Latent Class and Transition Analysis of Alzheimer’s Disease Data

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This study uses independent latent class analysis (LCA) and latent transition analysis (LTA) to explore accurate diagnosis and disease status change of a big Alzheimer’s disease Neuroimaging Initiative (ADNI) data of 2,132 individuals over a 3-year period. The data includes clinical and neural measures of controls (CN), individuals with subjective memory complaints (SMC), early-onset mild cognitive impairment (EMCI), late-onset mild cognitive impairment (LMCI), and Alzheimer’s disease (AD). LCA at each time point yielded 3 classes: Class 1 is mostly composed of individuals from CN, SMC, and EMCI groups; Class 2 represents individuals from LMCI and AD groups with improved scores on memory, clinical, and neural measures; in contrast, Class 3 represents LMCI and AD from individuals with deteriorated scores on memory, clinical, and neural measures. However, 63 individuals from Class 1 were diagnosed as AD patients. This could be misdiagnosis, as their conditional probability of belonging to Class 1 (0.65) was higher than that of Class 2 (0.27) and Class 3 (0.08). LTA results showed that individuals had a higher probability of staying in the same class over time with probability >0.90 for Class 1 and 3 and probability >0.85 for Class 2. Individuals from Class 2, however, transitioned to Class 1 from time 2 to time 3 with a probability of 0.10. Other transition probabilities were not significant. Lastly, further analysis showed that individuals in Class 2 who moved to Class 1 have different memory, clinical, and neural measures to other individuals in the same class. We acknowledge that the proposed framework is sophisticated and time-consuming. However, given the severe neurodegenerative nature of AD, we argue that clinicians should prioritize an accurate diagnosis. Our findings show that LCA can provide a more accurate prediction for classifying and identifying the progression of AD compared to traditional clinical cut-off measures on neuropsychological assessments.

Keywords: Alzheimer's disease, latent class analysis, latent transition analysis, neural markers, misdiagnosis

INTRODUCTION

The World Health Organization has identified Alzheimer’s disease (AD) as a public health priority, with ~30–35 million cases worldwide (World Health Organization, 2012). Alzheimer’s disease is a chronic neurodegenerative syndrome which causes severe progressive deterioration in cognitive impairment (Alzheimer Association, 2019). Impairments include detriments in memory, learning
ability, language, judgment, decision making, and disordered thinking (Alzheimer Association, 2019). Patients are diagnosed with AD after being assessed on multiple neuropsychological assessments, including memory, language functioning, personality, and behavioral changes. Assessments of specific biomarkers of AD are also being used to identify structural changes within specific brain regions as well as measure levels of Amyloid-β, tau, and phospho-tau (Alzheimer Association, 2019). Typically, the assessment of AD is based on clinical cut-off points for neuropsychological assessments and biomarkers. This technique allows a medical professional to identify those who have symptoms of AD. While clinical cut-offs are important for categorizing individuals with and without AD, it does not always contribute to our understanding of the progression of AD or identify individuals at risk of developing AD. Understanding the progression of AD is important to developing preventative interventions and earlier detection.

The Alzheimer's Disease Neuroimaging Initiative (ADNI) study has collected longitudinal data from more than 50 sites in North America on “…elderly individuals with normal cognition, mild cognitive impairment (MCI)” (Jack et al., 2008). In total, the ADNI project has collected data using 292 measurements (see http://adni.loni.usc.edu/data-samples/adni-data-inventory-for-a-full-list-of-items). These measurements include diagnostic assessments; neuropsychological assessments; bio-specimens; genetics; imaging—including different MRI and PET imaging techniques; demographic and medical history; and a participation record. Data were collected at 0, 6, 12, 24, and 36 months for participants in the normal cognition and mild cognitive impairment groups (Jack et al., 2008). However, the AD group's data were only recorded at 0, 6, 12, and 24 months (Jack et al., 2008). The main aims of the ADNI project are to improve early detection and track disease progression using biomarkers and advance early intervention, prevention, and treatment.

**AD Diagnosis**

The assessment and diagnosis of AD have primarily relied on cut-off scores on neuropsychological assessments. For example, the Clinical Dementia Rating Scale (Morris, 1997) can be used to categorize individuals into differing levels of severity ranging from normal cognitive functioning, questionable cognitive impairment, questionable impairment, very mild dementia, mild dementia, moderate dementia, and severe dementia (O'Bryant et al., 2008). An advantage of utilizing categories for the Clinical Dementia Rating Scale is that it reliably identifies individuals with mild cognitive impairment (Dura et al., 2013). This allows clinicians to use the results to identify patients who are suffering from differing degrees of dementia severity. However, utilizing the scale with cut-offs does not allow health professionals to track the progression of the disease or identify at-risk patients before the presentation of symptoms.

Other neuropsychological assessments such as the Functional Activities Questionnaire (Pfeffer et al., 1982), the Alzheimer’s Disease Assessment Scale (Mohs and Cohen, 1988), Clinical Dementia Rating Scale (Morris, 1997), Everyday Cognition Scale (Marshall et al., 2014), Montreal Cognitive Assessment (Nasreddine et al., 2005), the Mini-Mental State Exam (Folstein et al., 1975), and the Cognitive Change Index (Saykin et al., 2006; Rattanabannakit et al., 2016) have also been used to categorize cognitive impairment and AD. For instance, within the ADNI, participants are classified with AD if they obtain a score between 20 and 26 on the mini-mental state examination; a score between 0.5 and 1.0 on the global clinical dementia rating, a score between 1.0 and 9.0 for the summed box-score for the clinical dementia rating (Shaw et al., 2009). These standardized assessments are useful for diagnosing probable AD, with most yielding good sensitivity, specificity, and classification scores. That is, they reliably distinguish between individuals with mild cognitive impairment and AD—making them good diagnostic tools. However, utilizing these techniques is only useful for determining probable AD. The use of cognitive assessments only allows for a measure of current cognitive function and does not indicate if an individual may progress from mild cognitive impairment to severe cognitive impairment or AD. In the absence of objective diagnostic assessments for AD, a positive diagnosis is currently only determined through an autopsy (Perrin et al., 2009; Shaw et al., 2009). Recent advances in imaging techniques (i.e., MRI and PET) and acquiring cerebral spinal fluid have allowed researchers to identify potential biomarkers of AD and what structural changes occur within specific brain regions (e.g., hippocampus).

Shaw et al. (2009) collected cerebral spinal fluid from elderly individuals with normal cognitive functioning, mild cognitive impairment, and mild AD (classification were determined using the mini-mental state examination and the Alzheimer’s Disease Assessment Scale). The levels of Amyloid-β 1 to 42 peptide (Aβ 1−42), total tau (t-tau), and tau phosphorylated (p-tau) were assessed to determine potential biomarkers of AD. To gain more accurate cut-off points, models of the levels of Aβ 1−42, t-tau, and p-tau were determined from cerebral spinal fluid samples from autopsy-confirmed AD cases. The results indicated that Aβ 1−42 showed excellent sensitivity (96.4%) and specificity (76.9%) with a clinical cut-off of 192 pg/ml; t-Tau showed acceptable sensitivity (69.6%) and excellent specificity (92.3%) with a clinical cut-off of 93 pg/ml, and p-tau showed acceptable sensitivity (67.9%) and specificity (73.1%) with a clinical cut-off of 23 pg/ml. Further, the interaction between decreasing levels of Aβ and increasing levels of p-tau have recently been implicated with neuronal death, atrophy, and cognitive changes (Gomar et al., 2016; Veitch et al., 2019). These results suggest that Aβ 1−42 and p-tau are the most sensitive measures and best predictors of early diagnoses of AD.

The diagnosis of probable AD can also be assessed by measuring specific biomarkers (i.e., Aβ, t-tau, & p-tau). However, similar to the use of neuropsychological assessments, diagnosis relies on patients exceeding a clinical threshold for the levels of each biomarker. While biomarkers of AD appear to reliably distinguish between those diagnosed with (i.e., sensitivity) and without AD (i.e., specificity), some of the measures are still below the recommended threshold of 85% for sensitivity and specificity (Ronald and National Institute on Aging Working Group, 1998; Frank et al., 2003; Shaw et al., 2009). Again, the use of clinical cut-offs only provides clinicians with a measure to differentiate between mild cognitive impairment and probable AD based on particular biomarkers. Therefore, the use of cut-off
scores is essential for diagnosis but does not identify at-risk patients or to accurately track the progression of AD from mild cognitive impairment to pre-clinical AD, probable AD, and a final diagnosis of AD.

To promote the early detection of AD and to possibly identify at-risk individuals, research should not solely rely on clinical cut-off points, which are only useful once an individual presents with neuropsychological symptoms or biomarkers associated with probable AD. Secondly, there are criticisms of using cut-off points on continuous neuropsychological assessments because patients on either side of the cut-off are likely similar (Berlin et al., 2014; Petersen et al., 2019).

**Latent Class Analysis**

Instead, Latent Class Analysis (LCA) can be used to identify homogeneous subgroups of individuals who are externally heterogeneous to other sub-groups (Berlin et al., 2014; Eppig et al., 2017; Mooney et al., 2018; Petersen et al., 2019; Villeneuve et al., 2019; Zammit et al., 2019a). Latent class analysis can be used to identify homogenous subgroups of AD based on psychological assessments (e.g., Scheltens et al., 2016; Eppig et al., 2017; Zammit et al., 2019b). For example, Scheltens et al. (2016) identified eight cognitive subtypes of AD within their sample of probable AD patients (N = 938). The cognitive subtypes included patients with mild-memory impairment, moderate memory impairment, mild-visual/spatial-language impairment; moderate-visual/spatial impairment, mild-executive functioning impairment, moderate diffuse (cognitive impairment), and severe-diffuse (cognitive impairment). The authors suggest that the identification of cognitive subtypes highlights that AD is a heterogeneous to other sub-groups (homogeneous subgroups of individuals who are externally heterogeneous to other sub-groups). Instead, Latent Class Analysis (LCA) can be used to identify homogenous subgroups of AD based on psychological assessments (e.g., Scheltens et al., 2016; Eppig et al., 2017; Zammit et al., 2019b).

Zammit et al. (2019b) also used LCA to identify cognitive subtypes of AD within participants from the Rush Memory and Aging Project. Participants included in their study had no dementia at baseline; displayed signs of dementia at follow-up; were deceased at the time of the study, and had neuropathological data available. Neuropathological data were obtained from autopsies. Based on the neuropsychological outcomes at baseline (i.e., Episodic-, Semantic-, working-, and logical-memory; perceptual-and line orientation; and Perceptual Speed–Symbol Digits Modalities Test) latent class analysis was used to categories participants into 5 classes within two categories (i.e., impaired cognition and intact cognition). The impaired cognition classes included participants with mixed-domains impairment, memory-specific impairment, and frontal impairment. The intact cognition classes included participants with average cognition and superior cognition.

The aim of Zammit’s (2019b) study was to identify if neuropsychopathological evaluations at autopsy (i.e., Aβ, tau, hippocampal sclerosis, DNA-binding protein 43, Levy bodies, cerebral amyloid angiopathy, atherosclerosis, and arteriolosclerosis) were predicted by the five classes of cognitive impairment and intact cognition at baseline and if the neuropsychopathological measures differ between each class. Their results showed that baseline measurements on neuropsychological assessments were predictive of neuropathology measured at autopsy, suggesting that neuropsychopathological assessments are reliable for the assessment and prognosis of cognitive impairments associated with AD.

One of the main findings of Zammit et al. (2019b) study was that the biomarkers Aβ and Tau are strongly predictive of AD and can possibly be used as an early detector. Indeed, abnormal levels of Aβ and Tau were strongly associated with participants within the mixed-domains class, the memory-specific class, and the frontal impairment classes. With fewer abnormalities in the average cognition class and the superior cognition class. That is, abnormal Aβ and Tau were associated with impaired cognition but not intact cognition. One of the limitations of their study was that it did not account for individuals who might change classes from baseline to follow-up. For example, participants could progress from average intact cognition to memory-specific impairment. As such, the results are only capturing the class an individual belongs to at a single point in time.

Zammit et al. (2020) extended their previous work by using latent transition analysis to identify participants within the Rush Memory and Aging Project who transitioned from non-impairment to cognitive impairment. A second aim was to compare the classification of individuals within the LTA to the clinical criteria of MCI. The results showed that across three measurements (within 12 months) cognition remained relatively stable. That is, participants did not regularly change between the five classes of impairment; identified as mixed domains impairment, memory-specific impairment, frontal Impairment, average cognition, and superior cognition. However, of the 1,924 participants, 98 individuals did change membership class from time 1 to time 2 (n = 62) and from time 2 to time 3 (n = 37). A majority of the transitions were associated with a decline in cognitive impairment at both time points. These results identified that participants who changed classes had an 86% higher risk of developing AD than those who did not change status. Further, their study identified 541 participants with cognitive impairment at time 2, 10.5% of these participants progressed to developing dementia at time 3. While a majority of older adults cognition remains stable, those who are experiencing some level of cognitive impairment have an elevated risk of progressing to developing dementia. The authors provide evidence that using LTA is a robust tool to identify individuals at risk of cognitive decline, identifying risk factors for interventions to target.

Zammit’s (2020) study was not without limitations. Specifically, their LTA only used neuropsychological measures of episodic memory, semantic memory, working memory, and perceptual speed and orientation. With evidence suggesting that neurological biomarkers are significant and sensitive predictors of early diagnoses of AD (Shaw et al., 2009; Gomar et al., 2016; Veitch et al., 2019), it is important to identify if biological markers of AD can predict cognitive impairment transitions. However, their paper does highlight that LCA and LTA are at the forefront of research aiming to improve diagnostic methods and to identify individuals at risk of progressing toward AD. Zammit et al. (2020) also note the need to validate these methods through replication of their findings, and efforts to identify homogeneous classes of cognitive impairment using other neuropsychological measures of AD. As mentioned earlier, recent studies using
LTA have not included biomarkers of AD. Our study adds a novel contribution to this emerging area by identifying if neuropsychological measures and neurological biomarkers of AD are indicators of individuals transitioning from healthy individuals to individuals with mild cognitive impairment and AD.

**The Current Study**

To our knowledge, Latent Transition Analyses (LTA) has not been used to identify the neuropsychological and biomarkers associated with the progression of AD in terms of patients transitioning from one AD class to another. In the present study rather than using a set cut off point to diagnose individuals as Alzheimer's patients, Latent Class Analysis (LCA) was used to identify individuals that are more likely to develop dementia. In addition, the focus of the analysis was on the development of the individuals over time, that is, how an individual changes class membership over time. In total, the following three research aims were addressed in this study: (1) determine and describe the number of classes that best characterize individuals with respect to clinical measures and neurological biomarkers; (2) compare the classification results obtained from the LCA and the cut-off methods, to identify the misdiagnosed individuals and characterize these patients; and (3) explore the developmental course of individuals with respect to clinical and neural measures.

Below, we first describe the ADNI dataset, which we have utilized in the current study. Second, we provide details on our latent class analysis and latent transition analysis. Following that, we present the results from both latent class analysis and latent transition analysis, respectively. Finally, we discuss our results in terms of importance of our findings and clinical implications.

**METHOD**

**ADNI Dataset**

Data used in the preparation of this article were obtained from the Alzheimer's Disease Neuroimaging Initiative (ADNI) database (adni.loni.usc.edu). The ADNI was launched in 2003 as a public-private partnership, led by Principal Investigator Michael W. Weiner, MD. The primary goal of ADNI has been to test whether serial magnetic resonance imaging (MRI), positron emission tomography (PET), other biological markers, and clinical and neuropsychological assessment can be combined to measure the progression of mild cognitive impairment (MCI) and early Alzheimer's disease (AD).

The ADNI dataset includes 2,132 participants: 512 controls, and 333 with EMCI, 621 with LMCI, and 279 with SMC, and 367 AD patients. All participants were tested at 3 different times annually. In all participants, ADNI dataset includes the following measures for all participants: APOE4 = Apolipoprotein E4 gene; FDG = Fluorodeoxyglucose CDRSB = Clinical Dementia Rating Sum of Boxes; ADAS11 = Alzheimer’s Disease Assessment Scale (Cognitive Subscale), 11 item version; MMSE = Mini-Mental State Examination; RAViM = Rey Auditory Verbal Learning Test (Immediate word recall score); MOCA = The Montreal Cognitive Assessment; EcPtMm = Everyday Cognition-Participant Self Report (8 memory items); EcPtLg = Everyday Cognition-Participant Study Partner Report (9 language items); EcSPM = Everyday Cognition-Participant Study Partner Report (8 Memory items); EcSPlg = Everyday Cognition-Participant Study Partner Report (9 Language items); Hippc = Hippocampus volume; Entor = entorhinal cortex volume; Fusif = fusiform gyrus volume.

**Statistical Analysis**

A series of Latent Class Analysis (LCA), multivariate analysis of variance, and Latent Transition Analyses (LTA) were conducted. LTA is a longitudinal extension of LCA that explores changes in class membership over time by capturing individual movements in forward and backward directions across time points. This statistical method is based on Markov chain models (Kaplan, 2008) and uses an LCA model as a measurement model.

**Latent Class Analysis**

Latent Class Analysis (Lazarsfeld and Henry, 1968; Clogg, 1981) was employed to empirically identify the number of classes that best characterize individuals with respect to clinical and neural measures. LCA is a mixture model that classifies participants into optimal classes on the basis of shared characteristics that distinguish members of one class from another. Furthermore, unlike traditional cluster analysis, which is based on heuristic or distance procedures (Moustafa et al., 2018; Alashwal et al., 2019), this approach is a model-based statistical method that allows the LCA solution to be replicated with an independent sample (e.g., Nylund et al., 2007).

A commonly-used strategy to determine the optimal number of classes in LCA is to estimate a series of models by progressively increasing the number of classes and comparing the models through fit statistics and tests of significance and the quality of classification across models, as well as the usefulness and the interpretability of the latent classes (e.g., Muthén and Muthén, 2000; Vermunt and Magidson, 2002). To determine the optimal number of classes for the sample, each model was evaluated using three information criteria (IC), namely, the Akaike Information Criterion (AIC; Akaike, 1987), the Bayesian Information Criterion (BIC; Schwartz, 1978), sample size adjusted BIC (SBIC; Sclove, 1987), and the Lo-Mendell-Rubin likelihood ratio test (LMR; Lo et al., 2001). For AIC, BIC, and SBIC, a lower value indicates a better model. For the LRT, a significant p-value for a model with k classes followed by a non-significant p-value for a model with k + 1 classes indicates that the k class model is the best fitting model. The indices BIC, SBIC, and LMR have been shown to identify the appropriate number of groups within finite mixture models (e.g., Diaallo et al., 2016a,b; Diazlo et al., 2017). Furthermore, the entropy criterion was used to examine the quality of classification across models. The normalized entropy values ranged from 0 to 1 with values >0.80 representing a clear assignment of individuals to latent classes. Finally, class size was also considered when determining the optimal number of latent classes. Small classes (i.e., those that contain <5% of the sample) were considered spurious classes, as they are often associated with class over-extraction (Hipp and Bauer, 2006).
Latent Transition Analysis

LCA can be extended to accommodate longitudinal data through LTA. LTA is a type of Markov model that studies how individuals change membership in latent classes over time. LTA links LCA variables at different time points to each other using autoregressive models. A series of multinomial logistic regression, where the latent class variable at time t is regressed on the latent class variable at time t-1, is commonly used to estimate transitions over time in latent class membership.

Analytical Steps

The statistical analyses involved three steps. In the first step, we identified the optimal number of classes for each time point separately. In the second step, individuals were assigned to their most likely latent class (modal class assignment) and the latent class variable at time 1 is compared to the clinical diagnostic variable and misclassified patients are studied using multivariate analysis of variance. The third step involved exploring the developmental course of the patients with respect to clinical and neural measures. That is, transition probabilities were used to explore changes that had taken place in the latent classes. For this analysis, measurement invariance was assumed to ensure that the classes have the same meaning over time. Specifically, measurement model parameters were set to be equal over time. Hence, conditional item probabilities, item means, and item variances for the LCA were constrained to be equal at the three time points.

For this study, all models were estimated using a Full Information Maximum Likelihood (FIML) procedure available in Mplus 8.3 (Muthén and Muthén, 2019). FIML utilizes all available information during the estimation process and provides consistent and efficient population parameters (Enders, 2010). Furthermore, all LCA models with continuous indicators were estimated with residual variances of the outcomes constrained to be equal across classes and under local independence within classes assumption (i.e., indicators’ residual covariances within classes were constrained to zero). All models with two classes or more were estimated using 500 sets of random starting values, 50 iterations for each of these sets, and the 20 best sets of random starting values associated with the highest likelihood values were retained for the final optimization stage.

RESULTS

The first aim of this study was to determine the number of classes that best characterize patients with respect to clinical and neural measures. Table 1 provides an overview of patients’ characteristics with respect to clinical and neural measures. As these showed, there was substantial variability among the patients on their clinical and neural measures. This variability supports the value of using mixture methods to assess whether the patients can be grouped into different classes based on their clinical and neural measures.

Latent Class Analysis Results

Latent class models containing 1–7 classes at each time point were fitted to the data. The model fit statistics are available in Table 2. All LCA models converged at Time 1. The log-likelihood increased while no minimum was found for the ICs as their values decreased across the range of models considered. The LMR pointed to the three-class solution since the test of the two-class model against the three-class model has a p-value of 0.003, suggesting rejection, whereas the test of the three-class against the four-class has a p-value of 0.24. Further, an examination of the LCA models indicated that the four- and five-class models each included small classes that seemed to have splintered off from larger classes in the three-class model. Therefore, a three-class model was selected at time 1 based on the fit statistics (Muthén, 2004). The three-class model resulted in a log-likelihood value of −11010.52 with 60 parameters, an AIC of 22141.04, a BIC of 22481.49, a SBIC of 22290.86, and a high entropy value of 0.89. Moreover, the three-class solution satisfied the minimum class size required to be useful (each comprised at least 5% of the sample) and meaningful.

All models with fewer than seven classes converged at time 2. Consequently, only model results for classes between one and six were considered for further analysis. As in time 1, log-likelihood values increased, no minimum was found for the ICs but no solution was favored by the LRT. However, similar to the results at time 1, the results showed that the four- and five-class models each included small classes that seemed to have splintered off from larger classes in the three-class model. Hence, based on the interpretability and the usefulness of the classes, the three-class solution was also selected as the optimal number of classes at time 2. Fit indices for the three-class solution at time 2 were as follows: Log-likelihood = −6927.78, number of parameters = 60, AIC = 13975.57, BIC = 14303.75, SBIC = 14113.13, and entropy = 0.82.

Finally, all models with fewer than six classes converged at time 3, whereas models with six classes and more did not converge. Hence, only model results for classes between one and five were considered for further analysis. Similar to time 1, log-likelihood values increased, no minimum was found for the ICs whereas the LRT selected the three-class solution. Based on the interpretability and the usefulness of the classes, the three-class solution was also selected as the optimal number of classes at time 3. Fit indices for the three-class solution at time 3 were as follows: Log-likelihood = −6984.23, number of parameters = 60, AIC = 14088.45, BIC = 14415.74, SBIC = 14225.12, and entropy = 0.81.

Explanation of Latent Class Solutions

Here, we describe the latent class solutions at the three time points. Across the three time points, Class 1 showed a pattern of low means on CDRSB, ADAS11, EcPtMm, EcPtLg, EcSPM, EcSPlg, a pattern of high means on FDG, MMSE, RAVimD, MOCA, Hipc, Entor, Fusif, and selected category zero of with item probability >0.65. Class 1 is composed of 63% of the sample at time 1, 59% at time 2, and 65% at time three. In contrast, Class 3 showed a pattern of low means on FDG, MMSE, RAVimD, MOCA, Hipc, Entor, Fusif, a pattern of high means on CDRSB, ADAS11, EcPtLg, EcSPM. We, therefore, interpreted this class as the AD class. Class 3 is composed of 7% of the sample at time 1, 10% at time 2, and 6% at time three. Class 2, however, showed scores that overall were between Class 1 and Class 3. Class 2 was composed of 30% of the sample at time 1, 31% at
Descriptive statistics of the clinical and neural measures.

| Variables | Time 1 | | Time 2 | | Time 3 |
|-----------|--------|--------|--------|--------|--------|
|           | % N    | M     | SD     | % N    | M     | SD     |
| APOE4     | 1,726  |        |        | 1,210  |        |        |
| Zero      | 53.10  | 55.02  | 50.10  | 55.02  | 57.88  | 53.60  |
| One       | 37.00  | 38.26  | 38.01  | 38.26  | 40.00  | 39.20  |
| Two       | 9.90   | 8.72   | 11.89  | 8.72   | 13.12  | 17.19  |
| FDG       | 735    | 1.23   | 0.15   | 474    | 1.19   | 0.16   |
| CDRSB     | 1,990  | 1.69   | 2.39   | 1,342  | 2.28   | 2.62   |
| ADAS11    | 1,962  | 11.18  | 7.60   | 1,335  | 12.21  | 8.47   |
| MMSE      | 1,978  | 27.07  | 3.32   | 1,326  | 26.30  | 3.90   |
| RAVinD    | 1,966  | 36.33  | 13.47  | 1,340  | 32.41  | 13.56  |
| MOCA      | 1,121  | 23.70  | 4.86   | 1,329  | 22.56  | 5.19   |
| EcPlmM    | 1,144  | 1.98   | 0.72   | 537    | 2.12   | 0.73   |
| EcPlgLg   | 1,137  | 1.67   | 0.62   | 545    | 1.73   | 0.66   |
| EcSpm     | 1,135  | 1.98   | 0.97   | 557    | 2.56   | 1.03   |
| EcSplg    | 1,138  | 1.60   | 0.80   | 557    | 1.82   | 0.91   |
| Hipc      | 1,222  | 6676.92 | 1211.87 | 1,177  | 6609.89 | 1243.53 | 1,107  | 6625.67 | 1269.12 |
| Enter     | 1,170  | 3437.39 | 810.34  | 1,105  | 3397.00 | 815.24  | 1,032  | 3414.71 | 838.77  |
| Fusif     | 1,170  | 16942.23 | 2792.54 | 1,105  | 16886.15 | 2780.76 | 1,032  | 16877.08 | 2822.98 |

APOE4, Apolipoprotein E4 gene; FDG, Fluorodeoxyglucose; CDRSB, Clinical Dementia Rating Sum of Boxes; ADAS11, Alzheimer’s Disease Assessment Scale (Cognitive Subscale); 11 item version; MMSE, Mini-Mental State Examination; RAVinD, Rey Auditory Verbal Learning Test (immediate word recall score); MOCA, The Montreal Cognitive Assessment; EcPlmM, Everyday Cognition-Participant Self Report (8 memory items); EcPlgLg, Everyday Cognition-Participant Self Report (9 language items); EcSpm, Everyday Cognition-Participant Study Partner Report (8 memory items); EcSplg, Everyday Cognition-Participant Study Partner Report (9 language items); Hipc, Hippocampus volume; Enter, entorhinal cortex volume; Fusif, fusiform gyrus volume. N stands for number, and M is for mean.

Multivariate Analysis of Variance

Multivariate analysis of variance was used to compare the group for the three clinical outcome measures. The results of the multivariate analysis of variance with the 13 clinical and neural measures were between the mean of Class 1 without the 63 individuals and those from the AD individuals from the diagnoses variable without the 63 individuals (Table 4). The LCA classified the 63 individuals within Class 1 as their condition probability of belonging to this class was higher than those of belonging to the other two classes (with conditional probabilities of 0.65, 0.27, and 0.08 for Class 1, 2, 3, respectively). LTA will be used to study the development of these individuals over time.

Latent Transition Analysis Results

The LTA was conducted under the measurement invariance assumption. Consequently, measurement model parameters were set to be equal over time. Transition probabilities for the whole sample are presented in Table 5 and provide information on patient’s status at time 2 given their latent status at time 1, and patient’s status at time 3 given their latent status at time 2. The results showed most individuals stayed in the same class from time 1 to time 2 but some changes in class membership for some individuals were seen from time 2 to time 3. Individuals who were in the Class 1 at time 1 had a 0.99 probability of remaining there at time 2 (0.88 for Class 2 and 0.98 for Class 3, respectively). The probability (0.04) that individuals would move from the Class 2 to Class 1 by time 2 was not statistically significantly. Similarly, the probability (0.07) that individuals would move from the Class 2 to Class 1 by time 2 was not significantly different from zero. The probability (0.02) that individuals would move from the Class 3 to the Class 2 by time 2 was not statistically significant either. There was, however, a 0.10 probability that individuals would transition from Class 2 to Class
1 from time 2 to time 3. Multivariate analysis of variance showed that individuals who moved from Class 2 to Class 1 from time 2 to time 3 had significantly higher means than other individuals of Class 2 on FDG, MMSE, RAVimD, MOCA, EcPtMm, EcSPM. No significant mean difference was found between the two groups on EcSPLang. Entor, and Fusif. But these groups of individuals had significantly higher means than other individuals that individuals who moved from Class 2 to Class 1 from time 2 to time 3 had significantly higher means than other individuals that individuals who moved from Class 2 to Class 1 from time 2 to time 3. Multivariate analysis of variance showed that the LCA identified 63 individuals that were potentially misdiagnosed with AD. Indeed, based on the clinical and neural measures, it was deemed more probable that the misdiagnosed individuals be classified within the Class 1 instead of Class 2 or 3. Further, LTA did not show any significant change in class over time. Most individuals remained within their initial class (i.e., determined at baseline) and did not show a transition from Class 1 to Class 2, or from Class 2 to Class 3 between any time points. These results indicate that classifying individuals based on their cognitive and pathological parameters into different categories is an essential step toward understanding dementia and AD. To our knowledge, this is the first study to successfully use neuropsychological assessments and biomarkers of AD (e.g., Fluorodeoxyglucose, entorhinal cortex volume, and fusiform gyrus volume) to classify and predict individuals likely to transition from MCI to AD.

**DISCUSSION**

The aim of this study was to use LCA to identify and describe the number of classes that best characterize CN, SMC, EMCI, LMCI, and AD individuals with respect to clinical and neural measures. Our second aim was to compare the classification results obtained from the LCA to more traditional cut-off methods for classifying individuals with dementia. This can help us identify and characterize potentially misdiagnosed individuals. Finally, we used LTA to investigate changes in class membership over time. Our results showed that while there was substantial variability among individuals on their clinical and neural measures, the use of LCA with mixture methods to assess grouping individuals into optimal classes yields meaningful results. We confirm that using LCA, observing model fit indices, and entropy criterion were effective for selecting the optimal number of classes of individuals. Our results identified three classes of individuals with the following characterization: 37.5% of the individuals from Class 1 were CN, 20.10% with SMC, 19.60% with EMCI, 18.10% with LMCI, and 4.6% (63 individuals) with AD. Similar figures were 0.5, 0.9, 12.90, 53, and 32.60%; and 0.7, 0, 3.5, 28, and 67.80%, for Class 2 and Class 3, respectively. Further, our results showed that the LCA identified 63 individuals that were potentially misdiagnosed with AD. Indeed, based on the clinical and neural measures, it was deemed more probable that the misdiagnosed individuals be classified within the Class 1 instead of Class 2 or 3. Further, LTA did not show any significant change in class over time. Most individuals remained within their initial class (i.e., determined at baseline) and did not show a transition from Class 1 to Class 2, or from Class 2 to Class 3 between any time points. These results indicate that classifying individuals based on their cognitive and pathological parameters into different categories is an essential step toward understanding dementia and AD. To our knowledge, this is the first study to successfully use neuropsychological assessments and biomarkers of AD (e.g., Fluorodeoxyglucose, entorhinal cortex volume, and fusiform gyrus volume) to classify and predict individuals likely to transition from MCI to AD.

Similar to previous results, we identified multiple classes of cognitive impairment (Scheltens et al., 2016; Zammit et al., 2019b, 2020). For example, the results from our LCA identified 3 classes of individuals at each time point: Class 1, which is more healthy than the other classes representing 63, 59, and 65% of the sample at, respectively, time 1, 2, and time 3; Class

| Model          | Loglikelihood | #Free parameters | AIC       | BIC       | SBIC      | p LMR | Entropy |
|----------------|---------------|------------------|-----------|-----------|-----------|-------|---------|
| **Time point 1** |               |                  |           |           |           |       |         |
| Two-Class      | −12514.64     | 44               | 25117.28  | 25366.95  | 25227.15  | <0.001| 0.88    |
| Three-Class    | −11010.52     | 60               | 22141.04  | 22481.49  | 22290.86  | 0.003 | 0.89    |
| Four-Class     | −10133.10     | 76               | 20418.21  | 20849.44  | 20607.98  | 0.24  | 0.89    |
| Five-Class     | −9766.25      | 92               | 19716.50  | 20238.52  | 19948.22  | 0.189 | 0.87    |
| Six-Class      | −9492.06      | 108              | 19200.12  | 19812.93  | 19469.80  | 0.131 | 0.83    |
| Seven-Class    | −9279.09      | 124              | 18806.18  | 19509.77  | 19115.81  | 0.62  | 0.80    |
| **Time point 2** |               |                  |           |           |           |       |         |
| Two-Class      | −7936.22      | 44               | 15960.44  | 16201.10  | 16061.32  | <0.001| 0.81    |
| Three-Class    | −6927.78      | 60               | 13975.57  | 14303.75  | 14113.13  | <0.001| 0.82    |
| Four-Class     | −6513.27      | 76               | 13178.53  | 13594.23  | 13352.78  | 0.006 | 0.83    |
| Five-Class     | −6256.23      | 92               | 12696.45  | 13199.66  | 12907.38  | 0.004 | 0.75    |
| Six-Class      | −6087.00      | 108              | 12390.00  | 12980.73  | 12637.62  | 0.031 | 0.77    |
| **Time point 3** |               |                  |           |           |           |       |         |
| Two-Class      | −7936.56      | 44               | 15961.11  | 16201.12  | 16061.33  | <0.001| 0.79    |
| Three-Class    | −6984.23      | 60               | 14088.45  | 14415.74  | 14225.12  | 0.004 | 0.81    |
| Four-Class     | −6549.12      | 76               | 13250.23  | 13664.79  | 13423.35  | 0.514 | 0.78    |
| Five-Class     | −6374.21      | 92               | 12932.43  | 13434.26  | 13141.99  | 0.353 | 0.71    |

# number; AIC, Akaike Information Criterion; BIC, Bayesian Information Criterion; SBIC, Sample Size Adjusted BIC; p LMR, p- values for the Lo-Mendell-Rubin Likelihood ratio test for k vs. k+1 classes. LCA models converged at Time 1. The log-likelihood increased while no minimum was found for the ICs as their values decreased across the range of models considered.
2, which lies in between Class 1 and 3, representing ∼30% of the sample across the three time points; and Class 3, which include the least healthy individuals, representing 7, 10, and 6% of the sample at time 1, 2, and time 3, respectively. In other words, our LCA results reveal 3 classes that most likely match healthy individuals, individuals with MCI, and individual with AD, respectively. In comparison, Zammit et al. (2019b) classified participants into 5 classes within two categories (i.e., impaired and intact cognition) and Scheltens et al. (2016) classified participants into eight cognitive subtypes of AD. While there are differences in the number of classes identified with our results compared to others (Scheltens et al., 2016; Zammit et al., 2019b), it is important to note that the previous studies only utilized participants diagnosed with probable AD using neuropsychological assessments. Our research extends these findings by indicating that there are distinct classes of individuals who can be categorized as being healthy, experiencing MCI, and probable AD by using neuropsychological assessments and neurological biomarkers of AD. This has important clinical implications as individuals can be classified as experiencing different kinds of cognitive impairment early on (i.e., at baseline) and this categorization does not change significantly across time. Consistent with recent findings (i.e., Zammit et al., 2020), we also showed that using LCA to classify individuals with MCI or AD remains relatively stable over time (as indicated by LTA) and that LCA might better categorize and reduce the risk of misdiagnosis.

Our study has replicated previous findings that LCA and LTA can be used to identify homogeneous classes of cognitive impairment (Zammit et al., 2020). However, we have uniquely identified that neuropsychological measures of AD and the associated neurological biomarkers are indicators of an individual’s class membership and can predict their likelihood to transition between the healthy class (Class 1),
the MCI class (Class 2), and the AD class (Class 3). As mentioned earlier, recent studies using LTA have only utilize neuropsychological assessments of AD or MCI (e.g., Scheltens et al., 2016; Eppig et al., 2017; Zammit et al., 2019b, 2020). Our study adds a novel contribution to this emerging area by identifying that neurological biomarkers of AD can also be used to correctly classify individuals with MCI and AD and identify those at risk of transitioning from healthy cognitive function, to mild-cognitive impairment, and finally to AD.

A comparison of the classification results obtained from the LCA with the cutoff methods at Time 1 (baseline) revealed a group of 63 misclassified individuals. This group of individuals were classified as healthy individuals by the LCA, but were classified as AD by using clinical cut-off scores in neuropsychological assessments. The multivariate analysis of variance revealed that the misclassified individuals' scores on the clinical and neuropsychological assessments and neurological biomarkers were bounded between the mean of the Healthy individuals (i.e., Class 1) from the LCA without the 63 individuals and those from the AD individuals (i.e., Class 3) from the cut-off method without the 63 individuals. However, further analysis showed that it was more probable that the misclassified participants belonged to the healthy class (i.e., Class 1) rather than the MCI class (Class 2). As LCA takes into account several clinical and neural variables, longitudinal data, as well as also considers different groups of participants, it is likely to be more accurate than standard clinical cut-off methods, which often relies on one measure and does not compare data across different groups of participants.

There is a large discrepancy between the two methods of classification (i.e., clinical assessment vs. statistical), and perhaps reinforces the criticisms of using cut-off points on continuous neuropsychological assessments. Our results show that participants either side of the cut-off are similar (Berlin et al., 2014; Petersen et al., 2019). That is, based on clinical cut-off scores, some healthy participants were identified as similar to AD individuals, which resulted in misclassification. By using LCA, we have identified homogeneous sub-groups of individuals who are externally heterogeneous to other sub-groups (Berlin et al., 2014; Eppig et al., 2017; Mooney et al., 2018; Petersen et al., 2019; Villeneuve et al., 2019; Zammit et al., 2019a). For example, the healthy class (i.e., Class 1) is externally heterogeneous compared to the MCI class (Class 2) and the AD class (i.e.,

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**TABLE 3** | Cross tabulation of diagnostic variable and latent class variable at Time 1.

| Diagnostic variable at time 1 | Latent class solution at Time 1 |
|------------------------------|---------------------------------|
|                              | Class 1  | Class 2  | Class 3  |
| CN Frequency                 | 508      | 3        | 1        | 512      |
| Row %                        | 99.2%    | 0.6%     | 0.2%     |
| Column %                     | 37.5%    | 0.5%     | 0.7%     |
| SMC Frequency                | 273      | 6        | 0        | 279      |
| Row %                        | 97.8%    | 2.2%     | 0.0%     |
| Column %                     | 20.1%    | 0.9%     | 0.0%     |
| EMCI Frequency               | 266      | 82       | 5        | 353      |
| Row %                        | 75.4%    | 23.2%    | 1.4%     |
| Column %                     | 19.6%    | 12.9%    | 3.5%     |
| LMCI Frequency               | 245      | 336      | 40       | 621      |
| Row %                        | 39.5%    | 54.1%    | 6.4%     |
| Column %                     | 18.1%    | 53.0%    | 28.0%    |
| AD Frequency                 | 63       | 207      | 97       | 367      |
| Row %                        | 17.2%    | 56.4%    | 26.4%    |
| Column %                     | 4.6%     | 32.6%    | 67.8%    |

AD, Alzheimer's disease; CN, controls; EMCI, early-stage mild cognitive impairment; LMCI, late-stage mild cognitive impairment; SMC, subjective memory complaints.
3) classes were stable (with a probability that individuals in the Healthy (i.e., Class 1) and AD (i.e., Class 2) changes in the latent classes over time. The LTA results show a homogenous (i.e., after identifying misdiagnosed individuals) Class 3). Conversely, the AD class (i.e., Class 3) is relatively more heterogeneous compared to the healthy class and externally heterogeneous compared to the healthy class to transition to the MCI class (0.01 from time 1 to time 2 and 0.07 from time 2 to time 3). Furthermore, the results confirm the nature of AD as a progressive disease. Individuals with MCI (i.e., Class 3) are more likely to transition into the Healthy class or MCI class.

| Measures | The 63 patients (MD) | Diagnostics minus the 63 (DG) | Healthy minus the 63 (HT) | d | Contrasts |
|----------|----------------------|-----------------------------|--------------------------|---|-----------|
| FDG M (SD) | 1.26 (0.13) | 1.06 (0.15) | 1.29 (0.12) | 0.29*** | MD = HT > DG |
| CDRSBV (SD) | 1.69 (2.05) | 4.87 (2.45) | 0.52 (0.93) | 0.59*** | DG > MD > HT |
| ADAS11 (SD) | 1.14 (0.61) | 2.16 (0.90) | 0.75 (0.39) | 0.52*** | DG > MD > HT |
| MMSE (SD) | 2.70 (0.28) | 2.24 (0.34) | 2.87 (0.15) | 0.52*** | HT > MD > DG |
| RAVimD (SD) | 3.58 (1.26) | 2.17 (0.98) | 4.30 (1.13) | 0.36*** | HT > MD > DG |
| MOCA (SD) | 2.30 (0.43) | 1.70 (0.47) | 2.54 (0.30) | 0.48*** | HT > MD > DG |
| EcPtMm (MD) | 2.09 (0.74) | 2.27 (0.70) | 1.86 (0.66) | 0.05*** | DG > MD > HT |
| EcPtM (SD) | 1.77 (0.66) | 1.84 (0.66) | 1.60 (0.57) | 0.02*** | DG > MD > HT |
| EcSPM (MD) | 2.08 (0.83) | 2.99 (0.76) | 1.59 (0.61) | 0.41*** | DG > MD > HT |
| EcSPM (SD) | 1.70 (0.69) | 2.36 (0.79) | 1.31 (0.46) | 0.36*** | DG > MD > HT |
| HPCI (MD) | 0.70 (0.12) | 0.56 (0.11) | 0.72 (0.10) | 0.28*** | MD = HT > DG |
| Entor (MD) | 0.34 (0.07) | 0.27 (0.07) | 0.37 (0.07) | 0.26*** | HT > MD > DG |
| Fusif (MD) | 1.71 (0.25) | 1.49 (0.28) | 1.78 (0.26) | 0.16*** | HT > MD > DG |

***p < 0.001. All variables are defined in the text and in the captions of prior tables.

Table 5: Transition probabilities from Time 1 to Time 2 and from Time 2 to Time 3.

|        | Class 1 | Class 2 | Class 3 |        | Class 1 | Class 2 | Class 3 |
|--------|---------|---------|---------|--------|---------|---------|---------|
| T1     | 0.99a*  | 0.01    | 0.00    | T2     | 0.93*   | 0.07    | 0.00    |
| Class 2| 0.04p   | 0.88*   | 0.07    | Class 2| 0.10*   | 0.85*   | 0.06    |
| Class 3| 0.00    | 0.02    | 0.98*   | Class 3| 0.00    | 0.09    | 0.91*   |

*p < 0.05.

Class 3). Conversely, the AD class (i.e., Class 3) is relatively more homogenous (i.e., after identifying misdiagnosed individuals) and externally heterogeneous compared to the healthy class and MCI class.

The LTA results were used to examine the individuals’ transition probabilities with respect to clinical assessments, neurological measures, and neurological biomarkers to explore changes in the latent classes over time. The LTA results showed that individuals in the Healthy (i.e., Class 1) and AD (i.e., Class 2) classes were stable (with a probability >0.90 of staying in the same class over time). The results show that the transition from Healthy to AD classes was non-existent. This is consistent with the nature of the AD as a progressive disease. Individuals at early stages do not exhibit symptoms of AD. However, there is an insignificant probability for the individuals in the healthy class to transition to the MCI class (0.01 from time 1 to time 2 and 0.07 from time 2 to time 3). Furthermore, the results confirm the nature of AD as a neurodegenerative disease (Alzheimer Association, 2019). For example, at no time-point did individuals who are in the AD class show cognitive improvement by transitioning to the Healthy class or MCI class.

In contrast, individuals from the MCI class have non-zero probabilities moving to other classes over time. However, only the probability of transitioning to the Healthy class from time 2 to time 3 was significant (with a probability of 0.10). That is, some individuals classified within the MCI class (Class 2) showed cognitive improvement from time 2 to time 3. This further emphasizes the differences between MCI and AD. While AD is a regressive disease that does not allow cognitive improvements, MCI is not necessarily degenerative. As such, we should be cautious about suggesting individuals with MCI are on a progression toward AD. As we only observed movement from the MCI class into the healthy class, it is less probable that individuals with MCI will progress into AD. Therefore, based on our findings, any transition from the MCI class is likely to resemble cognitive improvement rather than decline. It is worth noting that we did not identify the characteristics that predict the movement of individuals from the MCI class. Further research is needed to investigate possible factors that may contribute to this movement, either to the Healthy or AD class.

**Limitations**

It is important to note that one main advantage of traditional rule-based diagnostic methods (as often used by most clinicians and doctors) is easy utilization in the everyday clinical setting. However, our methods used here are more complex and...
require applying analytical and statistical method to be able to reach a more robust diagnosis. Accordingly, because of its complexity (e.g., conducting the analysis and interpretations of results), it is expected that latent class analysis methods may not be widely used. However, we also agree that it is exactly LCA complexity over the traditional discrete diagnostic methods (e.g., surveys) that allow it to be a better predictor of class membership (e.g., an individual is healthy, has mild cognitive impairment, or has Alzheimer's disease). This is due to the fact that rule-based diagnostic methods are inherently additive, as they rely on discrete methods. However, LCA is a multivariate approach that attempts to find complex joint probability distributions that create a richer risk profile which is difficult to define using discrete rule-based decision tools and diagnostic methods.

**CONCLUSION**

In conclusion, this study demonstrated that latent class analysis can be used to classify participants within the ADNI project into three distinct classes: Healthy, MCI, or AD. We argue that LCA is a more suitable method for classifying individuals with SMC, MCI, and AD rather than using clinical cut-off measures. This is due to LCA's ability to create internally homogenous and externally heterogeneous sub-groups. This technique might help reduce the number of misclassifications of individuals incorrectly diagnosed with probable AD—as demonstrated by the misclassified individuals in our study. By using latent transition analysis, we showed that individuals classified as healthy or with AD had a high probability of staying in the same class over time. However, it was more probable for individuals to transition from the MCI class to the healthy class. Our results emphasize that AD is a neurodegenerative syndrome, with individuals within the AD class showing no evidence of cognitive improvement over time. However, individuals with MCI can show improvement over time. Therefore, we argue that LCA can be used to differentiate between individuals with AD and that this diagnosis remains stable across time and produces fewer misdiagnoses than using clinical cut-offs. Robust methods should be used to accurately diagnose patients and to identify individuals at a higher risk of developing AD. While using cut-off scores using traditional discrete diagnostic methods are quicker, our study has shown that LCA can provide a more accurate prediction for classifying individuals with SMC, MCI, and AD. While the time requirement to conduct LCA is burdensome, ensuring an accurate diagnosis for patients should be a prioritized. Especially given the severity and neurodegenerative nature of AD (Alzheimer Association, 2019). Using LCA and LTA can provide more accurate diagnoses and improve the outcomes for patients. Clinicians should consider alternative diagnostic methods for AD instead of relying solely on the clinical cut-off measures on neuropsychological assessments.

**DATA AVAILABILITY STATEMENT**

Publicly available datasets were analyzed in this study. This data can be found here: http://adni.loni.usc.edu.

**AUTHOR CONTRIBUTIONS**

All authors contributed to data analysis, writing, and editing the manuscript.

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**Conflict of Interest:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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