Bacterial Abundance qPCR

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Summary

- Objective quantify bacterial abundance in starting DNA
- Method bacterial abundance quantified using zymo kit
- Results
 - clean no template controls
 - $-R^2$ standard curve 0.99
 - Large fraction of samples (especially unmixed) with Ct values outside standard curve.
- Conclusion need to rerun samples, specifically unmixed samples, diluting so that Ct values are within standard curve.

Questions

• How much template was added to each reaction?

Objective

The proportion of pre and post exposure samples in individual titrations is dependent on the ratios at which the two samples were mixed. This assumes that the pre and post samples have equivalent proportions of bacterial to non-bacterial DNA. To validate this assumption the concentration of bacterial DNA was assayed using qPCR. Additionally, the concentration of bacterial DNA in the titrations was assayed.

Methods

- zymo qPCR assay https://www.zymoresearch.com/dna/dna-analysis/femto-bacterial-dna-quantification-kit
- 45 Samples all mixed and unmixed
- diluted samples need to find out how they were diluted
- triplicates per sample 135 reactions
 - three qPCR plates, one replicate of each sample ran on each plate
- 7 concentration standard curve for the assay
 - issue with fourth standard

Munging qPCR Data

```
colnames(bac_con) <- c("well", "sample_name",</pre>
                        "plate1_Ct", "plate1_quant",
                        "plate2_Ct", "plate2_quant",
                        "plate3_Ct", "plate3_quant")
bac_con <- bac_con %>% gather("id","value", -well, -sample_name) %>%
      separate(id, c("plate","var"), sep = "_") %>%
      spread(var, value)
bac_std <- read_excel(path = "../data/MixStudy_Nate_20160919.xls",</pre>
                      sheet = "QDNA_20160919",skip = 3, col_names = FALSE) %>%
      select(-X12, -X13, -X5,-X7,-X9) %>% filter(X2 %in% paste0("Std",1:7))
colnames(bac_std) <- c("well", "sample_name", "conc", "plate1", "plate2", "plate3")</pre>
bac_std <- bac_std %% gather("plate","Ct",-well, -sample_name, -conc) %>%
      mutate(conc = as.numeric(conc), Ct = as.numeric(Ct)) %>% filter(!is.na(Ct))
## Warning in eval(substitute(expr), envir, enclos): NAs introduced by
## coercion
## NAs introduced when converting samples with undetermined and omit Ct values.
```

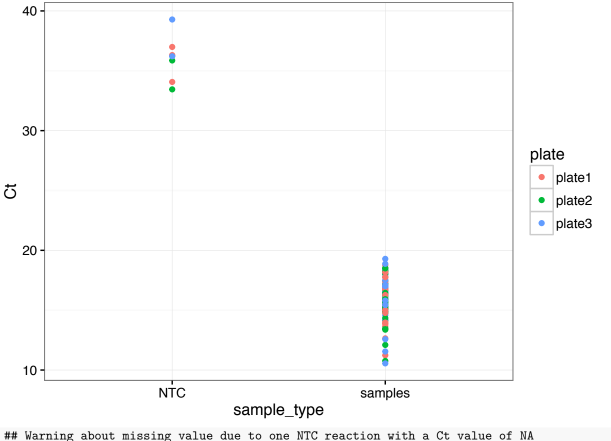
Results

NTC Check

For NTC (no template control) samples the Ct values were Comparison of Ct values between NTC and samples. NTC Ct values > 30, and sample Ct values < 20, excluding NTC from the rest of the analysis.

```
bac_con %>% mutate(sample_type = if_else(sample_name == "NTC", "NTC", "samples")) %>%
ggplot() + geom_point(aes(x = sample_type, y = Ct, color = plate)) + theme_bw()
```

Warning: Removed 1 rows containing missing values (geom_point).



```
## Warning about missing value due to one NTC reaction with a Ct value of NA
## plot shows okay to exclude NTC
bac_con <- bac_con %>% filter(sample_name != "NTC")
```

Analysis of Standard Curve

Fitting the standard curve using a linear model, Ct~log10(concentration).

Fitted model using the three plates as replicates and not fitting individual models for each plate. Using a random effect model is likely a better approach.

```
##
## Residuals:
                    1Q
                          Median
## -0.228775 -0.096812 0.006189 0.116039 0.235924
##
## Coefficients:
               Estimate Std. Error t value Pr(>|t|)
## (Intercept) 4.365381
                                      34.68 2.11e-13 ***
                           0.125876
## Ct
              -0.213182
                           0.004819 -44.24 1.16e-14 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 0.1501 on 12 degrees of freedom
## Multiple R-squared: 0.9939, Adjusted R-squared: 0.9934
## F-statistic: 1957 on 1 and 12 DF, p-value: 1.158e-14
std_fit2 <- lm(log_conc~plate/Ct, data = bac_std)</pre>
summary(std_fit2)
##
## Call:
## lm(formula = log_conc ~ plate/Ct, data = bac_std)
## Residuals:
       Min
                  1Q
                     Median
                                    30
                                            Max
## -0.19978 -0.07599 0.03538 0.06859 0.16901
##
## Coefficients:
                  Estimate Std. Error t value Pr(>|t|)
##
## (Intercept)
                  4.417411
                             0.189273 23.339 1.21e-08 ***
                                       1.312
## plateplate2
                  0.411203
                            0.313488
                                                  0.226
## plateplate3
                  -0.347403
                              0.260241 -1.335
                                                  0.219
## plateplate1:Ct -0.213979
                              0.006960 -30.743 1.36e-09 ***
## plateplate2:Ct -0.231415
                              0.010552 -21.930 1.97e-08 ***
## plateplate3:Ct -0.204254
                              0.006644 -30.741 1.36e-09 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 0.1347 on 8 degrees of freedom
## Multiple R-squared: 0.9967, Adjusted R-squared: 0.9947
## F-statistic: 487.5 on 5 and 8 DF, p-value: 1.029e-09
bac_fit <- bac_con %>% select(sample_name,plate, Ct) %>% add_predictions(std_fit2)
std_coef <- std_fit2 %>% coefficients()
coef_df <- frame_data(</pre>
      ~plate, ~intercept, ~slope,
      "plate1", std_coef[1], std_coef[4],
      "plate2",std_coef[1] + std_coef[2], std_coef[5],
      "plate3", std_coef[1] + std_coef[3], std_coef[6]
fit2_df <- coef_df %>% mutate(mod = "log_conc~plate/Ct")
std_fit3 <- lm(log_conc~plate*Ct, data = bac_std)</pre>
std_coef <- std_fit3 %>% coefficients()
## not sure if I need to add Ct
```

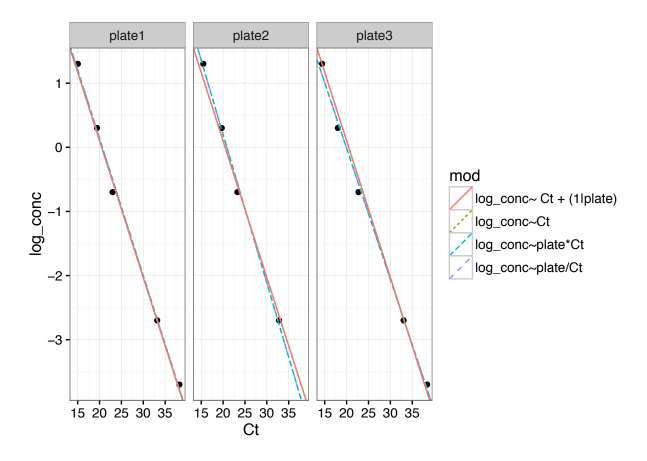
```
fit3_df <- frame_data(</pre>
      ~plate, ~intercept, ~slope,
      "plate1", std_coef[1], std_coef[4],
      "plate2",std_coef[1] + std_coef[2], std_coef[4] + std_coef[5],
      "plate3", std_coef[1] + std_coef[3], std_coef[4] + std_coef[6]
      ) %>%
      mutate(mod = "log_conc~plate*Ct")
summary(std_fit3)
##
## Call:
## lm(formula = log_conc ~ plate * Ct, data = bac_std)
## Residuals:
                     Median
                  1Q
                                    3Q
## -0.19978 -0.07599 0.03538 0.06859 0.16901
## Coefficients:
                   Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                   4.417411 0.189273 23.339 1.21e-08 ***
## plateplate2
                   0.411203
                              0.313488
                                       1.312
                                                  0.226
                              0.260241 -1.335
                                                  0.219
## plateplate3
                  -0.347403
## Ct
                  -0.213979
                              0.006960 -30.743 1.36e-09 ***
## plateplate2:Ct -0.017436
                              0.012641 -1.379 0.205
## plateplate3:Ct 0.009725
                              0.009623
                                       1.011
                                                  0.342
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 0.1347 on 8 degrees of freedom
## Multiple R-squared: 0.9967, Adjusted R-squared: 0.9947
## F-statistic: 487.5 on 5 and 8 DF, p-value: 1.029e-09
fit3_df
## # A tibble: 3 × 4
     plate intercept
                           slope
                                               mod
                <dbl>
                           <dbl>
##
      <chr>>
                                             <chr>
## 1 plate1 4.417411 -0.2139791 log_conc~plate*Ct
## 2 plate2 4.828613 -0.2314146 log_conc~plate*Ct
## 3 plate3 4.070008 -0.2042542 log_conc~plate*Ct
library(lme4)
## Loading required package: Matrix
## Attaching package: 'Matrix'
## The following object is masked from 'package:tidyr':
##
##
       expand
std_fit4 <- lmer(log_conc~ Ct + (1|plate), data = bac_std)</pre>
fit4_df <- tibble(plate = paste0("plate",c(1:3)),</pre>
                  slope = coef(std_fit4)$plate[1,2],
                  intercept = coef(std_fit4)$plate[1,1],
```

```
mod = "log_conc~ Ct + (1|plate)")
coef(std_fit4)
## $plate
##
          (Intercept)
                              Ct
## plate1
            4.365381 -0.2131823
            4.365381 -0.2131823
## plate2
## plate3
            4.365381 -0.2131823
##
## attr(,"class")
## [1] "coef.mer"
summary(std_fit4)
## Linear mixed model fit by REML ['lmerMod']
## Formula: log_conc ~ Ct + (1 | plate)
##
     Data: bac_std
##
## REML criterion at convergence: -1.9
## Scaled residuals:
##
       Min
                 1Q
                      Median
                                   30
## -1.52435 -0.64507 0.04124 0.77318 1.57198
## Random effects:
## Groups
                        Variance Std.Dev.
           Name
## plate
             (Intercept) 2.520e-17 5.020e-09
## Residual
                         2.252e-02 1.501e-01
## Number of obs: 14, groups: plate, 3
##
## Fixed effects:
##
               Estimate Std. Error t value
## (Intercept) 4.365381
                          0.125876
                                    34.68
## Ct
              -0.213182
                          0.004819 -44.24
##
## Correlation of Fixed Effects:
##
      (Intr)
## Ct -0.948
Comparing Regression Models
Model Matrix
std_fit$model
##
     log_conc
## 1
     1.30103 15.042
## 2 0.30103 19.425
## 3 -0.69897 22.977
## 4 -2.69897 33.186
## 5 -3.69897 38.270
## 6
     1.30103 15.481
## 7
     0.30103 19.697
```

8 -0.69897 23.272

```
## 9 -2.69897 32.773
## 10 1.30103 14.384
## 11 0.30103 17.992
## 12 -0.69897 22.717
## 13 -2.69897 33.033
## 14 -3.69897 38.407
std_fit2$model
##
      log_conc plate
## 1
       1.30103 plate1 15.042
      0.30103 plate1 19.425
## 2
## 3
    -0.69897 plate1 22.977
     -2.69897 plate1 33.186
     -3.69897 plate1 38.270
## 5
## 6
       1.30103 plate2 15.481
## 7
       0.30103 plate2 19.697
## 8
     -0.69897 plate2 23.272
     -2.69897 plate2 32.773
## 10 1.30103 plate3 14.384
## 11 0.30103 plate3 17.992
## 12 -0.69897 plate3 22.717
## 13 -2.69897 plate3 33.033
## 14 -3.69897 plate3 38.407
std_fit3$qr
## $qr
##
      (Intercept) plateplate2 plateplate3
                                                      Ct plateplate2:Ct
## 1
       -3.7416574 -1.0690450 -1.33630621 -92.647713076
                                                         -2.438037e+01
## 2
        0.2672612
                    1.6903085 -0.84515425
                                            -4.627304060
                                                           3.854875e+01
## 3
        0.2672612
                    0.1333828
                                            -0.748511122
                              1.58113883
                                                          -3.082758e-15
## 4
        0.2672612
                    0.1333828
                               0.22016954
                                           -30.791337969
                                                          -5.289791e+00
## 5
        0.2672612
                    0.1333828
                              0.22016954
                                             0.495148219
                                                           1.161455e+01
## 6
        0.2672612 -0.4582252
                               0.03419583
                                            -0.217992572
                                                           5.560054e-01
## 7
        0.2672612
                   -0.4582252
                               0.03419583
                                            -0.081070948
                                                           2.398987e-01
## 8
        0.2672612
                   -0.4582252
                               0.03419583
                                                          -2.814719e-02
                                             0.035033133
## 9
        0.2672612 -0.4582252 0.03419583
                                             0.343593937
                                                          -7.405120e-01
## 10
        0.2672612
                    0.1333828 -0.41228599
                                            -0.266174682
                                                          -9.114631e-02
## 11
        0.2672612
                    0.1333828 -0.41228599
                                            -0.148998871
                                                          -5.102175e-02
## 12
        0.2672612
                    0.1333828 -0.41228599
                                             0.004453376
                                                           1.524971e-03
## 13
        0.2672612
                                                           1.162492e-01
                    0.1333828 -0.41228599
                                            0.339482663
## 14
        0.2672612
                    0.1333828 -0.41228599
                                            0.514012266
                                                           1.760134e-01
##
      plateplate3:Ct
## 1
        -33.81736672
## 2
        -21.38798066
## 3
         40.01324792
## 4
        -13.34239940
## 5
         -6.07673005
## 6
        -13.99585865
## 7
          0.04184597
## 8
         -0.02320216
## 9
         -0.19607552
## 10
         -0.58345600
## 11
         -0.41237173
```

```
## 12
         -0.18832151
## 13
          0.30084294
          0.55566746
## 14
## attr(,"assign")
## [1] 0 1 1 2 3 3
## attr(,"contrasts")
## attr(,"contrasts")$plate
## [1] "contr.treatment"
##
##
## $qraux
## [1] 1.267261 1.133383 1.220170 1.330037 1.169554 1.118557
## $pivot
## [1] 1 2 3 4 5 6
##
## $tol
## [1] 1e-07
##
## $rank
## [1] 6
##
## attr(,"class")
## [1] "qr"
fit_coefs <- bind_rows(fit1_df, fit2_df, fit3_df,fit4_df)</pre>
ggplot(fit_coefs) + geom_point(data = bac_std, aes(y = log_conc, x = Ct)) +
      geom_abline(aes(slope = slope, intercept = intercept, color = mod, linetype = mod)) +
      facet_grid(.~plate) + theme_bw()
```



Samples outside of standard curve range.

Samples have Ct values lower then the standard curve samples and fall outside of the standard curve. This is not ideal. Only 2 of the 10 unmixed pre and post treatment samples have Ct values within the standard curve for all three plates, and none of the remaining 8 have Ct within the standard curve for 2 of the three plates. The linearity of the standard curve outside of the standard concentration range is unknown.

```
min_std <- bac_std %>% filter(conc == 20) %>% select(plate, Ct) %>% rename(std_ct = Ct)
bac_fit <- left_join(bac_fit, min_std) %>%
    mutate(relative_ct = if_else(Ct < std_ct, "outside","within")) %>%
    mutate(sam_type = if_else(grepl("\\(", sample_name), "unmixed", "titration"))
```

```
## Joining, by = "plate"
```

```
# ggplot(bac_std) + geom_point(aes(y = log_conc, x = Ct)) +

# geom_abline(aes(slope = std_slope, intercept = std_intercept), color = "grey80") +

# geom_point(data = bac_fit, aes(y = pred, x = Ct, shape = plate, color = relative_ct)) +

# facet_grid(plate~sam_type) +

# theme_bw()

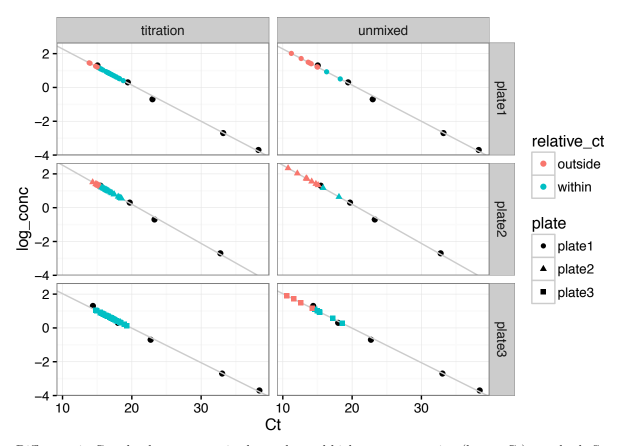
ggplot(bac_std) + geom_point(aes(y = log_conc, x = Ct)) +

geom_abline(data = coef_df, aes(slope = slope, intercept = intercept), color = "grey80") +

geom_point(data = bac_fit, aes(y = pred, x = Ct, shape = plate, color = relative_ct)) +

facet_grid(plate~sam_type) +

theme_bw()
```



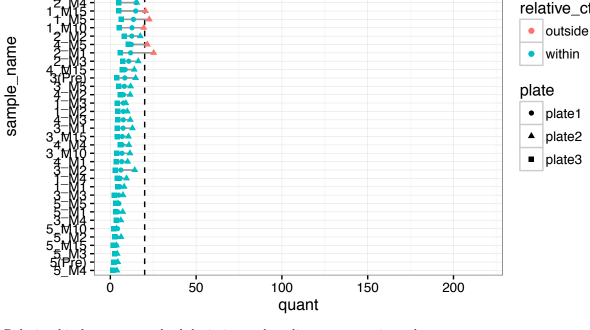
Difference in Ct value between unmixed samples and highest concentration (lowest Ct) standard. Samples with negative values are outside of the standard curve.

sample_name	plate1	plate2	plate3
1 Post	1.381	1.987	-0.572
1 Pre	0.029	0.384	-0.954
2 Post	3.810	4.720	3.829
2 Pre	0.056	2.111	-0.203
3 Post	2.395	3.384	2.836
3 Pre	-1.278	-0.336	-2.801
4 Post	0.107	0.705	-0.886
4 Pre	0.946	1.314	0.164
5 Post	1.153	2.067	1.806
5 Pre	-3.246	-2.623	-4.171

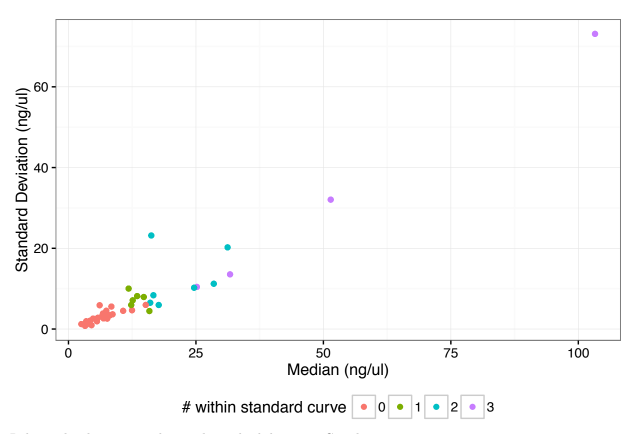
Variability of samples outside of the standard curve

The variability in concentration for the unmixed samples, especially post treatment biological replicates $2(7_2(Post))$ and $3(8_3(Post))$, is likely due to the sample concentration falling outside the range of the

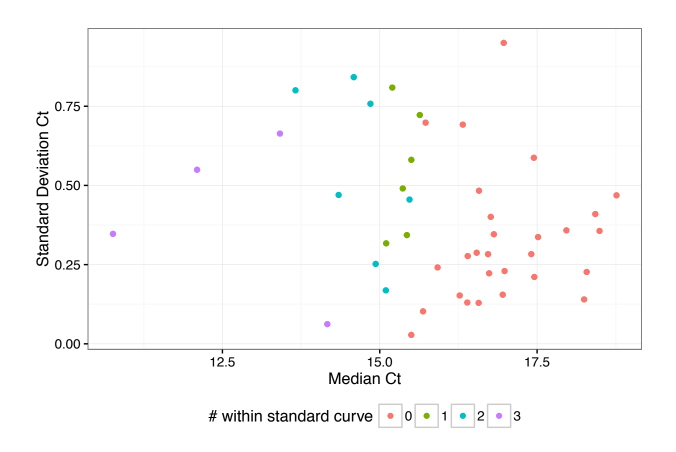
standard curve.



Relationship between standard deviation and median concentration values.



Relationship between median and standard deviation Ct values.



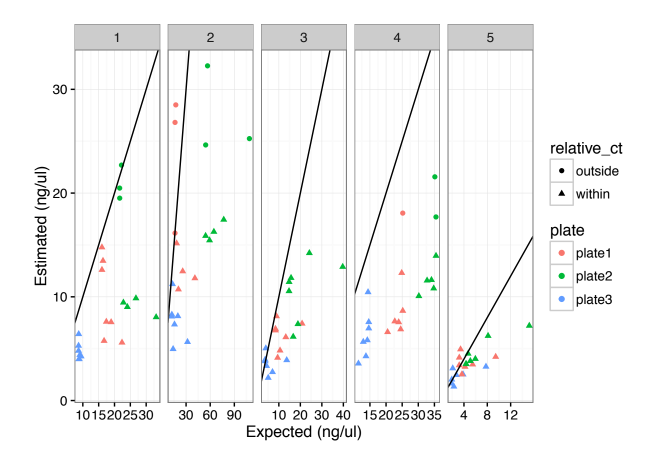
Tritration Concentration Deviation from Expected Value

geom_abline(aes(intercept = 0, slope = 1)) +
facet_grid(~bio_rep, scales = "free") +

geom point() +

```
bac_unmixed <- bac_fit %>% filter(sam_type == "unmixed") %>%
            mutate(sample_name = str_replace(sample_name, ".*_", ""),
                   sample_name = str_replace(sample_name, '\\('," "),
                   sample_name = str_replace(sample_name, '\\)',"")) %>%
            separate(sample_name,into = c("bio_rep","titration"),remove = FALSE) %>%
            mutate(titration_factor = if_else(titration == "Pre", 20, 0)) %>%
            select(bio_rep, plate,pred, titration) %>%
            spread(titration,pred)
bac_mixed <- bac_fit %>% filter(sam_type == "titration") %>%
      separate(sample name, into = c("bio rep", "titration"), remove = FALSE) %>%
      mutate(titration_factor = str_replace(titration, "M","") %>% as.numeric())
bac_fit2 <- left_join(bac_mixed, bac_unmixed) %>% select(-sample_name, -Ct, -std_ct, -sam_type) %>%
      mutate(exp_pred = Post*(2^-titration_factor) + Pre*(1-2^-titration_factor))
## Joining, by = c("bio_rep", "plate")
Expected Concentration - calculation based on concentration measurements for pre and post unmixed samples
Estimated Concentration - calculated using standard curve
bac_fit2 %>% ggplot(aes(x = 10^exp_pred, y = 10^pred, color = plate, shape = relative_ct)) +
```

labs(x = "Expected (ng/ul)", y = "Estimated (ng/ul)") + theme_bw()



Conclusions

The samples should be diluted and run again.

Caveats

• To reduce experimental design complexity, sample name confounded with well except for negative controls (No Template Control, NTC).