

HHS Public Access

Author manuscript *Neuroimage*. Author manuscript; available in PMC 2022 September 01.

Published in final edited form as:

Neuroimage. 2022 September ; 258: 119353. doi:10.1016/j.neuroimage.2022.119353.

Transfer Learning for Cognitive Reserve Quantification

Xi Zhu^{1,2}, Yi Liu³, Christian G. Habeck⁴, Yaakov Stern^{2,4}, Seonjoo Lee^{1,3}, Alzheimer's Disease Neuroimaging Initiative^{*}

¹Department of Psychiatry, Columbia University Irving Medical Center, New York, USA

²New York State Psychiatric Institute, New York, USA

³Department of Biostatistics, Columbia University Irving Medical Center, New York, USA

⁴Cognitive Neuroscience Division, Department of Neurology, Columbia University Vagelos College of Physicians and Surgeons, New York, USA

Abstract

Cognitive reserve (CR) has been introduced to explain individual differences in susceptibility to cognitive or functional impairment in the presence of age or pathology. We developed a deep learning model to quantify the CR as residual variance in memory performance using the structural MRI data from a lifespan healthy cohort. The generalizability of the sMRI-based deep learning model was tested in two independent healthy and Alzheimer's cohorts using transfer learning framework.

Structural MRIs were collected from three cohorts: 495 healthy adults (initially aged 20–80) from RANN, 620 healthy participants (age 36–100) from lifespan Human Connectome Project Aging (HCPA), and 941 subjects from Alzheimer's Disease Neuroimaging Initiative (ADNI). Region of interest (ROI)-specific cortical thickness and volume measures were extracted using the Desikan-Killiany Atlas. Cognitive reserve was quantified by residuals which subtract the predicted memory from the true memory. Cascade neural network (CNN) models were used to train RANN dataset for memory prediction. Transfer learning was applied to transfer the T1 imaging-based model from source domain (using RANN) to the target domain (HCPA or ADNI).

The CNN model trained on the RANN dataset exhibited strong linear correlation between true and predicted memory based on the chosen T1 cortical thickness and volume predictors. In addition, the model generated from healthy lifespan data (RANN) was able to generalize to an independent healthy lifespan data (HCPA) and older demented participants (ADNI) across different scanner types. The estimated CR was correlated with CR proxies such education and IQ across all three datasets.

Data Availability

Address correspondence to: Seonjoo Lee, PhD, 722 West 168th street, #211, New York, NY10032, Tel 646-774-5461, sl3670@cumc.columbia.edu.

^{*}Data used in preparation of this article were obtained from the Alzheimer's Disease Neuroimaging Initiative (ADNI) database (adni.loni.usc.edu). As such, the investigators within the ADNI contributed to the design and implementation of ADNI and/or provided data but did not participate in analysis or writing of this report. A complete listing of ADNI investigators can be found at: http://adni.loni.usc.edu/wp-content/uploads/how_to_apply/ADNI_Acknowledgement_List.pdf

Competing Interest Statement

The authors have declared no competing interest.

All data examined in the manuscript are available upon request in deidentified format.

The current findings suggest that the transfer learning approach is an effective way to generalize the residual-based CR estimation. It is applicable to various diseases and may flexibly incorporate different imaging modalities such as fMRI and PET, making it a promising tool for scientific and clinical purposes.

Keywords

Cognitive reserve; Transfer Learning; ADNI; HCP; MRI

Introduction

Approximately 15–20% of adults aged 65 or older suffer from significant cognitive decline resulting in mild cognitive impairment (MCI); among these, 11.3% adults later develop dementia due to Alzheimer's disease (AD) (Association, 2021). With the lack of an effective treatment strategy, there is a great need to identify factors that can slow the progression to dementia and maintain quality of life (Zissimopoulos, Crimmins, & St Clair, 2014).

Cognitive reserve (CR) has been introduced to explain individual differences in susceptibility to cognitive or functional impairment in the presence of age or diseaserelated brain changes (Stern, 2002). Individuals with high CR have greater resilience and maintained normal cognitive function longer when confronted with late-life neuropathology. Typical CR proxy measures include years of education (Meng & D'Arcy, 2012; Stern et al., 1994), premorbid IQ (Alexander et al., 1997), occupational achievement, and engagement in cognitively and socially stimulating activities (Scarmeas & Stern, 2003). These are thought to protect against functional impairment by promoting the ability to better compensate for brain changes. However, qualifying as a proxy for cognitive reserve does not imply sufficiency for the mechanism proper since a rigorous test of cognitive-reserve mechanisms requires both brain and cognitive measures. Proxy measures, while often used in the past and spoken about interchangeably with cognitive reserve, are not enough here since their effect on cognition could be fully mediated through brain structure, a clear case where better brain maintenance, and not better cognitive reserve, would be at play. An NIH-funded collaboratory for clarifying research definition for cognitive reserve and resilience has brought all this into clear focus over the last 3 years. (https://reserveandresilience.com/). Thus, proxy measures fail to provide the entirety of the construct; the same value of a proxy variable may reflect different experiences across people. In addition, most proxy measures only represent static cognitive reserve and cannot account for possible change over time. Lastly, these measures rely on the recollection of prior activities, which are an indirect proxy of CR. (Borenstein, Copenhaver, & Mortimer, 2006; Jones et al., 2011; Satz, Cole, Hardy, & Rassovsky, 2011). To address these limitations, a direct measure of CR based on unbiased current information is highly needed. One popular approach to quantify CR is to measure the residual variance between predicted cognitive performance based on an individual's level of brain status and neuropathology and the actual individual's performance (Reed et al., 2010). These residual-based measures offer a more precise measurement of CR (Bocancea et al., 2021). High-reserve individuals exhibit higher actual measured cognitive performance than that predicted.

Structural magnetic resonance imaging (sMRI) has been used as a measure of the regional brain atrophy underlying cognitive decline and dementia (Mueller, Schuff, & Weiner, 2006). Previous studies using the sMRI to calculate the residual variance operational measure of CR showed promising results (Reed et al., 2010; Zahodne et al., 2015; Zahodne et al., 2013) in older participants. Currently, most research in cognitive aging has used life-span data (Razlighi, Habeck, Barulli, & Stern, 2017; Salthouse, 2010; Taylor et al., 2017; Tucker-Drob, 2019). However, leveraging life-span brain and cognition data in quantifying cognitive reserve has not been done.

In addition, brain imaging data from multi-sites may have high variability due to different MRI sequences of different scanners, thus, limiting direct application of a previously trained model to new datasets acquired from different sites.

Traditional machine learning methods to mitigate the influence of variability across sites require a balanced sample from each site and assume the same distribution across training and test datasets. The performance of a predictive model declines when these assumptions are violated. Transfer learning is a machine learning technique that utilizes the knowledge gained from one task and applies it to a different but related task. It is a popular optimization approach that allows rapid progress or improved performance when modeling the second task. The sMRI obtained from various sites or scanners may represent similar brain properties but may exhibit different observational distributions. Thus, the transfer learning approach may be applied to improve the generalizability of the sMRI-based residual models.

In this study, we proposed a CR quantification framework that leverages a single-site, large scale lifespan data and uses transfer learning to handle scanner and site differences. We first built a deep learning model to quantify the CR as residual variance in memory performance using the sMRI data from a healthy lifespan cohort (age 20-80). Our first goal was to assess whether using lifespan data of healthy individuals, which shows more variability in cognition function, may enable better quantification of the relationship between sMRI and cognitive performance. Second, to test the generalizability of the sMRI-based deep learning model, we utilized the transfer learning approach to fine-tune the pre-trained deep learning model to an independent, healthy lifespan cohort: the Human Connectome Project-Aging cohort (HCPA). Third, to test whether the model generated from healthy lifespan data could generalize to older MCI or demented individuals, we used transfer learning again to fine-tune the model to fit data from participants in the Alzheimer's Disease Neuroimaging Initiative (ADNI). The ADNI datasets were acquired from different scanners and under different imaging conditions, so we could test whether the model is affected by different scanners. To validate our operationalization of CR in all three cohorts, we hypothesized that the estimated CR would correlate with education and IQ (i.e., a well-established CR proxy).

Method and Material

Participants

CR/RANN: 495 healthy adults (initially aged 20–80) were drawn from our ongoing studies at Columbia University Irving Medical Center: The Reference Ability Neural Network

(RANN) study and the Cognitive Reserve (CR) study (Stern, 2009; Stern et al., 2014). Demographic characteristics of these participants are summarized in Table 1.

Subjects were recruited primarily by randomized market mailing. An initial telephone screening determined whether participants met basic inclusion criteria (i.e., right-handed, English speaking, no psychiatric or neurological disorders, and normal or corrected-to-normal vision). Potentially eligible participants were further screened in person with structured medical and neuropsychological evaluations to ensure that they had no neurological or psychiatric conditions, cognitive impairment, or contraindication for MRI scanning. Global cognitive functioning was assessed with the Mattis Dementia Rating Scale (Lucas et al., 1998), on which a minimum score of 130 was required for retention in the study. In addition, participants who met diagnostic criteria for MCI were excluded. The studies were approved by the Internal Review Board of the College of Physicians and Surgeons of Columbia University. Initially, 561 participants were enrolled to CR/RANN study, and 532 participants had memory composite score, and 37 participants (7%) were further excluded due to missing T1images, resulting 495 participants. All 495 images passed QC.

CR/RANN Memory Tasks: all participants performed Selective Reminding Task (SRT) (Buschke & Fuld, 1974). Three memory measures were based on sub-scores of the SRT: the long-term storage sub-score, continuous long-term retrieval, and the number of words recalled on the last trial. The z-scores of each of the three measures were computed by subtracting the sample means followed by dividing by the sample standard deviation. The composite memory scores were computed as the average of the three z-scores.

HCPA: 620 healthy participants with available cognitive data (age 36–100) from the lifespan Human Connectome Project Aging were included in this study (Bookheimer et al., 2019). The demographic information for the participants was presented in Table 1. HCPA excludes participants who have been diagnosed and treated for major psychiatric disorders (e.g., schizophrenia, bipolar disorder) or neurological disorders (e.g., stroke, brain tumors, Parkinson's Disease). To be included in the current study, the following measurements had to be available: 1) T1-weighted MRI scans from 3T scanner, 2) years of education and recent occupation, and 3) Composite episodic memory score.

HCPA Memory Tasks: The cognitive and performance battery includes episodic memory measured by Picture Sequence Memory Test and Rey Auditory Verb al Learning Test (RAVLT). The z-scores of each of the three measures were computed by subtracting the sample means followed by dividing by the sample standard deviation. The composite memory scores were computed as the average of the three z-scores.

ADNI: 941 subjects, including 417 normal control (CN), 378 mild cognitive impairment (MCI), and 146 Alzheimer's disease (AD), were included from this study. The demographic information for the participants is presented in Table 1 and Table 3.

Detailed inclusion and exclusion criteria for the ADNI study can be found at adni.loni.usc.edu. To be included in the current study, the following measurements had

to be available: 1) T1-weighted MRI scans from 3T scanner, 2) years of education and recent occupation, and 3) Composite memory score (Crane et al., 2012). Written informed consent was obtained from all study participants according to the Declaration of Helsinki, and Ethical approval for data collection and sharing was given by the institutional review boards of the participating institutions in the ADNI.

ADNI Memory Tasks: ADNI memory was measured using modern psychometric approaches to analyze Rey Auditory Verbal Learning Test (RAVLT, 2 versions), AD Assessment Schedule – Cognition (ADAS-Cog, 3 versions), Mini-Mental State Examination (MMSE), and Logical Memory data. The composite scores were computed based on bifactor model (Crane et al., 2012). The computed data were downloaded from the ADNI website (UWNPSYCHSUM_03_26_20.csv).

Image Procedures

Neuroimaging data acquisition

RANN: Structural MRI scans were acquired on a 3.0T Philips Achieva scanner. T1weighted MPRAGE scan was acquired with a TE/TR of 3/6.5 ms and Flip Angle of 8° , in-plane resolution of 256×256 , field of view of 25.4×25.4 cm, and 165-180 slices in axial direction with slice-thickness/gap of 1/0 mm.

HCPA: Structural MRI scans were acquired from all sites using 3T Siemens Prisma scanner, and 32-channel Prisma head coil. T1-weighted images were acquired with 3D multi-echo magnetization prepared rapid gradient echo (MEMPRAGE) at 0.8 mm isotropic resolution (Harms et al., 2018). Other parameters include: TR/TI = 2500/1000, TE = 1.8/3.6/5.4/7.2 ms, flip angle of 8 deg, FOV of $256 \times 240 \times 166$ mm with a matrix size of $320 \times 300 \times 208$ slices, water excitation employed for fat suppression (to reduce signal from bone marrow and scalp fat), and up to 30 TRs allowed for motion-induced reacquisition.

ADNI: Structural MRI scans were acquired from all sites using 3T Philips, GE, and Siemens scanners. Since the acquisition protocols were different for each scanner, an image normalization step was performed by the ADNI. The imagining sequence was a 3-dimensional sagittal part magnetization prepared of rapid gradient-echo (MPRAGE). This sequence was repeated consecutively to increase the likelihood of obtaining at least one decent quality of MPRAGE image. Image corrections involved calibration, geometry distortion, and reduction of the intensity of non-uniformity applied on each image by the ADNI. More details concerning the sMRI images is available on the ADNI homepage (http://adni.loni.usc.edu/methods/mri-tool/mri-analysis/).

Neuroimaging data processing

Each subject's structural T1 scan was reconstructed using FreeSurfer v7.1.1 (http:// surfer.nmr.mgh.harvard.edu/). The accuracy of FreeSurfer's subcortical segmentation and cortical parcellation (Fischl et al., 2002) has been reported to be comparable to manual labeling. All T1 images went through an automated quality control through MRIQC (Esteban et al., 2017). For the multiple available T1 images at the same visit, we selected the images with the best quality for further analysis. For all images that passed quality

check, cross-sectional image processing was performed using FreeSurfer Version 7.1.1 (https://surfer.nmr.mgh.harvard.edu/). Region of interest (ROI)-specific cortical thickness and volume measures were extracted from the automated anatomical parcellation using the Desikan-Killiany Atlas (Desikan et al., 2006) for cortical and aseg atlas for subcortical ROIs. To test the robustness of the models (supplementary material), we also used an alternative Destrieux atlas (Destrieux, Fischl, Dale, & Halgren, 2010).

Brain memory prediction model

The memory prediction model was trained using the RANN dataset. An overview of the transfer learning method is presented in Figure 1. First, the RANN dataset was split into the training set (70%) and test set (30%) using a conditionally random method. The distributions of age and sex in the two sets were statistically identical. Cascade neural network models with all regional cortical thickness and volume from FreeSurfer as inputs were used to train the RANN dataset for memory prediction. The cascade neural network is a feedforward neural network involving connections from the input and every previous layer to the subsequent layer (Figure 2). The advantage of the model is that it accommodates the nonlinear relationship between input and output. The CNN outperforms the other common classical machine learning approaches for brain residual-based analysis and is more flexible and efficient to implement the transfer learning framework than other approaches. (Chen et al., 2020). The hyperparameters of the model, including numbers of hidden layers, numbers of neurons, penalty of regularization and types of activation function, were optimized through random search. The loss function of model optimization was specified as mean square error function optimized using gradient descent algorithm with an adaptive learning rate and constant momentum. A 10-fold cross-validation procedure was conducted within the training set to estimate the memory prediction model performance. To quantify model performance, metrics including Pearson's correlation coefficient (rho), mean absolute error (MAE) and cohen's f^2 between the predicted and true memory were calculated.

Transfer learning

To transfer the T1 imaging-based model from source domain (using RANN) to the target domain (HCPA or ADNI), we first randomly divided the whole HCPA or ADNI dataset into the tuning pool and test sets (tuning set 70%, test set 30%). The subset of the tuning pool was randomly selected to re-train the pre-trained model.

In the refined optimization procedure of the transfer learning, the optimal tuning sample, the regularization ratio (0 to 1), the loss function (i.e. mean square error), and the choice which layers were frozen (if the layer of the pre-trained model was frozen, the parameters in that layer were not updated in the fine-tuning process) were tested. The transfer learning process was optimized using an agile optimization process because it facilitated rapid prototyping and broad searching. After the tuning procedure, the transferred model was applied to the test set for model performance evaluation. We compared the performance of transfer learning approach with the TLCO, which is used for re-training the pre-trained memory prediction model by using a combination of the tuning and training sets with the site indicator. We optimized the tunning process by adopting an agile optimization method that exploit a time-saving optimizer called scaled conjugate gradient (SCG) algorithm for fast

optimization and the hyperparameter settings emulated as those of the training process in the target domain. We compared the performance of the optimized transfer learning with tuning procedure with the model applied pre-trained model without tuning. Since ADNI data was collected from multiple sites and multiple scanners, for the secondary analysis, we applied the transfer learning by the scanner manufacturers: GE, Siemens and Philips. The data in the three target domains were divided in to the tuning-pool and test sets (Siemens: tuning pool N=377, test set N=160; GE: tuning pool N=164, test set N=68; Philips: tuning pool N=124, test set N=48). Then, transfer learning was performed in the same pattern separately for three datasets. The transfer learning with cotrain (TLCO) method was used to compare the performance with transfer learning. TLCO integrated both training set from source domain and tunning set from target domain to tune the pre-trained CNN model. The TLCO approach accounts for intersite differences through statistical variance analysis. It employs statistical models to regress out site-specific differences by using statistical covariates. This approach requires the source domain data to be accessible and the data size from different sites to be balanced. The code of the transfer learning is available at https://github.com/XiZhu-CU/ Transfer-Learning-Submission.

Quantification of cognitive reserve

After establishing the memory prediction model, a person's predicted memory performance could be obtained. Structural brain features along with age and sex were included in the model as predictors (Reed et al., 2010). Race was not included in the model as a predictor because more than 93% of our targeted sample (ADNI) is non-Hispanic white. The impact of race on the model performance is presented in supplementary material. In addition, the estimated intracranial volume (eTIV) was extracted from each subject and used as a predictor. Cognitive reserve was quantified by residuals which subtract the predicted memory from the true memory. To validate our brain-based CR quantification, we performed correlation analyses between the residuals and several proxies of CR including education, occupation and IQ. For RANN and ADNI, we used National Adult Reading Test (NART) IQ, which reflects the crystallized intelligence. Occupational attainment variables (data, people, things) reflect the specific demands of an occupation. All RANN and ADNI findings were corrected for multiple comparison at p<0.01 (5 measures: education, IQ, data, people, things). Similarly, for HCPA, the NIH Toolbox was administrated provided the Crystallized composite scores which reflects the intelligence. The Crystallized Composite score is derived from performance on the Reading Recognition and the Picture Vocabulary tasks (Heaton et al., 2014). The HCPA findings were corrected for multiple comparison at p<0.025 (2 measures: education, IQ).

Results

Demographic Characteristics

Demographic and clinical characteristics are presented in Table 1. All three datasets significantly differed in age, sex, and education. Participants were older in ADNI compared with RANN and HCPA. Education was higher in HCPA subjects, compared with RANN or ADNI. IQ was not significantly different between CRNN and ADNI.

Training memory prediction modeling in the RANN dataset

The cascade neural network model (Figure 2) using 10-fold cross-validation on the RANN training set demonstrated significant linear correlation between true and predicted memory based on the chosen T1 cortical thickness and volume predictors for both training set (rho=0.6076, MAE=0.5856, cohen's f^2 =0.58) and independent test set (rho=0.3886, MAE=0.6980, cohen's f^2 =0.18) (Figure 3). After random search, the model performance improved in training set (rho=0.5578, MAE=0.5792, cohen's f^2 =0.45) and test set (rho=0.3963, MAE=0.6888, cohen's f^2 =0.19).

There was significant correlation of NART IQ with residuals for training set (NART IQ: rho=0.154, p-value =.004, cohen's f^2 =0.01). There was significant correlation between the residuals and both NART IQ (NART IQ: rho=0.169, p-value =.003, cohen's f^2 =0.03) and education (rho=0.2069, p-value=0.01, cohen's f^2 =0.04) for test set. Residuals were not associated with data, people or things.

Transfer learning to HCPA

The best model trained using RANN dataset (pre-trained model) was used in this analysis. First, we tuned the model using tuning set from target domain (HCPA). We found linear correlation and low MAE between true and predicted memory for tuning set (rho=0.4909, MAE=0.4101, cohen's f^2 =0.32) and test set (rho=0.4062, MAE=0.4107, cohen's f^2 =0.20). When we directly applied pre-trained model without tuning, the performance dropped in test set (rho=0.3099, MAE=0.5358, cohen's f^2 =0.11) (Figure 4). Second, the transfer learning with cotrain (TLCO) approach uses both training set from source domain (RANN) and tunning set from target domain (HCPA) to further tune the pre-trained model. The TLCO performed comparable with the transfer learning approach (Tuning set: rho=0.3872, MAE=0.4318, cohen's f^2 =0.18; Test set: rho=0.4474, MAE=0.3867, cohen's f^2 =0.25).

There was significant correlation of both IQ and education with residuals of the transfer learning model for both tuning set (IQ: rho=0.227, p-value <.001, cohen's f^2 =0.05; education: rho=0.255, p-value=0.0015, cohen's f^2 =0.07); IQ: rho=0.3612, p-value <.001, cohen's f^2 =0.15; education: rho=0.2798, p-value<.001, cohen's f^2 =0.09).

Transfer learning to ADNI

1. **Primary analysis**—We found strong linear correlation and low MAE between true and predicted memory for tuning set (rho=0.7385, MAE=0.4935, cohen's f^2 =1.2) and test set (rho=0.7117, MAE=0.5435, cohen's f^2 =1.03). When we directly applied the pre-trained model without tuning, performance dropped in test set (rho=0.5485, MAE=0.9259, cohen's f^2 =0.43) (Figure 5). The TLCO performed comparable with the transfer learning approach (Tuning set: rho=0.7187, MAE=0.5158, cohen's f^2 =1.1; Test set: rho=0.6684, MAE=0.5967, cohen's f^2 =0.81).

There was significant correlation between IQ, education, and residuals of the transfer learning model for both tuning set (IQ: rho=0.2025, p-value < .001, cohen's f^2 =0.04; education: rho=0.1698, p-value=0.0032, cohen's f^2 =0.03) and test set (IQ: rho=0.366, p-value < .001, cohen's f^2 =0.15; education: rho=0.255, p-value < .001, cohen's f^2 =0.07).

We further assessed the correlation of IQ and education with residuals separately within each diagnosis group (CN, MCI and AD). Correlations of NART IQ and education with residuals are presented in Table 4. Significant correlation between IQ and residuals was found in all three groups, while the significant correlation between education and residuals was found in CN and MCI.

2. Secondary analysis—We found strong linear correlation and low MAE between true and predicted memory in the Siemens, GE and Philips respectively for tuning set (Siemens: rho=0.6488, MAE=0.0.5349; GE: rho=0.7813, MAE=0.4700, cohen's f^2 =1.57; Philips: rho=0.8264, MAE=0.3850, cohen's f^2 =2.15) and test set (Siemens: rho=0.5909, MAE=0.5163, cohen's f^2 =0.54; GE: rho=0.6558, MAE=0.5932, cohen's f^2 =0.75; Philips: rho=0.5785, MAE=0.7179, cohen's f^2 =0.50). Using the transfer learning approach, the performance of the RANN pre-trained model could be reproduced in each target domain with a smaller amount of tuning data (Figure 6). The transfer learning approach always outperformed the TLCO. The results are shown in Table 3.

Significant and positive correlations between NART IQ, education and residuals were demonstrated in both tuning and test sets (Table 5).

Discussion

In this study, we built a deep learning model to quantify the CR as residual variance in memory performance using the sMRI data from a healthy lifespan cohort (age 20–80). Importantly, our study demonstrates that the pre-trained model constructed using the healthy lifespan data (RANN) from a single-site and a single sequence was able to generalize to two target datasets acquired with different age ranges, imaging protocols, and clinical status. These included healthy lifespan Human Connectome Project-Aging cohort (HCPA) and older MCI and demented participants from Alzheimer's Disease Neuroimaging Initiative (ADNI) across different scanner types. By tuning the models with relatively small sample sizes and the same T1 brain features, optimal transferred models were obtained with satisfactory prediction performance in both target cohorts. The estimated CR was also validated by showing significant correlation with CR proxies such as education and IQ across all three datasets.

We found that the cascade neural network (CNN) model trained on the RANN data demonstrated a linear correlation between true and predicted memory based on the T1 cortical thickness and volume predictors. The sMRI-based measure of CR was associated with CR the proxy measures of education and IQ. Previous studies have used sMRI from older healthy subjects (Sole-Padulles et al., 2009), older MCI, or patients with AD to quantify CR (van Loenhoud et al., 2017). However, patients with neurological diseases with aberrant cognition may lead to bias for the quantification of CR. We first demonstrated that using lifespan data of healthy individuals, enabled good quantification of cognitive performance. It is worth noting that the performance achieved by our model is also comparable to that of previous studies applying residual approaches on quantifying CR (Vieira, Pinaya, & Mechelli, 2017).

Second, to test the generalizability of the sMRI-based deep learning model, this study utilized the transfer learning approach to fine-tune the pre-trained deep learning model to an independent, healthy lifespan HCPA data. The transfer learning approach is an efficient and stabilized way to generalize the T1 imaging-based memory prediction model. Compared with the TLCO, the tuning methods with the transfer learning approach always provided lower MAE and a stronger correlation between the actual and predicted memory in all results.

Third, the model not only could generalize from healthy lifespan data to an independent healthy lifespan HCPA dataset, but also to an older demented participants from ADNI using transfer learning. Although the three datasets administrated different tests to assess memory, by tunning the models with relatively small sample size, prediction performance of the models were relatively comparable. Moreover, the models were robust across different scanners. When conducting retrospective multi-center imaging studies, such as ADNI, or applying models trained on one site to another, heterogeneous MRI data from different scanner hardware, and acquisition protocols will pose challenges in the evaluation and generalization of these trained models. Structured programs aimed at standardizing and harmonizing MRI acquisition in research settings (Weiner et al., 2017). However, data obtained in these selected frameworks might not be representative of real-world populations. In our work, CNN was trained, tested using the RANN dataset, then using transfer learning to fine-tune and test in another two datasets obtained by different MR protocols and scanners to capture the full spectrum of heterogeneity among data and provide a less dataset-specific approach. Through further training iterations, the pre-trained CNN network adjusted for data bias stemming from the differences in acquisition and reconstruction between different scanners. In fact, our approach overcomes the caveats of previous work, which obtained data from single-center datasets leading to a limited reproducibility of findings (Wen et al., 2020).

Interestingly, In the residual-based cognitive reserve (CR) or brain age gap (BAG) literature, a correlation between residuals and the outcome (e.g., age or cognition) has been observed (for example, (Anaturk et al., 2021; Le et al., 2018)). It is still an ongoing investigation in the cause of remaining correlation with age or cognition in the residual-based measures. It is potentially due to the choice of the imaging modality, mis-specified models, and many other unknown causes. Some studies suggested re-residualizing the outcome (e.g., age, cognition) to remove the correlation (Le et al., 2018), but it is also criticized by others (Butler et al., 2021), because it can inflate residual-based measure's relative quantity. At present, there are no established and validated remedies, and it is out of the scope of the current manuscript.

In the current study, we used a standard pipeline to process the raw MRI image and extracted the cortical thickness and volume measures from T1-weighed MRIs. Our ROIbased approach shown promising and robust results for the given sample sizes. Further deep learning studies with larger sample size may also consider using voxel-wise whole brain based approach as input. Moreover, despite progress on the interpretability of deep learning, deep neural networks are still considered, to a large extent, as black boxes, due to the difficulty of interpreting their inner networks. For example, even when an model allows detection of patients from controls with high levels of accuracy, it can be difficult to

establish the specific features that informed the classification decision (Cruz-Roa, Arevalo Ovalle, Madabhushi, & Gonzalez Osorio, 2013). However, our focus of this study was to better predict the memory measures, further studies may develop more interpretable deep learning models to better understand the underlying neural mechanism. Lastly, we only used sMRI to assess the feasibility for CR estimation across three studies. Future studies should consider adding other MRI modalities, such as, diffusion tensor imaging (DTI), PET, and CSF biomarkers together with sMRI to improve the power of prediction as well as the accuracy of the residual in estimating CR.

Conclusions

In conclusion, we have shown the general feasibility of using deep learning to quantify cognitive reserve by leveraging lifespan healthy data. Our findings showed that brain/ cognitive function across lifespan provided good brain-based quantification of CR. Moreover, transfer learning shows promises for building robust models that can be fine-tuned and generalized to independent healthy lifespan cohort and in patients with Alzheimer's disease, also robust across different scanners with different acquisition parameters. The residuals (CR) were significantly associated with NART IQ and education across different cohorts. The transfer learning method is applicable to various brain diseases or CR proxies and may flexibly incorporate different imaging modalities making it a promising tool for scientific and clinical purposes.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Funding Statement

This work was supported by the R01AG026158, R01AG038465, and R01AG062578. Dr. Zhu was supported by K01MH122774 and Brain and Behavior Research Foundation Grant 07040.

Data collection and sharing for this project was funded by the Alzheimer's Disease Neuroimaging Initiative (ADNI) (National Institutes of Health Grant U01 AG024904) and DOD ADNI (Department of Defense award number W81XWH-12-2-0012). ADNI is funded by the National Institute on Aging, the National Institute of Biomedical Imaging and Bioengineering, and through generous contributions from the following: AbbVie, Alzheimer's Association; Alzheimer's Drug Discovery Foundation; Araclon Biotech; BioClinica, Inc.; Biogen; Bristol-Myers Squibb Company; CereSpir, Inc.; Cogstate; Eisai Inc.; Elan Pharmaceuticals, Inc.; Eli Lilly and Company; EuroImmun; F. Hoffmann-La Roche Ltd and its affiliated company Genentech, Inc.; Fujirebio; GE Healthcare; IXICO Ltd.; Janssen Alzheimer Immunotherapy Research & Development, LLC.; Johnson & Johnson Pharmaceutical Research & Development LLC.; Lumosity; Lundbeck; Merck & Co., Inc.; Meso Scale Diagnostics, LLC.; NeuroRx Research; Neurotrack Technologies; Novartis Pharmaceuticals Corporation; Pfizer Inc.; Piramal Imaging; Servier; Takeda Pharmaceutical Company; and Transition Therapeutics. The Canadian Institutes of Health Research is providing funds to support ADNI clinical sites in Canada. Private sector contributions are facilitated by the Foundation for the National Institutes of Health (www.fnih.org). The grantee organization is the Northern California Institute for Research and Education, and the study is coordinated by the Alzheimer's Therapeutic Research Institute at the University of Southern California. ADNI data are disseminated by the Laboratory for Neuro Imaging at the University of Southern California.

Reference

Alexander GE, Furey ML, Grady CL, Pietrini P, Brady DR, Mentis MJ, & Schapiro MB (1997). Association of premorbid intellectual function with cerebral metabolism in Alzheimer's disease:

implications for the cognitive reserve hypothesis. Am J Psychiatry, 154(2), 165–172. doi:10.1176/ajp.154.2.165 [PubMed: 9016263]

- Anaturk M, Kaufmann T, Cole JH, Suri S, Griffanti L, Zsoldos E, . . . de Lange AG (2021). Prediction of brain age and cognitive age: Quantifying brain and cognitive maintenance in aging. Hum Brain Mapp, 42(6), 1626–1640. doi:10.1002/hbm.25316 [PubMed: 33314530]
- Association A. s. (2021). 2021 Alzheimer's disease facts and figures. Alzheimers Dement, 17(3), 327–406. doi:10.1002/alz.12328 [PubMed: 33756057]
- Bocancea DI, van Loenhoud AC, Groot C, Barkhof F, van der Flier WM, & Ossenkoppele R (2021). Measuring Resilience and Resistance in Aging and Alzheimer Disease Using Residual Methods: A Systematic Review and Meta-analysis. Neurology, 97(10), 474–488. doi:10.1212/ WNL.000000000012499 [PubMed: 34266918]
- Bookheimer SY, Salat DH, Terpstra M, Ances BM, Barch DM, Buckner RL, . . . Yacoub E (2019). The Lifespan Human Connectome Project in Aging: An overview. Neuroimage, 185, 335–348. doi:10.1016/j.neuroimage.2018.10.009 [PubMed: 30332613]
- Borenstein AR, Copenhaver CI, & Mortimer JA (2006). Early-life risk factors for Alzheimer disease. Alzheimer Dis Assoc Disord, 20(1), 63–72. doi:10.1097/01.wad.0000201854.62116.d7 [PubMed: 16493239]
- Buschke H, & Fuld PA (1974). Evaluating storage, retention, and retrieval in disordered memory and learning. Neurology, 24(11), 1019–1025. doi:10.1212/wnl.24.11.1019 [PubMed: 4473151]
- Butler ER, Chen A, Ramadan R, Le TT, Ruparel K, Moore TM, . . . Shinohara RT (2021). Pitfalls in brain age analyses. Hum Brain Mapp, 42(13), 4092–4101. doi:10.1002/hbm.25533 [PubMed: 34190372]
- Chen CL, Hsu YC, Yang LY, Tung YH, Luo WB, Liu CM, . . . Isaac Tseng WY (2020). Generalization of diffusion magnetic resonance imaging-based brain age prediction model through transfer learning. Neuroimage, 217, 116831. doi:10.1016/j.neuroimage.2020.116831 [PubMed: 32438048]
- Crane PK, Carle A, Gibbons LE, Insel P, Mackin RS, Gross A, . . . Alzheimer's Disease Neuroimaging, I. (2012). Development and assessment of a composite score for memory in the Alzheimer's Disease Neuroimaging Initiative (ADNI). Brain Imaging Behav, 6(4), 502–516. doi:10.1007/s11682-012-9186-z [PubMed: 22782295]
- Cruz-Roa AA, Arevalo Ovalle JE, Madabhushi A, & Gonzalez Osorio FA (2013). A deep learning architecture for image representation, visual interpretability and automated basal-cell carcinoma cancer detection. Med Image Comput Comput Assist Interv, 16(Pt 2), 403–410. doi:10.1007/978-3-642-40763-5_50 [PubMed: 24579166]
- Desikan RS, Segonne F, Fischl B, Quinn BT, Dickerson BC, Blacker D, . . . Killiany RJ (2006). An automated labeling system for subdividing the human cerebral cortex on MRI scans into gyral based regions of interest. Neuroimage, 31(3), 968–980. doi:10.1016/j.neuroimage.2006.01.021 [PubMed: 16530430]
- Destrieux C, Fischl B, Dale A, & Halgren E (2010). Automatic parcellation of human cortical gyri and sulci using standard anatomical nomenclature. Neuroimage, 53(1), 1–15. doi:10.1016/j.neuroimage.2010.06.010 [PubMed: 20547229]
- Esteban O, Birman D, Schaer M, Koyejo OO, Poldrack RA, & Gorgolewski KJ (2017). MRIQC: Advancing the automatic prediction of image quality in MRI from unseen sites. PLoS One, 12(9), e0184661. doi:10.1371/journal.pone.0184661 [PubMed: 28945803]
- Fischl B, Salat DH, Busa E, Albert M, Dieterich M, Haselgrove C, . . . Dale AM (2002). Whole brain segmentation: automated labeling of neuroanatomical structures in the human brain. Neuron, 33(3), 341–355. doi:10.1016/s0896-6273(02)00569-x [PubMed: 11832223]
- Harms MP, Somerville LH, Ances BM, Andersson J, Barch DM, Bastiani M, . . . Yacoub E (2018). Extending the Human Connectome Project across ages: Imaging protocols for the Lifespan Development and Aging projects. Neuroimage, 183, 972–984. doi:10.1016/ j.neuroimage.2018.09.060 [PubMed: 30261308]
- Heaton RK, Akshoomoff N, Tulsky D, Mungas D, Weintraub S, Dikmen S, ... Gershon R (2014). Reliability and validity of composite scores from the NIH Toolbox Cognition Battery in adults. J Int Neuropsychol Soc, 20(6), 588–598. doi:10.1017/S1355617714000241 [PubMed: 24960398]

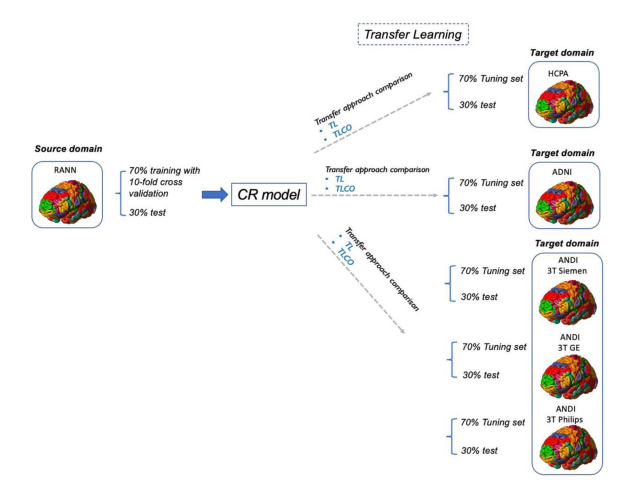
- Jones RN, Manly J, Glymour MM, Rentz DM, Jefferson AL, & Stern Y (2011). Conceptual and measurement challenges in research on cognitive reserve. J Int Neuropsychol Soc, 17(4), 593–601. doi:10.1017/S1355617710001748 [PubMed: 21411036]
- Le TT, Kuplicki RT, McKinney BA, Yeh HW, Thompson WK, Paulus MP, & Tulsa I (2018). A Nonlinear Simulation Framework Supports Adjusting for Age When Analyzing BrainAGE. Front Aging Neurosci, 10, 317. doi:10.3389/fnagi.2018.00317 [PubMed: 30405393]
- Lucas JA, Ivnik RJ, Smith GE, Bohac DL, Tangalos EG, Kokmen E, . . . Petersen RC (1998). Normative data for the Mattis Dementia Rating Scale. J Clin Exp Neuropsychol, 20(4), 536–547. doi:10.1076/jcen.20.4.536.1469 [PubMed: 9892057]
- Meng X, & D'Arcy C (2012). Education and dementia in the context of the cognitive reserve hypothesis: a systematic review with meta-analyses and qualitative analyses. PLoS One, 7(6), e38268. doi:10.1371/journal.pone.0038268 [PubMed: 22675535]
- Mueller SG, Schuff N, & Weiner MW (2006). Evaluation of treatment effects in Alzheimer's and other neurodegenerative diseases by MRI and MRS. NMR Biomed, 19(6), 655–668. doi:10.1002/ nbm.1062 [PubMed: 16986115]
- Razlighi QR, Habeck C, Barulli D, & Stern Y (2017). Cognitive neuroscience neuroimaging repository for the adult lifespan. Neuroimage, 144(Pt B), 294–298. doi:10.1016/j.neuroimage.2015.08.037 [PubMed: 26311605]
- Reed BR, Mungas D, Farias ST, Harvey D, Beckett L, Widaman K, . . . DeCarli C (2010). Measuring cognitive reserve based on the decomposition of episodic memory variance. Brain, 133(Pt 8), 2196–2209. doi:10.1093/brain/awq154 [PubMed: 20591858]
- Salthouse TA (2010). Selective review of cognitive aging. J Int Neuropsychol Soc, 16(5), 754–760. doi:10.1017/S1355617710000706 [PubMed: 20673381]
- Satz P, Cole MA, Hardy DJ, & Rassovsky Y (2011). Brain and cognitive reserve: mediator(s) and construct validity, a critique. J Clin Exp Neuropsychol, 33(1), 121–130. doi:10.1080/13803395.2010.493151 [PubMed: 20680883]
- Scarmeas N, & Stern Y (2003). Cognitive reserve and lifestyle. J Clin Exp Neuropsychol, 25(5), 625–633. doi:10.1076/jcen.25.5.625.14576 [PubMed: 12815500]
- Sole-Padulles C, Bartres-Faz D, Junque C, Vendrell P, Rami L, Clemente IC, ... Molinuevo JL (2009). Brain structure and function related to cognitive reserve variables in normal aging, mild cognitive impairment and Alzheimer's disease. Neurobiol Aging, 30(7), 1114–1124. doi:10.1016/ j.neurobiolaging.2007.10.008 [PubMed: 18053618]
- Stern Y (2002). What is cognitive reserve? Theory and research application of the reserve concept. J Int Neuropsychol Soc, 8(3), 448–460. Retrieved from https://www.ncbi.nlm.nih.gov/pubmed/ 11939702 [PubMed: 11939702]
- Stern Y (2009). Cognitive reserve. Neuropsychologia, 47(10), 2015–2028. doi:10.1016/ j.neuropsychologia.2009.03.004 [PubMed: 19467352]
- Stern Y, Gurland B, Tatemichi TK, Tang MX, Wilder D, & Mayeux R (1994). Influence of education and occupation on the incidence of Alzheimer's disease. JAMA, 271(13), 1004–1010. Retrieved from https://www.ncbi.nlm.nih.gov/pubmed/8139057 [PubMed: 8139057]
- Stern Y, Habeck C, Steffener J, Barulli D, Gazes Y, Razlighi Q, ... Salthouse T (2014). The Reference Ability Neural Network Study: motivation, design, and initial feasibility analyses. Neuroimage, 103, 139–151. doi:10.1016/j.neuroimage.2014.09.029 [PubMed: 25245813]
- Taylor JR, Williams N, Cusack R, Auer T, Shafto MA, Dixon M, . . . Henson RN (2017). The Cambridge Centre for Ageing and Neuroscience (Cam-CAN) data repository: Structural and functional MRI, MEG, and cognitive data from a cross-sectional adult lifespan sample. Neuroimage, 144(Pt B), 262–269. doi:10.1016/j.neuroimage.2015.09.018 [PubMed: 26375206]
- Tucker-Drob EM (2019). Cognitive Aging and Dementia: A Life Span Perspective. Annu Rev Dev Psychol, 1, 177–196. doi:10.1146/annurev-devpsych-121318-085204 [PubMed: 34046638]
- van Loenhoud AC, Wink AM, Groot C, Verfaillie SCJ, Twisk J, Barkhof F, ... Ossenkoppele R (2017). A neuroimaging approach to capture cognitive reserve: Application to Alzheimer's disease. Hum Brain Mapp, 38(9), 4703–4715. doi:10.1002/hbm.23695 [PubMed: 28631336]

- Vieira S, Pinaya WH, & Mechelli A (2017). Using deep learning to investigate the neuroimaging correlates of psychiatric and neurological disorders: Methods and applications. Neurosci Biobehav Rev, 74(Pt A), 58–75. doi:10.1016/j.neubiorev.2017.01.002 [PubMed: 28087243]
- Weiner MW, Veitch DP, Aisen PS, Beckett LA, Cairns NJ, Green RC, ... Alzheimer's Disease Neuroimaging, I. (2017). Recent publications from the Alzheimer's Disease Neuroimaging Initiative: Reviewing progress toward improved AD clinical trials. Alzheimers Dement, 13(4), e1–e85. doi:10.1016/j.jalz.2016.11.007 [PubMed: 28342697]
- Wen J, Thibeau-Sutre E, Diaz-Melo M, Samper-Gonzalez J, Routier A, Bottani S, . . . Lifestyle flagship study of, a. (2020). Convolutional neural networks for classification of Alzheimer's disease: Overview and reproducible evaluation. Med Image Anal, 63, 101694. doi:10.1016/ j.media.2020.101694 [PubMed: 32417716]
- Zahodne LB, Manly JJ, Brickman AM, Narkhede A, Griffith EY, Guzman VA, . . . Stern Y (2015). Is residual memory variance a valid method for quantifying cognitive reserve? A longitudinal application. Neuropsychologia, 77, 260–266. doi:10.1016/j.neuropsychologia.2015.09.009 [PubMed: 26348002]
- Zahodne LB, Manly JJ, Brickman AM, Siedlecki KL, Decarli C, & Stern Y (2013). Quantifying cognitive reserve in older adults by decomposing episodic memory variance: replication and extension. J Int Neuropsychol Soc, 19(8), 854–862. doi:10.1017/S1355617713000738 [PubMed: 23866160]
- Zissimopoulos J, Crimmins E, & St Clair P (2014). The Value of Delaying Alzheimer's Disease Onset. Forum Health Econ Policy, 18(1), 25–39. doi:10.1515/fhep-2014-0013 [PubMed: 27134606]

Page 15

Highlight:

- Quantification of cognitive reserve using brain measures for pre-symptomatic Alzheimer's patients can be estimated by leveraging lifespan data.
- Multi-center, multi-scanner, multi-sequence can affect the performance of the quantification.
- Leveraging lifespan data from a single site can improve the performance.
- Transfer learning allows the pre-trained network to successfully reconstruct the dataset acquired from different domains or age groups.





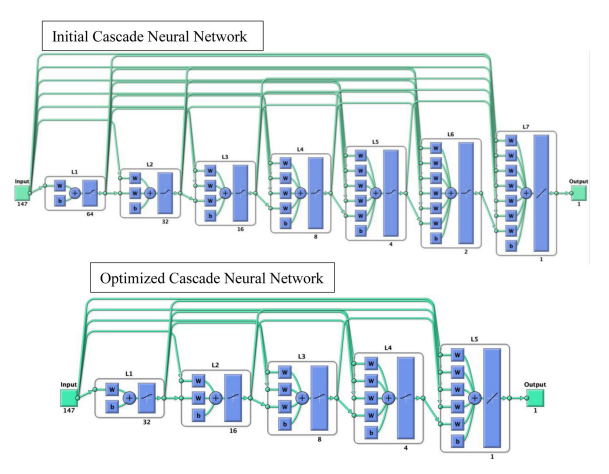


Figure 2:

the initial cascade neural network model used to train CR model (top). The cascade neural network is a feedforward neural network involving connections from the input and every previous layer to the subsequent layer. This network trained using RANN dataset has seven layers (L1 to L7). The number under each layer represents the number of neurons in that layer. The first layer has a weight coming from the input and each subsequent layer has weight coming from the input subsequent layer. The last layer is the network output, called as output layer. The output layer is also connected directly with the input layer beside with hidden layer. The hyperparameters of the model, including numbers of hidden layers and neurons, penalty of regularization, and types of activation function, were determined through random search. The optimized CNN model after random search (bottom) included 5 layers.

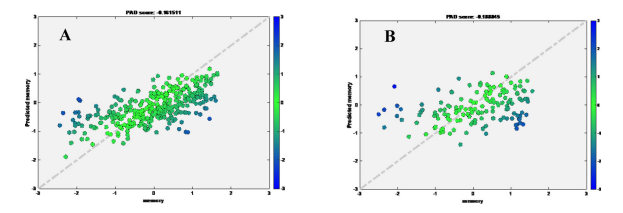


Figure 3.

Scatter plot for true memory (x axis) against predicted memory (y axis) in RANN dataset after random search.

A) Training set; B) Test set

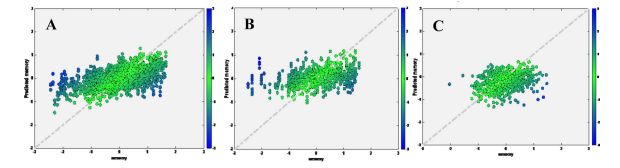


Figure 4.

Scatter plot for true memory (x axis) against predicted memory (y axis) in HCPA dataset, while applied pretrained model from RANN.

A) Tuning set using HCPA data; B) Test set after tuning using HCPA data; C) Test set if applying the pretrained model directly

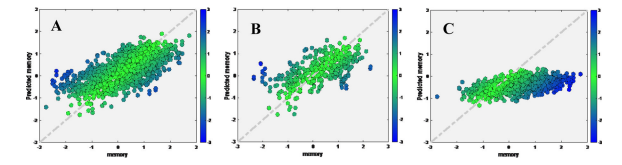


Figure 5.

Scatter plot for true memory (x axis) against predicted memory (y axis) in ADNI dataset, while applied pretrained model from RANN.

A) Tuning set; B) Test set after tuning; C) Test set if applying the pretrained model directly

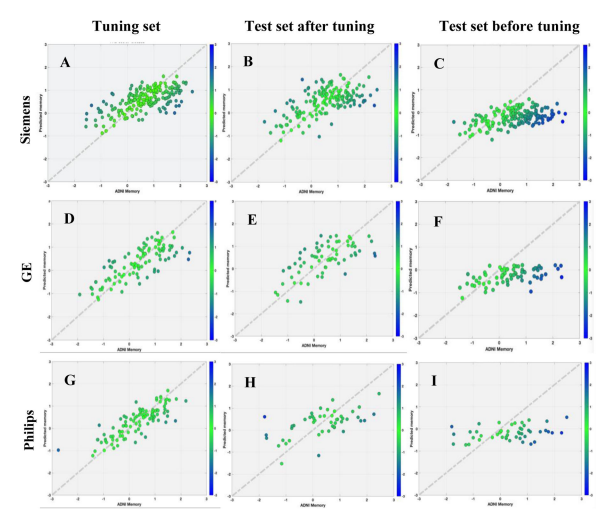


Figure 6.

Scatter plot for true memory (x axis) against predicted memory (y axis) in ADNI dataset by scanner types using random searching models.

Table 1.

Demographic Characteristics in RANN, HCPA, and ADNI Study

	RANN	НСРА	ADNI	р	
Total N	495	620	941		
Age				< 0.001	
- Mean(SD)	53.42 (16.90)	59.499 (15.171)	72.40 (7.21)		
- Median(Q1,Q3)	60.00 (38.00,67.00)	58.04 (46.92,70.98)	72.00 (67.00,77.00)		
Sex, n(%)				0.0065	
- Female	282 (57%)	358 (57.6%)	474 (50.4%)		
- Male	213 (43.0%)	263 (42.4%)	467 (49.6%)		
Memory				< 0.001	
- Mean(SD)	0.03(0.94)	0.01 (0.72)	0.45 (0.91)		
- Median(Q1,Q3)	0.13(-0.62,0.80)	-0.34 (-0.38, 0.37)	0.53 (-0.19,1.12)		
Education				< 0.00	
- Mean(SD)	16.20 (2.35)	17.46 (2.19)	16.42 (2.53)		
- Median(Q1,Q3)	16.00 (14.00,18.00)	18.00 (16, 19)	16 (15.00,18.00)		
IQ	NART IQ	Nih fluidcogcomp	NART IQ	0.512	
- N-Miss	5	2	17		
- Mean(SD)	117.02 (8.68)	120.85 (139.17)	116.35 (11.23)		
- Median(Q1,Q3)	119.20 (111.92, 124.00)	100 (91.00, 108.00)	119.44 (110.76,124.40)		
People					
- N-Miss	113	-	-		
- Mean(SD)	4.77 (2.28)	-	-		
- Median(Q1,Q3)	6.00 (3.00, 6.00)	-	-		
Data					
- N-Miss	113	-	-		
- Mean(SD)	1.73 (1.44)	-	-		
- Median(Q1,Q3)	1.00 (1.00, 3.00)	-	-		
Things					
- N-Miss	113	-	-		
- Mean(SD)	5.40 (2.40)	-	-		
- Median(Q1,Q3)	7.00 (2.00, 7.00)	-	-		
Diagnosis, n(%)					
CN	495 (100%)	620 (100%)	417 (44.3%)		
MCI	-	-	378 (40.2%)		
AD	-	-	146 (15.5%)		

Table 2.

Demographic Characteristics in ADNI dataset across three scanners

	GE (N=232)	Philips (N=172)	Siemens (N=537)	Total (N=941)	p-value
ADNI Memory					0.043
- Mean (SD)	0.38(0.95)	0.36(0.92)	0.52(0.90)	0.45(0.91)	
Diagnosis, n(%)					0.034
- CN	99(42.7%)	64(37.2%)	254(47.3%)	417(44.3%)	
- MCI	87(37.5%)	77(44.8%)	214(39.9%)	378(40.2%)	
- AD	46(19.8%)	31(18.0%)	69(12.8%)	146(15.5%)	
Age					0.803
- Mean (SD)	72.62(7.13)	72.50(6.86)	72.26(7.36)	72.40(7.21)	
Gender, n(%)					0.307
- Female	109(47.0%)	83(48.3%)	282(52.5%)	474(50.4%)	
- Male	123(53.0%)	89(51.7%)	255(47.5%)	467(49.6%)	
PT Education					0.665
- Mean (SD)	16.31(2.64)	16.38(2.58)	16.49(2.46)	16.42(2.53)	
NART IQ					0.365
- N-Miss	10	5	2	17	
- Mean (SD)	115.53 (11.55)	116.05 (11.31)	116.77 (11.07)	116.35 (11.23)	

Table 3.

Model performance for ADNI datasets by scanning manufacturers using random searching model

Manufacturers		Tuning set (TL)	Test set before tuning (TL)	Test set after tuning (TL)	Tuning set (TLCO)	Test set (TLCO)
Siemens	Rho	0.6488	0.4589	0.5909	0.7836	0.5904
	MAE	0.5349	0.8938	0.5163	0.5750	0.6244
GE	Rho	0.7813	0.5181	0.6558	0.5655	0.4636
	MAE	0.4700	0.8461	0.5932	0.6816	0.6731
Philips	Rho	0.8264	0.3138	0.5785	0.5039	0.5238
	MAE	0.3850	0.9285	0.7179	0.6406	0.6563

*transfer learning (TL), and the hybrid (TLCO) approaches

Table 4.

Pearson's correlation coefficient between IQ, education and residuals by group diagnosis in ADNI dataset

Group		IQ	p-value	Education	p-value
CN	Tuning set residuals	0.109	0.063	0.122*	0.036
	Test set residuals	0.302**	0.001	0.304**	0.001
MCI	Tuning set residuals	0.189**	0.002	0.148*	0.015
	Test set residuals	0.245*	0.011	0.112	0.244
AD	Tuning set residuals	0.355**	0.001	0.093	0.366
	Test set residuals	0.215	0.138	0.003	0.986

Table 5.

Pearson's correlation coefficient between IQ, education and residuals by scanner types in ADNI dataset

Manufacturer		IQ	p-value	Education	p-value
Siemens	Tuning set residuals	0.2364**	< .001	0.2171**	0.0020
	Test set residuals	0.1933*	0.0143	0.1858*	0.0186
GE	Tuning set residuals	0.2700**	0.0085	0.1999*	0.0461
	Test set residuals	0.0886	0.4791	0.3130*	0.0094
Philips	Tuning set residuals	0.1996*	0.0488	0.2808**	0.0047
	Test set residuals	0.5001**	<.001	0.3424**	0.0172