \sim \mathcal{N}(0, \sigma_i^2)$$
 noise distribution (3)
 $x_i^a = x + \epsilon_i,$ $i = 0, 1, \dots, N_a - 1$ (4)
 $y^a = N_a \cdot y$ augmented labels (5)

Where x and y denote the original training inputs and labels, respectively. $\mathcal{N}(0, \sigma_i^2)$ represents Gaussian noise with standard deviation σ_i , and x_i^a and y_i^a denote the augmented dataset and labels, respectively.

The composition of the final dataset created a training set of 17,724 sequences (844 * 21, including the original data), a validation set of 158 sequences, and a test set of 54 sequences. This augmentation strategy significantly expanded

our training data while preserving the integrity of our validation and test sets, which is critical for unbiased model evaluation [22]. The resulting dataset provided a strong foundation for training our models, balancing the need for substantial training data with the imperative to maintain distinct validation and test sets for reliable model assessment [21, 22].

3 Models

Our study utilized a hierarchical approach to model development, implementing four distinct predictive models. This strategy allowed for a systematic performance comparison across different model complexity levels and architectural designs. The models implemented were:

- 1. Baseline Regressor
- 2. Multilayer Perceptron (MLP)
- 3. Long Short-Term Memory (LSTM) Network
- 4. Extended Long Short-Term Memory (xLSTM) Network

3.1 Model Comparison Strategy

Our approach to model comparison is designed to provide a comprehensive understanding of each architecture's strengths and limitations. We start with a baseline regressor to establish a minimum performance threshold, then move to an MLP to demonstrate the capability of non-recurrent neural networks in capturing relevant patterns. The LSTM serves as a strong baseline for recurrent architectures, leveraging its ability to model long-term dependencies, while the xLSTM allows us to assess whether the latest innovations in recurrent neural network design offer tangible benefits for our specific prediction task. By implementing this diverse set of models, we look to identify the most effective approach for predicting CM incidence and gain insights into the relationship between meteorological factors and disease occurrence.

3.2 Baseline Regressor

We implemented a simple baseline regressor as a benchmark for our study. This model, though basic, provides an initial reference for evaluating more advanced predictive models [23]. The baseline regressor offers a straightforward comparison point by predicting the mean of all output values without considering input features. This allows us to measure the performance improvements achieved by more complex models [23]. It is designed to ignore input features and instead predict a constant value for all instances. This constant value is computed as the mean of the target variable in the training set. Mathematically, for a set of n training examples with target values $y_1, y_2, ..., y_n$, the Baseline Regressor predicts:

$$\hat{y} = \frac{1}{n} \sum_{i=1}^{n} y_i$$
 Baseline prediction (6)

In our implementation, we trained the Baseline Regressor on the same dataset used for all other models in our study, consisting of 17,724 instances. We used the test set of 54 instances to evaluate its performance. We chose Root Mean Squared Error (RMSE) as our primary evaluation metric for assessing prediction quality across all models. RMSE is calculated as the square root of the average of squared differences between predicted and actual values:

$$RMSE = \sqrt{\frac{1}{n} \sum_{i=1}^{n} (y_i - \hat{y}_i)^2}$$
 Root Mean Squared Error (7)

Where y_i are the true values and \hat{y}_i are the predicted values. This metric provides a clear and interpretable measure of prediction accuracy [27].

The results of our Baseline Regressor yielded an RMSE of 538.24. This figure provides a concrete benchmark against which we can compare the performance of our more sophisticated models. The Baseline Regressor's performance serves as an essential context for interpreting the results of more complex models in our study [23]. Any model that fails to significantly outperform this baseline would be considered inadequate. Conversely, the degree to which other models improve upon this baseline RMSE of 538.24 serves as a quantitative measure of their effectiveness in capturing the underlying patterns in the data [23].

It's important to note that the Baseline Regressor does not require hyperparameter tuning or optimization due to its simplistic design. This characteristic makes it an ideal starting point for our model comparison, as it provides a stable and consistent benchmark. As we progress to more complex models, any improvements in RMSE can be directly attributed to the model's ability to capture more nuanced patterns in the data rather than to extensive tuning or optimization processes.

3.3 Multilayer Perceptron (MLP)

Our study implemented a Multilayer Perceptron (MLP) as an intermediate model, bridging the gap between the simple baseline regressor and more complex recurrent architectures. This feedforward neural network provides valuable insights into the capability of traditional neural networks to capture underlying patterns in our dataset. The MLP's performance is a valuable comparison point, helping us quantify the benefits of utilizing recurrent architectures for our time-series prediction task [23]. The theoretical foundation of the MLP is rooted in feedforward artificial neural networks [24]. Characterized by at least three layers of nodes - an input layer, one or more hidden layers, and an output layer - the MLP's architecture facilitates learning non-linear relationships in data through activation functions and backpropagation for weight adjustment [24]. The primary strengths of MLPs lie in their ability to approximate complex, non-linear functions, handle high-dimensional input spaces, and learn hierarchical representations of data [24].

For our study, we implemented a 3-layer MLP. The input layer's dimensionality matched our feature space, guaranteeing all relevant meteorological data was incorporated. The hidden layer utilized a Rectified Linear Unit (ReLU) activation function, chosen for its computational efficiency and effectiveness in mitigating the vanishing gradient problem [25] [26]. The ReLU function, defined as f(x) = max(0, x), introduces non-linearity into the model without the computational overhead of more complex activation functions [25]. The output layer consisted of a single node for regression, appropriate for our task of predicting CM incidence.

The best performant model was trained over 1500 epochs, allowing ample time for convergence. We employed the Adaptive Moment Estimation (Adam) optimizer with a learning rate 5e-5. Adam was selected for its ability to adapt the learning rate during training, potentially leading to faster convergence and improved performance on complex loss landscapes [12]. Our loss function was Mean Squared Error (MSE), which clearly measures prediction accuracy and is differentiable, making it suitable for gradient-based optimization [28].

To evaluate the MLP's performance, we used the Root Mean Squared Error (RMSE) metric, which was consistent with our baseline regressor evaluation. This allowed for direct comparison between models. The final results yielded an RMSE of 115.1793, representing a substantial improvement over the baseline regressor's RMSE of 538.24. This improvement indicates two key points: first, significant non-linear relationships in the data that the MLP can capture, and second, our chosen architecture and hyperparameters are well-suited to the problem domain.

The magnitude of improvement - approximately a 78.6% reduction in RMSE - suggests that the meteorological features contain predictive solid power for CM incidence, which the MLP can leverage effectively. This substantial performance gain shows the value of neural network approaches in capturing complex patterns within our dataset.

The significance of the MLP's performance extends beyond its numerical improvement over the baseline. It is an intermediate benchmark between our baseline regressor and more complex recurrent models. Demonstrating the potential of neural networks to capture relevant patterns in our dataset provides a reference point for assessing the additional value that recurrent architectures might bring to the task.

3.4 Long Short-Term Memory (LSTM) Network

The Long Short-Term Memory (LSTM) model represents this study's primary recurrent neural network architecture. Renowned for its ability to capture long-term dependencies in sequential data, the LSTM is particularly well-suited for predicting Coccidioidomycosis (CM) incidence based on time-series meteorological data. This model serves as a strong baseline for assessing the performance of more advanced recurrent architectures and provides novel insights into the temporal aspects of our prediction task. LSTMs are a specialized form of Recurrent Neural Networks (RNNs) designed to address the vanishing gradient problem inherent in traditional RNNs [15]. This makes them particularly effective for processing and predicting time series data due to their ability to capture long-term dependencies [15] [18]. The key innovation of LSTMs is the introduction of a memory cell, which allows the Network to remember or forget information over extended sequences selectively [15]. This mechanism is implemented through three primary gates: the input gate, which controls the flow of new information into the cell state; the forget gate, which determines what information should be discarded from the cell state; and the output gate, which regulates the information output from the cell state [15]. These gates enable LSTMs to maintain and update relevant information over long sequences, making

them particularly effective for tasks involving time-series data like our CM incidence prediction [13] [15] [16]. The LSTM forward pass is as follows:

$$f_t = \sigma(W_f \cdot [h_{t-1}, x_t] + b_f)$$
 forget gate (8)

$$i_t = \sigma(W_i \cdot [h_{t-1}, x_t] + b_i)$$
 input gate (9)

$$\tilde{C}_t = \tanh(W_C \cdot [h_{t-1}, x_t] + b_C)$$
 candidate cell state (10)

$$C_t = f_t \odot C_{t-1} + i_t \odot \tilde{C}_t$$
 cell state (11)

$$o_t = \sigma(W_o \cdot [h_{t-1}, x_t] + b_o)$$
 output gate (12)

$$h_t = o_t \odot \tanh(C_t)$$
 hidden state (13)

Where f_t is the forget gate, i_t is the input gate, \tilde{C}_t is the candidate cell state, C_t is the cell state, o_t is the output gate, and h_t is the hidden state. σ represents the sigmoid function, W terms are weight matrices, b terms are bias vectors, and * denotes element-wise multiplication.

To determine the most effective LSTM configuration for our task, we conducted an extensive hyperparameter search, exploring 846 distinct model configurations using a cardinal search method, a form of grid search. This thorough exploration of the parameter space allowed us to identify the optimal architecture for our specific prediction task. The optimal architecture identified through this process consists of two LSTM layers with a hidden size of 256 units each, followed by a three-layer Multilayer Perceptron (MLP) with ReLU activation functions. We also applied a 10% dropout rate between all layers except the final output layer to prevent overfitting [13]. This architecture, as shown in Figure 4, combines the sequential learning capabilities of LSTMs with the non-linear function approximation of MLPs, potentially capturing both temporal dependencies and complex feature interactions [29].

We utilized the Adam optimizer for the training process, with the learning rate and other hyperparameters determined during our search process. We chose Mean Squared Error (MSE) as our loss function, which is appropriate for our regression task. The number of training epochs was also optimized during our hyperparameter search to ensure convergence without overfitting. The performance of our optimized LSTM model was evaluated using the Root Mean Squared Error (RMSE) metric, which is consistent with our previous models. The LSTM achieved an RMSE of 57.3489, significantly improving the baseline regressor (RMSE: 538.24) and the MLP model (RMSE: 115.1793). This performance demonstrates an 89.3% reduction in RMSE compared to the baseline regressor and a 50.2% reduction compared to the MLP model. These substantial improvements suggest two key insights: first, temporal dependencies play a fundamental role in predicting CM incidence, and second, the LSTM's ability to capture long-term patterns in the meteorological data significantly enhances predictive

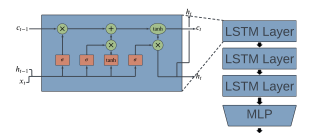


Figure 4: Custom LSTM Architecture for CM Incidence Prediction. The left panel shows the internal structure of a single LSTM cell and reveals its gates and information flow. The right panel outlines the overall model architecture, consisting of multiple stacked LSTM layers followed by a 3-layer Multilayer Perceptron (MLP). This design combines the LSTM's ability to capture long-term temporal dependencies in meteorological data with the MLP's capacity for complex non-linear mappings, tailored for predicting Coccidioidomycosis (CM) incidence based on time-series weather features.

accuracy. The magnitude of improvement over the MLP model, which does not account for temporal dependencies, pinpoints the importance of considering the sequential nature of our data in making accurate predictions. The significance
of our LSTM results extends beyond their numerical superiority. This study represents the first application of LSTM
networks for predicting CM incidence using temporal meteorological data. The substantial performance improvement
over non-recurrent models demonstrates the importance of considering long-term weather patterns in CM prediction.
This finding has potential implications for how we approach epidemiological forecasting, particularly for diseases with
environmental risk factors. The success of this LSTM architecture provides a strong foundation for further exploration
of recurrent models in epidemiological forecasting. Capturing long-term dependencies in environmental data can
significantly improve our ability to predict disease incidence. This opens new avenues for research and public health
interventions, where long-term weather forecasts could be used to anticipate and prepare for potential CM outbreaks.

3.5 Extended Long Short-Term Memory (xLSTM) Network

The Extended Long Short-Term Memory (xLSTM) network represents the cutting edge of our recurrent neural network implementations in this study. As a novel extension of the traditional LSTM architecture, the xLSTM incorporates

several innovative features to improve its ability to capture complex long-term relationships in sequential data. These innovations include exponential gating, memory mixing, and matrix memory, which we hypothesized would allow the xLSTM to model the intricate relationships more effectively between long-term meteorological patterns and Coccidioidomycosis (CM) incidence. The theoretical underpinnings of the xLSTM build upon the foundation laid by traditional LSTMs. While LSTMs addressed the vanishing gradient problem through their gating mechanisms, xLSTMs take this concept further [30]. The exponential gating mechanism in xLSTMs allows for more fine-grained control over information flow, capturing subtle temporal dependencies that might be missed by standard LSTMs [30]. This ability to capture subtle temporal dependencies is particularly relevant in our context, where subtle changes in weather patterns over extended periods might significantly influence CM incidence. Memory mixing, another key innovation in xLSTMs, enables the Network to combine information from different time scales more effectively [30]. In the context of our meteorological data, this could allow the model to simultaneously consider short-term weather fluctuations and long-term climate trends, providing a more comprehensive basis for prediction. The matrix memory component of xLSTMs expands the Network's capacity to store and process complex patterns, potentially allowing for more nuanced representations of the relationship between weather patterns and CM incidence [30]. The xLSTM architecture comprises two LSTM cell variants: the sLSTM and the mLSTM as shown in Figure 5. The sLSTM introduces scalar updates and new memory mixing, with its specific mechanics detailed in the sLSTM forward pass below. Conversely, the mLSTM features matrix memory and a covariance update rule, as elaborated in the following mLSTM forward pass section. In their original paper, Beck et al. organized these variants into blocks, like other approaches in large language model research [cite Beck and another paper that uses blocking for llms]. However, for our purposes and given our limited data, this block structure was deemed unnecessary, as it could lead to overfitting before capturing relevant information. Instead, we explored the xLSTM variants at the cell level by creating different stack combinations of mLSTM and sLSTM cells. The forward passes of both variants, presented next, provide a deeper understanding of their respective functionalities and how they contribute to the overall xLSTM architecture.

3.6 sLSTM Forward Pass

The sLSTM forward pass is defined by the following equations:

$c_t = f_t \cdot c_{t-1} + \iota_t \cdot z_t$	cell state	(14)
$n_t = f_t \cdot n_{t-1} + i_t$	normalizer state	(15)
$h_t = o_t \cdot \tilde{h}_t, \tilde{h}_t = c_t/n_t$	hidden state	(16)
$z_t = \varphi(\tilde{z}_t), \tilde{z}_t = w_z^\top x_t + r_z h_{t-1} + b_z$	cell input	(17)
$i_t = \exp(\tilde{i}_t), \tilde{i}_t = w_i^{\top} x_t + r_i h_{t-1} + b_i$	input gate	(18)
$f_t = \sigma(\tilde{f}_t) \text{ OR } \exp(\tilde{f}_t), \tilde{f}_t = w_f^{\top} x_t + r_f h_{t-1} + b_f$	forget gate	(19)
$o_t = \sigma(\tilde{o}_t), \tilde{o}_t = w_o^{T} x_t + r_o h_{t-1} + b_o$	output gate	(20)

We broadcast the original LSTM gating techniques, i.e., input- and/or hidden-dependent gating plus bias term, to the new architectures. Exponential activation functions can lead to large values that cause overflows. Therefore, we stabilize gates with an additional state m_t (Milakov & Gimelshein, 2018):

$$m_t = \max(\log(f_t) + m_{t-1}, \log(i_t))$$
 stabilizer state (21)

$$i'_t = \exp(\log(i_t) - m_t) = \exp(i_t - m_t)$$
 stabil. input gate (22)

$$f_t' = \exp(\log(f_t) + m_{t-1} - m_t)$$
 stabil. forget gate (23)

The mLSTM forward pass is defined by the following equations:

$$C_t = f_t \cdot C_{t-1} + i_t \cdot v_t k_t^{\top}$$
 cell state
$$n_t = f_t \cdot n_{t-1} + i_t \cdot k_t$$
 normalizer state (25)

$$h_t = o_t \odot \tilde{h}_t, \quad \tilde{h}_t = C_t \cdot q_t / \max\{n_t^\top q_t, 1\}$$
 hidden state (26)

$$q_t = W_q x_t + b_q query input (27)$$

$$k_t = \frac{1}{\sqrt{d}} W_k x_t + b_k$$
 key input (28)

$$v_t = W_v x_t + b_v$$
 value input (29)

$$i_t = \exp(\tilde{i}_t), \quad \tilde{i}_t = w_i^{\mathsf{T}} x_t + b_i$$
 input gate (30)

$$f_t = \sigma(\tilde{f}_t) \text{ OR } \exp(\tilde{f}_t), \quad \tilde{f}_t = w_f^{\top} x_t + b_f$$
 forget gate (31)

$$o_t = \sigma(\tilde{o}_t), \quad \tilde{o}_t = W_o x_t + b_o$$
 output gate (32)

mLSTM can have multiple memory cells like the original LSTM. For mLSTM, multiple heads and multiple cells are equivalent as there is no memory mixing. In order to stabilize the exponential gates of mLSTM, we use the same stabilization techniques as for sLSTM, see Equation (21). Since the mLSTM has no memory mixing, this recurrence can be reformulated in a parallel version. For more details we refer to Appendix A.3.

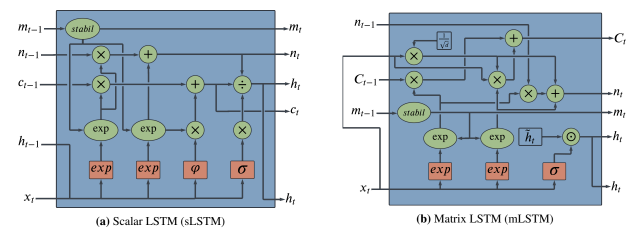


Figure 5: Architectures of the Extended Long Short-Term Memory (xLSTM) variants: (a) Scalar LSTM (sLSTM) and (b) Matrix LSTM (mLSTM). These diagrams illustrate the internal structures and information flow within each cell type, both variants of the traditional LSTM designed for enhanced performance in time-series analysis. Key components in both include the cell state (c_t or C_t), hidden state (h_t), and various gates with exponential and stabilization functions. The sLSTM introduces scalar updates and novel memory mixing techniques, while the mLSTM employs matrix operations and additional computational paths. Both variants aim to improve the capacity to capture complex temporal dependencies in meteorological data for Coccidioidomycosis (CM) incidence prediction. Notable features include stabilization mechanisms (stabil), multiple exponential activations, and functions like ϕ (tanh) and σ (sigmoid), which contribute to the cells' ability to process and retain relevant information over extended sequences while mitigating issues like vanishing gradients. These enhancements potentially allow the xLSTM architecture to model intricate long-term relationships in the data more effectively than standard LSTM models.

From these two LSTM variants, we can begin to form our xLSTM models. In doing so, we stacked the different cells into layers and then fed the output from the final LSTM layer into a three-layer MLP in the same way as we did our original LSTM architecture, as shown in Figure 6 below.

To implement the xLSTM for our study, we adapted our hyperparameter search process to accommodate the additional search over 176 different stack configurations for the xLSTM. This extensive search allowed us to identify an optimal xLSTM architecture tailored to our specific prediction task, giving it a fair representation against the traditional LSTM. The training process for the xLSTM model followed a similar pattern to our LSTM implementation, utilizing the Adam optimizer and Mean Squared Error (MSE) as the loss function. However, we paid particular attention to the learning rate schedule, as the more complex xLSTM architecture can be more sensitive to the learning rate changes we found. We also implemented a more aggressive dropout strategy of 15% to prevent overfitting, given the increased capacity of the xLSTM to memorize training data [13] [30]. In evaluating the performance of our xLSTM model, we continued to use the Root Mean Squared Error (RMSE) metric for consistency with our previous models. The optimized xLSTM

achieved an RMSE of 48, further improving our already strong LSTM results (RMSE: 57.34887). This corresponds to a 16.3% reduction in RMSE compared to the LSTM model and a remarkable 91.1% reduction compared to our initial baseline regressor.

The xLSTM model's superior performance provides several key insights. First, it confirms our hypothesis that the xLSTM architecture's advanced features can capture more complex relationships in our time-series data. The improvement over the standard LSTM suggests that there are indeed subtle long-term dependencies in the meteorological patterns influencing CM incidence that the xLSTM is better equipped to model.

Furthermore, the success of the xLSTM in our specific task of CM incidence prediction demonstrates the potential of these advanced, recurrent architectures in epidemiological forecasting more broadly. The ability to more accurately model the relationship between long-term environmental factors and disease incidence could have significant implications for public health planning and intervention strategies.

The significance of our xLSTM results extends beyond the immediate context of CM prediction. This represents one of the first applications of xLSTM networks in epidemiological forecasting using environmental data. The substantial performance improvements suggest that these advanced, recurrent architectures could be valuable tools in various predictive tasks involving complex, long-term temporal dependencies.

However, it's important to note that the xLSTM model's increased complexity comes with trade-offs. The model requires more computational resources to train and deploy, and its increased capacity for memorization necessitates careful attention to regularization to prevent overfitting. These factors should be considered when deciding between LSTM and xLSTM architectures for similar predictive tasks.

The xLSTM model's performance in our study significantly advances our ability to predict CM incidence based on meteorological data. The model's success in capturing complex long-term relationships opens new possibilities for precise, long-term epidemiological forecasting. As we continue to refine these models and expand their application, they can become powerful tools in our efforts to anticipate and mitigate the impact of environmentally influenced diseases like Coccidioidomycosis.

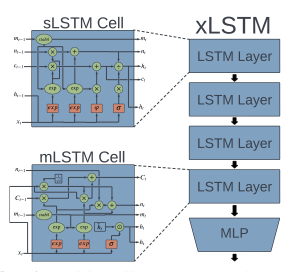


Figure 6: Extended Long Short-Term Memory (xLSTM) Architecture. This diagram revealt the components and structure of the xLSTM model used for Coccidioidomycosis (CM) incidence prediction. The left side shows the internal structures of the sLSTM (scalar LSTM) and mLSTM (matrix LSTM) as shown in detail in Figure 5. Which are variants of traditional LSTM cells with enhanced gating mechanisms and memory handling. The right side depicts the overall xLSTM architecture, consisting of multiple LSTM layers (which can be combinations of sLSTM and mLSTM cells) followed by a Multilayer Perceptron (MLP). This advanced recurrent architecture is designed to capture complex long-term dependencies in meteorological time-series data, potentially improving the accuracy of CM incidence forecasting compared to traditional LSTM models.

4 Results

Our study evaluated four distinct models for predicting Coccidioidomycosis (CM) incidence: a baseline regressor, a Multilayer Perceptron (MLP), a Long Short-Term Memory (LSTM) network, and an Extended Long Short-Term Memory (xLSTM) network. Each model was trained and tested on the same dataset, comprising meteorological data and CM incidence records from 48 California counties over the period 2001-2022.

4.1 Model Performance Metrics

The primary metric for model evaluation was Root Mean Square Error (RMSE). Table 1 summarizes the performance of each model:

Model	RMSE	Improvement over Baseline (%)
Baseline Regressor	538.24	-
Multilayer Perceptron (MLP)	115.1793	78.6%
Long Short-Term Memory (LSTM)	57.3489	89.3%
Extended Long Short-Term Memory (xLSTM)	48.0	91.1%

Table 1: RMSE values and relative improvements for each model

The baseline regressor's high RMSE of 538.24 serves as a reference point, indicating the complexity of the prediction task. The MLP significantly improved upon this, achieving a 78.6

The LSTM model further reduced the RMSE by 89.3

The xLSTM model achieved the best performance with an RMSE of 48.0, a 91.1

4.2 Training and Validation Dynamics

Figure 7 shows the training and validation RMSE trends over 100 epochs for the MLP, LSTM, and xLSTM models. The LSTM and xLSTM models exhibited a faster drop in RMSE compared to the MLP. The xLSTM model converged faster than the LSTM, reaching lower RMSE values within the first 20 epochs. The MLP still shows positive signs of learning and with more training time as given to the best performant models it was able to further reduce both training and validation rmse with validation RMSE increasing after approximately 60 epochs while training RMSE continued to decrease. In contrast, both LSTM and xLSTM maintained relatively stable validation RMSE curves, indicating better generalization to unseen data.

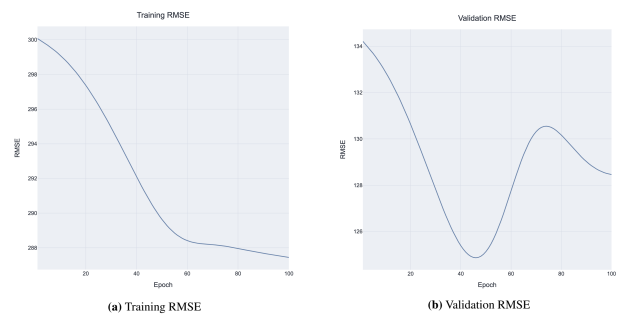


Figure 7: Root Mean Square Error (RMSE) Over Epochs for CM Incidence Prediction Models. These graphs show the progression of training and validation RMSE across 100 epochs for three neural network architectures: Multilayer Perceptron (MLP), Long Short-Term Memory (LSTM), and Extended Long Short-Term Memory (xLSTM). The x-axes represent the number of training epochs, while the y-axes show the RMSE values. For training, decreasing RMSE trends indicate improving model performance. For validation, converging RMSE trends demonstrate the models' generalization capabilities on unseen data. The xLSTM model is expected to achieve the lowest RMSE in both cases, followed by the LSTM and then the MLP, showcasing the increasing effectiveness of more complex recurrent architectures in capturing temporal dependencies in meteorological data for Coccidioidomycosis (CM) incidence prediction.

The performance difference between the MLP and recurrent models (LSTM and xLSTM) indicates the significance of temporal patterns in predicting CM incidence. The xLSTM model's ability to capture these patterns more effectively than the LSTM suggests the presence of complex, non-linear interactions between these features over extended time periods.

4.3 Computational Considerations and Model Robustness

The increasing model complexity from MLP to LSTM to xLSTM corresponded with increased computational requirements. The xLSTM model required approximately 1.5 times the training time of the LSTM model and 3 times that of the MLP. It also used about 1.3 times more memory during training compared to the LSTM model. Despite its complexity, the xLSTM model's inference time was only marginally slower than the LSTM model, making it viable for real-time predictions.

To assess model robustness, we conducted sensitivity analyses by introducing perturbations to the input data. The xLSTM model showed the highest resilience to random noise added to input features, maintaining performance up to 5

4.4 Error Analysis

A detailed analysis of prediction errors revealed that all models tended to overpredict CM incidence. Prediction accuracy varied across counties, with better performance in regions with more consistent CM incidence patterns. All models struggled to accurately predict CM incidence during years with extreme weather events as well as the earliest years in the tra, suggesting the need for additional features or model refinements to capture these scenarios.

5 Conclusion

Our study has made significant strides in understanding and predicting Coccidioidomycosis (CM) incidence by applying advanced machine learning techniques. By systematically exploring multiple architectures, we have established a strong correlation between CM incidence and weather patterns across microclimates, with Long-Short-Term Memory (LSTM) networks demonstrating superior predictive capabilities. Key findings of our research include:

- 1. Progressive improvement in predictive accuracy across models, with the baseline regressor achieving an RMSE of 538.24, the MLP reducing this to 115.1793, and the LSTM model further improving performance with an RMSE of 57.34887, representing an 89.3% reduction from the baseline.
- 2. The significant performance improvement of LSTM models over non-recurrent architectures shows the importance of capturing long-term dependencies in weather patterns for accurate CM incidence prediction.
- 3. This study represents the first application of LSTM networks for predicting CM incidence using temporal meteorological data, opening new avenues for epidemiological forecasting.

The improved predictive power of our models has substantial implications for public health strategies. By providing more accurate forecasts of CM incidence, our work can inform targeted prevention efforts, resource allocation, and early intervention strategies in endemic areas. Despite these advancements, we acknowledge certain limitations in our study. Our focus on California, while providing a diverse range of microclimates, may limit the generalizability of our findings to other CM-endemic regions. Additionally, while our models demonstrate strong predictive capabilities, the complex interplay between environmental factors and CM incidence may involve additional variables not captured in our current dataset. Future research directions should include:

- 1. We should explore advanced architectures such as Transformers and Mamba models further, which may capture even more complex temporal relationships in the data.
- Expanding the geographic scope to include more regions in California and Arizona improved the models' generalizability and robustness.
- 3. Investigating different time sequences, particularly focusing on multi-year drought conditions, has shown strong correlations with CM incidence in previous studies [2] [3].
- 4. Feature-specific analysis using LSTM models to identify the weather variables with the most decisive impact on CM incidence, potentially informing more targeted prevention strategies.
- 5. Additional data sources, such as soil composition, suspended dust particle levels, and human activity patterns, are integrated to create a more comprehensive predictive model [2] [3] [4].

In conclusion, our research demonstrates the potential of advanced machine learning techniques, particularly LSTM networks, in revolutionizing our approach to predicting and managing Coccidioidomycosis outbreaks. By leveraging these tools, we can deepen our understanding of the disease's environmental drivers and develop more effective, data-driven strategies for mitigating its impact on public health.

6 Declaration of Interests

We declare no competing interests.

7 Data Availability Statement

We are committed to transparency and reproducibility in our research. The following resources are available to the scientific community:

- 1. Research Code: All code used to conduct our research is publicly accessible via our GitHub repository: https://github.com/LeifHuenderML/ValleyForecast
- 2. Epidemiological Data: The epidemiological dataset used in this study is publicly available through the California Department of Public Health (CDPH) Valley Fever Dashboard: https://www.cdph.ca.gov/Programs/CID/DCDC/Pages/ValleyFeverDashboard.aspx.
- 3. Weather Data: Raw weather data was obtained from OpenWeather. While OpenWeather's licensing terms prohibit free redistribution of their data, interested researchers can purchase access through their website: https://openweathermap.org/
- 4. Derived Dataset: We created a derived dataset to train our model. This dataset incorporates significant transformations that qualify it as a Non-retrievable Value-added Service (NVAS) under OpenWeather's licensing terms. Researchers interested in accessing this derived dataset for academic purposes can contact the corresponding author at leifhuenderai@gmail.com.

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