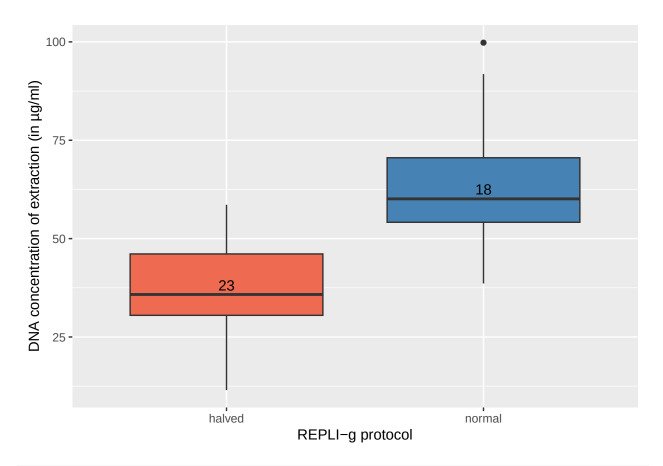
```
library(readxl)
## Warning: package 'readxl' was built under R version 4.1.3
library(ggplot2)
## Warning: package 'ggplot2' was built under R version 4.1.3
library(car)
## Warning: package 'car' was built under R version 4.1.3
## Loading required package: carData
## Warning: package 'carData' was built under R version 4.1.3
library(lme4)
## Warning: package 'lme4' was built under R version 4.1.3
## Loading required package: Matrix
library(emmeans)
## Warning: package 'emmeans' was built under R version 4.1.3
library(magrittr)
## Warning: package 'magrittr' was built under R version 4.1.3
library(DHARMa)
## Warning: package 'DHARMa' was built under R version 4.1.3
## This is DHARMa 0.4.6. For overview type '?DHARMa'. For recent changes, type news(package = 'DHARMa')
Metingen_DNA_conc <- read_excel("F:/UGent/2022-2023/Masterthesis/Metingen DNA conc z30.xlsx")
Metingen_DNA_conc %>% #Boxplots of the DNA concentration per extraction protocol
  ggplot() + aes(RepliG, conc_origineel_yg_ml, fill = RepliG) +
  geom_boxplot() +
  xlab('REPLI-g protocol') +
  ylab('DNA concentration of extraction (in µg/ml)') +
  theme(legend.position = "none") +
  annotate("text",
           x = 1:length(table(Metingen_DNA_conc$RepliG)),
           y = aggregate(conc_origineel_yg_ml ~ RepliG, Metingen_DNA_conc, median)[ , 2],
           label = table(Metingen_DNA_conc$RepliG),
           col = "black",
           vjust = -0.4) +
  scale_fill_manual('Storage', values=c('coral2','steelblue'))
```

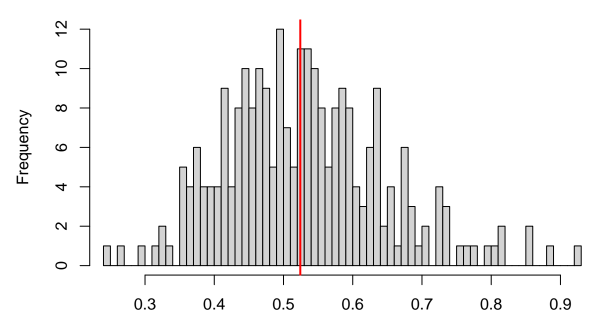


#linear model of the DNA concentration depending on the extraction protocol
lm.model1<-lmer(conc\_origineel\_yg\_ml~RepliG+(1|Species), data=Metingen\_DNA\_conc)
summary(lm.model1)</pre>

```
## Linear mixed model fit by REML ['lmerMod']
## Formula: conc_origineel_yg_ml ~ RepliG + (1 | Species)
##
      Data: Metingen_DNA_conc
##
## REML criterion at convergence: 322.6
## Scaled residuals:
                1Q Median
##
       Min
                                3Q
                                       Max
## -1.6990 -0.6317 -0.1627 0.5203 2.5908
##
## Random effects:
                         Variance Std.Dev.
##
  Groups
             Name
  Species (Intercept) 15.18
                                   3.896
   Residual
                         182.05
                                  13.493
## Number of obs: 41, groups: Species, 29
##
## Fixed effects:
##
                Estimate Std. Error t value
## (Intercept)
                  37.194
                              2.972 12.517
## RepliGnormal
                  26.242
                              4.437
                                     5.914
## Correlation of Fixed Effects:
```

```
##
              (Intr)
## RepliGnorml -0.651
anova(lm.model1, ddf="Satterthwaite",type=3)
## Warning in anova.merMod(lm.model1, ddf = "Satterthwaite", type = 3): additional
## arguments ignored: 'ddf', 'type'
## Analysis of Variance Table
         npar Sum Sq Mean Sq F value
## RepliG
          1
                6367
                       6367 34.973
emmeans(lm.model1, specs = pairwise ~ RepliG)
## $emmeans
## RepliG emmean SE df lower.CL upper.CL
## halved 37.2 3.09 31.3 30.9 43.5
## normal 63.4 3.57 29.4
                              56.1
                                       70.7
## Degrees-of-freedom method: kenward-roger
## Confidence level used: 0.95
##
## $contrasts
## contrast
              estimate SE df t.ratio p.value
## halved - normal -26.2 4.65 38 -5.638 <.0001
## Degrees-of-freedom method: kenward-roger
#test assumptions for lmer
testDispersion(lm.model1)
```

# DHARMa nonparametric dispersion test via sd of residuals fitted vs. simulated

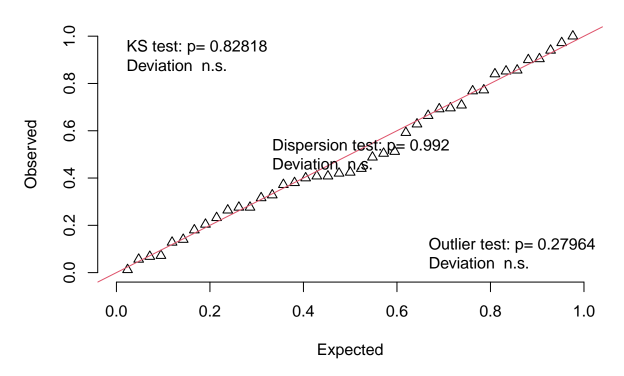


Simulated values, red line = fitted model. p-value (two.sided) = 0.992

```
##
## DHARMa nonparametric dispersion test via sd of residuals fitted vs.
## simulated
##
## data: simulationOutput
## dispersion = 0.98989, p-value = 0.992
## alternative hypothesis: two.sided
```

testUniformity(lm.model1)

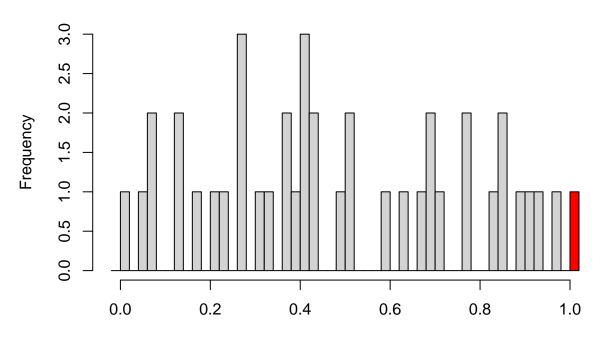
## QQ plot residuals



```
##
## One-sample Kolmogorov-Smirnov test
##
## data: simulationOutput$scaledResiduals
## D = 0.097756, p-value = 0.8282
## alternative hypothesis: two-sided

testOutliers(lm.model1)
```

#### Outlier test n.s.



Residuals (outliers are marked red)

```
##
   DHARMa outlier test based on exact binomial test with approximate
##
##
   expectations
## data: lm.model1
## outliers at both margin(s) = 1, observations = 41, p-value = 0.2796
## alternative hypothesis: true probability of success is not equal to 0.007968127
## 95 percent confidence interval:
   0.0006173169 0.1285540204
## sample estimates:
## frequency of outliers (expected: 0.00796812749003984 )
##
                                            0.02439024
simulationOutput <- simulateResiduals(fittedModel = lm.model1, plot = F)</pre>
residuals(simulationOutput, quantileFunction = qnorm, outlierValues = c(-7,7))
##
   [1]
       1.04504970 0.99445788 -0.19167090 -0.63106198 -0.32656093 0.03008408
   0.54755135 1.28155157 -1.58926756 0.42340472 -1.49085336 -0.23269275
## [19] -0.20189348 -0.59476585 -2.25712924 0.74544955
                                                     1.30468539 1.06251930
                  1.91103565 -0.47891373 7.00000000
## [25]
        0.01002668
                                                     0.32656093 -0.73227620
## [31] -0.82741832 -0.15096922 -0.30548079 -1.13589622 -0.91536509 -1.08031934
       0.73227620 0.50152740 -0.44544251 0.51293041 -1.46105627
```

### DHARMa residual

## QQ plot residuals 1.0 KS test: p= 0.82 Deviation n.s. 4 0.8 Observed 9.0 Dispersion test: p= 0.99 Deviation as 0.4 0.2 Outter test: p= 0.27964 Deviation n.s. 0.0 0.0 0.4 8.0 **Expected**

