Unraveling the heterogeneity in Alzheimer’s disease progression across multiple cohorts and the implications for data-driven disease modeling

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Abstract

Introduction: Given study-specific inclusion and exclusion criteria, Alzheimer’s disease (AD) cohort studies effectively sample from different statistical distributions. This heterogeneity can propagate into cohort-specific signals and subsequently bias data-driven investigations of disease progression patterns.

Methods: We built multi-state models for six independent AD cohort datasets to statistically compare disease progression patterns across them. Additionally, we propose a novel method for clustering cohorts with regard to their progression signals.

Results: We identified significant differences in progression patterns across cohorts. Models trained on cohort data learned cohort-specific effects that bias their estimations. We demonstrated how six cohorts relate to each other regarding their disease progression.

Discussion: Heterogeneity in cohort datasets impedes the reproducibility of data-driven results and validation of progression models generated on single cohorts. To ensure robust scientific insights, it is advisable to externally validate results in independent cohort datasets. The proposed clustering assesses the comparability of cohorts in an unbiased, data-driven manner.

KEYWORDS
Alzheimer’s disease, cohort study, data mining, data-driven, disease modeling, machine learning, sampling bias, statistical learning, translational research
In the last decade, understanding the progressive dynamics of Alzheimer’s disease (AD) and AD clinical syndrome,1 proved to be one of the fundamental challenges in our field.2,3 Comprehensive knowledge on AD progression opens crucial opportunities for medical intervention to counteract or delay impediments to activities of daily living.5 One path to facilitate this understanding manifests in the extraction of longitudinal progression signals from patient-level datasets collected in cohort studies. In this context, data mining and machine learning methods can be used to build mathematical models that elucidate and predict progression patterns hidden in the data. In the past, such progression models were used, for example, to approximate biomarker trajectories,5 to identify distinct progression subtypes,6 and to assess patient risk of progression toward more impaired disease stages.7 However, to demonstrate that progression patterns identified in one cohort generalize beyond the discovery dataset itself, it is imperative to externally validate them in an independent dataset.8 External validation data should originate from a separate cohort study independent from the training data used for building the model. Especially in the context of multifactorial and heterogenous diseases such as AD, external validation turns out to be a non-trivial undertaking.

The key limitation encountered in external validation manifests in the characteristics of clinical AD cohort data.9 By nature of the disease, AD cohorts are very heterogeneous with respect to their exhibited progression,10 for example, with respect to brain atrophy11 and age of disease onset.12 Furthermore, cohort study participants are recruited according to specific inclusion and exclusion criteria defined based on the goals of the study (e.g., selection of specific age ranges or risk factors). These specific sampling procedures shape potentially distinct statistical distributions from which each study’s participants are recruited and, in turn, inevitably introduce cohort-specific statistical biases into the collected dataset itself.13,14 These aspects potentially violate the fundamental assumption behind data mining and machine learning approaches that the participants of a validation dataset constitute a representative sample of the same population from which the original training data were drawn (Figure S1 in supporting information). Consequently, this indicates that training and validation data must be independently and identically distributed (i.i.d.) samples.15 As such, a well-trained model should show similar performance on a validation dataset that was drawn from the identical statistical distribution as the training data, while an overfitted model would fail such validation. However, on a validation dataset that is violating the assumption of being sampled from the same statistical distribution as the training data even a well-trained model would fail, because the validation data falls outside the domain of the model (Figure S1). In conclusion, data-driven models trained on cohort datasets cannot be expected to generalize appropriately beyond the statistical distribution from which this cohort’s participants were sampled.16,17

The heterogeneity found in AD cohort datasets, therefore, raises several important questions with respect to data-driven modeling of AD. First, it warrants an evaluation as to whether exhibited trends of disease progression are consistent across cohorts despite possible differences in their underlying populations. Further investigation should also determine whether progression models fitted on such datasets learn potential cohort-specific biases that could impede the generalizability of findings. Finally, as of now, there is no way to measure and express the general comparability between patient-level datasets on the level of disease progression. In the past, researchers mainly relied on comparing baseline study characteristics of their studied datasets.7,18,19 However, for obvious reasons, evaluating variable distributions at a singular time point is a very limited comparison in the scope of disease progression. Deriving a quantitative measure to compare longitudinal progression patterns across multiple clinical studies could aid researchers to better understand the landscape of existing studies and to identify datasets that might fulfill the i.i.d. assumption. Furthermore, it could be used to investigate whether the cause of a significant drop in prediction performance lies in systematic differences between the training and validation datasets (i.e., a probable violation of the i.i.d. assumption) or simply in an overfitted model.

In this work, we evaluated the heterogeneity of disease progression patterns encountered in six longitudinal clinical AD cohort studies. Relying on multi-state models (MSM),20 a well-established data mining approach in the AD field,21–24 we performed a systematic comparison of progression patterns extracted from these studies to assess whether discovered signals are robust. Furthermore, we investigated whether cohort-specific biases propagate into trained
progression models. Finally, we propose a novel method for clustering cohorts based on their exhibited progression patterns. This approach reveals the similarity of cohort studies in a data-driven and unbiased manner. It allows researchers to adequately understand and characterize performances measured via external validation of statistical and machine learning models developed on another cohort. In conclusion, our approach allows for better understanding of statistical differences that have previously been reported between various AD studies.13

2 | METHODS

2.1 | Data selection

Six longitudinal datasets stemming from the Alzheimer’s Disease Neuroimaging Initiative (ADNI),25 AddNeuroMed (ANMerge),26 Australian Imaging, Biomarker & Lifestyle Flagship Study of Ageing (AIBL),27 Japanese Alzheimer’s Disease Neuroimaging (J-ADNI),28 National Alzheimer’s Coordinating Center (NACC),29 and the Religious Orders Study and Rush Memory and Aging Project (ROSMAP)30 were used as training datasets for our progression models. All of these studies obtained ethical approval for human data collection and informed patient consent for data sharing. We excluded participants whose mild cognitive impairment (MCI) diagnoses were not attributed to AD. Information on the cohorts with respect to key variables, as well as the number of participants, can be found in Table S1 in supporting information.

2.2 | Progression models applied for statistical analysis

To extract disease progression patterns from the investigated datasets, we fitted one MSM per cohort using the msm R package.20 The states in our models represent the three commonly assessed stages for AD progression: cognitively unimpaired (CU), MCI, and AD. Consequently, transitions between states illustrate conversions from one clinical diagnosis stage to another. We modeled AD as an absorbing state, that is, we assumed that patients were not able to recover once deterioration was advanced enough to receive an AD diagnosis. However, because the classification of patients into CU, MCI, and AD in all cohorts had been performed based on clinical assessments, reversions from AD were observed in the data. These reversions were modeled as misclassifications. A graphical representation of the model can be seen in Figure S3 in supporting information. Each transition rate was estimated based on a set of covariates to account for the individual compositions of the cohorts. For determining the most informative combination of covariates, we performed a rigorous model selection using the Akaike’s information criterion (AIC). The choice of covariates was mainly limited by their availability across the cohorts (Figure S2 in supporting information). Ultimately, the selected covariates comprised participant’s age, biological sex, completed years of education, apolipoprotein E (APOE) ε4 status, and the Mini-Mental State Examination (MMSE). Likelihood-ratio tests comparing each MSM to a null model demonstrated that all models extracted progression signals from their training dataset (P < .05). To rule out potential overfitting of the models, we built 150 models on repeated bootstrap samples from each respective cohort and observed low variation in model estimates (Table S3 in supporting information). Application of interval censoring allowed for the inclusion of participants with missing intermediate visits while right censoring was used for individuals who did not receive an AD diagnosis during study runtime. More details on the methodology and model selection are presented in the supporting information.

2.3 | Comparison of data mined progression patterns across cohorts

To explore and assess the heterogeneity in disease progression trends across cohorts, we estimated several progression patterns using each cohort’s MSM: the state transition probabilities, probability of staying AD diagnosis free over time, and sojourn times (i.e., the expected time a participant spends in a considered state). All patterns were separately investigated for the CU and MCI states. For estimation of a cohort’s progression patterns starting in the CU state, we used the covariate values of patients at their first MCI diagnosis. Similarly, for estimating transitions from the MCI state, we relied on the covariate values of participants at their first MCI diagnosis. Where appropriate, uncertainty of estimates was quantified using 95% confidence intervals (CI). Differences between cohort-specific distributions of the aforementioned progression estimates were determined using Kruskal-Wallis and pairwise Mann-Whitney U tests employing a confidence level of 95%. P-values were corrected for multiple testing using the Bonferroni-Holm method.

2.4 | Evaluation of cohort biases in statistical models

The second set of analyses aimed at elucidating whether MSMs fitted to data from a single cohort would learn cohort-specific effects that reduce generalizability to other cohorts. Hazard ratios, for example, are covariate-specific parameters of a model that quantify the influence of covariates onto the transition risk between two states. Comparing these ratios, it becomes apparent whether models learned the same covariate influences from distinct cohorts. Furthermore, we used each cohort’s previously trained MSM to estimate the progression patterns for the same, combined set of participants from all cohorts. By fixing the data to be estimated across models, all variability in the progression patterns stems from the cohort-specific effects learned by the model. To evaluate the existence of these cohort-specific biases, we performed Kruskal-Wallis tests and pairwise Mann-Whitney
FIGURE 1  Probabilities to transition from one state to another are estimated for a 10-year period. Median probabilities are marked with white points. Statistical distributions are shown as box plots as well as superimposed kernel density estimates, resulting in violin plots. Because most deviations between depicted distributions were significant, we omit indication of significance for brevity. A-C, Transition probabilities starting from the cognitively unimpaired (CU) state. D-F, Transition probabilities starting from the mild cognitive impairment (MCI) state. AD, Alzheimer’s disease; ADNI, Alzheimer’s Disease Neuroimaging Initiative; AIBL, Australian Imaging, Biomarker & Lifestyle Flagship Study of Ageing; ANMerge, AddNeuroMed; J-ADNI, Japanese Alzheimer’s Disease Neuroimaging Initiative; NACC, National Alzheimer’s Coordinating Center; ROSMAP, Religious Orders Study and Rush Memory and Aging Project

U tests, again correcting for multiple testing using Bonferroni-Holm and assuming a confidence level of 95%.

2.5 Cohort similarity clustering

Whereas previous analyses focused on statistical differences between cohorts, we additionally developed an approach to cluster cohorts based on their global similarity across progression patterns. More specifically, each cohort’s MSM was used to calculate the log-likelihood of observing the actual transitions of all the participants of each other cohort. These pairwise log-likelihoods were afterward averaged across the number of participants per cohort to eliminate biases toward cohort size. This resulted in a pairwise similarity matrix between cohorts which was subsequently transformed into a symmetric distance matrix. Mathematical details can be found in the supporting information. The resulting distance matrix was then used in an agglomerative hierarchical clustering approach using average linkage.

3 RESULTS

3.1 Progression patterns differ across cohorts

Transition probabilities estimated for a 10-year period varied significantly between cohorts (Figure 1). While we observed in all cohorts that participants in the CU state were most likely to remain CU over the next 10 years, the proportions of probabilities showed evident differences (Figure 1A-C). We discovered a range of 25% difference between the maximum and minimum observed median probability to remain CU (J-ADNI, > 99%; ADNI, 75%). All observed differences between pairwise combinations of cohorts were significant ($P < .001$), with the exception of ROSMAP–NACC for remaining in the CU state ($P = .3$).

When investigating the estimated transition probabilities from the MCI state (Figure 1D-F), all cohorts exhibited their most probable transition toward the AD state. J-ADNI showed the highest median probability across cohorts with 85%, while ROSMAP held the lowest median probability with 50%, exposing a difference of 35% between them.
FIGURE 2  Average probability of staying AD diagnosis free over time for each cohort. Dashed lines indicate the standard errors of the
estimates. A, Starting from cognitively unimpaired. B, Starting from mild cognitive impairment. ADNI, Alzheimer’s Disease Neuroimaging Initiative; AIBL, Australian Imaging, Biomarker & Lifestyle Flagship Study of Ageing; ANMerge, AddNeuroMed; J-ADNI, Japanese Alzheimer’s Disease Neuroimaging Initiative; NACC, National Alzheimer’s Coordinating Center; ROSMAP, Religious Orders Study and Rush Memory and Aging Project

Additionally, compared to the other cohorts, ROSMAP showed a considerably higher median probability to revert from MCI back to CU of 23%. All pairwise differences across cohorts proved to be significant ($P < .001$). Numerical values for the transition probabilities are presented in Table S4 in supporting information.

In concordance with the transition probabilities, the probability of staying AD diagnosis free over time differed substantially across cohorts. Starting in the CU state (Figure 2A), the trajectories of cohorts deviated significantly after approximately 4 years. NACC and ROSMAP exhibited the steepest decline (respectively, 85% and 87% after 10 years), while the probability for ANMerge stayed relatively stable (99%). Considering the MCI state as a starting point, the probability of remaining AD diagnosis free exhibited a steeper decline (Figure 2B). After 10 years, the most extreme estimates were made for ROSMAP (48%) and J-ADNI (20%), while no significant differences were observed between J-ADNI and NACC (both 20%), as well as between AIBL and ADNI (both 42%). Ultimately, we discovered a maximum deviation of 14% for the CU state and 28% for the MCI state.

All pairwise comparisons between the cohorts’ sojourn time estimates turned out to be significant for the CU state ($P < .001$, with exception of ADNI–ROSMAP, $P < .05$; Figure 3A). Given their respective MSMs, ROSMAP displayed the shortest sojourn time with a median of 27.5 years, followed by ADNI (29.7 years), NACC (38.7 years), AIBL, ANMerge, and J-ADNI (all > 100 years). In the MCI state, again, most deviations were found to be significant ($P < .001$; Figure 3B). The only exception to this was ANMerge, which did not differ significantly from ADNI ($P = .9$) and AIBL ($P = .88$). The median sojourn time in the MCI state showed relatively lower values for J-ADNI (3.8 years) and NACC (3.1 years), while ADNI, AIBL, and ANMerge showed relatively higher values (7.7, 6.5, and 6.9 years, respectively). ROSMAP is placed in between with a median of 5 years. Detailed descriptions of the sojourn times distributions can be found in Table S5 in supporting information.

### 3.2 Comparison of cohort-specific models

In the second set of analyses, we explored the cohort-specific biases learned by our MSMs from their respective training datasets. We observed that the cohort-specific models learned significantly different relationships between covariate values and the disease progression. Non-overlapping CIs indicated significant differences in hazard ratios for the transition from CU to MCI between ROSMAP (CI: 1.05 to 1.1), NACC (1.0 to 1.04), and ADNI (0.86 to 0.99) regarding education level. With respect to the MMSE, significant differences were found for ROSMAP, NACC, J-ADNI, and ADNI (CIs: 0.60 to 0.67, 0.76 to 0.81, 0.11 to 0.58, and 0.76 to 0.98, respectively; Figure 4A). The influence of education in J-ADNI (CI: 1.15 to 1.92) differed significantly from ADNI (0.93 to 1.12), NACC (0.94 to 1.04), and ROSMAP (0.93 to 1.03) with respect to reverting from MCI to CU (Figure 4B). Regarding the conversion from MCI to AD, significant differences were discovered in the hazard ratios for age between ROSMAP (1.02 to 1.05) and NACC (1.00 to 1.01), for $\text{APOE}_4$ status between NACC (1.10 to 1.31) and ADNI (1.34 to 1.82), and for MMSE between NACC (0.83 to 0.87), ADNI (0.7 to 0.76), and ROSMAP (0.74 to 0.79; Figure 4C). In several cases, large CIs hampered the interpretation of the hazard ratios. The exact estimates of all hazard ratios are presented in Table S6 in supporting information.

When applying each MSM to the same set of data, the difference in the estimated progression patterns across models resembled the consequences of the learned cohort-specific biases (Figure 5). Numerical descriptions of the distributions in Figure 5 can be found in.
Sojourn times of cohort participants on a log10-scale. Because most deviations between depicted distributions were significant, we omit indication of significance for brevity and refer to the text. A, Occupying the cognitively unimpaired state. B, Occupying the mild cognitive impairment state. ADNI, Alzheimer’s Disease Neuroimaging Initiative; AIBL, Australian Imaging, Biomarker & Lifestyle Flagship Study of Ageing; ANMerge, AddNeuroMed; J-ADNI, Japanese Alzheimer’s Disease Neuroimaging Initiative; NACC, National Alzheimer’s Coordinating Center; ROSMAP, Religious Orders Study and Rush Memory and Aging Project.

FIGURE 3  

Clustering reveals overall similarity of studies

Figure 6 presents the results achieved by clustering the investigated cohorts based on the similarity of their progression patterns. ANMerge, AIBL, and NACC displayed close proximity indicating that their participants exhibited similar disease progression in combination with their trained MSMs. Furthermore, ADNI and J-ADNI formed a cluster that connected with the previously mentioned cluster in relatively high distance. ROSMAP was placed far from all other cohorts, constituting its own cluster.

3.3  Clustering reveals overall similarity of studies

We observed that naive pooling of datasets and training models on a combination of multiple, complete cohorts expectedly biases the estimates toward the cohort with the largest sample size (Figure S4 in supporting information). We also found differences between cohorts when extracting progression patterns for a cohort’s representative individual (Figure S5 in supporting information) and even when applying the same exemplary patients to each cohort’s specific MSM (Figure S6 in supporting information).

4  DISCUSSION

In this work, we explored the heterogeneity in AD progression across multiple, independent cohort datasets and the implications for data-driven approaches for progression modeling. Evident differences in mined progression patterns surfaced between six investigated cohorts. This finding raises concerns regarding the reliability of results discovered in single data resources and underlines the need for external validation. Furthermore, we demonstrated that models learn cohort-specific effects from their training dataset, which can impede model generalization. Last, we proposed a novel approach to identify similar cohort datasets that could help to find datasets that come closer to fulfilling the i.i.d. assumption. We demonstrated this approach by highlighting how six major AD cohorts relate to each other with regard to their exhibited disease progression.

4.1  Progression trends differ across cohort datasets

Analyzing the characteristic progression trends extracted from the investigated cohorts revealed substantial differences among them. The observation of lower variability in estimates for the CU state compared to the MCI state can be explained by the fact that only a fraction of the CU participants will eventually develop cognitive symptoms. Thus, a substantial amount of CU participants are expected to show no signals of AD progression at all. Overall, the discovered heterogeneity could likely stem from differences in the recruitment processes of cohort studies. Compositional shifts across sampled populations pose a critical confounder comparing cohort datasets and model performance. Here, statistical matching could potentially help to identify comparable subsets.

4.2  Data-driven models learn systematic biases present in cohort datasets

Using all cohort-specific MSMs to estimate progression patterns for the same set of participants revealed the presence of strong
FIGURE 4  Covariate hazard ratios learned per cohort-specific multi-state models. For readability, significant deviations are not indicated visually. Instead, we refer to the text for the corresponding evaluations. A, B, C, Impact on transition from cognitively unimpaired (CU) to mild cognitive impairment (MCI), MCI to CU, and MCI to Alzheimer’s disease (AD), respectively. ADNI, Alzheimer’s Disease Neuroimaging Initiative; AIBL, Australian Imaging, Biomarker & Lifestyle Flagship Study of Ageing; ANMerge, AddNeuroMed; J-ADNI, Japanese Alzheimer’s Disease Neuroimaging Initiative; NACC, National Alzheimer’s Coordinating Center; ROSMAP, Religious Orders Study and Rush Memory and Aging Project

cohort-specific effects that the models learned from their training datasets. The estimated covariate hazard ratios are an integral component of the cohort-specific progression signals and while we could observe commonalities in the directional influence of covariates, partially described by previous studies as well,\textsuperscript{7,21} the magnitude of these influences exposed several significant differences. With regard to education, even contradicting influences were found. Differences in such fundamental parameters of a model propagate into, and thereby bias, their estimates; this became apparent in the subsequently estimated progression patterns.

Naive pooling of data from several cohorts does not necessarily pose a solution for addressing the biases but leads to an overshadowing of signals in smaller cohorts by larger ones. Instead, more considerate methods must be applied, such as sampling the same number of participants from each cohort, weighting of subjects to favor smaller datasets, or ensemble techniques that combine
A) Transition Probabilities

B) Sojourn Times

C) Probability of No AD Conversion
dataset-specific models. Future work should explore these options in more detail.

4.3 Clustering allows assessment of cohort similarities

Our proposed approach to measure cohort similarity with regard to their global disease progression trends (informed by neuropsychological tests, biological sex, completed years of education, APOE ε4 status) elicited commonalities across cohorts that mirror the design of these studies. Finding ADNI and J-ADNI in one cluster together is reassuring as J-ADNI was designed as a complementary cohort to ADNI, and similar trends have been observed in both cohorts.28 Their use of equal eligibility criteria for participant recruitment counteracts the risk of sampling from two distinct populations. The distance we observe between them could be explained partially due to differences in ethnorracial composition31 and lifestyle.32 ROSMAP, on the other hand, is a special case in the landscape of AD cohorts. Its participants are exclusively recruited from religious orders, are considerably older, and hold a higher proportion of female participants compared to the other cohorts.13,30

Our proposed method enables a quantitative description of differences across cohorts and, subsequently, an evaluation of cohort similarity based not only on cross-sectional values of covariates but on their general progression. Consequently, it could help researchers to better understand and characterize progression signals between discovery and validation cohort originating from, for example, sampling of distinct statistical distributions.

4.4 Limitations

It is unknown how many of the CU participants per cohort would have eventually developed cognitive symptoms during their lifetime. While the models account for this factor using censoring, estimates based on the CU participants could be biased depending on the size of the participant fraction with prodromal AD.

One limitation of MSMs is the assumption that disease progression depends only on the current state of a participant. While this is a necessary and widely accepted assumption in the literature,7,21-24 there is no universal way to prove that it always holds true for all possible state transitions.

5 Conclusion

Applying machine learning and statistical modeling to single data resources can bias results and might render the generalizability of the models used infeasible. Ideally, it would be imperative that we go beyond single data resources and instead investigate and validate findings across the landscape of AD data we have at our disposal. In practice, however, external validation of data-driven machine learning models is often limited by the availability of semantically and statistically comparable datasets.13 For some investigations only single cohorts might be suitable. While results originating from such single-cohort investigations hold value as initial indications, they should be regarded as cohort-specific findings pending external validation, and
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CONFLICTS OF INTEREST

The authors have nothing to declare.

AUTHOR CONTRIBUTIONS

Colin Birkenbihl and Holger Fröhlich designed the study. Yasamin Salimi and Colin Birkenbihl implemented the methods and ran the experiments. Colin Birkenbihl wrote the manuscript. Holger Fröhlich and Yasamin Salimi revised the manuscript. Holger Fröhlich supervised the project.

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SUPPORTING INFORMATION
Additional supporting information may be found online in the Supporting Information section at the end of the article.