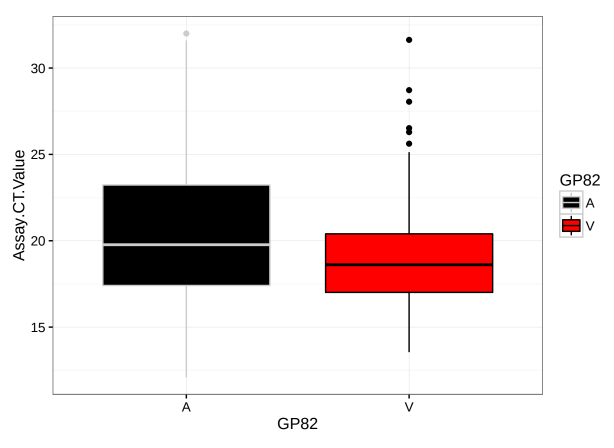
Ebola_GP82_analysis.r

max

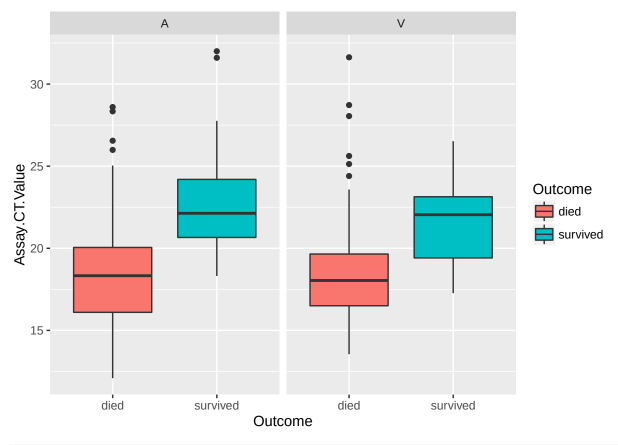
Sun Sep 18 17:08:38 2016

```
library(lme4)
## Loading required package: Matrix
library(ggplot2)
source("diehl_aux.r")
## Data preparation
raw <- read.csv("../data/GP82_raw_data.csv", header = TRUE) ## "all" data
EVD_GP82 <- read.csv("../data/EVD_case_fatality_GP82.csv", header = TRUE) ## clean data</pre>
coord_data <- read.csv("../data/coord_data_locations.csv", header = TRUE)</pre>
pop_level <- read.csv("../data/population_level.csv", header = TRUE)</pre>
EVD_GP82 <- merge(EVD_GP82, coord_data, by = "location")
EVD_GP82 <- merge(EVD_GP82, pop_level, by = "location")
EVD_GP82 <- subset(EVD_GP82, country.y == "GIN") ## restrict attention to Guinea
xtabs(~ Outcome + GP82, EVD_GP82)
            GP82
##
             A V
## Outcome
##
    died
             45 93
##
    survived 23 18
nrow(EVD_GP82)
## [1] 179
######
### Exploratory statistical analysis (ESA)
## Question 1: Do Ct values according to EBOV variant ?
# All data available
all_A <- raw$Assay.CT.Value[raw$GP82 == "A"]
sum(!is.na(all_A)) ## valid observations
## [1] 97
all_V <- raw$Assay.CT.Value[raw$GP82 == "V"]
sum(!is.na(all_V))
## [1] 216
```

```
t.test(all_A, all_V)
##
## Welch Two Sample t-test
## data: all_A and all_V
## t = 0.69147, df = 157.85, p-value = 0.4903
\#\# alternative hypothesis: true difference in means is not equal to 0
## 95 percent confidence interval:
## -0.6943069 1.4423291
## sample estimates:
## mean of x mean of y
## 20.05938 19.68537
# Just the data we'll use for further analyses
filtered_A <- EVD_GP82$Assay.CT.Value[EVD_GP82$GP82 == "A"]
sum(!is.na(filtered_A))
## [1] 68
filtered_V <- EVD_GP82$Assay.CT.Value[EVD_GP82$GP82 == "V"]</pre>
sum(!is.na(filtered_V))
## [1] 111
t.test(filtered_A, filtered_V)
##
## Welch Two Sample t-test
##
## data: filtered_A and filtered_V
## t = 1.8356, df = 112.02, p-value = 0.06907
\#\# alternative hypothesis: true difference in means is not equal to 0
## 95 percent confidence interval:
## -0.09118178 2.38791734
## sample estimates:
## mean of x mean of y
## 20.17765 19.02928
#######
ggplot(EVD_GP82) +
  geom_boxplot(aes(x = GP82, y = Assay.CT.Value, fill = GP82), colour = c("grey80", "black")) +
  scale_fill_manual(values = c("black", "red")) +
 theme_bw()
```



```
ggplot(EVD_GP82) +
  geom_boxplot(aes(x = Outcome, y = Assay.CT.Value, fill = Outcome))+
  facet_grid(~GP82)
```



```
#### Now let's look at odds ratios
(CrossTab_gp82 <- xtabs(~ GP82 + Outcome , EVD_GP82) )</pre>
```

```
## Outcome
## GP82 died survived
## A 45 23
## V 93 18
```

summary(CrossTab_gp82)

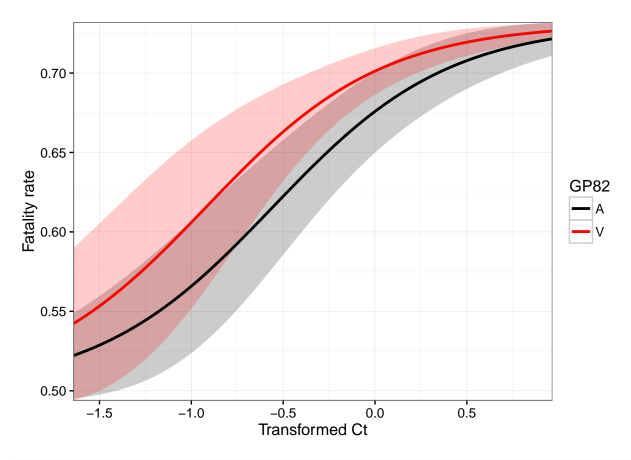
```
## Call: xtabs(formula = ~GP82 + Outcome, data = EVD_GP82)
## Number of cases in table: 179
## Number of factors: 2
## Test for independence of all factors:
## Chisq = 7.403, df = 1, p-value = 0.006511

a <- CrossTab_gp82["V", "died"]
b <- CrossTab_gp82["V", "survived"]
c <- CrossTab_gp82["A", "died"]
d <- CrossTab_gp82["A", "survived"]
SqrT <- sqrt(1/a + 1/b + 1/c + 1/d)
(rawOR <- (a*d)/(b*c))</pre>
```

[1] 2.640741

```
alpha <- .95
( lwrOR <- exp(log(rawOR) + qnorm((1 - alpha)/2) * SqrT) )
## [1] 1.295563
(uprOR \leftarrow exp(log(rawOR) + qnorm((1 + alpha)/2) * SqrT))
## [1] 5.382609
## Relative risk
m.RR \leftarrow log({a/(a+b)} / {c/(c + d)})
sdRR \leftarrow sqrt((1/a + 1/c) - (1/(a + b) + 1/(c+d)))
Z \leftarrow qnorm(.975)
CI.RR <- c(exp(m.RR - Z*sdRR), exp(m.RR + Z*sdRR))
c(exp(m.RR), CI.RR)
## [1] 1.266066 1.048446 1.528856
### Ok, but what happens if we try to correct for other variables?
### First, some data preparation
forFit <- EVD_GP82</pre>
forFit$Assay.CT.Value <- -standz(EVD_GP82$Assay.CT.Value) ## invert sign because of lower CT = higher v
forFit$pdensMN <- standz(EVD_GP82$pdensMN)</pre>
forFit$geconMN <- standz(EVD_GP82$geconMN)</pre>
forFit$tt50kMN <- standz(EVD_GP82$tt50kMN)</pre>
forFit$cumCases <- standz(log(EVD_GP82$cumCases))</pre>
forFit$GP82 <- ifelse(EVD_GP82$GP82 == "V", 1, 0) ## GP82-A as baseline
forFit$Outcome <- ifelse(EVD_GP82$Outcome == "died", 1 , 0)</pre>
simple_glm <- glm(Outcome ~ Assay.CT.Value + GP82, data = forFit, family = "binomial")</pre>
summary(simple_glm)
##
## Call:
## glm(formula = Outcome ~ Assay.CT.Value + GP82, family = "binomial",
       data = forFit)
##
## Deviance Residuals:
       Min 1Q Median
                                    3Q
                                            Max
## -2.2077 0.2806
                     0.4394
                                         1.8405
                                0.6210
##
## Coefficients:
                  Estimate Std. Error z value Pr(>|z|)
## (Intercept)
                    1.0189
                                0.3046 3.345 0.000824 ***
                                0.4308
                                         4.734 2.2e-06 ***
## Assay.CT.Value
                    2.0395
## GP82
                    0.7400
                                0.4028
                                        1.837 0.066199 .
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for binomial family taken to be 1)
##
```

```
Null deviance: 192.65 on 178 degrees of freedom
## Residual deviance: 156.94 on 176 degrees of freedom
## AIC: 162.94
##
## Number of Fisher Scoring iterations: 4
getOdds(simple_glm, hier = FALSE)
## Waiting for profiling to be done...
                               2.5 %
                                         97.5 %
##
                       est.
## (Intercept)
                  2.770133 1.5597542 5.187539
## Assay.CT.Value 7.686444 3.4580497 18.918033
## GP82
                  2.095999 0.9492025 4.647809
## (corrected) relative risks
simple.pois.glm <- glm(Outcome ~ Assay.CT.Value + GP82, data = forFit, family = "poisson")
exp(cbind(coefficients(simple.pois.glm), confint(simple.pois.glm)))
## Waiting for profiling to be done...
                                2.5 %
                                        97.5 %
##
## (Intercept)
                  0.665769 0.4891752 0.8805533
## Assay.CT.Value 1.654133 1.1447092 2.4290820
## GP82
                  1.196355 0.8425497 1.7240292
NewCt <- seq(min(forFit$Assay.CT.Value), max(forFit$Assay.CT.Value), length.out = 100)</pre>
mut <-c(0, 1)
newDat <- expand.grid(NewCt, mut)</pre>
names(newDat) <- c("Assay.CT.Value", "GP82")</pre>
Pred.logit <- predict(simple_glm, newdata = newDat, se.fit = TRUE, type = "response")
invlogit <- function(x) exp(x)/(1 + exp(x))
alpha <- 0.95
InvLogitPred <- data.frame(lwr = invlogit(Pred.logit$fit + qnorm((1-alpha)/2) * Pred.logit$se.fit) ,</pre>
                           mean = invlogit(Pred.logit$fit),
                           upr = invlogit(Pred.logit$fit + qnorm((1 + alpha)/2) * Pred.logit$se.fit))
toPlot_preds <- data.frame(newDat, InvLogitPred)</pre>
toPlot_preds$GP82 <- ifelse(toPlot_preds$GP82 == 1, "V", "A")
ggplot(toPlot_preds, aes(x = Assay.CT.Value, y = mean)) +
  # ggtitle("Predicted fatality rates") +
  geom_ribbon(aes(ymin = lwr, ymax = upr, fill = GP82), alpha = .2) +
  scale_x_continuous("Transformed Ct", expand = c(0, 0)) +
  scale_y_continuous("Fatality rate", expand = c(0, 0)) +
  scale_fill_manual(values = c("black", "red")) +
  geom_line(aes(colour = GP82), size = 1) +
  scale_color_manual(values = c("black", "red")) +
  guides(fill = FALSE) +
  theme_bw()
```



```
## Generalized linear mixed model fit by maximum likelihood (Laplace
     Approximation) [glmerMod]
   Family: binomial (logit)
## Formula: Outcome ~ Assay.CT.Value + GP82 + (cumCases | location)
##
     Data: forFit
##
##
                       logLik deviance df.resid
       AIC
                 BIC
      167.5
                       -77.8
##
               186.7
                                 155.5
                                            173
##
## Scaled residuals:
##
      Min
                1Q Median
                                3Q
                                       Max
## -3.5246 0.1808 0.2943 0.4408 2.1712
##
## Random effects:
                         Variance Std.Dev. Corr
   Groups
           Name
##
##
   location (Intercept) 0.2539
                                0.5039
##
             cumCases
                         0.4845
                                  0.6961
                                           1.00
## Number of obs: 179, groups: location, 17
##
## Fixed effects:
                  Estimate Std. Error z value Pr(>|z|)
##
## (Intercept)
                    0.9824
                               0.3862
                                        2.544
                                              0.0110 *
```

```
## Assay.CT.Value 2.1619
                               0.4935
                                       4.381 1.18e-05 ***
                   0.7973
                               0.4561 1.748 0.0804 .
## GP82
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Correlation of Fixed Effects:
               (Intr) A.CT.V
## Assay.CT.Vl 0.252
## GP82
              -0.668 -0.078
getOdds(model_Of)
## Computing profile confidence intervals ...
##
                       est
                               2.5 %
                                       97.5 %
## (Intercept)
                 2.670811 1.1461364 5.921091
## Assay.CT.Value 8.687964 3.5723089 25.581957
## GP82
                 2.219620 0.9042745 5.671575
RR <- ranef(model_Of, condVar = TRUE)</pre>
names(RR) <- " "
names(RR[[1]]) <- c("Location-specific intercept", "coefficient of cumulative number of cases")</pre>
lattice::dotplot(RR)
```

\$` `

