

Development of multimodal machine Learning methodology for studying the heterogeneity of brain ageing using neuroimaging and genetic data

Διπλωματική Εργάσια

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ΑΪΔΙΝΗ Ν. ΓΕΩΡΓΙΟΥ



Επιβλέπουσα: Κωνσταντίνα Νικήτα Καθηγήτρια

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Εθνικό Μετσόβιο Πολυτεχνείο Σχολή Ηλεκτρολόγων Μηχανικών και Μηχανικών Υπολογιστών Τομέας Συστημάτων Μετάδοσης Πληροφορίας και Τεχνολογίας Υλικών

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(Υπογραφή)

(Υπογραφή)

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. Κωνσταντίνα Νικήτα Ανδρέας Γεώργιος Σταφυλοπάτης Γεώργιος Στάμου Καθηγήτρια Καθηγητής Καθηγητής



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Απαγορεύεται η αντιγραφή, αποθήκευση και διανομή της παρούσας εργασίας, εξ ολοκλήρου ή τμήματος αυτής, για εμπορικό σκοπό. Επιτρέπεται η ανατύπωση, αποθήκευση και διανομή για σκοπό μη κερδοσκοπικό, εκπαιδευτικής ή ερευνητικής φύσης, υπό την προϋπόθεση να αναφέρεται η πηγή προέλευσης και να διατηρείται το παρόν μήνυμα.

Το περιεχόμενο αυτής της εργασίας δεν απηχεί απαραίτητα τις απόψεις του Τμήματος, του Επιβλέποντα, ή της επιτροπής που την ενέχρινε.

ΔΗΛΩΣΗ ΜΗ ΛΟΓΟΚΛΟΠΗΣ ΚΑΙ ΑΝΑΛΗΨΗΣ ΠΡΟΣΩΠΙΚΗΣ ΕΥΘΥΝΗΣ

Με πλήρη επίγνωση των συνεπειών του νόμου περί πνευματιχών διχαιωμάτων, δηλώνω ενυπογράφως ότι είμαι αποχλειστιχός συγγραφέας της παρούσας Πτυχιαχής Εργασίας, για την ολοχλήρωση της οποίας χάθε βοήθεια είναι πλήρως αναγνωρισμένη χαι αναφέρεται λεπτομερώς στην εργασία αυτή. Έχω αναφέρει πλήρως χαι με σαφείς αναφορές, όλες τις πηγές χρήσης δεδομένων, απόψεων, θέσεων χαι προτάσεων, ιδεών χαι λεχτιχών αναφορών, είτε χατά χυριολεξία είτε βάσει επιστημονιχής παράφρασης. Αναλαμβάνω την προσωπιχή χαι ατομιχή ευθύνη ότι σε περίπτωση αποτυχίας στην υλοποίηση των ανωτέρω δηλωθέντων στοιχείων, είμαι υπόλογος έναντι λογοχλοπής, γεγονός που σημαίνει αποτυχία στην Πτυχιαχή μου Εργασία χαι χατά συνέπεια αποτυχία απόχτησης του Τίτλου Σπουδών, πέραν των λοιπών συνεπειών του νόμου περί πνευματιχών διχαιωμάτων. Δηλώνω, συνεπώς, ότι αυτή η Πτυχιαχή Εργασία προετοιμάστηχε χαι ολοχληρώθηχε από εμένα προσωπιχά χαι αποχλειστιχά χαι ότι, αναλαμβάνω πλήρως όλες τις συνέπειες του νόμου στην περίπτωση χατά την οποία αποδειχθεί, διαχρονιχά, ότι η εργασία αυτή ή τμήμα της δεν μου ανήχει διότι είναι προϊόν λογοχλοπής άλλης πνευματιχής ιδιοχτησίας.

(Υπογραφή)

Γεώργιος Αϊδίνης

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15 Ιουλίου 2022

Περίληψη

Η νόσος Αλτσχάμερ είναι το αντιχείμενο ολοένα και περισσότερων μελετών, αφού αποτελεί μια από τισ σημαντικότερες νευροεκφυλιστικές ασθένειες. Η χρήση υπολογιστικών μεθόδων για την διάγνωση, την μελέτη αλλά και την αντιμετώπιση της νόσου γνωρίζει ραγδαία ανάπτυξη. Οι σύγχρονες μέθοδοι χρησιμοποιούν πολυτροπικά δεδομένα, με ιδιαίτερη έμφαση σε απειχονιστιχά χαι γενετιχά δεδομένα. Για την αποτύπωση, την επεξεργασία, τον μετασχηματισμό, αλλά και την κατηγοριοποίηση των δεδομένων εφαρμόζονται, ως επί το πλείστον, μέθοδοι μηχανικής μάθησης. Στο πλαίσιο της παρούσας διπλωματικής εργασίας, διερευνούνται εκτενώς διαφορετικές μέθοδοι ανάλυσης δεδομένων, μηχανικής μάθησης αλλά και βαθειάς νευρωνικής μάθησης, καθώς και οι μεταξύ τους συνδυασμοί για την κατηγοριοποίηση δεδομένων από το σύνολο δεδομένων Alzheimer's Disease Neuroimaging Initiative που προέρχεται από πάσχοντες από νόσο Αλτσχάιμερ, άτομα με ήπια νοητική διαταραχή, και φυσιολογικά άτομα. Το σύνολο δεδομένων περιέχει απειχονιστιχά χαι γενετιχά δεδομένα από 1302 συμμετέχοντες. Εξετάζονται μέθοδοι ανάλυσης δεδομένων, περιλαμβανομένων των Deep Canonical Correlation Analysis, Multiple Correspondence Analysis, Orthonormal Projective Non-Negative Matrix Factorisation και Factor Analysis of Mixed Data. Για την κατηγοριοποίηση των δεδομένων χρησιμοποιήθηχαν Support Vector Machines χαθώς χαι μέθοδοι Ensemble Classifiers. Τα μοντέλα που προέχυψαν από τους συνδυασμούς των παραπάνω μεθόδων, αξιολογήθηχαν ως προς την αχρίβεια, το F1 score, χαι την εξισορροπημένη αχρίβειά τους. Παρατίθεται συγκριτική αξιολόγηση και σχολιασμός των αποτελεσμάτων καθώς και προτάσεις μελλοντικής έρευνας.

Λέξεις Κλειδιά

Νόσος Αλτσχάιμερ, Ήπια Νοητική Διαταραχή, Βαθειά Νευρωνική Μάθηση, Μηχανική Μάθηση, Data Analysis, Classification, Deep Canonical Correlation Analysis, Non-Negative Matrix Factorization, Correspondence Analysis, Radiomics

Abstract

Alzheimer's Disease (AD) is the subject of an increasing number of studies, as it is the most common cause of dementia — a continuous decline in thinking, behavioral and social skills that affects a person's ability to function independently. The use of computational methods for the diagnosis, study, and treatment of Alzheimer's disease has enjoyed rapid growth. Currently, machine learning methods are applied for the visualization, processing, transformation, and classification of data related to the disease. Modern methods utilize multi modal data, with particular emphasis on imaging and genetic data. In this Thesis, the use of a multitude of data analysis and Machine and deep learning methods is investigated in order to classify data from the Alzheimer's Disease Neuroimaging Initiative dataset into AD patients, mild cognitive impairment patients, and cognitive normal people. The dataset contains imaging and genetic data from 1302 participants. in the present Thesis, the performance of Data Analysis methods including Deep Canonical Correlation Analysis, Multiple Correspondence Analysis, Orthonormal Projective Non-Negative Matrix Factorisation and Factor Analysis of Mixed Data is presented, along with the performance of classification methods including Support Vector Machines and Ensemble Classifier methods. Various combinations of these methods were evaluated in terms of accuracy, F1 Score and balanced accuracy. The results of the study are comparatively presented and future directions are discussed.

Keywords

Alzheimer's Disease, Mild Cognitive Impairment, Deep Learning, Machine Learning, Data Analysis, Classification, Deep Canonical Correlation Analysis, Non-Negative Matrix Factorization, Correspondence Analysis, Radiomics

στους γονείς μου

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Γεώργιος Αϊδίνης

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Κεφάλαιο 1

Εκτενής Ελληνική Περίληψη

1.1 Περιγραφή του προβλήματος

Οι νευροεχφυλιστικές ασθένειες αποτελούν μια ολοένα συχνότερη αιτία θανάτου στις ανεπτυγμένες χώρες. Οι ασθένειες αυτές, όπως η νόσος Αλτσχάιμερ, αλλά και διαταραχές όπως η Ήπια Νοητική Διαταραχή, είναι συνήθεις κυρίως σε άτομα μεγαλύτερης ηλικίας, με το ηλικιακό γκρουπ των 65 χρονών και άνω να είναι αυτό που επηρεάζεται περισσότερο. Υπολογίζεται ότι μόνο στις Η.Π.Α., 6.5 εκατομμύρια άτομα πάσχουν από την νόσο Αλτσχάιμερ σήμερα, ενώ παγκοσμίως ο αριθμός αυτός μπορεί να φτάνει τα 35 εκατομμύρια. Οι εκτιμήσεις αυτές αναμένεται να αυξηθούν σε 135 εκατομμύρια παγκοσμίως, και πολλές από αυτές τις περιπτώσεις είναι αδιάγνωστες ή δεν αντιμετωπίζονται καν. ([1],[2])

Κατά τη διάρχεια της φυσιολογιχής πορείας της γήρανσης, ο ανθρώπινος εγχέφαλος εμφανίζει αλλαγές, τόσο ανατομιχές όσο και λειτουργιχές, που φαίνεται να επιταχύνονται σε ασθενείς με τέτοιου είδους ασθένειες. Ο ανθρώπινος εγχέφαλος αναπτύσσεται μέχρι την ηλιχία των 25 ετών, και στην συνέχεια χάνει συνεχώς νευριχή μάζα, οδηγώντας σε εγχεφαλιχή ατροφία, μια κατάσταση που μελετάται εχτενώς, χάρη στην πρόοδο του τομέα της ιατριχής απειχόνισης. Το φαινόμενο της ατροφίας επιταχύνεται σε μεριχούς ασθενείς αρχετά σημαντιχά, και στην περίπτωση της νόσου Αλτσχάιμερ, οδηγεί σε αποσύνθεση και απώλεια νευρώνων, και τελιχά σε θάνατο.

Τα αποτελέσματα της νόσου Αλτσχάιμερ εμφανίζουν ιδιαίτερη ποιχιλλομορφία, και είναι εμφανή όχι μόνο κλινικά, αλλά και σε απεικονιστικές σαρώσεις και γενετικές έρευνες. Λόγω της χαμηλής επίγνωσης, η ΗΝΔ και η νόσος Αλτσχάιμερ συχνά αποδίδονται λανθασμένα στη γήρανση, ενώ λόγω της ανάγκης ενός έμπειρου ιατρού για τη διάγνωση, πολλές περιπτώσεις παραμένουν αδιάγνωστες. Ωστόσο, γίνεται μεγάλη έρευνα για τη μελέτη των συμπτωμάτων, των πιθανών αιτιών, καθώς και της θεραπείας της νόσου, με ορισμένες εκτιμήσεις να ανεβάζουν το συνολικό κόστος της έρευνας για την νόσο Αλτσχάιμερ σε δεκάδες δισεκατομμύρια δολάρια. Επιπλέον, για το συνολικό κόστος για την υγειονομική περίθαλψη που σχετίζεται με ασθενείς που πάσχουν από νόσο Αλτσχάιμερ για το έτος 2020 στις ΗΠΑ έχει εκτιμηθεί ότι είναι περίπου 300 δισεκατομμύρια δολάρια. ([2], [3])

Παρά τους σημαντικούς πόρους, δεν έχει βρεθεί σαφής αιτία για την νόσο Αλτσχάιμερ, δεν έχει βρεθεί οριστική μέθοδος πρόληψης και καμία μέθοδος θεραπείας δεν ήταν ευρέως επιτυχής.

Για την αντιμετώπιση του προβλήματος της διάγνωσης της νόσου Αλτσχάιμερ καθώς και

της ΗΝΔ, καθώς και για την πρόβλεψη και τη μοντελοποίηση των αντίστοιχων πορειών τους, έχει δημοσιευθεί μια πληθώρα μελετών. Ένα κοινό χαρακτηριστικό πολλών μελετών, είναι η χρήση μηχανικών μεθόδων και μεθόδων βαθιάς μάθησης, ειδικά στα αποτελέσματα μεθόδων νευροαπεικόνισης, αλλά και στα γενετικά και στα κλινικά δεδομένα που συλλέγονται από τους ασθενείς. Οι λόγοι πίσω από αυτή την προσπάθεια είναι οι πρόοδοι στην ιατρική απεικόνιση και η συγκλονιστική αύξηση των υπολογιστικών ικανοτήτων τα τελευταία χρόνια, καθιστώντας τις πιο πολύπλοκες και καλύτερες υπολογιστικές μεθόδους εφαρμόσιμες και πρακτικές. ([4], [5])

Οι περισσότερες μελέτες χρησιμοποιούν δεδομένα νευροαπεικόνισης που συλλέχθηκαν με τις μεθόδους απεικόνισης μαγνητικού συντονισμού και τομογραφίας εκπομπής ποζιτρονίων, που συλλέχθηκαν από τη βάση δεδομένων Alzheimer's Disease Neuroimaging Initiative. Τα δεδομένα απεικόνισης συχνά συνοδεύονται από βιοδείκτες, καθώς και με απλούς νουκλεοτιδικούς πολυμορφισμούς (SNPs), ακόμη και δεδομένα από κλινικές δοκιμές που πραγματοποιούνται από εξουσιοδοτημένους ιατρούς. ([6], [7], [8]).

Ενώ γενικά τα πιο πολύπλοκα μοντέλα που χρησιμοποιούν διαφορετικούς τρόπους δεδομένων είχαν καλύτερη απόδοση, δεν υπάρχει ενοποιημένη προσέγγιση και η ποικιλομορφία των εφαρμοσμένων μοντέλων είναι χαρακτηριστική. Ωστόσο, οι περισσότερες μελέτες επικεντρώνονται είτε στην ταξινόμηση εάν ένα υποκείμενο είναι ασθενής με νόσο Αλτσχάιμερ ή όχι, και στην πρόβλεψη της πορείας της νόσου, ενώ ορισμένες εργασίες επικεντρώνονται στην πρόβλεψη εάν ένας ασθενής με ΗΝΔ θα μετατραπεί σε Αλτσχάιμερ. [4]

Ένα κοινό πρόβλημα που έπρεπε να ξεπεράσουν οι περισσότερες μελέτες ήταν η υπερπληθώρα χαρακτηριστικών που είχαν τα δεδομένα, ενώ είχαν πολύ λίγα δείγματα. Αυτό είναι γνωστό ως το πρόβλημα της «κατάρας της διαστατικότητας» και συχνά συσχετίζεται με τα βιοϊατρικά δεδομένα. Τα δεδομένα χαρακτηρίζονται από πολύ μεγάλες διαστάσεις (ειδικά στην περίπτωση της νευροαπεικόνισης), αλλά λίγα δείγματα. Αυτό έχει τη συνέπεια να μην οπτικοποιούνται εύκολα τα δεδομένα, καθώς και την αναποτελεσματικότητα στην εκπαίδευση των μοντέλων, λόγω του μικρού αριθμού δειγμάτων. Για να αποφευχθούν τα προαναφερθέντα προβλήματα, έχουν χρησιμοποιηθεί τεχνικές ανάλυσης δεδομένων, από απλές πχ. Principal Component Analysis, ή περίπλοκες,όπως η χρήση βαθιάς μάθησης, για παράδειγμα νευρωνικών δικτύων. ([9], [10], [11])

Η εργασία αυτή επιχειρεί να δημιουργήσει μια συγκριτική ανάλυση των μεθόδων μηχανικής μάθησης και των αλγορίθμων που εφαρμόζονται στο πρόβλημα της πρόβλεψης εάν ένα άτομο είναι γνωσιακά φυσιολογικό, έχει ΗΝΔ ή νόσο Αλτσχάιμερ. Τα δεδομένα συλλέχθηκαν μέσω του συνόλου δεδομένων ADNI και πιο συγκεκριμένα, δεδομένων απεικόνισης που ελήφθησαν από σαρώσεις μαγνητικής τομογραφίας T1, μαζί με γενετικά δεδομένα με τη μορφή SNP.

1.2 Δομή της Διπλωματικής Εργασίας

Η εργασία αυτή χωρίζεται σε 8 κεφάλαια, συμπεριλαμβανομένων των κεφαλαίων της εισαγωγής. Το Κεφάλαιο 3 εισάγει ορισμένες θεωρητικές έννοιες για την αντιμετώπιση του προβλήματος που περιγράφεται στο εισαγωγικό κεφάλαιο, που αφορά τον ανθρώπινο εγκέφαλο και ορισμένες βασικές αρχές σχετικά με τις μεθόδους μηχανικής και βαθιάς μάθησης που χρησιμοποιήθηκαν. Το Κεφάλαιο 4 εξηγεί τη μεθοδολογία που χρησιμοποιήθηκε, μια σύντομη περιγραφή του συνόλου δεδομένων που χρησιμοποιήθηκε, τις παραμέτρους των μεθόδων, καθώς και τις μετρήσεις για την αξιολόγησή τους. Στο κεφάλαιο 5, παρουσιάζονται ορισμένα αποτελέσματα από την προσπάθεια βελτιστοποίησης των μοντέλων που χρησιμοποιήθηκαν αργότερα για τις εργασίες ταξινόμησης και στο κεφάλαιο 6 παρουσιάζονται τα αποτελέσματα της ταξινόμησης. Τέλος, στα κεφάλαια 7 και 8, εξάγονται συμπεράσματα από παρατηρήσεις που έγιναν για τα αποτελέσματα, συζητούνται μελλοντικές επεκτάσεις και κατευθύνσεις, καθώς και οι πρακτικοί περιορισμοί αυτής της εργασίας.

1.3 Περιεχόμενα της Διπλωματικής Εργασίας

1.3.1 Θεωρητικό υπόβαθρο - Ανθρώπινος Εγκέφαλος

Καθώς το ανθρώπινο σώμα γερνά, όλα τα όργανα υφίστανται επιδράσεις που σχετίζονται με την ηλικία, όπως και ο εγκέφαλος. Τόσο φυσιολογικά όσο και γνωστικά, υπάρχουν αρκετές αλλαγές που μπορούν να παρατηρηθούν ως μέρος της φυσιολογικής διαδικασίας γήρανσης του εγκεφάλου. [12]

Μια από τις κύριες λειτουργίες του εγκέφαλου που επηρεάζεται είναι αυτή της μνήμης, με τα άτομα μεγαλύτερων ηλικιών να εμφανίζουν συχνά την τάση να ξεχνούν ονόματα αντικειμένων ή προσώπων, να επαναλάμβάνουν ερωτήσεις, να τοποθετούν λανθασμένα αντικείμενα, να χάνονται και να δυσκολεύονται να ανακαλέσουν πληροφορίες γενικά. Επιπλέον, μπορεί να επηρεαστούν οι γλωσσικές δεξιότητες, όπως το λεξιλόγιο και οι γλωσσικές δεξιότητες, καθώς και η ικανότητα εκμάθησης νέων δεξιοτήτων και πολλαπλών εργασιών. [13]

Ενώ τα προαναφερθέντα συμπτώματα είναι πολύ παρόμοια με τα συμπτώματα που παρουσιάζουν η HNΔ και η νόσος του Αλτσχάιμερ, τα φυσιολογικά μοτίβα γήρανσης αποκλίνουν από αυτά της HNΔ και της νόσου Αλτσχάιμερ. Τα αποτελέσματα της φυσιολογικής γήρανσης του εγκεφάλου χαρακτηρίζονται από την περιστασιακή τους φύση, ενώ η HNΔ και η νόσος Αλτσχάιμερ εμφανίζουν πιο συνεπή συμπτώματα, τα οποία σε ορισμένες περιπτώσεις σταδιακά επιδεινώνονται, ή συνοδεύονται από άλλα συμπτώματα άνοιας, όπως σύγχυση, αλλαγές διάθεσης και άλλα. [14]

Η ήπια γνωστική εξασθένηση είναι μια κατάσταση ενός ατόμου που χαρακτηρίζεται από προβλήματα με τη μνήμη, τη γλώσσα, τη σκέψη ή την κρίση. Συνήθως παρατηρείται μεταξύ του σταδίου της φυσιολογικής γνωστικής έκπτωσης που συμβαίνει στους ανθρώπους λόγω της γήρανσης και της δραματικής πτώσης της γνωστικής ικανότητας που είναι εμφανής στα άτομα με άνοια. Τα άτομα με MCI έχουν απώλεια μνήμης ή άλλη απώλεια γνωστικής ικανότητας, που υπερβαίνει τη φυσιολογική μείωση λόγω ηλικίας και δεν έχουν άνοια. [15]

Η νόσος του Αλτσχάιμερ είναι μια νευροεχφυλιστιχή ασθένεια που επηρεάζει τον εγχέφαλο τόσο βιολογιχά όσο χαι γνωστιχά. Αναφέρθηχε για πρώτη φορά από τον A. Alzheimer το 1906, αλλά περιγράφηχε ως ασθένεια μόλις μετά το 1910, από τον E. Kraepelin. Η νόσος Αλτσχάιμερ είναι η πιο χοινή αιτία άνοιας, ένας όρος που χρησιμοποιείται για να περιγράψει μια ομάδα συμπτωμάτων που περιλαμβάνουν μείωση της μνήμης, του συλλογισμού χαι άλλων δεξιοτήτων σχέψης. Αναφέρεται ότι η νόσος Αλτσχάιμερ ευθύνεται για περίπου το 70% των περιπτώσεων άνοιας. ([16],[17],[18])

Τόσο η νόσος Αλτσχάιμερ όσο και η HN
Δ έχουν σημαντικά πιο επιδεινωμένα συμπτώματα

από αυτά της φυσιολογικής γήρανσης, με την νόσο Αλτσχάιμερ να είναι προοδευτική, δηλαδή τα συμπτώματα της εντείνονται με το πέρασμα του χρόνου, οδηγώντας στον θάνατο. Για την ΗΝΔ και την νόσο Αλτσχάιμερ δεν υπάρχει αποδεδειγμένα αίτια, αλλά υπάρχουν ενδείξεις περιβαλλοντικών και γενετικών παραγόντων κινδύνου. Και για τις δύο περιπτώσεις, η φυσική άσκηση, η κοινωνικοποίηση και η υγιής και ισορροπημένη διατροφή έχουν συσχετιστεί με χαμηλότερα ποσοστά εμφάνισης των ασθενειών. ([19],[20], [21], [22])

Ένα μεγάλο μέρος της μελέτης, της κατανόησης και της διάγνωσης των νευροεκφυλιστικών ασθενειών είναι τεχνικές απεικόνισης του εγκεφάλου, όπως η μαγνητική τομογραφία (Magnetic Resonance Imaging). Η τεχνική MRI λειτουργεί με τη μέτρηση του ενεργειακού σήματος των (συνήθως) πυρήνων υδρογόνου, ως αποτέλεσμα διέγερσης από εξωτερικούς παλμούς ραδιοσυχνοτήτων. Πιστεύεται ότι η λειτουργία του εγκεφάλου συσχετίζεται με δομικές και λειτουργικές συνδέσεις, και ως εκ τούτου ο προσδιορισμός τυποποιημένων και αξιόπιστων μορφών συσχετίσεων είναι ζωτικής σημασίας για την κατανόηση της σύνδεσης μεταξύ της αρχιτεκτονικής του εγκεφάλου και της λειτουργίας του. ([23], [24], [25], [26])

Προχειμένου να μελετηθεί απο γενετειχής άποψης μια ασθένεια, μελετώνται μεμονωμένες αλλαγές σε νουχλεοτίδια του DNA, αποχαλούμενες Single Nucleotide Polymorphisms. Για την νόσο του Αλτσχάιμερ, οι αλλαγές αυτές εστιάζονται σε νουχλεοτίδια που βρίσχονται στα γονίδια APOE, APP, PSEN1 και PSEN2. ([27], [28], [29], [30])

1.3.2 Θεωρητικό υπόβαθρο - Βασικές μέθοδοι

Η μέθοδος Canonical Correlation Analysis είναι ένα τυπικό εργαλείο πολυμεταβλητής στατιστικής ανάλυσης που χρησιμοποιείται για την ανακάλυψη και την ποσοτικοποίηση συσχετίσεων μεταξύ δύο συνόλων μεταβλητών. Στόχος αυτής της μεθόδου είναι να βρεθεί ένας μετασχηματισμός (προβολή) των δύο συνόλων μεταβλητών, έτσι ώστε να συσχετίζονται στο μέγιστο βαθμό (μετρούμενα με συσχέτιση). Στη συνέχεια, οι προβολές βρίσκονται πραγματοποιώντας μια κοινή ανάλυση συνδιακύμανσης των δύο μεταβλητών. [31]

Η μέθοδος Deep Canonical Correlation Analysis είναι μια επέκταση της μεθόδου CCA, που στοχεύει να ανακαλύψει και να μάθει συσχετισμούς μεταξύ δύο συνόλων μεταβλητών. Η μέθοδος μπορεί να μάθει πολύπλοκες, μη γραμμικές σχέσεις μεταξύ των δύο τυχαίων διανυσμάτων και να τα μετασχηματίσει ξανά μη γραμμικά για να τα συσχετίσει, ενώ η τυπική μέθοδος CCA δεν μπορεί. Προτείνεται η χρήση των βαθιών νευρωνικών δικτύων για την ταυτόχρονη εκμάθηση δύο βαθιών μη γραμμικών αντιστοιχίσεων δύο τυχαίων μεταβλητών, καθώς τα βαθιά νευρωνικά δίκτυα έχουν αποδειχθεί ότι είναι ικανά να αναπαριστούν με ακρίβεια και αξιοπιστία μη γραμμικές συναρτήσεις που μοντελοποιούν πολύπλοκα δεδομένα πραγματικού κόσμου. ([32], [33])

Η μέθοδος βασίζεται στο πέρασμα κάθε τυχαίου διανύσματος μέσω ενός νευρωνικού δικτύου, σχεδιασμένου και εκπαιδευμένου να μετασχηματίζει το τυχαίο διάνυσμα μη γραμμικά. Αυτό δημιουργεί μια αντιστοίχιση σε έναν υπερχώρο που τα αποτελέσματα συσχετίζεται καλύτερα με την αντιστοίχιση του αντίστοιχου (μετασχηματισμένου) τυχαίου διανύσματος.

Αν θ₁ και θ₂ είναι τα διανύσματα όλων των παραμέτρων των δύο δικτύων, τότε ο στόχος της μεθόδου είναι να βρεθούν οι τιμές των διανυσμάτων έτσι ώστε:

 $(\theta_1^*, \theta_2^*) = \operatorname{argmax}_{\theta_1, \theta_2} \{ corr(f_1(X_1; \theta_1), f_2(X_2; \theta_2)) \}.$



Σχήμα 1.1: Τα δύο παράλληλα δίκτυα, μαζί με τη διαδρομή πληροφοριών (βέλη)

Η μέθοδος Multiple Correspondence Analysis (MCA) είναι μια τεχνική ανάλυσης δεδομένων που χρησιμοποιείται για την ανάλυση της δομής ενός αριθμού εξαρτημένων κατηγορικών μεταβλητών σε ένα σύνολο δεδομένων. Είναι μια επέκταση της απλής μεθόδου Correspondence Analysis και είναι παρόμοια με τη γνωστή μέθοδο PCA. Η μέθοδος MCA εκτελείται σε έναν πίνακα δείκτη - που ονομάζεται επίσης Πλήρης Διαζευκτικός Πίνακας - ή σε έναν πίνακα Burt. Μπορεί επίσης να θεωρηθεί ως η μέθοδος PCA που εφαρμόζεται στον ΠΔΠ.

Η μέθοδος Non-Negative Matrix Factorization (NMF) είναι μια μη εποπτευόμενη, πολυμεταβλητή, αναλυτική μέθοδος για την κατά προσέγγιση παραγοντοποίηση ενός πίνακα V σε δύο πίνακες W,H υπό τον περιορισμό ότι τα στοιχεία τους είναι μη αρνητικά: V = WH, έτσι ώστε $H \ge 0$ και $W \ge 0$. [34]

Μια αξιοσημείωτη επέχταση είναι αυτή της μεθόδου Orthonormal Projective NMF (OP-NMF), όπου οι συντελεστές υπολογίζονται ως οι προβολές του πίναχα στα εχτιμώμενα στοιχεία W ($H = W^{\intercal}V$), ενώ διατηρείται η ορθοχανονιχότητα στα εχτιμώμενα στοιχεία ($W^{\intercal}W = I$). Ως αποτέλεσμα, όλα τα στοιχεία συμμετέχουν στην αναχατασχευή όλων των δειγμάτων δεδομένων, πράγμα που σημαίνει ότι η επιχάλυψη μεταξύ των εχτιμώμενων στοιχείων είναι σημαντιχά χαμηλότερη, έχοντας λιγότερες παραμέτρους προς εχμάθηση, ενώ διατηρείται υψηλή αραιότητα. [35]

Παρόμοια με την μέθοδο MCA, η μέθοδος Factor Analysis of Mixed Data (FAMD) είναι μια τεχνική ανάλυσης δεδομένων που χρησιμοποιείται για την ανάλυση της δομής των μικτών δεδομένων, που σημαίνει τόσο συνεχή αριθμητικά όσο και κατηγορικά δεδομένα. Χρησιμοποιείται επίσης για τη μείωση του αριθμού των διαστάσεων του συνόλου δεδομένων και τη βελτίωση της ερμηνευσιμότητας. Βασίζεται στις μεθόδους MCA και PCA.([36], [37])

Οι μηχανές διανυσμάτων υποστήριξης Support Vector Machines - SVM είναι μια οικογένεια μοντέλων που χρησιμοποιούνται για την κατηγοριοποίηση την παλινδρόμηση. Ο στόχος ενός SVM είναι να επιλέξει ένα υπερεπίπεδο (π.χ. μια ευθεία γραμμή σε δισδιάστατο χώρο, ένα επίπεδο σε τρισδιάστατο) που διαχωρίζει καλύτερα ένα σύνολο δεδομένων που αποτελείται από επισημασμένα δείγματα που ανήκουν σε μία από τις δύο κατηγορίες. Η μέθοδος στην οποία βασίζονται τα SVM για να επιτύχουν αυτόν τον στόχο είναι η επιλογή δύο παράλληλων υπερεπίπεδων που χωρίζουν τις δύο κατηγορίες έτσι ώστε η απόσταση μεταξύ τους (το περιθώριο) να είναι μέγιστη. [38]

Η μέθοδος Ensemble Learning είναι μια τεχνική συνδυασμού πολλών μοντέλων για τη βελτίωση της απόδοσης της εργασίας, όπως προβλήματα ταξινόμησης, παλινδρόμησης ή προσέγγισης κ.λπ. Αυτό επιτυγχάνεται με την εφαρμογή των (ίσως διαφορετικών) μοντέλων στα διαθέσιμα δεδομένα (ή ένα υποσύνολο αυτών) και να συνδυάσουν τα αποτελέσματα τους, προκειμένου να γίνει καλύτερη προσπάθεια επίλυσης του προβλήματος. ([39], [40])

Ένα τέτοιο σύνολο (Ensemble) δημιουργείται συνδυάζοντας είτε διαφορετικά μοντέλα είτε μοντέλα με διαφορετικές αρχικοποιήσεις παραμέτρων και διαμορφώσεις. Τέτοια μοντέλα μπορεί να είναι σχετικά απλά, όπως δέντρα αποφάσεων, απλοί ταξινομητές bayes ή SVM, ή πιο περίπλοκα, όπως πολυστρωματικά perceptrons, ή ακόμα και άλλα σύνολα συνολικά. Μια βασική πτυχή της επιλογής του βασικού μοντέλου είναι η δημιουργία αρκετής ποικιλίας απόψεων, δηλαδή η διαφοροποίηση μεταξύ των ίδιων των μοντέλων.

Μια μέθοδος εκμάθησης συνόλου που εφαρμόζεται στο πρόβλημα της ταξινόμησης είναι αυτή του Bagging ή Bootstrap Aggregating, όπου τα βασικά μοντέλα εκπαιδεύονται σε δείγματα που έχουν σχεδιαστεί από το αρχικό σύνολο δεδομένων, με αντικατάσταση, και τα βασικά μοντέλα είναι ταξινομητές του ίδιου τύπου. Τα αποτελέσματα των μεμονωμένων ταξινομητών συνδυάζονται σε στρατηγική ψηφοφορίας απλής πλειοψηφίας για να καθοριστεί το συνολικό αποτέλεσμα του συνόλου. Ένα άλλο αξιοσημείωτο παράδειγμα είναι αυτό του Adaboost, μια έκδοση του ενισχυτικού συνόλου, προσαρμοσμένη για το πρόβλημα της ταξινόμησης πολλαπλών κατηγοριών. Όπως προηγουμένως, τα δείγματα δεδομένων εκπαίδευσης με βοοτστραππεδ προέρχονται από μια αρχικά ομοιόμορφη αλλά συνεχώς εξελισσόμενη κατανομή, διασφαλίζοντας ότι τα δείγματα που είχαν προηγουμένως λανθασμένα επισημανθεί εμφανίζονται πιο συχνά και επομένως εκπαιδεύουν τους βασικούς ταξινομητές στις πιο δύσκολες περιπτώσεις. Οι βασικοί ταξινομητές συνδυάζονται με σταθμισμένη πλειοψηφία. ([40], [41])

1.3.3 Μεθοδολογία

Για την εργασία αυτή χρησιμοποιήθηκαν δεδομένα από το ADNI σύνολο δεδομένων. Τα απεικονιστικά δεδομένα έχουν τη μορφή 145 τιμών περιοχών ενδιαφέροντος Regions of Interest (ROI), που λαμβάνονται από σαρωτές MRI είτε 1,5T είτε 3T, χρησιμοποιώντας ακολουθίες με στάθμιση T1. Προκειμένου να ληφθούν οι τιμές έντασης ROI, χρησιμοποιήθηκε η μέθοδος MUlti-atlas region Segmentation utilizing Ensembles (MUSE) δημιουργώντας τις περιοχές ενδιαφέροντος. Τα γενετικά δεδομένα έχουν την μορφή 54 τόπων ευαισθησίας, με τη μορφή με τη μορφή αριθμού αλληλόμορφων γονιδίων (0,1,2) για κάθε SNP, που έχουν εντοπιστεί από γενετικές μελέτες της νόσου Αλτσχάιμερ.

Το σύνολο δεδομένων έχει 1302 συμμετέχοντες, (56,91% άνδρες), με μέση ηλικία 75,20 ετών. για τους άνδρες άνδρες και 74,36 ε. για τις γυναίκες. Το σύνολο δεδομένων έχει 433

συμμετέχοντες που χαρακτηρίζονται ως γνωσιακά φυσιολογικοί (Cognitive Normal - CN) (33,25%), 626 ασθενείς με HNΔ (48,07%) και 243 ασθενείς με νόσο Αλτσχάιμερ (18,66%). Πραγματοποιήθηκε γραμμική παλινδρόμηση για να αφαιρεθεί οποιαδήποτε τυχόν ανεπιθύμητη επίδραση που σχετίζεται με την ηλικία, το φύλο ή το μέγεθος του εγκεφάλου. Το μοντέλο της γραμμικής παλινδρόμησης εφαρμόστηκε στην ομάδα CN και ο μετασχηματισμός εφαρμόστηκε έπειτα σε ολόκληρο το σύνολο δεδομένων.

Στη συνέχεια, εφαρμόστηκαν τεχνικές ανάλυσης δεδομένων, όπως OPNMF στα δεδομένα απεικόνισης, MCA στα γενετικά δεδομένα ή FAMD σε ολόκληρο το σύνολο δεδομένων. Αυτές οι μέθοδοι εφαρμόστηκαν τόσο στα αρχικά δεδομένα, όσο και στην έξοδο της μεθόδου DCCA, προκειμένου να συγκριθούν αργότερα. Το αποτέλεσμα καθεμιάς από αυτές τις τεχνικές αποθηκεύτηκε στη συνέχεια για μεταγενέστερο πειραματισμό. Τέλος, κάθε ένα από τα αποτελέσματα συνδυασμών των μεθόδων τροφοδοτήθηκε σε SVM καθώς και σε ταξινομητές συνόλου (ensemble classifiers).

Παρακάτω φαίνονται σχηματικά οι διαδικασίες που εφαρμόστηκαν, καθώς και η σειρά με την οποία εφαρμόστηκαν:



Σχήμα 1.2: Επισκόπηση διεργασιών που εκτελέστηκαν στα δεδομένα



Σχήμα 1.3: Διάγραμμα βημάτων που ακολουθήθηκαν

Για την μέθοδο DCCA, το σύνολο δεδομένων για τη βελτιστοποίηση του μοντέλου χωρίστηκε σε τρία σύνολα, το σετ εκπαίδευσης, επικύρωσης και δοκιμής, με το τελευταίο να διατηρείται κρυφό από το μοντέλο κατά τη φάση εκπαίδευσης, προκειμένου να διασφαλιστεί μια ακριβής πρόβλεψη σε δεδομένα που δεν είχαν προηγουμένως χρησιμοποιηθεί. Ο διαχωρισμός μεταξύ των σετ ήταν 75% για το σετ εκπαίδευσης, 15% για το σετ επικύρωσης και 10% για το σετ δοκιμής, χωρισμένο τυχαία.

Συγκεκριμένα, οι υπερπαράμετροι που χρησιμοποιήθηκαν, περιλαμβάνουν τον αριθμό των

κρυφών επιπέδων καθώς και το μέγεθος τους, το μέγεθος του επιπέδου εξόδου, την παράμετρο κανονικοποίησης, τον ρυθμό εκμάθησης, καθώς και το μέγεθος παρτίδας. Αυτές οι υπερπαράμετροι επιλέχθηκαν μετά από εκτεταμένες δοκιμές με την καθεμία, με τις καλύτερες τιμές να αποθηκεύονται και να χρησιμοποιούνται για προπόνηση. Ο αριθμός της εποχής διατηρήθηκε στις 100 εποχές σε όλο τον πειραματισμό, κάτι που ήταν αρκετό για όλες τις περιπτώσεις για να σταθεροποιηθεί το σκορ ακρίβειας επικύρωσης.

Έγινε πειραματισμός με 3 ή 4 χρυφά επίπεδα, καθώς έγινε φανερό ότι λόγω της πολυπλοκότητας του προβλήματος χρειαζόταν ένα αρχετά μεγάλο δίχτυο και για τις δύο προβολές. Το μέγεθος των χρυφών στρωμάτων χυμαινόταν από 256 νευρώνες έως 1024 νευρώνες ανά στρώμα, με όλα τα χρυφά στρώματα να έχουν τον ίδιο αριθμό νευρώνων. Όσον αφορά το μέγεθος του στρώματος εξόδου, πειραματιστήχαμε με μεγέθη [10,50,100,150]. Επιπλέον, το ποσοστό εχμάθησης χυμαινόταν από 10^{-4} έως 10^{-2} και η παράμετρος ταχτοποίησης ήταν στην περιοχή από 10^{-4} έως 10^{-2} . Το μέγεθος παρτίδας που χρησιμοποιήσαμε ήταν είτε 500 δείγματα είτε 1000 δείγματα. Η συνάρτηση ενεργοποίησης ήταν μια έχδοση του CCA χρησιμοποιώντας μια μέθοδο βελτιστοποίησης χωρίς παραγώγους.

Για αυτό το σύνολο δεδομένων, εφαρμόσαμε το OPNMF στους χάρτες RAVENS των δεδομένων απεικόνισης, επειδή έχει παρατηρηθεί ότι επιτυγχάνει καλύτερα αποτελέσματα από τα MUSE ROI, ενώ μειώνει τις διαστάσεις των δεδομένων. Σε αυτή την εργασία, τα δεδομένα απεικόνισης περιορίστηκαν σε 30 στοιχεία. Ακόμα, εφαρμόστηκε MCA στα γενετικά δεδομένα, προκειμένου να μειώσουμε διαστατικά τα 54 SNP σε 10 παράγωγα στοιχεία. Τέλος, διερευνήθηκε η μέθοδος FAMD, όπου ο αριθμός των παραγώμενων στοιχείων τέθηκε στα 12.

Για τα μοντέλα κατηγοριοποίσης, χρησιμοποιήθηκαν SVM, με γραμμικό, πολυωνυμικό και με συνάρτηση ακτινικής βάσης για συναρτήσεις πυρήνα. Συγκεκριμένα, για τον γραμμικό πυρήνα πειραματιστήκαμε ως προς την ποινή κανονικοποίησης L2, με τις τιμές [0,00001, 0,0001, 0,001, 0,01, 0,1, 1, 10], ως προς την παράμετρο C (παράμετρος κανονικοποίησης) και ως προς την στρατηγική ταξινόμησης πολλαπλών κατηγοριών one-vs- rest. Όσον αφορά τον πολυωνυμικό πυρήνα, πειραματιστήκαμε με πολυωνυμικούς βαθμούς [2,3,4,5], ανεξάρτητες τιμές όρων [0,00001, 0,0001, 0,001, 0,01, 0,1, 1, 10], τιμές C των [0,00001, 0,00 0,001, 0,01, 0,01, 0,01, 0,1, 1, 10] και τέλος τιμές συντελεστών πυρήνα (τιμές g) [0,0001, 0,001, 0,01, 0,1, 1]. Τέλος, όσον αφορά τους πυρήνες συναρτήσεων ακτινικής βάσης, πειραματιστήκαμε με τιμές C των [0,0001, 0,001, 0,001, 0,1,01, 1].

Για τους κατηγοριοποιητές συνόλου, πειραματιστήκαμε με τις μεθόδους Bagging (Bootstrap Aggregating) και Adaboost. Και για τις δύο αυτές μεθόδους, πειραματιστήκαμε με τον βασικό ταξινομητή να είναι ένα δέντρο αποφάσεων ή ένα γραμμικό SVM.

Ο συντονισμός παραμέτρων και για τα δύο αυτά μοντέλα καθώς και για τους βασικούς ταξινομητές τους έγινε με αναζήτηση πλέγματος μαζί με διασταυρούμενη επικύρωση, χρησιμοποιώντας 5 πτυχές. Συγκεκριμένα, για το μοντέλο συνόλου ταξινομητή Bagging, οι παράμετροι με τις οποίες πειραματιστήκαμε ήταν ο αριθμός των εκτιμητών, με τιμές [5,10,15], τα μέγιστα δείγματα του συνόλου δεδομένων στα οποία θα μπορούσε να εκπαιδευτεί ένας εκτιμητής, με τιμές [60%, 80%, 100%]. Όσον αφορά το μοντέλο ταξινομητή συνόλου Adaboost, οι παράμετροι με τις οποίες πειραματιστήκαμε ήταν και πάλι ο αριθμός των εκτιμητών, με αυτούς τους [5,10,15,50] και τους SAMME και SAMME.R για τον αλγόριθμο ενίσχυσης. Το ποσοστό εκμάθησης για αυτό το μοντέλο διατηρήθηκε στο 1,0. Για τον ταξινομητή βάσης του δέντρου απόφασης, οι παράμετροι με τις οποίες πειραματιστήκαμε ήταν το κριτήριο του εκτιμητή, με αυτό να είναι είτε gini impurity είτε entropy, μαζί με το μέγιστο βάθος, με τις τιμές του να είναι [1,2,5]. Τέλος, για τον γραμμικό ταξινομητή βάσης SVM, πειραματιστήκαμε με την παράμετρο C, με τις τιμές της να είναι [0,00001, 0,0001, 0,001, 0,01, 0,1, 1, 10].

Για την αξιολόγηση των μοντέλων, χρησιμοποιήθηκαν οι μετρικές της ακρίβειας, της εξισορροπημένης ακρίβειας και του F1 Score:

 $Accuracy = \frac{True \ Positive \ + \ True \ Negative}{True \ Positive \ + \ False \ Positive \ + \ True \ Negative \ + \ False \ Negative}$

 $Balanced \ Accuracy \ = Avg(\frac{True \ Positive}{True \ Positive \ + \ False \ Negative} + \frac{True \ Negative}{True \ Negative \ + \ False \ Positive})$

 $F1 \quad Score = \frac{True \ Positive}{True \ Positive \ + \ \frac{1}{2}(False \ Positive \ + \ False \ Negative)}$

1.3.4 Βελτιστοποιήσεις

Πρώτα απ όλα, εκπαιδεύουμε τα δίκτυα DCCA στα ανεπεξέργαστα δεδομένα, δηλαδή τις 145 τιμές ROI (απεικονιστικά δεδομένα) και τα 54 SNP (γενετικά δεδομένα). Μετά από αυτό, λαμβάνουμε τα γενετικά δεδομένα που έχουν μετασχηματιστεί σε MCA (10 γενετικά στοιχεία) και τα ζευγαρώνουμε με τα αρχικά δεδομένα απεικόνισης, δηλαδή τις 145 τιμές ROI. Τέλος, πειραματιζόμαστε με τον αντίθετο συνδυασμό, που είναι τα δεδομένα απεικόνισης που έχουν μετασχηματιστεί με OPNMF (30 στοιχεία απεικόνισης) και τα ζευγαρώνουμε με τα αρχικά γενετικά δεδομένα, δηλαδή τα 54 SNP. Η μετρική αξιολόγησης για τις παραμέτρους των δικτύων ήταν η συσχέτιση μεταξύ των δύο όψεων εξόδου.

Για αυτούς τους πειραματισμούς, παρατηρήθηκαν τα παρακάτω. Είναι σαφές ότι όσο περισσότερους κόμβους έχει το επίπεδο εξόδου, τόσο καλύτερη είναι η συσχέτιση των μετασχηματισμένων δεδομένων. Επιπλέον, είναι σαφές ότι το κρυφό μέγεθος και ο αριθμός του στρώματος έχουν μικρή επίδραση στη συσχέτιση εξόδου. Μπορεί να σημειωθεί ότι περισσότερα κρυφά επίπεδα και περισσότεροι κόμβοι ανά κρυφό στρώμα φαίνεται να επιτυγχάνουν καλύτερα αποτελέσματα, αλλά η διαφορά είναι μικρή. Όσον αφορά τις παραμέτρους μεγέθους παρτίδας, στα πειράματά μας τα αποτελέσματα παραμένουν βασικά ίδια. Η μόνη αλλαγή που παρατηρείται όσον αφορά τον χρόνο εκπαίδευσης, καθώς όσο μικρότερο είναι το μέγεθος της παρτίδας, τόσο περισσότερος χρόνος χρειάζεται το μοντέλο ανά εποχή για να διατρέξει το σύνολο δεδομένων. Το ίδιο αποτέλεσμα μπορεί να παρατηρηθεί και με τον ρυθμό εκμάθησης. Τέλος, η παράμετρος κανονικοποίησης ακολουθεί την ίδια λογική, με την αλλαγή της να μην κάνει ουσιαστική διαφορά:

Parameter Action	Correlation (Negative Loss)	
Output Dimension Size \uparrow	\uparrow	
Hidden Layer Size ↑	\uparrow	
Learning Rate	Medium to low LR is best	
Batch Size	Stays basically the same	
Regularization Parameter	Stays basically the same	

Σχήμα 1.4: Συμπεράσματα από την βελτιστοποίηση των δικτύων DCCA. (η υψηλότερη συσχέτιση είναι επιθυμητή)

1.3.5 Αποτελέσματα

Παρουσιάζονται συνοπτικά τα αποτελέσματα, καθώς το πλήθος των διαφορετικών συνδυασμών για τα μοντέλα είναι αρκετά μεγάλος για το κεφάλαιο της περίληψης.

Για να συνοψίσουμε τα αποτελέσματα, παρουσιάζουμε έναν πίναχα με την καλύτερη ακρίβεια, το F1 Score και τις ισορροπημένες βαθμολογίες ακρίβειας για κάθε μοντέλο, μαζί με μερικές σημειώσεις. Το παρακάτω διάγραμμα απεικονίζει τα καλύτερα αποτελέσματα κάθε μοντέλου, ώστε να συγκρίνονται πιο εύκολα.

Model	Best Accuracy (%)	Best F1 Score (%)	Best Balanced Accuracy (%)	Notes
RAW DATA	55.48	54.00	56.65	145 ROIs (Scaled) and 54 SNPs (Balanced). Both AdaBoost DT.
DCCA-150-3	51.72	50.80	45.82	Output Dimension 150, 3 Hidden Layers, no scaling or balancing. Imaging Linear.
OPNMF	58.62	54.02	61.01	30 Imaging Components (After OPNMF) Balanced only. Imaging Bagging SVM.
OPNMF + DCCA-150-3	53.79	53.38	55.03	30 Imaging Components (After OPNMF) and 54 SNPs, then DCCA, then scaled and balanced. Both AdaBoost SVM.
МСА	59.59	57.85	59.41	145 ROIs (Scaled), 10 Genetic components, Balanced only. Both Poly SVM.
MCA + DCCA-150-3	52.05	50.10	53.37	145 ROIs and 10 Genetic components (After MCA), then DCCA, then scaled and balanced. Imaging Bagging SVM.
MCA + OPNMF	57.24	53.68	53.77	30 Imaging Components (After OPNMF) and 10 Genetic components (After MCA), Balanced only. Both AdaBoost SVM.
FAMD	55.17	44.42	50.87	10 Components, no scaling, no balancing. Both Poly / RBF SVM.

Σχήμα 1.5: Για κάθε μοντέλο, εμφανίζεται η καλύτερη ακρίβεια, το F1 score, και η εξισορροπημένη ακρίβεια που επιτεύχθηκε, μαζί με σημειώσεις που εξηγούν τον συνδυασμό δεδομένων και παραμέτρων που χρησιμοποιούνται. Με πράσινο χρώμα επισημαίνονται τα καλύτερα και τα δεύτερα καλύτερα (ανοιχτότερο πράσινο) αποτελέσματα που έχουν επιτευχθεί μεταξύ όλων των μοντέλων.



Σχήμα 1.6: Για κάθε μοντέλο, απεικονίζεται η καλύτερη ακρίβεια, F1 score και η εξισορροπημένη ακρίβεια.

1.3.6 Συμπεράσματα, συζήτηση και μελλοντικές επεκτάσεις

Αυτή η μελέτη προορίζεται για να είναι μια εχτενής σύγχριση των εφαρμογών των μεθόδων ανάλυσης δεδομένων, καθώς και των μεθόδων μηχανικής και βαθιάς μάθησης, που εφαρμόζονται στο πρόβλημα της ταξινόμησης γνωσιαχά φυσιολογιχών ατόμων / ατόμων με ${
m HN}\Delta$ / ατόμων με νόσο Αλτσχάιμερ. Από το έργο που επιτελέστηκε, υπάρχουν ορισμένα ξεκάθαρα μηνύματα. Πρώτα από όλα, οι τεχνικές ανάλυσης δεδομένων όπως το MCA και το OPNMF είναι καλύτερες σε σύγκριση με τα MUSE ROI επειδή τα στοιχεία OPNMF βασίζονται σε δεδομένα ενώ τα MUSE ROI προέρχονται από πρότυπα [42]. Το MCA είναι επίσης καλύτερο από τη ανεπεξέργαστη γενετική όψη, επειδή μετατρέπει τα γενετικά δεδομένα στον τύπο των δεδομένων απειχόνισης, χάνοντας χαι τις δύο όψεις να έχουν το ίδιο είδος. Επιπλέον, η μέθοδος DCCA, όπως αναφέρεται στην αρχική εργασία [43], δεν είναι ωφέλιμη για αυτό το πρόβλημα, τουλάχιστον όχι χωρίς περαιτέρω εχπαίδευση, ωστόσο αυξάνει τη συσχέτιση μεταξύ των απόψεων, σε συμφωνία με την αρχική εργασία. Οι μέθοδοι ensemble classifiers είναι ανώτερες από τις απλοϊκές μεθόδους μεμονωμένου ταξινομητή, όπως τα SVM. Τέλος, είναι σαφές ότι η χρήση μόνο γενετιχών δεδομένων δεν αρχεί για να δώσει αποτελέσματα υψηλότερης ποιότητας, χαθώς δείχνουν μόνο προδιάθεση για τη νόσο. Οι συνδυασμοί που πέτυχαν τα καλύτερα αποτελέσματα χρησιμοποίησαν απεικόνιση ή απεικόνιση μαζί με γενετικά δεδομένα, είτε δεδομένα απεικόνισης μετασχηματισμένα με OPNMF είτε γενετικά δεδομένα μετασχηματισμένα με MCA, συνοδευόμενα είτε από σύνολα Bagging SVM είτε με ένα πολυωνυμικό SVM πυρήνα ως ταξινομητές.

Τέλος, υπάρχουν πολλές πιθανές μελλοντικές κατευθύνσεις. Θα μπορούσε κανείς να εξερευνήσει, διαφορετικούς αλγόριθμους βελτιστοποίησης, συναρτήσεις ενεργοποίησης και γενικά διαφορετικές αρχιτεκτονικές για τα παράλληλα νευρωνικά δίκτυα της μεθόδου DCCA. Μια άλλη κατεύθυνση μπορεί να είναι η αύξηση δεδομένων ή η δημιουργία συνθετικών δεδομένων, προκειμένου να εμπλουτιστεί το σύνολο δεδομένων. Όσον αφορά τις τεχνικές ανάλυσης δεδομένων, η μέθοδος OPNMF είναι αυτή που θα μπορούσε να ωφεληθεί από τον πειραματισμό με διαφορετικό αριθμό στοιχείων, κάτι που δεν έγινε σε αυτή τη μελέτη λόγω περιορισμών στην υπολογιστική ισχύ. Επιπλέον, μπορούν να χρησιμοποιηθούν πιο σύνθετοι ταξινομητές, όπως ταξινομητές MLP και K-Nearest Neighbors. Αυτό μπορεί επίσης να επεκταθεί στις μεθόδους συνόλου, όχι μόνο για τα βασικά μοντέλα ταξινομητή, αλλά και για τις μεθόδους συνόλου. Τέλος, για να αντιμετωπιστεί το ζήτημα των διαφορετικών ως προς τη φύση και τον τύπο προβολών, θα μπορούσε κανείς να διερευνήσει τη χρήση των βαθιών αυτοκωδικοποιητών προκειμένου να αμβλυνθεί το πρόβλημα του χειρισμού κατηγορικών δεδομένων.

Κεφάλαιο 2

Introduction

A rapidly growing cause of death in the developed countries are neurodegenerative diseases. Diseases such as the Alzheimer's, as well as disorders such as Mild Cognitive Impairment are most prevalent on old people, with ages equal or older than 65 years old, being the group that's most affected by them. It is estimated that in the US alone, 6.5 million people are suffering from Alzheimer's Disease today, while globally this number can be as high as 35 million. Those estimations are expected to grow to 135 million globally, and many of them are undiagnosed and or untreated even today.([1],[2])

During the normal course of aging, the human brain displays changes, both anatomical as well as functional, that seem to be accelerated in patients with such diseases. The human brain is developed until the age of 25 years, and after that it continuously loses neural mass, leading to brain atrophy, a condition that is studied extensively, thanks to advances the field of medical imaging. The atrophy effect is sped up in some patients quite significantly, and in the case of the AD, it leads to neuronal decay and loss, and eventually death.

The effects of AD vary quite significantly, and are apparent not only clinically, but in imaging scans and genetic surveys as well. Because of low awareness, MCI and AD are often mistakenly associated with getting older, while due to the need of an experienced practitioner for the diagnosis, many cases go undiagnosed. However, a great deal of research is being done on studying the symptoms, potential causes, as well as the treatment of the disease, with some estimates putting the total cost of Alzheimer's research in the tens of billions USD. Furthermore, for the total cost for the healthcare related to AD patients for the year 2020 in the US has been estimated to be around 300 billion USD. ([2], [3]).

Despite the considerable resources, no clear cause has been found for AD, no definitive prevention method has been found, and no treatment method has been widely successful.

2.1 Contents of this Thesis

To tackle the problem of diagnosing Alzheimer's as well as MCI, and also predicting and modelling their respective courses, a plethora of studies have been published. A common characteristic that many have is the use of machine and deep learning methods, especially on neuroimaging, genetic and clinical data collected from patients of the diseases. The main drivers behind this effort are the advances in medical imaging and the staggering growth of computational abilities in recent years, making more complex and better methods applicable and practical. ([4], [5])

Most studies employ neuroimaging data collected with the Magnetic Resonance Imaging and Positron Emission Tomography methods, that were collected from the AD neuroimaging Initiative database. The imaging data are often accompanied with biomarkers, as well as Single Nucleotide Polymorphisms (SNPs), and even data from clinical tests performed by licensed practitioners. ([6], [7], [8]).

While in general more complex models employing different modalities of the data performed better, there is no unified approach, and the diversity of models is intriguing. However, most studies focus on either classifying whether or not a subject is a patient of AD and predicting the course of the disease, while some papers focus on predicting if a patient with MCI will convert to Alzheimer's. [4]

A common problem most studies had to overcome was the overabundance of features that the data had, while having too few samples. This is known as the 'curse of dimensionality' problem, and is associated with biomedical data. The data is characterised by very large dimensions (especially in the case of the neuroimaging view), yet not enough samples. This is due to the imaging techniques' post-collection data processing, resulting in very high dimensions, while the sample size is very low, due to the difficult, sometimes inaccessible, expensive and long-lasting nature of the technique. This has the adverse consequence of data not being easily visualised (since they cannot be interpreted in the three dimensional space humans are familiar with), along with the tendency of the models to overtrain and overfit on the low number of datapoints. To avoid the aforementioned problems, data analysis techniques have been used, with them being as simple as PCA, or as complex as employing deep learning, for example neural networks. ([9], [10], [11])

This study attempts to create a comparative analysis of machine learning methods and algorithms being applied to the problem of predicting whether a subject is cognitive normal, has MCI, or AD. The data was collected through the ADNI dataset, and more specifically, imaging data taken from T1-weighted MRI scans, along with genetic data in the form of SNPs.

In this thesis, we also explore data analysis techniques in order to tackle the dimensionality curse problem, as well as transformations and statistical analysis methods. To convert the exceedingly dissimilar imaging and genetic views, we experimented with a novel technique, called Deep Canonical Correlation Analysis, where neural networks are used to learn a transformation of the different views' features into a hyperspace that is better linearly correlated than the raw data, and thus potentially easier to perform the classification task.

Furthermore, we experimented with dimensionality reduction methods, such as Multiple Correspondence analysis, Orthonormal Projective Non-Negative Matrix Factorization, and Factor Analysis of Mixed Data. All of the possible combinations of the techniques were used, and their effect on the classification task was compared.

For the classification task in particular, we present the results from applying Support Vector Machines, which are widely regarded as a fairly simple, understandable, yet practical and capable model, as well as Ensemble Learning methods such as Bagging and Adaboost to the problem. For the ensemble methods, we experimented with Decision Trees and again Support Vector Machines as base model classifiers, and the yielded results were compared not only between them, but also to the single-classifier results.

2.2 Structure of the Thesis

This work is divided into 8 chapters, including the introduction chapters. Chapter 3 introduces some theoretical knowledge in order to tackle the problem described in the introductory chapter, pertaining to the human brain and some fundamentals about the machine and deep learning methods that were used. Chapter 4 explains the methodology that was used, with a brief description of the dataset that was used, the parameters of the methods, as well as the metrics to evaluate them. On chapter 5, we present some results from the attempt at optimizing the models that were later used for the classification tasks, and on chapter 6 the classification results are presented. Finally, on chapters 7 and 8, conclusions are drawn from observations made on the results, future extensions and directions are discussed, as well as the practical limitations of this work.
Κεφάλαιο 3

Theoretical Background

3.1 Human Brain

3.1.1 Aging

As the human body ages, all organs experience age-related effects, and so does the brain. Both physiologically and cognitively, there are several changes that can be observed as part of the normal brain aging process. [12]

Cognitively, memory (specifically episodic and semantic memory) is one of the core areas that are affected. Older people may be forgetting names of items or persons, having to repeat questions, misplacing items, getting lost, and having trouble recalling information in general. Additionally, language skills such as vocabulary and language skills may be affected, as well as the ability to learn new skills and multitask. [13]

Physiologically, the brain shrinks in the areas of the frontal lobe as well as the hippocampus, the areas that are generally thought to be linked with higher cognitive function and memory, at a rate of 5% per decade after the age of 40. This effect is due to the grey matter shrinkage, which is attributable to neuronal cell death. Additionally, cortical density decline is observed, meaning that the outer surface of the brain is becoming thinner. This effect is more pronounced in the frontal and temporal lobes. White matter also declines, with the myelinating regions of the frontal lobe being most affected by white matter lesions. Finally, the levels of neurotransmitters such as dopamine and serotonin see a steep decrease, an effect that has been associated with declines in cognitive and motor performance. ([44], [45])

While the aforementioned symptoms are very much similar to the symptoms that Mild Cognitive Impairment and Alzheimer's Disease exhibit, normal aging patterns of decline are divergent from the ones of MCI and AD. The effects of normal brain aging are characterized by their occasional nature, while MCI's and AD's ones are more consistent, and gradually worsen in some cases, and they are accompanied by other dementia symptoms, such as confusion, mood changes and others. Furthermore, the physiological changes of the brain are much more pronounced and significantly more severe. MCI and AD have a much more noticeable effect on the person's daily life, and some cases need assistance in order to perform normal daily tasks. [14]

3.1.2 Mild Cognitive Impairment

Mild Cognitive Impairment is a state of a person that is characterized by problems with memory, language, thinking or judgment. It is usually observed between the stage of normal cognitive decline that happens to humans due to aging and the dramatic fall in cognition that is apparent to people with dementia. People with MCI have memory loss or other cognitive ability loss, exceeding the normal decline due to age and are not demented. [15]

MCI's symptoms can manifest in many different functions of the human brain, including weaker memory, poor reasoning and judgment skills, visual perception and others. Frequently, MCI coexists with other illnesses or emotions, such as depression, anxiety, irritability and aggression, or apathy. [19]

The cause of the disorder is unknown; it is believed that MCI is caused by the same mechanisms that are thought to be responsible for the neuropathology of the early stages of Alzheimer's Disease, however that is unproven. Risk factors include age, family history of AD or dementia, genetic factors, and other medical conditions such as diabetes, high blood pressure, smoking, obesity, depression etc. [19]

People with diagnosed MCI have a significantly higher chance than that of cognitive normal population to develop AD or some form of dementia. Despite the fact that there is no standardized test for MCI, clinical characterization is achieved through the results of various tests (such as mental tests, neurological exams, patient family history, brain imaging and searching for biomarkers) and the informations that the patient provides. [20]

It is not exactly clear how to prevent MCI, but studies show that engaging in frequent physical activity, maintaining a healthy and balanced diet, engaging socially with others, being mentally active, reducing alcohol and not smoking may be mitigating factors to the risk of developing the condition. [20]

3.1.3 Alzheimer's Disease

Alzheimer's Disease is a neurodegenerative disease that affects the brain both biologically and cognitively. It was first reported by A. Alzheimer in 1906, but described as a disease only after 1910, by E. Kraepelin. AD is the most common cause of dementia, a term used to describe a group of symptoms that include decline in memory, reasoning, and other thinking skills. It is reported that AD is responsible for about 70% of dementia cases. ([16],[17],[18])

AD is a progressive condition, meaning the symptoms gradually appear and worsen over time. Early symptoms include short term memory loss, decline in conversational abilities and poor reasoning. As the disease progresses, patients have trouble recalling names, may have confusion and obsessive, repetitive or impulsive behaviour, serious problems with speaking and the use of language, and generally problems that require external assistance in their daily life. In later stages of the disease, the patients have trouble with even the most basic tasks, such as eating and moving, and require full time assistance. Gradually, the condition of the patients deteriorates, ultimately leading to death. [21]

The cause of AD is unknown, but genetic and environmental risk factors have been implicated. AD is linked with the formation and buildup of plaques (abnormal clusters of protein fragments) of the protein amyloid β , and neurofibrillary tangles (twisted strands of protein) of the tau protein. There are several hypotheses as to the disease's origin, yet none of them have been confirmed. There are two perhaps significant hypotheses, the amyloid and the cholinergic Hypothesis. [22]

AD is a multifactorial disease, being associated with several risk factors, such as age and gender, genetic factors, life style, coexistent or previous diseases, head injuries and environmental factors. The most important however is age, with most AD cases having a late onset that starts after 65 years of age. Normal brain aging is characterized by a reduction in brain volume and weight, a loss of synapses, and the enlargement of ventricles. These changes appear in AD patients as well, but more profound in general. There are two categories of AD based on the age that it appears, Early Onset AD which is generally familial and displays inheritance and has onset age that ranges from 30 to 60 years of age (1-6% of cases), and late onset AD, which is by far more common and has age of onset above 65 years. Genetic factors also play a significant role, with 70% of AD cases being related to genetic factors. The genes APP, PSEN-1, PSEN-2 and most importantly ApoE are associated with AD. ([22], [46])

In order to successfully diagnose the disease, a practitioner has to evaluate the person, with multiple tests if necessary. The diagnosis is based on medical history, advanced medical imaging of the brain (using CT or MRI or PET or SPECT), mental tests such as the MMSE, blood tests and psychological tests for depression, since depression can either be concurrent with Alzheimer's disease, an early sign of cognitive impairment, or even the cause. ([46], [47])

There's currently no cure for Alzheimer's disease, however there are certain medications available that can temporarily mitigate the symptoms. Since the cause of the disease is still unknown, there is no designated preventive roadmap. Despite that, frequent physical exercise, a healthy and balanced diet as well as staying mentally and socially active all have been linked to lower rates of AD.

3.1.4 Fundamentals of MRI (Imaging - ROIs)

A big part of studying, understanding and diagnosing neurodegenerative diseases are brain imaging techniques, such as Magnetic Resonance Imaging (MRI), Positron Emission Tomography (PET), Computerized Tomography (CT) and others. This work focuses on data collected with the method of MRI, which were used to recognize signs of MCI and AD.

The MRI technique works by measuring the energy signal of (typically) hydrogen nuclei, as a result of excitation by external radio frequency pulses. The MRI technique is frequently split into two different processes, based on the decay of the RF-induced nuclear magnetic resonance spin polarization, named T1 and T2, each producing different results depending on the signal and the tissue being imaged. Depending on the parameters of the process being used, MRI can imprint pictures of the anatomy of the human body, as well as its physiological processes. [23]

Oftentimes, to avoid examining every single point of space (commonly referred as a 'voxel' or volume-pixel), the scan is segmented in specified Regions Of Interest (ROIs). These regions are produced by segmenting the original image, either automatically, using Machine Learning methods or by employing previously computer computed brain atlases. Specifying however, the aforementioned regions, is quite complex, since there is a great variability of neuroanatomy between humans. It is believed that the brain's function is associated with structural and functional connectivities, and therefore identifying standardized and reliable is crucial for understanding the connection between the architecture of the brain and its function. ([24], [25], [26])

Because the MRI method produces a tremendous amount of data for each scan, frequently before the main task, preprocessing is applied to capture the desirable information, while maintaining ease of data manipulation. Such methods include data augmentations, feature selection, dimensionality reduction, etc.

3.1.5 Fundamentals of genetics (SNPs)

DNA in humans is arranged in chromosome pairs, with each cell having in its nucleus 23 pairs, 22 of which are autosomes and 1 pair being the sex chromosomes. In each of the 22 pairs of chromosomes, DNA is stored in identical copies, with specific chunks being characterized as genes. A gene contains genetic information in the form of long sequences of nucleotides (A,T,G,C).

If a change or variation in one or more nucleotide positions of a chromosome in the DNA sequence is found, it is called a Single Nucleotide Polymorphism (SNP - pronounced "snip"). If most humans have a specific certain nucleotide in a specific position of the genome, and a SNP occurs in some individuals in that exact position, then this position is said to have more than one allele. Because the DNA is stored in pairs of chromosomes, a person's DNA can contain in a specific position of the genome one SNP, two SNPs, or no SNPs at all. ([27], [28])

Observed SNPs can be associated with a disease, however, it may not always directly be the cause for that disease. An example of this is the APOE gene (chromosome 19 position q13.32), which has been determined to be a risk factor for AD, specifically the ε 4 allele. There are three versions of the gene in humans, ε 2, ε 3, and ε 4, with ε 3 being the most prevalent, the existence of the ε 4 variant being a risk factor, while having two ε 2 alleles being associated with lower probabilities of developing the disease. The disease however is also associated with other gene mutations, such as mutations in the genes APP, PSEN1, PSEN2 and others, which especially influence the early onset variant of the disease. ([28], [29], [30])

Recognizing the different SNPs that are contained in the human genome is the topic of studies such as Genome Wide Association Studies (GWAS), which are a part of the field of bioinformatics. Understanding the changes in the genome can help recognize how they translate in the phenotype, help with their treatment and their prevention. ([48])

3.2 Fundamentals of the Machine Learning and Deep Learning methods

3.2.1 Canonical Correlation Analysis

Canonical Correlation Analysis is a standard tool of multivariate statistical analysis used to discover and quantify associations between two sets of variables. The aim of this method is to find a transformation (projection) of the two sets of variables, such that they are maximally associated (measured by correlation). The projections are found by performing a joint covariance analysis of the two variables. [31]

This concept was introduced by C. Jordan (1875), but the method was initially described by H. Hotelling (1936). It has been used extensively in many fields, such as economics, medicine, psychology, etc., and has many extensions, such as the Kernel CCA. ([49], [50], [51])

Let $X \in \mathbb{R}^q$, and $Y \in \mathbb{R}^p$, two random vectors, and their respective covariances Σ_{11} and Σ_{22} , as well as the cross covariance Σ_{12} . The aim of CCA is to find vectors, such that the correlation $\rho(a, b) = corr(a^{\intercal}X, b^{\intercal}Y)$ is maximised.

The correlation $\rho(a, b)$ can also be written as follows:

 $\rho = \frac{a^{\mathsf{T}} \Sigma_{XY} b}{\sqrt{a^{\mathsf{T}} \Sigma_{XX} a} \sqrt{b^{\mathsf{T}} \Sigma_{YY} b}}$

This is achieved by setting the a, b parameters as follows:

$$a = \Sigma_{XX}^{-\frac{1}{2}}c$$
, and
 $b = \Sigma_{YY}^{-\frac{1}{2}}d$, where

c is an eigenvector of $\Sigma_{XX}^{-\frac{1}{2}} \Sigma_{XY} \Sigma_{YY}^{-1} \Sigma_{YX} \Sigma_{XX}^{-\frac{1}{2}}$,

and d is an eigenvector of $\Sigma_{YY}^{-\frac{1}{2}} \Sigma_{YX} \Sigma_{XX}^{-1} \Sigma_{XY} \Sigma_{YY}^{-\frac{1}{2}}$.

The vectors a, b are called canonical correlation vectors, and the indices $u = a^{\intercal}X$ and $v = b^{\intercal}Y$ are called canonical correlation variables.

This process may be repeated min(q, p) times, and find subsequent projections. However, the new vectors are subject to the constraint that they are to be uncorrelated with the previous ones, that is $a_i \Sigma_{XX} a_j = b_i \Sigma_{YY} b_j = 0, \forall i < j$.

CCA is implemented using Singular Value Decomposition on the correlation matrix.

3.2.2 Deep Canonical Correlation Analysis

Deep Canonical Correlation Analysis is an extension of the standard Canonical Correlation Analysis, created by G. Andrew, R. Arora, J. Bilmes and K. Livescu. As with normal CCA, DCCA is a method that aims to discover and learn associations between two sets of variables. [32]

In the case of DCCA, the method can learn complex, nonlinear relations between the two random vectors and transform them again non-linearly to correlate them, whereas the standard CCA cannot. The method is similar to the idea of Kernel CCA, where the optimal projections are found on the kernel-transformed random vectors, such that the resulting reproducing kernel hilbert space contains the variables in a manner that CCA can be impactful. [33]

However, the problem with KCCA is the computation complexity, as the kernel matrices become very large for real-world datasets, meaning that since it is a nonparametric method, the time required to learn the transformation scales poorly with the size of the data. Additionally, the KCCA method is also limited to the choice of the fixed kernel, meaning they can't be flexible for different types of datasets. [52]

To address these drawbacks, the use of deep neural networks is proposed, in order to simultaneously learn two deep nonlinear mappings of two random variables. In their paper, they focus on the performance metric of achieved correlation, and comparing it to the correlation of the standard method. [32]

The use of deep learning, meaning neural networks with more than two layers, is designated, since deep neural networks have been proven to be capable of representing accurately and reliably nonlinear functions that model complex real world data. The method is used to correlate different views of the same dataset, for example different modalities of a biomedical dataset.

The method relies on passing each random vector through a neural network, designed and trained to transform the random vector nonlinearly. That creates a mapping to a hyperspace that results is better correlated to the mapping of the respective (transformed) random vector.



 Σ χήμα 3.1: The two parallel networks, along with the information path (arrows)

In the figure above, the two neural networks are shown, consisting of 5 layers, with the red layer being the output layer, meaning the vectors that are maximally correlated, and the blue layer being the input layer, meaning the original random vectors.

If θ_1 is the vector of all parameters (W_i^1, b_i^1) of the first network, for each layer i, and respectively θ_2 is the vector of all parameters (W_i^2, b_i^2) of the second network for each layer, then the training goal is equivalent to finding the optimum parameters such that the correlation of the output of the networks $f_1(X_1; \theta_1), f_2(X_2; \theta_2)$ given two random vectors (views) (X_1, X_2) .

That is described as follows:

 $(\theta_1^*, \theta_2^*) = \operatorname{argmax}_{\theta_1, \theta_2} \{ corr(f_1(X_1; \theta_1), f_2(X_2; \theta_2)) \}$

Supposing that $H_1, H_2 \in \mathbb{R}^{o \times m}$ are the matrices that contain the respective outputs of the Neural Networks, for each of the training samples. The target then becomes $corr(H_1, H_2)$.

To train the networks, the computation of the gradient is needed, and its backpropagation in order to tune the networks parameters. The target is found using the same steps as the standard CCA, while the computation of the gradient, as well as its backpropagation is facilitated through singular value decomposition.

The authors of the original paper employed full-batch optimization, meaning that before every single weight update step, the network scanned the full dataset. They also used the Limited Memory Broyden–Fletcher–Goldfarb–Shanno (L-BFGS) optimization method. In order to initialise the parameter optimization for the two networks, they utilised a denoising autoencoder for each layer of the networks. The network proposes uses a non-saturating nonlinearity activation function, in the form of: If $g : \mathbb{R} \to \mathbb{R}$, and $g(x) = \frac{x^3}{3} + x$, then the function $s(x) = g^{-1}(x)$ is the activation function, maintaining a sigmoid shape, and unit slope at x = 0. ([53], [54])

3.2.3 Multiple Correspondence Analysis

Multiple Correspondence Analysis (MCA) is a data analysis technique used to analyse the structure of a number of dependent categorical variables in a dataset. It is an extension of simple Correspondence Analysis, and is similar to the well known method of Principal Component Analysis. [55]

MCA is used when a dataset contains variables that are described by nominal values, such as "Male" and "Female", or "Red", "Green", "Blue", etc. The variables can also contain quantitative values, split into categories. MCA is performed on an indicator matrix - also called a Complete Disjunctive Table - or on a Burt table. It can also be viewed as the PCA method applied to the CDT. [56]

Suppose there is a dataset containing only categorical variables, and its corresponding CDT, X. Let K be the number of the nominal variables, and each nominal variable has J_K levels and the sum of the J_K is equal to J. There are I observations. Then the table X is actually the $I \times J$ indicator matrix.

We indicate the sum of all entries to be N, and compute the probability matrix Z = fracXN. We also use the special vectors r, and c, which are the vector of the row totals of Z, and the vector of column totals of Z respectively.

Then, if

$$D_c = diag(c), D_r = diag(r),$$

we have the factor scores of the MCA are obtained from the following singular value decomposition:

$$M = D_r^{-\frac{1}{2}} (Z - rc^{\mathsf{T}}) D_c^{-\frac{1}{2}} = P \Delta Q^{\mathsf{T}},$$

where Δ is the diagonal matrix of the singular values, and the matrix of the eigenvalues is $\Lambda = \Delta^2$. MCA decomposes the matrix into coordinates (or scores) of the factor space, which can be found as follows:

 $F = D_r^{-\frac{1}{2}} P \Delta$, for the row coordinates and $G = D_c^{-\frac{1}{2}} Q \Delta$ for the column ones.

3.2.4 Non-Negative Matrix Factorization

Non-Negative Matrix Factorization (NMF) is a unsupervised, multivariate, analytic method for the approximate factorization of a matrix V into two matrices W, H under the constraint that their elements are non-negative: V = WH, such that $H \ge 0$ and $W \ge 0$. [34]

The method was created by P. Paatero and U. Tapper, and further developed by D. Lee and H. Seung ([57], [58]). It is used to reduce data dimensionality, perform clustering tasks and find underlying structures within the dataset. Because the resulting factorization contains non-negative elements, the method has the advantage of improved interpretability compared to other data dimensionality reduction methods, and has the ability to produce parts-based representation of the data, it has been applied in many different fields, such as machine learning, computer vision, signal processing, data mining, medical imaging etc. ([35], [59], [60], [61])

Lee and Seung's multiplicative update rule is the basis of the method's computation of the W and H matrices, and has the characteristics of being iterative and element based. However there are other ways, and it can be supplemented with additional constraints or regularizations, leading to many extensions.

One notable extension is that of Orthonormal Projective NMF (OPNMF), where the loading coefficients are estimated as the projection of the matrix onto the estimated components W ($H = W^{\intercal}V$), while maintaining orthonormality on the estimated components ($W^{\intercal}W = I$). As a result, all components participate in the reconstruction of all of the data samples, meaning that the overlap between the estimated components is significantly lower, having fewer parameters to be learned, while maintaining high sparsity. Additionally, this variant relies on the original update rule (and thus is computationally easier than the Projective NMF variant), and at the same time is able to generalize on unseen data without the need of retraining. [35]

3.2.5 Factor Analysis of Mixed Data

Similar to MCA, Factor Analysis of Mixed Data (FAMD) is a data analysis technique used to analyse the structure of mixed data, meaning both continuous numerical as well as categorical data. It is also used to in order to reduce the number of dimensions of the dataset, and improve interpretability. It is based on the methods of MCA and PCA.([36], [37])

Suppose there is a dataset containing both quantitative (numerical) and qualitative (categorical) variables. Let K_1 be the quantitative variables, Q the qualitative variables, and K_q the categories of the q^{th} variable. We can denote the overall number of categories of the qualitative variables as:

$$K_2 = \sum_q K_q$$

Let $K = K_1 + K_2$ be the total number of quantitative variables and indicator variables.

We assume that individuals have the same weight, and the diagonal metric of the weights of the individuals is:

$$D = \frac{1}{I}I_d$$

The quantitative variables are represented by a vector of length 1, and the qualitative ones by a cloud of datapoints N_q of its centered indicators. FAMD aims to look for a direction of v that maximizes the inertia (measure of weighted spread of the points) of the \mathbb{R}^I cloud. That goal is perfectly achieved by maximizing the following criterion:

$$\sum_{k \in K_1} r^2(k, v) + \sum_{q \in Q} \eta^2(q, v)$$
, where

 $\eta^2(q, v)$ is the squared correlation ratio between q and v, and $r^2(k, v)$ is the squared projection coordinate of variable k on v. [62]

FAMD's number of resulting components can range from 1 to $min(K_1, Q)$, where K_1 is the number of the quantitative variables and Q is the number of the categorical ones.

3.2.6 Support Vector Machines

Support Vector Machines are a family of models that are used for classification and regression analysis. The model was initially introduced by C. Cortes and V. Vapnik and has enjoyed considerable popularity since its creation, being one of the most widely used Machine Learning Techniques. They have been applied to many scientific fields, such as pattern recognition, image classification, biomedical research, petroleum exploration, etc.([38], [63], [64])

The goal of an SVM is to choose a hyperplane (e.g. a straight line in two-dimensional space, a plane in three-dimensional) that best separates a dataset consisting of labeled samples that belong to one of two classes. The method that SVMs rely on to achieve this goal is choosing two parallel hyperplanes that separate the two classes such that the distance between them (the margin) is maximal. The data points on the margin maximizing hyperplanes (supporting vectors) define the decision surface for the classification, as shown below: [65]



 $\Sigma\chi\eta\mu\alpha$ 3.2: Support Vectors are shown in orange, the decision boundary in blue, the datapoints in green (squares for one class, stars for the other), and the dash lines are the margin maximizing hyperplanes. [60]

SVMs can be used for multi-class classification as well, through a one-vs-one scheme or one-vs-rest approach, where decision boundaries are calculated between respective classes or a class and the rest of the dataset in each case, respectively. [66]

SVMs enjoy a variety of advantages, such as efficiency in both low and high dimensional spaces, memory efficiency, and being able to produce results in cases where the number of samples is less than the number of dimensions. [67]

However, one major drawback of SVMs is that the standard model does not work on datasets that are not linearly separable; that is one hyperplane cannot correctly divide the classes. To get around that, SVMs use kernels. Thus, the dataset is first non-linearly mapped through a kernel in a higher dimension space where the data is linearly separable, and then the original SVM algorithm is performed. In this approach, one can use the kernel trick to compute the transformations through the kernel function for the whole dataset. [65]

SMVs suffer from over-fitting issues, as well as being sensitive to parameter selection/tuning, such as the kernel function and regularization term choice, especially if the number of parameters is much greater than the number of available samples. Additionally, SVMs are not scale invariant, so scaling the dataset is highly recommended. ([67], [68])

3.2.7 Ensemble Learning

Ensemble learning is a technique of combining a multitude of models to enhance the performance of the task, such as classification problems, regression or approximation tasks etc. This is achieved by applying the (perhaps different) models to the data available (or a subset thereof) and combine their outputs, in order to make a better attempt at solving the problem. [39]

An ensemble is created by combining either different models, or models with different parameter initializations and configurations. Such models can be relatively simple, such as decision trees, naive bayes classifiers, or SVMs, or more complicated, such as multi layer perceptrons, or even other ensembles altogether. A key aspect of the base model selection is to create enough diversity of opinions, that is differentiation between the models themselves. [40]

The ensemble learning technique relies on two concepts: the way the dataset is used to train the base models (how the data is introduced to the models) and the way that the outcome of each base model is considered towards the combined outcome.

The method that is employed for training the base models can be as simple as dividing the dataset by the number of models and feeding each subset to each model, or as strategically complex as to involve feature selection along with data augmentation during the phase of training the base models. Another approach might be introducing different views of the dataset to different base models. [40]

Respectively, the method employed for combining the outcomes of the base models can be as simple as simple majority voting (for example the most voted class in a classification problem) or algebraic combiners, or more sophisticated and tailored to a specific problem strategies.

The idea behind ensemble learning is to enhance the decision taken with group knowledge; that is to reduce the likelihood of an unfortunate selection. While that is not guaranteed, there is empirical evidence that ensemble models achieve in general better performance than that of single models, and in some case better than that of the average of their base models. [40]

Another motive for employing ensemble methods is their ability to perform fairly well both in big data tasks and when there isn't adequate data for the successful training of a single model. The big data case is handled with dividing the dataset into many subsets, and training each model on a single subset, thus making the training phase much easier. On the other hand, with a strategy such as bootstrapping, different base models can be trained on different combination of samples of data, taken from the initial dataset, with replacement, and treated as if they were independently drawn. ([40], [69])



 Σ χήμα 3.3: Combining classifiers with different decision boundaries reduce error. [68]

One method of ensemble learning applied to the problem of classification is that of Bagging, or Bootstrap Aggregating, where the base models are trained on drawn samples from the initial dataset, with replacement, and the base models are classifiers of the same type. The individual classifiers' outcomes are combined in a simple majority voting strategy to determine the overall outcome of the ensemble. Another notable example is that of Adaboost, a version of the boosting ensemble, adapted for the problem of multiclass classification. As previously, bootstrapped training data samples are drawn from an initially uniform but continuously evolving distribution, ensuring that samples that were previously mislabeled are seen more often, and therefore training the base classifiers to the most difficult instances. The base classifiers are combined in a weighted majority voting manner. ([40], [41])

The different methods of ensemble classifiers have been applied to a number of fields, such as assessing the risk of developing cardiovascular disease as a long-term diabetes complication [70], error correcting output codes [71], feature selection [72], etc.

Κεφάλαιο 4

Methodology

4.1 Data Pipeline Overview

The data of this thesis were obtained from the Alzheimer's Disease Neuroimaging Initiative (ADNI) database, a global research study that focuses on understanding better how to prevent or delay the disease, as well as supporting the investigation on methods of treatment. The study collects imaging, genetic, clinical, biospecimen data from people with MCI, Alzheimer's, as well as from people that are Cognitively Normal (CN). The data was collected in 4 phases, ADNI1, ADNI2, ADNIGO and ADNI3. This study uses data from all three phases, however only uses the genetic and imaging views of the dataset. For more information on the dataset, please visit . ([73], [74])

The imaging data is in the form of 145 ROI values, acquired from scanners either 1.5T or 3T, using T1-weighted sequences, depending on the phase. The images collected by the scanners have been filtered through Quality Control, and have been preprocessed through intensity normalization and gradient un-warping. In order to acquire the ROI intensity values, the method of MUlti-atlas region Segmentation utilizing Ensembles (MUSE) was used creating the regions of interest. ROIs values' magnitude is analogous to the regional volume of brain tissue, depending on tissue type. The Regions that these values correspond to can be found in the appendix. ([75], [76])

As for the genetic data, each participant has values for 54 susceptibility loci, in the form of SNPs, that have been identified by AD genetics studies. These values have been filtered through QC, and as previously mentioned are int the form of number of alleles (0,1,2) for each SNP. ([77], [78], [79], [43])

The dataset has 1302 participants, (56.91% male), with mean age 75.20 y.o. for males and 74.36 y.o. for females. The dataset now has 433 CN participants (33.25%), 626 MCI patients (48.07%) and 243 AD patients (18.66%). Following that, Linear Regression was performed to remove any unwanted age, sex, or brain size related effect. The regressor was fitted on the CN group, and the transformation was applied to the entire sample. Subsequently, the data was transformed through experimentation, and the output of the method was stored, in order to be compared with the raw data. Afterwards, data analysis techniques were applied, such as OPNMF to the imaging data, or MCA to the genetic data, or FAMD to the whole dataset. These methods were applied both to the original data, as well as the output of the DCCA method, in order to be compared later. The



Σχήμα 4.1: Distribution of age for participants



Σχήμα 4.2: Distribution of classes for participants

result of each one of those techniques was then saved for later tasks. Finally, each one of the results of combinations of the methods was fed to SVM as well as ensemble classifiers.



Σχήμα 4.3: Data Pipeline Overview



Σχήμα 4.4: Data Pipeline Diagram

4.2 Linear Regression

Linear Regression is a statistical method that aims to learn the relationship between dependent and independent variables. In our case, the goal is to study the shrinkage effect that AD and MCI have on brain size, and therefore to study it properly we need to isolate that effect. In order to do that, any effects that age, sex and different cranium sizes have, must be removed. Finding out the pattern between the brain size, which is the 145 ROI values, and the age the person has, as well as his/her gender and cranium size, is necessary if we want to remove it.

LR learns the trend between the independent variables (age, gender, cranium size) and the dependent variable (ROI volume), so calculating the difference from the computed trend values and the real values is called the residual values. We use those residual values as a means to recognize how intense the effect of the disease on the participant's brain size is, since the other independent variables' effects have been subtracted, meaning that if there was no effect, the trend between age, gender and cranium size would accurately predict the brain size and there would be no error.

The model is trained on CN participants only, however all participants brain sizes are predicted using the trend line learned, to find out the difference between the real values and the predicted ones. We only keep the residual values, as they signify the difference from the trend line, which is equivalent to the intensity of the disease's effect.

The trend lines can be observed in the below figure for the ROI MUSE_Volume_48 which is translated to the left hippocampus area. It is clear that while the CN (green) population is centered around the x axis, meaning the brain size values are well predicted, the MCI population (blue) is lower than predicted, and the AD population (red) is much lower. This is expected, as the real brain size values are lower, since MCI and AD both cause a shrinkage of the brain.

Regression on CN for MUSE_Volume_48



Σχήμα 4.5: Linear Regression on CN for $MUSE_Volume_48$

4.3 DCCA model training

In addition to the previously mentioned techniques, DCCA was used in order to transform the two uncorrelated, different views into two that are more linearly correlated. The choice of hyperparameters, the methods and class functions, as well as the parameter optimization was subject to our experimentation.

As stated before, the DCCA model relies on two parallel networks, each taking as input a view of the dataset, and producing the output that are the views but nonlinearly transformed in order to use as subject for classification.

The dataset for the model optimization was split into three sets, the training, validation and the test set, with the last one being kept hidden from the model during the training phase, in order to ensure an accurate prediction on unseen data. The split between the sets was 75% for the training set, 15% for the validation set, and 10% for the test set, split randomly. This was done in order to ensure that the model had enough training samples, as well as validation samples to achieve good accuracy scores without overfitting, and enough test scores as to not skew the results.

The hyperparameters include the number of hidden layers as well as hidden layer size, output layer size, the regularization parameter, the learning rate, as well as the batch size. These hyperparameters were chosen after extensive testing with each one, with the best values stored and used for training. The epoch number was kept at 100 epochs along all experimentation, which was enough for all cases for the validation accuracy score to stabilize.

We experimented with 3 or 4 hidden layers, as it became apparent that due to the complexity of the problem, a big enough network was needed for both views. The original paper used the same architecture for both of the networks, a logic we followed in our study as well. The size of the hidden layers ranged from 256 neurons to 1024 neurons per layer, with all hidden layers having the same number of neurons. As for the output layer size, we experimented with sizes of [10,50,100,150]. One attempt was made with output layer size of 300 in order to observe the effect of large output layer size in the correlation metric, however the computation was extremely time-consuming. Furthermore, the learning rate ranged from 10^{-4} to 10^{-2} , and the regularization parameter being in the range of 10^{-4} to 10^{-2} . The batch size that we used was either 500 samples or 1000 samples.

The activation function was kept the same as the original paper, which was a sigmoid function, and the error metric was the error metric as defined from the paper, a version of CCA using a derivative-free optimization method. The optimizer function of the paper was the L-BFGS second-order optimization method, however due to it being more difficult to compute, the optimizer RMSProp was used, with similar results. The original paper initialized the neural network using a denoising autoencoder, but this was out of the scope of this study. [80]

The DCCA implementation we used was made with python by Z.Wu. The method is implemented with pytorch, which supports for multi-GPU training, however this study employed only CPU training. ([81], [82])

4.4 Data Analysis Techniques

In our experimentation with the dataset, it was noticed early that the nature of the different types of views of the dataset was an obstacle to the methods and the goals we wanted to achieve. We hypothesized that the difficulty to achieve better classification scores stemmed from the fact that the two views of data were different not only in type (numerical vs categorical), but in the difference of the dimensions as well, making the algorithms employed inefficient and/or not well suited for the task.

To remedy this situation, we experimented with data analysis techniques. Initially we tried OPNMF, to reduce the number of imaging dimensions, to more closely match the number of genetic dimensions. Then, we tried using MCA to transform the genetic data from categorical to numerical, and finally we tried creating a combining transformation of the two views using FAMD, as a benchmark for the other methods. All possible combinations of the methods were tested, in order to find the best possible mix.

A data analysis technique that this study explores is that of Orthonormal Projective Non-Negative Matrix Factorization. This method is used to dimensionally reduce a dataset, while still maintaining interpretability, due to the non-negative nature of the matrix decomposition. For this dataset, we applied OPNMF to the RAVENS maps of the imaging data, because it has been observed to achieve better results than the MUSE ROIs, while reducing the dataset dimensions. In this work, the imaging data was reduced into 30 components. [42]

The number of the resulting components was chosen in a way that was close to the number of the number of dimensions of the genetic data, close to the (later referenced) number of MCA components, while not too small in order to retain most of the information and minimize approximation error, and not too big in order for the method to have any use. It is worth noting that the OPNMF method rescales the data, a much needed action, since different ROIs have orders of magnitude different intensities, a feature that the SVM classifiers benefit from.

Another data analysis technique that is used in this study is Multiple Correspondence Analysis, which, as previously mentioned, analyses the structure of a number of dependent categorical variables, and performs dimensionality reduction if necessary. For this dataset, we applied MCA to the genetic data, in order to dimensionally reduce the 54 SNPs into 10 components. This had the added benefit of transforming the categorical data into numerical, which we hypothesized would greatly enhance the outcome of the classification.

The method was performed using mca, a package for python which is intended to be used along with pandas. [83]

The third and final technique we explored is Factor Analysis of Mixed Data, which combines multiple views of the dataset, to create a transformation along with a dimensionality reduction.

The number of the resulting components was chosen based on the fitting time, however it was quickly observed that above 12 components, the time needed for computing the resulting components was more than substantial, making the process of optimizing quite lengthy and not feasible. The method was based on the implementation of the python package prince, an opensource package developed by Max Halford. [84]

4.5 Classification

The task we tried to enhance the models for was the one of classification. To solve that, we chose the SVM family of models, and optimized it through parameter grid search, which is exhaustive search of all the different parameter combinations.

As mentioned previously, SVMs can have different kernels, in order to accommodate for non-linear datasets. We experimented with linear, polynomial, and radial basis function kernels, which are among the most commonly used. For each kernel, its specific parameters were optimized through grid search, and the results were evaluated on their ability to generalize through cross validation.

For all of the different cases, we used the python library Scikit-Learn, and specifically the sklearn.svm module. [85]

In particular, for the linear kernel we experimented with L2 normalization penalty, [0.00001, 0.0001, 0.001, 0.01, 0.1, 1, 10] C (regularization parameter) values, and one-vs-rest multi class classification strategy. As for the polynomial kernel, we experimented with polynomial degrees of [2,3,4,5], independent term values of [0.00001, 0.0001, 0.001, 0.01, 0.01, 0.1, 1, 10], C values of [0.00001, 0.0001, 0.001, 0.001, 0.01, 0.1, 1, 10], and finally kernel coefficient values (g values) of [0.0001, 0.001, 0.01, 0.1, 1]. Finally, as for the radial basis function kernels, we experimented with C values of [0.00001, 0.0001, 0.0001, 0.0001, 0.0001, 0.001

Each kernel was given 1000 iterations in order to converge, and the cross validation was done with 5 folds. The data was split into train and test splits, with respective sizes of 80% and 20% of the initial dataset, without shuffling, as the data was already ordered randomly. All of the combinations were run on all of the potential different combinations on views of every data analysis techniques (imaging + genetic views, only imaging, only genetic).

Furthermore, in addition to the simple method of the SVM models, we attempted to use the method of ensemble learning to further enhance the classification outcome. For that reason, we experimented with the methods of Bagging (Bootstrap Aggregating) and Adaboost. For both of those methods, we experimented with the base classifier being a decision tree or a linear SVM.

For all of the different combinations, we used the python library Scikit-Learn, and specifically the sklearn.ensemble module.

The parameter tuning for both of those models as well as their base classifiers was done with grid search along with cross validation, using 5 folds. Specifically, for the Bagging classifier ensemble model, the parameters that we experimented with were the number of estimators, with values of [5,10,15], the maximum samples of the dataset that an estimator could train on, with values of [60%,80%,100%]. As for the Adaboost ensemble classifier model, the parameters we experimented with were again the number of estimators, with those being [5,10,15,50], and SAMME and SAMME.R for the boosting algorithm. The learning rate for this model was kept at 1.0.

For the decision tree base classifier, the parameters we experimented with were the estimator criterion, with it being either gini Impurity or entropy, along with the max depth, with its values being [1,2,5]. Finally, for the linear SVM base classifier, we experimented with the C parameter, with its values being [0.00001, 0.0001, 0.001, 0.01, 0.1, 1, 10].

Before the classification task, if needed, balancing and scaling was applied. Balancing the dataset was done through random undersampling, while scaling was performed utilizing scikit-learn's preprocessing module, and more specifically the StandardScaler function. This function standardizes features by removing the mean and scaling to unit variance.

The metrics we chose for the classification task were those of accuracy, balanced accuracy, and F1 score. The implementation of the metrics that was used was again from Scikit-Learn. As for accuracy, it is the ratio of:

 $Accuracy = \frac{True \ Positive \ + \ True \ Negative}{True \ Positive \ + \ False \ Positive \ + \ True \ Negative} + \ False \ Negative}$

As for the balanced accuracy, it is the ratio of:

 $Balanced \ Accuracy \ = Avg(\frac{True \ Positive}{True \ Positive \ + \ False \ Negative} + \frac{True \ Negative}{True \ Negative \ + \ False \ Positive})$

And as for the F1 score, it is the ratio of:

F1 $Score = \frac{True \ Positive}{True \ Positive \ + \ \frac{1}{2}(False \ Positive \ + \ False \ Negative)}$



DCCA Optimizations

Before the classification, we must optimize the DCCA network parameters, and find the ideal values and combinations. We do that for the different cases of the views.

First of all, we train the DCCA networks on the raw data, meaning the 145 ROI values for the imaging view, and the 54 SNPs for the genetic view. After that, we take the MCAtransformed genetic data (10 genetic components), and pair them with the original imaging data, meaning the 145 ROI values. Finally, we experiment with the opposite combination, which is the OPNMF-transformed imaging data (30 imaging components) and pair them with the original genetic data, meaning the 54 SNPs.

As mentioned in the previous chapter, the parameter combinations we explored affect the number and sizes of the hidden layers, the size of the output layer of the network, the learning rate, the regularization parameter and the batch size. The following results are after exhaustive search of the different parameter combinations.

The metric we optimized for is the correlation between the two views, after being transformed by their respective DCCA network. Essentially we pass the data through their own trained network, transforming them and measuring how linearly correlated they have become. The total number of parameter combinations for each different case of the views is 288.

For each parameter, we plotted for the different values the correlation achieved by the worst combination of the parameter's value with all the other possible parameters values, the average correlation, as well as the best correlation. Since the implementation of the DCCA network employed the correlation as a Loss function (assigned with a negative signum, in order to be able to be minimized), we used that as the metric directly.

5.1 Original data: 145 ROI (Imaging) + 54 SNPs (Genetic)



Σχήμα 5.1: Output Layer Dimension size vs Achieved Correlation



Σχήμα 5.2: Hidden Layer size vs Achieved Correlation



Σχήμα 5.3: Batch size vs Achieved Correlation



Σχήμα 5.4: Learning Rate vs Achieved Correlation

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Σχήμα 5.5: Regularization Parameter vs Achieved Correlation

Based on the above figures, it is clear that the more nodes the output layer has, the better the correlation of the transformed data is, not only on the best case, but on average as well as on the worst case.

Furthermore, it is clear that the hidden layer size and number have little effect on the output correlation, since not only the best, but also the average and the worst cases, the correlation numbers seem to be the same. It can be noted that more hidden layers and more nodes per hidden layer do seem to be achieving better results, but the difference is minor.

As for the batch size parameters, in our experiments the results stay basically identical; the only change being noticed in terms of training time, since the lower the batch size, the more time the model takes per epoch to run through the dataset, hence more training time for the same number of epochs.

The same effect can be noticed with the Learning Rate, since the best case is more or less achieving the same results, however here we can observe that the average case benefits from a medium Learning Rate value of 0.001.

Finally, the Regularization parameter follows the same logic, with its changing not making a substantial difference, and on all accounts the results being identical.

To summarize the parameters' effects, we can create the following table:

Parameter Action	Correlation (Negative Loss)
Output Dimension Size ↑	Ŷ
Hidden Layer Size ↑	↑
Learning Rate	Medium to low LR is best
Batch Size	Stays basically the same
Regularization Parameter	Stavs basically the same

Σχήμα 5.6: Learned Conclusions from DCCA optimizations on 145 ROI (Imaging) and 54 SNPs (Genetic)

5.2 Transformed Genetic data: 145 ROI (Imaging) + 10 MCA components (Genetic)



Σχήμα 5.7: Output Layer Dimension size vs Achieved Correlation



Σχήμα 5.8: Hidden Layer size vs Achieved Correlation



Σχήμα 5.9: Batch size vs Achieved Correlation



Σχήμα 5.10: Learning Rate vs Achieved Correlation



Σχήμα 5.11: Regularization Parameter vs Achieved Correlation

As before, we notice the same patterns. Increasing the output layer size results in an increase in output correlation, hidden layer size and number of hidden layers seems to make a small difference, and for the rest of the parameters the effect seems to be negligible. We can sum up the parameter behaviour in the following table:

Parameter Action	Correlation (Negative Loss)
Output Dimension Size \uparrow	\uparrow
Hidden Layer Size ↑	\uparrow
Learning Rate	Medium to low LR is best
Batch Size	Stays basically the same
Regularization Parameter	Stays basically the same

Σχήμα 5.12: Learned Conclusions from DCCA optimizations on 145 ROI (Imaging) and 10 MCA Genetic components

5.3 Transformed Imaging data: 30 OPNMF components (I-maging) + 54 SNPs (Genetic)



Σχήμα 5.13: Output Layer Dimension size vs Achieved Correlation


Σχήμα 5.14: Hidden Layer size vs Achieved Correlation



Σχήμα 5.15: Batch size vs Achieved Correlation



Σχήμα 5.16: Learning Rate vs Achieved Correlation



Σχήμα 5.17: Regularization Parameter vs Achieved Correlation

Finally, in the case of the transformed through OPNMF imaging data combined with the raw genetic data, we can see that the parameter behaviour is again the same. Once again, output layer size increase correlates with better results, bigger hidden layer size, along with increasing the number of hidden layers improves the output correlation but only slightly, learning rate should be kept at a value of 0.001, and altering the other parameters has little to no effect.

Parameter Action	Correlation (Negative Loss)
Output Dimension Size ↑	\uparrow
Hidden Layer Size ↑	\uparrow
Learning Rate	Medium to low LR is best
Batch Size	Stays basically the same
Regularization Parameter	Stays basically the same

Σχήμα 5.18: Learned Conclusions from DCCA optimizations on 54 Imaging components and and 54 SNPs (Genetic)

Κεφάλαιο 6

Results

In this chapter, we introduce results the methods and combinations thereof we experimented with achieved.

First, the classification results of the raw views are presented, meaning the imaging data as is (145 ROIs) and the genetic data as is (54 SNPs), as a baseline. Those results are contrasted with the results of applying DCCA to those data. Following that, the classification results using MCA-transformed genetic data (10 genetic components) and the original imaging data are presented, and contrasted with DCCA applied on top of that. Afterwards, the respective results are presented after OPNMF (30 imaging components) and then with DCCA on top of that, using the combination of MCA and OPNMF, and finally after FAMD.

For the aforementioned combinations of methods, the classification results are presented before and after scaling (if needed) and balancing, to highlight the effect those techniques have on the task. Finally, to ensure that the effect of having both views is properly documented, we perform the same task with both views, as well as keeping only one view, testing imaging and genetic for every method.

6.1 Raw data vs DCCA

As mentioned before, we consider as the baseline results the SVM classification on the imaging and genetic data taken directly after LR on the ADNI dataset, and then compare them with the methods we experimented with.

6.1.1 Without scaling or balancing:



 $\Sigma\chi\eta\mu\alpha$ 6.1: Classification metric scores using Both views (Imaging and Genetic), on the SVM kernels (Linear, Polynomial, RBF), using raw data (3 left bar groups) vs using DCCA (3 right bar groups)



 Σ χήμα 6.2: Classification metric scores using only the Imaging view, on the SVM kernels (Linear, Polynomial, RBF), using raw data (3 left bar groups) vs using DCCA transformed imaging data, trained on both views (3 right bar groups)



 $\Sigma\chi\eta\mu\alpha$ 6.3: Classification metric scores using only the Genetic view, on the SVM kernels (Linear, Polynomial, RBF), using raw data (3 left bar groups) vs using DCCA transformed genetic data, trained on both views (3 right bar groups)



 Σ χήμα 6.4: Classification metric using Bagging on the imaging and genetic data.



 Σ χήμα 6.5: Classification metric using AdaBoost on the imaging and genetic data.



6.1.2 With scaling and balancing:

 $\Sigma\chi$ ήμα 6.6: Classification metric scores using Both views (Imaging and Genetic), on the SVM kernels (Linear, Polynomial, RBF), using raw data (3 left bar groups) vs using DCCA (3 right bar groups)



SVM

 $\Sigma\chi\eta\mu\alpha$ 6.7: Classification metric scores using only the Imaging view, on the SVM kernels (Linear, Polynomial, RBF), using raw data (3 left bar groups) vs using DCCA transformed imaging data, trained on both views (3 right bar groups)



 $\Sigma\chi\eta\mu\alpha$ 6.8: Classification metric scores using only the Genetic view, on the SVM kernels (Linear, Polynomial, RBF), using raw data (3 left bar groups) vs using DCCA transformed genetic data, trained on both views (3 right bar groups)



 Σ χήμα 6.9: Classification metric using Bagging on the imaging and genetic data.



 $\Sigma\chi\eta\mu\alpha$ 6.10: Classification metric using AdaBoost on the imaging and genetic data.

The following tables present the complete results for the raw data, as well as the DCCA transformed data, for each model, for each metric, for each view, and either with or without scaling and balancing. With green are highlighted the best values for each metric, depending on whether scaling and balancing were applied:

-														
						Ra	w Dat	ta						
Accuracy %				Original Data				Scaled and Balanced Data						
	Linear K	Poly K	RBF K	Bagging DT	Bagging SVM	AdaBoost DT	AdaBoost SVM	Linear K	Poly K	RBF K	Bagging DT	Bagging SVM	AdaBoost DT	AdaBoost SVM
Both Views	49.43	51.34	51.34	52.49	41	50.19	35.63	52.74	52.74	52.74	53.42	53.42	55.48	51.37
Imaging View	50.19	51.34	51.34	51.72	36.02	54.41	37.16	50	53.42	53.42	54.79	52.74	53.42	48.63
Genetic View	45.98	51.34	51.34	52.11	51.34	41.76	44.83	41.1	41.78	41.78	46.58	37.67	47.26	36.3
F1 Score				Original Data						C 1	ad and Balancad	Dete		
LA DOULS				Original Data						Scal	eu anu balanceu	Data		
LADIOIC	Linear K	Poly K	RBF K	Bagging DT	Bagging SVM	AdaBoost DT	AdaBoost SVM	Linear K	Poly K	RBF K	Bagging DT	Bagging SVM	AdaBoost DT	AdaBoost SVM
Both Views	Linear K 49.33	Poly K 34.83	RBF K 34.83	Bagging DT 50.66	Bagging SVM 42.26	AdaBoost DT 49.05	AdaBoost SVM 36.94	Linear K 49.42	Poly K 49.49	RBF K 49.49	Bagging DT 48.51	Bagging SVM 50.2	AdaBoost DT	AdaBoost SVM 45.29
Both Views Imaging View	Linear K 49.33 49.99	Poly K 34.83 34.83	RBF K 34.83 34.83	Bagging DT 50.66 50.47	Bagging SVM 42.26 36.02	AdaBoost DT 49.05 53.82	AdaBoost SVM 36.94 37.73	Linear K 49.42 47.8	Poly K 49.49 51.12	RBF K 49.49 51.12	Bagging DT 48.51 52.39	Bagging SVM 50.2 49.16	AdaBoost DT 54 51.13	AdaBoost SVM 45.29 47.39
Both Views Imaging View Genetic View	Linear K 49.33 49.99 41.28	Poly K 34.83 34.83 34.83	RBF K 34.83 34.83 34.83	Bagging DT 50.66 50.47 46.71	Bagging SVM 42.26 36.02 34.83	AdaBoost DT 49.05 53.82 38.39	AdaBoost SVM 36.94 37.73 40.74	Linear K 49.42 47.8 38.78	Poly K 49.49 51.12 40.71	RBF K 49.49 51.12 40.71	Bagging DT 48.51 52.39 36.82	Bagging SVM 50.2 49.16 37.13	AdaBoost DT 54 51.13 42.44	AdaBoost SVM 45.29 47.39 32.59
Both Views Imaging View Genetic View	Linear K 49.33 49.99 41.28	Poly K 34.83 34.83 34.83	RBF K 34.83 34.83 34.83	Bagging DT 50.66 50.47 46.71	Bagging SVM 42.26 36.02 34.83	AdaBoost DT 49.05 53.82 38.39	AdaBoost SVM 36.94 37.73 40.74	Linear K 49.42 47.8 38.78	Poly K 49.49 51.12 40.71	RBF K 49.49 51.12 40.71	Bagging DT 48.51 52.39 36.82	Bagging SVM 50.2 49.16 37.13	AdaBoost DT 54 51.13 42.44	AdaBoost SVM 45.29 47.39 32.59
Both Views Imaging View Genetic View	Linear K 49.33 49.99 41.28	Poly K 34.83 34.83 34.83	RBF K 34.83 34.83 34.83	Bagging DT 50.66 50.47 46.71	Bagging SVM 42.26 36.02 34.83	AdaBoost DT 49.05 53.82 38.39	AdaBoost SVM 36.94 37.73 40.74	Linear K 49.42 47.8 38.78	Poly K 49.49 51.12 40.71	RBF K 49.49 51.12 40.71	Bagging DT 48.51 52.39 36.82	Bagging SVM 50.2 49.16 37.13	AdaBoost DT 54 51.13 42.44	AdaBoost SVM 45.29 47.39 32.59
Both Views Imaging View Genetic View	Linear K 49.33 49.99 41.28	Poly K 34.83 34.83 34.83	RBF K 34.83 34.83 34.83	Bagging DT 50.66 50.47 46.71	Bagging SVM 42.26 36.02 34.83	AdaBoost DT 49.05 53.82 38.39	AdaBoost SVM 36.94 37.73 40.74	Linear K 49.42 47.8 38.78	Poly K 49.49 51.12 40.71	RBF K 49.49 51.12 40.71	Bagging DT 48.51 52.39 36.82	Bagging SVM 50.2 49.16 37.13	AdaBoost DT 54 51.13 42.44	AdaBoost SVM 45.29 47.39 32.59
Both Views Imaging View Genetic View Balanced Accuracy	Linear K 49.33 49.99 41.28	Poly K 34.83 34.83 34.83	RBF K 34.83 34.83 34.83	Bagging DT 50.66 50.47 46.71 Original Data	Bagging SVM 42.26 36.02 34.83	AdaBoost DT 49.05 53.82 38.39	AdaBoost SVM 36.94 37.73 40.74	Linear K 49.42 47.8 38.78	Poly K 49.49 51.12 40.71	Scal RBF K 49.49 51.12 40.71 Scal	Bagging DT 48.51 52.39 36.82	Bagging SVM 50.2 49.16 37.13 Data	AdaBoost DT 54 51.13 42.44	AdaBoost SVM 45.29 47.39 32.59
Both Views Imaging View Genetic View Balanced Accuracy	Linear K 49.33 49.99 41.28	Poly K 34.83 34.83 34.83 Poly K	RBF K 34.83 34.83 34.83 RBF K	Bagging DT 50.66 50.47 46.71 Original Data Bagging DT	Bagging SVM 42.26 36.02 34.83 Bagging SVM	AdaBoost DT 49.05 53.82 38.39 AdaBoost DT	AdaBoost SVM 36.94 37.73 40.74 AdaBoost SVM	Linear K 49.42 47.8 38.78 Linear K	Poly K 49.49 51.12 40.71	Scal RBF K 49.49 51.12 40.71 Scal Scal RBF K	Bagging DT 48.51 52.39 36.82 ed and Balanced Bagging DT	Bagging SVM 50.2 49.16 37.13 Data Bagging SVM	AdaBoost DT 54 51.13 42.44 AdaBoost DT	AdaBoost SVM 45.29 47.39 32.59 AdaBoost SVM
Both Views Imaging View Genetic View Balanced Accuracy Both Views	Linear K 49.33 49.99 41.28 Linear K 47.5	Poly K 34.83 34.83 34.83 Poly K 33.33	RBF K 34.83 34.83 34.83 84.84 84.84 84.84 84.84 84.84 84.84 84.84 84 84 84 84 84 84 84 84 84 84 84 84 8	Bagging DT 50.66 50.47 46.71 Original Data Bagging DT 49.09	Bagging SVM 42.26 36.02 34.83 Bagging SVM 39.41	AdaBoost DT 49.05 53.82 38.39 AdaBoost DT 47.58	AdaBoost SVM 36.94 37.73 40.74 AdaBoost SVM 34.48	Linear K 49.42 47.8 38.78 Linear K 54.96	Poly K 49.49 51.12 40.71 Poly K 54.78	Scal RBF K 49.49 51.12 40.71 Scal RBF K 54.78	ed and Balanced Bagging DT 48.51 52.39 36.82 ed and Balanced Bagging DT 54.59	Bagging SVM 50.2 49.16 37.13 Data Bagging SVM 54.46	AdaBoost DT 54 51.13 42.44 AdaBoost DT 56.65	AdaBoost SVM 45.29 47.39 32.59 AdaBoost SVM 52.96
Both Views Imaging View Genetic View Balanced Accuracy Both Views Imaging View	Linear K 49.33 49.99 41.28 Linear K 47.5 48.33	Poly K 34.83 34.83 34.83 Poly K 33.33 33.33	RBF K 34.83 34.83 34.83 RBF K 33.33 33.33	Original Data Bagging DT 50.66 50.47 46.71 Original Data Bagging DT 49.09 46.97	Bagging SVM 42.26 36.02 34.83 Bagging SVM 39.41 39.16	AdaBoost DT 49.05 53.82 38.39 AdaBoost DT 47.58 53.97	AdaBoost SVM 36.94 37.73 40.74 AdaBoost SVM 34.48 33.07	Linear K 49.42 47.8 38.78 Linear K 54.96 51.94	Poly K 49.49 51.12 40.71 Poly K 54.78 55.3	Scal RBF K 49.49 51.12 40.71 Scal RBF K 55.3	Barging DT 48.51 52.39 36.82 ed and Balanced Bagging DT 54.59 55.7	Bagging SVM 50.2 49.16 37.13 Data Bagging SVM 54.46 53.81	AdaBoost DT 54 51.13 42.44 AdaBoost DT 56.65 54.73	AdaBoost SVM 45.29 47.39 32.59 AdaBoost SVM 52.96 49.79

 $\Sigma\chi\eta\mu\alpha$ 6.11: For each model and classifier, the metric scores for raw data classification are presented. Highlighted green are the best performing models, for each metric.

							DCCA									
Accuracy %				Original Data						Scal	ed and Balanced	Data				
	Linear K	Poly K	RBF K	Bagging DT	Bagging SVM	AdaBoost DT	AdaBoost SVM	Linear K	Poly K	RBF K	Bagging DT	Bagging SVM	AdaBoost DT	AdaBoost SVM		
Both Views	47.13	50.19	50.19	49.81	42.15	47.13	46.36	48.63	47.95	47.95	43.15	50	39.04	41.78		
Imaging View	51.72	52.11	52.11	47.89	45.21	48.28	45.98	50	48.63	48.63	39.04	43.84	43.84	41.1		
Genetic View	44.44	51.34	51.34	52.49	42.15	50.96	45.59	37.67	36.3	36.3	36.3	39.04	38.36	39.73		
F1 Score				Original Data				Scaled and Balanced Data								
	Linear K	Poly K	RBF K	Bagging DT	Bagging SVM	AdaBoost DT	AdaBoost SVM	Linear K	Poly K	RBF K	Bagging DT	Bagging SVM	AdaBoost DT	AdaBoost SVM		
Both Views	46.4	38.16	38.16	38.81	42.63	42.1	46.59	A7 A	47.09	47.09	38.97	48.27	38.76	39.6		
		50.10	00.40					47.4								
Imaging View	50.8	47.76	47.76	42.31	45.15	43.36	46.01	49.01	48.08	48.08	38.58	42.63	43.9	38.05		
Imaging View Genetic View	50.8 40.35	47.76	47.76	42.31 40.54	45.15 41.43	43.36 46.28	46.01 44.78	49.01 36.73	48.08 35.12	48.08 35.12	38.58 35.25	42.63 38.82	43.9 38.12	38.05 39.43		
Imaging View Genetic View	50.8 40.35	47.76 34.83	47.76 34.83	42.31 40.54	45.15 41.43	43.36 46.28	46.01 44.78	49.01 36.73	48.08 35.12	48.08 35.12	38.58 35.25	42.63 38.82	43.9 38.12	38.05 39.43		
Imaging View Genetic View	50.8 40.35	47.76	47.76	42.31 40.54	45.15 41.43	43.36 46.28	46.01 44.78	49.01 36.73	48.08 35.12	48.08 35.12	38.58 35.25	42.63 38.82	43.9 38.12	38.05 39.43		
Imaging View Genetic View Balanced Accuracy	50.8 40.35	47.76 34.83	47.76 34.83	42.31 40.54 Original Data	45.15 41.43	43.36 46.28	46.01 44.78	49.01 36.73	48.08 35.12	48.08 35.12 Scale	38.58 35.25 ed and Balanced	42.63 38.82 Data	43.9 38.12	38.05 39.43		
Imaging View Genetic View Balanced Accuracy	50.8 40.35 Linear K	47.76 34.83 Poly K	47.76 34.83 RBF K	42.31 40.54 Original Data Bagging DT	45.15 41.43 Bagging SVM	43.36 46.28 AdaBoost DT	46.01 44.78 AdaBoost SVM	49.01 36.73	48.08 35.12 Poly K	48.08 35.12 <u>Scal</u> <i>RBF K</i>	38.58 35.25 ad and Balanced Bagging DT	42.63 38.82 Data Bagging SVM	43.9 38.12 AdaBoost DT	38.05 39.43 AdaBoost SVM		
Imaging View Genetic View Balanced Accuracy Both Views	50.8 40.35 Linear K 42.04	47.76 34.83 Poly K 33.73	47.76 34.83 RBF K 33.73	42.31 40.54 Original Data Bagging DT 33.8	45.15 41.43 Bagging SVM 37.05	43.36 46.28 AdaBoost DT 36.52	46.01 44.78 AdaBoost SVM 44.85	49.01 36.73 Linear K 49.97	48.08 35.12 Poly K 49.13	48.08 35.12 <u>Scal</u> <i>RBF K</i> 49.13	38.58 35.25 ad and Balanced Bagging DT 43.98	42.63 38.82 Data Bagging SVM 50.64	43.9 38.12 AdaBoost DT 39.74	38.05 39.43 AdaBoost SVM 43.38		
Imaging View Genetic View Balanced Accuracy Both Views Imaging View	50.8 40.35 Linear K 42.04 45.82	47.76 34.83 Poly K 33.73 40.45	47.76 34.83 <i>RBF K</i> 33.73 40.45	42.31 40.54 Original Data Bagging DT 33.8 35.61	45.15 41.43 Bagging SVM 37.05 42.85	43.36 46.28 AdaBoost DT 36.52 36.21	46.01 44.78 AdaBoost SVM 44.85 45.25	49.01 36.73 Linear K 49.97 51.37	48.08 35.12 Poly K 49.13 49.6	48.08 35.12 Scale <i>RBF K</i> 49.13 49.6	38.58 35.25 ad and Balanced Bagging DT 43.98 39.4	42.63 38.82 Data Bagging SVM 50.64 44.55	43.9 38.12 AdaBoost DT 39.74 43.77	38.05 39.43 AdaBoost SVM 43.38 42.28		

 $\Sigma\chi\eta\mu\alpha$ 6.12: For each model and classifier, the metric scores for DCCA transformed data classification are presented. Highlighted green are the best performing models, for each metric.

6.2 MCA vs MCA - DCCA

Moving on from the classification of the raw data versus the DCCA transformed data, we explore the effect that MCA has, and introduce the results of the classification. In this part, we present the classification results, (a) of the data after the genetic view has been transformed through MCA, and (b) of the DCCA transformed data after the genetic view has been transformed through MCA.

6.2.1 Without scaling or balancing:



Σχήμα 6.13: Classification metric scores using Both views (Imaging and Genetic), on the SVM kernels (Linear, Polynomial, RBF), using raw imaging and MCA transformed genetic data (3 left bar groups) vs using DCCA transformed raw imaging and MCA transformed genetic data (3 right bar groups)



 $\Sigma\chi\eta\mu\alpha$ 6.14: Classification metric scores using only the Imaging view, on the SVM kernels (Linear, Polynomial, RBF), using raw imaging data (3 left bar groups) vs using the DCCA transformed imaging data, trained on raw imaging data and MCA transformed genetic data (3 right bar groups).



 $\Sigma\chi\eta\mu\alpha$ 6.15: Classification metric scores using only the genetic view, on the SVM kernels (Linear, Polynomial, RBF), using MCA transformed genetic data (3 left bar groups) vs using the DCCA transformed genetic data, trained on raw imaging data and MCA transformed genetic data (3 right bar groups).



 Σ χήμα 6.16: Classification metric using Bagging on the MCA transformed imaging and genetic data.



 Σ χήμα 6.17: Classification metric using AdaBoost on the MCA transformed imaging and genetic data.



6.2.2 With scaling and balancing:

 $\Sigma \chi \eta \mu \alpha$ 6.18: Classification metric scores using Both views (Imaging and Genetic), on the SVM kernels (Linear, Polynomial, RBF), using raw imaging and MCA transformed genetic data (3 left bar groups) vs using DCCA transformed raw imaging and MCA transformed genetic data (3 right bar groups)



 $\Sigma\chi\eta\mu\alpha$ 6.19: Classification metric scores using only the Imaging view, on the SVM kernels (Linear, Polynomial, RBF), using raw imaging data (3 left bar groups) vs using the DCCA transformed imaging data, trained on raw imaging data and MCA transformed genetic data (3 right bar groups).



 $\Sigma\chi\eta\mu\alpha$ 6.20: Classification metric scores using only the genetic view, on the SVM kernels (Linear, Polynomial, RBF), using MCA transformed genetic data (3 left bar groups) vs using the DCCA transformed genetic data, trained on raw imaging data and MCA transformed genetic data (3 right bar groups).

The confusion matrices of the MCA and MCA - DCCA models are shown, because they performed exceptionally good:



 $\Sigma\chi\eta\mu\alpha$ 6.21: The Confusion Matrices for each class, per model, using both views (top row), only the imaging view (middle row), and only the genetic view (bottom row). The three left columns represent the CM of the raw imaging and MCA transformed genetic data classification, while the three right columns represent the CM of the DCCA transformed data classification.



 Σ χήμα 6.22: Classification metric using Bagging on the MCA transformed imaging and genetic data.



Σχήμα 6.23: Classification metric using AdaBoost on the MCA transformed imaging and genetic data.

The following tables present the complete results for the MCA transformed data, as well as the DCCA transformed data after MCA transformation on the genetic data, for each model, for each metric, for each view, and either with or without scaling and balancing. With green are highlighted the best values for each metric, depending on whether scaling and balancing were applied:

							MCA							
Accuracy %				Original Data						Scal	ed and Balanced	Data		
	Linear K	Poly K	RBF K	Bagging DT	Bagging SVM	AdaBoost DT	AdaBoost SVM	Linear K	Poly K	RBF K	Bagging DT	Bagging SVM	AdaBoost DT	AdaBoost SVM
Both Views	50.19	51.34	51.34	51.72	42.91	49.81	37.16	57.53	59.59	59.59	54.11	52.05	53.42	49.32
Imaging View														
Genetic View	49.04	51.34	51.34	51.72	49.81	48.28	49.81	37.67	30.14	30.14	31.51	35.62	30.82	38.36
F1 Score				Original Data						Scal	ed and Balanced	Data		
F1 Score	Linear K	Poly K	RBF K	Original Data Bagging DT	Bagging SVM	AdaBoost DT	AdaBoost SVM	Linear K	Poly K	Scal RBF K	ed and Balanced Bagging DT	Data Bagging SVM	AdaBoost DT	AdaBoost SVM
F1 Score Both Views	Linear K 49.99	Poly K 34.83	RBF K 34.83	Original Data Bagging DT 49.25	Bagging SVM 43.96	AdaBoost DT 49.62	AdaBoost SVM 36.25	Linear K 52.6	Poly K 57.85	<u>Scal</u> RBF K 57.85	ed and Balanced Bagging DT 50.29	Data Bagging SVM 48.15	AdaBoost DT 52.36	AdaBoost SVM 48.43
F1 Score Both Views Imaging View	Linear K 49.99	Poly K 34.83	RBF K 34.83	Original Data Bagging DT 49.25	Bagging SVM 43.96	AdaBoost DT 49.62	AdaBoost SVM 36.25	Linear K 52.6	Poly K 57.85	<u>Scal</u> RBF K 57.85	ed and Balanced Bagging DT 50.29	Data Bagging SVM 48.15	AdaBoost DT 52.36	AdaBoost SVM 48.43
F1 Score Both Views Imaging View Genetic View	Linear K 49.99 35.19	Poly K 34.83 34.83	RBF K 34.83 34.83	Original Data Bagging DT 49.25 36.96	Bagging SVM 43.96 35.01	AdaBoost DT 49.62 35.24	AdaBoost SVM 36.25 34.92	Linear K 52.6 37.21	Poly K 57.85 13.96	<u>Scal</u> <i>RBF K</i> 57.85 13.96	ed and Balanced Bagging DT 50.29 27.62	Data Bagging SVM 48.15 33.61	AdaBoost DT 52.36 30.6	AdaBoost SVM 48.43 38.45
F1 Score Both Views Imaging View Genetic View	Linear K 49.99 35.19	Poly K 34.83 34.83	RBF K 34.83 34.83	Original Data Bagging DT 49.25 36.96	Bagging SVM 43.96 35.01	AdaBoost DT 49.62 35.24	AdaBoost SVM 36.25 34.92	Linear K 52.6 37.21	Poly K 57.85 13.96	<u>Scal</u> RBF K 57.85 13.96	ed and Balanced Bagging DT 50.29 27.62	Data Bagging SVM 48.15 33.61	AdaBoost DT 52.36 30.6	AdaBoost SVM 48.43 38.45
F1 Score Both Views Imaging View Genetic View	Linear K 49.99 35.19	Poly K 34.83 34.83	RBF K 34.83 34.83	Original Data Bagging DT 49.25 36.96	Bagging SVM 43.96 35.01	AdaBoost DT 49.62 35.24	AdaBoost SVM 36.25 34.92	Linear K 52.6 37.21	Poly K 57.85 13.96	<u>Scal</u> RBF K 57.85 13.96	ed and Balanced Bagging DT 50.29 27.62	Data Bagging SVM 48.15 33.61	AdaBoost DT 52.36 30.6	AdaBoost SVM 48.43 38.45
F1 Score Both Views Imaging View Genetic View	Linear K 49.99 35.19	Poly K 34.83 34.83	RBF K 34.83 34.83	Original Data Bagging DT 49.25 36.96	Bagging SVM 43.96 35.01	AdaBoost DT 49.62 35.24	AdaBoost SVM 36.25 34.92	Linear K 52.6 37.21	Poly K 57.85 13.96	<u>Scal</u> RBF K 57.85 13.96	ed and Balanced Bagging DT 50.29 27.62	Data Bagging SVM 48.15 33.61	AdaBoost DT 52.36 30.6	AdaBoost SVM 48.43 38.45
F1 Score Both Views Imaging View Genetic View Balanced Accuracy	Linear K 49.99 35.19	Poly K 34.83 34.83	RBF K 34.83 34.83	Original Data Bagging DT 49.25 36.96 Original Data	Bagging SVM 43.96 35.01	AdaBoost DT 49.62 35.24	AdaBoost SVM 36.25 34.92	Linear K 52.6 37.21	Poly K 57.85 13.96	Scal RBF K 57.85 13.96 Scal	ed and Balanced Bagging DT 50.29 27.62 ed and Balanced	Data Bagging SVM 48.15 33.61 Data	AdaBoost DT 52.36 30.6	AdaBoost SVM 48.43 38.45
F1 Score Both Views Imaging View Genetic View Balanced Accuracy	Linear K 49.99 35.19 Linear K	Poly K 34.83 34.83 Poly K	RBF K 34.83 34.83 RBF K	Original Data Bagging DT 49.25 36.96 Original Data Bagging DT	Bagging SVM 43.96 35.01 Bagging SVM	AdaBoost DT 49.62 35.24 AdaBoost DT	AdaBoost SVM 36.25 34.92 AdaBoost SVM	Linear K 52.6 37.21 Linear K	Poly K 57.85 13.96 Poly K	<u>Scal</u> <i>RBF K</i> 57.85 13.96 <u>Scal</u> <i>RBF K</i>	ed and Balanced Bagging DT 50.29 27.62 ed and Balanced Bagging DT	Data Bagging SVM 48.15 33.61 Data Bagging SVM	AdaBoost DT 52.36 30.6 AdaBoost DT	AdaBoost SVM 48.43 38.45 AdaBoost SVM
F1 Score Both Views Imaging View Genetic View Balanced Accuracy Both Views	Linear K 49.99 35.19 Linear K 48.33	Poly K 34.83 34.83 Poly K 33.33	RBF K 34.83 34.83	Original Data Bagging DT 49.25 36.96 Original Data Bagging DT 47.17	Bagging SVM 43.96 35.01 Bagging SVM 40.35	AdaBoost DT 49.62 35.24 AdaBoost DT 48.08	AdaBoost SVM 36.25 34.92 AdaBoost SVM 35.59	Linear K 52.6 37.21 Linear K 57.24	Poly K 57.85 13.96 Poly K 59.41	<u>Scal</u> <i>RBF K</i> 57.85 13.96 <u>Scal</u> <i>RBF K</i> 59.41	ed and Balanced Bagging DT 50.29 27.62 ed and Balanced Bagging DT 56.38	Data Bagging SVM 48.15 33.61 Data Bagging SVM 54.67	AdaBoost DT 52.36 30.6 AdaBoost DT 53.98	AdaBoost SVM 48.43 38.45 AdaBoost SVM 49.88
FL Score Both Views Imaging View Genetic View Balanced Accuracy Both Views Imaging View	Linear K 49.99 35.19 Linear K 48.33	Poly K 34.83 34.83 Poly K 33.33	RBF K 34.83 34.83 RBF K 33.33	Original Data Bagging DT 49.25 36.96 Original Data Bagging DT 47.17	Bagging SVM 43.96 35.01 Bagging SVM 40.35	AdaBoost DT 49.62 35.24 AdaBoost DT 48.08	AdaBoost SVM 36.25 34.92 AdaBoost SVM 35.59	Linear K 52.6 37.21 Linear K 57.24	Poly K 57.85 13.96 Poly K 59.41	<u>Scal</u> <i>RBF K</i> 57.85 13.96 <u>Scal</u> <i>RBF K</i> 59.41	ed and Balanced Bagging DT 50.29 27.62 ed and Balanced Bagging DT 56.38	Data Bagging SVM 48.15 33.61 Data Bagging SVM 54.67	AdaBoost DT 52.36 30.6 AdaBoost DT 53.98	AdaBoost SVM 48.43 38.45 AdaBoost SVM 49.88

 $\Sigma\chi\eta\mu\alpha$ 6.24: For each model and classifier, the metric scores for MCA transformed data classification are presented. Highlighted green are the best performing models, for each metric.

						MCA	+ DC	CA						
Accuracy %				Original Data						Scale	d and Balancer	Data		
riccuracy /o	Linear K	Poly K	RBF K	Bagging DT	Bagging SVM	AdaBoost DT	AdaBoost SVM	Linear K	Poly K	RBF K	Bagging DT	Bagging SVM	AdaBoost DT	AdaBoost SVM
Both Views	46.36	50.96	50.96	44.44	45.59	46.74	49.81	40.41	37.67	37.67	45.89	41.1	45.21	42.47
Imaging View	42.91	47.89	47.89	46.36	46.74	47.13	41.38	50	41.48	41.48	43.15	52.05	45.21	45.21
Genetic View	39.08	51.34	51.34	51.34	43.68	43.68	42.53	40.41	34.93	34.93	31.51	33.56	38.36	38.36
F1 Score				Onininal Data										
				Original Data						Scale	d and Balanced	Data		
	Linear K	Poly K	RBF K	Bagging DT	Bagging SVM	AdaBoost DT	AdaBoost SVM	Linear K	Poly K	RBF K	Bagging DT	Bagging SVM	AdaBoost DT	AdaBoost SVM
Both Views	Linear K 46.75	Poly K 36.52	RBF K 36.52	Bagging DT 38.82	Bagging SVM 46.03	AdaBoost DT 41.42	AdaBoost SVM 49.71	Linear K 39.99	Poly K 37.69	RBF K 37.69	Bagging DT 35.49	Bagging SVM 40.55	AdaBoost DT 35.86	AdaBoost SVM 42.08
Both Views Imaging View	Linear K 46.75 41.7	Poly K 36.52 40.13	RBF K 36.52 40.13	Bagging DT 38.82 41.77	Bagging SVM 46.03 46.92	AdaBoost DT 41.42 39.33	AdaBoost SVM 49.71 41.1	Linear K 39.99 48.48	Poly K 37.69 40.4	RBF K 37.69 40.4	Bagging DT 35.49 42.82	Data Bagging SVM 40.55 50.1	AdaBoost DT 35.86 35.86	AdaBoost SVM 42.08 36.71
Both Views Imaging View Genetic View	Linear K 46.75 41.7 39.39	Poly K 36.52 40.13 34.83	RBF K 36.52 40.13 34.83	Bagging DT 38.82 41.77 35.5	Bagging SVM 46.03 46.92 43.45	AdaBoost DT 41.42 39.33 38.87	AdaBoost SVM 49.71 41.1 42.86	Linear K 39.99 48.48 40.14	Poly K 37.69 40.4 34.88	Scale RBF K 37.69 40.4 34.88	Bagging DT 35.49 42.82 29.45	Data Bagging SVM 40.55 50.1 33.73	AdaBoost DT 35.86 35.86 37.57	AdaBoost SVM 42.08 36.71 37.65
Both Views Imaging View Genetic View	Linear K 46.75 41.7 39.39	Poly K 36.52 40.13 34.83	RBF K 36.52 40.13 34.83	Bagging DT 38.82 41.77 35.5	Bagging SVM 46.03 46.92 43.45	AdaBoost DT 41.42 39.33 38.87	AdaBoost SVM 49.71 41.1 42.86	Linear K 39.99 48.48 40.14	Poly K 37.69 40.4 34.88	Scale RBF K 37.69 40.4 34.88	d and Balanced Bagging DT 35.49 42.82 29.45	Data Bogging SVM 40.55 50.1 33.73	AdaBoost DT 35.86 35.86 37.57	AdaBoost SVM 42.08 36.71 37.65
Both Views Imaging View Genetic View	Linear K 46.75 41.7 39.39	Poly K 36.52 40.13 34.83	RBF K 36.52 40.13 34.83	Bagging DT 38.82 41.77 35.5	Bagging SVM 46.03 46.92 43.45	AdaBoost DT 41.42 39.33 38.87	AdaBoost SVM 49.71 41.1 42.86	Linear K 39.99 48.48 40.14	Poly K 37.69 40.4 34.88	Scale RBF K 37.69 40.4 34.88	Bagging DT 35.49 42.82 29.45	Data Bagging SVM 40.55 50.1 33.73	AdaBoost DT 35.86 35.86 37.57	AdaBoost SVM 42.08 36.71 37.65
Both Views Imaging View Genetic View	Linear K 46.75 41.7 39.39	Poly K 36.52 40.13 34.83	RBF K 36.52 40.13 34.83	Bagging DT 38.82 41.77 35.5	Bagging SVM 46.03 46.92 43.45	AdaBoost DT 41.42 39.33 38.87	AdaBoost SVM 49.71 41.1 42.86	Linear K 39.99 48.48 40.14	Poly K 37.69 40.4 34.88	Scale <i>RBF K</i> 37.69 40.4 34.88	Bagging DT 35.49 42.82 29.45	Bagging SVM 40.55 50.1 33.73	AdaBoost DT 35.86 35.86 37.57	AdaBoost SVM 42.08 36.71 37.65
Both Views Imaging View Genetic View Balanced Accuracy	Linear K 46.75 41.7 39.39	Poly K 36.52 40.13 34.83	RBF K 36.52 40.13 34.83	Bagging DT 38.82 41.77 35.5	Bagging SVM 46.03 46.92 43.45	AdaBoost DT 41.42 39.33 38.87	AdaBoost SVM 49.71 41.1 42.86	Linear K 39.99 48.48 40.14	Poly K 37.69 40.4 34.88	Scale RBF K 37.69 40.4 34.88 Scale	d and Balancec Bagging DT 35.49 42.82 29.45 29.45	Data Bagging SVM 40.55 50.1 33.73	AdaBoost DT 35.86 35.86 37.57	AdaBoost SVM 42.08 36.71 37.65
Both Views Imaging View Genetic View Balanced Accuracy	Linear K 46.75 41.7 39.39	Poly K 36.52 40.13 34.83 Poly K	RBF K 36.52 40.13 34.83 RBF K	Bagging DT 38.82 41.77 35.5 Original Data Bagging DT	Bagging SVM 46.03 46.92 43.45 Bagging SVM	AdaBoost DT 41.42 39.33 38.87 AdaBoost DT	AdaBoost SVM 49.71 41.1 42.86 AdaBoost SVM	Linear K 39.99 48.48 40.14 Linear K	Poly K 37.69 40.4 34.88 Poly K	Scale RBF K 37.69 40.4 34.88 Scale RBF K	d and Balanced Bagging DT 35.49 42.82 29.45 d and Balanced Bagging DT	Bagging SVM 40.55 50.1 33.73 Data Bagging SVM	AdaBoost DT 35.86 35.86 37.57 AdaBoost DT	AdaBoost SVM 42.08 36.71 37.65 AdaBoost SVM
Both Views Imaging View Genetic View Balanced Accuracy Both Views	Linear K 46.75 41.7 39.39 Linear K 46.9	Poly K 36.52 40.13 34.83 Poly K 33.57	RBF K 36.52 40.13 34.83 RBF K 33.57	Bagging DT 38.82 41.77 35.5 Original Data Bagging DT 32.42	Bagging SVM 46.03 46.92 43.45 Bagging SVM 45.46	AdaBoost DT 41.42 39.33 38.87 AdaBoost DT 35.07	AdaBoost SVM 49.71 41.1 42.86 AdaBoost SVM 45.04	Linear K 39.99 48.48 40.14 Linear K 41.01	Poly K 37.69 40.4 34.88 Poly K 37.91	Scale RBF K 37.69 40.4 34.88 Scale RBF K 37.91	d and Balanced Bagging DT 35.49 42.82 29.45 d and Balanced Bagging DT 48.56	Data Bagging SVM 40.55 50.1 33.73 Data Bagging SVM 41.28	AdaBoost DT 35.86 35.86 37.57 AdaBoost DT 46.31	AdaBoost SVM 42.08 36.71 37.65 AdaBoost SVM 42.66
Both Views Imaging View Genetic View Balanced Accuracy Both Views Imaging View	Linear K 46.75 41.7 39.39 Linear K 46.9 37.47	Poly K 36.52 40.13 34.83 Poly K 33.57 34.63	RBF K 36.52 40.13 34.83	Original Data Bagging DT 38.82 41.77 35.5 Original Data Bagging DT 32.42 35.96	Bagging SVM 46.03 46.92 43.45 Bagging SVM 45.46 46.19	AdaBoost DT 41.42 39.33 38.87 AdaBoost DT 35.07 33.2	AdaBoost SVM 49.71 41.1 42.86 AdaBoost SVM 45.04 40.13	Linear K 39.99 48.48 40.14 Linear K 41.01 51.84	Poly K 37.69 40.4 34.88 Poly K 37.91 43.13	Scale RBF K 37.69 40.4 34.88 Scale RBF K 37.91 43.13	and Balance Bagging DT 35.49 42.82 29.45 cd and Balancee Bagging DT 48.56 43.62	Data Bagging SVM 40.55 50.1 33.73 Data Bagging SVM 41.28 53.37	AdaBoost DT 35.86 35.86 37.57 AdaBoost DT 46.31 46.31	AdaBoost SVM 42.08 36.71 37.65 AdaBoost SVM 42.66 46.27

 $\Sigma\chi\eta\mu\alpha$ 6.25: For each model and classifier, the metric scores for MCA - DCCA transformed data classification are presented. Highlighted green are the best performing models, for each metric.

6.3 OPNMF vs OPNMF - DCCA

Furthermore, we explore the effect that OPNMF has to the task of classification. To that extent, we compare the classification results of (a) the data after the imaging view has been transformed with OPNMF and (b) the DCCA transformed data, trained on the raw genetic data and the OPNMF transformed imaging data.

6.3.1 Without scaling or balancing:



 $\Sigma\chi\eta\mu\alpha$ 6.26: Classification metric scores using Both views (Imaging and Genetic), on the SVM kernels (Linear, Polynomial, RBF), using raw genetic and OPNMF transformed imaging data (3 left bar groups) vs using DCCA transformed raw genetic and OPNMF transformed imaging data (3 right bar groups)



 $\Sigma\chi\eta\mu\alpha$ 6.27: Classification metric scores using only the Imaging view, on the SVM kernels (Linear, Polynomial, RBF), using OPNMF transformed imaging data (3 left bar groups) vs using the DCCA transformed imaging data, trained on raw genetic data and OPNMF transformed genetic data (3 right bar groups).



 Σ χήμα 6.28: Classification metric scores using only the Imaging view, on the SVM kernels (Linear, Polynomial, RBF), using OPNMF transformed imaging data (3 left bar groups) vs using the DCCA transformed imaging data, trained on raw genetic data and OPNMF transformed genetic data (3 right bar groups).



 Σ χήμα 6.29: Classification metric using Bagging on the OPNMF transformed imaging and genetic data.



 Σ χήμα 6.30: Classification metric using AdaBoost on the OPNMF transformed imaging and genetic data.



6.3.2 With scaling and balancing:

 $\Sigma\chi\eta\mu\alpha$ 6.31: Classification metric scores using Both views (Imaging and Genetic), on the SVM kernels (Linear, Polynomial, RBF), using raw genetic and OPNMF transformed imaging data (3 left bar groups) vs using DCCA transformed raw genetic and OPNMF transformed imaging data (3 right bar groups)



 Σ χήμα 6.32: Classification metric scores using only the Imaging view, on the SVM kernels (Linear, Polynomial, RBF), using OPNMF transformed imaging data (3 left bar groups) vs using the DCCA transformed imaging data, trained on raw genetic data and OPNMF transformed genetic data (3 right bar groups).



 $\Sigma\chi\eta\mu\alpha$ 6.33: Classification metric scores using only the genetic view, on the SVM kernels (Linear, Polynomial, RBF), using raw genetic data (3 left bar groups) vs using the DCCA transformed genetic data, trained on raw genetic data and OPNMF transformed imaging data (3 right bar groups).



Σχήμα 6.34: Classification metric using Bagging on the OPNMF transformed imaging and genetic data.



Σχήμα 6.35: Classification metric using AdaBoost on the OPNMF transformed imaging and genetic data.

The confusion matrices of the OPNMF Bagging models are shown, because they performed exceptionally good:



 $\Sigma\chi\eta\mu\alpha$ 6.36: The Confusion Matrices for each class, with Bagging, for the OPNMF transformed imaging and genetic data.



 $\Sigma\chi\eta\mu\alpha$ 6.37: The Confusion Matrices for each class, with AdaBoost, for the OPNMF transformed imaging and genetic data.

The following tables present the complete results for the OPNMF transformed data, as well as the DCCA transformed data after OPMNF transformation on the imaging data, for each model, for each metric, for each view, and either with or without scaling and balancing. With green are highlighted the best values for each metric, depending on whether scaling and balancing were applied:

						PNM	IF							
Accuracy %				Original Data						Scale	d and Balanced	Data		
<u>needidey no</u>	Linear K	Poly K	RBF K	Bagging DT	Bagging SVM	AdaBoost DT	AdaBoost SVM	Linear K	Poly K	RBF K	Bagging DT	Bagging SVM	AdaBoost DT	AdaBoost SVM
Both Views	51.74	51.35	51.35	47.49	52.9	48.26	53.28	48.28	28.97	28.97	49.66	55.17	48.26	53.28
Imaging View	53.28	51.35	51.35	46.33	51.35	51.74	53.28	51.03	28.97	28.97	55.86	58.62	51.74	53.28
Genetic View														
F1 Score				Original Data						Scale	d and Balanced	Data		
	Linear K	Poly K	RBF K	Bagging DT	Bagging SVM	AdaBoost DT	AdaBoost SVM	Linear K	Poly K	RBF K	Bagging DT	Bagging SVM	AdaBoost DT	AdaBoost SVM
Both Views	Linear K 51.51	Poly K 34.85	RBF K 34.85	Bagging DT 46.33	50.03	AdaBoost DT 47.98	AdaBoost SVM 51.46	Linear K 44.16	Poly K 13.01	RBF K 13.01	Bagging DT 47.28	Bagging SVM 49.4	AdaBoost DT 47.98	AdaBoost SVM 51.46
Both Views Imaging View	51.51 53.16	Poly K 34.85 34.85	RBF K 34.85 34.85	Bagging DT 46.33 45.86	50.03 50.52	AdaBoost DT 47.98 51.46	AdaBoost SVM 51.46 51.46	Linear K 44.16 48.08	Poly K 13.01 13.01	RBF K 13.01 13.01	Bagging DT 47.28 53.45	Bagging SVM 49.4 54.2	AdaBoost DT 47.98 51.46	AdaBoost SVM 51.46 51.46
Both Views Imaging View Genetic View	Linear K 51.51 53.16	Poly K 34.85 34.85	RBF K 34.85 34.85	8agging DT 46.33 45.86	Bagging SVM 50.03 50.52	AdaBoost DT 47.98 51.46	AdaBoost SVM 51.46 51.46	Linear K 44.16 48.08	Poly K 13.01 13.01	RBF K 13.01 13.01	Bagging DT 47.28 53.45	Bagging SVM 49.4 54.2	AdaBoost DT 47.98 51.46	AdaBoost SVM 51.46 51.46
Both Views Imaging View Genetic View	Linear K 51.51 53.16	Poly K 34.85 34.85	RBF K 34.85 34.85	Bagging DT 46.33 45.86	Bagging SVM 50.03 50.52	AdaBoost DT 47.98 51.46	AdaBoost SVM 51.46 51.46	Linear K 44.16 48.08	Poly K 13.01 13.01	RBF K 13.01 13.01	Bagging DT 47.28 53.45	Bagging SVM 49.4 54.2	AdaBoost DT 47.98 51.46	AdaBoost SVM 51.46 51.46
Both Views Imaging View Genetic View	Linear K 51.51 53.16	Poly K 34.85 34.85	RBF K 34.85 34.85	Bagging DT 46.33 45.86	Bagging SVM 50.03 50.52	AdaBoost DT 47.98 51.46	AdaBoost SVM 51.46 51.46	Linear K 44.16 48.08	Poly K 13.01 13.01	RBF K 13.01 13.01	Bagging DT 47.28 53.45	Bagging SVM 49.4 54.2	AdaBoost DT 47.98 51.46	AdaBoost SVM 51.46 51.46
Both Views Imaging View Genetic View	Linear K 51.51 53.16	Poly K 34.85 34.85	RBF K 34.85 34.85	Bagging DT 46.33 45.86	Bagging SVM 50.03 50.52	AdaBoost DT 47.98 51.46	AdaBoost SVM 51.46 51.46	Linear K 44.16 48.08	Poly K 13.01 13.01	RBF K 13.01 13.01	Bagging DT 47.28 53.45	Bagging SVM 49.4 54.2	AdaBoost DT 47.98 51.46	AdaBoost SVM 51.46 51.46
Both Views Imaging View Genetic View	Linear K 51.51 53.16	Poly K 34.85 34.85	RBF K 34.85 34.85	Bagging DT 46.33 45.86 Original Data	Bagging SVM 50.03 50.52	AdaBoost DT 47.98 51.46	AdaBoost SVM 51.46 51.46	Linear K 44.16 48.08	Poly K 13.01 13.01	RBF K 13.01 13.01 <u>Scale</u>	Bagging DT 47.28 53.45 d and Balanced	Bagging SVM 49.4 54.2	AdaBoost DT 47.98 51.46	AdaBoost SVM 51.46 51.46
Both Views Imaging View Genetic View	Linear K	Poly K 34.85 34.85 Poly K	RBF K 34.85 34.85 	Bagging DT 46.33 45.86 Original Data Bagging DT	Bagging SVM 50.03 50.52 Bagging SVM	AdaBoost DT 47.98 51.46 AdaBoost DT	AdaBoost SVM 51.46 51.46 AdaBoost SVM	Linear K 44.16 48.08 Linear K	Poly K 13.01 13.01 Poly K	RBF K 13.01 13.01 Scale RBF K	Bagging DT 47.28 53.45 d and Balanced Bagging DT	Bagging SVM 49.4 54.2 Data Bagging SVM	AdaBoost DT 47.98 51.46 AdaBoost DT	AdaBoost SVM 51.46 51.46 AdaBoost SVM
Both Views Imaging View Genetic View Manced Accurat Both Views	Linear K 51.51 53.16 Linear K 48.64	Poly K 34.85 34.85 Poly K 33.33	RBF K 34.85 34.85 	Bagging DT 46.33 45.86 Original Data Bagging DT 41.57	Bagging SVM 50.03 50.52 Bagging SVM 44.51	AdaBoost DT 47.98 51.46 AdaBoost DT 46.72	AdaBoost SVM 51.46 51.46 AdaBoost SVM 45.3	Linear K 44.16 48.08 Linear K 52.55	Poly K 13.01 13.01 Poly K 33.33	RBF K 13.01 13.01 Scale RBF K 33.33	Bagging DT 47.28 53.45 d and Balanced Bagging DT 50.93	Bagging SVM 49.4 54.2 Data Bagging SVM 57.69	AdaBoost DT 47.98 51.46 AdaBoost DT 46.72	AdaBoost SVM 51.46 51.46 AdaBoost SVM 45.3
Both Views Imaging View Genetic View Ilanced Accura Both Views Imaging View	Linear K 51.51 53.16 Linear K 48.64 50.26	Poly K 34.85 34.85 Poly K 33.33 33.33	RBF K 34.85 34.85	Bagging DT 46.33 45.86 Original Data Bagging DT 41.57 42.9	Bagging SVM 50.03 50.52 Bagging SVM 44.51 45.78	AdaBoost DT 47.98 51.46 AdaBoost DT 46.72 48.71	AdaBoost SVM 51.46 51.46 AdaBoost SVM 45.3 45.3	Linear K 44.16 48.08 Linear K 52.55 54.79	Poly K 13.01 13.01 Poly K 33.33 33.33	RBF K 13.01 13.01 Scale RBF K 33.33 33.33	Bagging DT 47.28 53.45 d and Balanced Bagging DT 50.93 57.57	Bagging SVM 49.4 54.2 Data Bagging SVM 57.69 61.01	AdaBoost DT 47.98 51.46 47.98 40.00 40.00 40.00 40.72 48.71	AdaBoost SVM 51.46 51.46 40 AdaBoost SVM 45.3 45.3

 $\Sigma\chi\eta\mu\alpha$ 6.38: For each model and classifier, the metric scores for OPNMF transformed data classification are presented. Highlighted green are the best performing models, for each metric.

					0	PNN	1F + C	OCCA	L							
Accuracy %				Original Data						Scale	d and Balanced	Data				
	Linear K	Poly K	RBF K	Bagging DT	Bagging SVM	AdaBoost DT	AdaBoost SVM	Linear K	Poly K	RBF K	Bagging DT	Bagging SVM	AdaBoost DT	AdaBoost SVM		
Both Views	45.95	47.88	47.88	50.58	40.93	52.9	46.33	49.66	49.66	49.66	44.14	46.9	48.97	53.79		
Imaging View	48.65	49.42	49.42	50.97	47.49	43.24	50.19	49.66	49.66	49.66	43.45	50.34	47.59	51.03		
Genetic View	45.95	45.56	45.56	50.58	45.17	51.35	45.56	40.69	40.69	40.69	35.86	35.17	38.62	45.52		
E1 Score								Scaled and Balanced Data								
ra score				Original Data						Scale	d and Balanced	Data				
ri score	Linear K	Poly K	RBF K	Original Data Bagging DT	Bagging SVM	AdaBoost DT	AdaBoost SVM	Linear K	Poly K	Scale RBF K	d and Balanced Bagging DT	Data Bagging SVM	AdaBoost DT	AdaBoost SVM		
Both Views	Linear K 46.08	Poly K 47.25	RBF K 47.25	Original Data Bagging DT 42.38	Bagging SVM 40.56	AdaBoost DT 50.14	AdaBoost SVM 46.2	Linear K 46.52	Poly K 49.03	<u>Scale</u> RBF K 49.03	d and Balanced Bagging DT 43.68	Data Bagging SVM 42.04	AdaBoost DT 48.93	AdaBoost SVM 53.38		
Both Views Imaging View	Linear K 46.08 48.78	Poly K 47.25 48.36	RBF K 47.25 48.36	Original Data Bagging DT 42.38 48.27	Bagging SVM 40.56 47.33	AdaBoost DT 50.14 40.79	AdaBoost SVM 46.2 50.06	Linear K 46.52 45.55	Poly K 49.03 48.54	Scale <i>RBF K</i> 49.03 48.54	d and Balanced Bagging DT 43.68 42.13	Data Bagging SVM 42.04 46.74	AdaBoost DT 48.93 46.81	AdaBoost SVM 53.38 48.01		
Both Views Imaging View Genetic View	Linear K 46.08 48.78 44.72	Poly K 47.25 48.36 43.87	RBF K 47.25 48.36 43.87	Original Data Bagging DT 42.38 48.27 36.34	Bagging SVM 40.56 47.33 43.71	AdaBoost DT 50.14 40.79 46.94	AdaBoost SVM 46.2 50.06 43.69	Linear K 46.52 45.55 40.69	Poly K 49.03 48.54 40.7	Scale RBF K 49.03 48.54 40.7	d and Balanced Bagging DT 43.68 42.13 35.86	Data Bagging SVM 42.04 46.74 34.52	AdaBoost DT 48.93 46.81 38.28	AdaBoost SVM 53.38 48.01 45.52		
Both Views Imaging View Genetic View	Linear K 46.08 48.78 44.72	Poly K 47.25 48.36 43.87	RBF K 47.25 48.36 43.87	Original Data Bagging DT 42.38 48.27 36.34	Bagging SVM 40.56 47.33 43.71	AdaBoost DT 50.14 40.79 46.94	AdaBoost SVM 46.2 50.06 43.69	Linear K 46.52 45.55 40.69	Poly K 49.03 48.54 40.7	Scale <i>RBF K</i> 49.03 48.54 40.7	d and Balanced Bagging DT 43.68 42.13 35.86	Data Bagging SVM 42.04 46.74 34.52	AdaBoost DT 48.93 46.81 38.28	AdaBoost SVM 53.38 48.01 45.52		
Both Views Imaging View Genetic View	Linear K 46.08 48.78 44.72	Poly K 47.25 48.36 43.87	RBF K 47.25 48.36 43.87	Original Data Bagging DT 42.38 48.27 36.34	Bagging SVM 40.56 47.33 43.71	AdaBoost DT 50.14 40.79 46.94	AdaBoost SVM 46.2 50.06 43.69	Linear K 46.52 45.55 40.69	Poly K 49.03 48.54 40.7	Scale RBF K 49.03 48.54 40.7	d and Balanced Bagging DT 43.68 42.13 35.86	Data Bagging SVM 42.04 46.74 34.52	AdaBoost DT 48.93 46.81 38.28	AdaBoost SVM 53.38 48.01 45.52		
Both Views Imaging View Genetic View	Linear K 46.08 48.78 44.72	Poly K 47.25 48.36 43.87	RBF K 47.25 48.36 43.87	Original Data Bagging DT 42.38 48.27 36.34	Bagging SVM 40.56 47.33 43.71	AdaBoost DT 50.14 40.79 46.94	AdaBoost SVM 46.2 50.06 43.69	Linear K 46.52 45.55 40.69	Poly K 49.03 48.54 40.7	Scale RBF K 49.03 48.54 40.7	d and Balanced Bagging DT 43.68 42.13 35.86	Data Bagging SVM 42.04 46.74 34.52	AdaBoost DT 48.93 46.81 38.28	AdaBoost SVM 53.38 48.01 45.52		
Both Views Imaging View Genetic View Balanced Accuracy	Linear K 46.08 48.78 44.72	Poly K 47.25 48.36 43.87	RBF K 47.25 48.36 43.87	Original Data Bogging DT 42.38 48.27 36.34	Bagging SVM 40.56 47.33 43.71	AdaBoost DT 50.14 40.79 46.94	AdaBoost SVM 46.2 50.06 43.69	Linear K 46.52 45.55 40.69	Poly K 49.03 48.54 40.7	Scale <u>RBF K</u> 49.03 48.54 40.7 Scale	d and Balanced Bagging DT 43.68 42.13 35.86 d and Balanced	Data Bagging SVM 42.04 46.74 34.52 Data	AdaBoost DT 48.93 46.81 38.28	AdaBoost SVM 53.38 48.01 45.52		
Both Views Imaging View Genetic View Balanced Accuracy	Linear K 46.08 48.78 44.72 Linear K	Poly K 47.25 48.36 43.87 Poly K	RBF K 47.25 48.36 43.87 RBF K	Original Data Bagging DT 42.38 48.27 36.34 Original Data Bagging DT	Bagging SVM 40.56 47.33 43.71 Bagging SVM	AdaBoost DT 50.14 40.79 46.94 AdaBoost DT	AdaBoost SVM 46.2 50.06 43.69 AdaBoost SVM	Linear K 46.52 45.55 40.69 Linear K	Poly K 49.03 48.54 40.7 Poly K	Scale RBF K 49.03 48.54 40.7 Scale RBF K	d and Balanced Bagging DT 43.68 42.13 35.86 d and Balanced Bagging DT	Data Bagging SVM 42.04 46.74 34.52 Data Bagging SVM	AdaBoost DT 48.93 46.81 38.28 AdaBoost DT	AdaBoost SVM 53.38 48.01 45.52 AdaBoost SVM		
Both Views Balanced Accuracy Both Views	Linear K 46.08 48.78 44.72 Linear K 44.5	Poly K 47.25 48.36 43.87 Poly K 43.08	RBF K 47.25 48.36 43.87 RBF K 43.08	Original Data Bagging DT 42.38 48.27 36.34 Original Data Bagging DT 36.1	Bagging SVM 40.56 47.33 43.71 Bagging SVM 38.11	AdaBoost DT 50.14 40.79 46.94 AdaBoost DT 43.42	AdaBoost SVM 46.2 50.06 43.69 AdaBoost SVM 47.34	Linear K 46.52 45.55 40.69 Linear K 51.4	Paly K 49.03 48.54 40.7 Paly K 50.95	Scale RBF K 49.03 48.54 40.7 Scale RBF K 50.95	d and Balanced Bagging DT 43.68 42.13 35.86 d and Balanced Bagging DT 45.2	Data Bagging SVM 42.04 46.74 34.52 Data Bagging SVM 49.62	AdaBoost DT 48.93 46.81 38.28 AdaBoost DT 49.56	AdaBoost SVM 53.38 48.01 45.52 AdaBoost SVM 55.03		
Both Views Imaging View Genetic View Balanced Accuracy Both Views Imaging View	Linear K 46.08 48.78 44.72 Linear K 44.5 47.06	Poly K 47.25 48.36 43.87 Poly K 43.08 43.25	RBF K 47.25 48.36 43.87	Original Data Bagging DT 42.38 48.27 36.34 Original Data Bagging DT 36.1 41.55	Bagging SVM 40.56 47.33 43.71 Bagging SVM 38.11 46.86	AdaBoost DT 50.14 40.79 46.94 AdaBoost DT 43.42 34.14	AdaBoost SVM 46.2 50.06 43.69 AdaBoost SVM 47.34 53.33	Linear K 46.52 45.55 40.69 Linear K 51.4 51.45	Poly K 49.03 48.54 40.7 Poly K 50.95 50.95	Scale RBF K 49.03 48.54 40.7 Scale RBF K 50.95 50.95	d and Balanced Bagging DT 43.68 42.13 35.86 d and Balanced Bagging DT 45.2 45.2	Data Bagging SVM 42.04 46.74 34.52 Data Bagging SVM 49.62 52.71	AdaBoost DT 48.93 46.81 38.28 AdaBoost DT 49.56 49.17	AdaBoost SVM 53.38 48.01 45.52 AdaBoost SVM 55.03 48.01		

 $\Sigma\chi\eta\mu\alpha$ 6.39: For each model and classifier, the metric scores for OPNMF - DCCA transformed data classification are presented. Highlighted green are the best performing models, for each metric.

6.4 Application of MCA to the genetic data along with O-PNMF to the imaging data

Combining the two previous techniques, we now apply MCA to the genetic data, and OPNMF to the imaging data, and combine the two transformed views for the classification task.

6.4.1 Without scaling or balancing:



 Σ χήμα 6.40: Classification metric using both views (imaging and genetic) on the SVM kernels (Linear, Polynomial, RBF), using the MCA transformed genetic and OPNMF transformed imaging data.



 Σ χήμα 6.41: Classification metric using Bagging on the MCA and OPNMF transformed imaging and genetic data.



 $\Sigma\chi\eta\mu\alpha$ 6.42: Classification metric using AdaBoost on the MCA and OPNMF transformed imaging and genetic data.



6.4.2 With scaling and balancing:

 $\Sigma\chi\eta\mu\alpha$ 6.43: Classification metric using both views (imaging and genetic) on the SVM kernels (Linear, Polynomial, RBF), using the MCA transformed genetic and OPNMF transformed imaging data.



 $\Sigma\chi\eta\mu\alpha$ 6.44: Classification metric using Bagging on the MCA and OPNMF transformed imaging and genetic data.



 $\Sigma\chi\eta\mu\alpha$ 6.45: Classification metric using AdaBoost on the MCA and OPNMF transformed imaging and genetic data.

The following tables present the complete results for the MCA and OPNMF transformed data, for each model, for each metric, for each view, and either with or without scaling and balancing. With green are highlighted the best values for each metric, depending on whether scaling and balancing were applied:

					Ν	ICA	+ OPI	NMF						
Accuracy %				Original Data						Scale	d and Balanced	Data		
	Linear K	Poly K	RBF K	Bagging DT	Bagging SVM	AdaBoost DT	AdaBoost SVM	Linear K	Poly K	RBF K	Bagging DT	Bagging SVM	AdaBoost DT	AdaBoost SVM
Both Views	52.12	51.35	51.35	48.26	55.6	44.79	53.28	56.55	54.48	54.48	46.9	55.86	55.17	57.24
Imaging View														
Genetic View														
F1 Score				Original Data	-					Scale	ed and Balanced	Data		
	Linear K	Poly K	RBF K	Bagging DT	Bagging SVM	AdaBoost DT	AdaBoost SVM	Linear K	Poly K	RBF K	Bagging DT	Bagging SVM	AdaBoost DT	AdaBoost SVM
Both Views	E1 04			47.20	53.56									
	51.94	34.85	34.85	47.26	52.56	43.94	51.46	53.78	53.25	53.25	38.87	52.91	54.79	53.68
Imaging View	51.94	34.85	34.85	47.26	52.56	43.94	51.46	53.78	53.25	53.25	38.87	52.91	54.79	53.68
Imaging View Genetic View	51.94	34.85	34.85	47.26	52.56	43.94	51.46	53.78	53.25	53.25	38.87	52.91	54.79	53.68
Imaging View Genetic View	51.94	34.85	34.85	47.26	52.56	43.94	51.46	53.78	53.25	53.25	38.87	52.91	54.79	53.68
Imaging View Genetic View	51.54	34.85	34.85	47.26	52.50	43.94	51.46	53.78	53.25	53.25	38.87	52.91	54.79	53.68
Imaging View Genetic View	51.54	34.85	34.85	47.25		43.94	51.46	53.78	53.25	53.25	38.87	52.91	54.79	53.68
Imaging View Genetic View Balanced Accuracy	51.34	34.85	34.85	47.25		43.94	51.46	53.78	53.25	53.25	38.87	52.91	54.79	53.68
Imaging View Genetic View Balanced Accuracy	Linear K	34.85	34.85	0riginal Data Bagging DT	Bagging SVM	43.94	51.46	53.78 Linear K	53.25	53.25	38.87	52.91	54.79 AdaBoost DT	AdaBoost SVM
Imaging View Genetic View Balanced Accuracy Both Views	Linear K 49.03	34.85 Poly K 33.33	34.85	47.25 Original Data Bagging DT 42.35	Bagging SVM 45.5	43.94 AdaBoost DT 39.15	51.46 AdaBoost SVM 45.3	53.78 Linear K 58.18	53.25 Poly K 55.72	53.25 Scale RBF K 55.72	38.87 d and Balanced Bagging DT 49.58	52.91 Data Bagging SVM 57.86	54.79 AdaBoost DT 54.87	AdaBoost SVM 58.25
Imaging View Genetic View Balanced Accuracy Both Views Imaging View	Linear K 49.03	34.85 Poly K 33.33	34.85	47.26 Original Data Bagging DT 42.35	Bagging SVM 45.5	43.94 AdaBoost DT 39.15	51.46	53.78 Linear K 58.18	53.25 Poly K 55.72	53.25	38.87 d and Balanced Bagging DT 49.58	52.91 Data Bagging SVM 57.86	54.79 AdaBoost DT 54.87	53.68 AdaBoost SVM 58.25

 $\Sigma\chi\eta\mu\alpha$ 6.46: For each model and classifier, the metric scores for MCA and OPNMF transformed data classification are presented. Highlighted green are the best performing models, for each metric.

6.5 Application of FAMD to the dataset

Another method for dimensionality reduction is Factor Analysis of Mixed Data. FAMD combines both views, and creates a set of transformed components that contain information from both sets of variables.

6.5.1 Without scaling or balancing:



 $\Sigma\chi\eta\mu\alpha$ 6.47: Classification metric using both views (imaging and genetic) on the SVM kernels previously mentioned (Linear, Polynomial, RBF), using the FAMD transformed imaging and genetic data.



Σχήμα 6.48: Classification metric using Bagging on the FAMD transformed imaging and genetic data.



Σχήμα 6.49: Classification metric using AdaBoost on the FAMD transformed imaging and genetic data.



6.5.2 With scaling and balancing:

 $\Sigma\chi\eta\mu\alpha$ 6.50: Classification metric using both views (imaging and genetic) on the SVM kernels previously mentioned (Linear, Polynomial, RBF), using the FAMD transformed imaging and genetic data.



 Σ χήμα 6.51: Classification metric using Bagging on the FAMD transformed imaging and genetic data.



 Σ χήμα 6.52: Classification metric using AdaBoost on the FAMD transformed imaging and genetic data.

The following tables present the complete results for the FAMD transformed data, for each model, for each metric, for each view, and either with or without scaling and balancing. With green are highlighted the best values for each metric, depending on whether scaling and balancing were applied:

						E	AMD							
Accuracy %				Original Data						Seale	d and Palanco	Data		
Accuracy /6	linear K	Poly K	RRFK	Bagging DT	Bagging SVM	AdaBoost DT	AdaBoost SVM	linear K	Poly K	RRF K	Bagging DT	Bagging SVM	AdaBoost DT	AdaBoost SVM
Both Views	54.41	55.17	55.17	A9 A3	53.26	52.87	53.26	51 37	48.63	48.63	43.84	50.68	47.26	50.68
Imaging View	34.41	33.17	55.27	43.43	55.20	52.07	55.20	51.57	40.05	40.05	43.04	50.00	47.20	50.00
Genetic View														
F1 Score				Original Data						Scale	ed and Balanced	Data		
	Linear K	Poly K	RBF K	Bagging DT	Bagging SVM	AdaBoost DT	AdaBoost SVM	Linear K	Poly K	RBF K	Bagging DT	Bagging SVM	AdaBoost DT	AdaBoost SVM
Both Views	52.05	44.42	44.42	48.31	49.5	51.75	51.83	45.71	43.26	43.26	41.6	46.81	44.53	46.21
Imaging View														
Genetic View														
Balanced Accuracy				Original Data						Scale	ed and Balanced	Data		
	Linear K	Poly K	RBF K	Bagging DT	Bagging SVM	AdaBoost DT	AdaBoost SVM	Linear K	Poly K	RBF K	Bagging DT	Bagging SVM	AdaBoost DT	AdaBoost SVM
Both Views	45.82	50.87	50.87	43.52	42.33	46.57	45.98	53.88	51.24	51.24	45.41	52.66	49.73	53.77
Imaging View														
Genetic View														

 $\Sigma\chi\eta\mu\alpha$ 6.53: For each model and classifier, the metric scores for FAMD transformed data classification are presented. Highlighted green are the best performing models, for each metric.

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6.6 Results Summary

To summarize the results, we present a table with the best accuracy, F1 Score and balanced accuracy scores for each model, along with some notes. The diagram below visualizes the best results of each model, in order for them to be compared more easily.

Model	Best Accuracy (%)	Best F1 Score (%)	Best Balanced Accuracy (%)	Notes
RAW DATA	55.48	54.00	56.65	145 ROIs (Scaled) and 54 SNPs (Balanced). Both AdaBoost DT.
DCCA-150-3	51.72	50.80	45.82	Output Dimension 150, 3 Hidden Layers, no scaling or balancing. Imaging Linear.
OPNMF	58.62	54.02	61.01	30 Imaging Components (After OPNMF) Balanced only. Imaging Bagging SVM.
OPNMF + DCCA-150-3	53.79	53.38	55.03	30 Imaging Components (After OPNMF) and 54 SNPs, then DCCA, then scaled and balanced. Both AdaBoost SVM.
МСА	59.59	57.85	59.41	145 ROIs (Scaled), 10 Genetic components, Balanced only. Both Poly SVM.
MCA + DCCA-150-3	52.05	50.10	53.37	145 ROIs and 10 Genetic components (After MCA), then DCCA, then scaled and balanced. Imaging Bagging SVM.
MCA + OPNMF	57.24	53.68	53.77	30 Imaging Components (After OPNMF) and 10 Genetic components (After MCA), Balanced only. Both AdaBoost SVM.
FAMD	55.17	44.42	50.87	10 Components, no scaling, no balancing. Both Poly / RBF SVM.

 $\Sigma\chi\eta\mu\alpha$ 6.54: For each model, the best Accuracy, F1 Score, Balanced Accuracy achieved are shown, along with notes explaining the combination of data and parameters used. Highlighted in green are the best and second best (lighter green) results achieved among all models.



 Σ χήμα 6.55: For each model, the best Accuracy, F1 Score and Balanced Accuracy are plotted.

Κεφάλαιο 7

Discussion

Looking at the results, we can clearly see that the models that achieve superior results are the OPNMF and MCA models. The first model, employed the OPNMF technique in order to transform the 145 ROI values for the imaging view of the dataset into 30 imaging components. After that, without the genetic view, balancing on the imaging view was performed, and the classification was performed after training a Bagging ensemble classifier, using as a base model a linear SVM, achieving the highest balanced accuracy score, of 61.01%. The second model, utilized the MCA technique on the genetic view of the dataset, reducing the 54 (categorical) values of the SNPs into 10 genetic components (continuous). After that, along with the imaging view, balancing was performed, and the classification task was carried out using the model of SVM (single classifier). The polynomial kernel as well as the radial basis function were both used and both achieved identical results, which are the best in accuracy and F1 Score, 59.59% and 57.85% respectively.

Published studies attempt to either classify patients with AD or CN, AD or MCI, or predict the conversion rate of MCI to AD. This study attempts to classify simultaneously all three classes (AD, MCI, CN), but with the help of the confusion matrices, observations can be made for the performance of the models in the above tasks. These observations can not however lead to conclusions about the performance of the models trained in this thesis to the task of two class classification, since the models were trained for a three way classification. Looking at the highest accuracy model's (Bagging SVM ensemble classifier on OPNMF transformed imaging data) confusion matrix, we can deduce that the accuracy of the model at predicting AD vs CN is at 92.61%, which is in line with the results achieved by similar SVM approaches ([86], [87]). However, there is a margin to AD vs CN classification achieved scores compared to cutting-edge deep learning models, since they can achieve accuracy scores of 97% or more ([88], [89]). MCI vs CN classification based on the best models' confusion matrix achieved 65.80% accuracy, worse than the state of the art models published, that have an accuracy score of 86.57% [90]. Finally, AD vs MCI classification scores achieved by the best model was 66.88%, worse than the state of the art models, which achieved a score of 74.47% [91]. It has to be noted however, that the models trained in this thesis were not only significantly simpler than the state of the art ones, but also trained for the multi class classification task only.

It is immediately obvious that all models are achieving scores that are close to each other, with their best variations being around 50% and above. That however is not the

case for most variations of the models' parameters, as is easily observable through the figures of the fifth chapter (e.g. 6.11,6.12,6.24,6.25). Therefore, there is a clear benefit in using the imaging view, since many genetic-only models are quite far from the best models (achieving metric scores in the mid 30s). More importantly, no genetic-only model has been able to achieve better results than another model that included the imaging view. This is expected, since the genetic data do not contain any information about MCI patients, and more importantly, the existence of AD risk alleles are not guaranteed to be translated to having the disease. Furthermore, the information that is contained in the neuroimaging data is very important, and many times is a better indicator than the genetic data. Studies utilizing genetic only data have achieved prediction results that are worse than the results of models that used imaging data and/or genetic data ([92], [93], [94]). This is also evaluated by studies that have compared using the two views, and found that using only genetic data is relatively worse [95]

Nevertheless, the same cannot be said for the opposite; that is there is no clear benefit in including always using both the genetic and the imaging views, since models that have both views (imaging and genetic) are often outperformed by imaging-only models, yet they trade places depending on the model. For example, the two previously mentioned models (that achieve the best metric scores) are imaging-only (the case of the OPNMF model), and both views (the case of the MCA model).

There is no clear conclusion to be drawn about the benefit of the techniques of balancing and scaling. In the cases of (a) the data having no data analysis technique applied to them before classification, i.e. raw, (b) MCA-transformed genetic view models, (c) MCAtransformed genetic view and OPNMF-transformed imaging view models, balancing the dataset and applying scaling if needed seems to help, while in the case of the rest of the models it seems to worsen the results. This might be because information could be perhaps lost due to the method of standard scaling.

As for DCCA, it is clear that the technique achieves its target, which is to increase the linear correlation between the two views. However, one can clearly observe that it produces objectively worse results than equivalent no-DCCA methods. Even in the case of the original data, the DCCA transformed data produce results that are across the board worse in every metric. This is compounded by the fact that the DCCA networks not only took a considerable amount of time to train and optimize, but also the transformed outputs were clearly more computationally expensive to classify and therefore the fitting of the classifiers to the DCCA-transformed dataset took more time than that of their no-DCCA counterpart methods. The aforementioned facts can only lead us to conclude that the use of the method cannot be recommended for this type of problem, with the reservation that the model was not adequately trained or had not a nearly enough number of parameters. This however could very well go beyond the scope of this work, and is discussed in the next chapter. One detail that should be noted is that the DCCA method is not suitable for the great dissimilarity of the two views (the imaging being continuous and the genetic view being categorical). We attempted to remedy that dissimilarity with the MCA and OPNMF methods, which definitely improved the results of the models that utilized the DCCA method, however none of the resulting methods could achieve the results that the no-DCCA equivalent methods could achieve. While the model has the ability to non-linearly transform data in such a way that they are maximally correlated, this case is not a suitable case, as proven by the results. One reason might be that neural networks often require a much greater volume of data, and another might be that the two views are different in nature.

An interesting result is that of FAMD, in which both views are transformed into a reduced number of components. This method, as previously mentioned, is similar to performing PCA on the continuous imaging view while performing MCA on the categorical genetic view, but combined. In our case, we set the number of components to 10 due to long computational time, which proved to yield worse results than many of the other methods. It is entirely possible that the number of chosen components might not be enough to capture all of the information that the two views provide, and only achieve a substantial dimensionality reduction.

As can be seen by the extensive figures of the previous chapters, or more conveniently from the table's 6.54 notes, the ensemble classifiers were in general more successful in predicting the class. This aligns well with the empirical knowledge that ensemble methods of even simple (as decision trees and linear SVMs are) can improve upon the single method performance, even if the single method is more complex. The performance of the ensemble classifiers achieved not only better best-case results, but on average was better overall, with even the worst performing parameter combinations beating out the worst performing parameter combinations of the single classifier models of SVM. This was intuitively hypothesized, emprically proven, as well as expected as it has been shown in other studies. [96]

Moreover, comparing the different ensemble methods, it is not clear as to which method has the clear edge. The two methods have comparable results, with Bagging seemingly being in more models marginally better than AdaBoost. Furthermore, comparing the two methods for the base model classifier, which are the decision tree model and the linear kernel SVM, we can conclude that in nearly every case, the linear SVM base model is better, for both kinds of ensembles. While the difference between the performances of the two base models is not substantial, there is a clear trend. It is worth noting that both of those base models were chosen for their relatively simplistic nature, because of computational time limitations and as to create a basis for comparing single classifier models to ensemble classifiers.

Overall, ensemble methods result in higher classification performance compared to single classifier models. Concerning the single classifier model of the different SVM kernels, we can see that in most cases the polynomial and RBF kernels outperform the linear kernel. Additionally, the polynomial and RBF kernels seem to outperform in the best case the linear kernel, however the linear kernel seems to be more robust. That can be seen in many confusion matrices, where the polynomial and RBF kernels classify the dataset with very poor performance. There is however, a very interesting note to be added, concerning the models that had DCCA applied to the dataset before the classification task. In those cases, the linear kernel is better than the polynomial and RBF kernels. This might indicate that indeed the DCCA method achieves the goal of linearly correlating the two views, however that might be at the expense of information loss, as those models are beaten by simpler methods without having DCCA applied to the dataset. This alone might be indicative of the need for not only more computational time and power devoted to the training of the DCCA parallel neural networks, but also the need for bigger and more complex variations of the network explored. In agreement with the original paper, after we applied DCCA, the transformed views were more correlated than the original ones, so we succeeded at the goal of the DCCA method.

Another important aspect that can be observed is the success of the models in the classification of a specific class. We can see that most models (except from the models with distinctively poor overall performance) could reliably classify the patients that were suffering from AD from the CN people, while struggling to accurately decide for the case of the MCI class. The confusion matrices in the figures of the previous chapter highlight exactly that, with most models having a high percentage of the upper left portion of their confusion matrix well-defined, while the outer right and bottom part (which is the part of the MCI class) being quite confounded. As an example, one can observe the confusion matrices for the OPNMF method (no-DCCA), with the classifier being an ensemble classifier especially on using both or only the imaging view (6.36 Bagging, 6.37 Adaboost), but also the confusion matrices of the MCA method on both views, with scaling or balancing, using a single classifier model (6.21 SVMs). This is indicative of the strong performance of the model in AD vs CN classification, as well as the difficulty of the problem to distinguish CN vs MCI and MCI vs AD, as the MCI class is apparently blurring the lines of the other classes. Since the methods and the models were optimized and trained for the problem of multi-class classification on the classes of CN, MCI and AD aforementioned should not be taken as a fact, but rather as an indication of performance.

The utilization of the grid search and cross validation methods for the evaluation of the different model parameters' performance while producing extensive and very useful results, was limiting the number of models that could possibly be explored. Both the optimizations of the DCCA networks, as well as the classification methods training, took for every model time in the order of hours. As a result, only relatively simple methods such as Support Vector Machines and Decision Trees were selected to be explored, since training multilayer perceptrons and similar methods for each classification task and then optimizing them not only with every model and every view, but also with their respective grids of parameters was out of the question. Another limiting factor was the insufficient amount of data points that are inherently available to studies like these, which is due to the nature of the problem, since biomedical data and especially neuroimaging data are not only hard to collect, but pose storing, processing and visualization problems as well. Finally, the time frame for this study was not unbounded, and thus there had to be a selective process as to the direction of experimentation.

Finally, as to what the future directions might be for further research on the lessons learned from this work, there are many possible steps. One might be to explore, as mentioned previously, different optimization algorithms, activation functions and generally different architectures for the parallel neural networks of the DCCA method. Another direction might be data augmentation, or creation of synthetic data, as to enrich the dataset and unlock the potential of the DCCA method. Concerning the data analysis techniques, the OPNMF method is one that could benefit from experimenting with different number of components, something that wasn't done on this study due to computational power limitations. Furthermore, more complex classifiers can be used, such as MLPs and K-Nearest Neighbors classifiers. That can also be extended to the ensemble methods, not only for the base classifier models, but also for the ensemble methods, with other boosting methods and methods such as stacking being obvious candidates. Finally, to address the issue of the different in nature and type views, one could explore the use of deep autoencoders in order to alleviate the problem of handling categorical data.

Κεφάλαιο 8

Conclusions

This study was intended to be an extensive comparison of the applications of data analysis methods, as well as machine and deep learning methods, applied to the problem of CN / MCI / AD classification. From the work that was performed, there are some clear messages. First of all, data analysis techniques such as MCA and OPNMF are better compared to the MUSE ROIs because the OPNMF components are data driven while MUSE ROIs are derived from templates [42]. MCA is also better than the vanilla genetic view, because it transforms the genetic data into the type of the imaging data, making both views have the same kind. Furthermore, the method of Deep Canonical Correlation Analysis as stated in the original paper [43], is not beneficial to this problem, at least not without further tuning, however it increases the correlation between the views in agreement with the original paper. Ensemble classifier methods are superior to the simplistic single classifier methods such as Support Vector Machines. Finally, it is clear that using only genetic data is not sufficient to yield higher-quality results, as they only show predesposition for the disease. The combinations that achieved the best results used imaging or imaging along with genetic data, either OPNMF-transformed imaging data or MCA-transformed genetic data, accompanied with either Bagging ensembles of SVMs or a polynomial kernel SVM as classifiers.

- [1] 2022 Alzheimer's disease facts and figures. Alzheimers. Dement., 18(4):700-789, 2022.
- [2] Jeffrey Cummings, Carl Reiber Xai Parvesh Kumar. The price of progress: Funding and financing Alzheimer's disease drug development. Alzheimers Dement. (N. Y.), 4:330-343, 2018.
- [3] Sergio Grueso xa Raquel Viejo-Sobera. Machine learning methods for predicting progression from mild cognitive impairment to Alzheimer's disease dementia: a systematic review. Alzheimers. Res. Ther., 13(1):162, 2021.
- [4] Sayantan Kumar, Inez Oh, Suzanne Schindler, Albert M Lai, Philip R O Payne xau Aditi Gupta. Machine learning for modeling the progression of Alzheimer disease dementia using clinical data: a systematic literature review. JAMIA Open, 4(3):00ab052, 2021.
- [5] Taeho Jo, Kwangsik Nho xa Andrew J Saykin. Deep learning in Alzheimer's disease: Diagnostic classification and prognostic prediction using neuroimaging data. Front. Aging Neurosci., 11:220, 2019.
- [6] Ziyi Li, Xiaoqian Jiang, Yizhuo Wang xai Yejin Kim. Applied machine learning in Alzheimer's disease research: omics, imaging, and clinical data. Emerg. Top. Life Sci., 5(6):765–777, 2021.
- [7] Taeho Jo, Kwangsik Nho xa Andrew J Saykin. Deep learning in Alzheimer's disease: Diagnostic classification and prognostic prediction using neuroimaging data. Front. Aging Neurosci., 11:220, 2019.
- [8] Xiaofeng Zhu, Heung Il Suk xai Dinggang Shen. Low-rank dimensionality reduction for multi-modality neurodegenerative disease identification. World Wide Web, 22(2):907– 925, 2019.
- [9] G E Hinton xai R R Salakhutdinov. Reducing the dimensionality of data with neural networks. Science, 313(5786):504-507, 2006.
- [10] Andreas S Panayides, Amir Amini, Nenad D Filipovic, Ashish Sharma, Sotirios A Tsaftaris, Alistair Young, David Foran, Nhan Do, Spyretta Golemati, Tahsin Kurc, Kun Huang, Konstantina S Nikita, Ben P Veasey, Michalis Zervakis, Joel H Saltz xau Constantinos S Pattichis. AI in medical imaging informatics: Current challenges and future directions. IEEE J. Biomed. Health Inform., 24(7):1837–1857, 2020.

- [11] Katja Franke, Gabriel Ziegler, Stefan Klöppel, Christian Gaser και Alzheimer's Disease Neuroimaging Initiative. Estimating the age of healthy subjects from T1-weighted MRI scans using kernel methods: exploring the influence of various parameters. Neuroimage, 50(3):883–892, 2010.
- [12] 2022.
- [13] R Peters. Ageing and the brain. Postgrad. Med. J., 82(964):84-88, 2006.
- [14] Paul Denver xai Paula L McClean. Distinguishing normal brain aging from the development of Alzheimer's disease: inflammation, insulin signaling and cognition. Neural Regen. Res., 13(10):1719–1730, 2018.
- [15] Alzheimer's Disease and Dementia. https://www.alz.org/alzheimers-dementia/ what-is-dementia/related_conditions/mild-cognitive-impairment. Πρόσβαση: 2022-6-22.
- [16] Alzheimer's Disease and Dementia. https://www.alz.org/alzheimers-dementia/ what-is-alzheimers. Πρόσβαση: 2022-6-22.
- [17] Alzheimer's disease. https://www.nhs.uk/conditions/alzheimers-disease/. Πρόσβαση: 2022-6-22.
- [18] Francesca Fernandez και Jessica L Andrews. Genetics of dementia. Diagnosis and Management in Dementia, σελίδες 127–146. Elsevier, 2020.
- [19] Mild cognitive impairment Symptoms and causes. https://www.mayoclinic. org/diseases-conditions/mild-cognitive-impairment/symptoms-causes/syc-20354578., 2020. Πρόσβαση: 2022-6-22.
- [20] Ronald C Petersen, Glenn E Smith, Stephen C Waring, Robert J Ivnik, Eric G Tangalos xai Emre Kokmen. Mild cognitive impairment. Arch. Neurol., 56(3):303, 1999.
- [21] Alistair Burns και Steve Iliffe. Alzheimer's disease. BMJ, 338(φεβ05 1):β158, 2009.
- [22] Zeinab Breijyeh xa Rafik Karaman. Comprehensive review on Alzheimer's disease: Causes and treatment. Molecules, 25(24):5789, 2020.
- [23] R Brown. Magnetic Resonance Imaging : Physical Principles and Sequence Design. Wiley.
- [24] C Bohm, T Greitz, R Seitz xai L Eriksson. Specification and selection of regions of interest (ROIs) in a computerized brain atlas. J. Cereb. Blood Flow Metab., 11(2):A64– 8, 1991.
- [25] Tianming Liu. A few thoughts on brain ROIs. Brain Imaging Behav., 5(3):189–202, 2011.

- [26] Jimit Doshi, Guray Erus, Yangming Ou, Susan M Resnick, Ruben C Gur, Raquel E Gur, Theodore D Satterthwaite, Susan Furth, Christos Davatzikos xa. Alzheimer's Neuroimaging Initiative. MUSE: MUlti-atlas region Segmentation utilizing Ensembles of registration algorithms and parameters, and locally optimal atlas selection. Neuroimage, 127:186–195, 2016.
- [27] single nucleotide polymorphism / SNP. https://www.nature.com/scitable/ definition/snp-295/. Πρόσβαση: 2022-6-22.
- [28] Mohammed Amir Husain, Benoit Laurent xai Mélanie Plourde. APOE and Alzheimer's disease: From lipid transport to physiopathology and therapeutics. Front. Neurosci., 15:630502, 2021.
- [29] Rudolph E Tanzi. The genetics of Alzheimer disease. Cold Spring Harb. Perspect. Med., 2(10):α006296-α006296, 2012.
- [30] Fuhai Song, Guangchun Han, Zhouxian Bai, Xing Peng, Jiajia Wang xat Hongxing Lei. Alzheimer's disease: Genomics and beyond. Int. Rev. Neurobiol., 121:1–24, 2015.
- [31] Wolfgang Karl Härdle και Léopold Simar. Applied Multivariate Statistical Analysis. σελίδες 443-448, 2015.
- [32] Galen Andrew, Raman Arora, Jeff Bilmes και Karen Livescu. Deep Canonical Correlation Analysis. Proceedings of the 30th International Conference on Machine LearningSanjoy Dasgupta και David McAllester, επιμελητές, τόμος 28 στο Proceedings of Machine Learning Research, σελίδες 1247–1255, Atlanta, Georgia, USA, 2013. PMLR.
- [33] Shotaro Akaho. A kernel method for canonical correlation analysis. 2006.
- [34] Zhirong Yang xau Erkki Oja. Linear and nonlinear projective nonnegative matrix factorization. IEEE Trans. Neural Netw., 21(5):734–749, 2010.
- [35] Aristeidis Sotiras, Susan M Resnick xai Christos Davatzikos. Finding imaging patterns of structural covariance via Non-Negative Matrix Factorization. Neuroimage, 108:1– 16, 2015.
- [36] Sirine Sayadi, Estelle Geffard, Mario Südholt, Nicolas Vince και Pierre Antoine Gourraud. Secure distribution of factor analysis of mixed data (FAMD) and its application to personalized medicine of transplanted patients. Advanced Information Networking and Applications, Lecture notes in networks and systems, σελίδες 507–518. Springer International Publishing, Cham, 2021.
- [37] Jérôme Pagès. Chapter 3, Factorial Analysis of Mixed Data." Multiple Factor Analysis by Example Using R. The R Series, 1:67–72, 2015.
- [38] Sandeep Kumar Satapathy, Satchidananda Dehuri, Alok Kumar Jagadev και Shruti Mishra. Introduction. EEG Brain Signal Classification for Epileptic Seizure Disorder Detection, σελίδες 1–25. Elsevier, 2019.

- [39] Robi Polikar. Ensemble learning. Scholarpedia J., 4(1):2776, 2009.
- [40] B Efron. Bootstrap methods: Another look at the jackknife. Ann. Stat., 7(1):1–26, 1979.
- [41] R. Polikar. Ensemble learning. Scholarpedia, 4(1):2776, 2009. ρεισιον 186077.
- [42] Mca 1.0.3. https://pypi.org/project/mca/. Πρόσβαση: 2022-6-22.
- [43] G Andrew, R Arora, J Bilmes Xai K Livescu. Deep Canonical Correlation Analysis. JMLR: W&CP, 28, 2013.
- [44] L Svennerholm, K Boström xai B Jungbjer. Changes in weight and compositions of major membrane components of human brain during the span of adult human life of Swedes. Acta Neuropathol., 94(4):345–352, 1997.
- [45] Jessica S Damoiseaux. Effects of aging on functional and structural brain connectivity. Neuroimage, 160:32–40, 2017.
- [46] David S Knopman, Helene Amieva, Ronald C Petersen, Gäel Chételat, David M Holtzman, Bradley T Hyman, Ralph A Nixon xat David T Jones. Alzheimer disease. Nat. Rev. Dis. Primers, 7(1):33, 2021.
- [47] Guy G Potter xai David C Steffens. Contribution of depression to cognitive impairment and dementia in older adults. Neurologist, 13(3):105–117, 2007.
- [48] Panagiotis Katsonis, Amanda Koire, Stephen Joseph Wilson, Teng Kuei Hsu, Rhonald C Lua, Angela Dawn Wilkins xa Olivier Lichtarge. Single nucleotide variations: biological impact and theoretical interpretation. Protein Sci., 23(12):1650–1666, 2014.
- [49] C Jordan. Essai sur la géométrie à n n dimensions. Bull. Soc. Math. France, 3, 1875.
- [50] H Hotelling. Relations Between Two Sets of Variates. Biometrika, 28(3-4):321–377, 1936.
- [51] Shotaro Akaho. A kernel method for canonical correlation analysis. 2006.
- [52] R Arora xai K Livescu. Kernel CCA for multi-view learn- ing of acoustic features using articulatory measurements. Symp. on Machine Learning in Speech and Language Processing. 2012.
- [53] Hao Jun Michael Shi, Melody Qiming Xuan, Figen Oztoprak xai Jorge Nocedal. On the numerical performance of derivative-free optimization methods based on finitedifference approximations. 2021.
- [54] Pascal Vincent, Hugo Larochelle, Yoshua Bengio xa Pierre Antoine Manzagol. Extracting and composing robust features with denoising autoencoders. Proceedings of the 25th international conference on Machine learning - ICML '08, New York, New York, USA, 2008. ACM Press.

- [55] Hervé Abdi xa Dominique Valentin. Multiple Correspondence Analysis. Encyclopedia of Measurement and Statistics, 2007.
- [56] Brigitte Le Roux xa Henry Rouanet. *Multiple correspondence analysis*. Sage Publications, 2010.
- [57] Pentti Paatero xai Unto Tapper. Positive matrix factorization: A non-negative factor model with optimal utilization of error estimates of data values. Environmetrics, 5(2):111–126, 1994.
- [58] D D Lee xai H S Seung. Learning the parts of objects by non-negative matrix factorization. Nature, 401(6755):788–791, 1999.
- [59] A Cichocki, R Zdunek, A H Phan Xai S Amari. Non-negative Matrix and Tensor Factorizations: Applications to Exploratory Multi-Way Data Analysis. Wiley, New York, 2009.
- [60] Stefanos Zafeiriou, Anastasios Tefas, Ioan Buciu και Ioannis Pitas. Exploiting discriminant information in nonnegative matrix factorization with application to frontal face verification. IEEE Trans. Neural Netw., 17(3):683–695, 2006.
- [61] A Shashua και T Hazan. Non-negative tensor factorization with applications to statistics and computer vision; International Conference on Machine Learning. σελίδες 792–799, 2005.
- [62] Corinna Cortes xat Vladimir Vapnik. Support-vector networks. Mach. Learn., 20(3):273–297, 1995.
- [63] Raoof Gholami και Nikoo Fakhari. Support vector machine: Principles, parameters, and applications. Handbook of Neural Computation, σελίδες 515–535. Elsevier, 2017.
- [64] Farhad Maleki, Katie Ovens, Keyhan Najafian, Behzad Forghani, Caroline Reinhold και Reza Forghani. Overview of machine learning part 1: Fundamentals and classic approaches. Neuroimaging Clin. N. Am., 30(4):ε17–ε32, 2020.
- [65] 1.4. Support vector machines. https://scikit-learn.org/stable/modules/svm. html. Πρόσβαση: 2022-6-22.
- [66] Support Vector Machine, 2022.
- [67] Derek A Pisner και David M Schnyer. Support vector machine. Machine Learning, σελίδες 101–121. Elsevier, 2020.
- [68] Thomas G Dietterich. Ensemble Methods in Machine Learning. Multiple Classifier Systems, Lecture notes in computer science, σελίδες 1–15. Springer Berlin Heidelberg, Berlin, Heidelberg, 2000.
- [69] Leo Breiman. Bagging predictors. Mach. Learn., 24(2):123–140, 1996.

- [70] K. Zarkogianni, M. Athanasiou, A. C. Thanopoulou και K. S. Nikita. Comparison of machine learning approaches towards assessing the risk of developing Cardiovascular disease as a long-term diabetes complication. IEEE Journal of Biomedical and Health Informatics, ΠΠ:1–1, 2018.
- [71] Thomas G. Dietterich xai Ghulum Bakiri. Solving Multiclass Learning Problems via Error-Correcting Output Codes. CoRR, 55. AI/9501101, 1995.
- [72] Tin Kam Ho. The random subspace method for constructing decision forests. IEEE Transactions on Pattern Analysis and Machine Intelligence, 20(8):832–844, 1998.
- [73] ADNI. https://adni.loni.usc.edu/study-design/. Πρόσβαση: 2022-6-22.
- [74] ADNI. https://adni.loni.usc.edu/methods/mri-tool/mri-analysis/. Πρόσβαση: 2022-6-22.
- [75] C Davatzikos, A Genc, D Xu xai S M Resnick. Voxel-based morphometry using the RAVENS maps: methods and validation using simulated longitudinal atrophy. Neuroimage, 14(6):1361–1369, 2001.
- [76] Xiaohui Yao, Shannon L Risacher, Kwangsik Nho, Andrew J Saykin, Ze Wang, Li Shen xa. Alzheimer's Disease Neuroimaging Initiative. Targeted genetic analysis of cerebral blood flow imaging phenotypes implicates the INPP5D gene. Neurobiol. Aging, 81:213-221, 2019.
- [77] Iris E et al Jansen. Genome-wide meta-analysis identifies new loci and functional pathways influencing Alzheimer's disease risk. Nat. Genet., 51(3):404–413, 2019.
- [78] Brian W et al Kunkle. Genetic meta-analysis of diagnosed Alzheimer's disease identifies new risk loci and implicates Aβ, tau, immunity and lipid processing. Nat. Genet., 51(3):414–430, 2019.
- [79] J C et al Lambert. Meta-analysis of 74,046 individuals identifies 11 new susceptibility loci for Alzheimer's disease. Nat. Genet., 45(12):1452–1458, 2013.
- [80] https://github.com/Michaelvll/DeepCCA. Πρόσβαση: 2022-6-22.
- [81] PyTorch. https://pytorch.org/features/. Πρόσβαση: 2022-6-22.
- [82] Aristeidis Sotiras, Susan M. Resnick xai Christos Davatzikos. Finding imaging patterns of structural covariance via Non-Negative Matrix Factorization. NeuroImage, 108:1– 16, 2015.
- [83] Prince. https://pypi.org/project/prince/. Πρόσβαση: 2022-6-22.
- [84] Fabian Pedregosa, Gaël Varoquaux, Alexandre Gramfort, Vincent Michel, Bertrand Thirion, Olivier Grisel, Mathieu Blondel, Andreas Müller, Joel Nothman, Gilles Louppe, Peter Prettenhofer, Ron Weiss, Vincent Dubourg, Jake Vanderplas, Alexandre Passos, David Cournapeau, Matthieu Brucher, Matthieu Perrot xaı Édouard Duchesnay. Scikit-learn: Machine Learning in Python. 2012.

- [85] Taesic Lee xai Hyunju Lee. Prediction of Alzheimer's disease using blood gene expression data. Sci. Rep., 10(1):3485, 2020.
- [86] Qi Li και Mary Qu Yang. Comparison of machine learning approaches for enhancing Alzheimer's disease classification. PeerJ, 9:ε10549, 2021.
- [87] Morshedul Bari Antor, A. H. M. Shafayet Jamil, Maliha Mamtaz, Mohammad Monirujjaman Khan, Sultan Aljahdali, Manjit Kaur, Parminder Singh xa Mehedi Masud. A Comparative Analysis of Machine Learning Algorithms to Predict Alzheimer's Disease. Journal of Healthcare Engineering, 2021:1–12, 2021.
- [88] Saman Sarraf, Danielle D. DeSouza, John Anderson xat Ghassem Tofighi. DeepAD: Alzheimer's Disease Classification via Deep Convolutional Neural Networks using MRI and fMRI. 2016.
- [89] Ehsan Hosseini-Asl, Georgy Gimel'farb xai Ayman El-Baz. Alzheimer's Disease Diagnostics by a Deeply Supervised Adaptable 3D Convolutional Network, 2016.
- [90] Baiying Lei, Siping Chen, Dong Ni xai Tianfu Wang. Discriminative Learning for Alzheimer's Disease Diagnosis via Canonical Correlation Analysis and Multimodal Fusion. Frontiers in Aging Neuroscience, 8, 2016.
- [91] Classification of Alzheimer's Disease, Mild Cognitive Impairment, and Cognitively Unimpaired Individuals Using Multi-feature Kernel Discriminant Dictionary Learning. Frontiers in Computational Neuroscience, 11, 2018.
- [92] Taeho Jo, Kwangsik Nho xai Andrew J Saykin. Deep learning in Alzheimer's disease: Diagnostic classification and prognostic prediction using neuroimaging data. Front. Aging Neurosci., 11:220, 2019.
- [93] Eugene Lin, Chieh Hsin Lin xa Hsien Yuan Lane. Deep learning with neuroimaging and genomics in Alzheimer's disease. Int. J. Mol. Sci., 22(15):7911, 2021.
- [94] Kaida Ning, Bo Chen, Fengzhu Sun, Zachary Hobel, Lu Zhao, Will Matloff xai Arthur W Toga. Classifying Alzheimer's disease with brain imaging and genetic data using a neural network framework. Neurobiol. Aging, 68:151–158, 2018.
- [95] J Vijaya xai E Sivasankar. Computing efficient features using rough set theory combined with ensemble classification techniques to improve the customer churn prediction in telecommunication sector. Computing, 100(8):839–860, 2018.
- [96] Modeling of Artificial Intelligence, 11(3), 2016.