Subregional Basal Forebrain Atrophy in Alzheimer’s Disease: A Multicenter Study

Ingo Kilimann\textsuperscript{a,b,}\textsuperscript{*}, Michel Grothe\textsuperscript{a}, Helmut Heinsen\textsuperscript{c}, Eduardo Joaquim Lopez Alho\textsuperscript{c}, Lea Grinberg\textsuperscript{d,e}, Edson Amaro Jr.\textsuperscript{f}, Gláucia Aparecida Bento dos Santos\textsuperscript{f}, Rafael Emídio da Silva\textsuperscript{f}, Alex J. Mitchell\textsuperscript{g}, Giovanni B. Frisoni\textsuperscript{h}, Arun L.W. Bokde\textsuperscript{i}, Andreas Fellgiebel\textsuperscript{j}, Massimo Filippi\textsuperscript{k}, Harald Hampel\textsuperscript{l}, Stefan Klöppel\textsuperscript{m}, and Stefan J. Teipel\textsuperscript{a,b}

\textsuperscript{a}German Center for Neurodegenerative Diseases (DZNE) Rostock/Greifswald, Rostock, Germany \textsuperscript{b}Department of Psychosomatic Medicine, Rostock University Medical Center, Rostock, Germany \textsuperscript{c}Laboratory of Morphological Brain Research, Department of Psychiatry, Psychosomatic Medicine and Psychotherapy, University of Würzburg, Würzburg, Germany \textsuperscript{d}Department of Neurology, University of California San Francisco, San Francisco, USA \textsuperscript{e}Aging Brain Study Group, LIM-22, Department of Pathology, University of Sao Paulo Medical School, Sao Paulo, Brazil \textsuperscript{f}Department of Radiology, University of Sao Paulo, Medical School, Sao Paulo, Brazil \textsuperscript{g}Department of Psycho-oncology, University of Leicester, Leicester, UK \textsuperscript{h}LENEITEM Laboratory of Epidemiology, Neuroimaging and Telemedicine, IRCCS Centro San Giovanni di Dio, FBF, Brescia, Italy \textsuperscript{i}Cognitive Systems Group, Discipline of Psychiatry, School of Medicine and Trinity College Institute of Neuroscience (TCIN), Trinity College Dublin, Dublin, Ireland \textsuperscript{j}Department of Psychiatry, University Medical Center of Mainz, Mainz, Germany \textsuperscript{k}Neuroimaging Research Unit, Institute of Experimental Neurology, Division of Neuroscience, Scientific Institute and University “Vita-Salute” San Raffaele, Mailand, Italy \textsuperscript{l}Department of Psychiatry, Goethe University, Frankfurt, Germany \textsuperscript{m}Department of Psychiatry and Psychotherapy, Freiburg Brain Imaging, University Medical Center Freiburg, Freiburg, Germany

Abstract

Histopathological studies in Alzheimer’s disease (AD) suggest severe and region-specific neurodegeneration of the basal forebrain cholinergic system (BFCS). Here, we studied the between-center reliability and diagnostic accuracy of MRI-based BFCS volumetry in a large multicenter data set, including participants with prodromal ($n = 41$) or clinically manifest AD ($n = 134$) and 148 cognitively healthy controls. Atrophy was determined using voxel-based and region-of-interest based analyses of high-dimensionally normalized MRI scans using a newly created map of the BFCS based on postmortem in cranio MRI and histology. The AD group showed significant volume reductions of all subregions of the BFCS, which were most pronounced in the...
posterior nucleus basalis Meynert (NbM). The mild cognitive impairment-AD group showed pronounced volume reductions in the posterior NbM, but preserved volumes of anterior-medial regions. Diagnostic accuracy of posterior NbM volume was superior to hippocampus volume in both groups, despite higher multicenter variability of the BFCS measurements. The data of our study suggest that BFCS morphometry may provide an emerging biomarker in AD.

**Keywords**
Atrophy; biomarker; cholinergic system; dementia; European DTI Study on Dementia

**Introduction**
Postmortem brain studies show a significant reduction in markers of cholinergic function in the neocortex of patients with Alzheimer's disease (AD) compared to cognitively healthy controls (HC) [1–4]. A key role of cholinergic transmission in explaining cognitive impairment in AD is supported by the effect of anticholinergic pharmacological treatment on working memory and attention in HC [5]. The main cholinergic input to the cerebral cortex arises from the basal fore brain cholinergic system (BFCS) [6], consisting of the nucleus basalis Meynert (NbM) [7–9], the horizontal and vertical limb of the diagonal band of Broca [10], the nucleus subputaminalis (nucleus Ayala) [11], and a small pericommisural cluster of cholinergic perikarya (juxta-commisural cells).

Cholinergic nuclei appear to be particularly vulnerable to neurofibrillary pathology and amyloid-β accumulation [12–17]. Consistently, postmortem pathological studies of the basal forebrain in AD dementia patients have revealed a significant loss of cholinergic neurons, consistent with the cholinergic deficit in the neocortex [18]. In pre-dementia stages of AD like in mild cognitive impairment (MCI-AD), cholinergic dysfunction is present in the neocortex [19], but the role of neuron loss versus neuronal shrinkage in the BFCS is still unresolved [20].

The importance of cholinergic degeneration in AD has fueled efforts to determine the integrity of the BFCS in vivo using magnetic resonance imaging (MRI). Manual measurement of the thickness of the substantia innominata on a coronal section at the level of the anterior commissure suggests atrophy of this region in AD patients compared to HC [21–23], and this finding correlates with global cognitive impairment [23, 24]. This observation was reproduced using automated calculation of grey matter (GM) volume in a similarly defined region-of-interest (ROI) ventral to the anterior commissure [25]. However, these simple structural measures of anterior parts of the basal forebrain cover only a small part of the entire BFCS. Therefore, more detailed volumetric analysis of the BFCS has been developed based on maps of the basal forebrain nuclei in MRI standard space derived from postmortem MRI and histology data [26–28]. Using this approach, several studies reported significant reductions of BFCS volume in AD dementia compared to controls. These changes are mainly located in areas corresponding to the anterior lateral, anterior medial, and the posterior parts of the NbM, corresponding to the Ch4al, Ch4am, and Ch4p regions according to Mesulam's nomenclature [27, 29, 30].
BFSC morphometry in patients with MCI has documented atrophy in similar areas to those see in patients with AD dementia, albeit spatially more restricted [27, 31, 32]. Using a probabilistic map of the BFCS [28], it has been shown that subregional volumes of the cholinergic nuclei are differentially associated with global cognition and specific memory function in MCI patients. A significant volume reduction is found when compared to HC but appears restricted to posterior areas of the NbM [30].

In a recent single center study, the diagnostic use of BFCS volumetry appeared to be comparable to the diagnostic use of hippocampus volume, the most established structural imaging marker of AD to date [29]. Establishing BFCS volumetry as a potential future diagnostic marker requires demonstrating its stability and diagnostic accuracy in a multicenter setting. In the present study, we aimed to determine stability and diagnostic accuracy of BFCS volumetry in a large multicenter study on AD dementia and prodromal AD (MCI-AD), encompassing ten different MRI scanners across eight European sites. We used a newly created mask of BFCS subregions derived from histological sections and postmortem in cranio MRI to determine differential atrophy and diagnostic accuracy of BFCS subregions. An overview of the subregions of the BFCS mask in Montreal Neurological Institute (MNI) space can be seen in Supplementary Figure 1.

Material and Methods

Participants

Data sets of 323 participants from the European DTI Study in Dementia (EDSD) were analyzed. The EDSD is a newly established framework of nine European centers for clinical dementia research including Amsterdam (The Netherlands), Brescia (Italy), Dublin (Ireland), Frankfurt (Germany), Freiburg (Germany), Milano (Italy), Mainz (Germany), Munich (Germany), and Rostock (Germany). According to the Declaration of Helsinki, written informed consent was provided by all participants or their representatives and the study was approved by local ethics committees at each of the participating centers. Anatomical MRI scans from 134 patients with a diagnosis of clinically probable AD according to NINCDS-ADRDA criteria and 148 elderly cognitively HC were derived from eight centers, representing ten different MRI scanners (two center with two different MRI scanners) [33]. Additionally, 41 participants (from three centers) with MCI-AD were included in the study. We employed a scanner-matched subset of the cognitively HC for the AD group analysis (HC I; n = 138) and a scanner- and aged-matched subset (HC II; n = 42) for the MCI-AD group analysis. The number of participants per scanner varied from 13 to 46 with a median of 29. MCI-AD participants met the Petersen criteria for MCI and had objective cognitive impairment (above –1.5 standard deviation in the age and education years controlled Consortium to Establish a Registry of Alzheimer’s Disease (CERAD) testing battery) without dementia and either pathological levels of tau, phospho-tau, or amyloid-β in cerebrospinal fluid (CSF) analysis [34]. 26 participants of this group fulfilled criteria for the diagnosis of MCI due to AD with a high degree of certainty according to the National Institute on Aging-Alzheimer’s Association (NIA-AA) workgroups on diagnostic guidelines for AD (increased CSF levels of total-tau and phospho-tau levels and decreased amyloid-(3 1-42 levels) [35]. The remaining 15 participants fulfilling core clinical criteria of
MCI according to the NIA-AA workgroups on diagnostic guidelines for AD [35] and had either increased total-tau and increased phospho-tau and normal or decreased levels of amyloid-β 1-42 and normal levels of total-tau and phospho-tau. Beside the diagnosis of AD or MCI-AD, participants were free of any signs of neurological, psychiatric, or any medical conditions that may have interfered with cognition. In particular, there were no clinical or neuroimaging signs for cerebrovascular apoplexy, depression, or drug abuse and no clinical or blood testing signs for hypothyroidism.

The Mini-Mental State Exam (MMSE) and CERAD cognitive battery were used to assess the level of cognitive functioning [36, 37]. HC were required to score within one standard deviation of age and education adjusted norm in all subtest of the CERAD battery. Demographic characteristics are shown in Table 1. AD patients had significantly less years of education and were older than the control group participants. As it was to be expected, MMSE scores were significantly lower in patients with AD compared to HC.

MCI-AD participants scored significantly less in MMSE compared to HC. Age, years of education, and gender did not significantly differ between groups.

Postmortem Brain

The delineation and identification of the BFCS nuclei and its subregions followed the Mesulam nomenclature [38]. The transfer to MRI reference space was based on histological serial staining of a postmortem brain of a non-demented 56-year old man who died from myocardial infarction. Clinicofunctional structured evaluation was obtained from an informant by a trained nurse using a standardized and validated questionnaire. The review of the data involved a behavioral neurologist and two geriatricians and revealed no cognitive decline or psychiatric illnesses prior to death [39]. After scanning the brain in cranio (15 h postmortem) on a 3T MRI scanner (Philips Achieva) at the Radiologic Institute of Sao Paulo Medical School, the brain was removed from the skull and stored in formalin fixation (1 : 9) for three months. A second scan was done after this time period and did not reveal any pathological findings.

After the second scan was performed, the brain was stored in upgrading series of ethanol solutions. A third MRI scan was performed on the dehydrated brain on a 3T MRI scanner (Siemens Magnetom Verio). Afterwards the rostral parts anterior to the anterior horn and the dorsal parts of the frontal/parietal lobes of the brain were severed and the remaining parts of both hemispheres with the attached brainstem and the cerebellum were soaked in an 8% solution of celloidin. The tissue block was subjected to a slight vacuum in order to accelerate the concentration of celloidin from 8% to 16%. Chloroform vapors were used to harden the celloidin block before immersing it into a solution of 70% alcohol. Serial cutting in the coronal plane of the block was done at a thickness of 400 μm using a sliding microtome (Polycut, Cambridge Instruments, UK). Digital pictures of each newly appearing block surface were taken during sliding process using a digital single lens-reflex camera. More details on this process are described elsewhere [40, 41]. To identify and delineate cholinergic perikarya, celloidin sections were subsequently stained by a modified galloxyanin technique [42].
All parts of this procedure were approved by the Ethics committees of the Medical School of the University of Sao Paulo, Sao Paulo, Brazil, and the Medical Faculty of the University of Würzburg, Würzburg, Germany.

MRI Acquisition in vivo

The MRI acquisition was part of the European DTI Study on Dementia (EDSD). Data were obtained from ten different MRI scanners, including seven 3.0 Tesla and three 1.5 Tesla scanners. A sagittal high-resolution 3-dimensional gradient-echo sequence was performed for the anatomical studies with an isotropic spatial resolution of 1 mm³. In addition axial 2-dimensional T2-weighted sequences were performed to identify white matter lesions (fluid attenuation inversion recovery (FLAIR)). The exact imaging parameters for each scanner are described in Supplementary Table 1.

Postmortem MRI Scans

The postmortem scan in cranio and the scan of the formalin-fixed brain were performed on a 3.0 T MRI scanner (Achieva, Philips Medical Systems, Best, The Netherlands) using an eight channel head coil. For the anatomic study a sagittal 3-dimensional T1 Fast field echo (FFE) sequence with a spatial resolution of 1.0 × 1.0 × 1.0 mm³ (Field of view (FOV) 240 mm) was used (repetition time 7 ms, echo time 3.2 ms, flip angle 8°, number of slices 180).

The scan of the dehydrated brain was performed on a 3.0 T MR scanner (Magnetom Verio, Siemens Medical Solutions, Erlangen, Germany) using a sagittal high resolution, 3-dimensional gradient-echo sequence with a spatial resolution of 1.0 × 1.0 × 1.0 mm³ (MPRAGE, repetition time 2500 ms, echo time 4.81 ms, inversion time 1100 ms, flip angle 8°, number of slices 200). The brain was placed in an eight channel knee coil for scanning.

MRI Data Processing

Postmortem mapping of BFCS nuclei—The BFCS map was processed as shown in Fig. 1. Most parts of the magnocellular nuclei of the BFCS are located in the substantia innominata, ventral to the anterior commissure. The ventral pallidum and parts of the nucleus accumbens are located in the rostral substantia innominata. The caudal parts of the substantia innominata are occupied by the extended amygdala. We used the Mesulam nomenclature for subregions of the BFCS [6, 43]. The nuclei of the BFCS and their subregions were identified from the digital images of the histological sections and manually transferred into the corresponding slices of the MRI scan of the dehydrated brain. The MRI scan of the dehydrated brain was then transformed into the space of the postmortem in cranio scan using an initial 12-parameter affine transformation followed by a high-dimensional nonlinear registration [44] implemented in SPM8 software (Wellcome Trust Center for Neuroimaging). Next, the postmortem in cranio MRI was transferred into MNI standard space using the high-dimensional DARTEL (Diffeomorphic Anatomic Registration using Exponentiated Lie algebra) registration method [45]. The linear and non-linear transformations from alcohol through in cranio to MNI space were combined to spatially transform the basal forebrain mask into the MNI standard space. We used this map (i) to relate the voxel-wise group effects in local GM reduction to the anatomical position of the...
BFCS nuclei in MNI space and (ii) to automatically extract individual GM volumes for each BFCS subregion.

**In vivo MRI data processing**—For spatial normalization, the VBM8 toolbox (Version 414, http://dbm.neuro.uni-jena.de/vbm8/) implemented in SPM8 was used to create a customized DARTEL template. For better accuracy and quality of segmentation and to avoid favoring of one specific scanner in normalization and segmentation, we used a random subsample of $n = 54$ images, with the restriction to include scans from exactly 3 AD patients and 3 HC from 9 scanners [46]. The template was used for high-dimensional DARTEL normalization of the MPRAGE scans as implemented in VBM8. The images were segmented into GM and white matter and transformed to MNI space applying modulation for non-linear components only, thus accounting for global differences in head size and shape at this processing step. The GM segments were smoothed with a 4mm FWHM kernel for voxel-based analysis, considering the small size of the BFCS and previous experience in independent samples [27, 29, 30].

**Volumes of BFCS nuclei and hippocampus**—Individual GM volumes of the BFCS nuclei and the hippocampus were extracted automatically from the warped unsmoothed GM segments by summing up the modulated GM voxel values within ROI masks of the BFCS nuclei and the hippocampus, respectively.

The ROI mask for the hippocampus in MNI space was obtained by manual delineation of the hippocampus in the customized DARTEL template using the interactive software package Display (McConnell Brain Imaging Centre at the Montreal Neurological Institute) and a previously described protocol for segmentation of the medial temporal lobe [47].

**Statistical analysis**

We used voxel-based morphometry implemented in SPM8 to calculate the linear effect of diagnosis on voxel-wise volume. The search region was restricted to the BFCS mask and a FDR (false discovery rate) corrected statistical threshold of $p < 0.05$ and a cluster extension threshold of 5 voxels were applied.

In the ROI analysis, we determined the overall effect of BFCS atrophy using repeated measures ANOVA with diagnosis as between participants factor, age, gender, and years of education as covariates, scanner as co-factor and BFCS subregions as within participants factor. A significant overall effect of diagnosis was followed-up by pairwise linear models for each subregion controlling for age, education, gender, and scanner. The same parameters were applied to the comparison of hippocampus volumes between groups. We used receiver operating characteristics (ROC) analysis to determine the diagnostic accuracy of hippocampus and subregional BFCS volumes and compared the areas under the curves (AUCs) of different BFCS subregions and the hippocampus using a univariate binominal model implemented in ROCKIT software (Kurt Rossmann Laboratories, Chicago, Illinois) [48].
We employed variance component analysis to determine the effect of scanner on GM volumes. We determined the proportion of variance attributable to scanner relative to the variance attributable to scanner and error combined.

The ROI-based analyses were performed using SPSS 15.0 (IBM SPSS, Chicago, IL, USA), the variance component analysis was calculated using Proc Varcomp in SAS 9.2 (SAS, Cary, NC, USA).

Results

AD dementia participants

Using voxel-based analysis, we found significantly reduced volume in AD affecting virtually all cholinergic nuclei of the BFCS, including the vertical and horizontal limbs of the diagonal band, anterior medial and lateral, intermediate and posterior parts of the NbM as well as the nucleus subputaminalis (Fig. 2A). These findings remained significant after controlling for age, years of education, and gender as covariate and scanner as factor.

In the complementary ROI analysis, repeated measures ANOVA, controlling for age, gender, and scanner, revealed a significant overall effect of diagnosis across BFCS subregions \( F_{1}^{251} = 161, p < 0.0001 \). Pairwise follow-up analysis, controlling for age, years of education, gender, and scanner, showed volume reductions in patients with AD compared to controls in the hippocampus and in all subregions of the BFCS at \( p < 0.001 \) (Fig. 3A).

The areas under the ROC curves (AUC) differed significantly between subregions of the BFCS. The region projecting to localization of Ch4p showed the highest value with 0.88 (95% confidence interval (CI): 0.84–0.92) followed by the nucleus subputaminalis (0.80, 95% CI: 0.75–0.85) and the Ch3 (0.75, 95% CI: 0.70–0.81). The region corresponding to the localization of Ch2 showed the smallest AUC with a value of 0.70 (95% CI: 0.63–0.76). By contrast the hippocampal volume (bilaterally) yielded an AUC of 0.84 (95% CI: 0.78–0.88). Compared to the BFCS subregion with the highest AUC (Ch4p), AUCs of the other BFCS sub-regions as well as the hippocampus were significantly smaller \( p < 0.001 \) for the BFCS subregions; \( p = 0.005 \) for hippocampus).

The contribution of scanner to the variance of the data was estimated as \( \frac{v_{\text{scanner}}}{v_{\text{scanner}} + v_{\text{error}}} \), with \( v_{c} \) = variance component, and ranged from 0.17 for Ch4p to 0.49 for the region projecting to the localization of Ch2. For comparison, the contribution of scanner to the variance of hippocampus volume was 0.08 for the left and 0.11 for the right hippocampus in this model.

MCI-AD participants

Voxel-based analysis showed significant clusters of reduced volume in posterior and anterior lateral parts of the BFCS, whereas anterior-medial regions appeared relatively spared (Fig. 2B).
Complimentary ROI analysis revealed statistically significant volume reductions of all BFCS subregions with the exception of the region projecting to the localization of the Ch2 nucleus and the juxta-commisural cell cluster (Fig. 3B). Repeated measurements ANOVA controlled for age, gender, education, and scanner showed a significant overall effect for diagnosis ($F_{1}^{82} = 28, p < 0.0001$).

ROC analysis for the separation between MCI-AD and HC revealed a similar pattern of subregion-specific AUC values as for the classification of AD patients. The region corresponding to the Ch4p nucleus showed the highest value with an AUC of 0.86 (95% CI: 0.77–0.93), followed by N. subputaminalis with an AUC of 0.69 (95% CI: 0.57–0.79) and Ch4am_al (0.68; 95% CI: 0.56–0.78). All other regions showed only modest group separation with AUC values of 0.63 or below. Pair-wise comparisons showed that AUC of Ch4p was significantly higher than AUC of the other BFCS subregions ($p < 0.001$ for all subregions, except Ch4am_al; $p = 0.002$). The diagnostic accuracy of bilateral hippocampus volume (AUC = 0.77, 95% CI: 0.66–0.86) was lower than the value of Ch4p region. This difference was statistically significant ($p = 0.03$).

Discussion

In the present study, we examined atrophy of the BFCS in patients with AD using a newly created sub region specific map of the BFCS, based on postmortem MRI in cranio, in a large multicenter data set. In accordance with previous monocentric studies, we found significant volume reductions of all areas projecting to the location of BFCS subnuclei in patients with preclinical and clinical manifest AD compared to HC, which was most pronounced in the posterior NbM [27, 29, 30, 32]. In the pre-dementia stage of AD, the area corresponding to the location of Ch4p showed a similarly high volume reduction compared to age-matched controls, whereas the region of the anterior Ch2 nucleus was relatively spared. The diagnostic accuracy of basal forebrain volumetry measured by ROC analysis varied among subregions and yielded AUCs between 0.88 for Ch4p and 0.70 for Ch2 in the AD group. ROC based classification of the MCI-AD group showed a similar pattern of region-specific AUC values; group separation being most accurate for Ch4p region. The diagnostic accuracy of Ch4p was significantly higher compared to the other BFCS subregions in both diagnostic groups, and even compared favorably with the diagnostic accuracy of the hippocampus for classification of AD patients. Although the multicenter effect was stronger on the BFCS than on the hippocampus volumes, our data support the hypothesis that atrophy in the BFCS is an important morphologic biomarker in AD.

The Ch4p region is part of the isodendritic core, a network of interconnected nuclei affected in the very early AD stages [49]. Although profound loss of cholinergic neurons in the basal forebrain is a consistent finding in postmortem studies of end stage AD patients [3, 16, 50, 51], the underlying mechanism of regional BFCS atrophy in pre-clinical or mild clinical stages of AD is not resolved. In support of an ongoing neurodegenerative processes of the BFCS in pre-dementia stages of AD, BF cholinergic cells in MCI have been found to exhibit significant neurofibrillary pathology [20] as well as a reduction in trophic support due to a selective loss of receptors for the nerve growth factor [52–54]. Autopsy studies on persons who died in pre-dementia stages of AD (MCI) or mild AD stages are inconsistent in regard...
to atrophy of the BFCS. For example, choline acetyltransferase immunoreactivity in basal forebrain neurons and the vesicular acetylcholine transporter are not significantly reduced in MCI and mild dementia when compared to HC [55]. However, the number of participants was small ($n = 7$ MCI) and dystrophic cell shrinkage, which may be indicative of a neurodegenerative process prior to definite neuronal death, was not assessed. DeKosky et al. found elevated levels of choline acetyltransferase in the frontal cortex and hippocampus in 14 patients with MCI compared to HC [56]. Indeed, both a post-mortem study on the size of the Golgi apparatus in cholinergic neurons in MCI as well as a recent in vivo FDG-PET study on basal forebrain metabolism suggested an increase of metabolic activity in the BFCS in pre-dementia stages of AD [57, 58], possibly reflecting a compensatory adaptation.

George and colleagues employed a manual delineation of the substantia innominata in their MRI study with 19 participants with mild AD, 33 with MCI, and 27 elderly controls. They found a volume reduction of the substantia innominata in the AD group compared to MCI and controls but no significant differences between the MCI group and the control group [59]. Another study with 26 MCI, 12 AD patients, and 46 controls using manual tracing of the substantia innominata did show significant reduction in volume in the AD and MCI group compared to controls [60]. However, it is difficult for in vivo studies on the BFCS to offer a precise delineation of the complex structure of the scattered cholinergic cell groups in the absence of significant landmarks. Advanced imaging techniques using deformation-based morphometry in combination with cytoarchitectonic maps of the BFCS, based on dedicated histological examinations and postmortem MRI, have begun to provide access to the detailed study of the BFCS in vivo. Using these techniques, changes in subregions of the BFCS in pre-dementia and dementia stages of AD have been assessed using a single-person map derived from postmortem histological staining and MRI-based localization of the cholinergic fore brain [27] as well as using a probabilistic map of the magnocellular nuclei of the BFCS based on pooled data from 10 brain specimens [28, 30]. Studies using these maps demonstrated volume reductions in the BFCS both in AD and MCI, with the strongest and earliest effects located in the Ch4p region [27, 29, 30, 32]. However, limitations of these previous maps include the lack of subregional differentiation in the first map and the deviation from the Mesulam nomenclature for the identification of the BFCS subregions in the probabilistic map. A further limitation common to both maps is the reference to MRI of formalin fixed brains, which limits the accuracy of transformation of MRI data into MNI standard space. Therefore, for better differentiation of subregions of the BFCS, in this study we generated a new map based on postmortem histological staining, identification of subregions following the Mesulam nomenclature and high-dimensional transformation of BFCS labels into MNI standard space via in cranio postmortem MRI.

**Volumetric change: Subregional differences**

Our finding that the regions projecting to the localization of Ch4p shows the strongest changes in AD fits well with similar findings in independent samples of MCI patients [29, 30] and with neuropathological studies which showed that the posterior NbM region suffers the most severe neuronal loss in AD [20, 61]. Consistently, in MCI-AD participants, the volume reduction of the Ch4 region was more prominent in posterior parts, preserving anterior-medial regions of the NbM. Interestingly, cholinergic cells in the posterior NbM
subdivision project mainly to the temporal neocortex [6], which also shows the most prominent reduction in cortical cholinergic activity in AD [62].

Little is known about the function of the nucleus subputaminalis (NSP, Ayala's nucleus) and even less about its role in AD. The anatomic position of the NSP close to the external capsule and in proximity to cortical speech areas, the strong accentuation of this structure in the left hemisphere, and the fact that it has only been described in humans and anthropoid monkeys, but not in other species, gave rise to the hypothesis that the NSP might play an important role in speech processing [63]. In the future, our BFCS mask will provide the possibility to investigate a potential role of NSP changes for language impairment in AD and other neurodegenerative diseases.

The Ch3 region of the basal forebrain is located at the horizontal limb of the diagonal band of Broca, provides the major cholinergic source for the olfactory bulb [6], and is early and severely affected by AD [64]. These findings agree with the clinical observation that loss of smell is an early symptom in AD [65]. Our finding of significant volume reduction in the region projecting to the localization of Ch3 in patients with AD and MCI-AD compared to controls agrees with previous morphometric studies in independent samples [27, 30].

The Ch2 region corresponds to the vertical limb of the diagonal band of Broca and the cholinergic neurons project mainly to the hippocampus—similar to the Ch1 neurons—but also to cortical midline areas [6, 66]. Neuropathological examinations have shown that this region is the less affected compared to the more posterior BFCS nuclei in AD [16, 67], which is consistent with our *in vivo* findings in this and previous studies [29].

**Diagnostic accuracy**

The AUCs for the diagnostic separation of AD and HC ranged from 0.57 to 0.88. The highest group separation was achieved by using the volume of the Ch4p region both in AD and MCI-AD participants. The diagnostic accuracy of this region was significantly higher than the accuracy of hippocampus volumetry in both groups. To the best of our knowledge, there is no other study of the BFCS using subregional analysis of group discrimination. In monocentric MRI data from the Open Access Series of Imaging Studies (OASIS, [http://www.oasis-brains.org/](http://www.oasis-brains.org/)), a non-subregional map of the BFCS as a whole [27] reached a diagnostic accuracy of 0.81 AUC for the discrimination between 28 patients with AD and 96 HC, which was identical to the diagnostic value of hippocampus volume. However, in a sample of 69 participants in a pre-dementia stage of AD, the non-subregional map of the BFCS only achieved an AUC of 0.69 for the separation from HC and was significantly inferior to the diagnostic accuracy of hippocampus volume [29].

Variance component analysis revealed a considerably stronger effect of scanner on BFCS measurements than on hippocampus measurements. This result is not surprising, given the small size of the BFCS structures. Still, the data on group differences and diagnostic accuracy suggest that at least for Ch4p and NSP, the multicenter variance is offset by a strong involvement of these brain areas in AD and its value as a diagnostic test is unlikely to be diminished.
The interpretation of the anatomical findings is limited, because the histological examination of a single brain specimen for stereotactic mapping of the BFCS nuclei does not account for inter-individual differences in BFCS anatomy. However, MR images of dehydrated brains show a marked reduction in gray/white matter contrast, which reduces the accuracy of automated segmentation and spatial normalization procedures. Therefore, spatial variation in the location of stereotactically mapped BFCS regions across individuals may also reflect inter-individual differences in spatial normalization accuracy, in addition to anatomic variability. Although limited to a single brain specimen, in the present study we aimed to enhance the quality of the stereotactic mapping from histology into MNI space using a postmortem MRI in cranio, which showed image characteristics comparable to in vivo scans. This intermediate normalization step via the in cranio scan of the same subject allows high-dimensional spatial normalization from native to MNI standard space using the highly efficient DARTEL algorithm [45, 68]. This may significantly improve the spatial projection of the native localization of BFCS nuclei into MNI standard space when compared to the direct spatial normalization of the alcohol-fixed brain scans that has been used in previous cytoarchitectonic mapping studies [27, 28].

To determine the convergence of the location of the cholinergic space of the BF in MNI standard space across different processing pipelines and data sources, we compared our mask with a previously published probabilistic mask of the BFCS and found a very good spatial agreement between the masks, i.e., all center of gravity coordinates for the different BFCS subdivisions reported in [28] fell into the cholinergic space as defined by our mask, with exception of the coordinate for right Ch4p which was located slightly lateral of the space defined by our right Ch4p label.

In summary, we found significant and differential atrophy of BFCS subregions in AD and MCI-AD, most notable involving posterior parts of the NbM (Ch4p). Compared to AD dementia, the anterior-medial regions of the NbM were more preserved in MCI-AD. Diagnostic accuracy of BFCS subregional volumes was comparable to hippocampus volumetry for the most severely affected BFCS regions, despite higher multicenter variability. These data suggest that BFCS morphometry may provide an emerging biomarker in AD that may help in studying underlying neurobiological changes of specific cognitive impairments.

**Supplementary Material**

Refer to Web version on PubMed Central for supplementary material.

**Acknowledgments**

HH was supported by a research grant of the Katharina-Hardt-Foundation, Bad Homburg, Germany. ALWB has received funding from Science Foundation Ireland through the Stokes Programme. Dr. Grinberg is funded by the NIH (R01AG040311).

**References**


_J Alzheimers Dis._ Author manuscript; available in PMC 2015 January 01.


J Alzheimers Dis. Author manuscript; available in PMC 2015 January 01.


J Alzheimers Dis. Author manuscript; available in PMC 2015 January 01.


J Alzheimers Dis. Author manuscript; available in PMC 2015 January 01.


Fig. 1.
Postmortem MRI processing. Identification of the basal forebrain cholinergic system subregions. The upper part of the figure shows the postmortem processing with MRI scans of the brain in cranio and after formalin fixation. Afterwards the subregions were identified by histological staining and manually transferred into the MRI of the dehydrated brain in the alcohol space (color-coded mask at the lowest row of the figure). MRI scans were transferred first into the MRI space of the in cranio scan and then into MNI standard space. The combined transformation matrices were used to transfer the BFCS mask from alcohol space to MNI space via MRI in cranio space.
Fig. 2.
Volumetric changes projected onto the BFCS mask. Coronal section through the basal forebrain show the mask of the cholinergic basal forebrain in MNI space. The numbers on multislice line on top show the y-coordinate in MNI space (anterior to posterior position of the section). Red voxels show significant volume reduction in AD/MCI-AD group compared to HC (bright red areas show high T-values). Green voxels represent subregions or parts of subregions without significant reduction. (A: AD dementia compared to HC I; B: MCI-AD compared to HC II; $p < 0.05$ FDR corrected).
Fig. 3.
Bar diagram of BFCS and hippocampus volumes between AD and HC groups. Mean value of hippocampus and BFCS volumes in HC and AD/MCI-AD with error bars indicating standard deviation (data adjusted for head size). (A: AD dementia compared to HC I; B: MCI-AD compared to HC II).
Fig. 4.
ROC curves of BFCS subregions and hippocampus. ROC curves of the different subregions (left) and comparison of Ch4p and hippocampus volumes (right). (A: AD dementia compared to HC I; B: MCI-AD compared to HC II).
### Table 1

#### Participants' characteristics

<table>
<thead>
<tr>
<th></th>
<th>AD</th>
<th>HC I</th>
<th>MCI</th>
<th>HC II</th>
</tr>
</thead>
<tbody>
<tr>
<td>No. of participants (women)</td>
<td>134 (76)(^1)</td>
<td>138 (69)</td>
<td>41 (17)(^5)</td>
<td>42 (18)</td>
</tr>
<tr>
<td>Age (SD) in years</td>
<td>72.5 (8.3)(^2)</td>
<td>69.2 (5.9)</td>
<td>70.5 (6.4)(^6)</td>
<td>70.0 (5.2)</td>
</tr>
<tr>
<td>MMSE (SD)</td>
<td>20.7 (5.4)(^3)</td>
<td>28.8 (1.1)</td>
<td>25.7 (4.4)(^7)</td>
<td>28.7 (1.1)</td>
</tr>
<tr>
<td>Years of education (SD)</td>
<td>10.2 (3.4)(^4)</td>
<td>13.1 (3.8)</td>
<td>12.1 (2.9)(^8)</td>
<td>13.0 (3.1)</td>
</tr>
</tbody>
</table>

AD, Alzheimer's disease; HC I, cognitively healthy control group I (matched to AD group); MCI, (mild cognitive impairment); HC II, cognitively healthy control group II (matched to MCI group); MMSE, Mini-Mental State Exam;

1 not significantly different between groups, \(\chi^2 = 1.23, 1 \text{ df}, p = 0.27;\)

2 significantly different between groups, \(T = 3.81, 270 \text{ df}, p < 0.001;\)

3 significantly different between groups, Mann-Whitney-U=263, \(p < 0.001;\)

4 significantly different between groups, \(T = -6.7, 270 \text{ df}, p < 0.001;\)

5 not significantly different between groups, \(\chi^2 = 0.017, 1 \text{ df}, p = 0.90;\)

6 not significantly different between groups, \(T= -0.40, 81 \text{ df}, p = 0.83;\)

7 significantly different between groups, Mann-Whitney-U = 170, \(p < 0.001;\)

8 not significantly different between groups, \(T = 1.25, 81 \text{ df}, p = 0.215.\)