# WORKING EXAMPLE 4

In: A guide to test association between Polygenic Risk Scores and psychological and psychiatric traits: practical examples

Itziar Irigoien, Patricia Mas-Bermejo, Sergi Papiol, Neus Barrantes-Vidal, Araceli Rosa, and Concepción Arenas

### Working flow and code

In this example we simulate 10 PRSs, and a binary trait, with sex, clinical diagnosis (with 2 categories), age, and two Principal Components as covariates.

• data reading

```
dat <- read.table("WExample4.csv", header=TRUE, sep=";", dec=",")</pre>
names(dat) #
    [1] "Sex"
                       "Diagnostic" "Age"
                                                                 "PRS.1"
                                                   "Trait"
    [6] "PRS.2"
                      "PRS.3"
                                    "PRS.4"
                                                   "PRS.5"
                                                                 "PRS.6"
## [11] "PRS.7"
                       "PRS.8"
                                     "PRS.9"
                                                   "PRS.10"
                                                                 "PC1"
## [16] "PC2"
```

• do not forget to declare the categorical variables as factors

```
dat$Sex <- as.factor(dat$Sex)
dat$Diagnostic <- as.factor(dat$Diagnostic)
dat$Trait <- as.factor(dat$Trait)</pre>
```

#### 1. What full model should be considered?

First, given a particular PRS (named PRS.i), consider all the possible full models:

- $FM_{WI}$ : log(p/1-p) versus PRS.i + Sex + Diagnostic + Age + PC1 + PC2
- $FM_{Sex}$ : log(p/1-p) versus  $PRS.i + Sex + PRS.i \cdot Sex + Diagnostic + Age + PC1 + PC2$
- $FM_{Diagnostic}$ : log(p/1-p) versus PRS.i + Sex + Diagnostic + PRS.i · Diagnostic + Age + PC1 + PC2
- FM  $_{Sex/Diagnostic}$ : log(p/1-p) versus PRS.i + Sex + PRS.i · Sex + Diagnostic + PRS.i · Diagnostic + Age + PC1 + PC2

## 2. How to make a PRS ranking to find the important ones?

As is described in the paper, for each model, calculate the Tjur's coefficients of discrimination. If Nagelkerke's  $R^2$  is preferred, set statistic="PseudoR2" in function orderBin(), and calculate their sum, S.

According to S, list the PRSs in decreasing order:

```
# Order the PRSs
out <- orderBin(dat, yname="Trait", prsname = "PRS.", statistic = "D")
head(out)</pre>
```

```
## Model1 Model2 Model3 Model4 Sum
## PRS.7 0.1816453 0.1816453 0.2175212 0.2176745 0.7984862
```

```
## PRS.8 0.1657769 0.1657769 0.1921405 0.1999160 0.7236103

## PRS.1 0.1644174 0.1644174 0.1716650 0.1874602 0.6879599

## PRS.2 0.1648077 0.1648077 0.1710979 0.1799683 0.6806815

## PRS.3 0.1638651 0.1638651 0.1735185 0.1785984 0.6798472

## PRS.4 0.1581438 0.1581438 0.1715735 0.1741957 0.6620568

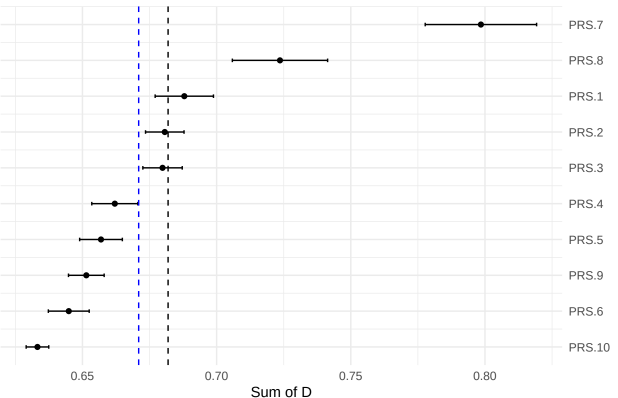
mainfilename <- "WExample4"

filename <- paste0(mainfilename, "_Ordered_PRS.csv")

write.csv2(out,file=filename)
```

Plot the sum of coefficients of discrimination coefficients D. Lines: in blue the median; in black the mean.

```
out <- data.frame(out)</pre>
n <- dim(out)[1]</pre>
select <- grep("Model", names(out), value=FALSE)</pre>
out$effect <- out$Sum</pre>
sds <- apply(out[, select], 1, sd)</pre>
out$lower <- out$effect - sds</pre>
out$upper <- out$effect + sds</pre>
out$rank <- n:1</pre>
ggplot(data=out, aes(y=rank, x=effect, xmin=lower, xmax=upper)) +
  geom_point() +
  geom_errorbarh(height=.1) +
  scale_y_continuous(name=NULL, breaks= n:1, labels=row.names(out), position="right") +
  labs(title='', x='Sum of D', y = 'PRS') +
  geom_vline(xintercept=mean(out$effect), color='black', linetype='dashed') +
  geom_vline(xintercept=median(out$effect), color='blue', linetype='dashed') +
  theme_minimal()
```



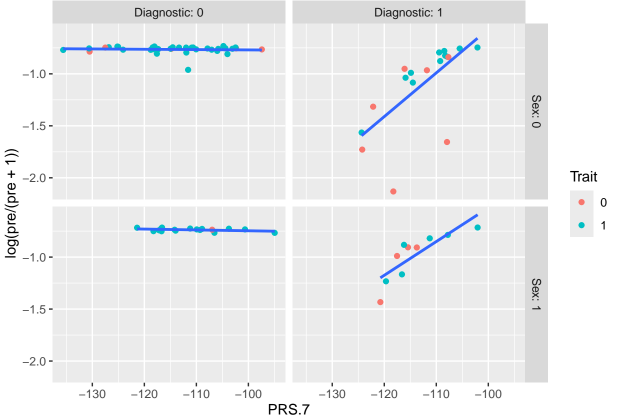
According to the obtained results, first PRS.7 is selected to analyse its association with the Trait.

### 3. Which model, of all the possible ones, should be used?

The following figure represents the scatter plot separated by Sex and Diagnostic groups.

```
# First candidate PRS.7
# Plot it
M <- glm(Trait ~ PRS.7*Sex + PRS.7*Diagnostic + Age + PC1 + PC2, data=dat, family=binomial())
pre <- M$fitted.values #predict(M, type='response')
ggplot(dat, aes(x=PRS.7, y=log(pre/(pre+1)))) +
    geom_point(aes(color=Trait)) +
    geom_smooth(method=lm, se=FALSE)+
    facet_grid(Sex ~ Diagnostic, labeller=label_both)</pre>
```

## `geom\_smooth()` using formula = 'y ~ x'



```
# Candidate FM Trait ~ PRS + Sex + Diagnostic + PRS*Diagnostic + C1 +C2
```

The plots suggest that the interaction between the PRS.7 and the diagnostic is relevant. Thus, we set the full model candidate (FM):  $log(p/1-p) \sim PRS + Sex + Diagnostic + PRS \cdot Diagnostic + Age + PC1 + PC2$ .

#### 5. For a binary trait, what steps should be followed for a correct analysis?

Check for overdispersion

```
#model
FM <- glm(Trait ~ Sex + PRS.7*Diagnostic + Age + PC1 + PC2, data=dat, family=binomial())
#Residual Deviance
FM$deviance</pre>
```

```
## [1] 67.46733
```

#### # Ratio

FM\$deviance/FM\$df.residual

```
## [1] 0.92421
```

Since this ratio is close to 1, there is not evidence of overdispersion.

#### ## [1] 0.3389642

With this p-value = 0.3389, we conclude that there is not evidence of overdispersion.

Based on the following table...

```
summary(FM)
```

```
##
## Call:
## glm(formula = Trait ~ Sex + PRS.7 * Diagnostic + Age + PC1 +
      PC2, family = binomial(), data = dat)
##
## Coefficients:
##
                     Estimate Std. Error z value Pr(>|z|)
## (Intercept)
                    1.953493 5.754567 0.339
                                                  0.7343
## Sex1
                     0.445672
                               0.698490
                                         0.638
                                                  0.5234
## PRS.7
                     0.008372
                               0.051088
                                         0.164
                                                  0.8698
## Diagnostic1
                                         1.500
                    16.714171 11.146266
                                                  0.1337
                     0.043589
                              0.065843
                                         0.662
                                                  0.5080
## Age
                               4.517987 -1.232
## PC1
                                                  0.2180
                    -5.565987
## PC2
                     3.784619
                               5.604881
                                         0.675
                                                  0.4995
## PRS.7:Diagnostic1 0.162405
                                0.097745
                                         1.662
                                                  0.0966 .
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for binomial family taken to be 1)
      Null deviance: 83.234 on 80 degrees of freedom
##
## Residual deviance: 67.467 on 73 degrees of freedom
## AIC: 83.467
## Number of Fisher Scoring iterations: 5
```

... the results show that PRS.7 is related with the log(p/1-p) in the following way:

- $log(p/1 p) = 1.953 + 0.008 \times PRS.7 + 0.446 \times Sex + 0.044 \times Age 5.566 \times PC1 + 3.785 \times PC2$ , if Diagnostic = 0.
- $log(p/1-p) = (1.953 + 16.714) + (0.008 + 0.162) \times PRS.7 + 0.446 \times Sex + 0.044 \times Age 5.566 \times PC1 + 3.785 \times PC2$ , if Diagnostic = 1.

Check whether the respective PRS coefficients under each group are significant or not.

```
summary(glht(FM, "PRS.7 = 0"))
```

```
##
##
     Simultaneous Tests for General Linear Hypotheses
##
## Fit: glm(formula = Trait ~ Sex + PRS.7 * Diagnostic + Age + PC1 +
##
       PC2, family = binomial(), data = dat)
##
## Linear Hypotheses:
##
              Estimate Std. Error z value Pr(>|z|)
## PRS.7 == 0 0.008372
                         0.051088
                                    0.164
## (Adjusted p values reported -- single-step method)
summary(glht(FM, "PRS.7 + PRS.7:Diagnostic1 = 0"))
##
##
     Simultaneous Tests for General Linear Hypotheses
##
## Fit: glm(formula = Trait ~ Sex + PRS.7 * Diagnostic + Age + PC1 +
##
       PC2, family = binomial(), data = dat)
##
## Linear Hypotheses:
                                  Estimate Std. Error z value Pr(>|z|)
## PRS.7 + PRS.7:Diagnostic1 == 0 0.17078
                                              0.08476
                                                        2.015
                                                                0.0439 *
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Adjusted p values reported -- single-step method)
```

That means that for those with Diagnostic=0, it seems that the PRS.7 is not related to the Trait with odds  $= \exp(0.0082) = 1.008$ , but for those with Diagnosis =1 the model indicates that the coefficient of PRS.7 is 0.0082 + 0.162 = 0.1702, so the odds increase  $\exp(0.1702) = 1.186$  for an incremental of one unit in PRS.7 with a p-value=0.0439.

It is also possible to compute a permutation test to assess whether the increase in the coefficient of determination D is significative.

In this particular case, it can be seen that the coefficient of discrimination of the FM model is 0.07 units bigger than the corresponding to the *Null Model NM*, but it is not statistically significant.

• Last step: We move to the next PRS.