Classification of Plant Leaf Disease Recognition Based on Self-Supervised Learning

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Abstract: Accurate identification of plant diseases is a critical task in agricultural production. The existing deep learning crop disease recognition methods require a large number of labeled images for training, limiting the implementation of large-scale detection. To overcome this limitation, this study explores the application of self-supervised learning (SSL) in plant disease recognition. We propose a new model that combines a masked autoencoder (MAE) and a convolutional block attention module (CBAM) to alleviate the harsh requirements of large amounts of labeled data. The performance of the model was validated on the CCMT dataset and our collected dataset. The results show that the improved model achieves an accuracy of 95.35% and 99.61%, recall of 96.2% and 98.51%, and F1 values of 95.52% and 98.62% on the CCMT dataset and our collected dataset, respectively. Compared with ResNet50, ViT, and MAE, the accuracies on the CCMT dataset improved by 1.2%, 0.7%, and 0.8%, respectively, and the accuracy of our collected dataset improved by 1.3%, 1.6%, and 0.6%, respectively. Through experiments on 21 leaf diseases (early blight, late blight, leaf blight, leaf spot, etc.) of five crops, namely, potato, maize, tomato, cashew, and cassava, our model achieved accurate and rapid detection of plant disease categories. This study provides a reference for research work and engineering applications in crop disease detection.

Keywords: plant disease recognition; self-supervised learning; deep learning; masked autoencoder; attention mechanism

1. Introduction

In modern agricultural practices, rapid and accurate identification of plant diseases is essential for safeguarding crop health, increasing yields, and reducing the use of chemical pesticides [1,2]. However, the variety of disease manifestations and the influence of environmental factors [3] make accurate disease identification a challenge.

Supervised deep learning algorithms have shown good recognition results in current agricultural disease recognition [4–6]. On the PlantVillage dataset, the recognition accuracy of different diseases can be up to 95%–99% [7,8]. Li et al. [9] added a coordinate attention (CA) module to the single-stage plant disease network YOLOv5s, as well as key feature weights to enhance the effective information of the feature maps, and the spatial pyramid pooling (SSP) module was modified to reduce the loss of feature information. Li et al. [10] proposed an improved algorithm for vegetable disease detection based on YOLOv5s which improves the CSP (Cross-Stage Partial), FPN (Feature Pyramid Network), and NMS (non-maximum suppression) modules in YOLOv5s, removes the influence of the external environment, and enhances the ability of multi-scale feature extraction. This improved the accuracy of detection of umbilical rot, gray mold, and cabbage anthracnose on tomato, cucumber, cabbage, and other vegetables. Muhammad et al. [11] recognized leaf spot, powdery mildew,
yellow wilt, and other diseases of cotton by meta-deep learning and the accuracy was 98.53%. Ma et al. [12] proposed a lightweight convolutional neural network (CNN) model that can be deployed on mobile terminals to accurately recognize gray leaf spot, common rust, and northern leaf blight of maize with 99.11% recognition accuracy. However, supervised deep learning relies heavily on manual labeling, so we had to find a more efficient solution.

Currently, self-supervised learning in agriculture is increasingly favored [13–15], and its learning advantages and good performance in classification tasks are exactly what is needed for plant leaf disease recognition [16]. Self-supervised learning does not require annotation in the traditional sense [17]. Instead, it extracts supervised signals from the input data itself, which means it can learn using unlabeled data. The characteristics of self-supervised learning are very effective when dealing with large amounts of unlabeled data, especially in areas such as disease identification where obtaining labeled data is costly or difficult [18,19]. In addition, mixing self-supervised and supervised methods can improve the performance of the system if there is a large amount of unlabeled data and a small amount of labeled data [20]. Self-supervised learning has been used by scholars for leaf segmentation [21], maturity detection [22,23], growth condition detection [24,25], seed identification [26], and other research directions in plants, which has injected new vigor and vitality into agricultural development.

An MAE (masked autoencoder) [27] is a type of self-supervised learning that is commonly used in the field of image processing. It is an autoencoder-based [28] structure specifically designed to efficiently learn a representation of an image. The key idea is to mask parts of the image and then train the model to reconstruct these masked parts. During the training process, the model learns to minimize the reconstruction error, and in this way the encoder is forced to learn the important feature representations in the image so that it can efficiently reconstruct the masked portions with the help of the decoder [29].

This study focuses on exploring and evaluating the application of self-supervised learning models to the task of plant disease recognition and classification. We employed state-of-the-art self-supervised learning architectures, such as MAE and attention modules (e.g., CBAM), which not only improve recognition accuracy but also enhance the model’s ability to understand disease features. Our experimental results show that through self-supervised learning, the model achieves significant performance improvement on several standard datasets, confirming the great potential of self-supervised learning in solving real-world problems. In addition, our study demonstrates the advantages of self-supervised learning in the case of dealing with limited labeled data, providing new perspectives for future research directions and agricultural practices.

2. Materials and Methods

2.1. Datasets

2.1.1. The Dataset We Collected

The dataset we collected was derived from the PLD dataset [30] and images taken and collected by the group at the Yuzi Organic Dry Farming Experimental Site (112°86'E, 37°76'N) (Figure 1). The dataset consisted of 3256 images divided into three categories: late blight, early blight, and healthy (Table 1). We processed our collected dataset by rotating, randomly cropping, and horizontally flipping the original images to increase the size of the dataset.

![Examples of samples from our potato disease dataset. (a) Early_Blight; (b) Healthy; (c) Late_Blight.](image-url)
Table 1. Summary of the potato dataset.

<table>
<thead>
<tr>
<th>Class Labels</th>
<th>Samples</th>
</tr>
</thead>
<tbody>
<tr>
<td>Early_Blight</td>
<td>1628</td>
</tr>
<tr>
<td>Healthy</td>
<td>1020</td>
</tr>
<tr>
<td>Late_Blight</td>
<td>1692</td>
</tr>
</tbody>
</table>

2.1.2. The CCMT Dataset

CCMT [31] is an open-source dataset for crop diseases, consisting of 88,010 images (23,672 for cashews, 21,768 for cassava, 15,402 for corn, and 27,168 for tomatoes), divided into 18 categories. The dataset includes 18 common leaf diseases such as rust, leaf blight, and leaf spot, as shown in Table 2. Unlike the PlantVillage dataset, the CCMT dataset images are sourced from field photos and have better adaptability to complex field environments.

Table 2. Summary of the CCMT dataset.

<table>
<thead>
<tr>
<th>CCMT Dataset</th>
<th>Class Labels</th>
<th>Samples</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cashew_anthracnose</td>
<td>4940</td>
<td>Maize_healthy</td>
</tr>
<tr>
<td>Cashew_healthy</td>
<td>7213</td>
<td>Maize_leaf_blight</td>
</tr>
<tr>
<td>Cashew_leaf_miner</td>
<td>4953</td>
<td>Maize_leaf_spot</td>
</tr>
<tr>
<td>Cashew_red_rust</td>
<td>6566</td>
<td>Maize_streak_virus</td>
</tr>
<tr>
<td>Cassava_bacterial_blight</td>
<td>5864</td>
<td>Tomato_healthy</td>
</tr>
<tr>
<td>Cassava_brown_spot</td>
<td>4733</td>
<td>Tomato_leaf_blight</td>
</tr>
<tr>
<td>Cassava_green_mite</td>
<td>4266</td>
<td>Tomato_leaf_curl</td>
</tr>
<tr>
<td>Cassava_healthy</td>
<td>3455</td>
<td>Tomato_septoria_leaf_spot</td>
</tr>
<tr>
<td>Cassava_mosaic</td>
<td>3450</td>
<td>Tomato_verticulium_wilt</td>
</tr>
</tbody>
</table>

2.2. Construction of the Model

2.2.1. The Masked Autoencoder Model

An MAE (masked autoencoder) [27] is a self-supervised learning algorithm mainly used for visual tasks. It is based on an encoder–decoder architecture (Figure 2) that trains a model by masking parts of the image and trying to reconstruct these masked parts. Blue and red squares are the encoder and decoder outputs respectively. MAE is a self-supervised learning algorithm, it does not require a large amount of labeled data for training. It generates the training signals through its own data structures. One of the main advantages of an MAE is its efficient learning process, which reduces the computational burden due to the fact that only some of the pixels need to be processed during the pre-training process. In addition, through the reconstruction task, an MAE is able to capture the global structure and local details of the image, which improves the model’s understanding of the image content.

After the MAE, ViT (Vision Transformer) [32] is used as the backbone structure. In ViT, images are segmented into patches and flattened into patch embeddings, which serve as inputs to the encoder. The encoder of the model consists of N consecutive blocks, each containing mainly of a Multihead Self-Attention (MSA) [33] and a Multilayer Perceptron (MLP) [34] block. ViT employs the Transformer [35] architecture, which is commonly used in natural language processing. This architecture relies on a self-attention mechanism to capture the global dependencies of the input data.

2.2.2. Convolutional Block Attention Module

A CBAM (convolutional block attention module) [36] is an attention mechanism module for convolutional neural networks (Figure 3). This module improves the performance and accuracy of the network by focusing on the most important features in the image [37]. A CBAM mainly consists of two modules, the channel attention mechanism (CAM) and spatial attention mechanism (SAM) modules.
global structure and local details of the image, which improves the model’s sensitivity to important features and thus improves performance.

2.2.3. Gate Recurrent Unit Module

A GRU (Gate Recurrent Unit) [38] is a type of recurrent neural network (RNN) [39]. It was proposed to solve problems such as long-term memory and gradient in backpropagation. Compared to LSTM, the use of a GRU can achieve comparable results [40] and is easier to train, which can largely improve the training efficiency, so it will often be preferred [41].

The GRU controls the flow of information through two gates (update gate and reset gate). The update gate helps the model decide how much of the previous memory to retain.
in the current state, while the reset gate decides how to combine new input information with past memories. At each time step, the GRU uses the gating mechanism to update its hidden state, and this updated hidden state captures information about the sequence so far. The structural design of the GRU helps to alleviate the problem of gradient vanishing that is common in traditional RNNs by making it capable of learning long-distance dependencies. The formulation of the GRU is as follows (Equation (1)).

\[
\begin{align*}
    z_t &= \sigma(W_z \cdot [h_{t-1}, x_t]) \\
    r_t &= \sigma(W_r \cdot [h_{t-1}, x_t]) \\
    \tilde{h}_t &= \tanh(W \cdot [r_t \odot h_{t-1}, x_t]) \\
    h_t &= (1 - z_t) \odot h_{t-1} + z_t \odot \tilde{h}_t
\end{align*}
\]

where \( x_t \) is the information entered at the current moment; \( h_{t-1} \) is the hidden state that represents the previous moment; \( h_t \) is the hidden state passed to the next moment; \( \tilde{h}_t \) is a candidate hidden state; \( r_t \) is a reset gate; \( z_t \) is the update gate; \( \sigma \) is the sigmoid function by which data can be changed to values in the range \( 0–1 \); and \( \tanh \) is the tanh function by which data can be changed to values in the range \([-1,1]\).

2.2.4. Improved Model Architecture

We improved the structure of the MAE model by combining the CBAM attention mechanism and GRU. First, we added a CBAM (Figure 4) for feature enhancement of the images before the image blocks enter the encoder. Second, we added the GRU module after the linear layer of the decoder to capture the sequential relationship between the diseased image blocks and enhance the processing of temporal information of the features passed from the encoder.

![Figure 4. Improved model architecture.](image)

When processing an image, ViT splits the image into a series of image blocks. These image blocks are feature-enhanced by the CBAM, and each image block is spread and converted into a series of vectors which are subsequently fed into the Transformer model.

The decoder consists of multiple decoding blocks, each containing a Multihead Self-Attention (MSA) [33] and a Multilayer Perceptron (MLP) [34] block. The work of the decoder begins with receiving the output of the encoder, which contains a compressed representation of the unmasked input and a mask token for the masked portion. The decoder uses this information to reconstruct the masked image block. During the training phase, the decoder compares the differences between the minimally reconstructed image and the original image to adjust its parameters. In this way, the decoder not only learns how to accurately fill in the masked portions, but also captures key features of the input
data without the need for external labeling. At the same time, our encoder design allows the model to learn complex features inside the image through a self-attentive mechanism, which can subsequently be used for various downstream tasks such as image classification, object detection, or image generation.

2.3. Test Platform and Parameters

The system software used for the test was Windows 11 (64-bit), the programming language was Python version 3.7, Pythorch version 1.14 was used as the processing framework, the deep learning computational framework of CUDA version 16.04 and the cuDNN version 7.6.3 configuration acceleration library were used, and PyCharm was used for debugging the program. The hardware configuration was NVIDIA GeForce RTX 4080 GPU graphics accelerated test process, 16 GB RAM, and 16-core (8P + 8E) 24-thread computer processor with a maximum RX frequency of 5.4 GHz. The size of the training image was adjusted to 256 × 256 pixels, and the batch size of the training was set to 64. The training process consisted of 100 iterations.

Both the CCMT dataset and our collected dataset were divided into training, validation, and test sets in the ratio of 8:1:1. All results are the average recognition accuracy obtained after 5 repetitions of training and testing.

2.4. Evaluation Indicators of Experimental Results

We used Accuracy, Precision, Recall, and F1 score as the evaluation metrics of the model. F1 score is a measure of the precision of a classification model. It is the reconciled average of model Precision and Recall. The formulas for calculating Accuracy, Recall, Precision and F1 score are given in Equations (2)–(5).

\[
\text{Accuracy} = \frac{\text{Number of Correct Predictions}}{\text{Total Number of Predictions}} \times 100\% \\
\text{Recall} = \frac{\text{True Positives}}{\text{True Positives} + \text{False Negatives}} \times 100\% \\
\text{Precision} = \frac{\text{True Positives}}{\text{True Positives} + \text{False Positives}} \times 100\% \\
F1 = 2 \times \frac{\text{Precision} \times \text{Recall}}{\text{Precision} + \text{Recall}} \times 100\% 
\]

3. Results

3.1. Comparison and Analysis of Different Improved Models

We added the GRU and CBAM to the MAE model for improving the accuracy and efficiency of the algorithm. Figure 5 shows the loss values of our model pre-trained with the PlantVillage dataset; the model with the addition of the GRU and CBAM has lower loss values and converges faster.

![Figure 5. Loss values of different methods in the pre-training process.](image_url)
Subsequently, we compared the accuracy of the improved model through experiments on two different datasets. The experimental results are shown in Table 3. On the self-built dataset, the integration of the GRU module resulted in a model with an accuracy of 98.81%, a recall of 98.02%, and an F1 score of 97.95%. Adding the CBAM slightly improves the performance with a 98.92% accuracy, 98.11% recall, and 98.10% F1 score. Combining both the GRU and CBAM performed best on this dataset with a 99.35% accuracy, 98.16% recall, and 98.17% F1 score. Compared to the model with only the GRU and CBAM added, the accuracy of combining both the GRU and CBAM is higher by 0.54% and 0.42%, respectively.

![Table 3. Performance of different improvements on CCMT and our collected dataset.](image)

<table>
<thead>
<tr>
<th>Dataset</th>
<th>Added Modules</th>
<th>Accuracy</th>
<th>Recall</th>
<th>F1</th>
</tr>
</thead>
<tbody>
<tr>
<td>Our dataset</td>
<td>GRU</td>
<td>98.81%</td>
<td>98.16%</td>
<td>98.17%</td>
</tr>
<tr>
<td></td>
<td>CBAM</td>
<td>98.92%</td>
<td>98.11%</td>
<td>98.10%</td>
</tr>
<tr>
<td></td>
<td>GRU + CBAM</td>
<td>99.35%</td>
<td>98.51%</td>
<td>98.62%</td>
</tr>
<tr>
<td>CCMT</td>
<td>GRU</td>
<td>95.29%</td>
<td>96.02%</td>
<td>95.43%</td>
</tr>
<tr>
<td></td>
<td>CBAM</td>
<td>95.17%</td>
<td>95.81%</td>
<td>95.1%</td>
</tr>
<tr>
<td></td>
<td>GRU + CBAM</td>
<td>95.61%</td>
<td>96.20%</td>
<td>95.52%</td>
</tr>
</tbody>
</table>

On the CCMT dataset, the GRU module achieved a 95.29% accuracy, 96.02% recall, and 95.43% F1 score. The model using the CBAM scored slightly lower compared to the GRU module, with a 95.17% accuracy, 95.81% recall, and 95.1% F1 score. The combination of the GRU and CBAM again provided the best results with a 95.61% accuracy, 96.20% recall, and 95.52% F1 score. Compared to the model with the addition of the GRU and CBAM only, the accuracy of combining both the GRU and CBAM was 0.32% and 0.44% higher. Therefore, we decided to use the GRU + CBAM model optimization method.

3.2. Comparison and Analysis of the Improved Model with Other Models

3.2.1. Comparison and Analysis on Our Collected Datasets

The performance of the model is first discussed on the dataset we collected, with the model’s loss function and accuracy curves (Figure 6). As the number of interactions increases, the accuracy of the model steadily increases while the loss continues to decrease. At the beginning of training, the accuracy rises rapidly and the loss decreases rapidly. Subsequently, we compared the improved model with other models on the self-constructed dataset, and three different models (ResNet50, ViT, and MAE) were selected for experiments on the self-constructed dataset (Figure 7). The accuracy of the ResNet50, ViT, and MAE models were 97.5%, 97.2%, and 98.2%, respectively, with Ours_MAE obtaining the highest accuracy (98.8%), which was 1.3%, 1.6%, and 0.6% higher than the other three models, respectively. The accuracy and loss curves of the four algorithms at different epochs are shown in Figure 8.
The normalized confusion matrix in Figure 9 evaluates the model’s performance in a multi-class plant disease recognition task. The results show that the model accurately predicted 99% of the early blight instances, and only 1% of the instances were misclassified as Late_Healthy. Overall, the model showed high accuracy on all three categories, especially in distinguishing early blight from late blight.

Figure 7. Accuracy of different models on our collected dataset.

Figure 8. Accuracy and loss of different models on our collected dataset. (a) Accuracy, (b) Loss.

Figure 9. Confusion matrix of the improved model on our collected dataset.
3.2.2. Comparison and Analysis on the CCMT Dataset

In order to evaluate the performance of the models, we selected three different models (ResNet50, ViT, MAE) to test with further experiments on the CCMT dataset, where batch_size was set to 64 and epoch was set to 100 during the training process, and Adma was chosen as the optimizer. Figure 10 shows the accuracy of the ResNet, ViT, and MAE models, which are 98.4%, 94.91%, and 94.81%, respectively. The accuracy of Ours_MAE reaches 95.61%, which is 1.2%, 0.7%, and 0.8% higher than other models, respectively. It can be seen that our model has improved in accuracy.

![Accuracy of different models on CCMT dataset](image)

**Figure 10.** Accuracy of different models on CCMT dataset.

The accuracy and loss curves of the four models under different epochs are shown in Figure 11 where Ours_MAE has the highest accuracy and a relatively smooth loss curve. The results show that our improved model achieves excellent performance on the CCMT dataset.

![Accuracy and loss of different algorithms on CCMT dataset](image)

**Figure 11.** Accuracy and loss of different algorithms on CCMT dataset. (a) Accuracy, (b) Loss.

In order to analyze the recognition accuracy of our model for different diseases, we analyzed the confusion matrices of the model’s classification predictions, and the normalized confusion matrices of Figure 12 evaluated the model’s performance in the task of recognizing multiple classes of plant diseases. Among them, Cashew_healthy, Tomato_healthy, Cassava_healthy, and Maize_healthy achieved a normalized true-positive rate of more than 0.97, implying that almost all truly healthy plants were correctly classified by the model. Of note, Maize_leaf_spot had a normalized value of 0.13 misclassified as Maize_leaf_blight, indicating some confusion between the two diseases. This provides ideas for further improvement of the model in the future.
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whether the disease is caused by different pathogenic factors such as viruses, bacteria, or which helps to recognize and understand complex patterns of disease.

The CBAM brings significant performance gains to the plant disease recognition task by providing fine-grained feature adjustment capabilities.

The MAE model has several advantages for image classification tasks such as plant disease recognition, including learning global and local features, effective use of unlabeled data, computational efficiency, etc. Once the model captures sufficient feature representations in the pre-training phase, it can be quickly adapted to a specific disease recognition task through a fine-tuning process.

Overall, the MAE model provides a powerful framework for plant disease recognition tasks, especially for situations where data labeling is costly or available labeled data are limited.

The CBAM automatically learns which regions of the image to focus resources on and which feature channels to focus on through channel and spatial attention mechanisms [45]. In disease recognition tasks, this means that the model can automatically highlight diseased spots or damaged regions and ignore background or irrelevant parts. At the same time, CBAM is able to adapt its attention dynamically to different disease types and plant species. The CBAM brings significant performance gains to the plant disease recognition task by providing fine-grained feature adjustment capabilities.

The GRU is a variant of a recurrent neural network (RNN) that is particularly suitable for processing sequence data; although it is not as common as a convolutional neural network (CNN) in processing image data, the GRU can effectively capture disease processes that evolve over time, such as the development and spread of spots, in specific application scenarios describing the development and spread of disease spots. For irregularly shaped disease features, the GRU can handle unstructured data through its recursive mechanism, which helps to recognize and understand complex patterns of disease.

In future research, we will focus on the specific causes of disease occurrence, i.e., whether the disease is caused by different pathogenic factors such as viruses, bacteria, or

Figure 12. Confusion matrix of the improved model on the CCMT dataset.

4. Discussion

Contrastive learning such as SimCLR [42] and MoCo [43] rely on a comparison of positive and negative sample pairs to learn the representation. These methods learn by maximizing the similarity of positive sample pairs and minimizing the similarity of negative sample pairs [44]. In contrast to this, which typically requires a large number of negative samples and a complex sample selection strategy, an MAE does not rely on sample pair comparisons but on reconstruction tasks, thus avoiding the challenge of needing large-scale negative samples and allowing for training at a much lower computational cost.

The MAE model has several advantages for image classification tasks such as plant disease recognition, including learning global and local features, effective use of unlabeled data, computational efficiency, etc. Once the model captures sufficient feature representations in the pre-training phase, it can be quickly adapted to a specific disease recognition task through a fine-tuning process.

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In future research, we will focus on the specific causes of disease occurrence, i.e., whether the disease is caused by different pathogenic factors such as viruses, bacteria, or
fungi. Exploration in this direction is important to improve the accuracy and usefulness of plant disease identification. In order to achieve this goal, we need to collect and organize detailed annotated datasets containing different disease causes. The dataset should include different plant species, disease types, and disease manifestations at different stages of development. This should be combined with plant pathology to discover more disease features related to specific pathogens, which can be used as an important basis for model training.

This study demonstrates the application of a self-supervised learning model to the task of plant disease recognition, in particular an approach to enhance model performance by integrating the GRU and CBAM. The results show that the model achieves excellent performance on both the self-built dataset and the CCMT dataset, with a high accuracy, recall, and F1 score. These results can be explained in several ways: firstly, regarding the improvement of the feature learning ability, the GRU module enhances the model’s ability to capture the time series and spatial structure, which is similar to the findings of Bi et al. [46], and the CBAM focuses on key regions in the image through the mechanism of attention, which has been similarly attempted by Alirezazadeh et al. [47], and these help to extract features that are closely related to disease identification. Secondly, regarding the optimization of data utilization, the self-supervised learning framework enables the model to learn useful representations from a large amount of unlabeled data, which reduces the dependence on large-scale labeled datasets, and thus improves the data utilization efficiency. Lastly, regarding the enhancement of the model generalization ability, by pre-training on large-scale datasets, the model learns more general image representations, which improves the model’s ability to generalize in the face of unknown disease types, and similar results were obtained by dong et al. [48].

5. Conclusions

By utilizing a large amount of unlabeled data, this study confirms the potential of self-supervised learning to reduce the reliance on expensive and specialized annotations. This finding is particularly important for the field of agriculture, where data acquisition is costly and specialized knowledge is required. This study not only provides a new technological path for automated identification of plant diseases, but also lays the foundation for further applications of deep learning in agricultural science. We believe that self-supervised learning will play an increasingly important role in the field of plant science with the continuous advancement of computational technology and data processing methods.

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Conflicts of Interest: The authors declare no conflicts of interest.

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