SAMsize

Setup

Normalize path and load libraries etc.

```
library(gsl)
library(dplyr)
library(magrittr)
library(cowplot)
library(viridis)
cbPalette <- c("#999999", "#E69F00", "#56B4E9", "#009E73", "#F0E442", "#0072B2", "#D55E00", "#CC79A7")
cols <- viridis(10)

## more packages
library(nlme)
library(quantgen)
library(tidyr)</pre>
```

Functions

```
countsize <- read.csv("data/SAM_cellcount.csv", header=T)
head(countsize)</pre>
```

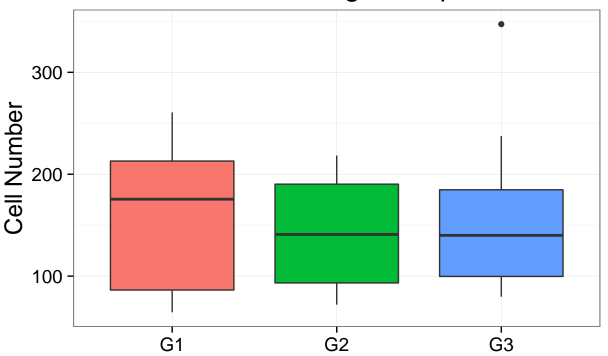
```
##
    Count_Cells Mean_Area_percell Genotype Plant Growth_Period BAK1 SDA1
## 1
             158
                          791.4494
                                       A554
                                                 1
                                                              G1 REF
                                                                       REF
## 2
             188
                          831.1223
                                       A554
                                                1
                                                              G2
                                                                  REF
                                                                       REF
                          947.1709
## 3
             199
                                       A554
                                                1
                                                              GЗ
                                                                  REF
                                                                       REF
             205
                          812.2780
                                       A554
                                                2
                                                              G1
                                                                  REF
                                                                       REF
## 4
                                       A554
                                                2
                                                              G2 REF
## 5
             137
                          732.2628
                                                                       REF
## 6
                          959.8991
                                       A554
                                                 2
                                                              G3
                                                                  REF
                                                                       REF
             228
       SAM_V
##
## 1 2502482
## 2 2502482
## 3 2502482
## 4 2502482
## 5 2502482
## 6 2502482
```

In the above SAM Cell count table, 14 Genotypes were collected for SAM cell counts in 3 growth periods, each period with 3 plants. There are also two factors, BAK1 and SDA1, that associated with some traits. From the data, we learned that Count_Cells is significantly correlated with SAM_V (r = 0.75, Pvalue < 0.01). But Count_Cells is not correlated with Mean_Area_percell (r = 0.03, Pvalue = 0.7).

We estimated BLUE values by fitting a linear mixed model, where **Genotype** as a fixed effect and **Plant** as a random effect.

```
csg1 <- mixed_model(data = subset(countsize, Growth_Period %in% "G1"),</pre>
                    model = Count_Cells ~ Genotype, random = ~1 | Plant)
csg2 <- mixed_model(data = subset(countsize, Growth_Period %in% "G2"),</pre>
                    model = Count_Cells ~ Genotype, random = ~1 | Plant)
csg3 <- mixed_model(data = subset(countsize, Growth_Period %in% "G3"),</pre>
                    model = Count_Cells ~ Genotype, random = ~1 | Plant, trait = "CS_G3")
cs <- merge(csg1, csg2, by="Genotype")</pre>
cs <- merge(cs, csg3, by="Genotype") %% set_names(c("genotype", "g1", "g2", "g3"))
lcs <- cs %>% gather(key="Growth", value="cellnum", 2:4)
#theme_set(theme_grey(base_size = 18))
p <- ggplot(lcs, aes(x=toupper(Growth), y=cellnum, fill = Growth)) +
    #opts(axis.text.x=theme_text(angle=90)) +
    theme_bw(base_size = 18) +
    \#theme(axis.text.x = element\_text(angle = 90, hjust = 1, size=12)) +
    geom_boxplot() +
    ggtitle("BLUEs of three growth periods") + xlab("") + ylab("Cell Number") +
    #ggtitle("Sequencing Depth") + xlab("") + ylab("Depth per cytosine site") +
    guides(fill=FALSE)
p
```

BLUEs of three growth periods



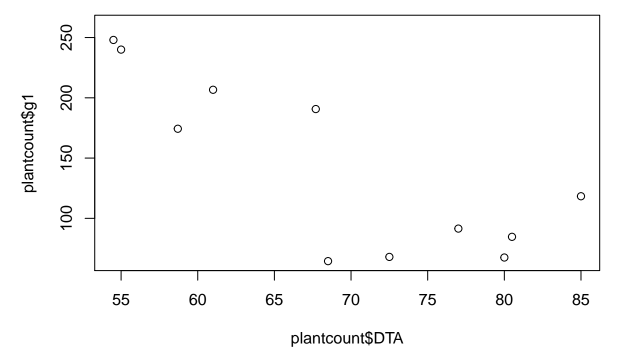
```
#samsize<-read.csv("~/Desktop/samsize.csv",header=T)
#samsize_unsummary<-read.csv("~/Desktop/samsize_unsummarized.csv",header=T)
plantstuff <- read.csv("data/plantstuff.csv",header=T) %>%
    select(Genotype,PlantH..cm.,EarH..cm.,mean_nodes,mean_dia_ear..mm.,mean_dia_below..mm.,DTA,SAM_V..um.
```

```
set_names(c("genotype","plant_height","ear_height","leaf_nodes","ear_width","stem_width","DTA","SAM_v
plantstuff$genotype <- toupper(plantstuff$genotype)
cs$genotype <- toupper(cs$genotype)

plantcount <- merge(plantstuff, cs, by="genotype")
head(plantcount)</pre>
```

```
##
     genotype plant_height ear_height leaf_nodes ear_width stem_width DTA
## 1
         A554
                     130.1
                                                              16.54333 58.7
                                 42.9
                                         12.66667
                                                   15.48333
## 2
          B57
                     135.7
                                  62.8
                                         12.33333
                                                   15.23333
                                                              16.24000 68.5
          B73
                                         14.00000
                                                   17.67667
## 3
                     169.3
                                 91.1
                                                              19.05667 67.7
## 4
       CML261
                     207.4
                                122.5
                                         21.00000
                                                   18.21000
                                                              18.76667 85.0
       CML322
                     136.6
                                 83.2
                                         17.00000
                                                   16.97667
                                                              17.13333 80.0
## 5
## 6
       CML333
                     169.8
                                109.1
                                         17.00000
                                                   14.00667
                                                              15.23000 77.0
                               g2
##
     SAM volume
                      g1
## 1 1164089.33 174.3333 145.3333 187.46636
## 2 -551935.71 64.5000 74.5000 79.79969
## 3 451176.36 190.6667 182.0000 161.50000
       92414.76 118.3333 144.3333 118.50000
## 5 -528356.95 67.5000 72.0000 89.13302
## 6 -104991.26 91.5000 113.7500 97.50000
```

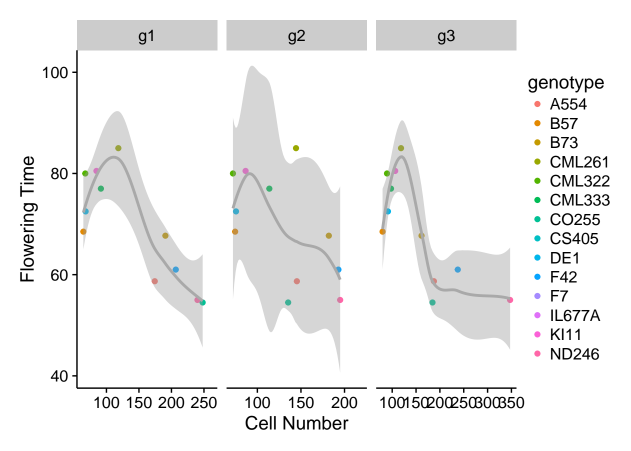
plot(plantcount\$DTA, plantcount\$g1)



```
lcs <- cs %>% gather(key="Growth", value="cellnum", 2:4)

p1 <- plantcount[, c("genotype", "g1", "g2", "g3")] %>%
    gather(key="Growth", value="cellnum", 2:4) %>%
    merge(plantcount[, 1:8], by="genotype")
```

```
#DTA, cell count
ggplot(p1, aes(y=DTA, x=cellnum))+
  geom_point(aes(color=genotype))+
  geom_smooth(color="dark grey")+
  facet_wrap(~Growth, scales="free_x")+
  xlab("Cell Number")+
  ylab("Flowering Time")
```



```
#cell count, SAM volume
ggplot(plantcount,aes(y=SAM_volume,x=cell_number))+
  geom_point(color=cols[1])+
  geom_smooth(method="loess",color="dark grey")+
  facet_wrap(~growth_period,scales="free_x")+
  ylab("SAM volume")+
  xlab("Cell Number")
#cell size, SAM volume
ggplot(plantcount,aes(y=SAM_volume,x=cell_size))+
  geom_point(color=cols[2])+
  geom_smooth(method="loess",color="dark grey")+
  facet_wrap(~growth_period,scales="free_x")+
  ylab("SAM volume")+
  xlab("Cell Size")
#cell size, cell number
ggplot(plantcount,aes(x=cell_number,y=cell_size))+
```

```
geom_point(color=cols[3])+
geom_smooth(method="loess",color="dark grey")+
facet_wrap(~growth_period,scales="free_x")+
xlab("Cell Number")+
ylab("Cell Size")

#more cells -> no diff plant height. smaller ear height, weakly smaller ear width & stem_width, shorter
summary(lm(data=plantcount,DTA~cell_number+SDA1+BAK1))
```