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MRI-Based Automated Computer Classification of Probable AD Versus Normal Controls

Simon Duchesne*, Member, IEEE, Anna Caroli, C. Geroldi, Christian Barillot, Senior Member, IEEE, Giovanni B. Frisoni, and D. Louis Collins

Abstract—Automated computer classification (ACC) techniques are needed to facilitate physician’s diagnosis of complex diseases in individual patients. We provide an example of ACC using computational techniques within the context of cross-sectional analysis of magnetic resonance images (MRI) in neurodegenerative diseases, namely Alzheimer’s dementia (AD). In this paper, the accuracy of our ACC methodology is assessed when presented with real life, imperfect data, i.e., cohorts of MRI with varying acquisition parameters and imaging quality. The comparative methodology uses the Jacobian determinants derived from dense deformation fields and scaled grey-level intensity from a selected volume of interest centered on the medial temporal lobe. The ACC performance is assessed in a series of leave-one-out experiments aimed at separating 75 probable AD and 75 age-matched normal controls. The resulting accuracy is 92% using a support vector machine classifier based on least squares optimization. Finally, it is shown in the Appendix that determinants and scaled grey-level intensity are appreciably more robust to varying parameters in validation studies using simulated data, when compared to raw intensities or grey/white matter volumes. The ability of cross-sectional MRI at detecting probable AD with high accuracy could have profound implications in the management of suspected AD candidates.

Index Terms—Accuracy, automated computer classification, magnetic resonance imaging (MRI), neurodegenerative diseases.

I. INTRODUCTION

NEURODEGENERATIVE diseases can be characterized by their gradual modification of the cellular environment leading to neuronal dysfunction, abnormal loss of brain tissue, and accompanying cognitive impairment. The factors and mechanisms underlying these processes will vary from one disease to another. In Alzheimer’s disease (AD), abnormal accumulation of insoluble A/β and Tau proteins into intra- and extra-cellular formations has been linked to cerebral tissue damage, leading to dementia [1]. The pattern of distribution of plaques and tangles has been characterized by Braak et al. [1] in six neuropathological stages, ranging from initial deposition in the medial temporal lobe to distribution throughout the cortex. This etiology implies that final AD diagnosis can only be confirmed at present by postmortem histopathological assessment. Structural images, such as T1, T2, and PD weighted magnetic resonance imaging (MRI) sequences acquired on standard clinical scanners of 1.0–3.0 T field strength, lack the resolution to measure directly microscopic changes due to probable AD in the cellular environment. Thus, it is clear that structural imaging cannot be used for deriving a final diagnosis; it can, however, image some of the resulting macroscopic disease-related effects that result in changes in shape, size, or image intensity of anatomical structures [2]. Specifically, the role of structural imaging in probable AD has shifted from one of exclusion of other possible causes of dementia to one of in vivo disease-specific marker that, it is hoped, can identify the disease at a very early stage [3], [4]. Various computer-aided techniques have been proposed in the past and include the study of texture changes in signal intensity [5], grey matter (GM) concentrations differences [6], atrophy of subcortical limbic structures [7]–[9], and general cortical atrophy [10]–[12]. These automated studies typically report group-level differences between probable AD patients and age-matched control subjects for various brain structures [2], often in a longitudinal setting.

The goal of our research into automated computer classification (ACC) techniques is to achieve individual classification. Further, we restrict our focus to cross-sectional or single time point assessment, as opposed to longitudinal acquisitions, since the former allows for a more timely alteration in therapy course that may represent a significant advantage when dealing with neurodegenerative diseases.

Within this context, the standard in terms of image analysis in probable AD remains manual segmentation of specific structures such as the hippocampus (HC) and parahippocampal gyrus (PHG), HC and PHG volumetric measurements have achieved individual classification rates between controls and probable AD patients as high as 93% when using linear combinations of volumes [13], [14]. Manual segmentation however is a time-consuming, center and expert-dependent technique [15], [16] that cannot be used in regular clinical practice.

ACC techniques on the other hand can be objective, quantitative, and practical. Previous reports on ACC mention the use of a variety of image features to fulfill the task of individual classification. The signal intensity was modeled, either directly [17], via texture analysis [5], [18], or via GM tissue classification [19]; classification based on the morphometry of specific brain structures has been proposed [9], [20], [21]; and lately cortical mantle thickness estimates have been investigated [12]. For these methods, the robustness to varying scanning parameters, repetition/echo time (TR/TE), field inhomogeneity and noise has yet to be studied and/or reported. The strength of any ACC
system should not lie solely in the optimality of its classifier but also in its choice of features; they must be representative of the disease process, and robust, that is, capable of providing the same information irrespective of noise, artifact, or intrinsic variation in the underlying image due to different parameter settings [22]. Real life datasets are rife with such differences. Sources of errors in MRI measurements have been aptly described and documented exhaustively elsewhere [23], and include, but are not limited to
- signal noise;
- intensity nonuniformity (INU);
- parameter changes (small changes in TR/TE);
- voxel dimension changes;
- missing slices;
- anatomical variability.

This assumes that all subjects to be assessed have been scanned on the same system, whose configuration (hardware, software) remained stable over time. It is important to note that there are no issues of scan–rescan effects for cross-sectional analysis, such as repositioning, as might be the case in longitudinal data [24], [25].

Our original contribution in ACC design is the combination of intensity and local shape descriptor [17] extracted from a large volume of interest (VOI), avoiding the pitfalls of structure segmentation and modeling de facto neighboring tissue interactions. Thus, the ACC assessment is based on estimating the individual subject’s propensity at exhibiting pathology-specific patterns of covarying tissue changes within the VOI; these can be expressed as areas of signal changes or tissue atrophy [17], [26] and related to biological phenomena. This is in line with recent findings in aging and probable AD [27]–[29]. It must be emphasized that we are not claiming to detect molecular changes per se on standard field strength MRI; rather, we are interested in detecting macroscopic pathology-related changes. Equally important is the fact that the VOI selected for this study is centered on the medial temporal lobes, as it represents the area most affected in probable AD; it does not address whole brain changes. This VOI is suitable for the classification of probable AD versus normal controls, but may not be appropriate for other classification tasks.

In this paper, our goal is to evaluate the robustness of our ACC methodology for probable AD versus controls differentiation [30] when presented with random and systematic errors in real and simulated datasets, such as changes in the acquisition parameters and varying INU. Our hypothesis is that the accuracy of our methodology will not be affected by realistic variations in the acquisition parameters. The focus is therefore on finding a workable solution to the problem of unequal scan quality found in a day-to-day clinical operation. This is a necessary first step before assessing the accuracy of our ACC methodology in multisite/multiscanner situations, itself a requirement before using such techniques in routine clinical use. These scenarios are outside the scope of this paper.

II. METHODS

A. Subjects

A total of 299 subjects were included in this study. The first cohort, or reference group, consisted in 149 young, neurologically healthy individuals from the International Consortium for Brain Mapping database (ICBM) [31], whose scans were used to create the nonpathological, reference space. The second cohort, or study group, consisted in 150 subjects: 75 patients with a diagnosis of probable AD and 75 age-matched normal controls (NC) without neurological or neuropsychological deficit. The AD subjects are individuals with mild to moderate probable AD [32] recruited among outpatients seen at the Centro San Giovanni di Dio Fatebenefratelli, The National Center for AD (Brescia, Italy) between November 2002 and January 2005. History was taken with a structured interview from a knowledgeable informant (usually the patient’s spouse) and was particularly focused on those symptoms that might help in the differential diagnosis of the dementias in order to avoid contamination of non-AD dementias in the study group. Laboratory examinations included complete blood count, chemistry profile, thyroid function, B12 and folic acid, and EKG. A neurologist performed structured neurological examination and a geriatrician performed the physical examination. A comprehensive neuropsychological battery was also administered. The MRIs were not used in the assessment of the subject’s condition. NC subjects were taken from an ongoing study of the structural features of normal aging. This study recruits outpatients attending the Neuroradiology Unit of the Citta’ di Brescia Hospital (Brescia, Italy) aged 40 and older and undergoing brain magnetic resonance (MR) scan for reasons other than cognitive impairment (usually headache and vertigo) and negative for major stroke, tumor, aneurysm, or other focal lesions. Incidental atrophy, white matter disease, and lacunes were not exclusionary criteria. Normality of cognitive functions was ascertained through neuropsychological evaluation and structured interview. Analysis of hippocampal shape for some of these probable AD and NC subjects has been published elsewhere [8].

The Ethics Committee of the IRCCS San Giovanni di Dio FBF (Brescia, Italy) approved the study and informed consent was obtained from all participants.

B. Images

The ICBM subjects from the reference group were scanned in Montreal, QC, Canada on a Philips Gyroscan 1.5-T scanner (Best, The Netherlands) using a T1-weighted fast gradient echo sequence (sagittal acquisition, TR = 18 ms, TE = 10 ms, 1 mm × 1 mm × 1 mm voxels, flip angle 30°), for all acquisitions.

MRI data for all subjects in the probable AD and NC study group were acquired in Brescia, Italy on a single Philips Gyroscan 1.0-T scanner (Best, The Netherlands) using a T1-weighted fast field echo sequence (sagittal acquisition). As the protocol set general guidelines for study parameters, there exist three combination of repetition and echo times (TR/TE) for both NC and probable AD groups, as described in Table I. Albeit the differences are small, the resulting images have slightly different contrasts between GM, white matter (WM), scalp tissue, and fat. Even though specified at 1.0 mm × 1.0 mm, there also co-exists small variations in the in-plane resolutions. While the slice thickness remained equal (1.3 mm), the in-plane pixel size ranged from slightly below (0.9 mm × 0.9 mm) to slightly above (1.1 mm × 1.1 mm) the requirement. The distribution of in-plane resolution parameters
TABLE I

<table>
<thead>
<tr>
<th>Set</th>
<th>NC</th>
<th>AD</th>
<th>TR (ms)</th>
<th>TE (ms)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>24</td>
<td>7</td>
<td>20</td>
<td>4.9</td>
</tr>
<tr>
<td>2</td>
<td>12</td>
<td>25</td>
<td>20</td>
<td>6.9</td>
</tr>
<tr>
<td>3</td>
<td>39</td>
<td>43</td>
<td>25</td>
<td>6.9</td>
</tr>
<tr>
<td>Totals</td>
<td>75</td>
<td>75</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

NC: Normal Controls; AD: probable Alzheimer’s Dementia

was approximately similar in either probable AD (34 below, 22 standard, 19 above; respectively, 45.3%, 29.3%, and 25.3% of total) or NC groups (41 below, 19 standard, 15 above; respectively, 54.7%, 25.3%, and 20% of total).

The use of a common referential target (from hereon, called the “target”) allows for interindividual comparisons. Rather than using a population average, we used a high-contrast, high-resolution target consisting in the average of 27 T1w MRI scans of the same individual [33], known as the Colin27 MRI average (see Fig. 1). This target exists within a Talairach-like stereotaxic space defined in the context of the ICBM project. Acquisition of the target images was done on the same scanner as the Colin27 MRI average.

C. Image Processing

All volumes (reference and study subjects, phantoms) were processed according to a similar pipeline, schematized in Fig. 2. The first step following image reconstruction (Fig. 2, item 1) consisted in the correction of the impact of INU (Fig. 2, item 2) due to scanner variations on global MRI data using a non-parametric estimation of the slow varying nonuniformity field [34] (Fig. 2, item 3). Next, prior to spatial alignment of the individual image with the target volume, we scaled its mean volume intensity to the target mean intensity and fixed the intensities within a [0–100] range (Fig. 2, item 1). Spatial alignment of individual images to the target volume (Fig. 2, item 1) was performed using a multistep procedure comparing intensity information from the slightly blurred subject image (FWHM=2 mm³) to the unblurred target [35], increasing at each step the number of degrees of freedom (DF) from a rigid to affine estimate (6, 7, 9, then 12 DF). Due to the difference in sequence parameters, the problem was treated as one of multimodality matching, and, therefore, mutual information was chosen as the objective cost function [36], [37]. This choice of cost function also has the advantage of reducing the impact of any intensity difference due to the parameters under study (e.g., TR/TE, INU, noise). This linear registration process was repeated twice, once (Fig. 2, item 5) to optimize the global brain orientation and size, and a second time (Fig. 2, item 6) to optimize the local alignment of the volume of interest (VOI) from the subject’s brain (Fig. 2, item 8) to the corresponding target VOI. The single VOI selected for this study encompassed both medial temporal lobes, and was oriented towards the long axis of the hippocampus (see Fig. 1). It measured $n = 90 	imes 90 	imes 50 = 405000$ voxels. Defined within the Colin27 target space, its extent captured the hippocampus and neighboring limbic structures (e.g., amygdala, parahippocampal gyrus), irrespective of normal interindividual variability. As a part of the alignment procedure, the data were resampled (Fig. 2, item 7) using trilinear interpolation onto a 1-mm isotropic grid [35]. The first feature of our model consists in the rasterized, resampled, scaled grey-level data from the linearly registered VOI, denoted $g$ (Fig. 2, item 10).

Nonlinear image registration (Fig. 2, item 9) was then performed to derive our shape estimator of local volume change, the second central feature of our ACC methodology. Nonlinear registration attempts to match image features from a subject volume to those of the target image at a local level, typically in a hierarchical fashion, with the aim of reducing a specific cost function. Whereas many nonlinear registration processes exist, the one chosen for this study was ANIMAL [35]. This algorithm attempts to match image grey-level intensity features at a local level (voxel) in successive blurring steps, by maximizing the mutual information of voxel intensities between the subject and target images. The result is a dense deformation field capturing the displacements required to align all voxels within the subject VOI with those of the target VOI. Our local volume change estimator is computed (Fig. 2, item 11) from the determinants of the Jacobian of the deformation fields. Our implementation of determinant follows the notation developed by Chung et al. [38]. If $U$ is the displacement field which matches homologous points between two images, then the local volume change of the deformation in the neighborhood of any given voxel is determined by the Jacobian $J$ [38], which is defined as

$$J(x,t) = \det \left( I + \frac{\partial U}{\partial x} \right) \quad (1)$$

where $I$ denotes the identity matrix and $(\partial U/\partial x)$ is the $3 \times 3$ displacement gradient matrix of $U$.

The rasterized determinant $\det(\nabla U)$, denoted $d$, is the second feature of our model (Fig. 2, item 11). It represents a biologically meaningful quantity, as it is an indicative measure of local brain...
tissue volume difference when compared to the target volume. When the change is near zero in the neighborhood of $x$, there is no local difference in volume between subject and target images. However, if the determinant is positive, the volume increases whereas when negative, the volume decreases after the deformation.

### D. Modeling and Classification

Our classification approach has been to create a model eigenspace based on grey-level intensity $g$ and determinant information $d$ from subjects in the ICBM database, the coordinate space in which we projected image data from our cohorts of patients. The classification was then based on the eigencoordinate distributions of the projected data.

Principal components analysis (PCA) (Fig. 2, items 12 and 13) is used to reduce the dimensionality of the input reference data and generate a linear variation model of covarying information for the ICBM subjects. A more thorough description of the PCA process was presented in a previous publication [17], based on the seminal work of Cootes and Taylor [39]. In short, we use as input data rasterized vectors of intensity $g_i$ and Jacobian determinants $d_i$ for all images $i$ belonging to the ICBM reference dataset ($N_{	ext{ICBM}} = 149$). Whereas a space built from young, neurologically healthy individuals may not be optimal to represent the study group, our primary goal was to create an independent basis for a comparative evaluation of the MRIs from probable AD and NC individuals and their subsequent individual classification, rather than a descriptive space that would be the best mathematical representation for aging and disease.

To make the PCA a zero-mean process modeling differences between subjects and the group mean, we first subtract the respective grey-level and determinant averages of the ICBM reference groups from each input vector

$$g_k = G_k - G_{	ext{ICBM}}$$  \(2\)

$$d_k = D_k - D_{	ext{ICBM}}.$$  \(3\)

To define the model, PCA is applied to the 149 $g_k$ and 149 $d_k$ reference input data vectors (see Cootes et al. [39] and Duchesne et al. [17], [40] for a more thorough description of the PCA process). The resulting ensemble of $p$ principal components, where $p = N - 1$, defines an allowable grey-level domain $G$ (Fig. 2, item 14) and allowable determinant domain $D$ (Fig. 2, item 15) as the spaces of all possible elements expressed by the determinant eigenvectors $\lambda_G$ and $\lambda_D$.

Most of the variation can usually be explained by a smaller number of modes, $l$, where $l \ll n$ and $l < p$. The total variance of all the variables is equal to

$$\lambda = \sum_{k=1}^{2n} \lambda_k$$  \(4\)

whereas for $l$ eigenvectors, explaining a sufficiently large proportion of $\lambda$, the sum of their variances, or how much these principal directions contribute in the description of the total variance.
of the system, is calculated with the ratio of relative importance of the eigenvalue \( \lambda_k \) associated with the eigenvector \( k \):

\[
\gamma_k = \frac{\lambda_k}{\sum_{j=1}^{p} \lambda_j}.
\]

(5)

The theoretical upper-bound on the dimensionality \( f \) of \( G \) and \( D \) is \( N - 1 \) however, we define restricted versions of these spaces denoted \( G* \) and \( D* \) (Fig. 2, items 16 and 17), using only the first \( k \) eigenvectors corresponding to a given ratio \( r \) for each space.

Following the notation of Duda et al. [41], we have defined two states of nature \( \omega \) for our study subjects: \( \omega_{NC} = NC \), and \( \omega_{AD} = \) probable AD. For the purposes of this work, the prior probabilities \( p(\omega_{NC}), p(\omega_{AD}) \) were known equal \( (p = 0.5; \ p = 0.5) \) since the compositions of the classification data sets were determined. It must be stated that they do not represent the normal incidence rates of probable AD in the general population.

Once the model eigenspaces \( G* \) and \( D* \) from reference data have been formed, we proceed with the task of projecting the rasterized vectors \( \mathbf{I}_i \) and \( \mathbf{d}_i \) for the subjects in the study group (Fig. 2, item 18). The projected data in the Domain \( G* \) forms the eigencoordinate vectors \( \gamma_{\omega}^i \); likewise, projected data into the Domain \( D* \) forms the eigencoordinate vectors \( \delta_{\omega}^i \).

The distribution of eigencoordinates along any principal component for a given population is assumed normally distributed; this normality was assessed via quantile plots and Shapiro-Wilke statistics. Under this assumption of normality, we can use the vectors \( \gamma_{\omega}^i \) and \( \delta_{\omega}^i \) as feature vectors in a system of supervised linear classifiers (Fig. 2, item 19). The \( k\)-dimensional real vectors \( \gamma_{\omega}^i \) and \( \delta_{\omega}^i \) are normalized to guard against variables with larger variance that might otherwise dominate the classification.

For the purposes of this study we wanted to test simple yet robust classifiers, namely the following.

1) Linear and quadratic discriminant analyses (LDA, QDA).

Both are used in machine learning to separate measurements of two or more classes of objects or events by either a linear or a quadratic surface. In QDA there is no assumption that the covariance of each of the classes is identical. Forward stepwise analysis using Wilk’s \( \lambda \) method with two different values of \( P \) (0.005; 0.01) was used to select the discriminating variables. Classification experiments were computed using the MATLAB Statistics Toolbox (The MathWorks, Natick, MA). The different parameter combinations are detailed in Table II.

2) Support vector machines (SVM). SVMs are a class of linear classifiers that attempt to achieve maximum separation (margin) between the two classes [42]. Experiments were run using three kernels (linear, quadratic, or polynomial of third degree), two optimization techniques [two-norm, soft margin (SM) and least squares (LS)], and different constraint sizes on the margin (bounding box). Classification experiments were performed using the MATLAB Statistics Toolbox. The different parameter combinations are detailed in Table II.

All classification experiments were run as leave one out trials. In this subset of cross-validation approaches, one subject is removed at a time from the study group; the remaining subjects are used as training data to estimate the classification function. The subject left out is then classified independently from the training group. The process is repeated \( \eta \) times, \( \eta \) being the number of subjects in the study group (\( N_{\text{Study}} = 150 \)). The leave-one-out cross-validation has been shown to give an almost unbiased estimator of the generalization properties of statistical models [43]. The leave-one-out experiments were performed using MATLAB’s Statistics Toolbox.

Results were obtained in terms of accuracy (proportion of all subjects correctly classified), sensitivity (proportion of individuals with a true positive result), specificity (proportion of individuals with a true negative result), positive predictive value (PPV) (true positive over all positive results), and negative predictive value (NPV) (true negative over all negative results). Four sets of experiments were conducted.

1) NC versus probable AD classification using linear discriminant analyses. Experiment 1 (LDA) was designed to assess the accuracy of individual classification using forward, stepwise linear discriminant leave-one-out analyses.

2) NC versus probable AD classification using quadratic discriminant analyses. Experiment 2 (QDA) was designed to assess the accuracy of individual classification using forward, stepwise quadratic discriminant leave-one-out analyses.
Fig. 4. Automated computer classification results for the three experiments using 14 different combinations of classifiers. The full list and parameters can be found in Table II. Results are given in terms of accuracy (proportion of all subjects correctly classified as being probable AD versus NC), sensitivity (proportion of individuals with a true positive result for probable AD), specificity (proportion of individuals with a true negative result for NC), PPV (true positive over all positive results), and NPV (true negative over all negative results). Using an SVM with linear kernel and least-square optimization (without constraint on boundary) gives the highest classification accuracy (92%, column 8). Gold standard for comparison is comprehensive clinical assessment.

3) NC versus probable AD classification using SVM. Experiment 3 (SVM) was designed to assess the accuracy of individual classification using linear and quadratic SVM under various condition.

4) Random permutations of ground truth. In order to test the robustness of the ACC methodology, Experiment 4 (Random) was designed so that random ground truth (probable AD, NC) was assigned to each of the 150 individual study subjects; the system then attempted to classify using this information. We expected that the system would be, at best, equal to chance in classifying group membership. To test this condition, we performed 1000 trials using the best classifier as selected following the previous LDA, QDA, and SVM experiments. Only mean accuracy will be reported.

### III. Experiments and Results

#### A. Study Group Demographics

There were no statistically significant differences for age between the 75 probable AD (mean = 73.3 years; SD = 8.4 years) and 75 NC individuals (mean = 73.3 years; SD = 4.6 years) (Student’s $t$-test, $DF = 148$, $P > 0.05$) (see Fig. 3).

#### B. Allowable Grey-Level and Determinant Domains

We set the variance ratio $r$ [see (5)] to 0.997, resulting in a PCA model composed of 112 $\lambda_G$ eigenvectors spanning Domain $G*$ and 144 $\lambda_D$ eigenvectors spanning Domain $D*$. We have not performed a sensitivity analysis of the classification results for different values of $r$.

#### C. NC Versus Probable AD Classification Using Linear Discriminant Analyses

The best accuracy result was 126/150 or 84% reached with P-to-enter set at 0.005, retaining seven variables in the final discriminant function (median over 150 leave-one-out trials); detailed results are reported in Fig. 4.
D. NC Versus Probable AD Classification Using Quadratic Discriminant Analyses

The best accuracy result was 125/150 or 83.3% reached with P-to-enter set at 0.01, retaining 12 variables in the final discriminant function (median over 150 leave-one-out trials); detailed results are reported in Fig. 4.

E. NC Versus Probable AD Classification Using SVM

The best accuracy result was 138/150 or 92% reached with a linear SVM using least-squares optimization; bounding box constraints did not influence the results. Further results on the other classifiers are reported in Fig. 4.

F. Random Permutation of Ground Truth

The mean accuracy of the linear SVM with least-squares optimization at classifying 1000 trials with randomly allocated ground truth was 49.9% (standard deviation 5.1%; maximum 68%; minimum 36%). Other results (specificity, sensitivity, PPV, NPV) not reported.

IV. DISCUSSION

A. Clinical Considerations

This paper reports results of an automated computer classification technique used in the cross-sectional assessment of probable AD versus normal controls. Cross-sectional methods are potentially useful for early diagnosis, while prospective or longitudinal techniques have limited value in this regard, unless the repeat scan is very close to the baseline, e.g., less than 2–3 months, but to date this is not routinely done.

The cross-sectional accuracy for clinical assessment of probable AD versus normal controls, based on neurological and neuropsychological assessment, but verified against histopathological gold standard, is 78% (22% error rate), as mentioned in a recent consensus report [44]. When combining neuropsychological testing (Mini-Mental State Examination [45]) with visual rating of atrophy in the medial temporal lobe as seen from MRI [46], Wahlund and colleagues reported 95% sensitivity to probable AD. Accuracy of cross-sectional assessment of medial temporal lobe atrophy when combined with cerebrospinal fluid proteomic markers is comparable [47], and so is a linear combination of manually defined volumes (hippocampus, parahippocampal cortex) between controls and probable AD patients (93%) [13], [14].

The proposed ACC methodology achieves a level of accuracy that is similar (92%, 8% error rate) to the previous studies, albeit in a larger cohort, based on MRI information alone, and in the presence of different TR/TE and in-plane resolutions. Further, the proposed technique is completely automated (and therefore reproducible), requires no external expertise and no expert time, and is completely objective. Since MRI is noninvasive, there are no complications related to the procedure, the only limit being contra-indications to MR scanning. Thus, such a technique could easily be implemented in a clinical setting and complement the initial assessment. This can be compared to other techniques, such as PET [48], [49] and SPECT [50], that offer sufficient specificity and sensitivity in the diagnosis of probable AD but are minimally invasive procedures with radiation dose limitations and therefore cannot be repeatedly performed on a single patient nor used as a screening mechanism for large populations. Other MRI-based techniques such as functional magnetic resonance imaging (fMRI) [51], MR spectroscopy [52], MR diffusion tensor [53], and MR magnetization transfer [54] show promise for the future, but are difficult to implement in a clinical setting without a dedicated research group for technical support, data acquisition and analysis. Finally, as compared to MRI, CT images lack the detailed soft-tissue information necessary for detecting subtle structure changes associated with the disease, especially at an early stage, even though late-stage measures show promise [55].

The paper presents a tool for classifying patients; the fact that the tool classified patients well here does not, strictly speaking, imply anything about the population of patients at large. Classical statistical theory tells us that the results of statistical tests on a representative sample can be used to infer properties of the population at large up to a point; but the generalizability of classifier results is unclear. Therefore whether our technique can say anything about the brain patterns that probable AD subjects have in general must be further justified by studies on larger cohorts. Neupathological confirmation is also required to replace the clinical evaluation as a gold standard in the evolution of ACC accuracy.

Biological interpretation of the information embedded in the classification function, in terms of local volume and intensity changes, remains to be performed. The separating hyperplane contains information that should be of biological importance in the study of probable AD. The differences in local volume changes should mirror the changes noticed in other reports, such as visual assessment [46], while differences in grey-level might reflect the intensity of neuronal loss induced by the neuropathological changes [56], which precede volume loss as visualized on MRI. This evaluation is beyond the scope of this manuscript.

B. Methodological Considerations

Our goal is to extract pathological grey-level and deformation patterns that are specific and sensitive to the task at hand, i.e., separating probable AD from NC. The registration process relies on point homology, which is of course approximate: in regions where there is complete homology, the displacement field will be nearly exact, up to the accuracy of the ANIMAL procedure; and in regions where it is not, the result will be noisy. This latter uncertainty will be rejected in the PCA modeling, as it is uncorrelated and noncovarying. Our results further suggest that the system will remain robust even in multisite acquisitions, where systematic differences exist due to the various hardware/software configurations, not to mention platform- or vendor-specific sequences that provide similar, but never equal, contrasts [25]. The work presented here (multiple acquisitions, single site) is a necessary step in the validation of the ACC technique, whereas evaluation of the procedure using multisite data will be the subject of a future manuscript.

Over determination in the creation of the reference space is not an issue as the reference group is composed of subjects taken from the ICBM study; these are completely independent of the study cohort. Further, by using leave-one-out cross-validation
trials, we ensured that there was no over learning in the classification stage while maximizing the amount of information available. Experiment 5 (Random permutation) was aimed at demonstrating that the resulting ACC classification is not an artefactual result of over determined classification.

In this paper, the results demonstrate that SVM are marginally superior to other supervised classification techniques (LDA/QDA) for the chosen task. We opted for a single classification stage using forward, stepwise regression rather than a two-stage univariate/multivariate classification scheme [57], since the latter may result in inclusion of strongly statistically significant but nonorthogonal features in the second stage, at the expense of variables with wider variance but better discriminative value when combined to other features.

There are similarities and differences between this methodology and other classification methods [21], principally Lao et al. [19]. Both reduce a high-dimensional input vector and attempt the classification on the reduction parameters, albeit using a different technique (PCA in this work; wavelet decomposition for Lao et al. [19]); both having some ability at representing spatial covariances in the data. The main difference between the two techniques resides in the spatial extent and choice of features. Lao et al. [19] use mass-preserved GM, WM and CSF whole brain concentration estimate maps, which are, effectively, trichotomized versions of the original intensity image. These, as discussed above, may prove sensitive to different acquisition parameters. Our modeling is centered on the medial temporal lobe. For intensity, our approach is more in line with that of Webb et al. [58] and sidesteps the issue of GM/WM classification, that may suffer from over and under determination of real GM and WM volumes and/or concentrations due to tissue classification problems. The latter GM/WM variability is demonstrated in the Appendix for different contrasts, and further discussed in the work of Brickman et al. [27]. Finally, the proposed technique explicitly models local volume changes, as opposed to its implicit formulation within the mass-preserving framework of the RAVENS maps [59].

It is proposed to use statistical criteria (Akaike, Bayes) at a later date to determine the optimum number of eigenvectors to be retained in the separating hyperplane. Multisquence information (e.g., T2) and/or multimodality (e.g., PET) may also benefit the model.

V. CONCLUSION

An immediate goal of image processing is to augment the physicians’ confidence in their diagnosis by offering quantitative assessments of brain images, especially in the context of pathologies such as neurodegenerative disorders. We have presented a methodology for computer-aided diagnosis and demonstrated using simulated and real data, that it appears robust to small variations in MRI acquisition parameters, as they present themselves in routine clinical use. Major strengths of this study include the number of subjects in each cohort, and the use of an ACC method that is completely automated (and, therefore, reproducible), requires no external expertise, and is completely objective. The ability of cross-sectional MRI at detecting probable AD with high accuracy is supported by the results: MR-based assessment could cut the diagnosis error by a factor of two. This could have profound implication in the management of suspected AD candidates.

VI. ROLE OF THE FUNDING SOURCES

The funding sources had no involvement in study design, collection, analysis, and interpretation of the data, writing of the report and in the decision to submit the paper for publication.

APPENDIX

A. Phantom Image Processing

In order to assess the sensitivity of various image features to changes in TR, TE, noise, and INU, sets of coregistered phantom images were produced using the MNI BrainWeb tool (see Table III) [33]. These phantoms, built using the target volume, were by definition coregistered with it. Each MRI phantom was simulated with different imaging parameters, mimicking those present in the real dataset, making it an ideal means of assessing ACC features sensitivity.

In order to assess the sensitivity of various features to changes in acquisition parameters, our strategy was to simulate different MRI phantoms [33], varying the image acquisition parameters for each simulation, mirroring the values found in the real dataset. In the first three phantoms, TR and TE were changed; in phantoms 4–6, TR/TE were identical, but noise was added to the phantom. In phantoms 7–9, TR/TE were identical, but three different INU fields were used, and finally, in phantoms 10–12, all parameters were changed (TR, TE, noise, INU). Table III summarizes the various parameters for each phantom.

Since it is only possible to change slice thickness and not in-plane resolution with the MNI BrainWeb, and since the original image from which the phantoms are built was acquired at 1 mm<sup>2</sup> isotropic resolution, we chose not to assess the variability of the measurements to different thickness/in-plane resolutions.

Different measures were then computed on each of the phantom image groups. These included the mean VOI T1-weighted intensity, in the native images (“unprocessed

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**TABLE III**

<table>
<thead>
<tr>
<th>Phantom</th>
<th>TR (ms)</th>
<th>TE (ms)</th>
<th>Noise (%)&lt;sup&gt;1&lt;/sup&gt;</th>
<th>INU Field&lt;sup&gt;2&lt;/sup&gt; (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
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<td>4.9</td>
<td>0</td>
<td>A</td>
</tr>
<tr>
<td>2</td>
<td>20</td>
<td>6.9</td>
<td>0</td>
<td>A</td>
</tr>
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<tr>
<td>5</td>
<td>20</td>
<td>4.9</td>
<td>0</td>
<td>B</td>
</tr>
<tr>
<td>6</td>
<td>20</td>
<td>4.9</td>
<td>0</td>
<td>C</td>
</tr>
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<td>5</td>
<td>A</td>
</tr>
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<tr>
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<td>20</td>
<td>6.9</td>
<td>5</td>
<td>B</td>
</tr>
<tr>
<td>12</td>
<td>25</td>
<td>6.9</td>
<td>5</td>
<td>C</td>
</tr>
</tbody>
</table>

<sup>1</sup>White additive Gaussian noise added to the ground truth of the BrainWeb phantom, in the given proportion relative to the signal at that location. A, B, and C represent 3 different INU fields added to the ground truth of the BrainWeb phantom, applied to the specified % level.
Fig. 5. Results from comparison of image features on the various Colin27 phantoms (see Table III for parameter settings). Mean value for each measure is scaled with respect to the mean value from Phantom 1. After subtraction of unity, deviations from 0 indicate sensitivity of the measurements to (A) changes in TR/TE parameters, (B) increasing levels of white additive Gaussian noise, and (C) the addition of various INU fields to the baseline phantom.

_intensity“) and after the global and local affine alignment to the target (“processed intensity”) as well as the mean VOI displacement (“determinant”), after nonlinear registration and determinant computation. For example, for the intensity, labeled int:

\[ \text{mean int} = \frac{1}{n} \sum_{i=1}^{n} \text{int}_i \]  

where \( \text{int}_i \) represents the intensity at voxel \( i \) within the VOI.

The total grey and white matter volumes \( v \) within the VOI was also assessed, following tissue classification of the locally aligned brain [60], for example, for GM

\[ v_{GM} = \sum_{i=1}^{n} \text{GM}_i \]  

where \( \text{GM}_i \) represents if voxel \( i \) has been classified as grey matter.

For each of these measures, its ratio with the base image (Phantom 1, most prevalent protocol in datasets) was calculated. Subtracting unity, any variation from 0 indicates sensitivity of the measurement to the variation in acquisition parameters. In some cases, the standard deviation ratio was also calculated in a similar fashion.

B. Experiments

Sensitivity of intensity, determinant and GM/WM volumes to changes in acquisition parameters were assessed by calculating the mean and standard deviation measures on each phantom and comparing them in turn to Phantom 1: mean variations due to TR/TE changes [Phantoms 1, 2, and 3; Fig. 5(a)]; given fixed TR/TE, mean variations due to 5%, 10%, and 20% noise [Phantoms 4, 5 and 6; Fig. 5(b)]; given fixed TR/TE, mean variations due to three example INU fields (generated automatically by the MNI BrainWeb phantom, see [61] for details) applied with 10% weight [Phantoms 7, 8, 9; Fig. 5(c)]; finally, mean and standard deviation variations when combining all factors—three TR/TE combinations, each with a different INU field at 10% weight, and 5% noise [Phantoms 10, 11, and 12; Fig. 6(a) and (b)].

C. Results

We make the following observations after analysis of these phantoms.

- **Unprocessed intensity**: mean unprocessed or native intensity within the VOI shows variations on the order of

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+3.2% to −16.2% when one changes TR/TE. For a given TR/TE combination, the impact of noise is smaller, with variations between −0.5% to −2.1%. There were variations of −1.1% to −14.8% when comparing the effect of the three different INU fields. Finally, when combining the three factors, the mean unprocessed intensity within the VOI varied from +1.6% to −19.5%, whereas its standard deviation varied from 0.7% to −17.6% with respect to that of Phantom 1.

• **Processed intensity**: After initial processing (INU correction, scaling and clamping, global and local linear alignment), the mean intensity within the VOI showed reduced variation, when compared to unprocessed results, of −29% to −0.02% when changing TR/TE parameters; sensitivity to noise was also reduced (−0.08% to 0.9%), as was the effect of INU (−0.08% to 12.6%). For all three parameters, the variation of the mean was 4.4% to 5.6%, whereas the standard deviation variation was 3.7% to 8.8%.

• **Grey and white matter**: volumes varied as well, indicating that the tissue classification process was not entirely immune to small contrast differences. Variations to TR/TE ranged from −2.9% to 0.6%; variations to noise from −0.9% to 2.6% and to INU from −15.7% to 10.6%. For all three parameters, variations ranged from −14.5% to 9.0%. There was no standard deviation on GM/WM volumes as they are single measurements for each phantom.

• **Determinant**: variations were 0.4% to 0.9% for TR/TE changes; −1.6% to 0.8% for noise variations and −9.7% to −29.7% for the different INU fields. Total variation for when all three parameters were combined was −8.7% to −10.1%. Standard deviation ratios for determinant were −4.8% to −6.1%, a 2.3% difference.

### Discussion

The results of our feature sensitivity validation on phantom data showed that the selected features (unprocessed or processed intensity, GM/WM volumes, and determinants) exhibited small to large variations when different contrasts, noise, and INU were present in the images, other variables (e.g., resolution, registration) remaining fixed.

GM/WM volume estimations were not immune to contrast differences. Whereas the contrast differences drive the classification process, one might have expected that the *a priori* spatial distribution knowledge embedded in the classification algorithm would have glossed over those differences. To be fair, this was single-modality classification (i.e., only T1-weighted images on input); it is assumed that multispectral images (e.g., with T2, PD) increase the accuracy of the classification. Finally, there was no estimation of partial volumes effects.

The mean intensity after processing (INU correction, scaling, clamping, global and local linear registration) was more robust than the unprocessed intensity. This seems to indicate that signal intensity, and possibly intensity-derived statistical or structural texture features (provided they are estimated in 3-D and are rotation-invariant), can act as valid features for ACC methodologies in the case of varying sequence parameters and acquisition irregularities. The smoothing effect on differences of the INU correction algorithm of Sled *et al.* was also highlighted in a recent multisite longitudinal study [25]. Determinant estimation showed sensitivity to INU fields but overall, displayed the most compact distribution, with nearly the smallest mean differences and the lowest standard deviation ratios of all measures. The use of geometrical information (e.g., Jacobian determinants) as opposed to intensity-based information, ensures that the technique is less sensitive to image transformations, an issue that was also mentioned by Thirion [62].

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### REFERENCES


