Suplementary Material 3: Code to Respond to Reviewers

Richard Meitern

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Introduction

The aim of this document is to respond to the reviewers comments. The comments are:

- 1) I would like to see whether indeed mothers are statistically more likely to stop having children after a twin birth (perhaps I missed that analysis and it was indeed presented)?
- 2) Directly test the odds that a mother terminate reproduction after a twin birth? In other words, use statistics to assess post-twinning reproductive outcomes compared to post-singleton reproductive outcomes.
- 3) Plot the real data as a cloud of points behind these lines- it is important to see the actual dataset in analyses like these.
- 4) Look at the analyses when limiting to opposite-sex offspring?
- 5) Based on analyses of Finnish data from 1953-1964 Eriksson and Fellman show that illegitimate maternities have a higher twinning rate than legitimate ones when maternal age and birth order are taken into account. They speculate that unmarried mothers are women who tend to become pregnant more easily than others owing to the effect of certain factors, e.g. a higher ovulation rate and to have relatively better physical qualifications for going through a twin pregnancy than mothers in general. It seems not unlikely that such mothers may remain single of the twin pregnancy, especially in earlier times?

```
#cleanup memory
gcstuff <- gc(verbose=FALSE); rm(gcstuff);</pre>
```

```
#get last birth adding function
source("./R/last_birth.R")

#simplified twinR summary tables
source("./R/twinR_summary.R")

#fix twinR compute predictions to do prediction with no lambda as well
source("./R/twinR_predictions.R")

#simple convenience functions
source("./R/utils.R")
```

```
## Identify number of CPU cores available for parallel computing,
## note: using a large number may lead RAM to max out, so you may have to adjust
## that according to your infrastructure:
```

```
nb_cores <- min(c(50L, parallel::detectCores() - 1))

## Set option in spaMM:
spaMM::spaMM.options(nb_cores = nb_cores)

## Registered S3 methods overwritten by 'registry':
## method from
## print.registry_field proxy
## print.registry_entry proxy</pre>
```

Data Import

The Estonian dataset has been formatted to include the same columns as the data_births_all dataset from the **twinR** package. The only difference is that the columns pop and monthly are excluded as these are constant.

```
#some labels
estLab <- "Estonian"
westLab <- "9 other European"</pre>
```

```
#Import and preprocess Estonian Data

data_births_monthly_EE <- readRDS("./data/data_births_all_EE.rds")

#the twink package expects population to be present

data_births_monthly_EE$pop <- "Estonia"

## Expand the birth level data for the fit of statistical models:
data_births_monthly_EE <- twink::expand_data(data_births_monthly_EE)

data_births_monthly_EE <- add_last_birth(data_births_monthly_EE)

data_births_monthly_EE_not_last <- data_births_monthly_EE[!data_births_monthly_EE$last,]

#make the aggregates

dmm_EE <- twink::aggregate_data(data_births_monthly_EE)

dmm_EE$prob_twin <- dmm_EE$twin_total / dmm_EE$births_total

dmm_EE_nl <- twink::aggregate_data(data_births_monthly_EE_not_last)

dmm_EE_nl $- twink::aggregate_data(data_births_monthly_EE_not_last)

dmm_EE_nl$prob_twin <- dmm_EE_nl$twin_total / dmm_EE_nl$births_total
```

```
##Import and pre-proccess twinR package data

## Filter the raw data to only keep data with monthly resolution:
data_births_monthly <- twinR::filter_data(twinR::data_births_all)

## Expand the birth level data for the fit of statistical models:
data_births_monthly <- twinR::expand_data(data_births_monthly)

data_births_monthly <- add_last_birth(data_births_monthly)
data_births_monthly_not_last <- data_births_monthly[!data_births_monthly$last,]</pre>
```

Fitting models

Comment 1: Logistic regression

We can test the hypothesis that mothers are statistically more likely to stop having children after a twin birth by fitting a logistic regression model with the response variable being the binary variable last and the predictor variable being the binary variable twin.

```
##
## Call:
## glm(formula = last ~ twin, family = binomial(link = "logit"),
##
      data = data_births_monthly_EE)
##
## Coefficients:
                 Estimate Std. Error
                                       z value
                                                  Pr(>|z|)
## (Intercept) -0.84910352  0.00340496 -249.3723 < 2.22e-16 ***
## twinTRUE
               0.35559579 0.02600496
                                       13.6741 < 2.22e-16 ***
## ---
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
##
## (Dispersion parameter for binomial family taken to be 1)
##
##
      Null deviance: 510562.5 on 417417 degrees of freedom
## Residual deviance: 510381.5 on 417416 degrees of freedom
## AIC: 510385.5
##
## Number of Fisher Scoring iterations: 4
```

```
#fit logistic regression Other European
logit_fit <- glm(last ~ twin,</pre>
                data = data births monthly,
                family = binomial(link = "logit"))
summary(logit_fit)
##
## Call:
## glm(formula = last ~ twin, family = binomial(link = "logit"),
##
       data = data_births_monthly)
## Coefficients:
##
                 Estimate Std. Error
                                         z value
                                                   Pr(>|z|)
## (Intercept) -1.38754400 0.00775251 -178.98007 < 2.22e-16 ***
## twinTRUE 0.45172685 0.05349239
                                         8.44469 < 2.22e-16 ***
## ---
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
##
## (Dispersion parameter for binomial family taken to be 1)
##
##
      Null deviance: 106259.4 on 105832 degrees of freedom
## Residual deviance: 106192.8 on 105831 degrees of freedom
```

Comment 2: Proportion of twins

Number of Fisher Scoring iterations: 4

AIC: 106196.8

##

The same test gives us the odds of mothers having more children after a twin birth.

```
# Extract coefficients
coef <- coef(logit_fit_EE)

# Calculate odds ratios
odds_ratios <- exp(coef)</pre>
```

For the Estonian dataset the odds of a birth event being the last one for a mother are 1.427 times higher for a twin birth than for a singleton birth.

```
# Extract coefficients
coef <- coef(logit_fit)

# Calculate odds ratios
odds_ratios <- exp(coef)</pre>
```

For the other dataset the odds of a birth event being the last one for a mother are 1.571 times higher for a twin birth than for a singleton birth.

```
#compare the precentage of twin births between last births and not last births
cat("Estonian dataset:\n")
```

Estonian dataset:

```
tble <- table(data_births_monthly_EE[c("twin", "last")])</pre>
tble
          last
##
## twin
            FALSE
                    TRUE
##
    FALSE 287874 123152
     TRUE
             3969
                    2423
cat("\n")
#now calcualte the % by hand
cat("Last twins: ", round(100 * tble[2,2] / sum(tble[,2]), 2), "%\n")
## Last twins: 1.93 %
cat("Prior twins: ",round(100 * tble[2,1] / sum(tble[,1]), 2), "%\n")
## Prior twins: 1.36 %
#compare the precentage of twin births between last births and not last births
cat("Western dataset:\n")
## Western dataset:
tblw <- table(data_births_monthly[c("twin", "last")])</pre>
tblw
##
          last
         FALSE TRUE
## twin
    FALSE 83276 20793
    TRUE
          1267
                  497
cat("\n")
#now calcualte the % by hand
cat("Last twins: ", round(100 * tblw[2,2] / sum(tblw[,2]), 2), "%\n")
## Last twins: 2.33 %
cat("Prior twins: ",round(100 * tblw[2,1] / sum(tblw[,1]), 2), "%\n")
## Prior twins: 1.5 %
# Combine the two tables into a 3-dimensional array
tbl \leftarrow array(c(tble, tblw), dim = c(dim(tble), 2))
#numeric problem workaround
tbl <- tbl / 1
# Perform the Mantel-Haenszel test
mantelhaen.test(tbl)
```

```
##
## Mantel-Haenszel chi-squared test with continuity correction
##
## data: tbl
## Mantel-Haenszel X-squared = 256.9326, df = 1, p-value < 2.22e-16
## alternative hypothesis: true common odds ratio is not equal to 1
## 95 percent confidence interval:
## 1.38715338 1.52045200
## sample estimates:
## common odds ratio
## 1.45227413</pre>
```

Comment 3: Original Data on Plots

The reviewers also asked to put the data on the Figure 1.

#use new base theme that displays also grid lines

source("./R/twinR theme.R")

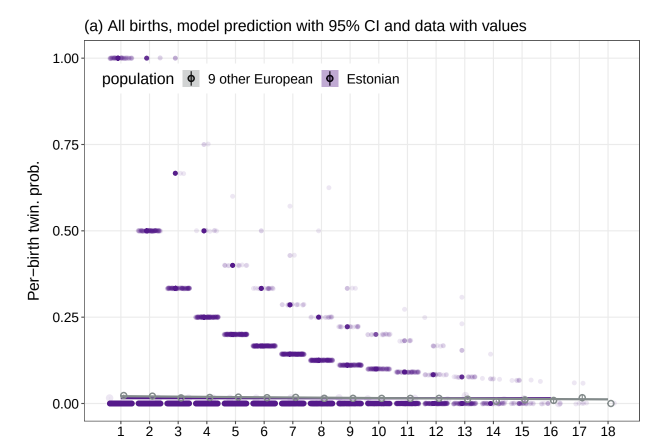
```
# import the function to do model fit and predictions
source("./R/fit_models.R")
## full data
## Estonia - mother level data
formula <- "cbind(twin total, singleton total) ~ 1 + births total"</pre>
dmm_EE_fit <- fitPredictions(dmm_EE, formula, predict = T)</pre>
## Warning in fitPredictions(dmm_EE, formula, predict = T): Pre-computed fit returned from file:
## ./data/predictions/cbind(twin_total,singleton_total)~1+births_totaldmm_EE_fit.rds
## If you want to re-run this step delete the file or change the saveDir!
## Warning in get_predictions(predDataFname, fit, dataset, args, save): Pre-computed predictions return
## ./data/predictions/cbind(twin_total,singleton_total)~1+births_totaldmm_EEdata_fig.rds
## If you want to re-run this step delete the file or change the saveDir!
## TwinR - mother level data
formula <- "cbind(twin_total, singleton_total) ~ 1 + births_total + (1|pop)"</pre>
dmm_orig_fit <- fitPredictions(dmm_orig, formula, predict=T)</pre>
## Warning in fitPredictions(dmm_orig, formula, predict = T): Pre-computed fit returned from file:
## ./data/predictions/cbind(twin_total,singleton_total)~1+births_total+(1_x_pop)dmm_orig_fit.rds
## If you want to re-run this step delete the file or change the saveDir!
## Warning in get_predictions(predDataFname, fit, dataset, args, save): Pre-computed predictions return
## ./data/predictions/cbind(twin_total,singleton_total)~1+births_total+(1_x_pop)dmm_origdata_fig.rds
## If you want to re-run this step delete the file or change the saveDir!
library(ggplot2)
#some nice colors
bc <- c("azure4", "purple4", "black", "navy", "darkgoldenrod2", "springgreen3", "gray")
```

generated.

```
fig2 EE plot data <- dmm EE fit$results</pre>
fig2_orig_plot_data <- dmm_orig_fit$results</pre>
infoTxt <- "model prediction with 95% CI and data with values"</pre>
fig2_ext_orig <- ggplot() +</pre>
  geom_line(data=fig2_EE_plot_data,
              aes(y = estimate, x=births_total, color=estLab), size = 1) +
  geom point(data=dmm EE[dmm EE$births total <17, ],</pre>
               aes(x=births_total, y=prob_twin, color=estLab, fill = estLab),
               alpha=0.1,
               position = position_nudge(x = -0.1)) +
    stat_summary(data=dmm_EE[dmm_EE$births_total <17, ],</pre>
                 aes(x=births_total, y=prob_twin, color=estLab, fill = estLab),
                 alpha=0.1,
                 position = position_jitter(),
                 fun.data=mean_se) +
    geom_ribbon(data=fig2_EE_plot_data,
                aes(y = estimate, x=births_total, ymin = lwr, ymax = upr,
                    color=estLab, fill = estLab),
                alpha = 0.3) +
    geom_line(data=fig2_orig_plot_data,
             aes(y = estimate, x=births_total, color=westLab), size = 1) +
  geom_point(data=dmm_orig[dmm_orig$births_total <19, ],</pre>
               aes(x=births total, y=prob twin, color=estLab, fill = estLab),
               alpha=0.1,
               position = position_jitter()) +
    stat_summary(data=dmm_orig[dmm_orig$births_total <19, ],</pre>
                 aes(x=births_total, y=prob_twin,
                     color=westLab, fill=westLab),
                 alpha=1,
                 shape = 1,
                 position = position_nudge(x = 0.1),
                 fun.data=mean_se) +
    geom_ribbon(data=fig2_orig_plot_data,
                aes(y = estimate, x=births_total, ymin = lwr, ymax = upr,
                    fill=westLab),
                alpha = 0.1) +
    ggplot2::scale_x_continuous(breaks = 1:18) +
    ggplot2::coord_cartesian() +
    labs(subtitle = paste0("(a) All births, ", infoTxt),
         y="Per-birth twin. prob.",
         x="Maternal total births")
## Warning: Using 'size' aesthetic for lines was deprecated in ggplot2 3.4.0.
## i Please use 'linewidth' instead.
## This warning is displayed once every 8 hours.
## Call 'lifecycle::last_lifecycle_warnings()' to see where this warning was
```

```
p2 <- fig2_ext_orig + base_theme(larger=8) + scale_color_manual(values=bc) +
    scale_fill_manual(values=bc) + guides(color="none") + labs(fill = "population")
p2</pre>
```

Warning: Removed 1 rows containing missing values ('geom_segment()').



The first plot clearly illustrates that we cannot plot the whole data itself on the plot, as there are women who only gave birth to twins. We can plot a subset (e.g. those with at least 10 children) but this would lose the point of the plot. There is just so much data that is semi-categorical that we cannot think of a way put into a plot so that it gives additional information.

Maternal total births

Comment 4: Dizygotic Twins

We can do the same analysis for dizygotic twins.

```
#re-import the data
dbm_EE <- readRDS("./data/data_births_all_EE.rds")

#take out mothers that had twins of single sex
single_sex_twin_mothers <- as.character(unlist(unique(
   dbm_EE[dbm_EE$twin & dbm_EE$cf_sex != "mixed", "maternal_id"])
))</pre>
```

```
#the twinR package expects population to be present
dbmEE wo single sex twins$pop <- "Estonia"</pre>
## Expand the birth level data for the fit of statistical models:
dbmEE_wo_single_sex_twins <- twinR::expand_data(dbmEE_wo_single_sex_twins)</pre>
dbmEE_wo_single_sex_twins <- add_last_birth(dbmEE_wo_single_sex_twins)</pre>
dbmEE_wo_single_sex_twins_not_last <- dbmEE_wo_single_sex_twins[!dbmEE_wo_single_sex_twins$last,]
#make the aggregates
dmm_EE_mixed <- twinR::aggregate_data(dbmEE_wo_single_sex_twins)</pre>
dmm_EE_mixed$prob_twin <- dmm_EE_mixed$twin_total / dmm_EE_mixed$births_total
dmm_EE_mixed_nl <- twinR::aggregate_data(dbmEE_wo_single_sex_twins_not_last)</pre>
dmm_EE_mixed_nl$prob_twin <- dmm_EE_mixed_nl$twin_total / dmm_EE_mixed_nl$births_total
Full Data
options(scipen = 99)
## Estonia
formula <- "cbind(twin_total, singleton_total) ~ 1 + births_total"</pre>
dmm EE mixed fit <- fitPredictions(dmm EE mixed, formula)
## Warning in fitPredictions(dmm_EE_mixed, formula): Pre-computed fit returned from file:
## ./data/predictions/cbind(twin_total,singleton_total)~1+births_totaldmm_EE_mixed_fit.rds
## If you want to re-run this step delete the file or change the saveDir!
## Warning in get_predictions(predDataFname, fit, dataset, args, save): Pre-computed predictions return
## ./data/predictions/cbind(twin_total,singleton_total)~1+births_totaldmm_EE_mixeddata_fig.rds
## If you want to re-run this step delete the file or change the saveDir!
knitr::kable(build_fit_summary.table(dmm_EE_mixed_fit$fit))
## Warning: Use of .data in tidyselect expressions was deprecated in tidyselect 1.2.0.
## i Please use '"Estimate" instead of '.data$Estimate'
## This warning is displayed once every 8 hours.
## Call 'lifecycle::last_lifecycle_warnings()' to see where this warning was
## generated.
## Warning: Use of .data in tidyselect expressions was deprecated in tidyselect 1.2.0.
## i Please use '"object" instead of '.data$object'
## This warning is displayed once every 8 hours.
## Call 'lifecycle::last lifecycle warnings()' to see where this warning was
## generated.
```

single_sex_twin_mothers),]

dbmEE_wo_single_sex_twins <- dbm_EE[!(dbm_EE\$maternal_id %in%</pre>

```
## Warning: Use of .data in tidyselect expressions was deprecated in tidyselect 1.2.0.
## i Please use '"name"' instead of '.data$name'
## This warning is displayed once every 8 hours.
## Call 'lifecycle::last_lifecycle_warnings()' to see where this warning was
## generated.

## Warning: Use of .data in tidyselect expressions was deprecated in tidyselect 1.2.0.
## i Please use '"value"' instead of '.data$value'
## This warning is displayed once every 8 hours.
## Call 'lifecycle::last_lifecycle_warnings()' to see where this warning was
## generated.
```

| Type | Variable | Value | Cond. SE | t-value |
|-----------------|-----------------------------------|---------|----------|---------|
| fixed effects | (Intercept) | -5.07 | 0.044 | -115 |
| | births_total | -0.0146 | 0.00843 | -1.73 |
| response family | binomial with logit link | | | |
| fit info | number of model parameters | 2 | | |
| | marginal log Likelihood | -11192 | | |
| | marginal AIC | 22388 | | |
| | conditional AIC (cAIC) | | | |
| data info | number of fitted observations (N) | 121779 | | |

No Last Births Data

```
## Estonia
formula <- "cbind(twin_total, singleton_total) ~ 1 + births_total"
dmm_EE_mixed_nl_fit <- fitPredictions(dmm_EE_mixed_nl, formula)</pre>
```

```
## Warning in fitPredictions(dmm_EE_mixed_nl, formula): Pre-computed fit returned from file:
## ./data/predictions/cbind(twin_total,singleton_total)~1+births_totaldmm_EE_mixed_nl_fit.rds
## If you want to re-run this step delete the file or change the saveDir!
```

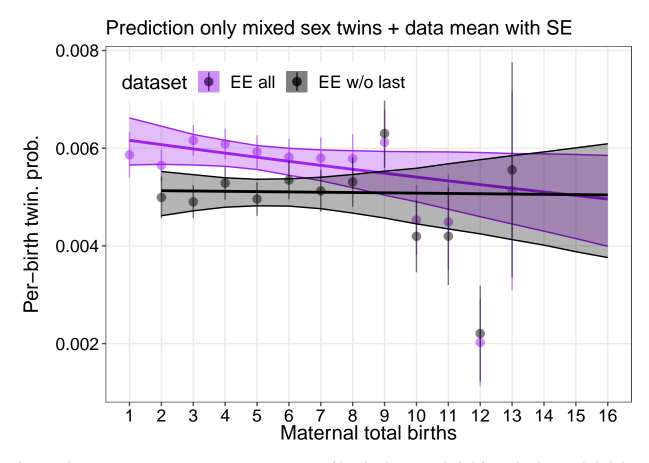
Warning in get_predictions(predDataFname, fit, dataset, args, save): Pre-computed predictions return
./data/predictions/cbind(twin_total,singleton_total)~1+births_totaldmm_EE_mixed_nldata_fig.rds
If you want to re-run this step delete the file or change the saveDir!

knitr::kable(build_fit_summary.table(dmm_EE_mixed_nl_fit\$fit))

| Type | Variable | Value | Cond. SE | t-value |
|-----------------|-----------------------------------|----------|----------|---------|
| fixed effects | (Intercept) | -5.27 | 0.0549 | -96 |
| | births_total | -0.00123 | 0.0112 | -0.11 |
| response family | binomial with logit link | | | |
| fit info | number of model parameters | 2 | | |
| | marginal log Likelihood | -7104 | | |
| | marginal AIC | 14212 | | |
| | conditional AIC (cAIC) | | | |
| data info | number of fitted observations (N) | 94662 | | |

Plot the Predictions

```
fig2_EE_plot_data <- dmm_EE_mixed_fit$results</pre>
fig2_EE_plot_data_nl <- dmm_EE_mixed_nl_fit$results</pre>
fig2_EE_plot_data_nl$births_total <- fig2_EE_plot_data_nl$births_total + 1</pre>
dmm_EE_mixed_nl_plot <- dmm_EE_mixed_nl</pre>
dmm EE mixed nl plot$births total <- dmm EE mixed nl plot$births total + 1
#colors
bc <- c("purple", "black")</pre>
fig2_ext_orig <- ggplot() +
  geom line(data=fig2 EE plot data,
              aes(y = estimate, x=births total, color="EE all"),
            linewidth = 1) +
    stat_summary(data=dmm_EE_mixed[dmm_EE_mixed$births_total <14, ],</pre>
                 aes(x=births_total, y=prob_twin, color="EE all",
                     fill = "EE all"), alpha=0.5,
                 fun.data=mean_se) +
    geom_ribbon(data=fig2_EE_plot_data,
                aes(y = estimate, x=births_total,
                    ymin = lwr, ymax = upr, color="EE all", fill = "EE all"),
                alpha = 0.3) +
    geom_line(data=fig2_EE_plot_data_nl,
              aes(y = estimate, x=births_total, color="EE w/o last"),
              linewidth = 1) +
    stat_summary(data=dmm_EE_mixed_nl_plot[dmm_EE_mixed_nl_plot$births_total <14,],</pre>
                 aes(x=births_total, y=prob_twin, color="EE w/o last",
                     fill = "EE w/o last"), alpha=0.5,
                 fun.data=mean_se) +
    geom_ribbon(data=fig2_EE_plot_data_nl,
                aes(y = estimate, x=births_total, ymin = lwr,
                    ymax = upr, color="EE w/o last", fill = "EE w/o last"),
                alpha = 0.3) +
    ggplot2::scale_x_continuous(breaks = 1:18) +
    ggplot2::scale_y_continuous(breaks = seq(0,0.03, by=0.002)) +
    ggplot2::coord_cartesian() +
    labs(subtitle = "Prediction only mixed sex twins + data mean with SE",
         y="Per-birth twin. prob.",
         x="Maternal total births")
p2 <- fig2_ext_orig + base_theme(larger=8) + scale_color_manual(values=bc) +
  scale_fill_manual(values=bc) + guides(color="none") + labs(fill = "dataset")
p2
```



The mixed sex twinning rate seem to remain constant if last births are excluded from the data and slightly decline if last births are included. The decline seems not to be significant.

Comment 5: Illegitimate Maternities

We can add a predictor to the model describing if the women had first birth or first conception before being married. We will do both to investigate if illegitimate maternities affect twinning rate.

```
cat("First birth before marriage:\n")

## First birth before marriage:
table(dmm_EE$illigimate_birth)

##

## FALSE TRUE
## 108483 17092

cat("First conception before marriage:\n")

## First conception before marriage:
table(dmm_EE$illigimate_conception)

##

## FALSE TRUE
## 89896 35679
With All Data
```

```
## Estonia
formula <- "cbind(twin_total, singleton_total) ~ 1 + births_total + illigimate_birth"
dmm_EE_fit_birth <- fitPredictions(dmm_EE, formula)</pre>
```

```
## Warning in fitPredictions(dmm_EE, formula): Pre-computed fit returned from file:
## ./data/predictions/cbind(twin_total,singleton_total)~1+births_total+illigimate_birthdmm_EE_fit.rds
## If you want to re-run this step delete the file or change the saveDir!
```

Warning in get_predictions(predDataFname, fit, dataset, args, save): Pre-computed predictions return
./data/predictions/cbind(twin_total,singleton_total)~1+births_total+illigimate_birthdmm_EEdata_fig.r
If you want to re-run this step delete the file or change the saveDir!

knitr::kable(build_fit_summary.table(dmm_EE_fit_birth\$fit))

| Type | Variable | Value | Cond. SE | t-value |
|-----------------|-----------------------------------|----------|----------|---------|
| fixed effects | (Intercept) | -4.14 | 0.0276 | -150 |
| | births_total | -0.00255 | 0.00504 | -0.506 |
| | illigimate_birthTRUE | -0.0759 | 0.0425 | -1.79 |
| response family | binomial with logit link | | | |
| fit info | number of model parameters | 3 | | |
| | marginal log Likelihood | -24445 | | |
| | marginal AIC | 48895 | | |
| | conditional AIC (cAIC) | | | |
| data info | number of fitted observations (N) | 125575 | | |

| Type Variable | Value | Cond. SE | t-value | |
|---------------|-------|----------|---------|--|
|---------------|-------|----------|---------|--|

Estonia

formula <- "cbind(twin_total, singleton_total) ~ 1 + births_total + illigimate_conception"
dmm_EE_fit_conception <- fitPredictions(dmm_EE, formula)</pre>

Warning in fitPredictions(dmm_EE, formula): Pre-computed fit returned from file:

./data/predictions/cbind(twin_total,singleton_total)~1+births_total+illigimate_conceptiondmm_EE_fit.

If you want to re-run this step delete the file or change the saveDir!

Warning in get_predictions(predDataFname, fit, dataset, args, save): Pre-computed predictions return

 $\verb| ## ./data/predictions/cbind(twin_total, singleton_total) \verb| ~1+births_total+illigimate_conceptiondmm_EE data_independent of the conception of the conce$

If you want to re-run this step delete the file or change the saveDir!

knitr::kable(build_fit_summary.table(dmm_EE_fit_conception\$fit))

| Type | Variable | Value | Cond. SE | t-value |
|-----------------|-----------------------------------|----------|----------|---------|
| fixed effects | (Intercept) | -4.15 | 0.0278 | -149 |
| | births_total | -0.00178 | 0.00502 | -0.356 |
| | illigimate_conceptionTRUE | -0.00944 | 0.0283 | -0.334 |
| response family | binomial with logit link | | | |
| fit info | number of model parameters | 3 | | |
| | marginal log Likelihood | -24446 | | |
| | marginal AIC | 48898 | | |
| | conditional AIC (cAIC) | | | |
| data info | number of fitted observations (N) | 125575 | | |

No Last Births Data

Estonia

formula <- "cbind(twin_total, singleton_total) ~ 1 + births_total + illigimate_birth"
dmm_EE_nl_fit_birth <- fitPredictions(dmm_EE_nl, formula)</pre>

Warning in fitPredictions(dmm_EE_nl, formula): Pre-computed fit returned from file:

If you want to re-run this step delete the file or change the saveDir!

Warning in get_predictions(predDataFname, fit, dataset, args, save): Pre-computed predictions return

./data/predictions/cbind(twin_total,singleton_total)~1+births_total+illigimate_birthdmm_EE_nldata_fi
If you want to re-run this step delete the file or change the saveDir!

knitr::kable(build_fit_summary.table(dmm_EE_nl_fit_birth\$fit))

| Type | Variable | Value | Cond. SE | t-value |
|---------------|-------------|-------|----------|---------|
| fixed effects | (Intercept) | -4.37 | 0.0338 | -129 |

| Type | Variable | Value | Cond. SE | t-value |
|-----------------|-----------------------------------|--------|----------|---------|
| | births_total | 0.018 | 0.00655 | 2.74 |
| | illigimate_birthTRUE | 0.0364 | 0.0547 | 0.665 |
| response family | binomial with logit link | | | |
| fit info | number of model parameters | 3 | | |
| | marginal log Likelihood | -15893 | | |
| | marginal AIC | 31792 | | |
| | conditional AIC (cAIC) | | | |
| data info | number of fitted observations (N) | 98183 | | |

```
## Estonia
formula <- "cbind(twin_total, singleton_total) ~ 1 + births_total + illigimate_conception"
dmm_EE_fit_nl_conception <- fitPredictions(dmm_EE_nl, formula)</pre>
```

```
## Warning in fitPredictions(dmm_EE_nl, formula): Pre-computed fit returned from file:
## ./data/predictions/cbind(twin_total,singleton_total)~1+births_total+illigimate_conceptiondmm_EE_nl_f
## If you want to re-run this step delete the file or change the saveDir!
```

Warning in get_predictions(predDataFname, fit, dataset, args, save): Pre-computed predictions return
./data/predictions/cbind(twin_total,singleton_total)~1+births_total+illigimate_conceptiondmm_EE_nlda

If you want to re-run this step delete the file or change the saveDir!

knitr::kable(build_fit_summary.table(dmm_EE_fit_nl_conception\$fit))

| Type | Variable | Value | Cond. SE | t-value |
|-----------------|-----------------------------------|--------|----------|---------|
| fixed effects | (Intercept) | -4.37 | 0.0344 | -127 |
| | births_total | 0.0174 | 0.00656 | 2.65 |
| | illigimate_conceptionTRUE | 0.0443 | 0.0355 | 1.25 |
| response family | binomial with logit link | | | |
| fit info | number of model parameters | 3 | | |
| | marginal log Likelihood | -15892 | | |
| | marginal AIC | 31790 | | |
| | conditional AIC (cAIC) | | | |
| data info | number of fitted observations (N) | 98183 | | |

Figures for Estonian mothers with and without Last Birth

dmm_EE_nl_plot\$births_total <- dmm_EE_nl_plot\$births_total + 1</pre>

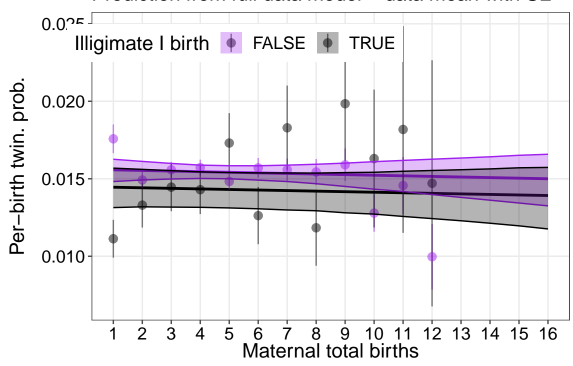
dmm_EE_nl_plot <- dmm_EE_nl</pre>

```
fig2_EE_plot_data <- dmm_EE_fit_birth$results
fig2_EE_plot_data_nl <- dmm_EE_nl_fit_birth$results
fig2_EE_plot_data_nl$births_total <- fig2_EE_plot_data_nl$births_total + 1</pre>
```

```
#colors
bc <- c("purple", "black")</pre>
fig2_ext_orig <- ggplot() +</pre>
  geom_line(data=fig2_EE_plot_data,
              aes(y = estimate, x=births_total, color=illigimate_birth),
            linewidth = 1) +
    stat_summary(data=dmm_EE[dmm_EE$births_total <13, ],</pre>
                 aes(x=births_total, y=prob_twin, color=illigimate_birth,
                     fill = illigimate_birth), alpha=0.5,
                 fun.data=mean_se) +
    geom_ribbon(data=fig2_EE_plot_data,
                aes(y = estimate, x=births_total, ymin = lwr, ymax = upr,
                    color= illigimate_birth, fill = illigimate_birth),
                alpha = 0.3) +
    ggplot2::scale_x_continuous(breaks = 1:18) +
    ggplot2::scale_y_continuous(breaks = seq(0,0.03, by=0.005)) +
    ggplot2::coord_cartesian() +
    labs(subtitle = "Prediction from full data model + data mean with SE",
         y="Per-birth twin. prob.",
         x="Maternal total births")
p2 <- fig2_ext_orig + base_theme(larger=8) + scale_color_manual(values=bc) +</pre>
  scale_fill_manual(values=bc) + guides(color="none") +
  labs(fill = "Illigimate I birth")
cat("\n")
```

p2

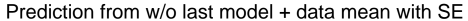
Prediction from full data model + data mean with SE

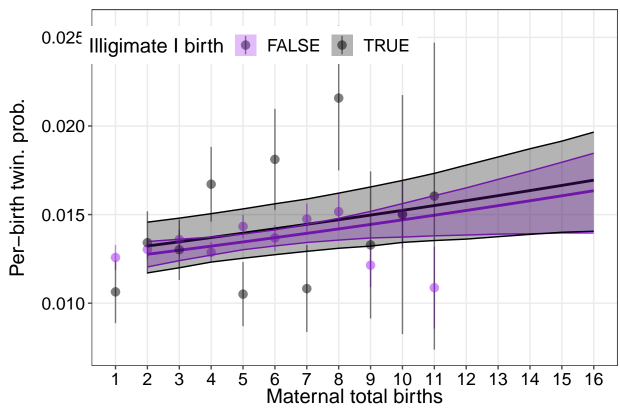


Illigimate Birth

```
cat("\n")
```

```
#colors
bc <- c("purple", "black")</pre>
fig2_ext_orig <- ggplot() +</pre>
  geom_line(data=fig2_EE_plot_data_nl,
              aes(y = estimate, x=births_total, color=illigimate_birth),
            linewidth = 1) +
    stat_summary(data=dmm_EE_nl[dmm_EE_nl$births_total <12, ],</pre>
                 aes(x=births_total, y=prob_twin, color=illigimate_birth,
                     fill = illigimate_birth), alpha=0.5,
                 fun.data=mean se) +
    geom_ribbon(data=fig2_EE_plot_data_nl,
                aes(y = estimate, x=births_total, ymin = lwr, ymax = upr,
                    color= illigimate_birth, fill = illigimate_birth),
                alpha = 0.3) +
    ggplot2::scale_x_continuous(breaks = 1:18) +
    ggplot2::scale_y_continuous(breaks = seq(0,0.03, by=0.005)) +
    ggplot2::coord_cartesian() +
    labs(subtitle = "Prediction from w/o last model + data mean with SE",
         y="Per-birth twin. prob.",
         x="Maternal total births")
p2 <- fig2_ext_orig + base_theme(larger=8) + scale_color_manual(values=bc) +
  scale_fill_manual(values=bc) + guides(color="none") +
  labs(fill = "Illigimate I birth")
cat("\n")
```





cat("\n")

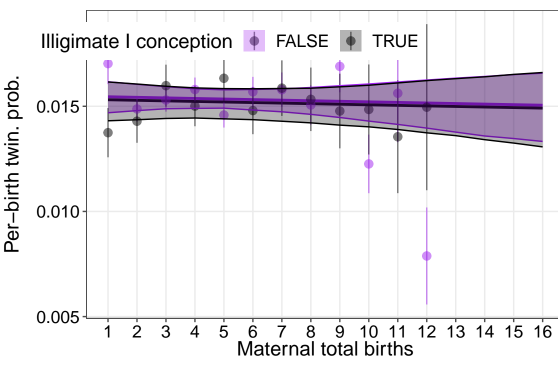
```
fig2_EE_plot_data <- dmm_EE_fit_conception$results
fig2_EE_plot_data_nl <- dmm_EE_fit_nl_conception$results
fig2_EE_plot_data_nl$births_total <- fig2_EE_plot_data_nl$births_total + 1</pre>
```

```
dmm_EE_nl_plot <- dmm_EE_nl
dmm_EE_nl_plot$births_total <- dmm_EE_nl_plot$births_total + 1</pre>
```

```
aes(x=births_total, y=prob_twin, color=illigimate_conception,
                     fill = illigimate_conception), alpha=0.5,
                 fun.data=mean_se) +
    geom_ribbon(data=fig2_EE_plot_data,
                aes(y = estimate, x=births_total, ymin = lwr, ymax = upr,
                    color= illigimate_conception, fill = illigimate_conception),
                alpha = 0.3) +
   ggplot2::scale_x_continuous(breaks = 1:18) +
   ggplot2::scale_y_continuous(breaks = seq(0,0.03, by=0.005)) +
   ggplot2::coord_cartesian() +
   labs(subtitle = "Prediction from full data model + data mean with SE",
         y="Per-birth twin. prob.",
         x="Maternal total births")
p2 <- fig2_ext_orig + base_theme(larger=8) + scale_color_manual(values=bc) +
  scale_fill_manual(values=bc) + guides(color="none") +
  labs(fill = "Illigimate I conception")
cat("\n")
```

p2

Prediction from full data model + data mean with SE



Illigimate conception

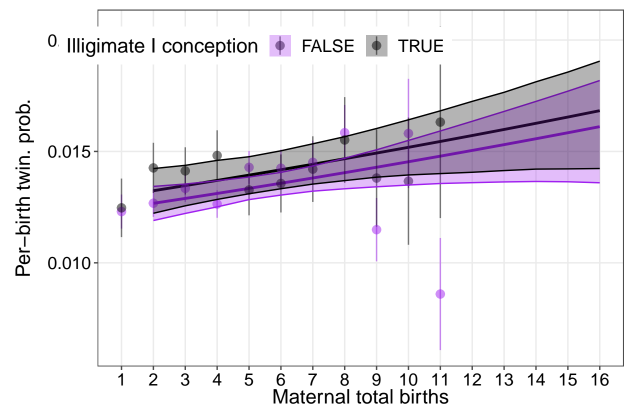
```
cat("\n")

#colors
bc <- c("purple", "black")
fig2_ext_orig <- ggplot() +
   geom_line(data=fig2_EE_plot_data_nl,</pre>
```

```
aes(y = estimate, x=births_total, color=illigimate_conception),
            linewidth = 1) +
    stat_summary(data=dmm_EE_nl[dmm_EE_nl$births_total <12, ],</pre>
                 aes(x=births_total, y=prob_twin, color=illigimate_conception,
                     fill = illigimate_conception), alpha=0.5,
                 fun.data=mean se) +
    geom_ribbon(data=fig2_EE_plot_data_nl,
                aes(y = estimate, x=births_total, ymin = lwr, ymax = upr,
                    color= illigimate_conception, fill = illigimate_conception),
                alpha = 0.3) +
   ggplot2::scale_x_continuous(breaks = 1:18) +
   ggplot2::scale_y_continuous(breaks = seq(0,0.03, by=0.005)) +
    ggplot2::coord_cartesian() +
   labs(subtitle = "Prediction from w/o last model + data mean with SE",
         y="Per-birth twin. prob.",
         x="Maternal total births")
p2 <- fig2_ext_orig + base_theme(larger=8) + scale_color_manual(values=bc) +
  scale_fill_manual(values=bc) + guides(color="none") +
  labs(fill = "Illigimate I conception")
cat("\n")
```

p2

Prediction from w/o last model + data mean with SE



```
cat("\n")
```

#END