Sperm size figures

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Wednesday, January 14, 2015

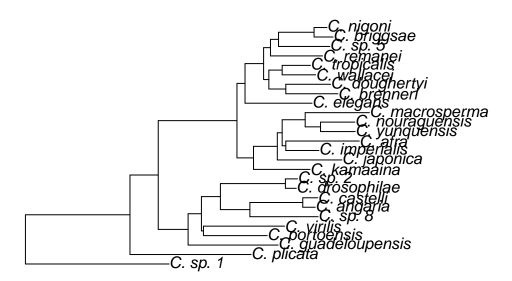
Sperm size figures

This is the output of an R markdown script - a fancy way of embedding r in a page so we can rerun/tune it as needed and then have the commands

I'm embedding, but hiding a bynch of stuff for readability - the actual code is in the .Rmd file.

Warning: package 'survival' was built under R version 3.1.3

Here's the tree from Kiontke et al 2011, figure 3 intial values I recieved had angaria and sp12 flipped - I've corrected it here Here's the tree:



Now read in the compiled data - we have oocytesize, spermsize, primary spermatocytesize, and oocytearea. The species names in the files are inconsistent - they are fixed now. For now, the oocyte area is using the area formulation as made by the initial sperm size formula. All response variables are logged unless otherwise noted.

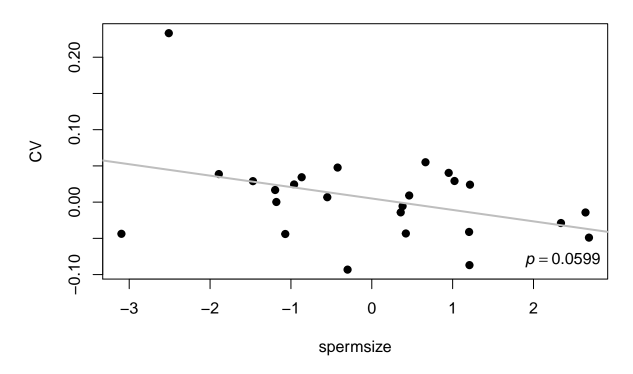
I have filtered out some of the data: Any oocytesize that is not stage -1, and any primaryspermatocyte data that is from hermaphrodites rather than males.

We now have clean data, clustered into species, strains or individuals. Variables ending in 1 are in species, 2 are strains and 3 are individuals. The non-clustered data is as is.

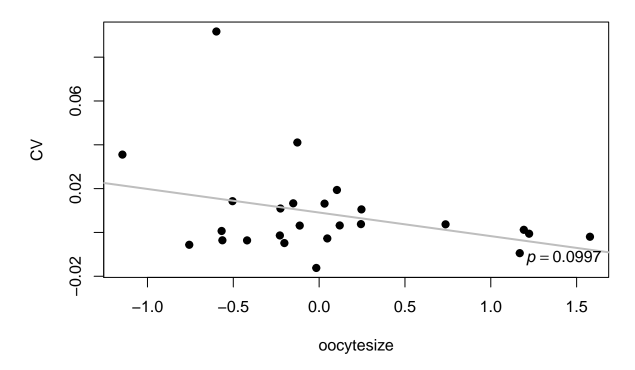
PIC plots - mean vs CV

Plotting the phylogenetic independent contrasts (PIC) of means of traits vs CV of traits. All means and CVs are grouped at strain level.

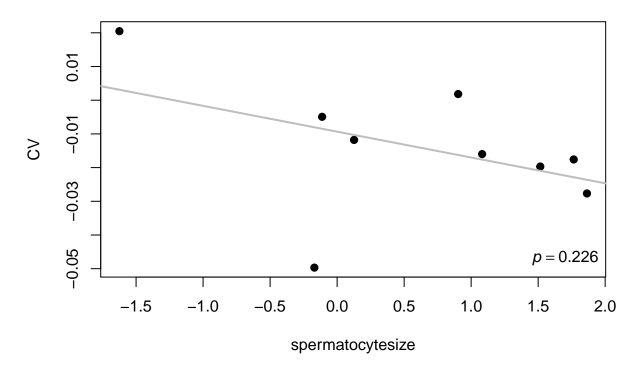
PIC of spermsize means vs CV



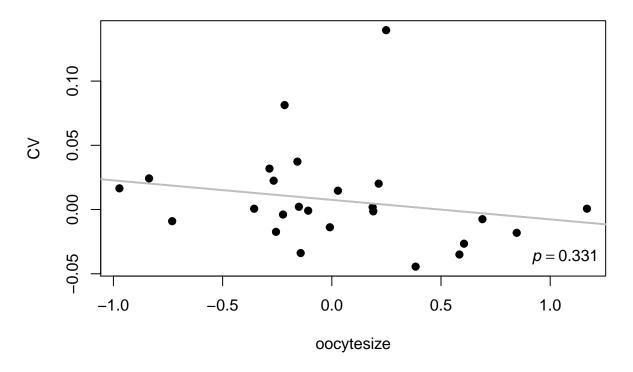
PIC of oocytesize means vs CV



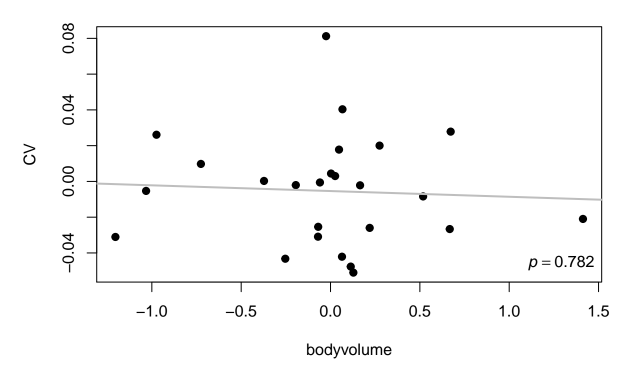
PIC of spermatocytesize means vs CV



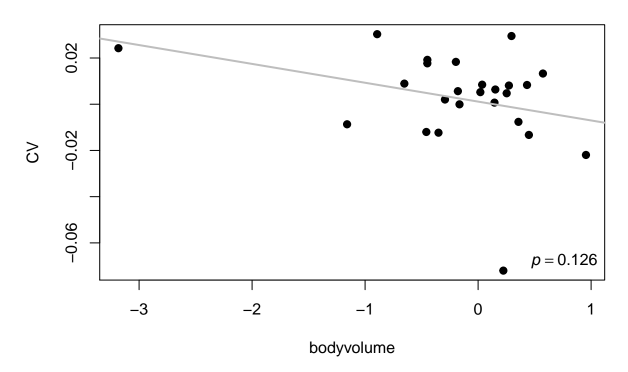
PIC of oocytesizearea means vs CV



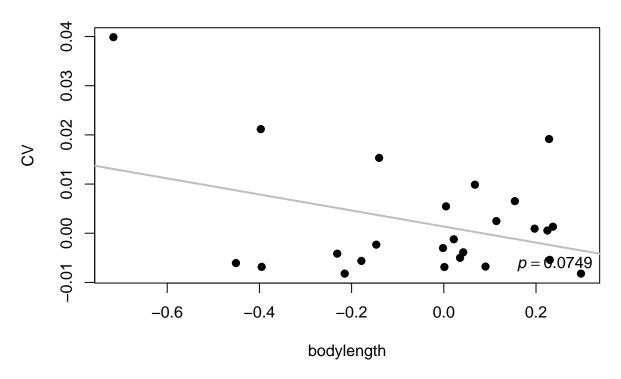
PIC cv bodyvolume male



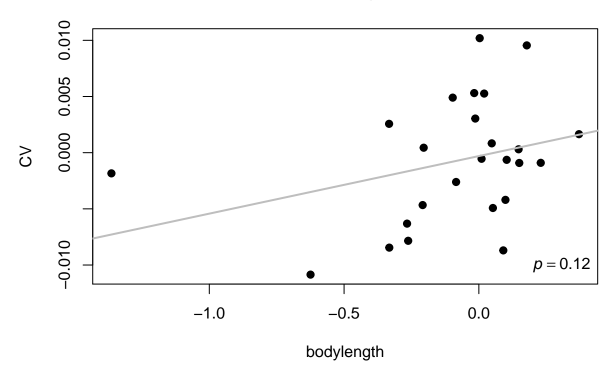
PIC cv bodyvolume female



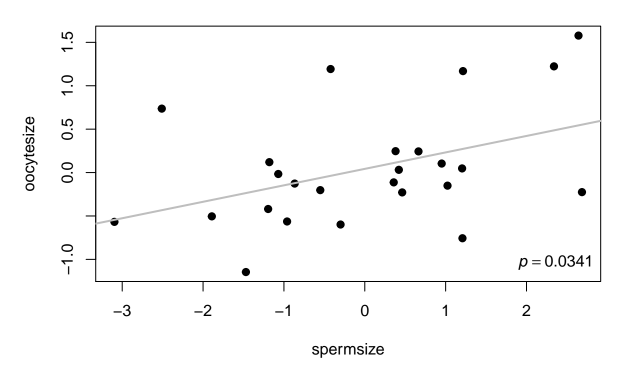
PIC cv bodylength male



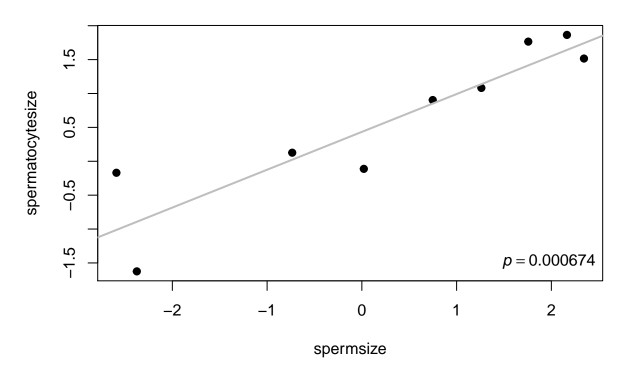
PIC cv bodylength female



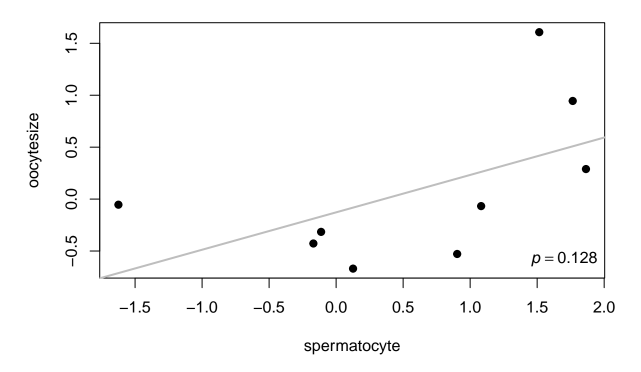
PIC of spermsize vs oocytesize



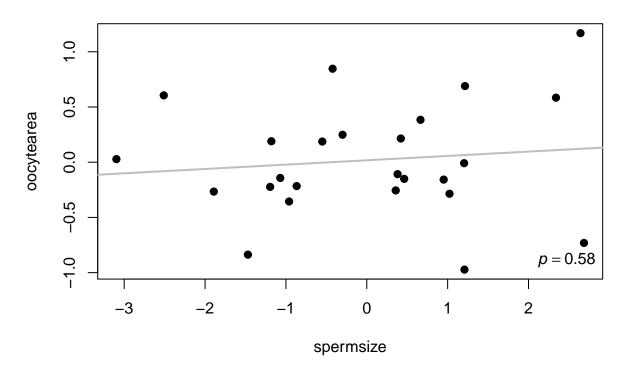
PIC of spermsize vs spermatocytesize



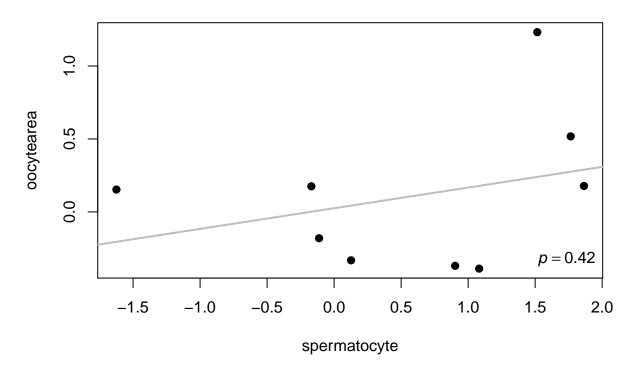
PIC of spermatocytesize vs oocytesize



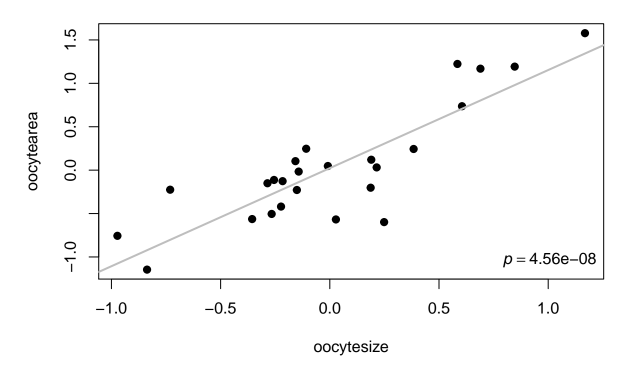
PIC of spermsize vs oocytearea



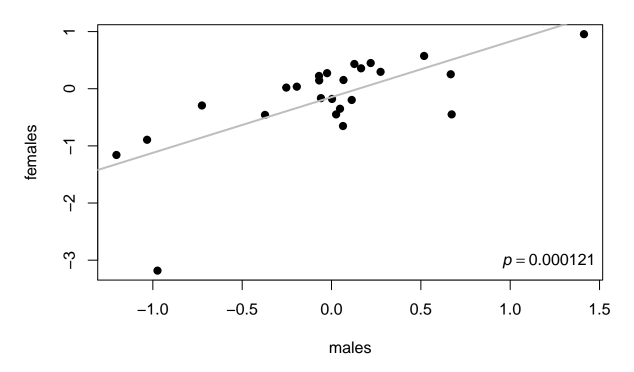
PIC of spermatocytesize vs oocytearea



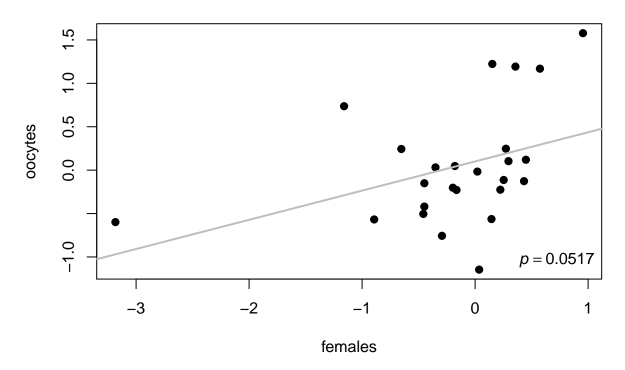
PIC of oocytesize vs oocytearea



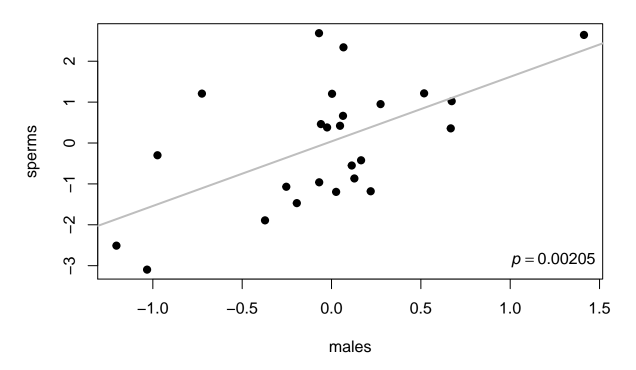
Body area male vs female



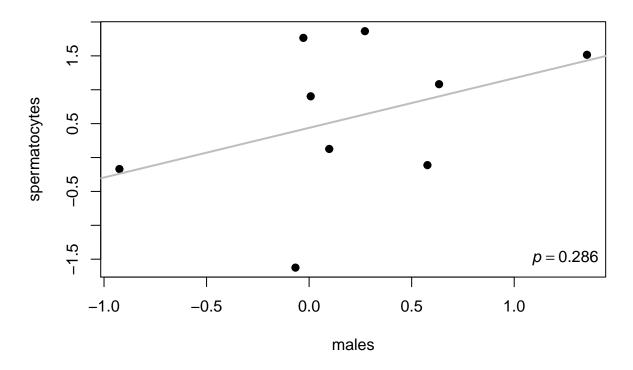
Body area female vs oocytesize



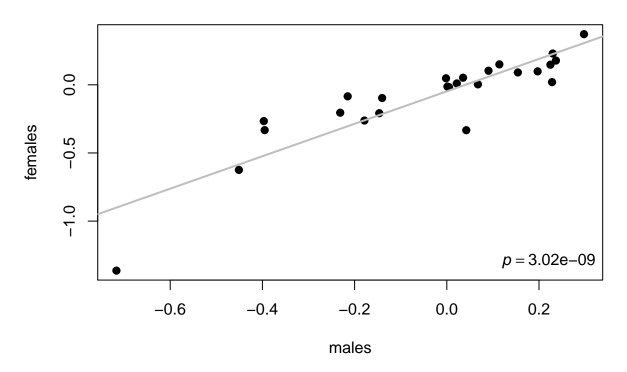
Body area male vs spermsize



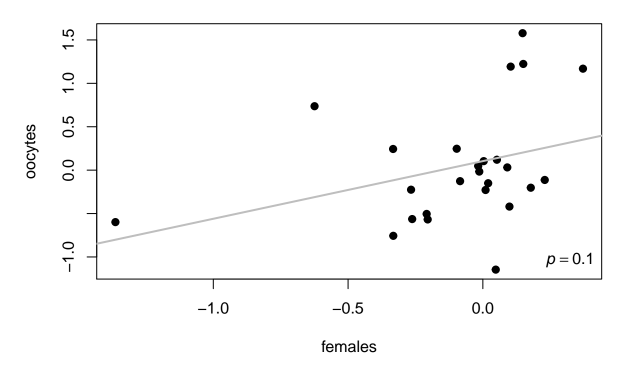
Body area male vs spermatocytesize



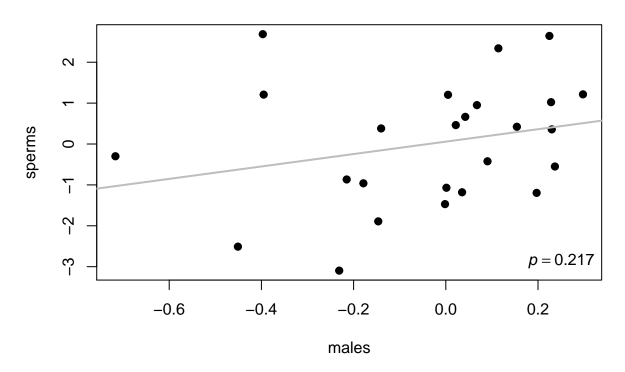
Body length male vs female



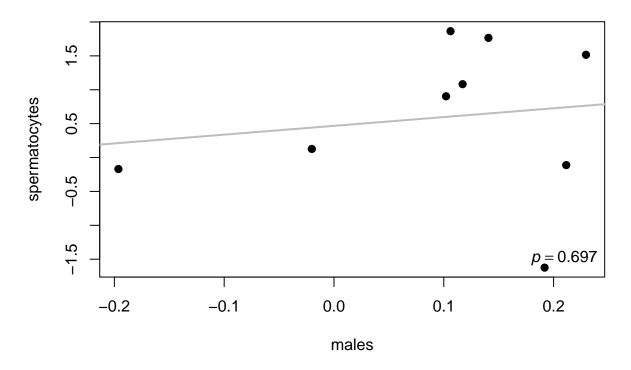
Body length female vs oocytesize



Body length male vs spermsize



Body length male vs spermatocytesize



PGLS - testing for correlation with evolutionary models

Output are summary tables from fits. The OU model won't fit the spermatocyte fits due to overparameterisation.

```
##
## pgls(formula = meansperm ~ meanooarea, data = compsperm, lambda = "ML")
##
## Residuals:
##
                1Q
                   Median
                                3Q
                                       Max
  -0.9643 -0.5988 0.4323
                            0.6861
##
                                    1.8263
##
## Branch length transformations:
##
## kappa [Fix]
                : 1.000
  lambda [ ML]
                : 0.000
      lower bound : 0.000, p = 1
##
##
      upper bound : 1.000, p = 0.11287
      95.0% CI
##
                 : (NA, NA)
## delta [Fix]
                : 1.000
##
## Coefficients:
               Estimate Std. Error t value Pr(>|t|)
## (Intercept) -6.09037
                           6.57159 -0.9268
                                             0.3812
```

```
## meanooarea 1.46171
                          0.94975 1.5390 0.1624
##
## Residual standard error: 1.057 on 8 degrees of freedom
## Multiple R-squared: 0.2284, Adjusted R-squared: 0.132
## F-statistic: 2.369 on 1 and 8 DF, p-value: 0.1624
##
## Call:
## pgls(formula = meansperm ~ meanoo, data = compsperm, lambda = "ML")
## Residuals:
##
      Min
               1Q Median
                               3Q
                                      Max
## -1.4586 -1.1960 0.6374 1.0989 1.4686
##
## Branch length transformations:
##
## kappa [Fix] : 1.000
## lambda [ ML] : 0.895
##
     lower bound : 0.000, p = 1
     upper bound : 1.000, p = 0.79696
##
     95.0% CI : (NA, NA)
## delta [Fix] : 1.000
##
## Coefficients:
               Estimate Std. Error t value Pr(>|t|)
## (Intercept) -11.31303
                           6.94067 -1.6300 0.14176
## meanoo
                1.50685
                           0.65983 2.2837 0.05177 .
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 1.315 on 8 degrees of freedom
## Multiple R-squared: 0.3946, Adjusted R-squared: 0.319
## F-statistic: 5.215 on 1 and 8 DF, p-value: 0.05177
##
## Call:
## pgls(formula = meansperm ~ meanprimsp, data = compsperm, lambda = "ML")
## Residuals:
      Min
               1Q Median
                               30
                                      Max
## -0.8244 -0.3185 -0.1789 0.1091 0.5614
## Branch length transformations:
##
## kappa [Fix] : 1.000
## lambda [ ML] : 0.000
##
     lower bound : 0.000, p = 1
     upper bound : 1.000, p = 0.0066141
##
     95.0% CI
               : (NA, 0.708)
## delta [Fix] : 1.000
## Coefficients:
              Estimate Std. Error t value Pr(>|t|)
## (Intercept) -2.63590
                          0.94589 -2.7867 0.0236793 *
```

```
0.17131 7.0873 0.0001033 ***
## meanprimsp
              1.21417
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 0.446 on 8 degrees of freedom
## Multiple R-squared: 0.8626, Adjusted R-squared: 0.8454
## F-statistic: 50.23 on 1 and 8 DF, p-value: 0.0001033
## Generalized least squares fit by REML
     Model: meansperm ~ meanooarea
##
     Data: fullpglsframe
##
         AIC
                  BIC
                         logLik
##
     54.95884 59.67106 -23.47942
##
## Correlation Structure: corMartins
## Formula: ~1
## Parameter estimate(s):
##
     alpha
## 6.841132
##
## Coefficients:
##
                  Value Std.Error
                                     t-value p-value
## (Intercept) -3.044212 3.398811 -0.8956698 0.3793
## meanooarea
              1.005070 0.498035 2.0180725 0.0549
##
## Correlation:
##
              (Intr)
## meanooarea -0.999
##
## Standardized residuals:
##
         Min
                     Q1
                                           QЗ
                                                     Max
                               Med
## -1.7775727 -0.6837636 -0.2886598 0.4237510
                                              2.2862305
## Residual standard error: 0.6334179
## Degrees of freedom: 26 total; 24 residual
## Generalized least squares fit by REML
    Model: meansperm ~ meanoo
##
     Data: fullpglsframe
         AIC
##
                  BIC
                         logLik
    50.30285 55.01506 -21.15142
##
## Correlation Structure: corMartins
## Formula: ~1
## Parameter estimate(s):
##
      alpha
## 5.469326
##
## Coefficients:
                  Value Std.Error t-value p-value
## (Intercept) -7.068653 3.346328 -2.112361 0.0452
## meanoo
               1.074887 0.329803 3.259179 0.0033
##
## Correlation:
```

```
(Intr)
## meanoo -0.999
##
## Standardized residuals:
          Min
                                Med
                                             QЗ
## -1.9084685 -0.7013875 -0.3474741 0.5038124 2.4034645
## Residual standard error: 0.5843052
## Degrees of freedom: 26 total; 24 residual
Some phylogenetic signal methods - lambda
## Loading required package: parallel
       Length Class
                         Mode
## lik
         1
              bm
                         function
## bnd
              data.frame list
## res 400
              -none-
                         numeric
## opt
              -none-
                         list
##
       Length Class
                         Mode
## lik
         1
              bm
                         function
## bnd
         2
              data.frame list
## res 400
              -none-
                         numeric
## opt
         8
              -none-
                         list
       Length Class
                         Mode
## lik
         1
              bm
                         function
## bnd
         2
              data.frame list
## res 400
              -none-
                         numeric
## opt
         8
              -none-
                         list
##
       Length Class
                         Mode
## lik
         1
              bm
                         function
## bnd
         2
              data.frame list
## res 400
              -none-
                         numeric
## opt
              -none-
                         list
Blombergs K
             K PIC.variance.obs PIC.variance.rnd.mean PIC.variance.P
## 1 0.3738911
                       2.289904
                                              3.977816
                                                                 0.035
   PIC.variance.Z
## 1
          -1.469807
            K PIC.variance.obs PIC.variance.rnd.mean PIC.variance.P
## 1 0.670685
                                             1.233429
                     0.4572932
                                                                0.002
   PIC.variance.Z
```

1

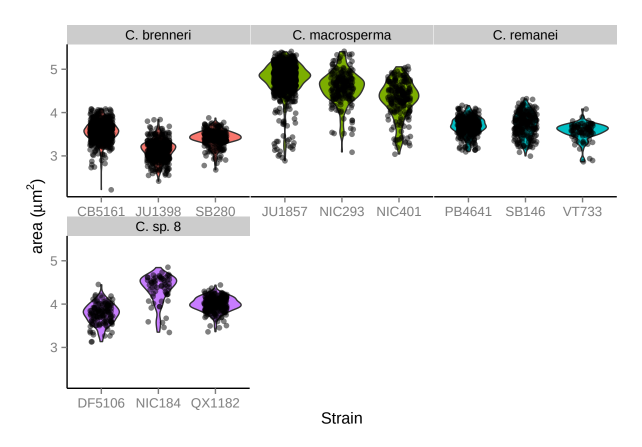
-1.950999

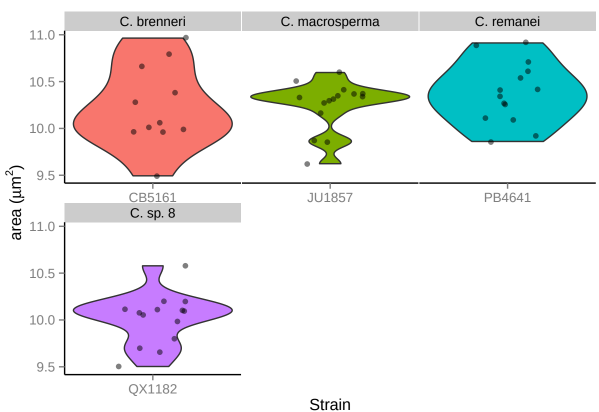
```
## [1] "Dropping tips from the tree because they are not present in the data:"
## [1] "C. sp. 1"
                         "C. portoensis"
                                           "C. virilis"
                         "C. angaria"
## [4] "C. sp. 8"
                                           "C. castelli"
## [7] "C. sp. 2"
                         "C. kamaaina"
                                           "C. imperialis"
## [10] "C. afra"
                         "C. nouraguensis" "C. doughertyi"
## [13] "C. wallacei"
                         "C. tropicalis"
                                           "C. sp. 5"
## [16] "C. nigoni"
           K PIC.variance.obs PIC.variance.rnd.mean PIC.variance.P
##
## 1 1.218587
                     1.696502
                                           2.558416
## PIC.variance.Z
## 1
         -1.816749
            K PIC.variance.obs PIC.variance.rnd.mean PIC.variance.P
## 1 0.5877787
                     0.2622437
                                          0.6080878
                                                              0.003
## PIC.variance.Z
## 1
         -1.781835
Morans I
## class: krandtest
## Monte-Carlo tests
## Call: as.krandtest(sim = matrix(res$result, ncol = nvar, byrow = TRUE),
      obs = res$obs, alter = alter, names = test.names)
##
## Number of tests: 1
## Adjustment method for multiple comparisons:
                                                none
## Permutation number:
                        999
   Test
                 0bs
                      Std.Obs Alter Pvalue
## 1 x -0.09396389 -1.458833 greater 0.942
## other elements: adj.method call
## class: krandtest
## Monte-Carlo tests
## Call: as.krandtest(sim = matrix(res$result, ncol = nvar, byrow = TRUE),
      obs = res$obs, alter = alter, names = test.names)
##
## Number of tests: 1
##
## Adjustment method for multiple comparisons:
                                                none
## Permutation number:
                        999
                 Obs Std.Obs Alter Pvalue
## Test
## 1
      x -0.01730539 0.632097 greater 0.194
## other elements: adj.method call
## class: krandtest
## Monte-Carlo tests
## Call: as.krandtest(sim = matrix(res$result, ncol = nvar, byrow = TRUE),
      obs = res$obs, alter = alter, names = test.names)
```

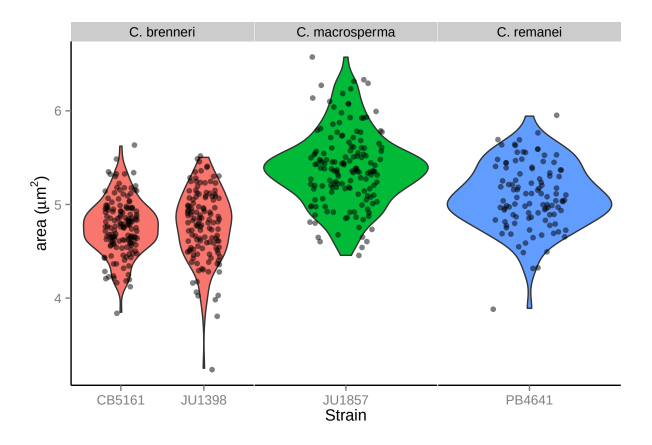
```
##
## Number of tests: 1
##
## Adjustment method for multiple comparisons:
                                                none
## Permutation number:
                        999
    Test
                Obs
                       Std.Obs
                                Alter Pvalue
       x -0.1482137 -0.9786967 greater 0.828
##
## other elements: adj.method call
## class: krandtest
## Monte-Carlo tests
## Call: as.krandtest(sim = matrix(res$result, ncol = nvar, byrow = TRUE),
##
       obs = res$obs, alter = alter, names = test.names)
##
## Number of tests: 1
## Adjustment method for multiple comparisons:
                                                none
## Permutation number:
                        999
    Test
                 Obs
                       Std.Obs Alter Pvalue
       x -0.03082416 0.2906171 greater 0.329
## other elements: adj.method call
abouheifs cmean
## class: krandtest
## Monte-Carlo tests
## Call: as.krandtest(sim = matrix(res$result, ncol = nvar, byrow = TRUE),
       obs = res$obs, alter = alter, names = test.names)
##
## Number of tests: 1
## Adjustment method for multiple comparisons:
## Permutation number:
                        999
## Test
                 Obs Std.Obs Alter Pvalue
## 1
       x -0.09396389 -1.465657 greater 0.944
## other elements: adj.method call
## class: krandtest
## Monte-Carlo tests
## Call: as.krandtest(sim = matrix(res$result, ncol = nvar, byrow = TRUE),
##
      obs = res$obs, alter = alter, names = test.names)
## Number of tests: 1
## Adjustment method for multiple comparisons:
## Permutation number:
                        999
## Test
                 Obs
                       Std.Obs Alter Pvalue
## 1
       x -0.01730539 0.6468218 greater 0.193
## other elements: adj.method call
```

```
## class: krandtest
## Monte-Carlo tests
## Call: as.krandtest(sim = matrix(res$result, ncol = nvar, byrow = TRUE),
##
       obs = res$obs, alter = alter, names = test.names)
## Number of tests:
## Adjustment method for multiple comparisons:
                                                none
## Permutation number:
                        999
    Test
                 Obs
                        Std.Obs Alter Pvalue
## 1
       x -0.1482137 -0.9847184 greater 0.827
## other elements: adj.method call
## class: krandtest
## Monte-Carlo tests
## Call: as.krandtest(sim = matrix(res$result, ncol = nvar, byrow = TRUE),
       obs = res$obs, alter = alter, names = test.names)
##
## Number of tests:
                    1
## Adjustment method for multiple comparisons:
## Permutation number:
                        999
                       Std.Obs Alter Pvalue
    Test
                 Obs
## 1
       x -0.03082416 0.2965012 greater 0.335
##
## other elements: adj.method call
```

figure 3A with violin plots







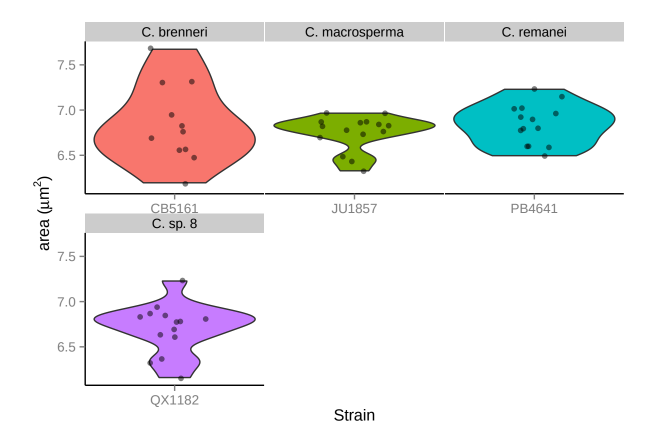
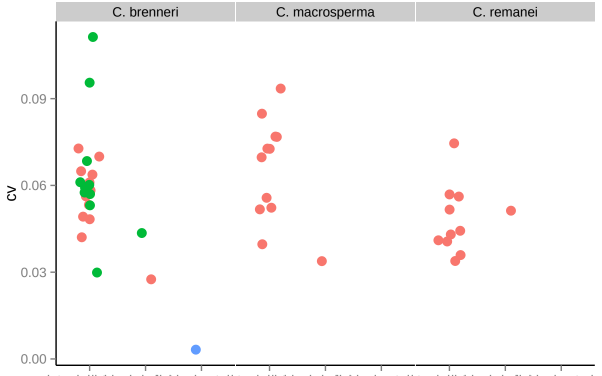
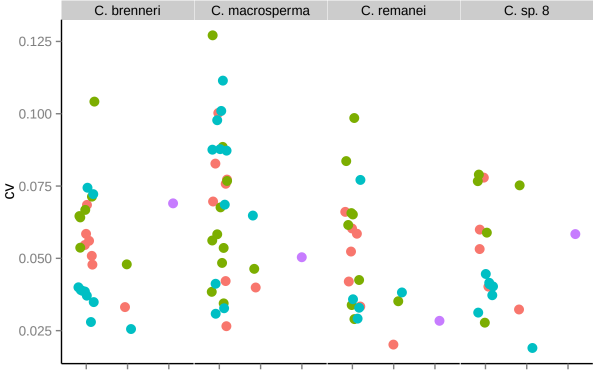


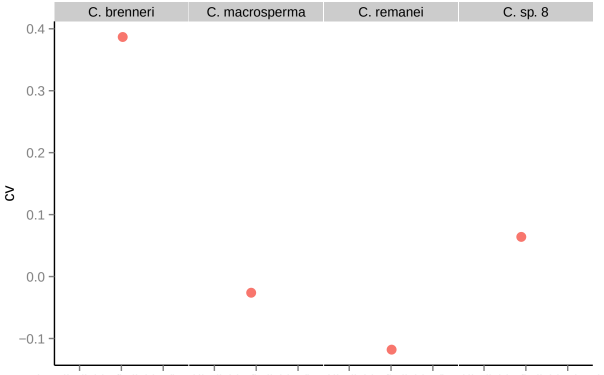
figure 3B



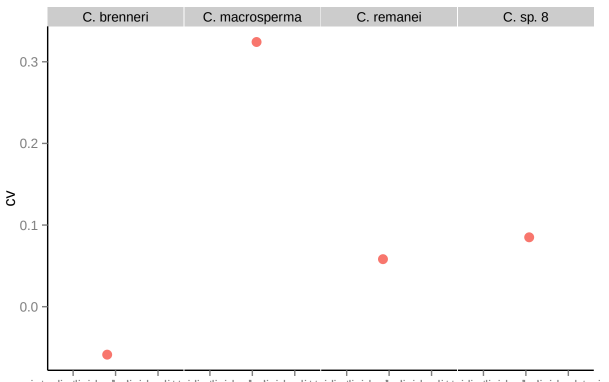
intra individuean individuean straintsa individuean individuean straintsa individuean individuean straints



intrabedixeenabedixee



intra biediweena betivietera listitaarbiediweena listitaarbiediwee



intra biediweduralbetiwielera listraarlisediweduralbetiwielera lis

figure mockup a we are starting agin here, as I've modified some of the variables etc above Easier to start with a clean slate Clean and read in data:

mean and sd

```
groupedmeans=ff %>% group_by(Species,sex) %>% summarise(mean=mean(area,na.rm=T),n=n(),sd=sd(area,na.rm=T))
limits <- aes(ymax = mean + 1.96*sd/sqrt(n), ymin=mean - 1.96*sd/sqrt(n))
dodge <- position_dodge(width=0.9)
ggplot(data=groupedmeans, aes(x=Species, y=mean, fill=sex)) + geom_bar(stat="identity", position=positi
    scale_fill_manual(values=c("gray","white"))+ylab( expression(paste("area (", mu, m^{2},")")))+xlab(
    theme(panel.grid.major = element_blank(), panel.grid.minor = element_blank(),
    panel.background = element_blank(),axis.line = element_line(colour = "black"),legend.position = "n</pre>
```

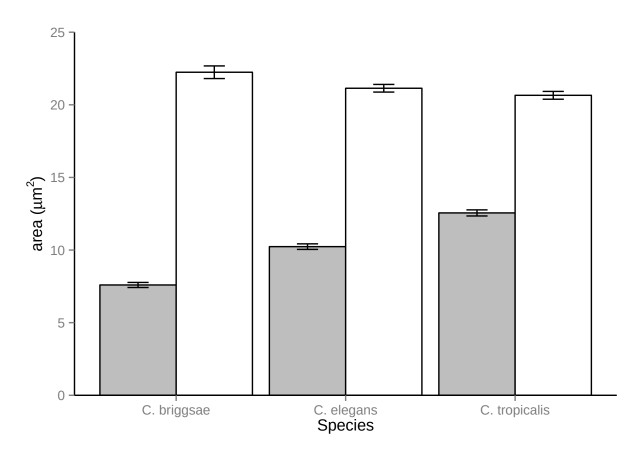


Figure mockup b group to get the data I want it's ugly, but it works

```
groupedstrains=ff %>% group_by(Species,Strain) %>%
    do(data.frame(meanM=mean(.[which(.$sex=="M"),]$area,na.rm=T),meanH=mean(.[which(.$sex=="H"),]$area,na.rm=T)
        sdM=sd(.[which(.$sex=="M"),]$area,na.rm=T),sdH=sd(.[which(.$sex=="H"),]$area,na.rm=T)
groupedstrains$sem=1.96*groupedstrains$sdM/sqrt(groupedstrains$nM)
groupedstrains$seh=1.96*groupedstrains$sdH/sqrt(groupedstrains$nH)
p=ggplot(data=groupedstrains, aes(x=meanM, y=meanH, colour=Species, ymin = meanH - seh,ymax=meanH + seh
p+geom_point(size=5)+coord_cartesian(ylim=c(0,15),xlim=c(0,33))+theme(panel.grid.major = element_blank(
    panel.background = element_blank(),axis.line = element_line(colour = "black"),legend.position = "n
    ylab(expression(paste("Hermaphrodite Sperm area (", mu, m^{2},")")))+xlab(expression(paste("Male Sgeom_smooth(aes(group=Species), method="lm",size=1.5,colour="black")
```

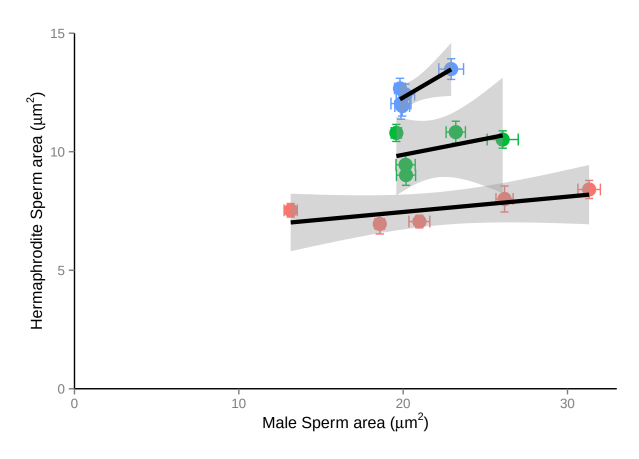
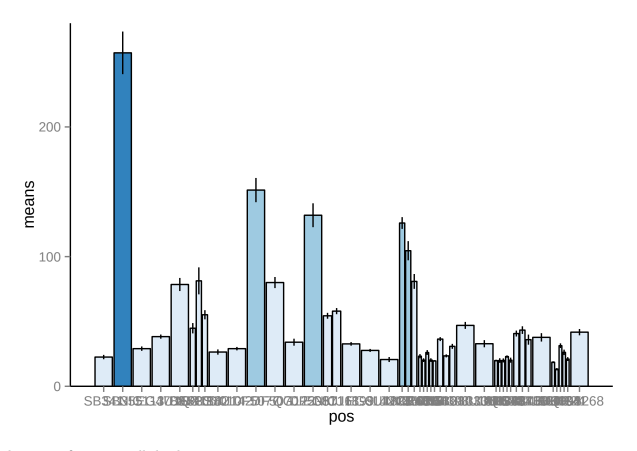


figure1a - bars need to rerun the above

```
ff<-spermsizes<-read.table("C:/Users/jeremy/Desktop/geiger/compileddata.csv",header=T,sep=",")
ff<-renameworms(ff)</pre>
spermsize2<-ff %>% group_by(Species,Strain) %>% summarise(means=mean(area,na.rm=T),cv=sd(area,na.rm=T)/
countspec=function(dataframe){
    holding=c()
    for(i in 1:nrow(dataframe)){
        holding=c(holding,sum(dataframe$Species==dataframe[i,]$Species))
    return(holding)
spermsize2$Species<-ordered(spermsize2$Species,levels=c(tr$tip.label))</pre>
spermsize2<-spermsize2[order(spermsize2$Species),]</pre>
spermsize2$Strain<-factor(spermsize2$Strain, levels = spermsize2$Strain)</pre>
spermsize2$specnum=countspec(spermsize2)
spermsize2$cols=cut(spermsize2$means,3)
findchange<-function(df){</pre>
    holding=c()
    for(i in 1:length(df$Species)){
        if(i==1){
            holding=c(holding,0)
        } else if(df$Species[i]==df$Species[i-1]){
            holding=c(holding,0)
        } else {
            holding=c(holding,1)
```

```
return(holding)
spermsize2$change<-findchange(spermsize2)</pre>
w<-(1-(spermsize2$specnum-1)*0.05)/spermsize2$specnum
pos \leftarrow 0.5 * (cumsum(w) + cumsum(c(0, w[-length(w)])))
gapsizes<-function(df,pos){</pre>
    pos1<-pos
    for(i in 1:length(df$specnum)){
        if(df$specnum[i]==1){
            pos1[i:length(pos1)]<-pos1[i:length(pos1)]+0.1
        } else if(df$change[i]==1){
            pos1[i:length(pos1)]<-pos1[i:length(pos1)]+0.1</pre>
        } else {
            pos1[i:length(pos1)]<-pos1[i:length(pos1)]+0.05
    }
    return(pos1)
}
pos<-gapsizes(spermsize2,pos)</pre>
ggplot(data=spermsize2,aes(x = pos, width = w, y = means, fill=cols,ymin=means-sem,ymax=means+sem)) + s
    geom_bar(stat = "identity",colour="black") + scale_x_continuous(labels = spermsize2$Strain, breaks
    theme(panel.grid.major = element_blank(), panel.grid.minor = element_blank(),
     panel.background = element_blank(),axis.line = element_line(colour = "black"),legend.position = "n
```

Warning: position_stack requires constant width: output may be incorrect



large tree first group all the data

```
spermsize1<-ff %>% group_by(Species) %>% summarise(means=mean(area,na.rm=T))
spermsize2<-ff %>% group_by(Species,Strain) %>% summarise(means=mean(area,na.rm=T))
spermsize3<-ff %>% group_by(Species,Strain,individual) %>% summarise(means=mean(area,na.rm=T))
```

function to fix the format of spermsizes, and fit tree

```
fitace <- function(spermsizes,tree) {
  holding<-spermsizes$means
  names(holding)<-spermsizes$Species
  return(fastAnc(tree,holding,CI=T))
}</pre>
```

function to choose a subset - works on any level, just chooses one from each species

```
subsetter<-function(spermsizes){
   x=lapply(split(spermsizes, spermsizes$Species),function(subdf) subdf[sample(1:nrow(subdf), 1),])
   do.call("rbind",x)
}</pre>
```

example usage fitace(subsetter(spermsizes2),tr) now need to loop and keep the values

```
looper<-function(spermsizes,reps,tree){
  outputdata=data.frame()
  outputCI=data.frame()
  for(i in 1:reps){
     outputdata=rbind(outputdata,fitace(subsetter(spermsizes),tree)$ace)
     outputCI=rbind(outputCI,fitace(subsetter(spermsizes),tree)$CI95)
  }
  names(outputdata)=seq(from=27,to=51)
  return(c(outputdata,outputCI))
}</pre>
```

ok we can call eg x=looper(spermsize3,10,tr) then mean and sd, etc. etc.

```
x<-fitace(spermsize1,tr)
AncSperm=x$ace
#or AncSperm=apply(x,2,mean) if it's bootstrapped
spermsize1$Species <- factor(spermsize1$Species, levels = c("C. sp. 1", "C. plicata", "C. guadeloupensis"
spermsize=spermsize1[order(spermsize1$Species),]$means
names(spermsize)<-spermsize1[order(spermsize1$Species),]$Species</pre>
plot(tr,label.offset=0,no.margin = TRUE,show.tip.label = F,x.lim=c(0,1),edge.width=1.5,edge.color=c(1))
#loads the graph into memory, then plots the anc. states on the nodes
#this is taken more or less from Nicos function
lastPP <- get("last_plot.phylo", envir = .PlotPhyloEnv)</pre>
    node <- (lastPP$Ntip + 1):length(lastPP$xx)</pre>
    XX <- lastPP$xx[node]</pre>
    YY <- lastPP$yy[node]
    for (i in 1:length(XX))
        if (round(AncSperm[i], digits=1)==59.6) #this is a horrible hack to fix the labels if they are too b
             points(XX[i],YY[i],col="red",lwd=round(AncSperm[i], digits=1)/max(round(spermsize,digit=1))*50,pci
             text(XX[i]+0.02,YY[i],labels=round(AncSperm[i], digits=1),col="black",cex=1.2,adj=c(1,0))
        }
         points(XX[i],YY[i],col="red",lwd=round(AncSperm[i], digits=1)/max(round(spermsize,digit=1))*50,pch=
        text(XX[i],YY[i],labels=round(AncSperm[i], digits=1),col="black",cex=1.4,adj=c(1,0))
    }
#now we label the tips
tip <- 1:lastPP$Ntip</pre>
XX <- lastPP$xx[tip]</pre>
YY <- lastPP$yy[tip]
points(XX,YY,col="blue",lwd=(round(spermsize,digit=1)/max(round(spermsize,digit=1))*50),pch=19)
text(XX+0.15,YY,labels=round(spermsize,digit=1),col="blue",cex=1.4)
text(XX+0.07,YY,labels=c("C. sp. 1","C. plicata", "C. guadeloupensis", "C. portoensis", "C. virilis", "C. sp. 1", "C. sp. 1",
```

