Detection of Alzheimer’s disease by displacement field and machine learning

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ABSTRACT

Aim. Alzheimer’s disease (AD) is a chronic neurodegenerative disease. Recently, computer scientists have developed various methods for early detection based on computer vision and machine learning techniques.

Method. In this study, we proposed a novel AD detection method by displacement field (DF) estimation between a normal brain and an AD brain. The DF was treated as the AD-related features, reduced by principal component analysis (PCA), and finally fed into three classifiers: support vector machine (SVM), generalized eigenvalue proximal SVM (GEPSVM), and twin SVM (TSVM). The 10-fold cross validation repeated 50 times.

Results. The results showed the “DF + PCA + TSVM” achieved the accuracy of 92.75 ± 1.77, sensitivity of 90.56 ± 1.15, specificity of 93.37 ± 2.05, and precision of 79.61 ± 2.21. This result is better than or comparable with not only the other proposed two methods, but also ten state-of-the-art methods. Besides, our method discovers the AD is related to following brain regions disclosed in recent publications: Angular Gyrus, Anterior Cingulate, Cingulate Gyrus, Culmen, Cuneus, Fusiform Gyrus, Inferior Frontal Gyrus, Inferior Occipital Gyrus, Inferior Parietal Lobule, Inferior Semi-Lunar Lobule, Inferior Temporal Gyrus, Insula, Lateral Ventricle, Lingual Gyrus, Medial Frontal Gyrus, Middle Frontal Gyrus, Middle Occipital Gyrus, Middle Temporal Gyrus, Paracentral Lobule, Parahippocampal Gyrus, Postcentral Gyrus, Posterior Cingulate, Precentral Gyrus, Precuneus, Sub-Gyral, Superior Parietal Lobule, Superior Temporal Gyrus, Supramarginal Gyrus, and Uncus.

Conclusion. The displacement filed is effective in detection of AD and related brain-regions.

INTRODUCTION

Alzheimer’s disease (AD) is an abnormal process of aging. AD is a special category of senile dementia (SD) which leads to short-term and long-term memory, thinking, and behavior (Dong et al., 2015a). Research on AD has attracted scholars from all over the world because of its importance and effect on the society. Symptoms of AD may become severe enough to
interfere with daily life, and to death (Goh et al., 2014; Hahn et al., 2013). There is neither cure nor treatment for AD. In 2006, more than twenty million people in the world suffered from this disease (Murphy et al., 2011). AD is predicted to influence 1 in every 85 people worldwide about thirty years later, and more than forty percent of prevalent cases need high level of care (Brookmeyer et al., 2007).

Now that the earth is growing into an aging society, AD has caused heavier burdens on families and society than before (Esposito et al., 2013; Kantanen et al., 2014). In China, AD accounts for more than half of SD, which extracts a total economic loss of more than eighty billion yuan every year, and is responsible for nearly sixty billion yuan in healthcare costs every year (Song & Wang, 2010). In United States, healthcare on people with Alzheimer’s disease currently costs roughly $100 billion per year and is predicted to cost $1 trillion per year by 2050 (Miller, Erlien & Piersol, 2012; Yuan, Machado & Arias-Carrion, 2014).

Nowadays, it is beneficial to develop early and accurate detection methods for AD, which is also necessary for the treatment and management for controlling the deterioration of AD (Zhang et al., 2015b). A 3D scan of the whole brain becomes acceptable and affordable with recent advances in neuroimaging technology (Zhang et al., 2015e; Zhang et al., 2014b), especially by the help of the most popular imaging technique: magnetic resonance imaging (MRI). With its high-resolution magnetic resonance (MR) images, the diagnostic accuracy of AD are greatly enhanced. MR images already play an essential role in detecting AD from normal elder controls (NC).

In this study, we proposed to employ displacement field (DF) to track the morphometry from normal brains to AD brains. The main advantage is that this method does not need to segment region-of-interest (ROI) beforehand, namely, it obtains the DF for the whole image. The classical level-set method was employed to estimate the DF.

This work is structured as follows: ‘State-of-the-Art’ provides the state-of-the-art. ‘Materials and Methods’ presents the materials and methodology. ‘Experiments and Results’ gives the experimental results. ‘Discussion’ discusses the results and the proposed method. ‘Conclusions’ concludes this work. The nomenclature in this work is listed in Table 7.

STATE-OF-THE-ART

In the past, most diagnosis work was carried out by measuring region of interest (ROI) of brain MR images, since researchers already know several typical AD-related regions and corresponding shape deformation (Anagnostopoulos et al., 2013; Kubota, Ushijima & Nishimura, 2006), such as the enlarged ventricles, the shrinkage of hippocampus (HC), and the shrinkage of cortex (Pennanen et al., 2004). Somehow, the ROI-based methods suffer from some shortcomings: (i) The ROI methods need a priori information and expert knowledge. (ii) The detection accuracy relies on the experiences of the interpreters (Yang et al., 2015). (iii) The mutual information among the voxels is difficult to implement (Lee, Park & Han, 2013; Xinyun, Wenlu & Xudong, 2011). (iv) We need to explore other potential regions that may be connected to AD (El-Dahshan et al., 2014). (v) Automatic segmentation of ROI is not feasible in practice, and examiners tend to segment the brain manually (Zhang et al., 2013).
Recently, a new type of methods, the “whole-brain analysis”, gets popularity since it considers all voxels in the brain as a whole. It does not need to segment the brain beforehand, and it does not need any biomarker for classification task. The main disadvantage is the curse of dimensionality, which can be solved as the high-speed computers are very cheap nowadays (Álvarez et al., 2009). The whole-brain analysis heavily relies on pure computation, and it can be accomplished by only computer scientists, after physicians help to label the data as either AD or healthy. Generally, the whole-brain analysis treats the whole brain as a ROI, and it consists of two stages: feature extraction and classification. We reviewed more than 10 literatures, and analyzed them in detail.

Scholars have presented various methods to extract efficient features for AD and other types of pathological brain detection.¹ Some abbreviations are modified to avoid conflict within this paper. In addition, various classification models and methods exist, nevertheless, only a few of them are suitable for MR images. El-Dahshan, Hosny & Salem (2010) extracted the approximation and detail coefficients of 3-level discrete wavelet transform (DWT). They used artificial neural network (ANN) and K-nearest neighbors (KNN) classifiers. Plant et al. (2010) used brain region cluster (BRC). They suggested to use information gain (IG) to evaluate the interestingness of a voxel, and applied clustering algorithm to identify groups of adjacent voxels with a high discriminatory power. They used support vector machine, Bayes statistics, and voting feature intervals (VFI), with the aim of pattern classification. Park (2012) employed manifold learning for classification. Their use of the first and second distance measure resulted in an 18% and 46% error rate for classifying between patients with AD and normal patients, respectively. Chaves et al. (2012) utilized large margin-based methodology for AD detection in SPECT and PET images. Their system yielded accuracy, sensitivity and specificity values of 90.67%, 88% and 93.33% (for PET) and 92.78%, 91.07% and 95.12% (for SPECT), respectively. Saritha, Joseph & Mathew (2013) was the first to use wavelet-entropy in pathological brain detection. They employed spider-web plots to reduce the size of feature space. They also used the probabilistic NN (PNN) for classification. Zhang et al. (2015a) suggested that removing spider-web-plot yielded the same classification performance. Savio & Grana (2013) presented a novel method that employed deformation-based morphometry (DBM) method. They tested five features, and found three features performed excellent as trace of Jacobian matrix (TJM), modulated GM (MGM), and Geodesic Anisotropy (GEODAN). Furthermore, they utilized Pearson's correlation (PEC), Welch’s $t$-test (WTT), and Bhattacharyya distance (BD), to measure the significance of voxel site. Kalbkhani, Shayesteh & Zali-Vargahan (2013) modeled the detail coefficients of 2-level DWT by generalizing autoregressive conditional heteroscedasticity (GARCH) statistical model, and the parameters of GARCH model were considered as the primary feature vector. They tested the KNN and SVM models. Wang et al. (2014) proposed to utilize the undersampling (US) technique on a three-dimensional image. They used singular value decomposition (SVD) to select features. Finally, they combined kernel SVM (KSVM) with decision tree (KSVM-DT). Zhou et al. (2015) followed Saritha’s method, and again employed wavelet-entropy for feature extraction. Naive Bayes classifier (NBC) was employed for abnormal brain detection. Harikumar & Kumar (2015) analyzed
the performance of ANN, in terms of classification of medical images, using wavelets as feature extractor. Their classification accuracy achieved 96%. Zhang et al. (2015c) employed discrete wavelet packet transform (DWPT). They utilized Tsallis entropy to get wavelet packet entropy features. They introduced a generalized eigenvalue proximal support vector machine (GEPSVM). Nazir, Wahid & Khan (2015) suggested to use filters for the removal of noises, and extracted color moments as mean features. Finally, they achieved an overall accuracy of 91.8%. Dong et al. (2015b) employed the eigen-brain (EB) to extract features, and then they employed Welch’s t-Test (WTT) for the aim of reducing features. They proposed to use SVM with radial basis function (RBF). Damodharan & Raghavan (2015) combined tissue segmentation and neural network for brain tumor detection. Zhang et al. (2015f) proposed a novel classification system that implemented 3D discrete wavelet transform (3D-DWT) to extract wavelet coefficients the volumetric image.

After reviewing the latest literature, two findings can be obtained: (1) The DWT based features are efficient. However, we presented a novel feature of displacement field, which is the first to be used in MR images. (2) SVMs were commonly used, compared with conventional decision tree, artificial neural network (Wang et al., 2015a; Zhang et al., 2015d), and other classifiers (Collins & Pape, 2011). Hence, we continue to use SVM. Besides, two variants of SVM are introduced in this study: generalized eigenvalue proximal SVM (GEPSVM) and twin SVM (TSVM), with the aim of augmenting the classification performance further.

**MATERIALS AND METHODS**

**Materials**

The open dataset was downloaded from “Open Access Series of Imaging Studies (OASIS)” (Ardekani, Figarsky & Sidtis, 2013), which consists of 416 subjects aged from eighteen to ninety-six. All subjects are right-handed. Then, we merely selected 126 samples (28 ADs and 98 NCs) from the dataset. The demographic statuses are reported in Table 1. Following common convention, clinical dementia rating (CDR) was interpreted as the target (label). Note that subjects either with missing records or under sixty years old were removed.

**Co-registration and brain-masking**

All three-dimensional MR brain images of each subject were motion-corrected, and coregistered to generate an averaged three-dimensional image, with the aim of increas-
ing signal-to-noise ratio. Afterwards, those three-dimensional images were spatially co-registered to the Talairach space and then were brain-masked.

Figure 1 offers an example of the preprocessing on 3D images with resolution of $1\text{ mm} \times 1\text{ mm} \times 1.25\text{ mm}$. The motion-correction procedure registered the images of three scans, and then generated an image in the original acquisition space with resampling to $1\text{ mm} \times 1\text{ mm} \times 1\text{ mm}$. Finally, the averaged image was normalized to the Talairach coordinate space, and brain-extracted (Fig. 1).

**Key slice selection**

Calculating the displacement field on the whole brain was time-consuming. Therefore, we introduced the “key-slice (KS) selection” method that picks up key slices containing structures indicative of AD from NC. The procedure was as follows: we established a criterion called “inter-class variance (ICV)” $\nu$ as

$$\nu(k) = \|\mu_A(k) - \mu_N(k)\|^2$$

where $k$ is the index of key slice, $\| \cdot \|^2$ represents the $l_2$-norm, $\mu_A$ and $\mu_N$ represents the mean of gray-level values of the $k$th slice of AD subjects and NC subjects, respectively, and $\nu$ denotes the ICV. We select the KSs whose ICVs are larger than half of maximum ICV, with $10\times$ undersampling factor (i.e., every ten slices).

Besides, the direction of key slice can be either sagittal, or axial, or coronal. Scholars found images along coronal direction give a well-defined view compared to those along sagittal and axial directions. Figure 2 shows an image along coronal direction has an advantage because it covers 3 most important AD-related regions, which are treated as indicative of AD. These AD-related regions consist of the cerebral cortex (CC), the HC, and the ventricle. When sagittal or axial directions are used, two or even more slices are needed to cover those tissues. Therefore, we chose the coronal direction for key slice selection, in order to use less slices.

**Shape registration**

The shape registration consists of both global and local registrations. The global registration estimates the rigid parameters, which was accomplished in ‘Co-registration and brain-masking’. Then, the local registration finds the “displacement field (DF)” between the moving and reference images (see Fig. 3).

The rigid registration is an essential preprocessing procedure before non-rigid registration. It can fix the deformation stemming from the position, move, and pose of patients. Then, the following non-rigid registration can reflect the shape deformation of diseases.

The problem of finding the non-rigid estimation can be thought of a motion estimation task between a normal brain (moving) and an AD brain (reference). Several types of solutions are available to solve this task, such as spline function based methods, phase-correlation methods, fluid methods, optical-flow methods, elastic methods, etc. Among the above methods, the first type is parametric, hence, this kind of solution needs
**Figure 1** Preprocessing of a specified subject.
Figure 2  Important regions between (A) an AD brain and (B) a normal brain. i, CC; ii, ventricle; iii, HC. (Pseudocolor enhancement is performed for enlarging contrast.)

Figure 3  Flowchart of displacement field.

to solve optimal spline-based function parameters (Petibon et al., 2013). The second type needs a mass of computation resources; it is difficult to determine the local search range, and it is impossible to guarantee to find global optimal points (Lu, Yang & Zhang, 2014). The other three methods are non-parametric. They find the DF by solving directly a predefined physical model using partial differential equation (PDE) (Yang et al., 2007).

The level-set motion estimation method (Huang et al., 2014; Lee, Sandhu & Tannenbaum, 2013; Vandemeulebroucke et al., 2012) is a rather novel method, which is formed on the basis of the level set evolution theory. The moving image $I_1$ morphs iteratively along its gradient direction, till it deforms close to the given reference image $I_2$. The displacement field is in the form of

$$\frac{dV}{dt} = (I_2 - I_1(V)) \frac{\nabla I_1(V)}{|\nabla I_1(V)|}$$

(2)

where $V$ represents the displacement field, and $I_1(V)$ the deformed image of $I_1$ by $V$.

The above equation can be solved by iterative algorithms described in reference (Kodipaka et al., 2007).
Figure 4 gives an example of the displacement field between a glioma brain and a normal one. Figure 4A presents a moving image while Fig. 4B is a reference image. Their overlap is shown in Fig. 4C. Apparently, the reference image is larger than the moving image, and they are not quite a match. Figure 4D shows the result of rigid-registration, and Fig. 4E shows the two images now match well except the cortex, the temporal and occipital lobe.
Table 2  Pseudocode of the region detection method.

| Region detection                                           |
|-----------------------------------------------------------|
| Step 1  Select a normal brain (I₁) and an AD brain (I₂) from the dataset |
| Step 2  For each key slice k                                |
|         Implement level-set displacement-field estimation between I₁ and I₂, and obtain the displacement field V. |
|         Move the points (x, y), that satisfies |V(x, y)| > 5, to the set R.                                 |
|         End                                                  |
| Step 3  Output R.                                          |

Figure 4F shows the result of non-rigid registration. Compare to Fig. 4B, they closely match each other, and Fig. 4G also suggests a nearly-perfect match. The displacement field is shown in Fig. 4H, with two enlarged areas in Figs. 4I and 4J.

The values of displacement field are complex, i.e., the real part represents the displacement field along the horizontal direction, and the imaginary part along the vertical direction. Those values of Cartesian system are transformed into polar coordinate system, viz., the magnitude-direction spaces. Hence, the inputs to the classifiers are two fold in this study: (1) the directions of the displacement vector of the whole brain and (2) the magnitude of the displacement field of the whole brain. We did not use the sign of the displacement vector, which were commonly used in literature (Das Gupta & Tamadapu, 2013; Kodipaka et al., 2007). The reason is the direction field already contains the information of sign of the displacement.

Method of region detection

Here, we proposed a visual interpretation approach based on displacement field to detect regions R that can distinguish AD and NC.

\[ R = \{ (x, y) | |V(x, y)| > T \} \]  

(3)

Here \( | \cdot | \) represents the magnitude, \( V(x, y) \) represents the displacement field at point of \( (x, y) \), \( T \) the threshold. From another point of view, Eq. (3) means we preserve the points whose displacement magnitude is larger than \( T \). In this work, \( T \) is assigned with a value of 5, which is defined empirically. A smaller threshold \( T \) may introduce more noises in the estimated displacement field; whereas, a larger threshold \( T \) will drop realistic deformation with short deformation magnitude. Hence, we believe the deformation with magnitude larger than 5 represents realistic deformation in the brain. In all, the visual interpretation is a three-step process (see Table 2).

Non-parallel support vector machine

Among all recent classifiers, the support vector machine (SVM) has gained popularity as the most excellent classifiers in small-size problem (Zhang & Wu, 2012). To further enhance the classification performance, two new variants of SVM were introduced:
**Generalized eigenvalue proximal SVM**

Original SVM has a limitation that two hyperplanes should be parallel (Wang et al., 2014). Mangasarian & Wild (2006) designed a generalized eigenvalue proximal SVM (GEPSVM). It drops the parallelism restrain on the two planes (remember the parallelism is necessary in original SVM), and requires each hyperplane should be as close as possible to one of the data sets and as far as possible from the other. The latest literature shows that GEPSVM yielded superior performance to canonical support vector machines in terms of sensitivity, specificity, precision, and accuracy (Khemchandani, Karpatne & Chandra, 2011; Shao et al., 2013).

Suppose samples are from either class 1 (denote by symbol $X_1$) or class 2 (denoted by symbol $X_2$), respectively. The GEPSVM finds the two optimal nonparallel planes with the form of ($w$ and $b$ denotes the weight and bias of the classifier, respectively)

$$w_1^T x - b_1 = 0 \quad \text{and} \quad w_2^T x - b_2 = 0. \quad (4)$$

To obtain the first plane, we deduce from Eq. (4) and get the following solution

$$(w_1, b_1) = \arg \min_{(w, b) \neq 0} \frac{\|w^T X_1 - o^T b\|^2}{\|w^T X_2 - o^T b\|^2} \quad (5)$$

$$U \leftarrow \begin{bmatrix} w \\ b \end{bmatrix} \quad (6)$$

where $o$ is a vector of ones with any possible dimension according to the context. Simplifying formula (5) gives

$$\min_{(w, b) \neq 0} \frac{\|w^T X_1 - o^T b\|^2}{\|w^T X_2 - o^T b\|^2}. \quad (7)$$

We include the Tikhonov regularization term to decrease the norm of the variables $U$ that corresponds to the first hyperplane in (4).

$$\min_{(w, b) \neq 0} \frac{\|w^T X_1 - o^T b\|^2 + t\|U\|^2}{\|w^T X_2 - o^T b\|^2} \quad (8)$$

where $t$ is a positive (or zero) Tikhonov factor. Formula (8) turns to the “Rayleigh Quotient (RQ)” in the following form of

$$U_1 = \arg \min_{U \neq 0} \frac{U^T P U}{U^T Q U} \quad (9)$$

where $P$ and $Q$ are symmetric matrices in size of $(p + 1) \times (p + 1)$

$$P \overset{\text{def}}{=} \begin{bmatrix} X_1 & -o \\ -o^T & X_1 \end{bmatrix} - o + tl \quad (10)$$

$$Q \overset{\text{def}}{=} \begin{bmatrix} X_2 & -o \\ -o^T & X_2 \end{bmatrix}. \quad (11)$$
Considering the stationarity and boundedness characteristics of RQ, the answer of (9) is deduced by figuring out a generalized eigenvalue problem (GEP) as

\[ PU = \lambda QU, \quad U \neq 0 \]  

(12)

here the global optimal result of (9) can be obtained at an eigenvector \( U_1 \) associated to the smallest eigenvalue \( \lambda_{\text{min}} \) of formula (12). Therefore, \( w_1 \) and \( b_1 \) can be deduced from formula (6), and utilized to from the plane as written in formula (4). Afterwards, a similar optimization problem is generated that is analogous to (7) by exchanging the symbols of \( X_1 \) and \( X_2 \). The eigenvector \( U_2^* \) corresponding to the smallest eigenvalue of the 2nd GEP will obtain the 2nd hyperplane approximate to samples of class 2.

**Twin support vector machine**

Jayadeva & Chandra (2007) provided a novel variant of standard SVM: the twin support vector machine (TSVM). The TSVM is similar to GEPSVM in the way that both obtain non-parallel hyperplanes. The difference lies in that GEPSVM and TSVM are formulated entirely differently. Each of the two quadratic programming (QP) problems in TSVM pair is formulated as a typical SVM. Reports have shown that TSVM is better than both SVM and GEPSVM (Nasiri, Charkari & Mozafari, 2014; Shao et al., 2014; Xu, Qi & Zhang, 2014). Mathematically, the TSVM is constructed by solving the two QP problems

\[
\min \frac{1}{2} (X_1 w_1 + o_1 b_1)^T (X_1 w_1 + o_1 b_1) + c_1 o_2^T q \\
\text{s.t. } -(X_2 w_1 + o_2 b_1) + q \geq o_2, q \geq 0
\]  

(13)

\[
\min \frac{1}{2} (X_2 w_2 + o_2 b_2)^T (X_2 w_2 + o_2 b_2) + c_2 o_1^T q \\
\text{s.t. } -(X_1 w_2 + o_1 b_2) + q \geq o_1, q \geq 0
\]  

(14)

here \( c_i \) \( (i = 1, 2) \) are positive parameters, and \( o_i \) \( (i = 1, 2) \) is the same as in formula (5). By this means, the TSVM constructed two hyperplanes. The first term in equations of (13)(14) is the sum of squared distances from the hyperplane to one class. The second term is the sum of error variables. Therefore, minimizing Eqs. (13) and (14) will force the hyperplanes approximate to data of each class, and minimize the misclassification rate. Finally, the constraint requires the plane to be at a distance of more than 1 from data in the other class. Another advantage of TSVM is that its convergence rate is four times faster than conventional SVM (Jayadeva & Chandra, 2007).

**Statistical analysis**

Generalization error was obtained by K-fold CV, and K was set to 10 because of two reasons: (1) to balance between computational cost and reliable estimates, and (2) for fair comparison since the common convention is to set K with the value of 10 (Zhang et al., 2014a).

For a 10-fold CV, the dataset is randomly divided into 10 mutually exclusively folds of nearly equal size. In each run, 9 subsets are used for training, and the rest for validation (see
Evaluation indicators.

| Indicator      | Explanation                  |
|----------------|------------------------------|
| TP             | True Positive                |
| FP             | False Positive               |
| TN             | True Negative                |
| FN             | False Negative               |
| Sensitivity (recall) | TP/(FN + TP)                  |
| Specificity    | TN/(FP + TN)                 |
| Accuracy       | (TN + TP)/(FN + FP + TN + TP) |
| Precision      | TP/(TP + FP)                 |

Fig. 5). Above procedure repeats 10 runs, such that each subset is used once for validation. The 10 results over validation set are combined together along the diagonal blocks in Fig. 5, with the aim of producing an individual out-of-sample evaluation. The 10-fold cross validation repeated 50 times, viz., a $50 \times 10$-fold cross validation was implemented.

Evaluation

We used four indicators to measure which algorithm performed the best. Those four indicators consisted of sensitivity (recall), specificity, accuracy, and precision (Table 3). In this work, a correctly detected AD was treated as TP. After the $50 \times 10$-fold cross validation, the final evaluation results are written in the form of “mean ± standard deviation (SD)”.

The whole proposed system

Remember that our aim contains two purposes. First, we need to develop a computer-aided diagnosis (CAD) system and report its performance. Second, we need to locate


**Table 4** Pseudocode of proposed method.

| Algorithm: proposed method |
|----------------------------|
| **Step A** | Input the ground-truth imaging data together with their labels. |
| **Step B** | Co-registration to Talairach Coordinate by Rigid Registration. |
| **Step C** | Pick up the key-slices by ICV (more than half of maximum), with $10 \times$ undersampling factor. |
| **Step D** | Produce displacement field for each key slice for each subject. |
| **Step E** | Submit the displacement field to the classifiers. |
| **Step F** | Report the classification performance based on a $50 \times 10$-fold cross validation. |
| **Step G** | Report the AD-related regions with the points whose displacement magnitude is larger than five. |

discriminant voxels that can detect AD from NC subjects; and (ii) The pseudocode is listed in Table 4.

**EXPERIMENTS AND RESULTS**

The programs were developed in-house using MATLAB (Version 2015a; Natick, MA), and run on an IBM laptop with 3 GHz Intel i3 dual-processor and 8 GB random access memory (RAM).

**Key-slice selection**

The curve of “inter-class variance” versus coronal slice index is shown in Fig. 6A. Ten coronal slices, from 60 to 150 with increasing steps of 10, were selected. The reason is that their corresponding ICVs were all higher than half of the maximum value. Figures 6B–6C shows respectively the axial and sagittal view of the selected key-slices, where the red lines represent the key-slices. The results are coherent with our formal work (Dong et al., 2015b).

**Displacement field**

Figure 7 shows the displacement field of an AD over the key-slices. Each column corresponds to a key slice. Here CI means coronal index. We show the moving image, the reference image, the overlap of both, the rigid registration, the overlap of rigid registration and the reference image, the non-rigid registration, the overlap of non-rigid registration and the reference image, and the displacement field for each key slice. From the row of displacement field, we can see how that a healthy brain is warped into an AD brain through the displacement fields (green arrows in the bottom row).

**Classification comparison**

The displacement fields were transformed into polar coordinate system, and their number was reduced by PCA as mentioned in the methodology section. Three classifiers were used: SVM, GEPSVM, and TSVM. Classification comparison results are provided in Table 5, together with the results of ten state-of-the-art methods.

**Region detection**

We implemented the AD-related region detection procedure to different AD subjects as ‘Method of region detection’ described. Figure 8 shows the related regions, which are labelled by green points. Some areas are slightly outside of the brain, such as Fig. 8A, this
Figure 6 Results of key-slice selection. (A) Curve of ICV against CI. (B) Key-slices from axial view. (C) Key-slices from saggital view.

is because the algorithm may also consider the distortion of background. This does not influence following labelling procedure.

Area labeling
Talairach Daemon software was used to provide the anatomical label information. Table 6 shows the results, where BA represented Brodmann area. Note that some areas (such as cerebral tonsil, declive, and pyramis) move due to the expansion and shrinkage of neighboring areas, so they are not included in Table 6. We only pick up the areas with changed volume, which are judged manually.

DISCUSSION
The results in Table 5 compare the proposed three classifiers (SVM, GEPSVM, and TSVM) with ten state-of-the-art methods. Plant’s results (Task 1 in Table 3 in Plant et al., 2010) presented the means together with 95% confidence intervals. Wang’s results (Table 7 in Wang et al., 2014) were obtained through a single K-fold cross validation analysis. Savio’s
CI varies from 60 to 150 with increase of 10. (Please zoom in to see the displacement field.)

(Table 5 in Savio & Grana, 2013) and Dong’s results (Table 9 in Dong et al., 2015b) gave the means with SD.

Among the proposed methods, the proposed “DF + PCA + TSVM” yields the accuracy of $92.75 \pm 1.77$, sensitivity of $90.56 \pm 1.15$, specificity of $93.37 \pm 2.05$, and precision of $79.61 \pm 2.21$. Additional to it, the proposed “DF + PCA + GEPSVM” offers the accuracy of $91.52 \pm 1.63$, sensitivity of $88.93 \pm 1.80$, specificity of $92.27 \pm 1.79$, and precision of $76.66 \pm 2.33$. The proposed “DF + PCA + SVM” obtains the accuracy of $88.27 \pm 1.89$, sensitivity of $84.93 \pm 1.21$, specificity of $89.21 \pm 1.63$, and precision of $69.30 \pm 1.91$.
In terms of average accuracy, the proposed “DF + PCA + TSVM” result is as large as 92.75%, better than nine approaches of AD prediction, e.g., BRC + IG + SVM of 90.00% (Plant et al., 2010), BRC + IG + Bayes of 92.00% (Plant et al., 2010), BRC + IG + VFI of 78.00% (Plant et al., 2010), MGM + PEC + SVM of 92.07% (Savio & Grana, 2013), GEODAN + BD + SVM of 92.09% (Savio & Grana, 2013), US + SVD-PCA + SVM-DT of 90% (Wang et al., 2014), EB + WTT + SVM of 91.47% (Dong et al., 2015b), EB + WTT + RBF-KSVM of 86.71% (Dong et al., 2015b), and EB + WTT + POL-KSVM of 92.36% (Dong et al., 2015b). Nevertheless, the average accuracy of the method
Table 5 Comparison of different methods.

| Existing methods                     | Accuracy | Sensitivity | Specificity | Precision |
|--------------------------------------|----------|-------------|-------------|-----------|
| BRC + IG + SVM (Plant et al., 2010)  | 90.00    | 96.88       | 77.78       | N/A       |
| BRC + IG + Bayes (Plant et al., 2010)| 92.00    | 93.75       | 88.89       | N/A       |
| BRC + IG + VFI (Plant et al., 2010) | 78.00    | 65.63       | 100.00      | N/A       |
| MGM + PEC + SVM (Savio & Grana, 2013)| 92.07± 1.12 | 86.67± 4.71 | N/A         | 95.83± 5.89 |
| GEODAN + BD + SVM (Savio & Grana, 2013)| 92.09± 2.60 | 80.00± 4.00 | N/A         | 88.09± 5.33 |
| TJM + WTT + SVM (Savio & Grana, 2013)| 92.83± 0.91 | 86.33± 3.73 | N/A         | 85.62± 0.85 |
| US + SVD-PCA + SVM-DT (Wang et al., 2014)| 90      | 94          | 71          | N/A       |
| EB + WTT + SVM (Dong et al., 2015b)  | 91.47± 1.02 | 90.17± 1.66 | 91.84± 1.09 | 75.93± 2.43 |
| EB + WTT + RBF-KSVM (Dong et al., 2015b) | 86.71± 1.93 | 85.71± 1.91 | 86.99± 2.30 | 66.12± 4.16 |
| EB + WTT + POL-KSVM (Dong et al., 2015b) | 92.36± 0.94 | 83.48± 3.27 | 94.90± 1.09 | 82.28± 2.78 |

| Proposed methods                        | Accuracy | Sensitivity | Specificity | Precision |
|------------------------------------------|----------|-------------|-------------|-----------|
| DF + PCA + SVM                          | 88.27± 1.89 | 84.93± 1.21 | 89.21± 1.63 | 69.30± 1.91 |
| DF + PCA + GEPSVM                       | 91.52± 1.63 | 88.93± 1.80 | 92.27± 1.79 | 76.66± 2.33 |
| DF + PCA + TSVM                         | 92.75± 1.77 | 90.56± 1.15 | 93.37± 2.05 | 79.61± 2.21 |

“DF + PCA + TSVM” is only less than one approach of “TJM + WTT + SVM” of 92.83% (Savio & Grana, 2013).

Why TSVM is better? There are two reasons. First, the non-parallel support vector machines provide more flexible and complicated hyperplanes than standard support vector machine. Second, the twin support vector machines formulate each of the two QP problems as a standard support vector machine, which makes it superior to generalized eigenvalue proximal support vector machine.

There were many other methods (Arbizu et al., 2013; Chaves et al., 2013; Cohen & Klunk, 2014; Dukart et al., 2013; Gray et al., 2012) proposed for detecting AD from NC, however, they dealt with images produced by other types of modalities: PET, SPECT, DTI, etc. Hence, it is inappropriate to compare the proposed methods with them. We will test our methods on SPECT and PET images in the future.
Table 6 shows that the displacement field finds the discriminant associated with the following regions reported in latest references: Angular Gyrus (Carbonell et al., 2014), Anterior Cingulate (Schultz et al., 2014), Cingulate Gyrus (Yu et al., 2014), Culmen (Almeida et al., 2011), Cuneus (Yokoyama et al., 2015), Fusiform Gyrus (Rieck et al., 2015), Inferior Frontal Gyrus (Eliaieova et al., 2014), Inferior Occipital Gyrus (Liu et al., 2012), Inferior Parietal Lobule (Tramutola et al., 2015), Inferior Semi-Lunar Lobule (Almeida et al., 2011), Inferior Temporal Gyrus (Narayan et al., 2015), Insula (Fletcher et al., 2015), Lateral Ventricle (Gonzalez-Marrero et al., 2015), Lingual Gyrus (Cooley et al., 2015), Medial Frontal Gyrus (Jensen et al., 2013), Middle Frontal Gyrus (Yoo et al., 2015), Middle Occipital Gyrus (Cai et al., 2015), Middle Temporal Gyrus (Han et al., 2015), Paracentral Lobule (Kang et al., 2013), Parahippocampal Gyrus (Shimoda et al., 2015), Postcentral Gyrus (Willette et al., 2015), Posterior Cingulate (Pekarya, Sattin & Lloyd, 2015), Precentral Gyrus (Wang et al., 2015c), Precuneus (Villeneuve et al., 2015), Sub-Gyrals (Ouyang et al., 2014), Superior Parietal Lobule (Yamashita et al., 2014), Superior Temporal Gyrus (Ramos et al., 2015), Supramarginal Gyrus (Redolfi et al., 2015), Uncus (Bangen et al., 2014).

Notwithstanding, some regions reported to be associated with AD are not interpreted by displacement field. Those areas include caudate nucleus (Montagne et al., 2015), claustrum (Pirone et al., 2015), lentiform nucleus (Dong et al., 2015b), subcallosal gyrus (Dong et al., 2015b), caudate nucleus (Montagne et al., 2015), claustrum (Pirone et al., 2015), lentiform nucleus (Dong et al., 2015b), subcallosal gyrus (Dong et al., 2015b).
Table 6  Discriminant areas with changed volume found by Talairach Daemon software.

| Regions                                | # of voxels | Reported by               |
|----------------------------------------|-------------|---------------------------|
| Angular gyrus                          | 33          | Carbonell et al. (2014)   |
| Anterior cingulate (BA-33, BA-32, BA-24)| 81          | Schultz et al. (2014)     |
| Cingulate gyrus (BA-32, BA-23, BA-24, BA-31) | 1,551       | Yu et al. (2014)          |
| Culmen                                  | 396         | Almeida et al. (2011)     |
| Cuneus (BA-18, BA-30)                   | 143         | Yokoyama et al. (2015)    |
| Fusiform gyrus (BA-18, BA-19, BA-20, BA-37) | 314         | Rieck et al. (2015)       |
| Inferior frontal gyrus (BA-13, BA-45, BA-47) | 320         | Eliasova et al. (2014)    |
| Inferior occipital gyrus                | 24          | Liu et al. (2012)         |
| Inferior parietal lobule (BA-2, BA-40)  | 311         | Tramutola et al. (2015)   |
| Inferior semi-lunar lobule              | 144         | Almeida et al. (2011)     |
| Inferior temporal gyrus (BA-20)         | 58          | Narayan et al. (2015)     |
| Insula (BA-44, BA-13)                   | 328         | Fletcher et al. (2015)    |
| Lateral ventricle                       | 33          | Gonzalez-Marrero et al. (2015) |
| Lingual gyrus (BA-18, BA-19)            | 184         | Cooley et al. (2015)      |
| Medial frontal gyrus (BA-6, BA-32)      | 53          | Jensen et al. (2015)      |
| Middle frontal gyrus (BA-6, BA-46)      | 144         | Yoo et al. (2015)         |
| Middle occipital gyrus (BA-19)          | 175         | Cai et al. (2015)         |
| Middle temporal gyrus (BA-19, BA-20, BA-21, BA-22, BA-37, BA-38, BA-39) | 485         | Han et al. (2015)         |
| Paracentral lobule (BA-5, BA-31)        | 161         | Kang et al. (2013)        |
| Parahippocampal gyrus (HC, BA-19, BA-30, BA-37) | 62          | Shimoda et al. (2015)     |
| Postcentral gyrus (BA-2, BA-3)          | 244         | Willette et al. (2015)    |
| Posterior cingulate (BA-23, BA-30)      | 323         | Pekarya, Satin & Lloyd (2015) |
| Precentral gyrus (BA-4, BA-6, BA-13, BA-43, BA-44) | 627         | Wang et al. (2015c)       |
| Precuneus (BA-7, BA-19, BA-31)          | 530         | Villeneuve et al. (2015)  |
| Sub-Gyral (Corpus, Callosum, BA-4, BA-13) | 2,358       | Ouyang et al. (2014)      |
| Superior parietal lobule                | 21          | Yamashita et al. (2014)   |
| Superior temporal gyrus (BA-13, BA-22, BA-38, BA-39, BA-41) | 462         | Ramos et al. (2015)       |
| Supramarginal gyrus (BA-40)             | 135         | Redolfi et al. (2015)     |
| Uncus (BA-20, BA-28, BA-34, BA-36, BA-38) | 112         | Bangen et al. (2014)      |

2015b), and subthalamic nucleus (De Reuck et al., 2014). Why are those areas not detected by our DF method? The reasons are tri-fold: (1) We only preserved the displacement field with magnitude longer than 5. Reducing the value of 5 may include more potential regions and noises. A feasible solution is to reduce from 5 to 3 and meanwhile develop robust anti-noise method. (2) Some literature used advanced imaging modalities, such as MRSI and fMRI for metabolism detection and function analysis. Therefore, we may also include these imaging-modality techniques. (3) The key-slice selection procedure may miss important regions. Hence, we will try to reduce the slice separation, although it will increase the computation burden.

The advantages of displacement-field are two-fold. On one hand, it reaches excellent classification performance, which was comparable to latest approaches (Dong et al., 2015b; Plant et al., 2010; Savio & Grana, 2013; Wang et al., 2014). On the other hand, it can directly
locate AD-related regions. The disadvantages of displacement field are: (i) The accuracy of displacement field estimation relies on the accuracy of rigid registration, which is a necessary pre-processing (see Fig. 3). (ii) The estimated DF may exist in the background (see Fig. 8A), which needs to be removed by masking technique.

| Acronym | Definition |
|---------|------------|
| (k) (GEP) (T) SVM | (kernel) (generalized eigenvalue problem) (twin) support vector machine |
| AD | Alzheimer’s disease |
| ANN | Artificial neural network |
| BA | Brodmann area |
| BD | Bhattacharyya distance |
| BRC | Brain region cluster |
| CAD | Computer-aided diagnosis |
| CC | Cerebral cortex |
| CDR | Clinical dementia rating |
| CSF | Cerebrospinal fluid |
| CI | Coronal index |
| CV | Cross validation |
| DBM | Deformation-based morphometry |
| DF | Displacement field |
| DWT | Discrete wavelet transform |
| GARCH | Generalized autoregressive conditional heteroscedasticity |
| GEODAN | Geodesic anisotropy |
| HC | Hippocampus |
| ICV | Inter-Class variance |
| IG | Information gain |
| KNN | K-nearest neighbors |
| MGM | Modulated GM |
| MMSE | Mini-mental state examination |
| MR(I) | Magnetic resonance (imaging) |
| NBC | Naive Bayes classifier |
| NC | Normal elder controls |
| OASIS | Open access series of imaging studies |
| PEC | Pearson's correlation |
| PNN | Probabilistic neural network |
| PSO | Particle swarm optimization |
| QP | Quadratic programming |
| ROI | Region of interest |
| RQ | Rayleigh quotient |
| SD | Standard deviation |
| SVD | Singular value decomposition |
| TJM | Trace of Jacobian matrix |
| US | Undersampling |
| VFI | Voting feature intervals |
| WTT | Welch’s t-test |
CONCLUSIONS

The contributions of this study consist of four points: (i) We proposed to use displacement field in the application of Alzheimer’s disease detection, and we proved its effectiveness. (ii) SVM and its two variants (GEPSVM and TSVM) were tested, and we proved TSVM performed better than SVM and GEPSVM. (iii) The proposed system achieved comparable sensitivity, specificity, and accuracy with ten state-of-the-art algorithms. (iv) The proposed computer-aided diagnosis (CAD) system can locate AD-related regions in the brain, which complies with 28 recent peer-reviewed articles.

Future research direction shall center in following aspects. First, the displacement field will be generalized to 3D, so key-slice selection can be discarded. Second, we shall try to embed kernels to classifiers. Third, spectroscopy method, like extracting the quantity of Docosahexaenoic Acid (DHA) (Tan et al.), may help to increase the classification accuracy. Forth, the proposed approach might be helpful in identifying brain mechanisms underlying endogenously defensive mechanism to neuroinjury and neurodegeneration (Yuan & Hou, 2015). Fifth, the least square technique (Hamedi, Salleh & Noor, 2015) may be applied to SVM and its variants, in order to further reduce training time. Sixth, the swarm-intelligence algorithm will be introduced to help to enhance the algorithm performance (Wang et al., 2015b). Finally, the stress field calculates the stress at every point within a particular region (Petit et al., 2015). It will be tested in future research.

ADDITIONAL INFORMATION AND DECLARATIONS

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Competing Interests
The authors declare there are no competing interests.

Author Contributions
• Yudong Zhang conceived and designed the experiments, analyzed the data, wrote the paper, prepared figures and/or tables, reviewed drafts of the paper.
• Shuihua Wang performed the experiments, contributed reagents/materials/analysis tools, wrote the paper.

Data Availability
The following information was supplied regarding data availability:
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REFERENCES
Almeida OP, Garrido GJ, Alfonso H, Hulse G, Lautenschlager NT, Hankey GJ, Flicker L. 2011. 24-month effect of smoking cessation on cognitive function and brain structure in later life. *NeuroImage* 55:1480–1489 DOI 10.1016/j.neuroimage.2011.01.063.
Álvarez I, Gorriz JM, Ramírez J, Salas-Gonzalez D, López M, Segovia F, Puntonet CG, Prieto B. 2009. Alzheimer’s diagnosis using eigenbrains and support vector machines. In: Cabestany J, Sandoval F, Prieto A, Corchado J, eds. *Bio-inspired systems: computational and ambient intelligence*. Berlin Heidelberg: Springer, 973–980.
Anagnostopoulos CN, Giannoukos I, Spenger C, Simmons A, Mecocci P, Soininen H, Kloszewska I, Vellas B, Lovestone S, Tsolaki M. 2013. Classification models for Alzheimer’s disease detection. In: Iliadis L, Papadopoulos H, Jayne C, eds. *Engineering applications of neural networks, Pt II*, Halkidiki, 193–202.
Arbizu J, Prieto E, Martínez-Lage P, Martí-Climent JM, García-Granero M, Lamet I, Pastor P, Riverol M, Gómez-Isla MT, Peñuelas I, Richter JA, Weiner MW. 2013. Automated analysis
of FDG PET as a tool for single-subject probabilistic prediction and detection of Alzheimer’s disease dementia. *European Journal of Nuclear Medicine and Molecular Imaging* **40**:1394–1405 DOI 10.1007/s00259-013-2458-z.

Ardekani BA, Figarsky K, Sidtis JJ. 2013. Sexual dimorphism in the human corpus callosum: an mri study using the OASIS brain database. *Cerebral Cortex* **23**:2514–2520 DOI 10.1093/cercor/bhs253.

Bangen KJ, Nation DA, Clark LR, Harmell AL, Wierenga CE, Dev SI, Delano-Wood L, Zlatar ZZ, Salmon DP, Liu TT, Bondi MW. 2014. Interactive effects of vascular risk burden and advanced age on cerebral blood flow. *Frontiers in Aging Neuroscience* **6**:Article 00159 DOI 10.3389/fnagi.2014.00159.

Brookmeyer R, Johnson E, Ziegler-Graham K, Arrighi HM. 2007. Forecasting the global burden of Alzheimer’s disease. *Alzheimer’s & Dementia* **3**:186–191 DOI 10.1016/j.jalz.2007.04.381.

Cai SP, Huang LY, Zou J, Jing LL, Zhai BZ, Ji GJ, Von Deneen KM, Ren JC, Ren AF, Alzheimer’s Disease Neuroimaging Initiative. 2015. Changes in thalamic connectivity in the early and late stages of amnestic mild cognitive impairment: a resting-state functional magnetic resonance study from ADNI. *PLoS ONE* **10**:e0115573 DOI 10.1371/journal.pone.0115573.

Carbonell F, Charil A, Zijdenbos AP, Evans AC, Bedell BJ, Alzheimer’s Disease Neuroimaging Initiative. 2014. Hierarchical multivariate covariance analysis of metabolic connectivity. *Journal of Cerebral Blood Flow and Metabolism* **34**:1936–1943 DOI 10.1038/jcbfm.2014.165.

Chaves R, Ramirez J, Gorriz JM, Alzheimer’s Disease Neuroimaging Initiative. 2013. Integrating discretization and association rule-based classification for Alzheimer’s disease diagnosis. *Expert Systems with Applications* **40**:1571–1578 DOI 10.1016/j.eswa.2012.09.003.

Chaves R, Ramirez J, Gorriz JM, Illan IA, Gomez-Rio M, Carnero C, Alzheimer’s Disease Neuroimaging Initiative. 2012. Effective diagnosis of Alzheimer’s disease by means of large margin-based methodology. *BMC Medical Informatics and Decision Making* **12**:17 DOI 10.1186/1472-6947-12-79.

Cohen AD, Klunk WE. 2014. Early detection of Alzheimer’s disease using PiB and FDG PET. *Neurobiology of Disease* **72**:117–122 DOI 10.1016/j.nbd.2014.05.001.

Collins MP, Pape SE. 2011. The potential of support vector machine as the diagnostic tool for schizophrenia: a systematic literature review of neuroimaging studies. *European Psychiatry* **26** (Supplement 1):P03–P194 DOI 10.1016/S0924-9338(11)73068-1.

Cooley SA, Cabeen RP, Laidlaw DH, Conturo TE, Lane EM, Heaps JM, Bolzenius JD, Baker LM, Salminen LE, Scott SE, Paul RH. 2015. Posterior brain white matter abnormalities in older adults with probable mild cognitive impairment. *Journal of Clinical and Experimental Neuropsychology* **37**:61–69 DOI 10.1080/13803395.2014.985636.

Damodharan S, Raghavan D. 2015. Combining tissue segmentation and neural network for brain tumor detection. *International Arab Journal of Information Technology* **12**:42–52.

Das Gupta A, Tamadapu G. 2013. In-plane dynamics of membranes having constant curvature. *European Journal of Mechanics—A/Solids* **39**:280–290 DOI 10.1016/j.euromechsol.2012.12.010.

De Reuck JL, Deramecourt V, Auger F, Durieux N, Cordonnier C, Devos D, Defebvre L, Moreau C, Caparros-Lefebvre D, Leys D, Maurage CA, Pasquier F, Bordet R. 2014. Iron deposits in post-mortem brains of patients with neurodegenerative and cerebrovascular diseases: a semi-quantitative 7.0 T magnetic resonance imaging study. *European Journal of Neurology* **21**:1026–1031 DOI 10.1111/ene.12432.

Dong Z, Liu A, Wang S, Ji G, Zhang Z, Yang J, Zhang Y. 2015a. Magnetic resonance brain image classification via stationary wavelet transform and generalized eigenvalue proximal support vector machine. *Journal of Medical Imaging and Health Informatics* **5**:1–9 DOI 10.1166/jmihi.2015.1357.
Dong Z, Phillips P, Wang S, Ji G, Yang J, Yuan T-F. 2015b. Detection of subjects and brain regions related to Alzheimer’s disease using 3D MRI scans based on eigenbrain and machine learning. Frontiers in Computational Neuroscience 66:1–15.

Dukart J, Mueller K, Barthel H, Villringer A, Sabri O, Schroeter ML, Alzheimer’s Disease Neuroimaging Initiative. 2013. Meta-analysis based SVM classification enables accurate detection of Alzheimer’s disease across different clinical centers using FDG-PET and MRI. Psychiatry Research-Neuroimaging 212:230–236 DOI 10.1016/j.pscychresns.2012.04.007.

El-Dahshan ESA, Hosny T, Salem ABM. 2010. Hybrid intelligent techniques for MRI brain images classification. Digital Signal Processing 20:433–441 DOI 10.1016/j.dsp.2009.07.002.

El-Dahshan ESA, Mohsen HM, Revett K, Salem ABM. 2014. Computer-aided diagnosis of human brain tumor through MRI: a survey and a new algorithm. Expert Systems with Applications 41:5526–5545 DOI 10.1016/j.eswa.2014.01.021.

Eliasova I, Anderkova L, Marecek R, Rektorova I. 2014. Non-invasive brain stimulation of the right inferior frontal gyrus may improve attention in early Alzheimer’s disease: a pilot study. Journal of the Neurological Sciences 346:318–322 DOI 10.1016/j.jns.2014.08.036.

Esposito R, Mosca A, Pieramico V, Cieri F, Cera N, Sensi SL. 2013. Characterization of resting state activity in MCI individuals. PeerJ 1:e135 DOI 10.7717/peerj.135.

Fletcher PD, Downey LE, Golden HL, Clark CN, Slattery CF, Paterson RW, Schott JM, Rohrer JD, Rossor MN, Warren JD. 2015. Auditory hedonic phenotypes in dementia: a behavioural and neuroanatomical analysis. Cortex 67:95–105 DOI 10.1016/j.cortex.2015.03.021.

Goh S, Dong Z, Zhang Y, DiMauro S, Peterson BS. 2014. Mitochondrial dysfunction as a neurobiological subtype of autism spectrum disorder: evidence from brain imaging. JAMA Psychiatry 71:665–671 DOI 10.1001/jamapsychiatry.2014.179.

Gonzalez-Marrero I, Gimenez-Llort L, Johanson CE, Carmona-Calero EM, Castaneyra-Ruiz L, Brito-Armas JM, Castaneyra-Perdomo A, Castro-Fuentes R. 2015. Choroid plexus dysfunction impairs beta-amyloid clearance in a triple transgenic mouse model of Alzheimer’s disease. Frontiers in Cellular Neuroscience 9:Article 00017 DOI 10.3389/fncel.2015.00017.

Gray KR, Wolz R, Heckemann RA, Aljabar P, Hammers A, Rueckert D, Alzheimer’s Disease Neuroimaging Initiative. 2012. Multi-region analysis of longitudinal FDG-PET for the classification of Alzheimer’s disease. NeuroImage 60:221–229 DOI 10.1016/j.neuroimage.2011.12.071.

Hahn K, Myers N, Prigarin S, Rodenacker K, Kurz A, Förstl H, Zimmer C, Wohlschläger AM, Sorg C. 2013. Selectively and progressively disrupted structural connectivity of functional brain networks in Alzheimer’s disease—revealed by a novel framework to analyze edge distributions of networks detecting disruptions with strong statistical evidence. NeuroImage 81:96–109 DOI 10.1016/j.neuroimage.2013.05.011.

Hamedi M, Salleh SH, Noor AM. 2015. Facial neuromuscular signal classification by means of least square support vector machine for MuCI. Applied Soft Computing 30:83–93 DOI 10.1016/j.asoc.2015.01.034.

Han PC, Caselli RJ, Baxter L, Serrano G, Yin JX, Beach TG, Reiman EM, Shi J. 2015. Association of pituitary adenylate cyclase-activating polypeptide with cognitive decline in mild cognitive impairment due to Alzheimer disease. JAMA Neurology 72:333–339 DOI 10.1001/jamaneurol.2014.3625.

Harikumar R, Kumar BV. 2015. Performance analysis of neural networks for classification of medical images with wavelets as a feature extractor. International Journal of Imaging Systems and Technology 25:33–40 DOI 10.1002/ima.22118.
Huang XJ, Dione DP, Compas CB, Papademetris X, Lin BA, Bregasi A, Sinusas AJ, Staib LH, Duncan JS. 2014. Contour tracking in echocardiographic sequences via sparse representation and dictionary learning. *Medical Image Analysis* 18:253–271 DOI 10.1016/j.media.2013.10.012.

Jayadeva KR, Chandra S. 2007. Twin support vector machines for pattern classification. *IEEE Transactions on Pattern Analysis and Machine Intelligence* 29:905–910 DOI 10.1109/TPAMI.2007.1068.

Jensen MM, Arvaniti M, Mikkelsen JD, Michalski D, Pinborg LH, Hartig W, Thomsen MS. 2015. Prostate stem cell antigen interacts with nicotinic acetylcholine receptors and is affected in Alzheimer’s disease. *Neurobiology of Aging* 36:1629–1638 DOI 10.1016/j.neurobiolaging.2015.01.001.

Kalbkhani H, Shayesteh MG, Zali-Vargahan B. 2013. Robust algorithm for brain magnetic resonance image (MRI) classification based on GARCH variances series. *Biomedical Signal Processing and Control* 8:909–919 DOI 10.1016/j.bspc.2013.09.001.

Kang K, Yoon U, Lee JM, Lee HW. 2013. Idiopathic normal-pressure hydrocephalus, cortical thinning, and the cerebrospinal fluid tap test. *Journal of the Neurological Sciences* 334:55–62 DOI 10.1016/j.jns.2013.07.014.

Kantanen M, Kiuru-Enari S, Salonen O, Kaipainen M, Hokkanen L. 2014. Subtle neuropsychiatric and neurocognitive changes in hereditary gelsolin amyloidosis (AGel amyloidosis). *PeerJ* 2:e493 DOI 10.7717/peerj.493.

Khemchandani R, Karpatne A, Chandra S. 2011. Generalized eigenvalue proximal support vector regressor. *Expert Systems with Applications* 38:13136–13142 DOI 10.1016/j.eswa.2011.04.121.

Kodipaka S, Vemuri BC, Rangarajan A, Leonard CM, Schmallfuss I, Eisenschchenk S. 2007. Kernel Fisher discriminant for shape-based classification in epilepsy. *Medical Image Analysis* 11:79–90 DOI 10.1016/j.media.2006.10.002.

Kubota T, Ushijima Y, Nishimura T. 2006. A region-of-interest (ROI) template for three-dimensional stereotactic surface projection (3D-SSP) images: initial application to analysis of Alzheimer disease and mild cognitive impairment. *International Congress Series* 1290:128–134 DOI 10.1016/j.ics.2005.11.104.

Lee W, Park B, Han K. 2013. Classification of diffusion tensor images for the early detection of Alzheimer’s disease. *Computers in Biology and Medicine* 43:1313–1320 DOI 10.1016/j.compbiomed.2013.07.004.

Lee J, Sandhu R, Tannenbaum A. 2013. Particle filters and occlusion handling for rigid 2D-3D pose tracking. *Computer Vision and Image Understanding* 117:922–933 DOI 10.1016/j.cviu.2013.04.002.

Liu ZY, Bai LJ, Dai RW, Zhong CG, Xue T, You YB, Tian J. 2012. Dysfunctional whole brain networks in mild cognitive impairment patients: an fMRI study. In: *Medical imaging 2012: biomedical applications in molecular, structural, and functional imaging*, vol. 8317, 6 pages.

Lu ZT, Yang W, Zhang MH. 2014. Robust medical image elastic registration using global optimisation strategy in frequency domain. *Imaging Science Journal* 62:42–47 DOI 10.1179/1743131X12Y.0000000043.

Mangasarian OL, Wild EW. 2006. Multisurface proximal support vector machine classification via generalized eigenvalues. *IEEE Transactions on Pattern Analysis and Machine Intelligence* 28:69–74 DOI 10.1109/TPAMI.2006.17.

Miller V, Erlien S, Piersol J. 2012. Identifying dementia in MRI scans using machine learning. Available at http://cs229.stanford.edu/proj2012/ErlienMillerPiersol-identifyingDementiaInMRIScansUsingMachineLearning.pdf.
Montagne A, Barnes SR, Sweeney MD, Halliday MR, Sagare AP, Zhao Z, Toga AW, Jacobs RE, Liu CY, Amezcua L, Harrington MG, Chui HC, Law M, Zlokovic BV. 2015. Blood–brain barrier breakdown in the aging human hippocampus. *Neuron* **85**:296–302 DOI 10.1016/j.neuron.2014.12.032.

Murphy MC, Huston J, Jack CR, Glaser KJ, Manduca A, Felmlee JP, Ehman RL. 2011. Decreased brain stiffness in Alzheimer's disease determined by magnetic resonance elastography. *Journal of Magnetic Resonance Imaging* **34**:494–498 DOI 10.1002/jmri.22707.

Narayan PJ, Lill C, Faull R, Curtis MA, Dragunow M. 2015. Increased acetyl and total histone levels in post-mortem Alzheimer’s disease brain. *Neurobiology of Disease* **74**:281–294 DOI 10.1016/j.nbd.2014.11.023.

Nasiri JA, Charkari NM, Mozafari K. 2014. Energy-based model of least squares twin Support Vector Machines for human action recognition. *Signal Processing* **104**:248–257 DOI 10.1016/j.sigpro.2014.04.010.

Nazir M, Wahid F, Khan SA. 2015. A simple and intelligent approach for brain MRI classification. *Journal of Intelligent & Fuzzy Systems* **28**:1127–1135 DOI 10.3233/IFS-141396.

Ouyang X, Sun XY, Guo T, Sun QY, Chen KW, Yao L, Wu X, Guo XJ. Alzheimer’s Disease Neuroimaging Initiative. 2014. Independent component analysis of DTI data reveals white matter covariances in Alzheimer’s disease. In: *Medical imaging 2014: biomedical applications in molecular, structural, and functional imaging*, vol. 9038, 7 pages.

Park H. 2012. Comparison of distance measures for manifold learning: application to Alzheimer’s brain scans. *Journal of the Korean Physical Society* **61**:1148–1155 DOI 10.3938/jkps.61.1148.

Pekarya AE, Sattin A, Lloyd RL. 2015. Ketamine modulates TRH and TRH-like peptide turnover in brain and peripheral tissues of male rats. *Peptides* **69**:66–76 DOI 10.1016/j.peptides.2015.04.003.

Pennanen C, Kivipelto M, Tuomainen S, Hartikainen P, Hänninen T, Laakso MP, Hallikainen M, Vanhanen M, Nissinen A, Helkala E-L, Vainio P, Vanninen R, Partanen K, Soininen H. 2004. Hippocampus and entorhinal cortex in mild cognitive impairment and early AD. *Neurobiology of Aging* **25**:303–310 DOI 10.1016/S0197-4580(03)00084-8.

Petibon Y, Ouyang J, Zhu X, Huang C, Reese TG, Chun SY, Li Q, El Fakhri G. 2013. Cardiac motion compensation and resolution modeling in simultaneous PET-MR: a cardiac lesion detection study. *Physics in Medicine and Biology* **58**:2085–2102 DOI 10.1088/0031-9155/58/7/2085.

Petit J, Castelnau O, Bornert M, Zhang FG, Hofmann F, Korsunsky AM, Faurie D, Le Bourlot C, Micha JS, Robach O, Ulrich O. 2015. Laue-DIC: a new method for improved stress field measurements at the micrometer scale. *Journal of Synchrotron Radiation* **22**:980–994 DOI 10.1107/S1600577515005780.

Pirone A, Magliaro C, Giannessi E, Ahluwalia A. 2015. Parvalbumin expression in the claustrum of the adult dog. An immunohistochemical and topographical study with comparative notes on the structure of the nucleus. *Journal of Chemical Neuroanatomy* **64–65**:33–42 DOI 10.1016/j.jchemneu.2015.02.004.

Plant C, Teipel SJ, Oswald A, Böhm C, Meindl T, Mourao-Miranda J, Bokde AW, Hampel H, Ewers M. 2010. Automated detection of brain atrophy patterns based on MRI for the prediction of Alzheimer's disease. *NeuroImage* **50**:162–174 DOI 10.1016/j.neuroimage.2009.11.046.

Ramos P, Santos A, Pinto NR, Mendes R, Magalhaes T, Almeida A. 2015. Anatomical regional differences in selenium levels in the human brain. *Biological Trace Element Research* **163**:89–96 DOI 10.1007/s12011-014-0160-z.
Redolfi A, Manset D, Barkhof F, Wahlund LO, Glatard T, Mangin JF, Frisoni GB, Alzheimer’s Disease Neuroimaging Initiative. 2015. Head-to-head comparison of two popular cortical thickness extraction algorithms: a cross-sectional and longitudinal study. PLoS ONE 10(3):e0117692 DOI 10.1371/journal.pone.0117692.

Rieck JR, Rodrigue KM, Kennedy KM, Devous MD, Park DC. 2015. The effect of beta-amyloid on face processing in young and old adults: a multivariate analysis of the BOLD signal. Human Brain Mapping 36:2514–2526 DOI 10.1002/hbm.22788.

Saritha M, Joseph KP, Mathew AT. 2013. Classification of MRI brain images using combined wavelet entropy based spider web plots and probabilistic neural network. Pattern Recognition Letters 34:2151–2156 DOI 10.1016/j.patrec.2013.08.017.

Savio A, Grana M. 2013. Deformation based feature selection for computer aided diagnosis of Alzheimer’s disease. Expert Systems with Applications 40:1619–1628 DOI 10.1016/j.eswa.2012.09.009.

Schultz SA, Larson J, Oh J, Koscik R, Dowling MN, Gallagher CL, Carlsson CM, Rowley HA, Bendlin BB, Asthana S, Hermann BP, Johnson SC, Sager M, LaRue A, Okonkwo OC. 2014. Participation in cognitively-stimulating activities is associated with brain structure and cognitive function in preclinical Alzheimer’s disease. Brain Imaging and Behavior Epub ahead of print.

Shao YH, Chen WJ, Zhang JJ, Wang Z, Deng NY. 2014. An efficient weighted Lagrangian twin support vector machine for imbalanced data classification. Pattern Recognition 47:3158–3167 DOI 10.1016/j.patcog.2014.03.008.

Shao YH, Deng NY, Chen WJ, Wang Z. 2013. Improved generalized eigenvalue proximal support vector machine. IEEE Signal Processing Letters 20:213–216 DOI 10.1109/LSP.2012.2216874.

Shimoda K, Kimura M, Yokota M, Okubo Y. 2015. Comparison of regional gray matter volume abnormalities in Alzheimer’s disease and late life depression with hippocampal atrophy using VSRAD analysis: a voxel-based morphometry study. Psychiatry Research-Neuroimaging 232:71–75 DOI 10.1016/j.pscychresns.2015.01.018.

Song YT, Wang JT. 2010. Overview of Chinese research on senile dementia in mainland China. Ageing Research Reviews 9:S6–S12 DOI 10.1016/j.arr.2010.08.007.

Tan Y, Ren H, Shi Z, Yao X, He C, Kang JX, Wan JB, Li P, Yuan TF, Su H. Endogenous Docosahexaenoic Acid (DHA) prevents abeta1-42 oligomer-induced neuronal injury. Molecular Neurobiology DOI 10.1007/s12035-015-9224-0.

Tramutola A, Triplett JC, Di Domenico F, Niedowicz DM, Murphy MP, Coccia R, Perluigi M, Butterfield DA. 2015. Alteration of mTOR signaling occurs early in the progression of Alzheimer disease (AD): analysis of brain from subjects with pre-clinical AD, amnestic mild cognitive impairment and late-stage AD. Journal of Neurochemistry 133:739–749 DOI 10.1111/jnc.13037.

Vandemeulebroucke J, Bernard O, Rit S, Kybic J, Clarysse P, Sarrut D. 2012. Automated segmentation of a motion mask to preserve sliding motion in deformable registration of thoracic CT. Medical Physics 39:1006–1015 DOI 10.1118/1.3679009.

Villeneuve S, Rabinovici GD, Cohn-Sheehy BI, Madison C, Ayakta N, Ghosh PM, La Joie R, Arthur-Bentil SK, Vogel JW, Marks SM, Lehmanna M, Rosen HJ, Reed B, Olichney J, Boxer AL, Miller BL, Borys E, Jin L-W, Huang EJ, Grinberg LT, DeCarli C, Seeley WW, Jagust W. 2015. Existing Pittsburgh Compound-B positron emission tomography thresholds are too high: statistical and pathological evaluation. Brain: A Journal of Neurology 138:2020–2033 DOI 10.1093/brain/awv112.
Wang S, Dong Z, Ji G, Zhang Y. 2014. Classification of Alzheimer disease based on structural magnetic resonance imaging by kernel support vector machine decision tree. Progress in Electromagnetics Research 144:171–184 DOI 10.2528/PIER13121205.

Wang XL, Du MY, Chen TL, Chen ZQ, Huang XQ, Luo Y, Zhao YJ, Kumar P, Gong QY. 2015c. Neural correlates during working memory processing in major depressive disorder. Progress in Neuro-Psychopharmacology & Biological Psychiatry 56:101–108 DOI 10.1016/j.pnpbp.2014.08.011.

Wang S, Zhang Y, Dong Z, Du S, Ji G, Yan J, Yang J, Wang Q, Feng C, Phillips P. 2015a. Feed-forward neural network optimized by hybridization of PSO and ABC for abnormal brain detection. International Journal of Imaging Systems and Technology 25:153–164 DOI 10.1002/ima.22132.

Wang S, Zhang Y, Ji G, Yang J, Wu J, Wei L. 2015b. Fruit classification by wavelet-entropy and feedforward neural network trained by fitness-scaled chaotic ABC and biogeography-based optimization. Entropy 17:5711–5728 DOI 10.3390/e17085711.

Willette AA, Modanlo N, Kapogiannis D, Alzheimer's Disease Neuroimaging Initiative. 2015. Insulin resistance predicts medial temporal hypermetabolism in mild cognitive impairment conversion to Alzheimer disease. Diabetes 64:1933–1940 DOI 10.2337/db14-1507.

Xinyun C, Wenlu Y, Xudong H. 2011. ICA-based classification of MCI vs HC. In: Natural computation (ICNC), 2011 seventh international conference on. 1658–1662.

Xu ZJ, Qi ZQ, Zhang JQ. 2014. Learning with positive and unlabeled examples using biased twin support vector machine. Neural Computing & Applications 25:1303–1311 DOI 10.1007/s00521-014-1611-3.

Yamashita K, Taniwaki Y, Utsunomiya H, Taniwaki T. 2014. Cerebral blood flow reduction associated with orientation for time in amnesic mild cognitive impairment and Alzheimer disease patients. Journal of Neuroimaging 24:590–594 DOI 10.1111/jon.12096.

Yang D, Deasy JO, Low DA, El Naqa I. 2007. Level set motion assisted non-rigid 3D image registration. In: Pluim JPW, Reinhardt JM, eds. Medical imaging 2007 conference, Proceedings of the society of photo-optical instrumentation engineers, vol. 6512. San Diego: SPIE, 651207–651211.

Yang G, Zhang Y, Yang J, Ji G, Dong Z, Wang S, Feng C, Wang Q. 2015. Automated classification of brain images using wavelet-energy and biogeography-based optimization. Multimedia Tools and Applications 74(9):1–17 DOI 10.1007/s11042-015-2649-7.

Yokoyama JS, Lee AK, Takada LT, Busovaca E, Bonham IW, Chao SZ, Tse M, He J, Schwarz CG, Carmichael OT, Matthews BR, Karydas A, Weiner MW, Coppola G, DeCarli CS, Miller BL, Rosen HJ. 2015. Apolipoprotein epsilon 4 is associated with lower brain volume in cognitively normal chinese but not white older adults. PLoS ONE 10(3):e0118338 DOI 10.1371/journal.pone.0118338.

Yoo SW, Han CE, Shin JS, Seo SW, Na DL, Kaiser M, Jeong Y, Seong JK. 2015. A network flow-based analysis of cognitive reserve in normal ageing and Alzheimer’s disease. Scientific Reports 5:Article 10057 DOI 10.1038/srep10057.

Yu Q, Peng Y, Mishra V, Ouyang A, Li H, Zhang H, Chen M, Liu S, Huang H. 2014. Microstructure, length, and connection of limbic tracts in normal human brain development. Frontiers in Aging Neuroscience 6:Article 00228 DOI 10.3389/fnagi.2014.00228.

Yuan TF, Hou GL. 2015. The effects of stress on glutamatergic transmission in the brain. Molecular Neurobiology 51:1139–1143 DOI 10.1007/s12035-014-8783-9.
Yuan TF, Machado S, Arias-Carrion O. 2014. Gut microbiota and brain function: a new target for brain diseases? CNS & Neurological Disorders-Drug Targets 13:733–735 DOI 10.2174/187152731305140717104045.

Zhang YD, Dong ZC, Ji GL, Wang SH. 2014b. An improved reconstruction method for CS-MRI based on exponential wavelet transform and iterative shrinkage/thresholding algorithm. Journal of Electromagnetic Waves and Applications 28:2327–2338 DOI 10.1080/09205071.2014.967365.

Zhang Y, Dong Z, Ji G, Wang S. 2015a. Effect of spider-web-plot in MR brain image classification. Pattern Recognition Letters 62:14–16 DOI 10.1016/j.patrec.2015.04.016.

Zhang Y, Dong Z, Phillips P, Wang S, Ji G, Yang J. 2015b. Exponential wavelet iterative shrinkage thresholding algorithm for compressed sensing magnetic resonance imaging. Information Sciences 322:115–132 DOI 10.1016/j.ins.2015.06.017.

Zhang Y, Dong Z, Wang S, Ji G, Yang J. 2015c. Preclinical diagnosis of Magnetic Resonance (MR) brain images via discrete wavelet packet transform with tssallis entropy and generalized eigenvalue proximal support vector machine (GEPSVM). Entropy 17:1795–1813 DOI 10.3390/e17041795.

Zhang Y, Wang S, Dong Z, Phillips P, Ji G, Yang J. 2015d. Pathological brain detection in magnetic resonance imaging scanning by wavelet entropy and hybridization of biogeography-based optimization and particle swarm optimization. Progress in Electromagnetics Research 152:41–58.

Zhang Y, Wang S, Ji G, Dong Z. 2013. An MR brain images classifier system via particle swarm optimization and kernel support vector machine. The Scientific World Journal 2013:Article 130134 DOI 10.1155/2013/130134.

Zhang Y, Wang S, Dong Z, Phillips P, Ji G, Yang J. 2015e. Exponential wavelet iterative shrinkage thresholding algorithm with random shift for compressed sensing magnetic resonance imaging. IEEJ Transactions on Electrical and Electronic Engineering 10:116–117 DOI 10.1002/tee.22059.

Zhang Y, Wang S, Phillips P, Dong Z, Ji G, Yang J. 2015f. Detection of Alzheimer's disease and mild cognitive impairment based on structural volumetric MR images using 3D-DWT and WTA-KSVM trained by PSOTVAC. Biomedical Signal Processing and Control 21:58–73 DOI 10.1016/j.bspc.2015.05.014.

Zhang Y, Wang S, Phillips P, Ji G. 2014a. Binary PSO with mutation operator for feature selection using decision tree applied to spam detection. Knowledge-Based Systems 64:22–31 DOI 10.1016/j.knosys.2014.03.015.

Zhang Y, Wu L. 2012. An MR brain images classifier via principal component analysis and kernel support vector machine. Progress in Electromagnetics Research 130:369–388 DOI 10.2528/PIER12061410.

Zhou X, Wang S, Xu W, Ji G, Phillips P, Sun P, Zhang Y. 2015. Detection of pathological brain in MRI scanning based on wavelet-entropy and naive bayes classifier. In: Ortuno F, Rojas I, eds. Bioinformatics and biomedical engineering. Granada: Springer International Publishing, 201–209.