### CS598 - Deep Learning for Healthcare Final Report: Reproducing Alzheimer's Disease Classification

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#### Abstract

This report details the approach we took to reproduce the paper along with an evaluation of the results of our reproduced code. This also includes an extension to assess three different pipeline model approaches: 1) CNN only, 2) CNN with GNN, 3) CNN with GNN including persistent homography features. Finally, our reproduction also extends the original concept by wrapping our code in a pyHealth code hosted on gitHub and requested to fold into the core pyHealth library. We reach similar conclusions to the paper, notably that CNN with GNN is sufficient to identify and classify Alzheimer's Disease on standard high resolution MRI images without the need for more computationally intensive processing such as Topological Data Analysis (TDA).

A 4-minute video summary of our reproduction is available at

https://mediaspace.illinois.edu/media/t/1\_pkrivzew

Full source code of our reproduction is also available at https://github.com/bozlingo/cs598-DLH-Project/blob/f5e59890f40e287a1311bd13dc93541cfae95fc4/CS598\_DLH\_Project\_Code\_Final.ipynb

Our pyHealth pull request is also available at https://github.com/sunlabuiuc/PyHealth/pull/353.

**Paper Selected:** "Back to the basics with inclusion of clinical domain knowledge - A simple, scalable and effective model of Alzheimer's Disease classification"; Bruningk S., Hensel F., Lukas L., Kuijs M., Jutzeler C., Rieck B.; Proceedings of Machine Learning Research; 2021. (1)

### 1. INTRODUCTION

Around 1 in 9 people (11.3%) aged 65+ in the US have Alzheimer's Disease (AD). MRI image diagnosis with existing machine learning models relies on complex architectures that are hard to interpret and costly to scale - limiting clinical availability. Down-sampling data has been considered to increase availability - but risks loss of key features.

We reproduced the paper "Back to the basics with inclusion of clinical domain knowledge - A simple, scalable and

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effective model of Alzheimer's Disease classification" by Bruningk et al:

- 1. Appling a 3D Convolutional Neural Network (CNN) on a subset of full-resolution MRI images to measure brain tissue atrophy, particularly in the hippocampus,
- 2. Employing Topological Data Analysis (TDA) to assess anatomical brain areas and measure loss of whole-brain tissue connectivity
- 3. Extension: Comparing Grad-CAM visualization for the CNN vs. GNN embeddings to understand how the different models assessed the MRI images
- 4. Extension: Comparison of 3 different pipeline models (CNN only, CNN+GNN, CNN+GNN with persistent homography features)
- 5. Extension: Expose our code as a pyHealth task for future

Our implementation is available at with sample image data available on request (due to ADNI patient confidentiality requirements). We tested our reproduction on a subset of 20 Azheimmer's positive and 20 clinically normal MRI subjects (downscaled from the 358 used in the paper for practicality reasons).

Development was undertaken using python in a Google Colab environment. We made extensive use of the sklearn and tensorflow.keras libraries for deep learning. We also used nibabel for MRI image manipulation and ghudi for topological data analysis. We used Grad-CAM and matplotlib for visualization.

Implementation note: While the paper source code was provided (1) we felt that there would be more learning in reproducing from scratch. The description below is 100% our own reproduction albeit heavily leaning on chatGPT to support implementation.

### 2. SOURCE DATA

Image data was sourced from the Alzheimer's Disease Neuroimaging Institute (ADMI) study Image and Data Archive (IDA) which is managed by the University of Southern California. MRI images were downloaded for 20 Alzheimer's Diagnosed (AD) and 20 Clinically Normal (CN) patients according to the following criteria:

Image Type	Purpose	Usage in ADNI	Key Features
MPRAGE	High resolution structural MRI	Brain volume, segmentation, atrophy detection	3d T-1 weighted, strong grey/white matter contrast
Localizer / 3-Plane Localizer	Quick positioning scan	Aligns patient for later scans	Fast, low resolution, 3 plane acquisition
B1-Calibration Head / Body	RF field uniformity correction	Optimizes image quality and signal homogeneity	Measures B1-field inhomogeneities
Double_TSE	T2-weighted image	White matter hyperintensities, lesions	High SNR, pathology detection
Circle Scout	Positioning scan	Confirms field of view	Circular FOV, low resolution
Axial PD-T2 TSE	Proton density and T2 contrast	White matter integrity, lesion detection	PD and T2 contrast, axial slices

Figure 1: ADNI MRI Image types - MPRAGE was selected

• ADNI 1, ADNI2, ADNI GO, ADNI 3 studies

Subject Age: 77 - 78 years
Modality: T-1 weighted MRI
Diagnosis: AD and CN subjects
Original (unprocessed file types)

 Image type: MPRAGE (high resolution T1 weighted structural image); used for brain volume, segmentation and atrophy detection.

Images were manually filtered using Description notes as we found that there was low adherence to the image type definitions below. Images with description MPRAGE or MPRAGE-REPEAT only were selected.

We used ChatGPT to guide data selection and processing with two key prompts but around 20 follow-on prompts. Generally we found the LLM to be helpful but imprecise in navigating the ADNI / LDA websites; the LLM was very helpful with Python coding.

- LLM Prompt: "How can I download MRI images from ADNI?" This provided procedural steps we could take to access the data; however we still needed selfexploration to figure this out
- LLM Prompt: "How do I convert MRI format to NIFTI format?" This more focused request gave us clarity on using dcm2niix to convert the files and then babel to load them in python.

# 3. GENERATING CODE FOR DATA PROCESSING

### (3.1) Processing MRI images

Images downloaded from IDA were initially in ADNI DI-COM (DCM) format. To allow input to clinica, the images were reformatted into NIfTI format using the dcm2niix tool:

Resulting .nii.gz formatted files were moved into a single "ADNI\_Input" folder along with the labelled ADNI\_output.csv file. These files were then opened sequentially using the nibabel (NeuroImaging in Python) library. Each image was then intensity normalized into a range [0,

1] and then center cropped into shape (120, 144, 120).

These cropped images were then broken into 4 nonoverlapping 3D patches of size (30, 36, 30) representing different segments of the cropped images. These patches are used as input to the 3D CNN model (model 1) to identify areas of lower voxel intensity corresponding to areas of the brain that atrophy with Alzheimer's Disease.

### (3.2) Left and Right Hippocampus Labelling

Alzheimer's Disease (AD) is known to affect the left hippocampus to a greater degree with asymmetrical atrophy of the left vs. right hippocampus being a known indicator typical AD and limbic-predominant AD (two other AD sub-types do not exhibit this trend). To provide this view we developed a left vs. right hippocampus labelling for each normalized and cropped image using the Cerebrum Atlas Mask from the McConnel Brain Imaging Center.

We sourced the symmetric MNI-ICBM152 template was used which provided cortical and subcortical labels in NIFTI format (10). This mask was provided in original MNI space with shape (394, 466, 378), so we used scipy.ndimage.zoom to sample down to (120, 144, 120) space which aligned with our normalized cropped images. Simply multiplying the atlas mask by the normalized cropped images provided a left vs. right hippocampus label within each image.

## (3.3) Persistent Homography Features for Topological Data Analysis (TDA)

Topological Data Analysis (TDA) is used by the paper for model 2. In this approach, we employ a mathematical model to extract a graph-analog structure from each of the normalized, cropped MRI images. In this approach, we select only voxels with a minimal intensity level R, allowing low intensity 'noise' voxels to be filtered and the remaining 'high intensity' voxels to be intensity scaled across the full [0, 1] range.

Topological features are then identified using a three dimensional vector where d=0 represents 'connected components', d=1 represents 'cycles or tunnels' and d=2 represents 'voids'. By looking at changes in these features as the intensity level R changes, the model is able to identify 'persistent homology features'. This method allows us to detect latent structures in the brain with a hypothesis that looking at changes in atrophy in these structures using GNN will identify deeper underlying indications of Alzheimer's Disease.

To develop the persistent homography features we first flattened the normalized cropped images to 1D with np.flatten(). These images were then loaded into the python Gudhi (Geometry Understanding in Higher Dimensions) library, creating a complex cubical form of the image. The paper refers to DIPHA to compute persistent homology; but we found that this was incompatible with Google Colab (must be run as a command line utility) and so used the equivalent

Gudhi.persistence() method to extract persistent homography features.

# 4. GENERATING CODE FOR ALGORITHM IMPLEMENTATIONS

To generate code we made extensive use of ChatGPT with specific requests to implement sections of the original paper text. We found this method worked well - but led to significant debugging which became circular when we didn't first stop to understand / interpret the code coming from the LLM:

• LLM Prompt: "Implement the following text from original paper" Resulting text was then run, debug and evaluated (around 50 prompts to get it right)

For model training, we split the input data into training and test sets using sklearn.stratifiedKFold with n\_splits = 2 and random state shuffling. Using these splits we executed two sequential modelling steps as follows.

We implemented the training code using ChatGPT with a direct prompt to the LLM to implement the relevant text from the original paper. We found this worked very well with the code working first time with no debugging

• LLM Prompt: "Can you implement the following training pipeline text from paper?"

## (4.1) 3D CNN model development (for cropped, normalized image patches)

Model 1 of the paper involves a standard 3D Convolutional Neural Network (CNN) applied to the (30, 36, 30) shape image patches established in section 3.1. Because full 3D MRI scans can be large and noisy, we pass the patches into the 3D CNN individually to extract deep embeddings and then train a logistic regression function on these patch embeddings.

To build the 3D CNN we established a build\_3d\_cnn function which is then trained over 50 epochs with batch size=8 to generate patch embeddings for both trainset and testsets:

- Layer 1: tensorflow.keras.layers.Conv3D with 32 filters on a (3, 3, 3) 3D kernel size and initialized using HeUniform for weight initialization. Followed by batch normalization, ReLu activation and then 30
- Layer 2: tensorflow.keras.layers.Conv3D with 64 filters on a (3, 3, 3) 3D kernel size and initialized, batch normalized as above. ReLu activation and 30
- Layer 3: tensorflow.keras.layers.Conv3D with 128 filters on (3, 3, 3) 3D kernel size and initialized, batch normalized as above. ReLu activation and 30
- GlobalAveragePooling3D followed by a fully connected layer with 128 neurons and ReLu activation and 40
- The model is then wrapped into a Keras functional API model with Adam optimizer (adaptive learning rate), Binary cross-entropy loss (ideal for binary classification) and AUC as the evaluation metric

To apply logistic regression on the patch embeddings we create a sklearn LogisticRegression model with balanced class weight and liblinear solver. We train the Logistic Regression with the patch embeddings from the trainset, and then use the test embeddings to develop a prediction.

### (4.3) GNN model development

In section 3.3 we developed persistent homography features based on the TDA method with the hypothesis that this would provide more precision around atrophy within the underlying brain structures. To incorporate these features in our analysis we performed two additional steps:

- 1. Concatenate persistent homography features with the trained patch CNN encodings. This was performed with a simple np.concatenate call, applying a max tensor length of 5,000 elements for scalability.
- 2. Perform Global/Topological Context Modelling with a Graph Neural Network (GNN) to balance interactions within each patch (CNN) and interactions across patches (GNN). To create the GNN network we created GN-NPatchAggregator class which established the network and allowed us to train over 50 epochs:
  - Layer 1: torch\_geometric.nn.GCNConv using the CNN patch embeddings shape as input and outputting a 64 channel hidden layer with ReLu activation.
  - Layer 2: torch\_geometric.nn.GCNConv using the resulting hidden layer as input and outputting a 64 channel hidden layer
  - GlobalMaxPooling to input into a linear regression classifier using binary cross entropy loss function.

### 5. GENERATING CODE FOR EXTENSIONS

We used ChatGPT extensively to ideate potential extensions, and to implement the extensions themselves. ChatGPT was surprisingly creative on these fronts and led to significant acceleration in our thinking for extension.

 LLM Prompt: "What extensions would you suggest to the paper?"

## (5.1) Adapting Grad-CAM to identify which brain areas affect the GNN decision

We asked ChatGPT to suggest potential extensions for the paper - and opted to support the following suggestion:

LLM Response: Try adapting Grad-CAM or attention-based explanations to visualize which graph nodes (i.e., brain regions or patches) influenced the GNN decision the most. Compare CNN Grad-CAM vs. GNN attention: Do CNN-based saliency maps highlight the same regions as what the GNN 'cares about'? This could be very revealing about the synergy between CNNs and GNNs.

This suggestion led us to create three separate model outouts:

1. CNN + Logistic Regression: Classify AD based on local patterns within each image patch. i.e., visualize predictions per subject and overlay with ground truth

- 2. CNN  $\rightarrow$  GNN: Combine local prediction from each patch with overall pattern recognition across the entire image
- 3. CNN + Persistence  $\rightarrow$  GNN: Incorporating the TDA output to incorporating structural elements

As described in section 6.2, these models were visualized using Grad-Cam (model 1), GNNExplainer (models 2 and 3).

### (5.2) pyHealth Extension

As a further extension we wrapped our code into a pyHealth task to aid future development. To do this we first defined a custom ADNI dataset class for loading and processing ADNI data, including label extraction. We submitted this as a pyHealth pull request available via .

We also completed a pyHealth task wrapper to run an end-to-end pipeline given inputs of ADNI image data, label csv and (optionally) hippocampus atlas mask. Details can be found on

## 6. ANALYZING AND VISUALIZING REPRODUCED RESULTS

We used ChatGPT extensively to implement our evaluation and visualization code. We found signficant difficulties in this area based on the code developed above which used a Sequence Object for the CNN model. Grad-Cam support for these objects is limited which led to a 2 week long circular debugging spiral until we explicitly rewrote the CNN model code from scratch. At this point the visualization worked as intended.

• LLM Prompt: "Use Grad-CAM to visualize the MRI patches and CNN categorization" Response was a solution to visualize one patch; we followed this up with a subsequent prompt to Concatenate all MRI patches and the CNN heatmap into one image for a global view.

### (6.1) Evaluating Performance

The paper evaluated performance based on 358 subjects on a high performance GPU. For our purposes, we evaluated performance on a scaled down dataset of 20 positive (AD) and 20 negative (CN) sample MRI images. Code was executed on Google Colab (T4 GPU with 16 GB of VRAM with 1 hr limit) with source data hosted on our local Google Drive for fast data access. The code executed end to end in 52 minutes, 43 seconds.

Model training was undertaken using 4 splits of the trainset using random selection without replacement. Results of these splits are shown in Figure 2.

Similar to the paper, we found the simplest 3D Convolutional Neural Network (3D CNN) model performed the best with 0.57 accuracy and 0.66 AUC. However, Precision and Recall measures were deemed unreliable due to likely lack of balanced AD positive samples in the splits.

Furthermore, we found that applying GNN to the CNN embeddings delivered reduced performance - suggesting that the the classification is predominately local within each

	3D CNN + Logistic Regression	3D CNN → GNN	CNN + Persistent Homography → GNN
Accuracy	0.57	0.43	0.43
AUC	0.67	0.66	0.54
Precision	-	0.43	0.43
Recall	-	1.0	1.0

Figure 2: Model performance comparison

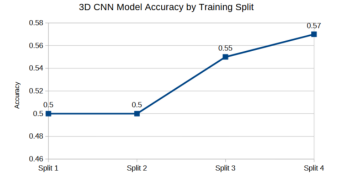


Figure 3: 3D CNN Model Training Accuracy Progression

image patch (rather than requiring pattern recognition across multiple patches.

Similar to the paper, we also found the Topological Data Analysis generated lower results than the simple 3D CNN suggesting that this method overly simplified the topology of the brain structure and therefore did not detect underlying patterns that would improve diagnosis.

Overall, our results were directionally correct - but fell short of the 86% accuracy presented in the paper. We believe this is due primarily to the reduced training dataset (238 subjects with multiple images over 10 years vs. our reduced 38 subject trainset with a single MRI image per subject). This is illustrated in figure 3 the learning rate of the model - where we saw an increasing rate of accuracy once the model saw two splits of data (20 images) - but also illustrating that the gradient of improvement was yet to flatten.

We also believe that the model results can be improved by using the Atlas Cerebral Mask to separately train the left and right hemispheres - supporting the observation that Alzheimer's disease tends to affect the left hippocampus. This is within the capabilities of our code - but required additional computational time outside of the Colab 1 hour allowance and so we do not publish results here.

### (6.2) Visualizing results for Interpretability

As with the paper, we leveraged Grad-CAM to visualize our results. To do this we had to first get the last 3d Convolution layer of the model and flatten to 2D. We then computed a heatmap that could be inputted into a matplotlib pyplot. As shown in Figure 4 below we found clear markers of Alzheimer's in the MRI patches - with a hypothesis that these would become sharper with larger training set and potentially focused training on the left hemisphere images.

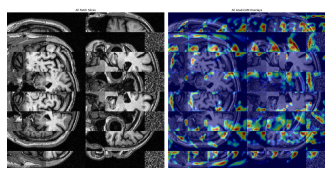


Figure 4: Original MRI image patches vs. GRAD-CAM categorization heatmap

### 7.ASSESSMENT OF REPRODUCTION

### 7.1 What was easy

Sourcing the ADNI data was relatively simple with the help of ADNI and the Image Data Archive (IDA). Reproducing the core 3D CNN and GNN models was also relatively straightforward using chatGPT. Google Colab helped enormously with code development and testing along with Google Drive to store image files in an easily accessible / low latency location.

### 7.2 What was difficult

Initial implementation required a steep learning curve to understand the different MRI image types. Sourcing the Cerebrum Atlas Mask was also ambiguous and required significant research and trial error.

During implementation we found the chatGPT model code utilized a Sequence datatype which led to significant issues with Grad-CAM. Removing this dependency (with a chatGPT prompt) resolved these issues. However, we still were unable to extract meaningful visualizations from the GNNExplainer library for models 2 and 3.

Finally, we found significant challenge around processing time for the models, and amount of image data required for the models to gain accuracy. We settled on a 38 subject sample, but estimate that this was around 1% of the scale utilized in the paper.

## 7.3 Recommendations for improving reproducibility

In retrospect we should have developed the code on our local machines with CUDA acceleration. This would have also allowed us to increase the amount of image data ingested which would have increased model accuracy. We also could have made further use of the cerebrum atlas mask to isolate the left hemisphere of the images which are known to exhibit more advanced atrophy with Alzheimer's Disease.

### 7.4 Author contributions

Tim Crosling drove the majority of the coding, source data development and report write-up. Bhavuk Jain provided support for ADNI image sourcing and preprocessing, along with quality checks throughout and analysis of the pyHealth extension options.

### **CONCLUSION**

While this project was challenging, we are happy with our paper selection on the basis that it tested multiple of the advanced concepts covered in this course. While Neuroimaging is certainly a complex field, we found a practical way to source real world MRI images from ADNI and many freely available tools to process and manipulate this data. ChatGPT was an invaluable guide in providing both initial source code generation and trouble shooting. However, we did also find that we needed to prompt ChatGPT on specific methodologies (e.g., we got into a multi-week black hole due to an initial prompt implementing the 3D CNN with a Sequence object, which was then incompatible with CAM-GRAD). From an output perspective, our project was limited by our decision to process only 40 subjects (10% of the volume covered by the paper). This was a necessary decision due to our resource constraints, but clearly has impacted the accuracy of our findings. However, our code modularity made it quite straightforward to implement our extensions and wrap the code in a pyHealth task - and we would suggest this is an improvement over the papers' authors who used a procedurally monolithic approach in their publicly available code. This is a rich topic and we see many potential extensions including: (1) Further visualization of the TDA structures detected to assess clinical applicability of the topological framework used (which would provide root understanding of the causes of Alzheimer's Disease), (2) Specialized extensions to provide diagnosis for other forms of Alzheimer's Disease which affect 14% of cases (e.g., hippocampal sparing subtype)

### References

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