Twin analysis Klaus Holst & Thomas Scheike February 26, 2017

Contents

Mets package

This document provides a brief tutorial to analyzing twin data using the mets package:

```
library("mets")
options(warn=-1)
```

The development version may be installed from *github*, i.e., with the devtools package:

```
devtools::install_github("kkholst/lava")
devtools::install_github("kkholst/mets")
```

Twin analysis, continuous traits

100006.1 100006 28.04282 52.57221

In the following we examine the heritability of Body Mass Index¹, based on data on self-reported BMI-values from a random sample of 11,411 same-sex twins. First, we will load data

```
data("twinbmi")
head(twinbmi)
           tvparnr
                         bmi
                                   age gender zyg
 100001.1 100001 26.33289 57.57974
                                         male DZ
 100002.1 100002 28.65014 57.04860
100003.1 100003 28.40909 57.67830
                                                 ΜZ
                                          male
                                          male
                                                 DΖ
 100004.1 100004 27.25089 53.51677
                                         male
 100005.1 100005 27.77778 52.57495
                                         male DZ
```

The data is on *long* format with one subject per row. we transpose the data allowing us to do pairwise analyses

male DZ

```
twinwide <- fast.reshape(twinbmi, id="tvparnr",varying=c("bmi"))</pre>
head(twinwide)
                                                    bmi2
          tvparnr
                      bmi1
                                 age gender zyg
 100001.1 100001 26.33289 57.57974 male DZ 25.46939
```

100002.1 100002 28.65014 57.04860 male MZ NΔ 100003.1 100003 28.40909 57.67830 100004.1 100004 27.25089 53.51677 male DΖ male DZ 28.07504 100005.1 100005 27.77778 52.57495 male DZ 100006.1 100006 28.04282 52.57221 male DZ 22.30936

Next we plot the association within each zygosity group

```
library("cowplot")
scatterdens <- function(x) {</pre>
    sp <- ggplot(x,</pre>
                 aes_string(colnames(x)[1], colnames(x)[2])) +
        theme_minimal() +
        geom_point(alpha=0.3) + geom_density_2d()
    xdens <- ggplot(x, aes_string(colnames(x)[1],fill=1)) +</pre>
        theme minimal() +
        geom_density(alpha=.5)+
        theme(axis.text.x = element_blank(),
```

- ¹ M. Korkeila, J. Kaprio, A. Rissanen, and M. Koskenvuo. Effects of gender and age on the heritability of body mass index. Int J Obes, 15(10):647-654, Oct 1991
- ² J. v. Hjelmborg, C. Fagnani, K. Silventoinen, M. McGue, M. Korkeila, K. Christensen, A. Rissanen, and J. Kaprio. Genetic influences on growth traits of BMI: a longitudinal study of adult twins. Obesity (Silver Spring), 16(4):847-852, Apr 2008

tvparnr twin id bmi Body Mass Index (kg/m²) age Age (years) gender Gender factor (male,female) zyg zygosity (MZ,DZ)

```
legend.position = "none") + labs(x=NULL)
        ydens <- ggplot(x, aes_string(colnames(x)[2],fill=1)) +</pre>
13
            theme_minimal() +
14
            geom_density(alpha=.5) +
15
            theme(axis.text.y = element_blank(),
                   axis.text.x = element_text(angle=90, vjust=0),
                   legend.position = "none") +
            labs(x=NULL) +
             coord_flip()
        g <- plot_grid(xdens, NULL, sp, ydens,
21
                       ncol=2,nrow=2,
22
                       rel_widths=c(4,1.4),rel_heights=c(1.4,4))
        return(g)
24
    }
```

We here show the log-transformed data which is slightly more symmetric and more appropiate for the twin analysis (see Figure 1 and 2)

```
mz <- log(subset(twinwide, zyg=="MZ")[,c("bmi1","bmi2")])</pre>
scatterdens(mz)
dz <- log(subset(twinwide, zyg=="DZ")[,c("bmi1","bmi2")])</pre>
scatterdens(dz)
```

The plots and raw association measures shows considerable stronger dependence in the MZ twins, thus indicating genetic influence of the trait

```
cor.test(mz[,1],mz[,2], method="spearman")
         Spearman's rank correlation rho
 data: mz[, 1] and mz[, 2]
 S = 165460000, p-value < 2.2e-16
 alternative hypothesis: true rho is not equal to 0
 sample estimates:
       rho
 0.6956209
```

```
cor.test(dz[,1],dz[,2], method="spearman")
```

```
Spearman's rank correlation rho
data: dz[, 1] and dz[, 2]
S = 2162500000, p-value < 2.2e-16
alternative hypothesis: true rho is not equal to 0
sample estimates:
     rho
0.4012686
```

Next we examine the marginal distribution (GEE model with working independence)

```
10 <- lm(bmi ~ gender + I(age-40), data=twinbmi)
estimate(10, id=twinbmi$tvparnr)
```

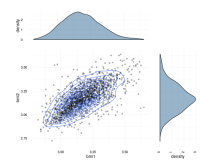


Figure 1: Scatter plot of logarithmic BMI measurements in MZ twins

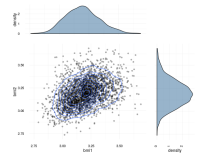


Figure 2: Scatter plot of logarithmic BMI measurements in DZ twins.

```
Estimate Std.Err 2.5% 97.5% P-value
(Intercept)
             23.369 0.05453 23.262 23.476 0.00e+00
             1.407 0.07322 1.264 1.551 2.35e-82
gendermale
I(age - 40)
             0.118 0.00479 0.108 0.127 2.00e-133
```

```
library("splines")
11 <- lm(bmi ~ gender*ns(age,3), data=twinbmi)</pre>
marg1 <- estimate(l1, id=twinbmi$tvparnr)</pre>
dm <- Expand(twinbmi,
             bmi=0,
             gender=c("male"),
             age=seq(33,61,length.out=50))
df <- Expand(twinbmi,</pre>
             bmi=0,
             gender=c("female"),
             age=seq(33,61,length.out=50))
plot(marg1, function(p) model.matrix(l1,data=dm)%*%p,
```

```
data=dm["age"], ylab="BMI", xlab="Age",
         ylim=c(22,26.5))
    plot(marg1, function(p) model.matrix(l1,data=df)%*%p,
         data=df["age"], col="red", add=TRUE)
    legend("bottomright", c("Male", "Female"),
15
           col=c("black","red"), lty=1, bty="n")
```

Polygenic model

Decompose outcome into

$$Y_i = A_i + D_i + C + E_i, i = 1, 2$$

- A Additive genetic effects of alleles
- D Dominante genetic effects of alleles
- C Shared environmental effects
- E Unique environmental genetic effects

Dissimilarity of MZ twins arises from unshared environmental effects only! $Cor(E_1, E_2) = 0$ and

$$Cor(A_1^{MZ}, A_2^{MZ}) = 1$$
, $Cor(D_1^{MZ}, D_2^{MZ}) = 1$,

$$Cor(A_1^{DZ}, A_2^{DZ}) = 0.5$$
, $Cor(D_1^{DZ}, D_2^{DZ}) = 0.25$,

$$Y_i = A_i + C_i + D_i + E_i$$

$$A_i \sim \mathcal{N}(0, \sigma_A^2), C_i \sim \mathcal{N}(0, \sigma_C^2), D_i \sim \mathcal{N}(0, \sigma_D^2), E_i \sim \mathcal{N}(0, \sigma_E^2)$$

$$\begin{split} \mathbb{C}\mathrm{ov}(Y_1,Y_2) = \\ \begin{pmatrix} \sigma_A^2 & 2\Phi\sigma_A^2 \\ 2\Phi\sigma_A^2 & \sigma_A^2 \end{pmatrix} + \begin{pmatrix} \sigma_C^2 & \sigma_C^2 \\ \sigma_C^2 & \sigma_C^2 \end{pmatrix} + \begin{pmatrix} \sigma_D^2 & \Delta_7\sigma_D^2 \\ \Delta_7\sigma_D^2 & \sigma_D^2 \end{pmatrix} + \begin{pmatrix} \sigma_E^2 & 0 \\ 0 & \sigma_E^2 \end{pmatrix} \end{split}$$

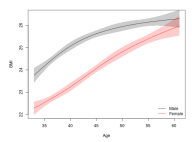


Figure 3: ...

```
dd <- na.omit(twinbmi)
10 <- twinlm(bmi ~ age+gender, data=dd,
          DZ="DZ", zyg="zyg", id="tvparnr", type="sat")
1 <- twinlm(bmi ~ ns(age,1)+gender, data=twinbmi,</pre>
         DZ="DZ", zyg="zyg", id="tvparnr", type="cor", missing=TRUE)
         _____
 Group 1
                     Estimate Std. Error Z value Pr(>|z|)
 Regressions:
                    4.005.
1.41143
   bmi.1~ns(age, 1).1
                               0.16354 25.00328
                                                 <1e-12
                               0.07285 19.37536
   bmi.1~gendermale.1
                                                 <1e-12
 Intercepts:
                     22.57414
                              0.07187 314.08431 <1e-12
   bmi.1
 Additional Parameters:
                      2.44584 0.01425 171.68385 <1e-12
   log(var)
                     atanh(rhoMZ)
 Group 2
                     Estimate Std. Error Z value Pr(>|z|)
 Regressions:
   bmi.1~ns(age, 1).1 4.08914 0.16354 25.00328 <1e-12
   bmi.1~gendermale.1 1.41143 0.07285 19.37536 <1e-12
 Intercepts:
                     22.57414 0.07187 314.08431 <1e-12
   bmi.1
 Additional Parameters:
   log(var)
                      2.44584 0.01425 171.68385 <1e-12
   atanh(rhoDZ)
                      0.29927 0.01848 16.19766 <1e-12
                    Estimate 2.5%
                                  97.5%
 Correlation within MZ: 0.65394 0.62750 0.67888
 Correlation within DZ: 0.29064 0.25715 0.32344
 'log Lik.' -29020.35 (df=6)
 AIC: 58052.71
 BIC: 58093.76
```

A formal test of genetic effects can be obtained by comparing the MZ and DZ correlation:

```
estimate(1,contr(5:6,6))
                        Estimate Std.Err 2.5% 97.5% P-value
 [1@atanh(rhoMZ)] - [4.... 0.483 0.0418 0.401 0.565 6.4e-31
  Null Hypothesis:
  [1@atanh(rhoMZ)] - [4@atanh(rhoDZ)] = 0
1 <- twinlm(bmi ~ ns(age,1)+gender, data=twinbmi,</pre>
         DZ="DZ", zyg="zyg", id="tvparnr", type="cor", missing=TRUE)
summary(1)
       -----
Group 1
                      Estimate Std. Error Z value Pr(>|z|)
 Regressions:
   bmi.1~ns(age, 1).1 4.08914 0.16354 25.00328 <1e-12
   bmi.1~gendermale.1 1.41143 0.07285 19.37536 <1e-12
 Intercepts:
   bmi.1
                      22.57414 0.07187 314.08431 <1e-12
```

Additional Parameters:				
log(var)	2.44584	0.01425	171.68385	<1e-12
atanh(rhoMZ)	0.78216	0.02290	34.15832	<1e-12
Group 2				
	Estimate S	td. Error	Z value	Pr(> z)
Regressions:				
bmi.1~ns(age, 1).1	4.08914	0.16354	25.00328	<1e-12
bmi.1~gendermale.1	1.41143	0.07285	19.37536	<1e-12
Intercepts:				
bmi.1	22.57414	0.07187	314.08431	<1e-12
Additional Parameters:				
log(var)	2.44584	0.01425	171.68385	<1e-12
atanh(rhoDZ)	0.29927	0.01848	16.19766	<1e-12
	Estimate 2.	5% 97.	5%	
Correlation within MZ:	0.65394 0.62750 0.67888			
Correlation within DZ:	0.29064 0.	25715 0.32	2344	

'log Lik.' -29020.35 (df=6)

AIC: 58052.71 BIC: 58093.76

Twin analysis, censored outcomes

Twin analysis, binary traits

Time to event

References

- [1] J. v. Hjelmborg, C. Fagnani, K. Silventoinen, M. McGue, M. Korkeila, K. Christensen, A. Rissanen, and J. Kaprio. Genetic influences on growth traits of BMI: a longitudinal study of adult twins. Obesity (Silver Spring), 16(4):847–852, Apr 2008.
- [2] M. Korkeila, J. Kaprio, A. Rissanen, and M. Koskenvuo. Effects of gender and age on the heritability of body mass index. Int J Obes, 15(10):647-654, Oct 1991.