Reworked analyses and figure drafts for paper1 after JEB reviews

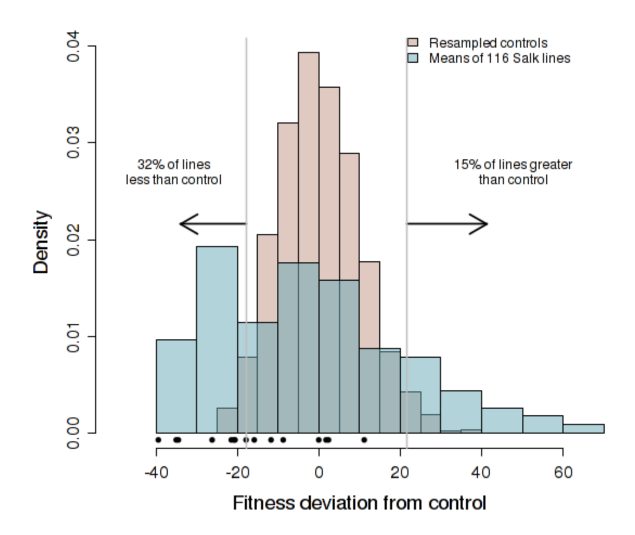
[1] "does gls exist TRUE"

1 Distribution of mutations

1.1 Distribution of mutations relative to control using means only.

Here are various figures emphasizing variation among mutant lines compared to controls. In this first figure, the distribution of line means is plotted with the distribution of all control plants as well as 116 resampled means of control replicates equal in size to the average number of reps per SALK line

This figure focuses on the resampled control means and compares to individual line means. This is the figure we placed in the ms

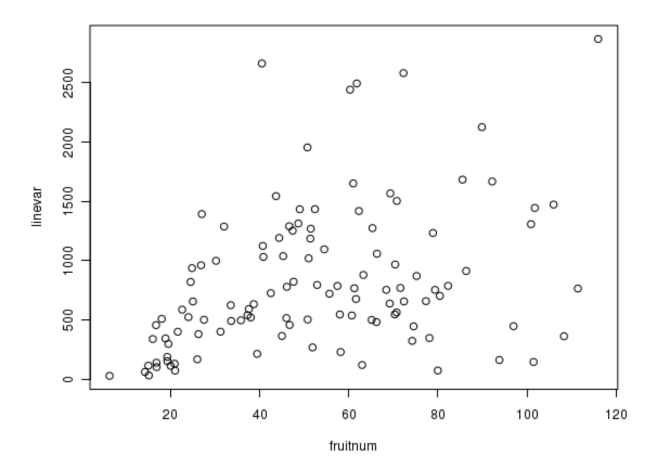


1.2 Additions suggested by JEB reviewer 1

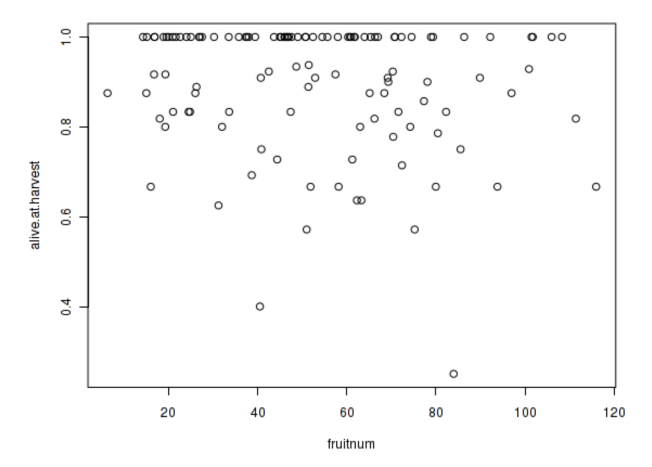
1.2.1 Some diagnostics

Does variance within a line depend on fruitnumber? This is the classic variance/mean correlation

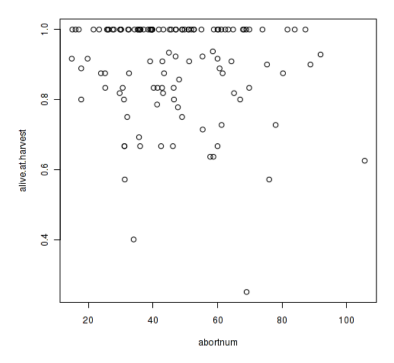
The answer in the figure below is that variance does depend on mean.



is there a reason to expect that survival rate in a line is related to average fruitnum for the line? Looks like if there is any pattern, it is the low fruitnumbers that have the high survival.

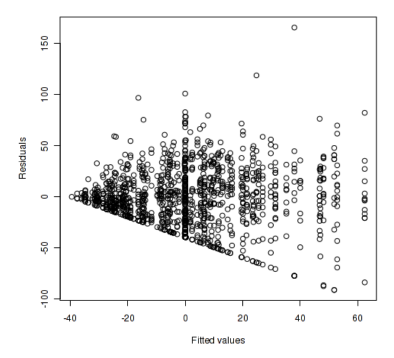


Does look like there is a similar pattern in abort/alive-at-harvest relationship



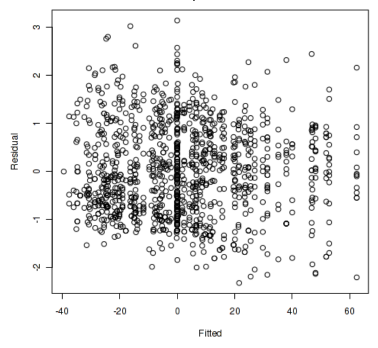
1.2.2 Crafting the best linear model to underly a testing framework

There is going to be heteroschedacity as established above. The figure below shows the residual by fitted response for a plain OLS anova: fitness as a function of line. Clearly there is terrible heteroscedacity



This figure shows the same plot using standardized residuals after using a weighting scheme that allows each line to have a separate variance (this does add tons of parameters to the model, however). You can see that the heteroschedacity disappears.

Standardized residuals (Pearson resids in R) versus fitted GLS model with separate variance estimates



Is the addition of all the extra parameters worth it? They dramatically improve residual performance. The following likelihood ratio tests suggest that the extra parameters still produces a much better model. "fit.gls" is a "standard anova", "fit.gls.id" is the same model with individual line variance weighting.

```
Model
                              AIC
                                        BIC
                                                logLik
                                                         Test
                                                               L.Ratio p-value
## fit.gls
                   1 118 10557.34 11141.99 -5160.671
## fit.gls.id
                   2 234 10392.93 11552.31 -4962.464 1 vs 2 396.4131
## Denom. DF: 1048
##
                numDF
                       F-value p-value
  (Intercept)
                      3336.923
                                < .0001
## SALK_Line
                        14.631
                  116
                                 < .0001
```

1.2.3 Multiple test approach using treatment contrasts

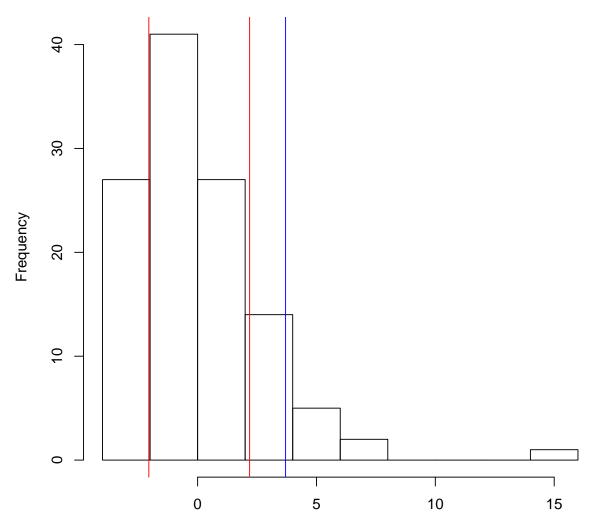
First, we could look at dummy contrast coding comparing each line to the control. Technically this could be thought of as a planned comparison. Hard for me to swallow this and I expect the same is true for reviewers.

I've produced a histogram of effect sizes calculated as t-statistics in comparison to the control. The interval between the blue lines indicates lines that are not different than controls after correcting the significance of these t-stats using Bonferroni. There is one line to the left of the blue interval and 11 to the right of the blue interval (?!). The interval among the

red line corresponds to no correction for multiple tests. In this case, 30 lines are lower than the red interval and 22 are larger.

```
## Warning in max(coef.lm[(coef.lm[, "bonsig"]) & (coef.lm[, 3] < 0), 3]): no non-missing arguments to max; returning -Inf
```

Distribution of effect sizes (t-stats) comparing to control



Effect size (t-statistic from analysis of regression coef)

```
## [1] "Numbers of lines that have more extreme fitness than control: bonferroni correct
##
## bonsig FALSE TRUE
```

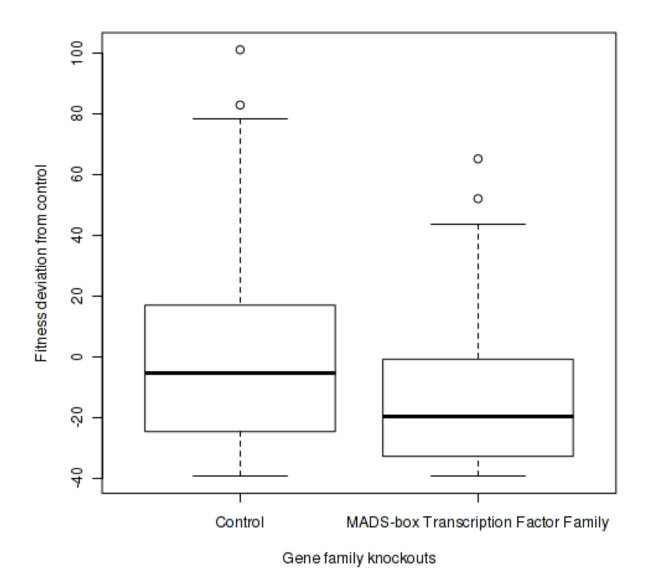
```
FALSE
              68
                   40
     TRUE
               0
                    9
##
  [1] "Numbers of lines that have more extreme fitness than control: no correction"
##
## sig
           FALSE TRUE
              41
                   27
     FALSE
              27
     TRUE
```

1.2.4 Multiple test approach using Dunnett's many to one test

The classic post-hoc test for comparing many treatments to a control is Dunnett's test. It corrects for multiple tests.

1.2.5 Specific comparisons among a-priori-chosen subsets of genes

Here is a comparison of the controls to all mads-box genes



Df Sum Sq Mean Sq F value Pr(>F)

GeneFamilyName 1 8841 8841 10.38 0.00146 **

Residuals 227 193427 852

--
Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1

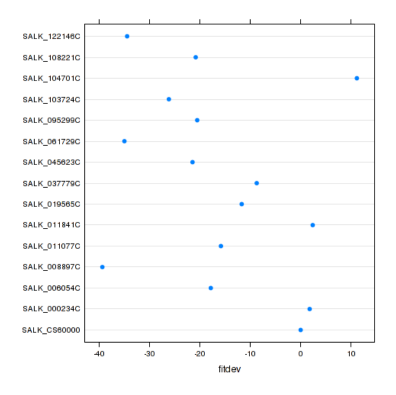
##

Welch Two Sample t-test

data: fitdev by GeneFamilyName

```
## t = 3.263, df = 224.06, p-value = 0.001275
## alternative hypothesis: true difference in means is not equal to 0
## 95 percent confidence interval:
## 4.930097 19.964495
## sample estimates:
## mean in group Control
-1.935837e-15
## mean in group MADS-box Transcription Factor Family
-1.244730e+01
```

Here is the distribution of fitness effects among mads box genes along with a test of among-line differences



```
## Df Sum Sq Mean Sq F value Pr(>F)

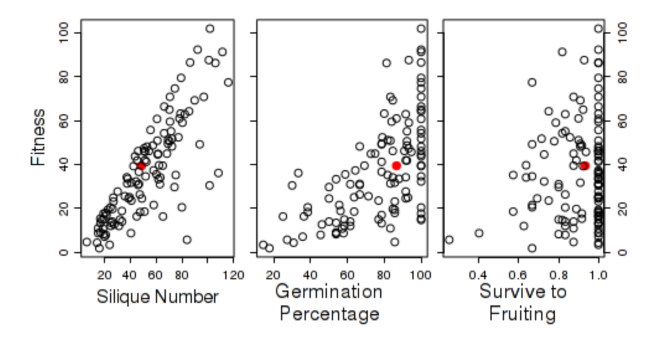
## SALK_Line    14    26332    1880.9    2.288    0.00616 **

## Residuals    214    175937    822.1

## ---

## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

1.3 Fitness components



2 Analysis of fitness

Taking all SALK lines and pooling them and comparing to the control:

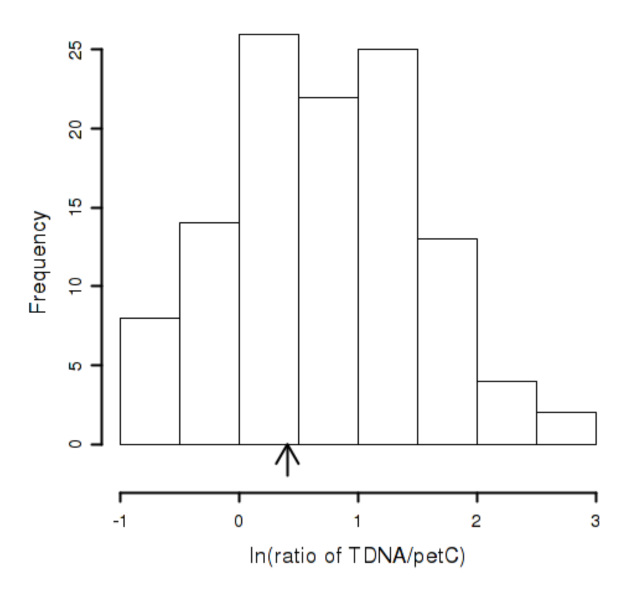
You can see from the resampled line distributions that salk lines and controls have similar mean fitnesses, with definite differences in variance among groups

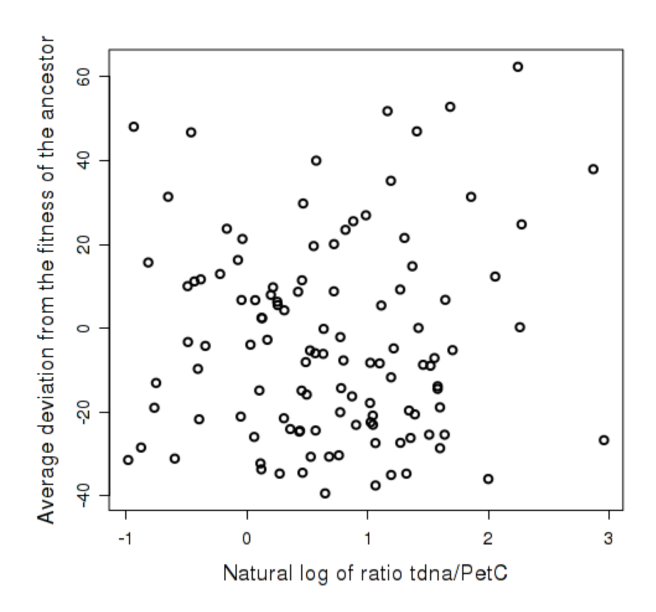
```
##
    Welch Two Sample t-test
##
##
## data: fitness by treat
## t = -0.56331, df = 157.04, p-value = 0.574
## alternative hypothesis: true difference in means is not equal to 0
## 95 percent confidence interval:
   -7.932356 4.411875
## sample estimates:
## mean in group control
                           mean in group treat
                39.39545
                                       41.15569
##
##
##
    Kruskal-Wallis rank sum test
##
          fitmerg$treat and fitmerg$fitness
## Kruskal-Wallis chi-squared = 1041.5, df = 737, p-value = 8.294e-13
```

```
## [1] "permutation typeI prob:"
## [1] 0.724
```

3 Effects of tdna insert number

Here is a plot that relates our total measure of fitness in the lines used in the original pilot study to the ratio of tdna to endogenous genes





```
(Intercept) -4.0472 2.8887 -1.401 0.164
log(area.ratio) 0.8829 2.6995 0.327 0.744

Residual standard error: 23.52 on 112 degrees of freedom
(3 observations deleted due to missingness)

Multiple R-squared: 0.0009542, Adjusted R-squared: -0.007966
F-statistic: 0.107 on 1 and 112 DF, p-value: 0.7442
```

Clearly no pattern there and a regression confirms.

Just to make sure, I also lumped the ratios into categories and looked for a pattern again: When focusing on medians, it looks a little bit like there might be some variation across categories

```
Df Sum Sq Mean Sq F value Pr(>F)
as.factor(ratiocat) 2 187 93.3 0.168 0.846
Residuals 111 61805 556.8
3 observations deleted due to missingness
```

Alas, no pattern there either. I might suggest that, at the least, we worry about copy number less than other factors when choosing lines for UnPAK projects

4 Fitness as a function of gene family size

In the original pilot study we chose genes from different families with differing sizes. Again, there does not seem to be a significant relationship between gene family size and reproductive output in the non-regulatory genes, but there does seem to be a slight pattern in the regulatory genes.

This figure is based on the number of genes in the Gene Family data on tair

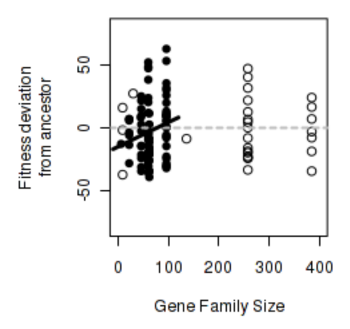
This is the same figure but with the means of each line plotted instead of all the reps for each line.

```
Call:
lm(formula = fitdev ~ FamilySize * Regulatory, data = fruit.by.line)

Residuals:
Min 1Q Median 3Q Max
-37.50 -17.46 -1.68 15.80 60.80

Coefficients:
```

```
Estimate Std. Error t value Pr(>|t|)
(Intercept)
                         0.031853 10.582719 0.003
                                                      0.998
FamilySize
                        -0.004502 0.038440 -0.117
                                                      0.907
Regulatoryyes
                       -16.152593 12.906961 -1.251
                                                     0.213
FamilySize:Regulatoryyes 0.188779 0.112813 1.673 0.097 .
Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
Residual standard error: 23.2 on 112 degrees of freedom
  (1 observation deleted due to missingness)
Multiple R-squared: 0.02922, Adjusted R-squared: 0.003212
F-statistic: 1.124 on 3 and 112 DF, p-value: 0.3427
Call:
lm(formula = fitdev ~ FamilySize, data = fruit.by.line, subset = Regulatory ==
    "yes")
Residuals:
   Min
          1Q Median 3Q
                                  Max
-34.700 -16.930 -1.639 15.799 60.804
Coefficients:
           Estimate Std. Error t value Pr(>|t|)
(Intercept) -16.1207 7.3637 -2.189 0.0313 *
FamilySize 0.1843
                      0.1057 1.743 0.0848 .
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
Residual standard error: 23.12 on 86 degrees of freedom
  (1 observation deleted due to missingness)
Multiple R-squared: 0.03414, Adjusted R-squared: 0.0229
F-statistic: 3.039 on 1 and 86 DF, p-value: 0.08484
```



And some analyses on the means of each line's fitness deviation from the ancestor: First OLS ancova. Then regressions for non-regulatory and then regulatory genes

```
summary(aov(fitdev~FamilySize*Regulatory,fruit.by.line))
                       Df Sum Sq Mean Sq F value Pr(>F)
FamilySize
                        1
                             306
                                   306.3
                                           0.569 0.452
Regulatory
                        1
                               1
                                     0.7
                                           0.001 0.972
FamilySize:Regulatory
                        1
                            1507
                                  1507.0
                                           2.800 0.097 .
Residuals
                           60277
                                   538.2
                      112
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
1 observation deleted due to missingness
summary(lm(fitdev~FamilySize,subset=Regulatory=="no",fruit.by.line))
Call:
lm(formula = fitdev ~ FamilySize, data = fruit.by.line, subset = Regulatory ==
    "no")
Residuals:
   Min
                 Median
                             3Q
             1Q
                                    Max
-37.504 -18.043 -1.843 16.262
                                47.859
```

```
Coefficients:
            Estimate Std. Error t value Pr(>|t|)
(Intercept) 0.031853 10.701052 0.003 0.998
FamilySize -0.004502 0.038869 -0.116
                                          0.909
Residual standard error: 23.46 on 26 degrees of freedom
  (1 observation deleted due to missingness)
Multiple R-squared: 0.0005156, Adjusted R-squared: -0.03793
F-statistic: 0.01341 on 1 and 26 DF, p-value: 0.9087
summary(lm(fitdev~FamilySize,subset=Regulatory=="yes",fruit.by.line))
Call:
lm(formula = fitdev ~ FamilySize, data = fruit.by.line, subset = Regulatory ==
   "yes")
Residuals:
         1Q Median 3Q
-34.700 -16.930 -1.639 15.799 60.804
Coefficients:
           Estimate Std. Error t value Pr(>|t|)
(Intercept) -16.1207 7.3637 -2.189 0.0313 *
FamilySize
           0.1843
                       0.1057 1.743
                                        0.0848 .
Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
Residual standard error: 23.12 on 86 degrees of freedom
  (1 observation deleted due to missingness)
Multiple R-squared: 0.03414, Adjusted R-squared: 0.0229
F-statistic: 3.039 on 1 and 86 DF, p-value: 0.08484
summary(lm(log(fitdev+40)~FamilySize,subset=Regulatory=="yes",fruit.by.line))
Call:
lm(formula = log(fitdev + 40) ~ FamilySize, data = fruit.by.line,
   subset = Regulatory == "yes")
Residuals:
   Min
            1Q Median
                        3Q
-3.7990 -0.5099 0.2162 0.6402 1.2325
```

```
Coefficients:

Estimate Std. Error t value Pr(>|t|)

(Intercept) 3.02358  0.27102  11.156  <2e-16 ***

FamilySize  0.00439  0.00389  1.128  0.262

---

Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1

Residual standard error: 0.8509 on 86 degrees of freedom

(1 observation deleted due to missingness)

Multiple R-squared: 0.01459,Adjusted R-squared: 0.00313

F-statistic: 1.273 on 1 and 86 DF, p-value: 0.2623
```

Here is a little more sophisticated analysis of the family size effect

```
fm1 <- lme(fixed=fitdev~FamilySize,random=~1|SALK_Line,method="ML",subset=Regulatory=="y
#plot(fm1)
vfix <- varFixed(~FamilySize)</pre>
fm2 <- lme(fixed=fitdev~FamilySize,random=~1|SALK_Line,method="ML",subset=Regulatory=="y
#plot(resid(fm2, type="pearson")~fitted(fm2))
fm3 <- lme(fixed=fitdev~1,random=~1|SALK_Line,method="ML",subset=Regulatory=="yes",na.ac
anova(fm3,fm1,fm2) #looks like fm1 is the best model
      Model df
##
                   AIC BIC
                                   logLik Test L.Ratio p-value
       1 3 7933.316 7947.407 -3963.658
## fm1
         2 4 7931.925 7950.713 -3961.962 1 vs 2 3.390619 0.0656
## fm2
         3 4 7985.516 8004.304 -3988.758
anova(fm1)
              numDF denDF F-value p-value
## (Intercept)
                1
                     722 1.105524 0.2934
                      86 3.460323 0.0663
## FamilySize
                 1
anova(fm2)
##
              numDF denDF F-value p-value
## (Intercept) 1 722 1.276404 0.2589
                      86 3.534842 0.0635
## FamilySize
                 1
anova(fm3)
              numDF denDF F-value p-value
## (Intercept) 1 722 1.100184 0.2946
```

Ok, here is the analysis of family size with line as random intercept. Terrible heteroscedacity, repaired using a fixed variance structure. No signal of family size in the final model.

Now, the joint categories approach: Two tests. The first assumes that the rows and columns are independent, but the expected values come from the marginal totals. The second assumes that the number of fitneses in each of the four categories is equal.

```
Pearson's Chi-squared test with Yates' continuity correction

data: tbl

X-squared = 2.3999, df = 1, p-value = 0.1213

[1] 6.025056e-07
```

Here is a test of the change in variance through time:

```
brks=c(0, seq(10, 150, 10), 166)
family.size.cat <- cut(fitmerg$FamilySize,breaks=brks)</pre>
sds <- with(fitmerg,tapply(fitdev,family.size.cat,sd))</pre>
brkmid <- (brks+(c(brks[-1],166)-brks)/2)[-length(sds)]
bartlett.test(fitmerg$fitdev,family.size.cat)
##
   Bartlett test of homogeneity of variances
##
##
## data: fitmerg$fitdev and family.size.cat
## Bartlett's K-squared = 38.058, df = 6, p-value = 1.095e-06
plot(sds~brkmid)
summary(lm(sds~brkmid))
##
## Call:
## lm(formula = sds ~ brkmid)
##
## Residuals:
      (0,10]
##
               (20,30]
                          (40,50]
                                    (50,60]
                                               (60,70] (90,100] (130,140]
      -7.426
                            2.664
                                     11.247
##
                -5.119
                                               -2.590
                                                          14.254
                                                                    -13.031
##
## Coefficients:
               Estimate Std. Error t value Pr(>|t|)
##
## (Intercept) 38.7467
                             7.4525
                                      5.199 0.00347 **
                -0.1638
                             0.1023 -1.602 0.17017
## brkmid
## ---
```

```
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 10.91 on 5 degrees of freedom
## (9 observations deleted due to missingness)
## Multiple R-squared: 0.339,Adjusted R-squared: 0.2069
## F-statistic: 2.565 on 1 and 5 DF, p-value: 0.1702
```

So not much change in variance, though not a lot of power either.

```
Welch Two Sample t-test

data: fitdev by Regulatory

t = 1.2283, df = 398.63, p-value = 0.2201

alternative hypothesis: true difference in means is not equal to 0

95 percent confidence interval:

-1.900829 8.231144

sample estimates:

mean in group no mean in group yes

4.215966 1.050808
```

5 Multiple environment experiment

5.1 Figures for the 1st multi-environment experiment

This figure is fitness deviation ignoring germination (we don't have per-sowed seed estimates (in other words, replicated) of germination for the first experiment, just average germination for that line for that germination effort. We do have those data for the second three treatments/time points

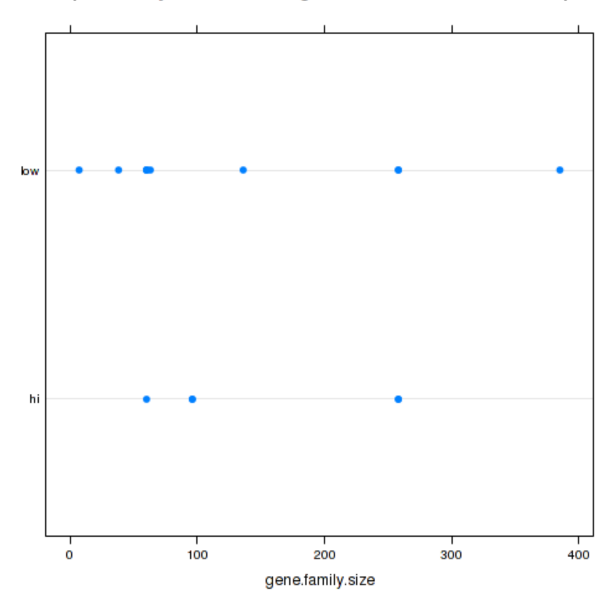
```
[1] "SALK_017933C"
[1] "SALK_033462C"
[1] "SALK_038957C"
[1] "SALK_042704C"
[1] "SALK_050488C"
[1] "SALK_059835C"
[1] "SALK_063722C"
[1] "SALK_063722C"
[1] "SALK_126600C"
[1] "SALK_126600C"
[1] "SALK_134535C"
[1] "SALK_150522C"
```

[1] "CS60000"

I'm going to try and address courtney's question about the gene family size of high and low lines

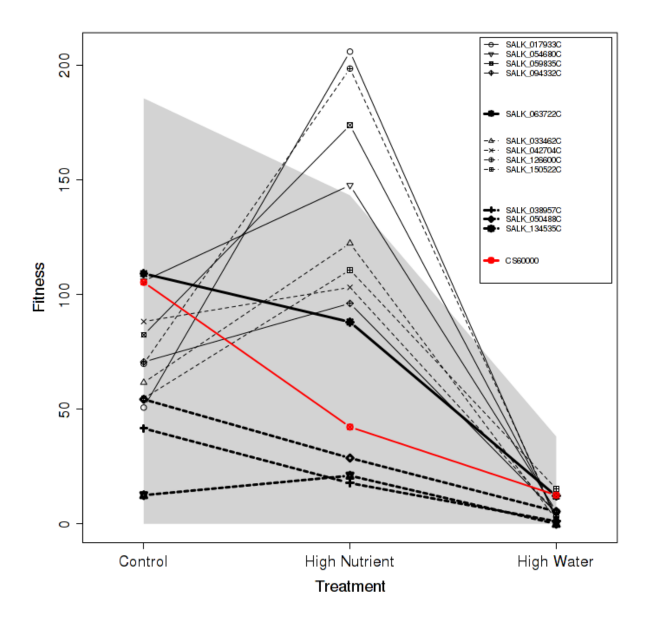
```
## [1] "SALK_Line" "fitdev" "gene.family.size" ## [4] "lohi"
```

Compare family sizes for the high and low lines chosen for exp2



Now here is the figure with straight mean fruit number per line:

```
[1] "SALK_017933C"
[1] 1
[1] "SALK_033462C"
[1] 1
[1] "SALK_038957C"
[1] 1
[1] "SALK_042704C"
[1] 1
[1] "SALK_050488C"
[1] 1
[1] "SALK_054680C"
[1] 1
[1] "SALK_059835C"
[1] 1
[1] "SALK_063722C"
[1] 1
[1] "SALK_094332C"
[1] 1
[1] "SALK_126600C"
[1] 1
[1] "SALK_134535C"
[1] 1
[1] "SALK_150522C"
[1] 1
[1] "CS60000"
[1] 2
```



5.1.1 Copy number

In the following figure line width is proportional to copy number category

5.2 Various tests of GxE

```
#MixedEffects
fit1 <- lmer(fitdevng~1+(1|SALK_Line),subset=SALK_Line!="CS60000",data=intresults)</pre>
```

```
fit2 <- lmer(fitdevng~treattype+(1|SALK_Line), subset=SALK_Line!="CS60000", data=intresult
fit3 <- lmer(fitdevng~treattype+treattype:SALK_Line+(1|SALK_Line),subset=SALK_Line!="CS6
anova(fit1,fit2,fit3)
refitting model(s) with ML (instead of REML)
Data: intresults
Subset: SALK_Line != "CS60000"
Models:
fit1: fitdevng ~ 1 + (1 | SALK_Line)
fit2: fitdevng ~ treattype + (1 | SALK_Line)
fit3: fitdevng ~ treattype + treattype:SALK_Line + (1 | SALK_Line)
          AIC BIC logLik deviance Chisq Chi Df Pr(>Chisq)
fit1 3 4643.4 4655.4 -2318.7 4637.4
fit2 6 4589.9 4613.9 -2288.9 4577.9 59.548
                                                3 7.343e-13 ***
fit3 50 4578.4 4778.1 -2239.2 4478.4 99.466 44 3.555e-06 ***
Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
#fit using OLS
fitaov1 <- aov(fitdevng~treattype*SALK_Line,subset=SALK_Line!="CS60000",data=intresults)
summary(fitaov1)
                    Df Sum Sq Mean Sq F value Pr(>F)
                     3 350733 116911 24.814 1.37e-14 ***
treattype
SALK_Line
                    11 191431 17403 3.694 5.42e-05 ***
treattype:SALK_Line 33 329706 9991 2.121 0.000476 ***
Residuals
                   353 1663151
                                4711
Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
63 observations deleted due to missingness
#fit using ML
fitaov1.glm <-glm(fitdevng~treattype*SALK_Line,subset=SALK_Line!="CS60000",data=intresul
anova(fitaov1.glm,test="Chisq")
Analysis of Deviance Table
Model: gaussian, link: identity
Response: fitdevng
```

```
Terms added sequentially (first to last)
                   Df Deviance Resid. Df Resid. Dev Pr(>Chi)
NULL
                                     400
                                            2535021
treattype
                    3
                        350733
                                     397 2184288 4.771e-16 ***
                                     386 1992857 2.789e-05 ***
SALK_Line
                   11 191431
                                     353 1663151 0.0001816 ***
treattype:SALK_Line 33
                        329706
Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
#now look in each environment and ask if there are line differences
for (trt in unique(intresults$treattype))
   fit.line <- glm(fitdevng~1+as.factor(SALK_Line), subset=SALK_Line!="CS60000", data=int
    fit.intercept <- glm(fitdevng~1, subset=SALK_Line!="CS60000", data=intresults[intresul
    cat(rep("-",30)); cat("\n")
    print (paste("Effect of including line in environment: ",trt))
    print(anova(fit.intercept,fit.line,test="Chisq"))
    cat(rep("-",30)); cat("\n")
  }
[1] "Effect of including line in environment: nutrient"
Analysis of Deviance Table
Model 1: fitdevng ~ 1
Model 2: fitdevng ~ 1 + as.factor(SALK_Line)
  Resid. Df Resid. Dev Df Deviance Pr(>Chi)
1
        98
              1417594
2
        87
              1125616 11 291978 0.02033 *
Signif. codes: 0 '*** 0.001 '** 0.01 '*' 0.05 '.' 0.1 ' ' 1
[1] "Effect of including line in environment: control"
Analysis of Deviance Table
Model 1: fitdevng ~ 1
Model 2: fitdevng ~ 1 + as.factor(SALK_Line)
Resid. Df Resid. Dev Df Deviance Pr(>Chi)
```

```
101 532531
2
        90
            439049 11 93482 0.05823 .
Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
[1] "Effect of including line in environment: highwater"
Analysis of Deviance Table
Model 1: fitdevng ~ 1
Model 2: fitdevng ~ 1 + as.factor(SALK_Line)
 Resid. Df Resid. Dev Df Deviance Pr(>Chi)
       95
              14671
1
2.
       84
              11170 11 3500.2 0.005812 **
Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
[1] "Effect of including line in environment: FIRST.EXP"
Analysis of Deviance Table
Model 1: fitdevng ~ 1
Model 2: fitdevng ~ 1 + as.factor(SALK_Line)
 Resid. Df Resid. Dev Df Deviance Pr(>Chi)
1
    103 219492
2
       92
              87315 11 132177 < 2.2e-16 ***
Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

Here is an analysis with Prior information and ecotypes included

```
results$genotype.cls = rep("control",dim(results)[1])
results$genotype.cls[results$fitdev<0] = "low"
results$genotype.cls[results$fitdev>0] = "high"
results$genotype.cls[results$SALK_Line=="CS60000"] = "control"
results$genotype.cls[grep("ECO",results$SALK_Line)] = "ecotype"

fit1 <- lm(fitness~treattype*genotype.cls,subset=treattype!="FIRST.EXP",data=results)
Anova(fit1,contrasts = list(treattype=contr.sum,genotype.cls=contr.sum),type=3)

Anova Table (Type III tests)</pre>
```

```
Response: fitness
                     Sum Sq Df F value
                                         Pr(>F)
                            1 49.774 7.596e-12 ***
(Intercept)
                     110881
                      44786
                             2 10.052 5.497e-05 ***
treattype
genotype.cls
                     490361 3 73.373 < 2.2e-16 ***
treattype:genotype.cls 330825 6 24.751 < 2.2e-16 ***
Residuals
                     895536 402
Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
fit2 <- lm(fitness~treattype*genotype.cls,subset=(treattype!="FIRST.EXP")&(genotype.cls!
Anova(fit2,contrasts = list(treattype=contr.sum,genotype.cls=contr.sum),type=3)
Anova Table (Type III tests)
Response: fitness
                     Sum Sq Df F value Pr(>F)
                     (Intercept)
treattype
                      44786 2 11.058 2.157e-05 ***
genotype.cls
                     490177 2 121.031 < 2.2e-16 ***
treattype:genotype.cls 328440 4 40.548 < 2.2e-16 ***
Residuals
                     759380 375
Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
```

This analysis just looks for GxE and main effects in the ecotypes. Not much signal Ecotype lines only

```
fitall <- glm(fitdev~treattype*SALK_Line,subset=treattype!="FIRST.EXP",data=intresults)
(anova(fitall,test="F"))
## Analysis of Deviance Table
##</pre>
```

```
## Model: gaussian, link: identity
##
## Response: fitdev
##
## Terms added sequentially (first to last)
##
##
##
                      Df Deviance Resid. Df Resid. Dev
                                                            F
                                                                 Pr(>F)
## NULL
                                        383
                                              3103317
## treattype
                       2
                           647029
                                        381
                                              2456288 59.5646 < 2.2e-16 ***
## SALK_Line
                                              2221538 3.6018 4.360e-05 ***
                      12
                           234750
                                        369
## treattype:SALK_Line 24
                           347732
                                        345
                                              1873806 2.6676 5.317e-05 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
#add in a classifier for early experiment performance
firstmns <- with(intresults[intresults$treattype=="FIRST.EXP",c("fitdev","SALK_Line")],</pre>
                aggregate(cbind(first.fitdev=fitdev), by=list(SALK_Line=SALK_Line), mean,
intresults <- merge(intresults, firstmns, all.x=T)</pre>
fitlo <- glm(fitdev~treattype*SALK_Line, subset=((treattype!="FIRST.EXP")&(first.fitdev<
(anova(fitlo,test="F"))
## Analysis of Deviance Table
## Model: gaussian, link: identity
##
## Response: fitdev
##
## Terms added sequentially (first to last)
##
##
##
                      Df Deviance Resid. Df Resid. Dev
                                                            F
                                                                 Pr(>F)
## NULL
                                        236
                                              1805730
## treattype
                       2
                           278944
                                        234
                                              1526786 25.0251 1.731e-10 ***
                                        227 1379812 3.7673 0.0007137 ***
## SALK_Line
                       7
                           146974
                                        ## treattype:SALK_Line 14
                           192704
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
fithi <- glm(fitdev~treattype*SALK_Line,subset=((treattype!="FIRST.EXP")&(first.fitdev>
(anova(fithi,test="F"))
```

```
## Analysis of Deviance Table
##
## Model: gaussian, link: identity
##
## Response: fitdev
##
## Terms added sequentially (first to last)
##
##
                       Df Deviance Resid. Df Resid. Dev
##
                                                             F
                                                                   Pr(>F)
## NULL
                                         146
                                                1232611
## treattype
                        2
                            428976
                                         144
                                                 803635 41.2297 1.228e-14 ***
## SALK_Line
                                                 785490 0.8720
                        4
                             18145
                                         140
                                                                  0.48267
## treattype:SALK_Line 8
                             98791
                                         132
                                                 686698 2.3738 0.02021 *
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
intresults$lohi <- factor(ifelse(intresults$first.fitdev<=0,"low","high"))</pre>
fitlowhi <- glm(fitdev~treattype*lohi,subset=(treattype!="FIRST.EXP"),data=intresults)</pre>
(anova(fitlowhi,test="F"))
## Analysis of Deviance Table
##
## Model: gaussian, link: identity
##
## Response: fitdev
##
## Terms added sequentially (first to last)
##
##
##
                 Df Deviance Resid. Df Resid. Dev
                                                              Pr(>F)
                                                       F
## NULL
                                    383
                                           3103317
## treattype
                  2
                       647029
                                    381
                                           2456288 52.4749 < 2.2e-16 ***
## lohi
                                           2387726 11.1209 0.0009381 ***
                   1
                       68562
                                    380
## treattype:lohi 2
                       57305
                                    378
                                           2330421 4.6475 0.0101397 *
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
Anova(lm(fitdev~treattype*lohi, subset=(treattype!="FIRST.EXP"), data=intresults), contrast
## Anova Table (Type III tests)
## Response: fitdev
```

```
##
                   Sum Sq Df F value Pr(>F)
## (Intercept)
                    23242
                           1 3.7699
                                        0.05293 .
## treattype
                   428976
                            2 34.7905 1.353e-14 ***
## lohi
                    15911
                          1 2.5808 0.10900
## treattype:lohi
                    57305
                            2 4.6475
                                       0.01014 *
## Residuals
                 2330421 378
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
fiteco <- glm(fitdev~treattype*SALK_Line,data=intresults.ecotypes)</pre>
(anova(fiteco,test="F"))
## Analysis of Deviance Table
##
## Model: gaussian, link: identity
##
## Response: fitdev
##
## Terms added sequentially (first to last)
##
##
##
                       Df Deviance Resid. Df Resid. Dev
                                                             F Pr(>F)
## NULL
                                          29
                                                 154737
                        2
                             18581
                                          27
                                                 136156 1.5574 0.2625
## treattype
## SALK_Line
                        9
                             23496
                                          18
                                                 112660 0.4376 0.8829
## treattype:SALK_Line
                       9
                             58971
                                                  53689 1.0984 0.4456
```

5.3 Tables for the MS

5.3.1 Gene families

```
famtable <- unique(fitmerg[!is.na(fitmerg$Regulatory),c("GeneFamilyName","FamilySize","R
linesfromfams <- with(unique(fitmerg[fitmerg$GeneFamilyName!="Control",c("SALK_Line","Ge
famtable <- merge(famtable,linesfromfams,all.x=T)
famtable <- famtable[order(famtable$Regulatory,-famtable$FamilySize),]
famtable$Function <- ifelse(famtable$Regulatory=="yes","Regulatory","Metabolic")
famtable <- famtable[,-which(names(famtable)=="Regulatory")]

require(xtable)

## Loading required package: xtable
##</pre>
```

5.3.2 SALK Line list

5.4 Test for effect of line on fitness for first exp.

```
summary(aov(log(fitness+1)~SALK_Line,data=fitmerg[grep("SALK.[0-9]+C",fitmerg$SALK_Line)

## Df Sum Sq Mean Sq F value Pr(>F)

## SALK_Line 115 604.2 5.254 2.991 <2e-16 ***

## Residuals 928 1630.4 1.757

## ---

## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1</pre>
```

6 Naturally occuring variants

We took the data from Cao et al 2010 and determined how many of the lines in this experiment also showed some sort of natural variation in gene function

```
suppressMessages(require(dplyr))
snp <- unique(read.csv(paste0(csvdir,"/phen-snp.csv"))[,1:2])
names(snp)[2] <- "snp.strains"
snp <- snp %>% group_by(Accession) %>% summarise(snp.strains.mn=sum(snp.strains))
sv <- unique(read.csv(paste0(csvdir,"/phen-sv.csv"))[,1:2])</pre>
```

```
names(sv)[2] <- "sv.strains"
sv <- sv %>% group_by(Accession) %>% summarise(sv.strains.mn=sum(sv.strains))
write.table(file="cao-digested.csv",sep=",",row.names=F,unique(merge(snp,sv)))
```

There are definitely lines that are knocked out in nature. The first table is the frequency of lines with no natural variants (false) versus variants for SNPs that should drastically alter gene function. The second is for the distribution of lines with large structural variants

```
with(unique(snp),table(snp.strains.mn>0))

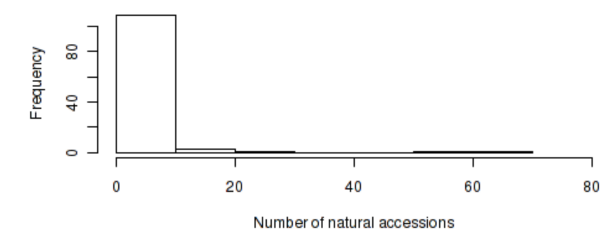
##
## FALSE TRUE
## 99 16

with(unique(sv),table(sv.strains.mn>0))

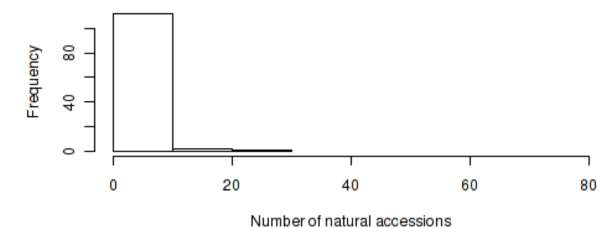
##
## FALSE TRUE
## 106 9
```

The following figure illustrates the distribution of the naturally occurring variants in our

Distribution of natural accessions with large effect SNPs



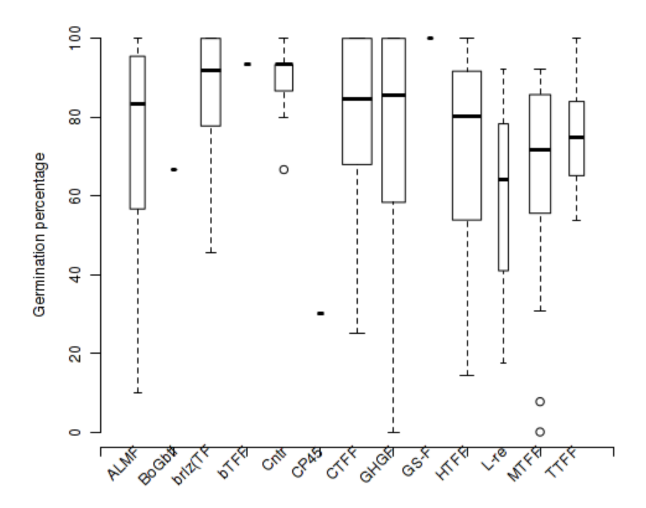
Distribution of natural accessions with large structural varian



collection of lines.

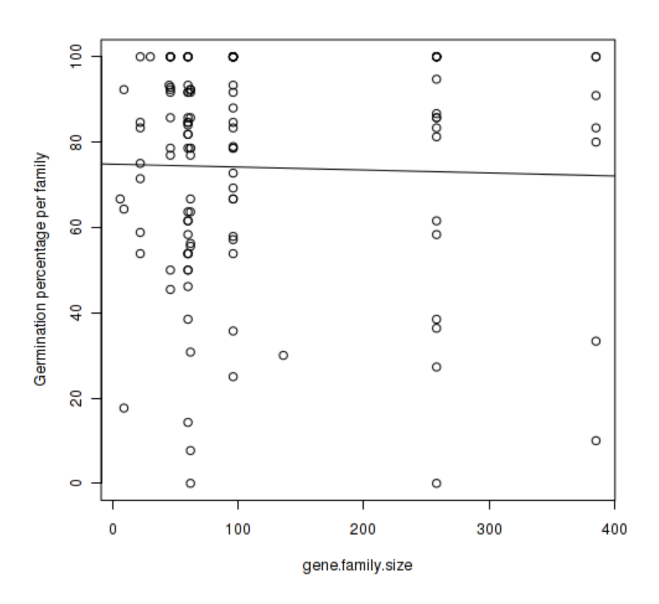
6.1 Some additional comparisons we have not emphasized

6.1.1 Germination rate as a function of gene-family



Gene Family

Though there is variation among families in germination rate, family size does not explain it.



```
Call:
lm(formula = percent ~ (gene.family.size), data = germ.gene.families)

Residuals:
    Min    1Q    Median    3Q     Max
-74.390 -16.074   8.212   18.926   27.838

Coefficients:
    Estimate Std. Error t value Pr(>|t|)
```

```
(Intercept) 74.817309 3.558032 21.03 <2e-16 ***
gene.family.size -0.006896 0.023795 -0.29 0.772
---
Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1

Residual standard error: 25.63 on 115 degrees of freedom
  (16 observations deleted due to missingness)

Multiple R-squared: 0.0007297, Adjusted R-squared: -0.00796
F-statistic: 0.08398 on 1 and 115 DF, p-value: 0.7725
```

And here's another test of the same hypothesis using permutation approach (may be less affected by unequal var).

```
[1] "typeI prob:"
[1] 0.008
```

6.2 Survival to harvest as a function of gene family

Here are analyses that examine the effect of gene family upon survival:

```
Df Sum Sq Mean Sq F value Pr(>F)
GeneFamilyName 12 0.1829 0.01524 0.758 0.692
Residuals 105 2.1122 0.02012
```