# Leveraging multimodal data to predict outcomes of antidepressant treatment

Preliminary results

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#### 1 Data

The data (n=98) we use has been provided by Emily Beaman. She processed relevant data from the CIMBI database. In addition we compute:

- a "MR" biomarker as the average of the thickness of the left lateral, right lateral, left medial, and right medial orbitol frontal cortex.
- a "PET" biomarker by combining the log binding potential from neocortex, hippocampus, caudate, and putamen via a latent variable model (adjusted for age, gender, and injected mass). This was done in a leave-one-out fashion i.e. the biomarker for individual *i* was obtained by fitting the model on all but the i-th individual and estimating the latent variable for the *i*-th individual based on its log-binding in the 4 regions.
- a "cognition" biomarker was obtained via a k-means algorithm on various cognitive outcomes (no leave-one-out here).
- the "outcomes" by computing the relative change in HAMD6 between week 8 (or 12) and baseline.

Individuals who had missing HAMD6 at both week 8 and 12 were excluded (n=13)

The corresponding R code is in the file 0-data-management. R available on Github.

# 2 Data analysis

After discussion with neuroscientist, we have identified 10 candidate biomarkers<sup>1</sup> for predicting recovery after SSRI treatment:

- MR\_OFCthick: thickness of the OFC brain region measured with MR.
- HAMD17: depression score at baseline.
- low\_hsCRP: high sensitivity CRP (I levels of inflammation in the body).
- lvpet: summary of the brain log-PET binding.
- cognitive cluster: summary of the cognition
- EEG vigilance: EEG signal (vigilance slope B1 bl)
- CATS\_scoretotal: ??
- CAR\_AUCi: Difference between the cortisol value and the cortisol value at wakeup cumulated over an hour.
- neuroticism:

Missing values: to simplify the analysis, we will assume that missing data occured completely at random. In particular, that it is not related to the outcome (patient did not leave the study because they fully recover or they were so seriously depressed that they could not stay in the study).

Association between recovery and biomarkers<sup>2</sup>: to assess whether the biomarker were associated with recovery we fitted a logistic regression with gender and age vs. a logistic regression with gender and age plus all biomarkers (as additive effects). A likelihood ratio test was used to compare the two models, i.e. assess the association over all biomarkers. Wald tests were used to test the association for each biomarker.

This was performed 4 times: using week 8 or week 12 as the outcome, using complete case analysis (excluding CATS, CAR, and neuroticism as biomarkers) or using multiple imputation (MI) to handle missing values. MI was performed using linear regression for continuous variables, logistic regression for binary variables, and a a proportional odds model for categorical variables based on all variables (outcome, age, gender, biomarkers) as suggested in Moons et al. (2006).

 $<sup>^{1}\</sup>mathrm{fMRI}$  is missing in the list

<sup>&</sup>lt;sup>2</sup>how does the recovery vary in average (i.e. at a population level) as a function of the biomarkers

Predictive value of the biomarkers<sup>3</sup>: to assess whether the biomarkers can be used to discriminate between patients who will recover and patients who won't, we tested whether the Area Under the Curve (AUC) of a logistic model with age, gender, and the biomarkers was greated than 0.5, and whether it was greater than a model with only age and gender. We also compared the performance with a random forest (default hyperparameters: mtry = 3, 500 trees, node size 1). The AUC was estimated via cross-validation (100 repetitions, test set of approximately 10%). An AUC was computed for each fold and then averaged acrossed folds (LeDell et al., 2015). The p-value was computed via a permutation test (2500 repetitions).

Calibration plot were also computed using the local polynomial regression fitting (loess).

Missing value was handled either using complete case analysis (excluding CATS, CAR, and neuroticism as biomarkers) or predicting the recovery probability for a given subject based on a logistic model containing all biomarkers for which the subject has available data.

The corresponding R code is in the file 1-prediction. R available on Github.

<sup>&</sup>lt;sup>3</sup>are the biomarkers useful to predict recovery for an individual

## 3 Results

#### 3.1 Descriptive statistics

The dataset contained 85 patients, 84 with the outcome at week 8 and 81 with the outcome at week 12. Some summary statistics are displayed below:

```
MR_OFCthick
                                                    HAMD17
                                                                    hsCRP
                  age
    sex
                    :18.24
male
     :25
             Min.
                                      :2.318
                                               Min.
                                                       :18.00
                                                                 Length:85
                              Min.
female:60
             1st Qu.:22.22
                              1st Qu.:2.510
                                               1st Qu.:20.00
                                                                 Class : character
             Median :23.93
                              Median :2.558
                                               Median :22.00
                                                                       :character
                                                                 Mode
             Mean
                    :27.22
                              Mean
                                      :2.573
                                               Mean
                                                       :22.76
             3rd Qu.:28.70
                              3rd Qu.:2.636
                                               3rd Qu.:25.00
             Max.
                     :57.31
                              Max.
                                      :2.889
                                               Max.
                                                       :31.00
                    cognitive_cluster EEG_vigilance
    lvpet
                                                              CATS_scoretotal
       :-0.82582
                            :1.00
                                                :-1.500000
Min.
                    Min.
                                        Min.
                                                              Min.
                                                                     : 0.00
                                                              1st Qu.:16.25
1st Qu.:-0.49196
                    1st Qu.:1.00
                                        1st Qu.: 0.000000
Median : -0.42337
                    Median:2.00
                                        Median : 0.000000
                                                             Median :23.50
Mean
       :-0.43227
                    Mean
                            :1.88
                                        Mean
                                                : 0.006173
                                                             Mean
                                                                     :30.42
3rd Qu.:-0.34832
                    3rd Qu.:3.00
                                        3rd Qu.: 0.000000
                                                              3rd Qu.:41.50
        :-0.09773
                                                : 1.500000
Max.
                            :3.00
                                        Max.
                                                             Max.
                                                                     :81.00
                    Max.
NA's
       :2
                    NA's
                            :2
                                        NA's
                                                :4
                                                              NA's
                                                                     :11
   CAR_AUCi
                     neuroticism
                                       Y_w8
                                                       Y_w12
Min.
       :-1070.30
                            : 67
                                   Mode :logical
                                                     Mode :logical
                    Min.
1st Qu.:
            76.69
                    1st Qu.:109
                                   FALSE:37
                                                     FALSE:22
Median:
          208.01
                    Median:119
                                   TRUE : 47
                                                     TRUE :59
Mean
       :
           180.26
                    Mean
                            :120
                                   NA's :1
                                                     NA's:4
3rd Qu.:
           382.60
                    3rd Qu.:133
Max.
        :
          768.90
                    Max.
                            :155
NA's
        :17
                    NA's
                            :24
```

The dataset contained many missing values. The pattern of the missing values is summarized on figure Figure 1. 48 patients had full data and the rest of the patients had between 1 and 4 missing data (number of red boxes per line). CATS, CAR, and neuroticm had a large number of missing data (11, 17, and 24) and this is why they were excluded from some analyses.

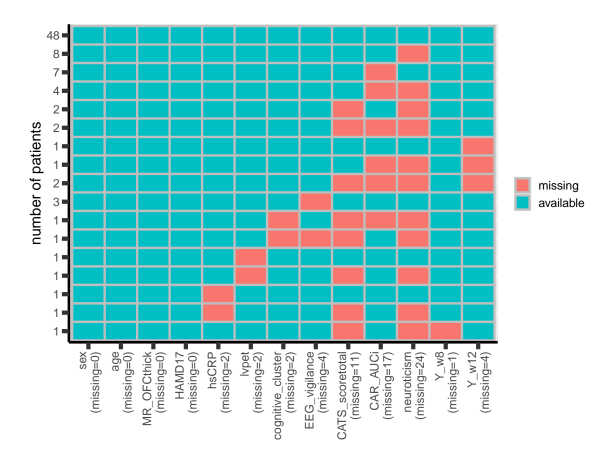


Figure 1: Missing data patterns

## 3.2 Association study (week 8)

Complete case: excluding CATS, CAR, and neuroticism, we fitted two logistic regressions (one with and one without the biomarkers) on the 75 patients with complete data. This likelihood ratio test showed evidence for an association between biomarkers and recovery:

```
Analysis of Deviance Table
```

Looking at the biomarker specific effects, high vigilance appeared to be associated with poor recovery: odd ratio 0.179 (unit?) adjusted p-value of 0.047. There was also a similar trend for cognitive cluster 3: odd ratio 0.15, adjusted p-value of 0.07.

```
p.value adjusted p-value
                      estimate
                                std.error
                                              odd ratio
(Intercept)
                    9.15328187 8.39872601 9.445388e+03 0.275782879
                                                                                   NA
female
                   -0.35932885 0.64097717 6.981447e-01 0.575073390
                                                                                   NA
                    0.05471804 0.04216889 1.056243e+00 0.194427297
                                                                                   NA
age
MR_OFCthick
                   -5.84569691 3.18471343 2.892318e-03 0.066424416
                                                                           0.23752311
HAMD17
                    0.16179531 0.09514669 1.175620e+00 0.089040138
                                                                           0.24208663
low_hsCRP
                    1.46954263 0.72733733 4.347246e+00 0.043337596
                                                                           0.19466776
                   -2.04888723 2.23330861 1.288782e-01 0.358921367
lvpet
                                                                           0.41535424
cognitive_cluster2 -0.81977323 0.69278301 4.405315e-01 0.236688505
                                                                           0.41535424
cognitive_cluster3 -1.89634154 0.75932143 1.501168e-01 0.012510207
                                                                           0.06993304
EEG_vigilance
                   -1.72062639 0.63876369 1.789540e-01 0.007066751
                                                                           0.04675833
```

Multiple imputation: as a sensitivity analysis, we now used all patients and all biomarkers and use multiple imputations (100 datasets) to handle missing value. Results are rather similar to the complete case analysis, with a slight tendency for stronger effects.

```
summary(pool(e.glm_impw8))[,c(1,2,3,5:6)]
```

```
estimate
                                       std.error
                                                        df
                                                               p.value
                 term
1
          (Intercept)
                       4.1516290163 8.6252902526 68.91234 0.631805406
2
            sexfemale -0.5825951700 0.6420666199 68.66383 0.367382059
3
                       0.0754554026 0.0456848728 68.79669 0.103163672
4
          MR_OFCthick -5.9114103612 3.0378756765 68.80810 0.055751906
5
               HAMD17
                       0.1808561403 0.0925544076 68.24495 0.054794466
6
             hsCRPlow
                       1.6868813295 0.7699177493 67.17518 0.031925680
7
                lvpet -2.6139562643 2.3770155497 68.85573 0.275301586
8
  cognitive_cluster2 -1.1511850446 0.7226909239 68.23190 0.115801761
   cognitive_cluster3 -2.7674590923 0.8726252184 68.19986 0.002273849
9
10
        EEG_vigilance -1.9950780957 0.6945417890 68.76199 0.005411043
11
      CATS_scoretotal -0.0001481392 0.0153985265 68.98892 0.992351959
12
                       0.0007498367 0.0009340191 66.90661 0.424929161
          neuroticism 0.0351547075 0.0211703226 68.99083 0.101339384
13
```

# 3.3 Association study (week 12)

Complete case: excluding CATS, CAR, and neuroticism, we fitted two logistic regressions (one with and one without the biomarkers) on the 72 patients with complete data. This likelihood ratio test showed no clear evidence for an association between biomarkers and recovery:

```
Model 1: Y_w12 ~ female + age
Model 2: Y_w12 ~ female + age + MR_OFCthick + HAMD17 + low_hsCRP + lvpet +
        cognitive_cluster2 + cognitive_cluster3 + EEG_vigilance
    Resid. Df Resid. Dev Df Deviance Pr(>Chi)
1     69     76.256
2     62     64.376     7     11.88     0.1046
```

This was confirmed when looking at the biomarker specific effects. We can also see that the biomarkers for which with have most evidence against the null (MR\_OFCthick and cognitive\_cluster2) differ from week 8

```
estimate std.error
                                             odd ratio
                                                          p.value adjusted p-value
                   15.1377108 9.66268612 3.751667e+06 0.11720467
(Intercept)
                                                                                 NA
                   -0.1631019 0.71162765 8.495046e-01 0.81871691
female
                                                                                 NA
                    0.1239538 0.07958311 1.131964e+00 0.11934255
age
                                                                                 NA
                   -8.2502682 3.58368566 2.611885e-04 0.02132536
MR_OFCthick
                                                                          0.1342289
HAMD17
                    0.1706117 0.10751351 1.186030e+00 0.11253840
                                                                          0.4384615
                    1.0809026 0.83834457 2.947339e+00 0.19728347
low_hsCRP
                                                                          0.5786588
                   -0.5703845 2.60457709 5.653080e-01 0.82665539
                                                                          0.9698902
lvpet
cognitive_cluster2 -1.4448166 0.79614533 2.357893e-01 0.06956004
                                                                          0.3344590
cognitive_cluster3 -0.8051151 0.88884433 4.470365e-01 0.36504179
                                                                          0.7385351
EEG_vigilance
                   -0.0668419 0.60067967 9.353431e-01 0.91139660
                                                                          0.9698902
```

Multiple imputation: as a sensitivity analysis, we now used all patients and all biomarkers and use multiple imputations (100 datasets) to handle missing value. Results are rather similar to the complete case analysis, but with a stronger evidence for an assocation between OFC thickness and recovery. Note that cognition and CATS are bordeline significant without adjustment for multiple comparisons.

```
summary(pool(e.glm_impw12))[,c(1,2,3,5:6)]
```

```
term
                          estimate
                                     std.error
                                                     df
                                                             p.value
          (Intercept) 16.760516575 9.890355948 65.99623 0.094859430
1
2
            sexfemale -0.898638556 0.731432568 65.79797 0.223597826
                  age 0.115967103 0.075427653 66.00500 0.128960677
3
4
          MR OFCthick -9.520173723 3.499858676 65.92820 0.008334242
5
               HAMD17
                       0.141151055 0.103171990 65.81972 0.175928909
6
                       1.004339184 0.857512761 65.03615 0.245782198
7
                lvpet -0.643880653 2.660297543 65.99493 0.809504882
  cognitive_cluster2 -1.906920071 0.887507143 65.52557 0.035364314
8
   cognitive_cluster3 -1.725228423 0.949783976 65.71480 0.073863526
9
10
        EEG_vigilance -0.382219479 0.630217817 66.01354 0.546270922
11
      CATS_scoretotal 0.037284123 0.020188426 66.02463 0.069257123
             CAR AUCi 0.001358667 0.001228576 65.78712 0.272803070
12
          neuroticism 0.017883788 0.023653524 65.95409 0.452297597
13
```

## 3.4 Predictive value (week 8)

Complete case: excluding CATS, CAR, and neuroticism, we assessed the predictive performance of two logistic regressions (one with glm\_ccw8 and one without the biomarkers glm0\_ccw8) as well as a random forest model (rf\_ccw8) on the 75 patients with complete data:

```
method metric
                         model
                                estimate
 1: internal
                 auc glm0_ccw8 0.6276901
                      glm_ccw8 0.8113343
 2: internal
                 auc
                       rf_ccw8 1.0000000
 3: internal
                 auc
4: internal
              brier glm0_ccw8 0.2387024
 5: internal
              brier
                      glm_ccw8 0.1748545
 6: internal
              brier
                       rf_ccw8 0.0000000
7:
                 auc glm0 ccw8 0.6058810
          cv
                      glm_ccw8 0.7187083
8:
          cv
                 auc
9:
                       rf_ccw8 0.5611190
          cv
                 auc
              brier glm0_ccw8 0.2565422
10:
          cv
                      glm_ccw8 0.2298009
11:
              brier
          cv
12:
              brier
                       rf_ccw8 0.4425000
          cv
```

After cross-validation, we observe that both the AUC and brier score of the random forest (with biomarkers) are similar to the logistic regression without biomarkers. This indicates poor predictive ability of the random forest that will not be considered further. The logistic model with biomarker has a higher AUC (i.e., better discrimination) and lower brier score (i.e., smaller discrepancy between prediction and observed outcome) compared to the logistic model without biomarkers. The permutation test confirmed that the logistic model with biomarkers was informative (p=0.003388554 for the AUC and p=0.003012048 for the brier score) while there was no clear evidence with the logistic model without covariates (p=0.1216114 for the AUC and p=0.8189006 for the brier score). This difference between the predictions from the two models (after cross validation) is illustrated in ??, as well as the calibration of the model with biomarkers.

Note that the average AUC estimated by the permutation test was 0.5, supporting that the proposed cross-validation procedure is unbiased (under the null).

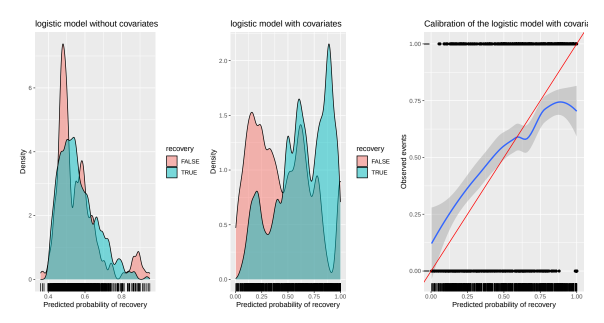


Figure 2: Distribution of the predicted probability of recovery according to the actual recovery for the logistic model without biomarkers (left) and with biomarkers (right). Calibration curve for the latter model: discrepancy of the smoother (in blue) from the identity line (in red) reflect poor calibration.

Full data: as a sensitivity analysis, we now used all patients and all biomarkers and modified the cross-validation procedure to handle missing data. We obtain slightly different results, but still in favor of the logistic model with biomarkers. The difference in AUC between the logistic models is similar to previously (about +0.1) but now the brier score is worse (+0.4 instead of 0.3) indicating poor calibration.

```
method metric
                      model
                             estimate
               auc glm0_w8 0.5991949
1: internal
2: internal
                     glm_w8 0.8878666
             brier glm0_w8 0.2409137
3: internal
  internal
             brier
                     glm_w8 0.1432597
5:
               auc glm0_w8 0.5574226
         cv
6:
                    glm_w8 0.6542563
         cv
               auc
7:
             brier glm0_w8 0.2581057
         cv
8:
             brier
                    glm_w8 0.3015577
```

#### 3.5 Predictive value (week 12)

Complete case: excluding CATS, CAR, and neuroticism, we assessed the predictive performance of two logistic regressions (one with glm\_ccw12 and one without the biomarkers glm0\_ccw12) as well as a random forest model (rf\_ccw12) on the 72 patients with complete data:

```
method metric
                          model estimate
 1: internal
                auc glm0_ccw12 0.6603774
                      glm_ccw12 0.7944389
 2: internal
                auc
                      rf_ccw12 0.0000000
 3: internal
                auc
              brier glm0_ccw12 0.1816008
4: internal
 5: internal
              brier
                      glm_ccw12 0.1493762
 6: internal
              brier
                       rf_ccw12 0.6554699
7:
                auc glm0 ccw12 0.6200357
          cv
                      glm_ccw12 0.6058274
8:
          cv
                auc
9:
                       rf_ccw12 0.2321845
          cv
                auc
              brier glm0_ccw12 0.2116598
10:
          CV
                      glm_ccw12 0.2404126
11:
              brier
          cv
12:
              brier
                       rf_ccw12 0.4660306
          cv
```

Random forests still seems to underperform but, compared to week 8, adding the biomarkers do not seems to greatly improve the AUC and lead to a worse Brier score.

**Full data**: as a sensitivity analysis, we now used all patients and all biomarkers and modified the cross-validation procedure to handle missing data. The results are in line with the complete case, with worse performances when adding the biomarkers.

```
method metric
                      model
                             estimate
               auc glm0_w12 0.6502311
1: internal
                    glm_w12 0.8936826
2: internal
             brier glm0_w12 0.1855812
3: internal
                   glm_w12 0.1172801
4: internal
             brier
5:
               auc glm0_w12 0.6017917
         cv
6:
               auc glm_w12 0.5762885
         cv
             brier glm0 w12 0.2181458
7:
8:
             brier glm_w12 0.2679212
```

## 4 Conclusion

There is evidence that some biomarkers (EEG, to a lesser extend cognition) are predictive of recovery at week 8. The corresponding gain in AUC was about +0.1 with a small improvement in brier score. These results were not seen at week 12 and the overall predictive performance was not great though (AUC of about 0.6 and brier score >0.2).

There was no evidence for non-linear effect or interaction between biomarkers (as assessed via a random forest model) probably due to the limited sample size.

Generally the results where robust to how missing data were handled (complete case or multiple imputation). Effects had a slight tendency to be stronger when not using complete case. The only exception is for the brier score at w8 which became worse compared to the complete case.

# 5 References

LeDell, E., Petersen, M., and van der Laan, M. (2015). Computationally efficient confidence intervals for cross-validated area under the roc curve estimates. *Electronic journal of statistics*, 9(1):1583.

Moons, K. G., Donders, R. A., Stijnen, T., and Harrell Jr, F. E. (2006). Using the outcome for imputation of missing predictor values was preferred. *Journal of clinical epidemiology*, 59(10):1092–1101.