

# University of Pisa

Master's degree in Artificial Intelligence and Data Engineering

Data Mining and Machine Learning project presentation

# ${\bf CogniPredictAD}$

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GitHub Repository: https://github.com/Tenshin000/CogniPredictAD/tree/main

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#### Abstract

This report introduces CogniPredictAD, a *Data Mining and Machine Learning* project designed to analyze clinical and biological parameters from the ADNI dataset, with the goal of predicting final clinical diagnoses (CN, EMCI, LMCI, AD) based on baseline data.

The analysis includes a detailed examination of preprocessing techniques: advanced missing data management, normalizations, and feature engineering. We assess the models based on their balanced accuracy and present the results through various evaluations.

Due to the ambiguity of the high predictivity of three values (CDRSB, LDELTOTAL, mPACCdigit), I built two families of models: with and without these features to reduce the risk of overfitting on ADNI.

The modeling phase includes Decision Tree, Random Forest, Extra Trees, XGBoost, LightGBM, CatBoost, Multinomial Logistic Regression, and Bagging. Hybrid sampling techniques, Grid Search for hyperparameter optimization, and cross-validation are applied. With the full set, the models achieve very high metrics (e.g., balanced accuracy  $\approx 0.926$ , ROC-AUC  $\approx 0.987$ ), while by removing the three dominant features, the LightGBM with hybrid sampling retains the best balanced performance (balanced accuracy  $\approx 0.721$ , ROC-AUC  $\approx 0.907$ ). For transparency, two optimized Decision Trees were also selected, and the trees and explanatory rules were saved.

The conclusions highlight good predictive performance on the ADNIMERGE dataset, but caution against possible sample bias and the need for external validation (by adding additional patients to the dataset or through data integration with similar datasets) before any clinical use.

## 1 Introduction

Early and accurate diagnosis of **Alzheimer's** disease (AD) is a clinical and social priority: intervening before cognitive impairment becomes severe and allows for the planning of therapies, treatments, and support strategies, and the testing of interventions that slow de-However, the disease is complex and multifactorial: clinical signs, cognitive tests, Cerebrospinal Fluid (CSF) biomarkers, genetics (e.g. APOE4), and neuroimaging measures interact in a nontrivial way. For this reason, Machine Learning (ML) techniques are particularly well-suited: they can integrate multimodal information, model nonlinear relationships, and identify combinations of features that improve the discrimination between cognitively normal (CN), mild cognitive impairment (MCI), and full-blown Alzheimer's subjects (AD).

A dataset widely used in the literature for these purposes is **ADNI**<sup>1</sup> (**Alzheimer's Disease Neuroimaging Initiative**), a multicenter longitudinal study that collects clinical, cognitive, genetic, CSF, and imaging data from USA and Canada. In this project, I

worked with the **ADNIMERGE.csv** tabular file, which is the merged version of the ADNI data and contains repeated visits over time, many clinical variables, biomarkers, and metadata. The notebooks show the entire process of building classification models and their employment.

In this project, baseline visits were selected (from 16,421 rows to 2,419 rows), extensive cleaning and imputation of missing features was performed, MRI volumes were normalized for ICV, and derived features (biological ratios and cognitive scores) were constructed. I divided the pipeline in two distinct sets with and without the three dominant cognitive features.

The modeling compared trees and ensembles, using hyperparameter optimization and sampling strategies. The **Random Forest** (with hybrid sampling) model was chosen as **Model1**. The **XGBoost** (with hybrid sampling) model, chosen as **Model2**, maintained good performance even when excluding the dominant features.

Furthermore, **XAIModel1** and **XAIModel2** represent the explainable Decision Trees for the dataset with and without the dominant features.

<sup>&</sup>lt;sup>1</sup>The Alzheimer's Disease Neuroimaging Initiative is a longitudinal, multicenter, observational study involving over 60 clinical sites in the United States and Canada. Launched in 2004 by the National Institute on Aging (NIA) in collaboration with the pharmaceutical industry, the initiative aims to develop and validate biomarkers to improve the diagnosis and monitoring of Alzheimer's disease.

## 2 Dataset

ADNIMERGE.csv is the ADNI merged table used as the main input in the notebooks: the copy used by the project contains 16,421 rows (representing visits) and 116 columns before any cleaning and selection, and incorporates repeat visits for each subject (VISCODE, EXAMDATE), identifiers (RID, PTID), and both the initial screening diagnosis (DX\_bl) and the more complete diagnosis assigned at the visit (DX).

The structure is mixed but rich: there are demographics (AGE, PTGENDER, PTEDUCAT, PTETHCAT, PTRACCAT, PTMARRY), genetics (APOE4), numerous cognitive and clinical scores (MMSE, CDRSB, ADAS11/13, LDELTOTAL, FAQ, MOCA, TRABSCOR, RAVLT..., mPACC...), CSF and PET biomarkers (ABETA, TAU, PTAU, FDG, also columns such as PIB and AV45), and MRI volumetric measures (Ventricles, Hippocampus, Entorhinal, Fusiform, MidTemp, WholeBrain, ICV). Some features are related to a single visit, while others are repeated but refer to measurements taken during the baseline visit and end with the suffix "\_bl".

ADNIMERGE.csv, however, isn't simply a concatenation: many variables are derived from source files. For example, the variable Hippocampus is derived from the sum of the left/right components (ST29SV + ST88SV) taken from the original  $FreeSurfer^2$  files.

## 3 Multiclass Problem

As I have seen, we have a different class distribution between DX-bl and DX.

- DX\_bl can be "CN", "SMC", "EMCI", "LMCI", and "AD". It indicates the screening diagnosis, i.e. the preliminary clinical judgment assigned during the first evaluation visit. It is a Screening diagnosis.
- DX can be "CN", "MCI", and "Dementia". It is instead the diagnosis assigned during the baseline visit (denoted

by *VISCODE* equal to "bl"), after a more in-depth clinical evaluation.

"AD" and "Dementia" are the same thing despite the different names.

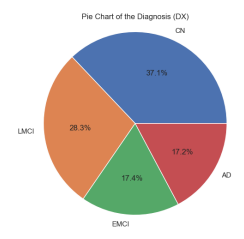
The acronym **SMC** refers to Subjective Memory Concern, i.e., cognitively normal (CN) subjects reporting perceived memory issues. Since predicting a subjective perception from objective data is not meaningful. So I reassign it based on the value it has in DX.

Furthermore, we divide MCI into EMCI (Early MCI) and LMCI (Late MCI), assuming that DX\_bl values accurately distinguish EMCI and LMCI when DX equals MCI. This is because the division into EMCI and LMCI reflects the degree of cognitive impairment and the risk of progression to dementia.

- EMCI: mild cognitive deficits, often detectable only with more sensitive tests.

  Lower or slower risk of progression to dementia.
- LMCI: more marked and evident impairment, greater impact on daily life. Higher risk of progression to Alzheimer's or other dementias.

Therefore, we decide to keep them in the diagnostic prediction.



Ultimately, our target variable will be a modified version of the *DX* column, which can now take on four values: "CN", "EMCI", "LMCI", and "AD." This makes our problem a multiclass problem.

<sup>&</sup>lt;sup>2</sup>FreeSurfer is an open-source software designed for the analysis and visualization of neuroimaging data, in particular structural (but also functional or diffusion) MRI scans, and provides a complete processing workflow.

## 4 Data Exploration

Explorations reveal that the dataset has 16,421 rows and 116 columns. However, these records represent the various visits, and we are only interested in the baseline ones. The dataset contains 2,419 useful patients (using "useful" means those who did not have a NULL baseline diagnosis) for the proposed problem.

Many columns contain significant percentages of missing cases. The diagnostic classes of the baseline sample are unbalanced, but not extremely unbalanced.

Demographic and risk analyses show bias in the ADNI sample. Ethnicity is heavily skewed toward white subjects, with high average levels of education, and many married individuals. There are more men than women, but overall the number is not disproportionate. All this, however, implies that models may perform worse on more heterogeneous clinical populations.

The *Data Exploration* was then divided into three parts:

- 1. the preliminary data exploration of the raw dataset;
- 2. the data exploration after splitting and preparing the actual dataset;
- 3. the data exploration after Preprocessing to select the classification models.

## 5 Data Preprocessing

We divided the *Preprocessing* into two phases. In the first phase, which involved preparing the dataset, we performed all the transformations and cleaning operations that did not involve the risk of *data leakage*, applying them to the entire dataset before splitting it into training and testing.

In the second phase, however, we applied the transformations that could introduce data leakage exclusively to the training set, with the sole exception of imputing missing values: in this case, the  $KNN^3$  Imputer was trained on the training set and then used for both training and testing, to ensure consistency and avoid

leakage. Preprocessing has in turn been divided into: Data Cleaning, Data Transformation, Outlier Detection and Data Reduction.

## 5.1 Data Preparation

- Selection of baseline visits only: The first clinical-operational choice was to work only on baseline visits (VISCODE == "bl"), because the goal is to predict the diagnosis based on the information collected at the first visit;
- SMC diagnosis management: Records with  $DX_-bl = SMC$  were realigned using the DX variable, but ultimately all have been classified as "CN". We already explained why in the reasons stated in the chapter "Multiclass Problem";
- Consolidating bl columns: Clean up duplicates and merge "baseline" values into "main" columns;
- Error handling: Deleted the row with RAVLT\_perc\_forgetting = -316.667 and RAVLT\_forgetting = -19;
- Text category cleaning (ethnicity, race, marriage): String values standardized to improve readability and avoid inconsistencies;
- Encoding of categorical variables: One-hot encoding (PTETHCAT, PT-MARRY), binary mapping (PTGEN-DER: Male=1, Female=0) and ordinal encoding for DX (CN=0, EMCI=1, LMCI=2, AD=3).
- Preliminary Feature Reduction: Removal of columns not relevant to the diagnosis;
- Splitting train/test: Separation into training and test sets while avoiding leakage.

### 5.2 Data Cleaning

• Handling missing values: Identifying percentages of missing values and using KNN Imputer for continuous variables;

<sup>&</sup>lt;sup>3</sup>K-Nearest Neighbors

• Numeric Value Conversion: Convert 5.5 cognitive scales and age from float to int, correcting for approximations due to imputation or format errors.

### 5.3 Data Transformation

- Creation of new CSF metrics: TAU/ABETA and PTAU/ABETA ratios more predictive than single measures according to the literature;
- MRI normalization to ICV (Intracranial Volume): Necessary to correct for differences due to gender and cranial size.

## 5.4 Outlier Detection

- Univariate Analysis: use  $IQR^4$  and Z-score threshold<sup>5</sup> for each column to find outliers;
- Multivariate Analysis: Constructs groups (EcogPt, EcogSP, Neuropsych, MRI, MRI/ICV, CSF, CSF/ABETA, mPACC), applies  $LOF^6$  and  $DBSCAN^7$  to normalized data ( $RobustScaler^8$ ). Only data points reported by both techniques and with a  $LOF\_score$  greater than 2 are kept to flag them as "extreme";
- Cleaning up problematic outliers: Outliers with values that were clearly out of range and therefore deemed highly unlikely were replaced with the mean by class.

### 5.5 Data Reduction

- Removal of redundant features: ADAS11, ADASQ4, EcogPtTotal, EcogSPTotal, mPACCtrailsB, and TAU were removed because they had a high correlation with other features and their informative value was low;
- Attribute Subset Selection: I apply four selection methods to the train: Pearson correlation ( $|r| \geq 0.6$ ), mutual information (top 25), SelectKBest with Kruskal-Wallis H-test (k=25), and Recursive Feature Elimination (RFE with Random Forest). It combines the results by counting how many times each feature appears and retains those selected at least three times, plus other features deemed useful even though they were counted less frequently.

## 5.6 Some Considerations

## 5.6.1 Correlation

The dataset contains groups of highly correlated variables (e.g., different neuropsychological scores, ECG components, and MRI volumetric measurements). Rather than eliminating them through aggressive reduction, I decided to retain them and rely on models that are intrinsically robust to correlation. This choice was motivated by two main reasons:

- 1. Clinical interpretability: Correlated variables can describe different facets of the same function or biomarker. Removing them would impoverish medical interpretation;
- 2. Complementary predictive value: Even correlated measures may contain specific variance useful for distinguishing clinical subgroups.

<sup>&</sup>lt;sup>4</sup>Interquartile Range is a statistical method to identify and handle outliers, which are extreme values that can negatively impact model accuracy. Outliers are then detected as data points falling outside a range defined by  $Q1-1.5 \cdot IQR$  and  $Q3+1.5 \cdot IQR$ .

<sup>&</sup>lt;sup>5</sup>Z-score: identifies data points that are a certain number of standard deviations away from the mean, typically flagging values with a Z-score greater than 3 or less than -3 as potential outliers.

<sup>&</sup>lt;sup>6</sup>Local Outlier Factor is based on a concept of a local density, where locality is given by k nearest neighbors, whose distance is used to estimate the density.

<sup>&</sup>lt;sup>7</sup>Density-Based Spatial Clustering of Applications with Noise identifies outliers as data points that do not belong to any dense cluster. These are points in low-density regions that are surrounded by empty space rather than other points, making them significantly different from the majority of the data.

 $<sup>{}^8</sup>x_i' = \frac{x_i - \operatorname{median}(X)}{Q_3(X) - Q_1(X)}$ 

### 5.6.2 Normalization

During the data preparation process, no global normalization or standardization was applied to all variables.

This choice was driven by one reason: I wanted to **preserve the clinical interpretability**. Maintaining variables in their original units facilitates the medical interpretation of the results and comparability with clinical reference values. Normalization would have made it more difficult to attribute direct clinical significance to the transformed values.

## 5.6.3 Binning

Binning was not applied because it would have reduced the useful information and discriminatory power of continuous variables. The models used already capture nonlinearities and thresholds, so prior discretization is unnecessary and could introduce artifacts.

## 6 Model Selection

The following classification algorithms were chosen:

- 1. **Decision Tree:** This model constructs a series of "if  $\rightarrow$  then" rules (split on individual features) to separate classes using a binary tree. Each leaf of the tree corresponds to a prediction. It was chosen because it is immediately interpretable  $(XAI^9)$ ;
- 2. Random Forest: Builds many different decision trees on subsamples of the data and averages their predictions. This reduces variance compared to a single tree and improves robustness to noise, outliers, and collinearity;
- 3. Extra Trees<sup>10</sup>: Similar to Random Forest but chooses more random splits, increasing diversity among trees and often reducing overfitting on noisy features. It was tested to compare with Random Forest and evaluate whether increased randomness improved generalization across the dataset;

- 4. **XGBoost:** A boosting algorithm that builds trees sequentially, each improving the errors of the previous one. It is highly efficient, regularized, and capable of capturing nonlinear interactions between variables, while also controlling overfitting;
- 5. **LightGBM:** A gradient boosting implementation designed to be very fast and scalable. It uses techniques (leaf-wise splitting, binning) that make it particularly efficient on heterogeneous datasets;
- CatBoost: A boosting variant that natively handles categorical variables and has robust default hyperparameters to reduce overfitting. It is suitable for working with clinical data;
- 7. Multinomial Logistic Regression: A linear model that estimates the probabilities of membership in each class using a linear combination of features. It requires feature standardization to function properly, which was ensured with StandardScaler in the pipeline. It was included as a simple and interpretable statistical baseline, useful for comparing whether the gain from complex models is consistent with a linear solution;
- 8. Bagging<sup>11</sup>: An ensemble approach that trains several models on different bootstrap samples drawn from the dataset, and then combines their outputs by averaging. The main effect is a reduction in model variance, leading to more stable predictions. It was selected because, in clinical datasets where variability and noise are substantial, bagging provides a straightforward way to assess how much predictive stability can be gained simply by aggregating multiple weak learners.

<sup>&</sup>lt;sup>9</sup>Explainable Artificial Intelligence

<sup>&</sup>lt;sup>10</sup>Extremely Randomized Trees

<sup>&</sup>lt;sup>11</sup>Bootstrap Aggregating

Table 1: Description of the final dataset attributes

Attribute	Description	Category
DX	Clinical diagnosis at the time of visit: CN, SMC, EMCI, LMCI, AD	Diagnosis
AGE	Participant's age at time of visit	Demographics
PTGENDER	Participant's gender (Male/Female)	Demographics
PTEDUCAT	Years of formal education completed	Demographics
APOE4	Number of APOE $\varepsilon 4$ alleles $(0, 1, \text{ or } 2)$ , a genetic risk	Demographics
	factor for Alzheimer's	
MMSE	Mini-Mental State Examination score (0–30, higher = better)	Clinical Scores
CDRSB	Clinical Dementia Rating - Sum of Boxes (0–18, higher = worse)	Clinical Scores
ADAS13	ADAS-Cog 13-item total score (higher = worse)	Clinical Scores
LDELTOTAL	Logical Memory II delayed recall total score	Clinical Scores
FAQ	Functional Activities Questionnaire – functional	Clinical Scores
	impairment score	Cilifical Scores
MOCA	Montreal Cognitive Assessment – global cognitive function (0–30)	Clinical Scores
TRABSCOR	Trail Making Test Part B – time in seconds (higher = worse)	Clinical Scores
RAVLT_immediate	RAVLT total immediate recall score (sum over 5 trials)	Clinical Scores
RAVLT_learning	Learning score (Trial 5 minus Trial 1 of RAVLT)	Clinical Scores
_	ing Percent forgetting from RAVLT (higher = worse)	Clinical Scores
mPACCdigit	Modified Preclinical Alzheimer's Cognitive Composite –	Composite Scores
mi 110 oaigi	Digit Symbol test	Composite Scores
EcogPtMem	Subject self-reported memory complaints (ECog)	ECogPT
EcogPtLang	Subject self-reported language difficulties (ECog)	ECogPT
EcogPtVisspat	Subject self-reported visuospatial difficulties (ECog)	ECogPT
EcogPtPlan	Subject self-reported planning difficulties (ECog)	ECogPT
EcogPtOrgan	Subject self-reported organizational issues (ECog)	ECogPT
EcogPtDivatt	Subject self-reported divided attention issues (ECog)	ECogPT
EcogSPMem	Informant-reported memory complaints (ECog)	ECogSP
EcogSPLang	Informant-reported language issues (ECog)	ECogSP
EcogSPVisspat	Informant-reported visuospatial issues (ECog)	ECogSP
EcogSPPlan	Informant-reported planning problems (ECog)	ECogSP
EcogSPOrgan	Informant-reported organization issues (ECog)	ECogSP
EcogSPDivatt	Informant-reported divided attention issues (ECog)	ECogSP
FDG	FDG PET SUVR – brain glucose metabolism	Biomarkers
PTAU/ABETA	CSF phosphorylated tau protein/A $\beta$ 42 ratio	Biomarkers
Hippocampus/ICV	Volume of hippocampus/Intracranial volume ratio from MRI	MRI
Entorhinal/ICV	Volume of the entorhinal cortex/Intracranial volume ratio from MRI	MRI
Fusiform/ICV	Fusiform gyrus volume/Intracranial volume ratio from MRI	MRI
${\rm MidTemp/ICV}$	Middle temporal gyrus volume/Intracranial volume ratio from MRI	MRI
Ventricles/ICV	Volume of ventricles/Intracranial volume ratio from MRI	MRI
${\it Whole Brain/ICV}$	Whole brain volume/Intracranial volume ratio from MRI	MRI

## 7 Hyperparameter Selection and Hybrid Sampling

## 7.1 Grid Search

To optimize the performance of the classifiers, a **Grid Search** procedure with layered cross-validation was adopted. Grid Search was chosen because it allows for a systematic and controlled exploration of the most relevant hyperparameters for each model, ensuring reproducibility and the ability to transparently compare the tested configurations.

## 7.2 Hybrid Sampling

The dataset also presented a slight imbalance in diagnostic classes, as already discussed in the "Multiclass Problem" chapter.

To address this problem, a **Hybrid Sampling strategy** was applied, combining:

- RUS<sup>12</sup> to reduce the number of instances in the majority classes, preventing the dataset from becoming excessively biased toward synthetic examples;
- 2. **SMOTENC**<sup>13</sup> to generate new synthetic examples of the minority classes, taking into account the mixed nature of the variables (continuous and categorical).

I kept the "old" dataset (the one obtained with Preprocessing) and the "new" one (the resampled one) and tried Hyperparameter Tuning on both.

# 7.3 The problem with CDRSB, LDELTOTAL, and mPACCdigit

The cognitive scores  $CDRSB^{14}$ ,  $LDELTO-TAL^{15}$  and  $mPACCdigit^{16}$  show exceptionally high predictive power compared to the rest. While this may be advantageous in terms of model accuracy, it also raises the concern of feature dominance: a small number of variables may disproportionately drive

the predictions, while many others contribute minimally. This imbalance can lead to a form of local overfitting, where models appear highly effective on the ADNI dataset but lose performance when applied to more heterogeneous clinical populations or external data.

However, this assumption cannot be verified, as it is equally possible that these three variables are genuine strong predictors of Alzheimer's diagnosis.

So the issue does not reflect a weakness of the cognitive measures themselves, but rather the possibility of dataset bias: the strength of these predictors may be tied to the specific characteristics of ADNI rather than to generalizable diagnostic patterns.

To address this, the modeling strategy should consider two complementary approaches:

- 1. building a predictive model that leverages these dominant variables;
- 2. building an alternative model that excludes them.

Hybrid Sampling will be applied to both datasets and then I will compare whether the standard models or the resampled models performs better on the test set.

## 7.4 Hyperparameter Optimization

In practice, I will end up with two models: one that can also use CDRSB, LDELTOTAL, and mPACCdigit, chosen from the standard and resampled models, and one that doesn't use CDRSB, LDELTOTAL, and mPACCdigit, chosen from the standard and resampled models.

Therefore, I need to run four Grid Searches. For each model, a large set of hyperparameter configurations was defined to explore, often with thousands of possible combinations. Optimization was conducted through five-fold cross-validation, with different metrics depending on the scenario: F1 Macro when the most

<sup>&</sup>lt;sup>12</sup>Random Under-Sampling

<sup>&</sup>lt;sup>13</sup>Synthetic Minority Over-sampling Technique for Nominal and Continuous features

 $<sup>^{14}\</sup>mathrm{Clinical}$  Dementia Rating - Sum of Boxes

<sup>&</sup>lt;sup>15</sup>Logical Memory II delayed recall total score

<sup>&</sup>lt;sup>16</sup>Modified Preclinical Alzheimer's Cognitive Composite – Digit Symbol test

predictive variables were present, Balanced Ac- is consistent i.e., worse values for these scores curacy in the most restrictive experiments, and F1 Macro.

 $F1 \; Macro^{17}$  assigns equal weight to each class and simultaneously punishes low precision or low recall. It is therefore suitable when a few strong features can "inflate" the accuracy without ensuring fairness between classes.

Balanced  $Accuracy^{18}$  is insensitive to prevalence and less dependent on precision. In the absence of dominant features, it directs the optimization to correctly retrieve all classes.

At the end of the Grid Search, the optimal parameters found for each model were used to instantiate the "final" versions of the classifiers and produce subsequent evaluations.

#### Classification 8

#### **Building Models** 8.1

From the Grid Search, the best estimators, their parameters, and scores were collected for each model. These best-estimators were saved and then retrained and evaluated via 5fold cross-validation, which produced several reports (confusion matrices, per-class metrics, accuracy, balanced accuracy, and ROC-AUC) and repeated evaluations for stability.

So we built four models for each algorithm seen before: one unsampled with the three feature, one sampled with the three feature, one unsampled without the three feature and one sampled without the three feature.

#### 8.2 **Explainability**

A method is called to generate  $SHAP^{19}$  summary plots on each model.

The three clinical scores dominate the **explanations when present**. SHAP plots calculated on models trained on the version with CDRSB, LDELTOTAL, and mPACCdigit clearly show that these three variables are the most important in absolute terms, far above the other features. The SHAP summary plots also highlight that the direction of the effect

bias the prediction toward more severe classes.

When the three scores are removed, biomarkers, demographics, and structural measures emerge. In the graphs produced on the version without those three scores, the feature ranking changes: MMSE, FAQ, MOCA, ADAS13, and EcogSPMem become extremely relevant. Also relevant are CSF (TAU/ABETA or PTAU/ABETA ratio), APOE4, and some MRI measures normalized for ICV (e.g., Hippocampus/ICV, Ventricles/ICV, WholeBrain/ICV).

The tree diagrams and exported rule files highlight that Decision Trees separate classes using thresholds on a few features (often one of the three clinical scores when present, otherwise a combination of biomarkers and cognitive tests).

The rules are therefore easily interpretable and useful for communicating "if  $\rightarrow$  then" statements to clinical staff.

The balancing effect  $(hybrid\ sampling)$  is visible in the importances and local explanations. Comparing the plots for models with and without hybrid sampling shows a modest change in the ranking: sampling tends to increase the relative importance of features that help identify less represented classes.

#### 8.3 Results

We evaluated the newly built models on the train dataset, on 5-fold-cross validation and on test dataset. However, for the final evaluation I mainly relied on the explainability and the results of the test set through the following statistics: Balanced Accuracy, F1 Score (macro), Accuracy, Precision (weighted), Recall (weighted), F1 Score (weighted), and ROC AUC (macro). The evaluation plots and confusion matrices are on the following pages, included with the evaluation tables of the statistics on the test dataset.

The following pages contain the graphs and tables that led to the final evaluation.

<sup>&</sup>lt;sup>19</sup>SHapley Additive exPlanations: A Python library for the explainability of Machine Learning models

The models ending with 1 were trained on the dataset with Hybrid Sampling, while those ending with 0 were not.

Although the models with CDRSB, LDELTOTAL, and mPACCdigit and those without CDRSB, LDELTOTAL, and mPACCdigit have the same names (e.g., XGBoost0, XGBoost1, Extra-Trees0, RandomForest1, and so on), they are actually distinct models born from distinct classifiers. Models with CDRSB, LDELTOTAL, and mPACCdigit are in the "results/all\_models/1" folder, and those without are in the "results/all\_models/2" folder.

### 8.4 Final Decision

For the dataset containing the three highly predictive cognitive scores (CDRSB, LDELTO-TAL, mPACCdigit), Random\_Forest1 (the RF trained with the hybrid sampling strategy) was chosen as the main model, and Decision\_Tree1 (the version with sampling) was chosen as the reference XAI model. This choice is motivated by very high test metrics (balanced accuracy, F1, ROC-AUC) that show the best sensitive tradeoff between classes.

Model1.pkl and XAIModel1.pkl are respectively Random\_Forest1 and Decision\_Tree1.

For the dataset where those three cognitive scores were removed, the model that maintained the best performance was **XGBoost1** (XGBoost with hybrid sampling), and, again, **Decision\_Tree1** was chosen as the XAIModel (the version built on the dataset without the three scores). This selection also comes from comparing the metrics on the test set.

Model2.pkl and XAIModel2.pkl are respectively XGBoost1 and Decision\_Tree1.

# 8.5 Comparison with the State of the Art

The ADNIMERGE.csv file is widely used in the scientific literature, with hundreds of studies explicitly citing it as the source of ADNI tabular data. Despite this, I have not come across many studies that have formulated the problem as a multiclass classification with the four labels CN, EMCI, LMCI, and AD. Most machine learning models proposed in the literature focus on binary tasks, such as CN vs. AD, CN vs. MCI, or MCI vs. AD. However, it is still possible to propose a comparison with the state of the art starting from these experimental settings.

Our results in short:

Model	Balanced Accuracy	F1 Score (weighted)	F1 Score (macro)	ROC AUC (macro)
Model1	0.9256	0.9198	0.9271	0.9256
Model2	0.7355	0.7210	0.7458	0.7355
XAIModel1	0.8988	0.8930	0.9015	0.8988
XAIModel2	0.6405	0.6521	0.6881	0.6405

- Kauppi et al. (medRxiv, 2020): Deeplearning risk-scoring pipeline using selected neurocognitive tests, achieving multiclass  $AUC \approx 0.984$ .
- Alatrany et al. (Scientific Reports, 2024): Multimodal, explainability-oriented approach on NACC. SVM reaches multiclass  $F1 \approx 90.7\%$ . Note that the data source (NACC vs. ADNI) and feature set are not directly comparable.
- Cuingnet et al. (NeuroImage, 2010): Historical ADNI benchmark (MRI-based). Multiclass performance typically 70–90% accuracy, lower than binary tasks.

Model1 is competitive with these works, Model2 favors robustness over dominant features, and XAIModels illustrate the trade-off between interpretability and accuracy. Model1 cannot be claimed superior without identical splits/preprocessing or formal statistical tests.

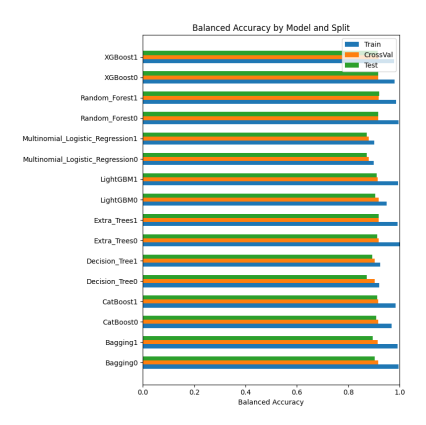


Figure 1: Evaluation of Balanced Accuracies with CDRSB, LDELTOTAL, and mPACCdigit (from folder results/all\_models/1)

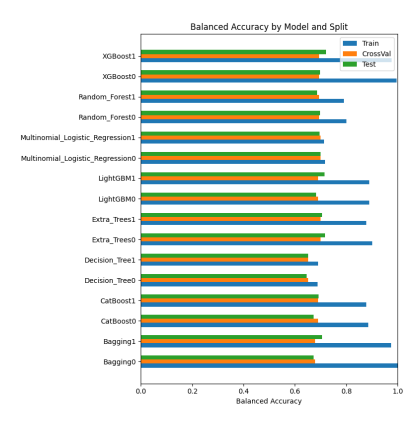


Figure 2: Evaluation of Balanced Accuracies without CDRSB, LDELTOTAL, and mPACCdigit (from folder results/all\_models/2)

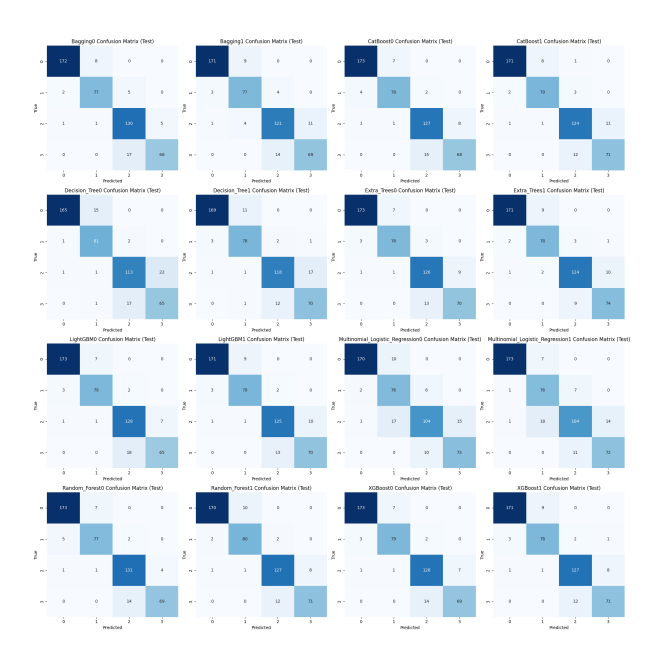


Figure 3: Confusion Matrix with CDRSB, LDELTOTAL, and mPACC digit (from folder results/all\_models/1)

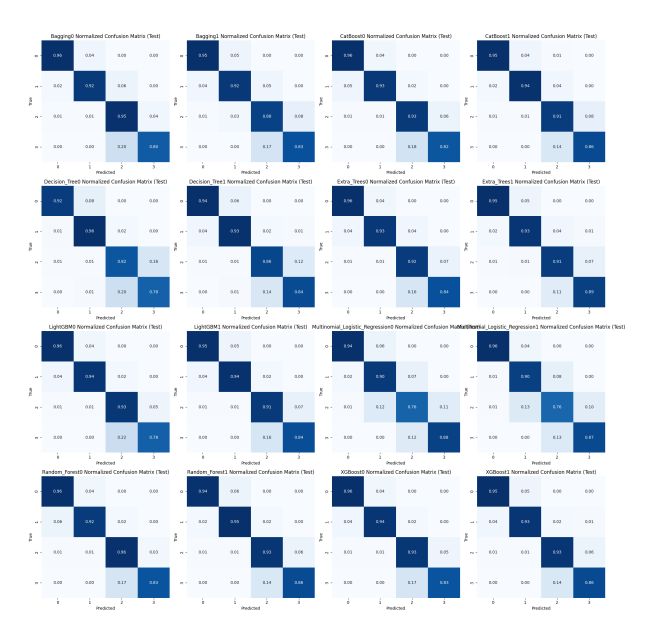


Figure 4: Normalized Confusion Matrix with CDRSB, LDELTOTAL, and mPACC digit (from folder results/all\_models/1)

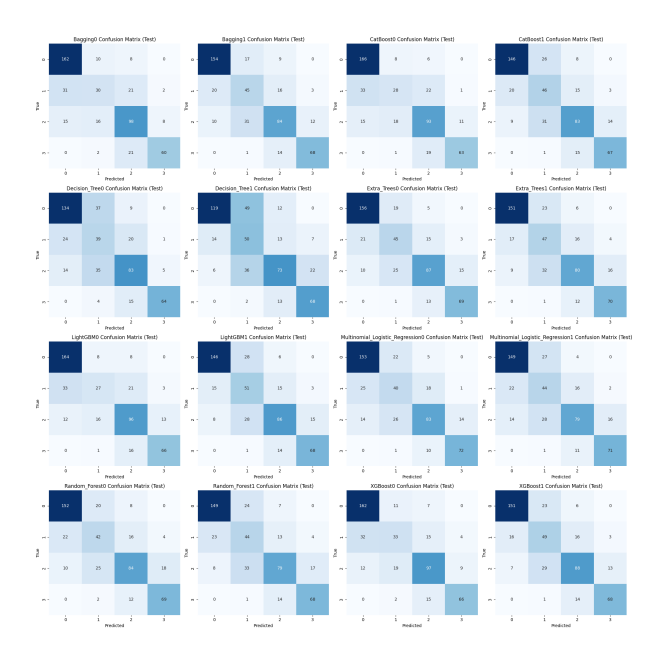


Figure 5: Confusion Matrix without CDRSB, LDELTOTAL, and mPACC digit (from folder results/all\_models/2)

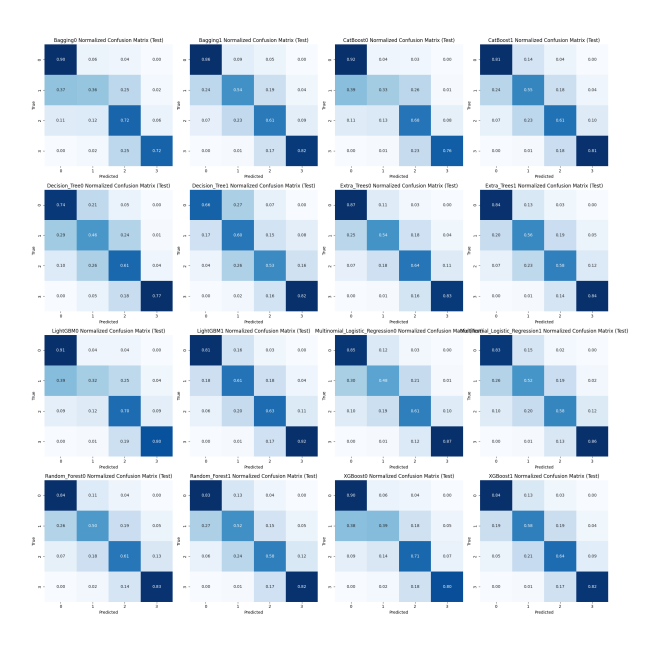


Figure 6: Normalized Confusion Matrix without CDRSB, LDELTOTAL, and mPACC digit (from folder results/all\_models/2)

 $M\_L\_R \ is \ for \ Multinomial\_Logistic\_Regression.$ 

## With CDRSB, LDELTOTAL, and mPACCdigit (from folder results/all\_models/1)

Model	Accuracy	Balanced	Precision	Recall	F1	F1	ROC
	· ·	Accu-	(weighted)	(weighted)	Score	Score	AUC
		racy			(weighted	) (macro)	(macro)
Random_Forest1	0.9256	0.9198	0.9271	0.9256	0.9258	0.9169	0.9865
$Extra\_Trees1$	0.9236	0.9188	0.9250	0.9236	0.9240	0.9143	0.9862
XGBoost0	0.9277	0.9168	0.9284	0.9277	0.9275	0.9180	0.9876
$Random\_Forest0$	0.9298	0.9163	0.9310	0.9298	0.9294	0.9205	0.9839
XGBoost1	0.9236	0.9153	0.9244	0.9236	0.9237	0.9138	0.9868
$Extra\_Trees0$	0.9236	0.9132	0.9240	0.9236	0.9236	0.9136	0.9884
CatBoost1	0.9194	0.9128	0.9205	0.9194	0.9197	0.9102	0.9875
LightGBM1	0.9194	0.9116	0.9202	0.9194	0.9195	0.9095	0.9871
CatBoost0	0.9215	0.9090	0.9219	0.9215	0.9212	0.9108	0.9887
LightGBM0	0.9194	0.9048	0.9207	0.9194	0.9189	0.9075	0.9843
Bagging0	0.9194	0.9041	0.9224	0.9194	0.9192	0.9079	0.9844
Bagging1	0.9050	0.8953	0.9062	0.9050	0.9053	0.8930	0.9835
$Decision\_Tree1$	0.8988	0.8930	0.9015	0.8988	0.8995	0.8862	0.9803
$M_LR1$	0.8781	0.8731	0.8831	0.8781	0.8785	0.8629	0.9806
$Decision\_Tree0$	0.8760	0.8722	0.8813	0.8760	0.8771	0.8615	0.9746
$M_L L_R 0$	0.8740	0.8720	0.8811	0.8740	0.8748	0.8598	0.9815

## Without CDRSB, LDELTOTAL, and mPACC digit (from folder results/all\_models/2)

Model	Accuracy	Balanced	Precision	Recall	F1	F1	ROC
		Accu-	(weighted)	(weighted)		Score	AUC
		racy			(weighted	<u>) (macro)</u>	$\frac{\text{(macro)}}{}$
XGBoost1	0.7355	0.7210	0.7458	0.7355	0.7392	0.7172	0.9071
$Extra\_Trees0$	0.7376	0.7172	0.7383	0.7376	0.7368	0.7140	0.9093
LightGBM1	0.7252	0.7163	0.7400	0.7252	0.7301	0.7098	0.9081
$Extra\_Trees1$	0.7190	0.7064	0.7285	0.7190	0.7211	0.6988	0.9068
Bagging1	0.7252	0.7059	0.7282	0.7252	0.7258	0.7043	0.9038
$M_LR0$	0.7190	0.6999	0.7188	0.7190	0.7172	0.6970	0.9071
XGBoost0	0.7397	0.6990	0.7287	0.7397	0.7314	0.7033	0.9126
$Random\_Forest0$	0.7169	0.6972	0.7173	0.7169	0.7159	0.6919	0.9053
$M_LR1$	0.7087	0.6959	0.7160	0.7087	0.7093	0.6900	0.9037
CatBoost1	0.7066	0.6930	0.7180	0.7066	0.7106	0.6894	0.9092
$Random\_Forest1$	0.7025	0.6869	0.7116	0.7025	0.7045	0.6809	0.9040
LightGBM0	0.7293	0.6821	0.7127	0.7293	0.7152	0.6827	0.9123
Bagging0	0.7231	0.6738	0.7138	0.7231	0.7131	0.6824	0.9040
CatBoost0	0.7231	0.6734	0.7089	0.7231	0.7101	0.6786	0.9093
$Decision\_Tree1$	0.6405	0.6521	0.6881	0.6405	0.6522	0.6357	0.8567
$Decision\_Tree0$	0.6612	0.6464	0.6904	0.6612	0.6726	0.6547	0.8380

## 9 Conclusions

## 9.1 Real World Applications

A CogniPredictAD application (in main.py) has been developed with customtkinter that allows you to select one of four previously saved models (Model1.pkl, Model2.pkl, XAIModel1.pkl, and XAIModel2.pkl), manually enter a set of clinical and cognitive measures, obtain a diagnostic prediction (labels 0, 1, 2, or 3 for CN, EMCI, LMCI, and AD, respectively), and display it. The user can confirm or dispute the diagnosis: both actions add a line to the data/NEWADNIMERGE.csv file (creating a folder/file if necessary), while an "Undo Last" command cancels the last saved entry. This application can be used by clinicians to collect data, evaluate prediction models, and ultimately help establish a diagnosis.

## 9.2 Final Considerations

One of the main limitations of the dataset is that, after filtering for baseline visits with nonzero diagnoses, only 2,419 patients remain, a size that limits the ability to generalize the results to external populations. While these metrics showed very high performance on the test set (for Model 1: Balanced Accuracy = 0.9198, F1 macro = 0.9169, ROC AUC (macro) = 0.9865), these metrics should be interpreted with caution.

Furthermore, three cognitive scores (CDRSB, LDELTOTAL, and mPACCdigit) provide a very strong diagnostic signal in this dataset: their presence largely explains the high effectiveness of Model 1 (Random Forest), while their removal leads to significantly lower metrics and a different distribution of important features, with XGBoost proving to be the best classifier in the pipeline without these variables. For these reasons, interpretable models (XAIModel1 and XAIModel2) and an alternative pipeline (Model2 and XAIModel2) that exclude the three scores were developed to assess the robustness and clinical plausibility of the predictions.

Furthermore, many columns have missing

Despite size limitations, the reliance on a few highly predictive cognitive scores, and the large amount of *NULL* values, the study retains methodological value and potential for application. The models can be useful as support tools (for example, risk stratification, imputation of missing diagnoses, or prioritized screening), not as a substitute for clinical assessment. Their use must be subject to external validation, calibration of the operating thresholds and post-deploy monitoring.

## 9.3 Improvements for Future Works

To improve the study, increasing the size and variability of the data is a priority: integrating future ADNI4<sup>21</sup> entries into ADNIMERGE and, if possible, compatible external cohorts would increase statistical power and generalizability.

In this context, we could also introduce a new feature that identifies patients based on their geographic area of origin. Since ADNI participants are from the United States and Canada, their area would be classified as North America. If, for example, we were able to integrate a European dataset compatible with ADNI for predicting Alzheimer's disease, we could analyze whether geographic area affects the likelihood of developing the disease. This approach could open the way to new and interesting lines of research.

It is also essential to harmonize the variables (mapping and units for imaging) and enrich the database with complementary modalities (genetics, PET, blood biomarkers) and longitudinal information, thus reducing reliance on a few cognitive scores.

However, CogniPredictAD is already a good support tool for doctors in helping with the diagnosis of Alzheimer's.

values, and the missingness pattern is often not  $MCAR^{20}$ . In fact, CSF and PET values are more often missing in healthy subjects or at certain visits. For example, ABETA, TAU, and PTAU have many missing values and are not so irrelevant in the diagnosis of Alzheimer's disease. This forces us to impute NULL values and potentially increase noise in the dataset.

<sup>&</sup>lt;sup>20</sup>Missing Completely At Random

<sup>&</sup>lt;sup>21</sup>ADNI4 is the most recent phase of the ADNI study and was initiated in 2022. Previous phases were ADNI1 (2004–2009), ADNIGO (2009–2010), ADNI2 (2011–2016), and ADNI3 (2016–2022).