# Variola louti

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### Life history analysis for Variola louti from Guam

Code provided for the analyis of size/age at maturity, size/age at sex change, spawning season and lunar spawning frequency. Bayesian growth model and extracting SST from ERDDAP for comparision in edge analysis are in separate R files.

1. Load packages, and load and clean data.

```
library(dplyr)
library(reshape2)
library(lme4)
library(boot)
library(magrittr)
library(plotrix)
library(lubridate)
library(plyr)
library(car)
library(lunar)
library(mgcv)
library(itsadug)
library(ggplot2)
library(ggpubr)
#VALO Reproduction Data & Cleaning
setwd("~/Documents/research/VALO/Vlouti")
valo_r<-read.csv("Reproduction_4_6_2021.csv")</pre>
valo_r<-valo_r %>% subset(Species=="Variola louti") #N=273
valo_r$ScientificName<-valo_r$Species</pre>
#load metadata and sampling data to link with reproduction information
setwd("~/Documents/github/TBP")
lhd<-read.csv("guam_7_22_2019.csv")</pre>
lhd$Date<-as.Date(lhd$FishedDate, format = "%m/%d/%Y")</pre>
lhd$Sex<-revalue(lhd$Sex, c("1 - Female"="F", "2 - Male"="M","3 - Unknown"="M"))
setwd("~/Documents/research/VALO/Vlouti")
lhd<-lhd%>% subset(ScientificName=="Variola louti")
```

```
lhd$SampleID=lhd$SampleTag
lhd<-lhd %>% select(SampleID, Length.cm., Weight.g., Sex, GonWeight, Date, Year, Month)
v_r<-left_join(valo_r, lhd)</pre>
#age data
age <- read.csv ("Age_VALO_March_2023.csv")
valo_ar<-left_join(age, v_r, by="SampleID")</pre>
#sub data sets for plotting use later
valo_ar$H_Sex<-as.factor(valo_ar$H_Sex)</pre>
valo_ar_f<-valo_ar %>%
  subset(H_Sex=="F") #female
valo_ar_m<-valo_ar %>%
  subset(H_Sex=="M" ) #male
valo_ar_po<-valo_ar %>%
  subset(H_Sex=="M_PO" ) #male with primary oocytes
valo_ar_t<-valo_ar %>%
  subset(H_Sex=="T" ) #transitional
```

#### ##Spawning Season

First assess spawning season, then can assess L50. Need to constrain data for L50 to within the spawning season. Since some spawning occurs all year for this species (spawning capable individuals found all year), all the data is used for determining L50. Also look at relationships with gonadosomatic index (GSI) by fish length and reproductive phase.

```
gsi<- v_r%>% mutate(GSI=GonWeight/Weight.g.*100)
gsi$Month<-month(gsi$Date, label=TRUE) #get monthly labels for summarizing and plotting
#separate sexes for analysis
gsi_m<-subset(gsi, H_Sex=="M") #males</pre>
gsi<-subset(gsi, H_Sex=="F") #females</pre>
gsi<- subset(gsi, GSI>=0.00001) #clean and remove NA and zeros
###GSI means and sd by reproductive phase###
gsi <- gsi [complete.cases(gsi$Length.cm.),] #subset data to get records that have Fork Lengths
gsi_phase<-gsi %>%
  dplyr::group_by(Phase) %>%
  dplyr::summarise(mean=mean(GSI), n=length(GSI), Sd=sd(GSI))
gsi_phase$SE <-gsi_phase$Sd/sqrt(gsi_phase$n) #adding standard error to the dataframe
gsi_phase_oocyte<-gsi %>%
  dplyr::group_by(Oocyte_Stage) %>%
  dplyr::summarise(mean=mean(GSI), n=length(GSI), Sd=sd(GSI))
#developing phase qsi and sd (VTI and VTII)
developing_gsi<-gsi %>% subset(Oocyte_Stage=="VTI" | Oocyte_Stage=="VTII")
developing_gsi<-developing_gsi %>%
  dplyr::summarise(mean=mean(GSI), n=length(GSI), Sd=sd(GSI))
developing_gsi$SE <-developing_gsi$Sd/sqrt(developing_gsi$n)</pre>
####GSI vs FL relationship########
gsi<- gsi[complete.cases(gsi$GSI),]</pre>
```

```
gsi <- gsi[order(gsi$Length.cm.),]</pre>
gsi_sc<-gsi %>% subset(Phase=="SC" |Phase=="AS")
gsi_sc$GFBW=gsi_sc$Weight.g.-gsi_sc$GonWeight
#####FL and GSI plots###
{par(mfrow=c(1,2),mar=c(4,4,3,3))}
plot(gsi$GSI~gsi$Length.cm., bty="l", ylab="GSI", xlab="Fork Length (cm)", pch=19, las=1,xaxs="i",yaxs=
gsiU<-gsi %>% subset(Phase=="U")
gsiD<-gsi %>% subset(Phase=="D"|Phase=="RG"|Phase=="RE")
gsiSC<-gsi %>% subset(Phase=="SC"|Phase=="U")
points(gsiU$GSI~gsiU$Length.cm., pch=21,col="black",bg="white", add=TRUE)
points(gsiD$GSI~gsiD$Length.cm., pch=21,col="black",bg="grey", add=TRUE)
text(21,9.5, "a")
plot(log(gsi_sc$GonWeight)~gsi_sc$GFBW, bty="l", ylab="log GW", xlab="GFBW", pch=19,xaxs="i",yaxs="i",x
axis(1, seq(200, 1400, by=200), labels = seq(200, 1400, by=200))
expr<-lm(log(gsi_sc$GonWeight)~gsi_sc$GFBW)</pre>
leng<-seq(100,1200,by=10)
lines(gsi_sc$GFBW,(predict(expr, list(x=leng))), lwd=2)
text(260,4.8, "b")}
summary(expr)
##
## Call:
## lm(formula = log(gsi_sc$GonWeight) ~ gsi_sc$GFBW)
##
## Residuals:
      Min
                1Q Median
                                3Q
                                       Max
## -1.2853 -0.5359 -0.1149 0.4344 1.7040
##
## Coefficients:
                Estimate Std. Error t value Pr(>|t|)
##
## (Intercept) 1.3859840 0.3274603 4.233 0.000131 ***
## gsi sc$GFBW 0.0015748 0.0004329 3.638 0.000778 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Residual standard error: 0.7213 on 40 degrees of freedom
## Multiple R-squared: 0.2486, Adjusted R-squared: 0.2298
## F-statistic: 13.23 on 1 and 40 DF, p-value: 0.000778
#####box plot spawning season#####
#female and sample sizes per month
guam_total<-as.data.frame(gsi %>%
                            dplyr::group_by(Month)%>%
                            dplyr::summarise(total=length(Length.cm.))) #female
male_total <- as.data.frame(gsi_m %>%
                            dplyr::group_by(Month)%>%
                            dplyr::summarise(total=length(Length.cm.))) #male
{par(mfrow=c(2,1),mar=c(3.5,3.5,1.5))}
```

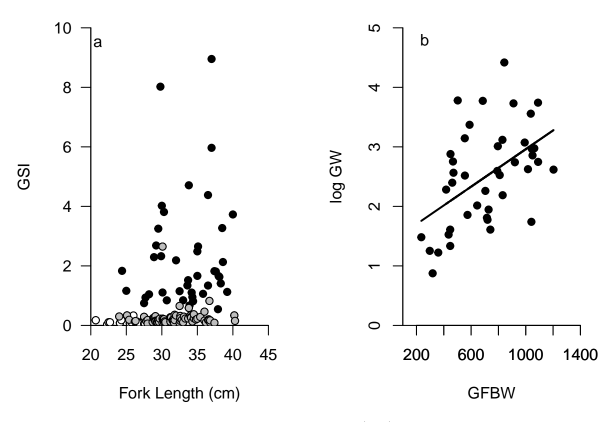


Figure 1: Female Guam Variola louti gonadosomatic index (GSI) versus fork length, and spawning capable and actively spawning female log gonad weight (GW) versus gonad free body weight (GFBW) (log GW = 1.3860 + GFBW \* 0.0016, df = 40, R2=0.23) for fish sampled from Guam between 2010 and 2017. Spawning capable and actively spawning females are black, immature females are white, and all other female reproductive phases are grey.

```
,1),mgp=c(2,.7,0))
boxplot(gsi$GSI~gsi$Month,bty="l",ylab="GSI",xlab="", ylim=c(0.0,10),frame=T,cex.axis=1.0, cex.lab=1.0,
par(xpd=TRUE)
text(x =guam_total$Month, y=11.2, label = guam_total$total , offset=0, cex = 1.0)#too add sample size a
text(0.3,9, "a", cex=1.0)
par(xpd=TRUE)
boxplot(gsi_m$GSI~gsi_m$Month,bty="l",ylab="GSI",xlab="", ylim=c(0,.5), frame=T,cex.axis=1.0, cex.lab=1
text(x =male_total$Month, y=0.56, label = male_total$total , offset=0, cex = 1.0)#too add sample size a
text(0.3,.45, "b", cex=1.0)}
```

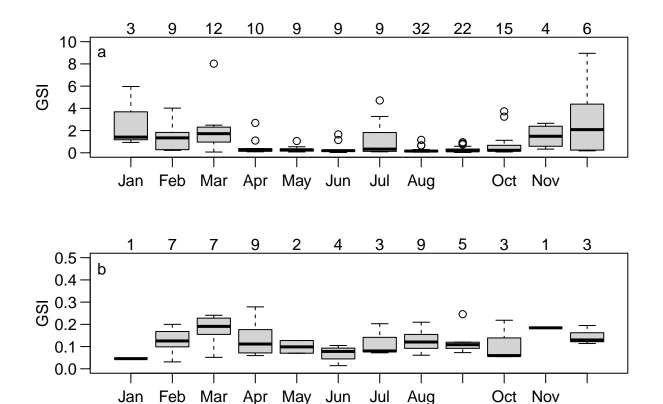


Figure 2: Median and interquartile range of gonadosomatic index (GSI) for mature female (a) and male (b) Variola louti by month

#### Proportion of Female Reproductive Phases per Month

```
###female histo reproductive phase per month frequency graph###
#number of females in each reproductive phase
temp<-gsi
temp <- temp[complete.cases(temp$Length.cm.),]
temp$PHASE<-factor(temp$Phase, levels=c( "AS", "SC", "D", "RG", "RE", "U"), ordered=TRUE)
tempAS<-subset(temp, PHASE=="AS")
tempSC<-subset(temp, PHASE=="SC")</pre>
```

```
tempD<-subset(temp, PHASE=="D")</pre>
tempRG<-subset(temp, PHASE=="RG")</pre>
tempRE<-subset(temp, PHASE=="RE")</pre>
tempU<-subset(temp, PHASE=="U")</pre>
#to compare the lengths of immature and regenerating females - this can be useful to look for outliers
#hist(tempRE$Length.cm.)
#hist(tempU$Length.cm.)
##percent of females that are immature
fem_total<-length(temp$PHASE)</pre>
u=length(tempU$Phase)
immaturef<-u/fem total
prop_month_cast <- dcast(temp, Month~PHASE, value.var="PHASE")</pre>
prop_month_cast$sum <- rowSums(prop_month_cast[2:7])# make sure it adds to 100%
prop_month_cast$AS<-(prop_month_cast$AS/prop_month_cast$sum)*100</pre>
prop_month_cast$D<-(prop_month_cast$D/prop_month_cast$sum)*100</pre>
prop_month_cast$SC<-(prop_month_cast$SC/prop_month_cast$sum)*100</pre>
prop_month_cast$RE<-(prop_month_cast$RE/prop_month_cast$sum)*100</pre>
prop_month_cast$U<-(prop_month_cast$U/prop_month_cast$sum)*100</pre>
prop_month_cast$RG<-(prop_month_cast$RG/prop_month_cast$sum)*100</pre>
# ###add in missing months of sampling if there are any, example below
# #install.packages("tibble")
 \textit{\# prop\_month\_cast} < -add\_row(\textit{prop\_month\_cast}, \textit{Month=6}, \textit{SC=0}, \textit{D=0}, \textit{RE=0}, \textit{RG=0}, \textit{U=0}, \textit{sum=0}) 
prop_month_cast<-prop_month_cast %>% arrange(Month)
prop_month_cast <- prop_month_cast[complete.cases(prop_month_cast$Month),]</pre>
counts.L <- t(prop_month_cast[2:7])</pre>
count.labL <- c("Jan", "Feb", "Mar", "Apr", "May", "Jun", "Jul", "Aug", "Sep", "Oct", "Nov", "Dec")</pre>
sample_n=prop_month_cast$sum
{par(mfrow=c(1,1),mar=c(4,5,2,11))}
sz<-barplot(counts.L,</pre>
             names.arg=count.labL, cex.lab=1.0, cex.axis=1.0,xaxs="i",yaxs="i",cex=1.0,ylab="Percent Fre
par(xpd=TRUE)
legend(15,80,bty="n",legend=c("Actively Spawning", "Spawning Capable", "Developing", "Regenerating", "Regr
        ,col=c( "firebrick", "tan3", "palegoldenrod", "mediumpurple2", "lightcyan2", "steelblue"),pch=15,
text(x = sz, y=104, label = sample_n , offset=10, cex = 1.0)}#too add sample size above plot
```

#### Lunar Spawning Assessment

```
####Lunar Spawning####
#uses libraries lunar, mgcv, itsadug

#subsetting data for functionally mature females and males
gsi<-gsi %>% subset(M_VT==1) #functionally mature females
gsi_m<-gsi_m %>% subset(Mature==1) #mature males
```

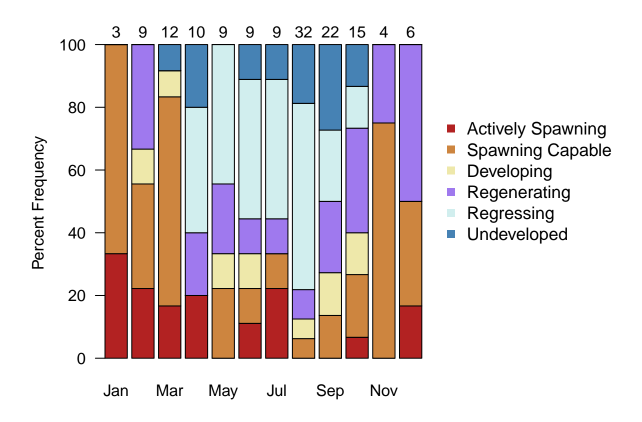


Figure 3: Trend in reproductive stages of female Variola louti by month for fish sampled from Guam between 2010 and 2017. Developing stages (Developing I & Developing II) were grouped together into "Developing." Numbers above each bar indicate aggregate monthly sample sizes.

```
gsi$lunar.phase<-lunar::lunar.phase(gsi$Date,shift=-10, name=8)</pre>
lunar_total<-as.data.frame(gsi %>%
                            dplyr::group_by(lunar.phase)%>%
                            dplyr::summarise(total=length(Length.cm.)))
gsi <-
 gsi %>%
 mutate(lunar day = lunar.phase(as.Date(Date), shift=-10) / 0.212769)
gsi$lunar_day<-round(gsi$lunar_day,0)</pre>
# GAM uses library mgcv
#testing two models and which fits the data better
mod1<-gam(log(GSI)~Length.cm.+s(lunar_day), data=gsi)</pre>
summary(mod1)
##
## Family: gaussian
## Link function: identity
## Formula:
## log(GSI) ~ Length.cm. + s(lunar_day)
## Parametric coefficients:
               Estimate Std. Error t value Pr(>|t|)
                         0.87047 -3.887 0.000171 ***
## (Intercept) -3.38393
                           0.02678 2.931 0.004082 **
## Length.cm. 0.07850
## ---
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
## Approximate significance of smooth terms:
                  edf Ref.df
                                F p-value
## s(lunar_day) 5.533 6.64 3.991 0.000896 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## R-sq.(adj) = 0.221 Deviance explained = 26.4\%
## GCV = 1.2379 Scale est. = 1.1608
                                        n = 121
mod2<-gam(log(GSI)~s(lunar_day), data=gsi)</pre>
summary(mod2)
##
## Family: gaussian
## Link function: identity
## Formula:
## log(GSI) ~ s(lunar_day)
##
## Parametric coefficients:
               Estimate Std. Error t value Pr(>|t|)
## (Intercept) -0.8485 0.1012 -8.38 1.51e-13 ***
## ---
```

```
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## Approximate significance of smooth terms:
                  edf Ref.df
                                 F p-value
## s(lunar_day) 5.331 6.428 3.865 0.00123 **
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## R-sq.(adj) = 0.168
                        Deviance explained = 20.5%
## GCV = 1.3089 Scale est. = 1.2404
#To look at models graphically
#plot.gam(mod1, shade=TRUE)
#plot.gam(mod2, shade=TRUE)
#use AIC to select best model
AIC(mod1, mod2)
##
              df
                      AIC
## mod1 8.532552 370.7134
## mod2 7.331324 377.6118
#to create a box plot of GSI by lunar day need to bin GIS by lunar day - used 3 day bins
gsitemp=gsi
brks \leftarrow seq(from=0,to=30,by=3)
gsitemp$binl <- cut(gsitemp[,"lunar_day"], breaks=brks, labels=brks[1:(length(brks)-1)],right=F)
gsitemp$binl<-as.factor((gsitemp$binl))</pre>
gsitemp<-gsitemp %>%
  dplyr::arrange(desc(lunar_day))%>%
  dplyr::group_by(binl)%>%
  mutate(Mean=mean(GSI, na.rm=T),lower=quantile(GSI,probs = 0.025, na.rm=TRUE), upper=quantile(GSI,prob
gsitemp$binl<-as.numeric(as.character((gsitemp$binl)))</pre>
gsilunarsum<-gsitemp %>%
  dplyr::group_by(binl) %>%
  dplyr::summarise(total=n())
gsilunarsum<-gsilunarsum[1:10,]
gsilunarsum$lunar_day<-gsilunarsum$binl
#make shift code to line up box plots on the x axis over the median lunar day for the bin
c(0,3,6,9,12,15,18,21,24,27)
## [1] 0 3 6 9 12 15 18 21 24 27
gsitemp1<-gsitemp%>%
ungroup()%>%
  select(GSI, binl)
10<-gsitemp1%>%
  subset(binl==0)
10<-10 %>%
  select(GSI)
13<-gsitemp1%>%
```

```
subset(binl==3)
16<-gsitemp1%>%
  subset(binl==6)
19<-gsitemp1%>%
  subset(binl==9)
112<-gsitemp1%>%
  subset(binl==12)
115<-gsitemp1%>%
  subset(binl==15)
118<-gsitemp1%>%
  subset(binl==18)
121<-gsitemp1%>%
  subset(binl==21)
124<-gsitemp1%>%
  subset(binl==24)
127<-gsitemp1%>%
  subset(binl==27)
xplace=as.numeric(c(0,3,6,9,12,15,18,21,24,27))
#lunar spawning figure#
{plot(gsitemp$Mean~gsitemp$lunar_day, ylab="GSI", xlab="Lunar Day", bty="n",frame=T,cex.axis=1.0,ylim=c
\#axis(1,at=c(seq(0,30,by=3)), cex.lab=1.0,cex.axis=1.0, las=1)
boxplot(10$GSI, 13$GSI, 16$GSI, 19$GSI, 112$GSI,115$GSI,118$GSI,121$GSI,124$GSI,127$GSI,
        at = c(1.5, 4.5, 7.5, 10.5, 13.5, 16.5, 19.5, 22.5, 25.5, 28.5),
        names = c("0", "3", "6", "9","12","15","18","21","24","27"),
        las = 2,
        boxwex=3, add=TRUE, bty='n', xaxt="n")
plot_smooth(mod1, view="lunar_day", lwd=2,
            transform = exp, se=FALSE,rug=FALSE, plot_all=NULL,bty='n',add=TRUE) }#plot best fit gam mo
## Summary:
## * Length.cm. : numeric predictor; set to the value(s): 31.8.
## * lunar_day : numeric predictor; with 30 values ranging from 0.000000 to 30.000000.
\#\# * NOTE : No random effects in the model to cancel.
##
```

### Size/age at Maturity

```
#L50 functions
# setup functions -----
data.setup <- function(dataframe,xname,yname){
  templ <- dataframe
  brks <- seq(from=min(templ[,xname]),to=max(templ[,xname]),by=2)
  templ$binl <- cut(templ[,xname], breaks=brks, labels=brks[1:(length(brks)-1)],right=F)
  templ <- dcast(templ, binl~templ[,yname])
  colnames(templ) <- c("binl","a","b")
  templ$binl <- as.character(templ$binl)
  templ$binl <- as.numeric(templ$binl)
  templ$sum <- rowSums(templ[2:3])
  templ$a.p <- templ$a/templ$sum
  templ$b.p <- templ$b/templ$sum</pre>
```

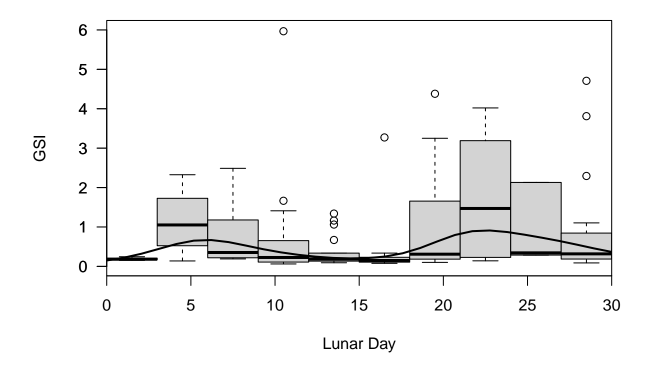
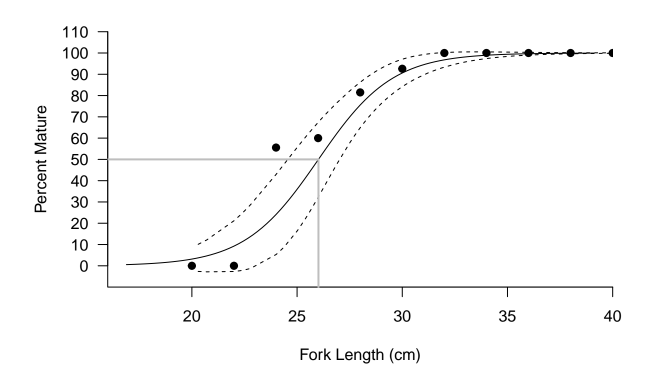


Figure 4: Generalized additive model GSI  $\sim$  Fork Length+s(Lunar Day) fit to raw data points (black line) by lunar day (3 day mean).

```
return(templ)
}
data.setup <- function(dataframe, xname, yname){</pre>
  templ <- dataframe
  brks \leftarrow seq(from=20,to=50,by=2)
  templ$binl <- cut(templ[,xname], breaks=brks, labels=brks[1:(length(brks)-1)],right=F)</pre>
  templ <- dcast(templ, binl~templ[,yname])</pre>
  colnames(templ) <- c("binl", "a", "b")</pre>
  templ$binl <- as.character(templ$binl)</pre>
  templ$binl <- as.numeric(templ$binl)</pre>
  templ$sum <- rowSums(templ[2:3])</pre>
  templ$a.p <- templ$a/templ$sum</pre>
  templ$b.p <- templ$b/templ$sum</pre>
  return(templ)
}
data.setup_age <- function(dataframe, xname, yname){</pre>
  templ <- dataframe
  brks <- seq(from=min(templ[,xname]),to=max(templ[,xname]),by=1)</pre>
  templ$binl <- cut(templ[,xname], breaks=brks, labels=brks[1:(length(brks)-1)],right=F)
  templ <- dcast(templ, binl~templ[,yname])</pre>
  colnames(templ) <- c("binl", "a", "b")</pre>
  templ$binl <- as.character(templ$binl)</pre>
  templ$binl <- as.numeric(templ$binl)</pre>
  templ$sum <- rowSums(templ[2:3])</pre>
  templ$a.p <- templ$a/templ$sum</pre>
  templ$b.p <- templ$b/templ$sum</pre>
  return(templ)
}
model.ci <- function(dataframe, xname, yname){</pre>
  templ <- dataframe
  glm.out <- glm(templ[,yname] ~ templ[,xname], family=binomial(link=logit), data=templ)</pre>
  newdata.l=data.frame(templ[,xname])
  colnames(newdata.1) <- xname</pre>
  templ <- predict(glm.out,newdata=newdata.1, type="response", se.fit=T)</pre>
  ci <- matrix(c(templ$fit+1.96*templ$se.fit,templ$fit-1.96*templ$se.fit),ncol=2)
  ci <- data.frame(ci)</pre>
  ci <- cbind(ci,newdata.1)</pre>
  ci <- ci[with(ci,order(ci[,3])),]</pre>
  return(ci)
est <- function(coef,p) (log(p/(1-p))-coef[1])/coef[2] #p is the proportion
boot.est <- function(dataframe, xname, yname){</pre>
  myfun <- function(dataframe, xname, yname){</pre>
    templ <- dataframe
    srows <- sample(1:nrow(templ),nrow(templ),TRUE)</pre>
    glm.out <- glm(templ[,yname] ~ templ[,xname], family=binomial(link=logit), data=templ[srows,])</pre>
    return(est(coef(glm.out),0.5))
```

```
bootdist <- replicate(1000, myfun(dataframe,xname,yname))</pre>
    boot.ci <- quantile(unlist(bootdist),c(.025,.975)) # 95% CI
    return(boot.ci)
}
repro_f<-subset(v_r, H_Sex=="F")</pre>
#repro_f<-subset(repro_f, Phase!="RE")</pre>
repro_f <- repro_f[complete.cases(repro_f$Length.cm.),]</pre>
\#repro_f < -repro_f \%\% subset(Month=="Apr"|Month=="May"|Month=="Jun"|Month=="Jul"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Aug"|Month=="Au
lbins.150vt <- data.setup(repro_f,"Length.cm.","M_VT")</pre>
glm.150vt <- glm(M_VT ~ Length.cm., family=binomial(link=logit), data=repro_f)</pre>
150.civt <- model.ci(repro_f,"Length.cm.","M_VT")</pre>
length(repro_f$Length.cm.)
## [1] 146
bcL_tlvt<-bootCase(glm.150vt, R=1000)</pre>
lrPerc <- function(cf,p) (log(p/(1-p))-cf[1])/cf[2] #p is the proportion</pre>
150vt<-lrPerc(coef(glm.150vt), 0.5)</pre>
150vt<-apply(bcL_tlvt, 1, lrPerc, p=0.5)
150civt<-quantile(150vt, c(0.25,0.975))
150vt <- est(coef(glm.150vt),.5)
#physiological maturity
lbins.150ca <- data.setup(repro_f,"Length.cm.","M_CA")</pre>
glm.150ca <- glm(M_CA ~ Length.cm., family=binomial(link=logit), data=repro_f)</pre>
150.cica <- model.ci(repro_f, "Length.cm.", "M_CA")
length(repro_f$Length.cm.)
## [1] 146
bcL_tlca<-bootCase(glm.150ca, R=1000)</pre>
lrPerc <- function(cf,p) (log(p/(1-p))-cf[1])/cf[2] #p is the proportion</pre>
150ca<-lrPerc(coef(glm.150ca), 0.5)</pre>
150ca<-apply(bcL_tlca, 1, lrPerc, p=0.5)
150cica <- quantile (150ca, c(0.25,0.975))
150ca <- est(coef(glm.150ca),.5)
#####Size at functional maturity####
plot(lbins.150vt$bin,lbins.150vt$b.p,pch=19,cex.axis=1.0, cex=1.0,cex.lab=1.0, bty="1",xlab="Fork Lengt"
          ylim=c(-0.1,1.1),yaxt="n",yaxs="i",xaxs="i",xlim=c(16,40))
axis(2,at=c(seq(0,1.1,by=.1)),labels=c(seq(0,110,by=10)), cex.lab=1.0,cex.axis=1.0, las=1)
curve((exp(coef(glm.150vt)[1]+coef(glm.150vt)[2]*x))/(1+exp(coef(glm.150vt)[1]+coef(glm.150vt)[2]*x)),a
lines(150.civt[,3],150.civt[,1],lty=2)
lines(150.civt[,3],150.civt[,2],1ty=2)
lines(c(150vt,150vt),c(-0.1,.5), lwd=2,col="grey",lty=1, font=2)
lines(c(16,150vt),c(.5,.5), lwd=2,col="grey",lty=1,font=2)
```

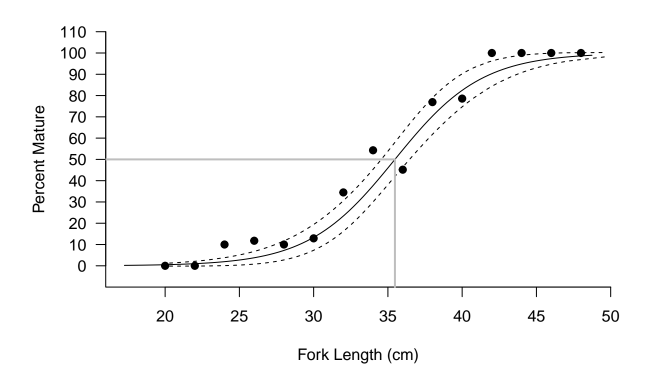


```
#age at maturity
valo_ar_f<-valo_ar %>%
  subset(H_Sex=="F")
valo_ar_f<-valo_ar_f[complete.cases(valo_ar_f$reage),] #make sure there are no blanks in ages
abins.a50vt <- data.setup_age(valo_ar_f,"reage","M_VT")</pre>
glm.a50vt <- glm(M_VT ~ reage, family=binomial(link=logit), data=valo_ar_f)</pre>
a50.civt <- model.ci(valo_ar_f, "reage", "M_VT")
coef(glm.a50vt)
                      reage
## (Intercept)
     -2.893769
##
                   1.103650
length(valo_ar_f$reage)
## [1] 134
bcL_tlvt<-bootCase(glm.a50vt, R=1000)</pre>
lrPerc \leftarrow function(cf,p) (log(p/(1-p))-cf[1])/cf[2] #p is the proportion
a50vt<-lrPerc(coef(glm.a50vt), 0.5)
a50vt<-apply(bcL_tlvt, 1, lrPerc, p=0.5)
a50civt<-quantile(a50vt, c(0.25,0.975))
a50vt <- est(coef(glm.a50vt),.5)
```

```
#physiological
abins.a50ca <- data.setup_age(valo_ar_f,"reage","M_CA")
glm.a50ca <- glm(M_CA ~ reage, family=binomial(link=logit), data=valo_ar_f)
a50.cica <- model.ci(valo_ar_f,"reage","M_CA")
length(valo_ar_f$reage)</pre>
```

## [1] 134

```
bcL_tlca<-bootCase(glm.a50ca, R=1000)</pre>
lrPerc \leftarrow function(cf,p) (log(p/(1-p))-cf[1])/cf[2] #p is the proportion
a50ca<-lrPerc(coef(glm.a50ca), 0.5)
a50ca<-apply(bcL_tlca, 1, lrPerc, p=0.5)
a50cica < -quantile(a50ca, c(0.25, 0.975))
a50ca <- est(coef(glm.a50ca),.5)
#####Size at sex change#####
v_r$SexID<-v_r$H_Sex
v_r$SexID<-revalue(v_r$SexID, c("F"=0, "T"=1, "M"=1, "M_P0"=1))</pre>
v r$SexID<-as.factor(v r$SexID)</pre>
v_r<-v_r[complete.cases(v_r$Length.cm.),]</pre>
#remove rows with blanks
v_r<-v_r[!(is.na(v_r$SexID) | v_r$SexID==""), ] #removed two rows with blank H_Sex
lbins.150sex <- data.setup(v_r, "Length.cm.", "SexID")</pre>
glm.150sex <- glm(SexID ~ Length.cm., family=binomial(link=logit), data=v_r)</pre>
150.cisex <- model.ci(v_r,"Length.cm.","SexID")</pre>
bcL_tlsex<-bootCase(glm.150sex, R=1000)</pre>
lrPerc \leftarrow function(cf,p) (log(p/(1-p))-cf[1])/cf[2] #p is the proportion
150sex<-lrPerc(coef(glm.150sex), 0.5)</pre>
150sex<-apply(bcL_tlsex, 1, lrPerc, p=0.5)
150cisex<-quantile(150sex, c(0.25,0.975))
150sex <- est(coef(glm.150sex),.5)</pre>
{plot(lbins.150sex$bin,lbins.150sex$b.p,pch=19,cex.axis=1.0, cex=1.0,cex.lab=1.0, bty="l",xlab="Fork Le
     ylim=c(-0.1,1.1), yaxt="n", yaxs="i", xaxs="i", xlim=c(16,50))
axis(2,at=c(seq(0,1.1,by=.1)),labels=c(seq(0,110,by=10)), cex.lab=1.0,cex.axis=1.0, las=1)
curve((exp(coef(glm.150sex)[1]+coef(glm.150sex)[2]*x))/(1+exp(coef(glm.150sex)[1]+coef(glm.150sex)[2]*x
lines(150.cisex[,3],150.cisex[,1],1ty=2)
lines(150.cisex[,3],150.cisex[,2],1ty=2)
lines(c(150sex,150sex),c(-0.1,.5), lwd=2,col="grey",lty=1, font=2)
lines(c(16,150sex),c(.5,.5), lwd=2,col="grey",lty=1,font=2)}
```

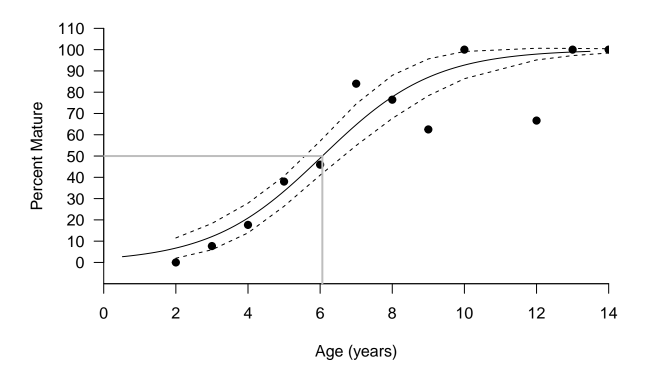


```
#Age at sex change
valo_ar$SexID<-valo_ar$H_Sex
valo_ar$SexID<-revalue(valo_ar$SexID, c("F"=0, "T"=1, "M"=1,"M_PO"=1))
valo_ar$SexID<-as.factor(valo_ar$SexID)
valo_ar<-valo_ar[complete.cases(valo_ar$SexID),]
valo_ar<-valo_ar[complete.cases(valo_ar$tength.cm.),] ###went from 273 to 257
valo_ar<-valo_ar[!(is.na(valo_ar$SexID) | valo_ar$SexID==""), ] #removed two rows with blank H_Sex
abins.a50sex <- data.setup_age(valo_ar,"reage","SexID")
abins.a50sex[12,1]<-14 #not sure why it dropped the last age
glm.a50sex <- glm(SexID ~ reage, family=binomial(link=logit), data=valo_ar)
a50.cisex <- model.ci(valo_ar,"reage","SexID")
length(valo_ar$Length.cm.)</pre>
```

#### ## [1] 229

```
bcL_tlsex<-bootCase(glm.a50sex, R=1000)
lrPerc <- function(cf,p) (log(p/(1-p))-cf[1])/cf[2]#p is the proportion
a50sex<-lrPerc(coef(glm.a50sex), 0.5)
a50sex<-apply(bcL_tlsex, 1, lrPerc, p=0.5)
a50cisex<-quantile(a50sex, c(0.25,0.975))
a50sex <- est(coef(glm.a50sex),.5)</pre>
```

```
#Age sex change
plot(abins.a50sex$bin,abins.a50sex$b.p,pch=19,cex.axis=1.0, cex=1.0,cex.lab=1.0, bty="l",xlab="Age (year,ylim=c(-0.1,1.1),yaxt="n",yaxs="i",xaxs="i",xlim=c(0,14))
axis(2,at=c(seq(0,1.1,by=.1)),labels=c(seq(0,110,by=10)), cex.lab=1.0,cex.axis=1.0, las=1)
curve((exp(coef(glm.a50sex)[1]+coef(glm.a50sex)[2]*x))/(1+exp(coef(glm.a50sex)[1]+coef(glm.a50sex)[2]*x
lines(a50.cisex[,3],a50.cisex[,1],lty=2)
lines(a50.cisex[,3],a50.cisex[,2],lty=2)
lines(c(a50sex,a50sex),c(-0.1,.5), lwd=2,col="grey",lty=1, font=2)
lines(c(0,a50sex),c(.5,.5), lwd=2,col="grey",lty=1,font=2)
```

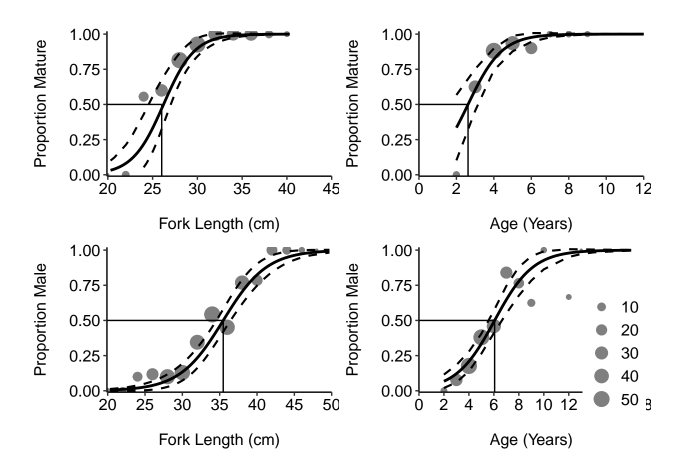


### Bubble Plot for Size/Age at Maturity and Size/Age at Sex Chnage

```
geom_segment(aes(x = 19.9, y = 0.5, xend =150vt, yend = 0.5))+ #to add L50 lines if wanted
  geom_segment(aes(x = 150vt, y = -0.01, xend = 150vt, yend = 0.5))+ #to add L50 lines if wanted
  theme_bw() +
  theme(
   panel.border = element_blank(),
   panel.grid = element_blank(),
   axis.title = element_text(size = 12),
   axis.title.y = element text(margin= margin(r= 15)), # move title away from axis,
   axis.title.x = element_text(margin= margin(t= 15)),
   legend.position = "none",
   # legend.text = element_text(colour="black", size=10, face="bold"),
   #legend.title = element_text(colour="black", size=10, face="bold"),
   axis.text = element_text(colour = "black", size = 12),
   axis.line = element_line(colour = "black")) +
  xlab('Fork Length (cm)') +
  ylab("Proportion Mature")+
  scale_x_continuous(name = "Fork Length (cm)",breaks=seq(20,45,5))
\#150bubble<-150bubble+theme(legend.position = c(.87,.2),
#legend.text = element_text(colour="black", size=12),
#legend.title = element_text(colour="black", size=12))
150bubble < -150bubble + scale y continuous(expand = c(0, 0), limits = c(-0.01, 1.02)) +
  scale_x = c(19.9,45), breaks = seq(20,45,5)
#Age at maturity
 a50bubble<- ggplot()+
  geom_smooth(data = valo_f, aes(x = reage, y = M_VT),
             method = "glm", method.args = list(family = "binomial"),
             se = FALSE, color="black") +
  geom_point(data=abins.a50vt, aes(x = binl, y =b.p, size=sum), alpha=0.5) +
  scale_size(range = c(.5, 5), name="Sample Size")+
  geom_line(aes(x=a50.civt[,3],y=a50.civt[,1], group=1), linetype="dashed", size= 0.75)+
  geom_line(aes(x=a50.civt[,3],y=a50.civt[,2], group=1), linetype="dashed", size= 0.75)+
  geom_segment(aes(x = 0, y = 0.5, xend = a50vt, yend = 0.5))+ #to add L50 lines if wanted
  geom_segment(aes(x = a50vt, y = -0.01, xend = a50vt, y end = 0.5))+ #to add L50 lines if wanted
  theme bw() +
  theme(
   panel.border = element_blank(),
   panel.grid = element_blank(),
   axis.title = element_text(size = 12),
   axis.title.y = element_text(margin= margin(r= 15)), # move title away from axis,
   axis.title.x = element_text(margin= margin(t= 15)),
   legend.position = "none",
   legend.text = element_blank(),
   legend.title = element_blank(),
   axis.text = element_text(colour = "black", size = 12),
   axis.line = element_line(colour = "black")) +
  xlab('Age (Years)') +
  ylab("Proportion Mature")+
  scale_x_continuous(name = "Age (years)",breaks=seq(0,14,2))
```

```
a50bubble<- a50bubble+ scale_y_continuous(expand = c(0, 0), limits = c(-0.01, 1.02))+
       scale_x_{continuous}(expand = c(0, 0), limits = c(0, 12), breaks=seq(0,12,2))
##length at sex change
   150sexbubble <- ggplot()+
       geom smooth(\frac{data}{data} = v r, \frac{data}{data} = v r, \frac{data}{
                              method = "glm", method.args = list(family = "binomial"),
                              se = FALSE, color="black") +
       geom_point(data=lbins.150sex, aes(x = bin1, y =b.p, size=sum), alpha=0.5) +
       scale_size(range = c(.5, 5), name="Sample Size")+
       geom_line(aes(x=150.cisex[,3],y=150.cisex[,1], group=1), linetype="dashed", size= 0.75)+
       geom_line(aes(x=150.cisex[,3],y=150.cisex[,2], group=1), linetype="dashed", size= 0.75)+
       geom_segment(aes(x = 19.9, y = 0.5, xend =150sex , yend = 0.5))+ #to add L50 lines if wanted
       geom_segment(aes(x = 150sex, y = -0.01, xend = 150sex, y yend = 0.5))+ #to add L50 lines if wanted
       theme_bw() +
       theme(
           panel.border = element_blank(),
          panel.grid = element_blank(),
           axis.title = element text(size = 12),
          axis.title.y = element_text(margin= margin(r= 15)), # move title away from axis,
           axis.title.x = element_text(margin= margin(t= 15)),
          legend.position = "none",
          legend.text = element_blank(),
          legend.title = element_blank(),
           axis.text = element_text(colour = "black", size = 12),
           axis.line = element_line(colour = "black")) +
       xlab('Fork Length (cm)') +
       ylab("Proportion Male")+
       scale_x_continuous(name = "Fork Length (cm)",breaks=seq(20,50,5))
   150sexbubble - 150sexbubble + scale_y_continuous(expand = c(0, 0), limits = c(-0.01, 1.02), breaks=seq
       scale_x_continuous(expand = c(0, 0), limits = c(19.9,50), breaks=seq(20,50,5))
#age at sex chage
   valo ar <- valo ar[!(valo ar$SexID == ""), ]</pre>
   valo_ar$SexID[valo_ar$SexID==' ']=NA
   valo_ar$SexID= droplevels(valo_ar$SexID)
   a50sexbubble <- ggplot()+
       geom_smooth(data = valo_ar, aes(x = reage, y = as.numeric(as.character(SexID))),
                              method = "glm", method.args = list(family = "binomial"),
                              se = FALSE, color="black") +
            geom_point(data=abins.a50sex, aes(x = bin1, y =b.p, size=sum), alpha=0.5) +
       scale_size(range = c(.5, 5), name=" ")+
       geom_line(aes(x=a50.cisex[,3],y=a50.cisex[,1], group=1), linetype="dashed", size= 0.75)+
       geom_line(aes(x=a50.cisex[,3],y=a50.cisex[,2], group=1), linetype="dashed", size= 0.75)+
       geom_segment(aes(x = 0, y = 0.5, xend =a50sex , yend = 0.5))+ #to add L50 lines if wanted
       geom_segment(aes(x = a50sex, y = -0.01, xend =a50sex, y yend = 0.5))+ #to add L50 lines if wanted
       theme_bw() +
       theme(
           panel.border = element_blank(),
```

```
panel.grid = element_blank(),
      axis.title = element_text(size = 12),
      axis.title.y = element_text(margin= margin(r= 15)), # move title away from axis,
      axis.title.x = element_text(margin= margin(t= 15)),
     legend.position = "none",
     legend.text = element_blank(),
     legend.title = element_blank(),
     axis.text = element text(colour = "black", size = 12),
     axis.line = element_line(colour = "black")) +
   xlab('Age (Years)') +
   ylab("Proportion Male")+
    scale_x_continuous(name = "Age (years)",breaks=seq(0,18,2))
  a50sexbubble<-a50sexbubble +
  scale_y_continuous(expand = c(0, 0), limits = c(-0.01, 1.02))+
    scale_x_{continuous}(expand = c(0, 0), limits = c(0, 18), breaks=seq(0,18,2))
  scale_y_continuous(limits = c(0.00, 1))
## <ScaleContinuousPosition>
## Range:
## Limits:
              0 --
 \#150bubble<-150bubble+theme(legend.position = c(.8,.25),
  #
                              legend.text = element_text(colour="black", size=10),
  #
                              legend.title = element_text(colour="black", size=10))
  a50sexbubble<- a50sexbubble+theme(legend.position = c(.87,.35),
                                legend.text = element_text(colour="black", size=12),
                                legend.title = element_text(colour="black", size=12))
  figure <- ggpubr::ggarrange(150bubble, a50bubble,150sexbubble, a50sexbubble,
                      labels = c("", "", "",""),
                      ncol = 2, nrow = 2,
                      hjust=-8.2,
                      vjust=0.6,
                      font.label = list(size = 12, color = "black", face = "plain", family = NULL))
  figure
```

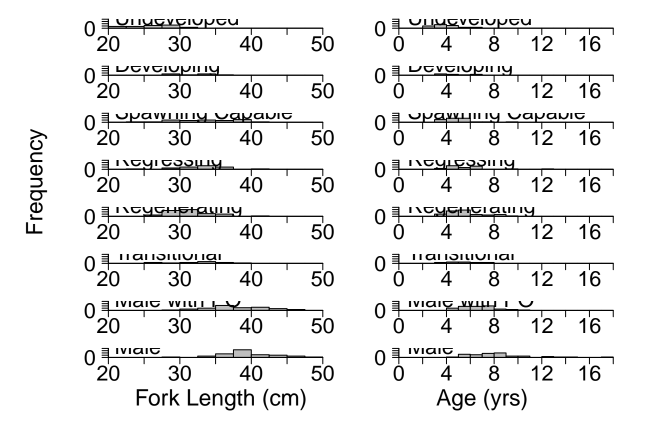


## Length/Age Frequency Distribution

```
###age freq distribution#######
valo_f_u<-valo_ar %>%
  subset(H_Sex=="F" & Phase=="U")
valo_f_d<-valo_ar %>%
  subset(H_Sex=="F" & Phase=="D")
valo_f_sc<-valo_ar %>%
  subset(H_Sex=="F" & Phase=="SC")
valo_f_rg<-valo_ar %>%
  subset(H_Sex=="F" & Phase=="RG")
valo_f_re<-valo_ar %>%
  subset(H_Sex=="F" & Phase=="RE")
valo_t<-valo_ar %>%
  subset(H_Sex=="T")
valo_t_po<-valo_ar %>%
  subset(H_Sex=="M_PO")
valo_m<-valo_ar %>%
  subset(H_Sex=="M")
#Plot it. I save it width=2900, height=3800, res=250 t see all the details in the figure that you cant
{par(mfcol=c(8,2), mar=c(1,5,2,1), oma=c(4,3,.4,.4))}
hist(valo_f_u$Length.cm., right=FALSE,main="" ,xaxs="i",yaxs="i", breaks=seq(20,50,2.5), ylab="", xlab=
axis(1,at=seq(20,50,by=5), labels = seq(20,50,by=5), cex=2, cex.lab=2, cex.axis=2)
```

```
box(bty = "1")
text(20,18.5, "Undeveloped", cex=2, cex.lab=2, cex.axis=2, pos = 4)
hist(valo_f_d$Length.cm., right=FALSE,main="" ,xaxs="i",yaxs="i",breaks=seq(20,50,2.5), ylab="",xlab=""
axis(1,at=seq(20,50,by=5), labels=seq(20,50,by=5),cex=2, cex.lab=2, cex.axis=2)
box(bty = "l")
text(20,18.5, "Developing",cex=2, cex.lab=2, cex.axis=2, pos = 4)
hist(valo_f_sc$Length.cm., right=FALSE,main="" ,xaxs="i",yaxs="i",breaks=seq(20,50,2.5), ylab="",xlab="
axis(1,at=seq(20,50,by=5), labels=seq(20,50,by=5), cex=2, cex.lab=2, cex.axis=2)
text(20,18.5, "Spawning Capable",cex=2, cex.lab=2, cex.axis=2, pos = 4)
box(bty = "l")
hist(valo_f_rg$Length.cm.,right=FALSE, main="" ,xaxs="i",yaxs="i",breaks=seq(20,50,2.5), ylab="", xlab=
axis(1,at=seq(20,50,by=5), labels = seq(20,50,by=5), cex=2, cex.lab=2, cex.axis=2)
text(20,18.5, "Regressing",cex=2, cex.lab=2, cex.axis=2, pos = 4)
box(bty = "1")
hist(valo_f_re$Length.cm.,right=FALSE, main="" ,xaxs="i",yaxs="i",breaks=seq(20,50,2.5), ylab="", xlab=
axis(1,at=seq(20,50,by=5), labels = seq(20,50,by=5), cex=2, cex.lab=2, cex.axis=2)
text(20,18.5, "Regenerating", cex=2, cex.lab=2, cex.axis=2, pos = 4)
box(bty = "l")
hist(valo_t$Length.cm.,right=FALSE, main="" ,xaxs="i",yaxs="i",breaks=seq(20,50,2.5), ylab="", xlab=""
axis(1,at=seq(20,50,by=5), labels = seq(20,50,by=5), cex=2, cex.lab=2, cex.axis=2)
text(20,18.5, "Transitional",cex=2, cex.lab=2, cex.axis=2, pos = 4)
box(bty = "l")
hist(valo_t_po$Length.cm.,right=FALSE, main="" ,xaxs="i",yaxs="i",breaks=seq(20,50,2.5), ylab="", xlab=
axis(1,at=seq(20,50,by=5), labels=seq(20,50,by=5), cex=2, cex.lab=2, cex.axis=2)
text(20,18.5, "Male with PO",cex=2, cex.lab=2, cex.axis=2,pos = 4)
box(bty = "1")
hist(valo_m$Length.cm.,right=FALSE, main="" ,xaxs="i",yaxs="i",breaks=seq(20,50,2.5), ylab="", xlab="Fo
axis(1,at=seq(20,50,by=5), labels = seq(20,50,by=5), cex=2, cex.lab=2, cex.axis=2)
text(20,18.5, "Male", cex=2, cex.lab=2, cex.axis=2,pos = 4)
box(bty = "l")
mtext(text="Fork Length (cm)", side=1,line=2,adj=0.22,outer=TRUE, cex=1.4)
mtext(text="Frequency",side=2,line=0,outer=TRUE, cex=1.4)
hist(valo_f_u$reage, right=FALSE, main="" ,xaxs="i",yaxs="i", breaks=seq(0,18, 1), ylab="", xlab="" ,yl
axis(1,at=seq(0,18,by=2), labels=seq(0,18,by=2), cex=2, cex.lab=2, cex.axis=2)
text(0.2,18.5, "Undeveloped", cex=2, cex.lab=2, cex.axis=2, pos = 4)
hist(valo_f_d$reage,right=FALSE, main="" ,xaxs="i",yaxs="i",breaks=seq(0,18, 1), ylab="",xlab="" ,ylim=
axis(1,at=seq(0,18,by=2), labels=seq(0,18,by=2), cex=2, cex.lab=2, cex.axis=2)
text(0.2,18.5, "Developing",cex=2, cex.lab=2, cex.axis=2,pos = 4)
box(bty = "l")
hist(valo_f_sc$reage, right=FALSE,main="" ,xaxs="i",yaxs="i",breaks=seq(0,18, 1), ylab="",xlab="" ,ylim
axis(1,at=seq(0,18,by=2), labels=seq(0,18,by=2), cex=2, cex.lab=2, cex.axis=2)
text(0.2,18.5, "Spawning Capable", cex=2, cex.lab=2, cex.axis=2, pos = 4)
box(bty = "l")
hist(valo_f_rg$reage,right=FALSE, main="" ,xaxs="i",yaxs="i",breaks=seq(0,18, 1), ylab="", xlab="" ,ylinest(valo_f_rg$reage,right=FALSE, main="" ,xaxs="i",breaks=seq(0,18, 1), ylab="" ,xlab="" ,ylinest(valo_f_rg$reage,right=FALSE, main="" ,xaxs="i",breaks=seq(0,18, 1), ylab="" ,xlab="" ,ylinest(valo_f_rg$reage,right=FALSE, main="" ,xaxs="i",breaks=seq(0,18, 1), ylab="" ,xlab="" ,xlab=" 
axis(1,at=seq(0,18,by=2), labels=seq(0,18,by=2), cex=2, cex.lab=2, cex.axis=2)
text(0.2,18.5, "Regressing",cex=2, cex.lab=2, cex.axis=2,pos = 4)
box(bty = "l")
hist(valo_f_re$reage,right=FALSE, main="" ,xaxs="i",yaxs="i",breaks=seq(0,18, 1), ylab="", xlab="" ,yli
axis(1,at=seq(0,18,by=2), labels = seq(0,18,by=2), cex=2, cex.lab=2, cex.axis=2)
text(0.2,18.5, "Regenerating",cex=2, cex.lab=2, cex.axis=2,pos = 4)
```

```
box(bty = "l")
hist(valo_t$reage,right=FALSE, main="" ,xaxs="i",yaxs="i",breaks=seq(0,18, 1), ylab="", xlab="" ,ylim=c
axis(1,at= seq(0,18,by=2) , labels = seq(0,18,by=2) ,cex=2, cex.lab=2, cex.axis=2 )
text(0.2,18.5, "Transitional",cex=2, cex.lab=2, cex.axis=2,pos = 4)
box(bty = "l")
hist(valo_t_po$reage,right=FALSE, main="" ,xaxs="i",yaxs="i",breaks=seq(0,18, 1), ylab="", xlab="" ,ylinaxis(1,at= seq(0,18,by=2) , labels = seq(0,18,by=2),cex=2, cex.lab=2, cex.axis=2 )
text(0.2,18.5, "Male with PO",cex=2, cex.lab=2, cex.axis=2,pos = 4)
box(bty = "l")
hist(valo_m$reage,right=FALSE, main="" ,xaxs="i",yaxs="i",breaks=seq(0,18,by=1), ylab="", xlab="Age (yraxis(1,at= seq(0,18,by=2) , labels = seq(0,18,by=2), cex=2, cex.lab=2, cex.axis=2 )
text(0.2,18.5, "Male", cex=2, cex.lab=2, cex.axis=2,pos = 4)
box(bty = "l")
mtext(text="Age (yrs)",side=1,line=2, adj=.8,outer=TRUE, cex=1.4)}
```



Male and female size and age differences

```
males<-rbind(valo_ar_m, valo_ar_po)
ks.test(valo_ar_f$Length.cm.,males$Length.cm., alternative = "greater")</pre>
```

```
##
## Asymptotic two-sample Kolmogorov-Smirnov test
##
```

```
## data: valo_ar_f$Length.cm. and males$Length.cm.
## D^+ = 0.64557, p-value < 2.2e-16
## alternative hypothesis: the CDF of x lies above that of y
ks.test(valo_ar_f$reage,males$reage, alternative = "greater")
##
## Asymptotic two-sample Kolmogorov-Smirnov test
## data: valo_ar_f$reage and males$reage
## D^+ = 0.49613, p-value = 1.101e-11
## alternative hypothesis: the CDF of x lies above that of y
t.test(valo_ar_f$Length.cm., males$Length.cm.)
##
## Welch Two Sample t-test
## data: valo_ar_f$Length.cm. and males$Length.cm.
## t = -12.805, df = 183.47, p-value < 2.2e-16
## alternative hypothesis: true difference in means is not equal to 0
## 95 percent confidence interval:
## -8.601041 -6.304349
## sample estimates:
## mean of x mean of y
## 31.15672 38.60941
t.test(valo_ar_f$reage, males$reage)
##
## Welch Two Sample t-test
## data: valo_ar_f$reage and males$reage
## t = -8.2563, df = 130.34, p-value = 1.457e-13
## alternative hypothesis: true difference in means is not equal to 0
## 95 percent confidence interval:
## -2.949576 -1.809281
## sample estimates:
## mean of x mean of y
## 4.656716 7.036145
max(males$reage, na.rm=TRUE)
## [1] 17
mean(males$reage, na.rm=TRUE)
```

## [1] 7.036145

mean(valo\_ar\_f\$reage, na.rm=TRUE)

## [1] 4.656716