Research on Voxel-Based Features Detection and Analysis of Alzheimer’s Disease Using Random Survey Support Vector Machine

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Alzheimer’s disease (AD) is a degenerative disease of the central nervous system characterized by memory and cognitive dysfunction, as well as abnormal changes in behavior and personality. The research focused on how machine learning classified AD became a recent hotspot. In this study, we proposed a novel voxel-based feature detection framework for AD. Specifically, using 649 voxel-based morphometry (VBM) methods obtained from MRI in Alzheimer’s Disease Neuroimaging Initiative (ADNI), we proposed a feature detection method according to the Random Survey Support Vector Machines (RS-SVM) and combined the research process based on image-, gene-, and pathway-level analysis for AD prediction. Particularly, we constructed 136, 141, and 113 novel voxel-based features for EMCI (early mild cognitive impairment)-HC (healthy control), LMCI (late mild cognitive impairment)-HC, and AD-HC groups, respectively. We applied linear regression model, least absolute shrinkage and selection operator (Lasso), partial least squares (PLS), SVM, and RS-SVM five methods to test and compare the accuracy of these features in these three groups. The prediction accuracy of the AD-HC group using the RS-SVM method was higher than 90%. In addition, we performed functional analysis of the features to explain the biological significance. The experimental results using five machine learning indicate that the identified features are effective for AD and HC classification, the RS-SVM framework has the best classification accuracy, and our strategy can identify important brain regions for AD.

Keywords: Alzheimer’s disease, RS-SVM, voxel-based features, gene-level, pathway-level

INTRODUCTION

Due to the development of medical technology, the world population has grown steadily, and the elderly population has increased rapidly. It is expected that this trend will continue to accelerate in the next few decades, and the occurrence of senile diseases and the social cost of aging are expected to increase. Alzheimer’s disease (AD) is a brain disease. It is also a progressive disease, meaning that it will get worse over time. It is believed that AD begins 20 years or more before
the onset of symptoms (Jiao et al., 2020b). The preclinical stage of AD is crucial for identifying early pathophysiological events and developing interventions for disease improvement. Given that changes in synaptic function occur early in the neurodegenerative process, functional MRI (fMRI) is particularly promising for detecting early changes in brain function (Agosta et al., 2017). MRI has aroused great interest in AD-related research due to its complete noninvasiveness, high availability, high spatial resolution, and good contrast between different soft tissues (Moradi et al., 2015).

Mild cognitive impairment (MCI), known as the early stage of AD, was a disease state of cognitive decline between normal elderly and dementia patients. MCI was divided into early mild cognitive impairment (EMCI) and late mild cognitive impairment (LMCI). Studies had pointed out that if MCI patients were not diagnosed early, the probability of developing AD could be as high as 80% after 6 years, and about two-thirds of AD patients were converted through MCI (Barnes and Yaffe, 2011; Lenhart et al., 2021; Vitali et al., 2021). Using linear mixed models, Vonk et al. (2020) analyzed 2,261 individuals with MCI and non-MCI and found that the neurodegeneration was associated with letter fluency and semantic fluency. Wang et al. (2017) introduced the linear regression classification to classify samples and obtained an accuracy of 97.51% (Zhang et al., 2014). Bi et al. (2020a; 2021b) applied the random forest to identify features associated with AD. Another study showed that the Flash Visual Evoked Potential-P2 latency had AD-specific pathological information (Arruda et al., 2020). Sabuncu et al. (2011) calculated the degree of atrophy of hippocampus and cortical areas and found that the specific cortical thinning and the reduction of hippocampal volume were accelerated in early AD. As the classic analysis methods, the machine learning algorithms brought new research sight to AD-specific biomarkers (Zhang et al., 2015a; Ji et al., 2021; Jiao et al., 2021; Wang et al., 2021). Zhuo et al. (2015) applied a group lasso support vector machine to obtain the AD-specific biomarkers. Patel et al. (2019) developed two XGBoost classification models to classify AD and healthy control (HC). Studies have proved that AD was closely related to brain atrophy and that brain atrophy was mainly reflected in the reduction of cortical surface area, thickness, and gray matter volume and, therefore, gray matter volume, cortical surface area, and average thickness contributed to the pathology of AD patients (Gullett et al., 2021; Lorenzo et al., 2021; Piersson et al., 2021; Talwar et al., 2021).

Despite many efforts, it is still challenging to determine effective AD-specific biomarkers for early diagnosis and prediction of disease progression and requires more research (Bi et al., 2020b; Zhang and Shi, 2020). In our study, we proposed a novel analysis framework based on the Random Survey Support Vector Machines (RS-SVM) for the early detection of AD conversion in MCI patients by using advanced machine learning algorithms and combining voxel-based data with standard neuropsychological test results. First, to obtain the voxel sets, we extracted the differences between AD and HC. Then, we applied the RS-SVM to identify important features that classified EMCI, LMCI, AD, and HC well. Subsequently, we applied several classical methods to construct the analysis frameworks and evaluate the accuracy of these features to classify with EMCI-HC, LMCI-HC, and AD-HC. The experiment results demonstrate that the identified features were effective in classifying AD, the RS-SVM framework performed well, and the identified regions and genes will further our understanding of AD.

**MATERIALS AND METHODS**

**Figure 1** illustrates the framework of a voxel-based three-level analysis for AD. The framework encompasses data processing (A), features extraction (B), RS-SVM construction (C), and the gene-level analysis using effective chi-square statistic (ECS) method (Li et al., 2019) and pathway-level analysis using the resulting genes (Bu et al., 2021) (D). The novation of this framework is to make the full use of voxel-based data.

**Imaging Data**

In this study, we downloaded and analyzed 1,426 participants with genotyping data and MRI scans from the Alzheimer's Disease Neuroimaging Initiative (ADNI) database (adni.loni.usc.edu). These data include 353 HCs, 273 EMCI, 504 LMCI, and 296 with AD. The characteristics of these participants, including average age and years of education, are shown in Table 1.

**Random Survey Support Vector Machines-Based Machine Learning Method**

**Data Processing**

MRI scans, using voxel-based morphometry (VBM), were aligned and normalized to a T1-weighted template image and the Montreal Neurological Institute (MNI) space, respectively. The gray matter density (GMD) maps were segmented, extracted, and smoothed with an 8-mm FWHM (full width at the half maximum) kernel. The Automatic Anatomical Labeling (AAL) atlas was employed to define the regions of interest (ROIs) and their coordinates (whole brain) (Tzourio-Mazoyer et al., 2002). We then down-sampled the resulting maps to a dimension of $61 \times 73 \times 61$ to reduce the data size for subsequent analysis in EMCI-HC, LMCI-HC, and AD-HC groups.

To extract the differences within the three groups, i.e., A, B, and C, we first performed the weighted process of the two sets of images separately and saved them as matrices $M$ and $N$ (i.e., $M$ for AD group and $N$ for HC group). Then, let $v_{m_i}^j$ and $v_{n_i}^j$ represent the vectors of two voxels ($v_{m_i}^j \in M$, $v_{n_i}^j \in N$), and we obtained a vector $V = \{v_{m_1}^1, v_{n_1}^1, v_{m_2}^2, v_{n_2}^2, \ldots, v_{m_5}^5, v_{n_5}^5\}$ ($k = 271, 633$). Since the voxels ($v_{m_i}^j = v_{n_i}^j$) in the two groups were meaningless for our research, we deleted these voxels and obtained 64,411 sets of different voxels. We used $V'$ to denote the voxel set.

**Feature Extraction**

The data sets of features were still too many for our final binary classification in EMCI-HC, LMCI-HC, and AD-HC groups. Therefore, we estimated the number of features by calculating the
similarity of the voxels is given in the following equation:
\[
\rho_i = \sqrt{(vm_i - vm_i')^2 + (vn_i - vn_i')^2}, \ (vm_i, vn_i), (vm_i', vn_i') \in V'
\]
where \(vm_i\) and \(vm_i'\) \((i, j = 1, 2, ..., 64411)\) are the values of AD group. \(vn_i\) and \(vn_i'\) \((i, j = 1, 2, ..., 64411)\) are the values of HC groups. \(\rho_i\) is the similarity between \((vm_i, vn_i)\) and \((vm_i', vn_i')\).

For the convenience of calculation, we divided the \(V'\) into ten groups and obtained 55 sets of similarity matrices. On this basis, we defined the number of minimal \(\rho_i\) as \(C_{\min}\) and the number of maximal \(\rho_i\) as \(C_{\max}\). Due to the value of \(C_{\min}\) and \(C_{\max}\) \((C_{\min} = 132, C_{\max} = 21)\), we defined that the number of features should be in \([C_{\max}, C_{\min}]\). Then, we extracted 64,411 features of all subjects from the original MR images to form a 649 \(\times\) 64,411 matrix as the initial data set.

**Random Survey Support Vector Machines Construction**

To extract the important feature, we proposed a single-kernel SVM model based on random survey. The goal of random survey was the establishment of a random experimental data set. Since the initial data set \(X\) was a two-dimensional matrix of 649 \(\times\) 64,411, we selected \(l\) column from the \(X\) randomly and constructed a single randomized experimental data set \(X' (l \in [C_{\max}, C_{\min}])\). At the same time, the set of columns corresponding to each column \(l\) in each extraction was \(R = \{r_1, r_2, ..., r_l\}\), which denoted the index of brain loci coordinates. The indices are extracted as follows:

\[
\begin{align*}
    u(k) &= \{u_1, u_2, ..., u_k\}, k = 64411 \\
    u'(k) &= \{u_p, u_q, ..., u_r\}, r, p, q \in [1, 64411] \\
    R &= \{r_1 = u_p, r_2 = u_q, ..., r_l = u_q\}, g \in [1, 64411]
\end{align*}
\]

After random extraction, we defined the training set-validation set-test set as 6:2:2. The training set was used as an input for training first. The validation set was applied to obtain the optimal hyperparameters and replaced the initial parameters. The remaining 20% was introduced as the test set to calculate the accuracy of the tuned model.

In the classification process of SVM, the input data \(X' = \{X'_1, X'_2, ..., X'_N, X'_M\}\) and the learning objective \(y = \{y_1, y_2, ..., y_N, ..., y_M\}\) were given, where \(N\) was the number of EMCI, LMCI, and AD samples, respectively, and \(M\) was the number of HC. The learning objectives were binary variables \(y = \{-1, 1\}\), where -1 represents EMCI, LMCI, and AD, respectively, and 1 represents HC in the three groups. The
feature set of the input data was regarded as the hyperplane \( D \) in decision boundaries to separate the learning targets by positive and negative classes, making the distance \( \epsilon_i \) between any sample and plane \( \geq 1 \). The hyperplane and the plane distance are defined as follows:

\[
D : w^T X' + b = 0
\]

\[
\epsilon_i = y_i(w^T X' + b), \, \epsilon_i \geq 1
\]

where \( w \) denotes the normal vector of the hyperplane and \( b \) denotes the intercept of the hyperplane. The decision boundary satisfying this condition actually constructed two parallel hyperplanes \( D_1, D_2 \) as interval boundaries to classify the samples (Eq. 4).

\[
w^T X'_i + b \geq +1 \Rightarrow y_i = +1
\]

\[
w^T X'_i + b \leq -1 \Rightarrow y_i = -1
\]

Based on Eq. 4, it could be derived that all samples above the upper interval boundary were positive and those below the lower interval boundary were negative. The distance between the two interval boundaries \( d = \frac{|b|}{|w|} \) was defined as the margin. Since our experimental data \( X' \) was selected randomly, there was hyperboloid in the feature set to separate positive and negative classes. Using nonlinear functions, the nonlinear separable problems from the original feature set were mapped to a higher dimensional Hilbert space \( H \). The hyperplane, using as the decision boundary, is defined as follows:

\[
w^T \phi(X') + b = 0
\]

where \( \phi : X' \mapsto H \) denotes the mapping function. Since the mapping function was complex, it was difficult to calculate the inner product. Therefore, the inner product of the mapping function was defined as kernel functions \( k(X'_1, X'_2) = \phi(X'_1)^T \phi(X'_2) \) to avoid the explicit operation.

### Parameter Determination

We used the original linear kernel function of the support vector machine first, and the penalty factor \( C \) and the kernel parameter gamma were set as default values (\( C = 1 \) and gamma = 0.5). Then, we applied the training data set and labels to train the model. Subsequently, the hyperparameters were optimized by grid search. The SVM could be transformed into an equivalent quadratic convex optimization problem to solve using the following equation:

\[
\min \frac{1}{2}||w||^2 + C \sum_{i=1}^{M} \epsilon_i
\]

\[
y_i(w^T X'_i + b) \geq 1 - \epsilon_i, \, \epsilon_i \geq 0
\]

### Evaluation Metrics

In this article, the samples were positive and negative, and the results classified had the following cases:

- True positive (TP): the positive sample was predicted as a positive sample.
- True negative (TN): the negative sample was predicted as a negative sample.
- False positive (FP): the negative sample was predicted as a positive sample.
- False negative (FN): the positive sample was predicted as a negative sample.

Let \( P \) denotes the positive sample and \( N \) denotes the negative sample. We then obtained the following equation:

\[
TP + FN = P
\]

\[
FP + FN = N
\]

The evaluation metrics used in our research are as follows:

- **Accuracy.** Accuracy was the number of correctly classified samples divided by the total number of samples (Eq. 8).

\[
ACC = \frac{TP + TN}{P + N}
\]

- **Precision.** Precision was the proportion of the samples that were actually positive (or negative) divided by samples classified as positive (or negative) (Eq. 9).

\[
precision = \frac{TP}{TP + FP}
\]

- **Recall.** Recall was the measure of coverage (Eq. 10).

\[
recall = \frac{TP}{TP + FN}
\]

- **Comprehensive evaluation indicators (F-Measure).** Accuracy and sensitivity sometimes needed to be considered together as given in the following equation:

\[
F = \frac{(\alpha^2 + 1) * P * R}{\alpha^2(P + R)}
\]

When \( \alpha = 1 \), Eq. 11 is transformed into the following equation as follows:

\[
F = \frac{2 * P * R}{P + R}
\]

### Model Comparison

We used the test set to evaluate the classification ability of 5 machine learning methods, including linear regression model, least absolute shrinkage and selection operator (Lasso) model, partial least squares (PLS) model, SVM model, and RS-SVM model. First, the initial default parameters were applied to each model to train and calculate the evaluation metrics. Then, the grid search algorithm was used to optimize the hyperparameters of the five models. Finally, the hyperparameters were introduced in each model to recalculate the evaluation metrics. The results were used to evaluate the pros and cons of the five models.
Since the RS-SVM model in this article was optimized based on the traditional SVM model, the other three evaluation models were described in detail in this section.

Linear regression model was a statistical analysis method that used regression analysis in mathematical statistics to determine the quantitative relationship between the interdependence of two or more variables.

Given a data set \( D = \{(x_1, y_1), (x_2, y_2), \ldots, (x_i, y_i)\} \), we learned that a linear model from this data set will reflect the correspondence between \( x_i \) and \( y_i \) as accurately as possible. The linear regression model, which was a function of linear combination of attributes \( x \), could be expressed as follows:

\[
 f(x) = w_1 x_1 + w_2 x_2 + \ldots + w_i x_i + b = W^T X + b \tag{13}
\]

where \( W = \{w_1, w_2, \ldots, w_j\} \) is column vector, indicating the weight of the corresponding attribute in the prediction result.

Eq. 13 was represented as the following equation:

\[
 f(x_i) = w_i x_i + b, f(x_i) \approx y_i \tag{14}
\]

Then it was to find a model such that \( \forall i \in [1, m] \) has \( f(x_i) \) as close to \( y_i \). Therefore, the sum of the squares of the difference between the predicted value and the real value of each sample is minimized and thus gives the following equation:

\[
 (w^*, b^*) = \arg\min_{(w, b)} \sum_{i=1}^{m} (f(x_i) - y_i)^2 \tag{15}
\]

where \((w^*, b^*)\) is the optimal parameter, and the minimum value of \((w, b)\) is taken for the above equation.

The Lasso model was a compression estimation method with the idea of reducing the variable set (decreasing order). By constructing a penalty function, it could compress the coefficients of variables and made some regression coefficients become 0, so as to achieve the purpose of variable selection.

Given \( n \) data samples \( \{(x_1, y_1), (x_2, y_2), \ldots, (x_n, y_n)\} \) where each \( x_i \in \mathbb{R}^d \) was a -dimensional vector, i.e., each observed data point was composed of the values of \( d \) variables, and each \( y_i \in \mathbb{R} \) was a real value. What we had to do was to find a map \( f : \mathbb{R}^d \rightarrow \mathbb{R} \) that minimized the sum of squared errors based on the observed data points. The optimization objective is given as follows:

\[
 \beta^* = \arg\min_{\beta} \frac{1}{n} \sum_{i=0}^{n} \left( y_i - \hat{y} - \beta^T (x_i - \bar{x}) \right)^2 \tag{16}
\]

where \( \beta \in \mathbb{R}^d \) is the optimized coefficient.

If Eq. 16 is expressed in matrix form, denoted by \( X = [x_1; x_2; \ldots; x_n]^T \), where each data point \( x_i \) was regarded as a column vector, then \( X \in \mathbb{R}^{n \times d} \), denoted as \( y = [y_1; y_2; \ldots; y_n]^T \), then the optimization objective in matrix form is given as follows:

\[
 \beta^* = \arg\min_{\beta} \frac{1}{n} \|y - X\beta\|^2 \tag{17}
\]

Lasso added the L1 regularization term (see Eq. 18) to make the model avoid over-fitting.

\[
 ||\beta||_1 = \sum_{j=1}^{d} |\beta_j| ; \quad 1 << j << d \tag{18}
\]

Then, the optimization objective function of Lasso is expressed as the following equation:

\[
 \beta^* = \arg\min_{\beta} \frac{1}{n} \|y - X\beta\|^2 + \lambda ||\beta||_1 \tag{19}
\]

PLS model was a many-to-many linear regression modeling method, i.e., there are multiple independent variables and multiple dependent variables. It found the best functional fit for a set of data by minimizing the sum of squared errors.

The general multivariate underlying model of PLS is given by the following equations:

\[
 X = TP^T + E \tag{20}
\]

\[
 Y = UQ^T + F
\]

where \( X \) is a \( n \times m \) prediction matrix, \( Y \) is a \( n \times p \) response matrix; \( T \) and \( U \) are \( n \times l \) matrices and both of them are the projections of \( X \) and \( Y \) in the higher dimensional space; \( P \) and \( Q \) are the orthogonal loading matrices of \( m \times l \) and \( p \times l \), respectively, and the matrices \( E \) and \( F \) are error terms, normally distributed random variables subject to independent and identical distributions. Decompose \( X \) and \( Y \) to maximize the covariance between \( T \) and \( U \).

### Gene-Level Analyses

We analyzed the voxel-based features using gene-level analysis. First, quality control (QC) was performed using the PLINK version 1.9 software\(^1\) (Purcell et al., 2007). We performed genome-wide association studies (GWASs) using the image data and genetic data in whole brain using the linear regression in PLINK. Age, gender, education, and the top 10 principal components from population stratification analysis were included as covariates. A total of 5,574,300 single-nucleotide polymorphisms (SNPs) were obtained by QC. We applied ECS method (Li et al., 2019) to assign SNPs to autosomal genes. Then the significant genes was obtained by Bonferroni correction (family-wise error rate \( p \)-value < 0.05).

### Pathway-Level Analyses

Using the resulting genes, we performed the pathway analysis to assess the biological significance of these features (Bu et al., 2021). KOBAS-I (Bu et al., 2021) pathway analysis tool (KOBAS; bioinfo.org) and the Kyoto Encyclopedia of Genes and Genomes database were applied to pathway analysis of the identified genes \( (P < 0.001) \).

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\(^1\)https://www.cog-genomics.org/plink/1.9/
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FIGURE 2 | The accuracy curves were obtained through ten experiments for five methods in three groups. (A) Prediction accuracy of EMCI-HC group. (B) Prediction accuracy of LMCI-HC group. (C) Prediction accuracy of AD-HC group.

TABLE 2 | Test results of different models.

| Group   | Model | Validation set |  | Test set |  |
|---------|-------|----------------|------------------|------------------|------------------|
|         |       | Accuracy | Precision | Recall | F-Measure | Accuracy | Precision | Recall | F-Measure |
| EMCI-HC | Linear regression | 0.67 | 0.67 | 0.67 | 0.67 | 0.73 | 0.73 | 0.73 | 0.73 |
|         | Lasso | 0.79 | 0.79 | 0.79 | 0.79 | 0.80 | 0.80 | 0.80 | 0.80 |
|         | PLS | 0.8 | 0.8 | 0.8 | 0.8 | 0.82 | 0.81 | 0.81 | 0.81 |
|         | SVM | 0.73 | 0.73 | 0.73 | 0.73 | 0.76 | 0.76 | 0.76 | 0.76 |
|         | RS-SVM | **0.86** | **0.86** | **0.86** | **0.86** | **0.86** | **0.86** | **0.86** | **0.86** |
| LMCI-HC | Linear regression | 0.62 | 0.62 | 0.62 | 0.62 | 0.78 | 0.78 | 0.77 | 0.77 |
|         | Lasso | 0.80 | 0.80 | 0.80 | 0.80 | 0.81 | 0.81 | 0.81 | 0.81 |
|         | PLS | 0.65 | 0.64 | 0.65 | 0.64 | 0.66 | 0.65 | 0.66 | 0.65 |
|         | SVM | 0.73 | 0.73 | 0.73 | 0.73 | 0.74 | 0.74 | 0.74 | 0.74 |
|         | RS-SVM | **0.85** | **0.85** | **0.85** | **0.85** | **0.85** | **0.85** | **0.85** | **0.85** |
| AD-HC   | Linear regression | 0.85 | 0.85 | 0.84 | 0.84 | 0.84 | 0.85 | 0.84 | 0.84 |
|         | Lasso | 0.85 | 0.85 | 0.85 | 0.85 | 0.85 | 0.85 | 0.85 | 0.85 |
|         | PLS | 0.91 | 0.92 | 0.91 | 0.91 | 0.91 | 0.91 | 0.91 | 0.91 |
|         | SVM | 0.87 | 0.87 | 0.87 | 0.87 | 0.87 | 0.87 | 0.87 | 0.87 |
|         | RS-SVM | **0.91** | **0.91** | **0.91** | **0.91** | **0.93** | **0.93** | **0.93** | **0.93** |

Bold fonts represented the model and experimental results in this paper.

RESULTS

In recent studies, machine learning was used to detect the subjects and brain regions of AD (Zhang et al., 2016) and the brain functional statuses of EMCI (Jiao et al., 2020a) and to identify AD and MCI (Zhang et al., 2015b; Wang et al., 2016). In this work, we applied a novel feature extraction method and SVM to obtain the features classified EMCI, LMCI, AD, and HC.

Comparison of the Five Methods

We employed the test set to evaluate the classification capability of the five methods, and the experiments were repeated 10 times with the selected parameter combination in each method. As shown in Figure 2, the RS-SVM model has the best prediction accuracy. The AD-HC group had more than 90% prediction accuracy, while the other four methods all peaked below 90%. The prediction accuracy of both the EMCI-HC and LMCI-HC groups exceeded 85%, while the peak values of the other four methods were all below 80%. The curves in Figure 2 also showed that RS-SVM had good stability. In ten replicates, the difference in accuracy was less than 10%. These analyses demonstrated the satisfactory classification ability and stability of the RS-SVM model.

Machine learning had been gradually maturing and has been applied to the classification and prediction of AD. We applied the validation set to obtain the optimal parameters and the test set to evaluate the classification capability of the five methods. The evaluation metrics of the five methods implemented in EMCI-HC, LMCI-HC, and AD-HC were shown in Table 2. As shown in Table 2, the RS-SVM has the best accuracy, precision, recall, and F-measure. Only the values of RS-SVM increase with the optimal parameters, and the values of other models are stable. In the AD-HC, EMCI-HC, and LMCI-HC groups, the F-measure of RS-SVM in the
Therefore, we performed GWAS of these features to analyze their and HC and were meaningful for the identification of AD. The identified features were excellent in the classification of AD obtained for all five models (all above 0.8). This proved that features were applied to the five models, good results were performance than single SVM. In addition, since the same scalable, and SVM combined with other schemes have better high to low. This also indicated that the RS-SVM model was of RS-SVM in the test set were 0.93, 0.86, and 0.85 from high to low. This also indicated that the RS-SVM model was easier affected by the characteristics of high-dimensional brain from MRI. Although the voxel-based research could solve AD that extracted the voxel-based ROI, which included the whole genome. All genes with conditional association P-values passing Bonferroni correction for family-wise error rate at 0.05 were extracted. We performed the gene-based association analysis by using P-values of 113 novel voxel-based features for identifying susceptibility genes of AD. There are 242 genes (corrected P < 0.001) associated with AD. These top 10 conditionally significant genes are shown in Table 3. Results of Gene-Level Genome-Wide Association Study
We performed the conditional gene-based association scans on whole genome. All genes with conditional association P-values passing Bonferroni correction for family-wise error rate at 0.05 were extracted. We performed the gene-based association analysis by using P-values of 113 novel voxel-based features for identifying susceptibility genes of AD. There are 242 genes (corrected P < 0.001) associated with AD. These top 10 conditionally significant genes are shown in Table 3. Studies have shown that CSMD1 (SNP: rs34464519, CorrectedP: 1.74556E-36) was related to AD (Stepanov et al., 2014; Li et al., 2020; Bi et al., 2021a). RBFOX1 (SNP: rs55642412, CorrectedP: 3.18755E-23) has been found to play a role in neuronal development (Raghavan et al., 2020). PTPRD (SNP: rs62538998, CorrectedP: 1.92988E-19) has been confirmed to be related to AD and MCI in previous studies (Huang et al., 2021). DLGAP2 (SNP: rs72507619, CorrectedP: 3.74049E-17) was found to be predominantly expressed in the brain and associated with a wide variety of neurological disorders (Linthorst et al., 2020). WWOX gene has been reported to be a potential mechanism that may be involved in the pathogenesis of AD, focusing on the cell death signaling pathway in neurons (Teng et al., 2012).

Results of Pathway-Level Genome-Wide Association Study
Detecting pathways may provide useful information about the pathogenic molecular mechanism underlying AD. In our work, 70 enriched pathways were identified. The top 10 significant pathways are shown in Table 4. Impaired insulin secretion was associated with higher risk of any dementia and cognitive impairment (Rönnemaa et al., 2008). Oxytocin signaling pathway was neuroprotective to many neurological disorders, such as AD (Almansoub et al., 2020). Vascular smooth muscle contraction was associated with the development of neurodegeneration in AD (Hald et al., 2016).

DISCUSSION
We proposed a voxel-based three-level analysis framework for AD that extracted the voxel-based ROI, which included the whole brain from MRI. Although the voxel-based research could solve the limitations of the research method based on the ROI, it was more easily affected by the characteristics of high-dimensional data. Feature selection in RS-SVM model solved the dimensional disaster caused by too many attributes. In this work, we identified 136, 141, and 113 MRI features for EMCI-HC, LMCI-HC, and AD-HC groups, respectively.

We performed RS-SVM model to identify important brain regions such as hippocampus, amygdala, angular gyrus, and calcarine sulcus for AD. The hippocampus was located in the midlimbic system of the brain and had an important impact on memory and cognitive function. Many studies had shown that abnormalities in hippocampal volume and function were closely linked to AD. Although many patients had not shown symptoms

TABLE 3 | Top 10 conditionally significant genes were obtained. Chr represents Chromosome; Gene represents the gene name; CorrectedP represents P-value generated by Bonferroni correction.

| No. | Chr | Gene | CorrectedP   |
|-----|-----|------|--------------|
| 1   | 8   | CSMD1| 1.74556E-36  |
| 2   | 16  | RBFOX1| 3.18755E-23  |
| 3   | 16  | CDH13| 1.07119E-20   |
| 4   | 9   | PTPRD| 1.92988E-19   |
| 5   | 8   | DLGAP2| 3.74049E-17   |
| 6   | 11  | CNTN5| 4.81388E-16   |
| 7   | 7   | MAGI2| 5.93057E-16   |
| 8   | 20  | MACROD2| 1.50704E-14  |
| 9   | 16  | WWOX| 1.64798E-14   |
| 10  | 3   | CNTN4| 1.87567E-13   |

TABLE 4 | Top 10 significant pathways.

| NO. | Pathways                          | CorrectedP-value | Gene                              |
|-----|-----------------------------------|------------------|-----------------------------------|
| 1   | Insulin secretion                 | 1.01E-06         | PLCB1, PRKCB, PRKCA, CREB5, RYR2, CHRM3, KNCMA1, RAPGEF4, CACNA1C |
| 2   | Oxytocin signaling pathway        | 4.80E-06         | PLCB1, PRKAG2, PRKCA, CACNB2, RYR3, RYR2, PRKCB, CACNA1C, ITPR2, CACNA2D |
| 3   | Salivary secretion                | 7.70E-06         | PLCB1, PRKCA, RYR3, PRKCB, CHRIM3, KNCMA1, PRKG1, ITPR2 |
| 4   | Vascular smooth muscle contraction| 7.94E-06         | PLCB1, CACNA1C, PRKCH, PRKCA, PRKCB, PRKCE, KNCMA1, PRKG1, ITPR2 |
| 5   | Calcium signaling pathway         | 1.48E-05         | PLCB1, PRKCB, ERBB4, PRKCA, RYR3, RYR2, CHRIM3, CACNA1C, ITPR2, PDE1A |
| 6   | Glutamatergic synapse             | 2.08E-05         | PLCB1, CACNA1C, PRKCA, GRIK2, PRKCB, DLGAP1, ITPR2, GRM7 |
| 7   | Morphine addiction               | 5.00E-05         | PRKCA, POE1A, PRKCB, PDE3A, GABBR3, PDE4D, PDE10A |
| 8   | Circadian entrainment            | 5.56E-05         | PLCB1, PRKCB, PRKCA, RYR3, RYR2, CACNA1C, PRKG1 |
| 9   | Pancreatic secretion              | 5.56E-05         | PLCB1, PRKCB, PRKCA, RYR2, CHRIM3, KNCMA1, ITPR2 |
| 10  | Aldosterone synthesis and secretion| 5.56E-05        | PLCB1, CACNA1C, PRKCA, CREB5, PRKCB, PRKCE, ITPR2 |

validationset were 0.91, 0.86, and 0.85 from high to low. In the AD-HC, EMCI-HC, and LMCI-HC groups, the F-measure of RS-SVM in the test set were 0.93, 0.86, and 0.85 from high to low. This also indicated that the RS-SVM model was scalable, and SVM combined with other schemes have better performance than single SVM. In addition, since the same features were applied to the five models, good results were obtained for all five models (all above 0.8). This proved that the identified features were excellent in the classification of AD and HC and were meaningful for the identification of AD. Therefore, we performed GWAS of these features to analyze their biological significance.
FIGURE 3 | ROC curve of five classification methods for three groups. (A) Prediction ROC of EMCI-HC group. (B) Prediction ROC of LMCI-HC group. (C) Prediction ROC of AD-HC group.

FIGURE 4 | An image showing the relation of genes and pathways.
obtained framework based on RS-SVM was optimal compared with the regression, Lasso, PLS, and SVM). This proved that the SVM, consistently outperforms other methods (i.e., linear studies demonstrate that altered levels of intracellular Ca ions (Ca\(^{2+}\)) to increase intracellular Ca\(^{2+}\) concentration. The studies demonstrate that altered levels of intracellular Ca\(^{2+}\) affect neurodegeneration (Gong et al., 2018; Nilipour et al., 2018). It has been shown that the expression of CACNA1C inhibits the hyperphosphorylation of Tau protein (Jiang et al., 2018).

Some pathways include insulin secretion, oxytocin signaling pathway, salivary secretion, vascular smooth muscle contraction, and AD closely related to genes (Figure 4).

In summary, our proposed framework based on RS-SVM performed well in features constructed, and the framework had good classification performance for EMCI-HC, LMCI-HC, and AD-HC groups. In particular, AD-HC was the best in terms of classification accuracy. Several pathogenic genes and abnormal subregions identified singing this framework are related to AD. Therefore, we speculate that the remaining genes identified could be regarded as the candidate genes for AD. The discoveries in this study provide new candidate genes for AD, and the constructed features can be regarded as a new indicator to distinguish AD from HC.

**DATA AVAILABILITY STATEMENT**

Rs-fMRI data were downloaded from the Alzheimer’s Disease Neuroimaging Initiative (ADNI) database (http://adni.loni.usc.edu/). Application for access to the ADNI data can be submitted at http://adni.loni.usc.edu/data-samples/access-data/.

**ETHICS STATEMENT**

The studies involving human participants were reviewed and approved by Data collection and sharing for this project was funded by the Alzheimer’s Disease Neuroimaging Initiative (ADNI). We applied the access from ADNI. The patients/participants provided their written informed consent to participate in this study. Written informed consent was obtained from the individual(s) for the publication of any potentially identifiable images or data included in this article.

**AUTHOR CONTRIBUTIONS**

XM, YWu, WL, and ZJ led and supervised research. XM, YWu, and WL designed the research and wrote the article. XM, YWu, WL, and ZJ performed data preprocessing and quality control. WL did gene-survey support vector machines. WL, YWa, and ZX performed features extraction and selection and random survey support vector machines. WL, YWa, and ZX performed features extraction and selection and random selections.

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