# NDVTransmissionTrial

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# Short summary

The transmission parameter  $\beta$  (unit =  $day^{-1}$ ), reproduction number R (dimensionless) and the infectious period (unit = day) for vaccinated and unvaccinated chickens was estimated. The vaccination reduced the infectious period and transmission parameter substantially and thus also the reproduction number. The reproduction number was, however, not smaller than 1. This indicates that the vaccination will reduce transmission speed, but cannot prevent an outbreak NDV in this setting. Additional measures such as compartimentalization can contribute to further reduction of spread and thus lower the reproduction number.

# Introduction

- Background trial
- Background transmission models

# Objectives

- Estimation of infectious period for vaccinated and unvaccinated
- Estimation of transmission parameter beta for vaccinated and unvaccinated
- Estimation of R0 for vaccinated and unvaccinated

### Material and methods

Four different assumptions were used to estimate the transmission parameter. An animal is positive if:

- 1. one or both samples are positive
- 2. both samples are positive
- 3. the ON sample is positive
- 4. the CLO sample is positive

For each of these assumptions we estimated the infectious period, the transmission parameter  $\beta$  the basic reproduction number  $R_0$ . The infectious period was estimated assuming that these are normally distributed. The transmission parameter  $\beta$  is estimated using the generalized linear model with a complementary loglog-link function:

```
glm(cbind(C, S-C) \sim 1, offset = log(I/N), family = binomial(link = "cloglog"), data = out.nona, na.action = na.omit)
```

The basic reproduction number  $R_0$  is than estimated by multiplying the  $\beta$  with the mean infectious period. The confidence interval is calculated by  $\log(R) = \log(\beta) + \log(T_{inf}) \pm Z_{0.05} \cdot (\log(SE(\beta)) + \log(SE(T_{inf}))$ , which thus assumes independence between the transmission parameter and the infectious period.

# Results

# Load and organize data

The data sets consist of two separate data files in 'xlsx' format. These files need to be combined to form one data set

with variables

#### names (cevadata) "Sample" [1] "Group" "Vaccinated" "Challenge" "bird.id" [6] "1.dpch" "2.dpch" "3.dpch" "4.dpch" "5.dpch" "7.dpch" "10.dpch" [11] "6.dpch" "8.dpch" "9.dpch" "NP.ELISA" [16] "11.dpch" "12.dpch" "13.dpch" "14.dpch"

Define a function reform.data to rearrange the data for analyses. The function takes the data and use the names of the sampledays, determine positivity based on one or all positive samples, and define the cutt-off value for positivity.

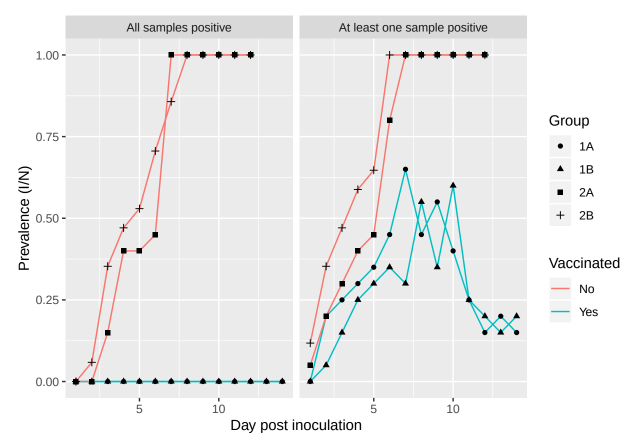
# Descriptive data

# Epidemic curves

```
epicurveplot
```

```
## Warning: Removed 4 rows containing missing values (geom_path).
```

## Warning: Removed 8 rows containing missing values (geom\_point).

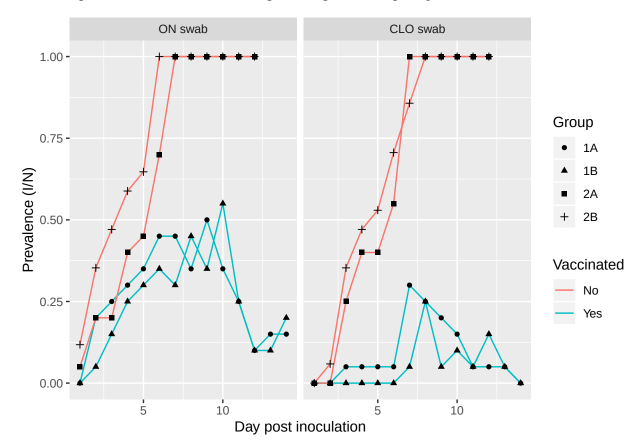


In some stable all chickens died, so that I/N was "NA" causing a message that these were not plotted. The vaccinated groups are found all to be positive in both swaps (CLO swab and NO swab), while non of the unvaccinated were positive.

The separate swabs represent the following epidemic curves:

```
epicurveplot
```

- ## Warning: Removed 4 rows containing missing values (geom\_path).
- ## Warning: Removed 8 rows containing missing values (geom\_point).



On first sight the curves are similar for the unvaccinated groups, but the CLO swab are less likely and later positive than the ON swab in the vaccinated groups.

# Transmission parameters

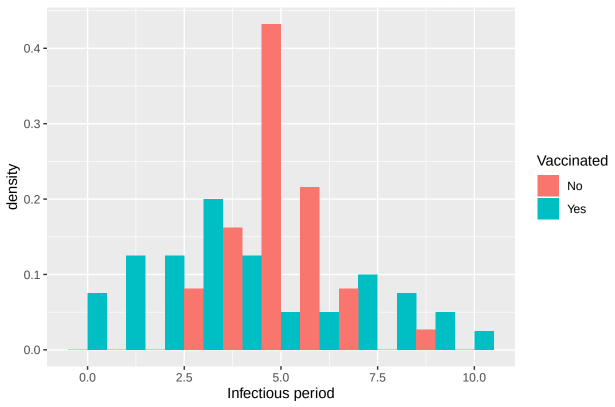
At the end of this section all results will be summarized.

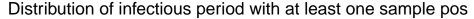
```
all.results <- NULL
```

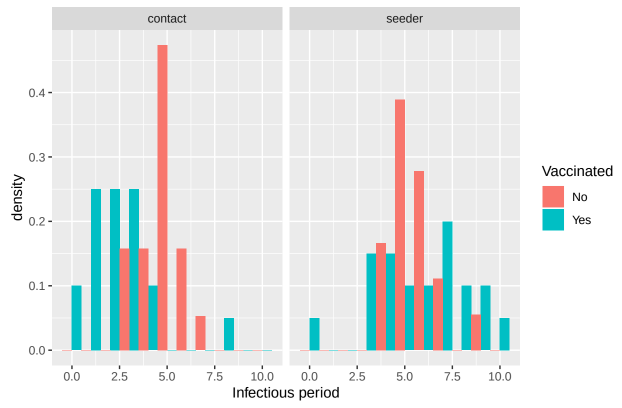
### One positive sample

### Infectious period

# Distribution of infectious period with at least one sample pos







Descriptive statistics for the infectious period.

### table.infper

```
## Group Mean SD
## 1 Overall 4.571429 2.232703
## 2 Unvaccinated 5.162162 1.213661
## 3 Vaccinated 4.025000 2.778004
```

From the table it already seems that the variances are not equal. Therefor we first test for equality of variance and than do the appropriate test to compare means.

```
#test for equal variances
fligner.test(out.indiv$infper, out.indiv$Vaccinated)

##
## Fligner-Killeen test of homogeneity of variances
##
## data: out.indiv$infper and out.indiv$Vaccinated
## Fligner-Killeen:med chi-squared = 13.666, df = 1, p-value = 0.0002184

#*Compare the mean infectious period
t.test(out.indiv$infper~out.indiv$Vaccinated,var.equal = FALSE)

##
## Welch Two Sample t-test
##
## data: out.indiv$infper by out.indiv$Vaccinated
## t = 2.3571, df = 54.253, p-value = 0.02205
```

```
## alternative hypothesis: true difference in means is not equal to 0
## 95 percent confidence interval:
## 0.170043 2.104281
## sample estimates:
##
   mean in group No mean in group Yes
            5.162162
                              4.025000
##
#Compare the sd infectious period
var.test(out.indiv$infper~out.indiv$Vaccinated)
##
##
   F test to compare two variances
##
## data: out.indiv$infper by out.indiv$Vaccinated
## F = 0.19087, num df = 36, denom df = 39, p-value = 2.188e-06
## alternative hypothesis: true ratio of variances is not equal to 1
## 95 percent confidence interval:
## 0.1000171 0.3673606
## sample estimates:
## ratio of variances
            0.1908662
##
```

The vaccinated group has a lower mean infectious period, but a larger variance.

Separate Seeder and contact birds Descriptive statistics for the infectious period.

#### table.infper

```
##
     Vaccination Challenge
                              Mean
                 Overall 4.571429 2.232703
## 1
        Overall
## 2
             No
                 contact 4.789474 1.084176
## 3
            Yes contact 2.300000 1.780006
## 4
             No
                   seeder 5.555556 1.247219
## 5
            Yes
                   seeder 5.750000 2.531382
```

From the table it already seems that the variances are not equal. Therefor we first test for equality of variance and than do the appropriate test to compare means.

```
#test for equal variances
#vaccinated vs unvaccinated
fligner.test(out.indiv$infper, out.indiv$Vaccinated)

##
## Fligner-Killeen test of homogeneity of variances
##
## data: out.indiv$infper and out.indiv$Vaccinated
## Fligner-Killeen:med chi-squared = 13.666, df = 1, p-value = 0.0002184

#seeder vs contact
fligner.test(out.indiv$infper, out.indiv$Challenge)

##
## Fligner-Killeen test of homogeneity of variances
##
## data: out.indiv$infper and out.indiv$Challenge
##
## Fligner-Killeen test of homogeneity of variances
##
## data: out.indiv$infper and out.indiv$Challenge
## Fligner-Killeen:med chi-squared = 0.085416, df = 1, p-value = 0.7701
```

```
#all four groups
fligner.test(out.indiv$infper, mapply(paste0,out.indiv$Challenge,out.indiv$Vaccinated))
##
##
  Fligner-Killeen test of homogeneity of variances
##
## data: out.indiv$infper and mapply(paste0, out.indiv$Challenge, out.indiv$Vaccinated)
## Fligner-Killeen:med chi-squared = 14.32, df = 3, p-value = 0.002501
Variances between vaccine groups are not equal. Between challenge groups there are no differences.
#Compare the mean infectious period
"vaccinated versus unvaccinated"
## [1] "vaccinated versus unvaccinated"
t.test(out.indiv$infper~out.indiv$Vaccinated,var.equal = FALSE)
   Welch Two Sample t-test
##
## data: out.indiv$infper by out.indiv$Vaccinated
## t = 2.3571, df = 54.253, p-value = 0.02205
## alternative hypothesis: true difference in means is not equal to 0
## 95 percent confidence interval:
## 0.170043 2.104281
## sample estimates:
## mean in group No mean in group Yes
##
            5.162162
                              4.025000
"seeder versus contact'
## [1] "seeder versus contact"
t.test(out.indiv$infper~out.indiv$Challenge,var.equal = FALSE)
##
## Welch Two Sample t-test
##
## data: out.indiv$infper by out.indiv$Challenge
## t = -4.781, df = 74.703, p-value = 8.542e-06
## alternative hypothesis: true difference in means is not equal to 0
## 95 percent confidence interval:
## -3.038918 -1.251231
## sample estimates:
## mean in group contact mean in group seeder
                3.512821
"vaccinated versus unvaccinated in contacts"
## [1] "vaccinated versus unvaccinated in contacts"
t.test(infper~Vaccinated ,
       data = out.indiv[out.indiv$Challenge == "contact", ],
       var.equal = FALSE)
##
   Welch Two Sample t-test
##
```

```
## data: infper by Vaccinated
## t = 5.3041, df = 31.643, p-value = 8.467e-06
## alternative hypothesis: true difference in means is not equal to 0
## 95 percent confidence interval:
## 1.533023 3.445924
## sample estimates:
## mean in group No mean in group Yes
            4.789474
                              2.300000
"vaccinated versus unvaccinated in seeders"
## [1] "vaccinated versus unvaccinated in seeders"
t.test(infper~Vaccinated ,
       data = out.indiv[out.indiv$Challenge == "seeder", ],
       var.equal = FALSE)
##
##
   Welch Two Sample t-test
## data: infper by Vaccinated
## t = -0.30486, df = 28.329, p-value = 0.7627
## alternative hypothesis: true difference in means is not equal to 0
## 95 percent confidence interval:
## -1.500277 1.111389
## sample estimates:
## mean in group No mean in group Yes
            5.55556
                              5.750000
#Compare the sd infectious period
var.test(out.indiv$infper~out.indiv$Vaccinated)
##
##
   F test to compare two variances
## data: out.indiv$infper by out.indiv$Vaccinated
## F = 0.19087, num df = 36, denom df = 39, p-value = 2.188e-06
## alternative hypothesis: true ratio of variances is not equal to 1
## 95 percent confidence interval:
## 0.1000171 0.3673606
## sample estimates:
## ratio of variances
##
            0 1908662
```

The vaccinated group has a lower mean infectious period, but a larger variance. The difference is not found in the seeder group, but is due to differences in the contact group.

#### Transmission parameters $\beta$

## Single term deletions

```
##
## Model:
## cbind(out.nona$C, out.nona$S - out.nona$C) ~ out.nona$Vaccinated +
##
       out.nona$Group
                       Df Deviance
                                       AIC
## <none>
                            81.620 168.00
## out.nona$Vaccinated 0
                            81.620 168.00
## out.nona$Group
                        2
                            85.748 168.13
In backward selection Group falls out, but vaccination stays in.
fit.empty <- glm(cbind(out.nona$C, out.nona$S-out.nona$C) ~1,
                 offset = log(out.nona$I/out.nona$N),
                 family = binomial(link = "cloglog"),
                 data = out.nona,
                 na.action = na.omit)
add1(fit.empty, ~. + out.nona$Vaccinated + out.nona$Group)
## Single term additions
##
## Model:
## cbind(out.nona$C, out.nona$S - out.nona$C) ~ 1
                       Df Deviance
## <none>
                            91.283 171.66
## out.nona$Vaccinated 1
                            85.748 168.13
## out.nona$Group
                        3
                            81.620 168.00
fit.vac <- glm(cbind(out.nona$C, out.nona$C-out.nona$C) ~out.nona$Vaccinated,
               offset = log(out.nona$I/out.nona$N),
               family = binomial(link = "cloglog"),
               data = out.nona, na.action = na.omit)
add1(fit.vac, scope = ~.+na.omit(out.nona$Group))
## Single term additions
##
## Model:
## cbind(out.nona$C, out.nona$S - out.nona$C) ~ out.nona$Vaccinated
                           Df Deviance
                                           AIC
## <none>
                                85.748 168.13
## na.omit(out.nona$Group) 2
                                81.620 168.00
fit.group <- glm(cbind(out.nona$C, out.nona$S-out.nona$C) ~out.nona$Group,
                 offset = log(out.nona$I/out.nona$N),
                 family = binomial(link = "cloglog"),
                 data = out.nona, na.action = na.omit)
add1(fit.group, ~.+out.nona$Vaccinated)
## Single term additions
##
## cbind(out.nona$C, out.nona$S - out.nona$C) ~ out.nona$Group
                       Df Deviance AIC
##
## <none>
                             81.62 168
## out.nona$Vaccinated 0
                             81.62 168
```

In forward selection both groups and vaccination are added as single terms to improve the empty model, but the addition of Group in the vaccine model or vaccination in the group model shows that Group should not be included in the model.

summary(fit.vac)

```
##
## Call:
## glm(formula = cbind(out.nona$C, out.nona$S - out.nona$C) ~ out.nona$Vaccinated,
       family = binomial(link = "cloglog"), data = out.nona, na.action = na.omit,
       offset = log(out.nona$I/out.nona$N))
##
##
## Deviance Residuals:
##
       Min
                 10
                      Median
                                   30
                                           Max
## -2.5373 -1.1985 -0.3804
                               1.0257
                                        3.2453
##
## Coefficients:
##
                          Estimate Std. Error z value Pr(>|z|)
## (Intercept)
                            0.1185
                                       0.1708
                                                0.694
## out.nona$VaccinatedYes -0.5181
                                       0.2146 - 2.414
                                                         0.0158 *
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for binomial family taken to be 1)
##
##
       Null deviance: 91.283 on 36 degrees of freedom
## Residual deviance: 85.748 on 35 degrees of freedom
## AIC: 168.13
## Number of Fisher Scoring iterations: 5
Vaccination lowers the infection parameter \beta.
beta.table
##
                             97.5
          beta
                    2.5%
## 1 1.1257870 0.7899902 1.547889
## 2 0.6705952 0.4431067 1.030285
Basic reproduction number R_0
mean.RO[, c(1,2,3,4)]
               Vaccinated
                               beta
                                       llbeta
                       No 1.1257870 0.8893607 1.7425939
## (Intercept)
                      Yes 0.6705952 0.2639445 0.6137075
mean.R0[, c(1,5,6,7)]
##
               Vaccinated
                            infper llinfper ulinfper
## (Intercept)
                       No 4.789474 4.301977 5.276970
                      Yes 2.300000 1.519892 3.080108
mean.R0[, c(1,11,12,13)]
               Vaccinated
                                 R
                       No 5.251570 3.3702032 8.183183
## (Intercept)
                      Yes 1.248118 0.5977554 2.606082
```

Based on the assumption of one of both samples required to be positive, we find that the transmission

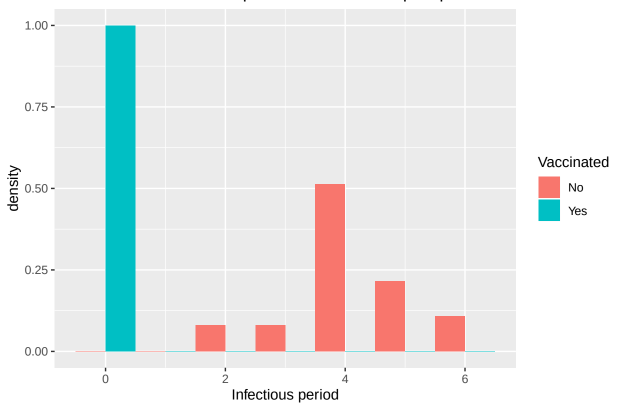
parameter  $\beta$  and the mean infectious period are lower, which results in a reproduction number R which is more than 4 times smaller, but is not lower than the threshold  $R \leq 1$ .

For all other options I have until now only used the overall infectious period!!!

## Both positive samples

### Infectious period

# Distribution of infectious period with both samples pos



Descriptive statistics for the infectious period.

# table.infper

```
## Group Mean SD
## 1 Overall 2.012987 2.221270
## 2 Unvaccinated 4.189189 1.023009
## 3 Vaccinated 0.000000 0.000000
```

Using the assumption that both swabs need a positive result, the vaccinated group is never infectious.

#### Transmission parameters $\beta$

Because there are no chickens that are set to infectious based on the criterium that both need to be positive, parameters can only be estimated for the unvaccinated group.

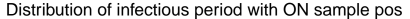
```
#test qlm
fit.full <- glm(cbind(out.nona$C, out.nona$S-out.nona$C) ~ out.nona$Group,
                offset = log(out.nona$I/out.nona$N),
                family = binomial(link = "cloglog"),
                data = out.nona, na.action = na.omit)
drop1(fit.full)
## Single term deletions
##
## Model:
## cbind(out.nona$C, out.nona$S - out.nona$C) ~ out.nona$Group
                  Df Deviance
##
                                 ATC
                       54.449 75.570
## <none>
## out.nona$Group 1
                       54.574 73.695
In backward selection again Group falls out.
fit.empty <- glm(cbind(out.nona$C, out.nona$S-out.nona$C) ~1,
                 offset = log(out.nona$I/out.nona$N),
                 family = binomial(link = "cloglog"),
                 data = out.nona,
                 na.action = na.omit)
add1(fit.empty, ~. + out.nona$Group)
## Single term additions
##
## cbind(out.nona$C, out.nona$S - out.nona$C) ~ 1
##
                  Df Deviance
                                 AIC
                       54.574 73.695
## <none>
## out.nona$Group 1
                      54.449 75.570
Also in forward selection, Group should not be included in the model.
summary(fit.empty)
##
## Call:
## glm(formula = cbind(out.nona$C, out.nona$S - out.nona$C) ~ 1,
       family = binomial(link = "cloglog"), data = out.nona, na.action = na.omit,
       offset = log(out.nona$I/out.nona$N))
##
##
## Deviance Residuals:
##
                                    3Q
       Min
                 1Q
                      Median
                                            Max
## -3.3345 -1.8171 -0.1445
                              1.3824
                                         4.4520
##
## Coefficients:
##
               Estimate Std. Error z value Pr(>|z|)
                 0.1469
                            0.1780
                                     0.825
## (Intercept)
##
## (Dispersion parameter for binomial family taken to be 1)
##
##
       Null deviance: 54.574 on 9 degrees of freedom
```

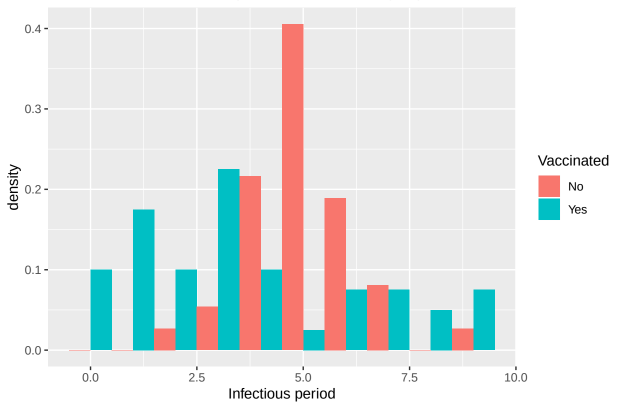
```
## AIC: 73.695
##
## Number of Fisher Scoring iterations: 5
Vaccination lowers the infection parameter \beta.
beta.table
                    2.5%
                             97.5
         beta
## 1 1.158205 0.8038566 1.606461
Basic reproduction number R_0
mean.R0[1, c(1,2,3,4)]
     Vaccinated
                             llbeta
                                      ulbeta
                    beta
## 1
             No 1.158205 0.9310305 1.860611
mean.R0[1, c(1,5,6,7)]
##
     Vaccinated infper llinfper ulinfper
             No 4.189189 3.859559 4.518819
## 1
mean.R0[1, c(1,11,12,13)]
     Vaccinated
                               11R
             No 4.691349 3.028026 7.268349
## 1
```

## Residual deviance: 54.574 on 9 degrees of freedom

# ON sample positive

### Infectious period





Descriptive statistics for the infectious period.

### table.infper

```
## Group Mean SD
## 1 Overall 4.311688 2.272604
## 2 Unvaccinated 5.054054 1.289831
## 3 Vaccinated 3.625000 2.742706
```

From the table it already seems that the variances are not equal. Therefor we first test for equality of variance and than do the appropriate test to compare means.

```
#test for equal variances
fligner.test(out.indiv$infper, out.indiv$Vaccinated) #conclude unequal variances

##
## Fligner-Killeen test of homogeneity of variances

##
## data: out.indiv$infper and out.indiv$Vaccinated
## Fligner-Killeen:med chi-squared = 12.14, df = 1, p-value = 0.0004936

#Compare the mean infectious period
t.test(out.indiv$infper~out.indiv$Vaccinated,var.equal = FALSE)

##
## Welch Two Sample t-test
##
## data: out.indiv$infper by out.indiv$Vaccinated
## t = 2.9604, df = 56.387, p-value = 0.004488
```

```
## alternative hypothesis: true difference in means is not equal to 0
## 95 percent confidence interval:
## 0.4621832 2.3959249
## sample estimates:
## mean in group No mean in group Yes
##
            5.054054
                              3.625000
#Compare the sd infectious period
var.test(out.indiv$infper~out.indiv$Vaccinated)
##
  F test to compare two variances
##
## data: out.indiv$infper by out.indiv$Vaccinated
## F = 0.22116, num df = 36, denom df = 39, p-value = 1.358e-05
## alternative hypothesis: true ratio of variances is not equal to 1
## 95 percent confidence interval:
## 0.1158917 0.4256676
## sample estimates:
## ratio of variances
##
            0.2211602
The vaccinated group has a lower mean infectious period, but a larger variance.
Transmission parameters \beta
#test qlm
fit.full <- glm(cbind(out.nona$C, out.nona$S-out.nona$C) ~ out.nona$Vaccinated +out.nona$Group,
                offset = log(out.nona$I/out.nona$N),
                family = binomial(link = "cloglog"),
                data = out.nona, na.action = na.omit)
drop1(fit.full)
## Single term deletions
##
## cbind(out.nona$C, out.nona$S - out.nona$C) ~ out.nona$Vaccinated +
##
       out.nona$Group
##
                       Df Deviance
                                      ATC:
## <none>
                            79.134 165.19
## out.nona$Vaccinated 0
                            79.134 165.19
## out.nona$Group
                            83.397 165.46
                        2
fit.vac <- glm(cbind(out.nona$C, out.nona$S-out.nona$C) ~ out.nona$Vaccinated,
               offset = log(out.nona$I/out.nona$N),
               family = binomial(link = "cloglog"),
               data = out.nona, na.action = na.omit)
drop1(fit.vac)
## Single term deletions
##
## Model:
## cbind(out.nona$C, out.nona$S - out.nona$C) ~ out.nona$Vaccinated
                       Df Deviance
```

83.397 165.46

## out.nona\$Vaccinated 1 91.087 171.15

## <none>

```
fit.group <- glm(cbind(out.nona$C, out.nona$S-out.nona$C) ~ out.nona$Group,
                 offset = log(out.nona$I/out.nona$N),
                 family = binomial(link = "cloglog"),
                 data = out.nona, na.action = na.omit)
drop1(fit.group)
## Single term deletions
##
## Model:
## cbind(out.nona$C, out.nona$S - out.nona$C) ~ out.nona$Group
                  Df Deviance
                                 AIC
## <none>
                       79.134 165.19
## out.nona$Group 3
                     91.087 171.15
In backward selection either group or vaccination should stay in the selection.
fit.empty <- glm(cbind(out.nona$C, out.nona$S-out.nona$C) ~1,
                 offset = log(out.nona$I/out.nona$N),
                 family = binomial(link = "cloglog"),
                 data = out.nona, na.action = na.omit)
add1(fit.empty, ~. + out.nona$Vaccinated + out.nona$Group)
## Single term additions
##
## Model:
## cbind(out.nona$C, out.nona$S - out.nona$C) ~ 1
                       Df Deviance
                                     AIC
                            91.087 171.15
## <none>
## out.nona$Vaccinated 1
                            83.397 165.46
## out.nona$Group
                        3 79.134 165.19
fit.vac <- glm(cbind(out.nona$C, out.nona$C-out.nona$C) ~out.nona$Vaccinated,
               offset = log(out.nona$I/out.nona$N),
               family = binomial(link = "cloglog"),
               data = out.nona, na.action = na.omit)
add1(fit.vac, scope = ~.+na.omit(out.nona$Group))
## Single term additions
##
## Model:
## cbind(out.nona$C, out.nona$S - out.nona$C) ~ out.nona$Vaccinated
##
                           Df Deviance
                                          AIC
                                83.397 165.46
## <none>
## na.omit(out.nona$Group) 2
                               79.134 165.19
fit.group <- glm(cbind(out.nona$C, out.nona$S-out.nona$C) ~out.nona$Group,
                 offset = log(out.nona$I/out.nona$N),
                 family = binomial(link = "cloglog"),
                 data = out.nona,
                 na.action = na.omit)
add1(fit.group, ~.+out.nona$Vaccinated)
## Single term additions
##
## Model:
## cbind(out.nona$C, out.nona$S - out.nona$C) ~ out.nona$Group
```

In forward selection either groups or vaccination are added as single terms to improve the empty model. We choose to use vaccination as this is the main interest.

```
summary(fit.vac)
```

```
##
## Call:
  glm(formula = cbind(out.nona$C, out.nona$S - out.nona$C) ~ out.nona$Vaccinated,
       family = binomial(link = "cloglog"), data = out.nona, na.action = na.omit,
##
##
       offset = log(out.nona$I/out.nona$N))
##
## Deviance Residuals:
                     Median
##
      Min
                 1Q
                                   30
                                           Max
## -2.4911 -1.1414 -0.4031
                               1.4409
                                        3.2962
##
## Coefficients:
##
                          Estimate Std. Error z value Pr(>|z|)
## (Intercept)
                            0.1487
                                       0.1686
                                                0.882 0.37758
## out.nona$VaccinatedYes -0.6159
                                       0.2165 -2.844 0.00445 **
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
  (Dispersion parameter for binomial family taken to be 1)
##
##
       Null deviance: 91.087
                             on 36
                                    degrees of freedom
## Residual deviance: 83.397
                             on 35 degrees of freedom
## AIC: 165.46
##
## Number of Fisher Scoring iterations: 4
```

Vaccination lowers the infection parameter  $\beta$ .

#### beta.table

```
## Vaccination beta 2.5% 97.5
## 1 Yes 1.1603785 0.8184697 1.5888716
## 2 No 0.6268027 0.4122417 0.9646928
```

The transmission parameter  $\beta$  is lower for the vaccinated group.

### Basic reproduction number $R_0$

```
mean.R0[, c(1,2,3,4)]
                                        llbeta
##
               Vaccinated
                                beta
                                                  ulbeta
                       No 1.1603785 0.9497346 1.843692
## (Intercept)
##
                      Yes 0.6268027 0.2226810 0.521099
mean.RO[, c(1,5,6,7)]
                             infper llinfper ulinfper
               Vaccinated
## (Intercept)
                       No 5.054054 4.638450 5.469658
                      Yes 3.625000 2.775042 4.474958
##
```

```
mean.R0[, c(1,11,12,13)]

## Vaccinated R 11R ulR

## (Intercept) No 5.670181 3.7331690 8.61224

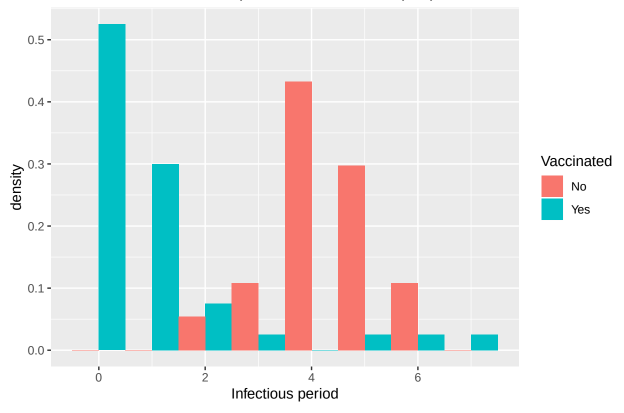
## Yes 1.661995 0.8244941 3.35021
```

Based on the assumption of ON samples required to be positive, we find that the transmission parameter  $\beta$  and the mean infectious period are lower, which results in a reproduction number R which is less than one thirth, but is not lower than the threshold  $R \leq 1$ .

# CLO sample positive

### Infectious period

# Distribution of infectious period with CLO sample pos



Descriptive statistics for the infectious period.

### table.infper

```
## Group Mean SD
## 1 Overall 2.571429 2.1547290
## 2 Unvaccinated 4.297297 0.9962392
```

#### ## 3 Vaccinated 0.975000 1.6406300

From the table it already seems that the variances are not equal. Therefor we first test for equality of variance and than do the appropriate test to compare means.

```
#test for equal variances
fligner.test(out.indiv$infper, out.indiv$Vaccinated)
##
##
   Fligner-Killeen test of homogeneity of variances
## data: out.indiv$infper and out.indiv$Vaccinated
## Fligner-Killeen:med chi-squared = 0.0073488, df = 1, p-value = 0.9317
#Compare the mean infectious period
t.test(out.indiv$infper~out.indiv$Vaccinated,var.equal = F)
##
   Welch Two Sample t-test
##
##
## data: out.indiv$infper by out.indiv$Vaccinated
## t = 10.829, df = 65.086, p-value = 3.346e-16
## alternative hypothesis: true difference in means is not equal to 0
## 95 percent confidence interval:
## 2.709625 3.934970
## sample estimates:
   mean in group No mean in group Yes
            4.297297
                              0.975000
t.test(out.indiv$infper~out.indiv$Vaccinated,var.equal = T)
##
   Two Sample t-test
##
## data: out.indiv$infper by out.indiv$Vaccinated
## t = 10.634, df = 75, p-value < 2.2e-16
## alternative hypothesis: true difference in means is not equal to 0
## 95 percent confidence interval:
## 2.699926 3.944669
## sample estimates:
## mean in group No mean in group Yes
##
            4.297297
                              0.975000
```

The vaccinated group has a lower mean infectious period, but no difference in variance found.

### Transmission parameters $\beta$

```
## Model:
## cbind(out.nona$C, out.nona$S - out.nona$C) ~ out.nona$Vaccinated +
       out.nona$Group
##
                       Df Deviance
                                      ATC:
## <none>
                            87.014 132.33
## out.nona$Vaccinated 0
                          87.014 132.33
## out.nona$Group
                           87.043 128.36
                        2
fit.vac <- glm(cbind(out.nona$C, out.nona$S-out.nona$C) ~ out.nona$Vaccinated,
               offset = log(out.nona$I/out.nona$N),
               family = binomial(link = "cloglog"),
               data = out.nona,
               na.action = na.omit)
drop1(fit.vac)
## Single term deletions
##
## Model:
## cbind(out.nona$C, out.nona$S - out.nona$C) ~ out.nona$Vaccinated
                       Df Deviance
##
                                      AIC
                            87.043 128.36
## <none>
## out.nona$Vaccinated 1 90.679 130.00
fit.group <- glm(cbind(out.nona$C, out.nona$S-out.nona$C) ~ out.nona$Group,</pre>
                 offset = log(out.nona$I/out.nona$N),
                 family = binomial(link = "cloglog"),
                 data = out.nona,
                 na.action = na.omit)
drop1(fit.group)
## Single term deletions
##
## Model:
## cbind(out.nona$C, out.nona$S - out.nona$C) ~ out.nona$Group
                  Df Deviance
                                 AIC
                       87.014 132.33
## <none>
                       90.679 130.00
## out.nona$Group 3
In backward selection either group or vaccination should stay in the selection.
fit.empty <- glm(cbind(out.nona$C, out.nona$S-out.nona$C) ~1,
                 offset = log(out.nona$I/out.nona$N),
                 family = binomial(link = "cloglog"),
                 data = out.nona,
                 na.action = na.omit)
add1(fit.empty, ~. + out.nona$Vaccinated + out.nona$Group)
## Single term additions
##
## Model:
## cbind(out.nona$C, out.nona$S - out.nona$C) ~ 1
##
                       Df Deviance
                                      ATC
                            90.679 130.00
## out.nona$Vaccinated 1 87.043 128.36
## out.nona$Group 3 87.014 132.33
```

```
fit.vac <- glm(cbind(out.nona$C, out.nona$C-out.nona$C) ~out.nona$Vaccinated,
               offset = log(out.nona$I/out.nona$N),
               family = binomial(link = "cloglog"),
               data = out.nona,
               na.action = na.omit)
add1(fit.vac, scope = ~.+na.omit(out.nona$Group))
## Single term additions
##
## Model:
## cbind(out.nona$C, out.nona$S - out.nona$C) ~ out.nona$Vaccinated
##
                           Df Deviance
                                           AIC
                                87.043 128.36
## <none>
## na.omit(out.nona$Group) 2
                                87.014 132.33
fit.group <- glm(cbind(out.nona$C, out.nona$S-out.nona$C) ~out.nona$Group,
                 offset = log(out.nona$I/out.nona$N),
                 family = binomial(link = "cloglog"),
                 data = out.nona,
                 na.action = na.omit)
add1(fit.group, ~.+out.nona$Vaccinated)
## Single term additions
## Model:
## cbind(out.nona$C, out.nona$S - out.nona$C) ~ out.nona$Group
##
                       Df Deviance
                                       ATC:
                            87.014 132.33
## out.nona$Vaccinated 0
                            87.014 132.33
In forward selection either groups or vaccination are added as single terms to improve the empty model. We
choose to use vaccination as this is the main interest.
summary(fit.vac)
##
## Call:
## glm(formula = cbind(out.nona$C, out.nona$S - out.nona$C) ~ out.nona$Vaccinated,
       family = binomial(link = "cloglog"), data = out.nona, na.action = na.omit,
       offset = log(out.nona$I/out.nona$N))
##
##
## Deviance Residuals:
                      Median
##
       Min
                                    3Q
                 1Q
                                            Max
## -3.1887 -1.0778 -0.8148
                               0.4695
                                         3.8361
##
## Coefficients:
##
                          Estimate Std. Error z value Pr(>|z|)
## (Intercept)
                           0.05745
                                      0.18170 0.316
                                                         0.7519
## out.nona$VaccinatedYes -0.54945
                                      0.29381 -1.870
                                                         0.0615 .
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for binomial family taken to be 1)
##
##
       Null deviance: 90.679 on 27 degrees of freedom
```

## Residual deviance: 87.043 on 26 degrees of freedom

```
## AIC: 128.36
##
## Number of Fisher Scoring iterations: 5
```

Vaccination lowers the infection parameter  $\beta$ , but this not statistically significant.

```
beta.table
```

```
## Vaccination beta 2.5% 97.5
## 1 Yes 1.0591299 0.7260748 1.483655
## 2 No 0.6113985 0.3386858 1.075261
```

The transmission parameter  $\beta$  is lower for the vaccinated group, but the confidence intervals overlap largely.

### Basic reproduction number $R_0$

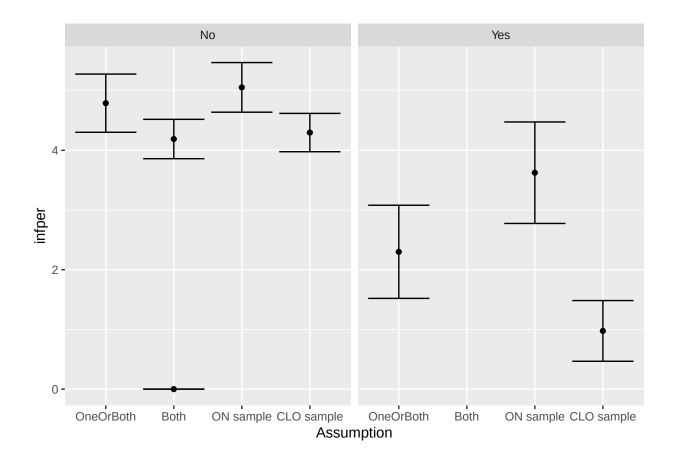
```
mean.R0[, c(1,2,3,4)]
##
                                beta
                                        llbeta
                                                   ulbeta
##
   (Intercept)
                        No 1.0591299 0.7690076 1.5713837
##
                       Yes 0.6113985 0.1955114 0.6207104
mean.R0[, c(1,5,6,7)]
                             infper llinfper ulinfper
##
               Vaccinated
##
   (Intercept)
                        No 4.297297 3.9762930 4.618302
##
                      Yes 0.975000 0.4665729 1.483427
mean.RO[, c(1,11,12,13)]
##
               Vaccinated
                                   R
                                            11R
                                                     ulR
                        No 4.4165020 2.8446651 6.856867
##
   (Intercept)
##
                      Yes 0.5258288 0.2349727 1.176715
```

Based on the assumption of CLO samples required to be positive, we find that the transmission parameter  $\beta$  and the mean infectious period are lower, although teh transmission parameter is not statistically significant. The reproduction number R is one nineth, and lower than the threshold R < 1 although the confidence interval incorporates 1.

### Graphic summary of results

### Infectious period

```
ggplot(data = all.results) +
  geom_point(aes(x = Assumption, y =infper))+
  geom_errorbar(aes( x = Assumption, ymin = llinfper, ymax = ulinfper))+
  facet_grid(.~Vaccinated)
```



# Conclusions