Early Detection of Alzheimer’s Disease Based on Clinical Trials, Three-Dimensional Imaging Data, and Personal Information Using Autoencoders

Abstract

**Background:** A timely diagnosis of Alzheimer’s disease (AD) is crucial to obtain more practical treatments. In this article, a novel approach using Auto-Encoder Neural Networks (AENN) for early detection of AD was proposed. **Method:** The proposed method mainly deals with the classification of multimodal data and the imputation of missing data. The data under study involve the MiniMental State Examination, magnetic resonance imaging, positron emission tomography, cerebrospinal fluid data, and personal information. Natural logarithm was used for normalizing the data. The Auto-Encoder Neural Networks was used for imputing missing data. Principal component analysis algorithm was used for reducing dimensionality of data. Support Vector Machine (SVM) was used as classifier. The proposed method was evaluated using Alzheimer’s Disease Neuroimaging Initiative (ADNI) database. Then, 10-fold cross-validation was used to audit the detection accuracy of the method. **Results:** The effectiveness of the proposed approach was studied under several scenarios considering 705 cases of ADNI database. In three binary classification problems, that is AD vs. normal controls (NCs), mild cognitive impairment (MCI) vs. NC, and MCI vs. AD, we obtained the accuracies of 95.57%, 83.01%, and 78.67%, respectively. **Conclusion:** Experimental results revealed that the proposed method significantly outperformed most of the state-of-the-art methods.

**Keywords:** Alzheimer’s disease, autoencoders, cerebrospinal fluid, early detection, magnetic resonance imaging, Mini-Mental State Examination, missing data, positron emission tomography

Introduction

Alzheimer’s disease (AD) is one of the neurodegenerative brain dysfunctions frequently observed in elderly people. Amyloid plaques, neurofibrillary tangles, and histopathologic changes are commonly used for characterizing this disease from dementia because of their association with neuronal loss and reduction in volume of the brain.[1] This disease starts with loss of memory, progressively, in the following cognitive functions, getting worse, and worse until patients lose their ability to remember recent events, and cannot recognize very familiar persons and things. In the upcoming years, the dominance and severity of this disease is expected to rise[2] due to the regular growth of the aged population all over the world. In this sequel, it could be one of the major causes of death in the near future. Nevertheless, there is not any thorough treatment for AD yet. Thus, early diagnosis of AD could be of a considerable help in increasing the patients’ survival rate. As such, many biomedical imaging techniques for early detection of AD are well developed and employed by the researchers including magnetic resonance imaging (MRI),[3-5] positron emission tomography (PET),[6,7] and others such as cerebrospinal fluid (CSF),[8] AD Assessment Scale–Cognitive behavior section,[9] and Mini-Mental State Examination (MMSE).[9]

Machine learning techniques are used on medical images of the brain for automatic diagnosis of AD in many studies such as Acharya et al. in[10] diagnosed AD with an accuracy of 99.48% on 33 patients and subjects using MRI images from the Harvard Brain Atlas. Wang et al. in[11] using 196 MRI images half train and half test, employed convolutional NN for a deep learning method for distinguishing AD from normal controls (NCs) and achieved 97.65% accuracy. In addition,

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Various approaches based on structural images have been proposed. Almost all of these CAD systems have three main steps, which are preprocessing, feature extraction, and classification. The procedure of the first step sets different images from different individuals, with brains of different sizes and shapes, at a comparable condition. At the second step, feature extraction algorithm converts the input data into small vectors. All the relevant information of the input data must be in these vectors. The classifier determines that the vectors are more similar to mild cognitive impairment (MCI) patient vectors, to AD patient vectors, or to NC vectors. Richer information can help to improve diagnostic accuracy.

Metrics of the entorhinal cortex have been used to discriminate AD patients and NCs but most of the studies on AD have used manual segmentation of the hippocampus in MR images. These studies have high efficiency in distinguishing between AD patients and NCs. Automatic hippocampal volume-measuring methods almost have equal results. Hippocampal volumes and entorhinal cortex metrics seem to be equally accurate in distinguishing between AD patients and NCs. Single tissue such as the hippocampus alone is not enough for the accurate diagnosis of the disease, and the combination of different structures has proven to be more useful for distinguishing AD patients from NCs. Therefore, using multivariate approaches, many variables simultaneously and observation of the essential patterns of different regions of the brain data can be analyzed. There were different techniques such as principal component analysis (PCA), artificial neural networks, fuzzy neural networks, partial least square, and support vector machine (SVM) to classify MRI data according to prior studies. Here, the SVM method utilizing autoencoder has been used for data analysis.

Combined techniques can use different modalities including MRI, PET, and other neurological data to diagnose AD/MCI patients from healthy people. Lahmiri and Shmuel used MRI images from the AD neuroimaging initiative (ADNI) dataset for distinguishing AD patients from NCs. They reported complete performance (100% accuracy) in distinguishing between the two groups. Magsood et al. reported a multiple classification using transfer learning on AD, while Beheshti et al. classified AD vs. NC with a great rate of accuracy using only MRI data. Spasov et al. classified progressive MCI vs. static MCI using combined MRI, APOe4 genetic, and cognitive measures include. Furthermore, automatic diagnosis of MCI using electroencephalogram spectral features is done. In this study, the feature extraction and feature combination were often performed independently. As investigated in the previous studies, there exist inherent relations between modalities of MRI and PET. Thus, finding the shared feature representation which combines the complementary information from modalities (e.g., PET, MRI, and CSF) is useful to enhance the diagnosis performance of AD and MCI patients from NCs.

The data used in this article are presented in the next section, preprocessing technique, feature selection, description of proposed methodology, and classification methods, which are shown in the “Experiments and Results” section. Proposed method and the experimental results in this work are provided in the “Discussion” section. The conclusions are presented in the final section.

The dataset was used in this article, which was obtained from various sources associated with the ADNI database (adni.loni.usc.edu). The foremost usages of determining sensitive and specific biomarkers associated with the early progression include the development of new treatments with monitoring their effectiveness and the reduction in the time and cost of clinical experiments. Moreover, it should be noted that the obtained dataset was compliant only with the first examination of each patient involving 705 patients’ images.

The demographic data of patients are summarized in Table 1. The MRI and positron emission tomography (PET) data were downloaded from the ADNI. A detailed description of PET protocols and acquisition can be found at www.adni-info.org. The CSF biomarker, the personal information, and the MMSE scores were downloaded from the ADNI website since July 2015.

### Materials and Methods

#### Feature extraction

The features of MRI images were extracted based on regions of interest (ROIs). The MMSE scores, PET extracted features, and CSF measures are obtained from the ADNI database. The volume and the voxel values of specific regions such as hippocampus, entorhinal cortex, temporal, parietal lobes, and ventricles are the main affected regions of the brain on which AD attacks and

| Table 1: Data of patients in the Alzheimer’s disease neuroimaging initiative database |
|--------------------------------|----|---|----------|--------|--------|-----------------|-------------|--------|
| **Count** | **Male** | **Female** | **Married** | **Widowed** | **Divorced** | **Never married** | **Average of age** | **Average of MMSE** |
| AD  | 156 | 76 | 80 | 127 | 18 | 8 | 3 | 74.89 | 23.32 |
| NC  | 211 | 110 | 101 | 142 | 38 | 17 | 14 | 75.91 | 29.13 |
| MCI | 338 | 215 | 123 | 269 | 39 | 24 | 6 | 74.51 | 27.05 |
| Total | 705 | 401 | 304 | 538 | 95 | 49 | 23 | 75.01 | 26.85 |

AD – Alzheimer’s disease; NC – Normal control; MCI – Mild cognitive impairment; MMSE – Mini-Mental State Examination
destroys several brain cells during the early stages of its progression.\cite{32} Figure 1 represents MRI samples of NCs and AD patients. The figure shows decreased gray matter volume in an AD patient compared.

In Figure 2, the segmented brain and corresponding labels containing the aforementioned ROIs are demonstrated. The ROIs included candidate input variables for diagnosis of Alzheimer’s disease, measures of regional cortical thickness.\cite{33} The ROI voxels’ values and their volumes for every MRI and PET images were extracted using SPM toolbox in MATLAB. Overall, the features MMSE, personal information, CSF biomarkers, and PET and MRI data have been used to classify NCs and MCI and AD patients.

**Autoencoder neural networks**

An autoencoder is a neural network consisting of at least three layers: an input layer, a hidden layer, and an output layer, as shown in Figure 3 (it can have multiple hidden layers though). The Neurons of the first layer represent the original input vector. The hidden layer can be seen as a high-level representation of the previous layer. The output layer is a specific representation of the input layer with the same dimensionality as the input one.\cite{34,35} The input layer sends features from MRI, PET, and CSF data to the hidden layers. Then, the hidden layer performs nonlinear transformations on the received data and imposes some optimization procedures to reconstruct the original instance.

The activation signals are propagated forward through the network which can be considered sigmoid function or hyperbolic tangent function to introduce nonlinearity for the network for complex relationships of the model.\cite{36}

By modifying the number of neurons at each hidden layer, we were able to perform the feature dimensionality reduction. The hidden layers of sparse autoencoder will be trained one at a time, and they will be stacked to form a complete neural network by removing the temporary output layer.\cite{37}

**Classification and evaluation**

One of the most common techniques for dimension reduction of data is PCA. It maps the data to a lower dimension while maintaining the variance of the data. Reduction of used storage space and computation time and elimination of correlated features are advantages of using PCA. Loosing some information of the original data, failing when covariance are not enough to define data, and indetermination of the number of principal components to keep information of data are disadvantages of using it.

SVM is one of the widely used classification algorithms in neuroimaging data.\cite{4,38,39} Classification efficiency of SVM in training high dimensional data has been proven. Moreover, SVM has been applied to voice activity detection, pattern recognition, classification, and regression analysis.\cite{40,41} It is used to separate a set of training data with a hyperplane that is maximally distant from the two classes. SVM is the most common and efficient classifier in binary classification. Here, SVM was used to distinguish between AD and MCI patients and NCs, pairwise.

One of the well-known evaluation measures is accuracy rate of a classification procedure, which computes the ratio between correctly classified samples and total samples. Sensitivity and specificity are the other evaluation metrics. Other widely used parameters to describe the performance of diagnostic procedures are positive predictive value (PPV), negative predictive value (NPV), area under the curve (AUC), and receiver operating...

![Figure 1: Magnetic resonance imaging sample, (a) a normal control and (b) a Alzheimer’s disease patient](image1.png)

![Figure 2: Sub-regions of medial surface of the human cerebral cortex](image2.png)
characteristic (ROC). The accuracy, sensitivity, specificity, PPV, and NPV are defined as below:

\[
\text{Accuracy} = \frac{TP + TN}{TP + TN + FP + FN} \tag{1}
\]

\[
\text{Sensitivity} = \frac{TP}{TP + FN} \tag{2}
\]

\[
\text{Specificity} = \frac{TN}{TN + FP} \tag{3}
\]

\[
\text{PPV} = \frac{TP}{TP + FP} \tag{4}
\]

\[
\text{NPV} = \frac{TN}{TN + FN} \tag{5}
\]

Where TP stands for the number of true positives (number of correctly classified AD or MCI patients). TN is the number of true negatives (number of correctly classified NC or MCI patients). FP stands for the number of false positives (number of NCs classified as AD or MCI patients or MCI classified as AD). FN is the number of false negatives (number of AD or MCI patients classified as MCI patients or NCs, wrongly).

Specificity and sensitivity were used to evaluate the rate of actual positives or negatives, which were identified correctly, for example, the percentage of AD or MCI patients or NCs. These measures show the detection capability of a method between AD, MCI, and NC patterns. These metrics were measured using K-fold cross-validation (with k = 10). K-fold cross-validation has been used to audit the partial accuracy of different multivariate analysis methods applied to the segmentation of brain dementia from AD. In the K-fold method, 10 selected sets of AD and MCI patients and NCs are sampled randomly. It holds out a set for testing purposes and trains the classifier with the remaining sets. This has to be done for all 10 sets, and an average value of the evaluation parameters is calculated. We train the SVM classifier until the cross-validation loss (error) should be less or equal to 0.05.

**Proposed method**

A method for early detection of AD was proposed. Data normalization, replacement of missing data using best possibility, and classification of AD, NC and MCI using SVM classifier on multimodal data (MRI and PET images of brain and CSF biomarker, MMSE scores, age, education, gender, and marriage information), and finally, reducing dimension of input data was the stages of the proposed method. Before doing so, proper features of the interested area of the images had to be extracted. Then, missing data were replaced using autoencoder (a type of neural network). Finally, classification was done using a SVM classifier. Figure 4 displays the diagram of the proposed algorithm of this article.

To distinguish between AD and MCI patients and NCs, a method for the integration and classification of the baseline MRI, PET, MMSE scores, personal information, and CSF data has been developed. The composite of extracted features has been used in this study. The measurements were combined as a long feature vector which was considered as the input of the classifier. This is called data concatenation. SVM classifier was used to differentiate between participants using all features.

In the first step of our work, the feature vector consisting of average voxel values from MR images and also volume of MR images of ROI using VBM-SPM were extracted. MRI of specific regions such as hippocampus, entorhinal cortex, temporal and parietal lobes, and ventricles could assess volumetric changes of brain structure and so on. They have been identified as ROI for AD diagnosis.

Because of the absence of some PET images and CSF measurement, compensating for missing data was necessary. One of the usual methods in dealing with missing data is filling missed data using average value of existing data. In this method, average of existing values of each column is computed and placed in every empty cell of vector of features in the corresponding column, as in Eq. 6.

\[
MDa_{ij} = \frac{1}{n} \sum_i ED_j \tag{6}
\]

Where \( MDa_{ij} \) refers to missing data element gained from average of \( ED_i \), which is existing data column, \( i \) is the row number of full data matrix, and \( j \) is element number \((i, j \in \{\text{missing data indices set}\})\).

To find the best value for missing data in each vector, after training an autoencoder using complete vectors
of features, missing data can be obtained by sending a vector having missing data (replaced in advance by average of the existing data) to the trained autoencoder. Then, by replacing the corresponding output cell to each cell having missing data, a good replacement for missed data could be found. This process can be shown in Eq. 7:

$$ MD_{ae}^i = MD_{a}^i \times AE $$

(7)

Where $MD_{a}^i$ is $i^{th}$ vector in data matrix containing missing data $MD_{ae}^i$ and is gained vector after multiplying autoencoder weights by $MD_{a}^i$.

Then, in Eq. 7, missing data taken from Eq. 6 are replaced by corresponding values of Eq. 7.

$$ MD_{a}^j = MD_{ae}^j $$

(8)

where $MD_{a}^j$ is averaged missing data, and $MD_{ae}^j$ is rebuilt missing data from Eq. 7. Therefore, $MD_{a}$ after Eq. 8 is the final processed data.

Where $MD_{a}$ is the vector from the Eq. 6, $MD_{a}L$ is the obtained vector from natural logarithm (output of logarithm on 1 will be 0, and +1 is to prevent this), and $MD_{a}L$s is scaled $MD_{a}L$ vector to range (0, 1).

In the third step, after data normalization using natural logarithm, an autoencoder using complete vectors of all data was made. In this step, missing data vectors were not used, and the structure was made using only perfect vectors of data.

In the fourth step, missing data were replaced using output of autoencoder for the corresponding vector.

In the fifth step, the PCA was used to classify patients.

In the final step, classification was done and proposed algorithm was evaluated using a 10-fold cross-validation method. Here, all data were split to 10 approximately equal parts. Nine parts were used as a training data set and one part was used as a test dataset.

The proposed method can be summarized as below:

1. Feature extraction and selection
2. Building missing data using the average of existing data
3. Normalizing data using natural logarithm
4. Making an autoencoder using complete vectors of data
5. Missing data imputation using prebuilt autoencoder
6. Using PCA to reduce dimension of data as input for classifier
7. Classification using SVM
8. Evaluation using 10-fold cross-validation.

**Experiments and Results**

In this part, experimental results using MRI, PET, MMSE, personal information, and CSF data have been represented. SVM classifier using linear kernel was evaluated to gain higher performance for AD diagnosis. The purpose of this work is to distinguish between NCs and MCI and AD patients. The method was tested to classify NC and AD data first. Then, the method was tested to classify NC and MCI data and finally was tested to classify MCI
and AD data. These evaluations were done using only MRI data and then using all the data. The performance of this method was calculated by means of 10-fold cross-validation.

SVM on the data has been applied with and without missing data imputation and with and without feature reduction using autoencoder. Its performance was compared with the performance of the standard SVM without missing data imputation. After the missing data imputation, the accuracy of classification was improved. In addition, dimension reduction was done independent of the classification procedure.

The proposed method was evaluated to distinguish between AD and MCI patients and NCs, and these results were compared with other methods. To this end, each group of participants is compared pairwise. For this part, totally 144 selected features are used (including 132 MRI, 1 MMSE, 4 personal information, 3 CSF, and 4 PET images).

Table 2 demonstrates the performance of proposed method for classifying AD and MCI patients and NCs, using SVM classifier on various datasets, including whole data, averaged missing data, rebuilt missing data, only MRI data, only MMSE scores, only demographic data. evaluate the performance of the proposed method were investigated including classification accuracy (ACC), sensitivity (SEN), specificity (SPE), PPV, NPV, and AUC. Table 2 meticulously shows that the combined data (took from various sources) achieved remarkably higher performance than that on the single dataset case. The highest accuracy (95.57%) was for combined data, as shown in the table.

From Table 2 and Figures 5-8, the ROCs, boxplots, and AUC values increased after recovering missing data using autoencoder. Using the rebuilt missing data method, the boxplots compressed inward more than that of the averaging method for estimating the missing data. Such compression depicts higher stability in the classification model for rebuilt missing data method. Moreover, the accuracy of

| Data                        | Classes | Accuracy (%) | Sensitivity (%) | Specificity (%) | PPV (%) | NPV (%) | AUC  |
|-----------------------------|---------|--------------|-----------------|-----------------|---------|---------|------|
| All data, averaged missing data | NC-AD   | 95.37        | 100             | 92.50           | 89.23   | 100     | 0.9625 |
|                             | MCI-AD  | 82.63        | 72.09           | 87.86           | 74.60   | 86.40   | 0.7988 |
|                             | NC-MCI  | 78.65        | 65.86           | 94.06           | 93.02   | 69.59   | 0.7990 |
| All data, rebuilt missing data | NC-AD   | 95.57        | 100             | 92.80           | 89.67   | 100     | 0.9640 |
|                             | MCI-AD  | 83.01        | 72.72           | 88.08           | 74.92   | 86.81   | 0.8030 |
|                             | NC-MCI  | 78.67        | 93.87           | 65.83           | 69.89   | 92.70   | 0.7983 |
| Only MRI data               | NC-AD   | 84.46        | 81.87           | 86.41           | 81.79   | 86.45   | 0.8407 |
|                             | MCI-AD  | 66.84        | 48.70           | 82.31           | 70.11   | 65.31   | 0.6539 |
|                             | NC-MCI  | 66.97        | 78.94           | 55.28           | 63.29   | 72.88   | 0.6707 |
| Only MMSE data              | NC-AD   | 91.83        | 99.21           | 87.81           | 81.59   | 99.52   | 0.9351 |
|                             | MCI-AD  | 78.92        | 64.14           | 88.38           | 77.81   | 79.44   | 0.7613 |
|                             | NC-MCI  | 70.26        | 93.14           | 56.88           | 55.81   | 93.41   | 0.7499 |
| Only demographic data       | NC-AD   | 60.61        | 61.82           | 60.44           | 21.82   | 89.76   | 0.6065 |
|                             | MCI-AD  | 66.11        | 46.54           | 73.79           | 41.07   | 77.86   | 0.5977 |
|                             | NC-MCI  | 55.86        | 68.43           | 44.59           | 52.57   | 61.14   | 0.5644 |

ACC – Classification accuracy; SEN – Sensitivity; SPE – Specificity; PPV – Positive predictive value; NPV – Negative predictive value; AUC – Area under the curve; AD – Alzheimer’s disease; NC – Normal control; MIC – Mild cognitive impairment; MMSE – Mini-Mental State Examination; MRI – Magnetic resonance imaging
NC versus MCI classification with autoencoder increased after recovering missing data. Yet, exploiting the average of missing data obtained drastically better results in terms of distinguishing between MCI and AD patients. Line 6 of the table is bold that shows best accuracy and other performance measures in distinguishing MCI patients from NCs. As in Figure 8, after reducing the dimensionality of the input vector for classification, the boxplots significantly compressed compared to that of the averaged missing data, which depicts less variance (i.e., higher stability) in the classification. Using PPV for missing data is more promising than other methods, and showed better prognosis capability with the proposed method.

**Discussion**

Figures 5-7 and data in Table 2 show that the AUC values increased after recovering missing data (≈0.964 for AD versus NC classification, and 0.803 and 0.799 values for AD versus MCI and for NC versus MCI classification, respectively). The boxplots after recovering missing data were clearly more compressed than that of the averaged missing data mode, which interprets more stability and reliability in the constructed classification model.

One important evidence is that utilizing the combination of different data sources can dramatically alleviate the weaknesses of each dataset. In this study, by combining five different data sources, we obtained high accuracy to distinguish AD and MCI patients and NCs from each other. The weakness can be the dependency of method on collecting several data sources and increasing missing data, a weakness that was handled well in this article using autoencoders.

In Table 3, the proposed method is compared to other methods. As shown in the table, proposed method outperformed other methods in most aspects of performance, and AUC of our method is above other methods. Due to normalization using natural logarithm and missing data imputation using autoencoder, rebuilt missing data method can estimate missed data more effectively. It also gained a higher performance in most aspects of AD diagnosis evaluation metrics. The true sensitivity rate using this method is 100%, which shows a perfect diagnosis of AD patients.

AUC = 1 means that the diagnostic test is perfect in the differentiation of diseased and nondiseased participants. This happens when the distribution of

| Method | Indicator, number of samples and data source | AD versus NC | AUC |
| --- | --- | --- | --- |
| Zhang et al., 2011 [42] | MRI, PET, CSF, MMSE, ADAS-Cog (202, ADNI) | 93.20 | 93.00 | 93.30 | 0.98 |
| Dai et al., 2013 [43] | MRI (83, OASIS) | 90.81 | 92.59 | 91.76 | 0.95 |
| Liu et al., 2016 [44] | MRI, PET (710, ADNI) | 94.65 | 95.03 | 91.76 | 0.95 |
| da Silva Lopes et al., 2010 [45] | EEG Signal (41,-) | 71.5 | 82 | 61 | - |
| Beheshi et al., 2017 [27] | MRI (186, ADNI) | 93.01 | 89.13 | 96.80 | 0.935 |
| Mishra et al., 2018 [46] | MRI (417, ADNI) | 89.15 | 85.06 | 92.53 | 0.93 |
| Khedher et al., 2015 [47] | MRI (818, ADNI) | 89.96 | 92.35 | 86.24 | 0.93 |
| Li et al., 2019 [48] | MRI (1457, ADNI) | 90.00 | 82.00 | 97.00 | 0.95 |
| Ben Ahmed et al., 2014 [49] | MRI (218, ADNI) | 87.00 | 75.50 | 100 | 0.85 |
| Zhou et al., 2018 [50] | MRI (507, ADNI) | 93.75 | 87.5 | 100 | - |
| Suk et al., 2014 [51] | MRI, PET, CSF, MMSE, ADAS-Cog (202, ADNI) | 93.05 | 90.86 | 94.57 | 0.95 |
| Maqsood et al., 2019 [21] | MRI (392, OASIS) | 89.66 | 100 | 82 | - |
| Saravanakumar and Thangaraj 2019 [52] | MRI (-), ADNI | 92.34 | 96 | 87.5 | - |
| Proposed method (EL) | MRI, PET, CSF, MMSE (705, ADNI) | 95.57 | 100 | 95.57 | 0.964 |

AD – Alzheimer’s disease; NC – Normal control; MRI – Magnetic resonance imaging; PET – Positron emission tomography; CSF – Cerebrospinal Fluid; MMSE – Mini-Mental State Examination; ADNI – Alzheimer’s Disease Neuroimaging Initiative; OASIS – Open Access Series of Imaging Studies; ADAS – Alzheimer’s Disease Assessment Scale
As can be seen, the AUC value in the proposed method is equal to 0.964 for the diagnosis of AD versus NC, which shows near excellent differentiation between diseased and nondiseased participants.

**Conclusion**

A novel method for distinguishing between AD and MCI patients and NCs was proposed. This has been done by missing data imputation and feature reduction using autoencoder, MR and PET images, MMSE scores, personal information, and CSF biomarkers using SVM.
with linear kernel. Data normalization was done using natural logarithm. The validity of the method was tested on the ADNI database. The accuracy of the method is tested using K-fold cross-validation (K = 10). To validate the effectiveness of proposed method, several tests are done on ADNI database, and it was compared with other AD diagnosis methods. The results on 705 baseline participants of ADNI show that the proposed method using multimodal data achieves a high accuracy for AD classification (accuracy = 95.57%) and also a significant improvement in area under the curve (AUC = 0.964). In addition, the boxplots are more compressed using rebuilt missing data by autoencoder. Using more modalities of data can help in improving classification metrics. In this article, an advanced method in machine learning, that is autoencoders, was used to overcome the limitation of small number of available individuals for training and testing classifier. It is useful in dealing with missing data for classification. The efficiency of this method was proved by the results of this article. In this article, only baseline data of the participants in ADNI database were used. As future work, longitudinal data can be used to predict the conversion from MCI to AD by finding the patterns of brain atrophy in multiple modalities, specifically in the ROI for AD in the MR and PET images.

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The dataset used in this article was obtained from various sources associated with the ADNI database (adni.loni.usc.edu). The National Institute on Aging, the National Institute of Biomedical Imaging and Bioengineering, the Food and Drug Administration (FDA), private pharmaceutical companies, and nonprofit organizations launched the ADNI in 2003, as a $60 million, 5-year public-private partnership. The primary goal of ADNI was to investigate the possibility of adapting the serial MRI, PET, other biomarkers, and clinical and neuropsychological estimations for evaluating the progression of early stages of AD.

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Conflicts of interest

There are no conflicts of interest.

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