## Sequencing: GLM for gene expression

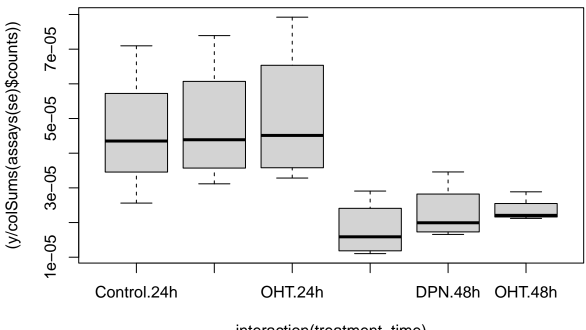
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#### Last compiled on 12 November, 2021

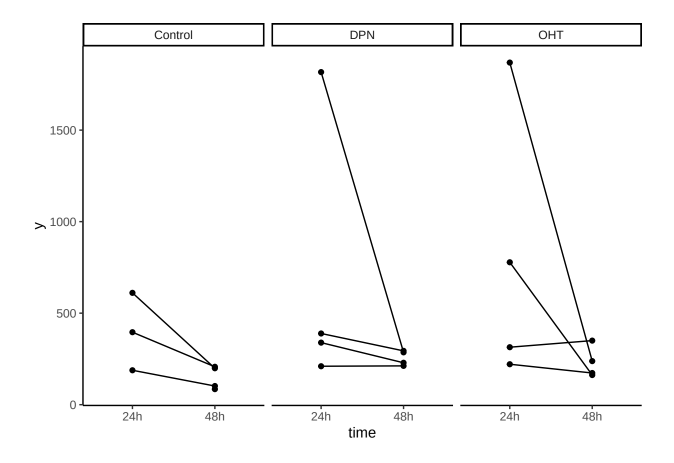
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suppressPackageStartupMessages(library(SummarizedExperiment))
se <- readRDS("data/seParathyroid.rds")</pre>
## extract data from one gene
y <- assays(se)$counts[5,]
## extract covariates for each sample
treatment <- colData(se)$treatment</pre>
table(treatment)
## treatment
## Control
           OHT
       DPN
##
```

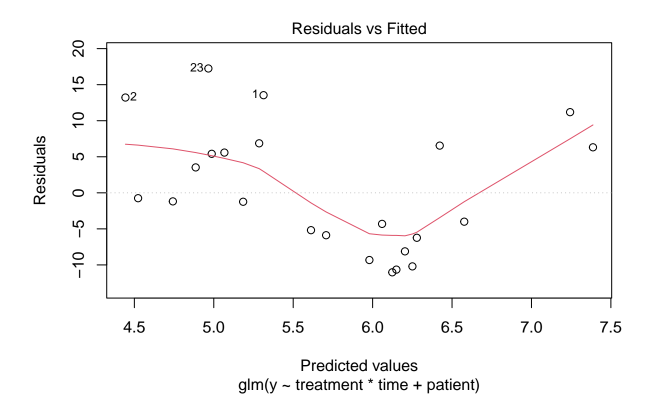
```
time <- colData(se)$time</pre>
table(time)
## time
## 24h 48h
## 11 12
patient <- colData(se)$patient</pre>
table(patient)
## patient
## 1 2 3 4
## 6 6 6 5
table(patient, treatment, time)
## , , time = 24h
##
##
       treatment
## patient Control DPN OHT
##
       1
             1 1
##
       2
              1 1
                      1
##
       3
             1 1 1
      4
             0 1 1
##
##
## , , time = 48h
##
       treatment
## patient Control DPN OHT
##
       1
           1 1 1
##
       2
             1 1 1
##
        3
             1 1 1
##
        4
               1 1 1
boxplot((y/colSums(assays(se)$counts)) ~ interaction(treatment, time))
```

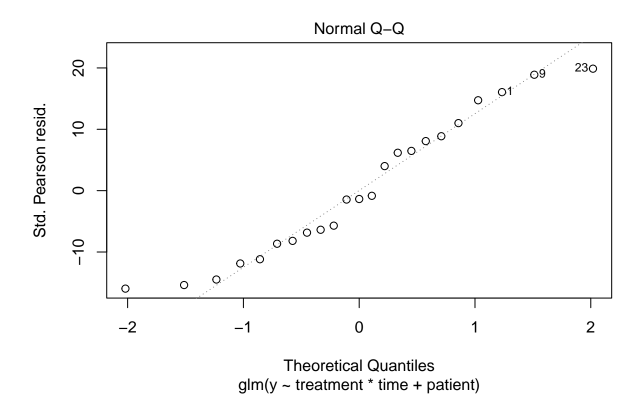


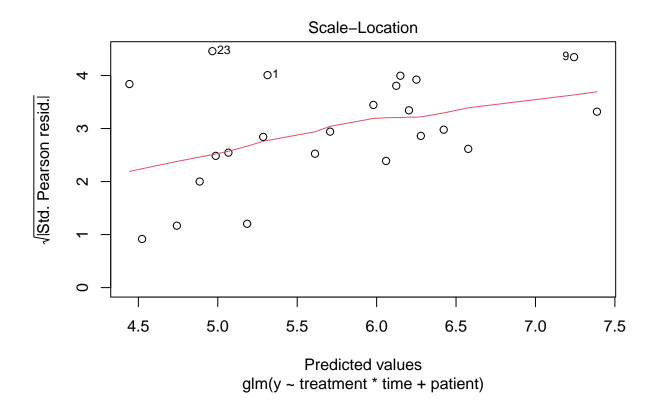
interaction(treatment, time)

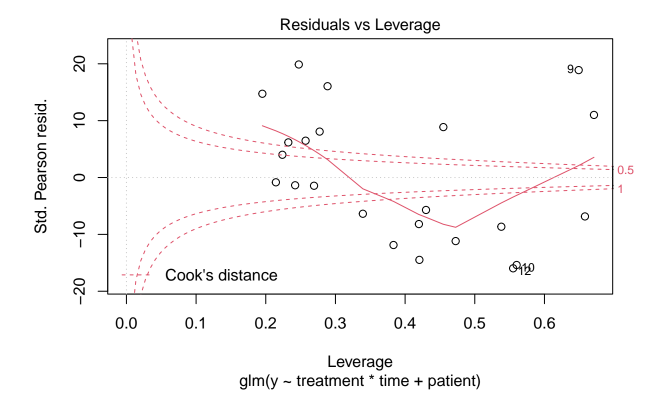


## 1 Poisson GLM







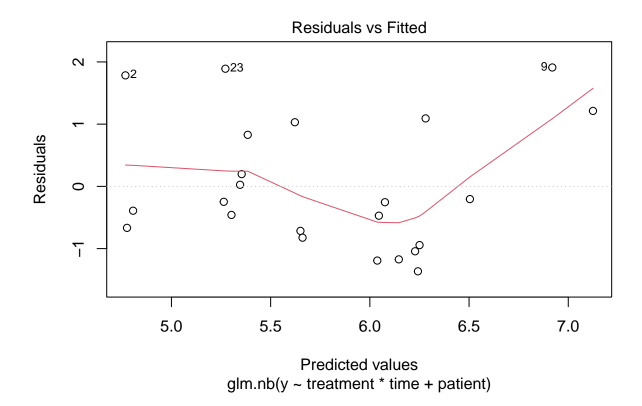


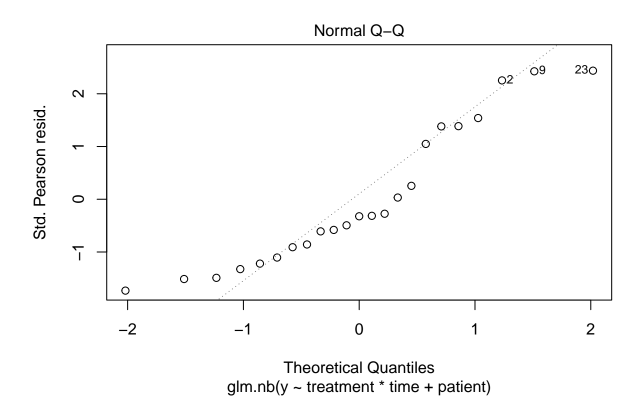
### 1.1 Check overdispersion

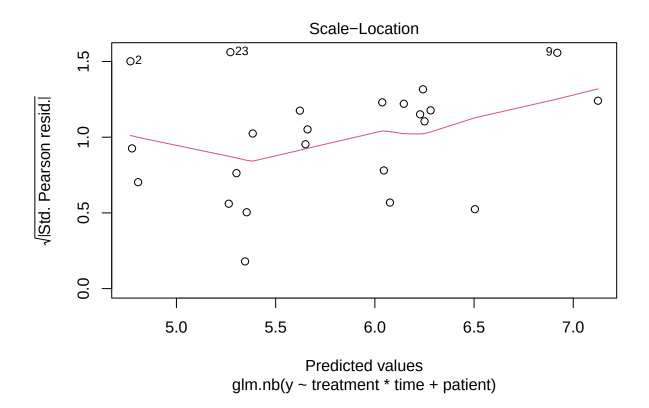
```
ePearson <- resid(m, type="pearson")
n <- length(y)
p <- length(coef(m))
sum(ePearson^2) / (n-p) # huge overdispersion.
## [1] 115.2631</pre>
```

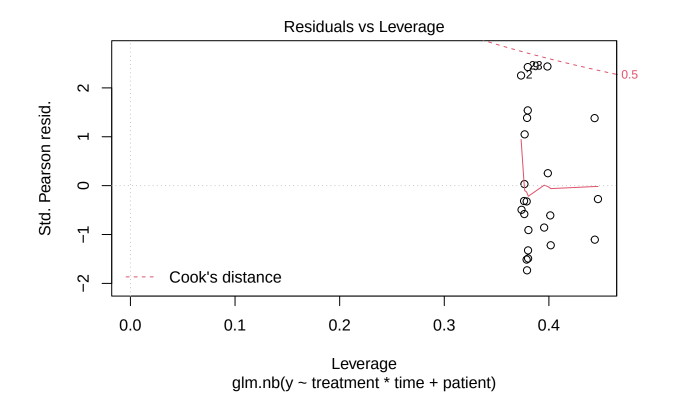
## 2 Negative binomial model

```
library(MASS)
mNB <- glm.nb(y ~ treatment*time + patient)
plot(mNB)</pre>
```









#### summary(mNB)

```
##
   glm.nb(formula = y ~ treatment * time + patient, init.theta = 5.797944761,
##
       link = log)
##
  Deviance Residuals:
##
       Min
                 1Q
                      Median
                                   3Q
                                            Max
## -1.7692
           -0.8726
                     -0.2641
                               0.8303
                                         1.5579
##
## Coefficients:
##
                         Estimate Std. Error z value Pr(>|z|)
                                    0.279514
## (Intercept)
                         5.622089
                                               20.114 < 2e-16 ***
## treatmentDPN
                         0.415319
                                    0.324036
                                                1.282 0.199945
## treatmentOHT
                         0.620294
                                    0.323896
                                                1.915 0.055479 .
## time48h
                        -0.854023
                                    0.325959
                                               -2.620 0.008792 **
## patient2
                         0.882139
                                    0.241916
                                                3.646 0.000266 ***
## patient3
                         0.038663
                                     0.242737
                                                0.159 0.873450
## patient4
                         0.008082
                                    0.258521
                                                0.031 0.975059
## treatmentDPN:time48h 0.162526
                                     0.440457
                                                0.369 0.712132
## treatmentOHT:time48h -0.124510
                                    0.440453
                                               -0.283 0.777417
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
```

```
## (Dispersion parameter for Negative Binomial(5.7979) family taken to be 1)
##
      Null deviance: 90.709 on 22 degrees of freedom
##
## Residual deviance: 23.560 on 14 degrees of freedom
## AIC: 308.63
##
## Number of Fisher Scoring iterations: 1
##
##
##
                 Theta: 5.80
##
            Std. Err.: 1.69
##
   2 x log-likelihood: -288.632
```

#### 2.1 Statistical inference

We will test seven different contrasts.

```
L <- matrix(0, nrow = length(coef(mNB)), ncol = 7)</pre>
rownames(L) <- names(coef(mNB))</pre>
colnames(L) <- c("DPNvsCON24", "DPNvsCON48",</pre>
                  "OHTvsCON24", "OHTvsCON48",
                  "DPNvsCONInt", "OHTvsCONInt",
                  "OHTvsDPNInt")
# DPN vs control at 24h
L[2,"DPNvsCON24"] <- 1
# DPN vs control at 48h
L[c(2,8),"DPNvsCON48"] <- 1
# OHT vs control at 24h
L[3,"OHTvsCON24"] <- 1
# OHT vs control at 48h
L[c(3,9),"OHTvsCON48"] <- 1
# DPN control interaction
L[8,"DPNvsCONInt"] <- 1
# OHT control interaction
L[9,"OHTvsCONInt"] <- 1
# OHT DPN interaction
L[c(9,8),"OHTvsDPNInt"] \leftarrow c(1, -1)
```

```
DPNvsCON24 DPNvsCON48 OHTvsCON24 OHTvsCON48 DPNvsCONInt
##
## (Intercept)
                                   0
                                              0
                                                          0
                                                                      0
                                                          0
## treatmentDPN
                                               1
                                                                      0
                                                                                   0
## treatmentOHT
                                   0
                                               0
                                                                                   0
                                                          1
                                                                      1
## time48h
                                   0
                                               0
                                                          0
                                                                      0
                                                                                   0
                                   0
                                              0
                                                          0
                                                                                   0
## patient2
                                                                      0
## patient3
                                   0
                                              0
                                                          0
                                                                      0
                                                                                   0
                                                          0
                                                                      0
                                                                                   0
## patient4
                                   0
                                              0
## treatmentDPN:time48h
                                   0
                                               1
                                                          0
                                                                      0
                                                                                   1
                                   0
                                                          0
## treatmentOHT:time48h
                                                                      1
                                                                                   Λ
                         OHTvsCONInt OHTvsDPNInt
## (Intercept)
                                    0
```

```
## treatmentDPN
                                    0
                                                 0
## treatmentOHT
                                    0
                                                 0
## time48h
                                    0
                                                 0
## patient2
                                    0
                                                 0
## patient3
                                    0
                                                 0
## patient4
                                    0
                                                 0
## treatmentDPN:time48h
                                                -1
## treatmentOHT:time48h
                                    1
                                                 1
```

#### 2.1.1 Wald test

```
beta <- matrix(coef(mNB), ncol = 1)
waldStats <- c()
for(ll in 1:ncol(L)){
   curL <- L[,ll,drop=FALSE]
   curWald <- t(curL) %*% beta %*% solve(t(curL) %*% vcov(mNB) %*% curL) %*% t(beta) %*% curL
   waldStats[ll] <- curWald
}</pre>
```

## [1] 1.64277658 3.75083592 3.66762230 2.75849551 0.13615681 0.07991173 0.46952521

```
pvalues <- 1-pchisq(waldStats, df=1)
pvalues</pre>
```

## [1] 0.19994494 0.05278111 0.05547928 0.09673917 0.71213195 0.77741707 0.49320567

#### 2.1.2 Likelihood ratio test

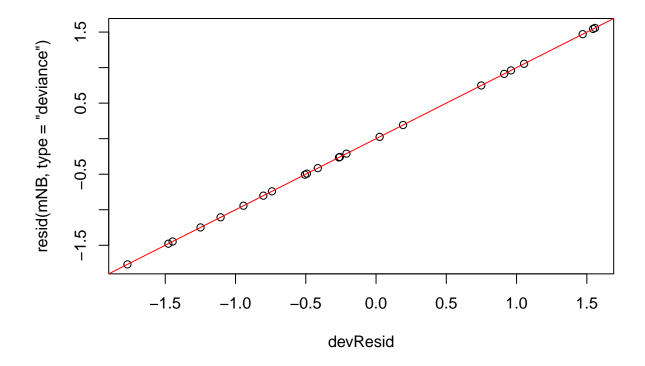
Implementing these contrasts using a likelihood ratio test is possible, but is not trivial. It would require a reparameterization of our model using the contrasts of interest. In this reparameterization, one variable may correspond to one contrast. We may then compare a full to an alternative model, dropping this variable, using a likelihood ratio test. While it is important to know that this is possible, we will not implement the reparameterization ourselves as it is considered outside the scope of this course.

#### 3 Residuals

## [1] 23.5601

#### 3.1 Deviance residuals

```
## residual deviance
sum(2*(dnbinom(x=y, mu=y, size=mNB$theta, log=TRUE) - dnbinom(x=y, mu=fitted(mNB), size=mNB$theta, log='
```

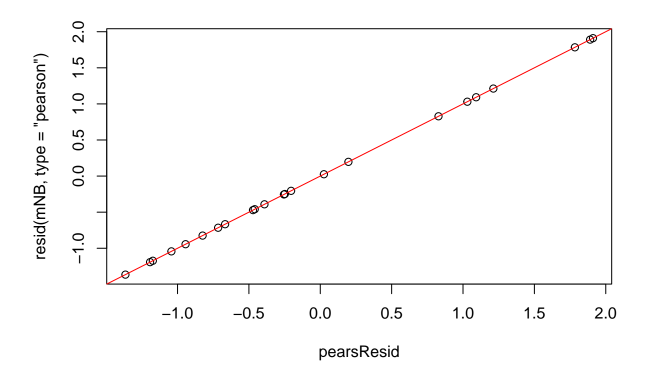


#### 3.2 Pearson residuals

```
pearsResid <- (y - fitted(mNB)) / sqrt(fitted(mNB) + 1/mNB$theta * fitted(mNB)^2)
range(pearsResid - resid(mNB, type="pearson"))

## [1] -9.888568e-10 7.026788e-10

plot(x=pearsResid, y=resid(mNB, type="pearson")); abline(0,1, col="red")</pre>
```



#### 3.3 Goodness-of-fit

```
X2 <- sum(pearsResid^2)
1-pchisq(X2, df=length(y) - length(coef(mNB)))</pre>
```

## [1] 0.04729423

## 4 Re-analysis upon basic normalization

A very simple normalization would use an offset to account for sequencing depth. Verify if our hypothesis test results remain upon using this basic normalization.

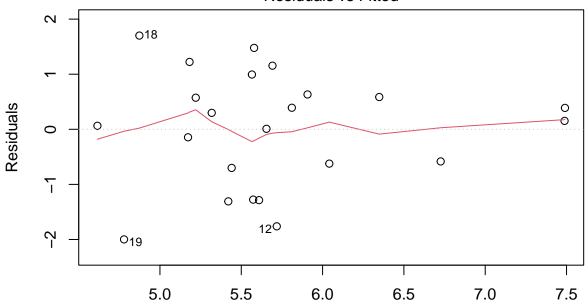
```
seqDepth <- colSums(assays(se)$counts)</pre>
```

#### 4.1 Statistical inference

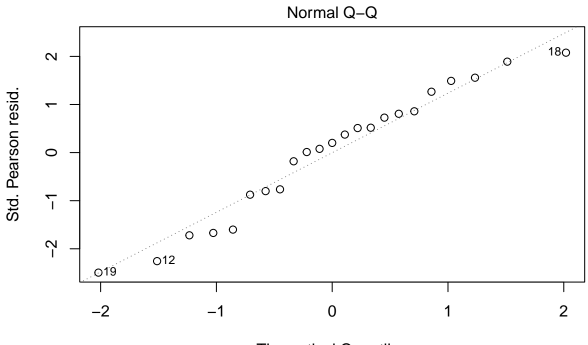
## 5 Negative binomial model, corrected for sequencing depth

```
library(MASS)
mNBOffset <- glm.nb(y ~ treatment*time + patient +</pre>
                 offset(log(seqDepth)))
plot(mNBOffset)
```

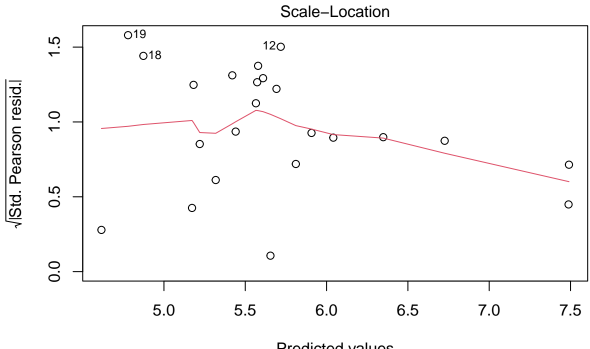
# Residuals vs Fitted



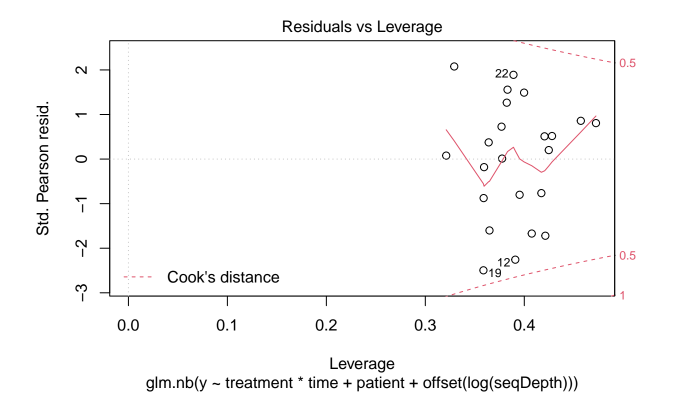
Predicted values glm.nb(y ~ treatment \* time + patient + offset(log(seqDepth)))



Theoretical Quantiles glm.nb(y ~ treatment \* time + patient + offset(log(seqDepth)))



Predicted values
glm.nb(y ~ treatment \* time + patient + offset(log(seqDepth)))



#### summary(mNBOffset)

```
##
   glm.nb(formula = y ~ treatment * time + patient + offset(log(seqDepth)),
       init.theta = 82.98036979, link = log)
##
##
  Deviance Residuals:
##
       Min
                 1Q
                      Median
                                    3Q
                                            Max
## -2.1746
           -0.6788
                      0.1520
                               0.5961
                                         1.6033
##
  Coefficients:
##
##
                         Estimate Std. Error z value Pr(>|z|)
                                      0.08217 -123.120 < 2e-16 ***
## (Intercept)
                         -10.11672
## treatmentDPN
                          0.09023
                                      0.09418
                                                 0.958 0.338022
## treatmentOHT
                          0.13403
                                      0.09362
                                                 1.432 0.152224
## time48h
                         -0.89874
                                      0.10008
                                                -8.980 < 2e-16 ***
                                                 6.925 4.37e-12 ***
## patient2
                          0.49680
                                      0.07174
## patient3
                         -0.27195
                                      0.07419
                                                -3.666 0.000247 ***
## patient4
                         -0.06219
                                      0.07888
                                                -0.788 0.430428
## treatmentDPN:time48h
                          0.15540
                                      0.13277
                                                 1.170 0.241820
## treatmentOHT:time48h
                                                 1.336 0.181560
                          0.17758
                                      0.13292
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
```

```
## (Dispersion parameter for Negative Binomial(82.9804) family taken to be 1)
##
      Null deviance: 422.284 on 22 degrees of freedom
##
## Residual deviance: 24.637 on 14 degrees of freedom
## AIC: 253.94
##
## Number of Fisher Scoring iterations: 1
##
##
##
                Theta: 83.0
##
            Std. Err.: 34.2
##
## 2 x log-likelihood: -233.939
```

#### 5.1 Wald tests

```
betaOffset <- matrix(coef(mNBOffset), ncol = 1)
waldStatsOffset <- c()
for(ll in 1:ncol(L)){
    curL <- L[,11,drop=FALSE]
    curWald <- t(curL) %*% betaOffset %*% solve(t(curL) %*% vcov(mNBOffset) %*% curL) %*% t(betaOffset) %
    waldStatsOffset[l1] <- curWald
}
waldStatsOffset

## [1] 0.91792044 6.84868664 2.04982086 10.79379449 1.36995308 1.78481023
## [7] 0.03194304

pvaluesOffset <- 1-pchisq(waldStatsOffset, df=1)
pvaluesOffset

## [1] 0.338021590 0.008870646 0.152224100 0.001018409 0.241819946 0.181559766
## [7] 0.858152733</pre>
```