Advantage of two scans in small populations demonstrated with a transgender dataset.

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Introduction

We have a dataset where FreeSurfer was used to determine cortical and subcortical brain anatomy in cis- and transgender population. For every participant two scans were administered (T1 and T2) and an average of both scans was computed. In this report we analyze the data from one scan and the average of both scans. Later on we show the advantage and increase in power obtained by administering 2 scans. One participant (P22) was removed from the analysis because no anatomical data was available for this participant.

Actual data

We first read in the data. In data all available measurements are stored (descriptive and anatomical) while in data hyp the anatomical results for every region of interest stored. Both files contain data of T1, T2 and the average.

```
# Read in data
data.all <- read.csv("../1.Data/Behzad_all.csv", sep=";", dec=",")
data.hyp <- read.csv("../1.Data/Behzad_hyp.csv", sep=";", dec=",")

# Check data
dim(data.hyp)

## [1] 140 80
dim(data.all)</pre>
## [1] 140 820
```

The regions we are interested in are the cerebellum, caudate, putamen, nucleus accumenbens, thalamus, fusiform, pre-central gyrus, post-central gyrus, frontal poles and inferior parietal gyrus. Here we list the variables we selected from FreeSurfer that comply with these regions.

```
# Regions of interest
names(data.hyp[,55:80])
##
    [1] "Tavg_LeftCerebellumWhiteMatter"
    [2] "Tavg_LeftCerebellumCortex"
##
    [3] "Tavg_RightCerebellumWhiteMatter"
##
       "Tavg_RightCerebellumCortex"
       "Tavg L caudalanteriorcingulate volume"
       "Tavg_L_caudalmiddlefrontal_volume"
##
       "Tavg_L_fusiform_volume"
##
    [7]
##
    [8] "Tavg L inferiorparietal volume"
    [9] "Tavg_L_postcentral_volume"
  [10] "Tavg_L_precentral_volume"
   [11] "Tavg_L_frontalpole_volume"
   [12] "Tavg_R_caudalanteriorcingulate_volume"
## [13] "Tavg_R_caudalmiddlefrontal_volume"
```

```
## [14] "Tavg_R_fusiform_volume"
## [15] "Tavg_R_inferiorparietal_volume"
## [16] "Tavg_R_postcentral_volume"
## [17] "Tavg_R_precentral_volume"
## [18] "Tavg_R_frontalpole_volume"
## [19] "Tavg_LeftThalamusProper"
## [20] "Tavg_LeftCaudate"
## [21] "Tavg_LeftPutamen"
## [22] "Tavg_LeftAccumbensarea"
## [23] "Tavg_RightThalamusProper"
## [24] "Tavg_RightCaudate"
## [25] "Tavg_RightPutamen"
## [26] "Tavg_RightAccumbensarea"
```

Descriptives

In this section population parameters are presented.

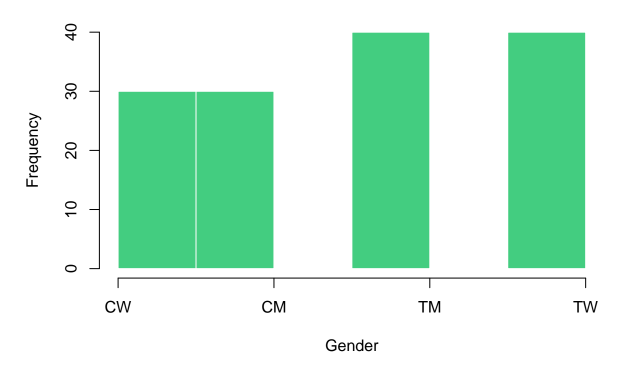
Gender

There were 60 cisgender and 80 transgender participants. One participant (P22) was removed from the analysis because no anatomical data was available for this participant.

```
ng1 <- sum(data.all[,2]==1)
ng2 <- sum(data.all[,2]==2)
ng3 <- sum(data.all[,2]==3)
ng4 <- sum(data.all[,2]==4)

paste("Cisgender women = ", ng1, ", cisgender men = ", ng2, ", transgender men = ", ng3, ", transgender
## [1] "Cisgender women = 30, cisgender men = 30, transgender men = 40, transgender women = 40"
hist(data.all[,2], xlab = "Gender", xaxt = "n", col = "seagreen3", border = "white", main = "Frequency axis(1,at=c(1:4),labels=c("CW", "CM", "TM", "TW"))</pre>
```

Frequency of gender



\mathbf{Age}

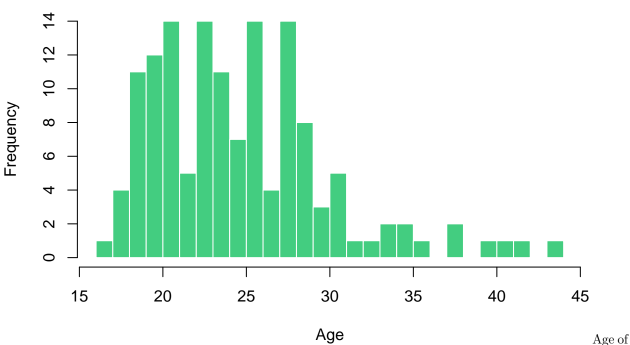
```
summary(data.all[,4])

## Min. 1st Qu. Median Mean 3rd Qu. Max.

## 16.00 21.00 24.00 25.18 28.00 44.00

hist(data.all[,4], xlab = "Age", col = "seagreen3", border = "white", main = "Histogram of age distributed in the seagreen of age distributed in the se
```

Histogram of age distribution



the participants ranged from 16 to 44. If we look at the distribution of age in the cis- and transgender group we see that the range is similar in both groups.

```
# Cisgender group
summary(data.all[data.all[,2]<3,4])</pre>
##
      Min. 1st Qu.
                     Median
                                Mean 3rd Qu.
                                                 Max.
##
     19.00
              23.00
                      25.00
                               25.92
                                       29.00
                                                41.00
# Transgender group
summary(data.all[data.all[,2]>2,4])
##
      Min. 1st Qu.
                     Median
                                Mean 3rd Qu.
                                                 Max.
     16.00
              20.75
                                       26.25
##
                      24.00
                               24.62
                                                44.00
```

Social-Economic Status

```
table(data.all[,5])
##
## 1 2 3
## 19 87 34
```

Level of education

22 76 28 14

```
table(data.all[,6])
##
## 2 3 4 5
```

Handedness

```
table(data.all[,7])
##
##
   1
        2
## 129 11
Gender identity
summary(data.all[,8])
##
     Min. 1st Qu. Median
                            Mean 3rd Qu.
                                            Max.
    0.000 0.000
                    5.000
                            5.036 10.000 10.000
table(data.all[,8])
##
## 0 1 2 3 7 8 9 10
## 49 4 13 4 1 9 16 44
# Cisqender women
summary(data.all[data.all[,2]==1,8])
     Min. 1st Qu. Median
##
                          Mean 3rd Qu.
                                            Max.
##
    7.000
           9.000
                   9.500 9.267 10.000 10.000
# Cisqender men
summary(data.all[data.all[,2]==2,8])
##
     Min. 1st Qu. Median
                            Mean 3rd Qu.
                                            Max.
##
      0.0
              0.0
                      0.0
                             0.9
                                     2.0
                                             3.0
# Transgender women
summary(data.all[data.all[,2]==3,8])
##
     Min. 1st Qu. Median
                            Mean 3rd Qu.
                                            Max.
    0.000
           0.000
                    0.000
                            0.375
                                   0.000
                                           3.000
# Transgender men
summary(data.all[data.all[,2]==4,8])
##
     Min. 1st Qu. Median
                            Mean 3rd Qu.
                                            Max.
##
    8.000
          9.000 10.000
                            9.625 10.000 10.000
Sexual orientation
summary(data.all[,9])
##
     Min. 1st Qu. Median
                            Mean 3rd Qu.
                                            Max.
    0.000
          0.000
                    5.000
                            4.957 10.000 10.000
table(data.all[,9])
##
## 0 1 2 3 7 8 9 10
## 50 12 7 1 1 9 14 46
```

```
# Cisqender women
summary(data.all[data.all[,2]==1,9])
      Min. 1st Qu. Median
##
                              Mean 3rd Qu.
                                              Max.
   0.0000 0.0000 0.0000 0.5333 1.0000
##
                                           2.0000
# Cisgender men
summary(data.all[data.all[,2]==2,9])
##
     Min. 1st Qu. Median
                              Mean 3rd Qu.
                                              Max.
     7.000
                             9.533 10.000
##
            9.250 10.000
                                           10.000
# Transgender women
summary(data.all[data.all[,2]==3,9])
##
     Min. 1st Qu. Median
                             Mean 3rd Qu.
                                              Max.
     8.000
            9.000 10.000
                             9.475 10.000 10.000
##
# Transgender men
summary(data.all[data.all[,2]==4,9])
##
      Min. 1st Qu.
                   Median
                              Mean 3rd Qu.
                                              Max.
##
     0.000
            0.000
                     0.000
                             0.325
                                     0.000
                                             3.000
```

Mental illnesses

There are no available measures for the cisgender group. ####Somatization

```
table(data.all[,22])
##
## #NULL! 0 1
## 60 67 13
```

Obsessive-compulsive disorder

```
table(data.all[,23])
##
## #NULL! 0 1
## 60 58 22
```

table(data.all[,25])

Among the cisgender participants 1 had a history of obsession.

```
## ## #NULL! 0 1 2
## 60 42 37 1
```

Among the cisgender participants 3 had a history of depression.

Anxiety

```
table(data.all[,26])

##
## #NULL! 0 1 2
## 60 54 25 1
```

Among the cisgender participants 1 had a history of a general anxiety disorder.

Hostility

```
table(data.all[,27])
##
## #NULL! 0 1 2
## 60 64 15 1
```

Phobic anxiety

```
table(data.all[,28])

##
## #NULL! 0 1
## 60 67 13
```

Paranoia

##

##

```
table(data.all[,29])
##
## #NULL! 0 1
```

Psychotism

60

46

67

34

13

```
table(data.all[,30])
##
## #NULL! 0 1
```

Global severity

60

```
table(data.all[,31])
....
```

```
## #NULL! 0 1
## 60 59 21
```

Cisgender group

Past psychiatric condition Out of 60 cisgender participants 3 had a history of depression, 1 reported a general anxiety disorder and 1 had a history of obsession.

Past medical condition Out of 60 cisgender participants 3 reported migraine, 1 reported left ear surgery, 1 participant had suffered from heart palpitations, 1 participants reported a history of meningitis, 1 participant reported they had asthma as a child and 1 participants reported favism.

Analysis of 1 measurement

```
bg <- 3
nd <- 28
ln <- nd-bg
# Object to save p-values of ANOVA
pan <- array(data=NA, dim = ln)
# Compute ANOVA for every predictor and save p-value
for(i in bg:nd){
  tempan <- aov(data.hyp[,i] ~ as.factor(data.hyp[,2]))</pre>
 pan[i-bg+1] <- unlist(summary(tempan))[9]</pre>
}
# FDR correction on p-values to correct for multiple testing
pancorr <- p.adjust(pan, method = "bonferroni")</pre>
round(pancorr,3)
## [1] 0.918 1.000 0.000 0.006 0.025 0.746 0.132 0.016 1.000 0.000 0.006
## [12] 0.312 0.029 0.012 0.008 0.000 0.018 0.000 0.000 0.039 0.116 1.000
## [23] 0.000 0.032 0.000 1.000
sum(pancorr < 0.05)</pre>
## [1] 17
names(data.hyp[,which(pancorr < 0.05) + bg - 1])</pre>
    [1] "T1_L_fusiform_volume"
   [2] "T1_L_inferiorparietal_volume"
##
##
   [3] "T1_L_postcentral_volume"
  [4] "T1_R_caudalanteriorcingulate_volume"
##
  [5] "T1 R fusiform volume"
##
   [6] "T1_R_inferiorparietal_volume"
##
##
   [7] "T1_R_precentral_volume"
##
  [8] "T1_R_frontalpole_volume"
##
  [9] "T1_LeftCerebellumWhiteMatter"
## [10] "T1 LeftCerebellumCortex"
## [11] "T1_RightCerebellumWhiteMatter"
## [12] "T1 RightCerebellumCortex"
## [13] "T1_LeftThalamusProper"
## [14] "T1_LeftCaudate"
## [15] "T1_RightThalamusProper"
## [16] "T1_RightCaudate"
## [17] "T1_RightPutamen"
```

Analysis of the average

```
bg <- 55
nd <- 80
ln \leftarrow nd-bg+1
# Object to save p-values of ANOVA
pan <- array(data=NA, dim = ln)
# Compute ANOVA for every predictor and save p-value
for(i in bg:nd){
  tempan <- aov(data.hyp[,i] ~ as.factor(data.hyp[,2]))</pre>
  pan[i-bg+1] <- unlist(summary(tempan))[9]</pre>
# FDR correction on p-values to correct for multiple testing
pancorr <- p.adjust(pan, method = "bonferroni")</pre>
round(pancorr,3)
## [1] 0.049 0.000 0.040 0.000 0.698 1.000 0.001 0.076 0.089 1.000 1.000
## [12] 0.015 1.000 0.009 0.053 1.000 0.193 0.035 0.000 0.073 0.008 1.000
## [23] 0.000 0.146 0.022 0.292
sum(pancorr < 0.05)</pre>
## [1] 12
names(data.hyp[,which(pancorr < 0.05) + bg - 1])</pre>
   [1] "Tavg_LeftCerebellumWhiteMatter"
  [2] "Tavg_LeftCerebellumCortex"
##
## [3] "Tavg_RightCerebellumWhiteMatter"
## [4] "Tavg_RightCerebellumCortex"
## [5] "Tavg_L_fusiform_volume"
##
  [6] "Tavg_R_caudalanteriorcingulate_volume"
## [7] "Tavg_R_fusiform_volume"
## [8] "Tavg_R_frontalpole_volume"
## [9] "Tavg_LeftThalamusProper"
## [10] "Tavg_LeftPutamen"
## [11] "Tavg RightThalamusProper"
## [12] "Tavg_RightPutamen"
```

Why are there less regions for which the difference is statistically significant when the average is used compared to when one measure is used?

Correlation between T1 and T2

```
bg <- 36
nd <- 108
ln <- nd-bg+1

# Object to save correlations
corrall <- array(data=NA, dim = ln)</pre>
```

```
# Compute ANOVA for every predictor and save p-value
for(i in bg:nd){
   corrall[i-bg+1] <- cor(x = data.all[,i], y = data.all[,i + ln + 1])
}
summary(corrall)

## Min. 1st Qu. Median Mean 3rd Qu. Max.
## 0.6334 0.7950 0.8243 0.8216 0.8466 0.9985</pre>
```

Simulations

Intro

Code

First we need to define the parameters of our simulations.

```
# variance/sd epsilon
seps <- 1
# Number of simulations
asim <- 10
# Effect size
delta <- 0.8
# Number of participants
n <- 30
n.1 <- n/2
           # in the first group
n.2 < - n/2
              # in the second group
# Level of statistical significance
alpha \leftarrow 0.05
# Correlation between first and second measurement
rho \leftarrow seq(0.01,0.99,0.01)
```

Then we prepare objects to store our results

```
# Number of simulations
pow.mean1<-vector("numeric",length(rho))
pow.mean2<-vector("numeric",length(rho))
pow.mean3<-vector("numeric",length(rho))
pow.mean4<-vector("numeric",length(rho))

## Loading required package: Matrix
##
## Attaching package: 'lmerTest'

## The following object is masked from 'package:lme4':
##
## lmer</pre>
```

```
## The following object is masked from 'package:stats':
##
##
# Loop over preset correlations between measure 1 and measure 2
for(i in 1:length(rho)){
  # Create objects to store power in for every simulations
  pow.1<-vector("numeric",asim)</pre>
  pow.2<-vector("numeric",asim)</pre>
  pow.3<-vector("numeric",asim)</pre>
  pow.4<-vector("numeric",asim)</pre>
  for(k in 1:asim){
    # Scenario 1: lower bound of power curve
    # two groups with equal amount of subjects, groups differ with an effect size delta
    # Construct a vector that determines in which group each subject falls
    x < -c(rep(1,n.1),rep(0,n.2))
    # Vector with observations in the set of participants
    y<-rnorm(n,0,seps)
    # Add an effect size to the first group
    y[1:n.1] < -y[1:n.1] + delta
    # Boolean of whether an effect is detected, this is later used to compute the power
    \label{local_pow.1} \begin{split} \text{pow.1[k]} <&-\text{summary}(\text{lm}(\text{y-x})) \\ \text{$^{\text{coef}[2,4]$}$} \end{aligned} \\ \text{$^{\text{alpha}}$} \end{split}
    # Scenario 2: upper bound of power curve
    # two groups with equal amount of subjects, twice as many as scenario 1, groups differ with an effe
    # Construct a vector that determines in which group each subject falls
    x2 < -c(rep(1,(n.1*2)),rep(0,(n.2*2)))
    # Vector with observations in the set of participants
    y2 < -rnorm(n*2, 0, seps)
    # Add an effect size to the first group
    y2[1:(n.1*2)]<-y2[1:(n.1*2)]+delta
    # Boolean of whether an effect is detected, this is later used to compute the power
    pow.2[k] < -summary(lm(y2~x2))$coef[2,4] <alpha
    # Scenario 3: two measurements for every subject, same amount of subjects as in scenario 1
    # two groups with equal amount of subjects, correlation between measurements, groups differ with an
    # Construct a vector that determines in which group each subject falls
    x3 < -c(rep(1,n.1), rep(0,n.2), rep(1,n.1), rep(0,n.2))
    # Vector with first observation of every participant
    y3 < -rnorm(n, 0, seps)
    # Factor to multiply second set of observations with to obtain results in line with predefined corr
    alpac<-sqrt(rho[i]^2/(1-rho[i]^2)*seps)</pre>
    # Construct second set of observations that are correlated with first set (y3)
    y3.2u<-alpac*y3+rnorm(n)
    y3.2 < -y3.2u/sqrt(var(y3.2u))
    # Add effect size to the first group of participants
    y3[1:n.1] < -y3[1:n.1] + delta
    y3.2[1:n.1] < -y3.2[1:n.1] + delta
```

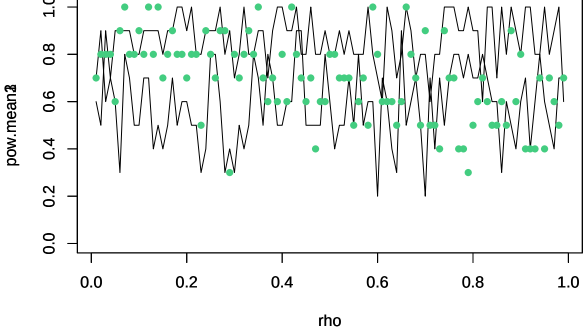
```
# Combine both observations in 1 vector
  y3o < -c(y3, y3.2)
  # Define subject numbers
  subject<-rep(1:n,2)</pre>
  # Construct mixed model
  mm<-lmer(y3o ~ x3 + (1 | subject))</pre>
  # Boolean of whether an effect is detected, this is later used to compute the power
  pow.3[k] <-summary(mm)$coef[2,5] <alpha</pre>
  # Scenario 4: What if we work with the average?
  y3m < -(y3+y3.2)/2
  # Boolean of whether an effect is detected, this is later used to compute the power
  pow.4[k] < -summary(lm(y3m-x)) $coef[2,4] <alpha
pow.mean1[i] <-mean(pow.1)</pre>
pow.mean2[i]<-mean(pow.2)</pre>
pow.mean3[i]<-mean(pow.3)</pre>
pow.mean4[i]<-mean(pow.4)</pre>
```

Results

```
# power of taking both measures into account
plot(rho,pow.mean1,ylim=c(0,1), type="l")
par(new = TRUE)
plot(rho,pow.mean2,ylim=c(0,1), type="l")
par(new = TRUE)
plot(rho,pow.mean3,ylim=c(0,1), type="p", col = "goldenrod3", pch = 2)
      \infty
      o
pow.mean3
     9.0
     0.4
      0.2
     0.0
                          0.2
            0.0
                                         0.4
                                                        0.6
                                                                       8.0
                                                                                      1.0
```

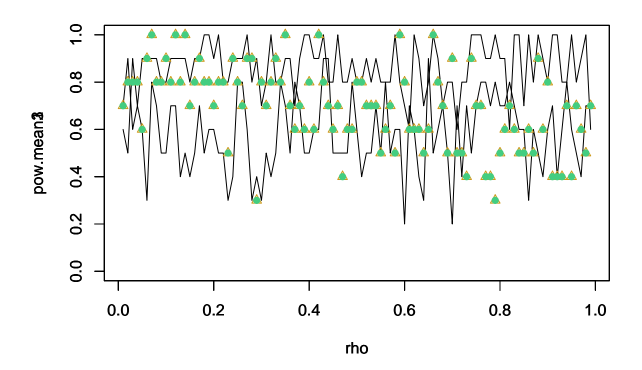
rho

```
# power of using the average
plot(rho,pow.mean1,ylim=c(0,1), type="l")
par(new = TRUE)
plot(rho,pow.mean2,ylim=c(0,1), type="l")
par(new = TRUE)
plot(rho,pow.mean3,ylim=c(0,1), type="p", col = "seagreen3", pch = 16)
```



If we overlay both plots we see that there is a lot of overlap. This can also be demonstrated by looking at the results that are exactly the same.

```
##
## FALSE TRUE
## 1 98
# Overlap of both plots
plot(rho,pow.mean1,ylim=c(0,1), type="l")
par(new = TRUE)
plot(rho,pow.mean2,ylim=c(0,1), type="l")
par(new = TRUE)
plot(rho,pow.mean3,ylim=c(0,1), type="p", col = "goldenrod3", pch = 2)
par(new = TRUE)
plot(rho,pow.mean3,ylim=c(0,1), type="p", col = "seagreen3", pch = 16)
```



Discussion